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Dye-sensitized solar cells using an anion-conducting polymer as a material of solid-state electrolyte

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Abstract—An anion-conducting polymer (ACP) has been used as an electrolyte material of solid-state dye-sensitized solar cells (DSCs) and the basic electrochemical properties of DSCs have been examined in this study. Initial photocurrent-voltage characteristics of the DSC prepared with ACP (DSC-S) were inferior to those of the DSC prepared with a general liquid electrolyte (DSC-Liq), but DSC-S exhibited better long-term stability. From AC impedance measurements, the relatively poor photocurrent-voltage characteristics of the as-fabricated DSC-S were anticipated to arise from larger resistance of charge transfer at electrolyte/TiO₂ electrode interface than that of the as-fabricated DSC-Liq.

Keywords; dye-sensitized solar cell; solid electrolyte; anion-conducting polymer

I. INTRODUCTION

Dye-sensitized solar cells (DSCs) are a promising candidate as an alternative to conventional silicon-based solar cells, because of their low cost, relatively high energy-conversion efficiency and a simple method for manufacturing, and the conversion efficiency of 11% for the DSCs has been achieved in some literatures [1, 2]. However, the long-term stability of the DSCs is extremely poor, because of leakage and volatilization of the liquid electrolyte involved. Hence, solidification of electrolyte is one of important techniques to improve these problems. Recently, several materials, such as CuSCN [3], 2,2’,7,7’-tetrakis (N,N’-di(4-methoxyphenyl)-amine)-9,9’-spirobifluorene (spiro-OMeTAD) [4], polyvinylidene difluoride (PVDF) [5], have been developed as a potential candidate of the solid-state electrolyte.

We have recently focused on anion-conducting polymers (ACP) as an electrolyte for various electrochemical devices, because of their high anion conductivity and improved long-term stability [6-8]. Therefore, ACP was employed as a solid-state electrolyte in fabricating DSCs, and fundamental electrochemical properties of the DSCs have been investigated in this study.

II. EXPERIMENTAL

A. Preparation of TiO₂ electrode modified with dye

A commercial TiO₂ powder (P-25, AEROXIDE®) was mixed with appropriate amounts of surfactant (Triton-X 100, SIGMA-ALDRICH®), pure water, polyethylene glycol (molecular weight: 20000) and acetylacetone. The slurry obtained was coated on an FTO substrate by spin coating (electrode area: 5 x 5 mm²). After drying in air, it was heat-treated at 500°C for 2 h. Then, it was soaked in dye solution {mixture of ruthenium dye ([RuL₂(NCS)₂]:2TBA, L: 2,2’-bipyridyl-4,4’-dicarboxylic acid, TBA: tetra-n-butylammonium (N-719), Solaronix), acetonitrile and tert-buthylalcohol} at 50°C for 12 h to ensure the adsorption of the dye on the surface of TiO₂.

B. Fabrication of solid-state DSC

The mixture of 0.05 M I₂, 0.1 M LiI, 0.5 M 4-tert-butylpyridine, 0.6 M 1,2-dimethyl-3-propylimidazolium iodide and acetonitrile was used as electrolyte solution. The electrolyte solution and ACP solution (AS-4, Tokuyama Corp.), which is 1-propanol containing a 5 wt% hydrocarbon-type ionic polymer with quaternary ammonium salts (ion-exchange capacity: 1.5 mmol g⁻¹ and ion conductivity: ca. 13 mS cm⁻¹), were dropped on a dye-modified TiO₂ film simultaneously, and they were dried at 50°C. The volume ratio of the ACP solution and the electrolyte solution dropped was 3:2. In assembling a DSC, a spacer (masking tape) was set between a counter electrode (ITO glass substrate coated with platinum by magnetron sputtering) and the dye-modified TiO₂.
film which had been coated with the dried mixture of ACP and electrolyte components. The solid-state dye-sensitized solar cell thus fabricated was denoted as DSC-S.

However, a large amount of precipitate was deposited when the ACP solution was mixed with the electrolyte solution, as shown in Fig. 1. Thus, the supernatant liquid and the precipitate were separated and dried at 50°C for 1 h. Viscosity of the supernatant liquid after drying was much larger than that before drying. The condensed supernatant liquid and the precipitate after the drying were coated on the dye-modified TiO₂ electrode as an electrolyte of a DSC, respectively. The additional two kinds of DSCs were fabricated by using these TiO₂ electrodes thus prepared, the separator and the counter electrode. The DSCs fabricated with the condensed supernatant liquid and the precipitate were denoted as DSC-Sup and DSC-Pre, respectively. For comparative purpose, another DSC was also fabricated only with the electrolyte solution, and this is referred to as DSC-Liq.

C. Characterization of DSCs

The current density (J)-voltage (V) measurements were performed under illumination of visible light generated from xenon lamp (100 mW cm⁻², cut with visible-light transmission filter). AC impedance characteristics of DSCs under open circuit conditions were also measured upon the illumination of visible light in the frequency range of 100 mHz ~ 200 kHz and at an AC amplitude of 10 mV.

III. RESULTS AND DISCUSSION

Figure 2 shows variations in J-V characteristics of DSC-Liq, DSC-Sup and DSC-S with time during their storage at RT under atmospheric conditions. Figure 3 shows variations in conversion efficiency (η) of DSCs with storage time. η is calculated by using the following equation:

$$\eta = \left( \frac{P_{\text{max}}}{I} \right) \times 100$$

where $P_{\text{max}}$ is maximum power density, $I$ is intensity of incident light. Since current density of as-fabricated DSC-Pre was extremely small over the whole voltage range, the J-V characteristics of DSC-Pre could not be evaluated at any storage periods. As-fabricated DSC-Liq showed the largest short-circuit current density ($J_{sc}$, ca. 7.40 mA cm⁻²) and η (ca. 3.68%) with relatively large open circuit voltage ($V_{oc}$, ca. 0.80 V), among all the DSCs tested. The $J_{sc}$ of as-fabricated DSC-Sup (ca. 5.05 mA cm⁻²) was slightly smaller than that of the as-fabricated DSC-Liq, while the $V_{oc}$ value (ca. 0.68 V) was the smallest among all the DSCs tested. On the other hand, as-fabricated DSC-S showed the smallest $J_{sc}$ (ca. 1.31 mA cm⁻²) and η (ca. 0.50%), while the $V_{oc}$ value (ca. 0.80 V) was comparable to that of DSC-Liq. The cell performances of DSC-Liq and DSC-Sup apparently reduced with an increase in the storage period at room temperature (RT) under

![Figure 2. J-V characteristics of (a) DSC-Liq, (b) DSC-Sup and (c) DSC-S.](image)

![Figure 3. Variations in conversion efficiency of DSCs with storage time.](image)
atmospheric conditions, and the \( J_{sc} \) values of DSC-Liq and DSC-Sol after 72 h was ca. 2.53 and 2.32 mA cm\(^{-2}\), respectively.

On the other hand, the \( J_{sc} \) and \( \eta \) values of DSC-S remained unchanged, irrespective of prolonging the storage time. These results indicated that the DSC-S showed the most excellent long-term stability among the DSCs fabricated.

AC impedance characteristics of these as-fabricated DSCs were measured at RT, as shown in Fig. 4. As for all the DSCs, two kinds of arcs were confirmed in high (~kHz) and medium (~Hz) frequency ranges, and they are referred to as Arc-h and Arc-m, respectively. In addition, both DSC-Sup and DSC-S had another complex impedance component in a low frequency range (~mHz).

To get information on such components in the AC impedance characteristics of the DSCs, electrolytic cells with a pair of Pt electrodes or a pair of dye-modified TiO\(_2\) electrodes (Pt cell-N or DST cell-N, N: Liq, Sup or S) were fabricated and
AC impedance characteristics of the Pt cell-N and the DST cell-N were measured at RT with illumination of visible light generated from xenon lamp (100 mW cm\(^{-2}\), cut with visible-light transmission filter), respectively, as shown in Figs. 5 and 6. As for all the Pt cells (Fig. 5), only one arc appeared in the high frequency range, and the size of the arc increased in the following order, Pt cell-Liq < Pt cell-Sup < Pt cell-S. The values of capacitance component calculated from these arcs by using the Eq. (2) were in the range of \(\sim 10^{-6}\) F and they are comparable to those of capacitance component calculated Arc-h of the DSCs (see TABLE 1).

\[
2\pi fRC = 1
\]

where \(f\) is a value of frequency at the top of the arc, \(R\) is a resistance value equal to the diameter of the arc. Therefore, the Arc-h of DSCs is considered to arise from the charge transfer resistance and capacitance of the Pt electrode/electrolyte interface. The size difference between the arc of Pt cells and the Arc-h of DSCs may be ascribed to slight variations in the experimental conditions. Another complex impedance component indicating Warburg impedance was confirmed for both the Pt cell-Sup and Pt cell-S in the low frequency range. However, such a component was hardly seen for Pt cell-Liq. On the other hand, DST cells had only an arc in the medium frequency range. However, the size of the arcs was larger than that of the Arc-m of all the DSCs, and the values of frequency at the top of arcs of DST cell-Ns (N: Sup and S) were larger than those of the Arc-m of DSC-N (N: Sup and S), respectively, indicating the certain changes in both resistance and capacitance component values. Such differences may arise from a slight change in illumination conditions form one to another cell, leading to a change in redox behavior of \(I_2/\text{I}^+\) at the electrode/electrolyte interface. Actually, the values of resistance calculated from the arcs of DST cell-Sup and -S are in the range of 300 – 400 \(\Omega\). Therefore, these values are closer to those calculated from the Arc-m of DSCs, rather than those of the Arc-h of DSCs (see TABLE 1 ). Thus, the Arc-m of DSCs can be considered to present the charge transfer resistance and capacitance of the dye-modified TiO\(_2\) electrode/electrolyte interface. Another complex impedance component, i.e. Warburg impedance, was also confirmed for all the DST cells in the low frequency range.

Based on these fundamental data, a possible equivalence circuit of DSCs is proposed as shown in the inset in Fig. 4.

### TABLE 1. Results of impedance measurements.

<table>
<thead>
<tr>
<th>Component</th>
<th>Cell</th>
<th>DSC-L</th>
<th>DSC-Sup</th>
<th>DSC-S</th>
</tr>
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<tbody>
<tr>
<td>(R_e(\Omega))</td>
<td>85.8</td>
<td>70.2</td>
<td>84.5</td>
<td></td>
</tr>
<tr>
<td>(\text{Arc-h} \quad R: R_{m}(\Omega))</td>
<td>2.3</td>
<td>7.6</td>
<td>15.6</td>
<td></td>
</tr>
<tr>
<td>(C: C_{m}(10^{-6}\text{F}))</td>
<td>4.9</td>
<td>3.5</td>
<td>2.0</td>
<td></td>
</tr>
<tr>
<td>(\text{Arc-m} \quad R: R_{m}(\Omega))</td>
<td>30.0</td>
<td>27.9</td>
<td>110.0</td>
<td></td>
</tr>
<tr>
<td>(C: C_{m}(10^{-6}\text{F}))</td>
<td>4.8</td>
<td>2.5</td>
<td>2.1</td>
<td></td>
</tr>
<tr>
<td>(Z_{\text{W}}(\Omega))</td>
<td>-</td>
<td>50.5</td>
<td>113.9</td>
<td></td>
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</table>

Here, \(R_e\), \(R_m\), and \(R_{m,TiO_2}\) represent resistance of the electrolyte, charge transfer resistance on the Pt electrode, charge transfer resistance on the dye-modified TiO\(_2\) electrode, respectively. \(Z_{\text{W}}\) stands for Warburg impedance including mass transfer effects in the diffusion layer near the electrode surface. \(C_m\) and \(C_{m,TiO_2}\) represent double layer capacitances on the Pt electrode and the dye-modified TiO\(_2\) electrode, respectively. The electrolyte resistance can be obtained as a distance between the origin and the intercept point, which is the closest to the origin, of Arc-h. Values of the each component in the equivalence circuit are listed in TABLE 1.

In Fig. 4, \(Z_{\text{W}}\) was observed in the low frequency range (\(\sim\) mHz) only for DSC-Sup and DSC-S, and the \(Z_{\text{W}}\) value of DSC-S was larger than that of DSC-Sup. Therefore, it was confirmed that the mass transfer rate in the diffusion layer near the surface of both the dye-modified TiO\(_2\) electrode and the Pt electrode of DSC-S was much slower than that of DSC-Liq. There was little difference in \(R_e\) value among three DSCs in Fig. 4, irrespective of the appearance of large differences in Figs. 5 and 6. Therefore, \(R_e\) may contain additional contact resistance between DSCs and lead wires. On the other hand, \(R_{m,TiO_2}\) values of DSC-S were larger than that of other two DSCs. These results indicate that the charge transfer speed of ions at the electrodes was the slowest when the electrodes were contacted with the solid polymer.

Several DSC-Ss were also fabricated by using solid-state electrolytes prepared at different mixing ratios (ACP solution: electrolyte solution = 1 : 8 or 8 : 1 in volume), and effects of the mixing ratios on \(J-V\) characteristics of as-fabricated DSC-Ss were measured. The \(J-V\) curves of the additionally fabricated DSC-Ss were shown in Fig. 7, together with that of the original DSC-S (ACP solution: electrolyte solution = 3 : 2.

Figure 7. \(J-V\) characteristics of DSC-Ss fabricated by using solid-state electrolytes prepared at different mixing ratios.
in volume). The performances of the additionally fabricated DSC-Ss were poor than that of the original DSC-S.

These results reveal that the mixing ratio of the ACP and electrolyte-components also largely influences on the performance. Therefore, solid-state DSCs can be further improved by simultaneous optimization of the electrode/electrolyte interface conditions and the electrolyte composition.

IV. CONCLUSIONS

A solid-state dye-sensitized solar cell (DSC-S) has been fabricated by using ACP as an electrolyte material, and its \( J-V \) characteristics and AC impedance properties have been investigated in this study. The \( J-V \) characteristics of as-prepared DSC-S were relatively inferior to as-prepared DSC-Liq and DSC-Sup, but the long-term stability was much better than that of DSC-Liq and DSC-Sup. In AC impedance measurements, the DSC-S showed the largest \( R_{Pt} \) and \( R_{TiO_2} \) values, indicating the slowest charge transfer rate of ions at the electrodes in the case of the direct contact with the solid polymer.

REFERENCES