

Mitigating the Effect of High Overpotential during Al Deposition on Aluminium-Graphite Battery Performance

Charan Mukundan^[a] and Jean-Francois Drillet*^[a]

This study investigates the impact of current density on electrode potential during aluminium (Al) dissolution/deposition step from/on an Al foil as well as the charge-discharge behaviour of aluminium-graphite batteries (AGB) in various AlCl_3 -based electrolytes. Preliminary experiments in a cell with graphite blocking electrodes evidenced higher chemical stability of 1:1.5 Urea: AlCl_3 electrolyte, followed by 1:1.5 TEA: AlCl_3 and 1:1.5 EMIMCl: AlCl_3 . In Al–Al symmetric cells, current densities above 1 mA cm^{-2} led to a notable rise in overpotential up to 100 mV during Al deposition in both TEA: AlCl_3 and Urea: AlCl_3 electrolytes mostly due to low surface area of native Al foil. Similar trend was observed in AGB full cells, where higher overpotentials during Al deposition caused ‘incomplete’ AlCl_4^- intercalation in natural graphite (NG), resulting in

capacity fade at current densities in the range between 0.5 and 5 Ag^{-1} . By adjusting the upper cut-off voltage (UCV) during charging step as a function of applied current value according to respective electrolyte stability, a significant improvement in specific capacity and energy density was achieved during charging and discharging steps. For instance at 1 Ag^{-1} , the specific energy density of AGB increased by 10% in EMIMCl: AlCl_3 , 48% in TEA: AlCl_3 , and an impressive 250% in Urea: AlCl_3 . During long-term cycling post-UCV adjustment, the capacities of AGB increased by 10%, 13%, and 27% for AGBs with EMIMCl: AlCl_3 , TEA: AlCl_3 and Urea: AlCl_3 , respectively with a negligible capacity fade of less than 1% for EMIMCl: AlCl_3 and TEA: AlCl_3 , and a 9% capacity fade for Urea: AlCl_3 .

1. Introduction

The global focus on reducing reliance on fossil fuels has led to an increased demand for renewable energy sources like solar, wind, and hydropower. However, a major challenge is related to the huge logistic required for large-scale energy storage and distribution.^[1,2] Rechargeable batteries meeting criteria such as high capacity, affordability, stability, and durability emerge as one of viable solutions for storing energy from fluctuating and unpredictable, renewable sources.^[3] Over the past few decades, the lithium-ion battery (LIB) market has grown significantly to address the rising need for energy storage in various applications, including consumer electronics, on-grid systems, and transportation.^[4] Yet, challenges such as limited resources and escalating costs of lithium and cobalt have prompted exploration into battery technologies based on more abundant metals like sodium, aluminium, zinc, magnesium, potassium, and calcium.^[5] Among these, aluminium stands out due to its high theoretical gravimetric energy (2.98 Ah g^{-1}), nearly 80% of that of lithium (3.86 Ah g^{-1}) due to its higher gram-equivalent weight (GEW) (8.97 vs. 6.94 g/eq for Li). The GEW is calculated

from ratio of molecular/atomic weight divided by the number of electrons involved in the reaction assuming that 1 gram of equivalent weight of active material can theoretically deliver 26.8 Ah .^[6] The lower the GEW value the higher the specific capacity. Due to the higher density of Al material (2.69 vs. 0.54 g cm^{-3} for Li), theoretical value of volumetric capacity of Al amounts to 8.04 vs. 2.06 Ah cm^{-3} for metallic Li. Moreover, Aluminium-graphite batteries (AGBs) rely on low-cost, abundant, non-flammable, and easily recyclable materials such as aluminium for the positive electrode and graphite for the negative electrode. It should be noted that the capacity of AGBs is primarily limited by the graphite material, if there is an excess of electrolyte, particularly with sufficient concentrations of Al_2Cl_7^- ions.^[7,8] However, AGB technology faces some challenging issues related to Al dendrite growth,^[9] Al passivation,^[10,11] and cost of EMIMCl: AlCl_3 chloroaluminate electrolyte^[12] as well as relatively low gravimetric and volumetric capacity at cell level.^[13]

A promising way to mitigate Al dendrite growth is to optimise the surface of the Aluminium electrode.^[9,14,15] Due to their exceptional chemical stability and high porosity ($> 80\%$), thick glass fiber (GF) separators, ranging from 260 to $650 \mu\text{m}$ in thickness, are commonly used in most published studies. However, this design promotes aluminium growth from the Al anode toward the cathode, leading to short circuits and their thickness strongly affects the volumetric cell density.^[16] Therefore, replacing the traditional GF separator with a less porous separator/membrane that functions as both dendrite barrier and electrolyte reservoir is crucial for improving AGB technology.^[12] In our recent work, we demonstrated for the first time that a polyether sulfone membrane (PES) of $160 \mu\text{m}$ in thickness can be a very promising separator for AGBs.^[17] To

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Supporting information for this article is available on the WWW under
<https://doi.org/10.1002/batt.202400718>

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prevent the passivation of the Al electrode, highly pure aprotic electrolytes such as EMIMCl:AlCl₃ chloroaluminate are required.^[18] While the EMIMCl:AlCl₃ ionic liquid mixture performs very well, there is considerable interest in exploring the feasibility of less corrosive and cheaper AlCl₃-based mixtures such as triethylamine hydrochloride (TEA),^[19,20] trimethylamine hydrochloride (TMA),^[21] urea,^[22,23] caprolactam,^[24] and acetamide.^[25,26]

The specific capacity of AGBs is largely determined by the nature and loading of the graphitic material, which must accommodate the considerably large AlCl₄⁻ anions (5.8 Å),^[27] in contrast to the much smaller Li⁺ ions (0.59 Å),^[2] as well as the constraints imposed by the electrolyte stability window (ESW).^[18] According to density functional theory (DFT) calculations, a 6x6 graphene sheet composed of 72 carbon atoms can host up to four AlCl₄⁻ intercalants, which corresponds to 18 carbon atoms per AlCl₄⁻.^[28] This arrangement allows for a maximum theoretical capacity of 124 mAhg⁻¹ when achieving full intercalation even up to *stage-1*, where each graphene layer is intercalated with AlCl₄⁻ ions to form graphite intercalated compounds (GIC). For comparison, Li⁺ intercalation up to *stage-1* can yield a maximum theoretical capacity of 372 mAhg⁻¹ by forming LiC₆ GICs.^[29] Due to the limited stability window of the EMIMCl:AlCl₃ electrolyte of 2.45 V vs Al/Al³⁺, the graphite cathode typically only reaches up to *stage-4* intercalation.^[30] Moreover first-principles calculations show that the large size of the AlCl₄⁻ anion (~5.28 Å)^[31] might significantly increase the graphene layer interlayer distance up to 8.8 Å.^[28,32] Surprisingly, experimental analysis indicates an increase in gallery height up to "only" 5.7 Å for AlCl₄-GIC, representing a 160% expansion compared to the pristine graphite gallery height of 3.5 Å.^[27]

More generally, graphite with higher crystallinity, as evidenced by its nearly perfect interplanar d-spacing of 0.3354 nm at room temperature and the low intensity of the D-band in Raman spectra, demonstrates superior charge-discharge performance in AGBs.^[33] Wang et al. showed that a cell with large flakes of natural graphite with I_d/I_g ratio of 0.09 and interplanar d-spacing of 0.3356 showed a capacity of 100 mAhg⁻¹ compared to only 75 mAhg⁻¹ for small graphite flakes with I_d/I_g ratio of 0.27 and interplanar d-spacing of 0.3371 nm.^[34] Further graphitic materials including graphene,^[35-38] kish graphite,^[34] and expanded graphite^[39-41] have been evaluated, whereas cost-effective, commercially available expanded graphite synthesised by chemical oxidation or high temperature exfoliation methods appears to be more suitable for industrial scaling of AGBs.^[41,42]

Another significant issue in AGB is the higher overpotential observed at the Al foil anode during aluminium electrodeposition step (charging) of AGBs at high current densities (> 1 Ag⁻¹). Various approaches for optimizing anode morphology have been successfully implemented, such as chemical treatment of the aluminium foil surface,^[9,43] coating with aluminium powder,^[44] removal of oxide layers through electropolishing,^[45,46] or using graphite layer as a substrate for aluminium deposition.^[16,47] In addition to the optimisation of Aluminium anode surface, electrolyte nature plays a major role. In a previous work, we reported on AGBs using a TEA-AlCl₃ electro-

lyte, where the capacity increased from 60 to 90 mAhg⁻¹ during long-term cycling at 1 Ag⁻¹.^[19] This improvement was primarily due to a continuous reduction in the aluminium anode overpotential with increasing cycles, enabling 'complete' charging of the graphite cathode up to 2.37 V vs Al/Al³⁺, and consequently, an increase in capacity over time. This study aims to explore the influence of both current density ranging from 0.1 to 5 Ag⁻¹ and electrolyte nature (EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃) on anode and cathode overpotential behaviour. By adjusting the upper cut-off voltage (UCV) during the charging step, an impressive increase in AGB cell performance in terms of specific capacity especially in TEA:AlCl₃ electrolyte was achieved.

2. Results and Discussion

2.1. Physical Characterisation of Pristine Natural Graphite Powder

The FESEM images in Figure 1a reveals that the natural graphite (NG) used in this study has a potato shaped morphology with an average particle size of 17 μm. The Raman spectra of the NG sample in Figure 1b shows characteristic peaks of D, G and 2D bands at around 1360, 1575, and 2710 cm⁻¹, respectively with an I_D/I_G ratio of 0.106 indicating a high graphitization degree of the material.

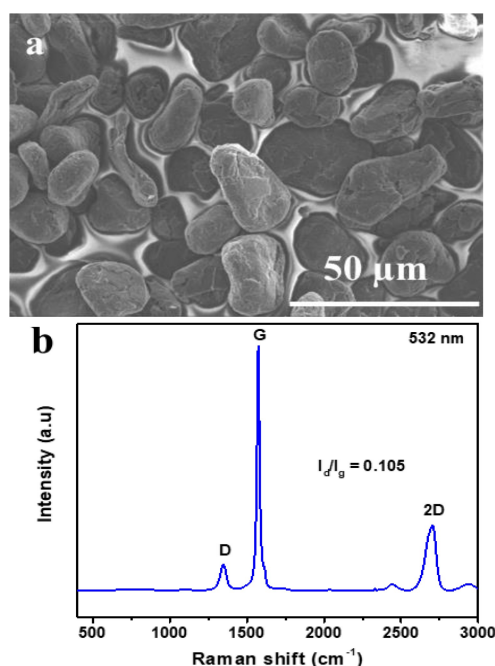


Figure 1. (a) SEM image, and (b) Raman spectra of the natural graphite.

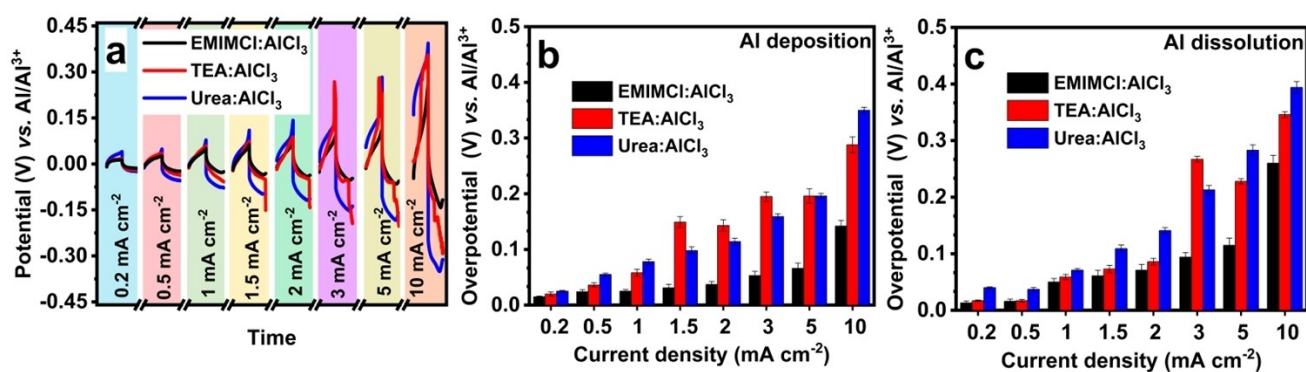


Figure 2. (a) Voltage vs time curve of 10th cycle of Al dissolution-deposition process, and average overpotential value at the end of (b) deposition and (c) dissolution step at different current densities in Al–Al symmetric cell.

2.2. Influence of the Electrolyte Composition on the Al Deposition–Dissolution Process

The influence of electrolyte nature (EMIMCl:AlCl₃, TEA:AlCl₃ and Urea:AlCl₃) on aluminium deposition and dissolution reactions was investigated in an Al–Al symmetric cell at current densities ranging from 0.2 to 10 mA cm⁻² as tabulated in table S1, with a fixed absolute capacity of 0.159 mAh (equivalent to 0.2 mAh cm⁻²). Figure 2a–c presents the whole voltage curve profiles during anodic (Al dissolution) and cathodic (Al deposition) scan at the 10th cycle step and the mean overpotential values of the vertex potential as bar diagram for the 10 cycles in the Al–Al symmetrical cell. Figure 2a, clearly shows that the best performance in terms of overpotential for both Al deposition and stripping was consistently achieved with EMIMCl:AlCl₃, followed by TEA:AlCl₃ and Urea:AlCl₃. Within Al deposition range of 1–3 mA cm⁻², however, overpotential values were about 30–50 mV higher in TEA:AlCl₃ than in Urea:AlCl₃. In summary as shown in Figure 2b&c, the overpotential values for both Al reactions increased on average from 25–30 mV at 0.2 mA cm⁻² up to 140–360 mV at 10 mA cm⁻². The overpotential of Al deposition reaction in TEA:AlCl₃ sharply increased up to 148, 195 and 288 mV at 1.5, 3 and 10 mA cm⁻², respectively. This rise can be attributed to the higher viscosity and lower ionic conductivity of the TEA:AlCl₃ electrolyte ($\eta=22.3$ mPa s and $\sigma=9.50$ mS cm⁻¹ at 20 °C)^[48] compared to EMIMCl:AlCl₃ ($\eta=15.7$ mPa s and $\sigma=16.35$ mS cm⁻¹ at 25 °C)^[48,49] In Urea:AlCl₃, the overpotential during aluminium deposition ranged from 25 to 98 mV at current densities of 0.2 to 1.5 mA cm⁻². However, at higher current densities between 2 and 10 mA cm⁻², the overpotential jumped from 141 up to 394 mV. The overpotential during aluminium dissolution followed a similar trend, as illustrated in Figure 2c. The significant variation in overpotential during Al dissolution/deposition process, particularly at elevated current densities, can be attributed to mass transport limitation due to high viscosity and low ionic conductivity of 1:1.5 Urea:AlCl₃ ($\eta=218$ mPa.s and $\sigma=1.6$ mS cm⁻¹ at 20 °C).^[50]

2.3. Influence of Electrolyte Nature and Electrochemical Stability on AlCl₄⁻ Intercalation/Deintercalation Behaviour in NG

The cyclic voltammograms (CV) of NG at a scan rate of 1 mV s⁻¹ (Figure 3a) showed significant differences in the anodic and cathodic peaks between ionic liquids (ILs) EMIMCl:AlCl₃ and TEA:AlCl₃ and deep eutectic solvent (DES) Urea:AlCl₃. In EMIMCl:AlCl₃ and TEA:AlCl₃, three major oxidation peaks during positive scan suggest multiple stages or pathways for AlCl₄⁻ anion intercalation into graphite matrix.^[30] Comparing the CV to

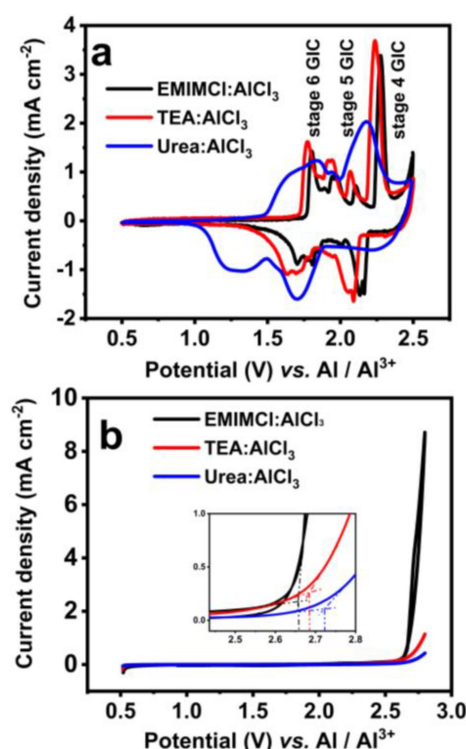


Figure 3. a) CVs obtained in a three-electrode configuration using natural graphite as working electrode in different electrolytes within a potential window of 0.5–2.5 V at a scan rate of 1 mV s⁻¹, b) Electrolyte stability window with glassy carbon rod as working and counter blocking electrode within a potential window of 0.5–2.8 V at a scan rate of 1 mV s⁻¹.

the results reported by Pan et al.,^[30] the oxidation peak between 1.7 to 2.0 V is associated to the intercalation of AlCl_4^- anions every sixth graphene layer, noted as *stage-6* intercalation. The subsequent peak from 2.0 to 2.1 V indicated *stage-5* intercalation, while distinct and predominant peak at 2.2 V (2.1–2.37 V) corresponds to *stage-4* intercalation process.^[30,51] Corresponding reduction peaks between 1.25–1.7 V and 2.25–1.75 V underscore the reversibility of AlCl_4^- intercalation processes. Although the redox peaks in EMIMCl: AlCl_3 and TEA: AlCl_3 are similar, the slight difference in offset potential (30 mV) of AlCl_4^- intercalation in TEA: AlCl_3 may be indicative of more favourable kinetics compare to that in EMIMCl: AlCl_3 . Conversely, in the Urea: AlCl_3 mixture, oxidation peaks shifted to lower potentials compared to those observed in ILs (EMIMCl: AlCl_3 and TEA: AlCl_3). The shift may be attributed to the differences in kinetics that arise from interactions between AlCl_4^- and the $[\text{AlCl}_2(\text{ligand})_2]^+$ species in Urea: AlCl_3 and is different from $\text{AlCl}_4^-/\text{Al}_2\text{Cl}_7^-$ interactions in ILs.^[25,50,52]

The presence of two oxidation peaks in the 1.4–2 V range and another between 2–2.27 V, alongside two reversible peaks in 1.85–1.5 V and 1.5–1.0 V, suggests distinct electrochemical reactions associated with AlCl_4^- anion intercalation-deintercalation process in NG structure in Urea: AlCl_3 . Similarly, the broader peaks observed during AlCl_4^- anion intercalation-deintercalation process in NG with Urea: AlCl_3 is likely due to ≈ 10 times higher viscosity and lower conductivity that obviously affect mass transport and consequently kinetics of AlCl_4^- anion processes.^[25] The stability window of 1:1.5 EMIMCl: AlCl_3 , TEA: AlCl_3 , and Urea: AlCl_3 electrolytes shown in Figure 3b was evaluated up to 2.8 V. At 0.5 mA cm^{-2} , lowest onset potential of electrolyte decomposition occurred at 2.65 V and 2.72 V for EMIMCl: AlCl_3 and TEA: AlCl_3 , respectively compared to 2.8 V for more chemically stable Urea: AlCl_3 mixture.

2.4. AGB Full-Cell Experiments

The optimal upper cut-off voltage (UCV) for the AGB cell in different electrolytes was determined at a current density of 0.1 A g^{-1} , aiming to achieve a similar coulombic efficiency (CE) of 98.5%, as illustrated in Figure S1. The UCV was set as 2.40 V for cells using EMIMCl: AlCl_3 and Urea: AlCl_3 while a slightly

higher UCV of 2.45 V was set for cells employing TEA: AlCl_3 . As shown in Figure S1, at a cut-off voltage of 2.45 V, the cells with EMIMCl: AlCl_3 and Urea: AlCl_3 exhibited a CE of 87.5% and 91%, respectively. Figure 4a illustrates the charge-discharge behaviour of the AGB cell during the 25th cycle at a current density of 0.1 A g^{-1} . The highest specific capacity and energy during discharge step was achieved in the cell with TEA: AlCl_3 ($\sim 95 \text{ mAh g}^{-1}$; 171 mWh g^{-1}) followed by EMIMCl: AlCl_3 ($\sim 90 \text{ mAh g}^{-1}$; 164 mWh g^{-1}) and Urea: AlCl_3 ($\sim 84 \text{ mAh g}^{-1}$; 140 mWh g^{-1}).

The superior performance and CE of cell with TEA: AlCl_3 compared to EMIMCl: AlCl_3 and Urea: AlCl_3 is also attributed to the higher cut-off voltage (by 50 mV) during charging step. The reason for enhanced ESW for TEA: AlCl_3 can be explained by the cation-anion interaction energy (ΔE_{int}) determined by density functional theory (DFT) calculations reported by Azimi et al.^[48] These calculations reveal that TEA: AlCl_3 forms stronger coulombic and hydrogen bonding interactions than EMIMCl: AlCl_3 facilitated by the formation of six hydrogen bonds between TEAH⁺ and chloroaluminate anions in contrast to only three hydrogen bonds for EMIM⁺. While the charge/discharge profiles of AGB with EMIMCl: AlCl_3 and TEA: AlCl_3 exhibit similar characteristics and plateaus, the profile of AGB with Urea: AlCl_3 is different signified by a lower cell voltage during both charging (220–230 mV) and discharging (250–280 mV) step. Nevertheless, the specific capacity of AGB with Urea: AlCl_3 amounts to 83 mAh g^{-1} which is comparable to its performance with other ILs, whereas cell discharge voltage is also lowering strongly affecting the specific cell energy density with respect to the mass of graphite (Figure 8b). During the charging step (Figure 4b), the overpotential of the Al anode at EoC, corresponding to the aluminium electrodeposition process, notably varied among the electrolytes, emphasizing the strong influence of the electrolyte nature. The overpotential values of Al anode at EoC in EMIMCl: AlCl_3 , TEA: AlCl_3 , and Urea: AlCl_3 were 0.06 V, 0.033 V, and 0.07 V, vs. Al/Al^{3+} respectively. Correspondingly (Figure 4c), the electrode potential of NG at EoC was measured as 2.34, 2.42 V, and 2.33 V vs. Al/Al^{3+} in EMIMCl: AlCl_3 , TEA: AlCl_3 , and Urea: AlCl_3 , respectively.

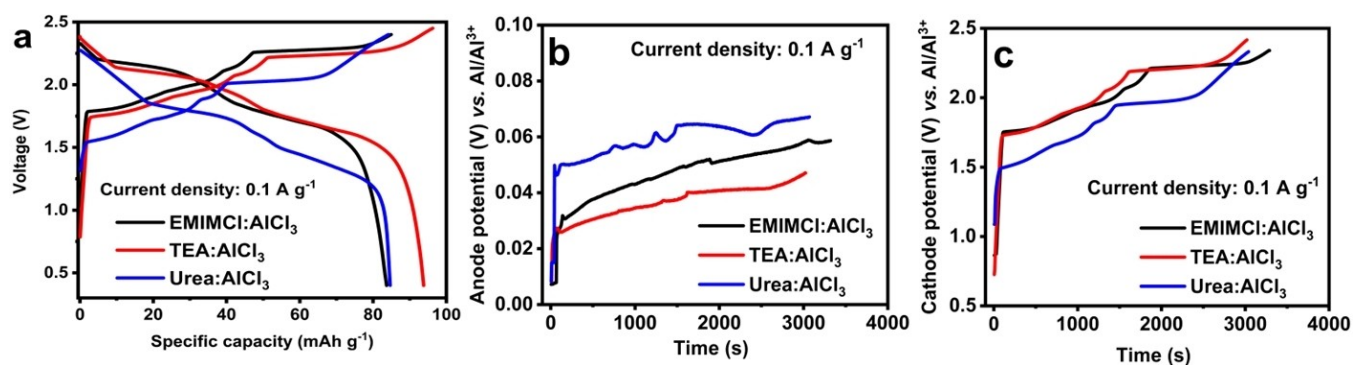


Figure 4. Comparison of a) 25th charge-discharge curves of AGB during galvanostatic cycling in different electrolytes at 0.1 A g^{-1} with respective b) anode and c) cathode potential curves during charging step.

2.5. Rate Capability

Figure 5 shows the rate capability of the AGB from a current density of 0.1 to 5 A g^{-1} . Notably, the AGB utilizing TEA: AlCl_3 demonstrates superior capacity retention, particularly at lower current densities. At 0.25 A g^{-1} , the specific capacities of the cells with EMIMCl: AlCl_3 , TEA: AlCl_3 , and Urea: AlCl_3 are 88.3, 94.4, and 84.9 mAh g^{-1} , corresponding to the capacity retention of 98, 97, and 94%, respectively, compared to their initial values at 0.1 A g^{-1} (88.7, 94.3, and 84.9 mAh g^{-1}). However, a decline in capacity is observed at 0.75 A g^{-1} , dropping to 81.3, 89.7, and 63.8 mAh g^{-1} (72, 90, and 75% capacity retention) for the respective electrolytes. This trend further intensifies at 1 A g^{-1} ,

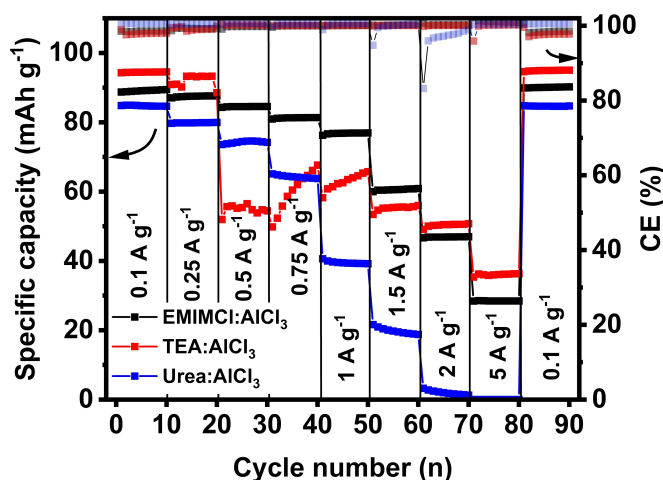


Figure 5. Comparison of specific capacity and coulombic efficiency (CE) of AGB in different electrolytes cycled at different current densities. The experiments were carried out after 25 activation cycles at 0.1 A g^{-1} shown in Figure 4a.

where capacity decreases to 64.8, 79.9, and 39.6 mAh g^{-1} . At higher current densities, cells with EMIMCl: AlCl_3 and TEA: AlCl_3 exhibit retention of 55 and 52% (at 2 A g^{-1}) and 39 and 32% (at 5 A g^{-1}), respectively. Conversely, cells using Urea: AlCl_3 show almost a complete capacity depletion at these high current densities.

Figure 6 illustrates the impact of electrolyte composition and current density on the charge-discharge behaviour (Figure 6a,c,e), as well as the potentials of the Al anode and NG cathode (Figure 6b,d,f) during the charging process. As expected, both electrolyte composition as well as current density exert a substantial influence on the potentials of the NG cathode and Al anode during the charging process (SI Table S2). These factors subsequently influence the overall capacity of AGBs, as demonstrated in Figure 6. With EMIMCl: AlCl_3 , a consistent and gradual increase in anode potential is observed as the current density rises from 0.1 A g^{-1} to 5 A g^{-1} , alongside a steady decrease in cathode potential from 2.333 V to 2.049 V (Figure 6a). The consistent potential changes on anode and cathode are reflected in the gradual decrease in capacity (especially from last charging plateau) from the charge-discharge curves (Figure 6b). In contrast, AGB with TEA: AlCl_3 (Figure 6c) shows a similar trend in cathode potential reduction which reduces from 2.42 V to 2.02 V, but the anode potential exhibits a more rapid increase, moving from 0.033 V to 0.359 V. This suggests that cell with TEA: AlCl_3 is less efficient at higher current densities which is evident by looking at abrupt transitions observed in the charge-discharge curves (Figure 6d). Cell using Urea: AlCl_3 electrolyte displays the most dramatic change as Al anode overpotential potentials increased significantly (Figure 6e), leading to AGB delivering almost no capacity at 5 A g^{-1} (Figure 6f). These variations are attributed to the diffusion limitations of AlCl_4^- intercalation/deintercalation with-

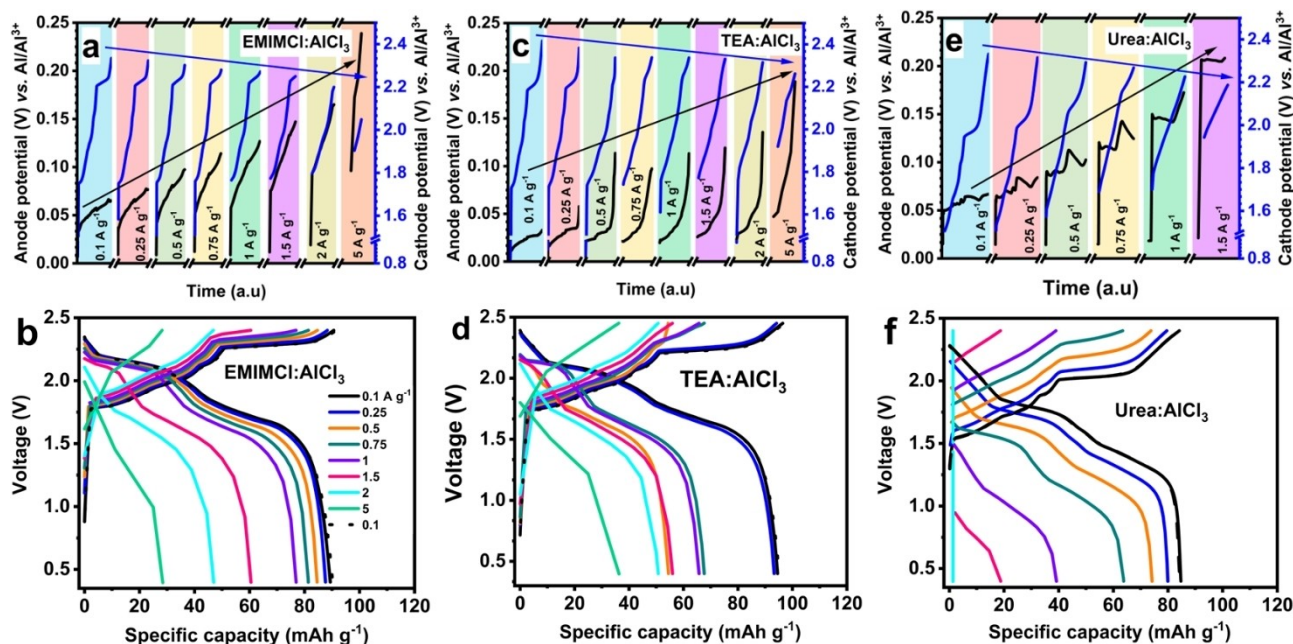


Figure 6. Comparison of potential of Al anode and natural graphite cathode during charging step at different current densities in a) EMIMCl: AlCl_3 c) TEA: AlCl_3 and e) Urea: AlCl_3 . Comparison of charge-discharge curves at different current densities in b) EMIMCl: AlCl_3 , d) TEA: AlCl_3 and f) Urea: AlCl_3 .

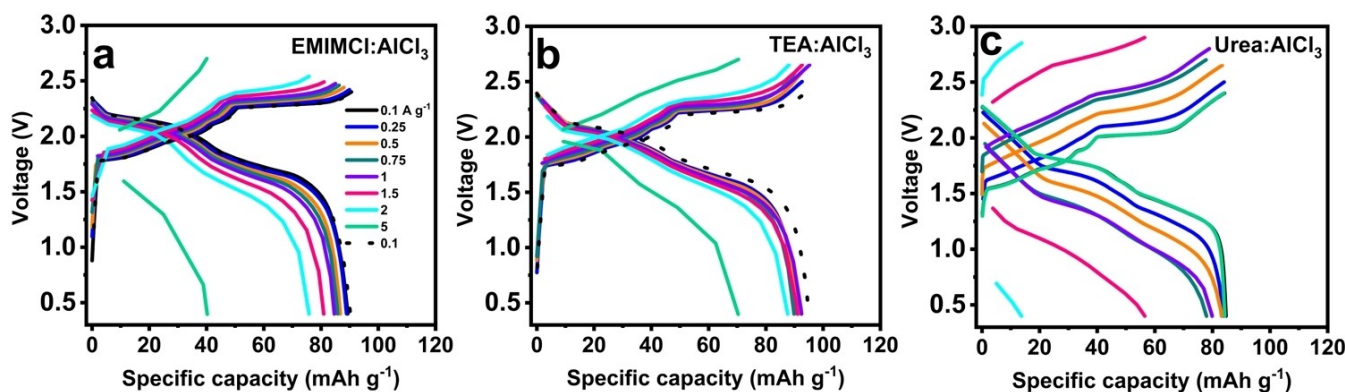


Figure 7. Comparison of charge-discharge curves at different current densities after adjusted UCV during charging step in a) EMIMCl:AlCl₃, b) TEA:AlCl₃ and c) Urea:AlCl₃.

in the graphite layers at higher current densities,^[53] as well as factors like ion-pair interactions, viscosity, and ion complex formation affecting ion mobility and conductivity of each electrolyte whose influence were also observed in Al–Al symmetric cells.^[48] The observed increase in the anode potential at EoC during the charging process with increase in current density in the AGB cell closely resembles the pattern of the Al electrode potential during the Al deposition process in the Al–Al symmetric cell (Figure 2a). Particularly noteworthy is the drastic surge in the Al electrode potential during Al deposition in TEA:AlCl₃ in both the AGB full cell and the Al–Al symmetric cell from the current densities of 0.5 A g⁻¹ and 1.5 mA cm⁻², respectively that confirms the poor efficiency of the Al deposition process in that electrolyte at higher current densities.

In contrast, the maximum achievable specific capacity of AGB in EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃ is realized only when the NG cathode approaches the potentials of 2.34 V, 2.417 V, and 2.332 V potential at EoC, respectively (Figure 4c), which is a determining factor for the overall capacity.^[54] Hence, a lower potential of NG cathode at EoC at higher current densities can be an indication of an ‘incomplete’ or partial charging process of NG, elucidating another possible reason for the observed capacity fade with an increase in current density. To mitigate the effects of increased overpotentials on the Al anode during charging process, the UCV of the AGB cell was meticulously adjusted for each current density (see Table S3).

Importantly, higher cut-off voltages did not affect the Mo current collector, which demonstrates electrochemical stability up to 2.5 V versus Al/Al³⁺.^[19] Moreover, the cathode potential consistently stayed below this threshold until the EoC, ensuring safe operation within the electrochemical stability limits of the materials involved. The impact of cut-off voltage adjustment on the specific capacity and the capacity retention (vs. at 0.1 A g⁻¹) especially at higher current densities from 0.5 to 5 A g⁻¹ is clearly observed in the AGB cell across all electrolytes, as shown in Figure 7 (a–c) and Figure S3. Figure 8a shows that at 0.5 A g⁻¹, the specific capacity of AGB with TEA:AlCl₃ increased substantially by ~36 mA h g⁻¹ compared to the average specific capacity before UCV adjustment at the same current density,

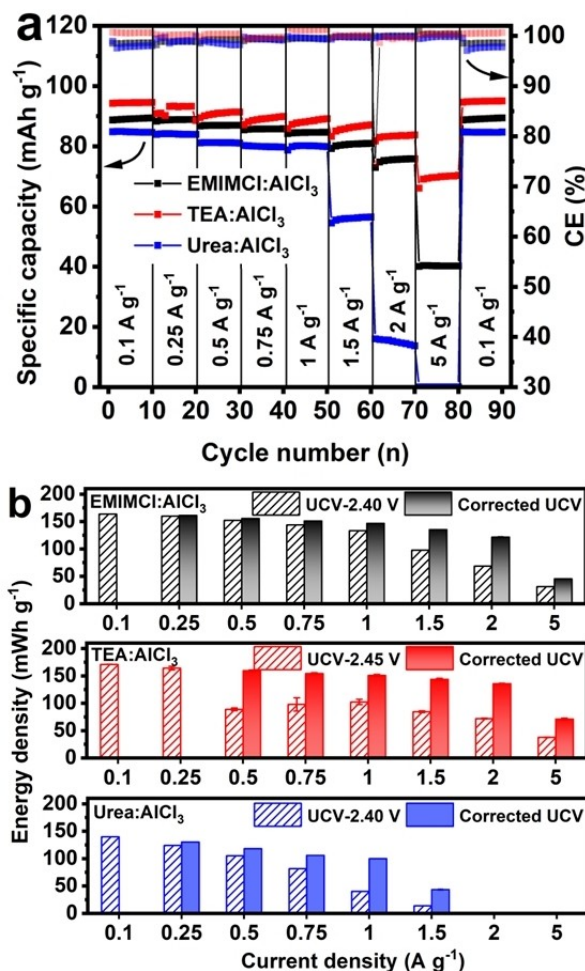


Figure 8. Comparison of a) specific capacity and coulombic efficiency (CE) of Al-NG AIB in different electrolytes cycled at different current density after UCV adjustment and b) energy density of NG cathode at different current densities between before and after UCV adjustment.

while the cell with EMIMCl:AlCl₃ and Urea:AlCl₃ showed smaller gains of ~2.5 mA h g⁻¹ and ~7 mA h g⁻¹, respectively. At 1 A g⁻¹, the capacity of AGB with EMIMCl:AlCl₃ increased by ~8 mA h g⁻¹, with TEA:AlCl₃ and Urea:AlCl₃ improved by ~25 mA h g⁻¹ and ~40 mA h g⁻¹, respectively. At 2 A g⁻¹, AGB

with TEA:AlCl₃ showed a significant increase by 33 mAh g⁻¹. Following the adjustment of UCV for different current densities, the cathode potential at EoC in EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃ remained consistent at 2.34 V, 2.417 V, and 2.332 V, respectively, across all current densities (Figure S2) indicating 'complete' charging of NG. Owing to substantial enhancements in AGB capacities across all electrolytes up to 2 Ag⁻¹ following UCV adjustments, an increase in the energy density of the NG cathode (reported energy densities were normalized to the mass of graphite) is evident, particularly at higher current densities, as depicted in Figure 8b. Notably, at 1 Ag⁻¹, the energy density of NG increased by 10% in EMIMCl:AlCl₃, 48% in TEA:AlCl₃, and an impressive 250% in Urea:AlCl₃ after the UCV adjustment. Despite encountering capacity fade at 2 Ag⁻¹ in Urea:AlCl₃, the energy density of NG witnessed a remarkable 300% increase in Urea:AlCl₃, 70% in TEA:AlCl₃, and 40% in EMIMCl:AlCl₃.

2.6. Long-Term Cycling

The long-term stability of AGBs was investigated in different electrolytes over a testing period of 1000 cycles at 1 Ag⁻¹ and the results are shown in Figure 9a. Initially, the cell with

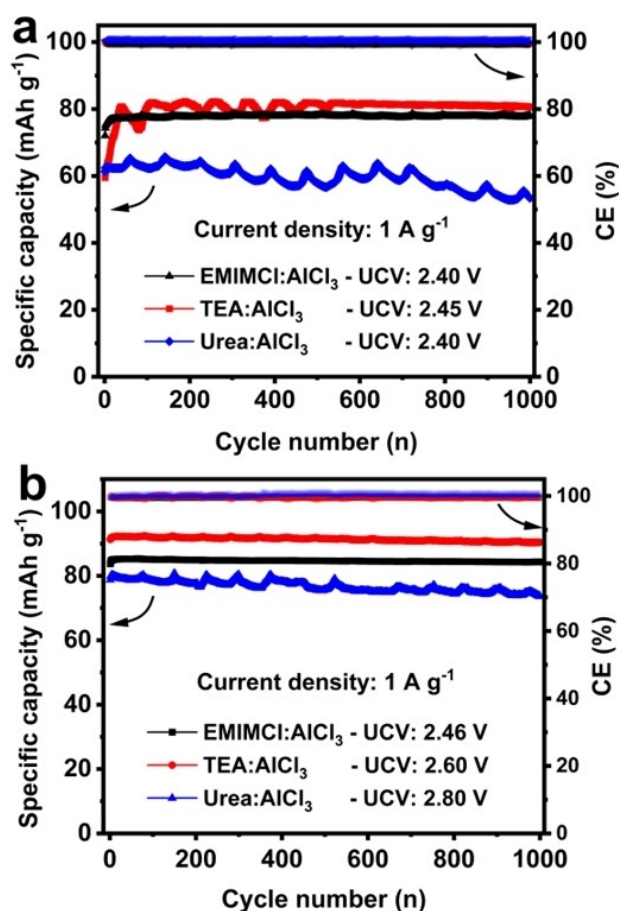


Figure 9. Comparison of specific capacities between AGBs with EMIMCl:AlCl₃, TEA:AlCl₃ and Urea:AlCl₃ electrolytes during long-term cycling at 1 Ag⁻¹ for 1000 cycles a) before and b) after UCV adjustment.

EMIMCl:AlCl₃ exhibited the highest capacity (78 mAh g⁻¹) during the first 50 cycles, maintaining this value consistently throughout the entire 1000 cycles. In contrast, the cell with TEA:AlCl₃ experienced a notable increase in capacity from 59 mAh g⁻¹ to 81 mAh g⁻¹ within the initial 60 cycles, stabilizing at 81 mAh g⁻¹ after 500 cycles. This change in capacity is attributed to a reduction in the aluminium anode overpotential (Figure S4), likely resulting from improvements in the aluminium-electrolyte interface. Conversely, the cell with Urea:AlCl₃ displayed a comparatively lower capacity of 62 mAh g⁻¹, with a 15% reduction to 52.7 mAh g⁻¹ after 1000 cycles, while cells with EMIMCl:AlCl₃ and TEA:AlCl₃ showed negligible fade. Throughout the long-term cycling, the CE remained consistently above 99% for all the cells.

Additionally, periodic fluctuations in capacity, which occurred approximately every 24 hours were attributed to day-night temperature variations in the lab. These fluctuation was estimated to 1.5, 6, and 15% for EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃, respectively due to the differences in the viscosity and transport properties of the electrolytes.^[55] Adjusting UCV resulted in a substantial capacity increase to 85, 92, and 79 mAh g⁻¹ corresponding to a capacity enhancements of 10, 13, and 27% with EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃, respectively (Figure 9b and Figure S3). Furthermore, the influence of temperature on capacity fluctuation was minimized to less than 2% in EMIMCl:AlCl₃ and TEA:AlCl₃, and less than 5% in Urea:AlCl₃ after UCV adjustment. The morphological changes in NG cathode after long-term cycling with EMIMCl:AlCl₃, TEA:AlCl₃ and Urea:AlCl₃ electrolytes and the impact of UCV adjustments were characterised by Raman spectra and XRD are shown in Figure 10a and Figure 10b, respectively. Before UCV adjustment, an increase in I_d/I_g ratio was observed from 0.105 (pristine NG) to 0.20, 0.45, and 0.13 cycled in EMIMCl:AlCl₃ (2.40 V), TEA:AlCl₃ (2.45 V) and Urea:AlCl₃ (2.40 V), respectively. However, a shift in peak corresponding to G band from 1575 cm⁻¹ to 1583 cm⁻¹ along with the shift of peak at 1614 to 1625 cm⁻¹ associated to D' band, is an indication for enhanced lattice disorder and greater AlCl₄⁻ intercalation-induced defects.^[22]

The XRD diffractogram (Figure 9b) show a similar change, revealing a peak shift of 2θ from 26.50° to 26.65° (only after cycling in ILs), corresponding to a decrease in interlayer spacing from 3.4 Å to 3.3 Å, and indicating compressive stress on the graphene layers. Moreover, the increased peak broadening and FWHM values suggest a rise in structural disorder and lattice distortions within the graphite matrix.^[21,56] The crystallite domain dimensions along the *c*-axis (L_c) of NG calculated using the Scherrer equation decreased from 52.13 nm (pristine NG) to 41.71 nm particularly after cycling in ILs EMIMCl:AlCl₃ and TEA:AlCl₃ (Figure S5). The observed decrease in L_c value can be attributed to the intercalation of larger EMIM⁺ (109.3 Å) and Et₃NH⁺ (128.7 Å) cations in NG leading to a lower L_c value.^[37] Zhang et al showed that the graphite with higher L_c requires more activation energy to enlarge the interlayer spacing between adjacent layers resulting in slower ion kinetics, compared to graphene materials with lower L_c values.^[51] Therefore the reduced L_c value of NG cycled in ILs aligns with the

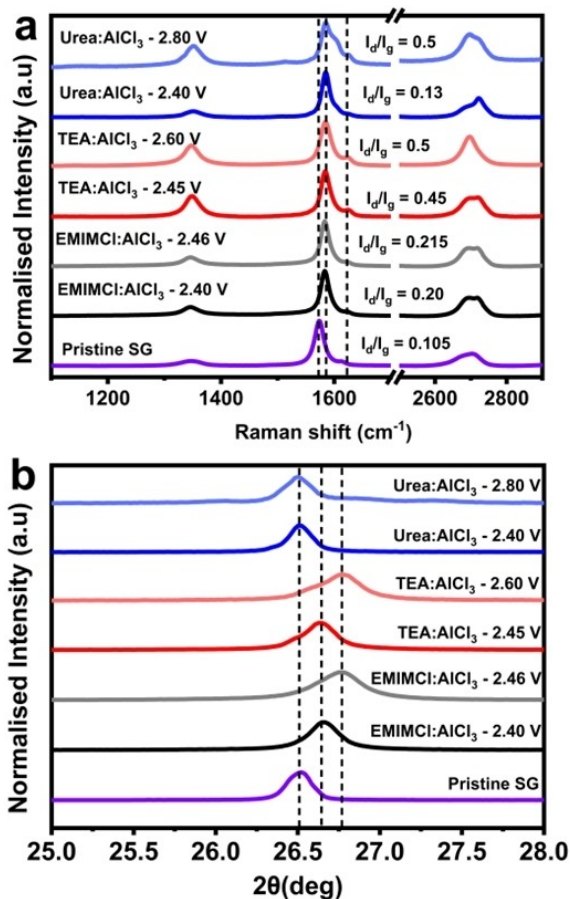


Figure 10. Comparison of a) Raman spectra and b) XRD diffractogram of NG cathode between AGBs with EMIMCl:AlCl₃, TEA:AlCl₃ and Urea:AlCl₃ electrolytes after long-term cycling at 1 Ag⁻¹ for 1000 cycles.

higher capacity observed, as this decrease in L_c value results in faster AlCl₄⁻ intercalation/deintercalation kinetics.^[51,57] In contrast, long-term cycling of NG with Urea:AlCl₃ resulted in minimal structural changes as evidence by a marginal increased in the I_d/I_g to 0.13 while no significant shift in the 2θ peak and L_c value. This suggest that AlCl₂(urea)₂⁺ cations do not significantly influence the graphite intercalation reaction.^[22] After UCV adjustments, the I_d/I_g ratio of NG increased to 0.215, 0.5, and 0.5 cycled in EMIMCl:AlCl₃ (2.46 V), TEA:AlCl₃ (2.60 V) and Urea:AlCl₃ (2.80 V), respectively, showing increased compressive strain as evidenced by a peak shift from 26.50° to 26.75° and a decrease in L_c values to 29.8 nm for NG cycled with EMIMCl:AlCl₃ and TEA:AlCl₃. The L_c value of NG cycled with Urea:AlCl₃ decreased to 34.75 nm, consistent with 27% increase in capacity, yet without a corresponding 2θ peak shift, highlighting the negligible influence of AlCl₂(urea)₂⁺ cations. These results show the distinct effects of electrolyte type (ILs and DES) and the effect of charging protocol (before and after UCV adjustment) on the NG morphology and AGB capacity after long-term cycling.

3. Conclusions

This study reports on the influence of current density on the electrode potential during the Al dissolution/deposition process and the charge-discharge behaviour of AGB in different 1:1.5 AlCl₃-based electrolytes. In the Al–Al symmetric cell, increasing current density above 1 mAcm⁻² resulted in a notable increase in overpotential up to 200 mV at 5 mAcm⁻² during the aluminium deposition process in both 1:1.5 TEA:AlCl₃ and Urea:AlCl₃. Similarly, higher overpotentials during Al deposition process were observed in AGB full-cell during charging step, resulting in incomplete AlCl₄⁻ intercalation in NG and subsequent capacity fade at higher current densities.

Adjusting the UCV in the AGB full-cell for each current density improved specific capacities up to 1 Ag⁻¹ and led to remarkable reduction in capacity fading less than 7% across all electrolytes. This shows a significant improvement compared to the fading rates of 14, 33, and 53% in EMIMCl:AlCl₃, TEA:AlCl₃ and Urea:AlCl₃, respectively, before UCV adjustment. The AGB cells exhibited substantial improvements in energy density, with a 48% increase observed with TEA:AlCl₃ and an impressive 250% increase with Urea:AlCl₃ at 1 Ag⁻¹ after UCV adjustment. Structural analyses through Raman spectroscopy and XRD provided insights into the changes within the graphite structure. These changes were attributed to the compressive stresses caused by cations (EMIM⁺ and Et₃NH⁺) in addition to the structural deformations induced by electrolyte interactions and AlCl₄⁻ intercalation at different UCVs. Systematic monitoring of anode overpotential during charging and discharging steps may provide valuable insights for optimizing anode morphology, such as increasing its active surface area for improved performance at high current densities.

Author Contributions

C.M. designed the work, conceptualized, and conducted the experiments, analysed the data, and wrote the manuscript. J.F.D supervised the project and reviewed the manuscript. All authors have approved the final version of the manuscript.

Acknowledgements

The authors would like to thank co-workers Ms. Melanie Thalheimer for her support for FESEM micrographs and Mr. Martin Eckert for valuable discussions. The authors acknowledge Federal Ministry of Education and Research for financial support (Grant No. 03XP0392 C ALBATROS project). Open Access funding enabled and organized by Projekt DEAL.

Conflict of Interests

The authors declare no conflict of interest.

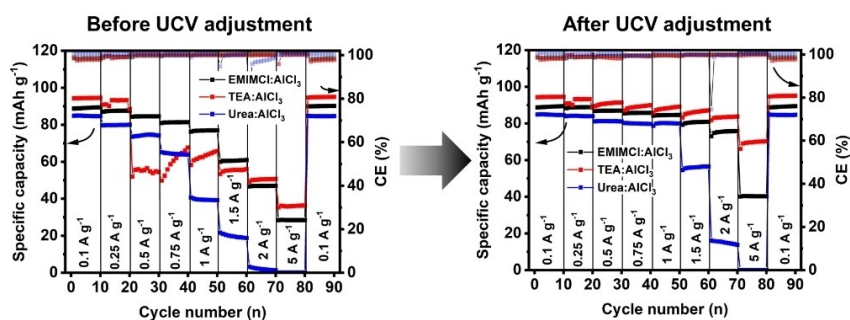
Data Availability Statement

The data that support the findings of this study are available in the supplementary material of this article.

Keywords: Aluminium graphite battery · Graphite electrode · Electrode overpotential · Chloroaluminate electrolyte · Aluminium ion battery

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Manuscript received: November 11, 2024
 Revised manuscript received: December 16, 2024
 Accepted manuscript online: January 1, 2025
 Version of record online: ■■■■■



This study shows the impact of current density on Al electrode potential during deposition/stripping reaction in a Al symmetric cell using 1:1.5 EMIMCl:AlCl₃, TEA:AlCl₃, and Urea:AlCl₃ electrolytes. By adapting

cut-off voltage value during charging step of Al-Graphite cell allowing maximum AlCl₄⁻ ion intercalation into the cathode, a clear increase in cell performance was observed during discharging step.

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Mitigating the Effect of High Overpotential during Al Deposition on Aluminium-Graphite Battery Performance

