Detachment Dynamics of Graphene Nanoribbons on Gold

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Supporting Information

ABSTRACT: Metal−surface physisorbed graphene nanoribbons (GNRs) constitute mobile nanocontacts whose interest is simultaneously mechanical, electronic, and tribological. Previous work showed that GNRs adsorbed on Au(111) generally slide smoothly and superlubrically owing to the incommensurability of their structures. We address here the nanomechanics of detachment, as realized when one end is picked up and lifted by an AFM cantilever. AFM nanomanipulations and molecular-dynamics (MD) simulations identify two successive regimes, characterized by (i) a progressively increasing local bending, accompanied by the smooth sliding of the adhered part, followed by (ii) a stick−slip dynamics involving sudden bending relaxation associated with intermittent jumps of the remaining adhered GNR segment and tail end. AFM measurements of the vertical force exhibit oscillations which, compared with MD simulations, can be associated with the successive detachment of individual GNR unit cells of length 0.42 nm. Extra modulations within one single period are caused by steplike advancements of the still-physisorbed part of the GNR. The sliding of the incommensurate moiré pattern that accompanies the GNR lifting generally yields an additional long-period oscillation: while almost undetectable when the GNR is aligned in the standard “R30°” orientation on Au(111), we predict that such feature should become prominent in the alternative rotated “R0°” orientation on the same surface, or on a different surface, such as perhaps Ag(111).

KEYWORDS: graphene nanoribbons, nanocontacts, manipulation, nanomechanics, moiré pattern

Understanding the adhesive and frictional properties of nanosystems, such as molecules, 3D nanoclusters or 2D adsorbates, polymeric chains, etc., on structurally well-characterized crystalline substrates1−18 is of key importance for fundamental sciences such as contact mechanics and nanotribology and for technological applications. Indeed, controlled nanomanipulation of deposited nano-objects, in terms of positioning, adhesion, depinning, and sliding, can be used to build molecular superstructures, to explore the influence of the environment on individual molecules, or to perform engineering operations at the ultimate, molecular, limit of fabrication for hi-tech nanodevices. Mostly due to the extremely large surface-to-volume ratio, the peculiar behavior of these nanosystems may present...
properties that vary dramatically with size. Atomically precise synthesis techniques, experimentally tuning their physical and chemical characteristics during the preparation procedure, may assemble structurally similar geometries, with quite distinctive features influencing the system in terms of adhesive and friction-related response.

With their strong resilient structure and the experimental possibility to be picked up at one edge, dragged laterally and/or lifted up vertically by atomic force microscopy (AFM) techniques, graphene nanoribbons (GNRs) physisorbed on Au(111) surfaces do represent an important actor in this exploration.

AFM-driven sliding displacements have been already exploited to probe the frictional properties of this interface at very low temperature, providing initial evidence for a weak length dependence of the static friction of armchair nanoribbons. Paralleling those experimental results, subsequent numerical simulations have confirmed a frictionally superlubric regime characterized by small friction that is strongly oscillating and basically periodic with no average increase upon increase of the GNR length. With a static friction trend dictated by the characteristic periodicity of the graphene–gold interface moiré pattern, the simulated dissipative behavior already highlighted the occurrence of different dynamical regimes, ranging from smooth sliding to multiple stick–slip friction depending on the vertical height of the lifted, and laterally pulled, end.

In the present work, by means of nonequilibrium atomistic MD simulations and AFM manipulation data, we investigate the detailed mechanisms of detachment of a 30 nm long armchair GNR from the Au(111) surface upon vertical lifting of one end. Depending on the actual configuration at the nanoribbon/gold interface, the combined theoretical and experimental analysis sheds light, at a molecular level, on the manner in which adhesion and lateral corrugation of the nanocontact determine the characteristic periodicities observed in the lifting force and its z-derivative during the progressive detachment process upon lifting.
RESULTS AND DISCUSSION

Detachment Dynamics at R30. Experiment. Seven carbon atoms wide \((n = 7)\) armchair GNRs are synthesized on a clean Au(111) surface at 4.8 K in UHV, as detailed in the Methods. We control the Au-covered AFM tip so that it picks up a \(\sim 25\) nm long GNR at one end, producing unilateral detachment as sketched in Figure 1a. Figure 1b shows a STM topography of the surface before performing the manipulation of that GNR aligned along the \([-1, 0, 1]\) direction of the gold substrate, also called R30. This is, within the herringbone reconstruction of Au(111), the spontaneous orientation adopted by the GNR. In the specific case, the GNR crosses two fcc-hcp terraces of the gold substrate, as highlighted in Figure S1. The tip is positioned at one GNR end, with a low bias voltage \(\sim 2\) mV. When the junction between the tip and the GNR is established, an abrupt increase in the tunneling current is detected. Subsequently, a dynamical AFM mode is turned on, with a vertical oscillation amplitude \(\sim 43\) pm. The tip is then slowly retracted while recording the frequency shift (Figure 1d) as well as the energy dissipation (see Figure S1). Figure 1c shows the STM topography taken after the full detachment of that GNR, followed by tip cleaning outside the scan area. Figure 1d shows the measured frequency shift, i.e., the variation of the resonance frequency of the AFM force sensor upon vertical lifting of the GNR, as a function of the height \(z\). This frequency shift is proportional to the derivative of the vertical force that the tip exerts on the GNR: \(dF_{\text{ext}}/dz\). This force gradient measurement with a stiff cantilever realizes a high sensitivity measurement while avoiding a typical mechanical instability, which can be caused in the quasi-static force measurement with a soft cantilever. After an initial steep increase of the frequency shift, due to the GNR curvature build-up, an oscillating signal is observed, with a main periodicity \(\sim 0.40\) nm.

Simulation and Theory. In order to shed light on the physical mechanisms that come into play in the detachment dynamics, we simulate this same system with nonequilibrium molecular dynamics (NEMD). We construct an \(n = 7\) armchair GNR, consisting of a stripe of alternating triplets and pairs of carbon hexagons, of width \(\approx 0.7\) nm and length \(\approx 30.2\) nm, where all the peripheral C atoms are passivated with hydrogens. All carbon atom coordinates are fully mobile. The Au(111) substrate consists of two unreconstructed layers, also fully mobile, on top of one rigid layer, with fcc stacking. The lifting effect of the AFM tip on the first row of three C atoms is simulated by means of a vertical spring with an effective elastic constant \(k_z = 1800\) N/m. One end of this spring moves vertically with constant velocity \(v_0 = 0.5\) m/s along the \(z\)-axis of Figure 1a. Although much larger than the experimental speed, \(\approx 1\) \(\mu\)m/s, we verified that the simulated lifting speed is still small enough to yield meaningful, speed-independent results. The spring is attached to the three C atoms in the first row. The in-plane coordinates \((x, y)\) of these three atoms are kept fixed during detachment, so that the lifting is vertical. The simulation proceeds until complete detachment of the GNR is achieved. To gain insight into the physics of this system, we are interested in comparing the time-evolution (or equivalently the evolution against the \(z\)-coordinate of the driving spring end) of the following quantities:

- the instantaneous vertical force acting along the pulling direction, namely \(F_{\text{ext}}(t) = 3k_z[v_0 - z_{\text{end}}(t)]\), where \(z_{\text{end}}(t) = \sum_{i=1}^{3} z_i(t)/3\) is the average \(z\)-coordinate of the lifted end of the GNR;
- the vertical force gradient \(dF_{\text{ext}}/dz\), directly comparable to measurements, which is proportional to the frequency shift \(\delta f(z)\), \(dF_{\text{ext}}/dz = \kappa \delta f(z)\), where \(\kappa = 0.15\) N m\(^{-1}\) Hz\(^{-1}\) is a conversion factor;
- the deviations of the GNR-substrate total adhesion energy \(\Delta E_{\text{adh}}(t)\) and the gold–gold total potential energy \(\Delta E_{\text{Au-Au}}(t)\) away from their linear behavior \(E(t) = E^0 - Pt\) where \(P\) and \(E^0\) are fitting parameters;
- the total variation of the intra-GNR carbon–carbon and carbon–hydrogen potential energy \(\Delta E_{\text{GNR}}(t) = E_{\text{GNR}}(t) - E_{\text{GNR}}(0)\). The value at \(t = 0\) corresponds to an unlifted relaxed GNR on gold;
- the horizontal advancement of the trailing end of the GNR along the \(x\)-axis \(\delta x_{\text{end}}(t) = x_{\text{end}}(t) - x_{\text{end}}(0)\). The quantity \(x_{\text{end}}(t) = \sum_{i=1}^{3} x_i(t)/3\) is obtained by averaging the instantaneous \(x\)-coordinate of the end row of C atoms, the last ones to get lifted up at complete peel-off.

These physical observables are conveniently reported as functions of the dimensionless vertical displacement \(z/a_{\text{GNR}}\), where \(a_{\text{GNR}} = 0.42\) nm equals one GNR unit cell length, and the reference coordinate \(z = 0\) is taken at the average vertical position of the C atoms of the fully relaxed GNR on the substrate in the unlifted configuration.

The detachment dynamics of the GNR occurs following two successive regimes. In the first regime at low lifting for \(z < 3\) nm, the vertical motion of the AFM initially builds up the GNR curvature, marked by an increase of the GNR bending energy, as shown in Figure S2. The bending energy rise is consistent with the initial upswing of the experimental frequency shift of Figure 1d. The GNR-Au(111) corrugation energy is small compared to the system bending elasticity, so that the first relevant, yet gradual, build-up of the GNR curvature results in a smooth sliding of the still attached GNR tail. For \(z \approx 3\) nm \(\approx 7a_{\text{GNR}}\) the frequency-shift profile reaches a plateau with a superimposed oscillation. This second regime corresponds to a steady peeling. Figure 2 shows in this second regime both the experimental and the theoretical force derivative curves. The simulated force derivative is smaller in magnitude and not identical in line shape to the experimental one. In that respect, one must recall that the force fluctuations represent only a small deviation relative to the total lifting force, a very large background quantity which is unaccessible experimentally but dominating in simulation.

The level of quantitative agreement between theory and experiment is limited by the use of simple two-body Lennard-Jones potentials to describe simultaneously adhesive interactions and the interface corrugation. The specific values of the LJ parameters adopted in the present work represent the best possible compromise to reproduce at the same time the correct order of magnitude of the adhesive forces for carbon/metal interactions and the crucial features of the lifting dynamics associated with lateral corrugation.

First, the main periodicity is equal to the GNR unit length \(a_{\text{GNR}}\), consistent with the observation that the vertical motion of the spring induces a sequence of discrete detachments of GNR sections with the size of one GNR unit cell, until complete pull-off. Second, within one single period of this vertical lifting the detachment force experiences significant and roughly periodic drops, that occur in coincidence with steplike slips of the still-physiosorbed section of the GNR, as illustrated.
by small steps advancement of the tail in Figure 2c. We verified that the force drops are enhanced by the gold mobility by repeating the entire detachment simulation with a substrate consisting of gold atoms frozen in their crystalline arrangement. A direct comparison between the results of the simulations with the mobile and the rigid substrate is provided in Figure S3 of the Supporting Information. The importance of the gold mobility, an element not discussed before, is also confirmed by the behavior of the fluctuations of the gold−gold potential energy of Figure 3b: they are small but not entirely negligible compared to the variations of the other energy terms of the system.

Focusing on just four \( a_{\text{GNR}} \) lifting periods for heights \( 15a_{\text{GNR}} < z < 25a_{\text{GNR}} \), Figure 3 provides further insight into the detachment. This zoomed-in detail shows that the main periodicity of \( F_{\text{ext}}(t) \) corresponds to the detachment of a full GNR unit cell. As in the experiments, that main period is additionally decorated by superimposed oscillating features. These finer force derivative features do not arise, as one might have initially suspected, from the progressive detachment of the \( n = 7 \) armchair GNR substructures (e.g., the alternating rows of two- and three-carbon hexagons). Instead, these secondary force derivative features actually arise from the slips, during the detachment dynamics, of the nanoribbon tail, against the lateral corrugation of the gold substrate. The intrinsic incommensurability between \( a_{\text{GNR}} = 0.42 \) nm and \( a_{\text{Au}} = 0.288 \) nm would in principle permit a frictionless, superlubric sliding. However, the trailing end interrupts the GNR−Au(111) moiré pattern, giving rise to an uncompensated region which breaks superlubricity causing a frictional barrier against sliding, capable of causing an unexpected stick−slip. That, together with the role played by the GNR bending elasticity, makes the recorded profile of the lifting force quite complex and rich. More information about this interplay of different length scales can be inferred from the behavior of the individual potential energy terms controlling the system dynamics. We note here that since the external spring drives the GNR away from the substrate at an average constant velocity, we expect the GNR−substrate potential energy to increase linearly in time on average. Likewise, the detachment of the GNR causes the substrate layers to relax around the detachment region, causing the Au−Au potential energy to
detachment/reattachment: the GNR is obtained even in the limit mechanical instabilities of the system that would, hence, be bending relaxations and tail slips along the upward and attachment of individual units of the GNR and consequent larger, and second, there are force peaks (in detachment) and 
\[ \Delta F \text{ along the upward cycle}, \] 
responsible for the total work of the external small portion of the histeretic cycle of Figure 4 but at a far smaller speed \[ \mu \text{m/s} \] 
than in simulations. As a result, the experiment is far closer to the adiabatic regime and finds significant dissipation practically only in connection to the detachments/reattachments of the GNR unit sections. The resulting experimental energy dissipation curve is reported in the Supporting Information as Figure S1. 
As a final observation, Figure 4c shows that the GNR internal energy variation \( \Delta E_{\text{GNR}} \) is systematically larger during reattachment than during detachment. This is due to an extra forced bending produced by the spring pushing the GNR down toward the substrate during the reattachment rather than pulling it up as during detachment. 

**Moiré Pattern Signatures: Detachment at R0.** In the previous section, we clarified the main features of the GNR dynamics, determined by the successive detachment of the GNR individual unit sections. By contrast, as pointed out in our previous works,\textsuperscript{12,13} the dependence of the static friction on the size of the GNPs and the sliding dynamics upon lateral pulling was strongly affected by other GNR characteristics, such as the moiré-pattern primary periodicity, which gives rise to regions of local registry match/mismatch between the GNR and the substrate. 
Surprisingly, no sign of this kind of feature has been observed so far either in the lithof experiment or in the simulations. We are therefore interested in understanding by
MD simulations whether this feature could naturally emerge by changing the relative strength between the adhesion of the GNR, which governs the detachment of the GNR unit cells, and the corrugation, which contrasts lateral motions and tunes the moiré pattern force contribution.12

In our model, one can, for example, reduce the characteristic distance $\sigma$ used in the Lennard-Jones (LJ) potential to describe the GNR/substrate interaction (see the Methods). Figure 5 shows the vertical force profile obtained by artificially modifying the LJ distance $\sigma$ from the regular value $\sigma = 0.342$ nm to a smaller one 0.274 nm (blue solid curve). The fitting function of eq 1 (yellow dashed curve) emphasizes the long-wavelength periodicity, now visible due to the detachment of moiré pattern units, at this artificially larger interaction strength.

$$F(z) = F_0 + F_1 \cos \left( \frac{2\pi}{\lambda_m} z + \varphi \right)$$

with $F_0$ being the average vertical force, $F_1$ the amplitude of the force oscillation, $\lambda_m$ its wavelength, and $\varphi$ its phase. The resulting $\lambda_m = 4.86$ nm corresponds to the moiré pattern wavelength in the R30 alignment.12 This is consistent with the subsequent detachment of regions with better adsorption energy, with good local registry between the GNR and the substrate, and regions of poorer adsorption energy with mismatched registry. Since by artificially enhancing the substrate corrugation the moiré pattern periodicity becomes evident in the force profile, a similar enhancement could, in principle, occur by choosing a different GNR orientation on the substrate. On Au(111), all GNR orientations different from R30 are disfavored by the herringbone reconstruction of Au(111), as mentioned above. Au(111) could, in principle, be replaced by a different substrate, such as perhaps Ag(111) or Cu(111) or others, that are not reconstructed, theoretically permitting different orientations and/or different adhesion energies and different moiré pattern periodicities.

For a qualitative impression of the kind of changes expected, we simulate the detachment of a GNR deposited along the [1 $\bar{1}$ 0], or R0, alignment, of our hypothetically unreconstructed Au(111). Both adhesion and corrugation are larger in that GNR alignment, compared to the real R30 alignment addressed above and earlier.12

Figure 6 shows the detachment traces from simulations of GNR/Au(111) for the R0 orientation. In this alignment, the substrate periodicity $a_{Au} = 0.499$ nm generates a moiré pattern with primary periodicity $\lambda_{m}^{R0} = 1.32$ nm due to its zigzag shape.12 With the main periodicity now determined by the moiré pattern, the signature of the detachment of the single unit cells of the GNR almost disappears. For this reason we express the quantities of interest as a function of $z/\lambda_{m}^{R0}$. In the R0 alignment, the dynamics consists of successive detachment of entire moiré pattern units, as indicated by peaks in the adhesive fluctuations, followed by multiple slip-like advancements of the physisorbed tail, that are accompanied by wide oscillations of the vertical force. Finding no way yet to realize...
the GNR R0 alignment in experiment, otherwise observed in other graphene/metal interfaces,24 the status of the results summarized in Figure 6 is that of an interesting theoretical prediction, deserving future experimental investigation.

CONCLUSIONS

We have described extensive nonequilibrium molecular dynamics simulations of detachment dynamics by vertical lifting of a graphene nanoribbon adsorbed on Au(111), along with experimental data of the tip-applied force derivative. The results show that the steady state of this vertical dynamics is characterized by the detachments of the individual physisorbed units of the GNR, with the GNR unit cell 0.42 nm periodicity, in both the experimental tip frequency shift and the simulated vertical-force profiles. These detachments are accompanied by an intermittent stick–slip-like, stop-and-go motion of the tail, which is responsible for extra vertical-force drops that are visible within the main periods.

A qualitatively similar phenomenology is observed during reattachment, where a sequence of unit cell attachments and tail slips still takes place. The frictional loop corresponding to one detachment/reattachment cycle also shows that a nonadiabatic partial detachment/reattachment of the GNR is an intrinsic source of energy dissipation. A non-negligible contribution to the detachment force evolution and to the dissipation comes from the deformability of the Au substrate. Theory overall parallels closely the experimental data, despite quantitative discrepancies due to the impossibility to reproduce in simulations the very low speed as well as the finest adsorption details of the experimental system. Importantly, we have shown that the detachment dynamics can be affected by different competing periodicities depending on the relative strength between the total adhesion and the corrugation of the interface. Enlarging our view, we have explored the simulated detachment for the R0 alignment. Here, the larger corrugation leads to an almost complete disappearance of the GNR unit-cell detachments in favor of moiré-pattern features, the detachment now involving moiré-pattern periods as entire units. Like the R30 case, the detachment of one complete unit is then followed by sudden multiple slips of the tail.

Further investigations of these observations could involve a theoretical and experimental investigation of this type of nanomanipulation either by changing the GNRs orientation on a possibly unreconstructed gold substrate or possibly by considering different substrates without reconstructions and/or larger corrugations, such as possibly silver, copper, or others.

METHODS

Experimental Setup. All measurements were performed with a commercially available Omicron low-temperature scanning tunneling microscopy (STM)/atomic force microscopy (AFM) system, operating in ultrahigh vacuum at 4.8 K. We used a tuning fork with a chemically etched tungsten tip as a force sensor.25 The resonance frequency and the mechanical quality factor are 23026 Hz and 19974, respectively. The high cantilever stiffness of 1800 N/m realizes a stable operation with a small amplitude of 43 pm.25 The frequency shift, caused by the tip–sample interaction, was measured with a commercially available digital phase-locked loop (Nanonis: OC-4 and Zurich Instruments: HF2-LI and HF2-PLL).24 For the STM measurement, the bias voltage was applied to the tip while the sample was electronically grounded. The tungsten tip of a tuning fork sensor was sharpened ex situ by focused ion beam milling technique and was then covered in situ with Au atoms by contacting to the sample surface. A clean gold tip was formed in situ by indenting the Au sample surface and applying a pulse bias voltage between tip and sample several times. Clean Au(111) surfaces were prepared in situ by repeated cycles of standard Ar+ sputtering (3 × 10−6 mbar, 1000 eV, and 15 min) and annealing at 750 K. As precursors to the GNRs synthesis, 10,10′-di-bromo-9,9’-dianthryl molecules were deposited on the substrate from a Knudsen cell crucible, heated resistively at 135 °C. Subsequently, the samples were annealed at 200 and 400 °C to synthesize graphene nanoribbons on Au(111).26,27 The STM topographic images were taken in constant current mode. Measured images were partially analyzed using the WSxM software.28

Theoretical Modeling. All MD simulations of detachment are performed using the LAMMPS package.29 The GNR atoms and the two gold mobile layers obey a T = 0 dissipative Langevin dynamics, with a damping parameter γ = 1.0 ps−1, which has been adjusted in order to highlight the stick–slip behavior of the GNR in the steady state. The force fields used to simulate the dynamics of the mobile gold layers and the GNR are the Embedded Atom Method (EAM) potential30,31 and the Adaptive Intermolecular Reactive Empirical Bond Order (AIREBO) potential,32 respectively.

The C–Au and the H–Au adhesive interactions are modeled via the Lennard-Jones (LJ) potentials of the form

$$V(r) = 4\varepsilon \left[ \sigma^12 \left( \frac{\sigma}{r} \right)^12 - \left( \frac{\sigma}{r} \right)^6 \right]$$

(2)

We adopted ε = 8 meV for the C–Au interaction and ε = 3.2 meV for the H–Au interaction. The adhesive energy per C atom that we obtain with these values of the LJ amplitudes for flat GNRs on gold is compatible with previous DFT estimates for graphene/metal interfaces33,34 and experimental results on graphite/graphite and graphene/silicon contacts.35,36 However, we found that the qualitative features of the dynamics are not significantly affected by variations of the LJ parameters in a suitable range of values. These more realistic energies are larger than those adopted in previous work.9,12,13 The adsorption distances for carbon and hydrogens are set by a common σ = 0.342 nm. This parametrization yields a good match between the shape of the experimental frequency shift profile and the simulated vertical force gradient for the R30 alignment (see Figure 2b). The same parameters were also used for the simulation of detachment in the R0 alignment.

As discussed above, for the calculations reported in Figure 5, the lateral corrugation was enhanced artificially by changing the value of σ to 0.274 nm, while keeping all other parameters unchanged.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.8b07894.

Comparison between the experimental frequency shift trace and the measured energy dissipation per cycle (Figure S1), behavior of the GNR internal energy variation (Figure S2), comparison between the detachment dynamics, in terms of the recorded vertical force profile and the GNR tail advancement (Figure S3), and comparison between the GNR dynamics with a “hard” and a “soft” virtual spring attached to the ribbon short edge (Figure S4) (PDF).

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REFERENCES


