Light soaking effect driven in porphyrin dye-sensitized solar cells using 1D TiO2 nanotube photoanodes

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Abstract
© 2020 Light soaking (LS) effect on porphyrin (GD2) dye-sensitized solar cells (DSSCs) using well-ordered TiO2 nanotube (TNT) photoanodes was studied with different lengths of TNTs and LS treatment time. The TNT array possessing longer length and larger tube diameter had improved photoelectrochemical (PEC) property by generating larger photocurrent, and besides provided a larger surface area to yield more dye loading. The LS effect on GD2-applied DSSCs using TNT photoanodes was triggered off in the first 5 min of the LS treatment, and gradually increased during 60 min LS treatment. The improved PEC property and dye loading by longer TNT array (22 μm) led to the noticeable enhancement rate of power conversion efficiency from 0.7% to 1.88% (168% increase) after 60 min LS treatment, which was more prominent compared to the enhancement of GD2-applied DSSCs using mesoporous TiO2 films. Especially, morphological property of well-ordered 1D TNT photoanodes offers more spaces to facilitate the cation exchange in electrolyte system, leading to enhancing electron injection and reducing recombination under the LS condition. Electrochemical impedance spectroscopy (EIS) results confirmed the contribution of well-aligned 1D TNT structure to significant LS effect in GD2-applied DSSCs. Bode phase and Nyquist plots in EIS results showed the elongated electron lifetime and diffusion length with an increase in TNT length, followed by improving power conversion efficiency with significantly increased Jsc.

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Light soaking effect driven in porphyrin dye-sensitized solar cells using 1D TiO$_2$ nanotube photoanodes

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**A B S T R A C T**

Light soaking (LS) effect on porphyrin (GD2) dye-sensitized solar cells (DSSCs) using well-ordered TiO$_2$ nanotube (TNT) photoanodes was studied with different lengths of TNTs and LS treatment time. The TNT array possessing longer length and larger tube diameter had improved photoelectrochemical (PEC) property by generating larger photocurrent, and besides provided a larger surface area to yield more dye loading. The LS effect on GD2-applied DSSCs using TNT photoanodes was triggered off in the first 5 min of the LS treatment, and gradually increased during 60 min LS treatment. The improved PEC property and dye loading by longer TNT array (22 μm) led to the noticeable enhancement rate of power conversion efficiency from 0.7% to 1.88% (168% increase) after 60 min LS treatment, which was more prominent compared to the enhancement of GD2-applied DSSCs using mesoporous TiO$_2$ films. Especially, morphological property of well-ordered 1D TNT photoanodes offers more spaces to facilitate the cation exchange in electrolyte system, leading to enhancing electron injection and reducing recombination under the LS condition. Electrochemical impedance spectroscopy (EIS) results confirmed the contribution of well-aligned 1D TNT structure to significant LS effect in GD2-applied DSSCs. Bode phase and Nyquist plots in EIS results showed the elongated electron lifetime and diffusion length with an increase in TNT length, followed by improving power conversion efficiency with significantly increased $J_{sc}$.

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1. Introduction

In the dye-sensitized solar cell (DSSC) research field, TiO$_2$ is a key material as it is employed as a hole blocking layer and light absorbing dye-sensitized active layer in the DSSCs owing to its chemical durability, non-toxicity, and abundance [1,2]. In order to improve the DSSCs’ photovoltaic performance, many studies on the fabrication and modification of TiO$_2$ photoanodes have been reported. A variety of structures and morphologies of TiO$_2$ such as mesoporous, nanorod, nanotube and hollow sphere have been studied for DSSCs [3–6]. In fact, the structure and morphology of TiO$_2$ are the important parameters governing dye loading as well as light harvesting capacity, and thus determining the concentration of photogenerated electrons. Besides, the electron injection, transport and recombination rate can be influenced by the electronic and physical properties such as porosity, morphology, crystallinity, and uniformity of the TiO$_2$ structure [3,7–10]. Compared with the conventional DSSCs consisting in mesoporous TiO$_2$ nanoparticles, vertically well-ordered TiO$_2$ nanotube (TNT)-based DSSCs showed improved charge collection by efficiently suppressing the recombination of photogenerated electron–hole pairs through minimizing the concentration of trapping sites, which are normally located in the grain boundaries of randomly aggregated TiO$_2$ particulate films [11–14]. Thus, a unique structural property of TiO$_2$ leads to the improved charge transport property, resulting in increased the power conversion efficiency [15–19].

To date, various Ru-containing dyes, i.e. N719, N3, Z907, and Black dye, have been applied to DSSCs with TNT photoanodes [20–22]. Dubey et al. compared the photovoltaic performance of DSSCs with TNT photoanodes adsorbed with three different dyes such as N3, Z907 and Black dyes, and reported Z907 to show more enhanced performance because of the presence of a long alkyl group suppressing agglomeration [22]. However, such Ru-based dyes act as an adverse factor limiting the commercialization of DSSCs due to the high cost and low abundance of the noble Ru metal. On the other hand, the employment of inexpensive Zn-based porphyrin dyes in DSSCs has attracted great attention in replacing expensive noble Ru-based dyes with the improved efficiency of 13% [23]. Porphyrin dyes have also been explored to develop DSSCs using TNT photoanodes by controlling
functional groups and molecular structures of porphyrin [24,25], where both porphyrin binding geometry and TNT morphology play a critical role in charge-collection efficiency [26]. Mozer et al. has found that light soaking (LS) treatment triggered the significant enhancement of short-circuit current density (Jsc), open-circuit voltage (Voc), and fill factor (FF) in the case of GD2 type porphyrin dye-applied DSSCs. This was mainly attributed to the improvement of charge injection and reduced electron recombination, induced by aligning porphyrin dye molecules adsorbed on the surfaces of mesoporous TiO2 [27,28]. Considering the alignment of porphyrin dye molecules on TiO2 film plays a critical role in the LS effect, the different sizes distribution in inner tube diameter and alignment of the tube diameter changes the accessibility of electrolyte to the TiO2/dye surface, which may affect the kinetics of ion exchange at the interface. Therefore, it is expected that the LS effect using well-ordered TNT nanotubes is faster compared to mesoporous films. This motivated us to employ the unique 1D-TNT structure to the porphyrin dye-applied DSSCs and investigate the characteristics of the DSSCs with different pre-illumination duration prior to measurement of the photovoltaic performance.

In this work, we presented a significant LS effect of the porphyrin (GD2) dye-sensitized solar cells using 1D-TNT arrays as a photoanode under the controlled light soaking treatment time. First, photovoltaic-electrochemical (PEC) measurement was performed to evaluate capability of TNT arrays with different lengths and diameters as a photoanode, followed by employing TNT arrays as a photoanodes to DSSCs. In addition to the typical photovoltaic performance study, we investigated the LS effect by evaluating dye loading of TNT photoanodes with different lengths and performing electrochemical impedance spectroscopy (EIS) measurement to understand electron transport and recombination in GD2 dye system on TNT photoanodes.

2. Experimental

2.1. Fabrication of TNT-based DSSCs

The TNT arrays (6 × 6 mm active device area), obtained from 1, 5, and 15 h of anodization at 60 V with ethylene glycol containing 0.5 wt % NaF and 5 wt% H2O, were integrated into a DSSC device. Prior to dye adsorption, TNT array was treated with 0.04 M of TiCl4 at 70 °C for 30 min followed by rinsing with water. TiCl4 treated TNT was calcined at 450 °C for 3 h and reheated at 450 °C for 30 min if not immediately used. For porphyrin-DSSCs, the TNT array was also soaked in 0.2 mM porphyrin dye GD2 ((E) 3-(5,10,15,20-tetra(3,5-dimethylphenyl)porphyrin-2-yl)propenylenemalonato zinc (II)) solution in anhydrous tetrahydrofuran for 2 h. The GD2 dye was synthesized according to published procedures [29,30]. A sandwich-type DSSC was assembled using the dye-sensitized TNT array as a photoanode and platinum-deposited fluorine-doped tin oxide (FTO) glass (Asahi, Rs ≤ 8 Ω sq.−1) as a counter electrode separated by a sealant (Surlyn 60 μm thickness, Solaronix). The electrolyte was a mixture of 0.1 M LiI, 0.6 M 1, 2-dimethyl-3-propylimidazoliumiodide (DMPImI), 0.03 M I2 and 0.5 M t-butylpyridine (tBP) in acetonitrile [31]. The electrolyte was injected to the cell through a hole drilled through the counter electrode with the aid of a vacuum.

2.2. Characterization of TNT-based DSSCs

The current-voltage (I-V) curves of the DSSCs were measured using a Keithley 2400 source measure unit under a calibrated AM 1.5 solar illumination (Oriel) at 100 mW cm−2 light intensity. LS treatment was performed by exposing DSSCs under 100 mW cm−2 simulated AM 1.5 light at open-circuit for 5–60 min. During the treatment, the temperature of the solar cell was not controlled. The LS treated DSSCs were contained under the dark until I-V measurements in Fig. 3 and Fig. 5. The adsorbed dye concentration was determined by measuring the absorbance of dye solution desorbed from the surface of the TNT array in basic solution. In order to desorb dyes, GD2 dye-adsorbed TNT arrays were immersed in a 0.1 M NaOH in water and ethanol with 1 to 1 volume ratio and 0.01 M tetrabutylammonium hydroxide (TBAOH) in tetrahydrofuran (THF) for about 40 min, respectively. The absorbance measurement was performed using UV–Vis spectrophotometer (Cary 300, Varian). Electrochemical impedance spectroscopy (EIS) measurements were performed by illuminating the DSSCs with 100 mW cm−2 calibrated AM 1.5 illumination at open-circuit conditions between
0.1 Hz and 100 kHz with an AC amplitude of ±10 mV using a Gamry Reference 600 instrument.

3. Result and discussion

3.1. Photoelectrochemical performance of TNT photoanodes

Fig. 1 shows top view SEM images of well-ordered TNTs prepared by anodization. By varying anodization time of 1, 5, and 15 h, the lengths of TNTs were tuned at ca. 3, 10, and 22 μm, respectively. The inner tube diameter of TNTs was also changed from 50 nm to ~100 nm with an increase in anodization time (see Table 1). The lengths of TNTs presented a significant increase with an increase in anodization time.

Photoelectrochemical measurements were carried out using a three-electrode system in 0.1 M Na2SO4 electrolyte under UV illumination. Fig. 2a shows the photocurrent-voltage curves of TNT arrays obtained by different anodization time. TNT arrays were named as TNT1, TNT5, and TNT15 by anodization time of 1, 5, and 15 h, respectively. For all the TNT array samples, the photocurrent density increased with an increase in applied voltages, while the dark current was negligible under the applied voltages ranging from −1 V to 1 V vs Ag/AgCl. Besides, overall photocurrent density was higher for the TNT arrays obtained by a longer anodization time because of an increase in the length of TNT. The longer TNT array provides the larger TiO2 surface area, which can be photoexcited and interfaced with electrolyte, and thus it contributes to producing more photogenerated electrons and transporting them to the counter electrode during photoelectrocatalysis [32–34]. Fig. 2b shows photocurrent response of TNT arrays at 1 V vs Ag/AgCl. The photocurrent responses of all the samples are reproducible during the repetitive on-off light irradiation cycles. They present a spike of photocurrent followed by decay until a stationary-state is saturated. The saturated photocurrent response of TNT array exhibits the obvious enhancement for a longer length of TNT array. This result suggests the longer TNT array has enhanced photoelectrochemical property in terms of larger inner tube diameter of TNT array. The longer TNT array with larger inner tube diameter provides the larger surface area to afford more anchoring sites for dye-uptake, and thus the increased dye adsorption by the longer TNT array contributes to the improvement of light harvesting efficiency of DSSCs.

3.2. Light soaking effect of GD2 porphyrin-applied DSSCs using TNT photoanodes

The light soaking (LS) treatment was performed by exposing DSSCs under the simulated AM 1.5 light source of 100 mW cm−2. In fact, a LS treatment accelerates the degradation of the DSSC performance. For example, the typical N719-DSSCs showed the dramatic decrease in Jsc and Voc during LS treatment [35–37]. Wang et al. reported overall efficiency of N719-DSSCs decreased nearly 35% after LS treatment where main reasons are due to the desorption of dye molecules and the contamination of interfaces between electrolyte and TiO2 surfaces by a LS treatment condition [38,39]. Fig. S1 and Table S1 present that photovoltaic performances of typical N719-applied DSSCs using TNT15 photoanodes have decreased after the LS treatment. Indeed, this LS treatment result on N719-applied DSSCs shows good agreement with the previous reports [37–39].

In contrast to the N719-applied DSSCs, GD2-applied DSSCs show the different tendency in photovoltaic performance with the LS treatment where the LS effect led to the increase in photocurrent by increasing electron injection rate from the excited dye molecules to the TiO2 [40]. The photovoltaic performances of the GD2-applied DSSCs with three different TNT lengths (3, 10, and 22 μm in length) were measured before and after 60 min LS treatment by AM 1.5 solar simulator (100 mW cm−2) as seen in Fig. 3. Before LS treatment, the DSSCs using 3, 10, and 22 μm long TNT arrays presented the short-circuit current densities (Jsc) of 1.36, 2.23, and 2.69 mA cm−2, respectively, while the Voc values were comparable at 0.48–0.49 V regardless of the lengths of TNT arrays. After 60 min of LS treatment, all DSSCs presented significantly enhanced photovoltaic performances. Therein, the DSSCs using 3, 10, and 22 μm long TNT arrays showed the Jsc of 1.6, 4.58, and 5.60 mA cm−2, the Voc of 0.56, 0.55, and 0.53 V, and the power conversion efficiency of 0.55, 1.45, and 1.88%, respectively. From these results in Fig. 3 and Table 2, the increase in Jsc with the increase in the lengths of TNT arrays can be attributed to a relatively larger surface area to absorb a larger amount of dyes (see Table 1) and an efficient cation exchange between dye molecule and electrolyte. It will be further discussed in the following section. In the meantime, the dye loading amount of mesoporous TiO2 films (in 3–4 μm thick) was approximately 10 times higher (0.9 × 10−7 mol cm−2) than that of TNT arrays, and thus the initial device performance of TNT-based GD2-DSSCs was much lower compared to the mesoporous TiO2 film-based devices [41]. Instead, LS effect of TNT-based GD2-DSSCs showed a relatively large magnitude in device performance, particularly, Jsc, compared to mesoporous TiO2-based GD2-DSSCs, indicating the migration of the chemical species to TiO2 surfaces is facilitated by more sparsely packed surfaces in the reduced dye loaded TNT-based devices [31]. Meanwhile, the big increase in TNT length from 3 μm thick to 22 μm thick resulted in the slight decrease in Voc. Although it was not a significant change, it can be due to the electron back reaction from conduction band (CB) of TNT to I3− by the increase in the surface area of the TiO2 and the longer travel distance of the photo-injected electrons [42,43]. In our work, however, the slight decrease in Voc was not a limiting factor of the overall device performance, while it is important to note that the Voc increase was observed from all TNT lengths after LS treatment. It indicates LS treatment facilitates photo-injection from dye to TiO2 and retards recombination between Li+ ions and electrons by more efficient cation exchange (see Fig. 6).

Mozer et al. demonstrated the origin of the lower Voc of the GD2-applied DSSCs compared to the Voc of the N719-applied DSSCs by investigating electron transport properties of the N719- and the GD2-applied DSSCs. It was found that a significantly reduced electron lifetime originates from different intermolecular interaction between GD2 porphyrin dye / I3− and N719 dye / I3− [44]. In order to investigate the main parameter affecting the efficiency before and after LS treatment, the enhancement degree of photovoltaic performances was considered. In Fig. 4, after 60 min LS treatment, the enhancement of Voc was increased to ~15%, which was comparable among different lengths of TNT arrays, while Jsc increased significantly with an increase in the lengths of TNT arrays. The enhancement of Jsc was from 1.36 to 1.60 mA cm−2 (18% increase) for 3 μm, from 2.33 to 4.58 mA cm−2 (96% increase) for 10 μm, and from 2.69 to 5.60 mA cm−2 (108% increase) for 22 μm, accompanying drastically enhanced efficiencies: from 0.37 to 0.55% (48% increase) for 3 μm, from 0.59 to 1.45% (146% increase) for 10 μm, and from 0.70 to 1.88% (168% increase) for 22 μm. Therefore, the major reason of the improved efficiency after LS treatment can be ascribed to the significant increase in Voc.

Table 1

<table>
<thead>
<tr>
<th>Sample</th>
<th>Anodization time (h)</th>
<th>TNT length (μm)</th>
<th>Pore diameter (nm)</th>
<th>Dye loading (× 10−7 mol cm−2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNT1</td>
<td>1</td>
<td>3.32 ± 0.06</td>
<td>50.01 ± 3.35</td>
<td>0.08</td>
</tr>
<tr>
<td>TNT5</td>
<td>5</td>
<td>10.45 ± 0.53</td>
<td>78.61 ± 5.55</td>
<td>0.57</td>
</tr>
<tr>
<td>TNT15</td>
<td>15</td>
<td>22.32 ± 0.85</td>
<td>98.67 ± 4.87</td>
<td>0.74</td>
</tr>
</tbody>
</table>
Fig. 5 exhibits the photovoltaic performances of the GD2-applied DSSCs using TNT1, TNT5, and TNT15 arrays with different lengths under LS treatment duration. The DSSCs were initially measured under AM 1.5 solar simulator (100 mW cm\(^{-2}\)), treatment was performed by changing the illumination time on the same sample under AM 1.5 solar simulator (100 mW cm\(^{-2}\)), and re-measured at identified LS treatment time. Wagner et al. found that the improvement in the photovoltaic performances of the GD2-applied DSSCs occurred within the first 60 min LS treatment and the improved photovoltaic performances lasted for 3 months [27]. In this work, it was found that Voc, Jsc, and power conversion efficiency of DSSCs using TNT5 and TNT15 increased drastically within 5 min, and subsequently the values became gradually saturating, indicating the LS effect was triggered off at the very beginning of the LS treatment. For DSSCs with TNT1, however, it showed a less profound enhancement in Jsc and power conversion efficiency compared to other two samples. It is possible that the short TNT is already light soaked during the first JV measurement, and therefore the change by subsequent light soaking is less. In addition to the adsorbed dye amount effect, there may be another possible factor affecting the significant LS effect on 1D TNT photoanodes. Wagner et al.
reported 26% improved efficiency in average with GD2-applied mesoporous TiO2 photoanode system after 1 h LS treatment, while this work with GD2-applied 1D TNT photoanode system shows 168% efficiency enhancement from 0.70 to 1.88% on TNT15 sample. It indicates that such significant LS effect in this work may be due to facilitation of dye molecule rearrangement on unique 1D TNT structure rather than on mesoporous TiO2 films.

Herein, efficient ion migration between the rearranged dye molecules and electrolytes on 1D TNT photoanodes can lead to the decrease in recombination with redox couples and the improved electron injection, followed by enhancing efficiency with an increase in Jsc and Voc. Specifically, the correlation between charge transport at the interface of TiO2/dye/electrolyte and TNT morphology (inner tube diameter & length) will be further discussed in interpreting EIS result below.

As seen in Fig. 6, Griffith et al. proposed surface Li⁺ species, initially associated with a carboxyl group of two carboxyl groups in GD2 dye, prevent electron injection from Zn, and besides lead to rapid recombination. However, under the illumination, the created dye cations induce the exchange of Li⁺ with DMPIm⁺ by removing Li⁺ ions located onto the TiO2 surfaces through Columbic repulsion. Then, the empty spaces by the removal of Li⁺ enhance electron injection and limit recombination process. Herein, widening interparticle channels inside TiO2 electrode materials can contribute to enhancing mass transport of chemical species in electrolytes [47]. For instance, compared to the conventional mesoporous TiO2 films, well-aligned tubular channels of 1D-TNT arrays offer longer mean free paths, leading to efficiently penetrating dye molecules and electrolytes inside TNT arrays [48]. As the mean free path can be explained by electron transfer resistance (Rtr) along with TiO2 network and electron diffusion length (L), the electron diffusion length (L) is proportional to the mean free path [49]. Schmuki et al. reported around 20 times longer diffusion lengths of TNT arrays than that of TiO2 nanoparticle films, indicating 1D TNT arrays possess longer mean free path [50]. In the meantime, the mesoporous TiO2 film system with shorter mean free path would result in slower cation exchange by

Fig. 5. open-circuit voltage (Voc), short-circuit current density (Jsc), fill factor (FF) and efficiency of the GD2-applied DSSCs using TNT1 (3 μm), TNT5 (10 μm), and TNT15 (22 μm) photoanodes of different TNT lengths with an increase in LS time.

Fig. 6. Schematic diagrams on charge injection and recombination mechanism through cation exchange in (a) Li⁺-less condition of 1D TiO2 nanotube and (b) Li⁺-rich condition of randomly oriented mesoporous TiO2 particulate films, driven by light soaking treatment.
At increased thickness influenced by the porosity of the films (see Fig. 1). Lower charge density with increased TNT length could contribute to longer lifetime. Another possibility is reduction in interfacial recombination between the Ti foil substrate and the electrolyte in using longer TNTs due to better passivation of this interface at longer anodising condition [5,51]. Jennings et al. reported that the electron diffusion length \( L \) in eq. (4) in TiO2 nanotube-applied N719-DSSCs was much longer than that of the tube lengths [52]. The longer electron lifetime in the longer TNT array can lead to increasing electron diffusion length from eq. (4), followed by higher \( J_{sc} \) and FF. 

\[

t = \frac{1}{2nf_{peak}} 
\]

\[
\tau = (D\tau)^{1/2} = d(R_{rec}/R_{tr})^{1/2} 
\]

where \( \tau \), \( f_{peak} \), \( R_{rec} \), and \( d \) represent the electron lifetime in TiO2, the peak frequency at the second semicircle in Fig. 7b, recombination resistance, electron transfer resistance and TiO2 tube thickness, respectively. \( L \) and \( D \) represent electron diffusion length and diffusion coefficient, respectively [53]. In Fig. 7b, the internal resistance of the first semicircle (R1) at high frequency range on the Nyquist plots is attributed to the redox reaction at the electrolyte and counter electrode interface, while semicircle (R2) at low frequency range is attributed to charge recombination between TiO2 electrons and acceptor species in the electrolyte. Herein, the second larger semicircles decreased significantly with the increase in TNT lengths and inner tube diameters, with the R2 values of the TNT1, TNT5, and TNT15 were 129.6, 54.48, and 37.10 \( \Omega \), respectively, which were correspondent to \( R_{rec} \). In the meantime, \( R_{tr} \) was calculated from the 45 degree slope at the transition region between the first and the second semicircles where the \( R_{tr} \) values of TNT1, TNT5, and TNT15 were 39.72, 37.08, and 27.99 \( \Omega \), respectively. According to eq. (4), the electron diffusion lengths for TNT1, TNT5, and TNT15 were calculated to 6.97, 11.87, and 24.00 \( \mu m \), respectively, and then the calculated electron diffusion lengths were approximately 20–110% longer than the TiO2 nanotube thicknesses, indicating that the applied TiO2 nanotube thicknesses were long enough to transfer electrons in the system, and also recombination process was much slower than diffusion through the TNT array [54]. In the meantime, inner tube diameter difference might also be correlated with the charge recombination resistance at the interface of TiO2/dye/electrolyte. With the length differences, the inner tube diameters of TNT1, TNT3, and TNT5 arrays were 50 nm, 78 nm, and 98 nm, respectively. With the larger tube diameter, a semicircle (R2) of TNT15 is smaller than those of TNT1, TNT5, and TNT15 were 39.72, 37.08, and 27.99 \( \Omega \), respectively. According to eq. (4), \( L \) in eq. (4)) in TiO2 nanotube-applied N719-DSSCs was much longer than that of the tube lengths [52]. 

4. Conclusions

Porphyrin (GD2) dye-sensitized DSSCs using TNT array as a photoanode showed considerably enhanced photovoltaic performances under light soaking (LS) treatment. After LS treatment, the enhancement of short-circuit current density \( (J_{sc}) \) showed the greatest increase.
from 2.69 mA cm$^{-2}$ to 5.60 mA cm$^{-2}$ (108% increase) with an increase in TNT length, attracting the significant increase of energy conversion efficiency from 0.70% to 1.88% (168% increase) for the GD2-applied DSSCs using 22 μm long TNT photoanode, which was more improved than that of TiO$_2$ nanoparticle-based DSSCs [27]. It was because of faster electron diffusion length, less recombination, and longer electron diffusion length, followed by significantly improving photovoltaic performance. Meanwhile, the photovoltaic performance of GD2-applied DSSCs using TNT photoanodes showed time-dependent tendency with different LS treatment periods. The enhancement degree of photovoltaic performance with LS treatment was dramatically prompted within the beginning stage of 5 min LS treatment, followed by gradually saturated photovoltaic performance with an increase in LS treatment time. The EIS results indicated that the enhanced photovoltaic performance after LS treatment was due to the elongated electron lifetime and diffusion length and the reduced internal resistance with an increase in the length of TNT photoanodes.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.susmat.2020.e00165.

References


