Electronic structure and excited state dynamics in a dicyanovinyl-substituted oligothiophene on Au(111)

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Dicyanovinyl (DCV)-substituted oligothiophenes are promising donor materials in vacuum-processed small-molecule organic solar cells. Here, we studied the structural and the electronic properties of DCV-dimethyl-pentathiophene (DCV5T-Me2) adsorbed on Au(111) from submonolayer to multilayer coverages. Using a multi-technique experimental approach (low-temperature scanning tunneling microscopy/spectroscopy (STM/STS), atomic force microscopy (AFM), and two-photon photoemission (2PPE) spectroscopy), we determined the energetic position of several affinity levels as well as ionization potentials originating from the lowest unoccupied molecular orbitals (LUMO) and the highest occupied molecular orbitals (HOMO), evidencing a transport gap of 1.4 eV. Proof of an excitonic state was found to be a spectroscopic feature located at 0.6 eV below the LUMO affinity level. With increasing coverage photoemission from excitonic states gains importance. We were able to track the dynamics of several electronically excited states of multilayers by means of femtosecond time-resolved 2PPE. We resolved an intriguing relaxation dynamics involving four processes, ranging from sub-picosecond (ps) to several hundred ps time spans. These show a tendency to increase with increasing coverage. The present study provides important parameters such as energetic positions of transport levels as well as lifetimes of electronically excited states, which are essential for designing organic-molecule-based optoelectronic devices.

Introduction

Semiconducting materials based on organic molecules are being explored as potential active materials in (opto)electronic devices such as organic photovoltaic cells, light emitting diodes, or thin film transistors.1–4 For device performance the properties of interfaces between the inorganic substrate and the organic material as well as between the active layers (donor/acceptor) are crucial. Thereby the orientation of the molecules is of particular importance for the light emission/absorption and charge transport properties. Intermolecular interactions compete with molecule/substrate interactions both defining the adsorption geometry and the arrangement of the molecular film and, accordingly, the electronic structure of the system. The electronic structure of interfaces crucially influences the functionalities of organic films.5–7 The energetic positions of molecular electronic states or bands (transport levels) with respect to the Fermi level of a (metallic) substrate yields quantitative information on the barriers for electron (hole) injection at the molecule–substrate interface. Additionally, the formation and decay dynamics of excitonic states both within the films and at interfaces (e.g. donor/acceptor-interfaces) are key processes governing the optoelectronic functionality of the films.

Oligothiophene derivatives, in particular dicyanovinyl (DCV)-substituted oligothiophenes, have been utilized in vacuum-processed small-molecule organic solar cells.5–13 Using these low-band gap donor materials promising efficiencies of up to 8.3% for a single junction and up to 9.7% for a triple junction cell have been obtained.14 The adsorption and electronic properties of bare non-substituted oligothiophenes adsorbed on noble metal surfaces are well-known.15–25 The substituted species DCV5T-Me2 (see Fig. 1(a)) is expected to show different structural properties that explain its larger photovoltaic efficiency. It combines an electron-rich donor backbone and two terminal
electron-poor acceptor moieties. On the Au(111) surface, this compound self-assembles in extended chains stabilized by intercalated Au adatoms coordinated via terminal CN-groups resulting in a redistribution of molecular orbitals.\textsuperscript{26}

In the present contribution we utilized complementary experimental techniques such as scanning tunneling microscopy/spectroscopy (STM/STS), atomic force microscopy (AFM), and two-photon photoemission (2PPE) spectroscopy to obtain a comprehensive picture of the adsorption and electronic properties of DCV5T-Me\textsubscript{2} on Au(111) from the submonolayer to the multilayer regime. We found that two-dimensional DCV5T-Me\textsubscript{2} molecular assemblies appear on the surface at submonolayer coverages with a structure similar to that adopted in multilayer growth. A combination of tunneling and photoemission spectroscopy of the first and second layer provided detailed insights into the energetic positions of affinity levels and ionization potentials. Additionally we utilized femtosecond time-resolved 2PPE to elucidate the dynamics of electronically excited states in DCV5T-Me\textsubscript{2} after optical excitation. We show that the submonolayer is strongly affected by the electronic coupling to the metallic states. With increasing coverage, the molecular states are better decoupled and the relaxation dynamics of the excited states slow down. These findings uncover the limit of ultra-thin films as functional building blocks for photovoltaic cells.

**Methods**

The Au(111) single crystal was prepared by Ar\textsuperscript{+} or Ne\textsuperscript{+} sputtering and annealing at 800 K. DCV5T-Me\textsubscript{2} molecules were evaporated from a Knudsen cell onto the substrate held at room temperature (around 300 K). STM/STS, AFM and 2PPE experiments were carried out in two different setups under ultrahigh vacuum conditions.

**STM/AFM experiments**

STM/AFM measurements were done on a custom-made low temperature scanning tunneling microscope, at 4.8 K, equipped with a tuning fork force sensor\textsuperscript{27} for simultaneous non-contact AFM and STM measurements. Tunneling spectroscopy was performed using lock-in techniques to measure the differential conductance of the junction, which allows us to localize the energy alignment of molecular-derived resonances. Spatial maps of the differential conductance at the resonance voltage were drawn with an open feedback loop. Non-contact AFM images were obtained by mapping relative changes in the resonance frequency of the sensor at a constant tip height over the sample. High resolution images with clear intramolecular contrast were obtained with Xe functionalized tips at the onset of repulsive tip-surface forces. AFM images were Laplace filtered to emphasize the intramolecular corrugation over the smooth molecular background.

Identification of molecular structures and orbitals from AFM and dI/dV maps was based on density functional theory (DFT) simulations of minimum energy molecular structure and orbital isosurfaces. We employed the Gaussian package, with the B3LYP exchange–correlation functional and the 6-31G basis set. Since DFT methods fail to reproduce level alignment correctly,\textsuperscript{28} the orbital level was strictly obtained by comparing STS and 2PPE spectra.

**2PPE experiments**

2PPE allows the investigation of occupied, unoccupied electronic and excitonic states\textsuperscript{29} in a pump–probe scheme using femtosecond (fs) laser pulses. While the pump pulse $h\nu_1$ excites an electron from below the Fermi level ($E_F$) to intermediate unoccupied states at energies $E - E_F = E_{\text{kin}} + \Phi - h\nu_2$ (with $\Phi$ the work function), the probe pulse $h\nu_2$ photoionizes the sample by lifting the excited electron above the vacuum level ($E_{\text{vac}}$). Photoelectrons are detected using a custom-built time-of-flight (TOF) spectrometer and are analyzed with respect to their kinetic energy ($E_{\text{kin}}$) (for details see ref. 30). In order to elucidate the origin of peaks in the 2PPE spectrum, viz. identify whether they originate from occupied initial or unoccupied intermediate states in the 2PPE process, the dependence of the electron kinetic energy $E_{\text{kin}}$ on the photon energy was investigated. When an unoccupied intermediate state, such as the LUMO, is probed in one-color 2PPE (pump and probe photon have the same energy) the change in $E_{\text{kin}}$ scales with that in photon energy $h\nu$ and $h\nu'$, i.e., $E_{\text{kin}} = 1\Delta h\nu (\Delta h\nu = h\nu - h\nu')$. On the other hand,
for an occupied initial state, e.g., the HOMO, the kinetic energy of the electron ejected scales with twice the photon energy $2\hbar \nu$. In two-color 2PPE (pump and probe photon have different energies, one in the visible the other in the UV regime. The latter is the frequency doubled visible pulse.) the energy of the ejected electron scales with three times the photon energy (i.e., $3\hbar \nu$ with respect to the photon in the visible regime) for occupied initial states and twice or once the photon energy ($2\hbar \nu$ or $1\hbar \nu$) for unoccupied intermediate states. This analysis is generally not applicable for transitions between bulk bands due to their strong perpendicular dispersion but holds in the case of surface and adsorbate-derived states. The assignment of 2PPE features to photoemission from occupied or unoccupied molecular orbitals is carried out via their energetic position with respect to the substrate Fermi level, i.e., the closed state to $E_F$ is the HOMO or the LUMO, respectively. Energetically lower and higher lying states are accordingly lower lying HOMOs (HOMO$-n$) and higher lying LUMOs (LUMO$+n$). Additionally, the STS results are used for the assignment. The 2PPE spectra are displayed with an energy axis representing the final state of the photoelectron, referenced to the Fermi level ($E_{\text{final}} - E_F = E_{\text{kin}} + \Phi$). The UV-spectra were fitted in two parts: the low energy part was fitted with an exponential function, for the secondary electron background, and a reasonable number of Gaussians, depending on the number of the peaks observed. The high energy part of the spectrum was fitted only with Gaussians since the secondary electron background is rather flat at high energies. The same fitting principle was applied to the correlated spectra. In time-resolved 2PPE, the pump–probe scheme enables us to study the population dynamics of excited electronic states on the time scale of the femtosecond laser pulses by delaying the probe pulse with respect to the pump pulse. For 2PPE experiments the DCV5T-Me$_2$ coverage was determined by temperature-programmed desorption and work function measurements.

Results and discussion

The properties of an organic film depend on the degree of electronic mixing between molecular and metal states at the organic–inorganic interface, and on the energy alignment of molecular states. Hence, we start by characterizing the regime of coverage below one monolayer.

Sub-monolayer coverage

We grew a sub-monolayer coverage of DCV5T-Me$_2$ molecules on Au(111) at room temperature and cooled down to 4.8 K for STM inspection. We found that DCV5T-Me$_2$ forms both two-dimensional molecular islands and extended chain structures (Fig. 1(a)). The latter were found to consist of coordination-bonded DCV5T-Me$_2$-Au structures in ref. 26. Here we study the two-dimensional assemblies, which dominate the aspect of the sample for coverages closer to one monolayer. We typically find domains with the extension of hundreds of nanometers. STM images show that the islands are composed of molecules adopting a bent configuration with the DCV moieties contacting the center of neighboring molecules (Fig. 1(b)). This molecular structure suggests the absence of coordinating Au adatoms in the layer structure, in opposition to the chains. In order to precisely determine the bonding geometry within the islands, we performed AFM measurements using a Xe functionalized tip. Fig. 1(c) shows a Laplace-filtered constant height AFM image of a molecular island, where the chemical structure of the DCV5T-Me$_2$ molecules is resolved. The thiophene rings are imaged as pentagonal structures with their sulfur atoms exhibiting the sharpest contrast.

The pentagon alternate their orientation along the molecular axis, as indicated in the molecular models superimposed to Fig. 1(c). The DCV groups are resolved as protrusion with brighter contrast localized at the terminal nitrogen atoms. AFM images further exclude that Au adatoms are forming part of the structure and, hence, mediate the intermolecular bonding, as it is the case for the chain structures. Instead, the DCV endgroups are arranged with a cyano end pointing towards a sulfur atom at the neighbor molecule. In spite of the large conformational flexibility of DCV5T-Me$_2$, all of the molecules appear with this shape. DFT calculations of the isolated molecule revealed that the shape observed here corresponds to the global minimum. This minimum energy configuration has a planar structure, thus enhancing its relative stability when forced to lay in a planar adsorption geometry on a metal surface.

Next, we investigated the electronic properties of DCV5T-Me$_2$ molecules in islands by combining STS and 2PPE measurements. Fig. 2(b) compares the STS spectra of unoccupied states (i.e., positive sample bias) acquired at three different positions within one DCV5T-Me$_2$ molecule. All spectra show a peak of differential conductance at 1.3 V with similar intensity at the three sites, but with a clear shoulder at 1.65 V only at the endgroups of the DCV5T-Me$_2$ molecule. Furthermore, the spectrum at the molecular center shows a second peak at 2.35 V, completely absent at the sides. This peculiar distribution of unoccupied molecular states is further confirmed by spatially mapping the corresponding $dI/dV$ signal at each resonance bias position (Fig. 2(c)–(e)). Supported by DFT calculations of the orbital shape (wave-function amplitude isosurfaces) we assign the two lowest resonances to molecular states derived from the LUMO and LUMO+1, respectively. The resonance at 2.35 V is highly localized at the central part of the molecule. We assign it to the LUMO+2 derived resonance, because it is also weighted at the center of the molecule. The resulting spectral structure indicates that the molecules retain their free-molecule character on the surface, and the degree of hybridization with the surface is relatively small.

We will demonstrate that using 2PPE enables us to also gain further insights into the energetic positions of occupied electronic molecular states as well as excitonic states. Fig. 3 shows 2PPE data of a sub-monolayer DCV5T-Me$_2$/Au(111).

The one-color 2PPE spectrum shows several peaks related to either occupied electronic states of the DCV5T-Me$_2$/Au(111) systems, such as d-bands (located around $-2.0$ eV and $-2.6$ eV) and the Shockley surface state (SS) (at $-0.48$ eV).
or occupied molecular resonances, or to unoccupied molecular resonances. In order to identify whether the peaks originate from occupied initial or unoccupied intermediate states in the 2PPE process, their energetic position is measured as a function of photon energy (see the inset of Fig. 3(a)). The peaks labeled as A, B, and C shift equally with the change in photon energy, resulting in photoemission from unoccupied electronic states located at 4.0 eV (state A), 3.6 eV (state B) and 3.4 eV (state C) with respect to \( E_F \). At \( E_{\text{final}} - E_F = 8.65 \) eV we observe a peak which shifts by twice the photon energy. This state is assigned to a low-lying occupied state, namely the HOMO (see below), with a binding energy of \(-1.0 \) eV. Note that the work function adapts a value of 5.4 eV in this low-coverage regime, i.e., a reduction by 50 meV compared to the bare Au(111) surface. In the two-color spectrum (Fig. 3(b)) two additional features are found, both shifting with twice the photon energy \((h\nu, \text{visible pulse})\) indicating that they originate from the photoemission of unoccupied states. They are located at 1.3 and 1.8 eV with respect to \( E_F \). Comparing with the STS data, we assign them to the LUMO and LUMO+1, respectively. However, in STS the LUMO+1 is found to be at 1.65 eV, i.e., 150 meV closer to \( E_F \). We admit that the feature overlaps with the very intense photoemission from the surface state leading to a higher uncertainty in the determination of the energetic position.

**Monolayer and higher coverages**

The growth of thicker films proceeds in a layer-by-layer fashion: only when the coverage is increased above one monolayer, self-assembled structures appear on top of a fully saturated first layer (Fig. 4(a)). The unit cell of the molecules in the second layer has the same dimension as in the first layer. However, high resolution STM images (Fig. 4(b)) suggest that the upper layer is shifted with respect to the underlying layer and that the molecules adopt a more asymmetric configuration, with their two ends appearing differently in the STM images. We expect that due to their large molecular flexibility, DCV endgroups and thiophene rings of second layer molecules rotate to bond and accommodate their structure over the first layer. However, the layer-by-layer growth indicates that the interaction between layers is weak, which is further confirmed by STS measurements.

In order to elucidate the evolution of molecular electronic properties as a function of coverage, we performed tunnel spectroscopy measurements on second layer DCV5T-Me₂ molecules (Fig. 4(c)). The spectra now show the effect of a less interacting layer. The broad resonance of the first layer, composed of both the LUMO and the LUMO+1, appears now as a multiple component peak with two main features at 1.78 V (over the whole molecule) and 2.14 V (at the sides), attributing...
to the LUMO and the LUMO+1. The higher alignment of the peaks (~0.5 V higher) and the sharper linewidth (~0.1 V and 0.2 V FWHM) prove that this molecular system is less electronically coupled to the metal surface due to reduced screening and less hybridization, respectively. It is thus a direct consequence of tunneling spectroscopy probing affinity levels instead of ground state unoccupied state alignment.\textsuperscript{36} Accompanying the LUMO state, a third component could be resolved in the spectra as a peak shifted by 180 meV. The energy separation between the LUMO and this feature coincides with the energy of totally symmetric (anti-symmetric) in-plane phonon of the DCV5T-Me\textsubscript{2} backbone, located at 179.4 meV in the Raman (infrared) spectrum.\textsuperscript{37} We thus attribute it to a vibronic feature excited by electrons during tunneling.\textsuperscript{38} In fact, the excitation of similar modes of the plain tetrathiophene (4T) molecules has been found to be crucial to explain the polaronic charge mobility in these molecular materials because they are highly coupled to 4T frontier orbitals.\textsuperscript{39}
(molecular Frenkel exciton). The photoemission intensity from this excitonic state increases drastically with increasing coverage as can be seen already in the 2PPE spectrum from 3 ML (see Fig. 5(c)). For a higher coverage (>6 ML) the spectrum is dominated by this feature, hiding completely the photoemission features associated with the LUMO and LUMO+1. As discussed below, most likely two excitonic states contribute to this broad photoemission signal. At a coverage of 3 ML we observed (see Fig. 5(d)) two new features located at 1.8 and 1.95 eV above \( E_p \) in the two-color 2PPE spectrum. According to the STS results (see Fig. 4(c)), we assign them to the LUMO and LUMO+1 of the molecules in the second decoupled layer.

Electronic states as a function of coverage

Merging together the STS and 2PPE data, we obtain a comprehensive picture of the energetic positions of the adsorbate-derived unoccupied and occupied electronic states. The energy levels and their assignments as a function of coverage are summarized in Fig. 6. Note that electronic states determined using 2PPE are transport states since in the 2PPE the unoccupied states (e.g. LUMOs) are populated via a metal-to-molecule electron transfer (creating a negative ion resonance) and not through an intramolecular charge transfer. Occupied electronic states are ionized, creating a positive ion. Thus we obtain the ionization potential or the electron affinity of the respective molecular state and for the quasi-particle (the exciton) the ionization potential is measured. The same is true for STS, since electron tunneling into unoccupied states leads to the creation of negative ion resonances; tunneling out of occupied states to the generation of positive ion resonances. Hence, both techniques allow the determination of transport levels and accordingly transport gaps. The lowest exciton state is generated by exciting an electron from the HOMO to the LUMO (the transport gap). The lowest exciton state is generated by exciting an electron from the HOMO to the LUMO (the transport gap).

Comparing the sub-monolayer and monolayer coverage, the main difference is a shift of the work function which affects all energy levels equally. Furthermore, the HOMO and HOMO−1 are not observed in 2PPE at sub-MLs, which is most likely due to the work function signal from a very small number of molecules. At 1 ML, with the LUMO level at 3.8 eV and the HOMO at 5.2 eV we obtain a transport gap of 1.4 eV. For comparison, DCV5T-Me2 in solution possesses oxidation and reduction potentials of ~3.75 and ~5.66 eV measured using an electrochemical technique of cyclic voltammetry corresponding to an energy gap of 1.91 eV, 0.5 eV larger than the gap measured in our study. This clearly indicates that the metallic substrate leads to a decrease of the gap size due to screening of molecular charge states. The excitonic state is located 0.6 eV below the affinity level associated with the LUMO. For comparison, in sexithiophene/Au(111) an energy difference of 0.9 eV has been found.

To clarify the origin of the higher lying unoccupied states labeled as A, B, and C we performed angular-resolved 2PPE measurements. Both states A and B show no dispersion (the signal of state C was too weak to resolve it). Additionally, they are not pinned to the vacuum level. Thus, we can exclude that they arise from image potential states (IPS), since IPS possess a free-electron like dispersion and are pinned to the vacuum level. Other possibilities are: (i) interface states (IS), (ii) charge-transfer excitons\(^41,42\) on the surface of the DCV5T-Me2 with the image band serving as an electron acceptor, or (iii) higher lying molecule-derived states. Considering possibility (i): IS have been observed at organic/metal interfaces, for instance at the interface of an imine-derivative at Au(111).\(^43\) There two delocalized unoccupied electronic states have been identified, which have been proposed to originate from a modified image potential at the interface. At the perylene-tetracarboxylic acid dianhydride (PTCDA) on the Ag(111) and Ag(100) interface,\(^44-46\) respectively, an interface state generated from a Shockley resonance has been suggested. Thereby the organic overlayer induced energetic shifts of a Shockley resonance. However, interface states disperse and do not reproduce the absence of angular dispersion found for states A and B. In both cases (ii) and (iii) one would expect no dispersion of the respective states, as we find here. Photoemission from an exciton (case ii) does not show dispersion since the photoionization process destroys the exciton (quasiparticle).\(^47\) The dispersion of molecular states (case iii) is highly improbable in flat-lying molecular layers. Hence, possibilities (ii) and (iii) have to be taken into account.

Going to higher coverages, i.e., above one ML, we observe pronounced shifts in energy for several molecular states, most dominant for the LUMO and LUMO+1 levels, where a shift of \( \approx 0.5 \) eV is found. This effect is attributed to electronic decoupling from the metallic substrate. In contrast, for the octathiophene (8T)/Au(111) no energetic shift of the LUMO and LUMO+1 in the first and second layer was found.\(^24\) In that case the molecules were arranged in a tilted geometry with respect to the surface plane, which allowed for intermolecular \( \pi-\pi \) interaction. With such adsorption geometry, already the adsorbate/substrate interaction in the first layer is not strong and accordingly, the electronic coupling. Therefore no significant decoupling

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**Fig. 6** Energy level diagram as a function of DCV5T-Me2 coverage. The blue levels are the ionization potentials (left axis) and the red ones are the electron affinities (right axis); \( E_F \) denotes the Au(111) Fermi level.
effect was observed contrary to the present system, where DCV5T-Me₂ adsorbs in a flat configuration. Hence, there is a stronger coupling with metal states. The first layer then acts as an isolating layer resulting in electronically decoupled molecular states in the second layer. The comparison between the two systems DCV5T-Me₂/Au(111) and 8T/Au(111) nicely pictures the effect of the adsorption geometry on the electronic structure of the adsorbate/substrate system (structure/property relationship).

**Excited state dynamics**

In the following, we investigate the excited state dynamics using femtosecond time-resolved 2PPE and show that they are also strongly influenced by the electronic coupling between the adsorbate and the surface at the interface. In the low-coverage regime (1–2 ML), the lifetimes of electrons in the molecular excited states, viz. the LUMO and LUMO+1 as well as the excitonic state possess values that are below the limit of our experimental resolution (τ < 10 fs). These ultrashort lifetimes point towards an efficient back transfer of electrons to the adsorbate and the surface at the interface. In the low-coverage regime (1–2 ML), the lifetimes of electrons in the molecular excited states, i.e., in the range of fs, have been found for several adsorbates. Contrarily, for higher coverages long lifetimes are found as can be seen in Fig. 7(a) for 20 ML DCV5T-Me₂ on Au(111). In this false color two-dimensional representation, the 2PPE intensity at a particular energy, $E_{\text{final}} - E_{\text{f}}$, is displayed as a function of time delay (Δt) between the two laser pulses $h\nu_1 = 2.4$ eV and $h\nu_2 = 4.8$ eV. Thereby positive time delays represent delayed UV pulses ($h\nu_2$, probe pulse). In the high coverage regime the excitation in 2PPE involves an intramolecular process, e.g. a HOMO–LUMO transition. The optical gap in a condensed DCV5T-Me₂ film is 1.69 eV as determined using UV/VIS absorption spectroscopy. The absorption spectrum shows a broad band between 730 and ≈ 450 nm, which is attributed to the $S_0 \rightarrow S_2$ (HOMO → LUMO) as well as the $S_0 \rightarrow S_3$-transition (HOMO → LUMO+1). Thus with a pump pulse photon energy of 2.4 eV (corresponding to 520 nm) we are able to induce both transitions. Therefore we assign the broad 2PPE signal to excitonic levels related to the LUMO and LUMO+1. As mentioned above with increasing coverage the 2PPE spectrum is dominated by this feature, which possesses a peak width much larger than the width of all other observed signals. This also underlines that only one excitonic state does not contribute to the signal.

The cross correlation (XC) curve for the energy region of the excited states is shown in Fig. 7(b). The solid line corresponds to a fit used to determine lifetimes. The fit model contains a Gaussian function representing the laser pulse duration convoluted with the response function of the intermediate state. The response function is not a single exponential due to different excitonic states involved in the excited state dynamics. A superposition of four exponential decays with different time constants $\tau_1$–$\tau_4$ results into agreement with the data. For a coverage of 20 ML the time constants are $\tau_1 = 130 \pm 13$ fs, $\tau_2 = 1 \pm 0.1$ ps, $\tau_3 = 11 \pm 1.1$ ps, and $\tau_4 = 390 \pm 39$ ps. Coverage-dependent measurements showed similar excited state dynamics. In all cases an exponential decay using four different time constants is needed to describe the data and the lifetimes increase with rising coverage. A coverage-dependent decay time indicates that the excited state decays via two relaxation channels, an intrinsic and a distance-dependent channel. The intrinsic decay rate corresponds to the decay in the bulk material and the distance-dependent (external) rate to quenching by the metal substrate.

So far neither time-dependent measurements on excited state dynamics of DCV5T-Me₂ both in the liquid or condensed phase nor electronic structure calculations are available in the literature. However we propose that the following processes may play a role. An initial hot exciton population which is highly delocalized within the first and second excitonic band structure localizes on an ultrafast time-scale ($\tau_1 = 130 \pm 13$ fs). The relaxation and localization of higher non-relaxed delocalized excitonic states on these short time-scales have also been observed in other organic semiconductors. Strong coupling of excited electrons to phonons within the film can explain the very effective energetic relaxation within the first picosecond. STS on the second decoupled DCV5T-Me₂ layer showed vibronic states (see Fig. 4(c)) indicating strong electron–vibration coupling with vibrational states of around 180 meV. The longer lived components ($\tau_3 = 11 \pm 1.1$ ps and $\tau_4 = 390 \pm 39$ ps) may be associated with the lifetime of the lowest lying excitonic state as well as polaron or electrons trapped into defect sites.

**Conclusions**

In summary, we have determined the electronic structure, i.e., the energetic position of affinity levels and ionization potentials of DCV5T-Me₂ adsorbed on a Au(111) surface as a function...
of coverage using scanning tunneling spectroscopy and two-photon photoemission (2PPE). Several unoccupied molecular electronic states shift to higher energies with respect to the Fermi level of Au(111) in the second adsorbate layer, indicating an efficient electronic decoupling from the metallic substrate. For molecular multilayers, we identified the presence of excitonic states, becoming stronger with the film thickness. Femtosecond time-resolved 2PPE pictured the excited state dynamics of the two lowest exciton bands as composed of four processes with different time scales. The quantitative determination of the energetic position of molecular electronic levels with respect to the Fermi level of an electrode and the elucidation of the charge carrier and exciton dynamics are important for the improvement and optimization of molecule-based device performance.

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References

28 Note that we utilized the DFT calculations just for visualizing of the molecular orbital shapes since they are used for assignment of the d/d' maps and not for their energy level alignment. In the latter case special types of functionals (range-separated DFT) have to be used as demonstrated e.g. for various thiophenes [B. M. Wong, et al., Phys. Chem. Chem. Phys., 2009, 11, 4498; T. Körzdörfer, et al., J. Chem. Phys., 2011, 135, 204107].