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Quantifying Image Charge Effects in Molecular Tunnel Junctions **Based on Self-Assembled Monolayers of Substituted** Oligophenylene Ethynylene Dithiols

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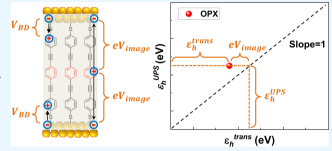
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ABSTRACT: A number of factors contribute to orbital energy alignment with respect to the Fermi level in molecular tunnel junctions. Here, we report a combined experimental and theoretical effort to quantify the effect of metal image potentials on the highest occupied molecular orbital to Fermi level offset, $\varepsilon_{\rm h}$, for molecular junctions based on self-assembled monolayers (SAMs) of oligophenylene ethynylene dithiols (OPX) on Au. Our experimental approach involves the use of both transport and photoelectron spectroscopy to extract the offsets, $\varepsilon_{\rm h}^{\rm trans}$ and $\varepsilon_{\rm h}^{\rm UPS}$, respectively. We take the difference in these quantities to be the image potential energy eV_{image} . In the theoretical approach, we use



density functional theory (DFT) to calculate directly eV_{image} between positive charge on an OPX molecule and the negative image charge in the Au. Both approaches yield $eV_{\rm image} \sim -0.1$ eV per metal contact, meaning that the total image potential energy is ~ -0.2 eV for an assembled junction with two Au contacts. Thus, we find that the total image potential energy is 25-30% of the total offset $arepsilon_{
m h}$, which means that image charge effects are significant in OPX junctions. Our methods should be generally applicable to understanding image charge effects as a function of molecular size, for example, in a variety of SAM-based junctions.

KEYWORDS: molecular junction, tunneling, charge transport, image charge, Fermi level-HOMO offset, single level model, photoelectron spectroscopy

INTRODUCTION

Relating the conductance behavior of molecules to their electronic structure is a central focus of the field of molecular electronics. 1-20 However, electronic phenomena at the interface between discrete molecules and metal electrodes generally yield substantial changes in orbital energies with respect to their values in isolated molecules, and thus, understanding these factors is critical. The presence of metal-molecule bond dipoles and metal image potentials are two recognized effects responsible for shifting of orbital energies when a molecule binds to a metallic substrate. 21-32 Additionally, for junctions based on molecular assemblies such as self-assembled monolayers (SAMs) rather than single molecules, intermolecular interactions also play a major role in energy level alignment.²⁸

In this paper, we seek additional clarity on factors affecting orbital energies in junctions based on SAMs of an oligophenylene ethynylene (OPE) dithiol^{33–37} and substituted derivatives (OPX = OPM, OPF, and OPN) on Au, as shown in Figure 1A,B. The original design principle underlying the OPX system was to exploit electroactive donating and withdrawing substituents on the central ring to manipulate the orbital level

alignment. However, we have previously demonstrated a pronounced highest-occupied molecular orbital (HOMO) level pinning in molecular junctions based on three of these molecules, OPE, OPF, and OPN.²⁸ The HOMO pinning, which appears to be a general phenomenon in molecules strongly coupled to metal electrodes, 26,38-45 is striking, as OPE, OPF, and OPN have gas-phase (isolated molecule) ionization energies $E_{\rm I}$ that differ by over 0.6 eV due to the different electroactive substituents on the central ring. Density functional theory (DFT) calculations reveal that the differences in $E_{\rm I}$ are nearly completely wiped out in crystalline monolayers of OPE, OPF, and OPN due to intermolecular interactions (specifically electrostatic interactions); the ionization potentials (IPs) for the three monolayers are approximately equal.²⁸ Thus, intermolecular interactions play a

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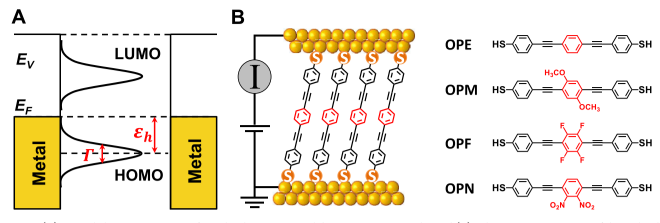


Figure 1. (A) Typical electronic structure of a molecular junction with key parameters ε_h and Γ and (B) Schematic representation of the molecular junctions based on OPE, OPM, OPF, and OPN with Au electrodes.

substantial role in the HOMO pinning and energy level alignment that occurs in close-packed OPX monolayers contacted by metal electrodes. We know from our prior theoretical work that the Au–S bond dipoles are also critical to energy level alignment in these systems, 46 but they are nearly equal across all the OPX derivatives in SAMs. 27,28

For molecules such as the OPX series where the bond dipoles at either contact are noninteracting (see Results and Discussion), the spectra of effects impacting orbital energies in molecular junctions are approximately independent. In such a case, one can write that the HOMO position ε_h with respect to the Fermi level E_f is the sum of four contributions

$$\varepsilon_{\rm h} + E_{\rm f} = E_{\rm I} + \Delta E_{\rm I_intermolecular} + {\rm BD} + eV_{\rm image}$$
 (1)

where $E_{\rm I}$ is the IP of the isolated molecule, $\Delta E_{\rm I_intermolecular}$ is the shift in $E_{\rm I}$ due to intermolecular interactions in a SAM, BD is the energy shift due to the molecule—metal bond that induces the bond dipole, 21,22 e is the elementary charge, and $V_{\rm image}$ is the image potential associated with the metal electrodes. 29,30 Please note that in eq 1, $\varepsilon_{\rm h}$ and $E_{\rm f}$ are taken as positive quantities simply for convenience ($\varepsilon_{\rm h} > 0$ is typical in molecular electronics), though in formal electron energy level diagrams, they would both be negative of the vacuum level $E_{\rm v}=0$ (Figure 1). With the sign convention used here, $eV_{\rm image} < 0$ reduces, $\varepsilon_{\rm h}$, that is, shifts the HOMO upward.

It is also critical to note that the absolute magnitude of $V_{\rm image}$ is anticipated to scale with the number of metal contacts, that is, $V_{\rm image}$ for a SAM-based molecular junction with two metal contacts is expected to be approximately twice as large as $V_{\rm image}$ for a SAM on a metal substrate (i.e., with one contact), as long as the relevant transport orbital (e.g., HOMO) is relatively spatially symmetric within the junction. In contrast, $E_{\rm I}$ and $\Delta E_{\rm I_intermolecular}$ do not depend on the number (or presence) of contacts, and BD, as we will explain below, is approximately the same for one vs. two metal contacts. The significance of this is described shortly.

Here, our particular focus is on determining the role of the image potential $V_{\rm image}$ on orbital energies in OPX molecular junctions. Ionization energies and electron affinities (EAs) of molecules near metal surfaces can be profoundly affected by the metal's image potential, $^{23,24,29-32}$ and this in turn can influence the HOMO (or LUMO) alignment with respect to the Fermi level. However, these effects depend on the proximity of the molecule to the metal surface and, crucially, on molecular orientation with respect to the surface. $^{47-49}$ It is

not immediately clear how big the image charge effect will be for a given molecular junction.

In this paper, we estimate $V_{\rm image}$ experimentally and theoretically. Our experimental approach involves extraction of the HOMO to Fermi level offset, $\varepsilon_{\rm h}^{\rm trans}$, from current–voltage (I-V) measurements on Au–OPX–Au junctions and subsequent comparison with $\varepsilon_{\rm h}^{\rm UPS}$ for OPX SAMs on Au obtained by ultraviolet photoelectron spectroscopy (UPS). The transport measurement involves two contacts to the SAM, while UPS probes the OPX SAM on one metal contact. Thus, $\varepsilon_{\rm h}^{\rm trans}$ is the experimental estimation of $\varepsilon_{\rm h}$ shown in Figure 1A from transport measurements, and $\varepsilon_{\rm h}^{\rm UPS}$ is the corresponding experimental value of $\varepsilon_{\rm h}$ for a SAM (one contact) obtained by UPS. As noted above, because to a first approximation only $eV_{\rm image}$ on the right hand side of eq 1 depends on the number of contacts, we can employ the difference between $\varepsilon_{\rm h}^{\rm trans}$ and $\varepsilon_{\rm h}^{\rm UPS}$ to provide an estimate of the image charge effect due to the second contact, that is, $eV_{\rm image} = \varepsilon_{\rm h}^{\rm trans} - \varepsilon_{\rm h}^{\rm UPS}$.

Our theoretical approach to $V_{\rm image}$ begins with DFT calculations of the electronic structure of a neutral junction to evaluate the relative location of the HOMO levels of the OPX derivatives with respect to the Fermi level of Au. The image charge effect is then estimated in a second step by first mirroring the distribution of the positive charge over the OPX backbone with respect to the metal surface and then by computing the Coulomb interaction between the two distributions of opposite sign; this is achieved by making the reasonable assumption that the interface dipole and the image charge have a weak mutual influence. Importantly, both experiments and theory demonstrate the image potential contribution to $\varepsilon_{\rm h}$ to be ~0.1 eV per contact for OPX, which translates to approximately 25-30% of the total magnitude of $arepsilon_{
m h}$ for complete, two-contact junctions. Note that this significant effect differs slightly from previous findings in which the image charge in single molecule junctions was found to dominate the energy level alignment. 23,24,29-32 Our results do not contradict the earlier findings, but rather indicate that image charge effects are system-specific and depend on the size and orientation of molecules with respect to the metal electrodes, as well as the electrode shape and total number of electrodes (up to 3 if there is a gate).

In summary, we present an experimental approach using junction conductance measurements and photoelectron spectroscopy to quantify image charge effects. The results are verified theoretically and are important for developing a

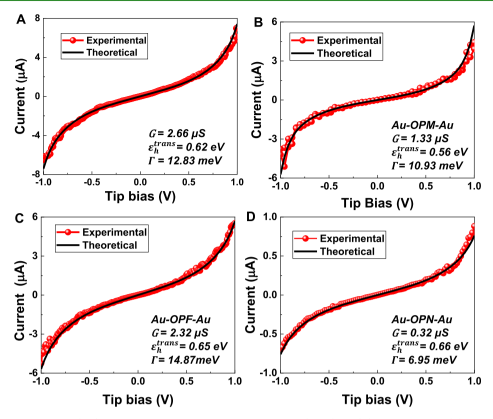


Figure 2. Experimental I-V curves (red) and single level model simulations via eq 2 (black) for (A) Au–OPE–Au, (B) Au–OPM–Au, (C) Au–OPF–Au, and (D) Au–OPN–Au junctions. The three extracted parameters for each junction, that is, low bias conductance G (1/R), the energy offset $\varepsilon_h^{\text{trans}}$, and coupling Γ, are listed in each panel. The N values used for calculating Γ are 80, 45, 55, and 35 for OPE, OPM, OPF, and OPN, respectively.

comprehensive picture of factors impacting energy level alignment in SAM-based molecular junctions.

METHODS

Materials. Gold nuggets (99.999% pure) were purchased from Mowrey, Inc. (St. Paul, MN). Evaporation boats and chromium evaporation rods were purchased from R. D. Mathis (Long Beach, CA). Silicon (100) wafers were obtained from WaferNet (San Jose, CA). Contact mode AFM tips (DNP SiN probes) were purchased from Bruker AFM Probes. The synthesis of the three-ring OPE and substituted derivatives (OPF and OPN) has been described elsewhere. S0,51 The synthesis of OPM is described in Section S1 in Supporting Information.

Monolayer Growth and Characterization. Preparation of conducting tips and template-stripped flat Au substrates is described in a previous publication. SAMs were formed by immersing clean template-stripped flat metal substrates into ethanol solutions of the molecules at a concentration of ~1 mM for 20 h. The thicknesses of the OPE, OPM, OPF, and OPN SAMs were characterized by ellipsometry (Figure S1) carried out on a VASE spectroscopic ellipsometer (J. A. Woolam Co., Inc.). The extracted molecular tilt angles from the normal direction amount to 33, 34, 30, and 34° for OPE, OPM, OPF, and OPN, respectively. The X-ray photoelectron spectroscopy (XPS) was used to characterize the difference of the surface coverage of SAMs (Figure S2) and the UPS was used to measure the HOMO-Fermi level offset of OPE, OPM, OPF, and OPN SAMs on the Au substrate. More details about XPS and UPS are described in Supporting Information.

Transport Measurements. Current—voltage measurements were completed by mounting the substrates in the AFM and bringing the metal coated tip into contact with the SAM under ~1 nN of applied compressive load. Voltages were applied to the tip with a Keithley model 236 source-measure unit operated in "DC mode". Voltage was

swept at the tip, with the bottom substrate grounded, and current–voltage characteristics (I-V) were recorded (V>0 corresponds to a positive voltage on the tip). All measured I-V curves (about 100 curves were collected for each OPX junction) crossed over from practically linear at low biases to gradually more nonlinear at higher biases. The inverse of the slope of the linear portion of the I-V characteristic was used to define a junction resistance R at low bias. Voltage sweeps to $\pm 1.2~V$ were applied to observe the pronounced nonlinear (I-V) behavior.

Theoretical Calculations. The methodology used to model the SAMs is the same as that used in our previous work. ²⁸ Additional information related with the computational details and the models employed to simulate the OPE-based SAMs and junctions are provided in Section S2 of the Supporting Information.

■ RESULTS AND DISCUSSION

Analysis of Transport Characteristics; Extraction of $\varepsilon_{\rm h}^{\rm trans}$. Figure 2 shows representative I-V characteristics for junctions based on OPX SAMs with Au contacts; such a complete dataset was not shown in our prior publication. To analyze these curves and to extract $\varepsilon_{\rm h}$ and the metal–molecule coupling Γ (see Figure 1), we employ the analytical single level model, which we have shown previously applies well to simple molecular tunnel junctions. In this model, the I-V characteristics of a symmetric junction are given as 42,52

$$I = GV \frac{\varepsilon_{\rm h}^2}{\varepsilon_{\rm h}^2 - (eV/2)^2}$$
 (2)

where the zero-bias conductance G of the junction can be expressed as follows

Table 1. Summary of the Experimental Results, Including Low Bias Conductance of the Junctions G, Estimated Conductance per Molecule G_m , Relative Difference in Molecular Coverage A, Energy Offset $\varepsilon_h^{\text{trans}}$, and Average Coupling Γ for the Junctions Made with SAMs of the OPE derivatives^a

quantity	OPE	OPM	OPF	OPN
G	$(3.00 \pm 0.71) \times 10^{-6}$	$(2.20 \pm 0.50) \times 10^{-6}$	$(2.56 \pm 0.62) \times 10^{-6}$	$(0.31 \pm 0.11) \times 10^{-6}$
A	1	0.57	0.69	0.43
G_{m}	$(3.75 \pm 0.88) \times 10^{-8}$	$(4.80 \pm 1.09) \times 10^{-8}$	$(4.66 \pm 1.12) \times 10^{-8}$	$(0.87 \pm 0.33) \times 10^{-9}$
$oldsymbol{arepsilon}_{ m h}^{ m trans}$	0.61 ± 0.06	0.56 ± 0.07	0.63 ± 0.07	0.67 ± 0.1
$arepsilon_{ m h}^{ m UPS}$	0.68 ± 0.1	0.65 ± 0.1	0.71 ± 0.1	0.74 ± 0.1
$\Gamma_{ m av}$	13.42 ± 2.20	13.94 ± 2.01	15.44 ± 2.75	7.09 ± 1.51

 ${}^a\epsilon_{
m h}^{\rm UPS}$ is the Fermi-HOMO offset of the OPE-based SAMs obtained by UPS. Units: G in S, $V_{
m t}$ in V, $\epsilon_{
m h}$ in eV, and Γ in meV obtained from eq 3 by taking N=80 for OPE as the reference junction⁵⁵ and N=45, 55, and 35 for OPM, OPF, and OPN, respectively. The averaged results from transport measurements were obtained from approximately 100 I-V curves for each OPX SAM.

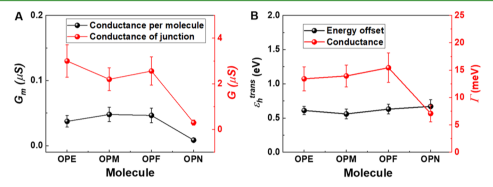


Figure 3. Plots of the comparison between (A) low bias conductance per molecule $G_{\rm m}$ and junction conductance G and (B) energy offset $\epsilon_{\rm h}^{\rm trans}$ and coupling Γ for OPE, OPM, OPF, and OPN junctions.

$$G = NG_0 \frac{\Gamma^2}{\varepsilon_h^2} \tag{3}$$

In eq 3, $\Gamma = \sqrt{\Gamma_s \Gamma_t} = \varepsilon_h \sqrt{G/NG_0}$ is the average interface coupling; Γ_s and Γ_t are determined by the molecular coupling to the substrate (s) and tip (t) ($\Gamma_s \approx \Gamma_t$ in symmetric junctions), $G_0 = 2e^2/h$ is the quantum conductance, and N is the number of molecules participating in the transport.

We have used eq 2 to fit the data in Figure 2 (see Supporting Information for the fitting method). Doing so, we reproduce the individual measured I-V curves for the OPX junctions extremely well, as shown by the black traces in Figure 2. We believe that the very good match between the experimental and simulated I-V curves indicates that the single level model applies well for OPX-based molecular junctions, as we have shown for other molecular systems. 41-45,53,54

The average values for the transport parameters extracted from approximately 100 I-V curves for each type of OPX SAM are compiled in Table 1. Evidently, substitution of the central ring with electron donating or withdrawing groups does influence the low bias conductance G of the OPX molecular junctions, but not to a large extent. Because substitution introduces steric interactions in the SAMs resulting in variations in the molecular packing density, the differences in molecular surface coverage across the OPX series were estimated by XPS (Figure S2). The C_{1s} signal normalized to the Au_{4f} signal for each OPX derivative was employed to calculate the relative surface coverage A of the SAMs; values of A are listed in Table 1. After correction for surface coverage, the conductance $G_{\rm m} = G/N$ per molecule also shows modest variation among the substituted OPX molecules. Figure 3A depicts the G and G_m data graphically and indeed only the OPN system appears to be appreciably different.

The estimated energy offsets $arepsilon_{
m h}^{
m trans}$ and interface couplings Γ extracted from the transport data are also listed in Table 1 and they are plotted in Figure 3B. It is evident that $\varepsilon_{
m h}^{
m trans}$ is essentially invariant across the OPX species, which is the signature of HOMO level pinning in molecular junctions. 28,42 Again, this is striking because the electron donating (OPM) and electron withdrawing (OPF and OPN) substituents on the central phenylene ring can shift the IP by as much as 0.6 eV for isolated (gas phase) OPX molecules.²⁸ Comparison of Figure 3A,B reveals that the weak dependence of the energy offset $\varepsilon_{\rm h}^{\rm trans}$ on the electroactive substituents cannot explain the much larger (by one order of magnitude) difference in conductance $G_{\rm m}$ between OPN and the other OPX molecules (OPE, OPM, and OPF). The cause of the conductance difference for OPN is Γ , which, as shown in Figure 3B, is significantly lower for OPN than for the other molecules. Note that $G \propto \Gamma^2$ via eq 3. Given the very similar $\varepsilon_{\rm h}^{\rm trans}$ values for the OPX junctions, the larger drop of the coupling (as well as the conductance) obtained for OPN is tentatively attributed to stronger localization of the HOMO level over the central ring, thus limiting the electrode-molecular interfacial coupling and hence the efficiency of charge transport.²⁸

Measurement of $\varepsilon_h^{\text{UPS}}$ and Estimation of the Image Charge Potential V_{image} . UPS is the standard experimental method used to obtain the energy of the HOMO for molecular thin films. 44,56-58 To obtain an independent verification of the energy level alignment $arepsilon_{
m h}^{
m trans}$ extracted from transport measurements and to understand the influence of the top electrode on $\varepsilon_{\rm h}$, that is, to estimate the image charge effect, we performed UPS on each OPX SAM. The spectra for OPE, OPM, OPF, and OPN SAMs on Au substrates are shown in Supporting Information, Figure S3. In these UPS spectra, the Fermi edge at zero binding energy is clearly evident and the HOMO-Fermi level offsets are determined using standard extrapolation protocols. 59,60 The resulting $\varepsilon_{\rm h}^{\rm UPS}$ values obtained by UPS are then compared to the corresponding $\varepsilon_{\rm h}^{\rm trans}$ values obtained from transport data, as shown in Figure 4 and Table 1. It is clear in

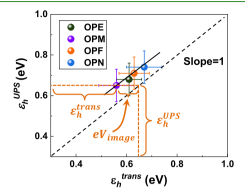


Figure 4. Correlation of ε_h^{trans} extracted from transport measurements (and the single level model) with ε_h^{UPS} from UPS measurements on SAMs for OPE, OPM, OPF, and OPN with Au contacts. The black dashed line shows the slope = 1 trend for perfect correspondence, and the black solid line shows the linear fit.

both Table 1 and Figure 4 that $arepsilon_{
m h}^{
m trans}$ is systematically less than $\varepsilon_{\rm h}^{\rm UPS}$ for each OPX molecule. This immediately implies an upward energy level shift of the OPX HOMOs in the twocontact molecular junctions versus the one-contact SAMs, that is, $eV_{\rm image}$ < 0, where $eV_{\rm image} \approx \varepsilon_{\rm h}^{\rm trans} - \varepsilon_{\rm h}^{\rm UPS}$. This result is consistent with expectations because the image charge stabilization should be additive, that is, greater (roughly twice the size) for two contacts than for one contact. Importantly, $\varepsilon_h^{\text{UPS}}$ also scales linearly with $\varepsilon_h^{\text{trans}}$ as shown by the fit to the data in Figure 4 with a slope = 0.9. The linear correlation with a slope ~1 is important in two respects. First, it is consistent with the straightforward interpretation that ${
m e}V_{
m image}$ is the only term that changes in eq 1 when comparing OPX SAMs vs. OPX junctions. If the data were not linear with slope \sim 1, we would be forced to conclude that either the value of $\mathrm{e}V_{\mathrm{image}}$ was different for each OPX molecule when adding a second contact (which is not supported by theory, nor expected, see below) or that the BD term for each OPX molecule was different in the two-contact junctions versus the one-contact SAM. Second, we note that there is significant uncertainty associated with individual values of $\varepsilon_{\rm h}^{\rm trans}$ and $\varepsilon_{\rm h}^{\rm UPS}$ for any given OPX molecule, as indicated by the error bars in Figure 4. However, the four measurements on four molecules minimize the overall error. From the four measurements, we conclude that $eV_{\text{image}} = -0.08 \pm 0.01$ eV for the OPX system.

As just noted, the bond dipole energy BD in eq 1 is not influenced by whether there are one or two contacts to an OPX SAM. A second bond dipole likely forms when the top contact is made to an OPX SAM, but it does not lead to a change in $arepsilon_{
m h}^{
m trans}$. This is because the charge transfer is localized at the contact and the vacuum level shift associated with the dipole is the mirror image of the shift at the first contact. This point is discussed further in the theory section. Note that the independence of BD on the number of contacts is a very different conclusion than we made about eV_{image} , which is expected to scale with the number of contacts because of the longer range of the charge-metal interaction that underlies the image charge effect, and the fact that the HOMO is symmetrically distributed in the junction.

We conclude from our experimental measurements that ${
m eV}_{
m image}$ associated with a single contact is ${\sim}{-}0.08$ eV for SAMbased OPX junctions. Given that the total energy level offset $arepsilon_{
m h}^{
m trans}$ is ~0.6 eV, $eV_{
m image}$ is clearly important as each contact provides \sim -0.08 eV, meaning that the total contribution of eV_{image} to $\varepsilon_{\text{h}}^{\text{trans}}$ is \sim -0.16 eV or 25% of the total offset from E_{f} . One can say that image charge effects are a significant factor though not the dominant factor—in overall energy alignment in OPX junctions. ^{23,24,29–32}

We next compare the experimental data to DFT calculations. Specifically, we are able to assess the role of interfacial charge transfer and the resulting Au-S bond dipoles, which determine BD. We also explicitly estimate eV_{image} .

DFT Calculations. Theoretical results for OPE, OPF, and OPN SAM-based junctions have been reported in a previous paper²⁸ and that previous study has been extended here to consider OPM and eV_{image} explicitly. Because the HOMO energy offset appears experimentally to be weakly affected by the actual areal density of the OPX derivatives, the SAMs were optimized with the same unit cell for each derivative, while imposing that the S atom is anchored on a Au bridge site and that the orientation of the backbone is normal to the surface (Figure S5); the latter constraint is motivated by the fact that the measured tilt angles are moderate and similar for all OPX compounds. The full theoretical methodology is described in ref 28. Table 2 collects the calculated HOMO offsets with

Table 2. Summary of the DFT Calculated Energy Offset $arepsilon_{h,DFT}^{SAM}$ and $arepsilon_{h,DFT}^{junction}$ in SAMs and Molecular Junctions, Respectively, and Amplitudes of the Image Charge Energies $(eV_{IP}$ and $eV_{EA})$ Calculated for the IP and EA. All Values are Given in eV.

quantity	OPE	OPM	OPF	OPN
$arepsilon_{ ext{h,DFT}}^{ ext{SAM}}$	0.61	0.63	0.64	0.65
$arepsilon_{ m h,DFT}^{ m Junction}$	0.54	0.56	0.56	0.59
$eV_{ m image}^{ m IP}$	-0.11	-0.11	-0.13	-0.14
$eV_{ m image}^{ m EA}$	0.13	0.13	0.12	0.12

respect to the Fermi level in one-contact OPX SAMs ($\varepsilon_{\rm h,DFT}^{\rm SAM}$) and two-contact molecular junctions ($\varepsilon_{h,DFT}^{Junction}$). Note that these values are extracted from the density of states of the SAM (central scattering region of the junction) projected onto C atoms as the energy at the maximum of the first intense band below the Fermi level (see Figures S8 and S9). The calculated $\varepsilon_{\rm h}$ values appear to be in quite good quantitative agreement with the corresponding experimental transport and UPS results, as shown in Figure 5A,B, in spite of the well-known limitations of DFT approaches in accurately predicting the energetic level alignment at hybrid interfaces. This can be qualitatively explained by a compensation error between the underestimation of the HOMO-LUMO gap of isolated molecules at the DFT/GGA level versus the neglect of electronic polarization effects in the solid state (due to the presence of electrodes and neighboring molecules) that strongly reduce the gap of isolated molecules (no explicit charge is considered when computing transmission spectra within the DFT/NEGF approach).61

Importantly, the DFT calculations show that anchoring the SAMs on Au or introducing a second Au slab to build up the junction (Figure S6) promotes a small charge transfer at each interface (between 0.12 and 0.14 lel, Figure S7), as computed using either the Hirschfeld or Voronoi charge population

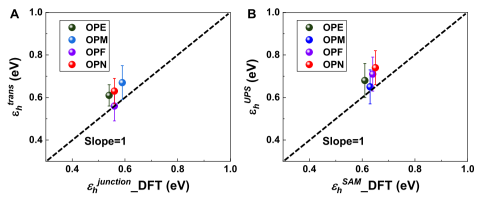


Figure 5. (A) Correlation of $\varepsilon_h^{\text{trans}}$ from transport measurements (and the single level model) with DFT calculated energy offsets $\varepsilon_{h\,\text{DFT}}^{\text{junction}}$ for SAM-based junctions of OPE, OPM, OPF, and OPN with two Au contacts. (B) Correlation of $\varepsilon_h^{\text{UPS}}$ from UPS measurements with the DFT calculated energy offsets $\varepsilon_h^{\text{SAM}}$ for OPE, OPM, OPF, and OPN SAMs on an Au substrate.

analysis (Table S1). The bond dipoles are symmetric (they point in opposite directions) and their associated electrostatic potentials, centered at each end of the OPX molecules, weakly overlap, and this means that the alignment of the HOMO level is controlled by a single electrode. In other words, the formation of a second bond dipole at the second Au contact/ SAM interface does not produce an additional electrostatic energy shift BD of the HOMO level (see eq 1). This is nicely demonstrated in Figure 6, which shows the planar averaged

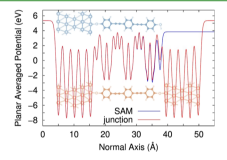


Figure 6. Calculated planar averaged potential in the Au/SAM vs Au/SAM/Au systems involving OPE, pointing to the weak coupling between the bond dipoles in the junction in view of the similar energy profile over the molecular backbone.

electrostatic potential across the Au/SAM and Au/SAM/Au systems involving OPE, highlighting the identical profile over the backbone and hence the independent character of the bond dipoles. BD is thus determined by one contact, that is, by the formation of the SAMs on Au. This is a crucial conclusion, because as discussed in the Introduction, only $eV_{\rm image}$ on the right hand side of eq 1 depends on the number of contacts, which allows us to estimate $eV_{\rm image}$ from the difference in experimental two-contact versus one-contact $\varepsilon_{\rm h}$ values, $eV_{\rm image} \approx \varepsilon_{\rm trans}^{\rm trans} - \varepsilon_{\rm h}^{\rm UPS}$.

In order to calculate $eV_{\rm image}$ theoretically, taking the difference between computed $e_{\rm h,DFT}^{\rm function}$ and $e_{\rm h,DFT}^{\rm SAM}$ values—in analogy with the experimental approach—is not valid, as the DFT-NEGF calculations described so far were performed on neutral systems; they do not take into account the influence of electronic polarization, due both to the creation of an image potential in the metallic electrode when a molecule is charged and to induced dipoles in molecules surrounding the charged molecule. Thus, $eV_{\rm image}$ has been explicitly calculated here for one electrode as the interaction energy between the

distribution of a positive charge over a molecule and its image (see Supporting Information for further details). We find that it leads to a shift of the IP and EA for all OPX compounds. The computed values of $\mathrm{eV}_{\mathrm{image}}^{\mathrm{IP}}$ and $\mathrm{eV}_{\mathrm{image}}^{\mathrm{EA}}$ reported in Table 2 are obtained with the RO(MP2) approach and are consistent with those previously computed for oligophenylene molecules.²⁷ They are also independent of the level of theory employed (Table S2) and of the molecular tilt angle (Table S3) and typically range between -0.11 and -0.14 eV, which is in good agreement with eV_{image} estimated from experiments. Thus, the DFT results support the conclusion that image effects are significant, but not dominant, in OPX junctions. Before closing this part, we note that our theoretical model used to estimate the image potentials neglects cooperative effects between neighboring molecules within the SAM that might lead to some depolarization effects.⁶² Nevertheless, it provides a first approximation to the magnitude of the image effect associated with one electrode, and it verifies that the chemical substituents on the OPX core do not significantly influence the magnitude of the potential.

CONCLUSIONS

Through a combination of experiments and quantum chemical calculations, we have investigated the magnitude of the image charge effect on the HOMO energy offsets in molecular junctions based on SAMs of a series of OPX molecules on Au. On the experimental side, we have proposed that the image potential energy $eV_{\rm image}$ can be estimated by taking the difference between the HOMO level offset determined from transport measurements, $\varepsilon_{\rm h}^{\rm trans}$, and the offset determined by UPS, $\varepsilon_{\rm h}^{\rm UPS}$, that is, $eV_{\rm image} \approx \varepsilon_{\rm h}^{\rm trans} - \varepsilon_{\rm h}^{\rm UPS}$. Using that approach, we find that eV_{image} is -0.08 ± 0.01 eV per metal contact, so that the total contribution of eV_{image} to ε_{h} for a two contact junction is -0.16 eV or about 25% of the total offset. DFT calculations produce similar numbers for eV_{image} . We thus conclude that image charge effects are significant in OPX junctions and influence the overall energy level lineup. Furthermore, the combined experimental and theoretical approaches employed here should be applicable to other SAM-based junctions, allowing for the impact of image potentials to be assessed generally in SAM-based molecular junctions as a function of molecular orientation and length, for example.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.1c16398.

Experimental and theoretical details and supplementary tables and figures (PDF)

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Notes

The authors declare no competing financial interest.

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