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Open-circuit voltage of organic solar cells: interfacial roughness makes the difference

Carl Poelking¹, Johannes Benduhn¹, Donato Spoltore¹, Martin Schwarze², Steffen Roland⁴, Fortunato Piersimoni⁴, Dieter Neher¹, Karl Leo², Koen Vandewal¹, Denis Andrienko¹

Organic photovoltaics (PV) is an energy-harvesting technology that offers many advantages, such as flexibility, low weight and cost, as well as environmentally benign materials and manufacturing techniques. Despite growth of power conversion efficiencies to around 19 % in the last years, organic PVs still lag behind inorganic PV technologies, mainly due to high losses in open-circuit voltage. Understanding and improving open circuit voltage in organic solar cells is challenging, as it is controlled by the properties of a donor-acceptor interface where the optical excitations are separated into charge carriers. Here, we provide an electrostatic model of a rough donor-acceptor interface and test it experimentally on small molecule PV materials systems. The model provides concise relationships between the open-circuit voltage, photovoltaic gap, charge-transfer state energy, and interfacial morphology. In particular, we show that the electrostatic bias generated across the interface reduces the photovoltaic gap. This negative influence on open-circuit voltage can, however, be circumvented by adjusting the morphology of the donor-acceptor interface.



¹ Max Planck Institute for Polymer Research, Ackermannweg 10, 55128 Mainz, Germany. ² Integrated Center for Applied Physics and Photonic Materials (IAPP), Technische Universität Dresden, Nöthnitzer Str. 61, 01187 Dresden, Germany. ³ Instituut voor Materiaalonderzoek (IMO-IMOMEC), Hasselt University, Diepenbeek, Belgium. ⁴ Institute of Physics and Astronomy, University of Potsdam, Karl-Liebknecht-Str. 24-25, 14476 Potsdam, Germany. ^{Se}email: koen.vandewal@uhasselt.be; denis.andrienko@mpip-mainz.mpg.de

mproving conversion efficiencies between photons and charges, phonons and charges¹, spins and charges², or triplet and singlet excitations³ is a central target of contemporary energy research. For example, the understanding of the conversion pathways in organic solar cells led to a more than tenfold increase in power conversion efficiency (PCE)^{4–15}, with current non-fullerene-acceptor-based devices having nearly 19 % PCE ^{16,17}.

Organic solar cells are actively studied due to their low cost, environmental friendliness, and ease of processing. In these cells, strongly bound excitons dissociate into charge-transfer (CT) states at donor-acceptor interfaces, which subsequently dissociate into free charge carriers, detected as photocurrent^{5,18}. Early optimization strategies targeted materials with small optical and large photovoltaic gaps, defined as the difference between the ionization energy (IE) of the donor and electron affinity (EA) of the acceptor, $\Gamma = IE_D - EA_A^{19}$. Small optical gaps helped to harvest a wider region of the solar spectrum, whereas larger photovoltaic gaps often correlated with larger open-circuit voltages, Voc. It was, however, soon realized that the abstraction of the density of states of a heterojunction to a single photovoltaic gap is an oversimplification. Accounting for the broadening of energy levels motivated additional optimization strategies²⁰⁻²³. Furthermore, detailed macroscopic models pointed out the importance of the charge-carrier mobility and non-geminate recombination coefficient, thus providing optimization criteria for the thickness of the active layer in order to achieve high shortcircuit currents and fill factors²⁴⁻²⁷. A whole new era in organic photovoltaic research started with the development of lowbandgap non-fullerene acceptors (NFAs)¹¹⁻¹⁵.

It was soon realized that long-range electrostatic interactions of a charge with surrounding molecular quadrupoles play a decisive role in defining the energetic landscape for electrons, holes and charge-transfer states^{28–31}. The crystal field created by molecular quadrupoles contributes to the ionization energy (IE) and electron affinity (EA) of the organic film, and therefore to the opencircuit voltage of the solar cell²⁸. This field also changes the energy offset between the charge-transfer and charge-separated states, favouring or disfavoring CT state dissociation^{29,31–34}. It is unfortunate that the electrostatic effects, which favor efficient CT state dissociation also lead to a reduction of the photovoltaic gap, energy of the charge-transfer state, $E_{\rm ct}$, and hence smaller $V_{\rm oc}$. As a result, in order to optimize solar cell efficiency, one has to appropriately balance the electrostatic contributions to the energy profiles^{32,33}.

In this paper, we use tailored material systems in order to disentangle the effects of morphology, molecular and device architecture on the microscopic energetics. In the first set of experiments, we study planar heterojunction solar cells. While the acceptor phase is always C₆₀, the donor phase of these cells is a mixture of chemically similar compounds, ZnPc:F₄ZnPc or ZnPc:F^{*}₄ZnPc, isomers of which are shown in Fig. 1. Due to fluorination, the quadrupole component along the normal of the molecular plane, $Q_{20} \equiv Q_{\pi\pi\nu}$ differs significantly for all three compounds. This difference allows us to accurately tune the electrostatic field in the intermixed crystalline phase. As a result, IEs and EAs of such mixtures can be continuously tuned, while the crystalline morphology of the donor phase is preserved^{31,35}. In the second set of experiments, an additional mixed donor:acceptor interlayer is deposited between the pure donor and acceptor layers. This interlayer is used to control the roughness of the donor-acceptor interface. We then show theoretically and prove experimentally that interfacial electrostatics can be used to tune the open-circuit voltage of an organic solar cell-by adjusting the roughness of the donor-acceptor interface. Moreover, we provide a detailed microscopic description of how V_{oc} , Γ , and $E_{\rm ct}$ depend on system composition, interface roughness, and energetic disorder.

Results

Ionization energy and photovoltaic gap. We begin by analyzing the density of states (DOS) of intermixed ZnPc:F₄ZnPc and ZnPc:F₄^{*}ZnPc co-crystals. In the absence of intermolecular interactions, the DOS of the binary mixture would only consist of two peaks corresponding to the ionization energies of ZnPc and F₄ZnPc (or ZnPc and F₄^{*}ZnPc) in the gas phase, IE₀. The composition dependence of the IE and EA is thus given solely by the solid-state contribution $\Delta_h(c)$ for holes and $\Delta_e(c)$ for electrons, IE(c) = IE₀ + $\Delta_h(c)$, EA(c) = EA₀ + $\Delta_e(c)$, where c is the fraction of ZnPc in the mixture.

The microscopic mechanism responsible for this dependence is the long-range superposition of quadrupolar fields that act on the charge carrier^{31,35}. ZnPc carries a net-negative out-of- π -plane quadrupole component, larger in magnitude than the corresponding negative quadrupole moment of F₄^{*}ZnPc, whereas the corresponding quadrupole component of F₄ZnPc is positive. In fact, this can be anticipated from the isopotential maps of the compounds, which are shown in Fig. 1. The long-range character of the charge-quadrupole interaction effectively results in a concentration-weighted average over both contributions, which can be used to tune ionization potential and electron affinity. This microscopic view is indeed supported by the good agreement between ionization energies measured by UPS^{31,35} and simulated using atomistically resolved models, as shown in Fig. 2.

To relate this composition dependence of the energetics of holes in the mixtures to photovoltaic observables, we first consider its effect on the photovoltaic gap $\Gamma(c) = \text{IE}_D(c) + \text{EA}_A = \Gamma_0 + \Delta_{h,D}(c) + \Delta_{e,A}$. Here, $\Gamma_0 = \text{IE}_{0,D} + \text{EA}_{0,A}$ is the sum of the gas-phase IE of the donor and EA of the acceptor. $\Delta_{e,A} \simeq 0.95$ eV for C_{60} denotes the solid-state correction to electron site energies in the acceptor component, as calculated from atomistic models. Note that we define the photovoltaic gap based on the mean of the *low*-energy Gaussian mode within the bimodal density of states of the mixed ZnPc:F₄ZnPc and ZnPc:F₄^{*}ZnPc systems. This low-energy contribution is due to the ZnPc component, and will eventually determine the chemical potential μ_h . The composition dependence of $\Gamma(c)$ is therefore inherited directly from $\Delta_{h,D}(c)$. As a result, $\Gamma(c)$ should display the same linear behavior in *c* as IE_D(*c*).

Electrostatic interfacial bias. To quantify the electrostatic heterogeneity of the donor-acceptor interface, we define an interfacial electrostatic bias for holes and electrons, $B_h = \Delta_{h(A)} - \Delta_{h(A)}$ $c_1\Delta_{h,D_1} - c_2\Delta_{h,D_2}, B_e = c_1\Delta_{e(D_1)} + c_2\Delta_{e(D_2)} - \Delta_{e,A}$, where $\Delta_{h(A)}$ is the solid-state contribution to the ionization energy of the donor molecule in the acceptor mesophase, i.e., when the donor molecule is completely surrounded (perfectly dispersed) by C_{60} . Similarly, $\Delta_{e(D)}$ is this contribution to the electron affinity of the acceptor molecule placed in the donor mesophase. As the donor mesophase is a mixture of two compounds, we weigh the corresponding contributions of D_1 and D_2 by their respective concentrations in the mixture. In other words, if D_1 is ZnPc then $c_1 = c$ and $c_2 = 1 - c$. In an ordered thin film, the solid-state contribution to the ionization energy and, therefore, the interfacial bias, are proportional to the $Q_{\pi-\pi}$ component of the quadrupolar quadripolar tensor^{31,33}. For the columnar molecular arrangement shown in Fig. 2, $Q_{\pi-\pi} \equiv Q_{20}$ is the component of the tensor perpendicular to the conjugated core, as shown in Fig. 1.

 B_h and B_e quantify the electrostatic asymmetry of the interface by comparing the external contributions for holes and electrons on the donor and acceptor side of the interface. They are defined



Fig. 1 Chemical structures and electrostatic surface potentials illustrating their distinct electrostatic architecture. Isopotential surfaces at -0.5 V and +0.5 V are shown in blue and red, respectively. Note that for F_4ZnPc , the material used is a mixture of different isomers, as the fluorine atoms can be in either of the two outer positions. IE₀ and EA₀ are gas-phase ionization energy and electron affinity, respectively, calculated at the B3LYP/6-311+g(d,p) level of theory. Q_{20} is the component of the quadrupole tensor in the direction normal to the molecular plane.



Fig. 2 Crystalline morphologies and corresponding ionization potentials. a Model of a crystalline binary ZnPc: F_4 ZnPc thin film, with species represented by (semi-transparent) and gray molecules, with periodicity in the *xy* plane. Deeper red implies more F_4 ZnPc molecules in a column. **b** Dependence of the ionization energy (IE) on the fraction of ZnPc in the ZnPc: F_4 ZnPc and ZnPc: F_4 ZnPc systems, *c*.



Fig. 3 Interfacial bias potential. Dependence of the interfacial bias potential for electrons (blue curves, filled circles) and holes (red curves, open circles) on the composition of the donor layer, *c*, for the ZnPc: F_4 ZnPc and ZnPc: F_4 ZnPc systems as obtained through simulations. The bias potential has a linear dependence on *c*, inherited from the solid-state contributions Δ_h and Δ_e to electron and hole site energies.

such that a positive interfacial bias pushes charge carriers away from the interface. To evaluate these quantities, we would need to explicitly simulate the donor–acceptor mixture. Instead, we approximate the corresponding solid-state contributions with their bulk values, $\Delta_{h(A)} \approx \Delta_{h,A}$ and $\Delta_{e(D)} \approx \Delta_{e,D}$. This approximation is reasonable at the donor–acceptor interface, where donor (acceptor) molecules have mostly donor (acceptor) nearest neighbors, and where the long-range charge-quadrupole interactions dominate the solid-state contribution.

Since the solid-state contributions to the electron and hole on the donor side are linear in *c*, the interfacial bias also linearly depends on the composition, as shown in Fig. 3. Remarkably, for F_4ZnPc , *B* becomes negative at $c \simeq 0.3$, which is also where the fill factor of the solar cell starts to deteriorate, see Table 1. This can be rationalized in that a negative *B*, i.e., a negative bias for charge push-out, leads to trap formation at donor/acceptor sites that protrudes into the opposite (acceptor/donor) domain. The electrostatic bias of the F_4^*ZnPc system is always positive. This cell indeed performs much better than the F_4ZnPc device.

Charge-transfer states. We now evaluate the energies of interfacial CT states. Generally speaking, the energy of these states can be written as

$$E_{\rm ct}(c) = \Gamma(c) - |\Delta_{eh}| + \chi(c, \rho). \tag{1}$$

Here, $|\Delta_{eh}|$ is an effective binding energy that results from the electron-hole attraction reduced by dielectric solvation^{29,30,33,36}. This binding energy can be assumed to be independent of composition, and amounts to 0.5 eV for the closest CT states in results obtained through simulations^{29,33}. $\chi(c, \rho)$ is the interfacial electrostatic bias, as discussed below.

The ionization energy and electron affinity are impacted by the interfacial morphology at the nanoscale^{29,37}. At a *rough* interface, donor molecules or domains can protrude into the acceptor phase. As a result, their gas-phase energy levels are dressed by the electrostatic field of the acceptor phase. This superposition of electrostatic fields in the donor and acceptor films across an

Table 1 Summary of the measurements for the ternary cells. For $ZnPc:F_4^ZnPc/C_{60}$ and $ZnPc:F_4ZnPc/C_{60}$ as well as the planar junction cell with an interlayer we report short-circuit currents, J_{scr} open-circuit voltages, V_{ocr} fill factors, power conversion efficiencies (PCE), charge-transfer energies E_{ct} and the corresponding fit parameters, as described in the "Methods" section.

| System | Thickness [nm] | J _{sc} [mA/cm ²] | <i>V</i> _{oc} [V] | FF [%] | PCE [%] | E _{ct} [eV] | λ [eV] | f [10 ⁻⁴ (eV) ²] |
|--|----------------|---------------------------------------|----------------------------|--------|---------|----------------------|---------------|---|
| ZnPc/ C ₆₀ | 10/40 | 3.33 | 0.47 | 66.4 | 1.04 | 1.08 | 0.24 | 0.50 |
| ZnPc:F ₄ ZnPc (3:1) / C ₆₀ | 10/40 | 3.17 | 0.48 | 63.3 | 0.97 | 1.13 | 0.34 | 2.54 |
| ZnPc:F ₄ ZnPc (2:1) / C ₆₀ | 10/40 | 3.18 | 0.48 | 65.2 | 1.00 | 1.13 | 0.35 | 2.35 |
| ZnPc:F ₄ ZnPc (1:1) / C ₆₀ | 10/40 | 3.14 | 0.51 | 65.0 | 1.03 | 1.16 | 0.41 | 5.63 |
| ZnPc:F ₄ ZnPc (1:2) / C ₆₀ | 10/40 | 3.09 | 0.53 | 65.3 | 1.06 | 1.20 | 0.50 | 19.4 |
| F ₄ ZnPc / C ₆₀ | 10/40 | 2.90 | 0.60 | 64.8 | 1.13 | 1.41 | 0.34 | 72.4 |
| ZnPc/C ₆₀ | 10/40 | 3.60 | 0.41 | 65.0 | 0.95 | 1.06 | 0.28 | 0.78 |
| ZnPc:F ₄ ZnPc (3:1) / C ₆₀ | 10/40 | 3.78 | 0.49 | 66.7 | 1.24 | 1.11 | 0.31 | 1.15 |
| ZnPc:F ₄ ZnPc (2:1) / C ₆₀ | 10/40 | 3.79 | 0.53 | 67.5 | 1.37 | 1.13 | 0.34 | 1.57 |
| ZnPc:F ₄ ZnPc (1:1) / C ₆₀ | 10/40 | 3.68 | 0.61 | 68.7 | 1.54 | 1.19 | 0.24 | 0.54 |
| ZnPc:F ₄ ZnPc (1:2) / C ₆₀ | 10/40 | 3.44 | 0.67 | 68.7 | 1.28 | 1.28 | 0.30 | 2.42 |
| F ₄ ZnPc / C ₆₀ | 10/40 | 1.78 | 0.89 | 36.9 | 0.59 | 1.49 | 0.18 | 7.89 |
| ZnPc/ ZnPc:C ₆₀ (1:1) / C ₆₀ | 10/2/40 | 3.63 | 0.47 | 65.37 | 1.11 | 1.10 | 0.29 | 1.86 |
| ZnPc/ ZnPc:C ₆₀ (1:1) / C ₆₀ | 10/4/40 | 4.15 | 0.49 | 66.30 | 1.36 | 1.16 | 0.35 | 16.8 |
| ZnPc/ ZnPc:C ₆₀ (1:1) / C ₆₀ | 10/6/40 | 4.62 | 0.52 | 67.02 | 1.60 | 1.17 | 0.33 | 23.9 |
| ZnPc/ ZnPc:C ₆₀ (1:1) / C ₆₀ | 10/8/40 | 5.07 | 0.53 | 67.23 | 1.79 | 1.17 | 0.32 | 31.7 |
| ZnPc/ ZnPc:C ₆₀ (1:1) / C ₆₀ | 10/10/40 | 5.57 | 0.52 | 66.56 | 1.91 | 1.17 | 0.32 | 42.4 |



Fig. 4 Open-circuit voltage and charge-transfer state energy of a planar junction solar cell. a Model of a planar junction solar cell with a rough interface and co-deposited donor materials. Geometrical parameters of a rough interface: period, λ , width, h, and a minimal separation between the donor and acceptor molecules, δ . **b** Dependence of the open-circuit voltage (V_{oc} , open symbols), and charge-transfer state energy (E_{ct} , closed symbols) on the concentration c of ZnPc in the ZnPc:F₄ZnPc and ZnPc:F₄^{*}ZnPc solar cells. The experimentally measured slopes (black lines and symbols) are best reproduced for $\rho = 1$.

intermixed region is triggered by the same long-range mechanism that gives rise to the composition dependence of site energies in the mixed donor phase.

The degree of interface roughness determines to what extent this bias actually impacts the CT energy of interfacial donor-acceptor pairs. In Supplementary Note 2 we quantify the amount by which E_{ct} is raised (B > 0) or lowered (B < 0) by the bias for a rough interface of width h, which is shown in Fig. 4(a),

$$\chi(c,\rho) = \frac{1}{2} \left(B_e(c) + B_h(c) \right) \frac{\rho}{1+\rho}.$$
 (2)

In the limit where the corrugation period λ of the profile significantly exceeds both the depth *h* of the intermixed region, and the interlayer spacing δ , the degree of nanoscale roughness can be captured by a single parameter $\rho = 2h/\delta$.

To validate this relation, the measured composition dependence of $E_{\rm ct}$ is shown in Fig. 4b. Experimental details are provided in the Methods section and Table 1. The slope is best fitted with $\rho = 1$, which corresponds roughly to two intermixed molecular layers. As we will see in the next section, the same roughness is required to fit the dependence of the open-circuit voltage on the composition of the donor. Moreover, we will also show that the interface roughness can be systematically varied in planar junction solar cells with an interlayer between the two phases.

Open-circuit voltage. With Γ and E_{ct} at hand, the compositiondependent open-circuit voltage, $V_{oc}(c) = \mu_e(c) - \mu_h(c)$ is calculated by evaluating the chemical potentials $\mu_e(c)$ and $\mu_h(c)$ of electrons and holes, respectively:

$$eV_{\rm oc}(c) \simeq \Gamma(c) + \alpha k_{\rm B} T \ln[n(c)p(c)] - \beta k_{\rm B} T \hat{\sigma}^2(c), \qquad (3)$$

where $\alpha \approx 1$ and $\beta \approx 1/2$ are parameters with a weak dependence on charge density; $\hat{\sigma} = \sigma/k_{\rm B}T \lesssim 5-6$ is the effective disorder strength of the donor (energetic disorder in the acceptor is negligible in the case of C₆₀), and *n* and *p* are the steady-state number densities (number of charges per number of available sites) of electrons and holes, respectively. The derivation is provided in Supplementary Note 1.

Evaluating *n* and *p* requires the simulation of the full dynamics of charge generation and recombination in a solar cell. Here, however, we are only interested in how the film composition affects V_{oc} . We therefore define n_0 and p_0 as the charge-carrier densities that would follow from an effectively zero electron-hole binding energy of CT states, and assume $n_0 = p_0 = 5 \times 10^{-6}$, as



Fig. 5 Charge-transfer state energy and open-circuit voltage dependence on interfacial roughness. a Co-deposited layer of ZnPc:C₆₀ sandwiched between pristine ZnPc and C₆₀ layers. **b** Variation of the charge-transfer (CT) energy (E_{ct} , blue curve and filled symbols) and open-circuit voltage (V_{oc} , red curve and open symbols) with interface roughness, represented here by the interlayer thickness *h*. Dots are measurements performed on planar heterojunction cells with a ZnPc/ZnPc:C₆₀ architecture. Solid lines are fits to Eqs. (6) and (3), with fit parameters $E_{ct,0} = 0.92 \text{ eV}$, $\chi_0 = 0.27 \text{ eV}$, $\delta = 1.43 \text{ nm}$ and $h_0 = 1.43 \text{ nm}$ and the $E_{ct} - eV_{oc}$ offset $\alpha k_B T \ln(n_0 p_0) = -0.66 \text{ eV}$. These fitted parameters are in very good agreement with the values predicted by simulations, as well as the interface roughnesses extracted from the composition-dependent measurements in Fig. 4.

was found to accurately describe the V_{oc} under AM1.5g conditions of well-functioning systems with B_{h} , $B_e > 0^{29}$. The steady-state charge-carrier densities can then be related to the charge push-out forces via a law of mass action, $np = n_0 p_0 \exp\left[-\frac{|\Delta_{eh}| - \chi_e(c,\rho) - \chi_h(c,\rho)|}{k_B T}\right]$ and the open-circuit voltage is obtained as

$$eV_{\rm oc}(c,\rho) \simeq \alpha E_{\rm ct}(c,\rho) + (1-\alpha)\Gamma(c) + \alpha k_{\rm B}T\ln[n_0p_0] - \beta k_{\rm B}T\hat{\sigma}^2(c)$$
(4)

where we have explicitly labeled the dependence on both composition *c* and roughness ρ . This relationship can finally be expressed in terms of the effective push-out energies $\chi_e + \chi_h = \chi$ simply via

$$eV_{\rm oc}(c,\rho) = eV_{\rm oc}(c,\rho=0) + \alpha\chi(c,\rho).$$
(5)

Two important observations to be made here are as follows: first, as $E_{\rm ct}(c, \rho = 0) + \alpha \Delta \chi(c, \rho)$ is the CT energy in the presence of charge push-out, the above relationship preserves correlations between the CT energy and $V_{\rm oc}$. Second, it adequately captures its low-temperature extrapolation, $V_{\rm oc} \rightarrow E_{\rm ct}$ for $k_{\rm B}T \rightarrow 0$ (for $\alpha \simeq 1$), in agreement with previous studies^{38,39}.

The measured open-circuit voltages as a function of composition are shown in Fig. 4. Note that the red (V_{oc}) and blue (E_{ct}) lines are not fits but theoretical predictions based on the expressions for $E_{ct}(c)$ and $\Gamma(c)$, Eqs. (1) and (5). The solid curves correspond to a surface roughness of $\rho = 1$, in which case the slope reproduces the experimental trends for both V_{oc} and E_{ct} . Note that here the increase in V_{oc} is correlated with the decrease in J_{sc} . This anticorrelation is due to the reduction of the driving force required to convert excitons into chare transfer states, or transfer the hole from the acceptor to the donor^{32,33}.

Of particular interest for solar-cell design is the correlation between the interfacial bias *B* and the open-circuit voltage. Due to symmetries of the electrostatic interaction sums for electrons and holes, the change of *B* and Γ with composition is anticorrelated: $\Delta B = -2\Delta\Gamma$. Meanwhile, $\Delta \chi \leq \Delta B/2$, as follows from Eq. (2). Two extremal situations can therefore be distinguished: (1) $\rho \sim 0$ (flat interface), where the change in open-circuit voltage follows precisely the change in photovoltaic gap, and (2) $\rho \gg 1$ (rough interface), where the change in the photovoltaic gap may be completely compensated by the change in charge-carrier density, thus effecting an approximately constant $V_{\rm oc}$. In systems such as ZnPc/C₆₀ with B > 0, interface roughness is therefore advantageous in that it enhances $V_{\rm oc}$, whereas it will be harmful in systems with B < 0, notably F₄ZnPc/C₆₀.

This conclusion can also be validated experimentally. To this end, we prepared planar heterojunction solar cells with an interlayer of mixed ZnPc:C₆₀ (1:1). By changing the thickness *h* of this interlayer, we effectively modify the roughness ρ of the donor-acceptor interface (as indicated by in situ evaporation studies⁴⁰), and thus the CT state energy:

$$E_{\rm ct}(h) = E_{\rm ct,0} + \frac{\chi_0}{1 + \frac{\delta}{h + h_0}}.$$
 (6)

 $E_{\rm ct,0}$ here denotes the CT state energy in the absence of any interface roughness, $\chi_0 = (B_e + B_h)/2$ is the interfacial bias for the $F_4 ZnPc/C_{60}$ system, δ is the center-of-mass distance between C_{60} and $F_4 ZnPc$ in an edge-on orientation, and h_0 is the interfacial roughness of the planar heterojunction with h = 0, i.e., without mixed interlayer. The experimentally measured $E_{\rm ct}$ and $V_{\rm oc}$ dependencies are shown in Fig. 5 (symbols) and are in excellent agreement with the trends suggested by Eqs. (6) and (3): This is illustrated by the fitted solid lines, which correspond to $E_{\rm ct,0} =$ $0.92 \, {\rm eV}, \ \chi_0 = 0.27 \, {\rm eV}$ and $\rho = \delta/h_0 = 1$. These are in turn very close to the simulated values ($E_{\rm ct,0} = 1.1 \, {\rm eV}, \ \chi_0 = 0.48 \, {\rm eV}$), while the surface roughness estimated from the fit agrees well with the earlier observation that $\rho \simeq 1$ also yields the best agreement between simulated and experimental trends in the compositiondependent setting, shown in Fig. 4.

Conclusions

To conclude, our model of a rough donor-acceptor interface accounts for the bending of the electrostatic potential and accurately captures the correlations between molecular architecture, film composition and open-circuit voltage. The linear dependence of the photovoltaic gap and charge-transfer state energies on composition is driven by the superposition of quadrupolar fields of ZnPc, F₄ZnPc, and F₄*ZnPc. The same superposition creates a composition-dependent electrostatic bias at the donor-acceptor interface. A positive interfacial bias helps to split charge-transfer states and avoid non-geminate recombination. As, however, the change of the photovoltaic gap and electrostatic interfacial bias with composition are anticorrelated, $\Delta B \sim - 2\Delta\Gamma$, an increased bias reduces the open-circuit voltage: For a perfectly planar interface, the change in voltage follows precisely the change in photovoltaic gap. The reduction of the open-circuit voltage that follows from a large positive bias can be avoided at interfaces with nanoscale roughness. In $ZnPc/C_{60}$ and F_4^*ZnPc/C_{60} with B > 0, interface roughness is advantageous in that it recovers the V_{oc} and suppresses charge recombination, whereas it is harmful in F_4ZnPc/C_{60} with B < 0.

Methods

Simulations. The microscopic calculations were performed on atomistic models of mixed ZnPc: F_4ZnPc and ZnPc: F_4^*ZnPc films of 10 nm thickness. Each thin-film layer consists of 17×6 molecules in an edge-on orientation as suggested by X-ray diffraction studies⁴¹. The intermixing of the two species is performed randomly, with the concentration of ZnPc denoted by *c*.

To appropriately model long-range effects in the energy calculations, periodic boundary conditions are used in the plane of the film, without, however, replicating the molecular excitation, or its polarization cloud. Based on this long-range embedding approach³⁷, we obtain the perturbative corrections $\Delta_{h,D}$ to hole (*h*) energy levels in the donor (*D*) film. The ionization energy (IE) results as IE = IE₀ + $\Delta_{h,D}$, where IE₀ denotes the IE of the respective compound in the gas phase³⁰. All calculations were performed using the VOTCA package^{42,43}.

Sensitive EQE_{PV} measurements. The measurements were performed according to previous works, reproduced here for completeness⁴⁴.

The light of a quartz halogen lamp (50 Ŵ) is chopped at 140 Hz and coupled into a monochromator (Cornerstone 260 1/4m, Newport). The resulting monochromatic light is focused onto the organic solar cell (OSC), its current at short-circuit conditions is fed to a current pre-amplifier before it is analyzed with a lock-in amplifier (7280 DSP, Signal Recovery, Oak Ridge, USA). The time constant of the lock-in amplifier was chosen to be 1 s and the amplification of the pre-amplifier was increased to resolve low photocurrents. The EQE_{PV} is determined by dividing the photocurrent of the OSC by the flux of incoming photons, which was obtained with a calibrated silicon (Si) and indium-gallium-arsenide (InGaAs) photodiode.

Electroluminescence measurements were obtained with an Andor SR393i-B spectrometer equipped with a cooled Si and cooled InGaAs detector array (DU420A-BR-DD and DU491A-1.7, UK). The spectral response of the setup was calibrated with a reference lamp (Oriel 63355). The emission spectrum of the OSCs was recorded at different injection currents, which correspond to applied voltages lower than or at least similar to the V_{oc} of the device at 1 sun illumination.

The low-energy tail of the EQE_{PV} spectrum is fitted with a Marcus equation^{38}

$$EQE_{PV}(E) = \frac{f}{E\sqrt{4\pi\lambda k_{B}T}} \exp\left[\frac{-(E_{ct} + \lambda - E)^{2}}{4\lambda k_{B}T}\right]$$
(7)

to obtain the charge-transfer state energy $E_{\rm CT}$, the relaxation energy λ , and the oscillator strength *f*.

In cases where E_{ct} is close to the optical E_{opt} of the donor and accordingly no distinct CT state absorption is observed, E_{ct} was obtained by fitting the high energy tail of the EL spectra³⁸.

Device preparation. The devices were fabricated according to our previous work. The description is reproduced here for completeness⁴⁵. All devices investigated in this work are constructed by a thermal evaporation vacuum system with a base pressure of less than 10^{-7} mbar. Before deposition, ITO substrates (Thin Film Devices Inc., USA) are cleaned for 15 min in different ultrasonic baths with NMP solvent, deionized water, and ethanol, followed by O2 plasma for 10 min. The organic materials are purified 1 or 2 times via thermal sublimation.

A first hole transporting layer is evaporated, consisting of 20 nm 9,9-bis[4-(N,N-bis-biphenyl-4-yl-amino)phenyl]-9H-fluorene (BPAPF) (Lumtec, Germany) doped at 10 wt% with NDP9 (Novaled, Germany). Subsequently, the active layer is evaporated. It comprises 10 nm of a blend of zinc-phthalocyanine (ZnPc) (ABCR, Germany) and tetrafluoro-zinc-phthalocyanine (F₄^{*}ZnPc) (BASF, Germany) or F₄ZnPc (synthesized by Dr. Beatrice Beyer, Fraunhofer FEP, Dresden) in varying ratios, from pure ZnPc to pure F₄^{*}ZnPc or F₄ZnPc (see main text for more information). An additional layer of ZnPc and Buckminster Fullerene (C60) (CreaPhys, Germany) in a 1:1 weight ratio is evaporated, the thickness is varied from 0 to 10 nm. Afterward a 40 nm C₆₀ layer is evaporated. The devices are finalized with 8 nm of Bathophenanthroline (BPhen) (Lumtec, Germany), used as electron contact, and finished with 100 nm of Al. The devices are defined by the area overlap of the bottom and the top contact with an active area of 6.44 mm². To avoid exposure to ambient conditions, the organic part of the device is covered by a small glass substrate, glued on top utilizing an epoxy resin (Nagase ChemteX Corp., Japan) cured by UV light. To hinder degradation, a moisture getter (Dynic Ltd., UK) is inserted between the top contact and the glass.

Current-voltage characteristics. Current-voltage characteristics are measured with a SMU (Keithley 2400) at standard testing conditions (16 S-150 V.3 Solar

Light Co., USA) with a mismatch corrected light intensity and under dark conditions. All results are summarized in Table 1.

Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding authors on reasonable request.

Code availability

The code VOTCA is available from the gitlab repository, https://gitlab.mpcdf.mpg.de/ votca/votca.git.

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Author contributions

C.P. performed simulations and prepared the first draft of the manuscript. J.B., D.S., M.S., S.R., F.P. performed experiments and contributed to manuscript writing. D.A. supervised the computational part of the work and contributed to the manuscript. K.V., K.L., and D.N. conceived the study, supervised the experiments and data analysis, and contributed to the manuscript. All authors contributed to the revision of the final version of the manuscript.

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Correspondence and requests for materials should be addressed to Koen Vandewal or Denis Andrienko.

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