Fully Atomistic Understanding of the Electronic and Optical Properties of a Prototypical Doped Charge-Transfer Interface

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Supporting Information

ABSTRACT: The current study generates profound atomistic insights into doping-induced changes of the optical and electronic properties of the prototypical PTCDA/Ag(111) interface. For doping K atoms are used, as K$_x$PTCDA/Ag(111) has the distinct advantage of forming well-defined stoichiometric phases. To arrive at a conclusive, unambiguous, and fully atomistic understanding of the interface properties, we combine state-of-the-art density-functional theory calculations with optical differential reflectance data, photoelectron spectra, and X-ray standing wave measurements. In combination with the full structural characterization of the K$_x$PTCDA/Ag(111) interface by low-energy electron diffraction and scanning tunneling microscopy experiments (ACS Nano 2016, 10, 2365−2374), the present comprehensive study provides access to a fully characterized reference system for a well-defined metal−organic interface in the presence of dopant atoms, which can serve as an ideal benchmark for future research and applications. The combination of the employed complementary techniques allows us to understand the peculiarities of the optical spectra of K$_x$PTCDA/Ag(111) and their counterintuitive similarity to those of neutral PTCDA layers. They also clearly describe the transition from a metallic character of the (pristine) adsorbed PTCDA layer on Ag(111) to a semiconducting state upon doping, which is the opposite of the effect (degenerate) doping usually has on semiconducting materials. All experimental and theoretical efforts also unanimously reveal a reduced electronic coupling between the adsorbate and the substrate, which goes hand in hand with an increasing adsorption distance of the PTCDA molecules caused by a bending of their carboxylic oxygens away from the substrate and toward the potassium atoms.

KEYWORDS: metal−organic interface, doping, density-functional theory calculations, X-ray standing wave, differential reflectance spectroscopy, electronic structure, optical properties

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doping can be exploited to tune the spin properties of a hybrid ferromagnetic metal—organic interface. The charge transfer between alkali metal atoms and organic molecules also strongly impacts the electronic properties of the latter. This is highly relevant for carrier injection in electronic devices, as the energy level alignment at the interfaces between the metal electrode and the active organic layer can be tuned by changing the stoichiometry of the dopant–host network. Consequently, alkali-metal doping has been shown to increase the efficiency of organic light-emitting devices. The doping-induced charge transfer also significantly influences the optical properties of organic molecules, where the nature of the formed intragap states depends on the amount of induced charges rather than the type of the dopant used.

Understanding the effect of alkali-metal doping on organic monolayers strongly interacting with metal substrates is particularly challenging. The reasons for that are twofold: First, the structure of the organic adsorbate layer can change significantly in the presence of alkali metal atoms, which can have a crucial impact on both the electronic and optical properties of the interface. Second, one is dealing with a complex interplay of charge transfer and potentially also covalent interactions occurring between the dopants and organic molecules, the dopants and the substrate, and the substrate and the adsorbed organic molecules. In fact, under certain circumstances alkali-metal atoms have been found to decouple adsorbate molecules from metal substrates, and for the sexiphenyl/Cu(110) interface at low doping concentrations they even result in a depopulation of formerly occupied states. As a consequence, to arrive at a conclusive picture, it is crucial to combine a variety of experimental techniques that provide information on the structural, optical, and electronic properties of the studied system with state-of-the-art quantum-mechanical calculations. The simulations, on one hand, allow an unambiguous interpretation of the experimental observations and, on the other hand, provide microscopic insight into charge-density rearrangements occurring at the interface.

K-doped perylene-3,4,9,10-tetracarboxylic diimide (PTCDA) on Ag(111) is a particularly well-suited system for studying the impact of doping on a molecular monolayer in the presence of interfacial charge transfer with a metal substrate. The reasons for this are (i) that PTCDA on Ag(111) is undisputedly the best investigated interface between a noble metal surface and an organic semiconductor molecule and (ii) that only specific phases of K$_x$PTCDA with well-defined stoichiometries exist on Ag(111), as known from a previous study. At comparably low K-doping levels (in the following referred to as K$_x$PTCDA/Ag(111)), the well-known herringbone structure of PTCDA on Ag(111) spontaneously converts into another highly ordered phase with two K atoms and one PTCDA molecule per adsorbate unit cell. The latter is characterized by a point-on-line (POL) epitaxial relation to the substrate. The potassium atoms are located between the carboxylic and anhydride oxygen atoms, as shown in the structural model in Figure 1b. At higher doping levels, a second phase transition occurs and the resulting structure is characterized again by rows of K atoms arranged between rows of PTCDA molecules. However, in this structural phase, the molecules are oriented essentially perpendicular to the direction of the potassium rows, and there are four potassium atoms per molecule (thus, this phase will be referred to as K$_x$PTCDA/Ag(111)). No intermediate phases (such as K$_x$PTCDA or K$_x$PTCDA$^-$) have been observed, which significantly eases the interpretation of the obtained spectra. Rather, at intermediate doping levels the above-mentioned stable phases coexist. Only at even higher potassium content has another well-ordered phase been detected, which is, however, unstable and degrades within an hour at room temperature by segregation of K, thereby forming the K$_x$PTCDA/Ag(111) phase with a relatively high defect concentration.

Here we present an in-depth study on how these structural and chemical changes of the adsorbate layer affect its optical and electronic properties. We start with a detailed discussion of the optical properties of the metal–organic interface obtained...
by differential reflectance spectroscopy (DRS) experiments. For the systems studied here, these, however, yield ambiguous results and do not allow arriving at a conclusive understanding of how the electronic structure of the interface changes upon K doping. To properly understand the observed trends, we therefore performed state-of-the-art simulations relying both on open boundary conditions for free molecules and on periodic ones for adsorbed PTCDA. These are combined with photoelectron spectroscopy and normal incidence X-ray standing wave (NIXSW) studies to obtain the energy level alignment of the occupied frontier orbitals and vertical adsorption distances, respectively. In this way, we are finally able to unambiguously assign the observed optical features and to achieve an in-depth atomistic understanding of the electronic and optical properties of the prototypical interfaces studied here.

RESULTS AND DISCUSSION

Optical Properties. Monolayer films of pristine and K-doped PTCDA were grown under ultra-high-vacuum conditions, as described in ref 19. During the growth of PTCDA and the subsequent deposition of K we performed DRS experiments in real time to determine the optical properties of the samples.28 From the DRS raw data, the imaginary part of the dielectric function of the adsorbate layer \( \varepsilon_{ads}^{\prime\prime} \) was extracted as described in the Methods section and, in greater detail, in the Supporting Information.

The \( \varepsilon_{ads}^{\prime\prime} \) spectra for PTCDA on Ag(111) at various K doping stages are shown in Figure 2 together with the results of previous DRS investigations for PTCDA on Au(111) and on mica.29,30 The latter are employed here as reference spectra, which can be used to identify the spectral features of the K-doped PTCDA films. The spectrum of the pristine PTCDA layer on Ag(111) (black dashed curve in Figure 2a) is significantly broadened and clearly red-shifted compared to that of pristine PTCDA on mica (dashed olive-green curve in Figure 2c). The latter is dominated by largely unperturbed molecular excitations comparable to those of PTCDA dissolved, for example, in dichloromethane \(^1\) or isolated in a xenon matrix.32 The broadening and shift of the spectrum on Ag(111) can be interpreted as an indication of a strong (chemical) interaction between the adsorbate layer and the substrate. It is well established that this results in a significant charge transfer from the Ag substrate to the PTCDA molecules and that, consequently, the band derived from the LUMO of PTCDA becomes partially filled \(^{24,26,33} \) (vide infra). In that context it is interesting to point out that the maximum of \( \varepsilon_{ads}^{\prime\prime} \) for PTCDA/Ag(111) in the displayed energy range is found at essentially the same energy as for the PTCDA monoanion on mica (dotted gray curve in Figure 2c).30 This fact will be the subject of a detailed discussion later in this paper.

Upon depositing K, the optical features sharpen and shift to higher energies with the strongest peak fitted at 2.19 eV and a low-energy shoulder remaining at 1.73 eV. That shoulder is essentially at the same energy as the peak for PTCDA/Ag(111). For an intermediate doping regime corresponding to \( K_{2}\text{PTCDA} \) (solid red curve in Figure 2a), the spectrum strongly resembles a superposition of the PTCDA on the Ag(111) spectrum and that of the PTCDA monolayer on Au(111) (solid orange curve in Figure 2c). Interestingly, for the latter it is well established that the adsorbate layer is essentially charge neutral with no charge transfer from the substrate.24–26,54 Thus, the evolution of the optical spectra at intermediate doping levels seems to imply that the deposited potassium oxidizes rather than further reduces the PTCDA layer, which contradicts chemical intuition. Exactly this behavior (namely, a reduction of the electron density on the adsorbate due to alkali-metal deposition) has, however, been observed by angle-resolved ultraviolet photoelectron spectroscopy (ARUPS) at low Cs-doping concentrations for the sexiphenyl/Cu(110) interface due to a Cs-induced decoupling of the adsorbate from the substrate.21

For the present interface the scenario of a K-induced oxidation is called into question by the evolution of the spectra upon additional K deposition depicted in Figure 2b: For \( K_{2}\text{PTCDA} \) (solid blue curve) one observes a further blue-shift and sharpening of the spectral features with a clearly resolved double-peak structure of the main maxima at 1.80 and 2.29 eV and their corresponding vibronic replica on the high-energy sides (vide infra). The final spectrum is strongly reminiscent of that of the PTCDA dianion on mica (solid blue curve in Figure 2c).30 This observation would be indicative of the \( a \ priori \) expected further reduction of the PTCDA layer by K doping (a trend also observed when increasing the Cs dose for the above-mentioned sexiphenyl/Cu(110) interface).21 Considering the increasingly sharper features upon K deposition, this chemical
reduction would actually go hand in hand with an electronic decoupling of the adsorbate layer from the Ag(111) substrate.

To resolve the conundrums arising from interpreting the DRS data (viz., the reduction vs oxidation of the PTCDA layer on Ag(111) upon K deposition) and to unambiguously identify the actual electronic structure of the interface, we performed simulations based on dispersion-corrected density-functional theory employing both periodic boundary conditions (when modeling extended interfaces) and open boundary conditions (when simulating molecular excitations). These simulations are backed up by additional ultraviolet photoelectron spectroscopy (UPS) and NIXSW experiments.

Density of States. A first quantity to analyze in order to understand how charges are rearranged upon K doping is the calculated total density of states projected onto the adsorbate layer (i.e., the PDOS), for the different cases (see Figure 3a–d). Figure 3a shows the PDOS of gas phase PTCDA; the PDOS of PTCDA upon adsorption on Ag(111) can be seen in Figure 3b.

The partial filling of the LUMO-derived band is clearly observed in the simulations and is also confirmed by the UPS experiments (Figure 3e). This indicates a metallic nature of the adsorbate layer (vide supra). Note that in the following we will label bands by the names of the orbitals in the neutral molecule that they are derived from. This is done to avoid confusions arising from using the same names for symmetry-inequivalent bands just as a consequence of different degrees of charging.

Interestingly, especially the LUMO, LUMO+1, and LUMO+2 derived bands for the PTCDA/Ag(111) interface broaden significantly compared to the gas phase molecule. The resulting peak width is a measure for the level of hybridization of the electronic states of the adsorbed molecules with the substrate continuum and for the coupling between molecules in the adsorbate layer (note that no vibronic finestructure is considered in the present calculations). A complication when quantifying the peak widths arises from the fact that all plotted DOSs contain an extrinsic Gaussian broadening, which is necessary to avoid artifacts due to the finite number of Ag layers in the slabs (instead of using a semi-infinite model for the Ag surface) and due to the finite k-point sampling (cf., Methods section). Nevertheless, applying the procedure described in detail in the Supporting Information, one can obtain an estimate for the intrinsic line width. In this way, for the LUMO-derived band in PTCDA/Ag(111) in Figure 3b a value of 0.26 eV is obtained for the FWHM.

The electron transfer from the substrate to the adsorbate for PTCDA/Ag(111) is also confirmed by the net charges residing on the PTCDA layer as derived from both a Löwdin and a Bader-type charge partitioning, the latter determined by means of a different plane-wave code in ref 19. Both approaches provide equivalent results, namely, the transfer of approximately one electron per PTCDA molecule (see Table 1). The seeming discrepancy between this value and the significantly above 50% filling of the LUMO-derived band (Figure 3) can be resolved by considering that at interfaces between acceptors bearing functional groups that strongly interact with the substrate one observes a delicate balance between charge forward and backward donation. In such a situation, the transfer of electrons to the LUMO is partially compensated by a back-transfer of electrons from the functional groups to the substrate involving strongly hybridizing molecular orbitals. This effect has been described previously also for the PTCDA/Ag(111) interface.

Another observation for PTCDA adsorbed on the Ag(111) surface is that the peak-to-peak HOMO–LUMO gap is reduced by 0.4 eV compared to an isolated PTCDA molecule.

| Table 1. Löwdin Charges (in Units of the Elementary Charge, e) of the PTCDA/Ag(111) Interface at Various Stages of Doping |
|------------------|------------------|------------------|
| System           | \(\rho_{\text{HOMO}}/e\) | \(\rho_{\text{LUMO}}/e\) | \(\rho_{\text{LUMO}^+}/e\) |
| PTCDA/Ag(111)    | \(-1.1\) (-1.1)   | \(+1.1\) (+1.1)   |
| \(K_2\)PTCDA/Ag(111) | +0.7 (+0.8)  | -1.3 (-1.7)  |
| \(K_4\)PTCDA/Ag(111) | +0.6 (+0.8) | -1.4 (-1.9) |

*Values for potassium are given per K atom; those for PTCDA and Ag per PTCDA molecule. As partitioning into atomic and molecular charges is not an unambiguous process, we also report (in parentheses) values obtained by the alternative partitioning into Bader charges (from the Supporting Information of ref 19).*
(Figure 3a). This is not due to screening effects at the metal surface, as these are not accounted for when merely looking at Kohn–Sham energies at the PBE level. It is also not related to the monolayer formation, as for a hypothetical free-standing monolayer we observe the same gap as for the isolated molecules (again bearing in mind that screening for excited states is not included in a ground-state calculation). Rather, the reduction of the gap can be interpreted as a consequence of the strong coupling between electronic and nuclear degrees of freedom in conjugated organic materials and the resulting molecular distortions in the charged molecules.40

Upon K doping all features in the calculated PDOS shift to higher binding energies and the LUMO-derived band becomes completely filled, resulting in no states at the Fermi level. This means that there is a metal-to-semiconductor phase transition of the PTCDA monolayer adsorbed on Ag(111) upon K doping, a phenomenon commonly observed when interfaces containing organic semiconductors are doped with alkali metals.13,18,41 This is consistent with the corresponding charges on the molecule (Table 1), which clearly show that the electron density on the molecule is further increased. The extent to which this is also a consequence of a reduced back-donation from deeper-lying orbitals involving the carboxylic oxygens is difficult to quantify, but the changes in adsorbate geometry suggest that this effect is also of some relevance. Interestingly, the width of the LUMO-derived peak in the DOS is significantly reduced to 0.14 eV for K2PTCDA/Ag(111), which amounts to nearly a factor of 2 compared to PTCDA/Ag(111) (see Supporting Information for more details) (Figure 3c).

Notably, a shift to higher binding energies and a complete filling of the LUMO-derived band is also seen in the experimental UPS data for K2PTCDA (Figure 3e). A similar observation has in fact been made in UPS studies of PTCDA on the Ag(110) surface upon potassium deposition in spite of the fundamentally different interface geometry at that metal surface, with K atoms substituting Ag atomic rows below the oxygens of the PTCDA molecule.42

For K2PTCDA/Ag(111) the trends already observed for K2PTCDA/Ag(111) prevail both in the simulations and in the experiments, with the shift toward higher binding energies being somewhat larger. Concomitantly, the net charge on the PTCDA molecules increases (Table 1). Also the peaks in the PDOS sharpen further, and the intrinsic line width of the LUMO-derived feature is reduced by another factor of 2 to 0.07 eV (see Supporting Information). At the same time, the splitting between the LUMO+1 and the LUMO+2 increases compared to lower doping stages. The scenario of charge transfer depicted here is additionally validated by the excellent agreement between the experimental and calculated changes of the surface work function with doping (vide infra).

The main conclusion from the above considerations is that at none of the doping stages is a K-induced reduction of the electron density in the PTCDA layer observed, as had been tentatively assumed based on the similar positions of the main peaks in the DR spectra of K2PTCDA/Ag(111) and PTCDA/Au(111) layers (vide supra). Thus, to reconcile the calculated PDOS and measured UPS data with the results of the DRS measurements, a more in-depth discussion of the optical properties of pristine and doped PTCDA layers is required.

**Simulated Optical Properties of Molecules and Adsorbate Layers.** Ideally, the simulations of the optical properties should account for the 2D extended interfaces including the metal substrate. For that, slab-type calculations employing periodic boundary conditions would be required. At the same time excitonic effects need to be accounted for, as they are highly relevant in molecular layers. Indeed, simulations including the necessary many-body effects also for infinitely extended crystalline systems are generally possible. They rely, for example, on solving the Bethe–Salpeter equation (BSE) and have been performed for molecular and polymer crystals.53–56 Recently, even implementations of the BSE for adsorbed molecules have been developed,47 building, however, on system-specific assumptions that do not apply here (e.g., that the adsorbate’s HOMO lies in the band gap of a semi-conducting substrate).

Consequently, BSE-type calculations on interfaces as complex as the present one are still beyond reach. Therefore, we resorted to a dual strategy: We first calculated optical properties including excitonic effects for neutral and charged PTCDA molecules in the gas phase using time-dependent density functional theory (TD-DFT). There, we tried to correlate the properties of the excitons with those of the molecular orbitals that dominate their description. Provided that the exciton is primarily represented by the excitation from a specific occupied to a specific unoccupied orbital, the optical properties of the interface can be directly correlated with the evolutions of the calculated densities of states. As a second step, we also simulated excitation spectra based on the independent particle approximation relying on the calculated Kohn–Sham orbitals of the extended interfaces. In these calculations the interaction with the substrate and with dopant atoms is directly considered, but excitonic effects are neglected.

Following that strategy, we report the molecular excitation energies and excited-state properties (obtained using TD-DFT) in Table 2. The corresponding orbital energies are contained in the Supporting Information together with the isodensity plots of the orbitals. Several relevant insights can be gained from those data: (i) The main absorption features of the neutral PTCDA molecule and the dianion are found at essentially the same energies (2.17 eV vs 2.25 eV). This occurs in spite of the fact that they are dominated by different single-particle excitations (the HOMO → LUMO transition in the neutral case and a transition from the now doubly occupied LUMO to the LUMO+2 in the dianion). Consequently, the similarity of the transition energies and oscillator strengths for those peaks are a mere coincidence. Therefore, the similarity of the peak positions in the measured optical absorption data of K2PTCDA/Ag(111) and PTCDA/Au(111) discussed above do not necessarily imply that the two films are in the same charge state. (ii) In the dianion there is a weaker peak at ~0.5 eV below the maximum. This peak is dominated by a LUMO to LUMO+1 excitation and can be associated with the lower energy maximum in the measured ε′′ spectra slope for K-doped PTCDA films in Figure 2. Indeed, the oscillator strength associated with this peak is almost an order of magnitude smaller than that of the LUMO to LUMO+2 excitation contributing to the main maximum. Simulations on a cluster consisting of a PTCDA molecule asymmetrically coordinated with two Na atoms yielded similar results (albeit with modified indices for the orbitals and excited states due to the explicit consideration of the dopants).48 We also calculate the most prominent Raman-allowed vibrational mode of the PTCDA dianion, which is at 0.20 eV. This is almost exactly the energy by which the vibronic replicas are shifted from the main peaks in the experimental spectra for the dianion on mica and for
Table 2. TD-DFT-Calculated Properties of the Lowest-Lying Optically Allowed Excited States of PTCDA Molecules at Various Charging Stages

<table>
<thead>
<tr>
<th>PTCDA</th>
<th>state</th>
<th>energy (eV)</th>
<th>oscillator strength</th>
<th>dominating contributions including their weights</th>
<th>orbital gap (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>neutral</td>
<td>S₁</td>
<td>2.17</td>
<td>0.48</td>
<td>0.71 H→L</td>
<td>1.50</td>
</tr>
<tr>
<td>monoanion</td>
<td>S₁</td>
<td>1.57</td>
<td>0.03</td>
<td>0.99 L₁→L₁</td>
<td>1.42</td>
</tr>
<tr>
<td></td>
<td>S₁</td>
<td>2.04</td>
<td>0.53</td>
<td>0.82 L₂→L₂</td>
<td>1.53</td>
</tr>
<tr>
<td></td>
<td>S₁</td>
<td>1.82</td>
<td>0.07</td>
<td>0.56 H₁→L₁</td>
<td>1.31</td>
</tr>
<tr>
<td>dianion</td>
<td>S₁</td>
<td>2.25</td>
<td>0.59</td>
<td>0.70 L→L+1</td>
<td>1.41</td>
</tr>
<tr>
<td></td>
<td>S₁</td>
<td>1.82</td>
<td>0.07</td>
<td>0.70 L→L+2</td>
<td>1.57</td>
</tr>
</tbody>
</table>

As discussed in the main text, H and L refer to the orbitals that have the same symmetry as the HOMO and LUMO of the neutral molecule (consequently, L for the dianion refers to the highest doubly occupied orbital), while L+1 and L+2 refer to the orbitals that have the same symmetry as the second and third unoccupied orbital. Importantly, the orbitals with a certain label have essentially the same shapes at different doping levels. The subscripts α and β refer to the two spin manifolds. There is no significant change in shape and, more importantly, no change in their order. The observation that the S₁ and S₁ coefficients in the description of the excitations in the neutral and dianion case appear to be 1/√2 smaller than S₁ for the anion is just a consequence of the fact that for a spin-restricted calculation a normalization factor of 1/√2 is included in the description of the singlet state in the Gaussian output. This is not the case for the spin-unrestricted calculation of the anion. In the characterization of the excited states only transitions with weights exceeding 0.13 are included.

K₄PTCDA/Ag(111) (see Figure 2). (iii) In the monoanion, the most prominent optical excitation (again dominated by a LUMO → LUMO+2 transition) is somewhat red-shifted compared to the neutral and dianion cases. This is reminiscent of the situation of PTCDA on Ag(111) in the absence of the potassium atoms. The calculated shift is, however, smaller than in the experiment. Some deviations between simulations and experiments are not surprising though, bearing in mind that upon adsorption on Ag(111) the PTCDA LUMOs are more than half-filled (i.e., the actual situation goes beyond monoanion formation). Moreover, on the surface the filling of the LUMO is partially compensated by a back-transfer of charges from the carboxylic oxygens (vide supra). None of these effects are captured in the calculations of the isolated monoanion.

Interestingly, all relevant optically allowed excited states in the neutral PTCDA molecule and the dianion (i.e., the systems most relevant for the present study) are dominated by individual single-particle excitations. Therefore, the properties of the excited states can be immediately associated with those of specific orbitals. The observation that the absolute values of the Kohn–Sham orbital gaps are much smaller than the excited-state energies can be traced back to the self-interaction error and lack of the derivative discontinuity in (semi)local functionals and has no bearing on the qualitative conclusions drawn here. What is important are the observed trends. For example, the HOMO → LUMO gap in the neutral PTCDA molecule (also reported in Table 2) is only 0.07 eV smaller than the LUMO → LUMO+2 gap in the dianion, which is fully consistent with an energetic difference of 0.08 eV between the respective states (S₁ for the neutral molecule and S₁ for the dianion).

To link these results to the calculated densities of states discussed in the previous section, it is worth pointing out that several trends observed in the band-structure calculations of the interfaces (Figure 3) are directly reflected in the shifts of molecular orbital energies (Table S1) (recalling that PTCDA is negatively charged by Ag(111) even without K). The HOMO → LUMO gap is strongly reduced upon charging both in the interface simulations and in the molecular calculations. Conversely, the LUMO+1 → LUMO+2 splitting increases upon K doping on the surface and likewise upon charging the individual molecules (cf. Figure 3 and Table S1). These correlations indicate that trends derived from single-particle densities of states of extended interfaces can be directly correlated with excitation processes within the PTCDA adsorbate layer. Consequently, it is also useful to calculate optical spectra (within the independent particle approximation) from the bands of the extended interfaces taking the hybridization of the molecular and substrate states into account.

The corresponding spectra obtained within the independent particle random phase approximation (IP-RPA) are shown in Figure 4.
the spectrum of PTCDA/Ag(111) (black curve in Figure 4) is characterized by a peak at 1.4 eV (1.75 eV in the experiments, Figure 2a). This peak narrows and shifts to higher energy for K₂PTCDA/Ag(111) (red curve in Figure 4). Both trends are consistent with the experimental observations (see Figure 2a) and prevail for K₄PTCDA/Ag(111) with a slightly larger shift of the main feature (see blue curve in Figure 4). Also the experimentally observed splitting into a weaker low-energy feature and a stronger high-energy feature for K₂PTCDA/Ag(111) and K₄PTCDA/Ag(111) is reproduced in the simulations. In passing we note that a fully quantitative agreement between theory and experiments is not expected within the independent particle approximation, and also the vibronic fine-structure observed in the experiments is not contained in simulations focusing on electronic excitations.

The decreasing line width of the spectral features after K doping can be directly traced back to the decreasing line widths of the relevant DOS peaks discussed in the previous section. The most relevant bands in this context are the ones derived from the molecular LUMO and LUMO+2 for the main peak and, additionally, the LUMO+1-derived band for the low-energy tail of the main feature.

These considerations allow us to unambiguously associate the DR spectra at various K-doping stages with specific charging states of the adsorbate layer. Particularly, they show that the similar peak positions for K₂PTCDA/Ag(111) and PTCDA/Au(111) are a coincidence rather than an indication for a reduction of the charge of the PTCDA layer due to K deposition. The reduced width of the spectral features upon K doping is an indication for a reduced coupling to the substrate, an aspect that shall be addressed more thoroughly in the following sections.

**Coupling between the PTCDA Layer and the Ag(111) Substrate at Various Stages of K-Doping: Band Structures and Adsorption Heights.** Given that optical transitions do not mix states with different k-vectors, studying the actual band structures of the interfaces provides insight beyond merely considering the densities of states. The calculated bands for PTCDA/Ag(111), K₂PTCDA/Ag(111), and K₄PTCDA/Ag(111) are shown as thin lines in Figure 5. Superimposed is a k-resolved density of states projected onto atom-centered functions of the molecules (KDOS(k, E)), in analogy to the evaluation of Löwdin charges. That quantity is highest for (k, E) values corresponding to states localized in the molecular regions; that is, it allows us to identify molecular contributions to the actual band structure. The KDOS could also be plotted by coloring each point in the band structure depending on the square of the wave function amplitude in the molecular region of the adsorbate layer. This would, however, generate a somewhat involved picture considering the fairly large number of bands due to using a slab of a finite thickness to represent the metal substrate. This complication can be avoided by representing the KDOS in Figure 5 as a color map adopting a Gaussian broadening for each state using a width of 0.11 eV, i.e., the same broadening that was applied when plotting the DOS in Figure 3 (for details see Methods section).

For PTCDA/Ag(111) one observes a massive energetic broadening of the PTCDA-derived LUMO (somewhat below E_F), LUMO+1, and LUMO+2 states. This is indicative of the strong hybridization with substrate states at virtually all k-points of the surface Brillouin zone. The hybridization is superimposed with a significant dispersion of the main features (red and yellow regions) of the bands, with the combination of both effects giving rise to the large width of the DOS features. For K₂PTCDA, the dominant KDOS features are much more clearly resolved in energy (see the increase in red regions in panel (b) compared to panel (a)). Note also the reduced energetic broadening indicating a reduced hybridization with the substrate. In K₄PTCDA/Ag(111), the molecule-derived bands become essentially flat, with the effect being most pronounced for the LUMO+1- and LUMO+2-derived states. This explains why in this system especially the LUMO+1- and LUMO+2-derived DOS peaks are well resolved and clearly separated (see also Figure 3). All these results confirm the reduced interaction between PTCDA and Ag due to K doping.

A difference in molecule–metal interaction should also be reflected in the vertical adsorption geometry of the adsorbate layer and, particularly, in the adsorption heights of the individual atomic species of the K₂PTCDA structure. The latter can be determined by NIXSW experiments with very high precision and chemical sensitivity. Details of the NIXSW experiments and the data analysis are discussed in the Methods section and the Supporting Information. The key results are illustrated in a structural model of the vertical adsorption geometry in Figure 6 together with the respective calculated values. All experimentally and theoretically determined adsorption heights for K₂PTCDA, K₄PTCDA, and literature data for PTCDA/Ag(111) are summarized in Table 3.

Before discussing the resulting model in detail, it is worthwhile mentioning that measured and calculated heights match well. When comparing the numbers, a general...
observation is that the calculated adsorption heights are consistently slightly smaller than the experimental ones. We attribute this at least in part to an inaccurate description of the vertical relaxation of the topmost Ag layer, which for K₂PTCDA/Ag(111) and K₄PTCDA/Ag(111) also arise from the modifications we had to make to the lattice constants to reconcile the point-on-line growth with periodic boundary conditions (see ref 19 and the Methods section). This Ag relaxation is indeed relevant here, as the true interaction is determined by the actual interatomic distances, while the quantities measured in NIXSW are the distances relative to the lattice planes of the Bragg reflection used, i.e., effectively relative to the (hypothetical) unrelaxed Ag(111) surface. An incorrectly modeled surface relaxation will, therefore, cause minor deviations between measured and calculated heights. Only for the K atoms do the deviations appear too large to arise from such effects. To test to what extent the very high experimental K position is a consequence of varying K-adsorption positions arising from point-on-line growth, we compared K atoms adsorbed on top and in hollow positions as the two limiting cases. There we, however, found differences in adsorption heights of only 0.02 Å. Another reason for the deviations could be that the adsorption heights measured for K₂PTCDA might be slightly influenced by a small amount of coexisting K₄PTCDA (see Supporting Information for more details) in such a way that the obtained values are systematically too large (cf. Table 3).

Independent of the above-mentioned deviations, the trends in the main geometrical changes upon K doping are consistent for simulations and experiments: (i) K deposition clearly increases the adsorption height of the perylene core of PTCDA above the Ag substrate, which again suggests a weakening of the coupling between the substrate and the adsorbate layer. (ii) While for PTCDA/Ag(111) especially the carboxylic oxygens bend toward the surface indicative of an additional bonding channel to the Ag substrate, upon K doping they are found at positions significantly above the perylene carbons. Interestingly, the anhydride oxygens also bend away from the surface upon K doping such that they come to lie above the K atoms. Note that the adsorption heights of the oxygen atoms are quantitatively the same for both oxygen species, but the differences in adsorption heights between O_{anhy} and O_{carb} are significantly smaller than in the case of PTCDA/Ag(111). The reasons for the upward bending will become apparent when analyzing the interfacial charge rearrangements (vide infra). Upon increasing the K concentration (i.e., for the K₂PTCDA/Ag(111) system), all K-induced geometrical changes are enhanced in the calculations (no experiments are available for that doping stage).

In passing we note that the rather strong lifting of the perylene core away from the Ag surface upon K doping is in sharp contrast to the situation observed upon exposing the PTCDA/Ag(110) interface to K atoms, where the measured adsorption height increases by only 0.05 Å.²¹ We attribute that difference to a significantly altered bonding geometry, as on Ag(110) the K atoms lie below the PTCDA molecules replacing Ag atoms in the topmost surface layer, while in the present case they lie above the top Ag layer.¹⁰ The latter results in a distinct upward bending of the carboxylic oxygens so that they lie 0.24 Å (0.31 Å) above the perylene core in the NIXSW experiments (DFT calculations). Importantly, the decoupling of the PTCDA molecules from the substrate upon K deposition is not a consequence of any tilting of the molecule, as has been observed when subjecting a 6P monolayer on Cu(110) to Cs doping.²² This can be inferred from both the STM images

Table 3. Calculated (Regular) and Measured (Bold) Average Adsorption Heights for Different Chemical Species in PTCDA/Ag(111), K₂PTCDA/Ag(111), and K₄PTCDA/Ag(111)*

<table>
<thead>
<tr>
<th>System</th>
<th>C_{pery} (Å)</th>
<th>C_{func} (Å)</th>
<th>O_{anhy} (Å)</th>
<th>O_{carb} (Å)</th>
<th>K (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTCDA/Ag(111)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>theory</td>
<td>2.73 (2.82)</td>
<td>2.71 (2.80)</td>
<td>2.75 (2.84)</td>
<td>2.58 (2.66)</td>
<td></td>
</tr>
<tr>
<td>experiment</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>K₂PTCDA/Ag(111)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>theory</td>
<td>2.85 (2.98)</td>
<td>3.10 (3.21)</td>
<td>3.16 (3.29)</td>
<td>3.17 (3.30)</td>
<td>2.74 (2.87)</td>
</tr>
<tr>
<td>experiment</td>
<td>3.12 (±0.02)</td>
<td>3.26 (±0.07)</td>
<td>3.36 (±0.05)</td>
<td>3.36 (±0.07)</td>
<td>3.23 (±0.03)</td>
</tr>
<tr>
<td>K₄PTCDA/Ag(111)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>theory</td>
<td>3.04 (3.10)</td>
<td>3.34 (3.41)</td>
<td>3.43 (3.50)</td>
<td>3.45 (3.51)</td>
<td>2.72 (2.79)</td>
</tr>
</tbody>
</table>

*All values are given relative to the (hypothetical) unrelaxed Ag(111) surface, as this is the quantity measured in X-ray standing wave experiments. Additionally, in parentheses we report the calculated distances relative to the relaxed top Ag layer. The values given in parentheses along with the measured adsorption heights are the experimental uncertainties. C_{pery} denotes the C atoms in the perylene core, C_{func} are those attached to the O atoms, O_{anhy} denotes the anhydride oxygens, O_{carb} are the carboxylic oxygens, and K are the potassium atoms. The adsorption heights of the hydrogen atoms were not measured. These distances are ca. 0.1 Å smaller than the ones reported in ref 62. This is mostly due to the fact that we here considered a five-layer Ag slab in which the two top layers were relaxed, while in ref 62 only three layers were considered, where just one was allowed to change its position during geometry relaxation. When repeating our geometry optimizations with that reduced-size model system, deviations between our calculations and those in ref 62 are at the most 0.04 Å; thus, the main reduction can be explained by a modified relaxation of the Ag surface. *Taken from ref 61; see also ref 60.

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and the high coherent fractions of the C atoms (see Supporting Information) in the NIXSW experiments.

To determine to what extent the reduced widths of the DOS features in K$_2$PTCDA/Ag(111) compared to PTCDA/Ag(111) are a consequence of geometric deformations and to what extent they are caused by the electronic interactions with the K atoms, we studied a model structure with all atoms in the same coordinates as in K$_2$PTCDA/Ag(111), but removed K. The resulting band structure and DOS, as well as additional intermediate models, are presented in the Supporting Information and testify that K atoms act both structurally by raising the molecule and electronically by weakening PTCDA−Ag and PTCDA−PTCDA interactions.

Charge Rearrangements. To better understand the interaction of the PTCDA layer with the K atoms, it is useful to analyze charge rearrangements, $\Delta \rho$, upon K doping. Here it should be noted that the purpose of the following gedanken experiment is not to mimic the actual growth process, but rather to analyze the way charges are redistributed due to the doping of K atoms into the PTCDA film on Ag.

$\Delta \rho$ is given by the difference of the charge densities of the interacting system and the subsystems (all in the geometry of K$_2$PTCDA/Ag(111)):

$$\Delta \rho = \rho(K_{2PTCDA}/Ag(111)) - \rho(PTCDA/Ag(111)) - \rho(K)$$

(2)

Here, $\rho(K_{2PTCDA}/Ag(111))$, $\rho(PTCDA/Ag(111))$, and $\rho(K)$ denote the charge densities of the K-doped PTCDA layer on Ag(111), the adsorbed system upon removing the K atoms, and the removed K atoms, respectively. An alternative scenario, where we first consider the formation of a hypothetical free-standing K$_2$PTCDA layer, which eventually adsorbs on the Ag(111) substrate, is discussed in the Supporting Information.

The charge rearrangements for the formation of the PTCDA/Ag(111) interface (with a herringbone arrangement of the molecules) have already been discussed extensively in the literature (e.g., in ref 49). There one primarily observes an electron transfer into the $\pi$-electron system of the PTCDA layer accompanied by an electron density reduction above the metal and in the $\sigma$-electron system, with the latter effect strongest in the region of the carboxylic oxygens. This is indicative of the charge back-donation process described earlier.

A similar situation is obtained when arranging the molecules on Ag(111) in the geometry they finally adapt in the K$_2$PTCDA/Ag(111) layer (see Supporting Information).

When adding the K atoms (cf., Figure 7a) to the adsorbed PTCDA layer, one observes the additional donation of electrons into the $\pi$-system of the PTCDA molecules especially in the molecular regions close to the K atoms. This is consistent with a further filling of the LUMO upon K doping. Additionally, one sees electron density accumulation above the topmost Ag layer especially directly below the K atoms. An interesting observation is that the most significant electron density depletion is not centered at the K atoms but localized in the region between them. This is also observed when potassium atoms are adsorbed on the bare Ag(111) surface in the absence of the PTCDA layer (see Supporting Information).

An additional feature is the accumulation of electron density on top of the K atoms, which is a common observation for K and heavier alkalis adsorbed on metals.63

Of particular interest are the charge rearrangements in the immediate vicinity of the oxygen atoms: There is electron density accumulation between both the carboxylic and
anhydride oxygens and the K atoms. Concomitantly, electron density is depleted between the oxygens and the neighboring carbon atoms. This is indicative of a modification of the bonding situation of the oxygens with a weakening of the bonds to the backbone of the PTCDA molecules and an interaction with the K atoms that goes clearly beyond a mere charge transfer. This observation reinforces the above conclusion that the direct chemical interaction between the anhydride groups of the PTCDA molecules and the K atoms plays a crucial role for the reduced coupling between PTCDA and Ag(111) upon K doping. Similar trends, albeit with more massive charge rearrangements, are also seen for the K₄PTCDA system (see Figure 7b).

As a final remark, we briefly mention work function changes, as a direct consequence of the charge rearrangements (especially those perpendicular to the metal surface). They are significant, as shown in Table 4 with K-doping triggering a work-function change of up to −1.6 eV. Notably, also for this observable, we obtain an excellent agreement between experiments and simulations, which supports the validity of the calculated charge rearrangements discussed in the preceding paragraphs of this section.

### CONCLUSIONS

The combination of the results from various complementary experimental and theoretical investigations shows that for the prototypical PTCDA/Ag(111) interface extensive geometrical reorderings occur upon potassium doping, which in turn result in massive modifications of the electronic and optical properties of the interface. In particular, K-deposition increases the net electronic charge of the PTCDA molecules, which results in a transition from a metallic to a semiconducting adsorbate layer. This behavior is in sharp contrast to the impact of (degenerate) doping in conventional semiconductors, where doping generates a highly conducting state.

Moreover, K deposition reduces the electronic coupling between the PTCDA layer and the Ag substrate. In the simulations, this is manifested in a reduction of the intrinsic widths of the DOS features associated with the adsorbate layer and a decreased dispersion and hybridization of the adsorbate-projected bands. In the experiments it gives rise to sharper spectral features upon increasing the K content in conjunction with an increasing adsorption distance of the perylene core and a pronounced upward-bending of the carboxylic O atoms. This implies that the strength of the electronic coupling between the molecules and the substrate can be tuned by the potassium dopants. Notably, in spite of the new K–O bonding channels and the associated massive charge rearrangements, we do not observe a stronger dispersion of the bands for K-vectors parallel to the substrate surface.

On more technical grounds we find that the observation that the main peak of the DRS spectrum of K₄PTCDA/Ag(111) is found at essentially the same position as the dominant peak in the spectrum of PTCDA/Au(111) is not a consequence of an actual chemical similarity between the two situations. Rather it is a mere coincidence arising from the specific positions of the unoccupied orbitals of PTCDA. This shows that such comparisons can be very misleading for complex surfaces and that, for situations as involved as the present one, only the combination of a manifold of complementary experimental techniques with state-of-the-art quantum-mechanical calculations allows a consistent, in-depth atomistic understanding of the actual situation. This understanding is, however, a vital requirement for a future rational design of complex metal–organic interfaces.

### METHODS

#### Sample Preparation.

The Ag(111) substrate was prepared by repeated Ar⁺-sputtering and annealing cycles according to ref 64. Each substrate preparation was checked by means of low-energy electron diffraction (LEED) prior to film deposition. The molecules were deposited under ultra-high-vacuum conditions (base pressure 10−10 mbar) from effusion cells with temperature-stabilized (±1 K) crucibles. The deposition rates were on the order of 0.1 ML per minute. The starting point of each doping experiment was a monolayer of PTCDA on Ag(111) with a coverage of 0.6 to 0.7 ML. Potassium was subsequently deposited from a commercially available dispenser source (SAES Getters and Alvacet), while the substrate was kept at room temperature. The evolution of the phases was followed by in situ optical spectroscopy and by distortion-corrected LEED experiments to ensure that the above-mentioned PTCDA/Ag(111), K₂PTCDA/Ag(111), and K₄PTCDA/Ag(111) phases are the only (ordered) ones that occur on the sample.²⁸

#### In Situ Optical Spectroscopy.

From the DRS raw data, shown in the Supporting Information, the imaginary part of the dielectric function of the adsorbate layer ε" was extracted by applying the following expression:²⁸

\[
\varepsilon''(d) \approx R(d) - R(0) \cdot \frac{\alpha^2 (1 - \varepsilon''(d))^2 + (\varepsilon''(d))}{8 \alpha d} \cdot \varepsilon''(d) \cdot \frac{1}{1 - \varepsilon''(d)}
\]

Here, ε"(d) and ε"(0) are the real and imaginary parts of the dielectric function of the bulk substrate. This approximation is valid for film thicknesses d much smaller than the photon wavelengths λ, and under the condition that |ε"(d)| ≪ |ε"(0)| holds (which is fulfilled for silver in the wavelength range considered here).³⁶ R(0) denotes the reference reflectance of the pristine Ag(111) substrate, while R(d) stands for the reflectance measured as a function of the effective PTCDA film thickness d. For the subsequent K doping R and hence ε" change as a function of the number of adsorbed K atoms, which itself is assumed to scale linearly with deposition time. For DRS, an actively cooled 100 W halogen lamp (Osram Xenophot) was operated with a stabilized power supply (Volkraft HPS-11530). The reflected light was spectrally analyzed by means of a monochromator (Acton Research SpectraPro SP2356) with a charge-coupled device attached (Princeton Instruments Spec-10 100BR, liquid-nitrogen-cooled CCD).

#### Photoelectron Spectroscopy Experiments.

XPS and UPS measurements were performed in Jena using a FOCUS 500 monochromatized X-ray (Al Kα emission) and a monochromatized

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**Table 4. Comparison between Experimentally Measured and Simulated Work-Function Changes with Respect to the Bare Substrate, ΔΦ_{KPTCDA}^{Ag(111)} (eV), and Relative to Undoped PTCDA/Ag(111), ΔΦ_{KPTCDA}^{PTCDA} (eV).**

<table>
<thead>
<tr>
<th>K₄PTCDA/Ag(111)</th>
<th>ΔΦ_{KPTCDA}^{Ag(111)} (eV)</th>
<th>ΔΦ_{KPTCDA}^{PTCDA} (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>x = 0</td>
<td>+0.16</td>
<td>-0.9</td>
</tr>
<tr>
<td>x = 2</td>
<td>-0.64</td>
<td>-0.80</td>
</tr>
<tr>
<td>x = 4</td>
<td>-1.42</td>
<td>-1.58</td>
</tr>
</tbody>
</table>

"A complication in the experiments is that the surface coverage is incomplete (d = 0.7 ML). In order to allow for a better comparison to the simulated values, the measured work-function changes were, thus, extrapolated to monolayer coverage (by dividing by 0.7). The overall agreement is excellent, although a significant amount of K not interacting with PTCDA may exist on the samples particularly in the K₄PTCDA film.
UV light source UVLS (He I resonance) combined with a toroidal-mirror monochromator TMM 304. The photoelectrons were analyzed using a PHOIBOS 150 hemispherical energy analyzer with a 3D-DLD detector attached. All parts were supplied by SPECS. The acceptance angle of the detector is ±7°. The Fermi edge region has been fitted by a convolution of the Fermi–Dirac function with a Gaussian representing the instrumental resolution, which was thereby estimated to be ~50 meV. For the determination of work functions, secondary electron cut-offs were measured with the sample biased at ca. –9 V. The UPS spectra were recorded at normal emission. The PTCDA on Ag(111) samples were transferred to the photoelectron spectroscopy setup prior to the potassium deposition using an ultra-high-vacuum shuttle with a base pressure of 10−10 mbar.

**Normal Incidence X-ray Standing Wave Experiments.**

NIXSW experiments utilize the standing wave that forms by the interference of an incident X-ray beam with an outgoing beam that is Bragg diffracted by a single crystal (the substrate). Scanning the photon energy through the Bragg condition shifts the nodes and antinodes of the standing wave in the direction perpendicular to the Bragg planes (in our experiment we use the (111) Bragg planes, which are aligned with the sample surface). This changes the amplitude of the X-ray field at the positions of a specific atomic species in a way that is characteristic for the (vertical) position of that atomic species relative to the nearest Bragg plane. These intensity profiles are parametrized by the coherently fraction $P^H$ and coherency position $P^H$ through

$$Y(E_{ph}) = 1 + R(E_{ph}) + 2 \sqrt{R(E_{ph})P^H \cos(k_{ph}d) - 2E_{ph}P^H}$$

The so-called yield, $Y$, is the intensity of the standing wave field at the species of interest, normalized by the intensity of the incoming beam. $E_{ph}$ is the photon energy of the incoming beam, and $R(E_{ph})$ and $k_{ph}$ are the reflectivity of the sample and the phase of the standing wave field at this energy, respectively. Since $R(E_{ph})$ and $k_{ph}$ follow from the dynamic theory of diffraction, the $E_{ph}$-dependence of $Y$ is purely determined by the parameters $P^H$ and $P^H$, which can thus be obtained by fitting the above equation to experimentally obtained yield curves. These two parameters can then directly be linked to the geometric structure of the sample, since $P^H$ represents the average position of the studied species relative to the nearest Bragg plane, in units of the Bragg spacing. The second parameter, $P^H$, indicates how well-ordered the species is, with a value of 1 meaning that all atoms of this species lie at precisely the same distance from the nearest Bragg plane and a value of 0 typically indicating complete disorder.

In this work, the amplitude of the X-ray interference field at the atomic species of interest was tracked by recording XPS spectra of the C 1s, O 1s, and K 2p photoemission signals as the incident beam energy was scanned through the Bragg condition. Subsequently, these XPS spectra were analyzed with core level models to obtain the yield profiles of individual chemical species using CasaXPS 2.3.16 PR 1.6. The fits of the yield curves were performed by means of TORRICELLI. All measurements were performed at beamline 109 of the Diamond Light Source, using a Scienta R4000 electron analyzer mounted at 90° relative to the incoming X-ray beam. Since in this geometry, nondipolar effects are expected to be minimal, no correction for nondipolar effects has been applied. The films were carefully checked for radiation damage prior to and after each XSW scan. Significant degradation can be excluded at least on the time scale of the experiments. For a more elaborate explanation of the experimental procedure, used core-level models, and the results of this NIXSW analysis refer to the Supporting Information.

**Density-Functional Theory Calculations.**

The optimized adsorption geometries were obtained in our previous work from means of DFT calculations using the Vienna ab Initio Simulation Package (VASP) in conjunction with GADGET. The GGA functional as proposed by Perdew, Burke, and Ernzerhof (PBE) was employed and the van der Waals interactions were included as described by the vdW-09 method. A complication arising from the experimentally observed point-on-line growth is that such a situation is a priori not compatible with the required periodic boundary conditions. This problem can, however, be fixed by choosing suitable supercells in combination with a minute deformation of the substrate (see Figure 1), which for the present system has been tested to have little impact on the obtained results. Consequently, while constructing the supercells for K\textsubscript{3}PTCDA/Ag(111) and K\textsubscript{4}PTCDA/Ag(111) we decided to keep the distances between the molecules at the values measured with high-resolution scanning tunneling hydrogen microscopy (STHM) and distortion-corrected LEED and instead slightly changed the Ag lattice constants in order to arrive at unit cells for which the epitaxial matrix contains only integer entries. For PTCDA/Ag(111), since the monolayer was commensurate with the substrate in the experiments, the PBE lattice constant was used. All slabs consisted of five layers of atoms, where the top two layers were allowed to relax. Please refer to ref 19 for more details of the geometrical optimization.

The structures obtained from VASP were further analyzed using the Quantum ESPRESSO suite, where ultrasoft pseudopotentials generated with the Rappe, Rabe, Kaxiras, and Joannopoulos scheme were used with a plane-wave cutoff of 567.35 eV (27 Ry). The density of states was calculated with Gaussian broadenings of 0.11 and 0.02 eV (FWHM) for PTCDA/Ag(111), K\textsubscript{3}PTCDA/Ag(111), and K\textsubscript{4}PTCDA/Ag(111), by using off-I Monkhorst–Pack k-points grids of $6 \times 6 \times 1$, $10 \times 6 \times 1$, and $6 \times 6 \times 1$, respectively. While analyzing the band structure (see Figure 5), in order to identify the molecular contributions to the fairly large number of bands depicted as thin lines, we superimposed a k-resolved density of states (KDOS) projected onto the atomic states belonging to the molecule: $\text{KDOS}(k, E) = \sum_{\text{species}} \sum_{\text{atom}} \sum_{\text{state}} \delta(E - E_{\text{atom}})$; $\text{atom}$ and $\text{state}$ are the atomic and molecular contributions to the fairly large number of bands, respectively. An additional summation over $k$ in the surface Brillouin zone produces the molecular DOS shown in Figure 3. Analyzing the KDOS is conceptually equivalent to plotting the bands with colors depending on the amplitude on a specific portion of the system, but it allows us to analyze cases characterized by dense bands (or even energy continuum; see ref 72 for an example). In practice, the $\delta$-function in the above equation was replaced by a Gaussian with the same width as used when plotting the DOS (0.11 eV FWHM); since absolute values of the KDOS depend on the broadening, we normalize it to the maximum value assumed by the broadening function (8.3/eV × 2 for spin multiplicity) so that a nondegenerate state $\text{state}$ fully belonging to the molecule has a maximum KDOS($k, E$) of 1. Figures 1 and 7 have been produced by using the program XCrySDen.

In order to calculate the optical absorption spectra of Figure 4 we adopted the Yambo software package to evaluate the independent particle response from PBE Kohn–Sham orbitals. Since Yambo can handle neither ultrasoft pseudopotentials nor PAW potentials, additional calculations of the electronic ground state were performed as a preliminary step using the same atomic positions and k-point meshes, but norm-conserving pseudopotentials (requiring a higher cutoff of 680 eV) were used.

The simulations of individual molecules employing open boundary conditions were conducted using Gaussian09, revision D.01, employing PBE exchange and correlation and a 6-31G(d,p) basis set. Geometries were optimized using standard convergence criteria, and excited-state properties were obtained employing TD-DFT using the same functional and the ground-state geometries.

Finally, we note that in Figure 3 we observe even a quantitave agreement between calculated Kohn–Sham energies and measured ionization energies. While semilocal DFT is typically capable of correctly describing trends, this quantitative agreement between theory and experiment is somewhat surprising considering that when solely considering Kohn–Sham orbital energies one suffers from self-interaction errors and from the fact that screening effects at the surface are neglected. Still, a quantitative agreement between Kohn–Sham orbital energies and UPS features is not uncommon, and we attribute that at least in part to a fortuitous cancellation of the above-mentioned errors for flat-lying adsorbates.
ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.7b05828.

Determination of the intrinsic width of PDOS peaks; plots showing the integrated local density of states; details of molecule-based simulations and those of adsorbed layer with calculated Kohn–Sham orbital energies; explanation of the LUMO splitting; K DOS plots for adsorbed monolayers without K atoms; additional charge rearrangement plots; details of experimental methods; extraction of dielectric function from the DRS; details of NIXSW measurements; comparison of experimental and calculated changes in work function (PDF)

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Notes

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