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A naphthalene diimide side-chain polymer as an electron-extraction layer for stable perovskite solar cells†

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Poly(*N*-(5-(5-norbornene-2-carbonyloxy)pentyl)-*N'*-*n*-hexyl-naphthalene-1,8:4,5-bis(dicarboximide)) has been synthesized by esterification of (*N*-(5-hydroxypentyl)-*N'*-*n*-hexyl-naphthalene-1,8:4,5-bis(dicarboximide)) with *exo*-5-norbornene-2-carboxylic acid, and has been polymerized using the first-generation Grubbs initiator. This side-chain polymer exhibits good transparency throughout the visible (absorption onset at ca. 400 nm), good solubility in common low- and medium-polarity organic solvents, good resistance to dimethylformamide, and appropriate electron affinity for use as an electron-extraction layer in lead-halide perovskite solar cells. The performance of this polymer in *n*-i-p perovskite solar cells was compared to that of several small-molecule naphthalene-1,8:4,5-bis(dicarboximide) derivatives and of SnO₂. Solar cells using the polymer exhibited open-circuit voltages of up to 1.02 V, short-circuit currents of over 21 mA cm⁻², and power conversion efficiencies (PCE) reaching 14% which stabilize at 13.8% upon 90 s of illumination. Meanwhile control SnO₂ devices exhibited a PCE of ca. 16%, and small-molecule devices gave PCE values of less than 10%. The devices employing the polymer exhibited improved long-term stability relative to the SnO₂ control devices under continuous illumination.

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Introduction

Lead-halide perovskite solar cells (PSCs) have received increasing attention since their debut in 2009.^{1,2} In recent years, further optimizations of the various layers and interfaces have allowed the emergence of PSCs with efficiencies exceeding 25%.³ Hole- and electron-transport materials (HTMs and ETMs, respectively) in PSCs serve not only to transport charges, but also to improve the stability of the cells and to reduce charge recombination. A variety of organic and inorganic ETMs have been incorporated into *n*-i-p (negative-intrinsic-positive) and *p*-i-n (positive-intrinsic-negative) PSCs.^{4–7}

The “regular” (*n*-i-p) architecture for PSCs requires the ETM to be deposited prior to the perovskite, and, therefore, for solution-processed cells, to be insoluble in the perovskite

precursor solution. Insoluble metal oxides (*e.g.*, SnO₂, TiO₂) offer one possible solution.^{8–10} Although many approaches require high-temperature (often 450–500 °C for TiO₂^{8,9}) sintering subsequent to casting the precursors from solution, limiting compatibility with flexible substrates, recently room-temperature methods for the deposition of efficient SnO₂ layers have been reported.¹¹ Nevertheless, these oxides can possess trap states that hinder charge extraction and TiO₂ can even act as a photocatalyst. There is also considerable interest in organic ETMs as alternatives that can be processed at low temperature, while also exhibiting good charge-transport abilities and transparency.^{12–15} These include both vacuum-processable and solution-processed materials;¹⁶ in some cases solution-processable ETMs have been crosslinked subsequent to deposition to prevent their dissolution during subsequent perovskite deposition.^{17,18} Although fullerenes have been widely used, non-fullerene electron acceptors with similar reduction potentials may be advantageous in terms of transparency, cost, and/or tunability of reduction potential. Rylene diimides have attracted attention as ETMs for *n*-i-p cells. These include perylene diimides, such as B (Fig. 1), and related thermally evaporable small molecules,¹⁹ solution-processable small molecules,²⁰ and solution-processable polymers.²¹

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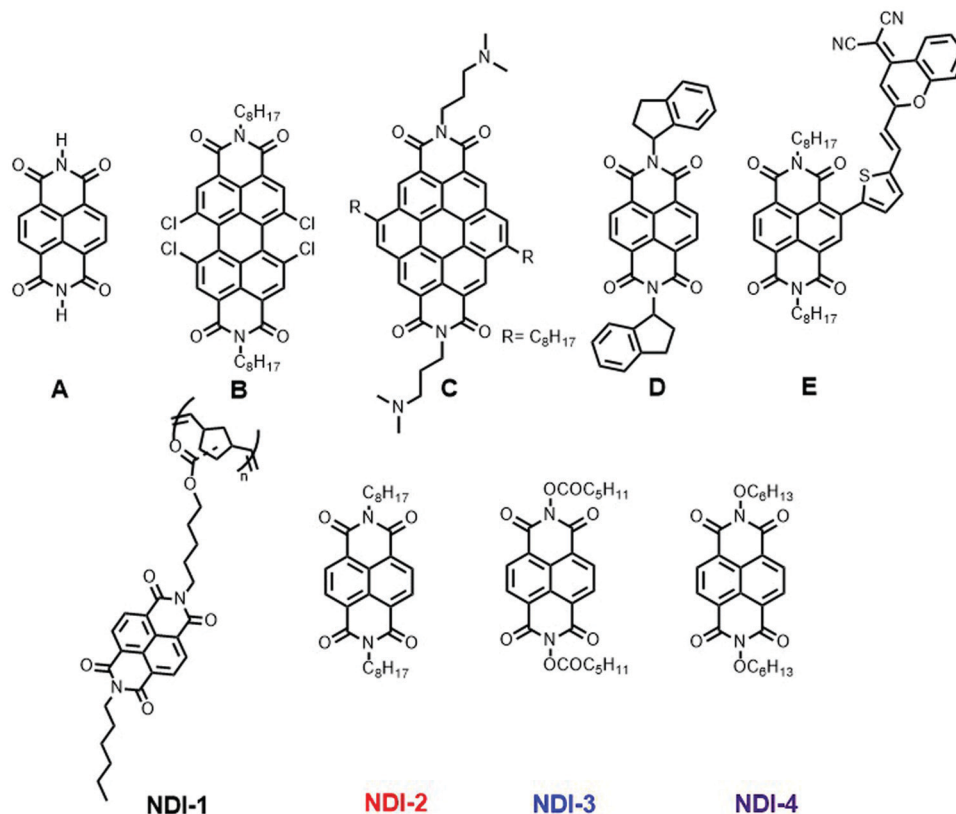


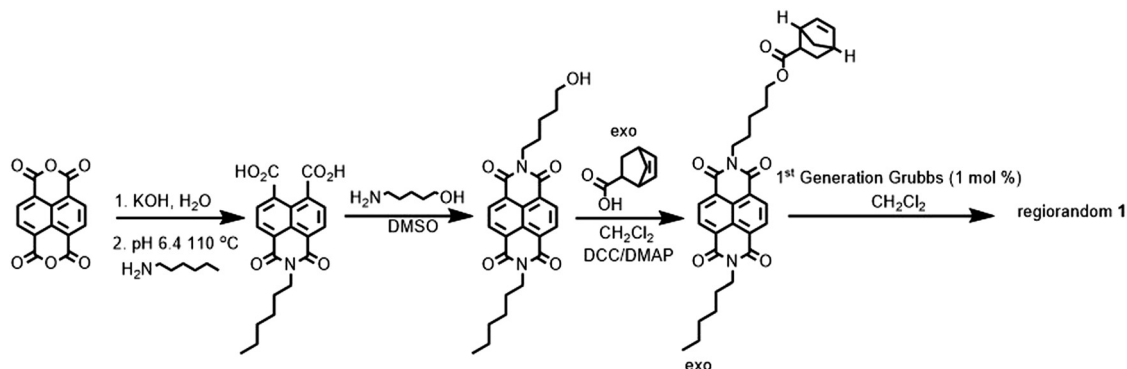
Fig. 1 Examples of rylene diimide ETMs reported in the literature (A–E) in the context of PSCs and NDI-based ETMs used in this work (1–4).

Naphthalene diimide (NDI) derivatives (such as D and E, Fig. 1) have been used in p-i-n devices,^{22–25} but there are few reports of their use in n-i-p devices. However, in one study an insoluble NDI (A, Fig. 1) was generated *via* high-annealing treatment of soluble solution-cast precursor for use in n-i-p devices.²⁶ A coronene diimide small molecule (C, Fig. 1) has also been used.²⁷ Unlike their perylene and coronene counterparts, naphthalene diimides (NDI) can exhibit minimal absorption within the visible spectrum.

Since polymeric HTMs such as PEDOT:PSS, P3HT, PTAA, and polyTPD^{28–31} have been reported to give impressive performances in PSCs (PCE > 19%), it may be beneficial to examine

electron-transporting polymers that are transparent in the visible, with appropriate electron affinities, electron mobility, solvent resistance, and solution processability. Moreover, such polymers may facilitate a “polymer sandwich” architecture in which both transport materials are polymeric, and which may be particularly useful for flexible devices.

In this work, we report a new NDI side-chain polymer (Fig. 1), which is transparent, solution processable, and thermally stable, and which can be used to fabricate n-i-p devices with efficiencies up to 14%. The utility of the polymer is illustrated by comparison to other NDI small molecules as well as SnO₂ as a reference.



Scheme 1 Synthesis of side chain NDI polymer (NDI-1)

Results and discussion

Synthesis

A naphthalene diimide polymer may help reduce aggregation and crystallization and, thus aid the formation of uniform continuous (*i.e.* pinhole free) thin films with reduced shunt paths between the transparent conducting oxide (TCO) and the perovskite. Although conjugated NDI polymers are well established and have been used in p-i-n cells, many absorb strongly throughout the visible spectrum, competing with perovskite layer light absorption.³² Accordingly, we chose to synthesize a side-chain polymer, which was expected to be colorless (NDI-1, Fig. 1). The polymer synthesis (Scheme 1) involves four steps, the first of which is an asymmetric imidization of the commercially available naphthalene tetracarboxy-1,8:4,5-dianhydride. The second step involves the imidization of *N*-hexyl-naphthalene-1,8-dicarboximide-4,5-di(carboxylic acid) with 5-amino-1-pentanol to afford the asymmetric hydroxyl-functionalized naphthalene diimide. Thirdly, the monomer is produced by esterification using *exo*-5-norbornenecarboxylic acid. Lastly, the regiorandom polymer NDI-1 ($M_n = 13.3$ kDa, $\bar{D} = 1.3$) is produced *via* ring-opening metathesis polymerization using the Grubbs first-generation initiator, Ru(=CHPh)(PCy₃)₂Cl₂ (Cy = cyclohexyl) (Fig. S1 and S2, ESI†). Several small-molecule NDI molecules (2–4) were synthesized for comparison according to published procedures,³³ and were chosen to exhibit a range of reduction potentials.

Characterization of the ETMs

Fig. 2A shows absorption spectra for NDI-1 in solution and in films, while Table 1 gives absorption maxima for NDIs 1–4. Both solution and films of the NDI derivatives exhibit absorptions onsets at wavelengths shorter than 400 nm. Although scattering is evident for films in the visible range, the transmittance for a 30 nm-thick film still exceeds 80% through the visible region (400–800 nm) (Fig. S7, ESI†). The electron affinities of ETMs are critical for efficient charge extraction from the perovskite; the reduction potentials were estimated from cyclic voltammetry (Fig. S3, ESI†, Table 1) and fall into a similar range to other molecular materials successfully used as ETMs in PSCs (such as C₆₀ (−0.98 V), PCBM (−1.1 V), and N2200 (−1.1 V)).^{32,34}

Table 1 Optical and thermal properties of NDI material used

Compound	$\lambda_{\max}^a$ (nm)	E_g^b (eV)	T_m^c (°C)	T_d^c (°C)	$E_{1/2 \text{ red}}^d$ (V)
NDI-1	381	3.1	—	384	−1.11
NDI-2	376	3.1	180	328	−1.14 ³³
NDI-3	376	3.2	—	274	−0.88 ³³
NDI-4	379	3.2	195	305	−0.96 ³³

^a UV-vis is measured in chlorobenzene. ^b Optical bandgap is determined from absorption edge. ^c DSC/TGA is done under nitrogen gas at 10 °C min^{−1} scan rate, and T_d is determined at 95% mass. ^d $E_{1/2 \text{ red}}$ vs. FeCp₂^{+/0} (CH₂Cl₂/0.1 M NBu₄PF₆); data for NDI-1 were acquired using the corresponding monomer.

The four NDI materials are highly soluble in common organic solvents (chloroform, dichloromethane, toluene), suggesting that they could be readily processed onto perovskite films for p-i-n devices without significant damage to the active layer. They exhibit varying resistance to *N,N*-dimethylformamide and dimethyl sulfoxide (DMF and DMSO respectively, Fig. S9 and S10, ESI†). The DMF resistance of NDI-1 is particularly promising for the fabrication of n-i-p device architectures; the small molecules are more easily removed by this solvent, although they are expected to be less soluble in concentrated solutions of perovskite precursors and when protected by the alumina interlayer. Thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) show that the naphthalene diimides examined here exhibit decomposition temperatures above 250 °C and no thermal transitions below 150 °C, which is compatible with temperatures needed for subsequent deposition and annealing of many hybrid perovskites used in PSCs (Fig. 2B and C).

Device architecture and performance

The materials were incorporated into n-i-p devices (Fig. 3). The NDI layers were spin-coated from chlorobenzene on fluorinated-tin-oxide (FTO) coated glass (optimally performing layers having thicknesses of *ca.* 10 nm). A thin mesoporous layer of alumina nanoparticles was found to be very helpful in achieving subsequent good coating of the ETM by the perovskite. Although primarily used in the present case to enhance the wettability of the perovskite solution on the NDI films, the alumina nanoparticles may play other roles, as discussed in the literature.^{35–37} For example, at least when used between oxide transport materials and perovskites, a mesoporous alumina

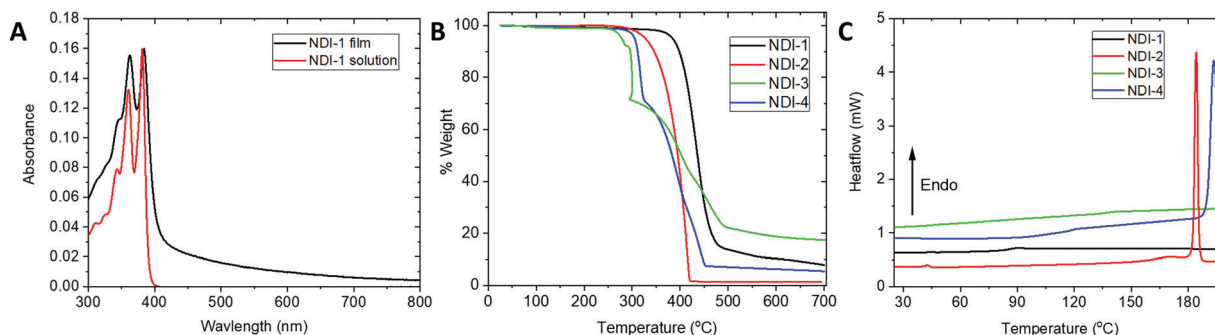


Fig. 2 Absorption spectra of NDI-1 in solution and thin film (A), TGA (B) and DSC (C) plots of NDI molecules used in this work. TGA and DSC were performed under nitrogen gas at 10 °C min^{−1} scan rate.

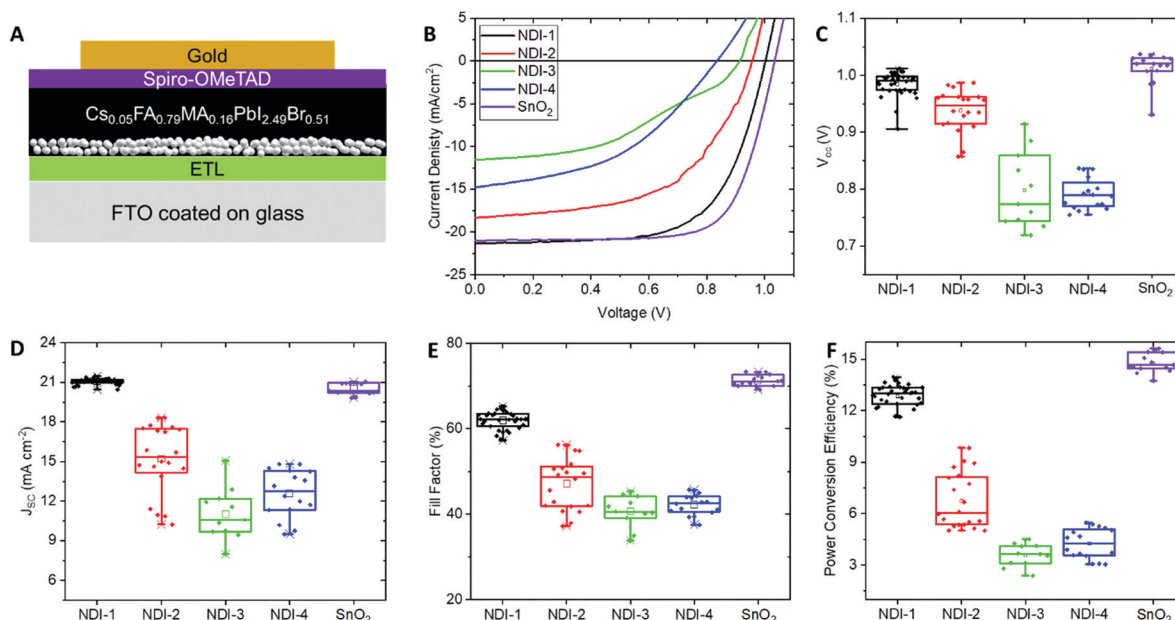


Fig. 3 Schematic representation of device architecture, white dots in active layer represent alumina nanoparticles (A). Reverse Scan J - V plots of optimized devices (B). Box plots of open-circuit voltage (C), short-circuit current (D), fill factor (E), and power conversion efficiency (F) of optimized devices.

layer leads to improved contact between the transport and active layers,³⁶ and in some cases the use of alumina leads to improved stability for the perovskite.³⁷ They may also help protect the ETM from dissolution in the perovskite casting solvent (primarily DMF). Fig. 3A schematically shows the structure of the resulting solar cell incorporating the insulating alumina scaffold.³⁸ Our control devices employed a SnO_2 ETL layer instead of the NDI-based material. A mixed-cation perovskite $\text{Cs}_{0.05}\text{FA}_{0.79}\text{MA}_{0.16}\text{PbI}_{2.49}\text{Br}_{0.51}$ (FA = formamidinium; MA = methylammonium) was then deposited *via* the solvent-quenching approach using chlorobenzene as an anti-solvent.^{39–42} Spiro-OMeTAD, with lithium bis(trifluoromethanesulfonyl)imide (LiTFSI) and 4-*tert*-butylpyridine (*t*BP) as additives, was then deposited on top as the hole transport layer, followed by evaporating 80 nm layer of gold as a counter electrode.

Fig. 3B–F show that optimized devices using the polymer were superior to all optimized devices incorporating NDI small molecules, the power conversion efficiencies following

the trend $\text{NDI-1} \gg \text{NDI-2} > \text{NDI-4} > \text{NDI-3}$, with the small molecules all exhibiting lower open-circuit voltage (V_{OC}), short-circuit current density (J_{SC}), and fill factor (FF) than the polymer. The photovoltaic parameters of the champion and average devices incorporating the small molecules (NDI-2, NDI-3, NDI-4), the polymer (NDI-1), and SnO_2 are summarized in Table 2.

Devices with NDI-1 exhibited comparable V_{OC} and J_{SC} to those using tin oxide; however, they fell short in terms of FF and consequently the power conversion efficiency (PCE, Fig. 3). This might be due to higher series resistance of the devices with a polymer ETL, as the conductivity of SnO_2 is likely higher than that of the undoped polymer. Nevertheless, the devices still exhibit a maximum power conversion efficiency of 14%. Both devices with organic and SnO_2 ETMs suffered from non-negligible hysteresis, likely due to the Spiro-OMeTAD/additive HTM. However, a maximum power point tracking test showed a maximum efficiency of 13.8% after 90 s of illumination,

Table 2 Photovoltaic parameters of champion and average devices of the various materials employed as electron transport layer in the n-i-p devices studied

Material		V_{OC} (V)	J_{SC} (mA cm^{-2})	Fill factor (%)	PCE (%)
SnO_2	Champion	1.03	21.0	74	15.6
	Average	0.99 (± 0.02)	20.9 (± 0.4)	73 (± 1.3)	15.0 (± 0.5)
NDI-1	Champion	1.02	21.5	65	14.0
	Average	0.98 (± 0.02)	21.1 (± 0.2)	62 (± 2.1)	12.8 (± 0.6)
NDI-2	Champion	0.98	18.3	56	9.8
	Average	0.94 (± 0.04)	15.2 (± 2.6)	47 (± 6)	6.7 (± 1.6)
NDI-3	Champion	0.91	11.6	43	4.5
	Average	0.79 (± 0.06)	11.0 (± 2.0)	41 (± 3.7)	3.6 (± 0.7)
NDI-4	Champion	0.84	14.8	46	5.5
	Average	0.79 (± 0.02)	12.6 (± 1.8)	42 (± 2.4)	4.3 (± 0.9)

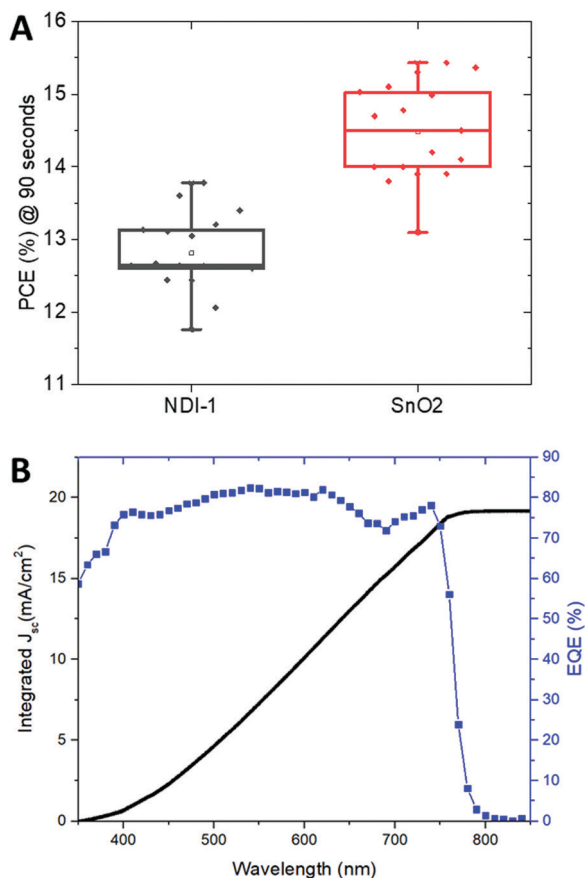


Fig. 4 (A) Box plots of the power density of NDI-1 and SnO₂ devices and (B) external quantum efficiency (EQE) and integrated short-circuit current plots for the NDI-1 devices.

representing only a 0.2% decrease from the reverse scan efficiency (Fig. 4A). Forward scan curves are presented in Fig. S5 (ESI[†]), and an example of maximum power point tracking (MPPT) evolution over time is given in Fig. S8 (ESI[†]). Fig. 4B shows external quantum efficiency and integrated current data in order to further characterize the efficiency of the perovskite absorber in the presence of NDI-1. Photoluminescence quenching of the perovskite was also measured and shown in the ESI[†].

Long-term stability

Both NDI-1 and SnO₂ based devices were stable for multiple months when stored in a dry box (relative humidity < 5%) in the dark, showing no changes in performance and indicating a high shelf life for the devices. To compare the stability differences between devices employing NDI-1 to their tin oxide counterparts, we encapsulated (see Experimental section) full devices and aged them in an environmental chamber at open circuit voltage at 60 °C, in air. The study was conducted over 290 hours where the tin oxide device efficiency degraded to <10% of the initial value, whereas that of the devices incorporating the NDI-1 polymer retained *ca.* 35% of the original performance, the degradation being dominated by a loss of FF. Fig. 5 shows the evolution of PCE over aging time for champion

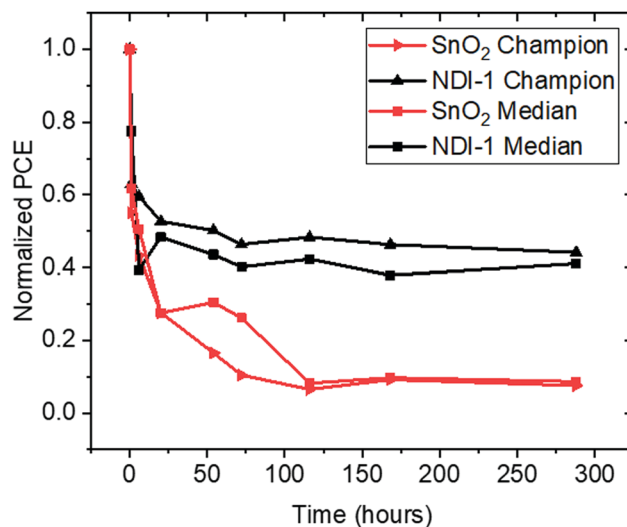


Fig. 5 Long term stability of devices employing SnO₂ and NDI-1.

and median devices. Although LiTFSI/tBP-doped Spiro-OMeTAD is the common HTM in both devices and has known instabilities that may ultimately limit the stability of the best devices here,^{43,44} those devices incorporating NDI-1 are more stable over the period of the study.

Conclusions

We demonstrate in this study a side chain NDI polymer that is superior in performance to several NDI small molecules in n-i-p PSCs. We achieved a champion device with 14% PCE using a undoped films of NDI-1. These devices exhibited greater stability than SnO₂-based control devices when aged under 1 sun conditions for almost 300 hours and 60 °C. This provides a foundation for designing other possible naphthalene diimides or other moieties to be used as dopant-free solution-processable electron-transport materials to replace metal oxide ETMs. This low temperature processing route open new avenues for polymer-based flexible solar cells and the optical transparency for incorporation in tandem devices.

Materials and methods

Commercially available precursors as well as the solvents were purchased from Sigma Aldrich and used without modifications. Detailed synthetic procedures and characterization data are presented in the ESI[†].

FTO coated glass slides (1 cm²) were cleaned by sonication in 2% Hellmanox in water, followed by deionized water, and ethanol for 15 min each. The slides were then dried and plasma etched for 10 min and used immediately. Organic ETM solutions were prepared by making 1.5 mg mL⁻¹ solutions in chlorobenzene,⁴⁵ which were spun at 2000 rpm for 30 s in air yielding layers of 10 nm thickness. When tin oxide was used as an ETM the layer was prepared by spin coating 10 mg mL⁻¹ solution in isopropyl alcohol of SnCl₂·5H₂O at 2000 rpm for 20 s

followed by annealing at 180 °C for 1 h. Plasma treatment for 1 min was done before SnO₂ nanoparticles were deposited by diluting commercially available precursor in 1:10 DI water then spin coating it at 2000 rpm for 20 s followed by annealing at 180 °C for 1 h. Alumina nanoparticles (<50 nm) were dispersed in isopropanol in 1:150 v/v ratio. 100 µL of this dispersion was then dynamically spun onto the ETL at 3000 rpm for 30 s and then dried at 60 °C for 1 min. The perovskite precursor was prepared by mixing 507.1 mg of PbI₂, 73.4 mg PbBr₂, 22.4 mg MABr, 172 mg FAI, with 41 µL of 1.5 M CsI solution (in DMSO) and 850 µL of a pre-mixed 4:1 DMF:DMSO. The perovskite was then spun using a two-step program by covering the film with 50 µL of the perovskite solution, and spinning that at 1000 rpm for 10 seconds, followed by 6000 rpm for 20 s. When 5 s were left to the end of the program, 260 µL of chlorobenzene was dropped to solvent quench the film. The films were rapidly moved to a hot hotplate to be annealed at 100 °C in the dark for 1 h where they turned black immediately. The Spiro-OMeTAD layer was prepared by preparing a 72 mg mL⁻¹ solution of Spiro-OMeTAD in chlorobenzene followed by the addition of a 17.5 µL LiTFSI solution (52 mg mL⁻¹ concentration in acetonitrile) and 28.8 µL of tBP. The mixed solution was spun dynamically at 2000 rpm for 30 s and devices were exposed to dry air in the dark for doping to take place overnight. Gold electrodes were evaporated using a metal sputter, using a shadow mask to fix the area at 0.16 cm², and evaporated at 0.1 nm s⁻¹ for the first 10 nm and 1 nm s⁻¹ afterwards.

Devices were measured under 1 sun illumination using a custom-built 16-device set-up in which the devices were light soaked for at least 90 s before measurements to ensure stabilization where the aperture size was fixed at 0.16 cm². Encapsulation was achieved by dispensing UV-curable epoxy resin from Nagase (XNR5516Z-B1) onto the edges of a cover glass with a recess in the middle. The cover glass was then gently pressed on to the solar cell devices and were illuminated from the cover glass side under UV light.

UV-vis-NIR measurements were done on a Cary 5000 spectrophotometer. PL spectra were done on a Horiba instrument. EQE data were recorded through a Keithley 2400 Source Meter under 300 W xenon lamp irradiation with an Oriel Cornerstone 260¹/₄ m monochromator. The monochromatic photon flux was quantified through a calibrated silicon cell (Pecell Technologies).⁴⁶ DSC and TGA experiments were performed on Mettler-Toledo TA instruments under nitrogen gas atmosphere and at a scan rate of 10 °C min⁻¹. Gel Permeation Chromatography (GPC) for polymer samples were performed using a Tosoh EcoSEC HLC 8320GPC system with TSKgel SuperHZ-L columns eluting CHCl₃ containing 0.25% triethylamine at a flow rate of 0.45 mL min⁻¹. All number-average molecular weights and dispersities were calculated from refractive index chromatograms using PStQuick Mp-M polystyrene standards. Electrochemical measurements were carried out under nitrogen atmosphere with 0.1 M tetra-*n*-butylammonium hexafluorophosphate in dry dichloromethane using a CH Instruments CHI620D Electrochemical Workstation CHI620D, and a conventional three-electrode cell with a glassy carbon working

electrode, platinum wire counter electrode, and an Ag wire coated with AgCl as the pseudo-reference electrode. Potentials were referenced to ferrocenium/ferrocene by using ferrocene as an internal standard. Cyclic voltammograms were recorded at a scan rate of 50 mV s⁻¹.

Conflicts of interest

There are no conflicts to declare.

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- 46 Small inconsistencies in *J*_{SC} between the integrated EQE data and the *J–V* data may result from imperfect EQE calibration.