



Cite this: DOI: 10.1039/d6ya00014b

# Deciphering solid–electrolyte interface in cellulose-montmorillonite nanocomposites for sodium batteries

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Electrolytes and their interphases are critical for emerging battery chemistries such as metal–sulphur and metal–oxygen, especially in the case of solid electrolytes, which offer attractive energy storage possibilities but involve drastic phase transitions and structural challenges. Therefore, developing improved electrolytes and interphases is a key to achieving sustainable battery performance. Here, we introduce a novel polymer composite electrolyte utilising abundant montmorillonite and cellulose nanocrystals (CNC), creating a stable interphase with the Na metal and alleviating common degradation issues. For example, this electrolyte exhibits a stability window of 2.3–5.3 V and a transference number of  $\sim 0.87$ , although its durability and performance need further improvement. FT-IR spectroscopy, XPS, and Raman spectroscopy provide valuable insights into the interfacial chemistry, as evidenced by a prominent hydroxyl stretching band associated with the CNC. While hydroxyl groups may compromise interfacial stability at the cathode, possibly causing cell degradation, they simultaneously enhance the sodium-ion mobility at the anode by facilitating favourable coordination with sodium metal. This dual function underscores the need for tuning functional groups in electrolyte design.

Received 19th January 2026,  
Accepted 22nd February 2026

DOI: 10.1039/d6ya00014b

rsc.li/energy-advances

## 1. Introduction

Recent incidents involving battery explosions highlight the critical importance of prioritizing safety in designing and deploying energy storage technologies. Solid-state batteries (SSBs), with their inherently safer design and non-flammable electrolytes, have the potential to significantly reduce the risk of such hazardous incidents.<sup>1–3</sup> The electrolyte, a central component in battery chemistry, is vital as it physically interacts with many other battery components. For example, it facilitates the movement of ions, effectively insulates electrons, and maintains stability when exposed to active materials of both highly reductive cathodes and strongly oxidative anodes. As a result, the mechanical, electrical, and chemical properties of the solid electrolyte have a critical impact on the safety and influence of various physicochemical processes within the cell. This, in turn, can lead to a greater stability of the electrochemical

system, especially when coupled with high-energy anodes such as Li, Na, and Si.<sup>4–6</sup> In this context, the use of composite polymer electrolytes with robust microscopic structures and non-flammability significantly improves operational safety, with some tolerance to overcharge/deep discharge. Moreover, solid inorganic electrolytes (SIEs) are often single-ion conductors, which leads to higher charge-transport efficiencies ( $10^{-3}$  S cm<sup>-1</sup>) than traditional liquid electrolytes, with lower transference numbers ( $\leq 0.5$ ). However, SIEs are brittle in nature, and maintaining good contact over a prolonged cycle life is difficult. Furthermore, they undergo volume changes, and a constant pressure needs to be maintained for proper contact, as elegantly demonstrated recently.<sup>7–10</sup>

The use of two-dimensional materials with large surface areas, high doping propensity, facile surface modification, and compatibility with layer-by-layer fabrication boosts ionic transport and promotes the thermal and mechanical stability of solid–polymer electrolytes, especially after composite formation.<sup>11–13</sup> Recently, two-dimensional (2D) boron nitride (BN) nanosheets, graphene oxides, and MXenes have been widely adopted as fillers and additives to enhance the ionic conductivity and mechanical strength of polymer electrolytes. However, the instability and poor processability of these nanomaterials remain the main limitations for their widespread application.<sup>14–16</sup> In this context, clay materials have

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emerged as attractive alternatives for hydrogels and semi-solid electrolytes owing to their large specific surface area, distinct physical and chemical adsorption properties, environmental friendliness, and cost-effectiveness.<sup>17,18</sup> Among these, aluminium magnesium silicate (montmorillonite or MMT) has drawn considerable research interest in battery applications. Its layered structure, featuring octahedral sheets sandwiched between two silica tetrahedral sheets, endows it with excellent chemical stability, robust mechanical properties, and environmental compatibility. Many of the above limitations could be alleviated by a composite polymer electrolyte using a suitable polymer matrix such as cellulose; hence, unsurprisingly, several recent results using MMT in various energy storage systems have shown promising outcomes.<sup>19–21</sup> For instance, He *et al.* synthesised a gel polymer electrolyte based on a polyacrylonitrile/organic montmorillonite membrane, which served as a host for LiPF<sub>6</sub> in ethylene carbonate.<sup>22</sup> Similarly, Kim *et al.* prepared a polymer composite electrolyte with PEO and LiClO<sub>4</sub> with different percentages of Li–MMT as a filler.<sup>23</sup> Zhao *et al.* synthesised a solid composite electrolyte based on a PEO matrix with LiTFSI and LLZTO in acetonitrile and MMT as a filler.<sup>24</sup> Dhatarwal *et al.* used a solution-cast method to prepare nanocomposite solid polymer electrolyte (NSPE) films incorporating MMT clay nanoplatelets based on a poly(ethylene oxide) and poly(methyl methacrylate) blend matrix with lithium tetrafluoroborate (LiBF<sub>4</sub>).<sup>25</sup> In comparison, Chen *et al.* prepared a montmorillonite composite poly(ether sulfone) ion-conducting membrane (MMT-M) where MMT, with high mechanical strength and negative charge, was used as a functional coating compounded on the matrix (PS-M) consisting of poly(ether sulfone) and sulfonated poly(ether ketone) for a zinc–iron flow battery.<sup>26</sup> However, in these studies, montmorillonite was used as a filler either to change the domain size or to increase the mechanical strength and thermal stability. None of the previous works has investigated MMT as a candidate for polymer composite electrolytes containing free mobile Na<sup>+</sup> ions within the interlayer. Also, the proposed electrolyte is flexible; thus, the application of pressure is not required, and it exhibits room temperature conductivity in the order of 10<sup>−5</sup>–10<sup>−6</sup> S cm<sup>−1</sup>, unlike its ceramic oxide counterpart. These features make MMT a promising candidate for sodium-based energy storage devices.

Accordingly, we recently reported clay–nanocellulose composite polymer electrolytes in which tuning the domain size and correlating structure–property relationships led to enhanced ionic conductivity.<sup>27</sup> In a subsequent study on LAPONITE<sup>®</sup>-nanocellulose composite polymer electrolytes, we showed that hydroxyl groups interfere (1.5% concentration increase) at the cathode–electrolyte interface, resulting in interfacial instability and capacity fading during cycling.<sup>28</sup> However, several questions related to the interfacial stability were elusive, including the precise role of hydroxyl functionalities in governing cathode–electrolyte interfacial chemistry, the dual role of the hydroxy groups, and degradation analysis of the coin cells coupled with the MoS<sub>2</sub> electrode during cycling. In a continuation, here we demonstrate the electrochemical properties and

solid–electrolyte interface issues, utilising MMT and CNCs to fabricate free-standing polymer nanocomposite films. The negatively charged CNC rods interact non-covalently with the positively charged edges of the MMT clay to presumably form a gel, as indicated by the zeta potential measurements.<sup>27</sup> The free-standing films of CNC/MMT composite, fabricated through the evaporation-induced self-assembly process, exhibit complex ordered structures, which can boost the sodium ion transport through their ordered network. These films have a wide electrochemical stability window of 2.3–5.3 V, and a high transference number of 0.87, facilitating their usage as a battery electrolyte. The as-prepared electrolyte has been coupled with MoS<sub>2</sub> free-standing sheets as cathodes and Na metal as anodes for the fabrication of Na cells.

## 2. Materials and methods

Few-layer MoS<sub>2</sub> nanosheets were prepared using the liquid-phase exfoliation method. Typically, bulk MoS<sub>2</sub> (3 g) (Sigma Aldrich) was stirred with *n*-butyl lithium in 1.5 M hexane (25 mL) (SRL) for 48 h under argon. The black slurry obtained was vacuum-filtered with hexane, and the dried powder was further dispersed in water and sonicated for 1 h for exfoliation. Lastly, the dispersed MoS<sub>2</sub> was centrifuged to obtain a few layers of exfoliated MoS<sub>2</sub> (~2.5 g). A typical electrode slurry was prepared using 85% MoS<sub>2</sub>, 5% PEDOT-PSS, and 10% CNC, which was coated onto a polyvinyl acrylate sheet with a uniform thickness using a doctor-blade. To obtain the free-standing films, the prepared composite films were immersed in acetone. Free-standing composite polymer electrolyte thin films with a composition of 2.5C:1M (C-CNC; M-MMT) were prepared by the casting method, as discussed elsewhere.<sup>27</sup>

### 2.1 Material characterization

FTIR analysis for the CNC–MMT polymer composites and MoS<sub>2</sub> electrodes was carried out in the 450 to 4000 cm<sup>−1</sup> range with a resolution of 2 cm<sup>−1</sup>, using a PerkinElmer ATR C117460 instrument. Raman analysis was performed with a confocal and resonance Raman instrument (LabRam, Horiba Scientific) with a spatial resolution of 1 μm, by placing the samples on top of quartz slides; a 40 mW, 405 nm excitation laser was used. X-ray photoelectron spectroscopy (XPS) measurements were performed with a Thermo Scientific–ESCALAB 250 xi instrument. Field-emission gun transmission electron microscope (FEG-TEM) images were acquired with a JEOL JEM-2100F instrument to observe a lattice resolution of 0.1 nm and point-to-point resolution of 0.19 nm with 200 kV acceleration voltage. Powder X-ray diffraction of the samples was recorded with a Smart Lab 9 kW Rigaku using a monochromatic Cu Kα source (λ = 1.5406 Å) at 45 kV and 200 mA.

### 2.2 Electrochemical characterisation

12 mm electrodes and 18 mm electrolyte discs were dissected using a cutter, followed by vacuum-drying at 70 °C, before fabricating the CR2032 cells inside an Ar-filled glove box. A



battery tester BCS 805 (Biologic, France) was used to measure galvanostatic charge/discharge, and linear sweep voltammetric studies of the MoS<sub>2</sub> electrode were carried out using the composite electrolyte. Electrochemical impedance studies were performed for the electrolyte at an open-circuit voltage using an AC amplitude of 5 mV (RMS value) in the range of 1 MHz–0.1 Hz with a Solartron 1470E electrochemical workstation. The electrochemical stability window of the 2.5C:1M thin film was investigated using linear sweep voltammetry (LSV). By placing the thin film between a stainless-steel working electrode and a metallic sodium counter (reference) electrode at a scan rate of 0.1 mV s<sup>-1</sup> between 2.3 to 6 V. For chronoamperometry measurements, a similar thin film of the composition 2.5C:1M was sandwiched between two freshly prepared Na-metal discs, and the current–time response was measured under a constant DC bias of 10 mV. The reversible ion transport through the 2.5C:1M thin film and the sodium electrodeposition behaviour were studied using galvanostatic cycling in a symmetric (Na/Na) cell configuration at room temperature by probing the reversible voltage response under a constant current density of ±0.1 mA cm<sup>-2</sup> for a sodium plating/stripping capacity of 0.1 mA h cm<sup>-2</sup>.

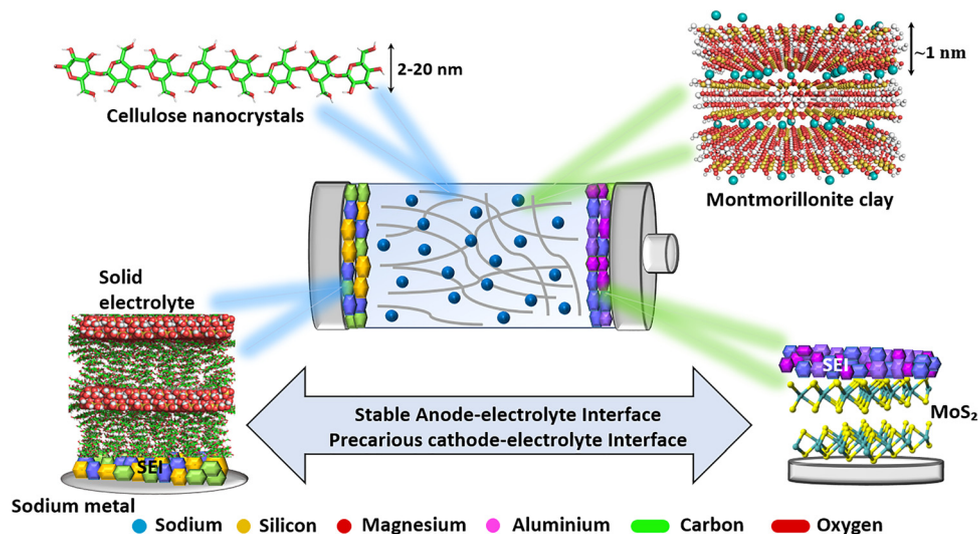
### 3. Results

The schematic in Scheme 1 illustrates that the hydroxy functional groups on CNC promote the formation of a stable anode–electrolyte interface with sodium metal, enhancing interfacial stability. In contrast, at the cathode side, these hydroxy groups interfere with solid–electrolyte interface (SEI) formation, leading to a heterogeneous and unstable cathode–electrolyte interface.

In order to gain insights into the structural changes in CNC–MMT composites and MoS<sub>2</sub> after exfoliation, we performed XRD and Raman spectroscopy to study the systematic changes in *d*-spacing. Accordingly, Fig. 1(a) and (b) present the XRD

patterns of the CNC–MMT composite and MoS<sub>2</sub> nanosheets, respectively. In the CNC–MMT composite, the characteristic (001) plane exhibits an interesting shift in  $2\theta$  from 7.2° to 6.1°, corresponding to an increase in *d*-spacing of 2.08 Å, which reveals that the one-dimensional (1D) CNC was successfully intercalated into the two-dimensional (2D) MMT nanosheets.<sup>29,30</sup> This was further observed from the cross-sectional SEM and small-angle X-ray scattering (SAXS) measurements.<sup>27</sup> Furthermore, while broadening is observed in CNC and MMT diffraction peaks, no significant changes in peak asymmetries occur, since the stacking disorder is perhaps negligible, preserving the integrity of the layered structure of the Montmorillonite. Similarly, the XRD pattern of the MoS<sub>2</sub> nanosheets (Fig. 1(b)) reveals a broadening of the (002) peak after exfoliation compared with that of the pristine sample, indicating the stacking of the 2D nanosheets along the *z*-direction. Interestingly, the (002) peak shifts to a lower angle, while the (100) peak shifts to a higher angle, suggesting an expansion along the (001) direction and in-plane compression in the synthesized MoS<sub>2</sub> nanosheets (6.15 Å). Additionally, a new (001) peak at  $2\theta \approx 6.4^\circ$  corresponds to a layer separation of approximately 5.5–6 Å during restacking, indicating the formation of the 1T phase. The broadness of the (002) peak and the presence of the (001) peak suggest that the nanosheets are randomly arranged during restacking, with a significant expansion of the interlayer spacing.<sup>31,32</sup>

Fig. 1c compares the Raman spectra for the pristine and exfoliated MoS<sub>2</sub> nanosheets. The spectra reveal the characteristic E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> bands at 377 and 402 cm<sup>-1</sup>, respectively, for pristine MoS<sub>2</sub>, corresponding to the in-plane and out-of-plane vibrations of sulphur atoms relative to molybdenum atoms. Furthermore, the exfoliated MoS<sub>2</sub> exhibits a peak at ~222 cm<sup>-1</sup> corresponding to the J<sub>2</sub> phonon mode, characteristic of metallic 1T-MoS<sub>2</sub>, which contributes to forming a super-lattice structure,<sup>33,34</sup> although the J<sub>1</sub> band is missing, as these are excitation-dependent. The exfoliated nanosheets exhibit a blue



Scheme 1 Illustration of the formation of the solid–electrolyte interface on the anode and cathode.



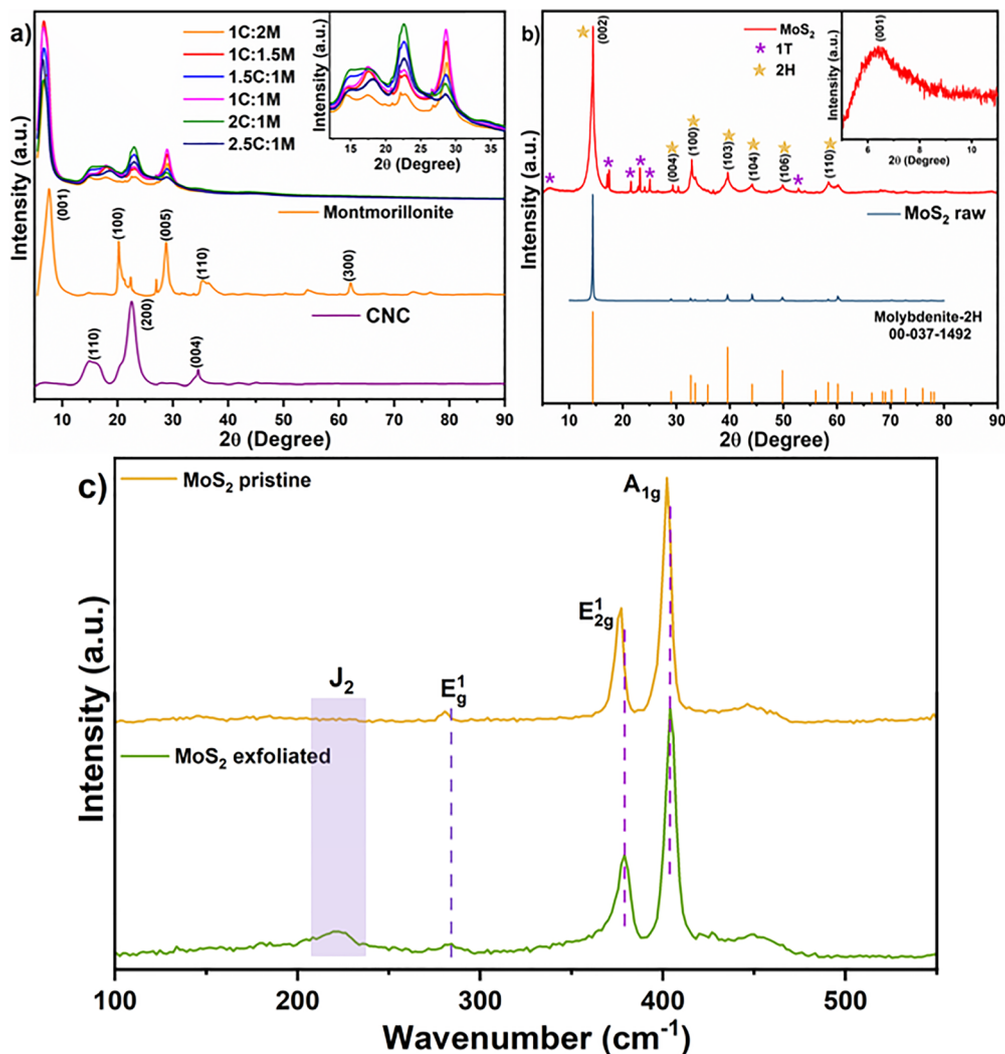


Fig. 1 (a) X-ray diffractograms of CNC–MMT composite electrolytes and (b) comparative diffractograms of pristine and exfoliated MoS<sub>2</sub> nanosheets. (c) Comparative Raman spectra of pristine and exfoliated MoS<sub>2</sub> nanosheets (C and M indicate CNC and MMT, respectively).

shift to 379 and 403 cm<sup>-1</sup> for these bands, while the intensity ratio ( $I_E/I_A$ ) for pristine and exfoliated samples is 0.46 and 0.4, respectively, suggesting that the particle size remains mostly unchanged after exfoliation.<sup>35,36</sup> The frequency, intensity, and width of the E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> bands are influenced by the thickness (layer count) of the MoS<sub>2</sub> flakes. Many studies indicate that exfoliation typically narrows the separation between these bands, with a common shift to 381 cm<sup>-1</sup> for E<sub>2g</sub><sup>1</sup> and 406 cm<sup>-1</sup> for A<sub>1g</sub>. However, some research demonstrates that, depending on the quality of the bulk MoS<sub>2</sub>, broad peaks for E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> can appear at 375 and 402 cm<sup>-1</sup>, respectively.<sup>37,38</sup> Interestingly, in such cases, exfoliation leads to a blue shift for both bands to 381 cm<sup>-1</sup> (E<sub>2g</sub><sup>1</sup>) and 405 cm<sup>-1</sup> (A<sub>1g</sub>), rather than reducing their separation. This study also identifies a similar blue shift. Instead of diminishing the band separation, the E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> modes shift to higher wave numbers. Nevertheless, the shift in the A<sub>1g</sub> band is less pronounced compared with the E<sub>2g</sub><sup>1</sup> band, yielding a band separation of 24 cm<sup>-1</sup> versus

26 cm<sup>-1</sup> in bulk MoS<sub>2</sub>. This shift may stem from lattice strain effects induced in MoS<sub>2</sub> nanosheets by interactions with *n*-butyl lithium (*n*-butyl Li), as Yang *et al.*<sup>39</sup> describe, or from defects developed during exfoliation. This interaction causes greater in-plane strain than out-of-plane strain, consistent with Raman spectra showing similar shifts in the E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> bands under in-plane lattice strain.

The deconvoluted Mo 3d spectrum of free-standing 2H-MoS<sub>2</sub> shows two prominent peaks at 232.3 and 229.1 eV, corresponding to Mo<sup>4+</sup> 3d<sub>3/2</sub> and Mo<sup>4+</sup> 3d<sub>5/2</sub>, respectively, while the exfoliated nanosheets show peaks at 232.18 and 229.02 eV, respectively, with a shift of 0.12 and 0.19 eV. The S 2p<sub>3/2</sub> and 2p<sub>1/2</sub> peaks of MoS<sub>2</sub> exhibit similar downshifts (Fig. 2b and d) as a result of the presence of the metallic 1T phase. It has been observed that the components from the 1T phase appear at a binding energy that is ~0.9 eV lower than their 2H counterparts. The prominent peaks at ~233 and 235 eV show oxidation of Mo, corresponding to Mo<sup>6+</sup>. Similarly, for S 2p,



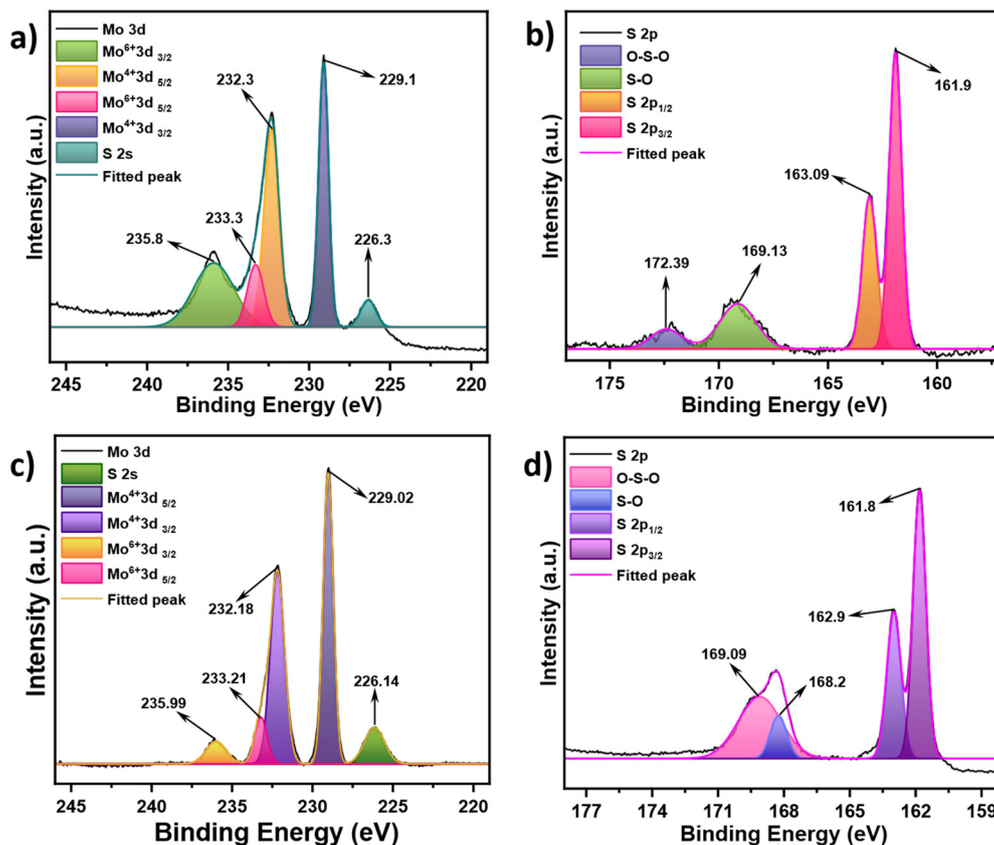


Fig. 2 X-ray photoelectron spectra (XPS) of (a) and (b) free-standing  $\text{MoS}_2$  (Mo: 12 meV and S: 19 meV shift) and (c) and (d) exfoliated  $\text{MoS}_2$  film.

peaks observed between 168 and 170 eV, after exfoliation, indicate that sulphur atoms might have oxidized or the shift is due to the interaction with PEDOT/PSS.<sup>40–42</sup>

The deconvoluted post-cycling Mo 3d spectrum (Fig. S1a) reveals an interesting shift in the peaks, providing valuable insights into the electronic interactions possible during cycling. For example, two prominent peaks at 228.8 and 231.2 eV correspond to  $\text{Mo}^{4+} 3d_{5/2}$  and  $\text{Mo}^{4+} 3d_{3/2}$ , respectively. Additionally, a  $\text{MoO}_3$  peak appears at 233.9, indicating that some  $\text{MoS}_2$  was oxidized during the cycling, which is attributed to interfacial reactions and the interference of hydroxyl groups. Moreover, the band separation between  $3d_{5/2}$  and  $3d_{3/2}$  is 3.2 eV and 2.4 eV before and after cycling, respectively, indicating a significant change in the electronic structure, which could be attributed to the oxidation.<sup>43</sup> Similarly, the S 2p spectrum (Fig. S1b) shows a shift in the peaks and the emergence of satellite peaks. The peak at 168.4 eV corresponds to the  $\text{S-O}_4$  bond, while 167.3 eV represents the S–O bond, 165.4 eV is attributed to the Mo–O–S bond, and 161 eV reveals the  $\text{S}^{2-}$  arising from  $\text{MoS}_2$ . The deconvoluted O 1s spectra (Fig. S1c) show several peaks corresponding to C–O bonds, mainly linking to the CNC. The peak at 535.6 eV corresponds to the C–O–C bond, 534.6 eV, the C–OH bond, and the peak at 530.7 represents the C=O/Mo–O bond.<sup>44</sup>

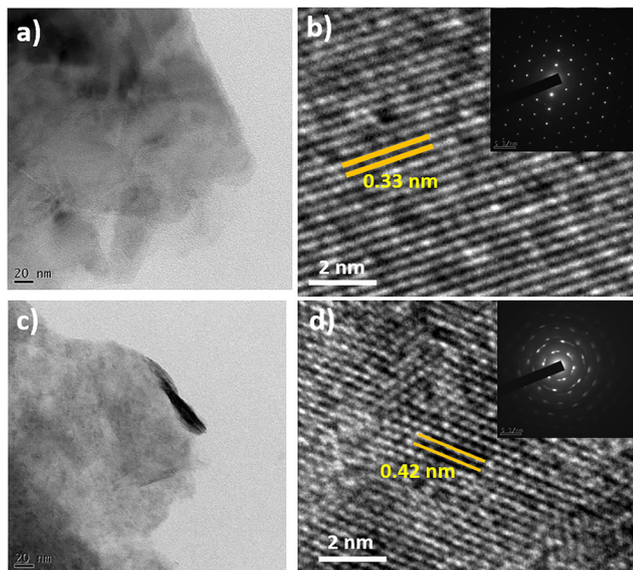
These observations collectively confirm that notable chemical and electronic modifications occur in the  $\text{MoS}_2$

electrode during cycling, with profound implications for the degradation of the cell. The emergence of  $\text{Mo}^{6+}$  and oxygenated sulphur species, along with the reduced Mo 3d spin-orbit splitting, clearly indicates partial oxidation of  $\text{MoS}_2$  and the formation of mixed Mo–O–S environments. The presence of S–O and  $\text{SO}_4^{2-}$  species further supports interfacial reactions with the electrolyte or hydroxyl groups. Overall, the XPS analysis demonstrates that cycling induces interfacial oxidation and structural reorganization within the  $\text{MoS}_2$  framework, also observed in post-cycling SEM, which may significantly influence its electrochemical behaviour and cell failure.

High-resolution transmission electron microscopy (HR-TEM) analysis on both bulk and exfoliated samples revealed typical 2D sheets consisting of thin nanosheets (Fig. 3). The bright and transparent contrast indicates the typical features shown by thin nanosheet structures, with the exfoliation of bulk  $\text{MoS}_2$  into a few monolayers with a  $d$ -spacing of  $\sim 0.42$  nm, in agreement with the XRD results shown in Fig. 3d. Also, the selected area electron diffraction (SAED) pattern in the inset shows a hexagonal structure with rings, suggesting that the exfoliated nanosheets retain the crystal structure of  $\text{MoS}_2$  with a slight rearrangement in the sheets during re-stacking. These nanosheets are in excellent agreement with the results of previous studies.<sup>45–47</sup>

Fig. S2(a–c) presents comparative SEM images of the pristine 2.5C:1M composite electrolyte and after cycling. The images





**Fig. 3** High-resolution transmission electron microscopic (HR-TEM) images of (a) and (b) pristine MoS<sub>2</sub> nanosheets and (c) and (d) exfoliated MoS<sub>2</sub> nanosheets resulting in an enlarged *d*-spacing of ~0.42 nm (Insets-SAED pattern).

show no visible cracks, suggesting the absence of dendrite formation. Corresponding EDX mapping images for the pristine and post-cycled 2.5C:1M samples (Fig. S3) confirm the uniform distribution of Si, Al, Mg, C, O, and Na throughout the composite polymer electrolyte. Furthermore, as shown in Tables S1 and S2, the oxygen atomic weight percentage increases notably from 33% in the pristine state to 38% after cycling, indicating the formation of hydroxyl groups. Likewise, Fig. S2(d and e) compares the SEM images of pristine and post-cycled MoS<sub>2</sub>. The pristine MoS<sub>2</sub> exhibits a sheet-like morphology, while it shows agglomeration post-cycling. EDX mapping (Fig. S4) further confirms the uniform distribution of Mo, S, and O; however, while oxygen is absent in the pristine sample, it appears in the post-cycled MoS<sub>2</sub>, highlighting oxygen incorporation at the interface due to hydroxyl groups during cycling.

Various characterisations have been utilised to probe the electrochemical performance of the as-prepared composite polymer electrolyte. For instance, Fig. 4a shows the linear sweep voltammogram (LSV), which is one of the fundamental requirements for analysing the electrochemical stability window for the charge–discharge processes. When the thin film electrolyte is placed between a stainless-steel working electrode and a metallic sodium counter (reference) electrode, the LSV profile spanning 2.3 to 5.5 V suggests robust operational stability. A slight increase in current (in  $\mu\text{A}$ ) was observed around ~2.7 V, which could be attributed to the oxidation of hydroxy groups or due to a soft short-circuit. With repeated LSV, we did not observe the small peak around ~2.7 V (Fig. S5), and a stability window of ~5.2 V was observed. Fig. S6 shows the Nyquist plot of bare CNC, which shows a conductivity of  $1.2 \times 10^{-5} \text{ S cm}^{-1}$  compared with that of the composite 2.5C:1M ( $9.2 \times 10^{-5} \text{ S cm}^{-1}$ ).<sup>27</sup>

The single-ion conductivity feature of the “as prepared” electrolyte (2.5C:1M) was investigated using chronoamperometry (Fig. 4b). The transference number for the Na<sup>+</sup> ion ( $t_{\text{Na}^+}$ ) was calculated using the well-known Bruce–Vincent method:<sup>48–50</sup>

$$T_+ = \frac{I^s(\Delta V - I^0 R_1^0)}{I^0(\Delta V - I^s R_1^s)}$$

where  $\Delta V$  is the potential applied across the cell,  $T_+$  is the cationic transference number,  $R_1^0$  and  $R_1^s$  are the initial and steady-state resistances of the passivating layers, respectively, and  $I^0$  and  $I^s$  are the initial and steady-state currents, 0.48 and 0.36 mA, respectively. The 2.5C:1M composite polymer electrolyte was sandwiched between Na metal discs, and the current–time response was measured under a constant DC bias potential of 10 mV. The measured initial current  $I^0$  reflects the movement of both cations and anions, while the final steady-state current  $I^s$  results exclusively from the motion of cations. The slight increase in the resistance ( $R_{\text{initial}} = 200 \Omega$  and  $R_{\text{ss}} = 275 \Omega$ ) upon polarization can be explained by the interaction between the ohmic polarization and sodium metal. Based on these results, the  $t_{\text{Na}^+}$  was calculated to be 0.87.

The initial sharp current spike corresponds to the charging of the electrical double layer at the electrode–electrolyte interface, along with the instantaneous faradaic reaction of electroactive species present directly at the electrode surface. This is followed by a rapid decay in current, which reflects the diffusion-controlled transport of reactants from the bulk solution to the electrode, typically described by the Cottrell equation, where the current decreases with  $t^{-1/2}$  as the diffusion layer expands over time. At longer times, the plot exhibits a steady-state region in which the current stabilizes and becomes nearly constant, indicating that the system has reached a dynamic balance due to thin-layer cell effects or persistent surface-controlled processes, such as adsorption or slow kinetics. Furthermore, the current jump observed at ~2000 s may arise from a transient interfacial instability, after which the system regains its structure and displays stable current behaviour.

The Na/2.5C:1M/Na symmetric cells (Fig. 4c), when cycled galvanostatically to study the compatibility with the Na electrode, provide the initial overpotential,  $\eta \approx 35 \text{ mV}$ , indicating a stable electrode–electrolyte interface. This overpotential then drops to a negligible value of 2.5 mV and remains stable for 130 h. After this, it slightly increases to ~4 mV for another 100 h, potentially due to small dendrite growth leading to the increase in solid electrolyte interface (SEI) resistance, which may result in a thickening of the SEI layer and voltage fluctuations. This cell exhibits stable stripping/plating for more than 300 h without ohmic losses associated with poor stripping or degradation of the composite polymer electrolyte.<sup>51–54</sup> Fig. S7 shows the Nyquist plot of symmetric 2.5C:1M, having a resistance of  $630 \Omega \text{ cm}^{-2}$ . This shows one diffused semi-circle attributed to charge-transfer resistance and interfacial reactions between the electrolyte and sodium metal.



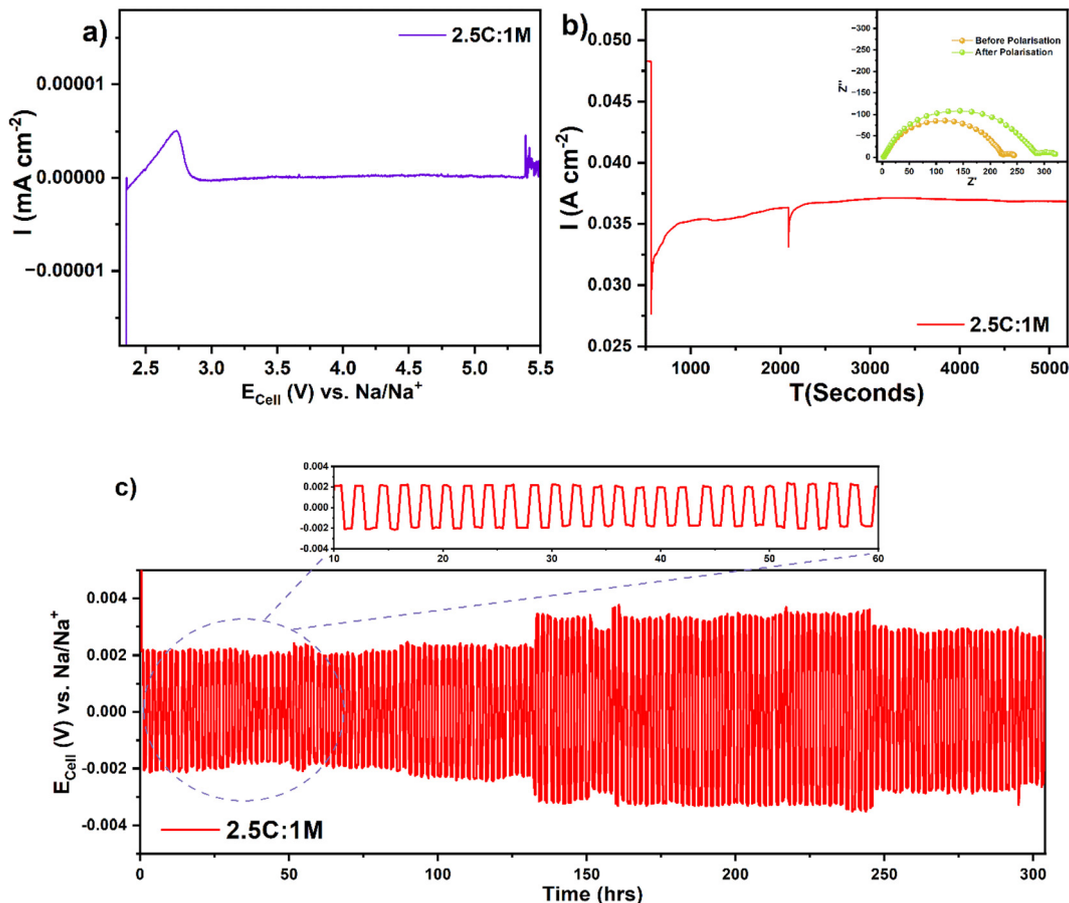


Fig. 4 (a) Linear sweep voltammetry (LSV) plot of the composite polymer electrolyte (CPE) (2.5C:1M) at a scan rate of  $0.1 \text{ mV s}^{-1}$ . (b) Chronoamperometric plot and (c) stripping/plating of the symmetrical cell (Na/CPE/Na) at a current density of  $0.1 \text{ mA cm}^{-2}$  with a capacity of  $0.1 \text{ mA h cm}^{-2}$  (inset: plating behaviour from 10–60 h).

The failed cells were disassembled, and the mechanism of degradation was investigated using several analytical techniques, such as Raman spectroscopy, XPS, and FT-IR spectroscopy, along with electrochemical methods.

## 4. Post-mortem analysis

Raman and FT-IR spectroscopy are powerful analytical techniques to study molecular vibrations, chemical compositions, and structural properties of materials. In crystalline substances, molecular vibrations can split due to intermolecular interactions within the lattice. By analyzing these split vibrations, interesting changes in properties, such as crystallinity, phase composition, shape, and orientation, can be determined, offering insights into structural variations within composite materials. In order to elucidate the interfacial reactions between the electrolyte and the  $\text{MoS}_2$  cathode, *ex situ* FT-IR and Raman spectroscopies were performed. Accordingly, Fig. 5a shows the comparative FT-IR analysis of the cycled symmetric and asymmetric 2.5C:1M cells with pristine composition. The films reveal that the peaks intensified in the range

of  $1700\text{--}1200 \text{ cm}^{-1}$ . The OH bending peak at  $1638 \text{ cm}^{-1}$  broadened and exhibited several new peaks, and at  $\sim 3300 \text{ cm}^{-1}$ , suggesting interfacial reactions between the Na metal and the electrolyte. Similarly, in Fig. 4b, the OH bending peak at  $1638 \text{ cm}^{-1}$  for the cycled  $\text{MoS}_2$  thin films broadened, and the free OH peak at  $3628 \text{ cm}^{-1}$  disappeared, while the intramolecular OH peak at  $3300 \text{ cm}^{-1}$  intensified, revealing the interaction of the  $\text{MoS}_2$  with the OH groups of the composite polymer electrolyte. The Si–O stretching peak at  $998 \text{ cm}^{-1}$  exhibits a slight blue shift to  $1005 \text{ cm}^{-1}$ , revealing the electronic coupling with  $\text{MoS}_2$ .<sup>55,56</sup>

Accordingly, Fig. 6a shows the Raman spectra of the comparative cycled symmetric and asymmetric composite polymer electrolyte with the pristine 2.5C:1M. New peaks emerged at  $100\text{--}400 \text{ cm}^{-1}$ , revealing the interaction between the Na metal and the electrolyte. The peaks in this range ( $144, 187, 214 \text{ cm}^{-1}$ ) correspond to the  $O_h$  symmetry of  $\text{AlO}_6$  or  $\text{MgO}_5\text{OH}$  octahedral sheets. The peaks in the range  $900\text{--}1200 \text{ cm}^{-1}$  correspond to the  $T_d$  symmetry of the  $\text{SiO}_4$  tetrahedral sheets. The peak at  $3648 \text{ cm}^{-1}$  is assigned to OH vibrations, while the broad envelope below  $3600 \text{ cm}^{-1}$  (centred at  $2676 \text{ cm}^{-1}$ ) is attributed to OH vibrations in  $\text{H}_2\text{O}$  molecules. These peak intensities have



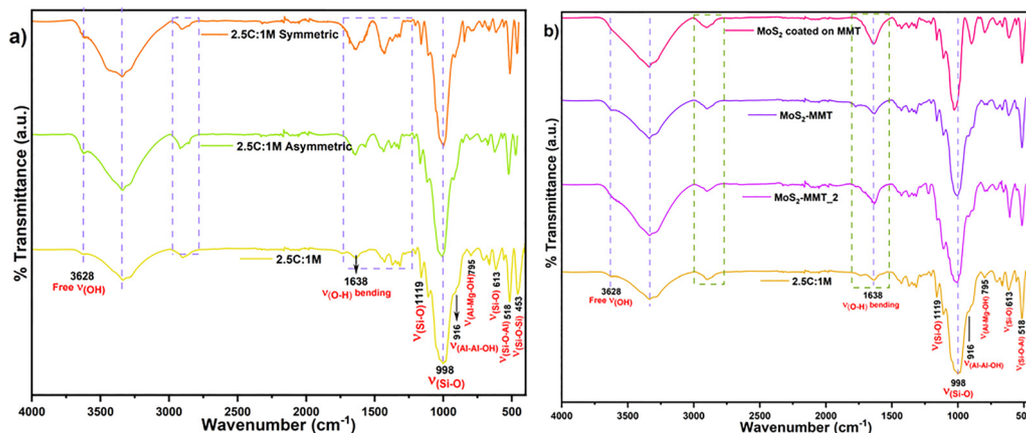


Fig. 5 (a) Comparative FT-IR (post-mortem) analysis of the cycled symmetric and asymmetric 2.5C:1M cells with the fresh composite polymer electrolyte and (b) cycled cells with MoS<sub>2</sub>-MMT thin films with the 2.5C:1M electrolyte.

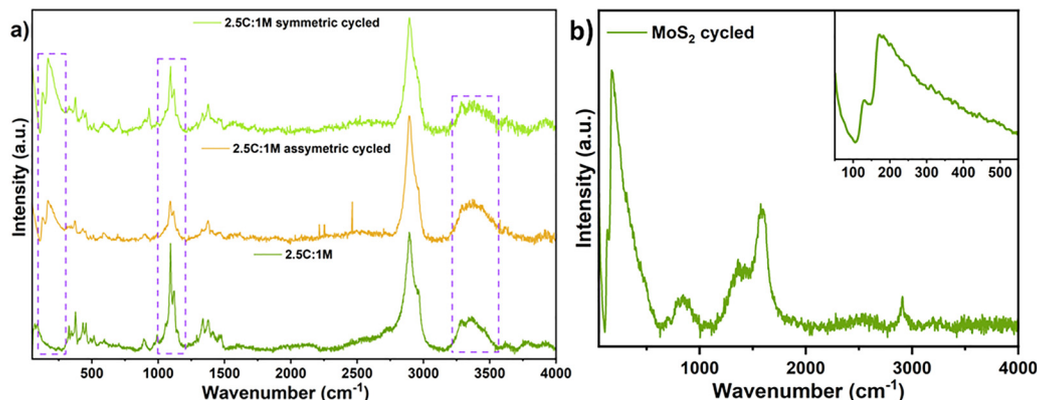


Fig. 6 Comparative Raman post-mortem analysis of the cycled symmetric and asymmetric 2.5C:1M with the fresh composite polymer electrolyte, showing the changes with cycling and (b) cycled MoS<sub>2</sub>-MMT thin film.

probably been reduced due to interfacial reactions between the Na metal and the electrolytes.<sup>57–59</sup>

Fig. 6b shows the cycled MoS<sub>2</sub> electrode. The E<sub>2g</sub><sup>1</sup>, A<sub>1g</sub> and J<sub>2</sub> bands (centred at ~377, 402 and 222 cm<sup>-1</sup>, respectively) merged with the O<sub>h</sub> symmetry of AlO<sub>6</sub> or MgO<sub>5</sub>OH octahedral sheets. Also, several other peaks emerged with cycling, centred around 800, 1300, 1650, and 2900 cm<sup>-1</sup>, indicating interfacial reactions.

## 5. Discussions

From the above results, it is evident that exfoliation of MoS<sub>2</sub> results in the formation of few-layered 2D nanosheets with a *d*-spacing of ~0.42 nm, through expansion in the (002) plane and contraction in the (100) plane, respectively, with an additional separation of 5.5–6 Å. Also, the crystallinity is retained after exfoliation of MoS<sub>2</sub>, as evident from Raman analysis. The composite polymer electrolyte shows a change in interlayer spacing of 2.08 Å, with the intercalation of CNC nanorods into the 2D MMT nanosheets. This design facilitates Na-ion

mobility, as evident from the high transference number of 0.87. Furthermore, we investigated the transference number using pulsed-field gradient NMR spectroscopy. However, due to the higher *T*<sub>2</sub> relaxation, the diffusion NMR of Na<sup>+</sup> ions was too short to observe. Also, this electrolyte provides an electrochemical stability window of 2.3–5.3 V, which implies this electrolyte could be useful for high voltage cathodes. *Ex situ* analysis reveals the presence of hydroxy groups at the electrode-electrolyte interface, which assists in stable cycling at the anode interface, as evident from galvanostatic cycling with an initial  $\eta = 35$  mV, and stable stripping/plating for more than 300 h. In contrast, these hydroxy groups could interfere at the cathode-electrolyte interface, which leads to cell failure by altering the crystallinity of MoS<sub>2</sub>, as observed by Raman analysis, as the characteristic E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> bands merged.<sup>60–62</sup> More interestingly, distinct changes in the morphology occur, as evident from the SEM images, pointing to the fact that agglomeration takes place in several locations, reducing the high surface area and disrupting the layer-by-layer structure. This is also in complete agreement with data obtained from XPS analysis, suggesting the role of surface oxygen in degradation.



Incorporation of 2D heterostructures or Janus-type architectures might overcome these issues as an ingenious strategy to achieve more stable electrode interfaces, as the dual-surface chemistry of such materials may facilitate controlled interactions with hydroxyl groups and potentially provide preferential Na-ion transport pathways.<sup>63–65</sup> However, these concepts alone cannot fully resolve the broader interfacial instabilities in solid-state systems. A more comprehensive approach is asymmetric electrolyte engineering, which aims to construct electrolytes capable of forming stable interphases at both anode and cathode sides, as well as across catholyte–anolyte boundaries.<sup>66–68</sup> In theory, multilayer solid electrolytes can integrate complementary functionalities across layers, thereby offsetting individual limitations and producing a Janus-like synergistic behaviour. Nevertheless, the inherent incompatibility between dissimilar electrolyte phases and their interfaces with electrodes remains a central challenge that must be directly addressed rather than assumed to be solved.

To mitigate these issues, asymmetric design strategies attempt to satisfy the distinct electrochemical and mechanical requirements of each electrode by assigning tailored roles to different electrolyte components. Such asymmetry can be introduced at both the macroscale and microscale.

- Macroscale asymmetry involves assembling bi- or multi-layer electrolytes composed of disparate materials, typically on the millimetre to centimetre scale, where each layer contributes a specific set of beneficial properties (*e.g.*, mechanical robustness, ionic conductivity, oxidative stability), ably supported by computational modelling and simulations.

- Microscale asymmetry, by contrast, focuses on fine-tuning the chemistry and structure of a single electrolyte phase at the micro- to nano-scale, to generate multifunctional behaviour while avoiding miscibility conflicts, interfacial incompatibility, or excessive interfacial impedance that often arise in macroscale multilayer systems.

Recent studies illustrate the conceptual value of such designs. For example, Tan *et al.* developed an asymmetric fire-retardant quasi-solid polymer electrolyte to address thermal runaway and interfacial instability in lithium metal batteries. A PVDF-HFP gel incorporating organophosphate flame retardants provided enhanced compatibility and thermal stability with high-voltage Ni-rich cathodes, while a polyether gel, synthesized *in situ*, formed a resilient interphase on the lithium metal anode, effectively suppressing dendrite formation.<sup>69</sup> Similarly, Chen *et al.* reported a Zn<sup>2+</sup>-conductive inorganic SSE based on 2D porphyrin paddlewheel framework MOF sheets; this layer promoted uniform Zn deposition, isolated the metal from bulk aqueous electrolyte, and facilitated Zn(H<sub>2</sub>O)<sub>6</sub><sup>2+</sup> desolvation, resulting in dendrite-free and hydrogen-free cycling. The complementary PAM hydrogel catholyte supplied H<sup>+</sup> carriers and water molecules that acted as lubricants to enable more facile Zn<sup>2+</sup> insertion/extraction.<sup>70</sup> Zhang *et al.* further demonstrated an asymmetric structural design using a tape-cast polyimide membrane, which provided resistance to lithium dendrite formation while maintaining compatibility with high-voltage cathodes.<sup>71</sup>

Together, these studies suggest that, while asymmetric electrolyte architecture is a promising framework, its success relies not merely on assembling multiple layers, but on rationally assigning distinct structural and chemical roles to each component. The approach must still contend with unresolved challenges, including phase compatibility, mechanical integrity, and long-term interfacial stability, before it can be considered as a broadly applicable strategy for next-generation solid-state batteries.

## 6. Conclusions

The CNC–MMT composite polymer electrolyte shows enhanced interfacial stability with Na metal, revealing an initial overpotential of ( $\eta$ ) 35 mV, inferring the formation of a stable electrode–electrolyte interface. This also shows a wide electrochemical stability window of 2.3–5.3 V with a high transference number of  $\sim 0.87$ . FT-IR and Raman spectroscopic analyses provide important clues to understanding the interfacial chemistry, since a pronounced hydroxyl stretching band is observed, attributed to the CNC component of the electrolyte. Furthermore, XPS and EDX analyses verify the presence of surface oxygen. In contrast, this hydroxyl functionality appears to disrupt interfacial stability at the cathode, as the E<sub>2g</sub><sup>1</sup> and A<sub>1g</sub> bands have merged, reducing the crystallinity and potentially contributing to cell failure during extended cycling. However, it simultaneously facilitates enhanced sodium-ion mobility at the anode interface, with stable cycling for more than 300 h; likely by promoting favourable ion coordination with sodium metal. This dual behaviour highlights the complex interplay between electrolyte chemistry and electrode interfaces, underscoring the need for precise functional group tuning in future electrolyte designs.

## Conflicts of interest

The authors declare no competing financial interest.

## Data availability

All the data supporting the findings of this study are fully available within the article. No additional datasets were generated or analysed beyond those reported in this manuscript.

Supplementary information (SI) is available. See DOI: <https://doi.org/10.1039/d6ya00014b>.

## Acknowledgements

R. K. P. acknowledges the Department of Science and Technology for an INSPIRE Faculty Award Grant [DST/INSPIRE/04/2016/002370] and the Core Research Grants [CRG/2020/006281 and CRG/2021/004759] from ANRF, Government of India. V. K. P. would like to thank the SERB for the J. C. Bose Fellowship (JCB/2020/000018). S. M. thanks the SAIF, IIT Bombay, for the HR-TEM measurements. C. T. acknowledges support from the



Department of Science and Technology, Government of India, for the INSPIRE fellowship.

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