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Intrinsic Efficiency Limits in Low-bandgap Non-fullerene Acceptor Organic Solar Cells

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In bulk heterojunction (BHJ) organic solar cells (OSCs) both the electron affinity (EA) and ionization energy (IE) offsets at the donor-acceptor interface should equally control exciton dissociation. Here, we demonstrate that in low-bandgap non-fullerene acceptor (NFA) BHJs ultrafast donor-to-acceptor energy transfer precedes hole transfer from the acceptor to the donor and thus renders the EA offset virtually unimportant. Moreover, sizeable bulk IE offsets of about 0.5 eV are needed for efficient charge generation and high internal quantum efficiencies, since energy level bending at the donor-NFA interface caused by the acceptors' quadrupole moments prevents efficient exciton-to-charge-transfer (CT) state conversion at low IE offsets. The same bending however, is the origin of the barrier-less CT state to free

charge conversion. Our results provide a comprehensive picture of the photophysics of NFA-based blends, and prove that sizeable bulk IE offsets are essential to design efficient BHJ OSCs based on low-bandgap NFAs.

The minimum driving force required for efficient charge separation in bulk heterojunction (BHJ) organic solar cells (OSCs) has remained controversial.¹⁻¹⁰ In BHJs with low-bandgap non-fullerene acceptors (NFAs) both the electron donor and acceptor harvest photons, preferably in complementary spectral regions. Consequently, two channels of charge generation are expected, electron transfer from the donor to the acceptor and hole transfer from the acceptor to the donor. In thin film BHJs, the former is driven by the electron affinity (EA) offset, the latter by the ionization energy (IE) offset.

In fullerene-based systems, the requirement of a minimal EA offset of around 0.3 eV was experimentally identified¹¹ and repeatedly confirmed.¹²⁻¹⁵ However, recent reports of efficient NFA-based BHJs imply that this requirement may not hold for NFAs. Indeed, a number of studies concluded that charge generation in NFA BHJs can be efficient even for zero offsets, contrary to fullerene-based systems.^{11,13-15} Moreover, negligible driving forces for charge separation via hole transfer^{8,16-20} and electron transfer²¹⁻²³ are often used as an explanation of low energy losses in high-performing NFA blends.²⁴

To determine IE and EA, characterization techniques such as ultraviolet photoelectron spectroscopy (UPS), photoemission spectroscopy in air (PESA), and cyclic voltammetry (CV) have been used. Notably, only CV supports the idea of zero IE^{8,16-20} and EA²¹⁻²³ offsets. However, CV does not provide reliable estimates of IEs and EAs of solid films.²⁵ PM6:Y6, for example, has zero offset according to CV, but 0.5 eV when measured by UPS.¹⁹ Apart from putting the necessity for a type-II heterojunction into question, efficient charge generation with zero energy offsets implies that the mechanisms of charge generation in NFA- and fullerene-based blends are different.²⁶ A few recent studies acknowledged that IE offsets are important for charge generation,^{9,27,28} demonstrating an increase of the charge generation efficiency with the IE offset, yet only for a very limited number of structurally-related systems.

A coherent study providing uniform characterization of a large NFA series as well as a model explaining the origin of the IE offset dependence in NFA-based systems and the recently reported barrier-less charge separation in NFA blends²⁹ is still missing, but needed for further optimization of energetic losses and internal quantum efficiency (IQE) of NFA BHJs.³⁰

Here, we demonstrate that the exciton-to-charge conversion efficiency (and, therefore, the IQE) of low-bandgap NFA-based BHJ solar cells increases with the donor-NFA IE offset, reaching its maximum for offsets of ~ 0.5 eV. Surprisingly, charge generation does not depend on the EA offset.

This is explained by ultrafast energy transfer from the donor to the acceptor. Indeed, by design, the photoluminescence (PL) of the higher bandgap donor overlaps with the absorption of the lower bandgap acceptor. This design principle broadens the BHJ's photoactive spectral region and thereby increases the maximum available short circuit current density, and it facilitates Förster resonance energy transfer (FRET) from the donor to the NFA.³¹⁻³⁴ Consequently, FRET is the major deactivation channel for donor excitons in NFA BHJs. In fullerene BHJs, FRET is limited by the low oscillator strength of fullerene molecules.³⁵

Furthermore, we demonstrate that bending of the energy levels at the donor-acceptor heterojunction, caused by the acceptors' quadrupole moments, increases the interfacial charge-transfer (CT) state energy. If this energy level bending is not counterbalanced by the IE offset, then it creates a barrier for exciton-to-CT state conversion. The same energy level bending however, leads to barrier-less CT state to free charge conversion, as recently observed experimentally.^{29,36} Importantly, the same energy level bending explains the presence of energetically high-lying CT states reported for many NFA-based systems.

A straightforward implication of our study is a clear design principle for NFA materials: high exciton-to-charge conversion yields can only be achieved for sizeable (~ 0.5 eV) bulk IE offsets, since only they provide sufficient driving force for dissociation of excitons to CT states, and further to separated charge carriers.

To reveal the impact of IE, EA, and FRET on charge generation in NFA BHJs, we studied representative donor-acceptor combinations: the small molecule electron donor DR3TBDTT (DR3),³⁷ the widely-used donor polymers PTB7-Th (PCE10) and PBDBT-SF (PCE13),³⁸ and the recently reported high-efficiency donor polymer PBDB-T-2F (PM6).³⁹ Ten small molecule NFAs were selected as acceptors, including Y6, one of the current state-of-the-art NFAs, as well as PC₇₁BM as reference.⁴⁰ Chemical structures of all materials are shown in Fig. 1 alongside their IE and EA values, determined by ultraviolet photoemission spectroscopy (UPS), and inverted photoemission spectroscopy (IPES) respectively, as well as the average internal quantum efficiencies (IQEs) of respective BHJs (details of UPS, IPES can be found in Supplementary Note 1).⁴¹ Since IE offsets are essential in our study, we confirmed the materials' IEs by another technique, PESA (Supplementary Note 1).^{41,42}

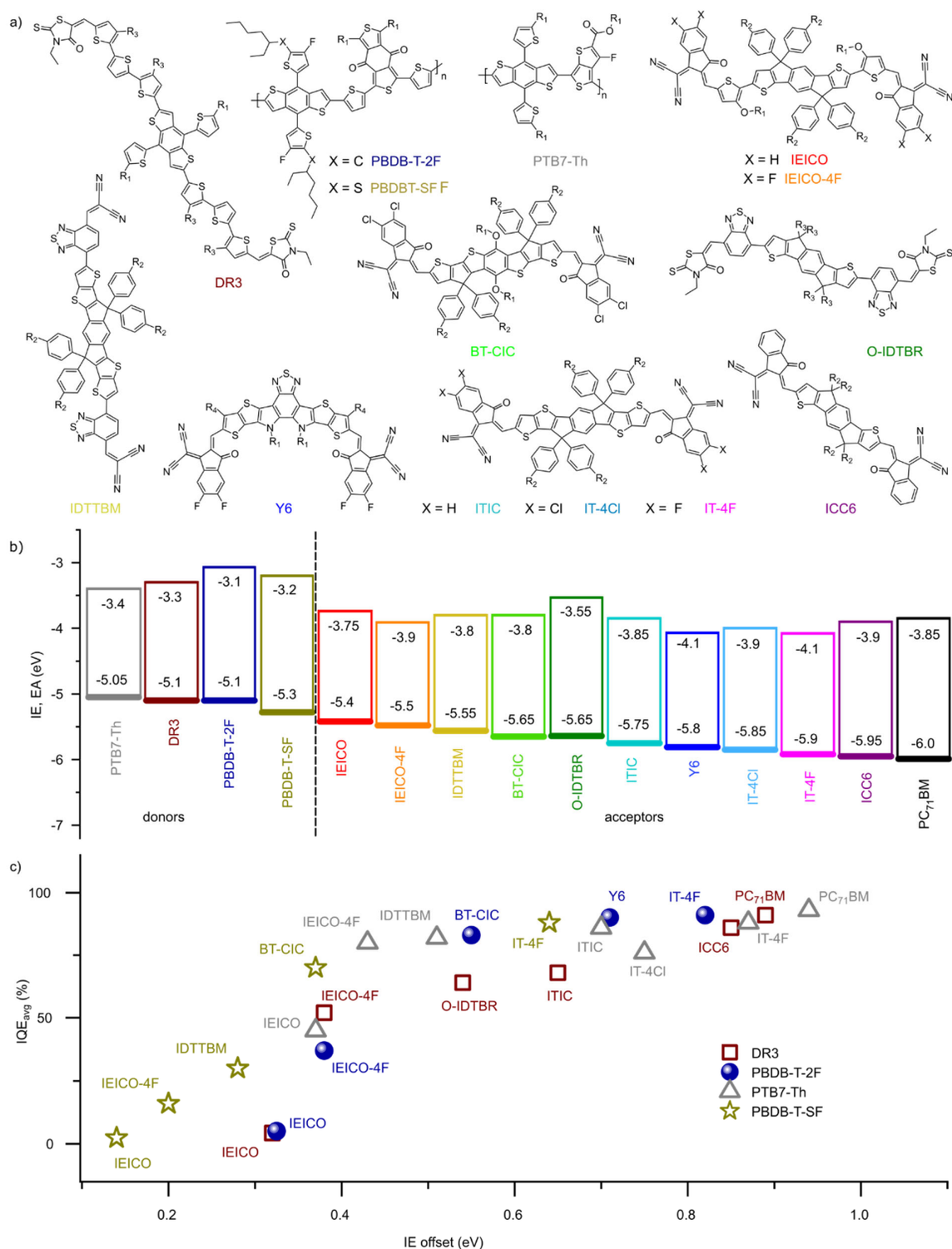


Fig. 1: Materials' structures, energy levels, and internal quantum efficiencies in photovoltaic devices. a) Chemical structures of the donor materials and non-fullerene acceptors investigated in this study. R₁: -CH(C₄H₉)(C₂H₅), R₂: -C₆H₁₃, R₃: -C₈H₁₇ and R₄: -C₁₁H₂₃. b) Ionization energy (IE, thick horizontal lines) determined by ultraviolet photoelectron spectroscopy (UPS) and electron affinity (EA, thin horizontal lines) by inverse photoemission spectroscopy (IPES). c) Spectrally averaged internal quantum efficiency (IQE_{avg}) of optimized bulk heterojunction donor-NFA devices as a function of the IE offset between donor and acceptor for the small molecule donor DR3 and the donor polymers PBDB-T-2F, PTB7-Th, and PBDB-T-SF. For OSC device data and preparation conditions, we refer to the Supplementary Note 2.

First, one intriguing observation is that the IQE averaged over the range of absorption of the BHJ, increases with the IE offset, as shown in Fig. 1(c). Second, IQEs are practically independent of donor or acceptor excitation (see IQE spectra in Supplementary Note 2). This implies that the same process governs the quantum efficiency, even though charge generation is expected to occur via electron transfer when exciting the donor, and hole transfer upon acceptor excitation, the former determined by the EA offset, the latter by the IE offset. In other words, the IE offset determines the IQE even when selectively exciting the donor, and despite large EA offsets in all systems. Third, the device fill factor, a measure of the field dependence of charge generation, increases with the IE offset as well, suggesting that in low offset systems, charge generation exhibits a stronger dependence on the external bias (see Supplementary Note 2).

To reveal the underlying reasons, we monitored the dynamics of excited states by transient absorption spectroscopy. Fig. 2 shows transient absorption (TA) spectra of selected thin film blends after selective donor or acceptor photoexcitation. Further TA data can be found in Supplementary Note 3. Clearly, donor and acceptor photoexcitation initially generate different excited states: donor excitation leads to photobleaching of the donor's ground state absorption (PB_D in Fig. 2), as well as to photo-induced absorption (PA). Excitation of the acceptor, on the other hand, initially bleaches the ground state absorption of the acceptor only (PB_A in Fig. 2), associated with PA features resembling the excitonic signatures of neat acceptor films. While donor and acceptor photoexcitation initially generate different excited states (see red lines for the initial TA spectra in Fig. 2), virtually similar TA spectra are observed after 300 ps (see blue lines in Fig. 2). We attribute this to charge generation, however not in the case of DR3:IEICO and PBDB-T-2F:IEICO, which we discuss below. To compare the excited state evolution after donor and acceptor excitation, we integrated the ground state bleach over the entire PB region (shaded areas in Fig. 2). Quantitatively, the relative photobleach area (indicating the overall excitation density) after charge generation is completed (probed at 300 ps) compared to the initial photobleach area (probed at 0.4 ps), does not depend on donor or acceptor excitation, but it increases with the IE offset (Fig. 2(i)).

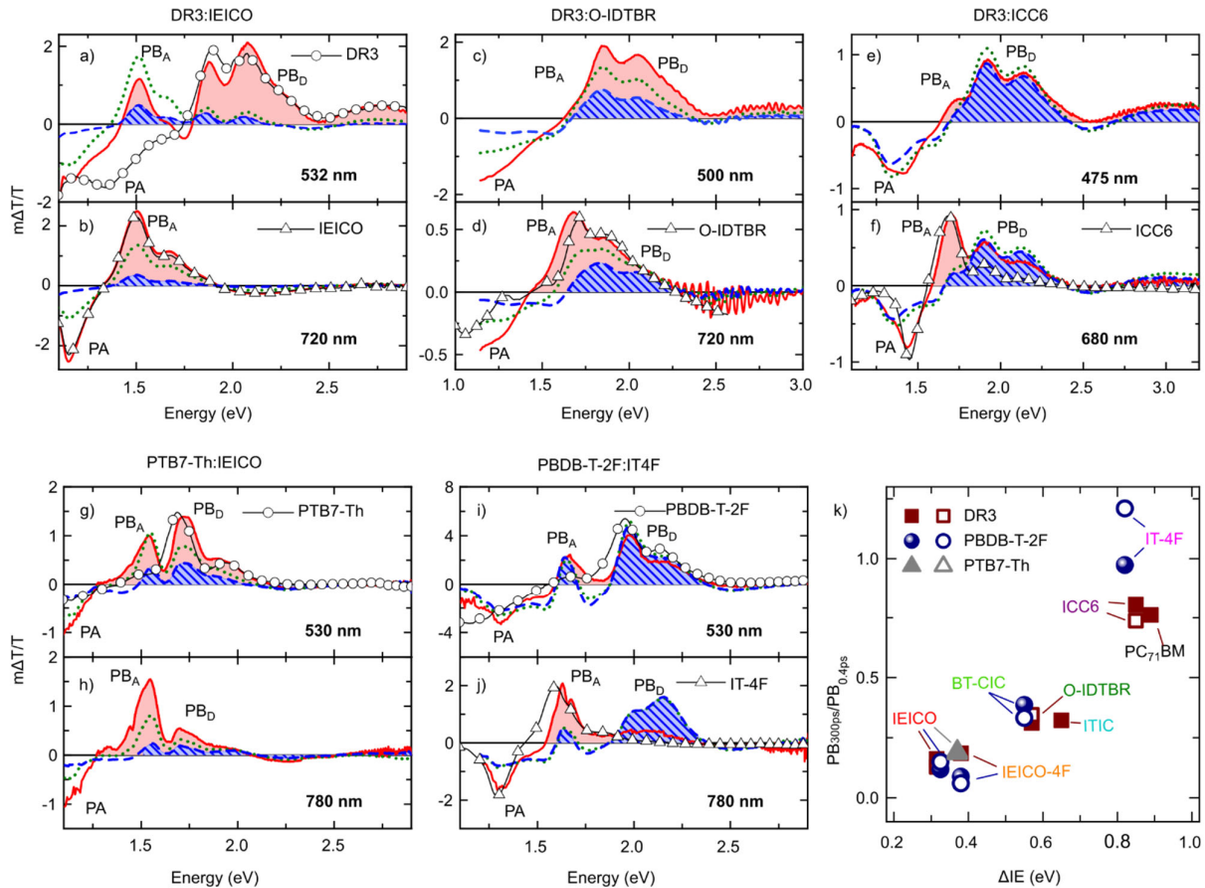


Fig. 2: The charge carrier generation yield is independent of donor or acceptor excitation, but it increases with the IE offset. (a- j) Picosecond-nanosecond transient absorption spectra of selected neat donors (\circ), neat acceptors (\triangle) and blends after 0.4 ps (red solid line), 30 ps (green dotted line) and 300 ps (blue dashed line): (a-b) DR3:IEICO, (c-d) DR3:O-IDTBR, (e-f) DR3:ICC6, (g-h) PTB7-Th:IEICO, and (i-j) PBDB-T-2F:IT-4F blend films after selective excitation of donor (a, c, e, g, i) and acceptor molecules (b, d, f, h, j) at the wavelength indicated in the figure. (k) Ratio of the integrated photobleach (after 300 ps) to the initial photobleach (0.4 – 0.7 ps) ($PB_{300ps}/PB_{0.4ps}$) of all investigated blends for absorbed fluences of $\sim 1.6\text{--}4 \mu\text{J}/\text{cm}^2$, with donor (open symbols) and acceptor (closed symbols) excitation, as a function of the IE offset (ΔIE). The integrated areas are shaded in the example spectra (a-j): unpatterned red for the initial bleach, and stripped blue for the bleach at 300 ps.

DR3:IEICO and PBDB-T-2F:IEICO blends are, however, very different. As shown in Fig. 2(a), within 30 ps after donor photoexcitation (see green line), the initially convoluted exciton-induced absorption signatures of donor and acceptor have evolved into exciton-induced absorption of the acceptor, while only marginal charge generation is observed. Besides, acceptor excitons created by direct photoexcitation of the acceptor subsequently decay to the ground state without undergoing charge transfer. We conclude that almost exclusively energy transfer (FRET) from the donor to the acceptor IEICO takes place, while hole transfer to the donor is inefficient. This is supported by

steady-state PL quenching measurements, which show poor PL quenching upon acceptor excitation, and importantly, acceptor PL upon donor photoexcitation (Supplementary Note 4), supporting FRET occurs in these systems. We note that PL quenching efficiencies in the other blends are larger. This low exciton quenching efficiency in DR3:IEICO and PBDB-T-2F:IEICO blends cannot be attributed to morphological differences between the blends. In fact, we have evidence that all donor-acceptor blends are intimately mixed from transmission electron microscopy (TEM) combined with electron energy-loss spectroscopy (EELS) and scanning transmission electron microscopy (STEM) (Supplementary Note 5), showing no indication of component demixing that could satisfactorily explain the inefficient exciton quenching. Secondly, in more intermixed as-cast DR3:IEICO blends, obtained without solvent vapor annealing, even lower PL quenching efficiencies were observed upon acceptor excitation and enhanced energy transfer upon donor excitation (Supplementary Figure 24).

The large overlap of donor photoluminescence and acceptor absorption, shown in Fig. 3(a) and (b) triggers efficient energy transfer in all donor:NFA systems. The FRET radii^{43,44} (Supplementary Note 6) are larger for the donor-NFA pairs than for the respective donor-donor pairs (see Fig. 3(c)). This implies that when approaching the donor-acceptor interface, an exciton in the donor is more likely to transfer to an acceptor than to reach the interface to undergo electron transfer. More precisely, for donor excitons at a distance of 1 nm from the interface, the calculated FRET rates are on the order of 10^{13} s^{-1} in all donor:NFA systems (Supplementary Note 6). This implies, FRET competes with donor-acceptor electron transfer at shorter distances, as suggested very recently.³⁴

Indeed, FRET from DR3 to NFA could clearly be observed in solution (Supplementary Figure 14). In films, however, it is instantaneously followed by hole transfer from the acceptor to the donor. We confirmed ultrafast hole transfer in diluted DR3:NFA (95:5) blends upon selective excitation of the acceptor, which resulted in a rise of the donor signal within the response time of our TA setup of a few hundred femtoseconds (Supplementary Figure 15) in line with a recent study.³⁶

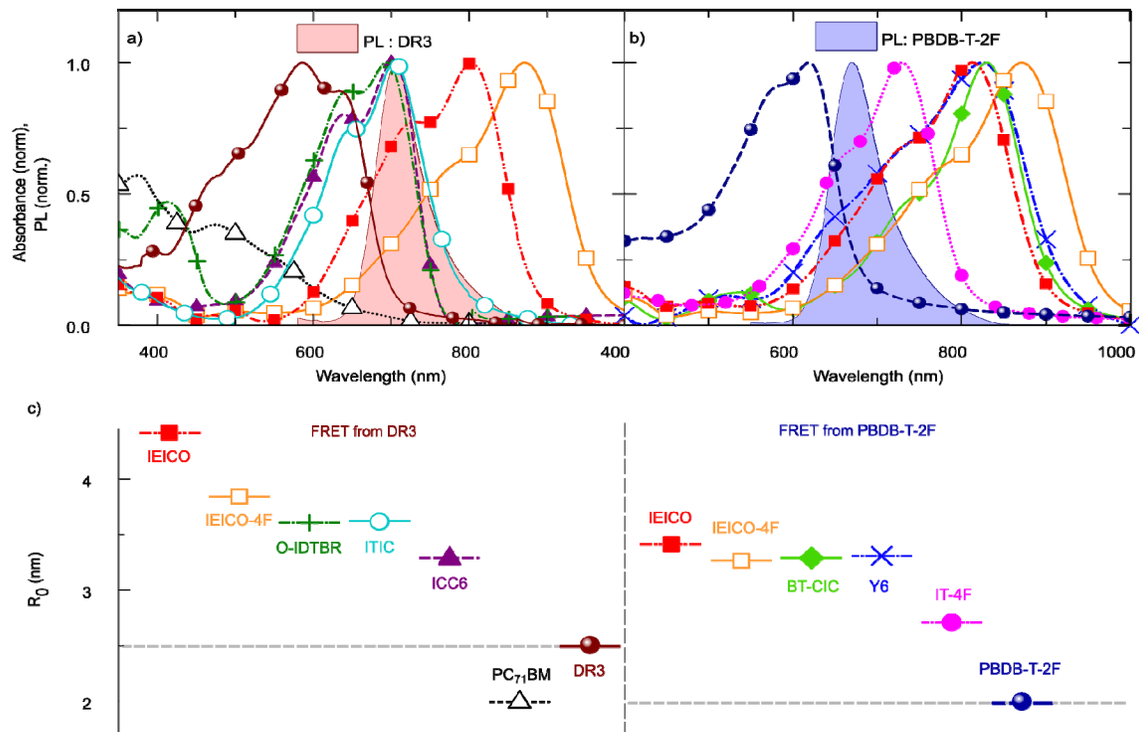


Fig. 3: Spectral overlap of donor photoluminescence and acceptor absorption facilitates ultrafast energy transfer. Normalized UV-Vis absorption spectra of (a) acceptors (continuous lines) used with DR3 as donor (dashed line) and (b) acceptors used with PBDB-T-2F as donor and photoluminescence spectra of DR3 and PBDB-T-2F (filled curves) as indicated by the legends. c) FRET radii for DR3- and PBDB-T-2F-based blends as well as within the donor phase.

As a result of FRET, charge transfer in blends is largely hole transfer from the acceptor to the donor, independent of donor or acceptor photoexcitation. This explains why donor and acceptor photoexcitation yield similar IQEs. Furthermore, it implies that the IQE depends on the driving force of the hole transfer reaction, governed by the difference between the singlet excited state energy of the acceptor and the blend's CT state.

The conversion efficiency of the acceptor exciton to free charges therefore controls the IQE. We identify the efficiency limit from the amount of acceptor exciton quenching efficiency determined by time-resolved photoluminescence (TRPL) experiments (Fig. 4(a-d)). The efficiency of exciton quenching closely follows the IQE, implying that apart from the acceptor singlet exciton to CT state conversion, CT to free charge separation and carrier extraction are efficient. We emphasize that in NFA-based blends TRPL experiments can provide an immediate measure of the free charge generation efficiency and thus device IQE, unlike in fullerene-based systems, in which ultrafast and efficient exciton quenching does not necessarily indicate high free carrier yields and IQE due to the occurrence of other loss channels such as geminate charge recombination. Finally yet importantly,

we stress that the correlation of PL quenching from TRPL and IQE cannot be explained by donor-acceptor demixing, as morphology does not follow energetic offsets and a coincidental correlation across such a large set of different donor-acceptor blends appears highly unlikely (see also STEM and EELS maps in Supplementary Note 5).

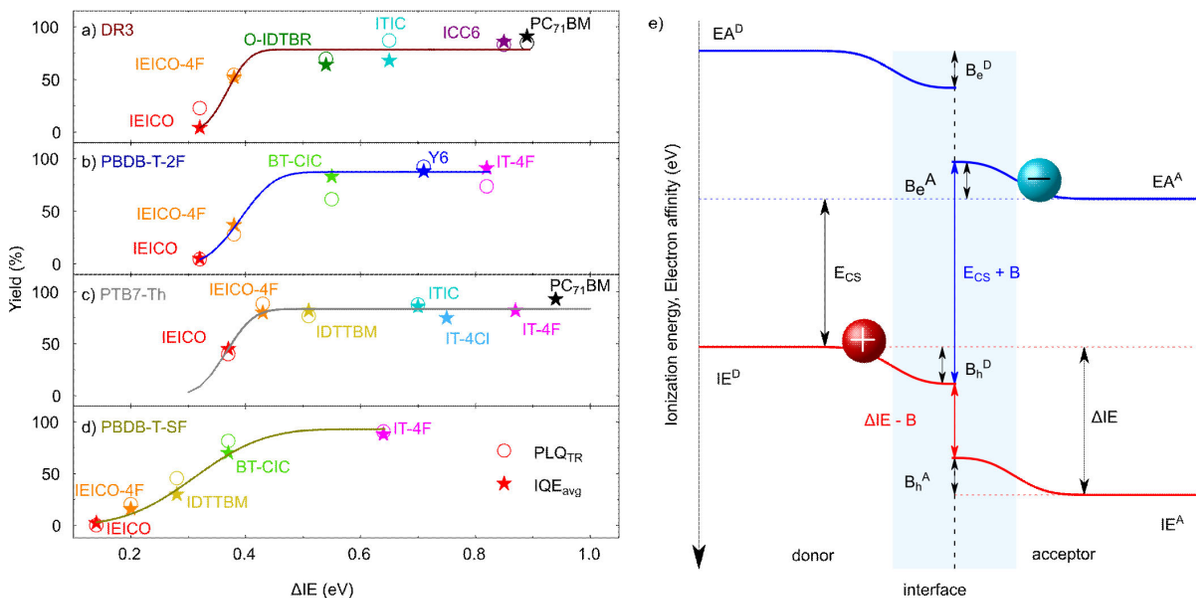


Fig. 4: The IQE follows the exciton quenching yield controlled by the IE offset and energy level bending at the D/A interface. Average internal quantum efficiency (IQE_{avg}, ★) and exciton lifetime quenching $1 - \tau_{blend}/\tau_{pristine}$ (PLQ_{TR}, ○) as a function of ionization energy offset (ΔIE) for (a) DR3-, (b) PBDB-T-2F-, (c) PTB7-Th- and (d) PBDB-T-SF-based blends. The dependence corresponds to the fraction of the interface, which exhibits an energy bending lower than ΔIE , assuming a Gaussian distribution of average energy level bending B and a standard deviation σ , we obtain $IQE = \frac{IQE_{max}}{2} (1 + \text{erfc} \frac{B - \Delta IE}{\sigma})$. The fits to this function (solid lines in (a-d)) yield $B = 0.37$ eV and $\sigma = 0.04$ eV for DR3, 0.37 eV and 0.05 eV for PTB7-Th, 0.39 eV and 0.06 eV for PBDB-T-2F and 0.31 eV and 0.13 eV for PBDB-T-SF. (e) Schematic of the energy level bending at the donor-acceptor heterojunction due to the electrostatic interaction of charges with quadrupole moments of surrounding molecules.

Since the transition from the acceptor singlet exciton to the interfacial charge transfer state is a hole transfer from the acceptor to the donor, it comes as little surprise that it depends on the IE offset. It is, however, surprising and unclear why a *system-independent* offset as large as 0.3 - 0.4 eV is required to obtain a 50% charge transfer yield and why the efficiency increase is less steep than that of a typical thermally-activated process.

To explain both observations, we depict an energy level diagram of a solar cell in Fig. 4(e). Here, the key difference to common diagrams is the bending of the ionization energy (IE) and electron affinity (EA) at the interface. This energy level bending is due to the gradual change in the electrostatic potential created by the quadrupole moments of the NFA and the donor polymer.⁴⁵⁻⁵¹ The energy

level is bent in the direction that corresponds to the (negative) sign of the in-plane quadrupole moment of the acceptor-donor-acceptor (A-D-A) NFA molecule. Because of energy level bending, the CT state energy increases, and the IE difference narrows by approximately the same amount at the interface. The energy level bending has no impact on the diffusion of excitons towards the interface, as both EA and IE in a material are bent in the same direction, leaving the energy gap within the acceptor (or donor) unchanged.

Energy level bending has a clear impact on the energetics of two-particle interactions of states involved in charge generation and separation: the excited state of the acceptor (A^*), the charge transfer state (CT), and the charge separated state (CS). For the acceptor exciton, $E^{A^*} = EA^A - IE^A - E_{\text{Coulomb}}^{A^*}$, where $E_{\text{Coulomb}}^{A^*}$ is the Coulomb interaction energy of the electron and hole, in the solid state. Different signs of energy level bending on the donor and acceptor sides increase the energy of the charge transfer state to $E^{\text{CT}} = EA_{\text{interface}}^A - IE_{\text{interface}}^D - E_{\text{Coulomb}}^{\text{CT}} = EA^A - IE^D + B - E_{\text{Coulomb}}^{\text{CT}}$. Here $E_{\text{Coulomb}}^{\text{CT}}$ is the Coulomb binding of a CT state at a perfectly flat interface (that is, without bending), and $B = B_e^A + B_h^D$ is the bias potential, as depicted in Fig. 4(e). Finally, the CS state is not affected by the energy level bending, since both electron and hole are apart from the interface, $E^{\text{CS}} = EA^A - IE^D$.

Next, we discuss the dissociation energy of the CT state, that is, the CT to CS transition, $\Delta E_{\text{CT} \rightarrow \text{CS}} = E^{\text{CS}} - E^{\text{CT}} = E_{\text{Coulomb}}^{\text{CT}} - B$. Clearly, the interfacial bias reduces the dissociation energy of the charge transfer state. In fact, the optimal condition is achieved for $B = E_{\text{Coulomb}}^{\text{CT}} \sim 0.4$ eV, which then provides barrier-less dissociation, as demonstrated very recently for Y6.²⁹ Interestingly, most of the reported blends provide sufficient bias to split the charge transfer state, which explains their high efficiency.

We now turn to the excited to charge transfer state transition. The driving force for this reaction, $\Delta E_{A^* \rightarrow \text{CT}} = E^{A^*} - E^{\text{CT}} = \Delta IE - B - (E_{\text{Coulomb}}^{A^*} - E_{\text{Coulomb}}^{\text{CT}})$, has three contributions. First, the offset between the ionization energies of the donor and the acceptor, $\Delta IE = IE^D - IE^A$. Second, the *negative* contribution of the bias potential, since the CT state energy is *increased* by the energy level bending (see Fig. 4 (e)). Third, the difference between the hole-electron binding energies of a Frenkel-type exciton and a charge transfer state. It might seem that the last term provides a large negative contribution to the driving force, since it takes a substantial amount of energy to pull an electron and hole apart from each other during an excited to charge transfer state transition, in a molecular dimer. However, polarizable force field calculations demonstrate that, *in a solid state*, strong dielectric stabilization of two charges in a CT state compensates Coulomb binding of a Frenkel exciton, that is, $E_{\text{Coulomb}}^{A^*} - E_{\text{Coulomb}}^{\text{CT}} \sim 0$. As a result, spontaneous dissociation of an excited state

has recently been observed experimentally, even in neat NFA films, that is, in the absence of an electron donor (hole acceptor).⁵²

The consequence is that $\Delta E_{A^* \rightarrow CT}$ is reduced to $\Delta IE - B$, which can be either positive or negative, depending on ΔIE . This is experimentally confirmed by electroluminescence (EL) spectroscopy of NFA-based solar cells: BHJs with large ΔIE , namely DR3:PCBM, DR3:ICC6, DR3:ITIC, PBDB-T-2F:IT-4F, exhibit a clear CT state EL emission observed at lower energy than the neat acceptor's EL, while no such CT state emission is observed in BHJs with low offsets (see Supplementary Note 7). We hypothesize that excitons reaching an interface that exhibits a negative $\Delta E_{A^* \rightarrow CT}$ have the chance to diffuse back to the bulk of the acceptor phase. Hence, the exciton lifetime as measured by TRPL gradually becomes similar to that of a neat acceptor film when ΔIE decreases (Fig. 4 (a), (b), (c)).

To further understand the gradual increase of the IQE with the IE offset (ΔIE), we note that *rough* interfaces provide a broad spread of available interfacial biases. Assuming a Gaussian distribution of B and integrating the fraction of excited to CT state transitions for which $\Delta IE - B$ is positive, we obtain $\text{IQE} = \frac{\text{IQE}_{\text{max}}}{2} (1 + \text{erfc} \frac{B - \Delta IE}{\sigma})$, where B is now the mean value of the bias potential. The fit to this simple expression describes the experimentally observed IQE vs. ΔIE dependence very well across all investigated systems, as shown in Fig. 4 (a-d). Remarkably, the fits provide similar average energy level bending B for DR3, PTB7-Th, and PBDB-T-2F as donors, precisely $B \sim 380 \pm 10$ meV, standard deviation of the bending distribution $\sigma \sim 50 \pm 10$ meV and $\text{IQE}_{\text{max}} \sim 80 \pm 4$ %. We note that for the PBDB-T-SF:NFA systems a smaller value of $B \sim 310$ meV with a broader distribution ($\sigma = 130$ meV) is observed. However, as for the other donors, the same minimum ionization energy offset of 0.5 eV is required to efficiently split CT states, that is, to compensate the energy level bending of ~ 0.5 eV. Within our model, the broader distribution of the bias potential B implies an (electrostatically) less ordered donor-acceptor interface.

The DR3-based systems exhibit a small additional increase in IQE with increasing IE offset, beyond the steep initial rise. This is most likely caused by incomplete charge separation, as indicated by the IE-offset dependent fraction of geminate recombination (Supplementary Note 3.4) possibly due to a larger CT state binding energy with DR3 than with conjugated donor polymers, related to lesser hole delocalization.

To summarize, the interfacial bias potential and associated interfacial energy level bending plays a dual role: it facilitates the dissociation of charge transfer states, but at the same time it reduces the driving force for the exciton to CT state conversion. For barrier-less CT state dissociation biases of at least 0.4 eV, that is the CT state Coulomb binding energy, are required.

We demonstrated that ultrafast energy transfer from either small molecule or polymer donors to low-bandgap acceptors in NFA-based solar cells competes with donor to acceptor electron transfer. Energy transfer funnels excitons from donor to acceptor, renders the electron affinity offset virtually unimportant, and leads to a charge generation efficiency exclusively dependent on the ionization energy offset. Furthermore, for small ionization energy offsets, an energy barrier for the acceptor exciton to charge transfer state transition exists. This barrier is due to the interfacial energy level bending by the electrostatic fields created by the NFA molecular quadrupole moments. As a result, IE offsets larger than the charge transfer state Coulomb binding energy are required for barrier-less hole transfer from the acceptor to the donor. In fact, IE offsets of 0.5 eV are needed to ensure close-to-unity charge generation efficiency. This sets an intrinsic limit to the minimum energy losses that can be tolerated in NFA-based systems without sacrificing the charge separation efficiency, in other words, ensuring close-to-unity quantum efficiency. It also pinpoints additional system parameters that can be used to further improve the device efficiency. Specifically, the largest fraction of losses that limit the V_{oc} is related to the electron-hole Coulomb binding of a CT state and the disorder of the interfacial electrostatic bias potential. Since Coulomb binding of the CT state in NFAs is of the order of 0.4 eV, and close-to-unity charge generation efficiency is achieved for 0.5 eV, we can, in principle, further reduce energetic losses by around 0.1 eV, when decreasing the disorder of the electrostatic bias. Further improvements can be achieved by reducing non-geminate (non-radiative) recombination of charges and charge transfer states. Additional improvements will require conceptually new designs that either decrease the Coulomb binding energy of the CT state or generate charges without passing through a CT state intermediate, for example, by tunneling directly from the excited to the CS state. Apart from the energy losses, the pragmatic way to improve overall OSC efficiency is to increase the absorption coefficient of the donor and acceptor and therefore push the EQE towards unity, by avoiding carrier recombination losses typical for thicker photoactive layers.

Author contributions

S.K. performed the transient spectroscopy experiments, data analysis, and prepared the first draft of the manuscript. J.G. conceived the study, analyzed the transient and steady-state spectroscopic experiments, carried out the FRET calculations, and contributed to the narrative of the manuscript. Y.F. performed the DR3, PCE10, and PM6-based solar cell device preparation and characterization, and the ellipsometry measurements and analysis. N.C. prepared and characterized PCE13-based devices and prepared thin film blends for TRPL experiments. C.S.D.C. prepared samples and characterized the steady-state absorption of thin films, PLQY, and PL quenching in DR3, PCE10, and PM6-based blends *via* TRPL. G.H. performed the UPS and LE-IPES experiments and data analysis. J.I.K. provided steady-state absorption, PL, and TRPL data of PCE13-based blends. A.M.

calculated the solid-state IEs, EAs, and bias potentials. W.L. performed quantum-chemical calculations of donor-acceptor dimers. A.H.B. and T.A.D.P. determined the CT state energies from EL spectra. R.-Z.L. prepared DR3-based devices and characterized their performance. W.Z. synthesized the O-IDTBR acceptor. Y.L. prepared thin films for steady-state characterization. Da.A. performed thin film imaging and EELS analysis. E.A. contributed to the development and maintenance of the ultrafast laser spectroscopy setups. S.H.K.P determined the IQE of PCE10-based devices. A.S. determined the IE and EA of IEICO-4F and IT-4F by UPS and LE-IPES, respectively. P.M.B. supervised the device preparation of DR3-based solar cells and synthesis of IDTTBM used in this work. S.D.W. oversaw the UPS/LE-IPES experiments and data analysis. I.M. supervised the synthesis of O-IDTBR used in this work. T.A. supervised the PBDB-T-2F donor device preparation and characterization. D.B. supervised the PTB7-Th donor device preparation and characterization. De.A. supervised the computational part of the work, conceived the model describing the energy level bending, and contributed to the narrative of the manuscript. F.L. conceived the study and supervised the steady-state and transient optical spectroscopy experiments and data analysis as well as the PBDB-T-SF donor device preparation and characterization and contributed to the narrative of the manuscript. All authors contributed to the revision of the final version of the manuscript.

Competing interests

The authors declare no competing interest.

Data availability

The datasets generated and analyzed during this study and presented in this Article are available online at <https://doi.org/10.6084/m9.figshare.c.5110076.v1> .

Code availability

The codes or algorithms used to analyze the data reported in this study are available from the corresponding authors upon reasonable request.

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Methods

Materials

PBDB-T-2F, IT-4F, Y6, IEICO-4F were purchased from Solarmer Materials Inc. DR3TBDTT (DR3), PTB7-Th, ITIC, ICC6 (IDIC), BT-CIC, IT-4Cl, IEICO and Phen-NaDPO were purchased from 1-Material Inc. IDTTBM⁵³ and O-IDTBR⁵⁴ were synthesized at KAUST. PC₇₁BM was obtained from Solenne BV, PEDOT:PSS from Heraeus and PBDB-T-SF and PFN-Br from Organtec Ltd. Chlorobenzene anhydrous 99.8%, carbon disulfide anhydrous ≥99% and 1,8-diiiodooctane 98% were purchased from Sigma-Aldrich, chloroform from VWR, 1-chloronaphtalene from Fluka and dimethydisulfide 99% from Alfa Aesar. All materials were used as received.

Full material names: IT-4Cl: 3,9-bis(2-methylene-((3-(1,1-dicyanomethylene)-chloro)-indanone))-5,5,11,11-tetrakis(4-hexylphenyl)-dithieno[2,3-d:2',3'-d']-s-indaceno[1,2-b:5,6-b']dithiophene, IT-4F: 3,9-bis(2-methylene-((3-(1,1-dicyanomethylene)-6,7-difluoro)-indanone))-5,5,11,11-tetrakis(4-hexylphenyl)-dithieno[2,3-d:2',3'-d']-s-indaceno[1,2-b:5,6-b']dithiophene, ITIC: 3,9-bis(2-methylene-(3-(1,1-dicyanomethylene)-indanone)-5,5,11,11-tetrakis(4-hexylphenyl)-dithieno[2,3-d:2',3'-d']-s-indaceno[1,2-b:5,6-b']dithiophene, Y6: (2,20 -((2Z,20 Z)-((12,13-bis(2-ethylhexyl)-3,9-diundecyl-12,13-dihydro-[1,2,5] thiadiazolo[3,4-e] thieno[2,30 ':4',50] thieno[20 ,30 :4,5]pyrrolo[3,2-g] thieno[20 ,30 :4,5] thieno[3,2-b]indole-2,10-diyl)bis(methanylylidene))bis(5,6-difluoro-3-oxo-2,3-dihydro-1H-indene-2,1-diylidene))dimalononitrile). ICC6: indacenodithiophene end capped with 1,1-dicyanomethylene-3-indanone. IEICO-4F: 2,2'-[[4,4,9,9-tetrakis(4-hexylphenyl)-4,9-dihydro-s-indaceno[1,2-b:5,6-b']dithiophene-2,7-diyl]bis[[4-((2-ethylhexyl)oxy)-5,2-thiophenediyl]methylidyne(5,6-difluoro-3-oxo-1H-indene-2,1(3H)-diylidene)]]bis[propanedinitrile], O-IDTBR: 5,5'-[(4,9-dihydro-4,4,9,9-tetraoctyl-s-indaceno[1,2-b:5,6-b']dithiophene-2,7-diyl)bis(2,1,3-benzothiadiazole-7,4-diylmethylidyne)]bis[3-ethyl-2-thioxo-4-Thiazolidinone, BT-CIC: 4,4,10,10-tetrakis(4-hexylphenyl)-5,11-(2-ethylhexyloxy)-4,10-dihydrodithienyl[1,2-b:4,5b']benzodithiophene-2,8-diyl)bis(2-(3-oxo-2,3-dihydroinden-5,6-dichloro-1-ylidene)malononitrile), IEICO: 2,2'-((2Z,2'Z)-((5,5'-(4,4,9,9-tetrakis(4-hexylphenyl)-4,9-dihydro-s-indaceno[1,2-b:5,6-b']dithiophene-2,7-diyl)bis(4-((2-ethylhexyl)-oxy)thiophene-5,2-diyl))bis(methanylylidene))bis(3-oxo-2,3-dihydro-1H-indene-2,1-diylidene))dimalononitrile, IDTTBM: indacenodithienothiophene 2-(benzo[c][1,2,5]thiadiazol-4-ylmethylene)malononitrile, PC₇₁BM: [6,6]-Phenyl-C71-butyric acid methyl ester. Phen-NaDPO: ((2-(1,10-phenanthroline-3-yl)naphth-6-yl)diphenylphosphine oxide), PEDOT:PSS: poly(3,4-

ethylenedioxythiophene):poly(styrenesulfonate), PFN-Br: Poly(9,9-bis(3'-(N,N-dimethyl)-N-ethylammonium-propyl-2,7-fluorene)-alt-2,7-(9,9-dioctylfluorene))dibromide. PBDB-T-2F: Poly[(2,6-(4,8-bis(5-(2-ethylhexyl-3-fluoro)thiophen-2-yl)-benzo[1,2-b:4,5-b']dithiophene))-alt-(5,5-(1',3'-di-2-thienyl-5',7'-bis(2-ethylhexyl)benzo[1',2'-c:4',5'-c']dithiophene-4,8-dione)], PBDB-T-SF: Poly[(2,6-(4,8-bis(5-(2-ethylhexylthio)-4-fluorothiophen-2-yl)-benzo[1,2-b:4,5-b']dithiophene))-alt-(5,5-(1',3'-di-2-thienyl-5',7'-bis(2-ethylhexyl)benzo[1',2'-c:4',5'-c']dithiophene-4,8-dione)], PTB7-Th: Poly([2,6'-4,8-di(5-ethylhexylthienyl)benzo[1,2-b;3,3-b]dithiophene]¹), DR3: (5Z,5'E)-5,5'-(((4,8-bis(5-(2-ethylhexyl)thiophen-2-yl)benzo[1,2-b:4,5-b']dithiophene-2,6-diyl)bis(3,3"-dioctyl-[2,2':5',2"-terthiophene]-5",5-diyl))bis(methaneylylidene))bis(3-ethyl-2-thioxothiazolidin-4-one).

Thin-film and device preparation

Organic photovoltaic (OPV) cells were fabricated using indium-tin-oxide (ITO)-coated glass substrate with a sheet resistance of 10 Ω .sq.⁻¹. Substrates were cleaned by sequential ultrasonication in dilute Extran 300 detergent solution, deionized water, acetone, and isopropyl alcohol for 10 min each. The substrates were then subjected to a UV-ozone treatment for 10 min. For normal architecture solar cells, a thin layer (\approx 30 nm) of PEDOT:PSS was spin-coated onto the UV-treated substrates and then dried on a heating plate at 150 °C for 10 min. For inverted solar cells, zinc oxide (ZnO) sol-gel was spin-coated on cleaned glass/ITO substrates with a spin speed of 4000 rpm for 30 seconds, yielding a thickness of around 20 nm. The as-cast films were then thermally annealed in air at 150 °C for 10 min and allowed to cool down to room temperature. Active layers were obtained by dissolving donor and acceptor in organic solvents (chloroform or chlorobenzene) inside the glovebox. The as-prepared solutions were stirred for at least 1h at 40-50 °C before being cast on the substrates. The active layers were spin-cast from the solutions at 50 °C for 45s, using a programmable spin-coater from Specialty Coating Systems (Model G3P-8). The active layers were then annealed (solvent vapor annealed (SVA) for all-small molecule cells and thermally annealed for polymer-based cells). For normal architecture cells, a layer of 5 nm of Phen-NaDPO (or PFN-Br for polymer-based cells) as the electron-transport layer (ETL) was spun from methanol solution (0.5 mg.mL⁻¹) on top of the BHJ layer. Next, the samples were placed in a thermal evaporator for evaporation of a 100 nm-thick layer of aluminum evaporated at 5 Å.s⁻¹ at a pressure of less than 2×10^{-6} Torr. For inverted architecture devices, 7 nm molybdenum(VI) oxide MoO₃ (Alfa) and 100 nm Ag (Kurt Lesker) were evaporated. The effective area of the tested solar cells is 0.1 cm². This effective area was determined

from the layout of ITO substrate and top contact mask. Details of active-layer preparation and optimization are given in Supplementary Note 2.

Device characterization

J - V characteristics of solar cells were measured in a nitrogen-filled glove box. Solar cells were illuminated with light from an Oriel Sol3A Class AAA solar simulator calibrated to 1 sun, AM1.5G, with a KG-5 silicon reference cell certified by Newport. A Keithley 2400 source meter was used to measure the J - V curves in the reverse direction, that is, from positive to negative bias with a dwell time of 10 ms. For each J - V curve 100 data points were collected. The solar cells' external quantum efficiency (EQE) was measured with a commercial setup (PV Measurement Inc.) at zero bias. Solar cells were illuminated with light supplied from a Xenon arc lamp in combination with a dual-grating monochromator. For each wavelength, the number of incident photons was calculated using a calibrated silicon photodiode. Calibration was performed by The National Institute of Standards and Technology (NIST). The internal quantum efficiency (IQE) of OPV devices was calculated using: $\text{IQE}(\%) = \text{EQE}(\%) / (100\% - \text{Reflectance}(\%) - \text{Parasitic Absorption}(\%))$. The reflectance spectra were measured in an integrating sphere using the same EQE setup, while the parasitic absorption spectra were obtained by transfer matrix modelling of the device stack.

Ultraviolet photoelectron (UPS) and low-energy inverse photoelectron (IPES) spectroscopy.

The substrate for film deposition was Au(100nm)/Cr(10nm)/Si(n-type), with the Au and Cr evaporated and sputtered within an Angstrom evaporation/sputter tool, respectively. Solutions of 2-5 mg·ml⁻¹ in chloroform were prepared and stirred overnight in a glove box and then spin-coated (5000 rpm for 1 minute) onto an Ar⁺ ion sputtered Au substrate and then transferred to UHV for analysis. The organic small molecule/polymer film thickness was 10-20 nm.

An in-house built UHV LE-IPES set-up was used following closely a design outlined by Yoshida⁵⁵ which was also developed from a prior design.⁵⁶ This operates in the Bremsstrahlung Isochromatic Mode (BIS). In brief, an incident electron beam (dispersion of 0.25-0.5 eV) is directed at 0° with respect to the sample plane, and the electron energy is swept slowly. The outgoing light from the surface was collected by placing an external focusing lens and an internal vacuum collimating lens with a detector and all further stray light shielded. This was conducted with a low energy electron source (Staib) operating in the range of 20-30 eV energy (BaO cathode Heatwaves), with a retarding bias of +20V applied to the sample in good electrical contact. As stated, the photons were collected outside of vacuum by a solid-state PMT detector (Hamamatsu R585) (0.5-1h, integration time ~10-

20s, 200 data points), facing the sample within a narrow wavelength window using a bandpass filter of 280 nm (Semrock) (10 nm width). This measurement was performed consecutively without either exposure to Al K α X-ray 1486.6 eV for XPS analysis or re-exposure of the surface to ambient atmosphere in a base pressure of 10⁻⁹ mbar to accurately join with UPS spectra. More information can be found in the Supplementary Note 1.

Optical bandgap from intersection of steady-state absorption and emission spectra

The thin-film UV-Vis absorption and photoluminescence spectra were measured with a Cary 5000 spectrophotometer and a Fluoromax-4 spectrofluorometer from HORIBA Scientific, respectively.

Transmission electron microscopy (TEM)

TEM studies in combination with EELS were performed on a Thermo Fischer (former FEI) Titan 80-300 TEM equipped with an electron monochromator and a Gatan Imaging Filter (GIF) Quantum 966. Details can be found in the Supplementary Information.

Transient absorption spectroscopy (TA)

Transient absorption (TA) pump-probe measurements were carried out with a home-built setup.⁵⁷ Two different configurations of the setup were used for either short delay, namely 100 fs to 8 ns experiments, or long delay, namely 1 ns to 300 μ s delays, as described below.

The output of a titanium:sapphire amplifier (Coherent LEGEND DUO, 4.5 mJ, 3 kHz, 100 fs) was split into three beams (2 mJ, 1 mJ, and 1.5 mJ). Two of them separately pumped two optical parametric amplifiers (OPA) (Light Conversion TOPAS Prime). TOPAS 1 generates tunable pump pulses, while TOPAS 2 generates signal (1300 nm) and idler (2000 nm). A fraction of the output signal of the titanium:sapphire amplifier was focused into a c-cut 3 mm thick sapphire window, thereby generating a white-light supercontinuum from 500 to 1600 nm. For short delay TA measurements, we used the output of TOPAS 1 as pump pulses, while the probe pathway length to the sample was kept constant at approximately 5m between the output of TOPAS 1 and the sample. The pump pathway length was varied between 5.12 and 2.6 m with a broadband retroreflector mounted on an automated mechanical delay stage (Newport linear stage IMS600CCHA controlled by a Newport XPS motion controller), thereby generating delays between pump and probe from -400 ps to 8 ns. Alternatively, for probing TA further in the blue to violet range of light, we used the 1300 nm (signal) output of TOPAS 2 to produce a white-light supercontinuum from 350 to 1100 nm. For the 1 ns to 300 μ s delay (long delay) TA measurement, the same probe white-light supercontinuum was used as for the 100 fs to 8 ns delays. Here the excitation pulse was provided by

an actively Q-switched Nd:YVO₄ laser (InnoLas piccolo AOT) frequency-doubled, providing pulses at 532 nm. The pump laser was triggered by an electronic delay generator (Stanford Research Systems DG535) itself triggered by the transistor-transistor logic (TTL) sync from the Legend DUO, allowing control of the delay between pump and probe with a jitter of roughly 100 ps.

Pump and probe beams were both focused on the sample, which was kept under a dynamic vacuum of $<10^{-5}$ mbar generated by a turbo-molecular pump (Pfeiffer Cube). The transmitted fraction of the white light was guided to a custom-made prism spectrograph (Entwicklungsbüro Stresing), where it was dispersed by a prism onto a 512 pixel NMOS (N-type metal-oxide-semiconductor) linear image sensor (HAMAMATSU G11608-512DA). The probe pulse repetition rate was 3 kHz, while the excitation pulses were mechanically chopped to 1.5 kHz (100 fs to 8 ns delays). Adjacent diode readings corresponding to the transmission of the sample with and without prior excitation were used to calculate $\Delta T/T$. Measurements were averaged over several thousand probes to obtain a reasonable signal-to-noise ratio. The chirp induced by the remaining transmissive optics was corrected with a home-built Matlab code. The delay at which pump and probe arrive simultaneously on the sample (referred as zero time) was determined from the point of maximum positive slope of the TA signal rise for each wavelength.

Time-resolved photoluminescence spectroscopy (TRPL)

For time-resolved PL experiments (TRPL), samples (films on quartz substrates unless stated) were excited with the wavelength-tunable output of an OPO (Radiantis Inspire HF-100), itself pumped by the fundamental of a Ti:Sa fs-oscillator (Spectra Physics MaiTai eHP) and a Modelocked Ti:Sa (Chameleon Ultra I from Coherent) at 820 nm. Typical pulse energies were in the range of several nano-joules (nJ). The samples were kept in a nitrogen-filled custom-made sample chamber and excited at 370 nm in the nitrogen atmosphere. The PL of the samples was collected by an optical telescope (consisting of two plano-convex lenses), focused on the slit of a spectrograph (PI Spectra Pro SP2300) and detected with a Streak Camera (Hamamatsu C10910) system with a temporal resolution of 1.4 ps. The data was acquired in photon counting mode using the Streak Camera software (HPDTA) and exported to Origin Pro 2019 for further analysis. The TRPL decays were analyzed with a sum of exponentials, and when more than one exponential was found, the amplitude weighted average lifetime was calculated.

Förster Resonance Energy Transfer (FRET) radii and rates

Transmittance and reflectance spectra for extinction coefficients were measured with a PerkinElmer LAMBDA 950 UV/Vis/NIR spectrophotometer equipped with a 150 mm InGaAs integrating sphere

and the spectra for PLQYs were measured with a FLS920-s Fluorescence Spectrometer by Edinburgh Instruments equipped with a 120 mm integrating sphere.

Sensitive EQE and electroluminescence (EL) spectroscopy

The high-dynamic range EQE spectra were collected at short-circuit conditions using focused monochromatic illumination from a Xenon arc lamp combined with a monochromator. The light was modulated by an optical chopper at 275 Hz. The device output current was measured as a function of the incident photon energy using a lock-in amplifier (Stanford Instruments SR 830). The lamp intensity was calibrated with Ge and Si photo-diodes.

Electroluminescence (EL) measurements were carried out on devices (active area of 0.1 cm²) biased with DC voltages (Keithley 2420) similar to or lower than V_{OC} at 1 sun illumination. The emission was collected by a collimator inside a nitrogen-filled glovebox and guided to a spectrograph (Princeton Instruments SP-2300) by an optical fiber. The spectrometer was equipped with a cooled Si CCD (Princeton Instruments PIX100BRX) and cooled InGaAs (Princeton Instruments PYR1024) camera. The system was wavelength calibrated with a Ne/Ar calibration light source (Princeton Instruments IntelliCal). The visible and infrared spectra were assembled at the 933 nm where they share the same relative sensitivity.

Computer simulations

Solid state ionization energies were evaluated in a perturbative way, starting from the gas-phase quantum chemical calculations (B3LYP/6-311g(d,p) level as implemented in the GAUSSIAN 09 package⁵⁸) and then taking into account solid-state contribution as a perturbation, where electrostatic and induction energies are the first- and second-order corrections. These contributions to site energies were calculated self-consistently using the Thole model^{59,60} using atomic polarizabilities and distributed multipoles using GDMA program⁶¹ for a cation and a neutral molecule. This approach, in combination with an aperiodic inclusion of charges to a neutral periodic morphology, is available in the VOTCA-CTP package. Further details are given in Supplementary Information.

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