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The impact of electrode conductivity on electrolyte interfacial structuring and its implications on the Na^{0/+} electrochemical performance*

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Molecular and ionic assemblies at electrode/liquid electrolyte interfaces, i.e., the electric double layer (EDL), define battery performance by directing the formation of stable interphases. An unstable interphase can hamper metal-cation diffusion, lead to continuous electrolyte consumption, and also promote nonuniform electrochemical processes like dendrite formation. The co-selection of electrolyte chemistry and initial cycling conditions together are generally considered for the design of desirable interphases. At the same time, the dielectric nature of the electrode material is largely ignored, notwithstanding the high unreliability of the assumption that the nature of the EDL and the mechanism of the interphase formation at metallic and semiconductive electrodes are identical. Here we show that the dielectric nature of the charged electrode greatly affects the interfacial metal-anion-solvent composition; therefore, different interphase chemistry will be formed, suggesting different initial cycling conditions need to be established on a case-by-case basis to form the desired interphase. This phenomenon correlates with the metal ion solvation chemistry and the adsorption of species at the electrified electrode due to the competition of van der Waals and coulombic interactions.

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Broader context

The well-established lithium-ion and the emerging sodium-ion battery technologies are at the forefront of energy storage solutions, but both face durability limitations associated with the irreversible processes at electrode/electrolyte interfaces. To overcome this, an ionically conductive and robust solid electrolyte interphase (SEI) needs to be formed, which is commonly achieved through optimisations of electrolyte chemistry and SEI formation protocols (i.e., initial cycling conditions, with specific current/voltage conditions). At the same time, the role of the electrode material in this process has received significantly less attention, notwithstanding it inherently affects the initial steps of the SEI formation. To address this knowledge gap, our present work explores the effects of the physicochemical properties of the electrode on the mechanism of the SEI formation for the ionic liquid and carbonate-based sodium electrolytes. Using a combination of experimental and theoretical tools, we demonstrate that the structure of the electrolyte-electrode interface and the properties of the SEI are substantially affected by the polarizability of the electrode, and explain these phenomena in the context of the ability of charged electrodes to adsorb electrolyte species. Our work shows that the SEI formation protocols should take all key factors into consideration including the electrolyte composition, cycling conditions, and electrode nature to produce high-performance, robust battery devices.

Lithium-ion batteries (LIBs) are the leading battery technology in the energy storage landscape; however, the demand to further reduce their cost and improve safety, and the growing concern related to lithium resources motivate the accelerated development of sodium-ion batteries (SIBs). An electric double layer (EDL) is an essential part of these batteries as with all electrochemical devices. This structure originates at the solidliquid interface and can extend up to a few nanometres from the electrode surface to the bulk electrolyte phase.^{1,2} Such interfacial behavior originates from different forces, e.g., local

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coulombic order and van der Waals forces. The EDL structure affects capacitive charge storage due to molecular/ionic packing near the electrode,³ as well as the mechanism of heterogeneous charge transfer, the chemistry of the solid electrolyte interphase (SEI) and its morphology in the batteries. 4-7 Both LIBs and SIBs suffer from deficient reversibility of redox processes which is associated with the formation of mechanically unstable and reactive interphases as well as hampered solid-state ion diffusion through these poor interphases. This leads to continuous electrolyte consumption and/or non-uniform metal deposition (i.e., dendrite formation in the case of metal anodes) causing battery failure and safety concerns. Understanding these processes is at the forefront of the development of stable and efficient LIBs/SIBs, 8-12 and requires a comprehensive analysis of the EDL structure and its potential-driven changes in different electrolytes.

Battery electrolyte engineering, e.g., the chemical structure of the solvent and salt, their ratio, presence of additives, etc., 13-16 and the application of the SEI formation protocols with specific current/voltage conditions are widely used to optimize the interphase chemistry and morphology to enable reversible charge transfer. 13,17-22 The structural changes and transport properties within the EDL were also explored with respect to these parameters. For example, the role of applied current/voltage in defining the interface composition and dynamics of ions within the EDL were examined in the case of different ionic liquids and carbonate-based electrolytes. 13,17,22,23

In contrast, the role of the electrode material in the structure and ion transport within the electrified interfaces is often neglected and thus the distinctions in the EDL at different electrodes, e.g., metal or semiconductor, in battery electrolytes are not well understood.

Here, we demonstrate that potential-driven structural changes at the interface with sodium electrolytes based on ionic liquids (ILs) and carbonates as solvents are greatly affected by the dielectric nature of the electrode material. This structural difference is mainly due to the different van der Waals forces between the electrode and the electrolyte, which is significant enough in the case of metallic electrodes promoting anionsolvated sodium clusters near the negatively charged metallic surface depending on the applied potential. This is contrasted by the preference of organic species, i.e., solvent molecules, on the semiconductive electrode. These dissimilarities, therefore, result in different interphase chemistries (anion-derived or solvent-derived), and the requirement for the use of different formation protocols, as discussed below.

Classical molecular dynamics (MD) simulations were conducted to study the difference in EDL structures of IL electrolytes in both metallic Au(111)||Au(111) and semi-metallic graphite (plane)|| graphite (plane) electrode models (Fig. 1a-d). Here, a neat IL, i.e., N-methyl-N-propylpyrrolidinium ([C₃mpyr]⁺) bis(fluorosulfonyl)imide ([FSI]-), and an equimolar mixture of the IL with sodium salt, i.e., 50 mol% Na[FSI] salt with a 50 mol% [C₃mpyr][FSI] IL were investigated. In what follows, the latter composition is also referred as the superconcentrated IL (for details, see the Methods section). As shown in Fig. 1a and Fig. S1 (ESI†), the charging of the neat IL leads to very similar interfacial changes for both

electrodes. Specifically, the calculated number density in Fig. 1a and the number of ions within the innermost layer (0.6 nm from the surface in this case) in Fig. 1c show that both Au and graphite electrode models demonstrate an increase in the interfacial population of the IL cation [C₃mpyr]⁺ (24 to 32 for graphite and 30 to 41 for gold), and a decrease in the number of anions [FSI] (24 to 1.5 for graphite and 27 to 5.4 for gold) when the potential is changed from 0.0 to -2.5 V vs. potential of zero charge (PZC). An exact opposite trend was seen on the positive electrode side. The angular distribution function (Fig. S2, ESI†) analyses the orientation of ions near the electrodes. The orientation of the [C₃mpyr]⁺ ring that is parallel to the negatively charged electrode surface (indicated by two peaks near 0 and 180 degrees) shows a higher probability near the Au(111) surface compared to the semi-metallic graphite (plane). Such observations are expected in the case of non-symmetrical [C₃mpyr]⁺ which appears closer to the Au(111) surface compared to the graphite plane (Fig. 1a), i.e., at a distance of $\sim 3.9 \text{ Å vs. } 4.7 \text{ Å for the nitrogen-atom of } [\text{C}_3\text{mpyr}]^+$, while the orientation of more symmetrical [FSI]- is less affected by the nature of the electrode (Fig. S2, ESI†).

In contrast to the neat IL, the interface of superconcentrated IL is much more susceptible to the dielectric nature of the electrode (Fig. 1b and d and Fig. S3, ESI†). Charging of the metallic Au(111) electrode increases the concentration of the molten-salt-like Na_xFSI_y clusters within the innermost layer (the x:y ratio of these clusters varies with the applied potential) (Fig. 1d), and promotes the migration of [C₃mpyr]⁺ into the bulk electrolyte phase. Importantly, charging of the semi-metallic graphite (plane) surface does not lead to similar interfacial changes with the superconcentrated IL as was seen for the Au(111) case. The composition of this graphite (plane)/superconcentrated IL electrolyte interface reorganizes like in the NaFSI salt-free IL, namely, the increase in the negative surface charge only attracts organic cations [C3mpyr], while the Na_xFSI_v clusters move away from the electrode surface (Fig. 1b and d). Besides, the positive potential on the graphite (plane) electrode accumulates at least twice less Na⁺ in the form of Na_xFSI_v at the interface compared to that on Au(111) (Fig. 1b and d).

Angular distribution analysis of the [C₃mpyr]⁺ and [FSI]⁻ orientations within the innermost layer of the superconcentrated IL shows similar changes to the applied charge as those seen for the neat IL (Fig. S4, ESI†). Also, after charging the electrode, the Na-FSI coordination in the innermost layer of the superconcentrated IL undergoes structural changes, viz. alternations in the ionic bond distance and the number of shared oxygens (Fig. S5, ESI†), which was also discussed in our earlier publications. 13,21,22

The classical MD can employ either surface charge (μC cm⁻²) or electrostatic potential (V vs. PZC) to model the electrified interfaces, which is done by a uniform charge distribution to each of the electrode atoms, i.e. the constant charge method (CCM). Herein, these two methods of charging produced similar outcomes (Fig. S6, ESI†). Also, certain modelling codes, in particular the constant potential method (CPM), enable surface charge fluctuation of the electrode, which might affect the energy barrier for certain ions to reach the surface.24 To estimate such an effect on the accuracy of our MD results with respect to

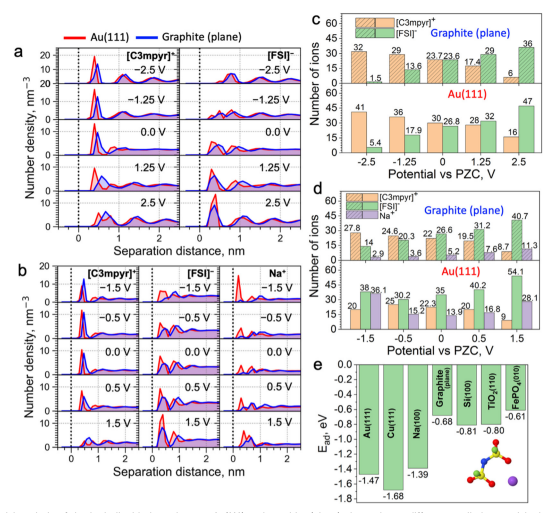


Fig. 1 Interfacial analysis of the ionic liquid electrolytes at Au(111) and graphite (plane) electrodes at different applied potentials through CCM MD simulations. Ion number density profiles of $[C_3mpyr]^+$, $[FSI]^-$, and Na^+ for (a) neat $[C_3mpyr][FSI]$ and (b) $[C_3mpyr][FSI]$ with 50 mol% NaFSI with (red) Au(111) and (blue) graphite (plane) electrodes at different applied potentials vs. PZC. The dashed line represents the position of the electrode surface. The number of different ions in the innermost interfacial layer (within 0.6 nm of the surface) for (c) neat [C3mpyr][FSI] and (d) [C3mpyr][FSI] with 50 mol% NaFSI at different potentials vs. PZC. (d) [C3mpyr][FSI] with 50 mol% NaFSI at different potentials vs. PZC. (e) Na-FSI complex adsorption energy (Ead) on different electrode materials through DFT calculations with Grimme D3 dispersion correction.

Au(111) and graphite (plane) systems, we conducted CPM MD simulations (Note S1, ESI†). It was found that the implementation of the CPM method does not alter the conclusions derived from the CCM MD method (Fig. 1).

Trends discussed above and demonstrated in Fig. 1 can be related to the difference in the adsorption energies of the electrolyte species on different electrodes. For example, as shown in Fig. 1e and Fig. S12, ESI,† the adsorption energy of the Na-FSI complex on different electrodes with explicit metallic nature calculated using the density functional theory (DFT) method is much stronger and almost twice that with semimetallic and semiconductive surfaces. This is caused by the higher electrical conductivity/electronic polarizability of metals forming strong multipole interactions, viz. dispersion forces or van der Waals interactions, with adsorbates (in this case the IL cations and/or salt complexes).25,26 This is also evident from the reported Lennard-Jones (LJ) potential parameters for modelling surfaces of different dielectric nature (Table S1, ESI†).

Indeed, the metal elements, such as Au and Cu, have much higher values of ε (good depth of the LJ potential) than elements like C and Si. This concept was further tested and confirmed in Fig. S13 (ESI†), where weak LJ parameters of the graphite were used for the calculations with the Au(111) electrode and vice versa. This allows for the formation of the Na_xFSI_v aggregate-rich interface for the graphite electrode and an IL cation-rich interface for Au when the same negative surface charge is applied. This also indicates the importance of using accurate LJ parameters to describe surface adsorption phenomena in classical MD.

Those results explain why the negative Au electrode can accommodate the Na_xFSI_v complexes easier than the semi-metallic graphite electrode, and the latter has a higher affinity toward the positively charged IL cation, e.g., $[C_3 \text{mpyr}]^+$ or $[P_{1222}]^+$, compared to neutral or negatively charged Na_xFSI_y clusters. This also explains why an additional energy barrier has been experienced for metal deposition from the IL electrolytes with semiconductive electrodes compared to that with metals. 27,28 Such strong dispersion forces of metals allow negatively-charged Na_xFSI_v complexes to remain near the anode surface, and subsequently reduce the number of [C₃mpyr]⁺ with the increase of Na⁺ in the clusters.

These differences in potential-driven EDL changes between metallic and semiconductive electrodes were also seen for the phosphonium-based superconcentrated IL electrolyte (Fig. S10 and S11, ESI†) and other IL electrolytes with different metal salt concentrations and chemistries. 22,29-32 Thus, such behaviour is likely to be general for the IL electrolytes, as metal cations move in the anion-coordinated clusters having a negative charge up to high salt concentrations.³³ However, we acknowledge that it would be useful to validate the generic nature of the phenomenon for a broader range of IL electrolytes. It is also noted though that the polarity of the IL cation can affect this interfacial phenomenon to some extent.³⁴

To further understand the EDL structure of neat and superconcentrated IL with different electrode materials, we performed differential capacitance measurements by recording electrochemical impedance spectra (EIS) at varying applied potentials within the non-faradaic region. Cyclic voltammetry data for the examined electrolytes, with and without the addition of 50 mol% NaFSI, are shown in Fig. S14 (ESI†). The non-faradaic region is affected by both the dielectric nature of the working electrode and the presence of the NaFSI salt, and this was considered for differential capacitance (C) measurements in Fig. 2. As shown in

Fig. 2a and b, the C-E curve for neat C3mpyrFSI exhibits a descending shape from the positive to the negative potential region, which was also reported for other ILs. 35,36 This phenomenon is likely caused by the difference in the ion size and shape. Specifically, the relatively symmetrical [FSI] pack is tighter per volume at the interface compared to [C₃mpyr]⁺ which results in an increase of charge accumulating at the interface and hence higher capacitance. When comparing neat [C3mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] with different surfaces, the shape of the C-E curve with a metallic gold electrode is significantly affected by the presence of the NaFSI salt (Fig. 2b). In contrast, with a semiconductive glassy carbon (GC) electrode, both neat and superconcentrated IL demonstrate differential capacitance curves of similar shapes (Fig. 2a). The C-E relationships for the GC electrode were recorded over a wider potential region compared to the gold system, which is due to the higher buildup of [C₃mpyr]⁺ at the interface and lesser faradaic contribution from the Na_xFSI_y decomposition around $-1.3 \text{ V} \text{ vs. ferrocene}^{0/+} (\text{Fc}^{0/+})^{28}$ Both neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] electrolytes on GC show higher capacitances than those on the metallic gold electrode, consistent with previous reports.35-37

To obtain a molecular-level understanding of the experimental differential capacitance trends (Fig. 2c and d), the interface for the same pyrrolidinium-based IL electrolytes with idealistic graphite (plane) and Au(111) electrodes was analyzed

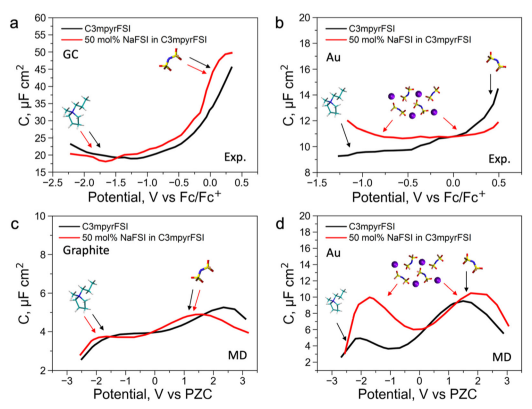


Fig. 2 Experimental and MD simulated differential capacitance for neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] with semiconductive carbon and metallic gold electrodes. (a) and (b) Experimentally obtained differential capacitance (potentials are reported against Fc^{0/+}) for neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C3mpyr][FSI] with (a) glassy carbon (GC) and (b) gold electrodes. (c) and (d) MD obtained differential capacitance (potentials are referred to as PZC) for neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] with (c) graphite (plane) and (d) gold (111) (Au) electrodes. Arrows show ions that will be present in the innermost layer at a certain applied potential.

by means of classical MD (for details, see Methods). As shown in Fig. 2b and c, the MD model represents only an EDL part (C_{EDL}) of experimental differential capacitance which is higher for a gold electrode than that for graphite (plane) regardless of the electrolyte nature. This is due to the tighter ionic packing resulting from greater interactions with metallic surfaces. 38 The higher experimental capacitance of the non-metallic GC electrode compared to the gold system can be explained by splitting the total experimental capacitance (C_{exp}) into two main parts, i.e., the EDL part (C_{EDL}) including Helmholtz/diffuse layer capacitance and a quantum capacitance (C_0) component related to the space charge region within the electrode material (eqn (1)).

$$\frac{1}{C_{\rm exp}} = \frac{1}{C_{\rm EDL}} + \frac{1}{C_{\rm Q}} \tag{1}$$

The $C_{\rm O}$ does not contribute much to recorded $C_{\rm exp}$ with a gold working electrode, as the metallic surface has a high density of states (DOS) near the Fermi level and the addition or removal of an electron does not affect the value of the electrochemical potential. In contrast, depleted DOS of the semiconductive surface creates an extra space charge region within the solid phase of the electrode with a substantial contribution to total experimental $C_{\text{exp.}}^{39}$ This fundamental impact of the quantum component (C_{O}) on the experimental differential capacitance C_{exp} was also seen for other IL-based capacitors with electrodes of different chemistries. 40,41 Besides, the presence of surface functional groups and differences in the morphology for the GC and gold electrodes can also play a role in the total value of experimental capacitance. 42-44 The MD simulated C-E curve for both neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] electrolytes results in a similar shape with the graphite (plane) electrode (Fig. 2c), whereas this is not the case for the Au(111) system (Fig. 2d). The shape of the MD differential capacitance with Au(111) is highly affected by the presence of the NaFSI salt in [C₃mpyr][FSI], which is due to the contribution of Na_xFSI_v to the innermost layer evolving at both positive and negative potential regions, as suggested by the MD studies (see Fig. 1a-d and discussions above). This further suggests that the interface of the superconcentrated IL electrolyte near the charged nonmetallic electrode is similar to that of the Na[FSI]-free IL, whereas for the metallic electrode the molten-salt-like Na_xFSI_v component dominates this region and [C₃mpyr]⁺ moves away to the bulk electrolyte phase at negative potential.

To investigate the mechanism of the interphase formation with the superconcentrated IL electrolytes in the light of these new findings, we conducted several symmetrical cell cycling experiments with 50 mol% NaFSI in [C3mpyr][FSI] and electrodes of different dielectric nature. Each cell was cycled under various current densities (leading to different surface charges) with a fixed depth of charge. This study included symmetrical cells with sodium iron phosphate (NFP) electrodes (non-metallic with low electric conductivity) commonly used as cathodes in sodium-ion batteries, as well as Na and Cu as two distinct metallic systems. Fig. 3a shows Na metal symmetrical cell cycling with a sodium salt superconcentrated IL at different current densities. In this

case, we used higher current density to control the initial cell voltage to mimic the higher negative potential values used in the MD simulations. Fig. 3b shows that the cell cycled at the highest rate used herein exhibits the lowest interphase resistance which correlates with the presence of a favorable SEI, and hence, better overall cycling performance. In contrast, the symmetrical cell cycling with the semiconductive NFP, under a similar protocol, leads to the most resistive interphase when the cell is preconditioned at the highest current density examined (Fig. 3c and d). We note here that the cathodes in these symmetric NFP cells were not fully sodiated and hence the electrochemical processes are expected to include Na insertion/de-insertion during cycling. A symmetric Cu/Cu electrochemical cell was also investigated using the same [Na][FSI] superconcentrated IL electrolyte (Fig. S15, ESI†). The electrochemistry, in this case, is likely to be quite complex but still influenced by the EDL layer and its forming interphase which we will also denote as the SEI. The Cu anode, which is often used to study electrochemical processes in sodium electrolytes, 45,46 was used herein to investigate the chemistry of the formed interphase. The use of the copper electrode facilitates discrimination between the electrode substrate and the formed SEI.

In our earlier studies, 13,21,22,47 we showed that an improved electrochemical behavior of the Na and Li anodes with superconcentrated ILs preconditioned at a high current density originates from the interface dominated by Na_xFSI_v or Li_xFSI_v clusters and the expulsion of the IL cation. This interface enables a uniform and dense deposition morphology and a desirable anion-derived inorganic-rich SEI. The self-heating of the electrode surface at a high rate might also contribute to a dendritefree morphology.¹⁷ In this work, we conducted an X-ray photoelectron spectroscopy (XPS) analysis of other electrode materials, viz. NFP and Cu, which were not previously considered in the context of the interface/interphase relationships.

The XPS data obtained for NFP suggests that low and high current cycling conditions have a different impact on the SEI composition/morphology as compared to those observed for Na and Li. According to the EDL discussion presented above, the SEI layer at more negative potentials on NFP is expected to be dominated by the IL cation and hence a rapid formation cycle could be detrimental to the creation of a low-resistive inorganicrich SEI layer. To investigate this phenomenon, we conducted XPS analysis on the cycled electrodes after different periods of Ar⁺ etching to analyze the chemistry of the deposition products at different depths into the SEI. The Cu surfaces were also analyzed and compared to determine the proportion of organic/ inorganic components in the deposition products after cycling at different rates.

Fig. S16 (ESI†) shows XPS data for neat [C₃mpyr][FSI], NaFSI, and 50 mol% NaFSI in [C3mpyr][FSI]. All recorded spectra confirm the elemental content and their oxidation states in examined compounds, 48 and that partial decomposition of [FSI] anions to NaF takes place upon drying of the NaFSI salt. 49 Keeping the NFP and Cu electrodes in contact with 50 mol% NaFSI in [C3mpyr][FSI] with no potential applied for 24 h results in the formation of polysulfides and some other oxidized species, which are collectively denoted as the initial/native SEI (Fig. S17, ESI†).22 Electrochemical

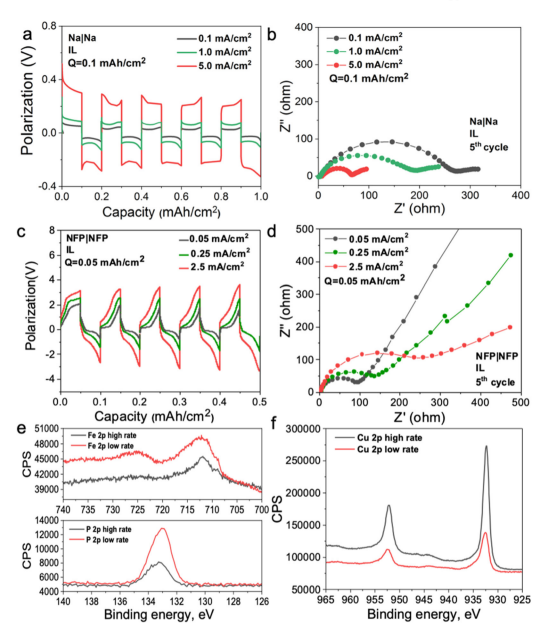


Fig. 3 Electrochemical and X-ray photoelectron spectroscopic characterization of symmetrical cells with different electrodes and 50 mol% NaFSI in the $[C_3 mpyr][FSI]$ electrolyte. (a) Na|Na symmetrical cell cycling with 50 mol% NaFSI in $[C_3 mpyr][FSI]$ at different rates (0.1, 1.0, and 5.0 mA cm⁻² with 0.1 mA h cm⁻² depth of charge) and (b) EIS recorded after the final discharge step. (c) NFP|NFP symmetrical cell cycling with 50 mol% NaFSI in [C₃mpyr][FSI] at different rates (0.05, 0.25, and 2.5 mA cm^{-2} with 0.05 mA h cm $^{-2}$ depth of charge) and (d) EIS recorded after the final discharge step. All electrochemical measurements were conducted at 50 °C. High-resolution (e) Fe 2p, P2p and (f) Cu 2p XPS data recorded for the NFP and Cu anodes after 5 complete cycles with 50 mol% NaFSI in $[C_3$ mpyr][FSI] at low and high current densities (fixed depth of charge). The cycling rates were 0.05 mA cm⁻²/0.05 mA h cm⁻² (low) and 2.5 mA cm⁻²/0.05 mA h cm⁻² (high) for NFP, and 0.05 μ A cm⁻²/0.05 μ Ah cm⁻² (low) and 2.5 μ A cm⁻²/0.05 μ Ah cm⁻² (high) for Cu.

cycling with 50 mol% NaFSI in [C₃mpyr][FSI] at a high rate produces a thicker SEI on the NFP surface as compared to the experiments at a low rate, as concluded from the comparisons of the intensities of the Fe 2p and P 2p spectra recorded for the cycled electrodes (Fig. 3e). The metallic Cu system demonstrates an opposite trend by forming a thinner SEI at the high rather than at the low rate (Fig. 3f). The thick SEI on the NFP surface at the high rate is likely associated with the formation of a porous organic-based film, 50,51 which increases the Na⁺ diffusion pathway, and increases the interfacial resistance with sluggish Na⁺ diffusion kinetics.

To obtain further insights into the composition of the SEI, we performed the XPS analysis of the cycled electrodes that were etched by Ar⁺. The F 1s spectra exhibit peaks characteristic of NaF (684.0 eV) and S-F at (687.4 eV),52 both being the decomposition products from the FSI anion but present at different ratios at the surface and within the SEI of NFP electrodes (Fig. S18, ESI†). Analysis of the Na 1s and O 1s spectra also suggests the presence of Na2O throughout the SEI (Fig. S18, ESI†).⁵² The XPS depth profiles for NFP cycled at the low rate indicate that relative concentrations of C and F decline,

while the contributions of O and Na increase upon increasing the Ar⁺ etching time (Fig. S19 and S20a, ESI†). These observations suggest that the inorganic SEI components Na₂O and NaF dominate the inner layer, while the surface layers majorly contain organic species formed through the electrochemically induced decomposition of the organic $[C_3mpyr]^+$ cations. Interestingly, a different trend was found for the NFP electrodes cycled at a high rate, with progressively increasing contribution of C and decreasing relative concentrations of Na, F and O with the duration of etching (Fig. S19 and S20b, ESI†). Hence, we conclude that inorganic species are major components of the outer layer of the SEI, while organic compounds are concentrated deeper into the SEI. Regarding the chemistry of the SEI on Cu anodes (Fig. S19-S21, ESI†), the low-rate cycling generates higher C % compared to that when the high current density is used. Both these systems show a decreasing trend for the C element, indicative of an organic-inorganic SEI layer on the Cu surface. Additionally, higher amounts of O, F, and S elements at high cycling rate suggest that there are more inorganic species (NaF, Na2O, Na2S, etc.) in the inner layer of the SEI, while there are barely inorganic species decomposed from the Na_xFSI_v clusters for the Cu electrode cycled at a low rate (Fig. S19b and S20e, f, ESI†).

Fig. 4 shows the proposed chemistry of the SEI formed at different current densities with superconcentrated IL electrolytes and electrodes of different dielectric nature. The non-metallic NFP electrode forms an organic-rich SEI near its surface at high cycling rate (Fig. 4a), which is due to the predominance of the IL cation at the interface (Fig. 1b and d). As a result, the [C₃mpyr]⁺ actively participates in the reduction processes and/or blocks Na⁺ intercalation. Both phenomena result in a more resistive interphase and therefore deteriorated performance (Fig. 3c and d). The observed

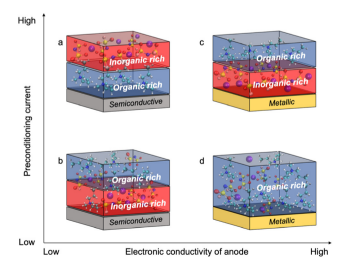


Fig. 4 Schematic relationships between the SEI composition, applied preconditioning current, and dielectric nature of the anode material. The semiconductive electrode cycled with 50 mol% NaFSI in [C3mpyr][FSI] at (a) high and (b) low rates. The metallic electrode cycled with 50 mol% NaFSI in [C₃mpyr][FSI] at (c) high and (d) low rates. The colour codes of atoms are as follows: nitrogen (blue), sulphur (yellow), sodium (purple), carbon (aqua blue), hydrogen (white), oxygen (red), and fluorine (pink).

inorganic-rich layer on the top of the organic-rich [C₃mpyr]⁺ decomposition layer under high current density cycling might come from the subsequent decomposition of the Na_xFSI_v clusters. This is observed in the MD simulations in the case of the semiconductive electrode where anion-solvated Na⁺ are present in the outer layer adjacent to [C₃mpyr]⁺. Meanwhile, the inorganic-rich film near the surface of the NFP forms during low current density cycling (Fig. 4b), which in turn leads to decreased interphase resistance (Fig. 3c and d).

The Cu anode with the superconcentrated IL electrolyte forms an inorganic-rich SEI near the surface when high current density cycling is applied (Fig. 4c), whereas the low-rate cycling leads to the organic-rich SEI (Fig. 4d). This behaviour is similar to that observed earlier for the Na and Li electrodes. 13,47 These observations correlate with the difference in the structure of the electrode/electrolyte interface for metallic and semiconductive surfaces with superconcentrated ILs discussed above.

To confirm that the interfacial resistance for the examined cells is affected by the EDL structuring near the electrode rather than its state of charge, we further expanded our understanding to the mono-cationic carbonate electrolyte, i.e., 1.0 M NaPF₆ electrolyte in ethylene carbonate (EC) and dimethyl carbonate (DMC) (1:1 vol.). In this case, Na is surrounded by a neutral solvent⁵³ which does not require a strong dispersion force for its retention at the electrified interface; hence, the formation protocols might be different from the ionic liquid electrolytes. Fig. 5a and b show Na symmetrical cell cycling with this electrolyte, where the cell cycled at a higher current density demonstrates lower interfacial resistance. A similar observation was reported for a Li symmetrical cell cycled with 1.2 M LiPF₆ in the same solvent system, where a high cell voltage was used instead of a high current density to control the reduction and oxidation rates during cycling. 19 Fig. 5c and d show that the interfacial resistance of the NFP symmetrical cells with the carbonate-based electrolyte decreases after cycling at a high current density, which is opposite to the case observed with the IL electrolyte (Fig. 3d and c). This also suggests that changes in the interfacial resistance of the NFP electrodes correlate with the EDL structure, and do not solely depend on the state of charge of the NFP, i.e., how much Na⁰ is in it. It was also shown that the use of high-rate formation cycling for the semiconductive full cell (graphite||NMC) and metallic symmetrical cell (Li||Li) with the same electrolyte leads to lower interfacial resistance and stable cycling behavior. 18-20

To interpret this phenomenon at a molecular level, the structure of electrified interfaces of Au(111) and graphite (plane) electrodes with 1 M NaPF₆ in EC: DMC (1:1 vol.) was investigated using MD simulations (extended discussions are provided in Note S2, ESI†). Two different electrode configurations were modelled, namely, Au(111)||Au(111) and graphite(basal plane)|| graphite(basal plane) (Fig. S23-S25, ESI†). In contrast to ILs, at the high negative/positive electrode charging, we can observe the increased Na⁺ and PF₆⁻ in the innermost layer near both Au(111) and graphite(basal plane) electrodes (Fig. 5e and f), and the most polar and strongly Na+ coordinated solvent molecule (EC in this case) saturate the majority of the electrified interface. The Au(111)

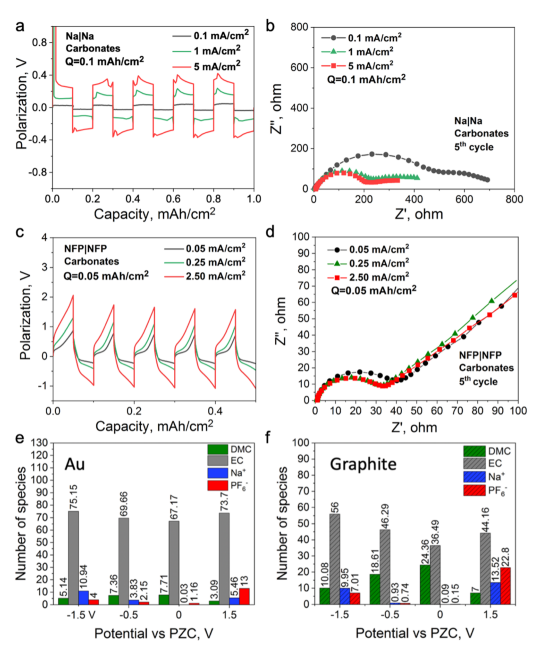


Fig. 5 Electrochemical and MD simulation data for the 1.0 M NaPF₆ in an EC: DMC (1:1 vol.) system with different electrodes and preconditioning. (a) Na|Na symmetrical cell cycling at different rates (0.1, 1.0, and 5.0 mA cm⁻² with 0.1 mA h cm⁻² depth of charge) and (b) recorded EIS after thefinal discharge step. (c) NFP|NFP symmetrical cell cycling at different rates (0.05, 0.25, and 2.5 mA cm⁻² with 0.05 mA h cm⁻² depth of charge) and (d) EIS after the final discharge step. All electrochemical tests were conducted at 30 °C. The number of interfacial species within 0.6 nm from (e) Au(111) and (f) graphite (plane) at different applied potentials derived from MD simulations

plane has a stronger affinity towards this highly polar EC solvent than the graphite (plane) electrode. Such a distinction compared with the IL system arises from the absence of a competing cationic species near the negative electrode, viz., the IL cation. This explains why both metallic and semiconductive electrodes form favourable SEI chemistry/morphology with the carbonate electrolyte after high current/voltage formation cycling. 18-20 However, apart from the optimum SEI chemistry/composition such aspects as optimum cell capacity and its retention should be also considered to design optimal formation protocols for batteries with different electrode and electrolyte materials.54 Moreover, the current densities and areal capacities used in this work should not be treated as the ideal cycling conditions for a given device. The electrode formation protocols are complex and will surely depend on the cell configuration, electrolyte composition, temperature, etc. Nevertheless, our data demonstrates that the formation conditions are at the very least dependent on the nature of the electrode and additional case studies are still required to optimize the performance of a given device.

In summary, the results presented herein suggest that the mechanism of interphase formation on battery electrodes,

especially with IL electrolytes, is significantly affected by the dielectric nature of the electrode material, which can cause different electrode/electrolyte interface structuring in response to different applied electrode charges. For achieving stable and low-resistance charge transfer, the interphase that is derived predominantly from the anion decomposition products appears favourable as these are likely to be insoluble, electronically insulating and ionically conductive.

The composition of the electrode/electrolyte interface depends on interfacial van der Waals and coulombic interactions, e.g., the charged semiconductive electrode is less likely to retain highly polar molecules or ions of the same charge near its surface owing to their low electronic polarizability and weak dispersion forces. This is especially important in the case of IL electrolytes because organic IL cations and anion-coordinated Li⁺/Na⁺ complexes compete near a negatively charged surface. The formation of an anion-derived interphase with IL electrolytes on a semiconductive electrode can occur at a low current density. In contrast, a higher current density leads to the formation of a thick organicrich interphase with high resistance. This is due to the increase the number of IL cations near the surface of the semiconductive electrode with an increase in the applied negative charging.

High negatively charged metallic electrodes can retain anions near their surface. This is due to the high electronic polarizability and strong dispersion forces in the case of metallic surfaces which require a high current density to enable the anionderived interphase to form a molten-salt-like rich interface.

In the case of carbonates electrolytes which only have alkali metal cations, the application of high current, i.e., the rapid formation cycling, for both metallic and semiconductive electrodes leads to a similar trend for the system investigated herein. Namely, this method enhances the formation of an anion-derived interphase from increased Na_x(PF₆)_v clusters attracted to the negative electrode. At the same time, the electrode nature affects the interfacial concentration of a specific solvent depending on its polarity, and this could be used for controlled SEI formation from the selected species to, especially, if only certain organic molecules are fluorinated. In this regard, the rational design of batteries requires not only the consideration of new electrolyte chemistries, the optimization of their transport properties and interphase formation protocols, but also the role of the dielectric nature of the electrode material. These factors should all be considered when designing electrode formation protocols for reversible charge transfer with stable/efficient interphases. These findings open new prospects for future implications in the field of electrochemical systems for controlled electrocatalysis, rare-metal recovery, and the design of EDL capacitors, batteries, etc.

Methods

Molecular dynamics simulations

All-atom classic molecular dynamic simulations were conducted with the GROMACS software package (version 2022.2).⁵⁵ The Canongia Lopes-Padua (CL&P) force field parameters were used

in this work for [C₃mpyr][FSI] and [P₁₂₂₂][FSI] ionic liquids, which were previously studied for bulk-phase investigation.^{56,57} The Lennard-Jones potential parameters (($\varepsilon = 22.1333 \text{ kJ mol}^{-1}$ and $\sigma = 0.2629 \text{ nm}$) and ($\varepsilon = 0.2305 \text{ kJ mol}^{-1}$ and $\sigma = 0.3412 \text{ nm}$)) for Au and graphite electrodes were adopted to describe van der Waals interaction at the interface.^{58,59} The bulk phase densities of [P₁₂₂₂][FSI] and 50 mol% NaFSI in [P₁₂₂₂][FSI] were calculated first and the computational details are described in the literature. 13 The simulation density errors for neat [P1222] [FSI] and 50 mol% NaFSI in $[P_{1222}][FSI]$ with 0.7 scaling factor were -4.43% and -1.13%at 333 K compared to experimentally obtained values of 1.2873 kg m⁻³ and 1.5232 kg m⁻³, respectively. The density errors for neat [C₃mpyr][FSI] and 50 mol% NaFSI in [C₃mpyr][FSI] with the same 0.7 charge scale factor are mentioned in the literature. 13 The interface models were constructed with the IL electrolytes confined between two Au(111) face-centered cubic or two graphite (plane) electrodes (Fig. S27, ESI†). The simulation box consisting of two electrodes and an electrolyte has the same the x and ydimension as that the of gold or graphite electrode, and the zlength was calculated based on the bulk phase density of electrolytes. An extra vacuum space twice the length of electrode separation was also introduced outside one side of the electrode to eliminate the artefact images resulting from the simulation in the slab geometry when applying periodical boundary conditions along the z-direction.⁶⁰ An annealing process was simulated at first upon changing the temperature from 393 K to 700 K and then to 393 K again before a long equilibration calculation at 393 K for a total of 15 ns using an NVT ensemble and the Nose-Hoover thermostat. The annealing procedure allows the system to gain sufficient dynamics to obtain a reasonable initial structure for equilibration calculation. Both energies and pressures were checked to determine whether the system reaches equilibrium. An additional 40 ns production run was performed at 393 K for structural and dynamics analyses. The trajectory file was written every 2 ps. The constant charge method (CCM) with a uniform distribution of point charges across the top layer of the electrode surface was used thoroughly unless it is specified somewhere else. The electrode potential was determined according to the Poisson equation,61 by calculating the electric double layer (EDL) potential drop between two electrodes $U_{\rm EDL}$ = $\psi_{\rm electrode} - \psi_{\rm bulk}$. The $U_{\rm EDL}$ value of the uncharged electrode was defined as the potential of zero charge (PZC). The electrode potential vs. PZC is defined as $U_{\rm EDL} = \psi_{\rm electrode} - \psi_{\rm bulk} - {\rm PZC}$. The surface charges used to achieve the electrode potential are given in Fig. S28 (ESI†). The MD differential capacitance (C_{diff}) was calculated by taking the derivative of the first order from the plot in Fig. S28 (ESI†), namely, $C_{\rm diff} = \frac{{
m d}\sigma}{{
m d}V}$. The σ is the surface charge of the electrode in μ C cm⁻² and V is the potential vs. PZC. The DC curve was obtained through the global fitting of the $C_{\text{diff}}V$ data points to a polynomic function of the fifth order, as described in the literature. 62 The customized Python code was used for trajectory analysis unless the required tool was available in the GROMACS package. 55,63

Density functional theory calculations

DFT calculations have been conducted to calculate the adsorption energy using the Gaussian Plane Wave scheme based on

CP2K version 6.1.0⁶⁴ with the Perdew-Burke-Ernzerhof (PBE) functional,65 TZVP MOLOPT basis set for all atoms,66 and Goedecker-Teter-Hutter pseudopotentials.⁶⁷ The orbital transform method was used with the full kinetic predictioner and conjugated gradient as a minimizer. The DFT-D3 Grimme dispersion correction with the PBE exchange correlation function was used for all calculations.⁶⁸ The SCF convergence factor was set up at 10^{-6} for geometry optimization and at 10^{-8} for the following scf energy calculations. The NaFSI adsorption energy on the slab of different electrode materials was calculated in a 2D periodic setup with z dimension 2.5 times longer than that of the largest x or y. Poisson solver Analytic and surface dipole correction along the z axis were applied to avoid improper treatment of periodicity.⁶⁹ The coordinate positions and cell dimensions of all four layers of slabs were initially optimized, and then, further fixed for geometry optimization with an absorbed NaFSI complex. All calculations have been done at the Γ point due to the large size of supercells resulting in a small Brillouin zone.

Materials

Na metal was purchased from Sigma-Aldrich. Sodium iron phosphate (NaFePO₄ or NFP) was synthesized and characterized in CIC energiGUNE as previously reported.⁷⁰ The electrode was prepared by mixing NFP, carbon black and the polyvinylidene fluoride (PVDF) binder at a weight ratio of 8:1:1 into the Nmethyl-2-pyrrolidone (NMP) solvent and cast onto an Al foil. Cu was purchased from Gelon LIB Co., Ltd. All the electrodes were punched into 8 mm diameter disks. The NFP mass loading was 2.0 mg cm⁻². Sodium bis(fluorosulfonyl)imide (NaFSI, 99.7%) and N-propyl-N-methylpyrrolidinium bis(fluorosulfonyl)imide ([C₃mpyr][FSI], 99.9%) were obtained from Solvionic Corporation. They were dried under vacuum using a Schlenk line at 50 °C before any electrolyte preparations. The superconcentrated ILs were prepared by adding 50 mol% NaFSI into [C3mpyr][FSI]. The water content of IL electrolytes used for battery cycling was tested by means of an 831 Karl Fisher coulometer with Hydranal Coulomat AG as the titrant and does not exceed 50 ppm. The procedure was conducted under an argon-filled glovebox with water and oxygen level <0.1 ppm. The organic ester electrolyte (1.0 M NaPF₆ in EC/DEC 1:1 by volume) was purchased from Kishida Chemical Co., Ltd.

Three-electrode setup measurements

Experimental differential capacitance curves were obtained by means of electrochemical impedance spectroscopy (EIS) in a three-electrode setup, namely, the working electrode was either a gold or glassy carbon electrode (BAS inc., Japan) with a surface area of 0.02 cm⁻², platinum wire as a counter electrode, and the Ag⁰/Ag⁺ reference electrode containing 10 mM silver triflate in [C₃mpyr][FSI] (eDAQ Pty Ltd, Australia). The reference electrodes were calibrated with the ferrocene redox couple (Sigma-Aldrich), and the potential reference scale at 0.0 V vs. Ag^0/Ag^+ was -0.361 V vs. Fc^0/Fc^+ for $[C_3mpyr][FSI]$. Prior to measurements, the working electrode was cleaned according to the method described previously.²⁹ The counter Pt wire was sonicated for 15 minutes with ethanol/deionized water, and dried under the same conditions as a working electrode with Ar flow initially and later in a 70 °C oven, and then Pt wire was annealed with a butane torch. The setup was equilibrated for 30 minutes at 50 °C before performing any electrochemical measurements. Cycling voltammetry tests were performed with a scan rate of 25 mV $\rm s^{-1}$ to define a non-faradaic region for all the following DC measurements.

Capacitance is extracted within a potential region of +0.70--1.85 V vs. Ag^0/Ag^+ for the GC electrode and of +0.85--1.85 Vvs. Ag⁰/Ag⁺ for the gold electrode with a step of 50 mV, and a chronoamperometry of 30 s was applied to reorganize the double layer at the fixed potential. EIS spectra were collected from a Biologic VSP potentiostat using the PEIS test from 400 kHz to 1 Hz with an amplitude of 10 mV. A relaxation step of 8 min was applied prior to switching to another potential. The capacitance was extracted by fitting the PEIS data to a circuit model with a constant phase element (CPE) in Fig. S29 (, ESI†). High frequencies (f), i.e., > 10 kHz, were ignored in the fitting as they are likely associated with the inductance/capacitance of the electrode connections.

The experimental capacitance value was extracted using $C = \frac{Q\omega_i^{q-1}}{\sin\frac{q\pi}{2}}$, where C is a differential capacitance (F); Q is a

component of CPE $(F \cdot s^{(a-1)})$; ω is the angular frequency (Hz), and $\omega = 2\pi f$; and a is a phase constant. All electrochemical measurements should be repeated several times to ensure that the observed trend is reproducible. The frequency (f) of 40 Hz related to capacitive behaviour (close to the 90-degree phase angle) is used for all calculations (Fig. S30, ESI†). The phase constants (a) calculated from the fitting of EIS data were ≥ 0.938 for the gold electrode or ≥ 0.824 for the glassy carbon electrode. The potential of zero charge (PZC) for superconcentrated [C₃mpy][FSI]-based electrolytes was previously measured by the electrode immersion method with the gold electrode as approximately -0.35 V vs. Fc/Fc^{+} .²² It should be recognised though that the direct conversion of $V \nu s$. PZC to $V \nu s$. Fc/Fc^+ within the theory of classical MD is not accurate as the calculated potential drop does not include the complete polarizability of the system. All measurements were conducted at 50 °C in an argon-filled glove box with H_2O and O_2 levels < 0.1 ppm.

Symmetrical cell cycling

R2032 symmetrical cells were assembled by putting two identical electrodes (Na||Na, NFP||NFP, and Cu||Cu) with 60 mL of superconcentrated ionic liquid electrolyte (50 mol% NaFSI in [C₃mpyr][FSI]) or an organic electrolyte (1.0 M NaPF₆ in EC/ DEC). The Solupor® polyethylene (19 mm in diameter) was used as a separator in all cells. The cells were then rested for 24 h before the charge/discharge test under 50 °C for IL and under 30 °C for carbonates. The areal capacity used for NFP||NFP is $0.05 \text{ mA h cm}^{-2}$ with 0.05, 0.25 and 2.5 mA cm⁻² and for Cu||Cu it is 0.05 μ Ah cm⁻² with 0.05, 0.25 and 2.5 μ A cm⁻² current densities. This low current for the Cu||Cu setup was chosen to adjust an equal amount of consumed Na and enable the same capacity per

cycle for each condition. The areal capacity used in Na||Na cell is 0.1 mA h cm^{-2} with 0.1, 1.0 and 5.0 mA cm $^{-2}$ current densities. EIS data for interfacial resistance was collected from the Biologic VSP potentiostat from 1 MHz to 100 MHz with 10 mV potential perturbation. EIS spectra were recorded after five cycles, and the electrodes were then taken from the disassembled cells and sent for XPS tests.

X-Ray photoelectron spectroscopic characterization

X-Ray photoelectron spectroscopy (XPS) analysis was performed using a Nexsa Surface Analysis System from ThermoFisher Scientific with a monochromatic Al Ka source (1486.6 eV). The X-ray spot size was set to 400 µm. The analysis chamber was maintained at a pressure of 1.0×10^{-8} bar or less. Survey scans were recorded at a pass energy of 200 eV and a step size of 1 eV, while high resolution data were obtained at a pass energy of 50 eV and a step size of 0.1 eV. Samples were loaded onto the holder inside the Ar-filled glovebox and left under vacuum in the glovebox antechamber for 10 min before being transported to the instrument without coming into contact with the ambient environment at any stage. The samples were kept in an ultra-high vacuum overnight before XPS measurements were carried out. No electrical contact between the sample and the instrument ground was present, and the samples were charge neutralised before the analysis. Collected spectral data were energy corrected by adjusting the maximum of the aliphatic C-C peak in C 1s spectra to 284.8 eV. When required, Ar⁺ etching was performed in a monoatomic regime at a pass energy of 420 eV at an estimated etching rate of 0.08 nm s⁻¹.

Author contributions

M. F. proposed the research. D. A. R. designed experiments, conducted MD and DFT modelling studies and three-electrode setup measurements including data analysis, interpretation, manuscript drafting and editing. J. S. conducted symmetrical cell cycling. P. V. C. conducted XPS measurements. All authors contributed to the discussion of the results, writing, and editing of the final manuscript.

Conflicts of interest

The authors declare no competing financial interest.

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