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13.9% Efficiency Ternary Nonfullerene Organic Solar Cells Featuring Low-structural Order

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ABSTRACT: The insufficient phase separation between polymer donors and non-fullerene acceptors (NFAs) featuring with low-structural orders disrupts efficient charge transport and increases charge recombination, consequently limits the maximum achievable power conversion efficiency (PCE) of organic solar cells (OSCs). Herein, an NFA IT-M has been added as the third component into the PBDB-T:m-INPOIC OSCs, and is shown to effectively tune the phase separation between donor and acceptor molecules, although all components in the ternary system exhibit low degrees of structural orders. The incorporation of 10 wt% IT-M into a PBDB-T:m-INPOIC binary host blend appreciably increases the length scale of phase separation, creating continuous pathways which increase and balance charge transport. This leads to an enhanced photovoltaic performance from 12.8% in the binary cell to 13.9% for the ternary cell with simultaneously improved open-circuit voltage, short-circuit current and fill factor. This work highlights the beneficial role of ternary components in controlling the morphology of the active layer for high performance OSCs.

Over the past few years, the development of non-fullerene acceptors (NFAs) has driven the impressive progress of organic solar cells (OSCs).^{1–4} The tunable energy levels and absorption spectra of NFAs can allow for control of complementary absorption and low voltage loss which are critical for high power conversion efficiency (PCE)^{5–8}, with over 16% PCE achieved for single-junction binary non-fullerene OSCs.^{9–11} Whilst the emergence of new electron donor and acceptor materials is the primary motivation to further advance OSCs, compositional and morphological optimization within the photoactive layer is vital to realize closer to the theoretical maximum PCE.^{10–12} The desired morphology of the photoactive layer should resemble nanoscale phase separated domains for efficient exciton diffusion and dissociation, of the order of the limited exciton diffusion lengths which are usually not more than 10 nm.^{15–17} Furthermore bicontinuous networks are favorable for charge carrier transport, collection and suppression of bimolecular recombination.^{12,18,19}

Most of the polymer donors, e.g. PTB7-Th and PBDB-T, exhibit low-structural order in the form of π - π stacking, due to the confinement of bulky conjugated repeating units in a macromolecular structure.^{20,21} The versatile chemical structures of NFAs endow this class of fascinating electron acceptors with very different molecular packing behaviors. For instance, COi8DFIC and INPIC-4F show a high tendency to crystallize into lamellae^{22–25}, whilst ITIC and IEICO series materials exhibit π - π stacking only^{26,27}. NFAs with similar chemical structures to conjugated polymers leads to good miscibility between donors and acceptors, especially with those of low structural orders; however this commonly results in insufficiently separated phases after solution casting, which impacts charge generation, transport and recombination.^{28–30} Whereas the fine phase separation and intimate contact between donors and acceptors benefits exciton dissociation, these morphologies are not ideal for charge transport.

Although good efficiencies have been achieved in those photovoltaic systems featuring low structural orders, e.g. PBDB-T:ITIC and PBDB-T:IT-M, further enhancement of performance has proved rather difficult. For example, thermal annealing of PBDB-T:IT-M blend films barely increases the structural orders of PBDB-T and IT-M, explained by their intrinsic low ability to self-organize. Consequently less than 10% PCE improvement has been obtained for annealed

devices compared with as-cast devices.³¹ Solvent vapor annealing (SVA) is another effective approach that has been demonstrated to reorganize molecular packing within blends and improve the efficiency of many fullerene-based OSCs.^{32,33} However, in non-fullerene OSCs featuring low-structural orders e.g. PTB7-Th:ITIC, only minor enhancement of molecular packing has been observed using a range of solvent or solvent mixture vapors to anneal devices, as such a PCE increase of only 10% can be achieved.³⁴

Ternary photovoltaic solar cells prepared by incorporating a third component into conventional binary solar cells have emerged as a promising strategy for realizing further improvements in efficiency.^{35–37} This method is favourable as it removes the time-consuming and expensive process of synthesizing new conjugated polymers. Whilst the primary advantage of the ternary strategy is to achieve complementary light absorption^{37–39}, it can also effectively regulate the morphology.^{40,41} Although literature reports have demonstrated reduced trap density and recombination in ternary systems compared to binary systems,^{42–45} less attention has been paid to tuning the phase separation and efficiency of non-fullerene OSCs featuring low structural orders.

In this work, we employ the non-fullerene acceptor IT-M as the third component to tune the domain size in PBDB-T:m-INPOIC blends, which have until now been inhibited by insufficient phase separation, as all components exhibit low-structural orders. The presence of an intermediate amount of IT-M enhances photon absorption in the ternary device. The appreciatively enlarged length scale of phase separation induced by the presence of 10 wt% IT-M facilitates increased and balanced charge mobilities with minimized trap-assisted recombination. As a result, the ternary OSC achieves a maximum PCE of 13.9% compared with 12.8% for the PBDB-T:m-INPOIC binary OSC, with the simultaneously increased device metrics of V_{oc} of 0.86 V, J_{sc} of 22.2 mA/cm² and FF of 71.3%. This work highlights the beneficial role of ternary components in mediating morphology of the active layer to improve device performance.

The chemical structures, energy levels of materials and schematic of the device structure used in this work are shown in **Figure 1**a-b.^{46,47} **Figure 1**c clearly shows the complementary absorption of the different components in the ternary system. To examine the possible Förster resonance energy transfer (FRET) between m-INPOIC and IT-M, we have measured the photoluminescence (PL) spectra of the individual components and their mixtures with different weight ratios. As shown in **Figure 1**d, IT-M and m-INPOIC exhibit distinct emission peaks at 765 and 873 nm respectively. The broad overlap between the emission spectrum of IT-M and the absorption spectrum of m-INPOIC (**Figure 1**c) should enable efficient energy transfer from IT-M to m-INPOIC. In the m-INPOIC:IT-M blend, the emission signal of IT-M is markedly quenched with a single emissive peak observed at 873 nm that is associated with m-INPOIC, suggesting efficient energy transfer from IT-M to m-INPOIC which is favorable for photovoltaic performance.⁴⁸ From this efficient energy transfer process we can also imply there is good miscibility between these two acceptors, with close mixing in the blend film.^{35,41,49}

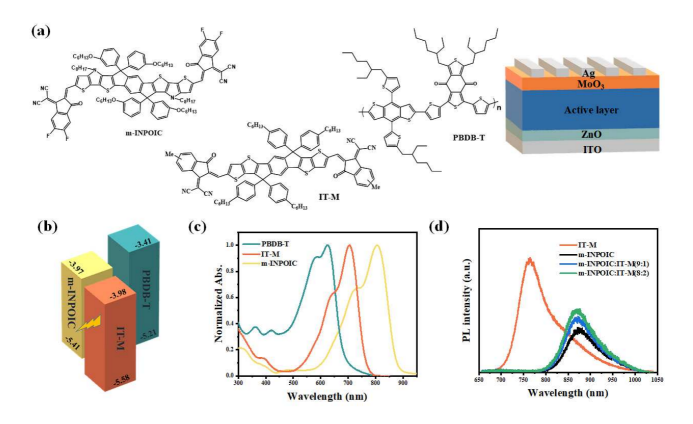


Figure 1. (a) Chemical structures of polymer donor and non-fullerene acceptors, and the device architecture used in this work. (b) Schematic energy diagrams of PBDB-T, IT-M and m-INPOIC, here the lightning bolt indicates the related energy transfer process. (c) Optical absorption spectra of neat films of PBDB-T, m-INPOIC and IT-M. (d) PL spectra of m-INPOIC, IT-M and their mixtures at the weight ratios of 9:1 and 8:2, excited with a 532 nm laser.

Transmission electron microscopy (TEM) was performed to understand the phase separated domain size within the binary and ternary blends. As shown in **Figure 2**a-c, the dark and bright

regions represent the acceptor and donor domains respectively, because of their different electron densities.⁵⁰ The TEM image of the PBDB-T:m-INPOIC binary film (see **Figure 2**a) shows fine mixing and homogeneous distribution of donors and acceptors, whilst the continuous domain networks are less pronounced. Upon the incorporation of IT-M (see **Figure 2**b and c), the distinction between dark and bright regions becomes more pronounced. This suggests increased phase separation leading to larger and purer domains within the ternary blend, which will help the formation of bicontinuous paths for efficient charge transport and reduced charge recombination.^{18,19} These larger domains will have higher charge mobility because of the much faster carrier transport within a single-phase domain than between domains.⁵¹

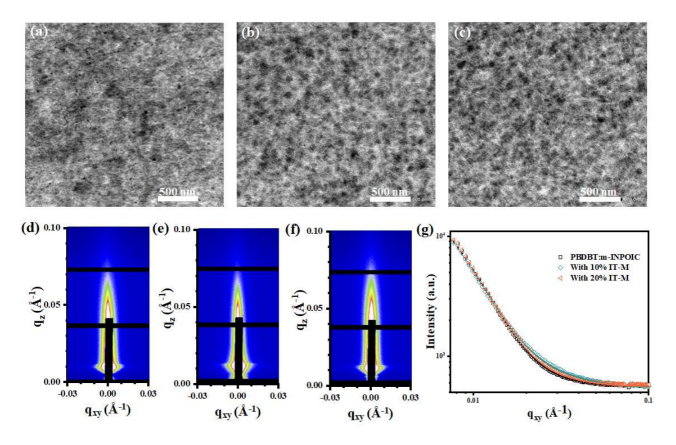


Figure 2. TEM images of (a) PBDB-T:m-INPOIC film, and its ternary blends with (b) 10%, (c) 20% IT-M. 2D GISAXS patterns of (d) PBDB-T:m-INPOIC, and its ternary blends with (e) 10% IT-M, (f) 20% IT-M. (g) 1D GISAXS profiles along the q_{xy} axis for PBDB-T:m-INPOIC blend films with different IT-M contents.

In order to quantify the phase separated domain sizes of donors and acceptors, grazing-incidence small-angle X-ray scattering (GISAXS) was employed. The corresponding 2D GISAXS patterns of binary and ternary blends and their 1D GISAXS profiles along q_{xy} axis are

shown in **Figure 2**d-g. A universal model (detailed in the supporting information) was used to fit the 1D profiles and the relevant fitting parameters are shown in **Table 1**. Here the correlation length (ξ) refers to the domain size of the PBDB-T-rich phase, η and D represent the correlation length and fractal dimension of acceptors and 2Rg (the product of η and D) is regarded as the domain size of acceptor aggregation. According to **Table 1**, the PBDB-T and m-INPOIC domain sizes in the binary blend are 13.3 and 32.1 nm respectively. Adding 10 wt% IT-M into binary blend leads to the growth of both PBDB-T and m-INPOIC domains, which are enlarged to 14.8 and 35.0 nm respectively. Further increased domain size and phase separation were observed when 20 wt% IT-M was added. Therefore, we conclude the addition of IT-M increases the domain size of both donors and acceptors. The ternary blend with 10 wt% IT-M shows the smallest fractal dimension of 2.6, suggesting loosely packed aggregates of m-INPOIC, which could extend to larger regions facilitating both exciton dissociation and charge transport.^{52,53} This quantitative observation is consistent with the more pronounced contrast and domain size within the TEM images of the ternary blends.

	ξ (nm)	η (nm)	D	2R _g (nm)
PBDB-T:m-INPOIC	13.3	13.9	2.8	32.1
With 10% IT-M	14.8	16.2	2.6	35.0
With 20% IT-M	16.0	16.7	2.9	39.7

Table 1. Fitting parameters of 1D GISAXS profiles for PBDB-T:m-INPOIC and its ternary films.

Due to the competition between phase separation and molecular ordering,²⁹ we further investigate the effect of the addition of the third component IT-M on the molecular packing and orientation via grazing incidence wide-angle X-ray scattering (GIWAXS) measurements. The 2D diffraction patterns and 1D profiles are shown in **Figure 3**. It is immediately apparent in the 2D GIWAXS patterns that the face-on π - π stacking of the blends is gradually enhanced upon the incorporation of IT-M. As shown in **Figure 3**d-e, the binary PBDB-T:m-INPOIC film shows a weak and broad π - π peak in the out-of-plane (OOP) direction and a sharp lamellar stacking in the in-plane (IP) direction, which is the character of preferential face-on orientation.⁵⁴ Deconvolution of this broad π - π peak gives two convolved peaks located at $q_z \approx 1.73$ Å⁻¹ and $q_z \approx 1.80$ Å⁻¹ (see in Figure S1), which can be assigned to PBDB-T and m-INPOIC components respectively.⁴⁷ With the addition of IT-M, no obvious diffraction peaks from IT-M were observed, which is consistent with the low structural order of IT-M.³¹ Meanwhile, stronger π - π stacking diffraction in the OOP direction and PBDB-T lamellar stacking in the IP direction were observed in Figure 3b and c. Deconvolution of these broad π - π peaks from 1.60 to 1.90 Å⁻¹ using multi- Gaussian peak fitting is shown in **Figure S1**. With the addition of IT-M, the π - π peak positions of PBDB-T and m-INPOIC slightly shift to higher q values, suggesting a tighter stacking distance. Comparing the intensity changes of the deconvoluted peaks, the main contribution to the enhanced π - π stacking comes from PBDB-T in the ternary blend, with negligible changes of m-INPOIC. This suggests enhanced ordering of PBDB-T, which is most pronounced with the addition of 20 wt% IT-M. We regard this enhanced ordering as resulting from the increased domain size within the ternary blends, allowing PBDB-T molecules to pack efficiently without the interruption of acceptor molecules. The fact that the π - π stacking intensity of m-INPOIC barely changes suggests the good miscibility between m-INPOIC and IT-M, which might form an alloy as observed in many ternary system.^{55,56} The increased packing of PBDB-T in the OOP direction within the slightly larger domains will increase the hole mobility (results presented in a later section) to facilitate carrier transport along the vertical direction towards the anode.57

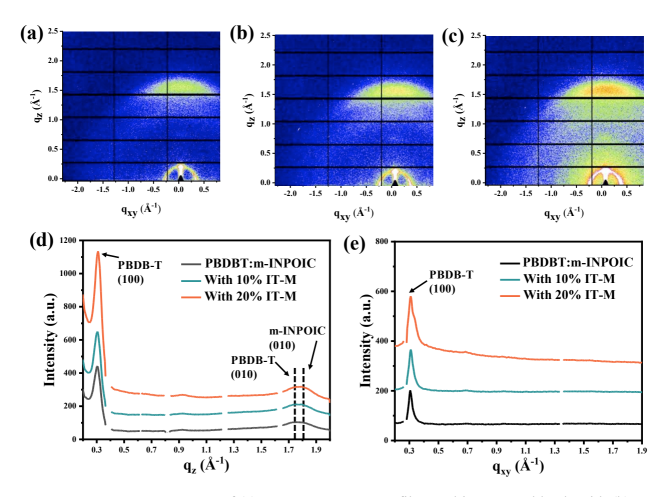


Figure 3. 2D GIWAXS patterns of (a) PBDB-T:m-INPOIC film, and its ternary blends with (b) 10 wt%, (c) 20 wt% IT-M. Corresponding (d) out-of-plane and (e) in-plane 1D profiles of GIWAXS patterns along the q_z - and q_{xy} -axis.

It is well accepted that the efficiency of OSCs is greatly related to the photon absorption of the photoactive layer.²⁷ Thus we further investigated the absorption spectra of the binary and ternary films where the overall donor/acceptor weight ratio was kept at 1:1. Referring to the absorption spectra in **Figure 4**a, the prominent peaks at 635 and 800 nm correspond to the absorption of PBDB-T and m-INPOIC respectively. With the addition of IT-M, the absorption is enhanced in the wavelength range from 500 to 700 nm. We attribute this to two reasons, firstly the stronger absorption of IT-M in this range, and secondly the enhanced molecular packing of PBDB-T (discussed above in the GIWAXS section) which will also enhance photon absorption.²⁹ With the addition of 20 wt% IT-M, however, the absorption at longer wavelengths reduces notably, which results from the reduced fraction of m-INPOIC in the blend.

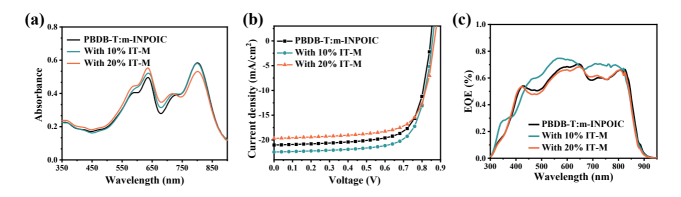


Figure 4. (a) Absorbance of PBDB-T:m-INPOIC films with different IT-M contents, (b) Champion J–V curves, and (c) EQE of devices fabricated with different IT-M contents.

Table 2. The photovoltaic parameters of PBDB-T:m-INPOIC OSCs with varying IT-M content obtained under simulated AM 1.5 G illumination at 100 mW/cm². The statistical data were obtained from over 20 individual devices.

	FF [%]	J _{sc} [mA cm ⁻²]	Cal. J _{sc} [mA cm ⁻²]	V _{oc} [V]	PCE _{avg} (PCE _{max}) [%]
PBDB-T:m-INPOIC	70.8±0.45	20.7±0.30	19.6	0.85±0.002	12.5±0.26 (12.8)
With 10% IT-M	71.3±0.66	22.2±0.24	21.0	0.86±0.003	13.7±0.16 (13.9)
With 20% IT-M	72.4±0.85	19.1±0.43	18.2	0.87±0.004	11.9±0.16 (12.2)

A series of inverted binary and ternary devices were fabricated to evaluate the effects of these morphological characteristics on device performance. The overall donor to acceptor ratio was kept constant at 1:1 in this work. The J-V curves of best-performing devices are shown in **Figure 4**b and the device metrics for each composition are summarized in **Table 2**. The control device refers to the PBDB-T:m-INPOIC binary cell, which exhibits a PCE_{max} of 12.8%, with a FF of 70.8%, a J_{sc} of 20.7 mAcm⁻² and a V_{oc} of 0.85 V. Incorporating 10 wt% IT-M into the PBDB-T:m-INPOIC blend leads to simultaneous improvements of the J_{sc} to 22.2 mAcm⁻², FF to 71.3% and V_{oc} to 0.86V, resulting in a PCE_{max} of 13.9%. The larger domain sizes in the ternary OSCs will facilitate charge carrier transport, on the other hand, the reduced D/A interface area and the longer distance that excitons have to migrate are harmful to efficient exciton dissociation.¹⁶ As a result, when 20 wt% IT-M was incorporated to generate domains with excessive sizes, the FF of the device further

increases to 72.4% and the V_{oc} increases to 0.87V, but the J_{sc} decreases abruptly to 19.1 mAcm⁻², leading to a low PCE_{max} of 12.2% only. In general, the V_{oc} of the device is associated with the offsets between the highest occupied molecule orbital (HOMO) of the donor and the lowest unoccupied molecule orbital (LUMO) of the acceptor.^{58,59} Here, the gradually increasing V_{oc} is in line with the IT-M content, suggesting that IT-M acts as an energy-level modulator to monotonically elevate the LUMO levels of the mixed acceptors due to its higher-lying LUMO.

The external quantum efficiency (EQE) spectra of the best-performing devices are shown in **Figure 4**c. When the binary blend had 10 wt% IT-M added, the EQE spectra of the device shows marked increase in the broad region between 450 and 800 nm, which is consistent with the enhanced absorption in this region. Significantly increased charge dissociation and collection as well as suppressed recombination account for the improved EQE values (discussed below). Further increasing the content of IT-M to 20% leads to slightly reduced photon to current efficiency from 450 to 650 nm. This indicates that excessive amount of IT-M begins to hamper current generation due to inefficient generation of free charges from excitons, because of sub-optimal phase separation.

To further investigate the effect of the varying degree of phase separation on exciton dissociation and collection, the photocurrent density (J_{ph}) as a function of the effective voltage (V_{eff}) were measured, as shown in **Figure 5**a.⁶⁰ Here $J_{ph} = J_L - J_D$, in which J_L and J_D are the photocurrent densities under illumination and dark respectively. $V_{eff} = V_o - V_a$, where V_o is the voltage when $J_L = J_D$ and V_a is the applied voltage. Assuming that all the generated excitons have dissociated and been collected by electrodes at large V_{eff} , a saturated photocurrent density (J_{sat}) will be reached. The maximum J_{sat} of the ternary device with 10 wt% IT-M is partly due to the efficient energy transfer from IT-M to m-INPOIC. The exciton dissociation possibility (P_{diss}) and charge collection efficiency (P_{coll}) are defined as J_{ph}/J_{sat} values under short-circuit condition and maximal power output condition respectively. As displayed in **Table 3**, the P_{diss} values of the ternary OSC with 10% IT-M content is higher than the binary OSC. But at the higher amount of 20 wt% IT-M, excessive phase separation and reduced D/A interfaces results in the inhibition of exciton dissociation. The dramatically increased P_{coll} for 10% IT-M is attributed to the more ordered molecular packing of

PBDB-T and the formation of bicontinuous networks, enabling efficient charge carrier transport with suppressed carrier recombination.

Additionally, we took J-V measurements of OSCs under different light intensity (P_{light}) to gain more insight into the charge recombination behavior of devices. The profiles of J_{sc} and V_{oc} as a function of light intensity (P_{light}) are described in **Figure 5**b and c. By analyzing the V_{oc} versus ln(P_{light}) plot, the dominant charge recombination mechanism can be distinguished. At the bimolecular recombination condition, the slope of V_{oc} versus ln(P_{light}) is close to KT/q, where K is the Boltzmann constant, T is the absolute temperature and q is the elementary charge. The slope will be larger than KT/q for trap-assisted recombination. The slope of V_{oc} versus ln(P_{light}) for the binary device is 1.32 KT/q, which decreases to 1.21 KT/q and 1.28 KT/q for ternary devices with 10% and 20% IT-M respectively, suggesting the most effectively suppressed trap-assistance recombination with the addition of 10% IT-M. Meanwhile, the slopes of log(J_{sc}) versus log(P_{light}) plots were analyzed to evaluate the degree of bimolecular recombination. As shown in **Figure 5**c, the slopes are 0.96, 0.98 and 0.96 for the binary and ternary devices with 10% IT-M and 20% IT-M, respectively. A slope close to unity signifies that bimolecular recombination dominates in these devices.

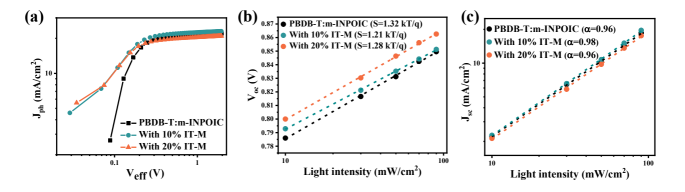


Figure 5. (a) Photocurrent density (J_{ph}) versus effective voltage (V_{eff}) curves. (b) Dependence of V_{oc} on light intensity and (c) dependence of J_{sc} on light intensity for the devices with different amounts of IT-M.

We finally investigated the effect of IT-M on carrier mobilities (μ) using the space charge limited current (SCLC) method. The hole- and electron- only devices were fabricated with the structures of ITO/PEDOT:PSS/active layer/MoO₃/Ag and ITO/ZnO/active layer/Ca/Ag,

respectively. The J^{1/2}-V curves are shown in **Figure S2**, and the mobilities are extracted and summarized in **Table 3**. The μ_h and μ_e of the PBDB-T:m-INPOIC binary device are 2.6×10⁻⁴ cm²V⁻¹s⁻¹ and 2.1×10⁻⁴ cm²V⁻¹s⁻¹ respectively. After the incorporation of 10% IT-M, the ternary device displays higher μ_h and μ_e of 2.8×10⁻⁴ cm²V⁻¹s⁻¹ and the hole and electron transport becomes balanced as determined by the μ_h/μ_e ratio of unity. The enhanced molecular order of PBDB-T and improved bicontinuous pathways account for these increased mobilities. As more IT-M was added, μ_h continuously increases to 4.2×10⁻⁴ cm²V⁻¹s⁻¹ and μ_e increases to 3.0×10⁻⁴ cm²V⁻¹s⁻¹, thus the μ_h/μ_e becomes unbalanced, which will increase charge accumulation and recombination, worsening device performance.

Table 3. J_{sat}, P_{diss}, P_{coll}, hole and electron mobilities of PBDB-T:m-INPOIC and its ternary OSCs with different contents of IT-M.

IT-M content	J _{sat} [mA cm ⁻²]	Pdiss	P _{coll}	Hole mobility (µh) [cm ² V ⁻¹ s ⁻¹]	Electron mobility (µe) [cm ² V ⁻¹ s ⁻¹]	μ_h/μ_e
0%	22.1	97.5%	83.3%	2.6×10 ⁻⁴	2.1×10 ⁻⁴	1.2
10%	22.9	98.3%	88.8%	2.8×10 ⁻⁴	2.8×10 ⁻⁴	1.0
20%	20.9	97.2%	86.9%	4.2×10 ⁻⁴	3.0×10 ⁻⁴	1.4

In summary, PBDB-T:m-INPOIC binary non-fullerene OSCs with a maximum PCE of 12.8% were enhanced to 13.9% PCE by employing 10 wt% IT-M (relative to the acceptor) as a ternary component, with simultaneously increased J_{sc}, FF and V_{oc}. Morphological studies show appreciably enlarged phase domain size via the addition of IT-M, helping to increase the molecular packing of PBDB-T, as well as creating continuous donor and acceptor networks for efficient charge transport towards the respective electrodes. The optimized morphology via tuning of the phase separation of the ternary system leads to increased light absorption and charge mobility, balanced charge transport and suppressed carrier recombination. We have there demonstrated that ternary component implementation is an effective strategy to prepare high-performance ternary non-fullerene OSCs consisting of components with low structural orders.

ASSOCIATED CONTENT

Supporting information

Materials, fabrication and characterization of OSCs, mobility calculation, GIWAXS and GISAXS fitting details and data.

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Notes

The authors declare no competing financial interest.

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