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Broader context

Rechargeable batteries are beginning to enter much wider use owing to the fact that their positive social and environmental impacts in terms of renewable energy integration, the proliferation of electric vehicles, energy security, and sustainable development are becoming increasingly understood and appreciated. The requirements for batteries intended for emerging applications include, but are not limited to, a high-level of safety, wide-temperature durability, and high power output, which are quite dissimilar from those intended for use under ambient conditions. Thus, unprecedented opportunities exist for materials science and technology research in this field. The use of Na secondary batteries with appropriate electrolytes, such as jonic liquids (ILs), is one of the most promising strategies to meet the demands for sustainability, safety, and performance, without increasing manufacturing difficulty or necessitating special cell designs. In this review, we provide an introduction to the fundamentals of ILs themselves as well as a variety of case studies demonstrating their feasibility as electrolytes for Na secondary batteries in concert with a variety of electrode materials. We hope that this review will provide a convenient basis for further improvement and optimisation of Na secondary batteries and help make them a commercial reality.

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Advances in sodium secondary batteries utilizing ionic liquid electrolytes[†]

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The development of Na secondary batteries that exhibit both sustainability and high energy density as potential successors to lithium-ion batteries for certain large-scale applications has received considerable research interest in recent years. However, although the importance of the electrolyte in such systems has long been largely overlooked, it is becoming increasingly recognized as a key consideration (along with the electrode material) for the non-incremental improvement of Na secondary batteries. Among the candidate electrolytes in this context, ionic liquids (ILs), which are liquids consisting entirely of ions, offer many unique advantages. In this review, the fundamental properties of ILs and the design strategies employed to facilitate their application in batteries are introduced. Comprehensive summaries of the recent advances in the development of positive and negative electrode materials for Na secondary batteries are then presented. Most of the IL-based systems discussed exhibit remarkably enhanced performances compared to those of batteries based on conventional electrolytes. Furthermore, reversible capacity, rate capability, recyclability, and coulombic efficiency are synergistically enhanced by combining IL electrolytes and elevated temperature conditions. Finally, the practical prospects and future challenges associated with the development of electrode materials fabricated from cheap, abundant elements; the efficient utilisation of Na metal as a negative electrode material; and considerations related to the solid-electrolyte interphase are also discussed

1. Introduction

A stable supply of energy is a vital global issue given the continuous increase in the demand for fossil fuels and the finiteness of these non-renewable resources. The present energy economy relies heavily on fossil fuels and is thus subject to problems in terms of future exhaustion as well as climate change caused by increasing atmospheric concentrations of greenhouse gases. Accordingly, increasing environmental awareness combined with energy safety concerns have led society to seek methods of generating, harnessing, and storing energy in a more sustainable and efficient manner.

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Energy storage systems (ESSs) are starting to play a transformative role in this context. Their application scope spans from personal and residential end-use to broader applications such as improving grid reliability and ameliorating the negative effects of insufficient power plants, thus fundamentally changing our utilisation of traditional centralised grids.^{1,2} ESSs can also compensate for the intermittency and variability of renewable energy sources and support their integration by providing power on demand and to schedule. The electrification of transportation represents an additional opportunity for utilising ESSs, minimising carbon footprints and reducing fossil fuel consumption. Thus, it is clear that the requirements of ESSs are diverse and more specialised than ever before. However, satisfying all these demands requires further advances and new combinations of different ESS technologies.³⁻⁶

Lithium-ion batteries (LIBs) based on non-aqueous electrolytes are currently at the forefront of ESS science owing to their

unmatched energy densities, energy efficiencies, and the vast range of power-to-energy ratios they can supply.^{7,8} The ongoing development of LIBs has realised commercially viable electric vehicles and a continuous reduction of cost, which fell below \$300 (kW h)⁻¹ for the first time in 2014.^{8,9} This cost reduction is primarily attributed to the economy of scale associated with increased production and engineering improvements at the cell and pack levels.^{8,9} Scaling up of contemporary LIBs for gridscale applications, however, still entails daunting challenges owing to the anticipated astronomic demand for Li and other critical elements such as Co, potentially leading to shortages much like those faced today with fossil fuels.¹⁰⁻¹² It may also entail races to exploit new reserves and geopolitical concerns owing to the uneven distribution of Li resources around the world.^{10,11} However, it is widely acknowledged that nonincremental improvements in battery performance parameters



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Fig. 1 Seawater and crust abundances of potential elements for use as charge carriers in secondary batteries and their theoretical gravimetric and volumetric capacities.³¹

including capacity, lifetime, affordability, safety, and fastcharging capability must be fulfilled at the active-material level because the performances of current LIBs are approaching their theoretical limits.¹² From the materials perspective, battery systems utilising earth-abundant elements as charge carriers and/or electrode materials have attracted considerable interest in recent years.^{13–22} Fig. 1 shows the crust and sea abundances of elements that can potentially act as charge carriers for secondary batteries, with each having its own advantages and disadvantages in scientific terms.

The chemical similarity of alkali metals allows the rational extrapolation of the huge body of knowledge pertaining to LIB technology accrued over the past decades, which is mainly applicable to the use and development of Na and potassium secondary batteries. For instance, dendrite-suppressing strategies developed for LIBs have been demonstrated to be applicable to



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His current research interests include fluorine compounds including ionic liquids and their derivatives as functional materials to be applied for energy storage and conversion. Na^{23-25} and K^{26} analogues. With respect to electrode materials, it is known that small alkali metal ions like Li⁺ tend to occupy octahedral sites, and larger ones prefer trigonal prismatic coordination.²⁷ The difference in preferred coordination and structural competition present an opportunity for exploring interesting new Na- or K-intercalation hosts.^{14,15} Rechargeable batteries fabricated using multivalent charge carriers (Mg,^{18,19} Ca,²⁰ Al,²¹ and Zn²²) have the potential to outperform current LIBs owing to their superior theoretical volumetric energy densities. However, despite several exciting advances achieved recently,^{28–30} research in this field remains at a nascent stage for the most part due to the lack of electrolytes that permit efficient metal deposition/dissolution combined with sufficient operating voltage windows and the lack of positive electrode materials with high capacities.^{18–21}

Na secondary batteries show enormous promise as candidates to meet the sustainability and performance requirements demanded by broad-scale applications for the following reasons:10 (i) Na is the most common alkali metal and the sixth most abundant element on Earth. The abundance of Na is approximately 24 000 ppm in Earth's crust (20 ppm for Li) and 10500 ppm in seawater (0.17 ppm for Li), making it a practically unlimited resource.³¹ The price of the Na-containing raw material typically used, Na₂CO₃, is therefore over 100-times lower than that of Li₂CO₃, and this gap is constantly expanding (Table S1, ESI[†]). (ii) Na is the lightest and smallest alkali metal next to Li, and it has a low redox potential (-2.71 V versus a)standard hydrogen electrode (SHE)), resulting in reasonably high cell voltages. (iii) The gravimetric capacity penalties ($\sim 15\%$)¹⁴ caused by the higher atomic mass of Na can be compensated for at the cell level by replacing the conventional and heavier Cu foil negative electrode current collector with an Al one, because Na does not alloy with Al.32 This replacement contributes to additional cost advantages for Na secondary batteries (USD 0.3 m⁻² and USD 1.2 m⁻² for Al and Cu foils, respectively).³³ (iv) The larger ionic radius of Na compared to that of Li leads to weaker solvation in polar solvents. Desolvation energetics are known to govern interfacial alkali metal transport, so this weaker interaction facilitates the intercalation of Na⁺.³⁴

Historically, ambient-temperature Na⁺ and Li⁺ intercalation chemistry was investigated in parallel in the 1970s,³⁵ but the higher energy density and subsequent commercial success of the latter led to research efforts being predominantly concentrated on LIBs.¹³ Furthermore, high-temperature Na-S batteries (along with the related Na-NiCl₂ batteries) employing Na metal negative electrodes and beta-alumina solid electrolytes (BASEs) have been explored worldwide.36,37 Both cells operate at 270-350 °C to ensure facile Na ion transport between the BASE and electrode materials and eliminate concerns over Na dendrite formation, as Na is molten in this temperature range (melting point, 98 °C). Since the 2000s, megawatt-hour scale Na-S cells have been widely used in Japan and other countries as a means to stabilise energy supply from renewables and thus improve supply reliability.^{36,37} Accordingly, Na secondary batteries do not fall into the category of "post LIBs" in the historical sense. Through intensive efforts in recent years and as a result of

leveraging knowledge of LIBs, phenomenal progress has been realised in the development of electrode and electrolyte materials for Na secondary batteries,^{38–45} which can now achieve comparable, or even better, performance characteristics than LIBs.^{46–50}

An electrolyte is an indispensable constituent of any battery, transferring and balancing charges in the form of ions between the electrodes.^{51,52} The electrolyte is positioned between, and in close interaction with, the positive and negative electrodes, and thus it has become increasingly recognised that electrolytes profoundly affect battery behaviour, instead of merely being a medium through which shuttle ions are transported. Of particular importance is the formation of a solid-electrolyte interphase (SEI) on the electrode surface upon decomposition of the electrolyte as the electrode works at a potential over the thermodynamic limit. Because every charge carrier must pass through this interphase to enter the electrode for electron transfer, the nature and thickness of the SEI largely dictates the reversibility and kinetics of the given electrode reactions.⁵² It has long been widely believed that such a passivation layer is indispensable, as once it is formed on the electrode, the electrolyte is prevented from continuous decomposition and the system is kinetically stabilised.53 The SEI concept was thoroughly elucidated for the successful operation of graphite, the universal negative electrode for LIBs, and it is now applied to Na analogues.

Organic-based electrolytes made from Na salts and organic solvents are most widely used for Na battery studies by virtue of their high ionic conductivity and good wettability, which parallel those featured in well-developed LIB technology. However, several studies have revealed that simply duplicating the strategies used in LIBs leads to SEIs with less homogeneity and inferior stability upon cycling.54-57 In addition, some established additives used in LIBs have been found to be ineffective or even detrimental to Na cell performance.⁵⁸ While a complete explanation for this is currently unavailable, the underlying factors are^{52,59} (i) differences in solvation structures and strengths, which affect the SEI evolution process; (ii) the higher redox potential (~ 0.3 V) of Na, which alleviates the reductive decomposition of the electrolyte; (iii) poorer polymerisation ability of Na; and (iv) more pronounced dissolution of SEI components due to the milder Lewis acidity of Na. The SEI properties are further complicated by the inherent reactivity of Na metal toward many electrolytes, highlighting the importance of choosing an appropriate electrolyte for the accurate evaluation of new electrode materials in half-cell configurations.^{52,59} Theoretical calculations of the ionic conductivity within the main SEI components have shown that Na ions must overcome a higher energy barrier to migrate across as compared to the case of Li ions.^{58,60,61} Based on this understanding, it is clear that Na batteries require their own optimised electrolytes, and the development of electrolytes that provide higher electrochemical stability could be key to solving SEI issues and thus realising high-performance Na batteries.

While LIBs function well for consumer electronics at room temperature, the demand for batteries that withstand and operate under harsh environmental conditions is growing.^{62,63}

The larger spectrum of applications expected for next-generation batteries includes all-climate electric cars, high-altitude drones, and robotics, as well as task-specific areas such as the aerospace industry, thermal reactors, oil and gas exploration, all of which call for batteries that can work safely over a wide temperature range and/or withstand dramatically dissimilar temperatures during charge and discharge. Apart from the flammability and volatility of organic electrolytes,^{8,63} their high susceptibility to thermal history and temperature fluctuation limits their use in these emerging applications.^{64,65} Indeed, they have been the cause of many battery faults and even explosions.⁶⁶ Other cell concepts, such as water-based technologies like lead-acid and nickel–metal hydride batteries, are also limited to operation between -20 and 55 °C.⁶⁷

For Na secondary batteries to be truly competent as a complementary technology or successor in some applications to LIBs, it is imperative to address the needs of these unprecedented sectors. Accordingly, alternative electrolytes are thought to be a key enabler for expanding the operational temperature window.^{62,63,68-74} The exploitation of electrolytes with greater chemical stabilities also has a further benefit, *i.e.*, the feasibility of Na metal negative electrodes. These have been intensively researched in the past few years as a means to achieve higher energy densities.^{25,73,74} Na metal outperforms other negative electrode candidates in terms of redox potential and specific capacity and plays an essential role in combination with many high-capacity but initially unsodiated positive electrode materials to form Na-S, Na-Se, Na-O2, and Na-CO2 batteries. Nevertheless, uncontrolled dendrite growth of Na metal upon cycling and continuous reforming of SEIs, which are interrelated, lead to rapid battery degradation and severe safety hazards.

Among the numerous candidates for next-generation electrolytes, ionic liquids (ILs), which are liquids consisting entirely of ions,⁷⁵⁻⁷⁸ have attracted considerable interest for energy storage devices owing to their numerous unique properties such as negligible vapour pressure, low flammability, high thermal and chemical stabilities, wide liquid-temperature range, broad electrochemical window, and intrinsic ionic conductivity. These properties provide a strong platform for developing batteries with additional functionalities and/or enhanced performances, *i.e.*, high-level safety and wide-temperature durability, for the above-mentioned large-scale battery systems and strategic applications. The absence of molecular species from IL electrolytes leads to elimination of parasitic reactions and their complicated interplay with shuttle ions and other cell components.52 This could contribute to lessening SEI formation and improving coulombic efficiency, which are prerequisites for developing long-lasting batteries. For example, an average coulombic efficiency greater than 99.95% is required to achieve reasonable capacity retention (>60%) over 1000 cycles. Furthermore, it has been demonstrated that Na metal and alloybased negative electrode materials can provide stable cycling in IL electrolytes but not in conventional electrolytes.^{79–81} Moreover, the tunable "solvent" properties of ILs allow the use of strategies to suppress the corrosion of Al current collectors and the dissolution of active materials. In conjunction with

Although ILs may initially appear to be an uneconomical choice of electrolyte, the amount of electrolyte required in a cell is limited (1-1.3-times the total pore volume of the composite electrodes and separator) and thus its contribution to the entire cost is not always significant.88,89 Furthermore, their cost is expected to decrease as more experience and knowledge pertaining to their synthesis, manufacture, and recycling are gained.^{10,90,91} It is also important to bear in mind that the enhanced durability and/or additional functionalities of a battery can, in terms of performance-to-cost ratio, compensate for or even eliminate cost concerns arising from the use of IL electrolytes. Fig. 2 shows a Ragone plot for various ESSs. It can be seen that Na secondary batteries with IL electrolytes simultaneously exhibit good specific energy (approaching that of LIBs) and power (exceeding that of a supercapacitor) at optimal temperatures, which is difficult for conventional devices.

Several reviews on the versatile roles of ILs in energy-related fields have been published.^{92–101} In this review, we focus on the development of IL electrolyte materials for Na secondary batteries.



Fig. 2 Ragone plot showing sodium secondary batteries with IL electrolytes^{164,180} in comparison with various energy storage systems (Li-ion, ^{428,429} Li-S, ^{430,431} Li-O₂, ^{432,433} Na-ion, ^{49,50} Na-S, ⁴³⁴ Al and Mg batteries, ^{29,435} and supercapacitors ^{436,437}). The operation temperature of each device is room temperature unless otherwise stated. Specific energy and power densities were calculated based on the weight of the positive electrode material.

The first section presents an introduction to the fundamental properties and design strategies of IL electrolytes. We then present a comprehensive summary on the negative and positive electrode materials investigated. By exploiting the thermal and chemical stabilities of ILs, the operational window of Na batteries has been effectively extended to the intermediate temperature range (90-150 °C), accompanied by greatly facilitated electrode kinetics without sacrificing stable cycleability. Elevated temperature operation has also been demonstrated to be beneficial for overcoming the electrochemical inactivities of certain electrode materials previously considered inapplicable under ambient conditions.^{102,103} In addition to the above-mentioned specific applications involving intermediate temperature range, recovering waste heat generated from a battery could be valid to create such an environment. The practical prospects and future challenges associated with the development of electrode materials for Na secondary batteries, the efficient utilisation of Na metal as a negative electrode material, and specific considerations pertaining to the SEI are also discussed.

2. IL electrolytes

ILs can comprise a wide variety of compounds, and thus the possible number of combinations of cation and anion for ILs is virtually countless.^{104–108} In this section, some performance-relevant properties of ILs as electrolytes for Na secondary batteries are summarised by sectionalising them into physico-chemical, thermal, and electrochemical properties. The handling of ILs, especially for electrochemical applications, is also summarised here. Comparisons with organic electrolytes or analogous Li systems will be frequently highlighted for a better understanding of Na systems.

Details on other ILs, including protic ILs, deep eutectic solvents, and task-specific ILs, all of which have great research and application importance but fall outside of the scope of this review, can be found elsewhere in the literature.^{93,104,107–110}

2.1 Ionic species for IL electrolytes

Fig. 3 summarises the ionic species used for ILs, especially those of interest for application to Na secondary batteries, along with their abbreviations, which will be used hereafter.79,84,105-107,111-135 Although a number of ILs appear to be of value for secondary batteries, the number of truly useful ILs for this purpose are actually quite limited. In early studies of ILs, fluorocomplex anions (PF_6^- and BF_4^-) were widely used to obtain salts with low melting points.136,137 PF6-based ILs are still extensively used for organic synthesis, but their low stability against hydrolysis and high viscosities prevent their use as electrolytes for modern ESSs. Although there are some examples of research into the hydrolytically more stable BF₄⁻ in Na secondary batteries, the synthesis of highly pure BF4-based ILs is highly expensive and time-consuming because of their hydrophilicity, and thus they are less attractive for practical use. Sulfonylamide-based (also known as sulfonylimide-based) ILs are widely applied as electrolytes owing to their facile synthesis (they are hydrophobic



Fig. 3 Ionic species in ILs of interest for sodium secondary battery applications. Abbreviations for specific ions are also shown. Cations: C_nC_1 im⁺ = 1-alkyl-3-methylimidazolium, C_nC_1 pyrr⁺ = *N*-alkyl-*N*-methylpyrrolidinium, C_nC_1 pip⁺ = *N*-alkyl-*N*-methylpiperidinium, N_{nnnn}^+ = tetraalkylammonium, AS[*m*·*n*]⁺ = azoniaspiro[*m*,*n*]nonane, and P_{nnnn}^+ = tetraalkylphosphonium. The symbols "*n*" and "*m*" in the abbreviations indicate the number of carbon atoms in the alkyl chain ("H" and "i4" denote hydrogen and isobutyl groups, respectively). The alkoxy group is described in the form of "*nOn*" (*e.g.* "(101)" means methoxymethyl group). Anions: BF₄⁻ = tetrafluoroborate, PF₆⁻ = hexafluorophosphate, N(CN)₂⁻ = dicyanamide, FSA⁻ = bis(fluorosulfour)amide, and TFSA⁻ = bis(trifluoromethylsulfonyl)amide.

in most cases) and relatively high ionic conductivities (*e.g.*, 8.8 mS cm⁻¹ for [C₂C₁im][TFSA]¹³⁸ and 15.4 mS cm⁻¹ for [C₂C₁im][FSA]^{139,140}).

The cationic species in ILs can be varied more than the anionic species by extending their organic frames. Although melting points typically decrease upon the introduction of bulky or long substituent groups, viscosity and ionic conductivity usually increases and decreases, respectively. Thus, asymmetric cations with short alkyl chains, such as C_3C_1 pyrr⁺ and C_2C_1 im⁺, are generally preferred. Non-aromatic pyrrolidinium-based cations are reductively more stable than imidazolium-based cations, and thus widely utilised for secondary battery applications.¹⁴¹ In a recent study, the physicochemical properties of a range of cationic structures (ammonium, pyrrolidinium, piperidinium, and morpholinium) in the forms of FSA- and TFSA-based ILs

were systematically studied, revealing the preferable features of ether-functionalised cations.¹³¹

Two additional categories are briefly considered here; inorganic ILs and solvate ILs. Inorganic ILs are also electrochemically attractive media at elevated temperatures. In a series of Na[TFSA]–M[TFSA] ($M = Li^+, K^+, Rb^+$, and Cs^+) binary systems, Na[TFSA]–Cs[TFSA] exhibited the lowest eutectic temperature of 110 °C at a Na[TFSA]/M[TFSA] molar ratio of 0.07:0.93.¹⁴² Introduction of FSA⁻ further reduces the eutectic temperatures to between 52 and 76 °C with the FSA⁻ salts of Li⁺, K⁺, Rb⁺, and Cs⁺.¹⁴³ This system realises even lower eutectic temperatures approaching ambient conditions upon introduction of a third component to form a tertiary system. The lowest eutectic temperature of 36 °C was observed for the Na[FSA]/K[FSA]/Cs[FSA] system at a molar ratio of 40:25:35.¹⁴⁴

Solvate ILs are another category in this field,^{95,145,146} although the number of solvate ILs investigated for Na secondary batteries is currently limited. Solvate ILs are formed by Lewis basic species such as oligoethers interacting with metal salts at equimolar concentrations. A representative example is [Li(G3)][TFSA] (G3 = triethylene glycol dimethyl ether, also called triglyme) which has a melting point of 23 °C.¹⁴⁷ The glyme molecules in such solvate ILs coordinate to the metal centre and their behaviour is different from those of pure glymes or diluted solutions, with the coordinating glymes having considerably higher oxidation resistances and thermal stabilities than free glymes. The Na⁺ analogues [Na(G3)][TFSA], [Na(G4)][TFSA], and [Na(G5)][TFSA] (G4 = tetraglyme and G5 = pentaglyme) are not room-temperature ILs,¹⁴⁸ but their electrochemical application to Na secondary batteries has been studied in both neat and diluted forms with several electrode materials.149

2.2 Thermal properties

The thermal properties of IL electrolytes are typically investigated using thermogravimetry (TG) and differential scanning calorimetry (DSC). In general, ILs have higher thermal stabilities than typical organic solvents,^{141,150} but the determination of IL thermal decomposition temperatures by TG requires special attention because the results are highly dependent on the atmosphere, cell material, scan rate, sample amount, impurities, and method. This is related to the fact that the measured thermal decomposition of ILs is often kinetically controlled (see details on short-term and long-term stability in a previous review¹⁵⁰). Thus, comparisons of thermal decomposition temperatures determined under different conditions are usually meaningless. In contrast to organic solvents, which boil at relatively low temperatures, ILs can extend the upper operating temperatures of batteries. In general, ILs are less stable in air than in inert atmospheres, as water and O₂ gas facilitate decomposition.¹⁵⁰

Systematic studies on the thermal decomposition temperatures of ILs containing Na salts have revealed that the addition of Na[BF₄], Na[ClO₄], Na[N(CN)₂], and Na[TFSA] salts to [C₄C₁pyrr][TFSA] IL somewhat decreases the decomposition temperature to 330–360 °C (~400 °C for the neat salt).^{119,127} A clearly different case is that for ILs supplemented with Na[PF₆], which decompose at a lower temperature than the others

(weight loss starts around 100 °C). The Na[BF₄]-[C₂C₁im][BF₄] IL system does not ignite even upon contact with a flame and is thermally stable up to 380 °C regardless of $Na[BF_4]$ concentration (0.10–0.75 mol dm⁻³).¹¹⁸ The thermal stabilities of FSA-salts are somewhat controversial as they are unstable against hydrolysis, even for a trace amount of water. Slow decomposition rates sometimes enable observation of melting, even when the melting point is higher than the decomposition temperature (e.g. T_d = 70 °C and T_m = 130 °C for neat Li[FSA]).¹⁴³ A systematic thermogravimetric study has shown that the thermal decomposition temperatures of alkali metal FSA salts increase with increasing size of the alkali metal cation.¹⁴³ The thermal decomposition temperatures of some FSA-based ILs, such as [N4411] [FSA], [AS(4.5)] [FSA], and [N6111] [FSA], containing Na[FSA] are dominated by that of Na[FSA] (~130 °C).¹²² Another study revealed the higher thermal decomposition temperatures but essentially the same trend (493 °C for neat $[N_{2(20201)3}]$ [FSA] and 271 °C for Na[FSA]- $[N_{2(20201)3}]$ [FSA] with 55 mol% Na[FSA]).

Furthermore, the decomposition temperature in the presence of the electrode material is more important in view of battery applications. Electrode materials in the charged state are highly reductive (negative electrode) or oxidative (positive electrode) and thus their contact with electrolytes at high temperature can cause thermal runaway. For example, 1.0 mol dm^{-3} Li[PF₆]-EC/ DMC exothermically decomposes in the presence of Li metal and a charged positive electrode, $Li_{1-x}CoO_2$, whereas 0.32 mol kg⁻¹ Li[TFSA]-[C₃C₁pip][TFSA] exhibits negligible heat evolution under the same conditions, demonstrating the advantage of IL electrolytes in terms of thermal stability.¹⁵¹ Although few studies on the thermal properties of ILs for Na secondary batteries are available, some studies on organic electrolytes with various electrode materials (hard carbon (HC),152-155 $Na_xSn_{,}^{156} Na_{0.5}Ni_{0.5}Mn_{0.5}O_2,^{157} Na_{0.35}CrO_2,^{158} Na_{0.58}FeO_2,^{159}$ revealed the high reactivity of charged electrodes toward organic electrolytes (and that the presence of Na salts affects decomposition temperature). The thermal stability of a $Na_3V_2(PO_4)_3/$ $Na_3V_2(PO_4)_3$ symmetric cell with a $Na[BF_4]-[C_2C_1im][BF_4]$ IL electrolyte was compared with that with a 1 M Na[ClO_4]-PC electrolyte after charging,87 and the resulting DSC curves demonstrated the high stability of the IL system. The thermal stability of electrolytes is also related to the stability of the SEI, and the formation of a high-stability SEI at elevated temperatures was demonstrated in Li-containing ILs.¹⁶⁰ These factors must be taken into account in future studies on the application of IL electrolytes for Na secondary batteries.156

Fig. 4 shows a phase diagram for a Na[FSA]–[C₃C₁pyrr][FSA] system.¹¹⁶ Mixing of a Na salt with organic salts usually leads to a certain type of eutectic system (or one that appears to be eutectic) such as Na[FSA]–[C₂C₁im][FSA],¹²⁴ Na[FSA]–[N₄₄₁₁][FSA],¹²² Na[FSA]–[As(4.5)][FSA],¹²² Na[FSA]–[C₃C₁pyrr][FSA],¹³⁰ Na[TFSA]–[C₄C₁im][TFSA],¹¹⁷ Na[TFSA]–[C₄C₁pyrr][TFSA],¹¹⁷ Na[TFSA]–[C₄C₁pyrr][TFSA],¹¹⁷ Na[TFSA]–[C₄C₁im][TFSA],¹¹⁷ and Na[TFSA]–[C₂C₁im][TFSA] ($0 \le$ Na[FSA] ≤ 50 mol%) contrasts with that of Na[TFSA]–(C₂C₁im][TFSA] at room temperature ($0 \le$ Na[TFSA] $\le \sim 20$ mol%), indicating the



Fig. 4 Phase diagram for the Na[FSA]–[C₃C₁pyrr][FSA] system. $T_{m_o(1)}$ and $T_{m_e(1)}$ are the melting onset and final temperatures, respectively; $T_{m_o(2)}$ and $T_{m_o(2)}$ are the melting onset and final temperatures for the metastable state, respectively; T_g is the glass transition temperature. T_{tr_o} and T_{tr_e} are the onset and final temperatures for the solid-solid phase transition. The scale at the top is the concentration of Na[FSA] at 25 °C. Reproduced with permission.¹¹⁶ Copyright 2015, American Chemical Society.

ability of FSA⁻ to provide low melting points. In such diagrams, the drop in the liquidus line is often steep, which is partly explained by the existence of a crystallinity gap, where only the glass transition is observed and melting behaviour disappears (see the $0.2 \le x(Na[FSA]) \le 0.5$ range in Fig. 4). The origin of this crystallinity gap is not yet fully understood; it may be either thermodynamically or kinetically stable. However, long-term ageing at low temperatures (in the liquid range) often does not lead to crystallisation of ILs.¹⁶¹ ILs are structurally frozen into a glassy state at the glass transition temperature, and the change in heat capacity can be observed by DSC analysis. The glass transition temperature is correlated with the fluidity (i.e., the reciprocal of viscosity) of ILs,^{162,163} and thus fluid ILs tend to have low glass transition temperatures. The glass transition temperatures in IL systems containing Na-salts typically increase as the amount of Na salt is increased, as shown in Fig. 4. This is related to the low fluidity observed at high Na[FSA] molar ratios. An increase in glass transition temperature tends to be observed upon increasing the organic cation size (e.g. -77.3 °C vs. -75.8 °C for Na[TFSA]-[C₂C₁im][TFSA]¹¹⁷ and Na[TFSA]-[C4C1im][TFSA] (20 mol% Na[TFSA]]) or replacing FSA⁻ with TFSA⁻ (-91 °C for Na[FSA]-[C₂C₁im][FSA]¹²⁴ ν s. -77.3 °C for Na[TFSA]– $[C_2C_1im]$ [TFSA]¹¹⁷ 20 mol% Na salt).

Fig. 5 summarises the verified temperature range of IL electrolytes for Na secondary batteries (the highest limit of each IL system is not its thermal decomposition temperature) and the operating temperature ranges of various applications using secondary batteries. The available temperature range of



Fig. 5 Temperature range of ILs verified for sodium secondary batteries. The highest limit of each IL system does not indicate its thermal decomposition temperature. The operating temperature ranges of various applications and commercial LIBs are included for comparison. See Fig. 3 for abbreviations of each ionic species. ^{62,63,87,127,128,149,164,166,237,268,289,316,321,438}

commercial LIBs is included for comparison. Although several studies on FSA-based ILs have explored low-temperature operation $(-30 \degree C)$, ^{164–166} most works on Na secondary batteries were performed at room temperature (25 °C). However, several studies have been performed above room temperature, with many cases concerning electrochemical tests above the melting point of the IL or improving the performances of electrode materials that do not work well at room temperature. Intermediate-temperature operation (up to 150 °C) has also been attempted with ILs, exploiting the use of waste heat and hot environments, in order to enhance Na⁺ ion transport and electrode reactions.^{63,164,167} For example, an increase in temperature from 25 to 85 °C leads to an increase in ionic conductivity from 1.9 to 17.3 mS cm^{-1} for Na[FSA]–[C₃C₁pyrr][FSA] (30 mol% Na[FSA]). Further details of performance at intermediate temperatures will be presented in the following sections on positive and negative electrodes.

2.3 Physicochemical properties

Fig. 6 summarises the molar concentration dependency of viscosity, ionic conductivity, and Na⁺ ion transport number for selected ILs used for Na secondary batteries (see Table S2, ESI[†]). In general, ILs exhibit higher viscosities than organic electrolytes at similar Na contents. This arises from the strong coulombic interactions between cations and anions, and thus dilution of ILs with a solvent leads to a decrease in viscosity.^{168–170} Statistical data have shown that an increase in ion volume increases the viscosity of ILs,¹⁷¹ whereas the introduction of ether groups tends to decrease viscosity.^{131,172} High viscosity is physically not preferable as it makes the handling of IL electrolytes, such as their impregnation into composite electrodes and separators, difficult.

In addition, viscosity (η) is related to molar ionic conductivity (λ) *via* the Walden rule (or fractional Walden rule with the α parameter between zero and unity corresponding to the slope of the Walden plot (eqn (1)) (log(η^{-1}) *vs.* log(λ))):^{162,173–175}

$$\eta \cdot \lambda = \text{const.} \text{ (or } \eta^{\alpha} \cdot \lambda = \text{const.} \text{)}$$
 (1)

The Walden rule is generally held for both pure ILs and ILs containing alkali metal cations. The α value indicates the ratio of B parameters in the Vogel-Tammann-Fulcher (VTF) equation for viscosity and molar ionic conductivity, the details of which will be discussed later. This implies that addition of a Na salt to organic ILs decreases the ionic conductivity (σ) of the resulting inorganic-organic hybrid ILs because the viscosity increases upon adding the Na salt owing to increased coulombic interactions (Fig. 6a and b). This is in contrast to cases of organic solutions that exhibit a maximum ionic conductivity at a certain Na molar concentration.¹⁷⁶ Although this is not directly regarded as a disadvantage in terms of Na⁺ ion transport, the bulk resistance, which always exists during battery operation, certainly increases by adding a Na salt. Although the ionic conductivities of Na-containing ILs are highly dependent on the ionic structures, the values for common ILs are comparable to those for organic solutions at similar concentrations (e.g., 3.6 mS cm⁻¹ for Na[FSA]–[C₃C₁pyrr][FSA] at 0.98 mol dm⁻³,¹¹⁶ 3.9 mS cm⁻¹ for Na[TFSA]–[C₂C₁im][TFSA] at 0.70 mol dm⁻³,¹¹⁷ 8.5 mS cm⁻¹ for Na[FSA]–[C₂C₁im][FSA] at 1.1 mol dm⁻³,¹²⁴ 11.8 mS cm⁻¹ for Na[BF₄]-[C₂C₁im][BF₄] at 0.75 mol dm⁻³,¹¹⁸ 7.98 mS cm⁻¹ for Na[PF₆]-PC at 1.0 mol dm^{-3} ,¹⁵² and 6.2 mS cm⁻¹ for Na[TFSA]-PC at 1.0 mol dm⁻³¹⁵²). Na ion



Fig. 6 Dependency of (a) viscosity, (b) ionic conductivity, and (c) Na⁺ ion transport number on Na salt concentration for IL electrolytes used in sodium secondary batteries. See Tables S2 and S3 (ESI[†]) for the data represented here.^{79,84,116–124} Other related data are also available in the literature.^{125–135}

transport in the steady state is governed by Na ion conductivity (σ_{Na^+}) , as expressed in eqn (2):

$$\sigma_{\mathbf{N}\mathbf{a}^+} = \sigma \cdot t_{\mathbf{N}\mathbf{a}^+} \tag{2}$$

where t_{Na^+} is the transport number of Na⁺ formulated as follows (eqn (3)) and determined by the AC–DC method.^{177,178}

$$t_i = \frac{|z_i|c_i\mu_i}{\sum\limits_i |z_i|c_i\mu_i} \left(\sum\limits_i t_i = 1\right)$$
(3)

It should be noted that t_i is not simply derived from the ratio of mobilities. Such a simple form can be applied in limited cases such as that of a 1:1 single salt in a solution (*e.g.*, NaPF₆ in propylene carbonate). However, IL electrolytes for Na secondary batteries containing organic cations are usually composed of more than three constituent ions, and thus concentration has to be taken into account. The simple ratio of the diffusion coefficients of constituent ions does not always indicate the transport number in ILs as ILs are highly concentrated systems (*i.e.*, not infinitely diluted systems) and the frame of reference has to be carefully considered.¹⁷⁵ The term 'apparent transport number' is also used in several reports, and is derived by considering the different environments of Na⁺ in ILs in the initial and final states during polarisation.^{116,179,180}

The t_{Na^+} values for several Na⁺-containing ILs have been reported (see Fig. 6c for Na salt concentration dependency)^{79,84,116,119,125,126,135,180} and tend to increase with increasing Na salt concentration in the Na[FSA]–[C₃C₁pyrr][FSA]^{79,116} and Na[FSA]–[C₂C₁im][FSA]¹⁸⁰ systems (*e.g.*, 0.13 (10 mol% Na[FSA]) to 0.35 (50 mol% Na[FSA]) for Na[FSA]–[C₂C₁im][FSA]). The Na⁺ ion transport number also clearly increases with increasing temperature. The resulting Na⁺ ion conductivity exhibits a maximum at a certain Na[FSA] fraction (*e.g.* at 20 mol% Na[FSA] for Na[FSA]–[C₃C₁pyrr][FSA] and 30 mol% Na[FSA] for Na[FSA]–[C₂C₁im][FSA]).¹⁸⁰ Molecular dynamics simulations have indicated that clustering of Na⁺ and FSA⁻ ions occurs at high Na[FSA] concentrations,^{79,181} enabling site exchange and/or indicates a structural diffusion mechanism for Na⁺, leading to high transport numbers. Clustering of Na⁺ and FSA⁻ ions and rapid exchange of the Na⁺ ion between different coordination environments have also been suggested in this and analogous IL systems by Raman spectroscopic,¹⁸⁰ NMR spectroscopic,^{79,182} and atomic force microscopic¹⁸² studies.

The fluidity (η^{-1}) of an IL usually deviates from Arrhenius behaviour as it approaches the glass transition temperature (T_g) , and the rate at which the transport properties change with temperature near the glass transition is termed 'fragility'.¹⁶² In the same way, such temperature dependence of viscosity and ionic conductivity for ILs is best fitted with the VTF equations as follows (eqn (4) and (5)).^{183–186} Inorganic–organic hybrid IL systems including Na⁺-containing ILs also follow this behaviour.^{116,122,124,125,130}

$$\eta(T) = A_{\eta} \sqrt{T} \exp\left(\frac{B_{\eta}}{T - T_{0\eta}}\right)$$
(4)

$$\sigma(T) = \frac{A_{\sigma}}{\sqrt{T}} \exp\left(-\frac{B_{\sigma}}{T - T_{0\sigma}}\right)$$
(5)

The A_{η} , B_{η} , $T_{0\eta}$, A_{σ} , B_{σ} , and $T_{0\sigma}$ fitting parameters are determined by mathematical fitting.

The experimentally observed glass transition temperature is always higher than the ideal glass transition temperature $(T_g > T_0$, where T_0 is the ideal glass transition temperature in the VTF equation) by the empirical approximation $T_0/T_g \approx 0.75$.¹⁸⁷

The polarity of a solvent is regarded as an indicator of its ability to dissolve and stabilise ions and is briefly mentioned here because it is an important concept in the design of electrolytes based on ILs for secondary batteries. Polarity is usually discussed based on physical parameters such as relative permittivity, dipole moment, and refractive index. Dielectric spectroscopy has revealed that the relative permittivities of ILs containing BF_4^- , PF_6^- , $SO_3CF_3^-$, and $TFSA^-$ fall between 11 and $15.^{188-191}$ However, such macroscopic parameters are not always effective for assessing the polarity of ILs because the solubility of inorganic salts is largely different (even though the relative permittivities of the ILs above are similar).

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ILs are not homogeneous dielectrics and local interactions apparently play more important roles. Polarity based on local interactions is evaluated by spectroscopic methods using solvatochromic dyes.^{192–197} The empirical $E_{\rm T}(30)$ and $\lambda_{\rm Cu}$ parameters are determined by the π - π * absorption band of Reichardt's betaine dye and the d-d absorption band of Cu²⁺ in [Cu(acac)(tmen)] (acac = acetylacetone and tmen = N,N,N',N'tetramethylethylenediamine), respectively. The $E_{\rm T}(30)$ parameter exhibits good correlation with the acceptor number of solvents and generally decreases in ILs as the alkyl chain length on the cation increases and as the form of the cation changes from aromatic (imidazolium and pyridinium) to non-aromatic (pyrrolidinium and tetraalkylammonium).¹⁹⁸⁻²⁰³ Anionic structures have a minor effect on the acceptor number of cations and accordingly $E_{\rm T}(30)$. Conversely, the $\lambda_{\rm Cu}$ parameter correlates with the donor number of solvents and is thus affected by the anionic structures in ILs. The resulting donor ability has the trend $[CF_3CO_2]^- > [SO_3CF_3]^- > [FSA]^- \approx [TFSA]^- > [BF_4]^- >$ [PF₆]⁻.^{204,205} There are other indicators known as Kamlet–Taft parameters (π^* , α , and $\beta^{206-209}$) that are widely applied in discussing the polarity of ILs.^{197,202,205} The π^* parameter is related to the dipolarity/polarisability of solvents, and those for ILs are generally higher than those for typical aprotic organic solvents and increase from nonaromatic to aromatic cationbased ILs. TFSA-based ILs have smaller π^* parameter values compared to those of other ILs. The α and β parameters correlate with hydrogen bonding donor and acceptor abilities, respectively, but are less important for electrolytes for secondary batteries. The correlation of polarity and the molar conductivity ratio, $\Lambda_{\rm imp}/\Lambda_{\rm NMR}$, has been discussed in previous works.^{205,210} $\Lambda_{\rm imp}$ and $\Lambda_{\rm NMR}$ are the molar conductivities obtained by AC impedance and NMR spectroscopies, respectively. This correlation provides basic insight for considering ion structures in IL electrolytes for secondary batteries because the $\Lambda_{\rm imp}/\Lambda_{\rm NMR}$ ratio is the proportion of ions (charged species) that contribute to ionic conduction from all the diffusing species on the time scale of the measurement.²¹⁰

The $\lambda_{\rm Cu}$ parameter, which is predominantly affected by the anionic structure, exhibits good correlation with the $\Lambda_{\rm imp}/\Lambda_{\rm NMR}$ ratio; an increase in the donor ability of the anion decreases the number of ions contributing to ion conduction. The $E_{\rm T}(30)$ values show a somewhat more complicated correlation with the $\Lambda_{\rm imp}/\Lambda_{\rm NMR}$ ratio. The major dependency is due to the difference in the cation backbone structures for TFSA-based ILs, and the $\Lambda_{\rm imp}/\Lambda_{\rm NMR}$ ratio increases with decreasing $E_{\rm T}(30)$, *i.e.*, decreasing the acceptor number of cations. Na ions in ILs are surrounded by the anions therein, and thus polarity values based on anions, such as those of $\lambda_{\rm Cu}$, are considered to reflect IL behaviour. However, direct evidence of correlation between such parameters and Na⁺ ion properties have not been adequately investigated from static and dynamic viewpoints. Thus, further work in the future, including computational studies, is required in this field.

2.4 Electrochemical properties

The term 'electrochemical window' is used to indicate the potential range from the reductive to oxidative limits of electrolytes and is one of the most important properties of electrolytes for secondary batteries. It is theoretically limited by the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) (see the literature²¹¹ on the correct use of these terms) of the molecule constituting the electrolyte, but is very often kinetically dominated in practical situations, as is clearly demonstrated by the importance of the SEI.^{45,212} In terms of Na secondary batteries, an electrochemical window from the Na⁺/Na redox potential up to ~5 V is ideal for operation with high energy density, and many ILs containing BF_4^- , TFSA⁻, or FSA⁻ satisfy this demand (or provide windows close to this).

Fig. 7 shows the electrochemical windows of several ILs. A study combining molecular dynamics (MD) simulations and density functional theory (DFT) calculations indicated that the electrochemical stabilities of ILs should be discussed based on the stabilities of ions combined with counter ions in a certain structural model. Thus, considering the stability of individual ions in a vacuum is not always sufficient when considering the electrochemical stabilities of ILs (Fig. 7a).²¹³ This study also indicated the higher instability to reduction of TFSA⁻ compared to that of $C_3C_1pyrr^+$, and this has been experimentally confirmed.²¹⁴ Thermodynamic assessment based on DFT data has demonstrated that the electrochemical window is independent of the alkyl chain length for 1-alkyl-3-methylimidazolium cations and that BF_4^- and PF_6^- complex anions provide more electrochemically stable ILs than TfO or TFSA anions.²¹⁵

X-ray photoelectron spectroscopy (XPS) can provide information for discussing the oxidative and reductive resistances of ILs. The direct comparison of energy gaps revealed by XPS and voltammetry requires special care owing to differences in the properties of the bulk material and the electrode surface as well as differences in the Madelung potentials of the bulk and free surface.²¹⁶ A study combining ultraviolet photoemission, inverse photoemission, and near-edge X-ray absorption fine structure spectroscopy indicated that mainly cations contribute to the top of the occupied states and the bottom of the unoccupied states of [C₂C₁im][BF₄] and [C₂C₁im][PF₆].²¹⁷ Additional studies have confirmed that the molecular orbital energies of imidazolium-based ILs with [BF₄]⁻, [PF₆]⁻, [TfO]⁻, and [TFSA]⁻ are significantly affected by the electrostatic Madelung potential among the ions. For ILs with $[BF_4]^-$ and $[PF_6]^-$, both the highest occupied and lowest unoccupied states are derived from the imidazolium cation, i.e., the cation determines the band gap. Conversely, the highest occupied states for ILs with [TfO]⁻ and [TFSA]⁻ are determined by contributions from both the cation and anion.^{218,219}

Cyclic and linear sweep voltammetry have revealed that pure ILs with typical fluoroanions have electrochemical windows of 4 to 5 V (see Fig. 7b for the relationship between electrochemical windows for typical ILs and the average discharge potentials of selected electrode materials for Na secondary batteries^{107,111,141,220,221}). Although the three-electrode cell is a common setup for this type of measurement (typically, Pt, W, Cu, Al, and glass-like carbon are used as working electrode materials), two-electrode cells (usually coin-type cells) are often



Fig. 7 (a) Electrochemical windows for selected ILs as derived by a combination of MD and DFT calculations. Solid green bars indicate the potential range in which the IL is stable. The calculated Li metal Fermi level is also indicated. The 95% confidence interval for each limit is given by the black dumbbells. The potential scale is relative to the vacuum level. Reproduced with permission.²¹³ Copyright 2011, American Chemical Society. (b) Electrochemical windows of typical ILs used for sodium secondary batteries.

used for ILs containing shuttle ions such as Li⁺ and Na⁺ combined with Li and Na metal counter electrodes. The limit potentials for pure ILs are often compared with respect to the Ag⁺/Ag couple or more preferably the Fc⁺/Fc couple, as based on IUPAC recommendations.²²² In the cases of Li⁺- or Na⁺containing ILs, the electrochemical windows are usually discussed based on the Li⁺/Li or Na⁺/Na redox couple. Special attention has to be paid for the two-electrode cell because of the possible large polarisation on the Li or Na metal counter electrodes, which causes deviation from the Li⁺/Li or Na⁺/Na redox potential (see below). The addition of Li or Na salts to ILs often extends their reductive limits, regardless of the anion structure.121,223-226 Although aromatic cations including alkylimidazolium and alkylpyridinium suffer from instability to reduction in pure IL forms compared to nonaromatic- or linear alkyl-ammonium cations, the use of FSA⁻ as a counter anion greatly improves stability, even for aromatic cases; the difference due to the cationic structure is minimised for FSA-based ILs. This phenomenon has been interpreted in terms of both differences in SEI properties and the double-layer structure model. The former is based on the observation that FSA⁻ and TFSA⁻ reductively produce a radical anion, and the radical anion from FSA⁻ has unusually low reactivity to form a stable surface film.²²⁷ The latter is based on a specific double-layer model where FSA⁻ acts as an anchor to pull Li⁺ toward and exclude organic cations from the interface.^{224,226}

Residual water in ILs can have a significant effect on the electrochemical window.²²¹ A significant decrease in the electrochemical window (over 1 V) is observed with increasing water content (15, 58, and 273 ppm) for a Pt electrode in a $[N_{3111}]$ [TFSA] IL, but not for a glass-like carbon electrode. Practical active electrode materials also produce different behaviours compared to Na and Li metal electrodes; both cathodic and anodic decomposition of ILs is enhanced in the presence of certain electrode materials such as nano-sized active materials, especially graphitised carbon, and at elevated temperatures.^{103,228}

An interesting behaviour of IL electrolytes is the low corrosivity of Al electrodes at high potentials. Unlike PF₆-based electrolytes, which instantaneously form a fluorine-containing passivation film on Al,²²⁹ TFSA⁻ and FSA⁻ corrode Al current collectors in organic solutions.^{140,230–233} This is detrimental for practical applications because Al is commonly used as a current collector. However, Al corrosion is minimised in Li-based IL electrolytes.^{124,234–236} The same behaviour has also been observed in Na systems; Al corrosion also occurs in 1 mol dm⁻³ Na[TFSA]-PC solution,¹⁵² but not in FSA- or TFSA-based ILs,^{237,238} indicating that, in ILs, a stable passivation film is formed on Al electrodes at high potentials.

Na metal deposition–dissolution behaviour is another factor to consider when assessing the suitability of electrolytes for Na secondary batteries. In addition to the general importance of the behaviour of negative electrodes,⁷⁴ this consideration is important for the possible use of Na metal as a negative electrode or counter electrode in two-electrode cells. However, the large polarisation of Na metal sometimes makes the accurate evaluation of target electrodes difficult when using a half-cell configuration.^{86,124,239–242} Fig. 8 summarises some information on Na metal deposition–dissolution.

The reaction of Na metal in $[C_2C_1im][FSA]$ and $[C_2C_1im][TFSA]$ ILs has been investigated.¹²¹ Fig. 8a shows the appearances of $[C_2C_1im][FSA]$ and $[C_2C_1im][TFSA]$ ILs before and after immersion of Na metal for four weeks. The transparent colourless $[C_2C_1im][TFSA]$ IL turns brown after four weeks, whereas $[C_2C_1im][FSA]$ shows no change, suggesting the high effectiveness of FSA⁻ against reduction. Just 10 mol% FSA⁻ is effective for suppressing the decomposition of ILs. The continuous increase of interfacial resistance for a TFSA⁻ system has been studied by impedance spectroscopy. As discussed above, both the SEI layer and double-layer structure can contribute to the differences between FSA⁻ and TFSA⁻,^{121,223-226} but the time dependence of interfacial resistance seems to correlate with the former in this case. The presence of Na[FSA] is also a key factor for stabilising interfacial resistance.

Table 1 lists selected data for Na metal deposition–dissolution tests, mainly summarising coulombic efficiency (as obtained by cyclic voltammetry and repetitive DC polarisation tests), cycle dependence, and water effects. Cyclic voltammetry tests provide information on Na metal deposition–dissolution behaviour.



Fig. 8 (a) Photographs of ILs (top) before and (bottom) after Na metal immersion for four weeks. (A) Na[FSA]– $[C_2C_1im][FSA]$ (10:90 in mol) and (B) Na[TFSA]– $[C_2C_1im][TFSA]$ (10:90 in mol). Reproduced with permission.¹²¹ Copyright 2016, American Chemical Society. (b) Photographs of Na metal electrodes after alternate 50-cycle Na deposition–dissolution at 0.2 mA cm⁻² in Na[FSA]– $[C_2C_1im][FSA]$ (30 mol% Na[FSA]) at 0, 25, and 90 °C. Reproduced with permission.²⁴⁶ Copyright 2016, The Electrochemical Society. (c) Na/Na symmetric cell polarisation profiles for the Na[FSA]– $[C_3C_1pyrr][FSA]$ IL at 25 °C (5 mol% Na[FSA], 25 mol% Na[FSA], and 50 mol% Na[FSA]) and a summary of overpotential trends for this system. Reproduced with permission.⁷⁹ Copyright 2016, American Chemical Society.

Repetitive cathodic and anodic polarisation in Na/Na or Al/Na cells with Na metal deposited on the Al is also used to observe long-term behaviour. Na metal dissolution does not occur from deposited Na metal in alkylimidazolium-based ILs without FSA⁻.^{121,243} However, it is observed in alkylpyrrolidinium, tetraalkylammonium-, and tetraalkylphosphonium-based ILs, even those without FSA⁻, indicating the strong influence of the cation structure on Na metal deposition–dissolution behaviour.^{132,243} In the presence of FSA⁻, high coulombic efficiencies are observed in most cases.^{79,121,122,124,125,130,135,168,169,244} An increase in measurement temperature typically improves the coulombic efficiencies of Na metal deposition–dissolution.^{122,124,245} The higher coulombic efficiencies at 90 °C are considered to arise from the suppression of dendritic Na metal deposition at temperatures near the melting

point of Na metal (98 °C) because diffusion of the Na atoms at the metal surface becomes faster. This is clearly demonstrated by the different appearances of the Na/Na symmetric cell deposition–dissolution test beaker cells operated at 0, 25, and 90 °C shown in Fig. 8b.²⁴⁶ After 50-cycle deposition–dissolution at 0.2 mA cm⁻², dendrites are clearly observed on the Na metal electrode surfaces operated at 0 and 25 °C, whereas those operated at 90 °C maintain their shiny flat appearances and show no dendrite formation.

The effects of Na salt concentration on Na metal deposition– dissolution behaviour have been investigated in Na[FSA]– $[C_3C_1pyrr]$ [FSA] IL electrolyte.⁷⁹ Higher salt concentrations result in lower interfacial impedance, leading to a lower polarisation potential and enabling relatively high current charging (Fig. 8c). This study concluded that the interfacial layers formed

Table 1 Selected results of Na metal deposition-dissolution tests in ILs^a

IL	Na salt concentration, coulombic efficiency, and some comments	Ref
Na[TFSA]–[C ₄ C ₁ pyrr][TFSA]	0.1 mol dm ⁻³ , $\eta^{\rm CV}$ = 75%	243
	0.4 mol dm^{-3} , $n^{\text{CV}} \approx 30\%$	425
Na[TFSA]–[N ₂₁₃₃][TFSA]	0.1 mol dm ⁻³ , reversible in CV	243
Na[TFSA]–[N ₆₂₂₂][TFSA]	0.1 mol dm ⁻³ , $\eta^{CV} = 57\%$	243
Na[TFSA]-[N ₁₍₂₀₂₀₁₎₃][TFSA]	2.0 mol kg ⁻¹ , at 50 and 70 °C; reversible in CV at 80 °C 10 mol%, $\eta^{CV} \approx 0\%$	120
Na[TFSA]–[C ₂ C ₁ im][TFSA]	$10 \text{ mol}\%, \eta^{CV} \approx 0\%$	121
Na[TFSA]–[C ₄ C ₁ im][TFSA]	$0.1 \text{ mol } dm^{-3}, \eta^{CV} \approx 0\%$	243
Na[TFSA]–[C ₄ C ₁ C ₁ im][TFSA]	0.1 mol dm ⁻³ , $\eta^{\circ} \approx 0\%$	243
Na[TFSA]–[C ₄ C ₁ im][SO ₃ CF ₃]	0.1 mol dm ⁻³ , $\eta^{\rm CV} \approx 0\%$	243
Na[TFSA]–[C ₃ C ₁ pyrr][FSA]	0.25, 0.75, and 1.35 mol kg ⁻¹ , $\eta^{CV} \approx 40-60\%$; η^{CV} decreases to ~30% after 5 cycles	130
Na[FSA]-[C ₃ C ₁ pyrr][FSA]	20 mol%; η^{DC} = 94% at 80 °C; reversible in CV at 80 °C	245
	5–50 mol%, reversible in CV; η^{CV} = 70% (5 mol%), unstable after cycling; η^{CV} = 30% (30 mol%),	79
	stable after cycling; low overpotential at high Na[FSA] concentrations	
	50 mol%, low overpotential at high H_2O contents (up to 2000 ppm) after cycling	168
	50 mol%, low overpotential at high H_2O contents (500 ppm)	169
Na[FSA]–[C ₂ C ₁ im][FSA]	30 mol%, η^{DC} = 69% at 25 °C, η^{DC} = 96% at 90 °C	124
	10 mol%, reversible in CV	121
Na[FSA]–[P _{1i4i4i4}][FSA]	45 mol%, η^{CV} = 93% at 50 °C, η^{CV} decreases to 53% after 20 cycles	244
	45 mol%, at 50 °C, decrease of polarization after cycling	125
Na[FSA]–[P _{111i4}][FSA]	2.3 mol dm^{-3} , at 50 °C, stable cycling in DC	135
	42 mol%, at 50 $^{\circ}$ C, low overpotential at high H ₂ O contents (up to 500 ppm) after cycling	169
	42 mol%, at 50 °C, decrease of polarization after cycling	125
Na[FSA]–[N ₂₍₂₀₂₀₁₎₃][FSA]	55 mol%, at 50 °C, decrease of polarization after cycling	125
Na[FSA]–[N ₄₄₁₁][FSA]	30 mol%, $\eta^{\rm CV}$ = 32% at 25 °C, $\eta^{\rm DC}$ = 67% at 25 °C, $\eta^{\rm DC}$ = 92% at 90 °C	122
Na[FSA]–[N ₆₄₁₁][FSA]	$30 \text{ mol}\%, \eta^{\text{CV}} = 36\%$ at 25 °C, $\eta^{\text{DC}} = 79\%$ at 25 °C, $\eta^{\text{DC}} = 92\%$ at 90 °C	122
Na[FSA]–[AS(4.5)][FSA]	30 mol%, $\eta^{\rm CV}$ = 35% at 25 °C, $\eta^{\rm DC}$ = 71% at 25 °C, $\eta^{\rm DC}$ = 90% at 90 °C	122
Na[DCN]-[C ₄ C ₁ pyrr][DCN]	0.5 mol dm ⁻³ , $\eta^{\rm CV}$ = 74%	247
Na[FSA]–K[FSA]	56 mol%, 90 °C, reversible in CV	316
	45 mol%, 67 °C, reversible in CV	426
Na[TFSA]–Cs[TFSA]	10 mol%, 150 °C, reversible in CV	237

as room temperature ~ 25 °C) unless otherwise mentioned. See Fig. 3 for abbreviation of ionic species in $C_4C_1C_1$ im⁺: 1-butyl-2,3-dimethylimidazolium. CV: cyclic voltammetry, DC: DC repetitive polarization for Na metal deposition–dissolution, η^{CV} and η^{DC} : coulombic efficiencies of Na metal deposition–dissolution based on cyclic voltammetry and repetitive DC polarization.

in the higher-concentration electrolytes are more important to the deposition-dissolution behaviour than the ionic conductivity of the electrolyte itself. The influence of water on Na metal deposition-dissolution behaviour has been investigated in FSA-based ILs.^{168,169} Polarisation and impedance tests revealed that addition of water at the ppm level (~ 500 ppm) provides positive effects such as smooth morphology and low interfacial resistance as well as contributing to the formation of an ideal SEI.¹⁶⁹ Another study revealed that the interfacial resistance of a Na/Na symmetric cell with a Na[FSA]-[C₃C₁pyrr][FSA] electrolyte containing 2000 ppm water is lower than that for one containing 20 ppm water after 20 cycles of Na metal depositiondissolution.¹⁶⁸ Such observations suggest that water can act as an effective additive in practical Na secondary batteries with FSAbased ILs. DCA-based ILs also improve Na metal depositiondissolution behaviour.247

2.5Handling of IL electrolytes

ILs sometimes require special handling. One of the most important considerations for IL electrolytes is controlling water content. Even hydrophobic ILs that form a two-phase state with water can contain water up to a few wt%.¹⁷⁰ For example, water-saturated [C2C1im][TFSA] and [C3C1pyrr][FSA] ILs (twophase conditions) contain 19940 and 18000 ppm water, respectively.^{221,248} The removal of water from hydrophobic ILs is much easier than that from hydrophilic ILs, with evacuation

at elevated temperatures reducing the water content to below 30 ppm. Some BF₄⁻-based ILs are hydrophilic, and removal of water by simple evacuation to the level required for nonaqueous batteries is very troublesome, even at elevated temperatures. The use of molecular sieves can be effective in such cases, but contamination by other components is inevitable as ILs are difficult to separate by distillation.²⁴⁹ Purging dry gas through ILs at elevated temperature has also been reported to be effective for water removal.^{250,251}

There are now many commercially available ILs that need special attention in terms of impurity control. ILs are often contaminated by alkali metal (or silver) and/or halide ion impurities during synthesis. These ions have high polarities and can be removed by washing with water if the ILs are hydrophobic. Passing ILs through activated alumina or activated carbon can sometimes work effectively for purification, but, again, it has to be performed carefully because of possible contamination.^{252,253} Thus, pre-treatment of the starting materials used for the synthesis is more effective for obtaining high-grade ILs than post-synthesis purification of the ILs obtained.^{254,255} Another important process for battery testing using IL electrolytes is impregnation of the IL into the separator and electrode materials. Highly viscous ILs are sometimes resistant to soaking into porous materials. In such cases, treating such porous materials in ILs under vacuum at elevated temperatures (e.g., ~ 80 °C) sometimes works for impregnation

of ILs and derives the true performance of the IL electrolytes to improve the results of battery tests.^{164,180,248}

2.6 Current limitation of IL electrolytes

Cost is undoubtedly a serious issue for most applications of ILs. Battery-grade ILs are generally very expensive due to the lack of large-scale production methods and the need for fluorinated anions for high electrochemical stability.94 Kuzmina described economical aspects of ILs in 2016 and mentioned that it is very difficult to truly estimate the industrial price of ILs at the current stage.²⁵⁶ The problem of the estimation based on catalogue prices was also critically pointed out. Within the nine issues to the commercial use of ILs proposed by Joglekar et al.,²⁵⁷ the lack of scale-up studies is quite serious for practical applications. This makes cost estimation of ILs very difficult and leads to apathy from the industrial side. Furthermore, battery-grade ILs need to be extremely pure, which leads to a further increase in cost. Such pure ILs are not widely synthesized at industrial levels and thus estimation of price is difficult even for lithium ion batteries. Recovery and recycling of ILs are widely mentioned for synthetic applications, but they are not very hopeful for electrolyte applications. In general, quaternary ammonium cations are more expensive than ternary ammonium cations, and a fluorine-containing anion is more expensive than the one without fluorine. Alkylammonium and alkylphosphonium are more cost-effective than dialkylimidazolium.²⁵⁸ One of the most important steps for the application of ILs, including electrolyte applications, is collecting data for true estimation of each purpose. In contrast to TFSA salts, FSA salts can be synthesized without the costly electrofluorination and therefore could potentially achieve a lower price through further improvement in the synthetic process^{259,260} and mass production^{109,261} in the near future.

As mentioned in the previous section, high viscosity of ILs, especially at low temperature, potentially causes problems in handling such as impregnation onto porous separators and electrodes. Purification including dehydration requires many efforts for ILs. Some ILs are not always environmentally benign, and harmful decomposition products must be taken into account.²⁶² Solubility of inorganic salts into some ILs is very low, which intrinsically makes their use as electrolytes difficult. From the viewpoint of their high thermal stability, disposal of ILs after their use in batteries would be energy-consuming compared to conventional organic electrolytes.

3. Electrode materials investigated in IL electrolytes

3.1 Positive electrode materials

The prospective advantages of Na secondary batteries are their economic feasibility and their ability to act as sustainable energy sources for large-scale applications such as ESSs and electric vehicles. Thus, research into electrode materials for Na secondary batteries should comply with the principle of the utilisation of earth-abundant elements.^{49,263,264} Positive electrode

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Positive Electrode Materials Investigated in ILs



Fig. 9 Positive electrode materials for sodium secondary batteries investigated in IL electrolytes. The relationship between theoretical capacity and working potential.

materials have been widely studied as a means to maximise the performance of Na secondary batteries with respect to power and energy densities, efficiency, cycleability, and safety.

Various polyanionic compounds, layered transition-metal oxides, and organic compounds have been reported as positive electrode materials for Na secondary batteries.^{14,15,264–266} Several compounds exhibit excellent performance and some unique electrochemical results in IL electrolytes, including improved cycleability, high rate capability, and enhanced electrochemical activity at elevated operating temperatures.^{103,164,180,266–268} These discoveries were enabled by exploiting the electrochemical and thermal stabilities of IL electrolytes.

Fig. 9 illustrates the relationship between theoretical capacity and working potential for positive electrode materials applied in IL electrolytes. The gravimetric theoretical capacities (mA h g^{-1}) and working potentials (V vs. Na⁺/Na) are presented in Fig. 9. Most of the studies on positive electrode materials utilising IL electrolytes have been carried out using polyanionic compounds and O3- and P2-type layered transition-metal oxides. The "O3" and "P2" nomenclature developed by Delmas et al.²⁶⁹ is used to indicate the Na-O coordination (O for octahedral and P for prismatic) and repetition stacking period in the unit cell (2 or 3). These materials are highly reversible based on Na-intercalation with limited volumetric change. The structural properties of selected representative polyanionic compounds $(Na_2FeP_2O_7 \text{ and } Na_3V_2(PO_4)_3)$ and O3-type and P2-type layered transition-metal oxides are shown in Fig. S1 (ESI†). A comprehensive list of positive electrode materials tested in IL electrolytes is provided in Table S4 (ESI⁺) along with their structural and electrochemical characteristics. Table 2 shows their detailed electrochemical results in designated IL electrolytes including mass loading, operating temperature, cut-off voltage, and current density. In this section, the electrochemical properties of polyanionic compounds and layered oxide materials

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Table 2 Specifications of positive electrode materials in IL electrolytes^d

Cell configuration	Electrolyte	Mass loading/ mg cm ⁻²	Temp. °C	Temp./ Discharge capacity/ $^{\circ}C$ mA h g ⁻¹ (a) mA g ⁻¹	Rate capability/ 1 mA h g $^{-1}$ (a) mA g $^{-1}$ Cycleability	¹ Cycleability	Ref.
$Na/Na_2FeP_2O_7$	56 mol% Na[FSA]–K[FSA]	Ι	06	91 @ 10	59 @ 2000	91% after 1000 cycles	295
$Na/Na_2FeP_2O_7$	30 mol% Na[FSA]–[C ₂ C ₁ im][FSA]	1	25	89 @ 1	17 @ 2000	97% after 300 cycles	180
$Na/Na_2FeP_2O_7$	30 mol% Na[FSA]-[C ₂ C ₁ im][FSA]		90	93 @ 10	83 @ 2000	93% after 1500 cycles	180
Na/Na _{1.56} Fe _{1.22} P ₂ O ₇	20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA]	2	25	90 (a) 11.8	47 (a) 1180	98.5% after 300 cycles	298
Na/Na _{1.56} Fe _{1.22} P ₂ O ₇	20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA]	2	06	0 0 70	64 (a) 4720	71% atter 3000 cycle	298
$Na/Na_4NI_3(PO_4)_2(P_2O_7)$ $Na_V(PO_1)/Na_V(PO_1)$	10 III01% Na[1F3A]-[C4C1pyIT] F3A] 0.4 mol dm ⁻³ No[BF]-[C C im][BF]	8.7	07	63 (a) 10 63 (a) 0.1			202
Na,V,(PO.),/Na,V,(PO.), Na,V,(PO.),/Na,V,(PO.),	0.4 mol dm ⁻³ Na[RF.]-[C.C.im][RF.]		67 80	83 @ 0.1	$\sim 1.5 \ @ 2.0$		87
$AC^{a}/Na^{2}V_{2}(PO_{1})$	0.3 mol dm ⁻³ Na[TFSA]-[C.Hnvrrl[TFSA]	1.6	40	00 @ 0.1 70 @ 118	~ 25 (a) 1180	93 mA h o ⁻¹ after 100 cvcles	12.8
$HC^{b//N}$ $BC^{c}(PO.)$			5 10	85 @ 11 7	60 @ 234	75% after	408
Na/Na,V.(PO,),@C	20 mol% Na[FSA]–[C,C,im][FSA]		-20	58 @ 11.7	82.7 @ 11 700	/ 2 /0 attCl	164
$Na/Na_3V_2(PO_4)_3(a)C$	20 mol% Na[FSA]-[C,C,im][FSA]	. 6	25	100 @ 11.7	41 (a) 2340	99.6% after 200 cycles	164
Na/Na ₃ V ₂ (PO ₄) ₃ @C	50 mol% Na[FSA]–[C2C1im][FSA]	3	90	$104 \ {\textcircled{a}} \ 11.7$	82.7 (a) 11 700	89.2% after 5000 cycles	164
$Na/Na_3V_2(PO_4)_3$ OCNF ^c	40 mol% Na[FSA]–[C_2C1im][FSA]	8.5	25	105 @ 11.7	51 @ 1170	94 mA h g^{-1} after 200 cycles	283
$Na/Na_3V_2(PO_4)_3$ (BOF)	40 mol% Na[FSA]–[C ₂ C ₁ im][FSA]	8.5	06	107 @ 11.7	77 @ 11 700	104 mA h g^{-1} after 200 cycles	283
$Na/Na_3V_2(PO_4)_3$ (D		Ι.	25	107 @ 50		104 mA h g^{-1} after 50 cycles	286
$Na/t-NaFePO_4$	0.5 mol dm ⁻³ Na[TFSA]-[C ₄ C ₁ pyrr][TFSA]		50	125 @ 7.8	75 @ 155	83% after 100 cycles	84
Na/t-NaFePO ₄	1 mol dm ⁻³ Na[BF ₄]–[C ₄ C ₁ pyrr][TFSA]	2.5	75	152 @ 7.8	91 @ 155	90% after 100 cycles	119
Na/t-NaFePO ₄	42 mol% Na[FSA]- $[N_{2(20201)3}]$ [FSA]		20	97 @ 15.5	66 (a) 155	71% atter 100 cycles	289
Na/t-NaFePO4	45 mol% Na[FSA]-[P _{1i4i4i4}][FSA]	1	50	113 (a) 15.5	88 (a) 155 84 (a) 155	12% atter 100 cycles	289
Na/t-INAFEPO4	25 III01% Nd[F5A]-[F1114][F5A]	,	00	10/ (B) 101	64 (J) 100	55% after 100 cycles	202
Na/m-NarerO4 Na/NaVODO - AD-0 h	20 III01% Na[F3A]-[C2C1IIII][F3A]	°, c	0.0	c.c1 (ق) /01 60 @ 5	CC1 (P) 70	0/% aller 100 cycles	502
Na/NaVOFO4-AD-0 II Na/NaVODO -AD-0 h	20 III01% Na[F3A] [0301Py11] F3A]	1	C7	00 (a) 5 101 @ F			502
Na/NaCrO.	56 mol% NafFSAl_KfFSAl	4	06	77 @ 15	77 @ 723 58 @ 150	20% after 100 cycles	316
Na/NaCrO.	10 mol% Na[TFESA]_Cs[TFESA]		150	94 @ 10	50 @ 130 66 4 @ 100	$83.4 \text{ mA h} \text{ o}^{-1}$ after 10 cycles	237
Na/NaCrOs Na/NaCrOs	20 mol% Na[FSA]-[C.C. wurl[FSA]		951 25	92 @ 20	00.4 @ 100	03.4 IIIA II & AUGI 10 CYCICS	245
Na/NaCrO.	20 mol% Na[FSA]-[C.C.mvrrl[FSA]		80	<u>106 ක 20</u>	62. @ 200		245
HC ^b /NaCrO,	20 mol% Na[FSA]-[C ₃ C ₁ Dyrr][FSA]	13.5 (HC 5.2)	25	27 A h @ 2.7 A	26 A h @ 10.8 A		411
$HC^{b}/NaCrO_{2}$	20 mol% Na[FSA]–[C ₃ C ₁ pyrr][FSA]	13.5 (HC 5.2)	60	28 A h @ 2.7 A	27 A h @ 10.8 A	87% after 500 cycels	411
$HC^{b}/NaCrO_{2}$	20 mol% Na[FSA]–[C ₃ C ₁ pyrr][FSA]	13.5 (HC 5.2)	06	28 A h @ 2.7 A	27 A h @ 10.8 A		411
$Na/NaMnO_2$	10 mol% Na[TFSA]–[C ₄ C ₁ pyrr][TFSA]	2^{-4}	25	130 @ 9.7		81% after 10 cycles	133
$Na/Na_{0.44}MnO_2$	[Na(G5)1][TFSA]		60	110 @ 12.7	Ι	95% after 30 cycles	149
$Na/Na_{0.44}MnO_2$	1 mol dm ⁻³ Na[ClO ₄]-[C ₄ C ₁ pyrr][TFSA]		50	109 @ 6.4	82 @ 127	80% after 100 cycles	127
$HC^{\nu}/Na_{0.44}MnO_{2}$	1 mol dm ⁻² Na[FSA]-[C ₃ C ₁ pyrr][FSA]	1.5	25	120 @ 30	70 @ 1000	97% after 100 cycles	86
$Na/P2-Na_{2/3}Fe_{1/3}Mn_{2/3}O_2$	20 mol% Na[FSA]–[C ₃ C ₁ pyrr][FSA]	I	00 -	227 (a) 20	65 @ 500	64% after 200 cycles	318
$Na/P2-Na_{2/3}Fe_{2/3}Mn_{1/3}O_2$	2.3 mol dm \sim Na[FSA]–[P _{111i4}][FSA]	Ι	50	130 (a) 17.4	110 (a) 34.8		135
$Na/U3-Na_{2/3}Fe_{2/3}Mn_{1/3}U_2$	2.3 mol dm ⁻ Na[FSA]-[P _{111i4}][FSA]	,	00	16/ (a) 1/.4	130 (a) 34.8	5/% atter 100 cycles	135
Na/FZ-Na2/3N11/3N112/3O2	I III0I UIII INALIFAA]-[U4U1PyI1][IFAA] O E mol dm ⁻³ No[ECA]-DEMIE-[TECA]	c I	00 9 1 0	220 (a) 10 205 (a) 20		200 IIIA II g allef 14 cycles 70% offer 350 moles	019 121
Na/Na = Mn = CO = CO	0.5 mol dm ⁻³ Na[FSA]-MOFMDID_[TFSA]	[v	сл С7 С7	203 (a) 20 117 (a) 20	00 @ 400 20 @ 400	20% after 350 cycles	131
Na/P2-NaNa Ni Ni Fe Mn O.	0.2 mol dm ⁻³ Na[TFSA]-[C.C. nvrr][TFSA	1	57	11/ @ 20 150 @ 10	001 m 07		409
Na.SB-C/P2-NaNao 6Nio 225 C0.114110.66 C2	0.2 mol dm ⁻³ NalTFSA]-	A] 1 (Na.SB-C 1.9)	25	120 @ 10 120 @ 10			409
$AC^{a}/Na_{0.67}Mn_{0.89}Mg_{0.11}O_{2}$	0.3 mol dm ⁻³ Na[TFSA]-		40	57 @ 100		59 mA h g^{-1} after 100 cycles	128
$Na_{2.55}V_6O_{16}/Na_{0.6}CO_{0.1}Mn_{0.9}O_2$				125 (a) 50	60 @ 600	76% after 100 cycles	410
Na/P2-Na _{0.45} Ni _{0.22} Co _{0.11} Mn _{0.66} O ₂	0.4 mol dm ⁻³ Na[TFSA]-[C ₄ C ₁ pyrr][TFSA]	A] 2.5	20	210 @ 23		80% after 100 cycles	267
$Na/P2-Na_{2/3}Mn_{0.8}Fe_{0.1}Ti_{0.1}O_2$	2.3 mol dm ⁻⁷ Na[FSA]-[P _{111i4}][FSA]	Ι	20	$176 \ (a) \ 17.6$	125 @ 88	92% after 50 cycles	125
Na/P2-Na _{2/3} MII _{0.8} F6 _{0.1} I10.1O2 Na/P2-NaMnFFaT1O_	45 mol% Na[FSA][P _{1i4i4i4}][FSA] 55 mol% Na[FSA][N ₋₆₆₂₀₀ .][FSA]		000	101 (a) 1/.6 153 (a) 17 6	60 (a) 88 30 (a) 88	58% after 50 cycles	125
Na/calix[4]quinone	0.3 mol dm ^{-3} Na ^{[TFSA]-[C₂C₂000]₃[L²C₂Dyrr][TFSA]}	A] 0.58	25	406 @ 20	20 m 20	99.7 after 300 cycles	380
Na/5,7,12,14-pentacenetetrone			25	245 @ 30	I		380
		1		1			

^{*a*} AC = activated carbon. ^{*b*} HC = hard carbon. ^{*c*} CNF = carbon nanofiber. ^{*d*} See Fig. 3 for abbreviation of ionic species in ILs.

tested with IL electrolytes in Table 2, including their structural properties, are reviewed.

3.1.1 Polyanionic compounds. In recent years, polyanionic compounds have been attracting considerable attention as high-performance positive electrode materials for Na secondary batteries. Polyanionic compounds build stable molecular frameworks with $(XO_4)^{n-}$ or $(X_mO_{3m+1})^{n-}$ (X = S, P, Si, As, B, etc.) tetrahedral units and MO_6 (M = transition metal) octahedral units.^{270,271} These units are connected via oxygen bridging with the edges or corners of the polyhedral being shared, resulting in a variety of polyanionic compounds that exhibit one-, two-, and three-dimensional structural ordering for facile Na⁺ diffusion paths.²⁷²⁻²⁷⁴ For example, a three-dimensional Na₃V₂(PO₄)₃ crystal structure is formed by corner-sharing of octahedral VO₆ units and tetrahedral PO₄ units, and this material is well known as a Na Super Ionic CONductor (NASICON).87,275-277 Polyanionic compounds usually exhibit high thermal stabilities (owing to the strong X-O covalent bonds) and strong resistance to natural oxidation but low electronic conductivities.271,278-280 However, poor electronic conductivity results in poor electrochemical performance, but it can be overcome by surface modifications, carbon coating, and elevation of operating temperature using IL electrolytes, as reported in recent studies.

NASICON-type Na₃V₂(PO₄)₃ crystallises in the rhombohedral space group $R\bar{3}c^{281,282}$ and is the polyanionic material that has been most extensively studied using IL electrolytes owing to its open structure for facile Na⁺ transport, high thermal stability, and high energy density. Na₃V₂(PO₄)₃ can provide two redox reactions based on V⁴⁺/V³⁺ and V³⁺/V²⁺. The V⁴⁺/V³⁺ reaction is based on a two-electron reaction (for two V atoms) with a theoretical capacity of 117 mA h g⁻¹ showing an average potential of 3.4 V *vs.* Na⁺/Na (eqn (6)). The V³⁺/V²⁺ reaction is based on a

one-electron reaction (for one V atom) with a theoretical capacity of 55 mA h g^{-1} showing an average potential of 1.6 V vs. Na⁺/Na (eqn (7)).^{87,164,283,284} This double-redox system facilitates the fabrication of symmetric Na₃V₂(PO₄)₃ cells, and is often used for testing new electrolytes.

$$\operatorname{Na}_{3}\operatorname{V}_{2}(\operatorname{PO}_{4})_{3} \rightleftharpoons \operatorname{Na}_{2}(\operatorname{PO}_{4})_{3} + 2\operatorname{Na}^{+} + 2e^{-} \qquad (6)$$

$$Na_4V_2(PO_4)_3 \rightleftharpoons Na_3V_2(PO_4)_3 + Na^+ + e^-$$
(7)

Fig. 10 shows the electrochemical behaviour of a Na₃V₂(PO₄)₃ electrode in different IL electrolytes. The first study on the electrochemical performances of positive electrodes in Na secondary batteries assembled using an IL electrolyte was performed using Na₃V₂(PO₄)₃ symmetric cells with a Na[BF₄]–[C₂C₁im][BF₄] IL electrolyte.⁸⁷ The Na₃V₂(PO₄)₃ symmetric cell exhibits a lower discharge capacity in a Na[BF₄]–[C₂C₁im][BF₄] IL than in 1 mol dm⁻³ Na[ClO₄]-PC at 25 °C, but the rate performance is improved at 80 °C, as shown in Fig. 10a.⁸⁷ The high viscosity and low conductivity of the IL and low electronic conductivity of Na₃V₂(PO₄)₃ at 25 °C were identified as the obstacles to rate performance.

A more recent study on Na₃V₂(PO₄)₃ revealed that Na₃V₂(PO₄)₃ combined with carbon nanofibers exhibits good rate performance in IL electrolytes at 25 °C.²⁸³ In this study, a symmetric cell comprising a Na₃V₂(PO₄)₃/carbon nanofiber positive electrode with a 40 mol% Na[FSA]–[C₂C₁im][FSA] IL electrolyte exhibited a high rate capability of 48.3 mA h g⁻¹ at 100C (1C = 117 mA g⁻¹) and 25 °C, as shown in Fig. 10b.²⁸³ Carbon nanofibers efficiently improve the electronic conductivity of Na₃V₂(PO₄)₃. Owing to the high thermal, chemical, and electrochemical stabilities of Na₃V₂(PO₄)₃, it has become the standard material for assessing new IL electrolytes. For example, the



Fig. 10 Electrochemical behaviour of NASICON-type $Na_3V_2(PO_4)_3$ in ILs. (a) Rate performance of $Na_3V_2(PO_4)_3$ symmetric cells with a 0.4 mol dm⁻³ $Na[BF_4] - [C_2C_1im][BF_4]$ IL and 1 mol dm⁻³ $Na[ClO_4]$ -PC organic solvent electrolytes. Reproduced with permission.⁸⁷ Copyright 2010, The Electrochemical Society. (b) Rate performance of CNF-incorporated $Na_3V_2(PO_4)_3$ symmetric cells with a 40 mol% $Na[FSA] - [C_2C_1im][FSA]$ IL electrolyte. Reproduced with permission.²⁸³ Copyright 2019, American Chemical Society.

protic IL Na[TFSA]–[C₄Hpyrr][TFSA] (0.3 mol dm⁻³) has been studied as an electrolyte for Na secondary batteries.¹²⁸ This protic IL electrolyte exhibits more stable cycle performance than that with a 0.3 mol dm⁻³ Na[TFSA]–[C₄C₁pyrr][TFSA] aprotic IL. A gelled 0.8 mol kg⁻¹ Na[FSA]–[C₃C₁pyr]][FSA]-PVdF-*co*-HFP (PVdF-*co*-HFP = polyvinylidene fluoride-*co*-hexafluoropropylene) IL gel electrolyte showed good cycle performance with 92% capacity retention and 99.9% coulombic efficiency over 150 cycles.²⁸⁵ Another study on Na₃V₂(PO₄)₃ with Na[PF₆]–[C₄C₁im][TFSA] showed improved cycling performance over that achieved with an organic solvent electrolyte.²⁸⁶ Cycleability results will be considered in more detail later in Section 4.3.

Olivine-type LiFePO₄ is one of the most well studied and commercialised materials for LIBs because of its costeffective, facile synthesis and high theoretical capacity of 170 mA h g^{-1} .^{273,287} In the same manner, Fe-based phosphates (NaFePO₄) have been investigated as potential positive electrode materials using IL electrolytes. Olivine NaFePO₄, which has the highest theoretical capacity of 155 mA h g^{-1} among the Fe-based phosphates, has two polymorphs, i.e., triphylite and maricite phases, both of which crystallise in the orthorhombic space group Pnma.²⁸⁸ However, triphylite-phase NaFePO₄ is not thermally stable and can only be prepared by electrochemical/ chemical extraction of Li from LiFePO4 and substitution with Na.²⁸⁸ Fig. 11 shows the electrochemical behaviours of triphylite-NaFePO₄ and several layered metal oxides in ILs. Triphylite NaFePO₄ was assessed using [C₄C₁pyrr][TFSA] IL electrolytes containing Na[BF₄], Na[ClO₄], Na[PF₆], or Na[TFSA] and a conventional 1 mol dm⁻³ Na[ClO₄]-EC/DEC (1:1, v/v) organic electrolyte at 25, 50, and 75 °C.^{84,119} At 25 °C, 1 mol dm⁻³ Na[ClO₄]-EC/ DEC exhibited the highest rate performance and cycleability, but at the elevated temperature of 50 °C, the cycleability became poorer. A significant improvement in reversible capacity, rate performance (Fig. 11a), and cycleability were observed in 1 mol dm⁻³ Na[BF₄]–[C₄C₁pyrr][TFSA]. The discharge capacity was further improved to 152 mA h g^{-1} (at 0.05C) at 75 °C in 1 mol dm⁻³ Na[BF₄]-[C₄C₁pyrr][TFSA] electrolyte.^{84,119}

Recently, cycle performance improvement was reported for a 1 mol dm⁻³ Na[FSA]–[P₁₁₁₁₄][FSA] IL electrolyte with a capacity retention of 95% over 100 cycles (50 °C, C/2).²⁸⁹ In this study, it was suggested that electrolyte conductivity is not the main factor for determining cell performance and is not significantly related to capacity. Above all, the SEI determines the cycle stability, as is discussed in more detail in the section on negative electrodes below. XPS measurements confirmed the existence of IL-derived decomposition products like NaOH, Na₂S, and NaF on the Na metal surface, which provided stable cycle performance in a half-cell.²⁸⁹

Maricite-NaFePO₄ can be prepared by the conventional solid-state method. It was previously believed that maricite-NaFePO₄ was electrochemically inactive.^{288,290} However, nanosized maricite-NaFePO₄ transforms into an amorphous FePO₄ phase during the first charge (desodiation) and shows unexpected charge-discharge performance.²⁹¹ Another study on maricite NaFePO₄ using ILs revealed that ball-milled maricite-NaFePO₄ shows reversible (de)sodiation properties,



Fig. 11 Electrochemical behaviours of triphylite-NaFePO₄ and layered metal oxides in ILs. (a) Charge-discharge curves for a Na/1 mol dm⁻³ Na[BF₄]-[C₄C₁im][TFSA]/NaFePO₄ cell at various current rates and 50 °C. Reproduced with permission.¹¹⁹ Copyright 2014, American Chemical Society. (b) Charge-discharge curves for Na/2.3 mol dm⁻³ Na[FSA]-[P₁₁₁₁₄][FSA]/O3- and P2-type Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ cells. Reproduced with permission.¹³⁵ Copyright 2017, Elsevier. (c) Charge-discharge curves for a Na/Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂/10 mol% Na[TFSA]-[C₄C₁pyrr][FSA] cell. Reproduced with permission.²⁶⁷ Copyright 2014, Elsevier.

as confirmed by X-ray diffraction (XRD) and X-ray absorption near edge structure (XANES) spectroscopy.^{102,103} Further details will be presented in Section 4.1.

Pyrophosphate compounds, such as Na₂FeP₂O₇, Na_{1.56}Fe_{1.22}P₂O₇, and Na₄Fe(PO₄)₂P₂O₇, offer stable three-dimensional diffusion paths for Na ions composed of corner-sharing FeO₆ units.²⁹² Na₂FeP₂O₇ crystallises in the triclinic space group $P\bar{1}$.^{293,294} Its theoretical capacity estimated from the one-electron reaction based on the Fe²⁺/Fe³⁺ redox couple reaction is 97 mA h g⁻¹ (eqn (8)).²⁹⁴

$$Na_2FeP_2O_7 \rightleftharpoons NaFeP_2O_7 + Na^+ + e^-$$
 (8)

Na₂FeP₂O₇ in Na[FSA]–[C₂C₁im][FSA] with various salt concentrations at 25 to 90 °C and 56 mol% Na[FSA]–K[FSA] at 90 °C show adequate reversible capacities.^{180,295} In particular, Na₂FeP₂O₇ in a 30 mol% Na[FSA]–[C₂C₁im][FSA] IL electrolyte shows an excellent rate capability of 83 mA h g⁻¹ at 2000 mA g⁻¹ and 90 °C but only 17 mA h g⁻¹ at 25 °C.^{180,295} Na_{2-x}Fe_{1+x/2}P₂O₇ compounds (where 0 < *x* < 0.44) were prepared by a solid-state method and evaluated. Among them, Na_{1.56}Fe_{1.22}P₂O₇ (where *x* = 0.44) is introduced here, as it showed the highest theoretical capacity (118 mA h g⁻¹) among the Na_{2-x}Fe_{1+x/2}P₂O₇ compounds.^{296,297} Notably, the theoretical capacity was increased (~20 mA h g⁻¹) by the substitution of Na with Fe to form Na_{1.56}Fe_{1.22}P₂O₇, and it is estimated that 1.22 equivalents of Na is reversibly (de)sodiated based on the Fe²⁺/Fe³⁺ redox couple reaction (eqn (9)).^{296,297}

$$Na_{1.56}Fe_{1.22}P_2O_7 \rightleftharpoons Na_{0.34}Fe_{1.22}P_2O_7 + Na^+ + e^-$$
 (9)

However, the actual reversible capacity of $Na_{1.56}Fe_{1.22}P_2O_7$ was limited to 85 mA h g⁻¹ in the organic solvent electrolyte. Using 20 mol% Na[FSA]–[C₃C₁pyrr][FSA] IL electrolyte at 90 °C, a reversible capacity of 108 mA h g⁻¹ was obtained, which is more than that obtained in the organic electrolyte by approximately 15% under the same conditions.²⁹⁸ Furthermore, superior cycle performance was reported with this IL electrolyte at 25, 50, and 90 °C. This improvement was achieved owing to the high cycleability of the Na metal counter electrode in the IL electrolyte and the small volume change of $Na_{1.56}Fe_{1.22}P_2O_7$ during (de)sodiation (only 1.9%) as compared to those of the triphylite-NaFePO₄ (17.6%), $Na_3V_2(PO_4)_3$ (8.3%), and $Na_2FeP_2O_7$ (2.6%).^{288,293,297–299}

 Na_2MnSiO_4 is a low-cost and environmentally benign material composed of only earth-abundant elements. In a recent study, carbon-coated Na_2MnSiO_4 was prepared by a sol-gel method and assessed in a 20 mol% $Na[FSA]-[C_3C_1pyrr][FSA]$ IL.³⁰⁰ This material has theoretical capacities of 278 mA h g⁻¹ and 125 mA h g⁻¹ based on a two-electron reaction and one-electron reaction, respectively.^{300,301} However, the obtained reversible capacities of 70, 94, and 125 mA h g⁻¹ at 25, 50, and 90 °C, respectively, suggested that the redox activity based on only the one-electron reaction, which corresponds to 0.9 equivalents of Na being reversibly (de)sodiated based on the Mn^{3+}/Mn^{2+} redox couple reaction.

Vanadium-based phosphate (NaVOPO₄) crystallises in the monoclinic space group $P2_1/c$. NaVOPO₄ has a high operating potential of 3.6 V vs. Na⁺/Na with a large theoretical capacity of 145 mA h g⁻¹ based on the V⁵⁺/V⁴⁺ redox couple reaction.³⁰² The charge–discharge behaviour of NaVOPO₄ was investigated using 20 mol% Na[FSA]–[C₃C₁pyrr][FSA] IL electrolytes at 25 and 90 °C. Reversible capacities of 60 and 101 mA h g⁻¹ were obtained at 25 and 90 °C, respectively.³⁰³

3.1.2 Transition-metal oxides. LIB usage has been promoted by the intensive utilisation of transition-metal oxides such as Li_xMO_2 (M = Co, Ni, Mn).^{304–309} Such transition-metal oxides as positive electrode materials exhibit excellent performance in terms of high capacity, energy density, and cycle performance.^{305,308,309} Accordingly, a number of transition-metal oxides for Na secondary

batteries have been explored.^{14,36,266,310–313} In this section, representative transition-metal oxide compounds that have been investigated with IL electrolytes are discussed, including O3- and P2-type layered transition-metal oxides. Na_{0.44}MnO₂ with an S-shaped tunnel structure will also be discussed.

The structures of O3- and P2-type layered transition-metal oxides are shown in Fig. S1c and d (ESI[†]), respectively. Despite the success of layered transition-metal oxide compounds as positive electrode materials in LIBs, the number of studies on their use in IL electrolytes for Na secondary batteries is still limited. In terms of cost and resource sustainability, Ni and Co layered transition-metal oxides are not particularly attractive materials for Na secondary batteries. Thus, the layered transition-metal oxides for Na secondary batteries investigated in IL electrolytes mostly contain Mn, Fe, and Cr elements instead of Co and Ni.

The first report on reversible Na insertion and extraction concerned O3-type NaCrO₂.³¹⁴ The theoretical capacity of this material is 125 mA h g⁻¹ for the one-electron reaction based on the Cr⁴⁺/Cr³⁺ redox couple.³¹⁵ O3-type NaCrO₂ exhibits numerous two-phase regions in its charge–discharge plateaus up to 3.5 V. The upper cut-off voltage was 3.6 V because further desodiation causes an irreversible phase transition. The material was assessed with IL electrolytes comprising 56 mol% Na[FSA]–K[FSA] at 80 °C. The Na/Na[FSA]–K[FSA]/NaCrO₂ cell exhibited a specific discharge capacity of 77.3 mA h g^{-1.316} This capacity was improved to 101.4 mA h g⁻¹ at 150 °C using 10 mol% Na[TFSA]–Cs[TFSA].²³⁷ Furthermore, the use of NaCrO₂ in a wide temperature range (–20 to 90 °C) with 20 mol% Na[FSA]–[C₃C₁pyrr][FSA] IL electrolyte has been reported (see a later Section 4.1 for the temperature dependence of the performance).¹⁶⁶

Recently, binary and ternary mixed transition metal oxides have been investigated using IL electrolytes. O3 and P2-type NaFe_{1-x}Mn_xO₂ consisted of earth-abundant elements and was expected to deliver a high reversible capacity. O3- and P2-type $Na_{2/3}Fe_{2/3}Mn_{1/3}O_2$ was assessed in a 0.5 mol dm⁻³ Na[PF₆]-EC/ DMC organic solvent electrolyte. The P2-type Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ and O3-type Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ delivered the first discharge capacities of 151 and 157 mA h g^{-1} , respectively, over the voltage range 4.2-1.5 V at C/10. However, the capacity of both P2- and O3-Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ faded within 10 cycles.³¹⁷ Interestingly, the capacity and cycle performance were improved in measurements using 2.3 mol dm⁻³ Na[FSA]-[P_{111i4}][FSA]¹³⁵ only for O3-type Na2/3Fe2/3Mn1/3O2, which exhibited excellent performance using the IL electrolyte at 50 °C. The capacity of O3-type $Na_{2/3}Fe_{2/3}Mn_{1/3}O_2$ was improved to 184 mA h g⁻¹ over the voltage range of 4.0-1.75 V, while P2-type Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ provided a capacity of 130 mA h g^{-1} because a narrower voltage range was applied (Fig. 11b). The rate capability and cycle performance were improved for both P2-type and O3-type Na_{2/3}Fe_{2/3}Mn_{1/3}O₂ upon using IL electrolytes.¹³⁵

A material with a different Fe/Mn ratio, P2-Na_{2/3}Fe_{1/3}Mn_{2/3}O₂, was investigated in a Na[FSA]–[C₃C₁pyrr][FSA] IL electrolyte, and showed a high discharge capacity of 227 mA h g⁻¹ at a current density of 20 mA g⁻¹ over the voltage range 2.0–4.3 V. With a wider cut-off voltage of 1.5–4.2 V, P2-Na_{2/3}Fe_{1/3}Mn_{2/3}O₂

exhibited a very high discharge capacity of 317 mA h g⁻¹ in the IL, but this capacity exceeds its theoretical capacity of 261 mA h g⁻¹, indicating unfavourable side reactions.³¹⁸

A recent report on P2-Na_{2/3}Ni_{1/3}Mn_{2/3}O₂ revealed stable cycling involving an O redox reaction in a 1 mol dm⁻³ Na[TFSA]–[C₄C₁pyrr][TFSA] IL electrolyte. The discharge capacity of 220 mA h g⁻¹ fell to 200 mA h g⁻¹ after 14 cycles in the IL electrolyte at 60 °C, whereas it decreased to 160 mA h g⁻¹ after 14 cycles in 1 mol dm⁻³ Na[PF₆]-EC/DMC at 25 °C.³¹⁹

The ternary transition metal oxide P2-Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ provides a good example of the superiority of IL electrolytes (Fig. 11c). Compared to the case in an organic electrolyte, higher reversibility of the P2–O2 phase transition was observed for P2-Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ using 10 mol% Na[TFSA]–[C₄C₁pyrr][FSA] above 4.2 V, which is attributable to the wide electrochemical window of the IL. The cell exhibited a capacity of approximately 220 mA h g⁻¹ at 12 mA g⁻¹ and 20 °C. This is because the electrochemical stability of the IL electrolyte is higher than that of the organic electrolyte. Furthermore, the low solubility of Mn in the IL electrolyte at potentials below 2.2 V helps to maintain its good cycleability.²⁶⁷ A discussion of the cycleability of the materials is presented in Section 4.3.

 $Na_{0.44}MnO_2$ is an attractive material for Na secondary batteries in terms of its low cost and the earth-abundance of its constituent elements. It crystallises in the orthorhombic space group *Pbam* and has wide tunnels for Na⁺ diffusion.³²⁰ The initial charge desodiates 0.26 equivalents of Na⁺ from $Na_{0.44}MnO_2$ (eqn (10)). After the initial charge, the charge– discharge reaction is based on (de)sodiation with 0.46 equivalents of Na⁺ (eqn (11)). The theoretical capacity is calculated to be 127 mA h g⁻¹ based on the Mn⁴⁺/Mn³⁺ redox couple reaction.

 $Na_{0.44}MnO_2 \rightarrow Na_{0.18}MnO_2 + 0.26Na^+ + 0.26e^-$ (10)

 $Na_{0.18}MnO_2 + 0.46Na^+ + 0.46e^- \rightleftharpoons Na_{0.64}MnO_2$ (11)

Electrochemical measurements were performed using a Na/ [Na(G5)][TFSA]/Na_{0.44}MnO₂ cell. The cell showed a reversible charge–discharge capacity of 110 mA h g⁻¹ over a voltage range of 2.0–4.0 V at 60 °C.¹⁴⁹ This electrochemical performance was improved using [Na(G5)][FSA] mixed with the molecular solvent hydrofluoroether (HFE). [Na(G5)][FSA] with HFE was found to exhibit lower viscosity and higher ionic conductivity than Na[(G5)₁]–[TFSA] with HFE. As a consequence, Na_{0.44}MnO₂ in [(NaG5)][FSA] with HFE showed a reversible charge–discharge capacity of 113 mA h g⁻¹ at 30 °C and was stable for 50 cycles (coulombic efficiency 99%) without capacity degradation.³²¹

3.2 Negative electrode materials

In this section, the performances of negative electrode materials in IL electrolytes will be discussed. Table 3 shows electrochemical results of negative electrode materials in IL electrolytes. The various types of negative electrodes are classified based on their sodiation/desodiation behaviour. They are divided into carbon materials, titanium-based compounds, alloy materials, and conversion-based materials. Each section begins with a brief description of the material characteristics and electrochemical behaviour in organic electrolytes and their performances in ILs. Finally, based on the insight accumulated from the studies reported, future prospective investigations are suggested.

3.2.1 Carbon materials. Carbon materials have long been the focus of electrode materials in electrochemistry research owing to their low-cost, good electrical conductivities, and chemical robustness. Despite delivering a high reversible capacity (~ 372 mA h g⁻¹ for LiC₆) as a negative electrode in LIBs,³²² graphitic carbon generally shows very limited capacity (\sim 35 mA h g⁻¹ for the first charge (sodiation) with the formula NaC₆₄)³²³ in Na secondary batteries using conventional esterbased electrolytes owing to the thermodynamic instability of Na-inserted graphite intercalation compounds (GICs).^{324,325} This limitation has been circumvented using co-intercalation phenomena through the formation of $(Na(G2)C_{20})$ GICs. A reversible capacity approaching 100 mA h g^{-1} with stable cycleability over 1000 cycles was recently achieved for a graphite electrode.³²⁶ This result inspired researchers to investigate new electrolytes alongside new carbon materials.

To date, hard carbon (HC) has been the most extensively studied carbonaceous negative electrode material for Na secondary batteries. HC prepared by the pyrolysis of sugar demonstrated a high storage capacity of 300 mA h g⁻¹ with excellent capacity retention and rate capability.³²⁷ The esterbased solvent EC/PC provided lower overpotential and charge transfer resistance than EC/PC with fluoroethylene carbonate (FEC) as an additive. Overall, this study proposed that the performance is highly dependent on the electrolyte. The importance of electrolytes is readily understood by such studies. Therefore, many researchers have recently investigated ILs with carbon materials.

Fig. 12 shows the electrochemical behaviours of negative electrode materials in ILs. The electrochemical properties of semi-graphitic HC prepared by pyrolysing olive pits were investigated with 0.5 mol dm⁻³ Na[TFSA]-[C4C1pyrr][TFSA] with addition of 2% vinyl ethylene carbonate (VEC) and 0.3 mol dm⁻³ Na[TFSA]-[C4Hpyrr][TFSA] at 30 °C.³²⁸ This carbon material showed good cycleability with Na[TFSA]-[C4C1pyrr][TFSA] but the capacity faded quickly in the case of 0.3 mol dm⁻³ Na[TFSA]–[C₄Hpyrr][TFSA] owing to the instability of the $[C_4Hpyrr]^+$ cation below 1 V with the Na metal as a counter electrode. The electrochemical performance of commercial HC powders was investigated with a 20 mol% Na[FSA]-[C3C1pyrr][FSA] IL in the wide temperature range -10 to 90 °C, as shown in Fig. 12a.³²⁹ This HC showed a high capacity of 277 mA h g^{-1} at 20 mA g^{-1} and remarkable rate capability at 90 °C (Fig. 12b) owing to the high ionic conductivity and low charge transfer resistance of the IL. It also showed a capacity of 193 mA h g^{-1} at 20 mA g^{-1} and 25 °C, which is important for room temperature applications. Although only a limited number of studies conducted with ILs is available, the above-mentioned studies demonstrate the high capacity, excellent rate capability at elevated temperature, and superior cycleability of carbon materials achievable in ILs.

3.2.2 Titanium-based compounds. Another class of frequently investigated negative electrode materials is Ti-based materials, which are known for their chemical stability,³³⁰ low costs, and

Cell configuration	Electrolytes	Mass loading/ [–] mg cm ^{–2} °	Temp./ °C	1st cycle discharge capacity @ current (col. eff.)/mA h $\rm g^{-1}$ @ mA $\rm g^{-1}$	Rate capability/ mA h g^{-1} (current)	Cycleability	Ref.
Na/3-D CF	Na[FSA]–[C ₃ C ₁ pyrr][FSA]	-	25 60	$235 \ (a) \ 30 \ (40\%)$	115 (1000 mA g^{-1}) —	85% (250 cycles) 80% (250 cycles)	397
Na/semi-graphitic HC Na/HC	0.5 M Na[TFSA]-[C ₃ C ₁ pyrr][TFSA] + 2% VEC 20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA]		30 25 25	©	i e	100% (55 cycles) 	328 329
Na/anatase TiO ₂ Na/Nb doped rutile TiO ₂	10 mol% Na[TFSA]-[C ₃ C ₁ pyrr][FSA] Na[FSA]-[C ₃ C ₁ pyrr][FSA] Na[FSA]-PC Na[FSA]-PC		90 25 30 30 30 30 30		230 (1000 mA g ⁻¹) 47 (3350 mA g ⁻¹) —	84% (500 cycles) 98% (80 cycles) 77% (350 cycles) 97% (100 cycles)	336 341
Na/TiO ₂ -Fe ₃ O ₃ Na/FeTiO ₃ -C	Na[rSA]-PC 20 mol% Na[rSA]-[C ₃ C ₁ pyrr][rSA] 20 mol% Na[rSA]-[C ₃ C ₁ pyrr][rSA] 20 mol% Na[rSA]-[C ₃ C ₁ pyrr][rSA]	0.8-1.3	60 90 10 10 10 10 10 10 10 10 10 10 10 10 10	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	$\stackrel{-}{\stackrel{-}{\stackrel{-}{\stackrel{-}{\stackrel{-}{\stackrel{-}{\stackrel{-}{\stackrel{-}$	91% (350 cycles) 80% (300 cycles) — 113% (2000 cycles)	337 338
Na/TiO2-C	Na[\Fr6_FLC/DMC 20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA] 20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA] Na[DF.]-FC/DMC	1-2	25 90 25	150 $(@ 10 ())$ 143 $(@ 10 ())$ 149 $(@ 10 (45\%))$ 114 $(@ 11 (-))$	— — — — (2000 mA g ⁻¹)	79% (1000 cycles) 	228
Na/Sn film Na/Sn-Fe film Na/Sn-Cu film	56 mol% Na[FSA]-K[FSA] 56 mol% Na[FSA]-K[FSA] 56 mol% Na[FSA]-K[FSA] 56 mol% Na[FSA]-K[FSA]		06 06	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	— — 70 (847 mA g ⁻¹)	66% (30 cycles) 43% (100 cycles) 100% (1000 cycles)	427 355 353
Na/Sn-Ni film Na/P-AB	56 mol% Na[FSA]-K[FSA] 0.25 M Na[FSA]-[C ₃ C ₁ pyrr][FSA] Na[PF ₆]-PC Na[PF ₆]-EC/DEC		06	$ \begin{array}{c} 605 @ 84.7 (86.4\%) \\ 1595 @ 125 (72.3\%) \\ 1768 @ 125 (72.3\%) \\ 1755 @ 125 (77.3\%) \\ 1755 @ 125 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1654 & 0.26 (77.3\%) \\ 1754 & 0.26 (77.3\%)$		56% (100 cycles) 93% (80 cycles) 26% (80 cycles) 30% (80 cycles)	354 80
Na/black P	Na[FFS/=LC/FC/JDEC Na[FFSA]-[C ₍₁₀₂₀₁)C ₁][FSA] Na[TFS1]-[C ₂ C ₁ im][FSA] Na[TFSA]-PC		30 30 30	(~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~			362
Na/CuP ₂ -C Na/V D_5D	20 mol% Na[FSA]-[C ₃ C ₁ pyrr][FSA] 20 mol% No[EsA]-[C ₂ C ₂ mm+]EsA]	,	25 90 25	73.7%) 70.2%) 80%)	$50 (8000 \text{ mA g}^{-1}) \\ 368 (8000 \text{ mA g}^{-1}) \\ 155 (7000 \text{ mA g}^{-1}) \\ 155 (7000 \text{ mA g}^{-1}) \\ 151 (7000 \text{ mA g}^{$	— 71% (200 cycles) 54% (100 cycles)	369 271
Na/V4F7-3F Na/V4P7	20 moi% natraaj-[C3C1Pyt1][r5A] Na[PF6]-EC/DEC 20 mol% Na[FSA]-[C3C1pyt1][FSA]	-	25 25 25 25	(86%) (88%) ()	155 (2000 mA g ⁻¹) 360 (8000 mA g ⁻¹) —	24% (100 cycles) 48% (100 cycles) SCD –	371
Na/InP Na/CuP2 Na/GeP	Na[FSA]–[C ₃ C ₁ pyrr][FSA] Na[FSA]–[C ₃ C ₁ pyrr][FSA] Na[FSA]–[C ₂ -C ₂ mrr][FSA]	0.8–1 0.8–1 0.8–1 0.8–1	30 30 30	470 @ 100 (—) 746 @ 50 (78.7%) 758 @ 50 (72.7%)		— 105% (100 cycles) SCD	368 368 368
Na/GeP ₅ Na/SiP Na/Sn-P,	Nat ESA C ₃ C ₁ pyr1[ESA] Nat ESA C ₅ C ₁ pyr1[ESA]		0 30 30 30 30	$\begin{array}{c} 660 \\$		SCD SCD	308 308 308 308 308 308
Na/Sh2 ₃ -graphene	Na[FSA]-[C ₃ C ₁ Pyrr][FSA] Na[ClO ₄]-PC Na[ClO ₄]-PC Na[ClO ₄]-EC/DEC Na[ClO ₄]-EC/DEC Na[ClO ₄]-EC/DEC Na[FSA]-[C,C,mrr][FSA]			$\begin{array}{c} 670 \\ 670 \\ 976 \\ 1007 \\ 1057 \\ 1008 \\ 100 \\ 82.5\% \\ 1100 \\ 100 \\ 82.6\% \\ 1100 \\ 10$	$ \begin{array}{c} 520 & (1000 \text{ mA } \text{g}^{-1}) \\ 250 & (3000 \text{ mA } \text{g}^{-1}) \\ 230 & (3000 \text{ mA } \text{g}^{-1}) \\ \hline \\ \hline \\ 340 & (1500 \text{ mA } \text{g}^{-1}) \\ 55 & (1500 \text{ mA } \text{g}^{-1}) \\ 150 & (1500 \text{ mA } \text{g}^{-1}) \\ 150 & (1500 \text{ mA } \text{g}^{-1}) \\ \end{array} $	112% (200 cycles) 	395 395 373 373
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Table 3 Specifications of negative electrode materials in IL electrolytes^a

Energy & Environmental Science

Cell configuration	Electrolytes	Mass loading/ mg cm ⁻²	Temp./ °C	Mass loading/ Temp./ 1st cycle discharge capacity $^{(1)}$ mg cm $^{-2}$ $^{\circ}$ C current (col. eff.)/mA h g $^{-1}$ $^{(2)}$ mA g $^{-1}$	Rate capability/ mA h g^{-1} (current) Cycleability	Cycleability	Ref.
Na/SnO ₂ -graphene	Na[ClO ₄]-PC/FEC Na[FSA]-[C ₃ C, pyrt][FSA] Na[ClO ₄]-PC/FEC Na[ClO ₄]-EC/PC Na[ClO ₄]-EC/PC	1-1.2	60 60 25 25	$ \begin{array}{c} 610 @ 50 (50\%) \\ 760 @ 50 (75\%) \\ 505 @ 20 () \\ 500 @ 20 () \\ 505 @ 20 () \\ 505 @ 20 () \end{array} $	$\begin{array}{c} 120 \ (1500 \ \text{mA} \ \text{g}^{-1}) \\ 420 \ (1500 \ \text{mA} \ \text{g}^{-1}) \\ 197 \ (1000 \ \text{mA} \ \text{g}^{-1}) \\ 310 \ (1000 \ \text{mA} \ \text{g}^{-1}) \\ 237 \ (1000 \ \text{mA} \ \text{g}^{-1}) \end{array}$	43% (100 cycles) 95% (100 cycles) 73% (100 cycles) 71% (100 cycles) 88% (100 cycles)	374
Na/Na2DBQ-OMC	Na[FSA]-[C3/C, IDYTT][FSA] Na[CIO4]-EC/PC/FEC Na[FSA]-[C3/C, IDYTT][FSA] Na[CIO4]-EC/PC Na[FSA]-EC/PC Na[FSA]-[C3/C, IDYTT][FSA] Na[FSA]-[C3/C, IDYTT][FSA] Na[FSA]-[C3/C, IDYTT][FSA]	0.3-0.6	25 60 22 22 60	$\begin{array}{c} 410 & @ 20 () \\ 505 & @ 20 () \\ 600 & @ 20 () \\ \\ 145 & @ 25 (43\%) \\ 277 & @ 25 () \end{array}$	$\begin{array}{c} 148 \ (1000 \ \text{mA g}^{-1}) \\ \hline & \\ 330 \ (1000 \ \text{mA g}^{-1}) \\ \hline & \\ & \\ & \\ 175 \ (300 \ \text{mA g}^{-1}) \end{array}$	99% (100 cycles) 59% (100 cycles) 96% (100 cycles) 38% (50 cycles) 88% (300 cycles) 88% (300 cycles)	384
a See Fig. 3 for abbrevia	a See Fig. 3 for abbreviation of ionic species in ILs. b mAh cm $^{-2}$.						

insignificant toxicity. Recent studies using organic electrolytes have shown promising performance for Na secondary batteries. The high capacity of 521 mA h g⁻¹ and good capacity retention of 88% after 200 cycles was achieved for NiTiO₃ (prepared by a hydrothermal process) in 1 mol dm⁻³ Na[PF₆]-EC/DMC (1:1, v/v).³³¹ The hexagonal crystal structure of NiTiO₃, which features zigzag open tunnels and mesoporosity, enabled this high rate performance.

A NASICON-type NaTi₂(PO₄)₃ (NTP) was investigated in 1 mol dm⁻³ Na₂SO₄ aqueous electrolytes.³³² Owing to the high Na⁺ conductivity of the NASICON structure and the nanoparticle conformation of NTP, high rate capability (103 mA h g⁻¹ at 2C) was attained. Another study on an NTP/graphene composite prepared by the sol-gel method revealed the reversible insertion/ extraction of Na⁺ involving the Ti⁴⁺/Ti³⁺ redox couple.³³³

The charge–discharge mechanism of anatase-TiO₂ was found to be rather interesting. It proceeds with the insertion of Na⁺ into the TiO₂ lattice, distorting the structure and subsequently forming Na_{0.69}TiO₂, Ti metal (Ti⁰), and O₂ by reduction.³³⁴ Although the formation of Ti metal is unfavourable according to the Gibbs energy calculation,³³⁵ the larger size of Na⁺ causing strain in the lattice might be responsible for triggering the disproportionation reaction. The aforementioned studies highlight the attention-grabbing properties of Ti-based compounds. Accordingly, the following studies highlight the performance of Ti-based compounds with IL electrolytes.

The charge-discharge behaviour of commercial anatase-TiO₂ in 10 mol% Na[TFSA]-[C₃C₁pyrr][FSA] has been reported.³³⁶ Even at 25 °C, TiO₂ showed good rate capability by providing 47 mA h g^{-1} at 2C (1C = 335 mA g^{-1}) (Fig. 12c). The electrochemical impedance spectroscopy (EIS) results, obtained at the fully desodiated state of the TiO₂/Na cell, are shown in Fig. 12d. The semicircle gradually decreases in size from the 1st to the 30th cycle and reaches a stable value of approximately 70 Ω , suggesting a stabilised electrode/electrolyte interface. TiO₂/C prepared by heat treatment of citric acid with commercial anatase-TiO2 exhibited remarkable capacity retention of 79% after 1000 cycles and excellent rate performance at 90 °C in 20 mol% Na[FSA]-[C₃C₁pyrr][FSA].²²⁸ To increase the reversible capacity, Fe₂O₃ was incorporated into TiO_2 by heat treatment of commercial TiO_2 and $Fe(NO_3)_3$. A high reversible capacity of 386 mA h g^{-1} at 10 mA g^{-1} and 80% capacity retention after 300 cycles at 200 mA g^{-1} were achieved at 90 °C using 20 mol% Na[FSA]-[C₃C₁pyrr][FSA].³³⁷ An FeTiO₃/ C electrode prepared by a two-step heat treatment method from the same precursor as that used above and sucrose as the carbon source showed no discernible capacity fading over 2000 cycles at 90 °C, which is the best cycleability for negative electrodes in ILs currently reported.338

In addition to fabricating composite materials, doping with aliovalent ions has been demonstrated to be an effective technique for improving the electrical conductivities of materials, which has a positive effect on the performance of batteries.^{339,340} For example, Nb-doped rutile-TiO₂ (Ti_{0.94}Nb_{0.06}O₂) prepared by a sol–gel method was investigated as a negative electrode material for Na secondary batteries using 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA]

Table 3 (continued)



Fig. 12 Electrochemical behaviours of hard carbon (HC) and Ti-based negative electrode materials in ILs. (a) Galvanostatic charge–discharge profiles of HC at -10 to 90 °C at a current density of 20 mA g⁻¹ in 20 mol% Na[FSA]–[C₃C₁pyrr][FSA]. (b) Rate capabilities of the HC electrode at 25, 50, and 90 °C, corresponding to (a). Reproduced with permission.³²⁹ Copyright 2015, Elsevier. (c) Rate capability of anatase-TiO₂-based electrodes in 10 mol% Na[TFSA]–[C₃C₁pyrr][FSA]. After two formation cycles at 0.01C (0.1 to 2.0 V), various C rates (1C = 335 mA g⁻¹) were applied to the electrode. (d) Impedance spectra for the 1st, 15th, and 30th cycles recorded for a fully charged (*i.e.*, desodiated) TiO₂/Na cell. Reproduced with permission.³³⁶ Copyright 2016, Elsevier.

and 1 mol dm⁻³ Na[FSA]-PC.³⁴¹ A high capacity retention of 97% was observed with Na[FSA]–[C₃C₁pyrr][FSA] at 60 °C. Flammability tests demonstrated no ignition of Na[FSA]– [C₃C₁pyrr][FSA] even at 300 °C, whereas explosive combustion of Na[FSA]-PC was observed at 150 °C, demonstrating the benefit of ILs in terms of safety as well as performance.

3.2.3 Alloy materials. Alloy materials are known for their outstanding theoretical capacities. They are generally costeffective, easy to prepare or obtain, and environmentally benign. However, their high capacity is inevitably accompanied by huge volume changes, causing heavy stress in the active material, which results in pulverisation of electrodes. To counter this issue, nanostructuring,³⁴² forming intermetallic compounds, surface modification with Al₂O₃,³⁴³ and fabricating composites with carbon materials³⁴⁴ are well-employed strategies. Moreover, electrolyte and interfacial properties also play a key role in improving their performances.^{342,345,346} Silicon has been intensively investigated for LIBs owing to its large Li⁺ ion accommodation (3579 mA h g^{-1} for Li₁₅Si₄). A comparative study of SEIs formed in 1.2 mol dm⁻³ Li[FSA]-[C₃C₁pyrr][FSA] and 1 mol dm⁻³ Li[PF₆]-EC/DEC (1:1, v/v) using Si negative electrodes revealed the high stability of the SEI formed, which was attributed to the volume change suppression and negligible mass change during delithiation observed by electrochemical quartz crystal microbalance analysis.³⁴⁷ Although there are many studies of Si with Li, limited studies have been performed with Na. Theoretically, Na forms NaSi at the maximum capacity (~954 mA h g⁻¹) but in practical cases, a reversible capacity of ~ 300 mA h g⁻¹ (Na_{0.29}Si) has been recorded so far.³⁴⁸ Numerous materials have been investigated as candidates for alloy materials in Na secondary batteries. These have been summarised in several review articles.^{349–351} Most of these studies adopted organic electrolytes, and there is a very limited number of reports using ILs. This section addresses the reports on alloy-based materials with ILs.

Fig. 13 shows the charge–discharge behaviours of alloy and phosphide-based negative electrode materials for Na secondary batteries. Sn exhibits high capacity (~847 mA h g⁻¹ for Na₁₅Sn₄), low cost (~20.8 \$ kg⁻¹),³⁵² and non-toxicity. Several intermetallic Sn–M compounds (M = Fe, Ni, Cu) with 56 mol% Na[FSA]–K[FSA] IL at 90 °C have been investigated for Na secondary batteries.^{353–355} The intermetallic compounds were synthesised by electrodeposition followed by annealing for different periods of time. The annealing time was optimised with respect to capacity and cycleability. Longer annealing times caused the evolution of inactive intermetallic compounds, which results in increased buffering at the cost of capacity. The charge–



Fig. 13 Electrochemical behaviours of alloy- and conversion-based negative electrode materials in ILs. (a) Charge–discharge curves for the 1st, 10th, 100th, and 1000th cycles of an annealed Sn-Cu film at a current density of 84.7 mA (g-Sn) ⁻¹ and 90 °C in 56 mol% Na[FSA]–K[FSA]. Reproduced with permission.³⁵³ Copyright 2014, Elsevier. (b) Rate capabilities of a CuP₂/C composite in 20 mol% Na[FSA]–[C₃C₁pyrr][FSA] at 25 and 90 °C. Reproduced with permission.³⁶⁹ Copyright 2018, Wiley.

discharge curves for an Sn–Cu film annealed for 4 h show four plateaus, even after 1000 cycles at 90 °C, which highlights the excellent cycleability of this system (Fig. 13a). This result was attributed to the buffering effect of the inactive component Cu_3Sn and the randomly dispersed less electrochemically active component Cu_6Sn_5 .

Phosphorus is another alloyable material, providing a large theoretical capacity in Na secondary batteries ($\sim 2596 \text{ mA h g}^{-1}$) owing to the formation of Na₃P. Red P with carbon composites

has been extensively investigated in the past decade for Na secondary batteries with organic electrolytes^{356–358} but there is only one report concerning its use in an IL.⁸⁰ The red P/carbon composite exhibited stable cycleability in 0.25 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] but the capacity faded quickly in an organic electrolyte. The reason for this behaviour is discussed below in the SEI layer section. Black P, being an extraordinary conductor of electricity and thermodynamically stable under ambient conditions, is widely applied in optoelectronics,



Fig. 14 Comparison of the electrochemical performances of Sb_2S_3 in IL and organic electrolytes. SEM micrographs of (a) the as-prepared Sb_2S_3 / graphene electrode and the electrode after the first charge–discharge cycle in (b) 1 mol dm⁻³ Na[ClO₄]-PC/FEC and (c) 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] electrolytes at 25 °C. Effects of temperature on (d) reversible capacity at 50 mA g⁻¹, (e) capacity retention at 1500 mA g⁻¹ (compared to values at 50 mA g⁻¹), and (f) capacity retention after 100 cycles for Sb_2S_3 /graphene electrodes measured in 1 mol dm⁻³ Na[ClO₄]-PC/FEC and 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA]. Reproduced with permission.³⁷³ Copyright 2017, American Chemical Society.

electronics,³⁵⁹ and batteries.^{360,361} Black P was synthesised by heating red phosphorus at 800 °C under high pressure and the effect of electrolyte additives (FEC and VC) in achieving good SEIs was investigated. In another study, black P prepared by the ball-milling of red P showed the best cycleability at 30 °C in ILs among the three different electrolytes: 1 mol dm⁻³ Na[TFSA]– $[C_{(10201)}C_1pyrr]$ [FSA], 1 mol dm⁻³ Na[TFSA]– $[C_2C_1im]$ [FSA], and 1 mol dm⁻³ Na[TFSA]-PC.³⁶² These results indicate the potential of ILs with alloy-based negative electrodes for better cycleability and safer performance.

3.2.4 Conversion-based materials. Conversion based materials are a class of materials that provide high capacities with tunable properties such as electrical conductivity and air stability. Typically, the conversion reaction proceeds in the following way (eqn (12)):

$$\mathbf{M}_{a}\mathbf{X}_{b} + (b \times c)\mathbf{N}\mathbf{a} \rightleftharpoons a\mathbf{M} + b\mathbf{N}\mathbf{a}_{c}\mathbf{X}$$
(12)

M can be both an inactive or active metal (Sn, In) or metalloid (Si, Ge, Sb) for sodiation. X is an electronegative non-metal, *e.g.*, P, O, S, or F. The reaction potential *vs.* Na⁺/Na is found to be generally governed by the ionicity of the MX bond between the metal M and the non-metal X (F (~1.1 V) > O (~0.9 V) > S (~0.7 V) > N (~0.6 V) > P (~0.4 V) > H (~0.25 V)).^{335,363,364} Overall electrochemical performance is highly dependent on the choice of elements, morphology, and operation conditions. The main challenge for conversion-based materials is to attain good cycleability and rate performance. The selection and combination of metal and non-metal as well as optimisation of the electrolyte must be performed in order to achieve good performance.

Metal phosphides are emerging as a separate class of negative electrodes for Na secondary batteries. In addition to showing high capacities, they exhibit low overpotential as conversion-based materials due to the low electronegativity of P, as mentioned above. The sodiation of metal phosphides usually results in the formation of Na₃P and separation of metal nanoparticles, which improves electronic conduction. However, the large volume change causes electrode pulverisation, which causes severe capacity deterioration. Similar to the situation with alloybased materials, carbon composites, nanostructuring, and morphology and electrolyte optimisation are some of the strategies used to improve performance. Although transitionmetal phosphides have been highly investigated in organic electrolytes, 365-367 there are limited reports using ILs. A variety of binary phosphides (GeP₅, GeP, SiP, LaP, InP, and CuP₂) were evaluated in 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] at 30 °C.³⁶⁸ The best cycleability was observed for InP, which was attributed to the provision of electronic conductivity and maintenance of interparticle contact by the adjustment of NaIn morphology in response to volume expansion by Na₃P.

A CuP₂/C carbon composite was investigated using 20 mol% Na[FSA]–[C₃C₁pyrr][FSA] at 25 and 90 °C.³⁶⁹ Owing to the high ionic conductivity of the IL and enhanced electrode kinetics at 90 °C, good rate capability was achieved (Fig. 13b). Earlier studies with organic electrolytes revealed the reversible formation of CuP₂³⁷⁰ but in this case, the product remained amorphous after

the first sodiation. The authors proposed that the chargedischarge mechanism can be changed by changing the electrolyte and operating conditions. An early transition-metal phosphide, V₄P₇, investigated recently,³⁷¹ shows a peculiar insertion-based mechanism as revealed by ex situ XRD. Moderate capacities of 270 and 470 mA h g⁻¹ at 25 and 90 $^\circ$ C, respectively, were observed using 20 mol% Na[FSA]-[C3C1pyrr][FSA] as the electrolyte. Moreover, high rate capability and good cycleability were also attained at 90 °C. In the same study, a V₄P₇ P composite (V₄P₇/5P) was also investigated in 20 mol% Na[FSA]-[C₃C₁pyrr][FSA] and 1 mol dm^{-3} Na[PF₆]-EC/DEC (1:1, v/v). Stable cycleability over 100 cycles was observed for the IL, whereas the capacity faded after six cycles for the organic electrolyte. This observation was supported by EIS and SEM observations. After 10 cycles, a significant decrease in impedance and a similar particle size compared to the pristine state were observed in the IL case, whereas large impedance and enlargement of particle size were observed for the organic electrolyte.

As discussed above, the metal and non-metal can be varied to tune the properties. Although sulfides and oxides have larger overpotentials than phosphides, they also show high capacities and rate capabilities. Usually, the reaction proceeds with the formation of Na_2X (X = S, O) and separation or alloying of the metal. Recently Sn- and Sb-based materials have become popular owing to their alloying properties and relative abundance. Sb₂O₃ prepared by electrostatic spray deposition provided a reversible capacity of 331 mA h g⁻¹ with stable cycleability and good rate capability using 1 mol dm⁻³ Na[PF₆]-EC/DEC/PC (4:4:2 by vol) with 5% FEC.³⁷² The sodiation product NaSb(OH)6 was observed by ex situ XRD and transmission electron microscopy (TEM) and appeared to be amorphous, which is a common phenomenon in conversion-based materials. The sulfide analogue Sb₂S₃ was investigated using 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] IL and various organic electrolytes at 25 and 60 °C.³⁷³ High first cycle discharge capacities of 710 and 660 mA h g^{-1} were delivered with 1 mol dm⁻³ Na[ClO₄]-PC/FEC and 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] at 25 °C, respectively.

Fig. 14a shows SEM images of the pristine Sb₂S₃ and those after charge-discharge in 1 mol dm⁻³ Na[ClO₄]-PC/FEC (Fig. 14b) and 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] (Fig. 14c). It can be clearly seen that after charging and discharging in 1 mol dm⁻³ Na[ClO₄]-PC/FEC, the particles are covered by thick SEI layers, whereas in Fig. 14c, the morphology is retained, demonstrating formation of thin and robust SEI layers. In Fig. 14d-f, the capacity, rate capability, and cycleability are compared at 25 °C and 60 $^{\circ}C$ for 1 mol dm⁻³ Na[ClO₄]-PC/FEC and 1 mol dm⁻³ Na[FSA]– $[C_3C_1pyrr]$ [FSA]. It can be observed that at 60 °C, the 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] clearly outperforms the 1 mol dm⁻³ Na[ClO₄]-PC/FEC in all performance aspects. The large difference in the rate performance at 60 °C is attributed to the elevated side reaction in 1 mol dm^{-3} Na[ClO₄]-PC/FEC. Conversely, 1 mol dm⁻³ Na[ClO₄]-PC/FEC delivered higher capacity and better rate performance than 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] at 25 °C due to the high viscosity in the IL system, which restricts Na ion diffusion.

A SnO₂/graphene composite prepared by supercritical-CO₂-assisted synthesis was analysed in similar organic and IL electrolytes.³⁷⁴ High rate capability with the incorporation of EC in PC at 25 °C and degradation of rate with the addition of FEC were demonstrated. At the elevated temperature of 60 °C, high reversible capacities of 600 and 330 mA h g⁻¹ at 20 and 1000 mA g⁻¹, respectively, were obtained with 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA].

3.3 Summary and future investigations

3.3.1 Positive electrode aspect. Vanadium-based fluoride phosphates ("fluorophosphate" is the wrong nomenclature because there are no P–F bonds) such as $Na_3V_2(PO_4)_2F_3$ and $NaVPO_4F$ are good candidate materials in terms of high energy density and have been vigorously studied in organic solvents, but not in IL electrolytes. Fe- and Mn-based materials are excellent candidates in terms of economic and resource considerations. Na_2FePO_4F has a high theoretical capacity of 124 mA h g⁻¹ and

high operation voltage of 3.5 V, as does Na_2MnPO_4F .^{375,376} Thus, these materials would be promising if their cycle performances and rate capacities could be improved.

Prussian blue analogues have not yet been tested using IL electrolytes. For example, $Na_4Fe(CN)_6$, which is made from naturally abundant resources, could be an interesting material that simultaneously achieves good capacity and moderate working potential.³⁷⁷

Finally, organic compounds are very attractive materials for Na secondary batteries in terms of sustainability and diversity.³⁷⁸ One of the most important advantages of organic compounds is their chemical designability and tunable functionality. Thus, various redox-active organic electrodes are currently known. Nevertheless, organic electrode materials for Na secondary batteries have not been well investigated in IL electrolytes. A study on the organic electrode material poly((3,4)-ethylenedioxythiophene)-biopolymer in 20 mol% Na[TFSA]–[C₄C₁pyr][TFSA] and Na[FSA]–[C₂C₁im][FSA] IL electrolytes indicated their stable electrochemical



Fig. 15 Temperature effects on the charge–discharge behaviours of selected positive electrode materials in ILs. Charge–discharge curves of (a) a Na/ 20 mol% Na[FSA]–[C₂C₁im][FSA]/carbon coated Na₃V₂(PO₄)₃ cell at -30 to 25 °C. Reproduced with permission.¹⁶⁴ Copyright 2018, Wiley-VCH; (b) Na/ 20 mol% Na[FSA]–[C₃C₁pyrr][FSA]–NaCrO₂ cell at -20 to 90 °C. Reproduced with permission.¹⁶⁶ Copyright 2015, The Electrochemical Society of Japan. Charge–discharge curves for a Na/30 mol% Na[FSA]–[C₂C₁im][FSA]–maricite NaFePO₄ cell at (c) 25 °C and (d) 90 °C. Reproduced with permission.¹⁰³ Copyright 2018, Elsevier.

performance.³⁷⁹ In another recent study, calix[4]quinone and 5,7,12,14-pentacenetetrone organic-based positive electrode materials were investigated in a 0.3 mol dm⁻³ Na[TFSA]-[C₃C₁pyrr][TFSA] IL electrolyte.³⁸⁰ A Na/calix[4]quinone halfcell exhibited the very high revisable capacity of 406 mA h g^{-1} at 20 mA g^{-1} and excellent cycleability with a stable capacity retention of 99.7% after 300 cycles. A Na/5,7,12,14-pentacenetetrone half-cell with a 0.3 mol dm⁻³ Na[TFSA]–[C₂C₁pvrr][TFSA] IL electrolyte exhibited similarly good performance with a capacity of 245 (30 mA g^{-1}). The authors insisted that these improvements were achieved using the IL electrolyte because it overcame problems associated with the poor dissolution of quinone compounds.³⁸⁰ Thus, IL electrolytes may be useful to overcome problems associated with organic compounds such as suppressing the dissolution of active materials that typically occurs in aprotic solvents and ameliorating the low intrinsic electronic conductivity by elevated temperature operation.378,381-383

3.3.2 Negative electrode aspect. Finding a material that simultaneously provides high capacity, low voltage, high cycleability, and high rate capability is a challenging task. However, combining materials with already known properties may be a solution. For example, HC is the most extensively studied and suitable negative electrode material, but its reversible capacity is only moderate. Incorporating other high-capacity materials like

P and Sn into HC may be an effective approach to creating candidate materials. Organic compounds are versatile, renewable, low-cost, and abundant. However, there is only one report on the quinone-based material 2,5-disodum-1,4-benzoquinone (Na₂DBQ) with ordered mesoporous carbon (OMC) as a negative electrode material for Na secondary batteries containing ILs³⁸⁴ but there are many active organic compounds such as disodium terephthalate^{385,386} that provide excellent performance in organic electrolytes. Therefore, organic compounds may be a prospective class of materials for future investigation.

There are many candidate materials for Ti-based compounds. For example, Na titanates, especially $Na_2Ti_3O_7^{387,388}$ and $Na_2Ti_6O_{13}$,³⁸⁹ show lower redox potentials than other Ti-based compounds and are highly investigated as negative electrode materials in organic electrolyte systems. Furthermore, the SEI layer should be studied in detail along with ILs. In the study on red P mentioned in this review,⁸⁰ it included a large amount of binder (~20 wt%) to facilitate good cycleability, but the amount and type of binder also influence the decomposition of the electrolyte. Therefore, an investigation into the SEI layer should be performed without (or with a minimal amount of) binder. In addition, the surface morphology of the SEI layer should be inspected by characterisation tools like atomic force microscopy (AFM), TEM, and cross-sectional focused ion beam SEM.



Fig. 16 Charge–discharge curves for (a) Na/1 mol dm⁻³ Na[PF₆]-EC/DEC (1:1, w/w)/Na₄Ni₃(PO₄)₂(P₂O₇)/C, cut-off voltages: 3.0–4.9 V, (b) Na/10 mol% Na[TFSA]–[C₃C₁pyrr][FSA]/Na₄Ni₃(PO₄)₂(P₂O₇)/C, cut-off voltages: 3.0–5.1 V, and (c) *ex situ* X-ray diffraction patterns for a Na₄Ni₃(PO₄)₂(P₂O₇) electrode during charge–discharge. Reproduced with permission under the Creative Commons Attribution 4.0 International License.²⁶⁸



Fig. 17 Electrochemical performance of carbon-coated $Na_3V_2(PO_4)_3$ in an IL. (a) Mass loading dependence of rate performance for Na/40 mol% Na[FSA]–[C_2C_1 im][FSA]/carbon-coated $Na_3V_2(PO_4)_3$ cells at 25 and 90 °C. Reproduced with permission.²⁸³ Copyright 2019, American Chemical Society. (b) Superfast rate performance test results for a Na/50 mol% Na[FSA]–[C_2C_1 im][FSA]/carbon-coated $Na_3V_2(PO_4)_3$ cell at 90 °C, and (c) electrochemical impedance spectra for Na/50 mol% Na[FSA]–[C_2C_1 im][FSA]/carbon coated $Na_3V_2(PO_4)_3$ cells. Reproduced with permission.¹⁶⁴ Copyright 2018, Wiley-VCH.

Recently, there have been reports on volume change suppression of Si-based negative electrodes for LIBs in ILs.^{390–393} Similar results were reported for $V_4P_7/5P$ in Na secondary batteries,³⁷¹ but an understanding of the key mechanism behind the phenomenon is lacking.

4. Benefits of using IL electrolytes

As previously stated, organic electrolytes such as EC, PC, DEC, DMC, and 1,2-dimethoxyethane (DME) coupled with the Na salts $Na[ClO_4]$ and $Na[PF_6]$ are routinely used for Na secondary battery studies and to achieve good performance. However, the safety issues surrounding secondary batteries are mostly concerned with the volatility and flammability of organic electrolytes. As Na secondary batteries are mainly intended as large-scale batteries for ESSs and electric vehicles, the safety problems become more significant as the battery size increases. In addition to the safety benefits, recent studies have revealed that ILs bring excellent cycle performance and superior rate capability. This is because IL electrolytes can form more stable SEIs that enable more facile Na diffusion and can also form a stable passivation layer on Al current collectors, preventing Al corrosion.³⁹⁴ Furthermore, ILs enable operation at elevated temperatures, which can facilitate Na diffusion in the electrode and electrolyte. In this section, some recent electrochemical discoveries and interesting accomplishments obtained by utilising IL electrolytes are presented.

4.1 Thermal and electrochemical stability

The use of ILs extends the temperature limits and cut-off voltage range upon electrochemical measurements. This flexibility in testing conditions is advantageous not only for conducting more comprehensive evaluations of well-developed materials but also for identifying the optimal working environments for new materials.

Fig. 15 shows the temperature effects on the charge–discharge performances of some positive electrode materials. In this context, $Na_3V_2(PO_4)_3$ and $NaCrO_2$ were evaluated in the wide temperature

range -30 to 90 °C using 20 mol% Na[FSA]–[C₂C₁im][FSA] and Na[FSA]–[C₃C₁pyrr][FSA] IL electrolytes, respectively, as shown in Fig. 12a and b.^{164,166} The Na₃V₂(PO₄)₃ cell provided stable reversible performance across the whole temperature range and achieved capacities of 100.8, 78.1, and 57.6 mA h g⁻¹ at 25, -10, and -20 °C, respectively (Fig. 15a).¹⁶⁴ Charge–discharge testing of the NaCrO₂ cell was performed between -20 and 90 °C, and it also exhibited very stable reversible capacities (Fig. 15b). With a decrease in temperature, the polarisation of the cell increases and the reversible capacity decreases as a consequence of the increase in internal cell resistance, which is due to the reduced ionic conductivity of the electrolyte, retarded Na diffusion in the electrolytes could support a comprehensive operation temperature range for Na secondary batteries (Fig. 5).

Research on maricite-NaFePO4 has yielded interesting electrochemical results. As noted earlier, maricite-NaFePO₄ is generally considered to be an electrochemically inactive material because of its structural arrangement, which features FeO₆-FeO₆ (edgesharing) and PO₄-FeO₆ (corner-sharing) units that hinder effective diffusion of Na⁺.²⁸⁸ The elevation of operating temperature possible using IL electrolytes enhanced Na⁺ diffusion in this structural frame, which effectively allows maricite-NaFePO4 to be electrochemically active. Sodiation/desodiation into/from maricite-NaFePO4 were realised upon elevating the operation temperature.^{102,103} The charge-discharge of maricite-NaFePO₄ in a Na[FSA]-[C2C1im][FSA] IL electrolyte demonstrated a drastic increase in reversible capacity from only 23 mA h g^{-1} at 25 °C (Fig. 15c) to 107 mA h g⁻¹ at 90 °C (Fig. 15d). The *ex situ* XRD patterns of desodiated maricite-NaFePO₄ continuously shifted to higher 2θ angles, indicating a decrease in *d*-spacing. Fe K-edge XANES spectra demonstrated the reduction of Fe³⁺ to Fe²⁺ by sodiation.¹⁰³

Fig. 16 illustrates the electrochemical behaviours of a $Na_4Ni_3(PO_4)_2(P_2O_7)/C$ electrode in organic and IL electrolytes. The Ni-based polyanionic compound $Na_4Ni_3(PO_4)_2(P_2O_7)/C$ was expected to exhibit a high operating voltage based on its Ni^{3+}/Ni^{2+} redox activity. Unfortunately, when a 1 mol dm⁻³



Fig. 18 Cycle performances of selected electrode materials in ILs. (a) Cycle performance of a Na/30 mol% Na[FSA]–[C₂C₁im][FSA]/Na₂FeP₂O₇ cell at 90 °C. Reproduced with permission.¹⁸⁰ Copyright 2016, Elsevier. (b) Cycle performances of Na/1 mol dm⁻³ Na[PF₆]-PC/Na₃V₂(PO₄)₃ and Na/0.1, 0.25, 0.5, and 0.75 mol dm⁻³ Na[PF₆]–[C₂C₁im][TFSA]/Na₃V₂(PO₄)₃ cells. Reproduced with permission.²⁸⁶ Copyright 2018, Elsevier. (c) Cycle performances of Na/0.5 mol dm⁻³ Na[PF₆]-PC/Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ and Na/10 mol% Na[TFSA]–[C₄C₁im][TFSA]/Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ cells. Reproduced with permission.²⁶⁷ Copyright 2014, Elsevier. (d) Cycling performances and coulombic efficiencies of an Sn₄P₃ electrode in 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA], 1 mol dm⁻³ Na[FSA]–[C₂C₁im][FSA], and 1 mol dm⁻³ Na[FSA]-PC at 30 °C. Reproduced with permission.³⁹⁵ Copyright 2017, American Chemical Society. (e) Capacity and coulombic efficiency for three-dimensional CF electrodes measured at 100 mA g⁻¹ in 1 mol dm⁻³ Na[CIO₄]-EC/DEC (1:1, v/v) and 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] at 25 and 60 °C. Reproduced with permission.³⁹⁷ Copyright 2018, Elsevier.

Na[PF₆]-EC/DEC organic solvent electrolyte was used, the cell could only be charged to 4.9 V. The first charge capacity was approximately 97 mA h g^{-1} , which involves 2.3 equivalents of Na based on the capacity during the desodiation process.²⁶⁸ However, only 0.9 equivalents of Na were reversibly sodiated into the composite, corresponding to a coulombic efficiency for the first cycle of about 40% (Fig. 16a). This may indicate electrolyte decomposition at such a high potential. Conversely, this material could be charged up to 5.1 V in 10 mol% Na[TFSA]-[C3C1pyrr][FSA] IL and the composite was able to undergo deeper (de)sodiation; 2.7 and 1.3 equivalents of Na were desodiated and sodiated, respectively, during the first cycle (Fig. 16b), suggesting the possible high stability of the IL at high potentials. The ex situ XRD patterns (Fig. 16c) indicate that neither phase change nor irreversible structural reorganisation occurred during the charge-discharge process, confirming the robustness of this material. Although maricite-NaFePO4 and Na4Ni3(PO4)2(P2O7)/ C may appear to be less appealing for practical use owing to the limited achievable capacity, these studies highlight the possibility of using IL electrolytes for electrode studies with new materials.

4.2 Rate performance and power density

Rate capability and power density are significantly improved by the elevated temperature operation available using ILs owing to the improved interfacial process between electrodes and electrolytes as well as ionic conductivity, as mentioned above. Fig. 17 shows the mass loading and temperature effects on the electrochemical behaviour of Na3V2(PO4)3 in 40 mol% Na[FSA]- $[C_2C_1im]$ [FSA]/carbon.^{164,283} The synergy of Na₃V₂(PO₄)₃ and the IL electrolyte leads to reasonable rate performance at 25 °C. The limited performance in terms of low electronic conductivity at room temperature was easily overcome by carbon coating the electrode material. However, the rate performance dramatically deteriorates at 25 °C as mass loading of the electrode becomes large because the rate performance is very sensitively related to the applied geometric current density (mA cm^{-2}). For example, in the cases of 3.0 mg cm^{-2} at 10C and 6.0 mg cm⁻² at 5C, the capacity retentions are similar (68.3% and 63.2%, respectively, at 25 °C) because the same geometric current density of 3.53 mA cm⁻² is applied with doubled mass loading (Fig. 17a). Rate performance is maintained even with a high mass loading of 6 mg cm⁻² for $Na_3V_2(PO_4)_3$ at 90 °C (Fig. 17a). The capacity retentions at 90 °C for 3.0 mg cm⁻² at 10C and 6.0 mg cm⁻² at 5C are 96.1% and 97.3%, respectively.²⁸³ Thus, the true rate performance is dependent on geometric current density. Intermediate temperature operation can certainly improve the rate performance, as evidenced in Fig. 17b, which shows that the discharge capacities increase after adjusting the cut-off voltages, giving capacity retentions of 69%, 58%, 36%, and 19% at 500C, 1000C, 1600C, and 2000C, respectively. This level of rate performance could never be achieved by organic solvent electrolytes around room temperature.¹⁶⁴ The EIS measurements shown in Fig. 17c reveal that the interfacial resistance, which is indicated by the semicircle (characteristic

frequencies of 60–700 Hz), drastically decreases as the operating temperature increases compared to the bulk resistance (Fig. 17c, inset).¹⁶⁴

4.3 Cycle performance and coulombic efficiency

Numerous studies have reported that IL electrolytes exhibit improved cycling performance over a wide temperature range.^{84,164,180,267,283,286} It is widely acknowledged that IL electrolytes can provide a uniform and robust SEI layer on reactive Na metal (see Section 2.4 for details).

Fig. 18 shows the cycle performances of selected electrode materials in IL electrolytes. Cycling tests of a Na/NaFePO₄ cell initially confirmed the better cycle performance using a Na[TFSA]– $[C_4C_1pyrr]$ [TFSA] IL (Na[FSA] = 0.1, 0.5, and 1 mol dm⁻³) electrolyte than that using a 1 mol dm⁻³ Na[ClO₄]-EC/DEC organic solvent electrolyte at 50 °C.⁸⁴ The capacity retentions after 100 cycles were 62% and 87% with the organic and IL electrolytes, respectively. Furthermore, Na[TFSA]– $[C_4C_1pyrr]$ [TFSA] IL electrolyte containing 0.5 mol dm⁻³ Na[TFSA] salt gave better cycle performance than that with 0.1 and 1 mol dm⁻³ Na[TFSA].⁸⁴ The authors reported that the organic solvent electrolyte at 50 °C presented undesirable electrode–electrolyte interactions and evaporation of the electrolyte, whereas the thermally stable IL electrolyte provided higher Na⁺ supply as the temperature increased without the above-mentioned drawbacks.⁸⁴

The extended cycle performance of a Na/Na[FSA]–[C₂C₁im][FSA]/ Na₂FeP₂O₇ cell was assessed at 90 °C (Fig. 18a).¹⁸⁰ A high capacity retention of 93% was attained for 1500 cycles in this cell with an average coulombic efficiency of 99.9%. Another study using Na₃V₂(PO₄)₃ with the same IL electrolyte also revealed very stable cycleability. The capacity retention was 99.9% after 300 cycles at 25 and 90 °C at 1C (117 mA g⁻¹). Furthermore, a faster current cycle test at 20C and an elevated temperature of 90 °C revealed a capacity retention of 89.2%, even after 5000 cycles.¹⁶⁴ This extremely good cycle performance was due to the combination of the superior performance of this positive electrode material and stable Na metal deposition–dissolution at the counter electrode in the IL electrolyte at 90 °C (see Fig. 8b for Na metal deposition–dissolution behaviour).²⁴⁶

A study on $Na_3V_2(PO_4)_3$ in $Na[PF_6]-[C_4C_1im][TFSA]$ ($Na[PF_6] =$ 0.1, 0.25, 0.5, and 0.75 mol dm^{-3}) IL electrolytes indicated that optimising the salt concentration of the IL electrolyte could improve capacity and cycle performance at the same time.²⁸⁶ As shown in Fig. 18b, 0.25 mol dm⁻³ Na[PF₆]-[C₄C₁im][TFSA] exhibited a discharge capacity of 107.2 mA h g⁻¹ and retained 104 mA h g^{-1} after 40 cycles at a current density of 50 mA g^{-1} .²⁸⁶ The discharge capacity and capacity retention with the IL were superior to those with 1 mol dm^{-3} Na[PF₆]-PC. The authors suggested that the IL electrolyte forms a better SEI on $Na_3V_2(PO_4)_3$ than the organic solvent electrolyte and demonstrated that the concentration of Na[PF₆] is an important parameter for electrochemical performance.²⁸⁶ It should be noted that the (de)sodiation process can be promoted by supplying a sufficient amount of Na⁺ while excessive Na⁺ can negatively affect electrochemical performance owing to the decreased ionic conductivity of the electrolyte.

Another study on Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ reported that the cycleability with an IL electrolyte was superior to that with an organic solvent electrolyte at 20 °C,²⁶⁷ which clearly indicates that IL electrolytes present more stable cycling behaviour, not only at elevated temperature but also at room temperature, than the organic solvent electrolyte. Cycling of Na045Ni022Co011Mn066O2 in 0.5 mol dm⁻³ Na[PF₆]-PC and 10 mol% Na[TFSA][C₄C₁pyrr][FSA] at 12 mA g^{-1} also revealed improved cycleability and coulombic efficiency with the IL electrolyte (Fig. 18c).²⁶⁷ The low initial coulombic efficiency was improved after several cycles for both cells. The Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ cell with the organic solvent electrolyte exhibited a capacity retention of only 40% (approximately 90 mA h g^{-1}) after 100 cycles with an average coulombic efficiency of 97.8%, whereas the cell with the IL electrolyte showed 80% capacity retention (discharge capacity of 177 mA h g^{-1}) after 100 cycles with a coulombic efficiency of 98.7%. The authors claimed that the rapid capacity fading in both electrolytes was due to the irreversibility of the P2-O2 phase transition of Na_{0.45}Ni_{0.22}Co_{0.11}Mn_{0.66}O₂ in the organic solvent electrolyte. However, the dissolution of Mn ions in the IL electrolyte is restricted, making the Na045Ni022Co011Mn066O2 with IL much more electrochemically reversible and stable.²⁶⁷

With respect to negative electrode materials, Sn_4P_3 is a special case in which remarkable cycleability (112% for 200 cycles) and high reversible capacity were achieved with 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] at 30 °C (Fig. 18d).³⁹⁵ The charge–discharge mechanism here was similar to that for InP, and the reason for the better performance in the IL was proposed to be the homogeneity and stability of the SEI layer. Moreover, Sn_4P_3 exhibits a reversible capacity of 220 mA h g⁻¹ at 0 °C and high rate capability at 90 °C in the IL, which denotes its applicability in a wide temperature range.³⁹⁶ The P-rich phase SnP_3 was also investigated using 1 mol dm⁻³ Na[FSA]– [C₃C₁pyrr][FSA] at 30 °C, but severe capacity degradation was observed owing to the insufficient buffering power of SnP_3 , which is due to its low Sn content.³⁹⁵

The electrochemical performance of three-dimensional CFs prepared by high-speed homogenisation of graphene nanosheets and carbon nanospheres was analysed using 1 mol dm⁻³ Na[ClO₄]-EC/DEC (1:1, v/v) and 1 mol dm⁻³ Na[FSA]–[C₃C₁pyrr][FSA] IL at 25 and 60 °C (Fig. 18e).³⁹⁷ A capacity retention of 85% was observed after 250 cycles with the IL electrolyte at 25 °C, while only 70% capacity retention was observed with the organic electrolyte at room temperature. At 60 °C, the performance was significantly improved in the IL compared to that in the organic electrolyte. Moreover, the self-discharge in the organic electrolyte (15%) exceeded that in IL (8%) after five days of rest. Robust SEI layer formation owing to the decomposition of FSA⁻ was identified as the main reason for the improved cycleability.

4.4 Solid electrolyte interphase layer

Extensive efforts have been focused on the improvement of negative electrode materials in recent decades, but the investigation of electrode/electrolyte interfacial properties is equally important for ensuring the overall performance and safety of batteries. The SEIs formed by reductive decomposition of



Fig. 19 Hard X-ray photoelectron spectra of C1s, F1s, and P1s regions for P/AB/PANa electrodes (as-prepared (black), after first (red), and 20th (blue) cycles) in (a) 1.0 mol dm⁻³ Na[PF₆]-EC/DEC and (b) 0.25 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] electrolytes. Reproduced with permission.⁸⁰ Copyright 2017, Elsevier

conventional ester-based electrolytes are considered not sufficiently stable, leading to severe capacity degradation.³⁹⁸ The success of conventional HC negative electrodes depends on complicated electrolyte formulations and the introduction of SEI forming additives. Thus, for the development of negative electrodes, exploring compatible electrolytes is indispensable.

The performance and safety of a battery are highly dependent on SEI layer properties³⁹⁹ which are sensitive to parameters such as the salt concentration of the electrolyte,⁴⁰⁰ temperature,^{160,401} and the binder used.⁴⁰² Investigating the physical and chemical properties of SEI layers is very important for understanding the degradation mechanisms in batteries. However, such a study is often hindered because of the complex structure and nanoscale dimensions of the SEI layer, as well as the limited characterisation tools available. Moreover, owing to the higher reactivity and higher standard redox potential of Na than Li, transferring the knowledge of interfacial properties from LIBs is not always applicable. In addition, the solubility of SEIs in conventional electrolytes for Na secondary batteries aggravates self-discharging properties. Accordingly, self-discharge in Na secondary batteries using conventional 1 mol dm⁻³ Na[PF₆]-EC/DEC has been confirmed, whereas minimal self-discharge has been observed in corresponding LIBs.⁴⁰³ Therefore, it is imperative to investigate the interfacial properties for Na secondary batteries with other types of electrolytes.

Na metal is generally used as the counter electrode in halfcell configurations. However, the electrochemical behaviour of the Na electrode itself is generally not well investigated, causing uncertainty upon interpreting data and identifying the origin of cell degradation.⁴⁰⁴ The SEI layer formed on Na metal in ester-based EC/DEC and ether-based diglyme and tetraglyme with 1 mol dm^{-3} Na[PF₆] was investigated by XPS, and formation of a thin and stable SEI was observed in the diglyme case,⁴⁰⁵ whereas a thick and highly resistive film was formed in EC/DEC. In-depth XPS studies identified the formation of a uniform and solvent-impermeable layer consisting of inorganic components in glyme-based electrolytes containing Na2O and NaF, contrasting with the results for the mixed organic (RCH₂OCO₂Na) and inorganic SEI formed in EC/DEC. It was found that a mixed organic-inorganic SEI is prone to solvent permeation, resulting in dendrite growth.

Recently, the electrochemical properties of Na metal were studied in IL electrolytes (see Section 2.4 for details). Typically, a large interfacial resistance, presumably resulting from the surface film on the Na metal, is observed in the IL electrolytes.^{86,124,164} Such impedance can hinder the evaluation of the true performance of the target electrode in half-cell configurations. A systematic study on Na/Na symmetric cells using ILs showed that the overall cell resistance dramatically decreases with increasing operating temperature but increases with the Na salt content of the electrolyte.¹⁶⁴ The effects of electrolyte on the composition of the SEI layer formed on HC were extensively investigated by Eshetu et al.406 The anion of the electrolyte salt was reported to govern the characteristics of the SEI formed on HC. The organic content

decreases in the order $Na[PF_6] > Na[ClO_4] > Na[TFSA] >$ Na[FTFSA] > Na[FSA]. The effect of electrolyte on the thermal stability of the SEI and corrosion of the Al current collector were also studied by the same group. For a fixed solvent (EC/DMC), the onset temperature of the first exothermic peak (related to SEI cracking) follows the sequence $Na[ClO_4] < Na[PF_6] <$ $Na[TFSA] \approx Na[FTFSA] < Na[FSA].^{407}$ For the corrosion of the Al current collector, the Al dissolution or anion decomposition increases in the order $Na[PF_6] < Na[ClO_4] < Na[TFSA] <$ Na[FTFSA] < Na[FSA] in EC/DEC. However, Na[FSA] showed stability towards decomposition in the different solvents in the order EC/DEC < EC/DEC + 5% NaPF₆ < [C₄C₁pyrr][FSA]. This result shows that ILs impart the highest thermal stability to the SEI and suppress Al corrosion.

IL-derived SEIs can be distinctly different from those formed in organic electrolytes. Accordingly, the performances and interfacial properties of HC in a 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] IL and a conventional organic electrolyte (1 mol dm⁻³ Na[ClO₄]-EC/DEC (1:1, v/v)) were compared at 25 °C. The SEIs formed in the HC/HC symmetric cell showed a lower charge transfer resistance in the IL than that in the organic electrolyte. A dense SEI enriched with Na2CO3 and NaCO2R species was confirmed to form in the organic electrolyte by scanning electron microscopy (SEM) and XPS measurements. These species are considered to be the resistive components in SEI layers.³⁹⁷ Similar species were also observed using 1 mol dm^{-3} Na[ClO₄]-EC/DEC (1:1, v/v) in a three-dimensional carbon framework (CF), whereas polyolefin $((CH_2)_n)$ and S=C=O derived from FSA anions were found in the SEI layer formed in the IL.³⁹⁷

For alloy-based electrodes, a robust SEI layer is widely acknowledged as a crucial factor for realising good cycleability as these materials undergo significant volume change during sodiation/desodiation. Fig. 19 shows the compositional evolution of SEIs on red phosphorus cycled in organic electrolytes (Fig. 19a) and an IL (Fig. 19b) as revealed by a combination of hard X-ray photoelectron spectroscopy (HAXPES) and timeof-flight secondary ion mass spectroscopy (TOF-SIMS).⁸⁰ The TOF-SIMS analysis shows that the SEI layer generated in the IL electrolyte is composed of a uniform mixture of organic and inorganic components (with the inner surface primarily covered by inorganic species), whereas mainly inorganic species are observed in the organic electrolyte formed under the same conditions. Moreover, it can be clearly seen that the SEI composition dynamically changes with cycle number in the organic electrolyte, while the composition remains stable in the IL electrolyte. The homogeneous and complete coverage of the SEI layer on red phosphorus was considered to be the reason behind the improved reversible capacity and capacity retention in the IL electrolyte.

4.5 Full cell performance

Despite the fact that full cell studies are required to confirm the actual feasibility of Na secondary batteries using IL electrolytes, they are still in their infancy. There are several reports of full cell tests with ILs using coin-cells^{86,128,380,408-410} and one report for a prismatic cell.411

Although full cell tests were carried out using coin-cells, it is worthy of mention that the cycle performance was notably improved using IL electrolytes. A full cell with Na₃V₂(PO₄)₃ and HC positive and negative electrodes in 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] achieved the high energy density of 368 W h kg⁻¹ with good capacity retention of 75% after 100 cycles, while the test using a 1 mol dm^{-3} Na[ClO₄]-EC/PC (1:1, v/v) organic electrolyte exhibited a capacity retention of 57% after 100 cycles at room temperature.⁴⁰⁸ Another study on the combination of Na_{0.44}MnO₂ positive and HC negative electrode materials in a 1 mol dm⁻³ Na[FSA]-[C₃C₁pyrr][FSA] IL electrolyte achieved a capacity retention of 97% after 100 cycles, which is better than that in a 1 mol dm^{-3} Na[ClO₄]-EC/DEC (1:1, v/v) organic solvent electrolyte, which achieved a capacity retention of 61% after 100 cycles at 25 °C.86 The authors demonstrated that the SEI components generated from the organic solvent electrolyte hinder Na⁺ transport and desolvation of Na⁺.^{86,412} Superior safety was confirmed using a 0.5 mol dm⁻³ Na[FSA]-[C4C1pyrr][TFSA] IL electrolyte compared to that with a 1 mol dm⁻³ Na[ClO₄]-EC/PC (1:1, w/w) using accelerating rate calorimetry analysis for a full cell with Na0 6C00 1Mn0 9O2 positive and Na_{2.55}V₆O₁₆ negative electrodes. This analysis indicated that this full cell was stable up to 250 °C. In addition, the performance of the full cell with an IL electrolyte was superior to that with an organic electrolyte at room temperature.410

A full cell test with organic electrode materials showed a very interesting electrochemical result. Calix[4]quinone was used as the positive electrode material and disodium terephthalate was used as a negative electrode material. This full cell was prepared in a mass ratio of 1:2.5 (calix[4]quinone:disodium terephthalate) with 0.3 mol dm⁻³ Na[TFSA]-[C₃C₁pyrr][TFSA] in a 2032-type coin cell, and stable cycleability was observed compared to that achieved with 0.3 mol dm⁻³ Na[TFSA]-DME.³⁸⁰ This is because quinone compounds easily dissolve in organic electrolytes but not in ILs.

A large-sized prismatic full cell was fabricated using a NaCrO₂ positive electrode and a HC negative electrode with a 20 mol% Na[FSA]-[C₃C₁pyrr][FSA] electrolyte.⁴¹¹ This prismatic full cell was designed to have a capacity of 27 000 mA h with a gravimetric energy density of 75 W h kg⁻¹ and a volumetric energy density of 125 W h dm^{-3} . The discharge capacity was 27 300 mA h at 2700 mA at 25 and 60 $^\circ$ C, which is equal to the designed capacity, with high coulombic and energy efficiencies of 99.8% and 93.2%, respectively, at 25 °C and 99.8% and 96.9%, respectively, at 60 °C. Furthermore, excellent cycleability was achieved at 60 °C, with a capacity retention of 87% after 500 cycles at 10 000 mA. This large-sized full cell is comparable with prismatic lithium-ion batteries developed in the early 2000s in terms of energy density, power density (rate capability), energy efficiency, and cycleability.413,414

Conclusions and perspectives

This article comprehensively reviewed the use of ILs as electrolytes in Na secondary batteries in view of their large-scale

applications in devices such as electric vehicles and EESs. Although Na secondary batteries have a long history, their rapid progress over the last decade has revealed the potential of a variety of electrode and electrolyte materials including ILs. Research in this field has certainly advanced by exploiting the knowledge base of Li analogues, but several studies have confirmed different chemistries that are unique to Na secondary batteries.

There are a number of advantages for ILs over organic electrolytes. In particular, their non-volatility and wide liquidphase temperature range are very attractive for large-scale and task-specific applications. Further research is required to identify ideal ILs and develop their formulation in terms of meeting growing demands for suitable potential windows, fast ionic conduction, and workable temperature extremes. Ion and solvation structures, Na salt form and concentration, cell configuration, and operating protocols have to be further optimised to derive optimal performances for both the positive and negative electrodes simultaneously. The merits of FSA-based ILs are remarkable: allowing the formation of robust SEIs, relatively high ionic conductivity, and great thermal stability.

New anionic species with similar or superior functionalities are an interesting target for future studies. Although not fully considered in this review, IL gel polymer electrolytes prepared by incorporating ILs into polymer hosts may be used for constructing safe all-solid-state Na secondary batteries owing to their excellent mechanical strengths and the low probability of leakage.415-418 Moreover, deep eutectic solvents (DESs) and some complex-salt based electrolytes that behave like ILs could be a cheaply obtainable alternative.149,419-421 The mixing of water or organic solvents with ILs to a certain extent is also a method to ameliorate the high viscosity of ILs without sacrificing their safety.^{168,238} Such ideas should be intensively explored for practical applications. Recent advances using ILs and their analogues as electrode pretreatment agents,⁴²² reaction media for electrode material synthesis, and for the recycling of spent batteries^{423,424} show the multiple roles of ILs in energy-related fields and that many exciting opportunities are waiting to be discovered.

Electrode materials without expensive minor metals are required for Na secondary batteries because large-scale applications are envisioned. In this sense, Fe-based positive electrode materials are preferable, but Mn- and V-based compounds are also good candidates. Electrode design for IL electrolytes can be different to that for organic electrolytes, as the performance of the electrode material in ILs is better than that in organic electrolytes in some cases and worse in other cases. Various factors can affect such behaviour, but one certainty is the difficulty in evaluating true performance in a half-cell configuration because of the large polarisation at the Na counter electrode. This is an intrinsic problem for research into Na secondary batteries and a sophisticated solution is necessary for reliable and efficient screening of electrode materials.

In addition to the influence of the counter electrode, polarisation caused by the lower ionic conductivity of ILs and thus underestimated capacity and inferior rate capability are also often observed with incautious use of ILs. These issues can be overcome *via* modifying electrode–electrolyte interfacial properties by operation at elevated temperatures, which is a practically viable feature of thermally stable ILs that allows electrode materials to deliver greatly improved performance as a consequence of better SEI formation at both negative and positive electrodes, favourable ion transport, and stable Na metal deposition–dissolution without dendrite growth. Particularly, relatively stable cycleability over thousands of cycles has readily been attained for various materials in half-cell using ILs at elevated temperatures, confirming that these beneficial effects are universal regardless of electrode chemistry.

The high cost of ILs is another issue for the construction of practical cells. Undoubtedly, ILs show promise for batteries that operate under harsh environmental conditions or that must withstand widely dissimilar temperatures upon charge and discharge. Other specific applications for which the unique advantages of ILs could be used must be explored in more depth in order to make them truly competitive, even though the price of ILs may become more affordable in mass production than that today.

Furthermore, improved academic understanding is still required for the development of Na secondary batteries, including those made with ILs. The bulk properties of electrolytes, interfacial properties between the electrode materials and the IL, and the effects of binders and conductive additives are all valid targets of fundamental research. Thus, further systematic studies should be undertaken along with those concerning commercial opportunities.

Conflicts of interest

There are no conflicts to declare.

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