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Electro-polymerisation and characterisation of PEDOT in Lewis basic, neutral and acidic EMImCl-AlCl₃ ionic liquid

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A B S T R A C T

This work studied the electro-polymerisation of 3,4-ethylenedioxythiophene (EDOT) and its electrochemical behavior in Lewis acidic, neutral and basic chloroaluminate ionic liquid 1-ethyl-3-methylimidazolium chloride aluminum chloride (EMImCl-AlCl₃) by cyclic voltammetry. It was found that the electro-polymerisation on vitreous carbon only occurs in Lewis neutral EMImCl-AlCl₃ as a dark blue-violet film whereas the electro-polymerisation in a Lewis acidic or basic compositions is not possible due to the interactions between the conductive polymer and the ionic liquid as well as the potential stability limits of the electrolyte. PEDOT films synthesised in Lewis neutral ionic liquid were tested in monomer-free Lewis acidic, basic and neutral EMImCl-AlCl₃ and show different doping and de-doping behavior for chloride ionic species. The PEDOT films in a Lewis neutral composition showed higher doping levels due to the higher potential stability window, up to 2.6 V vs. Al/Al(III) than in a Lewis acidic and basic solutions. Furthermore, it was shown that the doping and de-doping levels are predefined during the electro-polymerisation of PEDOT. The anion doping and de-doping reaction reached 97% reversibility in the neutral composition, which suggests that PEDOT is a suitable electrode material to store charged species in this media and could be used in rechargeable energy storage devices.

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1. Introduction

Conductive polymers such as poly-(3,4-ethylenedioxythiophene) (PEDOT) have high potential as electrode materials for high performance energy storage devices due to their low costs, low weight, high electrochemical stability and reversibility as well as conductivity of up to 1000 S cm⁻¹ [1]. PEDOT is characterised by its ability to store a large amount of charges, working as capacitor and battery at the same time. The charge storage occurs due to the doping of anions into the conductive polymer backbone while the polymer is oxidised. The oxidation causes positive charged centers (sulfur or carbon) in the polymer structure, which are compensated by the doping anions [2,3].

High performance energy storage devices require high capacities and high cell potentials, which can only be reached in non-aqueous systems like ionic liquids. The combination of a conductive polymer electrode in an ionic liquid electrolyte offers the advantages of a stable potential window from 4.5 to 6 V, high chemical stability, conductivity and safe electrode reactions [4,5]. PEDOT films can be formed by a simple electrochemical synthesis due to their low oxidation potential in aqueous (≈ 1 V vs. SHE) and non-aqueous (≈ 1.5 V vs. Al/Al(III)) solutions. The electropolymerisation of PEDOT films in several ionic liquids by cyclic voltammetry (CV), chronocoulometry (CA) and chronopotentiometry (CP) has been already reported in several studies (Table 1).

A great variety of ionic liquids have been tested in which the polymerisation resulted in films with different electrochemical and structural characteristics. PEDOT films obtained in BMIm-BF₄, EMIm-AlCl₃ and EMIm-PFSix show a very high cycle stability and no degradation or detachment from their substrates. PEDOT can be doped by anions (p-doping) and cations (n-doping) either in the oxidised or reduced state [1]. However, PEDOT films in BMMAm-TFSA and BMPy-OTF show only anion doping. Electro-polymerised EDOT appears as agglomerated surface structure in...
aqueous, organic and also ionic liquid electrolytes. Highly porous structures were obtained in EMIm-PSFI offering a high number doping positions and consequently potential high storage capacities.

The doping reaction of PEDOT in a chloroaluminate ionic liquid like 1-ethyl-3-methylimidazolium chloride-aluminum chloride (EMImCl-AlCl₃) (Equation (1)) [17] is suggested because the doping anions Al₂Cl₇, AlCl₄ or Cl⁻ present in the electrolyte have a smaller ion radius than the organic anions like TFSI^−, TFSA^− and OTF^− with complex and bulky structures, which can be reversibly doped into the conductive polymer. In addition, solutions with the imidazolium cation EMIm⁺ are characterised by a high conductivity (order of 10 mS cm⁻¹) and low viscosity (10–50 mPa s at 20 to 30 °C) among other ionic liquids [4,18]. The type of the chloroaluminate doping anion X⁻ depends on the Lewis acidity of the ionic liquid.

\[ \text{EDOT}^{1/2} + X^{-} \xrightarrow{\text{doping}} \text{[EDOT]}_{1/2} + e^{-} \]  
\[ X^{-} = \text{Cl}^{-}, \text{AlCl}_{4}^{-}, \text{Al}_{2}\text{Cl}_{7}^{-} \quad |\alpha| = \text{degree of doping: 0 to 1.} \]

This study investigates the electro-polymerisation of EDOT in EMImCl-AlCl₃ with different Lewis acidities by cyclic voltammetry. Furthermore, the influence of the doping anion species on the conductive polymer performance was investigated in monomer free Lewis acidic, neutral and basic EMImCl-AlCl₃. The open question of whether or not the doping and de-doping levels of the conductive polymer are predefined during the electro-polymerisation or the levels are formed during cycling in the electrolyte with present doping anions is addressed.

### Table 1

| Name ionic liquid (abbreviation) | Methods/Properties | Ref. |
|---------------------------------|--------------------|-----|
| 1-Butyl-3-methylimidazolium-tetrafluoroborate (BMIm-BF₄) | CV, CA | High polymerisation or the levels are formed during cycling in the conductive polymer | [17] |
| 1-Butyl-3-methylimidazolium-trifluoromethanesulfonate (BMIm-TSFI) | CV | Only anion doping, difficult electro-polymerisation, nodular structure | [11] |
| 1-Butyl-1-methylpyrrolidinium-trifluoromethanesulfonate (BMPy-TSFI) | CV | Only anion doping | [11] |
| 1-Ethyl-3-methylimidazolium-aluminum chloride (EMIm-AlCl₃) | CV, CA, CP | Highly stable films, granular and porous structure | [13,14] |
| 1-Ethyl-3-methylimidazolium-trifluoromethanesulfonate (EMIm-TSFI) | CV | Only anion doping, highly porous structure | [11] |
| 1-Ethyl-3-methylimidazolium-trifluoromethanesulfonate (EMIm-OFT) | CV | Limited reversibility anion doping, fast film growth | [11,12,15] |
| 1-Ethyl-3-methylimidazolium-bis(trifluoromethysulfonyl)imide (EMIm-TFSI) | CV, CA | Cation-doping, highly conductive films, compact and open structure films | [7,9,16] |
| 1-Ethyl-2,3-dimethylimidazolium-bis(trifluoromethysulfonyl)imide (EMMMIm-TFSI) | CV, CA | Medium to low ion exchange rate | [9] |
| 1-Octyl-3-methylimidazolium-bis(trifluoromethanesulfonate) (OMIm-TFSI) | CV | Fast polymerisation, high degree of conjugation | [6] |

### Table 2

| Molar ratio (\(\chi\)) of EMImCl to AlCl₃ and the resulting Lewis acidity and dominant anion species in a binary mixture of EMImCl-AlCl₃ ionic liquid [19]. | \(\chi(\text{EMImCl})\) (mol%) | \(\chi(\text{AlCl}_3)\) (mol%) | Lewis acidity | Predominant anions in solution |
|-----------------------------------------------|---------------------------------|----------------------------|-----------------|-----------------------------|
| 33.3 | 66.7 | Acidic | AlCl₄⁻, AlCl₂⁻ |
| 50 | 50 | Neutral | AlCl₄⁻ |
| 66.7 | 33.3 | Basic | AlCl₄⁻, Cl⁻ |

### 2. Experimental

#### 2.1. Ionic liquid preparation

A Lewis acidic, neutral and basic imidazolium based ionic liquid was prepared using the two components: 1-ethyl-3-methylimidazolium chloride (EMImCl; Merck, purity ≥ 98%, water < 1%) and aluminum chloride (AlCl₃; Alfa Aesar, anhydrous, ultra-dry, packed in glass ampoule under argon gas, metal basis 99.99%). The Lewis acidity of the ionic liquid was controlled by the molar ratio of EMImCl to AlCl₃ (Table 2).

The components EMImCl and AlCl₃ were dried before use in a vacuum oven at 60 °C for 24 h and subsequently transferred into a glove box (MBraun, Workstations UNIlab Plus/Pro - SP/DP) with nitrogen atmosphere (<0.5 ppm water and <0.5 ppm oxygen).

The amount of AlCl₃ (\(m_A\)) (Equation (2)) and EMImCl (\(m_E\)) (Equation (3)) for the synthesis of a Lewis acidic, neutral and basic ionic liquid was calculated based on the molar ratio (\(\chi\)), molar mass (\(M\)) of AlCl₃ and EMImCl and density (\(\rho\)) as well as volume (\(V\)) of the binary solution.

\[ m_A = \frac{\chi_A \cdot M_A \cdot \rho \cdot V}{\chi_A \cdot M_A + \chi_E \cdot M_E} \]  
\[ m_E = (\rho \cdot V) - m_A \]  

The density of the solution (\(\rho\)) (Equation (4)) was determined by using the reported density (\(\alpha\)) and corrected by a temperature dependent density coefficient (\(b\)) at 30 °C (Table 3) [20].

\[ \rho = \alpha + b \cdot (T - 60) \]
The calculated amount of EMImCl and AlCl3 was weighted inside the glove box with a scale (Ohaus, ± 0.001 g) and transferred into a clean and dry beaker, respectively. The ionic liquid was prepared by adding slowly AlCl3 to EMImCl under continuous stirring and cooling in a Peltier controlled cooling device with ceramic-coated beads (Techne, Bibby Scientific, N° 1°C, 0–40 °C) (Fig. 1a). The ionic liquid reached temperatures between 20 °C and 40 °C during the mixing. Afterwards, the solution was transferred into a glass bottle and stirred for further 24 h, resulting in translucent yellowish solutions (Fig. 1b). The electro-polymerisation solutions were prepared using the previous synthesised Lewis acidic, neutral and basic solutions for electro-polymerisation.

2.2. Electro-polymerisation of EDOT in liquid ionic

The electro-polymerisation of the monomer EDOT was performed in a PTFE-cell with a vitreous carbon disk working electrode (Micro-to-Nano, high purity, surface area 0.8 cm²), which faced a counter electrode of the same material and area at an inter electrode-distance of 2 mm. The cell was equipped with an aluminium-tip reference electrode (<1 mm) located very close to the working electrode (≈ 0.5 mm). The vitreous carbon electrode surfaces were polished with aluminium oxide suspension (Alfa Aesar, particle size 0.3 μm), cleaned with ethanol (Fluka, absolute, analytical reagent grade) and dried under vacuum at 60 °C for 24 h before use in the glove box. The tip of the reference electrode was polished with very fine abrasive paper inside the glove box in order to remove possible oxides on the aluminum surface.

The Lewis acidic, neutral and basic ionic liquid electrolyte each containing 0.1 mol dm⁻³ EDOT were tested as polymerisation solution. The polymerisation was performed by cyclic voltammetry (BioLogic SP-150 potentiostat and Ivium Vertex) from −0.5 V to 2.6 V vs. Al/Al(III) for 20 cycles, 100 mV vs. Al/Al(III) in a monomer-free LEwIS acidic EMImCl-AlCl3 ionic liquid electrolyte with 0.1 mol dm⁻³ EDOT by cyclic voltammetry.

3. Results and discussion

3.1. Electro-polymerisation of EDOT

The electro-polymerisation of EDOT on vitreous carbon substrates was performed in Lewis acidic, basic and neutral EMImCl-AlCl3 ionic liquid electrolyte with 0.1 mol dm⁻³ EDOT by cyclic voltammetry.

3.1.1. Electro-polymerisation of EDOT in Lewis acidic EMImCl-AlCl3

In a first step, the potential stability window of the monomer-free Lewis acidic EMImCl-AlCl3 was determined by cyclic voltammetry. The aluminium deposition and dissolution occurs from −0.2 to −0.5 V and −0.2 V to 0.1 V vs. Al/Al(III) on a carbon substrate, respectively, in a monomer-free Lewis acidic EMImCl-AlCl3. The decomposition of the cation species EMIm⁺ on a carbon electrode was observed below −0.5 V vs. Al/Al(III) and the oxidation of the chloroaluminate anion to chlorine gas takes place over ≈ 1.5 V vs. Al/Al(III) (Fig. 2, dashed line).

The addition of 0.1 mol dm⁻³ EDOT to the Lewis acidic EMImCl-AlCl3 caused a dark brown and very viscous solution immediately (Fig. 1c). It is assumed that a stable complex is formed between EDOT and AlCl3 [13]. The solution was tested to electro-polymerise EDOT on a carbon electrode of 0.8 cm² surface area from 0 V to 2 V vs. Al/Al(III) (Fig. 2, solid line).

The cyclic voltammetry of the Lewis acidic ionic liquid with 0.1 mol dm⁻³ EDOT showed a steady increasing current at anodic potentials until 1.5 V vs. Al/Al(III). It is assumed that the current slope is caused by the high viscosity of the ionic liquid solution which increases the electrical resistance of the electrolyte. The solution decomposed to chlorine gas at applied potentials higher than 1.5 V vs. Al/Al(III). A polymer film was not deposited on the vitreous carbon substrate, instead EDOT and AlCl3 formed a stable complex in solution, preventing a further electro-polymerisation on the vitreous carbon substrate. However, a small peak around 1 V vs. Al/Al(III) in the backward scan is visible, which might be an indication of a de-doping reaction of a thin polymer film attached on the electrode surface.

Fig. 1. (a) Synthesis of EMImCl-AlCl3 ionic liquid in cooling bath. (b) Resulting Lewis basic, neutral and acidic solution. (c) Image of the solutions in (b) after addition of 0.1 mol dm⁻³ EDOT and resulting Lewis basic, neutral and acidic solutions for electro-polymerisation.
3.1.2. Electro-polymerisation in Lewis basic EMImCl-AlCl$_3$

The potential stability window of a monomer-free Lewis basic EMImCl-AlCl$_3$ is similar to a Lewis acidic composition shown in the previous section. The anodic and cathodic limits are below 1.5 V and 0.8 V vs. Al/Al[III], respectively (Fig. 3, dashed line).

The addition of 0.1 mol dm$^{-3}$ EDOT to the light-yellow Lewis basic EMImCl-AlCl$_3$ ionic liquid did not cause any color changes but the electro-polymerisation of EDOT on the electrode using this solution was not observed. The liquid decomposed to chlorine gas from 1.5 V vs. Al/Al[III] before EDOT could have been polymerised, although the anodic stability potential window is $\approx 0.5$ V wider with EDOT than the bare Lewis basic ionic liquid (Fig. 3, solid line). In addition, the conductivity of a Lewis basic chloroaluminate ionic liquid is lower (2.07 mS cm$^{-1}$ at 21.4 $^\circ$C [20]) than a Lewis acidic (14 mS cm$^{-1}$ at 22.2 $^\circ$C [20]) composition, which is represented in the lower current density of the cyclic voltammogram (Fig. 3, solid line).

3.1.3. Electro-polymerisation in Lewis neutral EMImCl-AlCl$_3$

The monomer-free Lewis neutral solution has the widest potential stability window of 4 V from -2 V to +2 V vs. Al/Al[III] (Fig. 4, inset CV).

The Lewis neutral ionic liquid with 0.1 mol dm$^{-3}$ EDOT changed its color from translucent yellowish two dark-orange within 48 h. The cyclic voltammogram of the electro-polymerisation from $-0.5$ to $-2.6$ V vs. Al/Al[III], recorded from 0 V vs. Al/Al[III], shows a continuous growth of the polymer film on the vitreous carbon surface with every cycle (Fig. 4, main CV). The two first cycles show a clear peak $>2.0$ V vs. Al/Al[III], which is related to the nucleation of a polymer film on the vitreous carbon surface. Cycle three to twenty are characterised by the growth of polymer on previously polymerised polymer. The anodic potential stability window of the Lewis neutral ionic liquid with 0.1 mol dm$^{-3}$ EDOT seems to reach up to 2.8 V vs. Al/Al[III], which is $\approx 0.8$ V wider than the bare Lewis neutral ionic liquid.

The predominant anion in the Lewis neutral chloroaluminate ionic liquid is AlCl$_4^-$ and the doping of the anion into the conductive polymer occurs simultaneously during the polymerisation of EDOT on the vitreous carbon electrode surface. The oxidation/polymerisation and doping process is characterised by four anodic peaks at 0.75 V, 1.1 V, 1.7 V and 2.2 V vs. Al/Al[III], which overlap each other. The de-doping reaction shows two clear peaks at 1.6 V and 0.9 V vs. Al/Al[III].

Anodic ($Q_a$) and cathodic ($Q_c$) transferred charges from cycle 1 to 20 increased from 35 mC to 74 mC and 19 mC–66 mC, respectively, due to polymer growth and the formation of more doping positions in the polymer films, which allows a higher number of transferred doping anions. If it is assumed that the cathodic transferred charges indicate the de-doping of AlCl$_4^-$, the same proportion of anodic charges ($Q_a$) will correspond to the initially amount of doped charges AlCl$_4^-$ ($Q_{dop} = Q_{dop}$). The difference between the transferred charges of the doping reaction and the measured overall anodic value corresponds to the charge of the polymerisation reaction ($Q_{pol} = Q_c - Q_{dop}$). The ratio $Q_{pol}/Q_a$ decreases from 45% to 10%, indicating that the polymerisation reaction reaches a limit and the doping reaction becomes dominant. At the same time, the ratio $Q_a/Q_c$ increases from 54% to 89%, which suggest a reversible doping and de-doping of AlCl$_4^-$ into PEDOT.
3.2. Characterisation PEDOT films

The doping and de-doping behavior of PEDOT were studied in monomer-free Lewis acidic, basic and neutral ionic liquid by cyclic voltammetry (Fig. 5). The PEDOT films were previously electro-polymerised in Lewis neutral EMImCl-AlCl₃ with 0.1 mol dm⁻³ EDOT by cyclic voltammetry for 20 cycles, at 100 mV s⁻¹ and 25 °C. The PEDOT films were rinsed once with acetonitrile after the electro-polymerisation in order to remove residual polymerisation solution.

The cyclic voltammogram of the PEDOT film in Lewis acidic ionic liquid (Fig. 5a) shows three doping processes at 0.5 V, 1.5 V and 2 V as well as two de-doping peaks at 1.2 V and 0.8 V vs. Al/Al(III) (Table 4). The doping anions in this Lewis acidic ionic liquid are AlCl₄⁻ and Al₂Cl₇⁻, showing a doping/de-doping reversibility (Qc/Qa) of 43% into the polymer film. On the PEDOT surface a viscous and dark orange-brown film was formed after a few cycles. The color and consistence of the film was comparable with Lewis acidic EMImCl-AlCl₃ solution with EDOT (Fig. 1c). It is assumed that this viscous film increased the electrolyte resistance, which was observed as increasing slope of the cyclic voltammogram.

The PEDOT film in Lewis neutral EMImCl-AlCl₃ (Fig. 5b) shows one pair of doping and de-doping peaks at higher potentials. Two clear separated de-doping peaks are shown at 1.6 V and 0.8 V vs. Al/Al(III), respectively. The peak at 0.8 V vs. Al/Al(III) was observed for all PEDOT films characterised in Lewis acidic, basic and neutral EMImCl-AlCl₃. The doping peaks in the Lewis neutral composition appear at 1.2 V, 1.7 V and 2.3 V vs. Al/Al(III) (Table 4).

The anodic peak at 1.2 V vs. Al/Al(III) describes a lower doping level, whereas the anodic peak at 1.7 V vs. Al/Al(III) shows an intermediate doping level. The increase of the energy level due to oxidation and the gradual formation of more doping points enables consequently a higher degree of anion doping. The doping peak at 2.3 V vs. Al/Al(III) is the maximum value for PEDOT in Lewis neutral EMImCl-AlCl₃. An even higher doping potential of PEDOT cannot be investigated in this solution. A further increase of the anodic potential would cause the decomposition (>2.6 V vs. Al/Al(III)) of the Lewis neutral ionic liquid and degradation of the PEDOT film. Furthermore, the formation of a higher degree of doping contributes also to higher conductivity of PEDOT, which can be observed as increasing peak current density at higher potentials.

The PEDOT film in Lewis basic EMImCl-AlCl₃ (Fig. 5c) is characterised by a wide cathodic peak at ≈ 0.8 V vs. Al/Al(III) with...
157 mA s transferred de-doping charge (Table 4). In contrast, the transferred doping charges are 29 mA s between around 0.8 V and 1.5 V vs. Al/Al(III). The cyclic voltammogram clearly indicates that the de-doping is dominant, pointing to an inefficient doping and de-doping of both the Cl\(^-\) ions of the Lewis basic ionic liquid and AlCl\(_4\) ions doped in the previously polymerised PEDOT film. The viscosity of the Lewis basic composition (\(\approx 306\) mPa s at 20.7 °C [21]) is 27 to 19 times higher than a Lewis acidic (\(\approx 11.5\) mPa s at 20.0 °C [21]) and neutral solution (\(\approx 16\) mPa s at 20.0 °C [21]). Therefore, the anion doping might be mass transport controlled. Furthermore, the PEDOT film shows a low stability at high anodic potentials. The decomposition starts already at 1.8 V vs. Al/Al(III).

3.2.1. Reversibility of PEDOT in Lewis neutral EMImCl-AlCl\(_3\)

The reversibility of the doping and de-doping reaction in the Lewis neutral composition was studied by cyclic voltammetry with 240 cycles (Fig. 6).

The cyclic voltammogram is characterised by two main anodic and cathodic peaks, whose current density decreases while the number of cycles increases. However, the reversibility of the doping and de-doping reaction (\(Q_c/Q_a\)) reached 97%. It is assumed, that the decrease of current density was caused by an initial structural change of the PEDOT film during doping and de-doping with AlCl\(_4\) [22,23]. The high reversibility shows that the doping anion AlCl\(_4\) does not remain trapped into the conductive polymer, which makes it suitable for a charge storage material in energy storage devices.

![Fig. 7. Cyclic voltammograms of PEDOT films polymerised from -0.5 V to (a) 1.5 V, (b) 2.0 V and (c) 2.6 V vs. Al/Al(III) in Lewis neutral EMImCl-AlCl\(_3\) with 0.1 mol dm\(^{-3}\) EDOT at 20 cycles. Cyclic voltammograms of PEDOT films polymerised to (d) 1.5 V, cycle 15, (e) 2.0 V, cycle 7 and (f) 2.6 V, cycle 15 in monomer-free Lewis neutral EMImCl-AlCl\(_3\) at 100 mV s\(^{-1}\) and 25 °C.](image-url)
3.2.2. Formation of doping and de-doping levels

The formation of the doping and de-doping levels of the polymer was studied by cycling and increasing the anodic polymerisation potential window in monomer-free Lewis neutral EMImCl-AlCl₃ from −0.5 V to 2.6 V vs. Al/Al(III). This experiment refers to the question of whether the doping and de-doping levels of the conductive polymer are predefined during the electropolymerisation or whether the levels are formed during cycling in the monomer-free ionic liquid electrolyte in the presence of doping anions.

The cyclic voltammograms of the electro-polymerisation obtained from −0.5 V to 1.5 V (Fig. 7 a), 2.0 V (Fig. 7 b) and 2.6 V (Fig. 7 c) vs. Al/Al(III) show an increasing amount of doping and de-doping levels. PEDOT polymerised from −0.5 V to 1.5 V vs. Al/Al(III) is characterised by one clear doping level at 0.85 V and de-doping level at 0.8 V vs. Al/Al(III), whereas PEDOT obtained from −0.5 V to 2.0 V vs. Al/Al(III) shows merged doping levels without clear peaks between 0.8 V and 1.8 V and de-doping levels between 1.6 V and 0.3 V vs. Al/Al(III). PEDOT polymerised in the widest polymerisation window from −0.5 V to 2.6 V vs. Al/Al(III) has four doping levels at 0.75 V, 1.1 V, 1.7 V and 2.2 V vs. Al/Al(III), which overlap each other and two clear de-doping levels at 1.6 V and 0.9 V vs. Al/Al(III).

The PEDOT film obtained from −0.5 V to 1.5 V vs. Al/Al(III) (Fig. 7a) shows, in monomer-free Lewis neutral EMImCl-AlCl₃, from −0.5 V to 2.6 V vs. Al/Al(III) (Fig. 7d) the same doping and de-doping levels observed already during polymerisation plus an additional doping wave at 1.75 V vs. Al/Al(III) (Table 5). The doping and de-doping levels in monomer-free Lewis neutral EMImCl-AlCl₃ of PEDOT polymerised from −0.5 V to 2.0 V vs. Al/Al(III) (Fig. 7e) has the same doping potential window comparable during polymerisation (Fig. 7b). However, the de-doping level appears as wide but clear peak at 0.9 V vs. Al/Al(III) in monomer-free ionic liquid (Table 5). The PEDOT film obtained from −0.5 V to 2.6 V vs. Al/Al(III) (Fig. 7c) shows three doping levels at 0.5 V, 1.5 V and 2 V as well as two de-doping peaks at 1.2 V and 0.8 V vs. Al/Al(III) (Fig. 7 f, Table 5).

The PEDOT films characterised in monomer-free Lewis neutral EMImCl-AlCl₃ show clearly that the doping and de-doping levels are predefined during electro-polymerisation. Doping or de-doping levels of PEDOT films in monomer-free ionic liquid, which are higher than the initial polymerisation potential window, were not observed. The characteristic doping and de-doping levels are similar to the doping and de-doping levels, which are formed during polymerisation in a certain polymerisation potential window. Additional doping levels are not formed during cycling in monomer-free Lewis neutral EMImCl-AlCl₃.

The formation of the doping and de-doping levels is in accordance with the generation of energetic conducting states of the polymer, which increase with the polymerisation potential window. Clear doping and de-doping levels were formed at polymerisation windows from −0.5 V to 1.5 and 2.6 V vs. Al/Al(III). The polymerisation potential window from −0.5 V to 2.0 V indicates the formation on an intermediate energetic level because of the merged wide doping and de-doping peak.

The polymerised PEDOT appear as dull dark blue-violet films on the vitreous carbon substrates. The SEM images of the PEDOT surfaces (Fig. 8) show porous structures of agglomerated granules (≤1 μm), which are similar for electro-polymerised PEDOT in ionic liquids reported in the literature [11,12,24].

| Polymerisation potential window /V | Doping level /V | De-doping level /V |
|-----------------------------------|----------------|------------------|
| −0.5 to 1.5                       | 0.85 | 1.75 | 0.8 |
| −0.5 to 2.0                       | 0.8 | 1.8 | 0.9 |
| −0.5 to 2.6                       | 0.5 | 1.5 | 2.0 | 0.8 | 1.2 |

Fig. 8. SEM images of PEDOT surfaces at (a,b,c) 1000-fold and (d,e,f) 5000-fold magnification obtained in Lewis neutral EMImCl-AlCl₃ with 0.1 mol dm⁻¹ EDOT from −0.5 V to (a,d) 1.5 V, (b,e) 2.0 V and (c,f) 2.6 V vs. Al/Al(III) at 100 mV s⁻¹, 20 cycles and 25 °C.
The polymerisation potential window has a clear influence on the density of the agglomerated granules. The PEDOT film obtained at the lowest polymerisation potential window from –0.5 V to 1.5 V vs. Al/Al(III) shows the lowest granule density (Fig. 8 a and d). The increase of the polymerisation potential window to 2.0 V (Fig. 8 b and e) and 2.6 V vs. Al/Al(III) (Fig. 8 c and f) causes an increase of the granule density.

The increase of grain density results from the growth of the chain length of PEDOT with a wider polymerisation potential window [1], forming a cross-linked network of polymer chains. It is assumed that a highly cross-linked polymer network generates more accessible doping levels for AlCl4–, which are accessible at higher potentials.

4. Conclusions

In this work, the electro-polymerisation of EDOT in EMImCl-AlCl3 ionic liquid with different acidities was studied. The polymerised PEDOT films were characterised by cyclic voltammetry in EDOT free Lewis acidic, basic and neutral EMImCl-AlCl3. It has been shown that:

(a) A Lewis basic solution decomposes at around 1.8 V vs. Al/Al(III) before EDOT can be polymerised. In a Lewis acidic solution, the excess of the oxidising compound AlCl3 causes an interaction with EDOT to a stable and highly viscous complex, which also prevents the electro-polymerisation. The electro-polymerisation of EDOT is only feasible in Lewis neutral EMImCl-AlCl3 with AlCl4– as the predominant doping anion.

(b) PEDOT films synthesised in Lewis neutral ionic liquid show different doping and de-doping behavior in monomer-free Lewis acidic, basic and neutral EMImCl-AlCl3. PEDOT films in a Lewis neutral composition describe a higher doping level due to the higher potential stability window over 2.6 V vs. Al/Al(III) than the Lewis acidic and basic solutions.

(c) The doping and de-doping reaction of AlCl4– into the PEDOT film reaches a reversibility of 97% in a Lewis neutral EMImCl-AlCl3.

(d) The doping and de-doping levels are predefined during electro-polymerisation and do not seem to form during cycling in ionic liquid with present doping anions.

(e) A wider polymerisation potential window generates a cross-linked surface structure and higher doping and de-doping levels.

As a general conclusion, the conductivity of the conductive polymer PEDOT as electrode material and a chloroaluminate ionic liquid as electrolyte has a potential use for rechargeable energy storage devices like batteries and capacitors with high stability, reversibility and high energetic performance.

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