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Effective elastic properties of a van der Waals molecular monolayer at a metal surface

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Adsorbing anthracene on a Cu(111) surface results in a wide range of complex and intriguing superstructures spanning a coverage range from 1 per 17 to 1 per 15 substrate atoms. In accompanying first-principles density-functional theory calculations we show the essential role of van der Waals interactions in estimating the variation in anthracene adsorption energy and height across the sample. We can thereby evaluate the compression of the anthracene film in terms of continuum elastic properties, which results in an effective Young’s modulus of 1.5 GPa and a Poisson ratio =0.1. These values suggest interpretation of the molecular monolayer as a porous material—in marked congruence with our microscopic observations.

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Understanding how materials respond to stress and, subsequently, having the ability to predict the resultant deformation (strain) are fundamental to virtually all practical applications of materials. In-depth understanding of materials-specific stress-strain relationships impacts a wide range of scientific fields as varied as the geological deformation of rock1 during plate tectonics and the posture-induced deformation of lumbar disks resulting in herniation.2 It also has significance for applications at the nanoscale: e.g., in the development of molecular-scale coatings, knowledge of the elastic properties of the employed materials is essential for predicting their ability to maintain integrity under the deformation of their substrate. Yet investigations of molecular overlayers at solid surfaces have generally not addressed issues related to the elastic properties of films, with a few exceptions such as cantilever measurements of magnetostriction.3–5 Currently the formation of molecular networks on metal surfaces garners significant attention because of the patterns of unprecedented size and versatility that are produced.6–10 The present study aims to complement the advances in creating molecular surface patterns by developing a framework for understanding their elastic properties.

The focus of this study is a system that was chosen because of the simplicity of the geometry of its components and of their interaction with one another: anthracene on Cu(111). Anthracene consists of three fused benzene rings (Fig. 1 inset) and adsorbs flat on Cu(111), similar to other acenes.11–17 Dispersive or van der Waals (vdW) interactions play an important role in adsorption of acenes (such as benzene) on low-corrugation metal surfaces. In a prior study18 some of us addressed the physisorption of benzene and found that with density-functional theory (DFT) calculations in the generalized gradient approximation, a negligible adsorption energy of ~20 meV results,19 which favors the ring centered over a hollow site compared to an atop site by ~7 meV. In contrast, in similar calculations that also account for vdw interactions,20 markedly higher adsorption energies of 0.55 eV and 0.53 eV at hollow and atop sites, respectively, are found. We extend such calculations to anthracene on Cu(111) and find substantially higher binding energies of 0.95 eV and 0.93 eV at hollow and bridge sites, respectively, [see Figs. 1(a) and 1(b)]. There is a similar difference in adsorption configuration of anthracene on Cu(111) (a) at a hollow site and (b) at a bridge site. Each adsorption configuration is threefold rotational degenerate. Panel (c) shows an STM image of an anthracene superstructure. The unit cell is indicated as a white dotted parallelogram. The superstructure can be described by a geometric motif resembling a fish and its reflection (red solid polylines and green dotted polylines). The inset shows a schematic of anthracene. Imaging parameters: current =20 pA, bias =–3.1 V, and size =90 × 100 Å2.

FIG. 1. (Color online) Adsorption configuration of anthracene on Cu(111) (a) at a hollow site and (b) at a bridge site. Each adsorption configuration is threefold rotational degenerate. Panel (c) shows an STM image of an anthracene superstructure. The unit cell is indicated as a white dotted parallelogram. The superstructure can be described by a geometric motif resembling a fish and its reflection (red solid polylines and green dotted polylines). The inset shows a schematic of anthracene. Imaging parameters: current =20 pA, bias =–3.1 V, and size =90 × 100 Å2.
In this study we address anthracene coverages on Cu(111) that are complete: no bare surface areas remain. We find a large range of adsorbate superstructures which distinguish themselves in geometry, fraction of molecules adsorbed at hollow vs bridge sites and total coverage. In particular, our measurements show that higher coverage leads to an increasing fraction of the molecules in less-favorable sites. We show that this reduction in average adsorption energy with lateral compression can be interpreted as elastic behavior of the monomolecular film.

All measurements were performed in a low-temperature scanning tunneling microscopy (STM) chamber built on site. Sample preparation involved the typical cycles of sputtering and annealing followed by cooling of the bare surface to \( \approx 80 \) K. Deposition of anthracene through a leak valve proceeded in three incremental steps, each time dosing for \( \approx 1 \) min at \( 2 \times 10^{-9} \) torr (an order of magnitude above the chamber base pressure) and imaging to check the coverage. All coverages shown coexist on the resultant surface. Modeling of the adsorption of anthracene involves a nonself-consistent approach in which the RPBE functional\(^{23}\) was used self-consistently to obtain electron densities, which in turn were used to evaluate the nonlocal correlation energies through a first-principles DFT treatment for vdW interactions.\(^{20}\) All calculations use the projector augmented wave method\(^{24}\) implemented in the grid-based real-space code GPAW (Ref. 25) and were corroborated using vdW-DF enabled DACAPO.\(^{26}\) The supercell was a \( 5 \times 5 \) Cu(111) slab of five layers topped by 21 Å of vacuum, with an anthracene molecule placed parallel to the Cu(111) surface. The Brillouin zone was sampled only at the \( \Gamma \) point. Wave functions, electron densities, and potentials were represented on a 0.16 Å real-space grid. We calculated the adsorption of anthracene at a hollow and a bridge site, to find (besides the mentioned difference in energy) equilibrium heights of the molecule’s carbon atoms above the top-layer substrate atoms: 3.49 Å and 3.52 Å, respectively, a small difference, in good agreement with data obtained on benzene (see Ref. 18 and references therein) but in disagreement with the x-ray standing-wave measurements on pentacene/Cu(111) of Ref. 27. In addition to the adsorption geometries discussed above and shown in Figs. 1(a) and 1(b), we investigated a set of alternate configurations which, however, turned out to have prohibitively high energies.

Despite the relative simplicity of the adsorption system, we find highly polymorphic patterns, many of which are characterized by large and complex unit cells. Figure 1(c) shows as an example a \((\sqrt{5} \times \sqrt{5})\) superstructure with a unit-cell area of 957 Å\(^2\), corresponding to 170 substrate atoms.

Given 11 molecules per unit cell, not every molecule in the unit cell interacts directly with every other; consequently, it may be useful to describe the unit cells in terms of the underlying geometric motif that contains the intermolecular geometry. Figure 1(c) highlights how in this superstructure six molecules surround a seventh one in a shape that resembles a “fish.” Both the central molecules and one of the “fin” molecules adsorb at less favorable bridge sites, with the remaining molecules adsorbing at the optimal hollow sites. The unit cell consist of a fish shape and an “antifish” shape (with a few shared molecules) connected by a glide-reflection line. In total we can identify five different surface patterns, all of which can be found on the same sample in the same preparation.

The \((1 \times 1)\) superstructure, shown in Fig. 2(a), is the least dense one observed; its unit cell contains only one molecule, which is adsorbed at a hollow site. All molecules are aligned in parallel and are adsorbed at equivalent adsites. There are 17 substrate atoms in the unit cell, giving a coverage of \(1/17\) ML = 0.059 ML with the monolayer (ML) referred to the substrate atom density.

Figure 2(b) shows a \((2 \times 2)\) superstructure with a unit-cell size of 631 Å\(^2\), in which six molecules form a symmetric hexagon surrounding a seventh; similar to the structures of Figs. 1(c), 2(c), and 2(d), the surrounding molecule is adsorbed at a bridge site. The density of this film is 16 substrate atoms per molecule, corresponding to coverage of 0.0625 ML.
The $^{10}_{-6}^{19}$ superstructure of Fig. 2(c) consists of rows of the fish motif of Fig. 1(c) arranged in head-to-tail fashion; between these rows two parallel molecules are found adsorbed at the optimal hollow site. At a size of 1159.9 Å$^2$ the unit cell comprises 13 molecules, of which nine come to rest on optimal hollow sites. The coverage is 0.0631 ML.

Figure 2(d) repeats the superstructure of Fig. 1(c); it corresponds to coverage of 0.065 ML. Finally Fig. 2(e) shows a $^{8}_{-3}^{12}$ superstructure with a unit-cell size of 507 Å$^2$ that features alternating bands of molecules equally distributed between hollow and bridge sites. There are six molecules in each unit cell for a total coverage of 0.067 ML, the densest coverage observed.

In total, we observed five different phases that span a compression of the surface area per molecule by more than 10%. The broad geometric diversity of the surface patterns may appear astonishing; however, similar observations have been made in the study of the statistical mechanics of granular media, even as conventional ones as Basmati rice spread out on a vibrating plate. Bates and Frenkel showed that if compressed from the coverage of Fig. 2(c) to 0.065 ML. Finally Fig. 2(b) shows a $^{8}_{3}^{12}$ superstructure with a unit-cell size of 507 Å$^2$ that features alternating bands of molecules equally distributed between hollow and bridge sites. There are six molecules in each unit cell for a total coverage of 0.067 ML, the densest coverage observed.

In the different superstructures a varying fraction of the copper interlayer spacing is mainly accommodated by reduction in pore volume rather than requiring transverse expansion of the object. If a porous material is deformed, a relatively small amount of matter has to deform compared to the overall volume of the object, and a relatively low Young’s modulus results. This is also found here, suggesting that the closer packing of the molecules observed in our STM experiments can indeed be interpreted as compression of a porous material, revealing a remarkable congruence of macroscopic and microscopic behavior. It should be pointed out that we have treated the anthracene molecules as hard, noninteracting particles; the interactions evaluated here stem exclusively from the variation in the vdW interaction of anthracene across the substrate.
In summary, we have investigated the adsorption geometry of anthracene on Cu(111) and found—despite the adsorption system’s comparative simplicity—a large number of highly ordered superstructures with coverage varying by >10%. The polymeric nature of the anthracene film coincides with the behavior of macroscopic granular media of similar aspect ratio. Modeling of the adsorption behavior of the system in first-principles DFT with inclusion of vdW interactions provides us parameters which we use further to evaluate properties of the superstructures in terms of continuum elastic theory. Based exclusively on the lateral variation of the vdW interactions between the adsorbates and the substrate, the monolayer turns out to have ~20% of the Young’s modulus of the molecular bulk and a Poisson ratio of ~0.1.

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