Nanoelectrical analysis of single molecules and atomic-scale materials at the solid/liquid interface

Peter Nimalraj1*, Damien Thompson2,3, Agustín Molina-Ontoria4, Marilyne Sousa1, Nazario Martín4, Bernd Gotsmann1 and Heike Riel1

Evaluating the built-in functionality of nanomaterials under practical conditions is central for their proposed integration as active components in next-generation electronics. Low-dimensional materials from single atoms to molecules have been consistently resolved and manipulated under ultrahigh vacuum at low temperatures. At room temperature, atomic-scale imaging has also been performed by probing materials at the solid/liquid interface. We exploit this electrical interface to develop a robust electronic decoupling platform that provides precise information on molecular energy levels recorded using in situ scanning tunnelling microscopy/spectroscopy with high spatial and energy resolution in a high-density liquid environment. Our experimental findings, supported by ab initio electronic structure calculations and atomic-scale molecular dynamics simulations, reveal direct mapping of single-molecule structure and resonance states at the solid/liquid interface. We further extend this approach to resolve the electronic structure of graphene monolayers at atomic length scales under standard room-temperature operating conditions.

The elemental properties of carbon nanostructures are environment dependent. Interpreting the synergism between electronically active nanomaterials and their local chemical domain is pivotal to both scientific and technological interests. Scanning probe microscopy operated at cryogenic conditions with functionalized metal tips1,2 in combination with inorganic decoupling platforms3–4 is a widely adopted technique to examine the intramolecular structure of organic materials. In addition to probing matter under ultrahigh-vacuum conditions, scanning tunnelling microscopy (STM) operated at the solid/liquid interface5–11 has previously been demonstrated to record real-time molecular dynamics12–14, register ultrafast chemical reactions15, probe the structure of supra-molecular architectures16–18 and chemical-field-effect transistors19, and perform spectral analysis of molecules20–24 and real-space visualization of biomolecules2 at room temperature. It would be highly attractive to capitalize on principal findings from these two research disciplines to probe the precise electronic states in liquids and quantitatively describe the functionality of single molecules and two-dimensional nanostructures. The experimental challenge lies in addressing the electrochemical congestion present at the solid/liquid electrical interface at room temperature, which stems from the high mobility of organic molecules on metals, the dynamics of the solvent that serves as a liquid sheath and the detrimental effects of the metallic platform due to mixing of metal bands with discrete molecular electronic states25. A potential strategy is to employ nonpolar, highly viscous liquid sheaths with low affinity towards the substrate and to engineer a spacer layer that electrically disconnects the organic material of interest from metal-induced perturbations. The bottom-up construction of the spacer is crucial as it requires an optimal thickness that suppresses dominant interfacial charge transfer whilst channelizing a small population of electrons to tunnel through the remnant barrier to excite the adsorbed organic material.

Here we report real-space imaging of the intrinsic electronic structure of low-dimensional nanostructures adsorbed on a wide-bandgap organic insulating spacer layer in an electrochemically inert liquid environment at room temperature. The key advance lies in sandwiching the delicate conductive organic analyte between the ultrathin insulating organic membrane, which electronically decouples the material of interest from the metal, and the top-liquid sheath, which protects the molecules adsorbed on the spacer-coated gold and the STM tunnel gap against external contamination. In the case of single molecules, this multicomponent architecture based on the bottom-spacer and top-liquid sheath also serves to reduce single-molecule mobility, as discerned from molecular dynamics simulations. This reduction in the motion of molecules is essential for resolving single-molecular structure and performing consistent spectroscopic analysis. We further show that the organic spacer layer combined with in situ spectroscopy can sense energy-level shifts (energy resolution ~50 meV) in chemically tailored molecules, which are rationalized using density functional theory (DFT) calculations. Importantly, the organic spacer serves as an intermediate cushioning layer that prevents metal-induced topological defects in single-layer graphene, thus opening a route towards engineering ultraflat two-dimensional nanostructures on metal supports.

Organic spacer/gold interaction in liquids

n-C60H32 (solubilized in 1-phenyloctane) spray-deposited on Au(111) (Methods) at room temperature26 forms close-packed two-dimensional lamellar structures27,28. The formation of ordered alkyl-organic layers and their lower dielectric constant (~2) in comparison with inorganic spacers (NaCl, ε:~6) make them...
excellent electronic decoupling platforms. The organic layers are homogeneous with an overall thickness of ∼4 Å, confirmed by ex situ high-sensitivity ellipsometry (Supplementary Section 1.1). At the nanoscale the n-C16H33 units are seen to adsorb in a fully extended conformation with their backbone axis parallel to the underlying Au(111) substrate, recorded in the liquid-cell set-up (see schematic Fig. 1a) as shown in the in situ STM image (Fig. 1b) recorded in 1-phenyloctane solvent (for spacer-layer-stability tests see Supplementary Section 1.2). The average length of an individual n-C16H33 unit is (3.75 ± 0.05) nm, calculated from the in situ STM topographs and is in agreement with previous STM reports on identical systems.25,26 The van der Waals binding energy between the alkyl-spacer and Au(111) calculated from molecular dynamics structures (Fig. 1c, for details see Supplementary Section 8.1) is (−2.6 ± 0.3) eV nm−2, which corresponds to (−119 ± 14) kcal mol−1, indicating a strong adsorption of the long-chain alkanes on Au(111), in agreement with the highly ordered structures shown by in situ STM (ref. 25). The inset in Fig. 1b is a high-resolution in situ STM snapshot of the n-C16H33 units in a silicone oil ((Si(CH3)3O)n)n environment at room temperature, arranged in an energetically favourable flat-lying configuration. The intermolecular spacing is ∼0.5 nm and the packing energy between adjacent alkane molecules is calculated to be (−49 ± 6) meV per carbon, based on atomic-scale modelling (Supplementary Section 8.1). This corresponds to (−729 ± 89) meV nm−2, indicating a tightly packed monolayer.27 The in situ STM data confirm that the long-range geometric ordering and intramolecular configuration of the n-C16H33 spacer layer remains unperturbed at the interface with silicone oil, reflecting the robustness of the organic spacer architecture. The preservation of organic material integrity by silicone oil, its electrochemical inertness (dielectric constant: 2.5) and density (0.93 g ml−1 at 25 °C), and the minimization of surface contamination stemming from surrounding hydrocarbons and moisture over time (Supplementary Fig. 2d,e) justify the use of silicone oil as a good liquid sheath model system for in situ probing of soft materials and for imaging atomic-scale features of planar surfaces13 under a wide working energy range at room temperature.28

Tracking C60 dynamics on solid surfaces in silicone oil

The geometry, intermolecular mechanics and electronic structure of C60 molecular ensembles directly coupled to noble metals in low-density liquids have been discussed at length.24,29,30 Here, we focus on the structure, dynamics and energetics of single C60 molecules adsorbed on spacer-coated Au(111) in the high-density liquid silicone oil. Of particular interest is the interaction strengths between C60 and the spacer layer as well as the subtle interplay between the spacer layer and silicone oil in damping molecular surface diffusion. Following the evaporation of 1-phenyloctane solvent, the alkyl-spacer-coated Au(111) dosed with pristine C60 molecules (99.9% purity, purchased in powder form from Sigma-Aldrich and solubilized in tetrahydrofuran) is encapsulated in a liquid cell with silicone oil as schematically represented in Fig. 1a (for C60 solution synthesis see Supplementary Section 2). The large-area in situ STM image (Fig. 1d) shows a random distribution of C60 molecules with no apparent preferential adsorption site on the spacer layer. The C60 molecules reveal the typical caged structure with a diameter of (6.5 ± 0.5) Å (measured over ∼80 individual C60 units) close to the hard-sphere diameter value of 7.1 Å for a single C60 molecule.20 The spread in the measured diameter value originates from the electronic structure disparity present between the adsorbate and substrate, which is markedly large in the case of a high-conductance C60 molecule adsorbed on a near-insulating alkyl-spacer layer. This difference in electronic structure is evident in Fig. 1d where the C60 molecules are better resolved than the underlying spacer layer imaged at a common tunnel-current set point. Note it has not been possible to resolve the multicomponent organic architectures in silicone oil medium simultaneously; however, we observed that the underlying alkyl-spacer structure becomes more prominent when imaged independently (generally at higher negative sample bias and tunnel-current set points) revealing sub-molecular image contrast (Fig. 1b, inset). The variations in the local composition of the spacer layer probably originating from the corrugations in the underlying metallic surface are captured in the in situ STM snapshot (Fig. 1d, regions colour coded in green and yellow).

Atomic-scale simulations of film formation support the in situ STM data, indicating that the C60 molecules adsorb directly on the spacer-coated Au(111). We employ a modified gold van der Waals potential31 and model the assembly of the spacer layer from a starting structure of clusters of n-C16H33 molecules suspended over a bare Au(111) surface (for details see Supplementary Section 8.2). There is negligible deviation in molecular adsorption energies (−16 ± 1) kcal mol−1 for C60 adsorbed at different sites on the spacer layer, with molecule methylene and methyl regions showing comparable, weak van der Waals attraction for C60. The computed structures indicate that the spacer layer is sufficiently stable to prevent the exchange of alkanes for C60 on gold (Supplementary Section 8.2). Taken together, the in situ STM electronic imaging and atomic-scale modelling data show that the spacer layer separates the C60 molecule at an ideal distance from Au(111), which suppresses direct charge transfer from the metal reservoir but allows a small population of tunnelling electrons.

Interestingly, the isolated C60 molecules as shown in the spatially magnified three-dimensionally represented in situ STM image (Fig. 1e) were stationary and in a kinetically near-frozen state. This is typical of C60, behaviour previously observed only under cryogenic temperatures (4–8 K; refs 32,33). Damping of molecular motion at room temperature has been reported previously for single C60 molecules trapped on engineered porous organic templates4, graphene Moiré superlattices5 and even macromolecules36 on alkyl-coated surfaces. Imaging a single molecule under different positive biasing conditions with a fixed tunnel-current set point exhibited only a bias-dependent profile variation (Fig. 1fg). This is reminiscent of an electronic effect but reveals no traces of sub-molecular structure. In control experiments, a C60 molecule placed in direct contact with Au(111) in a silicone oil environment was imaged (Supplementary Fig. 3b). The structure of the C60 molecules remains unchanged but a comparable drop in mobility was observed that contradicts previous in situ STM observations of C60 dynamics on defect-free Au(111) (ref. 24) in low-density solvents (n-tetradecane)13, where the molecules exhibited high mobility at room temperature. To resolve this puzzling finding of reduced molecular mobility in silicone oil, we rely on molecular dynamics simulations of neat silicone oil (described in detail in Supplementary Section 8.3) that highlight the rigidity of the silicone oil, with a low root mean square fluctuation of (1.2 ± 0.3) Å, computed in the atomic positions, and a large favourable intermolecular packing energy of (−125.6 ± 4.3) kcal mol−1. The computed silicone oil structures (Supplementary Fig. 20) show that the individual molecular strands are highly intertwined through van der Waals contacts and the presence of this top layer of silicone oil (Fig. 1h) reduces the computed mobility of the C60 on both spacer-coated and bare gold (for details see Supplementary Section 8.5). The in situ STM observations and molecular dynamics simulations of C60 dynamics on bare and spacer-coated gold and previous reports of C60 dynamics on gold in low-density liquids13,24 confirm that silicone oil plays a dominant role in reducing molecular motion.

In situ mapping of C60 molecular energy levels

Dielectric spacers are known to weaken metal/molecule interaction energies and thus preserve free-molecular structure3,27,38 recorded by
ultrahigh-vacuum-based STM/scanning tunnelling spectroscopy (STS). However, mapping electronically decoupled organic electronic structures at the solid/liquid interface at room temperature has not been demonstrated with high spatial and energy resolution until now. We focus here on the electronic influence exerted by the spacer layer on the C60 energy states in liquids using local-point probe in situ STS. Several tips were tested during the STS measurements under silicone oil to confirm reproducibility of the spectral data. To record the spectral data directly, the STM tip is positioned above a well-resolved individual C60 unit (as shown in Fig. 1f) and the voltage is swept with the feedback loop shut off to eliminate tip–sample height fluctuations during measurements (see Methods for in situ STS parameters). Figure 2a shows a typical differential conductance (dI/dV) spectrum of an individual C60 molecule adsorbed on spacer-coated Au(111) that reflects the energetic positions of the frontier molecular orbitals with a discernable low-conductance gap region. The dI/dV spectrum exhibits well-structured molecular resonance peaks centred around −1.1 eV and +0.9 eV with respect to the Fermi level, which is in contrast with the weak spectral signal of C60 adsorbed directly on Au(111) in silicone oil (Supplementary Fig. 3b) and with previous in situ STS reports of C60 adsorbed directly on Au(111) in an n-tetradecane environment13. The peak positions correspond to the highest occupied molecular orbital (HOMO)- and the lowest unoccupied molecular orbital (LUMO)-derived resonances of the C60 molecule. The observed energetic positions are consistent with previous STS observations where lateral electrostatic interactions between C60 and other organic complexes were used to elevate the C60 molecule locally above the Au(111) surface16. In particular, the localization of the LUMO-derived peak at 0.9 eV above the Fermi edge tends to dominate the charge transport in pristine C60 (refs 39,40), which confirms minimal charge transfer at the metal/molecule electrical interface16,42 as a result of the spacer layer.

The overall spectral line shape of the dI/dV spectrum recorded on a specific C60 molecule reveals no significant variation, but slight fluctuations in the molecular-orbital-associated peak positions were observed as measured on several individual C60 molecules. We attribute this fluctuation to two key reasons: variation in the tunnelling conditions between each measured molecule in silicone oil; and spacer-layer-induced alterations in the binding geometry and molecular orientation. To map the spread in the molecular orbital peak positions quantitatively we performed a series of in situ STS measurements over ~80 individual C60 molecules adsorbed on the spacer layer. We derived a conductance gap (energy difference between the HOMO- and LUMO-derived molecular resonance peaks) of (2.2 ± 0.3) eV at room temperature, which is close to the previously reported ΔE value of 2.6 eV where the coupling between C60 and the underlying metal was reduced14. It also compares reasonably well to our DFT calculations that predict a ΔE value of 2.8 eV for a single C60 molecule. The calculated frontier molecular orbital wavefunctions are shown in Fig. 2b and the associated experimentally
observed peak positions are indicated by the green arrows (details of the DFT calculations are given in Supplementary Section 8).

**Detecting energy-level shifts in chemically modified C\textsubscript{60}**

To determine whether the proposed metal/molecule weakening mechanism applies beyond pristine molecules, we performed additional experiments on chemically modified C\textsubscript{60} molecules. The interest in functionalizing pristine C\textsubscript{60} molecules with molecular moieties is driven by the potential application of the C\textsubscript{60} cage to act as a stable chemical anchor\textsuperscript{44} on the contact metal. This stability of the anchor group ensures well-defined molecular conductance\textsuperscript{46}, a germane feature in engineering single-molecule circuits. Of particular interest is the electronic coupling between the C\textsubscript{60} anchor and the chemically linked molecular moiety. C\textsubscript{60} molecules functionalized with exTTF-p-cyclophane (chemical structure shown in Fig. 3a) were deposited from solution (see Supplementary Section 4 for full details on the synthesis and characterization of the functionalized C\textsubscript{60} molecules) onto spacer-coated Au(111) by the same procedure used for pristine C\textsubscript{60}. The functionalized C\textsubscript{60} molecules (hard-sphere diameter \(\sim 0.68\) nm) resolved with nanometre-scale spatial resolution (in situ STM image, Fig. 3c, inset) showed no preferential adsorption site on the spacer layer but exhibited similar damping in molecular motion in the silicone oil medium as in the case of the pristine C\textsubscript{60} molecules. Molecular dynamics simulations indicate a reduction in mean square deviation value from 0.25 to 0.15 nm\(^2\) for functionalized C\textsubscript{60} adsorbed on spacer-coated Au(111) in the presence of silicone oil (computed structure shown in Fig. 3b).

The molecular dynamics calculations provide a quantitative description of functionalized C\textsubscript{60} mobility on spacer-coated Au(111) and substantiate the *in situ* STM observations. Molecular dynamics models were used to calculate the physisorption energies between the functionalized C\textsubscript{60} molecules and the spacer layer, thus providing a deeper understanding of the adsorbate/substrate interfacial interactions. A physisorption energy of \((-27 \pm 4\) kcal mol\(^{-1}\)) is calculated for functionalized C\textsubscript{60} on spacer-coated Au(111), which is greater than the \((-16 \pm 1\) kcal mol\(^{-1}\)) calculated for pristine C\textsubscript{60}. This confirms the participation of the functional groups in the physisorption on the spacer layer (Fig. 3b). Simulations reveal an energetic penalty of \((24 \pm 9\) kcal mol\(^{-1}\)) for penetration of the functionalized C\textsubscript{60} through the spacer layer and direct adsorption on bare gold (for details see Supplementary Section 8.4), indicating that the molecules will prefer to physisorb directly on the spacer layer and remain decoupled from gold.

The spectroscopic measurements were performed only on well-spaced functionalized C\textsubscript{60} molecules (Fig. 3c, inset), to eliminate parallel conduction pathways occurring through the free end of the anchored molecular moiety. Figure 3c shows a representative \(dI/dV\) spectral signal acquired over a single functionalized C\textsubscript{60} molecule. Two distinct molecular resonance peaks in an energy span of \(\pm 1.5\) eV are visible in the \(dI/dV\) spectra centred around \(-0.5\) and \(+0.7\) eV, which correspond to the HOMO and LUMO peak positions, respectively. The DFT-calculated HOMO and LUMO eigenstates in the gas-phase molecule (indicated by the green arrow) show the HOMO localized at the interface between the C\textsubscript{60} anchor and the functional group whereas the LUMO is localized on the C\textsubscript{60} lobe, comparable to previous DFT calculations on C\textsubscript{60}-terminated molecular dumbbells\textsuperscript{49}, in which the LUMO remains unperturbed on molecular functionalization. Also the LUMO-derived peak position for the functionalized C\textsubscript{60} molecule with respect to the Fermi edge is equivalent to the LUMO position of pristine C\textsubscript{60} but there is a significant shift in the HOMO peak position closer to the Fermi edge resulting in a decreased conductance gap for the functionalized C\textsubscript{60} molecule. Having measured more than \(\sim 65\) isolated functionalized C\textsubscript{60} molecules, we derived a \(\Delta E\) value of \((1.1 \pm 0.2\) eV), which is close to our DFT-calculated \(\Delta E\) value of 0.9 eV for a functionalized C\textsubscript{60} molecule. The observed reduction in \(\Delta E\) for a functionalized C\textsubscript{60} molecule in comparison with pristine C\textsubscript{60} (\(\Delta E = 2.2 \pm 0.3\) eV, above) suggests good electrical coupling between the anchor (C\textsubscript{60}) and the core of the chemically linked molecular moiety (exTTF-p-cyclophane) with minimal functionalization-induced symmetry disruption to the C\textsubscript{60} cage. This ability to detect and discern the origin of the electronic shifts on C\textsubscript{60}-anchor-based molecular complexes in a selective manner is possible only by weakening the tendency of the anchor (C\textsubscript{60}) molecular orbitals to hybridize with gold\textsuperscript{46}. If allowed to occur, this hybridization results in significant broadening and clouding of intrinsic spectral features (see Supplementary Fig. 12b for *in situ* STS measurements of functionalized C\textsubscript{60} on bare gold).

**Atom-scale analysis of graphene in silicone oil**

Tuning metal/molecule electronic interactions by inserting insulating organic spacers at the metal/molecule interface is not limited to single molecules. In addition we can electronically decouple single-layer graphene from metals and record its unique band structure in liquids. Liquid-phase-exfoliated graphene (see Supplementary Section 6 for sample preparation) in low-boiling-point solvents was spray-deposited onto spacer-coated Au (111) and placed directly in a liquid cell packed with silicone oil. The single-layer graphene flakes possess well-defined edges with a lateral flake size ranging from 0.2 to 1.0 \(\mu\)m and a characteristic thickness of 0.35–0.5 nm based on *in situ* STM and intermittent-contact atomic force microscopy (AFM; Methods). The atomic plane of carbon atoms of graphene is evident from the high-resolution constant-current *in situ* STM image (Fig. 4a). The graphene crystal structure can be distinctly imaged in silicone oil at room temperature with a resolution on par with graphene resolved using cryogenic STM (refs 47,48). The hexagonal lattice of carbon atoms can also be resolved at high tunnel-current set points (\(\sim 1.8\) nA), reflecting the structural stability of the interface between the graphene and the spacer layer, previously observed for graphene on two-dimensional insulating platforms\textsuperscript{67}. Molecular dynamics simulations (Supplementary Section 8.6) revealed a binding energy of \(-1.0\) and \(-3.8\) eV nm\(^{-2}\) for graphene on spacer-coated and bare gold, respectively, in good agreement with dispersion-corrected DFT (ref. 49). Summing over the graphene/spacer and spacer/gold interfaces...
yields a net energy of $-4.3 \text{ eV nm}^{-2}$, indicating a preference of $\sim 500 \text{ meV nm}^{-2}$ for graphene adsorption on the spacer layer rather than on bare gold. This implies that a single layer of graphene can lie on the spacer layer without disrupting its geometric order (see Supplementary Section 8.6 for binding-energy calculations). Large-area \textit{in situ} STM images show that the overall graphene landscape does not suffer from a high density with topological defects (Supplementary Fig. 14a) when compared to single-layer graphene placed directly in contact with Au(111) as shown in Fig. 4c, which reveals regions where the lattice symmetry is clearly disrupted in addition to the presence of local-point defects on the atomic lattice consistent with previous reports of graphene adsorbed on noble metals\textsuperscript{46–50}.

The electronic disentanglement of graphene from the metal through spacer-layer mediation was verified by \textit{in situ} STS measurements. We then compared the spectral signatures between graphene adsorbed on the spacer layer and graphene directly in contact with Au(111). Figure 4b shows a well-resolved \(dI/dV\) spectrum (acquired over the sites indicated in the \textit{in situ} STM image, Fig. 4a) with a distinctive line shape resembling the energy spectrum of graphene, similar to previous STS observations of graphene on non-metallic substrates\textsuperscript{31–33}. The minimum in each of the directly measured spectral curves corresponds to the charge-neutral Dirac point \((E_D)\) located at positive bias energy slightly above the Fermi level. To quantify the spatial variation of the energy landscape, \(dI/dV\) spectra were acquired over a monolayer graphene area of 100 nm\(^2\) under identical tunnelling conditions, which yields a mean \(E_D\) of \((48 \pm 14) \text{ mV} \) (Dirac point shift histogram; Fig. 4b, inset) indicative of minimal substrate-induced doping. The spectral data for graphene on spacer-coated gold are remarkably different from the spectral signature for graphene on bare gold (Supplementary Fig. 15) where two significant dips at \(E_F\) and \(E_D\) now located at $\sim \pm 350 \text{ mV}$ are discernible. This observation of \(E_D\) occurring at higher positive bias energy ($\sim \pm 350 \text{ mV}$), with respect to the Fermi edge, clearly points to alignment of the Fermi levels occurring through charge transfer between the gold surface and graphene. This shift in \(E_D\) is comparable to previous analytical modelling reports of metal doping on graphene\textsuperscript{54} and low-temperature STS (ref. 48) observations of graphene in contact with metals.

In addition to mapping the variations in electronic structure by detecting the shifts in \(E_D\) for graphene, the bonding configuration of in-plane atoms in graphene can be clearly resolved in liquids at room-temperature imaging conditions. The atomic positions of the sp\(^2\)-bonded carbon atoms and the respective bond lengths can be precisely identified on the basis of high-resolution \textit{in situ} STS images. Figure 4d shows a three-dimensional \textit{in situ} STM image of a locally magnified region on single-layer graphene adsorbed on spacer-coated gold imaged under silicone oil. The C–C bond distance measured over a well-resolved atomic hexagon as seen in Fig. 4d is $\sim 1.42 \text{ Å}$. An average C–C bond distance (measured over all atomic hexagons over an area of 100 nm\(^2\) of single-layer graphene) yields a value of $(1.38 \pm 0.12) \text{ Å}$. Imaging over the same graphene lattice region (Fig. 4d) and incrementing the bias in small steps revealed that the local electronic contrast over the atomic hexagons is tunable and is dependent on the energy of the tunnelling electrons (Fig. 4e). These observations highlight that useful information from bond length fluctuations to atomic-scale contrast can be extracted from two-dimensional materials in a liquid medium at room temperature. Resolving in real-space the atomic positions in graphene immersed in liquids using a non-functionalized STM probe reflects the versatility of this technique. Importantly, the inherent nature of silicone oil, with its rigid molecular packing structure and electrochemical inertness, provides the necessary high stability during imaging at room temperature by minimizing liquid-induced thermal fluctuations, preventing tip–apex contamination and parasitic faradaic current at the STM junction.

**Outlook**

We provide compelling evidence for the singular read-out of integral mono-molecular and atomic-material electronic structure on an insulating organic layer using \textit{in situ} STS and STS. The engineering of an organic membrane with ångström-scale thickness control, scalable to industrial needs, can be extended to printed electronics where a percolating organic nanomaterial network is requisite. The findings deepen our understanding on the dependence of molecular electronic states of pristine and chemically tailored carbon particles on their interfacial structure in complex environments, thus suggesting a reliable route towards chemical sensing of complex analytes. It is expected that the spatial resolution capability can be further advanced by chemically terminating the scanning probe tip–apex\textsuperscript{1,2,23,51}. We draw particular attention to the potential of...
Figure 4 | Atom-scale mapping of graphene electronic structure. a High-resolution three-dimensional in situ STM topographic image of a graphene monolayer adsorbed on the organic spacer-coated Au(111) surface (I = 15 pA, V = 0.8 V, scan bar, 2 Å). b, dI/dV spectral curves acquired over the locations indicated in a revealing a characteristic graphene linear band structure. The green arrow indicates the Dirac point (E_D; feedback loop parameters I = 150 pA, V = 1.5 V). The inset shows the histogram of fluctuations in the measured Dirac point over a graphene area of 100 nm². c, Wide-angle in situ STM image of topological defects on a monatomic graphene adsorbed directly on Au(111) in silicone oil environment (I = 40 pA, V = 1 V, scan bar, 2 nm). d, High-resolution in situ STM of graphene on spacer-coated Au(111) where the atomic positions are well resolved (scan area: 0.75 nm × 0.8 nm). The red and blue spheres marked on the graphene lattice represent the two atoms per unit cell depicted as A and B. e, Bias-dependent imaging of the graphene atomic hexagons in silicone oil (scan area: 0.75 nm × 0.8 nm). The in situ STM/STS measurements on monolayer graphene adsorbed on the spacer-coated and bare gold were performed in a silicone oil environment at room temperature.

Methods

Preparation of gold substrate. One hundred nanometres of Au (99.99%, FHR Anlagenbau GmbH) was sputter-deposited (Cluster system CS320S, Von Ardenne, Dresden) in an argon atmosphere onto CaF₂(111) substrates (2 cm × 2 cm, pre-heated at 500 °C for 3 h) at a rate of 10 Å s⁻¹ and a base pressure of 2 × 10⁻⁷ mbar. The dominant (111) texture for the resulting metal films was confirmed using high-resolution X-ray diffraction.

Spacer-layer fabrication. n-C₃₆H₇₃ (Triacantane, 98%) was purchased from Sigma-Aldrich and used without any additional purification procedure. A near-saturated solution of this compound was prepared in 1-phenyloctane solvent (Sigma-Aldrich) and subjected to gentle bath sonication for 1 min. Half a millilitre of this solution was spray-coated (Dual action, Evolution Airbrush, FHR Anlagenbau GmbH) was sputter-deposited (Cluster system CS320S, Von Ardenne, Dresden) in an argon atmosphere onto CaF₂(111) substrates (2 cm × 2 cm, pre-heated at 500 °C for 3 h) at a rate of 10 Å s⁻¹ and a base pressure of 2 × 10⁻⁷ mbar. The dominant (111) texture for the resulting metal films was confirmed using high-resolution X-ray diffraction.

In situ STM/STS and AFM measurements. STM measurements were performed in constant-current mode using a Digital Instruments Nanoscope IIIa scanning tunnelling microscope (Multimode System, scanner model: E-Scanner). Nanoscale calibration of the scanner was performed using a freshly cleaved highly oriented pyrolytic graphite surface (x, y axes) and a Au(111) on mica substrate (z axis). Bias voltages imply sample voltage with respect to the STM probe. For the STM probe, mechanically cut Au wire (0.25 mm, Good Fellow GmbH) was used. The quality of the tip was constantly monitored by imaging and acquiring dI/dV spectra over clean Au(111)-mica samples as reference. Low tunnel-current set points were employed to avoid tip-induced motion of the single molecules. For the in situ STS measurements the feedback loop is turned off (~30 s) after ensuring the tip is positioned above the molecule of interest and the voltage is swept while the current is recorded. The tip drift rate is ~1 nm min⁻¹ and it takes ~2 s to record a single spectral curve. Each dI/dV spectrum thus recorded (on individual and non-interconnected pristine fullerenes and functionalized fullerene molecules) was averaged over 20 voltage sweeps. To check the reproducibility of the differential conductance spectra reported for the single molecules we thoroughly analyse each forward and reverse voltage sweep (tip stability and image resolution is constantly monitored) for each individual case and the molecular structure was verified before and after acquisition of the spectral curve. A custom-built liquid-cell holder was used for in situ STM and STS measurements. AFM measurements were performed on the same system with an interchangeable optical head in intermittent-contact mode using n-type silicon tips with aluminium coating on the detector side (Nanosensors, 200–500 kHz resonant frequency). The in situ STM/STS set-up is located in a noise-free laboratory (Supplementary Section 3).

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Author contributions

P.N. designed and performed the in situ STM/STS experiments. M.S. carried out the ellipsometry measurements and data analysis. A.M.-O. and N.M. synthesized the functionalized C60 molecules and performed the electrochemical characterization. D.T. designed and performed the molecular dynamics simulations and DFT calculations. All authors contributed and commented on the manuscript and analysis of the data.

Additional information

Supplementary information is available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to P.N.

Competing financial interests

The authors declare no competing financial interests.