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7.35%–Efficiency rear–irradiated flexible dye–sensitized solar cell by sealing liquid electrolyte in a groove

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We present here the feasibility of assembling rear–irradiated flexible dye–sensitized solar cell employing transparent Ni–Se alloy counter electrode along with groove stored TiO2 and liquid electrolyte. The flexible device with NiSe counter electrode and anode at a groove depth of 36 µm yields a maximum efficiency of 7.35%.

Dye–sensitized solar cell (DSSC) is an electrochemical device converting solar energy into electricity with no emissions.1–7 The state–of–the–art DSSCs are always assembled on fluorine–doped tin oxide (FTO) conductive glass substrates,8,9 leading to inferiorities in frangibility, high weight, unyieldiness, and high cost. One solution to this impasse is the employment of flexible substrates such as Ti foil,10 tin doped indium oxide (ITO)/poly(ethylene naphthalate) (PEN),11 or ITO/poly(ethylene terephthalate) (PET) substrates for flexible DSSC assembly. Due to a fact that the ITO/PEN or ITO/PET substrates suffer devitrification and deformation at calcination temperature exceeding 150 ºC, therefore, conductive plastic supported TiO2 can not realize the crystal transition from rutile to anatase. That is not surprising to obtain unsatisfactory power conversion efficiencies from flexible cell devices.12–14 Although there are emerging low–temperature techniques for anatase TiO2 nanocrystallites,15 the adhesiveness between TiO2 film and flexible substrate is still unsatisfactory, leading to a high interfacial resistance for electron transportation. An ideal flexible DSSC comprises a Ti foil supported TiO2 anode, Pt coated ITO/PEN (or ITO/PET) counter electrode (CE), and liquid electrolyte containing I3–/I− redox couples. Viewed from this point, irradiation from rear side is the sole technique for driving this system.16 However, traditional Pt CE has metallic luster and therefore can reflect most of incident light, resulting in a low light intensity for dye irradiation and electron generation. Another remaining problem is the leakage and evaporation of liquid electrolyte. Although the utilization of quasi–solid–state and full–solid–state electrolytes can well reserve redox couples,17–20 the relatively low charger transfer kinetics within the electrolytes has been considered a tremendous limit for efficiency enhancement. By addressing these two issues, the explanations of cost–effective and transparent CEs as well as a new sealing strategy for liquid electrolyte seem to be crucial for achieving cell devices with high efficiency and good stability.

Inspired by a container for liquid storage, in which the bottom and surrounding sides are blocked, leaving top cover open, in the current work, we have successfully designed transparent nickel selenide (Ni–Se) alloys on flexible ITO/PEN substrates and etching of Ti foil for reserving dye–sensitized TiO2 and liquid electrolyte. The TiO2 film is consolidated within the etched groove and liquid electrolyte is stored in the nanopores of TiO2 layer. Therefore, the leakage issue for liquid electrolyte can be well solved by sealing it in a groove surrounded by Ti foil and Ni–Se CE cover in comparison with interface sealing in a traditional flexible DSSC device. A maximum efficiency of 7.35% is recorded under simulated air mass 1.5 (AM1.5) global sunlight by optimizing Ni/Se ratio and groove depth.

Fig. 1 Schematic diagram for the flexible DSSC device comprising of ITO/PEN supported Ni–Se alloy and Ti foil reserved liquid electrolyte and dye–sensitized TiO2.

Fig. 1 presents schematic diagram of processes for etching, TiO2 coating, dye sensitization, electrolyte injection, and cell assembly. The depth of the groove is controlled by adjusting etching time. In the current work an etching speed of 3 µm cm−1 is utilized to create grooves. Like a container, the injected liquid electrolyte is reserved by etched groove and covered by Ni–Se alloy electrode.
As shown in Fig. 2a, X-ray diffraction (XRD) gives characteristic diffraction peaks at 2θ = 33.2°, 44.7°, 50.5°, and 61.8° for (210), (111), (311), and (400) facets, respectively. The result indicates that NiSe alloy (PDF#88-1711) has been successfully synthesized by the electrochemical deposition reported here. Lattice fringes can be observed in high-resolution transmission electron microscopy (HRTEM) photograph, as shown in Fig. 2b, suggesting that the resultant alloy has a good crystallinity. Moreover, the detection of lattice distortions reveals the formation of defects by alloying Ni and Se, promoting the I$_3^-$ reduction process. As shown in Fig. 2c, the resultant flexible Ni–Se alloy electrodes have high optical transparency in visible region (>70%), suggesting a good dye excitation and electron generation at rear irradiation. Fig. 2d compares J–V curves of the DSSCs employing various CEs irradiated from rear side and the detailed photovoltaic parameters are summarized in Table S1. The cell device with NiSe alloy CE yields an η of 7.35% (J$_{sc}$ = 17.52 mA cm$^{-2}$, V$_{oc}$ of 0.69 V, and FF of 60.8%) under simulated AM1.5, which is superior to the flexible DSSCs in literatures. Ni is a typical transition metal, having unfilled valence in d orbital. Therefore, the alloying of Ni with Se can accept electrons to form coordinated intermediates, which is prerequisite for robust CE materials. Due to a fact of having remarkable catalytic activities for Se and Ni, they have been widely used as electrocatalysts in driving fuel cells. To our knowledge, the measured η is in a very high level for liquid electrolyte based flexible DSSCs with Pt–free CEs. The η follows an order of NiSe > Ni$_{1/2}$Se > Ni$_{1/3}$Se > Ni$_{1/4}$Se > Ni$_{1/5}$Se > Ni$_{1/6}$Se > Ni$_{1/7}$Se because of a synergistic effect of catalytic activity, charge–transfer ability, and transparency. In order to investigate the dependence of TiO$_2$ thickness on photovoltaic parameters, the Ti foil has been etched to obtain various groove depths with a range of 0–66 µm. Fig. 2e exhibits the characteristic J–V curves of cell devices with FeSe CE and anodes at different TiO$_2$ film thickness. Different from traditional DSSCs, the incident light penetrate into the cell device from rear side (CE side), therefore thick TiO$_2$ is expected to increase N719 dye loading. From Table S2, it is apparent that the J$_{sc}$ has a peak value at TiO$_2$ thickness of 36 µm (Fig. S1). The elevated dye loading and therefore photogenerated electrons density on conduction band of TiO$_2$ have a promotion on J$_{sc}$ in a thickness range of 0–36 µm. However, a much higher TiO$_2$ thickness such as higher than 36 µm can cause two issues: (i) no light irradiation for adsorbed N719 dyes in the deep, (ii) the diffusion of I$_3^-$ species to groove bottom. A direct result is the decrease of interconversion between I$_3^-$ and I$^-$ and therefore dye recovery kinetics. If not specified, the thickness of TiO$_2$ layer is 36 µm. As a reference, the ITO/PEN supported Pt electrode was also prepared by the same cyclic voltammetry method, giving a comparable η of 2.75% (J$_{sc}$ = 6.03 mA cm$^{-2}$, V$_{oc}$ of 0.682 V, and FF of 66.9%), as shown in Fig. S2a. The low power conversion efficiency is mainly attributed to the high reflection toward incident light (Fig. 2c) because the pristine Pt electrode has similar catalytic activity with Ni$_{0.8}$Se alloy CE (Fig. S2b and Fig. S2c).

Fig. 3 (a) CV curves of various CEs for I$_3^-$ redox species recorded at a scan rate of 50 mV s$^{-1}$. (b) Bode and (c) Nyquist EIS plots and (d) Tafel polarization curves of the symmetric dummy cells from two identical CEs. The insets show magnified Nyquist EIS plots and an equivalent circuit.

A golden rule in evaluating the function of a CE for DSSC application is to compare catalytic activity for the reduction of I$_3^-$ ions. Fig. 3a displays cyclic voltammogram (CV) curves of Ni–Se alloy electrodes recorded at a scan rate of 50 mV s$^{-1}$, giving two pairs of oxidation/reduction peaks (Ox/Red), point current density of Red$_1$ and Ox$_{1}$/|Red$_1$| ratio are two key parameters for assessing the catalytic activity and reversibility of a CE toward I$_3^-$ redox couples. Due to a fact that the Red$_1$ peaks have similar positions, therefore the catalytic activities of alloy electrodes are NiSe > Ni$_{1/2}$Se > Ni$_{1/3}$Se > Ni$_{1/4}$Se > Ni$_{1/5}$Se > Ni$_{1/6}$Se > Ni$_{1/7}$Se > Ni$_{1/8}$Se > Ni$_{1/9}$Se > Ni$_{1/10}$Se > Ni$_{1/11}$Se > Ni$_{1/12}$Se > Ni$_{1/13}$Se > Ni$_{1/14}$Se > Ni$_{1/15}$Se > Ni$_{1/16}$Se > Ni$_{1/17}$Se > Ni$_{1/18}$Se > Ni$_{1/19}$Se. From the outward extension of the peak positions in stacking CV curves of NiSe alloy electrode along with linear relationships between square root of scan rate and peak current densities, as shown in Fig. S3, we can conclude that the redox reaction is a diffusion-controlled mechanism on alloy CE. From the Red$_1$ peak current density (J$_{red1}$), we can obtain diffusion coefficient (D$_{a}$) of I$_3^-$ redox couples at electrode surface by Randles–Sevcik theory ($J_{red1} = 2.69 \times 10^5 n^{3/2} A D_a^{1/2} v^{1/2} C^{lim}$), where D$_{a}$ = 2.82 × 10$^{-6}$ cm$^2$ s$^{-1}$ > Ni$_{1/11}$Se (1.97 × 10$^{-6}$ cm$^2$ s$^{-1}$) > Ni$_{1/12}$Se (1.80 × 10$^{-6}$ cm$^2$ s$^{-1}$) > Ni$_{1/13}$Se (1.31 × 10$^{-6}$ cm$^2$ s$^{-1}$) > Ni$_{1/14}$Se (1.25 × 10$^{-6}$ cm$^2$ s$^{-1}$) > Ni$_{1/15}$Se (7.42 × 10$^{-7}$ cm$^2$ s$^{-1}$) > Ni$_{1/16}$Se (7.29 × 10$^{-7}$ cm$^2$ s$^{-1}$), where n is number of electron participating in Red$_1$ reaction, A is active area, v is scan rate, and C is I$_3^-$ concentration. Notably, Ox$_{1}$/|Red$_1$| of alloy electrodes are calculated to be 0.995 for NiSe, 1.03 for Ni$_{1/2}$Se, 1.12 for Ni$_{1/3}$Se, 1.21 for Ni$_{1/4}$Se, 1.28 for Ni$_{1/5}$Se, 1.30 for Ni$_{1/6}$Se, and 1.32 for Ni$_{1/7}$Se. Higher peak current density and D$_{a}$ along with lower Ox$_{1}$/|Red$_1$| suggest that the NiSe alloy CE presents higher catalytic activity and better reversibility, which are paramount prerequisites for a robust CE in efficient DSSC. Bode EIS plots from symmetric dummy cells in Fig. 3b provides direct evidence to calculate the average reduction time of I$_3^-$ species. The electrons collected by alloy CE
from external circuit participate in reduction reaction of $I_3^- \rightarrow 3I_2$, therefore lifetime ($\tau$) of the electrons at electrolyte/CE interface can be utilized to assess the reaction kinetics of CE material. The electron lifetime can be obtained by $\tau = 1/2f$,\textsuperscript{34} giving average electron lifetime of NiSe (15.9 $\mu$s) $< \text{Ni}_1\text{Se}$ (19.1 $\mu$s) $< \text{Ni}_0\text{Se}$ (34.0 $\mu$s) $< \text{Ni}_0\text{Se}$ (61.9 $\mu$s) $< \text{Ni}_1\text{Se}$ (155.5 $\mu$s) $< \text{Ni}_2\text{Se}$ (235.4 $\mu$s) $< \text{Ni}_0\text{Se}$ (289.6 $\mu$s), where $f$ is frequency of the second peak. A short $\tau$ means that the reflux electrons (from external circuit to CE) can be facilely collected by alloy film and are involved in the $I_3^-$ reduction reaction ($I_3^- + 2e^- \rightarrow 3I_2$). To demonstrate the charge-transfer ability of alloy electrodes, Nyquist electrochemical impedance spectra (EIS) spectra in Fig. 3c are recording in symmetric dummy cells. After being fitted by Zview software, the related charge-transfer resistances ($R_{ct}$) obeys an order of NiSe (0.21 $\Omega$ cm$^2$) $< \text{Ni}_1\text{Se}$ (0.35 $\Omega$ cm$^2$) $< \text{Ni}_0\text{Se}$ (0.37 $\Omega$ cm$^2$) $< \text{Ni}_1\text{Se}$ (1.07 $\Omega$ cm$^2$) $< \text{Ni}_2\text{Se}$ (1.52 $\Omega$ cm$^2$) $< \text{Ni}_3\text{Se}$ (2.10 $\Omega$ cm$^2$) $< \text{Ni}_0\text{Se}$ (2.74 $\Omega$ cm$^2$). Tafel polarization measurements are utilized to reconfirm the catalytic activity and charge-transfer ability of the metal selenide alloy electrodes. As shown in Fig. 3d, the exchange current density ($J_0$) of NiSe film with a thickness of 36 $\mu$m provides $J_0 = RT/nF\pi f_{\text{ct}}$ derived from the slopes for anodic or cathodic branches in an order of NiSe $> \text{Ni}_1\text{Se} > \text{Ni}_0\text{Se} > \text{Ni}_3\text{Se} > \text{Ni}_2\text{Se} > \text{Ni}_1\text{Se} > \text{Ni}_0\text{Se}$.\textsuperscript{35} Additionally, the limiting diffusion current density ($J_{\text{lim}} = 2nFCD_{\text{ct}}/l$) generated from the intersection of cathodic branch with $Y$-axis also has the same sequence to $J_0$.$\textsuperscript{36}$ Till now, we can make a conclusion that the results from CV curves, lifetime analysis, EIS spectra, and Tafel polarization measurements are highly consistent.

Fast start–up, multiple start/stop cycling, and high stability are three prerequisites for the applications of solar panels in movable power sources and photovoltaic curtain walls. A critical issue in solving these problems is to explore efficient electrocatalyst for CE.$\textsuperscript{37,38}$ Fig. 4a shows on–off (start–stop) switch on the cell with NiSe electrode by alternatively irradiating the device with intensities of 100 and 0 mW cm$^{-2}$. An abrupt increase in current density and no delay in time suggests that the resultant NiSe alloy CE is vigorous in reducing $I_3^-$ species. As a reference, the related on–off switch of standard Pt electrode based DSSC is also provided, in which a lower current density means a lack of motivation in driving a DSSC. Additionally, start/stop cycling can be employed to evaluate the multiple start–up capability of a DSSC. After eight start/stop cycles, the cell still remain properties in its initial level, which is an essential prerequisite for the application of DSSCs in power sources. Under durable irradiation over 600 min, as shown in Fig. 4b, 101.3% and 99.8% of initial photocurrent densities are remained for the cells with NiSe and Pt electrodes, respectively. The result suggests relatively good stabilities for the solar cells with NiSe alloy and pristine Pt electrode, which is consistent with normalized efficiency in Fig. S4. Although the 600 min-test is far from to evaluate the long–term stability of a real cell device, the presented results demonstrate the potential realization of robust flexible DSSCs reported here.

### Conclusions

In summary, we have demonstrated that electrochemical deposition technique of Ni–Se alloys is an effective strategy for designing cost–effective and transparent CE materials and enhancing the photovoltaic performances of flexible DSSCs, whereas the seal of liquid electrolyte in etched groove of Ti foil can significantly enhance the electrolyte retention. NiSe alloy CE exhibits superior electrocatalytic activity toward $I_3^-$ species and charge–transfer ability. The DSSC assembled by NiSe alloy CE and Ti foil reserved TiO$_2$ film with a thickness of 36 $\mu$m provides an impressive power conversion efficiency of 7.35% for irradiation from rear side. Moreover, the virtues on fast start–up, multiple start/stop capability, and high stability motivate the potential applications of such flexible DSSCs in movable power sources and photovoltaic curtain walls. The research presented here is far from being optimized but the new concept along with cost–effective synthesis and scalable materials promise the CEs and anodes to be strong candidates in robust flexible DSSCs.

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