

Molecular Quadrupole Moments Promote Ground-State Charge Generation in Doped Organic Semiconductors

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The role of local environmental interactions on the generation of free charge carriers in doped organic layers is investigated. Via a classical micro-electrostatic model, a dual effect of molecular quadrupole moments of host and dopant molecules on doping is demonstrated. Namely, electrostatic interactions ease ionization of the dopant by altering the energy level alignment between the host and the dopant and reduce the barrier for charge dissociation by flattening the energy landscape around the ionized dopants. These results indicate that tailoring molecular quadrupole moments of the host and/or dopant are an attractive strategy toward improved doping efficiency in organic semiconductors.

1. Introduction

Molecular doping has proven to be a game-changer in the field of organic optoelectronics as it namely allows the precise alignment of the energy levels at the interfaces of multi-layer devices and the minimization of Ohmic losses at the electrodes.^[1] As a result, molecular doping has laid the foundations for the development of efficient organic devices including organic light-emitting diodes, organic field effect transistors, and organic solar cells.^[2] The last years witnessed a considerable advancement of the research on doped organic semiconductors.^[3] These efforts led to a better understanding of the relationships between structure and properties in doped materials,^[4a] as well as to the emergence of novel strategies to dope organic semiconductors, including light-activated dopants,^[4b] anion exchange processing,^[4c] and the use of Lewis acids complexes as molecular dopants.^[4d]

Recent studies have shown that the charge generation in doped organic semiconductors consists of two elementary

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steps. [1b,4,5] The first step involves either a hybridization of the host–dopant frontier molecular orbitals or a ground-state integer charge transfer (CT) process between the host and the dopant. [1a,6] For p-type doping, the latter is favored when the energy difference between the ionization potential (IP) of the host and the electron affinity (EA) of the dopant (i.e., the host–dopant gap $\Gamma_{\rm hd} = {\rm IP}_{\rm host} - {\rm EA}_{\rm dop}$) is smaller in magnitude than the Coulomb binding energy between an electron and its geminate hole sitting on nearest neighbor molecules $(V_{\rm eh})$. [7] In such conditions, a bound CT state with an electron

added to the dopant and a hole left on the host molecule is generated. This is the ionization step. The second step, namely, the charge dissociation, then consists of the spatial migration of the hole (for p-doping) away from the ionized acceptor that requires overcoming the Coulomb binding of the CT pair, which usually amounts to several hundreds of meV. The energetic and kinetic aspects of both steps need to be concomitantly optimized in order to maximize the overall charge generation efficiency, which calls for a control of the interactions between the molecules at the microscopic level.

Tuning these microscopic interactions can be achieved by molecular and material engineering. For instance, Warren et al. demonstrated that the energy levels in an organic semiconductor and the Fermi energy can be simultaneously tuned by taking advantage of molecular quadrupolar interactions. [1c] This result was achieved by co-evaporating a host mixture between zinc-phthalocyanine (ZnPc) and its eight times fluorinated counterpart (F₈ZnPc) together with the p-type dopant F6-TCNNQ. Such mixtures are very interesting because they provide an effective and practical approach to finely control the p-doping efficiency of F6-TCNNQ upon tuning the host energy levels, which are a function of the ZnPc:F₈ZnPc molar ratio. [1c,9]

Here, in the wake of the very recent experimental work reported by Warren et al., [1c] we investigate the role of environmental interactions on the charge generation mechanism in doped binary (ZnPc:F6-TCNNQ and F₈ZnPc:F6-TCNNQ) and ternary (ZnPc:F₈ZnPc:F6-TCNNQ) blends. Based on an in-depth atomistic modeling of electrostatic and dielectric phenomena in molecular solids, [10] we show that charge-quadrupole interactions affect both the ionization step (by reshuffling the energy levels of the dopant and the host) and the charge dissociation step (by creating a favorable energy pathway for the hole). Interestingly, in addition to long-range electrostatics phenomena characterizing the energy landscape of ordered molecular films, [10] we observe that the substitution



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of a host molecule with F6-TCNNQ has a significant shortrange effect, namely, the IP of the host molecules next to a dopant impurity strongly differs (up to 0.4 eV) from the IPs of the host molecules further away. This electrostatic contribution is due to the difference in quadrupole moment between F6-TCNNQ and the host molecules, and it significantly affects the host-dopant gap Γ_{hd} , which on average amounts to 0.63 eV for ZnPc and 0.86 eV for F₈ZnPc. When accounting for screened electron-hole Coulomb interactions ($V_{\rm eh} \approx 0.65$ eV for all samples), we find an overall energy barrier that is close to zero for the ionization step in a binary ZnPc:F6-TCNNQ blend. In F₈ZnPc:F6-TCNNQ, this barrier is 0.2 eV larger, in line with the poorer ionization efficiency observed in literature for the latter blend. Most interestingly, the explicit calculation of the energy profile for charge separation reveals that the quadrupole moments of dopant impurities can positively impact this crucial second step. We believe these results pave the way toward a full control of both steps (ionization and charge dissociation) in the doping of organic semiconductors through the proper engineering of molecular quadrupolar moments.

2. Results

2.1. Binary Blends

First, we calculated the gas-phase IP and EA of the host (ZnPc and F_8 ZnPc) and p-dopant (F6-TCNNQ) molecules (see chemical structures in **Figure 1**a and Figure S2, Supporting

Information) at the evGW many-body perturbation theory level.[11] The obtained values, reported in Figure S3 in the Supporting Information, are in excellent agreement with the experimental ones,^[12] and indicate an increase in both the molecular IP and EA by ≈0.5 eV upon fluorination, as a result of the strong, inductive, electron-withdrawing effects induced by the fluorine atoms.^[13] The huge energy difference in the gasphase between the EA of F6-TCNNQ (4.44 eV) and the IP of the host (6.41 eV for ZnPc and 6.91 eV for F₈ZnPc) would completely impede ground-state CT. The situation changes dramatically when accounting for environmental solid-state effects on the energy levels of both the host and dopant molecules. [10,14] In molecular solids, the IP and EA values depend strongly on collective, long-range, electrostatic effects associated with the varying charge density distributions in the neutral and ionized molecules. These include induction effects, describing the dielectric screening of the polarizable medium to an added charge, and electrostatic interactions, brought about by the molecular electrical multipole moments.[10] In our case, electrostatic interactions are led by the quadrupole term, since the molecules of interest are centrosymmetric and have no dipole moment. [9a]

We first consider the case of a F6-TCNNQ dopant as a single substitutional impurity in two pristine lattices, the first made of ZnPc molecules and the second of F_8 ZnPc ones. We specifically considered films (2D slab calculations) of the two compounds, composed of molecular stacks whose axes lie in the film plane, leading to an edge-on molecular orientation with respect to the slab normal (Figure S4, Supporting Information). [9a] This molecular orientation has been proposed from our previous

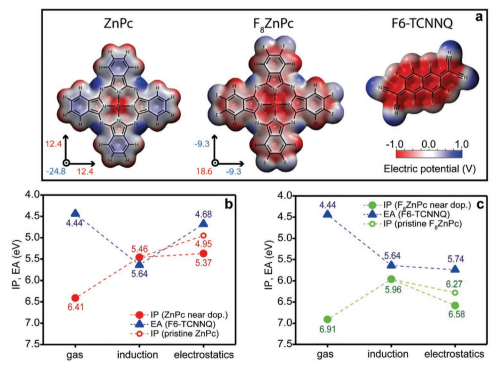


Figure 1. a) Molecular structures and electrostatic properties (electrostatic potential (ESP) color maps and molecular quadrupole components expressed in Debye-Å) of ZnPc, F_8 ZnPc, and F6-TCNNQ, as obtained from DFT calculations at the PBE0/cc-pVTZ level of theory. b,c) Effect of induction and electrostatic interactions on the frontier energy levels of the dopant (blue triangles) and the host molecules, either next to the dopant (full circle) or in the pristine film (open circle). Specifically, b) binary blend composed of ZnPc (host, red dots) and F6-TCNNQ (dopant, blue triangles) and c) binary blend composed of F_8 ZnPc (host, green dots) and F6-TCNNQ (dopant, blue triangles). Energy levels are referred to a common vacuum level.

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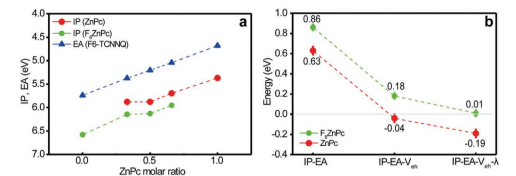


Figure 2. a) Effect of the solid-state interactions on the IPs of ZnPc (red circles) and F_8 ZnPc (green circles) and the EA of F6-TCNNQ (blue triangles) in ZnPc:F8ZnPc blends doped with F6-TCNNQ as a function of ZnPc molar ratio. The data correspond to averages over different microscopic hosts-dopant configurations. b) Energies of the CT state where the electron is localized on the F6-TCNNQ and the hole either on the nearest-neighbor ZnPc or F_8 ZnPc. The energy was calculated by considering different contributions, namely, 1) the Γ_{hd} energy difference, 2) the e-h Coulomb interaction (V_{eh}), and 3) the relaxation energy of the CT state (λ). The last two contributions make the CT state more stable.

grazing-incidence wide-angle X-ray scattering analysis and is in line with literature studies. [1b,c,9a] Electrostatic and induction phenomena are described here with a classical micro-electrostatic (ME) model of atomistic resolution, see the Experimental Section. [15] Our calculations reveal large environmental effects on charge energetics, as shown in Figure 1b,c and reported in Table S2 in the Supporting Information. In line with earlier findings, we notice that induction interactions reduce the host–dopant gap by $\approx\!\!2$ eV irrespective on the host material, as a result of the similar dielectric susceptibilities of organic solids. [10]

On top of induction, the electrostatic contribution further affects the energy levels with an amplitude and sign that depend on the host material. Namely, these interactions are determined by the component of the traceless quadrupolar tensor perpendicular to the molecular plane, which is negative in ZnPc and positive in F₈ZnPc because of the inverted polarity of C-F versus C-H bonds (Figure 1a and Table S1, Supporting Information). In Figure 1b,c, we distinguish between the IP of a host molecule in the pristine host material (open circle) and an host molecule next to the dopant along the π -stacking axis (full circle), which we consider the most important direction for both charge motion and electrostatic interactions. We notice that the sign of the electrostatic shifts of the energy levels of both the dopant (EA, blue triangle) and of its neighboring host molecule (IP, full circle) is the same as the pristine host (IP, open circle). Electrostatic shifts are negative in the ZnPc host (Figure 1b) and positive in F₈ZnPc (Figure 1c). This determines a 1 eV difference in the EA of the F6-TCNNQ depending on the host material, in line with recent findings by some of us.^[14] Interestingly, we also observe that the levels of host molecules strongly depend on their position with respect to the dopant, being the IP of the molecule near the dopant significantly different (0.3-0.4 eV) from that of a host molecule in the pristine host or at infinite distance from the dopant. The position dependence of the host levels can be ascribed to the quadrupolar interaction with the dopant, characterized by an electrostatic layout that is markedly different from that of the replaced host molecule. As it originates from the quadrupole of the single dopant molecule, this effect is rather short-ranged and has an appreciable impact only on the nearest-neighbor molecules.

2.2. Ternary Blends

We now turn our attention to the energy levels of ternary blends composed by a mixed ZnPc:F8ZnPc host, doped with a F6-TCNNQ molecule. ME calculations were carried out for different host mixing ratios, ranging from pure ZnPc to pure F₈ZnPc. For the 1:1 ratio, we opt for an alternating ZnPc:F₈ZnPc mixed-stack motif, which is expected to be the most stable structure for π -stacked molecules with opposite quadrupoles. A similar alternating packing was recently resolved for a 1:1 co-crystal of pentacene and perfluoropentacene. [16] Crystalline supercells of blends with different ratio were created in a similar way but with a different alternating periodicity (one ZnPc molecule every two F₈ZnPc molecules, and vice versa). Our calculations for the energy levels in undoped ZnPc:F₈ZnPc blends of different composition (Figure S5 and Table S3, Supporting Information) are in substantial agreement with the similar ones reported by Schwarze et al.[9a]

The energy levels of host and dopant molecules in ternary blends are shown in Figure 2a. Since different microscopic configurations can be generated upon introducing a dopant in the ZnPc:F₈ZnPc blends (for the 1:1 ratio, the dopant can be placed both at ZnPc and F₈ZnPc site; more configurations are possible for other host compositions), the figure displays the average over the possible microstructures, which is representative of the specific values reported in Figure S6 in the Supporting Information. Calculations in Figure 2a show that the energy levels of host molecules and dopant evolve linearly with the ZnPc molar ratio, as all the molecules probe an effective quadrupolar field resulting from the additive contributions sourced by the two host molecules. Most importantly, this translates into a host–dopant gap Γ_{hd} that does not significantly vary with the host mixing ratio, leading to the conclusion that replacing a significant fraction of ZnPc molecules by electron-poor F₈ZnPc should not be overly detrimental to ionization, at least from an energetic point of view. We note that this was assumed in the statistical approach successfully used by Warren et al. to model the ternary blends.^[1c]

On average, when accounting for the whole range of host chemical composition, Γ_{hd} ranges from 0.63 eV for ZnPc to 0.86 eV in F_8 ZnPc (Figure 2b). This energy difference is largely

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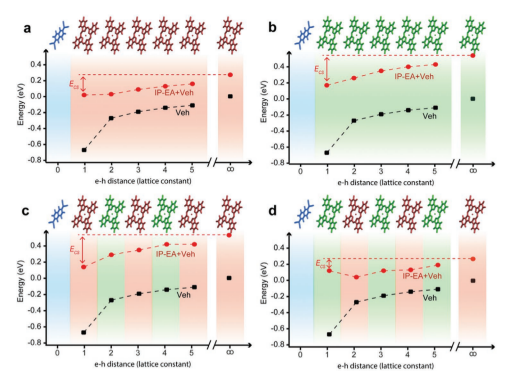


Figure 3. Energies of CT states (red dots), with the electron localized on the F6-TCNNQ and the hole on host molecules, as a function of the electron–hole distance along the π -stacking direction. The screened Coulomb binding energies, $V_{eh}(r_{eh})$ (black squares) and the energy barrier for charge separation (E_{cs}) are also shown. The calculations are reported for a) ZnPc:F6-TCNNQ and b) F_8 ZnPc:F6-TCNNQ binary blends, and two 1:1 configurations of c,d) ZnPc:F6-TCNNQ ternary blends. These results highlight the striking dependence of the charge separation energy profile on the morphology and composition of the host material.

compensated by the screened electron-hole Coulomb binding energy $V_{\rm eh}$ (≈ 0.65 eV), making the overall ionization process barrier-free for ZnPc, while still significantly activated (energy barrier ≈0.2 eV) for F₈ZnPc (Figure 2b). This is in good agreement with the high ionization efficiency experimentally measured in the ZnPc:F6-TCNNQ blends.[1c] Conversely, the energy to ionize the dopant in the F₈ZnPc host remains positive and larger than the thermal energy, which points to a low ionization efficiency, in line with our previous experimental results.^[1c] In addition to $V_{\rm eh}$, we considered the role played by structural reorganization upon charging,[14] quantified by $\lambda = \lambda^+ + \lambda^- = 0.15$ eV, being λ^+ and λ^- the hole and electron reorganization energies of host and dopant, respectively. This contribution, which we computed in gas-phase, represents an upper estimate of the relaxation energy in the solid state where molecular geometries are constrained. Structural reorganization further stabilizes the CT state, hence making the ionization step energetically more favorable (Figure 2b). When accounting for this contribution in full (the geometric relaxation might be partly hindered by steric constraints in the solid), the ionization remains considerably more favorable (by ≈200 meV) in ZnPc compared to F₈ZnPc. Considering the typical error bar (≈100 meV) on the calculated energies and other approximations used in the model (i.e., the neglect of dynamic disorder), a limited electron transfer between the tail states of the dopant and F₈ZnPc molecules cannot be excluded from our simulations, as it could not be ruled out on the basis of our previous electron paramagnetic resonance (EPR) data.[1c]

2.3. Charge Separation

Moving on further, we discuss the results of ME calculations explicitly targeting the electron-hole separation energy profile along the π -stacking direction. Figure 3 and Table S5 in the Supporting Information report the energies of the CT states and the Coulomb binding energies $V_{
m eh}$ as a function of the hole distance from the F6-TCNNO anion for the ZnPc:F6-TCNNO and F₈ZnPc:F6-TCNNQ binary blends and for ternary blends with 1:1 ZnPc:F₈ZnPc composition, distinguishing the two cases where the dopant sits at the molecular site of ZnPc or F₈ZnPc. Remarkably, our calculations highlight that the energy barrier for charge separation (E_{cs}) of the CT state is significantly lower than the mere Coulomb binding energy $V_{\rm eh} \approx 0.65$ eV (compare red and black lines in Figure 3). This is essentially due to the effect of the quadrupolar field of the F6-TCNNQ dopant that destabilizes CT states with charges localized on neighboring molecules (see Figure 2, and Table S4 and Figure S7, Supporting Information). The magnitude of this effect depends however on the host composition and on the dopant location, with E_{cs} ranging from 0.37 eV in F₈ZnPc (Figure 3b) down to 0.15 eV in the ZnPc:F₈ZnPc blend when the impurity substitutes a ZnPc molecule (Figure 3d). In the latter case, and similarly in ZnPc:F6-TCNNQ (Figure 3a), the energy profile for charge separation is rather flat, rationalizing the possibility for a thermally activated charge separation observed for these two systems.[1c] This energy level bending effect sourced by electrostatic intermolecular interactions, similar to those previously



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reported in the context of photoinduced charge separation at donor–acceptor interfaces,^[17] provides a decisive contribution to holes to escape the Coulomb well of their parent dopant.

3. Conclusion

This work delved into the effect of the environmental solid-state interactions on the generation of free charge carriers by doping organic semiconductors. We have carried out our investigations on a model system comprising ZnPc:F₈ZnPc blends, as a host, p-doped with F6-TCNNQ. This ternary blend allowed to control doping efficiency by simply varying the ZnPc molar ratio. [1c] Our theoretical work discloses the fundamental role of chargequadrupolar interactions emerging at different length scales on both 1) the ionization and 2) the charge dissociation steps of molecular doping. Our modeling reveals that the presence of a substitutional F6-TCNNQ molecule in the host lattice strongly influences the IP of the nearest-neighbor molecules without significantly affecting those further apart, ultimately controlling both steps of the doping process. As for the ionization step, this short-range electrostatic contribution significantly increases (up to 0.4 eV) the host-dopant gap. After considering the electron-hole Coulomb interaction, the energy barrier for the formation of a bound CT state between nearest-neighbor molecules becomes close to zero for ZnPc:F6-TCNNQ and is ≈0.2 eV larger for F₈ZnPc:F6-TCNNQ, in line with the larger ionization efficiency measured in the former blend.[1c] The charge dissociation step is expected to be very inefficient on the sole consideration of the Coulomb binding energy of the generated holes to their ionized acceptors, with $V_{\rm eh}$ as large as \approx 0.65 eV. Remarkably, however, this is largely alleviated by the shortrange perturbation of the quadrupole moment of the dopant impurity that, destabilizing CT states between nearest-neighbor molecules, results in a more favorable energy landscape for charge dissociation, with residual effective energy barriers in the range 0.15-0.37 eV depending on the host composition and local micro-structural configuration.

On a more general vein, our findings highlight that the control of electrostatic interactions in thin films is essential to foster both the ionization and the charge dissociation steps in molecular doping. This control requires to simultaneously master the molecular electrostatic moments and the supramolecular organization. Molecular ternary blends of variable composition offer a versatile platform for dissecting the different effects at play, paving the way for the development of new organic semiconductors with improved doping efficiency based on molecular electrostatics engineering.

4. Experimental Section

Gas-phase ground state geometry optimizations were carried out at density functional theory (DFT) level with the Gaussian16^[18] suite, using the PBE0 functional and the cc-pVTZ basis set. The point group symmetry (D_{4h} for ZnPc and F₈ZnPc, and C_{2h} for F6-TCNNQ) was retained during the optimizations. Molecular quadrupole moments are defined as

$$Q_{ij} = \int \rho(\mathbf{r}) \left(r_i r_j - \frac{1}{3} |\mathbf{r}|^2 \delta_{ij} \right) d\mathbf{r}$$
 (1)

where i and j label Cartesian components and $\rho(r)$ is the molecular charge density, including nuclei. Gas-phase GW calculations were performed starting from Kohn–Sham eigenstates calculated for the optimized molecular structures with the ORCA package. [19] A partial self-consistent scheme on the eigenvalues (evGW) was used, along with the cc-pVXZ basis set (where X=D, T). Then, quasiparticle energy levels IP $_g$ and EA $_g$ were extrapolated to the complete basis set limit. [14] The universal Weigend Coulomb fitting set of functions was used as auxiliary basis in the resolution of identity (RI-V) approach. evGW calculations were performed with the Fiesta package. [11a]

ME calculations were parametrized with ESP atomic charges and polarizability tensor calculated with DFT at the PBE0/cc-pVTZ level of theory. ESP charges were computed for all the neutral species, for the positively charged ZnPc and F_8ZnPc and for the negatively charged F6-TCNNQ. The polarizability of charged species was set equal to that of neutral ones. Our ME scheme, implemented in the MESCal code, $^{[1S]}$ relies on a point-charge description of the molecular charge distributions and on tensor dipole polarizabilities distributed at atomic sites. The solid-state charge carrier transport levels were calculated as

$$IP = IP_g + \Delta^h \tag{2}$$

$$\mathsf{E}\mathsf{A} = \mathsf{E}\mathsf{A}_\mathsf{g} - \Delta^\mathsf{e} \tag{3}$$

Environmental solid-state effects entailed a polarization and an electrostatic contribution

$$\Delta^{x} = \Delta_{l}^{x} + \Delta_{E}^{x} \tag{4}$$

where x labels the charged excitation (electron or hole). The polarization Δ_1^x term describes the dynamical response of the surrounding dielectric medium to the presence of a charge carrier. The electrostatic Δ_E^x term corresponds to the energy to charge a molecule in the electrostatic field of all other molecules in the neutral crystal. [16]

CT states with localized charges were modeled as classical electronhole pairs. The binding energy V_{eh} of ion pairs was computed as the sum of screened Coulomb interactions between a hole residing on a host molecule (ZnPc or a F_8 ZnPc) and an electron on the F6-TCNNQ dopant

$$V_{\rm eh} = \frac{1}{\varepsilon_r} \sum_{j,i} \frac{\delta q_i^{\rm e} \delta q_j^{\rm h}}{r_{ij}} \tag{5}$$

medium, where $\varepsilon_r=3.2$ is the computed dielectric constant of the medium, $\delta q_i^e(\delta q_j^h)$ is the difference between the atomic charge of the anion (cation) compared to the neutral molecule, and r_{ij} is the interatomic distance.

The structural relaxation energies λ^\pm upon charging were computed at the DFT level (PBE0/cc-pVTZ) by comparing the vertical and the adiabatic IP and EA of the hosts and of the dopant, respectively. In F6-TCNNQ, the dihedral angles involving the dicyanovinyl groups were kept frozen during the optimizations, in order to get rid of low-frequency contributions to $\lambda=\lambda^++\lambda^-$. Specifically, $\lambda^+=20$ meV for ZnPc, $\lambda^+=37$ meV for F8ZnPc, and $\lambda^-=131$ meV for F6-TCNNQ.

Molecular stacks were build following the recipe described in ref. [9a], starting from the crystallographic structure of CuPc, [20] replacing the Cu atoms with Zn ones and replicating the unit cell along the three directions. The convergence of the electrostatic energy $\Delta_{\xi}^{\rm E}$ was checked with calculations considering pure ZnPc films of 3 (5 × 17 × 3 supercell) and 5 (5 × 17 × 3 supercell) molecular layers stacked along the film normal direction z. $\Delta_{\xi}^{\rm E}$ of a central ZnPc molecule in each layer was computed accounting for periodic replica in the x-y plane within a cut-off distance R ranging from 120 to 300 Å. The electrostatics converged (within 20 meV) in the three middle layers for a 5 × 17 × 5 supercell with a cut-off of 300 Å (Figure S1, Supporting Information). Therefore, all subsequent calculations employed a 5 × 17 × 3 stack with 300 Å in-place cut-off for the computations of the $\Delta_{\xi}^{\rm E}$ terms. On the other hand, polarization energies $\Delta_{1}^{\rm K}$ were calculated



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for spherical clusters of increasing radii extracted from the bulk structure and centered at the molecule of interest and extrapolated for an infinite system.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest

The authors declare no conflict of interest.

Author Contribution

A.P. and G.L. contributed equally to this work. A.P. and G.L. conceived and designed the project with significant input from D.B., G.D., and M.R.. G.L. and A.P. carried out all the DFT and ME calculations. D.B. and G.D. supervised the work and contributed with fruitful discussion. The first manuscript draft was prepared by A.P. and G.L. All the authors reviewed the manuscript and participated actively in the final discussion of the results.

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charge generation, molecular doping, organic semiconductors, quadrupole moment

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