Extending the Low-Temperature Operation of Sodium Metal Batteries Combining Linear and Cyclic Ether-Based Electrolyte Solutions

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Extending the low-temperature operation of sodium metal batteries combining linear and cyclic ether-based electrolyte solutions

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Nonaqueous sodium-based batteries are ideal candidates for the next generation of electrochemical energy storage devices. However, despite the promising performance at ambient temperature, their low-temperature (e.g., < 0 °C) operation is detrimentally affected by the increase in the electrolyte resistance and solid electrolyte interphase (SEI) instability. Here, to circumvent these issues, we propose specific electrolyte formulations comprising linear and cyclic ether-based solvents and sodium trifluoromethanesulfonate salt that are thermally stable down to –150 °C and enable the formation of a stable SEI at low temperatures. When tested in the Na||Na coin cell configuration, the low-temperature electrolytes enable long-term cycling down to –80 °C. Via ex situ physicochemical (e.g., X-ray photoelectron spectroscopy, cryogenic transmission electron microscopy and atomic force microscopy) electrode measurements and density functional theory calculations, we investigate the mechanisms responsible for efficient low-temperature electrochemical performance. We also report the assembly and testing between –20 °C and –60 °C of full Na||Na3V2(PO4)3 coin cells. The cell tested at –40 °C shows an initial discharge capacity of 68 mAh g⁻¹ with a capacity retention of approximately 94% after 100 cycles at 22 mA g⁻¹.

Lithium-ion batteries (LIBs) have been extensively employed in portable electronics and electric vehicles because of their high energy density and long cycle life1–3. Nevertheless, they inevitably suffer from severe energy/power losses in cold environments, especially when the temperature drops below –20 °C4,5. Such poor low-temperature performance limits their application in aeronautics/space missions, polar expeditions and many military and civil facilities in cold regions, in which a battery operating temperature below –40 °C is required4,6.

Searching for a system with appealing electrochemical energy storage features beyond Li-based technologies would be promising for addressing the challenges associated with low-temperature operation. As an alkali metal, sodium (Na) stands out because it shares many chemical and physical properties with Li while being substantially more naturally abundant7–9. Na has a lower first ionization energy than Li (495.8 vs. 520.2 kJ mol⁻¹)10, which may contribute to improved chemical/electrochemical reactivity and facilitate electrochemical

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reactions in cold environments. Na metal plays a crucial role as an anode material for Na batteries because of its low electrode potential (~2.714 V vs. standard hydrogen electrode) and high theoretical specific capacity (1166 mAh g⁻¹)₁⁻⁻¹⁴. Nevertheless, investigation of Na batteries at low temperatures has been limited, and in particular, an understanding of the behavior of Na metal as an electrode is largely lacking.¹⁵⁻¹⁸

The enabling of low-temperature battery operation highly depends on the nature of the electrolyte.¹⁹ The electrolyte resistance rapidly increases as the temperature drops because of the relatively high freezing/melting points of nonaqueous carbonate solvents and the reduced solubility of conducting salts.¹⁹ Moreover, the solid electrolyte interphase (SEI) formed at ambient temperature might not be able to maintain the same protective capabilities under cold conditions to enable efficient cycling. Meanwhile, the low-temperature structural and compositional evolution of the SEI formed on the Na metal electrode still remains elusive.

One viable solution to circumvent these problems is to formulate electrolytes targeted at low-temperature operation using solvents with low melting points and salts capable of forming a stable SEI. Here, applying such an electrolyte strategy, we demonstrate that an electrolyte solution comprising an acyclic ether solvent and a compatible Na salt can extend the Na metal operating temperature to -40 °C. The trifluoromethanesulfonate (OTf) salt is found to play a critical role in enabling the formation of a stable SEI at low temperatures. Furthermore, adding a cyclic ether solvent to prepare a binary-ether electrolyte solution can expand the thermostability temperature threshold down to ~150 °C. We demonstrate stable Na metal plating/stripping in symmetric cells at low temperatures down to -80 °C, exhibiting a low overpotential of -150 mV for over 750 h. This performance expands the low-temperature operational capability of alkali metal electrodes in nonaqueous electrolyte solutions (see Supplementary Fig. 1 and Supplementary Table 1 for a comparison against the state of the art). Coupled experimental characterizations (e.g., X-ray photoelectron spectroscopy, cryogenic transmission electron microscopy, and atomic force microscopy) and density functional theory calculations allow for an understanding of the mechanistic features that enable efficient low-temperature electrochemical performance. Full Na||Na₅V₂(PO₄)₃ coin cells are also assembled and are tested between -20 °C and -60 °C. The cells tested at -40 °C and -60 °C show initial discharge capacities of ~68 and 39 mAh g⁻¹, respectively, with capacity retentions of ~94% and 91% after 100 cycles at 22 °C.

**Results**

Screening of single-solvent electrolytes at low temperatures

We screened different electrolyte solutions at -35 °C using a variety of salt-solvent combinations. Ethers with low melting points, including diethylene glycol dimethyl ether (DEGDME), 1,2-dimethoxyethane (DME), and 1,3-dioxolane (DOL), were chosen as candidate solvents (Supplementary Table 2). A variety of conducting salts, including sodium hexafluorophosphate (NaPF₆), sodium trifluoromethanesulfonate (NaOTf), sodium perchlorate (NaClO₄), sodium bis(trifluoromethanesulfonyl)imide (NaTFSI) and sodium bis(fluorosulfonyl)imide (NaFSI), were investigated as candidate salts. The salt concentration in the solvent was kept at 1 M for the solubility screening of these single-solvent electrolytes at low temperatures. The results (Supplementary Table 3) suggest that the solubility of NaPF₆ cannot reach a 1 M concentration in all three solvents at -35 °C, while the other four salts present good solubility in DEGDME. Therefore, while 1 M NaPF₆-DEGDME has been shown to be a suitable electrolyte for the Na metal anode under ambient conditions,¹ it may not be suitable for low-temperature operation. Moreover, DOL cannot easily dissolve Na salts except for NaTFSI. The electrolyte consisting of 1 M NaPF₆-DEGDME was further investigated across a range of temperatures in Na||Na and stainless steel||stainless steel symmetric cells, respectively (Supplementary Fig. 2). A comparison of the electrochemical impedance spectroscopy (EIS) spectra of the 1 M NaPF₆-DEGDME tested in Na||Na symmetric cells shows that the electrode-electrolyte interfacial resistance at -20 °C is significantly higher than that at 20 °C (Supplementary Table 4). Meanwhile, the total electrolyte resistance measured in the stainless steel||stainless steel cells (cell configuration is shown in Supplementary Fig. 3) increases with decreasing temperature. This increase in electrolyte resistance is especially significant at a temperature below -20 °C (Supplementary Table 5), which could be mainly due to the precipitation of NaPF₆ in DEGDME at low temperatures. In summary, eight of the fifteen single-solvent electrolytes are thermally stable at ~35 °C at 1 M concentration, and these were chosen for further electrochemical evaluation with metallic Na.

**Electrochemical behavior of single-solvent electrolytes**

Galvanostatic cycling measurements were performed in symmetric Na||Na cells at a current density of 0.5 mA cm⁻² with a cycling capacity of 0.5 mAh cm⁻² at ~20 °C (Fig. 1a–c) and +20 °C (Supplementary Figs. 4 and 5). Among the eight electrolyte candidates, the cell with 1 M NaOTf-DEGDME presents the most stable cycling with the smallest average overpotential (~10 mV) for 600 h (300 cycles) at +20 °C (the electrochemical behavior for different electrolytes at -20 °C is summarized in Supplementary Table 6). The electrolytes with NaTFSI salt all show much higher overpotentials than the other electrolytes. Specifically, the cell with NaTFSI-DOL shows an increasing overpotential that rapidly reaches 1 V, and the cells with 1 M NaTFSI-DEGDME and 1 M NaTFSI-DME failed quickly, reaching the protection voltage (5 V) at 32 and 44 h, respectively.

When tested at -20 °C, the cell with 1 M NaOTf-DEGDME still displays the highest stability, with smooth and steady voltage plateaus as well as the lowest overpotential of ~16 mV (Fig. 1a, b). In comparison, unstable and fluctuating voltage spikes are observed for the other electrolyte formulations (Fig. 1b, c). Because of the better performance of NaOTf-DEGDME at both +20 °C and -20 °C, it was selected for further testing in symmetric cells at ~40 °C (Fig. 2a). At such a low temperature, the initial overpotential of 40 mV only rises to 50 mV even after 500 h of cycling at 0.5 mA cm⁻² (capacity: 1 mAh cm⁻²), corresponding to a less than 0.2% increase per cycle. Even at a current density of 1 mA cm⁻², an average overpotential of 100 mV can be maintained for over 300 h. Regarding the coulombic efficiency (CE) evaluation (Fig. 2b), the low average value of the CE at ~40 °C compared to that at +20 °C suggests that a significant Na inventory loss occurs at ~40 °C.

**Physicochemical characterization of single-solvent electrolytes**

The morphology of the Na metal surface after Na plating/stripping in different electrolytes was characterized using scanning electron microscopy (SEM). NaTFSI, NaClO₄, and NaFSI salts all result in irregular surfaces (pores and/or cracks) of the deposited Na (Supplementary Figs. 6—9). In contrast, NaOTf-DEGDME enables relatively smooth surfaces at both +20 °C and -20 °C, where an inhomogeneous texture can be observed on the surface and within the cross-section. This microstructure becomes more distinct at ~40 °C (Fig. 2c).

The chemical composition of the SEI formed on the Na metal electrode in NaOTf-DEGDME was revealed by X-ray photoelectron spectroscopy (XPS), as shown in Supplementary Fig. 10. The binding energies of all elements were calibrated with respect to the C 1s signal at 284.8 eV. In the C 1s XPS profile, the peak with a binding energy of 293.4 eV is assigned to CF₃, and the peaks at 288.7 eV and 286.6 eV are ascribed to O=C=O and O=C–O, respectively.¹⁻² In the O 1s spectrum, the peaks at 533.0 eV and 531.2 eV correspond to CH₂=O–/CH₂–O– groups³⁶⁻⁷ and Na–O=Na (e.g., RCHO(Ona)³, respectively, and the peak at 536.3 eV is assigned to Na KLL. For S 2p, the doublets at 169.6 eV and 167.2 eV (based on 2p₃/₂) are assigned to SO₄²⁻ and SO₃²⁻, respectively.
For F Is, the peaks at 689.3 eV and 684.1 eV are assigned to C–F and NaF, respectively. These XPS spectra at different temperatures reveal similar compositions of the SEI surface, suggesting the existence of \(-\text{CF}_3\)-containing compounds, \(\text{Na}_2\text{SO}_4\), \(\text{Na}_2\text{SO}_3\), \(\text{Na}_2\text{CO}_3\), NaF, C–O–Na and organic debris. \(\text{Na}_2\text{SO}_3\) can only be detected on the surface at +20 °C. In addition, \(\text{Na}_2\text{S} (\text{S}^2-: 161.1 \text{ eV})\) cannot be detected in the inner SEI at −20 °C and −40 °C (Supplementary Figs. 11–13). The absence of species with lower oxidation states at low temperatures suggests that the parasitic reactions between the electrolyte and Na metal electrode are suppressed. Importantly, the SEI composition formed in NaOTF-DEGDME shows high consistency across these temperatures. For the other systems tested, the discussion of the Na surface morphology (Supplementary Figs. 6–9) and chemical composition of the SEI (Supplementary Figs. 14–22, Supplementary Tables 7 and 8) is detailed in Supplementary Notes 1 and 2, respectively.

The distribution of chemical species within the SEI formed in 1 M NaOTF-DEGDME at different temperatures was further analyzed by XPS depth profiling (Fig. 2d). At all temperatures, the O and C contents decrease with increasing depth, revealing that the organic debris is mainly present in the upper layer of the SEI. For inorganic components, NaF and \(\text{Na}_2\text{SO}_4\) are the two major species. The NaF/\(\text{Na}_2\text{SO}_4\) ratio increases with increasing depth, suggesting that NaF dominates the inner part of the SEI. At ~40 °C, the Na content exhibits a rapid increase with increasing etching time due to the formation of a thinner SEI layer at a lower temperature.

**Theoretical investigation on single-solvent electrolytes**

To help rationalize the performance of different salts, we performed first-principles computations within density functional theory (DFT) to evaluate their reduction potentials. Figure 3 shows the computed reduction potentials for the different salts and compares them to the SEI quality. Supplementary Table 9 collects the calculated reduction potentials of salts and solvents with or without the Na cation. Salts such as NaTFSI, which have a low reduction potential close to the electrolyte reduction potential, do not allow fast formation of an inorganic-compound-rich SEI capable of preventing electrolyte decomposition. This is well consistent with the Na stripping/plating behavior in all NaTFSI-based electrolytes, which show high over-potentials (Supplementary Table 6). In contrast, salts with high reduction potentials such as NaFSI and NaClO4 (2.37 V and 4.96 V vs. Na/Na⁺, respectively) promptly react with Na metal to form an inorganic SEI layer, but the strong driving force for the reaction leads to the formation of SEIs with a nonuniform morphology.

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*Fig. 1 | Electrochemical cycling of Na metal electrodes in various electrolyte formulations at −20 °C. a Galvanostatic cycling of Na||Na symmetric cells operating in eight different electrolyte formulations with a 1 M salt concentration at 0.5 mA cm⁻² and 0.5 mAh cm⁻². b, c Corresponding enlarged voltage profiles at the 50th cycle and 100th cycle for four of the formulations and the remaining four formulations.*
Figs. 6 and 7). NaOTf exhibits an intermediate reduction potential (1.02 V vs. Na/Na+) leading to the formation of an inorganic protective layer before the electrolyte is decomposed, but with a moderate driving force, leading to a mild reaction and formation of a smooth SEI film (Fig. 2c). For the sake of comparison, we also provide the results for NaPF6, which is known to form a stable SEI at room temperature23. NaPF6 also exhibits a moderate reduction potential (0.75 V vs. Na/Na+), confirming our speculation that a salt with an intermediate reduction potential helps the formation of a uniform SEI on the Na metal electrode surface. Note that NaPF6 may not be a suitable low-temperature salt because of solubility issues and the fact that the performance of NaOTf comes from the combination of a low-temperature solubility and a moderate reduction potential.

Design of binary-solvent electrolytes for temperatures below −40 °C

Motivated by the electrochemical behavior of the NaOTf-DEGDME electrolyte at low temperatures, this electrolyte solution was selected for further formulation of electrolytes that can operate below −40 °C. DOL, which is a cyclic ether with a melting point of −95 °C, was introduced into DEGDME to produce binary-solvent electrolytes (Supplementary Fig. 23) to further improve the low-temperature properties of the electrolyte. However, the Na+ solvation ability of DOL is lower than that of DEGDME32. Therefore, the solvent volume ratio (DEGDME:DOL = 8:2, 5:5 or 2:8 volume fraction) and salt concentration (0.5 M or 1 M) were evaluated in the screening of binary-solvent electrolytes (Supplementary Table 10).

The introduction of DOL extends the low-temperature operating range of the NaOTf-DEGDME system, in which a higher DOL volume fraction leads to better Na cycling stability in symmetric cells (Fig. 4a and Supplementary Fig. 24). Additionally, a lower salt concentration (0.5 M) can accommodate an increased DOL proportion, further enhancing the cell cycling performance (Fig. 4a). At −80 °C, the symmetric cell overpotential (~35 mV) in 0.5 M NaOTf-DEGDME/DOL (2:8) is less than half that (~75 mV) in 0.5 M NaOTf-DEGDME/DOL (5:5) and six times less than that (>200 mV) in 1 M NaOTf-DEGDME/DOL (5:5). Thus, both the NaOTf salt concentration and DEGDME volume fraction should be carefully chosen to formulate an electrolyte solution that enables good low-temperature electrochemical energy storage performance. Replacing DEGDME with DME leads to an increased over-potential and/or an asymmetric voltage profile (Supplementary Fig. 25), which also occurs with the substitution of NaOTf by NaClO4 (Supplementary Fig. 26).

Physicochemical and electrochemical characterization of binary-solvent electrolyte solutions

To understand the mechanism of the enhanced performance at low temperatures, the electrolyte resistance was investigated at different temperatures (Fig. 4b). The resistance of the electrolyte solution was measured by EIS in stainless steel||stainless steel cells (see Methods for details). The symmetric cell configuration is shown in Supplementary Fig. 3 and the resistance evaluation based on EIS is show in Supplementary Fig. 27. The resistance of the 0.5 M NaOTf-DEGDME/DOL (2:8) electrolyte solution at +20 °C is 11.6 Ohm, and it increases by ~3.6 times when the temperature decreases to −80 °C (Supplementary Table 11). This difference is almost half that for 0.5 M NaOTf-DEGDME/DOL (5:5) and six times less than that (>200 mV) in 1 M NaOTf-DEGDME/DOL (5:5). Thus, both the NaOTf salt concentration and DEGDME volume fraction should be carefully chosen to formulate an electrolyte solution that enables good low-temperature electrochemical energy storage performance. Replacing DEGDME with DME leads to an increased over-potential and/or an asymmetric voltage profile (Supplementary Fig. 25), which also occurs with the substitution of NaOTf by NaClO4 (Supplementary Fig. 26).
Figure 4c shows that 0.5 M NaOTf-DEGDME/DOL (2:8) gives the slowest decrease in the ionic conductivity with temperature. At −80 °C, the ionic conductivities of 0.5 M NaOTf-DEGDME/DOL (2:8), 0.5 M NaOTf-DEGDME/DOL (5:5) and 1 M NaOTf-DEGDME are 1.05 mS cm⁻¹, 0.44 mS cm⁻¹, and 0.16 mS cm⁻¹, respectively. The ionic conductivities of the other four single-solvent systems are shown in Supplementary Fig. 28. The small change in the electrolyte resistance and ionic conductivity of NaOTf-DEGDME/DOL (2:8) could be attributed to the relatively low dynamic viscosity of the solvents (Supplementary Fig. 29). Note that both binary-solvent and single-solvent electrolytes show good homogeneity without salt precipitation across temperatures (Supplementary Fig. 30). Differential scanning calorimetry (DSC) further confirms the thermal stability of 0.5 M NaOTf-DEGDME/DOL (2:8), revealing no phase transition even down to −150 °C (Fig. 4d). Similarly, no phase transition is detected until −140 °C for 0.5 M NaOTf-DEGDME/DOL (5:5). In comparison, 1 M NaOTf-DEGDME displays a second-order phase transition at −126 °C during cooling and a first-order phase transition at −64 °C (melting point of DEGDME) during subsequent heating.

Based on the resistance and thermal behavior of the electrolyte solutions, we carried out prolonged Na||Na symmetric cell cycling at −80 °C (Fig. 4e and Supplementary Fig. 31). The cells using 0.5 M NaOTf-DEGDME/DOL (2:8) display a small overpotential of ~50 mV, which does not increase for over 2000 h at 0.2 mA cm⁻² with a
0.1 mAh cm$^{-2}$ cycling capacity (Supplementary Fig. 31). Even at a higher current of 0.5 mA cm$^{-2}$ with a higher capacity of 0.25 mAh cm$^{-2}$, stable operation for over 750 h can still be achieved (Fig. 4e). The enlarged voltage profile (Fig. 4f) further suggests that the overpotential at 0.5 mA cm$^{-2}$ is initially 147 mV (after the stabilization process) and slightly increases to 150 mV after 500 h, revealing a smooth voltage profile without any spikes. In comparison, 0.5 M NaOTf-DEGDME/DOL (2:8) shows large spikes (>4000 mV) and an asymmetric voltage profile during cycling.

Ex situ postmortem physicochemical characterization of sodium metal electrodes

An improved CE with an average value of ~97.6% at ~40 °C is observed in 0.5 M NaOTf-DEGDME/DOL (2:8) (Fig. 5a) compared to the single-solvent case (~92.2%, Fig. 2b) using Na|stainless steel cells at ~40 °C at 0.5 mA cm$^{-2}$ and 0.5 mAh cm$^{-2}$. Ex situ atomic force microscopy (AFM) measurements were carried out to investigate the surface roughness and mechanical properties of the SEI formed on copper foil at 0.5 mA cm$^{-2}$ (0.5 mAh cm$^{-2}$) for the 1st cycle at ~40 °C (Fig. 3b) and Supplementary Fig. 32. In 0.5 M NaOTf-DEGDME/DOL (2:8), the height of the SEI formed at the electrode surface is uniform with a deviation of 50 nm (Fig. 5a, Supplementary Fig. 32), and the average Young’s modulus is ~1.2 GPa, as shown in Supplementary Fig. 32. The better uniformity and mechanical property of the SEI contributes to more efficient plating/stripping behavior on copper foil (Supplementary Fig. 33) in the binary-solvent electrolyte compared to the single-solvent case. The SEI morphology after Na stripping was investigated via SEM (Supplementary Fig. 34), in which a smoother surface when using 0.5 M NaOTf-DEGDME/DOL (2:8) can be noted compared to the irregular and porous morphology obtained using the 1M NaOTf-DEGDME electrolyte solution.

The binary-solvent system can also alter the composition and microstructure of the Na electrode surface in comparison to the single-solvent system. Specifically, a gradual increase in the volume fraction of DOL results in the lack of formation of irregular morphologies such as those observed in NaOTf-DEGDME at low temperatures (Fig. 5d and Supplementary Figs. 35 and 36). Regarding the composition, ex situ postmortem XPS measurements of the Na metal electrodes cycled using the NaOTf-DEGDME/DOL electrolyte at ~80 °C (Supplementary Figs. 37–40) suggest inorganic components (NaF, Na$_2$SO$_4$, and Na$_2$SO$_3$) in the SEI similar to those formed in the single-solvent counterpart. Nevertheless, XPS depth profiling analysis reveals a difference in the distribution of elements and inorganic components in binary-solvent electrolytes (Fig. 5e, f and Supplementary Fig. 41) compared to the single-solvent counterpart (Fig. 2d). Specifically, the NaF/Na$_2$SO$_4$ ratio is higher on the surface (~1:1) when using the binary-solvent electrolytes (Fig. 5f and Supplementary Fig. 41b) than those (1:8.9 at ~20 °C and 1:3.3 at ~40 °C) in the single-solvent system (Fig. 2d). Taking into account the AFM investigations, assuming that the higher percentage of the NaF inorganic component in the binary-solvent electrolyte contributes to a higher Young’s modulus (Fig. 5e and Supplementary Fig. 32b) and thus to a better mechanical strength compared to the single-solvent system is plausible.

Cryo-TEM further indicates the crystalline nature of the SEI (likely containing Na$_2$SO$_4$ and Na$_2$CO$_3$) via identification of lattice fringes and analysis of electron diffraction patterns. Specifically, the SEI growth in the binary-solvent system (Fig. 5g, h) shows a mosaic structure, consisting of an amorphous matrix with Na-rich nanocrystallites dispersed within, similar to that in the single-solvent system (Supplementary Fig. 42). These data are consistent with the XPS experimental findings on the SEI for both cases. However, the crystallite size and overall quantity of crystallites within the SEI are larger in the binary-solvent electrolyte, as shown by the TEM images as well as fast Fourier

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**Fig. 5 | Electrochemical and physicochemical characterizations of Na metal electrodes cycled in 0.5 M NaOTf-DEGDME/DOL (2:8) at low temperatures.**

a CE as a function of cycle number in Na|stainless steel cells at ~40 °C at 0.5 mA cm$^{-2}$ and 0.5 mAh cm$^{-2}$. b AFM topography of the SEI formed on copper foil after Na plating/stripping at 0.5 mA cm$^{-2}$ (0.5 mAh cm$^{-2}$) with a cutoff voltage of 1.5 V after the 1st cycle at ~40 °C (Cu foil is fully desodiated). c Young’s modulus (determined by AFM) of the SEI in Fig. 5b. d SEM of the Na metal electrode surface after cycling at ~80 °C (Inset: corresponding cross-sectional SEM image). e Contents of C 1s, O 1s, F 1s, S 2p and Na 1s elements in the SEI at ~80 °C. f Contents of the main SEI inorganic components, including NaF, Na$_2$SO$_4$ and Na$_2$SO$_3$, at ~80 °C. g Cryo-TEM image of the SEI after cycling at ~40 °C. Visible lattice fringes from crystalline regions are overlaid with different colors. The red lattice fringes have a d-spacing of 3.0 Å, the blue have a d-spacing of 3.5 Å, and the green have a d-spacing of 3.8 Å. These three fringes likely correspond to the labeled planes shown. h SAED pattern of the SEI formed at ~40 °C. The positions of the rings expected to be most intense from Na$_2$O, Na$_2$SO$_4$, Na$_2$CO$_3$, and NaF are labeled; the large variety of crystalline phases likely present makes fully indexing the structures difficult. These crystalline phases also give rise to extra rings that are not marked, which likely account for the reflections that fall between marked lines. Note that the presence of Na$_2$O may be attributed to the transfer process for these samples.
NaF-rich SEI with a uniform morphology and an improved Young's modulus is the main reason for the long-term stability of the cell performance. We speculate that the cell voltage hysteresis is also temperature sensitive. At ~60 °C, ~42% of the specific capacity obtained at ~20 °C can be retained. The capacity loss from the temperature stepping is reversible, and the capacity is fully recovered as the temperature rises back to ~20 °C (Fig. 6a). In addition, the CE increases with decreasing temperature. It increases from an average value of 91.3% at 0 °C to 99.9% at ~60 °C, and then decreases to 99.5% when the temperature increases back to ~20 °C (Supplementary Fig. 43a). The charge–discharge voltage profiles from 0 °C to ~80 °C (Fig. 6b) suggest that the cell voltage hysteresis is also temperature sensitive. At ~60 °C, the cell voltage hysteresis is approximately four times that at ~40 °C and five times that at ~20 °C. For long-term cycling (Fig. 6c), a cell at ~20 °C exhibits an initial discharge capacity of 92 mAh g⁻¹ and a capacity retention of 94% after 100 cycles, with an average CE of 98.1%. At ~40 °C, the initial capacity decreases to 68 mAh g⁻¹ with a capacity retention of 94% after 100 cycles, with an average CE of 98.1%. At ~60 °C, the capacity is further decreased to 63 mAh g⁻¹ with a capacity retention of 91% after 100 cycles, and a further average CE of 99.6% can be achieved. Even at ~60 °C, a similar CE (99.5%) can still be achieved, with a capacity retention of 91% (decay rate < 0.089% per cycle) after 100 cycles. The enlarged CE data are shown in Supplementary Fig. 43b. Figure 6d shows the rate performance of the cells at currents up to 110 mA g⁻¹ (1 C) at ~40 °C and ~60 °C, which further confirms the ability of Na metal to be cycled in the full cell configuration using the designed electrolyte under extremely cold conditions. The corresponding CE data at different rates are detailed in Supplementary Fig. 43c, which shows that the CE at different temperatures increases with increasing current rate.

Good cycling performance is also achieved at higher mass loadings of Na₃V₂(PO₄)₃ up to 3.0 mg cm⁻² at ~40 °C (Supplementary Figs. 44 and 45). Moreover, an anode of Na@MXene (Na metal incorporated into an MXene electrode) is employed for pairing with Na₃V₂(PO₄)₃ at ~40 °C, showing a stable performance (Supplementary

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**Fig. 6** | Electrochemical energy storage performance of Na||Na₃V₂(PO₄)₃ coin cells using the 0.5 M NaOTf-DEGDME/DOL (2:8) electrolyte solution at low temperatures. a Temperature-dependent galvanostatic cycling of cells at 22 mA g⁻¹ (equivalent to 0.2C, 1C = 110 mA g⁻¹), based on the active material Na₃V₂(PO₄)₃ down to ~60 °C with voltage cutoffs of 2.3 and 3.8 V. Note that the duration of cycling at each temperature was kept the same for comparison. b Galvanostatic charge–discharge voltage profiles at 22 mA g⁻¹ from 0 °C to ~80 °C. c Long-term galvanostatic cycling of cells at 22 mA g⁻¹ at ~20 °C, ~40 °C and ~60 °C. d Rate cycling performance (up to 110 mA g⁻¹, equivalent to 1C) of cells at ~40 °C and ~60 °C.

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**Testing of full Na metal coin cells at temperatures ≤0 °C**

Sodium metal batteries comprising Na₃V₂(PO₄)₃ as the cathode (active material loading: ~2 mg cm⁻²) and Na metal as the anode in the 0.5 M NaOTf-DEGDME/DOL (2:8) electrolyte were tested at 22 mA g⁻¹ (equivalent to 0.2C, 1C = 110 mA g⁻¹) and low temperatures down to ~80 °C (Fig. 6 and Supplementary Fig. 43). Galvanostatic cycling (Fig. 6a) shows that the discharge capacity decreases as the cell is sequentially cooled. At ~60 °C, ~42% of the specific capacity obtained at ~20 °C can be retained. The capacity loss from the temperature stepping is reversible, and the capacity is fully recovered as the temperature rises back to ~20 °C (Fig. 6a). In addition, the CE increases with decreasing temperature. It increases from an average value of 91.3% at 0 °C to 99.9% at ~60 °C, and then decreases to 99.5% when the temperature increases back to ~20 °C (Supplementary Fig. 43a). The charge–discharge voltage profiles from 0 °C to ~80 °C (Fig. 6b) suggest that the cell voltage hysteresis is also temperature sensitive. At ~60 °C, the cell voltage hysteresis is approximately four times that at ~40 °C and five times that at ~20 °C. For long-term cycling (Fig. 6c), a cell at ~20 °C exhibits an initial discharge capacity of 92 mAh g⁻¹ and a capacity retention of 94% after 100 cycles, with an average CE of 98.1%. At ~40 °C, the initial capacity decreases to 68 mAh g⁻¹ with a capacity retention of 94% after 100 cycles, while a higher average CE (99.6%) can be achieved. Even at ~60 °C, a similar CE (99.5%) can still be achieved, with a capacity retention of 91% (decay rate < 0.089% per cycle) after 100 cycles. The enlarged CE data are shown in Supplementary Fig. 43b. Figure 6d shows the rate performance of the cells at currents up to 110 mA g⁻¹ (1 C) at ~40 °C and ~60 °C, which further confirms the ability of Na metal to be cycled in the full cell configuration using the designed electrolyte under extremely cold conditions. The corresponding CE data at different rates are detailed in Supplementary Fig. 43c, which shows that the CE at different temperatures increases with increasing current rate.

Good cycling performance is also achieved at higher mass loadings of Na₃V₂(PO₄)₃ up to 3.0 mg cm⁻² at ~40 °C (Supplementary Figs. 44 and 45). Moreover, an anode of Na@MXene (Na metal incorporated into an MXene electrode) is employed for pairing with Na₃V₂(PO₄)₃ at ~40 °C, showing a stable performance (Supplementary
Figs. 46 and 47), which demonstrates a proof-of-concept full cell with less Na metal (2 mAh g⁻¹) employed on the anode side.

In summary, we have formulated low-temperature electrolytes comprising acyclic/cyclic ethers (DEGDME/DOL) and NaOTf salt, which extend the operating temperature limit for Na metal batteries. In particular, a single-solvent electrolyte of 1 M NaOTf-DEGDME enables stable Na metal cycling down to −40 °C, showing a low overpotential of 100 mV at 1 mA cm⁻² with a capacity of 1 mAh cm⁻² for over 300 h. Theoretical calculations show that NaOTf has a moderately high reduction potential (1.02 V vs. Na/Na⁺) compared to the other commonly used Na salts, leading to the formation of an inorganic protective layer before the electrolyte solvent is decomposed, but with a moderate driving force. This could contribute to a moderate reaction with Na metal and thus the formation of a smooth and uniform SEI. To further decrease the temperature threshold, DOL was introduced to form binary-solvent electrolytes. An optimized electrolyte consisting of 0.5 M NaOTf-DEGDME/DOL (2:8 in volume ratio) exhibits a resistance increase of only 3.4 times from +20 °C to −80 °C. The good electrochemical performance could be ascribed to the formation of a robust NaF-rich SEI (with high NaF content in the upper layer) with uniform morphology and mechanical strength, revealed by a suite of characterizations, including SEM, AFM, XPS and cryo-TEM. As a proof of concept, sodium metal batteries were assembled comprising Na₃V₂(PO₄)₃ as the cathode and Na metal as the anode and thus the formation of a smooth and uniform SEI. To further demonstrate stable Na cycling with a small overpotential of 50 mV for over 2000 h at −80 °C. The good electrochemical performance could be ascribed to the formation of a robust NaF-rich SEI (with high NaF content in the upper layer) with uniform morphology and mechanical strength, revealed by a suite of characterizations, including SEM, AFM, XPS and cryo-TEM. As a proof of concept, sodium metal batteries were assembled comprising Na₃V₂(PO₄)₃ as the cathode and Na metal as the anode and thus the formation of a smooth and uniform SEI.

**Methods**

**Materials**

Metallic sodium (Na) chips (average thickness of 1 mm) were made from rolling-pressed Na metal cubes (99.9% trace metals basis, Sigma–Aldrich) in an argon (Ar)-filled glove box (O₂ < 0.6 PPM, H₂O < 0.1 PPM, Mbraun), whose oxidized surfaces were cut off using a knife. Sodium hexafluorophosphate (NaPF₆, Sigma–Aldrich), sodium trifluoromethanesulfonate (NaOTf, Beantown Chemical), sodium perchlorate (NaClO₄, Sigma–Aldrich), sodium bis(trifluoromethanesulfonyl)imide (NaTFSI, Solvionic) and sodium bisfluorosulfonimide (NaFSI, Solvionic) were dried in glass vials for at least an hour to attain thermal equilibrium. The electrolytes were prepared in the glove box (O₂ < 0.6 PPM, H₂O < 0.1 PPM, Mbraun) before being used. Diethylene glycol dimethyl ether (DEGDME, anhydrous, H₂O < 50 ppm, Sigma–Aldrich), 1,2-dimethoxyethane (anhydrous, H₂O < 30 ppm water, Sigma–Aldrich) and 1,3-dioxolane (anhydrous, H₂O < 30 ppm, Sigma–Aldrich) were directly used as solvents for preparing electrolytes.

To synthesize Na₃V₂(PO₄)₃, 0.364 g vanadium oxide (V₂O₅, Alfa Aesar) and 0.54 g oxalic acid (H₂C₂O₄, anhydrous, Alfa Aesar) were dissolved in 20 mL distilled water at 70 °C. Sodium dihydrogen phosphate (NaH₂PO₄, 0.72 g, Alfa Aesar), 0.2 g glucose (anhydrous, Alfa Aesar) and 50 mL 1-propanol (Alfa Aesar) were added and mixed, followed by drying at 70 °C. The dried sample was ground and annealed at 400 °C for 4 h and then 750 °C for 8 h under an Ar gas atmosphere with a heating rate of 5 °C min⁻¹. The annealed powder was mixed with carbon black (Super C65, CNERGY) and polyvinylidene fluoride (PVDF, Hsv 900, Arkema) in a weight ratio of 6:3:1 using 1-methyl-2-pyrrolidone (NMP, anhydrous, Sigma–Aldrich) to form a slurry. The slurry was cast onto aluminum foil (99.9% purity, 16 µm thickness, MTI) and dried at 70 °C in a vacuum oven, followed by cutting the specimen into a cathode with a diameter of 12 mm (average thickness of 60–160 µm). The active material loading was up to 3 mg cm⁻². To synthesize Ti₃C₂ MXene, 3 g Ti₃AlC₂ powder (98%, 400 mesh, Jilin 11 Technology Co., Ltd. China) was added into 30 mL hydrofluoric acid (HF) solution (40%, Alfa Aesar) under stirring for 24 h to obtain Ti₃C₂ MXene. The obtained Ti₃C₂ MXene was washed with deionized (DI) water until a pH value of 6–7 was reached for the solution, which was then freeze-dried. Then, 0.25 g as-obtained Ti₃C₂ MXene was weighed and added into a 50 mL cyclohexylmethylammonium bromide (CTAB) solution (0.20 wt %, 99%, Alfa Aesar) under stirring at 40 °C for 24 h. The collected Ti₃C₂ MXene was washed with DI water and dried at 70 °C. After that, 0.25 g as-obtained Ti₃C₂ MXene was weighed and dispersed in 40 mL NaOH solution (1 M) under stirring (500 rpm) at room temperature for 72 h, and then, the precipitate was collected and washed with DI water and freeze-dried. The electrode slurry composed of the as-synthesized Ti₃C₂ MXene and PVDF in a weight ratio of 9:1 (dispersed in NMP) was cast on Cu foil (99.8% purity, 9 µm thickness, MTI). The wet coating was dried at 70 °C in a vacuum oven, followed by cutting into an anode with a diameter of 12 mm (average thickness of 100 µm). The active material loading was up to 2 mg cm⁻².

**Physicochemical characterizations**

Dynamic viscosity (or absolute viscosity) measurements were conducted using a viscometer (microVISCTM, RheoSense), which was specifically calibrated for low-temperature measurement (down to −40 °C), in the glove box (O₂ < 0.6 PPM, H₂O < 0.1 PPM, Mbraun). Before testing, each sample (100 µL) was isothermally stored at the target temperature for over 10 h. Scanning electron microscopy (SEM) images were obtained using a field emission gun environmental SEM system (XL 300 ESEM-FTG). Cryogenic transmission electron microscopy (cryo-TEM) analysis was conducted using an FEI Tecnai F30 TEM operating at 300 kV. All images were recorded at a low dose rate between 1 and 2 × 10² electrons Å⁻²s⁻¹. X-ray photoelectron spectroscopy (XPS) analysis was performed via a PHI VersaProbe II scanning XPS microprobe with a 0.47 eV system resolution using a monochromatic 1486.7 eV x-ray source. Differential scanning calorimetry (DSC) was performed using a Netzsch DSC 204 F1 Phoenix. Both the samples and reference were tested between −150 °C and +20 °C at a controlled rate of 10 °C min⁻¹. Atomic force microscopy (AFM) was conducted on copper foils after Na plating/stripping using a Bruker’s Dimension Icon AFM with an RTESP-150 probe.

**Electrochemical characterizations**

A temperature chamber (MC-812, ESPEC) was employed to ensure stable temperature environments between −80 °C and +20 °C. Tested coin cells were maintained at a specified temperature for at least an hour to attain thermal equilibrium. The electrolytes were prepared in the glove box (O₂ < 0.6 PPM, H₂O < 0.1 PPM, Mbraun). The mass of Na metal was weighed using a balance (PA84C, OHAUS), and the volumes of solvents were measured using pipettes (100–1000 µL, High Performance Single-Channel Pipettor, VWR). Galvanostatic Na|Na symmetric cycling was conducted in 2023-type coin cells, which were assembled with two identical Na chips (average thickness of 1 mm, diameter of 12 mm) and a separator (Celgard 2400, thickness of 24 µm, diameter of 15 mm, average porosity of 39%) filled with 40 µL electrolyte. Asymmetric coin cells were also assembled using stainless steel foil (thickness of 0.5 mm, 316 stainless steel) as the working electrode and freshly cut Na as the counter electrode with a Celgard 2400 separator and 40 µL electrolyte. CE measurements were performed in asymmetric coin cells by plating and stripping at a current density of 0.5 mA cm⁻² for over 1 h (0.5 mAh cm⁻² deposited or stripped per half cycle). Sodium metal cells were assembled with Na₃V₂(PO₄)₃ cathodes and Na metal or...
Na@MXene anodes with the tested electrolytes. To prepare Na@MXene electrodes, 2 mAh cm⁻² Na was deposited onto the Ti₃C₂MXene electrode using asymmetric Na@MXene cells at 0.5 mA cm⁻² in the tested electrolyte with a Celgard 2400 separator. After Na deposition, the Na@MXene cells were disassembled, and the Na@MXene electrodes were collected. Both symmetric cells and sodium metal batteries were tested using a standard battery tester (CT2001A, Wuhan LANHE Electronic Co., Ltd.). Electrochemical impedance spectroscopy (EIS) was conducted using an electrochemical workstation (VMP3, Bio-Logic Science Instruments) at a scanning frequency from 1 MHz to 0.1 Hz with an AC amplitude of 5 mV and 10 points per decade. The average open-circuit voltage time before EIS measurement was ~2 h.

The total resistance of the electrolyte solution (Rₑ) was measured through EIS in stainless steel/stainless steel cells. A polymer membrane (Celgard, average thickness of 25 µm, porosity of 39%, diameter of 15 mm) was sandwiched between two pieces of 316 stainless steel (thickness: 0.5 mm) in a 2032-type coin cell with 40 µL electrolyte added (Supplementary Fig. 3). The ionic conductivity (δ) of the electrolyte solution was calculated based on the equation below:

$$\delta = L/(A R_e)$$

where $R_e$ is the resistance value from EIS fitting curve using the equivalent electrical circuit, $L$ is the distance between the two pieces of stainless steel, and $A$ is the product of the membrane area and porosity.

TEM samples were prepared by carrying out a single Na plating/stripping cycle (0.25 mA cm⁻² for 2 h) using a Cu TEM grid as the working electrode. The coin cell containing the grid was disassembled in an Ar-filled glove box, and the grid was washed with 0.5 mL DEGDME and vacuum dried inside the glove box. Next, the grid was loaded into a Gatan cryo-specimen holder inside the glove box. The Cu TEM grid was quickly transferred (~10 s air exposure) from the argon environment into the TEM load lock, where it was brought to vacuum. Next, the temperature of the sample was decreased using liquid nitrogen and maintained at ~175 °C during imaging. The XPS samples were transferred into the chamber via a sealed Ar-filled vessel to avoid exposure to air. Regarding the changes in the electrolyte resistance (Fig. 4b), the averages and deviations were based on three consecutive EIS measurements.

First-principles computations

Density functional simulations were carried out using the Gaussian 16 package. We used the B3LYP functional with the D3 version of Grimme (empirical dispersion = g3d) and a 6-31+G basis set. Solvent effects were included with the polarization continuum model (PCM) using DEGDME as the solvent (dielectric constant = 7.39). The reduction potential was computed using the thermodynamic cycle described by Han et al. and Borodin et al. The reduction potential, $E_{red}$, was calculated based on the equation below:

$$E_{red} = \left[ \frac{\Delta G_{\text{red}} + \Delta G_{\text{sol}}(S^\text{+}) - \Delta G_{\text{sol}}(S)}{F} \right] - 1.8$$

where $\Delta G_{\text{red}}$ is the free energy of electron affinity computed in the gas phase, $\Delta G_{\text{sol}}(S^\text{+})$ and $\Delta G_{\text{sol}}(S)$ are the Gibb’s free energy for the solvation process of charged and neutral species, respectively, and $F$ is the Faraday constant. To convert the potential to the Na/Na’ reference according to previous works, 1.8 V was extracted from the equation. The International Union of Pure and Applied Chemistry recommends a value of 4.42–4.44 V on the absolute potential scale.

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The data that support the findings of this study are available within the text, including the Methods and Supplemental information. Raw datasets related to the current work are available from the corresponding author on reasonable request.

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Author contributions

C.W. and W.L. conceived the idea. C.W. conducted the electrolyte formulation, material characterization and electrochemical evaluation. J.L. synthesized the NaMXene electrodes. H.Z. and H.X. performed AFM characterization and analysis. A.C.T., P.P.S. and M.T.M. carried out the cryogenic transmission electron microscopy and corresponding analysis. S.P.-P. and G.H. conducted theoretical calculations. C.W., A.C.T., M.T.M., G.H. and W.L. cowrote the paper. All the authors discussed the results and commented on the manuscript.

Competing interests

The authors declare no competing interests.
