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Shuttle effect quantification for redox ionic liquid electrolyte correlated to the coulombic efficiency of supercapacitors

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The use of redox active electrolyte is a good opportunity to increase the energy density of supercapacitors, the main limitation of this technology. The addition of redox molecules allows the storage of charge in the electrode and in the electrolyte. The key to keep the increase of charge is to avoid the shuttle effect of the redox molecule. Indeed, once the molecule is oxidized or reduced, it diffuses across the cell to react at the surface of the opposite electrode and the stored charge is lost. Is this shuttle effect however damageable for the device? This study proposes to answer this question by quantifying the shuttle effect and correlating it to the decrease of Coulombic efficiency of supercapacitors.

Introduction

In their field, supercapacitors have a lower energy density than batteries (20 times lower) but a much greater maximum power (10 times).[1] However, the search for devices with both high energy density and high maximum power remains an important challenge for all scientists in this field.

In the field of electrochemical double layer capacitor (EDLC), the state of the art offers an alternative based on redox molecules dissolved in the electrolyte to enhance the storage of electrical charges. As far as we know, the pioneering work is that of Frackowiak et al. devoted on the development of a redox active aqueous electrolyte with potassium iodide (KI) and vanadyl sulfate (VOSO$_4$).[2] Device performance was great but the high-cost ion selective membrane (Nafion) used to avoid the crossover of redox species somehow tempered the success of this proof of concept. More recently, Boettcher et al. investigated redox-based EDLC supercapacitor using viologen and halides couples in an aqueous electrolyte for the negative and the positive electrodes, respectively.[3] After testing various halides and viologens, reported performance was beyond the state-of-the-art, including a low self-discharge despite the absence of any ion-selective membrane.

The redox active electrolyte approach, although very attractive, has two identified limitations: (1) the self-discharge is usually strong and (2) shuttle effects affect the performances.[4] Redox ionic liquids are good opportunities to address these two issues.[5] Some of us were the first to develop redox ionic liquids while not considering them for operation in devices. It is only from the work of Rochefort et al. that the first applications of redox ionic liquids are considered for batteries and supercapacitors.[6]

However, depending on the nature of the redox ionic liquid, the shuttle effect may still remain, especially if the molecule has no particular adsorption or confinement within the porosity of the electrode material. In such a case, the scientific question is: how many times does the redox shuttle have to go back and forth before highly impacting the coulombic efficiency? To answer this question, we propose the study of the shuttle effect of a redox ionic liquid, 4-methylimidazolium-2,2,6,6-tetramethylpiperidine 1-oxyl (TEMPOIm)$^-$ TFSI$^-$ in this study, the number of shuttles of the redox molecule is correlated to the decrease of coulombic efficiency of the device. To calculate the number of shuttles, the Fick’s law is used, which means that the redox molecule diffuses linearly in a continuous medium. First, the diffusion of TEMPO moieties is studied to confirm the diffusional behaviour of the molecules. Then the diffusion is correlated with the properties of the electrolyte that depend on its viscosity and ionic conductivity.
Results and Discussion

Diffusion coefficient of TEMPO moieties

(TEMPOIm)- TFSI- is electrochemically characterized and compared to commercial TEMPO. Figure 2 exhibits the cyclic voltammetry (CV) profiles of these molecules solubilized in NEt4+BF4- at 0.5 mol·l-1 in acetonitrile (Figure 2a) and in BMIm+-TFSI- ionic liquid (Figure 2c). In both electrolytes, the TEMPO (black line) is oxidized as TEMPO+ at the potential of 0.29 V vs Fc/Fc+, then reduced back as TEMPO at 0.22 V vs Fc/Fc+ by a reversible single electron transfer step. Similarly, the (TEMPOIm)2+ TFSI- at 0.33 V vs Fc/Fc+ and reduced back as (TEMPOIm)+ TFSI- at 0.26 V vs Fc/Fc+. The difference of redox potential between TEMPO and (TEMPOIm)+ TFSI- about 0.04V could be explained by an impact of the imidazolium grafted on the TEMPO. The imidazolium is positively charge so the addition of a second positive charge by the oxidation of (TEMPOIm)+ TFSI- needs a greater energy leading to a higher potential.

CV shapes are characteristics of the diffusional behavior of the redox molecules.\[7\] The potential difference between current peaks of oxidation and reduction, ΔEpp, is 70 mV for all CVs, proving the reversibility of these redox systems.\[8\] Figures 2b and 2d compare the values of current peak in oxidation depending on the scan rate, for NEt4+BF4- at 0.5 mol·l-1 in acetonitrile and BMIm+-TFSI- respectively, with TEMPO (plain square) and (TEMPOIm)+ TFSI- (open square). Current values form linear lines between 2.5·10-2 V·s-1 and 1 V·s-1. The apparent diffusion coefficient, Dpp, proportional to the line slope, is greater for TEMPO compared to (TEMPOIm)+ TFSI- in both electrolytes. This discrepancy can be assigned to molecule size difference. TEMPO is a smaller molecule than (TEMPOIm)+ TFSI-, and, as such, diffuses faster than the larger one.\[9\]
The size of the molecule is not the sole factor influencing the diffusion coefficient of redox molecules. Indeed, the physico-chemical properties of the electrolyte (conductivity, viscosity) are critical parameters for characterizing the diffusion properties of redox molecules.

**Influence of viscosity and molar conductivity on the apparent diffusion coefficient**

To characterize the properties of bulk electrolytes, five concentrations of BMIm⁺ TFSI⁻ were chosen from 1 mol·l⁻¹ in acetonitrile to pure ionic liquid (which corresponds to 3.4 mol·l⁻¹). The concentration of (TEMPOIm)⁺ TFSI⁻ was increased (compared to 2·10⁻³ mol·l⁻¹ previously) to observe the Faradaic contribution of the redox ionic liquid in the device. All solutions include (TEMPOIm)⁺ TFSI⁻ at 0.1 mol·l⁻¹.

To characterize these electrolytes, an ultramicroelectrode (UME) is used as working electrode. This electrode has a much smaller diameter than an usual macroelectrode (10 µm in our case against 3 mm). With UME, the EDLC contribution is drastically reduced and does not interfere with measured signal. Then, at the potential of the redox couple, an increase in current is visible up to a plateau due to the steady state (all species in the diffusion layer have reacted), Figure 3a. From this plateau, the limit current is extracted for each concentration of BMIm⁺ TFSI⁻ and used to calculate the apparent diffusion coefficient of the (TEMPOIm)⁺ using the Stokes-Einstein equation (Eq. 1).

\[ i_l = 4 \cdot z \cdot F \cdot D_{app} \cdot C \cdot r \]  

(Eq. 1)

\( n \) corresponds to the number of electrons exchanged, \( F \) is the Faraday constant in C·mol⁻¹, \( D_{app} \) is the apparent diffusion coefficient in m²·s⁻¹, \( C \) represents the concentration of redox ionic liquid in the electrolyte in mol·m⁻³ and \( r \) is the radius of the ultramicroelectrode in m.

**Figure 3.** (a) CV with Pt UME (Ø 10 µm) at 0.1 V·s⁻¹, a Pt wire as counter electrode and a silver wire as reference. The electrolytes are composed of (TEMPOIm)⁺ TFSI⁻ at 0.1 mol·l⁻¹ with various concentrations of BMIm⁺ TFSI⁻ in acetonitrile. (b) Apparent diffusion coefficient (solid squares) and molar ionic conductivity (open circle) depending on the BMIm⁺ TFSI⁻ concentration. (c) Apparent diffusion coefficient and viscosity of the solutions (white circle) depending on the BMIm⁺ TFSI⁻ concentration. (d) Apparent diffusion coefficient as a function of the inverse of viscosity.
The Figure 3a shows the CV measured with a platinum UME at 0.1 V·s⁻¹ for each electrolyte. The current is zero for the applied potential from 0.2 V to 0.6 V vs Ag/Ag⁺ for tested solutions. This potential zone corresponds to the formation of the electrochemical double layer. Then, in the whole electrolyte series, the current increases with a significant slope between 0.6 V and 0.8 V vs Ag/Ag⁺. This step corresponds to the electronic transfer during the oxidation of (TEMPOIm)⁺. Above 0.8 V vs Ag/Ag⁺, the slope is decreasing and the curve could be considered as a plateau. It corresponds to the mass transfer of redox molecules to the surface of the UME. Since the current is not quite constant, the regime is not fully stationary. Limit current of the plateau (measured at the potential of 1.3 V vs Ag/Ag⁺) increases when BMIm⁺ TFSI⁻ concentration decreases. Apparent diffusion coefficients, D_app, were calculated from these limit current values using (Eq. 1). As such, the apparent diffusion coefficient decreases while increasing BMIm⁺ TFSI⁻ concentration, as shown in Figure 3b. Figure 3b and Figure 3c depicts the diffusion coefficients as a function of the BMIm⁺ TFSI⁻ concentration. Figure 3b also shows that D_app and the molar conductivity (open squares) which decrease the same way as the salt concentration increases. Figure 3c compares D_app with the viscosity of the electrolyte. The viscosity gradually increases from 0.7 and 4.6 mPa·s for diluted solutions with salt concentrations from 1 to 2.5 mol·l⁻¹). The viscosity for pure BMIm⁺ TFSI⁻ is 44 mPa·s. When the diffusion coefficient is expressed as a function of the inverse of viscosity, Figure 3d, a straight line with a slope of 3.6·10⁻⁷ cm²·mPa⁻¹·s is obtained. The diffusion coefficient is therefore correlated with both viscosity and molar conductivity. Table 1 summarizes the physical characteristics of each BMIm⁺ TFSI⁻ solutions depending on its concentration. The solvation radius (r_solv) is calculated from apparent diffusion coefficient values determined at corresponding limit current values (UME), (Eq. 2), where k_ө is the Boltzmann constant and T (K) the temperature.[12]

\[ r_{solv} = \frac{1}{k_B T \cdot \pi \cdot n \cdot \eta} \]  

(Eq. 2)

Overall, the value of the r_solv decreases with the diffusion coefficient and the increase in viscosity. However, the order of magnitude of the solvation radius is the same (about 4 nm).

In this section, the critical correlation between diffusion coefficient and viscosity is highlighted. The next section is about the charge storage with respect to the characteristics of prepared electrolytes. The objective is to determine if one of them is leading to better charge storage (in the sense of a larger quantity of charges). For this purpose, the parameter studied is the electrical charge Q normalized by the electrode mass.

### Impact of the diffusion coefficient of (TEMPOIm)⁺ TFSI⁻ on the electrical charge of a porous electrode

The electrical charge of the working electrode is measured in a three-electrode cell to limit the redox shuttle effect by applying a large distance between the working electrode and the counter electrode to measure the impact of the physical and chemical properties of the electrolyte on the electrical charge. The study of redox shuttles is reported in the next section using two-electrode cell with controlled thickness separating the electrodes. CVs are performed under inert atmosphere with a cell composed of self-supporting porous carbon electrodes. The counter electrode has a loading at least three times greater than that of the working electrode.

<table>
<thead>
<tr>
<th>[BMIm⁺ TFSI⁻]</th>
<th>[BMIm⁺ TFSI⁻] (%wt)</th>
<th>D_app (cm²·s⁻¹)</th>
<th>η_app (mPa·s)</th>
<th>Λ_app (S·m²·mol⁻¹)</th>
<th>r_solv (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 mol.1⁻¹</td>
<td>30</td>
<td>5.1·10⁻⁷</td>
<td>0.75</td>
<td>4.0·10⁻¹</td>
<td>5.6</td>
</tr>
<tr>
<td>1.5 mol.1⁻¹</td>
<td>45</td>
<td>3.0·10⁻⁷</td>
<td>1.2</td>
<td>2.6·10⁻¹</td>
<td>5.9</td>
</tr>
<tr>
<td>2 mol.1⁻¹</td>
<td>60</td>
<td>2.3·10⁻⁷</td>
<td>2.1</td>
<td>1.7·10⁻¹</td>
<td>4.5</td>
</tr>
<tr>
<td>2.5 mol.1⁻¹</td>
<td>75</td>
<td>1.1·10⁻⁷</td>
<td>4.6</td>
<td>8.3·10⁻¹</td>
<td>4.1</td>
</tr>
<tr>
<td>3.4 mol.1⁻¹ (pur)</td>
<td>100</td>
<td>1.6·10⁻⁷</td>
<td>44</td>
<td>1.2·10⁻¹</td>
<td>3.1</td>
</tr>
</tbody>
</table>

[a] Measurement at 20 °C.
Figure 4. (a) CVs with porous carbon based electrodes and a silver wire as reference. The supporting electrolyte is BMIm\textsuperscript{+} TFSI\textsuperscript{-} at 1 mol·l\textsuperscript{-1} in acetonitrile (dashed line) and with the addition of 0.1 mol·l\textsuperscript{-1} of (TEMPOIm)\textsuperscript{+} TFSI\textsuperscript{-} (green line). (b) CVs with porous carbon based electrodes with various concentrations of BMIm\textsuperscript{+} TFSI\textsuperscript{-} in acetonitrile. The electrolytes contain (TEMPOIm)\textsuperscript{+} TFSI\textsuperscript{-} at 0.1 mol·l\textsuperscript{-1}. (c) Apparent diffusion coefficient (solid squares) and electrical charge (open circle) depending on the BMIm\textsuperscript{+} TFSI\textsuperscript{-} concentration. (d) Viscosity (solid squares) and electrical charge (open circles) of the solutions depending on the BMIm\textsuperscript{+} TFSI\textsuperscript{-} concentration.

Figure 4a details the CV measured in 1 mol·l\textsuperscript{-1} of BMIm\textsuperscript{+} TFSI\textsuperscript{-} in acetonitrile (dotted black) electrolyte before and after addition of 0.1 mol·l\textsuperscript{-1} of (TEMPOIm)\textsuperscript{+} TFSI\textsuperscript{-} (light green). The Faradaic contribution (0.6 V to 1.2 V vs Ag/Ag\textsuperscript{+}) comes along with the capacitive contribution (0 V to 0.6 V vs Ag/Ag\textsuperscript{+}). This superposition is supported by considering the purely capacitive curve when no redox molecule is present in the solution (dashed black). An increase of 0.7 mA is observed thanks to the Faradaic contribution. CVs corresponding to (TEMPOIm)\textsuperscript{+} TFSI\textsuperscript{-} at 0.1 mol·l\textsuperscript{-1} solutions with the various BMIm\textsuperscript{+} TFSI\textsuperscript{-} concentrations are shown in Figure 4b. The current is normalized by the mass of active material in the working electrode. At concentrations of 1 and 1.5 mol·l\textsuperscript{-1} of BMIm\textsuperscript{+} TFSI\textsuperscript{-}, CVs are almost identical and the Faradaic contribution is supposed to be the same. A slight shift in redox potential about 60 mV is observed for the electrolyte at 2 mol·l\textsuperscript{-1} (blue curve), Figure 4b. The fact that there is only one curve with a redox potential shift suggest a shift of the reference Ag/Ag\textsuperscript{+} during the measure. On the other hand, when the concentration of BMIm\textsuperscript{+} TFSI\textsuperscript{-} was increased, the current peak and the peak-to-peak separation corresponding to the Faradaic contribution decreased. For pure ionic liquid, no current peak is visible (Figure 4a dashed curve).

The charge calculated from CV measurements in the three-electrode cell is plotted in Figure 4c as a function of BMIm\textsuperscript{+} TFSI\textsuperscript{-} concentration and compared to the apparent diffusion coefficient. The observed charge and the diffusion coefficient decrease while BMIm\textsuperscript{+} TFSI\textsuperscript{-} concentration increases that corresponds to an increase in electrolyte viscosity (Figure 4d).

The Faradaic contribution in the most viscous electrolyte (BMIm\textsuperscript{+} TFSI\textsuperscript{-} containing (TEMPOIm)\textsuperscript{+} TFSI\textsuperscript{-}) is almost zero. However, it is in this electrolyte that the diffusion coefficient is the lowest, and therefore also the one for which the redox shuttle effect should be the lowest. The effect of the redox shuttle can be quantified by calculating the number of shuttles back and forth between the two opposite electrodes. The objective is to correlate the number of shuttle back and forth made by the redox active ion with the coulombic efficiency of the device. It should allow to estimate the charge loss by the device because of these shuttles.

Quantification of the number of shuttles of the (TEMPOIm)\textsuperscript{+} and its impact on coulombic efficiency

It has been shown that the redox molecule (TEMPOIm)\textsuperscript{+} has a diffusional behavior (Figure 2). When the molecule is oxidized at a concentration gradient of oxidized and reduced molecules is established from the electrode-electrolyte interface toward the electrolyte bulk. This concentration gradient is limited to a certain distance from the electrode assigned as the diffusion layer δ. The time needed for molecules to diffuse from
one electrode to the other one can be calculated. In this case, the diffuse layer corresponds to the distance between the electrodes i.e. the thickness of the separator. From the diffusion coefficient of the redox molecule and the thickness of the separator, the time, $t_1$, to diffuse from one side of electrode to the other side is calculated with the (Eq. 3):

$$
\delta = \sqrt{\pi \cdot D_{app} \cdot t_1}
$$

(Eq. 3)

Knowing the time of diffusion from one electrode to the other, the number of shuttle back and forth made by the redox cation (TEMPOIm)$^+\,$ can be calculated and correlated to the performance of the device. Among the main performance indicators for a supercapacitor are Coulombic efficiency, power, energy, energy efficiency and lifetime. Coulombic efficiency is particularly affected by the redox shuttle effect, so it is on this parameter that we focused. The setup used is a Swagelok cell with two porous carbon electrodes and a separator in between of a defined thickness about 130 µm.

**Figure 5.** (a) The time of a single back and forth shuttle is calculated according to the diffusion coefficient (log-log scale) for various separator thicknesses. The orange rectangle illustrates the value scales for laboratory systems of diffusion coefficients and separator sizes used.\(^{[13]}\) (b) Electrolyte diffusion coefficient as a function of BMIm$^+$ TFSI$^-$ concentration diluted in acetonitrile with respect to the corresponding time for a single shuttle (one back and forth) of the redox molecules, calculated for a separator thickness of 130 µm (experimental value).

The relationship between the diffusion time, $t_2$ ($t_2 = 2 \times t_1$), required for a back and forth shuttle of the redox molecules and the apparent diffusion coefficient is depicted on Figure 5a as calculated from Eq. 2. It shows in log-log scale the time of a single back and forth shuttle depending on diffusion coefficient values and separator thicknesses. For example, for a diffusion coefficient of $10^{-8}$ cm$^2$·s$^{-1}$ and a separator of 130 µm, the time for a back and forth shuttle of the molecule between the electrodes is $1.10^{-4}$ s ($t_2 = 2 \times (130 \times 10^{-4})^2 / (\pi \times 10^{-8})$). As the relationship (Eq. 2) shows, the time for a back and forth shuttle is inversely proportional to the diffusion coefficient. The thicker the separator, (i.e. the distance between the two electrodes), the longer the time required for a back and forth shuttle. In this figure, the orange rectangle corresponds to the usual thicknesses of separators used for supercapacitors: from 25 µm (Celgard) to 500 µm (Whatman glass fiber) and typical values of diffusion coefficient observed for a redox molecule (as an example for TEMPO) in ionic liquid media such as BMIm$^+$ TFSI$^-$ or solvent such as ACN.\(^{[13-14]}\) Figure 5b shows the time for a back and forth shuttle through a 130 µm thick separator versus diffusion coefficients in the usually reported range. The greater the BMIm$^+$ TFSI$^-$ concentration, the longer the diffusion time. For pure BMIm$^+$ TFSI$^-$, a calculated duration of almost 7000 s is out of scope when using a supercapacitor device.

By performing galvanostatic measurements, it is possible to distinguish the Faradaic contribution from the purely capacitive contribution and to extract a characteristic time from it. This time corresponds to the total time of diffusion of the redox molecule $t_c$. From this time and the time required for a single back and forth shuttle of a redox molecule, $t_2$, the number of back and forth shuttle of the molecule can be calculated. Calculation details are given in the supplementary information section. This number of shuttles has been correlated to the Coulombic efficiency of the device.
Correlation between the Coulombic efficiency of a supercapacitor and the number of back and forth shuttles of a redox shuttle

To determine the impact of the redox shuttle effect on the performance of a supercapacitor, galvanostatic measurements were conducted using a two-electrode Swagelok device. For this purpose, porous carbon electrodes coated on an aluminum current collector (see details in experimental part) and a 130 µm thick cellulose separator were used. Measurements in galvanostatic mode were performed for currents between 1.5 A·g⁻¹ and 0.01 A·g⁻¹. Figure 6 shows the galvanostatic curves for current densities at 0.5 A·g⁻¹, 0.05 A·g⁻¹, 0.02 A·g⁻¹ and 0.01 A·g⁻¹. The total diffusion time, \( t_c \), considered for calculating the number of back and forth shuttles of the redox ion is the charging time between 1.4 V and 2.8 V, Figure 7a. The voltage of 1.4 V corresponds to the change in slope on the galvanostatic curves, the Figure 7a is given as an example of it. The number of redox shuttles is determined and reported in Figure 7b for the BMIm⁺ TFSI⁻ electrolyte diluted to 1 mol·l⁻¹. In this electrolyte, a single shuttle is observed for an applied current density of 0.2 A·g⁻¹. The applied current is quite low for operating a supercapacitor: the tests reported in the literature can go up to more than 10 A·g⁻¹, however for such high currents, Faradaic reactions do not have time to occur in the present case. For a current density of 0.03 A·g⁻¹, the number of calculated shuttles is 11.
Figure 7. (a) and (b) Galvanostatic measurements for a device with two porous carbon electrodes. The electrolyte is containing (TEMPOIm)$^+$ TFSI at 0.1 mol·l$^{-1}$ in BMIm$^+$ TFSI at 1 mol·l$^{-1}$ in acetonitrile with current densities of (a) 0.03 A·g$^{-1}$ and (b) 0.03, 0.05, 0.07, 0.1 and 0.2 A·g$^{-1}$. (c) and (d) Coulombic efficiency associated to the galvanostatic measurements (c) with and (d) without (TEMPOIm)$^+$ TFSI$^-$ at 0.1 mol·l$^{-1}$.

Coulombic efficiency is the ratio of discharge time to charging time. Less than 100% means that a part of the charges stored are not recovered during discharge. Coulombic efficiencies of the devices are shown as a function of the applied current in Figure 7 for electrolytes with different concentrations of BMIm$^+$ TFSI$^-$ (7d : “without redox”) and with the addition of 0.1 mol·l$^{-1}$ of (TEMPOIm)$^+$ TFSI$^-$ (7c). The loss of Coulombic efficiency begins at greater currents in the presence of redox molecules, 70% at 0.06 A·g$^{-1}$ for the 1 mol·l$^{-1}$ BMIm$^+$ TFSI$^-$ electrolyte, Figure 7d, than without redox molecule, 73% at 0.03 A·g$^{-1}$ for the 1.5 mol·l$^{-1}$ BMIm$^+$ TFSI$^-$ electrolyte. The addition of (TEMPOIm)$^+$ TFSI$^-$ speed-up the Coulombic efficiency fading.

For all applied currents corresponding to 4 shuttles, the Coulombic efficiency dropped down to 70 % ± 5 %. This means that a 30 % loss of Coulombic efficiency was observed after 4 shuttles of the redox molecules. This observation could be extended to any redox molecule with diffusional behavior (no adsorption at the electrode surface as an example).

To go further, the different impacts of the shuttle effect on the device were categorizing: (i) protection (ii) degradation (iii) or competition favourable or (iv) unfavourable to the device. These impacts are depicted on Figure 7. The term protection is used to mean that the coulombic efficiency of the device is not degraded at all with the addition of redox ionic liquid compare to a degradation with redox free electrolyte (CE$\text{redox} = 100\%$). The redox ionic liquid provides a full protection by avoiding degradation of electrolyte at same potential. The second case presented is the opposite: the addition of redox shuttle degrades the electrolyte but no degradation is observed with redox free electrolyte (CE$\text{redox} = 100\%$). When these two phenomena are observed, there is a competition. If the coulombic efficiency is better with the addition of redox ionic liquid, what means there is degradation of the electrolyte but the RIL protects in part the device (CE$\text{redox} < 100\%$ and CE$\text{redox} \gg$ CE$\text{no redox}$). In the other case, the RIL precipitates the degradation of the RIL, a situation to avoid (CE$\text{no redox} < 100\%$ and CE$\text{redox} \ll$ CE$\text{no redox}$). The last case that could be observed is no effect of the addition of RIL (CE$\text{redox} \approx$ CE$\text{no redox}$), the coulombic efficiency is similar with and without RIL. Statistically speaking, our case is more corresponding to a decrease of the coulombic efficiency with the addition of RIL but with also a degradation of the electrolyte (18 measures on 31 in totality). So far, the redox shuttle is damaging the device in our conditions. One step further is to find which conditions are needed to protect the device with the addition of RIL. The protection of overcharge by using RIL has been study for batteries$^{[17]}$ but could be a great advantage also for the use of supercapacitor containing electroactive species.
Figure 8. Classification of different impacts of the redox shuttle on the coulombic efficiency (CE) of supercapacitor and statistics of measurements realized. The proportion \((y/x)\) on the left are: \((x)\) the number of coulombic efficiencies calculated for different current density and \((y)\) electrolyte and the number of CE corresponding to the aligned case.

Conclusions

In this study, the TEMPO molecule functionalized with an imidazolium ion has a diffusional behavior leading to a loss of stored redox charges because of a shuttle effect between electrodes during the charge of the device. The impact of the shuttle effect on the Coulombic efficiency is quantifiable. In this case, 4 shuttles result in a 30% loss of Coulombic efficiency.

This observation can be generalized. The redox shuttle effect is highly dependent on the diffusion coefficient of the redox molecule. The diffusion coefficient reflects the viscosity and molar conductivity of the electrolyte. It is therefore useful to determine the conditions needed to limit the redox shuttle effect and the loss of Coulombic efficiency. This also allows applying our conclusions to the use of other redox molecules and other supporting electrolytes as long as there are no specific interactions between the molecule and the electrode. The charging (or diffusion) times given are for a diffusion/separator size layer of 130 \(\mu\)m.

The use of a viscous medium such as BMIm\(^+\)TFSI avoids the redox shuttle effect. The diffusion coefficient is limited \((10^{-8} \text{ cm}^2\cdot\text{s}^{-1})\) which generates a very long diffusion time \(6840\) s for a distance of 130 \(\mu\)m. This time constant is way too long to be realistic in a supercapacitor and any redox shuttle effect will never be encountered. However, the use of a viscous electrolyte is detrimental to the power of the device.

A diffusion coefficient of \(10^{-7} \text{ cm}^2\cdot\text{s}^{-1}\) is a good compromise, the charging times are faster than in the present case and the current density limit is 0.03 A·g\(^{-1}\) to avoid the Coulombic efficiency fading. For diffusion coefficients from \(2\cdot10^{-7}\) to \(5\cdot10^{-7}\) cm\(^2\)·s\(^{-1}\), it is necessary to avoid slow cycling (current density < 0.08 A·g\(^{-1}\)) which would allow the diffusion of redox molecules and the loss in Coulombic efficiency.

Experimental Section

Synthesis of ionic liquids

The 4-hydroxy-TEMPO, bromotrimethylsilane (TMSBr) and methylimidazole (MIm) were provided by Sigma Aldrich. The precursors 1-butyl-3-methylimidazolium chloride and lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) supplied by Iolitec. For the synthesis of 1-methyl-3-imidazolium-TEMPO bis(trifluoromethanesulfonyl)imide, Figure 1, 4-hydroxy-TEMPO was dissolved in a minimum volume of anhydrous acetone under argon atmosphere. The bromotrimethylsilane (5 eq.) was added drop per drop. The mixture was stirring during 24 h at room temperature. The excess of bromotrimethylsilane was removed as well as the sub-product trimethylsilanol under vacuum. The obtained product was a brown batter. It was solubilized in dichloromethane before the addition of the methylimidazole. The slurry was stirring during 24 h at 45 °C. The solvent was removed with rotating evaporator. Afterwards, the metathesis was performed by solubilizing separately the obtained product and LiTFSI (1.2 eq.) in water. The two solutions were mixed and stirred during 2h at room temperature. Then, liquid-liquid extraction was performed to collect the organic phase. The aqueous phase was washed with dichloromethane three times. The organic layer was dried over magnesium sulfate and the solvent was
removed with rotating evaporator. The obtained product was a little viscous red liquid. The \((\text{TEMPOIm})^+\) \(\text{TFSI}^-\) is characterized by DSC, TGA and IR in supplementary information.

The butylmethylimidazolium bis(trifluoromethanesulfonyl)imide were also synthesized. The components are separately dissolved in water then mixed and stirred during 12 h. The result ionic liquid is extracted with dichloromethane. The organic phase is washed with pure water three times to remove the sub product LiCl and dried with magnesium sulfate before removing the solvent with a rotary evaporator. The obtained product is still a transparent liquid.

**Conductivity and viscosity measurement**

To determine the molar ionic conductivity, a conductivity cell with a cell constant is used to measure the impedance of each solution. The cell constant, \(k\), is 1.8 cm\(^{-1}\). To determine the ionic conductivity, \(\sigma\), in S·cm\(^{-1}\), of the solutions, the calculation \(\sigma = \frac{k}{R}\) is carried out with the electrical resistance \(R\) (Ohms) extracted from the impedance measurements. The molar conductivity \(\Lambda\) corresponds to the ionic conductivity divided by the concentration of the solution. All measurements are made in a glove box at 20 °C with an Ametek Princeton Applied Research potentiotstat.

The viscosity of the solutions is measured with a Couette rheometer (cylinder) at a controlled temperature of 20 °C. The viscosity measurements correspond to an average of 4 measurements made for the shear gradients of 10, 20, 40 and 50 s\(^{-1}\).

**Electrochemical characterization**

Electrochemical characterizations with three-electrode cell were performed in a glove box using an Ametek Princeton Applied Research potentiostat. The working electrode used was glassy carbon, the counter electrode was a platinum rod and the reference electrode was a silver wire. The working electrode was polished with alumina 0.3 µm and cleaned with water then ethanol before each experiment. The apparent diffusion coefficient was determined by using ultramicroelectrode (UME) in platinum with a diameter of 10 µm. The reference and counter electrode were a silver wire and a platinum wire respectively. The UME was polished with alumina 0.3 µm then rinsed with ultrapure water and ethanol before each experiment. The cyclic voltammetry were performed at 100 mV·s\(^{-1}\) with a sweep potential from 0.2 V to 1.3 V vs Ag/Ag\(^+\).

For cyclic voltammetry presented figure 3, three-electrode cell composed of self-supporting porous carbon electrodes were used. The electrodes were constituted of PICACTIF as activated material, acetylene black and polytetrafluoroethylene (60 wt % dispersion in water) as binder with a ratio of 75:15:10 respectively. Firstly, powders were dispersed in ethanol, the polytetrafluoroethylene was added. The ethanol was removed at 50 °C, then the mixture is spread out on a glass plate until the thickness reach 150 µm. Diameter of the self-supported film (electrodes) was 6 mm for the working electrode and 10 mm for the counter electrode. Electrodes were pressed into a stainless steel grid at 10 T.

For device experimentation, a two-electrode Swagelok was used with porous carbon electrode. The electrodes were constituted of PICACTIF as activated material, acetylene black and polyvinylidene fluoride as binder with a ratio of 75:15:10 respectively. Firstly, an ink was formulated by dissolving the polymer PVdF in NMP during 1 h at room temperature. Then the two carbons were added and the ink was stirred during 1 h at room temperature followed by 1 h in ultrasonic bath. Secondly, the ink was coated on a current collector. For the working electrode, the ink was coated on aluminium foil with a doctor blade at 200 µm. The dried thickness is about 60 µm (± 10µm). For the counter electrode, the ink was coated on copper foil with a doctor blade at 500 µm. The dried thickness is about 130 µm (± 10µm). Finally, the coating was dried at 90 °C during 12 h. The coating was punched at a diameter of 9.5 mm to make electrodes. The separator used is Whatmann cellulose paper previously dried under vacuum at 80 °C during 12 h. The thickness of the separator is about 130 µm.
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