Research Article

Application of Diaminium Iodides in Binary Ionic Liquid Electrolytes for Dye-Sensitized Solar Cells

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1. Introduction

Dye-sensitized solar cells (DSCs) have been extensively studied due to their high light-to-electric energy conversion yield. The electrolyte is one of the critical components of DSCs and has the remarkable effect on the stability under long-term thermal and light-soaking dual stresses [1–3]. Various functionalized imidazolium iodide salts [4–9] and the additives containing basic N atoms, such as pyridine [10, 11], benzimidazole [12, 13], aminotriazole [14], have been evaluated as electrolytes in DSCs with some success. Compared with the imidazolium iodides and pyridine derivatives, diaminium iodides serve as the iodide resource of I3−/I− redox couples and contain basic N atoms. However, their application in the electrolytes of DSCs has not been reported.

In this paper, N-(2-hydroxyethyl)ethylenediaminium iodides (HEEDAI) and N-(2-hydroxyethyl)piperazinium iodides (HEPI) were synthesized, and their thermal properties were characterized by thermogravimetric analysis and differential scanning calorimeter. The influence of HEEDAI and HEPI on I3−/I− redox behavior in binary ionic liquid was investigated. The result revealed that HEEDAI can suppress the recombination between I3− and the injected electrons in TiO2 conduction band and be used as the alternative of 4-tert-butylpyridine in the electrolyte of dye-sensitized solar cells. The electrolyte C, 0.15 mol·L−1 I2, HEEDAI and MPII with mass ratio of 1 : 4, gave the short-circuit photocurrent density of 9.36 mA·cm−2, open-circuit photovoltage of 0.67 V, fill factor of 0.52, and the corresponding photoelectric conversion efficiency of 3.24% at the illumination (air mass 1.5, 100 mW·cm−2, active area 0.25 cm2).

2. Experimental

2.1. Materials. All chemical reagents were commercially available and used without further purification. 1-methyl-3-propylimidazolium iodide (MPII) was synthesized according to our previous report [15].

2.2. Measurement. The melting and freezing points of HEEDAI and HEPI were characterized by differential scanning calorimeter using a Thermal Analysis DSC821 with heating rate at 10°C·min−1, under a nitrogen flow rate of about 30 mL·min−1. And their thermal stability was analyzed on a TG209F3 Tarsws NETZSCH thermogravimetric balance at a heating rate of 10°C·min−1 under nitrogen atmosphere.

The measurements of cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS) for binary ionic liquid and the fabrication of DSCs were measured.
Scheme 1: The synthesis reaction of HEEDAI and HEPI.

Figure 1: Thermogravimetric analysis curves of HEEDAI.

Figure 2: Thermogravimetric analysis curves of HEPI.

disk ultramicroelectrode (CHI107); the auxiliary electrode and the reference electrode were a 1.0 mm radius Pt disk electrode (CHI102). A slow scan rate at 5 mV·s⁻¹ was used to obtain steady-state current-voltage curves. The EIS measurement was carried out with a two-platinum-black-electrode conductance cell (DJS-1C, REX, cell constant is 0.974). After every EIS measurement, the conductance cell was washed with anhydrous ethanol and immersed in distilled water overnight. EIS spectra were obtained by applying sinusoidal perturbations of ±10 mV over the bias 0 V at frequencies from 1 to 10 kHz. The TiO₂ paste was printed on transparent conducting glass sheets (TEC-8, LOF).

Figure 3: Differential scanning calorimeter curves of (A) HEEDAI and (B) HEPI.

Figure 4: Steady-state voltammograms of solutions A–D at 25°C. Solution component: (A) m(HEEDAI) : m(MPII) = 1 : 9; (B) m(HEPI) : m(MPII) = 1 : 9; (C) m(HEEDAI) : m(MPII) = 1 : 4; (D) m(HEEDAI) : m(MPII) = 1 : 1, and 0.15 mol·L⁻¹ I₂.
and sintered in air at 450°C for 30 min. The film was ~10 μm thick, which was determined by profilometer (XP-2, AMBIOS Technology Inc. USA). A 4 μm thick light scattering layer was used. After cooling to 80°C, the TiO₂ films were immersed in anhydrous ethanol solution with 5.0 × 10⁻⁴ mol·L⁻¹ cis-dithiocyanate-N,N′-bis-(4-carboxylate-4-tetraethylammonium carboxylate-2,2′-bipyridine) ruthenium (II) (N719) overnight. The excess of N719 dye in TiO₂ films was rinsed off with anhydrous ethanol before assembly. The counterelectrode was platinized by spraying H₂PtCl₆ solution to 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 TiO₂ films. The gap between the two electrodes was sealed by thermal adhesive (Surlyn, Dupont). The electrolyte was filled from a hole made on the counter electrode, and the hole was later sealed by a cover glass and thermal adhesive films. The active area of DSCs was 0.25 cm². The photovoltaic performances of DSCs were measured with a solar simulator (Class AAA, Oriel, Newport, USA, Air Mass 1.5, 100 mW·cm⁻²) and a Keithley 2420 source meter controlled by Testpoint software. The irradiation intensity was calibrated with standard crystalline silicon solar cell (Oriel, Newport, USA).

2.3. Synthesis Procedure. HEEDAI and HEPI were synthesized by the neutralization reaction of N-(2-hydroxyethyl)ethylenediamine/N-(2-hydroxyethyl)piperazine and HI using the stoichiometric amounts of reactants (Scheme 1).

N-(2-hydroxyethyl)ethylenediamine (45.00 g, 0.43 mol, HEEDA) was dissolved in the equal weight of deionized water. Then hydriodic acid (122.81 g, 0.43 mol HI, 45% mass fraction) was slowly dropped to the above solution with magnetic stirring at room temperature. Subsequently, the reaction mixture was stirred for 30 min, and the deionized water was removed by rotary evaporator at 90°C and dried in vacuum at 80°C for 6 h. A red viscous liquid, HEEDAI, was obtained. Yield: 94.26 g (94%).

The synthesis of HEPI was similar to that of HEEDAI. A white solid, HEPI, was obtained with the reaction of N-(2-hydroxyethyl)piperazine (45.54 g, 0.35 mol, HEP) and hydroiodic acid (99.50 g, 0.35 mol HI, 45% mass fraction). Yield: 86.70 g (96%).

3. Results and Discussion

3.1. Thermal Analysis of HEEDAI and HEPI. Figures 1 and 2 showed thermogravimetric analysis curves of HEEDAI and HEPI. The decomposition temperatures of HEEDAI and HEPI were 345°C and 325°C. This result revealed that the thermal stability of HEEDAI was superior to that of HEPI. Because the pK_b1 of HEEDA and HEP were 4.4 and 4.8, respectively [19], the basicity of HEEDA was more intense than that of HEP, and the N–H bond energy in the resulting iodide salt, HEEDAI, was higher than that in HEPI. The N–H bond break of HEEDAI required higher decomposition temperature than HEPI.
Figure 3 showed differential scanning calorimeter curves of HEEDAI and HEPI. HEEDAI had a glass transition temperature at −28 °C, a crystallization peak followed by a broad melting transition between −10 °C and 170 °C. And the melting point of HEPI was 94 °C, while the freezing points of HEEDAI and HEPI were not observed. It was indicated that the two iodide ionic liquids were transferred from the solid state when refrigerated from high temperature to low temperature. Due to the large excess of I⁻ relative to I₃⁻, only the cathodic steady-state diffusion current of I₃⁻ was measured in solutions A–D and the apparent diffusion coefficient (D_app) of I₃⁻ was listed in Table 1. Compared with solution A and B, D_app value of I₃⁻ in binary ionic liquid of HEEDAI and MPII was higher than that in HEPI and MPII. This was because the molecular size of HEPI was larger than that of HEEDAI. From solutions A, C, and D, the diffusion coefficients of I₃⁻ increased with the increase of MPII. This was because MPII had lower macroscopic viscosity than that of HEEDAI. It was noteworthy that the diffusion coefficients of I₃⁻ in binary ionic liquid of HEEDAI and MPII, the mass ratio of 1:4 and 1:9, were higher than pure MPII solution (1.88 × 10⁻⁸ cm²·s⁻¹) [20].

Figure 5 showed the EIS spectra of the solutions A–D at 25°C. Experimental data were represented by symbols and the solid lines corresponded to the fit using the equivalent circuit. The parameters obtained by fitting the experimental spectra with the equivalent circuit were listed in Table 2. With the increase of HEEDAI, the viscosity of solution increased, and Rₛ and Rₛ increased. This result was consistent with the diffusion coefficient of I₃⁻ in binary ionic liquid of HEEDAI and MPII. Moreover, the values of Rₛ and Rₛ in binary ionic liquid of HEPI and MPII were higher than that in binary ionic liquid of HEEDAI with MPII.

### Table 2: Parameters obtained by fitting the experimental spectra with the equivalent circuit.

<table>
<thead>
<tr>
<th>Solution</th>
<th>Rₛ/Ohm</th>
<th>Rₛ/Ohm</th>
<th>Yₒ/μF·cm⁻¹</th>
<th>n</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A) m(HEEDAI) : m(MPII) = 1:9 Without TBP</td>
<td>440</td>
<td>120</td>
<td>33</td>
<td>0.8</td>
</tr>
<tr>
<td>(B) m(HEEDAI) : m(MPII) = 1:9</td>
<td>549</td>
<td>156</td>
<td>25</td>
<td>0.8</td>
</tr>
<tr>
<td>(C) m(HEEDAI) : m(MPII) = 1:4</td>
<td>769</td>
<td>340</td>
<td>32</td>
<td>0.8</td>
</tr>
<tr>
<td>(D) m(HEEDAI) : m(MPII) = 1:1</td>
<td>1209</td>
<td>303</td>
<td>46</td>
<td>0.8</td>
</tr>
</tbody>
</table>

Component of solutions A–D was the same as that in Figure 5. Rₛ: serial resistance (resistance of solutions); Rₛ: charge-transfer resistance; CPE: constant phase element; Yₒ: constant phase angle element; n: a constant ranging from 0 ≤ n ≤ 1.

### Table 3: Photovoltaic performance of DSCs with electrolytes A–G.

<table>
<thead>
<tr>
<th>Electrolyte</th>
<th>J_sc/(mA·cm⁻²)</th>
<th>V_oc/V</th>
<th>FF</th>
<th>η (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A) m(HEEDAI) : m(MPII) = 1:9 Without TBP</td>
<td>6.88</td>
<td>0.68</td>
<td>0.46</td>
<td>2.16</td>
</tr>
<tr>
<td>(B) m(HEEDAI) : m(MPII) = 1:9, 0.5 mol·L⁻¹ TBP</td>
<td>6.64</td>
<td>0.69</td>
<td>0.45</td>
<td>2.06</td>
</tr>
<tr>
<td>(C) m(HEEDAI) : m(MPII) = 1:4 Without TBP</td>
<td>9.36</td>
<td>0.67</td>
<td>0.52</td>
<td>3.24</td>
</tr>
<tr>
<td>(D) m(HEEDAI) : m(MPII) = 1:4, 0.5 mol·L⁻¹ TBP</td>
<td>8.96</td>
<td>0.66</td>
<td>0.52</td>
<td>3.06</td>
</tr>
<tr>
<td>(E) m(HEEDAI) : m(MPII) = 1:1 Without TBP</td>
<td>5.76</td>
<td>0.63</td>
<td>0.50</td>
<td>1.80</td>
</tr>
<tr>
<td>(F) m(HEEDAI) : m(MPII) = 1:1, 0.5 mol·L⁻¹ TBP</td>
<td>5.68</td>
<td>0.64</td>
<td>0.48</td>
<td>1.74</td>
</tr>
<tr>
<td>(G) m(HEPI) : m(MPII) = 1:9 Without TBP</td>
<td>7.20</td>
<td>0.65</td>
<td>0.53</td>
<td>2.44</td>
</tr>
</tbody>
</table>

The electrolyte component of A–G: (A) m(HEEDAI) : m(MPII) = 1:9; (B) m(HEEDAI) : m(MPII) = 1:9, 0.5 mol·L⁻¹ TBP; (C) m(HEEDAI) : m(MPII) = 1:4; (D) m(HEEDAI) : m(MPII) = 1:4, 0.5 mol·L⁻¹ TBP; (E) m(HEEDAI) : m(MPII) = 1:1; (F) m(HEEDAI) : m(MPII) = 1:1, 0.5 mol·L⁻¹ TBP; and 0.15 mol·L⁻¹ LiCl; (G) m(HEPI) : m(MPII) = 1:9 and 0.15 mol·L⁻¹ LiCl. Illumination: AM 1.5, 100 mV·cm⁻², active areas of DSCs: 0.25 cm².

3.3. Photovoltaic Performances of DSCs. Table 3 and Figure 6 showed the photovoltaic performance of DSCs. The electrolyte C gave the short-circuit photocurrent density (J_sc) of 9.36 mA·cm⁻², open-circuit photovoltage (V_oc) of 0.67 V, fill factor (FF) of 0.52, and yielded an overall photoelectric conversion efficiency (η) of 3.24%. From Table 3, it was interesting that the addition of 0.50 mol·L⁻¹ 4-tert-butylpyridine (TBP) in binary ionic liquid of HEEDAI and MPII cannot improve the photoelectric conversion efficiency of DSCs. This was because HEEDAI, similar to TBP, contains the basic N atoms. These N atoms can be adsorbed on the TiO₂ surface to suppress the recombination between I₃⁻ and the injected electrons in TiO₂ conduction band. Therefore, HEEDAI can be used as the alternative of TBP in the electrolyte of DSCs and has lower volatility than TBP.

Compared with electrolytes C and A, the short-circuit photocurrent density (J_sc) and fill factor (FF) of electrolyte C were higher than those of electrolytes A. This result was because the HEEDAI content in electrolyte C was higher than electrolyte A, which can suppress the recombination between I₃⁻ and the injected electrons in TiO₂ conduction band. Compared with electrolytes E and C, the short-circuit photocurrent density (J_sc) and fill factor (FF) of electrolyte E were lower than those of electrolyte C. This result was because the diffusion coefficient of I₃⁻ in electrolyte E is lower those electrolyte C. Moreover, there were no obvious differences of photoelectric conversion efficiency between electrolytes A and G.
4. Conclusion

HEEDAI and HEPI were synthesized, and their thermal properties were characterized by the thermogravimetric analysis and differential scanning calorimeter. The influence of HEEDAI and HEPI on $I_3^-/I^-$ redox behavior in binary ionic liquid was investigated by cyclic voltammetry using a Pt disk ultramicroelectrode and electrochemical impedance spectroscopy. It was found that HEEDAI had a glass transition temperature at $-28^\circ$C, the melting point of HEPI was 94$^\circ$C, and the freezing points of HEEDAI and HEPI were not observed. And HEEDAI can suppress the recombination between $I_3^-$ and the injected electrons in TiO$_2$ conduction band and be used as the alternative of TBP in the electrolyte of DSCs. Moreover, The electrolyte C, 0.15 mol·L$^{-1}$ I$_2$, HEEDAI and MPII with mass ratio of 1:4, gave the short-circuit photocurrent density of 9.36 mA·cm$^{-2}$, open-circuit photovoltage of 0.67 V, fill factor of 0.52, and the corresponding photoelectric conversion efficiency of 3.24%.

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References
