Quasi one-dimensional band dispersion and surface metallization in long-range ordered polymeric wires

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On-surface covalent self-assembly of organic molecules is a very promising bottom-up approach for producing atomically controlled nanostructures. Due to their highly tuneable properties, these structures may be used as building blocks in electronic carbon-based molecular devices. Following this idea, here we report on the electronic structure of an ordered array of poly(para-phenylene) nanowires produced by surface-catalysed dehalogenative reaction. By scanning tunnelling spectroscopy we follow the quantization of unoccupied molecular states as a function of oligomer length, with Fermi level crossing observed for long chains. Angle-resolved photoelectron spectroscopy reveals a quasi-1D valence band as well as a direct gap of 1.15 eV, as the conduction band is partially filled through adsorption on the surface. Tight-binding modelling and ab initio density functional theory calculations lead to a full description of the band structure, including the gap size and charge transfer mechanisms, highlighting a strong substrate–molecule interaction that drives the system into a metallic behaviour.

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A major challenge in modern surface science is to create ordered arrays of covalently linked organic nanostructures. By doping molecular electronic bands into highly conductive states, these structures may be promising for use as elemental building blocks in electronic carbon-based molecular devices such as organic field-effect transistors, light-emitting diodes, photovoltaics and sensors. Despite its exceptional physical properties, graphene’s lack of a bandgap severely limits its potential for creating such devices. Engineering the gap in graphene by using nanostructuring, for example, creating graphene nanoribbons (GNRs) of narrow width, has been proposed as a feasible route towards carbon-based electronics. Thus, the GNRs’ bandgap can be tuned by altering their lateral size or by modifying their edge termination (armchair versus zigzag). An emerging bottom-up approach for producing such carbon nanostructures, exploits covalent linking (polymerization) of precursor molecules on metal surfaces. In producing such carbon nanostructures, exploits covalent linking (polymerization) of precursor molecules on metal surfaces.

Polymers are typically demonstrated by measuring the valence band structure along the chains (Supplementary Fig. 3, Supplementary Note 2). However, it is possible to obtain an additional organometallic phase at high coverage, above 0.9 monolayer, which co-exists with the organometallic phase described above and fully dominates the surface at 1 monolayer. STM images and low-energy electron diffraction patterns characterising these two phases are provided and discussed in the Supplementary Figs 1 and 2 and Supplementary Note 1. When this high-coverage phase is annealed at 500 K, alternating rows of polymers and bromine atoms are formed, as shown in the STM images presented in Fig. 1b. The measured internal periodicity of 4.4 ± 0.2 Å agrees with the expected phenyl–phenyl spacing, which is a characteristic of PPP. The polymer chains produced from high coverage are packed into domains and aligned along the two <1–1–2> and <1–1–12> substrate directions, a detail that complicates measurement by surface-averaging techniques such as photoelectron spectroscopy. While a single-domain orientation dominates in Fig. 1b, both orientations are observed in longer range STM images, and cover the surface with equal probability (Supplementary Fig. 4). The substrate periodicity along <1–1–12> is 4.43 Å, which matches with the lattice parameter of the PPP. Thus, the polymer is here fully commensurate with the surface and may be grown to long size with little or no strain arising from lattice mismatch (Supplementary Note 3).

The results of DFT optimization of the surface structure are presented in Fig. 1c (see Methods section for details). The GGA–PBE formulation was used to approximate the exchange-correlation functional potential. The van der Waals (vdW) interactions were described using the vdW density functional (vdW-DF) together with the optB86b exchange functional. This approach yields a very good description of the vdW forces between molecules and surfaces. The PPP polymer (C atoms grey) is adsorbed in a flat geometry, 2.2 Å above the copper surface. Such a short distance (cf. 3.35 Å for inter-planar spacing in graphite) points to a significant hybridization of the electronic levels of the polymer with the copper interface states. Bromine atoms (green spheres) are adsorbed on the short bridge (SB) site of the underlying Cu lattice (orange spheres). The centre of each phenyl ring overlaps the hollow (H) site, which is the preferred site for benzene adsorption on Cu(110)42. An STM image calculated using the Tersoff–Hamann approximation at a bias voltage of 200 mV is an excellent match to the experimental data as presented in Fig. 1d,e.

Results

An ordered and commensurate polymeric phase on Cu(110). A systematic investigation of the dBB/Cu(110) interface as a function of coverage and annealing temperature allows us to identify an unreported structural arrangement, which was used as a starting point for the formation of an ordered polymer phase via thermal activation. In previous work, Di Giovannantonio et al. demonstrated that the thermal treatment of vacuum-deposited dBB on Cu(110) leads to the formation of PPP chains. This process is understood to be an Ullmann coupling reaction, which is summarized in Fig. 1a: depositing the molecules at room temperature leads to an organometallic phase, where Br–C bonds are replaced by C–Cu–C bridges. Subsequent annealing above 460 K promotes C–C bonding, resulting in the formation of coaligned π-conjugated PPP chains with a lattice parameter of 4.4 Å as measured by STM. In that case, the PPP chains were aligned along the <1–1–10> direction, for which the interatomic Cu distance of the surface is 2.55 Å. This leads to the incommensurability between the polymer and the substrate (Supplementary Fig. 3, Supplementary Note 2).

In the current study, the thermal treatment of vacuum-deposited dBB on Cu(110) leads to the formation of PPP chains. This has been demonstrated by measuring the valence band structure along the chains (Supplementary Fig. 3, Supplementary Note 2). However, it is possible to obtain an additional organometallic phase at high coverage, above 0.9 monolayer, which co-exists with the organometallic phase described above and fully dominates the surface at 1 monolayer. STM images and low-energy electron diffraction patterns characterising these two phases are provided and discussed in the Supplementary Figs 1 and 2 and Supplementary Note 1. When this high-coverage phase is annealed at 500 K, alternating rows of polymers and bromine atoms are formed, as shown in the STM images presented in Fig. 1b. The measured internal periodicity of 4.4 ± 0.2 Å agrees with the expected phenyl–phenyl spacing, which is a characteristic of PPP. The polymer chains produced from high coverage are packed into domains and aligned along the two <1–1–2> and <1–1–12> substrate directions, a detail that complicates measurement by surface-averaging techniques such as photoelectron spectroscopy. While a single-domain orientation dominates in Fig. 1b, both orientations are observed in longer range STM images, and cover the surface with equal probability (Supplementary Fig. 4). The substrate periodicity along <1–1–12> is 4.43 Å, which matches with the lattice parameter of the PPP. Thus, the polymer is here fully commensurate with the surface and may be grown to long size with little or no strain arising from lattice mismatch (Supplementary Note 3).

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Local spectroscopy and tight-binding modelling. Using STS, confinement of unoccupied molecular states has been observed previously in individual polythiophene chains, and more recently, in 7-AGNRs and GNR heterojunctions. We recorded differential conductivity maps as a function of chain length, to build the k-resolved band structure associated with the conduction band of the infinite polymer. STS measurements obtained on PPP oligomers with N = 6 (26 Å), N = 10 (44 Å) and N = 22 (97 Å) phenyl rings are presented in Fig. 2a. As expected, the local density of states (LDOS) is spatially modulated along the...
chains, leading to standing wave patterns which depend strongly both on the bias voltage and the oligomer length. For \( N = 6 \), the lowest unoccupied molecular orbital (LUMO; indicated by a blue segment in Fig. 2a) is slightly above the Fermi level. As the length of the chain is increased, the LUMO progressively moves below the Fermi level, reaching \(-0.4 \text{ eV}\) for the \( N = 22 \) polymer.

A simple TB model (see Methods section for details) reproduces our results. The long-range modulations observed in the LUMO, LUMO + 1 and LUMO + 2 orbitals are in agreement with the local conductance maps recorded for the first three states in the polymer chains, as shown in Fig. 2b,c for the \( N = 6 \) case. Using a single C–C resonance integral \( \beta = -3.6 \text{ eV} \), an excellent fit to experimental binding energies (Supplementary Fig. 4 and Supplementary Note 3) as a function of oligomer length is observed as presented in Fig. 2d. However, the absolute position of these levels is not well described by the model and has to be manually shifted by 1.9 eV towards occupied states. Defining \( k_{\parallel} = 2\pi n / N \) \((p < N, a = 4.4 \text{ Å})\) as the wave vector associated to the 1D-confined states allows us to build the conduction band dispersion for the infinite polymer, as shown in Fig. 2e. A parabolic fit (black dashed line) to the bottom of the conduction band implies an effective mass of \( 0.15 \pm 0.02 \text{ m}_e \) for charge carriers in the vicinity of the Fermi level. This value is in agreement with those obtained for AGNRs\(^{30}\). As the band crosses the Fermi level at \( k_F = \pm 0.12 \text{ Å}^{-1} \), the PPP polymers are metallic and thus represent an array of conducting nanowires. None of the occupied states were resolved by STS, and thus we were unable to extract the bandgap of the PPP.

**Band structure from angle-resolved photoelectron spectroscopy.**

ARPES has previously been used to reveal quantum well states in molecules\(^{46,47}\) or to reconstruct the discrete electronic orbitals of self-assembled individual molecules at surfaces\(^{48,49}\). In addition, evidence of changes in the density of states of polymers, especially as a function of doping, have been observed by ultraviolet photoelectron spectroscopy (UPS)\(^{50}\). Nevertheless, direct evidence of a \( k \)-resolved band structure associated with the C–C covalent bonding and a long-range delocalization of charge carriers have been reported to date only on graphene materials\(^{31}\). Here ARPES intensity maps were measured along the \(<1–1 2>\) axis, parallel to the chain direction of one domain (therefore probing roughly half of the monolayer). The incident photons were \( p \)-polarized, with an energy of \( h\nu = 35 \text{ eV} \). The spectral intensity from the second domain, which is rotated by \( 70.52^\circ \) with respect to the first, does not contribute to the ARPES signal because its \( k \)-points are not accessible in this experimental geometry, except at the \( \Gamma \) point. Comparison of surfaces both with and without polymers permits the identification of two dispersive bands, on both sides of the \( 3d \) states of the substrate,
Parabolic dispersion with (blue solid line), is obtained from a tight-binding model using the same resonance integral as the Hückel model. The black dashed curve represents an energy range bands it is not possible to resolve the molecular dispersion in the dSupplementary Fig. 5). Due to the high intensity of Cu 3
geometry of the phenyl rings (Supplementary Note 4,
pvector parallel to p-orbital axis) to s-polarization (having only in
s-polarization (having an out of surface plane component of the polarization
vector parallel to π-orbital axis) to s-polarization (having only in
surface plane component of the polarization vector) consistent
with the π-character of the molecular orbitals and the flat-lying
geometry of the phenyl rings (Supplementary Note 4,
Supplementary Fig. 5). Due to the high intensity of Cu 3d
bands it is not possible to resolve the molecular dispersion in the
energy range \(E - E_F\) between \(-2\) and \(-5\) eV.

Calculation of band structure and ARPES spectral intensity
using the Hückel (tight-binding) models provides an understanding
of the observed dispersions. The valence band structure of the infinite free polymer, calculated in the TB model using a resonance integral of \(\beta = -3.5\) eV, is presented in Fig. 3b (red dashed lines). Here the molecular band was rigidly shifted by 0.1 eV to match the data. Three bands are easily identified, two of them dispersing in opposite phase and crossing at the edge of the first Brillouin zone (BZ) of the system, that is, for \(k_{||} = \frac{\pi}{d} = 0.71\) Å\(^{-1}\), consistent with the PPP periodicity of \(d = 4.4\) Å. The theoretical ARPES intensity (colour scale on Fig. 3b) was deduced from the Fourier-transform of each molecular orbitals calculated for a \(N=20\) polymer, according to the Fermi’s golden rule as proposed in the cases of sexyphenyl on the Cu(110) surface\(^{41}\) and PTCDA on the Ag(110) surface\(^{42}\) (Supplementary Figs 6 and 7, Supplementary Note 5). It appears that, as usually observed in ARPES, the spectral intensity does not follow the periodicity of the reciprocal lattice, due to cross-sectional effects in the photoelectron emission\(^{31,32}\). In particular, the ARPES intensity of the top band at the gamma point of the first BZ is equal to zero, and thus no signature of this band was observed in normal emission. This phenomenon leads to an apparent band periodicity of \(2.84\) Å\(^{-1}\), which does not appear to match the BZ size (1.42 Å\(^{-1}\)). Therefore, knowledge of the orbital topology and a rudimentary model of the electronic structure are necessary to explain the appearance of experimental dispersion curves.

High-resolution measurements taken in the second BZ close to the top of the occupied molecular band show that the molecular spectral intensity disperses up to a binding energy of \(-1.4\) eV at \(k_{||} = 1.42\) Å\(^{-1}\), while a small portion of a higher energy band dips below the Fermi level. A direct bandgap of 1.15 eV is found between the minimum of this band and the top of the valence band described above. This result is in agreement with the

**Figure 2 | Conduction band structure deduced by scanning tunnelling spectroscopy.** (a) Experimental dI/dV measurements recorded along a polymer for three lengths (N correspond to the number of phenyl rings). The vertical axis is energy, the horizontal axis is position along the polymer. The blue marks show the bottom of the bands. (b) Schematic representation of the free sexiphenyl molecule and its LUMO, LUMO + 1 and LUMO + 2 states calculated using a tight-binding model built from Hückel orbitals. (c) Experimental topography (bottom) and STS conductance maps for the LUMO conduction band. Experimental points (blue) are extracted from data shown in (d), using the relation \(k_{||} = 2pN/Na\) \((p = 1, 2, \ldots)\). The theoretical band (blue solid line), is obtained from a tight-binding model using the same resonance integral as the Hückel model. The black dashed curve represents a parabolic dispersion with \(m^* = 0.15 \pm 0.02 m_e\). In (d) and (e), absolute positions of calculated bands have been shifted by 1.9 eV towards occupied states to match the experimental results.
Figure 3 | Angle-resolved photoelectron spectroscopy measurements. (a) ARPES intensity map measured along the <112> direction, parallel to the polymers chains for 1ML-PPP/Cu(110). The bottom part is displayed in second derivative. Arrows indicate bands due to the presence of polymers. (b) Tight-binding modelling of the band structure of the N = 20 PPP polymer using \( \beta = -3.5 \) eV. (c) High-resolution ARPES intensity map recorded close to the Fermi level showing the HOMO–LUMO bandgap. Measurements (d) and tight-binding modelling (e) of the ARPES constant energy map at -1.8 eV. (f) Zoom on the measured (top) and calculated (bottom) ARPES constant energy maps. The transverse resonance integral used in the calculations is \( \beta' = -0.15 \) eV.

DFT calculations. In Fig. 4a (right part), we show that our local STS measurements (blue squares) for different chain lengths are in remarkable agreement with the \( k \)-resolved conduction band structure obtained from ARPES (green squares). Red and blue solid lines present the band structure calculated in the TB model using a single C–C resonance integral \( \beta = -3.5 \) eV in agreement with the expected value for graphene materials \(^{54}\). However, the valence and conduction bands are independently shifted in energy, respectively, by 0.1 and 1.9 eV toward occupied states, since our TB model leads to a bandgap of 2.9 eV. So far, the precise value of the gap (1.15 eV) and the partial filling of the unoccupied molecular states cannot be understood in the framework of this simple yet powerful method. DFT computation of the electronic band structure for the model appearing in Fig. 1c is depicted at the left Fig. 4a. The bands originating from the occupied and unoccupied molecular states can be distinguished from those originating from the copper substrate by plotting the projection of the band structure onto the top layer of the system, which contains only H, Br and C atoms. The result of this projection is shown as red circles in Fig. 4a. The conduction band as well as several valence bands can be clearly identified. However, the projected band structure contains signatures of hybridization with the substrate, which are not captured in the TB model. A comparison between the measured and calculated electronic density of states (DOS) is shown in Fig. 4b. The DOS of pristine Cu(110) (filled
blue line) is flat between 0 and $-2\, \text{eV}$, and originates from copper $s$–$p$ bands. The additional spectral intensity arising from the PPP overlayer appears clearly as peaks corresponding to the top and bottom of the valence and conduction molecular bands, respectively. The separation between these peaks corresponds to a bandgap of 1.15 $\text{eV}$. The calculated gap is $\sim 0.90\, \text{eV}$, which is slightly smaller than the experimental value. This is considered as a common drawback of DFT calculations, which neglect correlation effects. Overall, both the size of the gap and the minimum of the conduction band below the Fermi level are captured by these simulations, in agreement with the finding of metallic character in the PPP nanowires.

**Discussion**

The gap for freestanding PPP is expected to be $2.9\, \text{eV}$ on the basis of the TB approach\(^5^5\). Assuming the PPP polymer as being a 3$p$-type AGNR with $p = 1$ and $w = 2.4\, \text{Å}$ (Supplementary Fig. 9, Supplementary Note 7), we expect a bandgap of 2.45 $\text{eV}$ from DFT calculations corrected at 4.1 $\text{eV}$ including GW corrections\(^6\). We have also carried out DFT calculations in the gas phase using B3LYP/6–31g(d) that yield a bandgap of 3.05 $\text{eV}$ (Supplementary Note 8). In another study, STS measurements were performed on isolated PPP chains grown on Cu(111) using a different precursor\(^3^2\). The bottom of the conduction band was found to be 0.9 $\text{eV}$ above the Fermi level for long chains, in the unoccupied electronic states, that is, shifted by 1.3 $\text{eV}$ compared with our measurements. Thus, understanding the interaction between the polymer and the substrate appears to be of major importance so as to explain both the bandgap closing and the filling of unoccupied molecular states, as emphasized very recently\(^3^4\).

Figure 5a dissects the essential mechanisms that should be taken into account in the energy level alignment and gap modifications in organic semiconductor/metal interfaces. Considering that the PPP polymer is adsorbed onto a surface, the resulting position of the molecular levels depends on the substrate work function as indicated on the left side of Fig. 5a. The Cu(111) work function (4.94 $\text{eV}$) is larger than that of Cu(110) (4.48 $\text{eV}$). Consequently molecular levels are shifted to lower binding energies on the Cu(110). This effect may be enhanced by an additional reduction of the work function $\Delta \phi$, due to the compression of the electron cloud inside the metal substrate induced by the proximity of the molecular layer\(^5^6–5^8\). In our case, the short distance of 2.2 $\text{Å}$ between the polymer and the surface may explain the large shift observed in the present work as compared with the report on Cu(111)\(^3^2\). However, it has also been shown that certain physisorbed molecules experience Fermi level pinning: in this case, charge transfer from the substrate to the LUMO creates a new surface dipole, which increases the work function until the LUMO lies with the Fermi energy, inhibiting its occupation\(^5^9,6^0\). For chemisorbed molecular materials, substantial molecule/substrate hybridization leads to a direct charge transfer, which favours LUMO occupation. This mechanism has been recently identified as being responsible for the metallization of pentacenequinone and pentacenetetrone molecules adsorbed on coinage metal surfaces\(^5^4\). In addition, several works have also reported a strong reduction of the HOMO–LUMO gap induced by the proximity of molecules with a metallic substrate, due to charge screening effects\(^5^9,6^0\). To test the impact of the substrate on the electronic properties, the band structure was recalculated in a geometry where the separation between thePPP and the substrate is increased from 2.2 to 8 $\text{Å}$ (Fig. 5b). It is clear that: (i) the valence and conduction molecular bands are better defined, having lost the signatures of hybridization with the substrate; (ii) the full molecular band structure is shifted upwards, with the top of the valence band grazing the Fermi level; (iii) the gap has increased to $\sim 1.4\, \text{eV}$ but still remains much lower than expected.

**Figure 4 | Band-structure of PPP polymers grown on Cu(110).** (a) DFT band structure (black lines, red circles correspond to the projection on the molecular layer), tight-binding modelling (red and blue solid lines), ARPES (green squares) and STS (blue squares) experimental dispersion curves. The absolute positions of the valence and conduction bands in the tight-binding model have been independently shifted towards the occupied states by 0.1 and 1.9 $\text{eV}$, respectively. (b) k-integrated photoemission DOS on the Cu(110) substrate (left panel, blue), on PPP/Cu(110) (left panel, yellow) and corresponding DFT DOS (right panel, yellow).
from gas phase calculations. In general the local density approximation and generalized gradient approximation (GGA) underestimate band gaps of GNRs and organic polymers in gas phase as compared with GW calculations. Nevertheless, the new position of the conduction band is above 1 eV as observed for PPP/Cu(111). Finally, we discuss the influence of the Br atoms on the electronic structure. To this end, we calculated the band structure both with (Fig. 5c) and without (Fig. 5d) Br atoms at surface. When Br is not present at surface the conduction band minimum (CBM) of the polymer is only slightly lowered from \(-0.36\) eV to \(-0.40\) eV. This can be understood by the fact that Br atoms adsorbed on Cu(110) act as electron acceptors, depleting some electron density from the surface and thereby slightly impeding the charge transfer from the Cu to PPP. In addition, copper end atoms terminating each PPP chain may have a higher contribution to the PPP doping due to their stronger C–Cu bonding. Nevertheless, such a doping should strongly depend on the PPP length. Hence, from our STS study showing no evidence of modification of the LUMO filling as function of the chain lengths we conclude that this process is negligible. We therefore conclude that hybridization between molecular \(\pi\)-states and copper states is the essential mechanism that permits partial filling of the LUMO, the lowering of the bandgap and results in metallic behaviour for the PPP nanowires. This effect is enhanced by the fact that PPP polymers are aligned in such a way as to be commensurate with the Cu(110) substrate, which maximizes here the molecule/surface interaction in comparison to the PPP/Cu(111) interface.

In conclusion, we combine here molecular self-assembly and covalent bonding processes to synthesize long-range ordered arrays of PPP and characterized their full \(\pi\)-band structure. We demonstrate the quasi-1D metallic character of these chains grown in a commensurable way on the Cu(110) surface. A simple Hückel TB model provides a good understanding of both ARPES and STS measurements and allows estimating effective inter- and intrachains resonance integrals. In addition, DFT calculations evidence the strong hybridization existing between the polymer and the substrate, which explains both bandgap reduction and metallization of the chains. Despite the substantial hybridization, bands characterizing the PPP wires remain strongly quasi-1D leading to conducting polymeric nanowires. A future systematic investigation of the electronic properties as a function of the chemical nature of the substrate and its geometry, as well as the role of the halogen species could be useful to tune the bandgap and the metallicity. This work is of particular relevance in the context of covalent organic nanostructures design for potential applications in nanoelectronic devices. Thus, the complete band dispersion of the polymer may be considered as the spectroscopic fingerprint of the polymerization process of such molecules as dBB on surfaces where the Ullman reaction can be achieved like gold, silver or copper and also as a recent promising approach on semiconductors.
Methods
Sample preparation. The experiments were carried out in ultrahigh vacuum at base pressures of 10^-10 mbar or better. The Cu(110) crystal was prepared by repeated cycles of sputtering (Ar + 1 keV) and annealing (750 K). 1,4-dibromobenzene molecules (Sigma Aldrich, purity 98%) were sublimed onto the surface (held at room temperature), using a leak valve at partial pressures of 10^-8 mbar. The PPP polymer presented here was obtained by reaching the saturation coverage of the clean Cu(110) surface. Under these conditions, the monolayer organic layer was accu-

rately after t = 9 min of deposition corresponding to 7 Langmuir. After deposition, the sample was post-annealed between 450 and 550 K for 5 min, to fully polymerize the system29.

STM/STS and ARPS measurements. STM and STS experiments were carried out using an Omicron LT-STM at a temperature of 5 K. STM images were recorded at constant tunneling current (0.2 nA) and constant bias voltage (capped to the sample). The dI/dV spectra were recorded in the open feedback loop mode (V

tunnel = 2 V) using the lock-in technique (peak to peak modulated voltage V

mod = 30 mV, f = 1.100 Hz). Spectra were normalized by subtracting a background corresponding to the clean surface. ARPS experiments were performed at the CASSIOPEA beamline (see Supplementary Note 1). The polymerization process and the orientation of the polymers were carefully analysed using the characteristic low-energy electron diffraction and X-ray photoelectron spectroscopy signatures (respectively, shown in Supplementary Figs 1 and 2, Supplementary Information and ref 26). The data presented in Fig. 3 were measured at 30 K using a Scienta R-4000 high-resolution hemispherical electron analyser, with linearly polarized 35 eV photons (for the experimental geometry see Supplementary Fig. 5).

Tight binding and first-principle band-structure calculations. The free-mole-

cule orbitals and energy levels were obtained using a standard Hückel model. The hopping integral between two first neighbour carbon atoms was assumed to be constant and equal to β. The Coulomb integral x was also assumed to be the same for all atoms and was adjusted to rigidly shift the band structure. STS differential conductivity has been obtained for a N = 6 oligomer by taking the square modulus of the TB-wave functions as a function of the quantized energies. Calculated ARPS intensity maps were obtained by taking the square modulus of the Fourier-transform of the calculated orbitals, according to the method presented in refs. 46,48. The calculated ARPS spectral intensity shown in Fig. 3e was obtained by calculating orbitals of three chains of 20 phenyl rings with an interchain coupling constant β of -0.15 eV.

DFT calculations were performed using the plane-wave pseudopotential code VASP30-33. We used the GGA formulation to approximate the exchange-correlation functional potential proposed by Perdew, Burke and Ernzerhof (PBE).

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Author contributions
Y.F.R., D.M., F.R., J.L.D., G.C. and D.F.P. conceived the project. Y.F.R., G.V., L.C. and L.M. participated to the elaboration of the material. G.V., Y.F.R., M.S., M.D.G., G.G. did the sample preparation and ARUPS measurements at the synchrotron. P.L.F. and F.B. assisted with ARUPS measurements on the CASSIOPEE beamline. G.V. with the help of M.S. performed the STS experiments under the guidance of B.K. Preliminary results were obtained by Y.F.R., L.C., M.D.G. and G.C. G.V. did the tight-binding modelling of STS and ARUPS data. L.L. and V.M. did the DFT calculations at surface. J.L.D. did the gas phase DFT calculations. Y.F.R. coordinated the work and wrote the manuscript with the help of G.V. All the authors commented critically on the manuscript.

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