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Surface modification of TiO$_2$ by ionic liquid electrolyte in dye-sensitized solar cells using a molecular insulator

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A factor that limits the performance of dye-sensitized solar cell (DSC) based on ionic liquid is the disappearance of electron density due to electron recombination at the Dyed-TiO$_2$/electrolyte interface. Adding an additive to the electrolyte can affect processes related to kinetics of the electron recombination. In this paper a novel solid state additive, 1,2-dimethylimidazolium-3-propylsulfonate (DMImBS), was synthesized by autoclave and directly applied to a DSC based on an ionic liquid electrolyte to form a molecular insulator layer. It was found that the presence of DMImBS in the ionic liquid electrolyte enhances both the open circuit photovoltage ($V_{oc}$) and fill factor ($FF$) without significant penalty in photocurrent ($J_{sc}$), as well significantly increasing the power output of the DSC. This is suggested to be mainly due to the passivation of surface states, which effectively reduces electron recombination and restrain the Li$^+$ intercalation. The device based on DMImBS also showed outstanding stability under prolonged light soaking at 50 °C, maintaining 100% photovoltaic efficiency.

1. Introduction

Dye sensitized solar cells (DSC) attract immense attention in scientific research and industrial applications as low production cost solar cells$^{1-4}$. Three components are generally present in DSC: a nanocrystalline mesoporous TiO$_2$ thin film coated on a conducting glass substrate with a monolayer of dye as a sensitizer, an electrolyte containing $I^-/I_3^-$ and a platinum counter electrode. Electrons generated by photo-excited dye molecules are injected into the conduction band of the TiO$_2$ and transported from the injection sites to the contact electrode under light irradiation. Finally, electrons are collected at the transparent conducting glass substrate interface and then pass through the external circuit. It is believed that electron transfer at the Dyed-TiO$_2$/electrolyte interfaces and redox charge transfer at the counter electrode plays an important role in determining the photovoltaic performances of DSC$^5$. Additives in the electrolyte can affect processes related to kinetics of the electron transport and recombination, for this reason studies have been made on additives to improve the DSC efficiency and stability$^6$$^{12}$. Electron transfer dynamics at the Dyed-TiO$_2$/electrolyte interface is the key factor effecting the short circuit photocurrent ($J_{sc}$) and open circuit photo voltage ($V_{oc}$), which is related to photoelectric conversion efficiency ($\eta$)$^{13}$$^{17}$. Various additives have been added to the electrolyte solution, such as 4-tert-butylpyridine (TBP)$^{18}$$^{19}$, N-methylbenzimidazol (NMBI), benzimidazol (BI)$^{20}$$^{21}$ and lithium iodide (LiI)$^{22}$$^{23}$, to improve the device efficiency. To the best of our knowledge, adsorption of TBP, NMBI or BI on the TiO$_2$ surface can elevate the conduction band edge ($V_{CB}$) of TiO$_2$ to increase $V_{oc}$$^{24}$. Raising $V_{CB}$ would also decrease the electron injection efficiency ($\eta_i$).
and explain the reduced $J_{sc}$ observed in DSC using these additives. The electron transfer is improved by laying Li$^+$ in the TiO$_2$ crystal. In addition, the presence of LiI causes a positive shift in the $V_{CB}$ of TiO$_2$ which explains the increase in $J_{sc}$ and decrease in $V_{OC}$.

Electrolyte, as the redox mediators transport layer, is an important component in DSC. Organic solvents, such as γ-butyrolactone, acetonitrile and 3-methoxypropionitrile, are conventionally used as DSC electrolyte solvents but they are usually poisonous and volatile, which results in the restriction of DSC industrialization. The air and water stable room temperature liquid solvents (RTILs) are attractive as solvents due to their chemical and thermal stability, negligible vapor pressure, nonflammability, high ionic conductivity and wide electrochemical window$^{28}$. Furthermore, the physical and chemical properties of RTILs can be changed by varying the nature of cations and anions. During past decades, ionic liquid electrolytes based on imidazolium compounds have been applied in DSC but $\eta$ is still lower than when organic solvent electrolytes are used. Some works reported that electron recombination is more apparent using ionic liquid electrolyte compared to using organic solvent electrolyte$^{29,30}$; however it is necessary to find a more effective additive to restrain the electron recombination in DSC based on ionic liquid electrolyte

Recently, large efforts have been made to select and prepare new materials in order to achieve higher performance in DSC. In this paper, we report on the characterization of a zwitterionic imidazolium propylsulfonate, 1,2-dimethylimidazolium-3-propylsulfonate (DMImBS), as a novel solid state additive in 1-methyl-3-propylimidazolium iodide (MPII) based electrolyte. The alkyl chain of DMImBS is suggested to form the insulation layer which restrains the electron recombination and intercalation of Li$^+$. The negligible vapor pressure of DMImBS as a solid material is good for the stability of DSC.

2. Experimental

2.1 Materials

Anhydrous lithium iodide, 1,2-dimethylimidazole, and 1,4-butanesultone were purchased from Aldrich and used as received. The synthesis of zwitterionic-type salt having an imidazolium cation is shown below in scheme 1$^{31,32}$. Their structures were confirmed by 1H-NMR spectroscopy (Bruker DMX 400 spectrometer using D$_2$O as solvents).

![Scheme 1 Synthesis and characterization of 1,2-dimethylimidazolium-3-propylsulfonate (DMImBS)](image)

The mixture of 1,2-dimethylimidazole (9.61 g, 0.1mol) and 1,4-butanesultone (13.62 g, 0.1mol) was placed in a 60 mL Teflon-lined, stainless steel autoclave, and was heated in an oven at 100 °C for 7 h. The resulting DMImBS was decanted and dried under vacuum at 70 °C for 6 h. Crude products were washed with 1,1,1-trichloroethane before drying, using a weight ratio of 1:2 for crude products to 1,1,1-trichloroethane. 1 H NMR spectra of DMImBS were measured on a Bruker DAX 300 spectrometer using D$_2$O as solvent. 1H-NMR (D$_2$O, 400MHz): δ1.59 (m, 2H), 1.79 (m, 2H), 2.42 (s, 3H), 2.78 (t, 2H), 3.59 (s, 3H), 4.00 (t, 2H), 7.15 (s, 1H), 7.20 (s, 1H).

The standard electrolyte (electrolyte A) contains 0.1 mol/L iodine (I$_2$, 99%, Aldrich) and 0.1 mol/L anhydrous lithium iodide (LiI, 99%, Aldrich) dissolving in MPII. To prepare electrolyte B and electrolyte C, 0.2mol/L DMImBS and 0.5 mol/L TBP was added into the standard electrolyte, respectively. Electrolyte A, electrolyte B and electrolyte C were used to assemble Device A, Device B and Device C, respectively.

2.2 DSC assembly

TiO$_2$ (anatase) colloidal solution was prepared by hydrolysis of titanium tetraisopropoxide$^{33}$. The TiO$_2$ paste was printed on transparent conducting glass sheets (TEC-15, LOF) and sintered in air at 450 °C for 30 min. The film was 10 µm thick, which was determined by a profilometer (XP-2, AMBIOS Technology Inc. USA). A 4-µm-thick light scattering layer was used. After cooling to 80 °C, the TiO$_2$ films were immersed in an anhydrous ethanol solution with 5.0×10$^{-4}$ M cis-dithioiocyanate-N,N′-bis(4-carboxylate-4-tetrabutylammonium carboxylate-2,2′-bipyridine) ruthenium (II) (N719) for 12 h. Excess N719 dye in TiO$_2$ films was rinsed off with anhydrous ethanol before assembly. The counter electrode was platinumized using H$_2$PtCl$_6$ solution on transparent conducting glass and fired in air at 410 °C for 20 min. Then, it was placed directly on the top of the dye-sensitized TiO$_2$ films. The gap between the two electrodes was sealed by thermal adhesive (Surlyn, Dupont). The electrolyte was then filled from a hole made on the counter electrode and the hole was later sealed by a cover glass and thermal adhesive films. The total active electrode area of the DSC was 0.25 cm$^2$.

2.3 Measurements

The adsorption process of DMImBS was investigated by fourier transform infrared spectroscopy (FTIR). Electrochemical impedance spectroscopy (EIS) is performed to investigate the electron recombination at the Dyed-TiO$_2$/electrolyte interface modified with and without DMImBS. The photovoltaic performances of DSC were measured with a Keithley 2420 digital source meter (Keithley, USA), under a 450 W Xenon lamp (Oriel, USA) with a filter (AM 1.5). Electrochemical impedance spectroscopy (EIS) measurements were carried out on 1M6ex (Germany, Zahner Company), at perturbation amplitude of 5 mV within the frequency range from 3 MHz to 5 mHz in the dark. Intensity modulate photocurrent spectroscopy and intensity modulate photo voltage spectroscopy (IMVS/IMPS) measurements were also performed on an IM6ex using light emitting diodes (λ=610 nm) driven by Export (Germany, Zahner Company). The LED provided both dc and
ac components of the illumination. A small ac component is 10% or less than that of the dc component and the frequency range was 3 kHz to 0.3 Hz. The IR spectra for all the samples were measured using a NICOLET8700 FTIR spectrometer (USA, Thermo Fisher Scientific). The Raman spectra were measured by LABRAM-HR (Franch, YJ).

The samples for ATR-FITR and UV-vis were prepared by immersing TiO$_2$ films (5 µm) in a 0.2 mol/L DMImBS glycol solution (DMImBS/TiO$_2$) and a 0.5 mmol/L N719 ethanol solution (N719/TiO$_2$) for 12 h respectively. These samples were then rinsed with ethanol three times and dried under vacuum at room temperature. The samples for ATR-FITR and UV-vis were prepared by immersing TiO$_2$ films (5 µm) in a 0.2 mol/L DMImBS glycol solution (DMImBS/TiO$_2$) and a 0.5 mmol/L N719 ethanol solution (N719/TiO$_2$) for 12 h respectively. These samples were then rinsed with ethanol three times and dried under vacuum at room temperature.

The DMImBS/TiO$_2$ sample for Raman spectra were prepared by immersing TiO$_2$ powder in a 0.2 mol/L DMImBS glycol solution for 12 h, then rinsed with ethanol three times and dried under vacuum at room temperature.

3. Results and Discussion

Figure 1 shows the Raman spectra of TiO$_2$ and DMImBS/TiO$_2$. The anatase TiO$_2$ has Raman-active fundamentals in the vibrational spectrum. The peaks at 638 cm$^{-1}$, 395 cm$^{-1}$ and 512 cm$^{-1}$ are attributed to Eg mode, Bg mode and Ag mode respectively. In the high wavenumber region of the Raman spectrum of DMImBS/TiO$_2$ sample, new bands are obtained at 2879 cm$^{-1}$ and 2949 cm$^{-1}$ which are assigned to the C–H stretching of the methyl and methylene groups. $^{34,35}$ Imidazolium ring stretching modes give rise to weak bands at 1420 cm$^{-1}$ and 1335 cm$^{-1}$. $^{36}$ The peaks at 226 cm$^{-1}$ and 307 cm$^{-1}$ are assigned to $\nu$(N–C) and bending vibration of imidazolium. $^{37}$ The symmetric stretching mode of SO$_3^-$ is between 1025 and 1080 cm$^{-1}$, this peak is observed at 1031 cm$^{-1}$ for DMImBS. $^{38}$ The result indicate that DMImBS adsorb on the TiO$_2$ surface.

The ATR-FITR spectra of DMImBS/TiO$_2$, N719/TiO$_2$ and DMImBS/N719/TiO$_2$ films are compared in Figure 2. The band at 1229 cm$^{-1}$ and 1291 cm$^{-1}$ are attributed to asymmetric sulfonyl (O=S=O) stretching vibrations of the sulfonate groups ($\nu_{as}$ (R-SO$_3^-$)). $^{1120}$ cm$^{-1}$ and $^{1040}$ cm$^{-1}$ can be assigned to symmetric sulfonyl stretching vibrations ($\nu_s$ (R-SO$_3^-$)). $^{39}$ Sulfonate groups are very sensitive to changes in local coordination geometry; a symmetry change of sulphate and sulfonate groups may result in splits or shifts in the asymmetric vibrations. $^{40}$ The spectra of DMImBS/TiO$_2$ samples predict that adsorption of DMImBS on the TiO$_2$ surface occurs. Comparison of DMImBS/N719/TiO$_2$ with DMImBS/TiO$_2$ shows that the intensity of the $\nu_s$(R-SO$_3^-$) band is diminished relative to other bands when co-adsorption with N719 is added. The change upon adsorption to TiO$_2$ surface is suggested to be due to a change in the local environment of the sulfonate head. $^{41}$ Kevin $^{42}$ reported that the adsorbed surfactant was surface-bound via electrostatic and hydrogen bonding interactions between the sulfate head group and the oxide surface. In the DMImBS/TiO$_2$ sample, small peaks between 1400 cm$^{-1}$ and 1550 cm$^{-1}$ are assigned to $\nu$(C=N) of imidazole group. The band at 1662 cm$^{-1}$ is attributed to $\nu$(C$=$C). The $\nu$(C$=$N) peak of imidazole group is observed in the DMImBS/N719/TiO$_2$ sample. $^{43,44}$ Due to the weak electrostatic adsorption between imidazolium cation and TiO$_2$ surface, the intensity of characteristic peak is very weak. From the IR spectroscopic data we further confirm that the DMImBS adsorbed on the TiO$_2$ surface and the sulfonate group is predominant adsorption group. $^{45,46}$

The N719/TiO$_2$ shows strong bands at 1600 cm$^{-1}$ and 1381 cm$^{-1}$, due to $\nu_{as}$(-COO$^-$) and $\nu_s$(-COO$^-$), which are attributed to the carboxylate group anchored to the TiO$_2$ surface. The bands at 1724 cm$^{-1}$ are due to the $\nu$(C=O) and $\nu$(C=O) of the free carboxylic acid of N719 $^{47,49}$. The intensity at 1724 cm$^{-1}$ relative to that at 1381 cm$^{-1}$ indicates that the relative amounts of unstable and stable type dye molecules on the TiO$_2$ surface. $^{50}$ The intensity of these characteristic peaks do not significantly change between the N719/TiO$_2$ and DMImBS/N719/TiO$_2$ sample, which indicates that dye adsorption is not influenced by co-adsorption with DMImBS.
Figure 3. Current density-voltage characteristics of Device A, Device B and device C under AM 1.5 simulated sunlight (100 mW cm\(^{-2}\)) illumination.

<table>
<thead>
<tr>
<th>Device</th>
<th>(V_{oc}/\text{V})</th>
<th>(J_{sc}/\text{mA cm}^{-2})</th>
<th>(FF)</th>
<th>(\eta/%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.63±0.01</td>
<td>12.35±0.44</td>
<td>0.57±0.02</td>
<td>4.42±0.12</td>
</tr>
<tr>
<td>B</td>
<td>0.66±0.00</td>
<td>11.28±0.35</td>
<td>0.65±0.01</td>
<td>4.83±0.08</td>
</tr>
<tr>
<td>C</td>
<td>0.67±0.00</td>
<td>10.29±0.44</td>
<td>0.67±0.02</td>
<td>4.62±0.09</td>
</tr>
</tbody>
</table>

*The results are the average values obtained from 5 cells with standard deviation.

\(I-V\) curves of Device A, Device B and Device C measured under full air mass 1.5 AM (intensity of the solar simulator: 100mW/cm\(^2\)) are predicted in Figure 3. The \(J_{sc}, V_{oc}\) and fill factor (FF) of Device A are 12.56 mA cm\(^{-2}\), 0.63 V and 0.57, respectively, yielding an overall the photovoltaic conversion efficiency (\(\eta\)) of 4.53%. The corresponding \(I-V\) parameters of Device B with DMImBS as additive are 11.68 mA cm\(^{-2}\), 0.66 V, 0.64 and 4.93%. Table 1 illustrates the average characteristic performance parameters of the DSCs with standard deviation based on five cells of different electrolyte. In general, compared with Device A, Device B has a higher \(V_{oc}\) and FF. Using DMImBS as an additive enhances the power conversion efficiency without significant penalty in photocurrent. In addition, Device B containing DMImBS displays better \(\eta\) compared to Device C containing TBP as a classic additive.

DSC is a multi-phase complicated photo-electrochemical system where two phases inside the cell contact each other to form several interfaces. Photoelectric energy conversion of DSC’s is related to the processes of electron transfer that take place on the contact interfaces\(^{31}\). EIS is a powerful frequency-domain technique used to identify and study the kinetics of interface electron transfer in DSC. EIS measures the current response to a modulated applied bias superimposed on a constant applied voltage, it can be evaluated using resistance and capacitance elements as an equivalent circuit. At the Dyed-TiO\(_2\)/electrolyte interface electrons are transferred to the electrolyte to react with \(I_2\), the difficulty of this reaction can be denoted with a reaction resistance (\(R_{ct}\)). The capacitance of TiO\(_2\) film can be denoted with a chemical capacitance (\(C_p\)). The Dyed-TiO\(_2\)/electrolyte interface can therefore be represented by a composite component in which \(R_{ct}\) is connected in parallel with \(C_p\). Near the counter electrode/electrolyte interface a double electrode layer is formed between the counter electrode surface and the electrolyte, so transfer impedance in the reaction between the electrons and the electrolyte can also be denoted with a composite component consisting of parallel circuit. The \(R_{ct}\) is connected in parallel with \(C_p\). By fitting the measurement impedance data, electron transfer resistance and capacitance at the Dyed-TiO\(_2\)/electrolyte and the counter electrode/electrolyte interface can be deduced. The fitting results of EIS measurements under various applied potential at equal intervals are plotted in Figure 4.

\[
R_{ct} = R_0 \exp\left(-\frac{eV}{k_{B}T}\right) \tag{1}
\]

where \(R_0\) is constant, \(e\) is the electron charge, \(V\) is the bias voltage, \(k_{B}\) is the Boltzmann constant, \(T\) is the temperature and \(\beta\) is the transfer coefficient which corresponds to the reciprocal value of diode quality factor. Comparison of device A and device B shows that \(R_{ct}\) increases when DMImBS is added to the electrolyte. The \(V_{oc}\) is determined by the difference between
the electron quasi-Fermi level in the TiO\textsubscript{2} film under illumination and the redox potential of the electrolyte in the dark. In DSC, photoelectrons easily transfer from the TiO\textsubscript{2} film to the electrolyte and recombine with \textit{I}\textsubscript{1}. Electron recombination plays a significant role in reducing the electron density in the TiO\textsubscript{2} film, a problem which becomes more prevalent when using ionic liquid electrolytes instead of liquid electrolytes\textsuperscript{26}. The increase of \textit{R}\textsubscript{ct} indicates that suppression of electron recombination occurs as a means of increasing electron density. This result indicates that addition of DMImBS is effective in retarding the back reaction between \textit{I}\textsubscript{1} and electrons in TiO\textsubscript{2} film.

Due to the small radius of TiO\textsubscript{2} particles, there is no space charge layer in the nanoporous TiO\textsubscript{2} film. Research shows that the TiO\textsubscript{2} film has two main types of capacitance: chemical capacitance inside the film and Helmholtz double electric layer capacitance at the interface of Dye-TiO\textsubscript{2}/electrolyte (significant only under high bias voltage)\textsuperscript{54-56}. The capacitance of TiO\textsubscript{2} film can be denoted with \(C_{\mu}\) which is related to the concentration of electrons and the chemical potential of the TiO\textsubscript{2} film\textsuperscript{57}. Figure 4 (b) shows a strong exponential increase of \(C_{\mu}\) with increasing carrier density or bias voltage in the solar cell.

More efforts of electron recombination achieve by the EIS results. the kinetics of electron transfer to \(I_{1}\) are usually discussed in terms of the electron lifetime (\(\tau_{n}\))\textsuperscript{58}, which can be calculated directly from EIS response (\(\tau_{n} = R_{ct} \times C_{\mu}\)). The greater \(\tau_{n}\) gets, the slower the recombination between electrons and \(I_{1}\) will be. As shown in Figure 5, at the identical electron energy levels, Device B has a consistently longer apparent recombination lifetime \(\tau_{n}\) than that of Device A; further supporting evidence that slower charge recombination occurs in Device B relative to Device A. It is generally believed that electrons may undergo recombination reactions with \(I_{1}\) on the Dye-TiO\textsubscript{2}/electrolyte interface through the following two ways: direct recombination via conduction band or recombination via surface states. If electrons undergo recombination reactions via the TiO\textsubscript{2} conduction band, \(\beta\) should equal to 1\textsuperscript{59}. The values of \(\beta\) are 0.24 for both Device A and Device B, which is calculated from the slope of the straight line as illustrated in Figure 4(a). Both values of \(\beta\) are less than 1, indicating that electrons primarily undergo recombination reactions via the surface state rather than via the TiO\textsubscript{2} conduction band. Since DMImBS only covers the surface of the TiO\textsubscript{2}, it can be inferred that electron recombination via surface state may play an importance role in DSC.

![Figure 5](image-url)  
**Figure 5** The lifetime versus applied potential in Device A and Device B.

To further investigate the kinetics of electron transfer at the Dye-TiO\textsubscript{2}/electrolyte interface are generally discussed according to the results of IMVS/IMPS measurements. The \(V_{oc}\) of DSC are dependent on the gap between the TiO\textsubscript{2} Fermi energy level and the redox potential in the electrolyte. As shown in Figure 6, lithium perchlorate (LiClO\textsubscript{4}) is used as the lithium salts in the electrolyte, and the redox potentials of each electrolyte are maintained throughout the experiment with the same \(\textit{I}^{+}/\textit{I}_{3}\) ratio. Improvement of the \(V_{oc}\) therefore depends on the suppression of recombination and the TiO\textsubscript{2} conduction band shift, they are both related to TiO\textsubscript{2} Fermi energy level. As we know, a major feature of nanoporous TiO\textsubscript{2} film is that a large number of excess electrons can be stored within the network. The charge buildup on the surface of TiO\textsubscript{2} particles causes the variation of potential across the Helmholtz layer, resulting in the TiO\textsubscript{2} conduction band edge movement\textsuperscript{18}. Positive charge buildup causes the band edges to shift downward toward positive electrochemical potentials, which leads to the decrease in \(V_{oc}\). The difference between the TiO\textsubscript{2} conduction band edge and the quasi-Fermi level depends on the photo induced charge (\(Q\))\textsuperscript{55}. For a given \(Q\), the increase in \(V_{oc}\) is attributed to an upward shift of the TiO\textsubscript{2} conduction band whereas a decrease in \(V_{oc}\) is attributed to a downward shift of the TiO\textsubscript{2} conduction band. Figure 6 (a) depicts the dependence of \(V_{oc}\) on \(\ln (Q)\) in TiO\textsubscript{2} at open circuit following Equation (2)\textsuperscript{18}

\[
V_{oc} = V_{0} + m_{0} \ln Q
\]  
(2)

where \(V_{0}\) is the vertical intercept and \(m_{0}\) is the slope rate, that is, \(m_{0} = dV_{oc}/d \ln Q\). Along with the increase of LiClO\textsubscript{4} consistency in the electrolyte, the TiO\textsubscript{2} conduction band shows a gradual positive shift and decrease in \(V_{oc}\) at constant \(Q\). Adsorption of Li\textsuperscript{+} on the TiO\textsubscript{2} surface leads to a positive shift of the TiO\textsubscript{2} conduction band potential\textsuperscript{60}. Employing DMImBS alone as an additive negatively shifts the conduction band and increases the \(V_{oc}\) slightly at constant \(Q\). As previously mentioned in the IR data, sulfate is the major adsorption group, anchoring to the TiO\textsubscript{2} surface. The conduction band shift is suggested to be due to the negative charge of sulfate. Moreover, addition of DMImBS in the electrolyte containing LiClO\textsubscript{4} causes an obviously upward
shift of the TiO₂ conduction band. The conduction bands shift back to the negative potential is attributed to a reduction in the amount of adsorbed Li⁺ on the TiO₂ surface.

Due to the small size of TiO₂ particles and effect of electrolyte shielding, the driving force for electron transport in the TiO₂ film is the electron concentration gradient in film, but not electric field action. The electron transit time (τ₁) is associated with electron transport in porous TiO₂ film, which can be obtained directly from IMPS. The relation between τ₁ and J_{sc} can be described following the exponential distribution in Equation (3).

\[ \tau_1 \propto J_{sc}^{-\alpha} \]

where the parameter \( \alpha \) is related to the steepness of the trap-state distribution. As predicted in Figure 6(b), \( \tau_1 \) is a function of the \( J_{sc} \) in double logarithmic form and \( \tau_1 \) decrease with the increase of \( J_{sc} \) in devices. At the constant of \( J_{sc} \), the \( \tau_1 \) of device with LiClO₄ slightly increases when compared with the device without additive. In the presence of DMImBS, TiO₂ conduction band undergoes a negative shift, which harms the electron injection. The condition of grain boundaries between TiO₂ particles can influence electron transport, DMImBS molecules cover the surface of joints between TiO₂ particles, hindering the electron transport and thus increasing \( \tau_1 \). The slope of the ln(\( \tau_1 \)) versus ln(\( J_{sc} \)) plot, which reflects the trap-state distribution of TiO₂ surface, is described by \( \alpha \) in Equation (3). The decrease of \( \alpha \) is ascribed to the widening of the exponential conduction band and corresponds to a steeper trap state distribution, which may slow down the electron transport process. The values of \( \alpha \) are 0.56, 0.56, 0.51, 0.58 and 0.55 for the X1, X2, X3, X4 and X5, respectively. The steeper trap state distribution with addition of LiClO₄ is attributed to the Kopidakis et al. reported that the Li⁺ intercalated into TiO₂ electrodes. However, the trap state distribution obviously gets shallow with addition of DMImBS, even with the concentration increase of LiClO₄. Trap state distribution doesn’t change with presence of DMImBS alone. So the variety of trap state distribution is not directly derived from adsorption of DMImBS on the surface. There should be some interaction between Li⁺ and DMImBS. We presumed that the addition of DMImBS restrain the intercalation of Li⁺ and improve the cell stability in the aging process.

The counter electrode is an equally important component of DSC. Normally, fluorine-doped tin oxide glass is coated with platinum to afford more reversible electron transfer. The counter electrode/electrolyte interface is a catalytic interface and its role is to transfer electrons arriving from the external circuit back to the electrolyte. The \( R_{CE} \) is an important parameter for the performance of DSC, and it depends on the electrochemical reaction rate at the counter electrode/electrolyte interface. The capacitance of the counter electrode/electrolyte interface can be denoted by a Helmholtz double electric layer capacitance. Figure 7 presents the \( R_{CE} \) and \( C_p \) of Device A and Device B at the counter electrode/electrolyte interface. At a given dark current, there are very small differences of \( R_{CE} \) and \( C_p \) between Device B and Device A. This result rules out the effect from the electron transfer process occurring at the counter electrode with DMImBS. In other words, DMImBS does not influence the catalytic activity of the counter electrode/electrolyte interface.
In conclusion, we have identified DMImBS as a novel solid additive which can improve performance of DSC based on ionic liquid electrolyte. DMImBS successfully adsorbed on TiO$_2$ surface when DMImBS was added to MPII-based ionic liquid electrolyte. The alky fridge is formed with two functional group adsorption. The electron recombination is effectively restrained by the DMImBS modification. Adsorption of dye molecular was not influenced by the co-adsorption of DMImBS. The intercalation of Li$^+$ in TiO$_2$ film was restrained by the adsorption of DMImBS. Therefore, modification of the interface improved the stability of solar cell. The $V_{oc}$ and FF of the DSC were significantly improved without significant decrease in $J_{sc}$. Use of DMImBS as an additive substantially increased the power output of the DSC due to suppression of interfacial recombination, whilst achieving a more stable interface which improves long-term stability of the DSC.

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