



## A solid-state p–n tandem dye-sensitized solar cell†

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The development of p–n tandem dye-sensitized solar cells (t-DSCs) offers the potential for substantial open-circuit voltages, holding great promise for a wide range of applications, particularly in the fields of photovoltaics and photoelectrochemical devices. Most reported t-DSCs are liquid-based, and suffer from unsatisfactory stability due to the leakage of liquid electrolytes and photovoltage that is limited to the energy difference of the two utilized semiconductors. In this study, we present the first realization of a solid-state p–n tandem dye-sensitized solar cell that incorporates both p-type and n-type solid-state dye-sensitized solar cells (ssDSCs) by using a transparent indium-doped tin oxide (ITO) back contact for both sides. Notably, this tandem system shows a remarkable open-circuit voltage of 1.4 V, surpassing the constraints of its liquid-based counterparts. Although the performance variations between p-ssDSCs and n-ssDSCs hint at challenges related to charge recombination and the efficiency of p-ssDSCs, this study underscores the significant potential inherent in solid-state tandem configurations.

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## Introduction

Solar cells, with conversion of abundant and clean solar energy into electricity, have emerged as a prominent technology in the race to reduce our dependence on fossil fuels and combat climate change. Among the various photovoltaic technologies, dye-sensitized solar cells (DSCs) have garnered significant attention due to their cost-effectiveness, ease of fabrication, and tunable optical properties.<sup>1–4</sup> With the development of traditional n-type DSCs (n-DSCs) and p-type DSCs (p-DSCs) based on liquid electrolytes, the concept of p–n tandem DSCs has been proposed to improve cell efficiencies and also realized based on sharing a redox couple in liquid electrolytes.<sup>5–16</sup>

However, the liquid p–n tandem DSCs (t-DSCs) have been challenged by leakage of electrolyte and stability issues. Moreover, the theoretical photovoltage (cumulative in a series design) of the liquid p–n tandem DSC is also limited by the energy difference between the conduction band (CB) of n-type semiconductors, such as TiO<sub>2</sub>, used in n-DSCs and the valence band (VB) of p-type semiconductors, such as NiO, used in p-DSCs, typically *ca.* 1 V (Fig. 1A).<sup>9,10</sup> To overcome these hurdles, we previously proposed the concept and design principle of p–n tandem solid-state dye-sensitized solar cells (t-

ssDSCs)<sup>17</sup> in which a hole transport material (HTM) and electron transport material (ETM) are used to replace the liquid electrolyte to extract holes and electrons from the dye-sensitized photoanode and dye-sensitized photocathode, respectively. If both photoelectrodes are then connected by a transparent conductor, it has been proposed that the photovoltage ( $V_{oc}^{ss}$ ) of p–n t-ssDSCs should be able to overcome the photovoltage limit in liquid p–n t-DSCs ( $V_{oc}^{liquid}$ ) since the two half cells do not need to share the same redox couple, possibly reaching up to 2 V with suitable materials and device optimization (Fig. 1B).<sup>17</sup> However, until now, the concept has not been realized to show that such a device really could surpass the limitations of liquid t-DSCs.

Owing to the development of solid-state n-type DSCs (n-ssDSCs)<sup>18–20</sup> and recent good progress on solid-state p-type DSCs (p-ssDSCs),<sup>21–26</sup> the time is ripe to combine p- and n-type ssDSCs to realize the first p–n-type solid-state tandem device. Conventional designs of ssDSCs feature a metal back contact, meaning incident light is obstructed from one side of the photoelectrode stack. The incorporation of transparent counter electrodes in ssDSCs offers a dual benefit: it not only facilitates the penetration of incident light, a crucial aspect for various DSC applications like windows<sup>27,28</sup> or agrivoltaics,<sup>29–31</sup> but it also unlocks the potential for the fabrication of a tandem device. While semi-transparent n-ssDSCs have been demonstrated,<sup>32–35</sup> a transparent back contact has not been realized in p-ssDSCs until now. Inspired by the work using indium doped tin oxide (ITO) as a back contact in n-ssDSCs<sup>32</sup> and the transparent conductive layer in tandem solar cells,<sup>36–39</sup> we herein present the first realization of a p–n t-ssDSC that incorporates both a p-ssDSC and n-ssDSC connected by a transparent ITO conductive layer in a series configuration. The resulting p–n t-ssDSC is semi-transparent and boasts an open-circuit voltage of 1.4 V,

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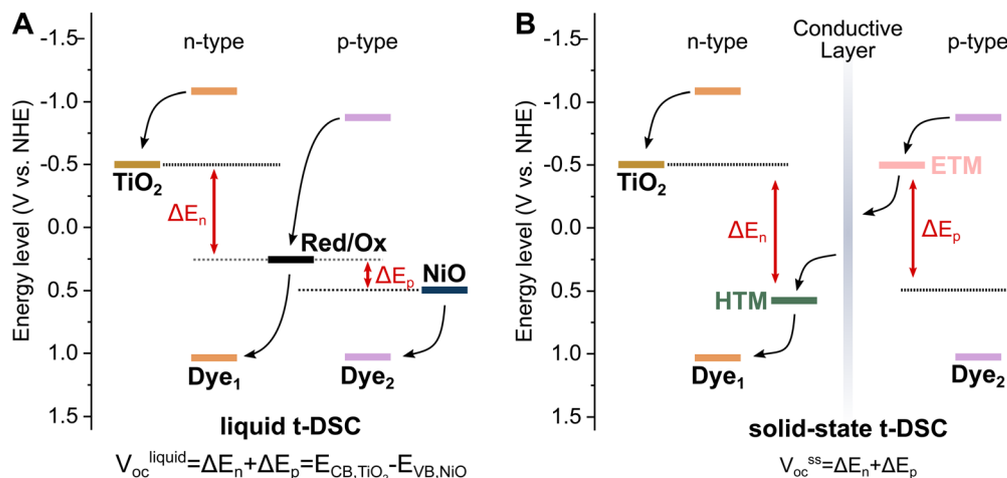


Fig. 1 Potential diagram for a (A) liquid and (B) solid-state p–n tandem DSC.  $\Delta E_n$ : theoretical photovoltage of an n-DSC;  $\Delta E_p$ : theoretical photovoltage of a p-DSC;  $E_{CB,TiO_2}$ : the conduction band of  $TiO_2$ ;  $E_{VB,NiO}$ : the valence band of  $NiO$ .

surpassing the theoretical voltage limitation of its liquid counterpart sharing one redox couple.

## Experimental section

### Materials

All chemicals were purchased from Sigma-Aldrich and used directly unless stated otherwise. Fluorine-doped tin-oxide coated glass (FTO, TEC15) was purchased from Pilkington.  $TiO_2$  paste (Dyesol, 30NR-T) was purchased from Dyesol. D35 was purchased from Dyenamo. Spiro-OMeTAD (sublimed grade) was purchased from Borun New Material Technology. The  $NiO$  screen printing paste<sup>10</sup> and PB6 photosensitizer<sup>24</sup> were lab-made and available from a previous study.

### Device fabrication

**Preparation of  $NiO$  films for p-ssDSCs.** Fluorine-doped tin oxide (FTO) glass substrates were etched with 2 M hydrochloric acid and zinc powder and masked with tape to obtain the desired electrode pattern. Afterwards, the etched FTO substrates were cleaned in a sonication bath in the following order for 30 min each: 2% detergent in water, deionized water, acetone, and ethanol. The compact  $NiO$  blocking layer ( $NiO$ -BL) was sputtered onto the cleaned FTO substrates by reactive magnetron sputtering (CS730S, Von Ardenne) according to the literature.<sup>40</sup> In short, a  $Ni$  target was used and the deposition was carried out for 250 s at a power of 500 W with a processing pressure of 6 mTorr in an atmosphere of  $Ar$  (25 sscm) and  $O_2$  (25 sscm) which resulted in a thickness of 80–100 nm compact  $NiO$ . The film was annealed at 450 °C for 1 h in an ambient atmosphere and cooled down to room temperature. One layer of  $NiO$  screen printing paste was then deposited on the blocking layer and dried on a hotplate at 115 °C for 5 min. The screen-printed  $NiO$  film was then sintered at 450 °C for 30 min in an ambient atmosphere and cooled down to 60 °C overnight. Afterwards, the electrode was sensitized in a 0.2 mM PB6 in *tert*-butanol/acetonitrile (1 : 1 v/v) solution overnight. The sensitized films

were rinsed with dichloromethane and dried with an  $N_2$  flow in order to eliminate physically adsorbed dye molecules on the surface.

Subsequently, atomic layer deposition (ALD) was employed to deposit  $ZnO$  on the sensitized electrodes. The films were masked with tape to obtain the desired device structure (see the ESI, Fig. S1†). First, 40 cycles of a stop-flow sequence were used to accommodate the mesoporous nature of the  $NiO$  film before 600 cycles of the “normal” sequence were used. The precursors for ALD were diethylzinc (DEZn) and water ( $H_2O$ ) and the temperature in the chamber was 120 °C (stabilization time of 1 h before the start of the deposition process). The stop flow involved a sequence of (1) reduced carrier gas flow, (2) reduced pumping speed through a bypass constriction, (3) precursor injection, (4) precursor exposure, (5) restored pumping speed, and (6) restored carrier gas flow and purging. For the DEZn precursor, the durations of steps 1 to 6 were 2, 2, 0.2, 19.8, 2, and 4 s. During this sequence, the carrier gas flow was reduced from 50 to 10 standard cubic centimeters per minute (scm) in each of the five unused precursor lines of the system. The TMA line used a flow of 150 scm during the injection step. In the  $H_2O$  half-cycle, the duration of steps 1 to 6 was the same, that is, 2, 2, 0.2, 19.8, 2, and 4 s, while the  $H_2O$  carrier gas flow was increased to 200 scm and the other precursor line flows were reduced to 10 scm.

To obtain an extra layer of  $ZnO$ , a  $Zn(acac)_2$  solution (0.33 M in methanol, filtered using a 0.45  $\mu m$  syringe filter) was spin-coated onto the  $ZnO$  ALD layer (90  $\mu L$ , 3 s acceleration to 2000 rpm, then 200 s at 2000 rpm). Three layers were deposited and after each layer, the electrode was put on the hotplate for 5 min at 200 °C. After the final layer, the film was put on the hotplate for an additional 15 min at 200 °C. An illustration of the device layout can be found in the ESI, Fig. S1.†

**Preparation of  $TiO_2$  films for n-ssDSCs.** The n-ssDSCs were prepared on the basis of published work by Chiang *et al.*<sup>32</sup> and Xu *et al.*<sup>41</sup> A compact  $TiO_2$  blocking layer was deposited by spray pyrolysis with an airbrush onto etched and cleaned FTO glass



substrates on a hotplate at 450 °C. Before deposition, the samples were ozone treated for 15 min and preheated for 15 min at 450 °C. For an area of 100 cm<sup>2</sup>, 7 mL of a solution of 0.6 mL titanium diisopropoxide bis(acetylacetonate), 0.4 mL acetylacetonate and 9 mL pure ethanol was used. The films were left to anneal on the hotplate for 30 min and then cooled down to room temperature slowly under ambient air. For the mesoporous TiO<sub>2</sub> layer, a diluted TiO<sub>2</sub> paste (1 : 3 ethanol to Dyesol, 30NR-T w/w) was used. Before deposition *via* spin-coating (150 μL, 10 s acceleration to 3000 rpm, then 30 s at 3000 rpm), the films were ozone treated for 10 min. The spin coating procedure was repeated the desired times and after each layer, the films were allowed to dry on a hotplate at 115 °C for 5 min. Following the sintering process of the TiO<sub>2</sub> film on a hotplate at 500 °C for 30 min, the film was allowed to cool to room temperature. Subsequently, it was submerged in a 0.02 M aqueous TiCl<sub>4</sub> solution at 70 °C for 30 min. After rinsing with deionized water, the film was annealed on a hotplate at 500 °C in an ambient atmosphere for 30 min. Once it cooled down to 60 °C, the film was placed in a 0.2 mM D35 in *tert*-butanol/acetonitrile (1 : 1 v/v) solution overnight for sensitization.

After rinsing the sensitized electrodes with ethanol and allowing them to dry, a chlorobenzene solution containing 150 mM SPIRO-OMeTAD, 200 mM 4-*tert*-butylpyridine and 20 mM LiTFSI was applied to create the HTM film. This was achieved by allowing 80 μL of the solution to permeate the sensitized electrode for 30 s and then spin-coating for 30 s at 2000 rpm. Following this, the cells were left in the dark in an air environment for 3 days. An illustration of the device structure can be found in the ESI, Fig. S1†

### Preparation of the transparent back contact for p-ssDSCs and n-ssDSCs

For p-ssDSCs and n-ssDSCs, the transparent ITO back contact electrode was deposited on the ZnO and SPIRO of the respective NiO and TiO<sub>2</sub> films. The deposition was done *via* direct-current magnetron sputtering from a ceramic ITO (90/10, purity 99.99%) sputtering target in a deposition system, CS730 (Von Ardenne). The depositions were carried out for 120 s at a power of 640 W, corresponding to 2 W cm<sup>-2</sup>, in an Ar + O<sub>2</sub> atmosphere (1.8% O<sub>2</sub>) at a total pressure of 0.6 Pa which resulted in an ITO layer with a thickness of 200–250 nm.

For better contact with the crocodile clamps used during device characterization, a small spot of silver paste was deposited on the contacts.

### Assembly of p–n t-ssDSCs

The p-ssDSC and n-ssDSC were pressed together with clamps with the p-ssDSC on the top, avoiding contact between the FTO layers of the two films (ESI, Fig. S1†).

### Solar cell characterization

Absorption spectra were recorded with a diode array spectrophotometer (Agilent 8453) with a home-made film holder.

The solar cells were evaluated under an irradiation intensity of 100 mW cm<sup>-2</sup>, utilizing a Wave Labs SINUS-70 solar

simulator that included UV and IR range extenders. In the *J*–*V* measurements, a black mask with a 0.2 cm<sup>2</sup> aperture area was employed and the voltage was screened in 0.01 V increments with a voltage settling time of 0.1 s. IPCE measurements (ESI, Fig. S6†) were carried out using a setup consisting of a xenon lamp (Spectral Products ASB-XE-175), a monochromator (Spectral Products CM110), and a LabJack U6 data acquisition board.

## Results and discussion

The fabricated p–n tandem solid-state dye-sensitized solar cell in our study incorporated well-established materials, such as D35<sup>33,42–44</sup> and PB6<sup>22–24</sup> dyes for sensitizing TiO<sub>2</sub> and NiO films, respectively (Fig. 2A). The configuration of the p–n t-ssDSCs, energy alignment of each component, and the individual p-ssDSC and n-ssDSC are illustrated in Fig. 2B and 3. On the basis of our previous work on p-ssDSCs,<sup>22,23</sup> we used a compact NiO layer on FTO glass to avoid short circuit with the back contact onto which a PB6-sensitized mesoporous NiO layer was deposited, as shown in Fig. 2. As an electron transport material (ETM), ZnO was chosen since previous studies<sup>23,45</sup> show the highest photovoltage with ZnO while maintaining a decent photocurrent (see the ESI, Table S1†). With the hopes of achieving a good pore filling, ZnO was deposited *via* atomic layer deposition (ALD) where we first used a stop-flow sequence to accommodate the mesoporous nature of the NiO film to deposit a thin layer of ZnO (~6 nm) before we deposited approx. 100 nm ZnO with “normal” ALD. To protect the film from short-circuits caused by the sputtering of the ITO back contact,<sup>32</sup> we added an extra layer of spin-coated Zn(acac)<sub>2</sub> that was transformed to ZnO upon heat exposure at 200 °C.<sup>23</sup> We found that the combination of ZnO from ALD with the extra layers from Zn(acac)<sub>2</sub> solution yielded the best performing cells (ESI, Fig. S2†). This is likely due to a combination of a good pore filling from the ALD with the extra protection of the additional ZnO layer during the consequent sputtering process. Aging was also found to be an important part to get good working cells, and storage in the dark in ambient air improved cell performance significantly (ESI, Fig. S2†), corresponding with previous literature.<sup>23,45</sup>

Inspired by previous work<sup>33,41,46</sup> the n-ssDSC was fabricated from a D35-sensitized mesoporous TiO<sub>2</sub> layer on a compact TiO<sub>2</sub> blocking layer (Fig. 3). D35 was chosen over other dyes due

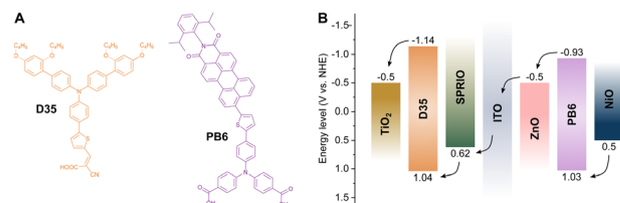


Fig. 2 Dye structure (A) and energy level diagram (B) of the components in the p–n t-ssDSC stack. For the energy levels of PB6<sup>24</sup> and D35<sup>44</sup> dyes, the LUMO energy was obtained from the reported ground state oxidation potential (HOMO) and E<sub>0–0</sub> transition energy. Spiro-OMeTAD<sup>45</sup> energy levels were taken directly from the literature.



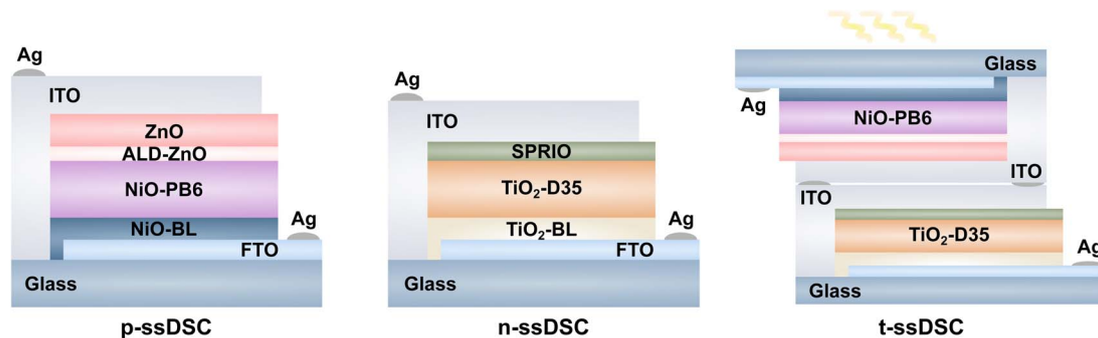


Fig. 3 Device structure of p-ssDSC, n-ssDSC and p-n t-ssDSCs.

to its favorable absorption wavelength with PB6 in comparison with other commercial dyes and exhibits typically a good photovoltage and current. The hole-transporting material (HTM), a lithium doped Spiro-OMeTAD solution (SPIRO), was spin-coated on the dye-sensitized film.<sup>41</sup> The transparent ITO back contact was then directly deposited on the HTM.

To assemble the p-n tandem cell, the p-ssDSC cell was placed on the top due to its lower photocurrent compared to the n-ssDSCs. Even the current published record photocurrent for p-ssDSCs, with a short-circuit photocurrent density ( $J_{sc}$ ) of 1 mA  $\text{cm}^{-2}$ , is still considerably lower than for n-ssDSCs.<sup>25</sup> The t-ssDSCs are therefore illuminated from the p-type side since the overall

photocurrent is limited by the cell with the lowest current in a series connection, which limits the overall performance in our case (Fig. 3). We additionally adjusted the thickness of the  $\text{TiO}_2$  in the n-ssDSC to match the photocurrent when shaded by the p-ssDSC on top to suppress unnecessary photo annihilation (ESI, Fig. S3 and S4<sup>†</sup>). Two layers of the spin-coated *meso*- $\text{TiO}_2$  solution proved to be ideal to match the  $J_{sc}$  of the p-ssDSC of around 0.8 mA  $\text{cm}^{-2}$  when shaded by the p-ssDSC on top. This is a critical step in optimizing the performance of the tandem cell to reduce losses and photo annihilation. The resulting solar cells are clearly semi-transparent with a pink color for the p-ssDSC and yellow color for the n-ssDSC (insets in Fig. 4A).

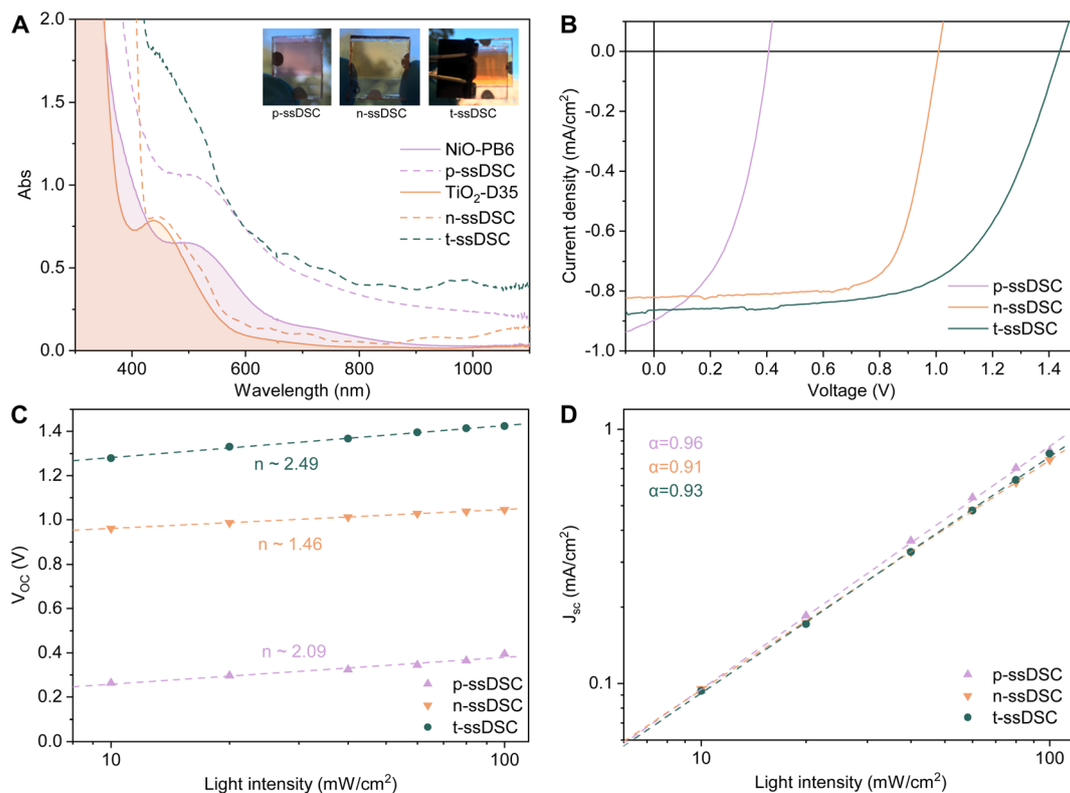


Fig. 4 (A) Absorption spectra of dye-sensitized NiO and  $\text{TiO}_2$  films as well as the fabricated solar cell devices, with pictures of the ss-DSCs as insets. (B)  $J$ - $V$  curves of the champion cells. Note that the p-ssDSC is illuminated from the glass side and that the n-ssDSC is shaded by an overlaying p-ssDSC film during measurement. Light intensity dependence of the (C)  $V_{oc}$  with the light ideality factor  $n$  and (D)  $J_{sc}$  with the power law factor  $\alpha$ .



The tandem device showed an orange color but also remained semi-transparent. Absorption spectra were recorded to evaluate the optical properties of all fabricated solar cells. The absorption spectra were obtained for both the dye-sensitized films and the complete solar cells and are shown in Fig. 4A. In the dye-sensitized film, we can clearly see the characteristic and complementary absorption of PB6 and D35 on film and in the final devices, as well as in the IPCE (ESI, Fig. S6†). Upon comparing the absorption spectra of the pure sensitized films to those of the functioning solar cell, we observed that the dyes remained stable during the cell fabrication process. In the absorption spectra of the n-ssDSC, we can clearly see the contributions of the Spiro-OMeTAD HTM which absorbs below 400 nm. Furthermore, it is evident that there is a baseline shift in the absorption spectra for the PB6-based solar cell. This shift can primarily be attributed to the scattering of light by the ZnO/ITO interfacial layer, a phenomenon that could also be visually perceived.

The  $J$ - $V$  curves of the champion devices are presented in Fig. 4B, with corresponding photovoltaic performance parameters detailed in Table 1. In the tandem cell, we achieved a power conversion efficiency (PCE) of 0.75%, a  $J_{sc}$  of  $0.86 \text{ mA cm}^{-2}$  and an exceptional open-circuit voltage ( $V_{oc}$ ) of 1.437 V. Notably, the  $J_{sc}$  values of the p- and n-ssDSCs in the tandem configuration align remarkably well, a critical requirement to suppress the unnecessary photo annihilation for tandem devices in series. In this setup, the overall current is governed by the lowest current cell, while the voltages are additive. Our findings clearly demonstrate that the  $V_{oc}$  of the tandem device (1.437 V) is the sum of the individual  $V_{oc}$  values of the p-ssDSCs (0.407 V) and n-

ssDSCs (1.000 V), and is not limited by the difference of the valence band of the photocathode and conduction band of the photoanode like in the liquid-state tandem cell. It is also worth noting an intriguing observation in the tandem device – there is a subtle increase of the current when compared to the n-ssDSC (Fig. 4B). This discrepancy, observed across all fabricated tandem devices, appears to be linked to the scattering effect introduced by the ZnO/ITO layer of the p-ssDSC and the ITO layer of the n-ssDSC. This phenomenon can boost the light absorption by n-ssDSCs and contributes to the amplified current in the tandem configuration.

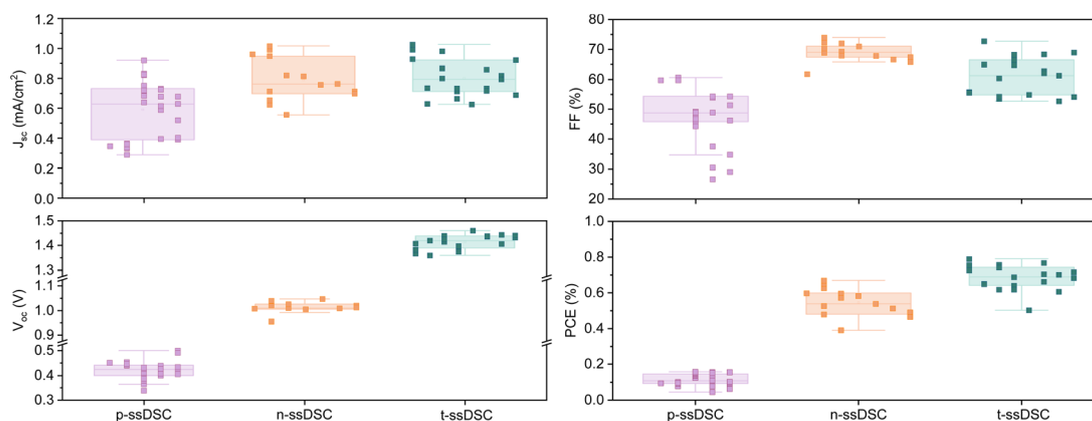
To further investigate the fabricated solar cells, light intensity dependencies were recorded. When looking at the dependency of the  $V_{oc}$  with light intensity in Fig. 4C, we observe that the p-ssDSC shows a high light ideality factor ( $n$ ) of 2.09 which is indicative of trap assisted recombination that is typically also reflected in a lower fill factor (FF). The light ideality factor of the n-ssDSC is significantly lower at 1.46. Notably, the p-ssDSCs without the extra spin-coated ZnO layer in addition to ALD showed a higher FF (ESI, Fig. S2†) which indicates that additional transport losses may arise from the extra ZnO layer, despite the higher PCE of the p-ssDSCs with the extra ZnO layer.

Interestingly, the increase of  $J_{sc}$  with light intensity ( $I$ ), that is generally described by  $J_{sc} \propto I^\alpha$ , shows a quasi linear behavior for all solar cells (Fig. 4D). A value of  $\alpha = 1$  indicates that all separated free carriers are extracted before recombination, and  $\alpha < 1$  indicates that  $J_{sc}$  depends on bimolecular recombination (*i.e.*, carriers recombine before being extracted).<sup>47</sup> Notably, the obtained  $\alpha = 0.96$  from the slope of the  $\log(J_{sc})$  versus  $\log(I)$  plot for the p-ssDSC is much better than in a previous study with NiO–PB6 cells that only used a ZnO layer from  $\text{Zn}(\text{acac})_2$ .<sup>24</sup> This indicates that the ALD ZnO layer achieves a good pore filling and reduces the Langevin recombination between the photo-generated charge carriers, likely due to better charge extraction and transporting abilities.

The spread of  $J$ - $V$  parameters of the manufactured devices for p-ssDSC, n-ssDSC and the tandem devices is shown in Fig. 5. It is visible that the p-ssDSCs exhibit higher photovoltaic performance deviation in general and against the sample position (vertically aligned points) compared to the n-type cells.

**Table 1** Metrics for the champion devices. Note that the p-ssDSC was measured from the FTO side and that the n-ssDSC was covered by p-ssDSC during measurement

Device	PCE (%)	FF (%)	$J_{sc}$ ( $\text{mA cm}^{-2}$ )	$V_{oc}$ (V)
p-ssDSC	0.166	45.6	0.89	0.407
n-ssDSC	0.595	71.9	0.82	1.000
t-ssDSC	0.766	62.0	0.86	1.437



**Fig. 5**  $J$ - $V$  parameters of the manufactured p-ssDSCs, n-ssDSC and t-ssDSCs under 1 sun illumination.



This significant discrepancy, particularly in the  $J_{sc}$  and FF of the p-ssDSC devices, also suggests potential issues related to solar cell fabrication leading to transport losses and recombination.

Furthermore, the shunt resistance of the p-ssDSC devices is notably lower, measuring  $2.7 \text{ k}\Omega \text{ cm}^2$ , as compared to the n-ssDSCs with a shunt resistance of  $15 \text{ k}\Omega \text{ cm}^2$ . A low shunt resistance is likely caused by the presence of pin-hole contacts within the p-ssDSCs due to the manufacturing, providing an alternative path for photo-generated current. This also contributes to the observed variations in device performance depending on spot selection in the p-ssDSCs.

The series resistances for the p-ssDSC ( $156 \Omega \text{ cm}^2$ ) and n-ssDSC ( $200 \Omega \text{ cm}^2$ ) are similar. As expected, the series resistance of the tandem cell ( $400 \Omega \text{ cm}^2$ ) is close to the sum of the series resistances of the individual devices. However, there is still some additional resistance originating from purely pressing the two cells together in the tandem device.

Despite the challenges and limitations associated with p-ssDSCs in general (see the ESI, Table S1†), and also limited efficiency of the overall device in this study, our results highlight that the open-circuit voltage of a p-n t-ssDSC can surpass the energy difference between the NiO valence band and TiO<sub>2</sub> conduction band, approximately 1 V (Fig. 1), which is the theoretical photovoltage limit of p-n liquid t-DSCs sharing one redox couple.<sup>17</sup> This achievement confirms the potential of solid-state tandem configurations and their capacity to harness the combined open-circuit voltages of the constituent devices.

## Conclusion

In summary, we report the first realization of solid-state p-n tandem dye sensitized solar cells (t-ssDSCs) constructed from a PB6 dye-based p-ssDSC and D35 dye-based n-type DSC with indium tin oxide (ITO) as the back contact. With a good match of dye absorption between PB6 and D35, the solid-state tandem DSC system yielded a short-circuit photocurrent density of up to  $0.86 \text{ mA cm}^{-2}$  with a remarkably high open-circuit voltage of up to 1.4 V, which surpasses the theoretical voltage limitations of its liquid counterpart sharing one redox couple. This work represents a significant milestone in the advancement of p-n t-ssDSCs. These innovations hold great promise across a range of applications within the fields of photovoltaics and photo-electrochemical devices, paving the way for sustainable and efficient energy conversion technologies.

## Conflicts of interest

The authors declare no competing financial interest.

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