1	Single-junction organic solar cells with over 19% efficiency enabled by a
2	refined double-fibril network morphology
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29 Abstract

In organic photovoltaics, morphological control of donor and acceptor domains on the 30 nanoscale is key for efficient exciton diffusion and dissociation, carrier transport, and suppression 31 of recombination losses. To realise this, here, we demonstrated a double-fibril network based on 32 ternary donor: acceptor morphology with multi-length scales constructed by combining ancillary 33 34 conjugated polymer crystallizers and non-fullerene acceptor filament assembly. Using this approach, we achieved an average power conversion efficiency of 19.3% (certified 19.2%). The 35 36 success lies in the good match between the photoelectric parameters and the morphological characteristic lengths, which utilizes the excitons and free charges efficiently. This strategy leads 37 to enhanced exciton diffusion length (hence exciton dissociation yield) and reduced recombination 38 rate, hence minimizing photon-to-electron losses in the ternary devices as compared to their binary 39 counterparts. The double-fibril network morphology strategy minimizes losses and maximizes the 40 power output, offering the possibility towards 20% power conversion efficiencies in 41 single-junction organic photovoltaics. 42

43 Main

Organic semiconductors offer the advantage of high optical absorption and tunable energy 44 levels, enabling thin-film solar cells with high light-to-electron conversion efficiencies over a wide 45 range of wavelengths¹⁻⁴. Desipte recent progresses, the performance of organic solar cells (OSCs) 46 is still limited by non-ideal exciton and charge transport, which depend not only on the electronic 47 structure of organic semiconductors but also on the nanostructure that is formed by material 48 crystallization and phase separation in a bulk heterojunction (BHJ) setting⁵⁻⁸. A suitable sized 49 phase-separated morphology that balances crystalline region and mixing domain on the nanoscale 50 is therefore needed to further push the power conversion efficiency (PCE) of OSCs, however it is a 51

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nontrivial endeavor since the thin-film deposition is a non-equilibrium process that depends on multiple factors including the chemical structures and processing conditions⁹. 53

A useful strategy to manipulate thin-film morphology is the introduction of a third component 54 which can selectively interact with donor or acceptor to induce extra morphology control 55 handles^{6,10-15}. And by using enthalpy dictated crystallization, a crystallization induced 56 phase-separated morphology with suitable size scale can be achieved^{16,17}, creating an intertwined 57 donor (D)-acceptor (A) bicontinuous network with domains on the tens of nanometers scale, such 58 that efficient exciton and carrier transport can be achieved. Meanwhile, the size of the mixing 59 domain needs to be refined to enhance the carrier diffusion before on board the crystalline 60 transport phase, which dictates the efficiency of carrier transport at conditions close to open circuit, 61 hence influencing the fill factor (FF) of the device. 62

In this work, a bicontinuous double-fibril network morphology (DFNM) is demonstrated using 63 a carefully designed ternary mixture, in which the phase separation, crystallinity, and specific 64 fibril widths are collectively manipulated. The resulting nanostructure matches well with the 65 requirements for efficient exciton and free charge transport with ~40 nm exciton diffusion length, 66 ~105 nm free charge diffusion length, and ~2700 nm free charge drift length (estimated at short 67 circuit condition) 18,19 . The matured crystallization of polymer donor and non-fullerene acceptor 68 (NFA) fibrils in the blended thin film leads to a reduced mixing zone and suppressed 69 recombination. An extraordinary FF of ~82% is recorded and a breakthrough in average PCE of 70 19.3% with a maximum value of 19.6% is obtained in the optimized single-junction OSCs. 71 Similar improvements are seen in parallel experiments with a similar morphology, showing the 72 bright future of the DFNM perspective. 73

Materials and device performance 74

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75	The chemical structure and absorption spectra of materials used in this research are shown in
76	Fig. 1a and Supplementary Fig. 1. L8-BO primarily absorbs light between 550 and 900 nm ²⁰ .
77	PM6 and D18 in combination are used as donors to absorb short-wavelength photons from 400-
78	700 nm and to manipulate material crystallization forming a high-quality crystalline donor phase.
79	The highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital
80	(LUMO) energies are -5.20 eV/ -3.06 eV for PM6, -5.24 eV/ -2.95 eV for D18, and -5.67
81	eV/-3.92 eV for L8-BO measured by ultraviolet photoelectron spectroscopy (UPS) and inverse
82	photoemission spectroscopy (IPES) ²¹ , respectively (Fig. 1b and Supplementary Fig. 2). The
83	current density-voltage $(J-V)$ curves and performance parameters of the devices are shown in
84	Fig. 1c, Supplementary Fig. 3, and Table 1. The device based on PM6:L8-BO achieved a
85	maximum PCE of 18.2% with an open-circuit voltage (V_{oc}) of 0.888 V, a short-circuit current
86	density (J_{sc}) of 25.7 mA cm ⁻² , and a FF of 79.9%, while D18:L8-BO showed a V_{oc} of 0.907 V, a
87	$J_{\rm sc}$ of 24.9 mA cm ⁻² , a FF of 78.5%, and a PCE of 17.7%. The best mixing weight ratio of
88	PM6:D18:L8-BO=0.8:0.2:1.2 offers a maximum PCE of 19.6% with a V_{oc} of 0.896 V, a J_{sc} of
89	26.7 mA cm ⁻² , and a FF of 81.9%. Such device obtained a certified PCE of 19.2% subject to the
90	calibration procedures of the National Photovoltaic Industry Metrology and Testing Center
91	(NPVM, China, Supplementary Fig. 4), which is a breakthrough from the previous record ^{4,22-25} .
92	The highly crystalline polymer D18 is used as the seeding agent for the moderate crystalline
93	polymer PM6 in order to retain the advantageous PM6:L8-BO morphology framework while
94	enhancing the crystallinity of the blend ^{20,26} . An optimum amount of D18 added into PM6:L8-BO
95	raises J_{sc} and FF simultaneously (Fig. 1d, Supplementary Fig. 5a and Supplementary Table 1),
96	Meanwhile, the glass-encapsulated devices exhibited ~95% of the initial PCE after 2520 hours of
97	storage in the glove box at 25 °C. More importantly, the ternary device showed better

98 illumination stability than that of binary devices, indicating that the introduction of D18 can 99 better balance the photochemical and morphology stability of the blend system (shown in 100 Supplementary Fig. 6). The external/internal quantum efficiency (EQE/IQE) results show an 101 increase in EQE of the ternary device due to more efficient exciton diffusion, which contributes 102 to improved J_{sc} and PCE (see Fig. 1e, Supplementary Figs. 5b,c and 7, and Supplementary Note 103 1 for details.) The statistics of solar cell devices were shown in Fig. 1f and a jump in PCE value 104 is recorded in the ternary mixture.

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Morphology details of double-fibril network

To understand the origin of the encouraging device performance, we investigated the film 106 microstructure. As shown by transmission electron microscopy (TEM) investigation (Fig. 2a and 107 108 Supplementary Fig. 8), a fibrillar network structure is seen in all blended films. The fibril texture in primitive PM6:L8-BO blends is roughly defined with a smaller fibril diameter and more 109 sparse fibril distribution. D18 is a strongly crystalline polymer and of slightly low solubility than 110 111 PM6, which precipitate out in an earlier film growth stage and induce crystallization. Adding D18 into PM6 helps to improve the crystallinity of the blends by forming a high-quality fibril 112 network with increased fibril diameter and distribution, as evidenced by atomic force microscope 113 (AFM) and scanning electron microscope (SEM) characterizations (Supplementary Figs. 9 and 114 10). The Fourier transformation yields an inter-fibril distance ~25 nm, a suitable length scale of 115 116 phase separation to balance exciton splitting and carrier transport. With this phase image, we 117 then looked into the detailed phase construction of the blended thin-films. Photo-induced force microscope (PiFM) was employed that uses localized near field infrared imaging to identify the 118 119 chemistry associated with thin-film morphology (see Fig. 2b, Supplementary Figs. 11 and 12, and Supplementary Note 2 for details). The corresponding PiFM phase images of 120

PM6:D18:L8-BO blends were shown in Fig. 2b and Supplementary Fig. 12a. Interestingly, both 121 PM6 and L8-BO formed fibril network structures in the blend. The overlay of the two images 122 (Supplementary Fig. 12b) yields an entangled DFNM with donor and acceptor domain in close 123 contact. Such observation provides a good understanding of the OSCs' morphology, indicating 124 that crystallization-induced morphology can result in better phase connectivity and finer sizes. 125 Line profiles across PiFM images were shown in Fig. 2b, full width at half maximum (FWHM) 126 of the peaks was used to estimate the fibril diameter, yielding average values of 22.1 and 22.6 127 nm for acceptor and donor fibrils. In combination with the inter fibril distance value, it can be 128 concluded that the mixing domain volume is low, indicating low geminate recombination and 129 high FF as discussed later in more detail. 130

Grazing incidence X-ray diffraction (GIXD) was used to investigate the thin film 131 crystallinity²⁷. The 2D diffraction patterns and 1D line cuts are summarized in Fig. 2c,d and 132 Supplementary Fig. 13a-c. Both donor and acceptor show good crystalline order in neat films 133 (show in Supplementary Fig. 13a and Supplementary note 3). In blended films, both conjugated 134 polymers and NFA are preferentially face-on oriented according to π - π stacking peak locations, 135 which facilitate carrier transport. The overlapping of lamellar and π - π stacking peaks were used 136 to evaluate the structure order of the thin film (shown in Supplementary Fig. 14 and 137 138 Supplementary Table 2). The D18:L8-BO film exhibits high crystallinity and large crystal size, where the PM6:L8-BO film shows weak crystallinity and small crystal size, and the 139 PM6:D18:L8-BO film lies in-between and is close to PM6:L8-BO film. The (021) diffraction 140 peak in L8-BO was used to label the NFA crystallization feature in blends (Supplementary Figs. 141 142 13d,e and 15 and Supplementary Table 3). It is found that D18 retains a good crystallinity in ternary blends as evidenced by its (001) diffraction intensity and crystal coherence length (CCL). 143

D18's presence also promotes the crystallization of L8-BO, judging from its (021) peak area 144 enhancement. The small amount of D18 in ternary blend promotes the crystallization of both 145 donor polymer and NFA molecules. We ascribe such effect to the heterogeneous nucleation 146 seeding mechanism^{28,29}, which helps to increase the fibril content and crystallite quality in a 147 blended thin-film. 148

Resonant soft X-ray scattering (RSoXS) is used to investigate the phase separation at the 149 energy of 285.2 and 286.7 eV related to C 1s $\rightarrow \pi^*$ transition (see Supplementary Figs. 16 and 150 17, and Supplementary Note 4 for details) by taking advantage of energy-dependent chemical 151 sensitivity³⁰⁻³⁴. The D18:L8-BO blends show a broad scattering hump with the center around 152 0.0098 Å⁻¹ (64.1 nm) at both photon energy (Supplementary Fig. 17a), where the PM6:L8-BO 153 blends show a better-resolved scattering hump at 285.2 eV located at 0.0131 Å⁻¹ (47.8 nm) 154 (Supplementary Fig. 17b). Introducing D18 into PM6:L8-BO leads to the enhancement of 155 RSoXS with a slightly reduced hump center ($q = 0.0130 \text{ Å}^{-1}$, 48.2 nm) (Supplementary Fig. 17c). 156 NK-NEXAFS was conducted to further clarify the chemical origin of the contrast (as shown in 157 Supplementary Fig. 17d). From the NK-NEXAFS of the nitrogen-containing material D18 and 158 L8-BO, the absorption of the blend film at 400 eV is only from L8-BO (Supplementary Fig. 159 S16b). Therefore, the phase separation scales of L8-BO in the three blend films are 68.8 nm 160 161 (D18:L8-BO), 49.0 nm (PM6:L8-BO), and 51.8 nm (PM6:D18:L8-BO), respectively, which is consistent with the result of 285.2 eV in CK-RSoXS. Thus, D18 could induce a better-defined 162 fibril structure in ternary blends by improving the fibril diameter, as demonstrated by AFM and 163 PiFM characterization. It is noted that the inter-fibril distance is only slightly larger than the 164 diameter of the fibril, thus the mixing region in-between the fibrils is small (summarized in 165 Supplementary Table 4). Therefore, the charges in the mixing zone could effectively diffuse to 166

the fibril network and be collected by electrodes. Scattering anisotropy is seen in RSoXS (Supplementary Fig. 17), which is parallel with a polarized electrical field at 285.2 eV. Thus, the L8-BO fibril/mixing zone interface is "edge-on" oriented^{35,36}. A schematic illustration of this DFNM is shown in Fig. 2e. The closely packed D-A fibrils in the mixture matrix enjoy an efficient exciton splitting and carrier transport, as will be shown later.

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Exciton diffusion and photophysical analysis

To further clarify the the exciton diffusion properties in the films based on the optimized DFNM, we carried out exciton annihilation method to measure exciton diffusion constants (see Supplementary Table 5 and Supplementary Note 5 for details). The results show that the addition of D18 to PM6:L8-BO increased the exciton diffusion length in the donor phase by $\sim 11\%$, steming from the optimized morphology with increased in fibril crystallinity and diameter.

Next, we carried out transient absorption spectroscopy (TAS) measurements to look into the 178 exciton and carrier dynamics within DFNM (see Supplementary Figs. 18–21, and Supplementary 179 Note 6 for details)^{37,38}. In blended films (Supplementary Figs. 19–21) with the decay of L8-BO 180 bleach and stimulated emission (SE) peak at ~840 nm, D18 bleach peak in 550-650 nm rises, 181 indicating the hole transfer process. The kinetics of 600 nm in three blends are extracted 182 (Supplementary Fig. 22) and fitted with a double exponential decay function (Supplementary 183 Table 6). The hole transfer in PM6:L8-BO is faster compared to D18:L8-BO (smaller τ_1 value), 184 and exciton diffusion time is longer in D18:L8-BO compared to PM6:L8-BO (larger τ_2 value). 185 Such observation implies the differences in morphology, in which PM6:L8-BO forms a good 186 mixing zone and D18:L8-BO has a larger NFA fibril diameter. The ternary blends show a τ_1 187 value close to that of PM6:L8-BO and a τ_2 value close to that of D18:L8-BO. Thus, the 188

advantages of the two binary blends are naturally collected in the ternary blend, leading to superior performance in J_{sc} and FF. The kinetics of the polaron state was shown in Supplementary Fig. 23. The ternary blend film shows the slowest decay, originating from the improved morphology that surpresses recombination.

193 **Device physics**

The photocurrent (J_{ph}) as a function of effective voltage shows the ternary blends have a 194 higher capability in both charge generation and transport. (see Supplementary Fig. 24, 195 Supplementary Table 7 and Supplementary Note 7 for details). The charge transport properties 196 are further evaluated through space charge limited current (SCLC) method, as shown in 197 Supplementary Fig. 25 and Supplementary Tables 8 and 9. Both hole and electron mobilities are 198 on the order of 10⁻³ cm² V⁻¹ s⁻¹. For PM6:L8-BO devices, the lower hole mobility sets the 199 fundamental limit, hence limiting the performance. The introduction of D18 leads to high and 200 balanced transportation (Fig. 3a) that can reduce the likelihood of bimolecular and trap-assisted 201 recombinations. Transient photovoltage (TPV) measurements were performed to investigate the 202 carrier dynamics under different Voc conditions³⁹. As shown in Supplementary Fig. 26a, the 203 lifetimes of the ternary devices show an enhancement in a broad operating range, indicating less 204 free charge recombination in the ternary blends than the binaries. Combining the lifetime values 205 with the extracted mobility values, the carrier diffusion and drift length at short circuit^{18,19} can be 206

estimated via
$$L_{\rm d} = \sqrt{\frac{k_{\rm B}T}{q}\mu\tau}$$
 and $L_{\rm dr} = \frac{\mu\tau V_{\rm bi}}{d}$, respectively (see Fig. 3b, Supplementary Table

10, and Supplementary Note 8 for details). All devices show high free charge drift length from close to 1000 nm to above 2000 nm (way above device thickness), indicating efficient charge collection at short circuit conditions and negligible loss of IQE, EQE, and J_{sc} due to free charge

collection losses. The difference between the diffusion lengths (Fig. 3b) could explain the 211 enhanced FF in the ternary device. Moreover, in the ternary device, a balanced diffusion length 212 of ~105 nm and drift length (at short circuit condition) of ~2700 nm are obtained, which in 213 combination with the typical photogeneration depth, guarantees an efficient ambipolar charge 214 transport in ternary blends. Moreover, transient photocurrent (TPC), TPV, and Mott-Schottky 215 methods were used to probe the density of states (DoS) and recombination mechanisms. We find 216 that the ternary device possesses the lowest recombination coefficient and the highest occupied 217 density of states of LUMO, as shown in Fig. 3c,d, and that trap assisted recombination has 218 limited contribution (see Supplementary Figs. 27-29, Supplementary Table 11, and 219 Supplementary Note 9 for details). 220

So far, the key findings here are: i) exciton diffusion and charge generation are enhanced; ii) bimolecular recombination is suppressed; iii) charge transport is optimized; iv) high density of states is obtained in the ternary devices; and v) trap-assisted recombination is negligible in all devices, which are consistent with improved EQE, IQE, J_{sc} , and FF in the ternary devices.

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Exciton diffusion, optical, and electrical simulation

To further support experimental results, we here perform quantitative device modelling, using 226 a coupled exciton diffusion, optical transfer matrix, and drift diffusion model based on 227 gpvdm⁴⁰⁻⁴². Specifically, the effects of exciton diffusion efficiency and free charge 228 recombination rate constant on IQE/EQE and FF are modelled (see Supplementary Figs. 30-32, 229 230 Supplementary Tables 12–13, and Supplementary Note 10 for details). By modelling exciton 231 diffusion within a single, spherical, acceptor domain whose size is determined by the phase separation length, and using exciton diffusion lengths from Supplementary Table 12, we 232 demonstrate that exciton collection is more efficient from the ternary and PM6:L8-BO structures 233

 $(\sim 97\%)$ than from the lowest performing D18:L8-BO structure ($\sim 94\%$). Further simulations based on optical and electrical models also demonstrated the highest IQE/EQE, J_{sc} , and FF in the ternary blends (see Supplementary Fig. 32), which is a collective effect by the efficient exciton diffusion and suppressed recombination, further supporting the claim that enhanced exciton dissociation yield and reduced free charge recombination are important reasons for the performance enhancement in the ternary device.

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Formation mechanism of DFNM

While the conjugated polymer shows a high tendency to form crystalline fibrils, the NFA 241 molecule crystallization behavior is a curious property to investigate. Conventional NFAs that 242 rely on π - π stacking to form ordered structures have a high tendency to form 2D brick wall 243 packing or 3D network packing^{43,44}. The Y6 molecules with a banana geometry show a different 244 packing motif whereby they align their molecular long axes to form a polymer-like structure and 245 extend along the body center diagonal in the lattice plane⁴⁵. The L8-BO with a long branched 246 side-chain gives slightly stronger alkyl chain interactions and adopts a more condensed packing 247 with the long axis along the face center half diagonal, forming smaller domains between 248 conjugated backbones in crystals. A careful examination of the single crystal yields more 249 interesting information in which multiple dimer packing motifs are seen (Fig. 4a-c) and 8 250 monomer packing is needed to form a complete period. More importantly, several linear 251 molecular assemblies intertwined together to form a strong assembly with a high aspect ratio 252 (Fig. 4b,c) that can be of high tolerance to local structure defects. These properties make L8-BO 253 forming needle-type single crystals (Fig. 4d). In the TEM diffraction investigation, it is 254 confirmed that the molecular long axis is aligned with needle packing length (Fig. 4e). Such 255 nature is preserved to form fibrils in blended thin films (Fig. 4f). 256

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Material selection rules for constructing DFNM

DFNM enables efficient exciton and free charge transport in the blended thin films studied 258 herein. To confirm that, we try to correlate the fundamental photophysical properties of donor 259 and acceptor materials in the blend with the structural features. A large group of acceptor 260 materials was studied to look into the detailed photophysical property change versus material 261 nature. The representative exciton decay kinetics as a function of excitation density of L8-BO is 262 shown in Supplementary Fig. 33, from which the exciton diffusion length can be estimated. As 263 shown in Fig 5a, Supplementary Figs. 34 and 35, and Supplementary Table 14, we see that 264 PCBM acceptors have a low diffusion length around 5 nm, ITIC families have a diffusion length 265 of 10–20 nm, Y6 based NFAs show a much larger diffusion length \sim 40 nm, which is well above 266 the fibril diameter and should guarantee a highly efficient exciton splitting. The exciton diffusion 267 in the donor phase is also investigated as complementary to acceptors (Fig. 5b, Supplementary 268 Figs. 36–39, and Supplementary Table 14). The conventionally used donor materials such as 269 270 PM6, PM7, and PTQ10 show an exciton diffusion length around 14 nm, which is much lower compared to that in the high-efficiency Y6 acceptor family. Thus, enhancing the exciton 271 diffusion of polymer donors can be a viable approach in improving photo harvesting. We noticed 272 that several high crystalline donor polymers have higher exciton diffusion lengths compared to 273 the above-mentioned donor polymers, and thus designed the binary donor polymer blends to 274 improve exciton diffusion. Adding a secondary, high crystalline donor (20 wt.%) such as D18 275 could improve the exciton diffusion length to around 20 nm, which better matches the polymer 276 fibril's diameter and catches up with that in NFAs. We believe this property could result in better 277 photo management in devices, leading to high J_{sc} and FF. 278

Last, solar cell performances in these NFA-based systems (chemical structure of materials were summarized in Supplementary Fig. 40) are used to establish the correlation between

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photophysical features and morphological structures. As shown in Fig. 5c, Supplementary Figs. 281 41–43, and Supplementary Tables 15–17, conventional ITIC analogs using the ternary blends do 282 not help to improve device performance, and obvious FF drop is recorded in several blends. We 283 ascribe these results to unoptimized NFA morphology in which domain connection is an issue 284 (no similar fibril network channels are formed). In the Y6-based NFA blended system, the 285 ternary blends with improved exciton diffusion length and donor-acceptor crystallinity do help to 286 improve the J_{sc} and FF, resulting in higher PCEs. In addition, we ran experiments on more 287 blending ratios and different material systems to monitor the device performances and 288 characteristics (see Supplementary Figs. 44-49, Supplementary Tables 18-21, and 289 Supplementary Note 11 for details). The results show that an appropriate reduction of the mixing 290 zone size can increase exciton utilization. The high-quality double fibril transport channel 291 ensures the efficient diffusion of carriers and is no longer a shortboard that restricts the 292 performance of the device. 293

294 Outlook

From the above-discussed observations, the successful demonstration of adding D18 into the 295 PM6:L8-BO blends provides a showcase of average efficiency of 19.3% (certified 19.2%), 296 associated with very high J_{sc} and FF. The device performance improvement was achieved by 297 DFNM. Such a morphological scheme requires both high-quality D-A crystallites to form a 298 299 multi-length scaled dual fibril network, facilitating exciton diffusion and charge carrier transport. This morphology also reduces the mixing zone content and feature size, from which exciton 300 splitting and carrier generation are more efficient. Meanwhile, the observation of NFA fibril-type 301 302 network crystallization is encouraging, which provides insights for the next generation material design. Moreover, the synergetic combination of D18 and PM6 adds up the morphological and 303

photophysical advantages of the donor phase network to allow balanced exciton/free charge 304 transport and recombination, contributing to the improvement of J_{sc} and FF. To sum up, the 305 306 advantage of DFNM has been demonstrated, showing a bright future for its application in OSCs.

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Author contributions 321

F.L. and Y.S. conceived and directed this project. L.Z. fabricated and characterized the OPV 322 devices. L.Z. and M.Z. conducted the certification. J.X. processed and analyzed the single-crystal 323 data. C.L. synthesized L8-BO. M.Z. and T.H. carried out the TPV, TPC, impedance 324 characterization and analyzed the data. TA results and corresponding analysis were provided by 325 G.Z. and H.Z. W.Z. carried out the GIXD and RSoXS measurement and assisted with data analysis. 326

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327	J.S. conducted the AFM measurements. J.Y., R.C.M., and J.N. conduct the drift diffusion
328	simulation and analysis. Yecheng Zou conducted the TEM measurements. Yongming Zhang, X.X.
329	Z.Z., and R.Z. contributed to the fruitful discussion of this project. L.Z. and M.Z. wrote the
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331	manuscript. This manuscript was mainly prepared by F.L., Y.S., L.Z., M.Z. and J.X., and all
332	authors participated in the manuscript preparation and commented on the manuscript.

333 **Competing interests**

The authors declare no competing interests.

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Table 1 | Photovoltaic parameters of OSCs under the illumination of AM 1.5 G, 100 mW/cm².

Blends	Device Area (mm ²)	$V_{ m oc}$ (V)	$J_{\rm sc}~({ m mA~cm^{-2}})$	FF (%)	PCE (%)
D10.1 0 DO2	5.2°	$0.907~(0.908\pm0.001)$	$24.9~(24.3\pm 0.5)$	$78.5\;(78.5\pm1.0)$	17.7 (17.4 ± 0.2)
D18:F8-RO _*	14.2 ^d	$0.907~(0.908\pm0.001)$	$24.8\;(24.2\pm 0.5)$	77.6 (77.4 \pm 1.3)	17.5 (17.1 ± 0.2)
	5.2°	$0.888~(0.888\pm0.001)$	$25.7~(25.5\pm0.3)$	79.9 (79.5 \pm 1.1)	$18.2~(18.0\pm 0.1)$
PM6:L8-BO ^a	14.2 ^d	$0.888~(0.889\pm0.001)$	$25.6~(25.6\pm 0.5)$	78.8 (78.4 ± 1.4)	17.9 (17.6 ± 0.2)
	5.2°	$0.896~(0.896\pm0.001)$	$26.7~(26.6\pm 0.3)$	81.9 (81.1 ± 0.8)	$19.6~(19.3\pm 0.1)$
PM6:D18:L8-BO ^b	14.2 ^d	$0.897~(0.897\pm0.001)$	$26.7~(26.3\pm 0.4)$	81.2 (80.4 ± 1.3)	$19.4~(19.0\pm 0.2)$
	5.2	0.891 ^e	26.7°	80.8 ^e	19.2 ^e

^a1:1.2 (weight ratio); ^b0.8:0.2:1.2 (weight ratio); ^cthe average parameters were calculated from 50 devices, with an aperture area of 3.2 mm²; ^dthe average parameters were calculated from 20 devices, with an aperture area of 10.4 mm². The values outside the parentheses denote the best optimal photovoltaic parameters, and the values inside the parentheses represent the average photovoltaic parameters and their standard deviations. ^cCertified by National Photovoltaic Industry Metrology and Testing Center (NPVM), China. The designated illumination area was 3.084 mm².

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Fig. 1 | Materials and device performance. Chemical structures (a) and energy level lineup (b) of PM6, D18, and L8-BO. (c) J-V curves of the devices based on D18:L8-BO, PM6:L8-BO, and PM6:D18:L8-BO (0.8:0.2:1.2) with a device area of 5.2 mm². (d) PCE, J_{sc} , and FF values of devices based on PM6:D18:L8-BO with different donor ratios. Error bars represent the standard error of the mean (n=50). (e) EQE curves of the devices based on PM6:L8-BO, D18:L8-BO, and PM6:D18:L8-BO. Inset is the change curves in EQE of the ternary device relative to the binary
 devices. (f) Histogram of PCE counts for 50 devices based on PM6:L8-BO, D18:L8-BO, and
 PM6:D18:L8-BO.

Fig. 2 | Morphology of thin films. (a) TEM image of PM6:L8-BO and PM6:D18:L8-BO blend 352 films. (b) PiFM image at the wavenumber of 1532 cm^{-1} (representing L8-BO) and 1648 cm^{-1} 353 (representing PM6) and the line profile to obtain the fibril width for PM6:D18:L8-BO blended 354 film. The fibril width is in statistic from the full width at half maximum (the distance between 355 the two adjacent dashed lines in the graph) (c) 2D GIXD patterns of PM6:L8-BO and 356 PM6:D18:L8-BO blends. (d) In-plane (IP) and out-of-plane (OOP) line cut profiles of the 2D 357 GIXD data based on binary and ternary blend films. (e) The scheme of DFNM with the charge 358 transfer process. Robust fibrils in network surrounded by tiny ones. Such a hierarchical fibril 359 network could form a good connection via chain segments with the mixing region, leading to a 360 multi-length scaled morphology. 361

Fig. 3 | Device physics characterization. Carrier mobilities (a) and carrier diffusion length (b) of 362 the binary and ternary devices (μ_e : electron mobility, μ_h : hole mobility, L_e : electron diffusion 363 length, L_h : hole diffusion length). Error bars represent the standard error of the mean (n=5). (c) 364 The relationship between charge density and non-geminate recombination rate coefficient of the 365 binary and ternary devices. The data points correspond to different light intensity and the filled 366 symbol correspond to values at 1 sun intensity. (d) The derived occupied LUMO DoS from the 367 capacitance spectra of the binary and ternary devices, exhibiting an exponential shape, where N is 368 the total density per unit volume, and δ is the energetic disorder parameter. 369

Fig. 4 | Single-crystal structure of L8-BO and the scheme of DFNM. (a) The single twisted chain of L8-BO with three different dimer stacking modes. Each dimer is composed of two adjacent L8-BO. Top view (b) and side view (c) stacking diagram of three L8-BO twisted chains. (d) The optical micrographs of L8-BO single crystal. (e) Transmission electron microscopy (TEM) selected-area electron diffraction (SAED) patterns of L8-BO single crystal. The insert is the TEM bright-field (BF) image of L8-BO single crystal with labeled spindle axis direction. (f) The scheme of DFNM.

Fig. 5 | Exciton diffusion length and device parameters. (a) Exciton diffusion length in different 377 acceptors (PC₆₁BM, PC₇₁BM, ITIC, IT4F, IDIC, Y6, BTP-eC9, and L8-BO). (b) Exciton diffusion 378 length in different single or double blend donors (PM6, PM7, PTQ10, D18, D18-Cl, and 379 PBT1-C-2Cl; PM6:D18=4:1, PM6:D18-Cl=4:1, PM6:PBT1-C-2Cl=4:1, PM7:D18=4:1, and 380 PTQ10:D18=4:1). (c) Violin plots of PCE, FF, J_{sc} , and V_{oc} distribution of the binary (red) and 381 ternary (blue) devices. System list: A: PM6:ITIC and PM6:D18:ITIC; B: PM6:IT4F and 382 PM6:D18:IT4F; C: PM6:IDIC and PM6:D18:IDIC; D: PM6:IDTT-C10-TIC 383 and PM6:D18:IDTT-C10-TIC; E: PM6:Y6 and PM6:D18:Y6; F: PM6:BTP-eC9 and 384 PM6:D18:BTP-eC9; G: PM6:L8-BO and PM6:D18:L8-BO; H: PM7:Y6 and PM7:D18:Y6; I: 385 PTQ10:Y6 and PTQ10:D18:Y6; J: PM7:L8-BO and PM7:D18:L8-BO; K: PTQ10:L8-BO and 386 PTQ10:D18:L8-BO; L: PM6:Y6 and PM6:D18-Cl:Y6; M: PM6:Y6 and PM6:PBT1-C-2Cl:Y6; N: 387

PM6:L8-BO and PM6:D18-Cl:L8-BO; O: PM6:L8-BO and PM6:PBT1-C-2Cl:L8-BO. A-D: non-DFNM, E-O: DFNM. (n=50 for PM6:L8-BO, D18:L8-BO, and PM6:D18:L8BO, n=20 for the remaining conditions)

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483	Meth	ods		
484	Devic	re Fabrication. Organic solar cell devices with indium tin oxide (ITO)		
485	/poly((3,4-ethylenedioxythiophene):poly(styrene sulfonate) (PEDOT:PSS)/active		
486	layer/	layer/PNDIT-F3N/Ag structures were fabricated according to the following procedure. Patterned		
487	ITO g	ITO glass substrates were sequential cleaned by ultrasonication in detergent, deionized water,		
488	acetor	acetone, and isopropyl alcohol for 20 min each and then dried under a dry oven. The precleaned		
489	substr	substrates were treated in an ultraviolet-ozone chamber for 15 min, then a ~20 nm thick		
490	PEDO	PEDOT:PSS (Clevious P VP AI 4083 H. C. Stark, Germany) thin film was deposited onto the ITO		
491	surfac	surface by spin-coating and baked at 150 °C for 15 min. The solution of PM6:L8-BO (1:1.2, wt.,		
492	14.3 1	14.3 mg mL ^{-1} in total), D18:L8-BO (1:1.2, wt., 13.2 mg mL ^{-1} in total), and PM6:D18:L8-BO		

493	(0.8:0.2:1.2, wt., 14.3 mg mL ^{-1} in total) in chloroform with 1,4-diiodobenzene (DIB) as a solid
494	additive (the content of DIB is 50% of the total mass of donor and acceptor) was stirred at 25 $^{\circ}$ C
495	for 2 hours in advance and then spin-coated on top of the PEDOT:PSS layer. The optimum
496	thickness of the active layer is 120 nm (statistical variations within 5 nm). The prepared films were
497	treated with thermal annealing at 85 °C for 5 min. After cooling to room temperature, a \sim 5 nm
498	thick of PNDIT-F3N (0.5 mg mL ^{-1} in methanol with 0.5% acetic acid, v/v) was spin-coated on the
499	top of the active layer. Then, those samples were brought into an evaporating chamber and a 150
500	nm thick silver layer was thermally evaporated on the PNDIT-F3N layer at a base pressure of 1 \times
501	10^{-6} mbar. The evaporation thickness was controlled by SQC-310C deposition controller
502	(INFICON, Germany). Ten devices were fabricated on one substrate and the active area of each
503	device was 0.052 cm^2 defined by a shadow mask.













- [101]

4.60 nn

€.61 nm

** (222)

(202)

