Nanotribological Properties of the h-BN/Au(111) Interface: A DFT Study

Merve Baksi,† Daniele Toffoli,‡ Oguz Gulseren,§ and Hande Ustunel*†§

†Department of Physics, Middle East Technical University, Dumlupinar Blv 1, 06800 Ankara, Turkey
‡Dipartimento di Scienze Chimiche e Farmaceutiche, Università degli Studi di Trieste, Via L. Giorgieri 1, I-34127 Trieste, Italy
§Department of Physics, Bilkent University, 06800 Ankara, Turkey

Supporting Information

ABSTRACT: Understanding the quantum-mechanical origins of friction forces has become increasingly important in the past decades with the advent of nanotechnology. At the nanometer scale, the universal Amontons–Coulomb laws cease to be valid, and each interface requires individual scrutiny. Because of the well-known lubricating properties of two-dimensional materials, a significant amount of research has been performed in an effort to understand interfaces they form with one another. However, the interfaces between these two-dimensional materials and metals red from a tribological point of view, important for such applications as friction force microscopy, have yet to be thoroughly investigated. In the current work, we present a detailed density functional theory investigation of the hexagonal BN/Au(111) interface. Because of a good agreement between their characteristic lengths, a high level of commensurability is achieved in a suitably constructed model between the bulk surfaces of the two materials. As a result of our calculations, we find that the corrugation in the potential energy surface and the lateral forces in this interface are low compared to other similar interfaces. The friction coefficient falls rapidly with increasing load down to 0.005 for the largest loads considered. In contrast, Au clusters (n = 1, 4, 13, and 19) sliding on the h-BN surface exhibit much larger lateral forces, indicating strong size and edge effects. The reduction of energy corrugation in going from the Au4 to the Au19 cluster may already indicate a decreasing trend with increasing size even at this very small scale.

1. INTRODUCTION

Two-dimensional materials have generated a great deal of interest due to their potential use as lubricants in nanoscale systems. In particular, graphene,1 h-BN,2 and MoS2,3 have been predicted to exhibit exceptional performance in reducing friction and wear. In addition to providing a lubricating buffer layer between two surfaces in relative motion, two-dimensional materials have been observed to display very low friction forces while sliding on metal substrates.4 Ultralow friction, often observed in a similar study.8 Materials have been observed to display very low friction forces while sliding on metal substrates.8 In addition to providing a lubricating buffer layer between two surfaces in relative motion, two-dimensional materials have been observed to display very low friction forces while sliding on metal substrates.8 However, even for components in perfect interfacial registry, such as bilayers of some two-dimensional materials, very low friction forces have been observed. A striking example of this is the fluorographene/fluorographene bilayer interface5,6 where the energy corrugation, defined as the difference between the maximum and the minimum of the potential energy surface (PES), was calculated to be 0.31 meV/atom. In contrast, the graphene/graphene interface was found to display a corrugation of 6.47 meV/atom in a similar study.7

The experimental characterization of the tribological behavior of interfaces is performed with the use of atomic force microscopy (AFM)9–11 or friction force microscopy (FFM)12,13 techniques. The coating materials on the surfaces of the AFM/FFM tips can be chosen from a large selection including SiO2,10 Pt,14 and Cu.15 Whereas the tribological properties of interfaces between 2D materials have been extensively studied, the friction forces between these materials and metal substrates have been investigated in only a limited number of studies.16,17 Furthermore, as in the case of the graphene/Cu(111) interface,18 most asymmetric interfaces present a lattice mismatch, which calls for either large simulation cells or the application of strain. The h-BN/Au(111) interface under investigation in the present work requires neither due to an extraordinarily high match between the two lattices, given a suitable choice of the simulation cell. We are therefore able to decouple the effect of the interaction from strain effects.

The theoretical basis of friction at the atomic level between two periodic surfaces was established by an array of seminal works. Building on the treatments in 1928 by Prandtl19 and in
1929 by Tomlinson,20 the earliest theory was developed by Sokoloff21 for two periodic surfaces in contact, whose atoms interact with simple springs. These models identified two regimes of relative motion, stick–slip and continuous sliding, which emerge through an interplay between the corrugation and the rigidity of the materials in contact. In a more modern treatment, Zhang and Tomanek illustrated the calculation of the lateral friction forces and eventually the friction coefficient for a model system of a single Pd atom sliding on a Pd surface using density functional theory (DFT).22 More recently, Cahangirov et al. presented a straightforward way of translating the DFT PES data to stick–slip criteria,23 establishing the link between phenomenological parameters such as the stick–slip threshold and atomistic DFT results. The conclusion from this body of work is that a lower corrugation correlates with lower friction forces and motion in the continuous sliding regime. Experimental setups designed to measure friction forces often involve contact and sliding between a coated AFM tip and the material of interest. It is, however, rather difficult to induce relative lateral motion between two sheets of material. Some techniques for overcoming this difficulty are summarized in the review by Baykara et al.24 In this respect, DFT calculations are of utmost utility since they can easily map the PES between surfaces of virtually any kind. For instance, the DFT work by Wang et al.8 identified the graphene/MoS2 contact as an extremely low-friction surface due to the lack of charge density fluctuations during sliding.

In this work, we aim to investigate the static limit of the friction forces between the Au(111) surface and h-BN. We approach this problem from two different directions. The first part of our results deals with the atomically perfect interface between the bulk Au(111) surface and an infinitely repeated periodic h-BN sheet. In the second part, we focus on the friction forces between Au clusters (n = 1, 4, 13, and 19) and a perfect h-BN sheet. The motivation here is to understand the effect of imperfections such as tip edges or protrusions which usually act to increase the friction forces. We finally present a summary of our most important conclusions.

2. COMPUTATIONAL DETAILS

All of the calculations presented in this work were performed using the open-source code suite Quantum ESPRESSO.25 The simulation cell used in the bulk interface calculations between the Au(111) surface and the h-BN sheet, shown with a red parallelogram in Figure 1, was composed of 12 Au atoms (4 atomic layers) and 4 BN pairs. Because of the exceedingly small energy differences between different configurations, all bulk calculations were performed within tight thresholds. Integrals over the Brillouin zone were computed using a 12 \times 20 \times 1 Monkhorst-Pack k-point mesh along with a cold-smearing scheme of a width of 0.01 Ry. The kinetic energy and augmentation charge thresholds for the plane-wave expansion were 60 and 600 Ry, respectively. Going from a cutoff energy of 50 Ry and a k-point mesh of 17 \times 10 to a cutoff energy of 60 Ry and k-point mesh of 20 \times 12 changes the corrugation in the total energy by only about 0.04 meV for the entire simulation cell. We therefore conclude that our results are well converged for the calculation parameters used. Geometry optimizations were performed using the Broyden–Fletcher–Goldfarb–Shanno (BFGS) algorithm28–31 until all atomic forces were lower than about 4.0 \times 10^{-5} nN. During geometry optimizations involving the Au slab, the bottom two layers of the slab were fixed to mimic bulk behavior. The convergence criterion for electronic energies within a single self-consistent field step was kept at a conservative value of 0.5 \times 10^{-12} eV/atom. The vacuum between periodic images along the direction normal to the surface was kept at \sim 13 Å. Larger vacuum values were tested and were found to yield very similar results. The weak interaction between the h-BN sheet and the underlying Au surface was modeled by using the vdW-DF232 functional with the C09 exchang partner.33 The ion–electron interaction was modeled by using ultrasoft pseudopotentials.

The figures of merit we consider here for quantifying interfacial friction are the magnitude of the corrugation in the PES of interaction between Au and h-BN and that of the lateral forces in the sliding direction as a function of vertical load. The corrugation is defined as

$$\Delta E = E_{\text{max}} - E_{\text{min}}$$

where \(E_{\text{max}}\) and \(E_{\text{min}}\) are the maximum and minimum values of the PES within an entire spatial domain or along a certain direction. We constructed the PES by means of calculating the adhesion energy of the h-BN layer to the Au surface on an 11 \times 11 grid of relative positions as shown in Figure 1. At each point of the grid, a partial geometry optimization was performed where the atoms were allowed to move along the normal axis (z-axis as defined in Figure 1) while kept fixed.
along the plane of the sheet (the xy-plane). This procedure yields the PES under zero load.

In preparation for the finite load analysis, we performed single-point self-consistent field (SCF) calculations at a number of evenly spaced points, \( \{x_i\} \), along the directions labeled Path 1 and Path 2 in Figure 1. For each set of such calculations, the vertical distance between the two surfaces were kept fixed. We scanned vertical distances ranging from 2.8 to 3.6 Å in steps of 0.1 Å. At each vertical distance, we calculated the component of the total lateral force \( F_N(x_i) \) along the sliding path and the total vertical load \( F_N \) for all \( \{x_i\} \). Once the data were collected, we obtained the lateral force corresponding to the desired load, \( F_{N0} \), via a spline interpolation at each point \( x_i \). The lateral forces \( F_N(x_i) \) were calculated via two methods: using the total Hellmann–Feynman forces on the h-BN atoms, obtained directly at the end of the SCF run (the direct method) and by means of differentiating the total energy surface with respect to the sliding coordinate for the desired load (the derivative method). We then calculated the average of the lateral force over the total sliding direction using

\[
F_L = \frac{1}{N} \sum_{i=1}^{N} F_L(x_i)
\]

(2)

where the prime refers to a condition imposed on the sum where only negative values of \( F_L \) are considered. In other words, only those positions that result in a force opposing the motion are included in the calculation of the average lateral force. \( N \) is then the total number of points included in this sum. As a simple check, the lateral forces were estimated by a third method, which we refer to as the Min/Max method. This method involves a simple division of the energy difference between the maximum and the minimum point of the energy landscape by the distance between them. Results from all three methods were seen to agree well as presented later in the text.

Once the average lateral force, \( F_L \), over a certain distance along the sliding path was calculated for a given load \( F_{N0} \), the static friction coefficient was calculated within the standard definition via

\[
\mu = \frac{F_L}{F_N}
\]

(3)

Because the bulk calculations are performed for an infinite interface, we present our results in terms of the intensive normal pressure \( \sigma_N \)

\[
\sigma_N = \frac{F_N}{A}
\]

(4)

where \( A \) is the surface area of the supercell described above.

As a means of quantifying size and edge effects, we also investigated the lateral forces on Au_{11}, Au_{13}, and Au_{19} clusters as they slide along the armchair direction on a single h-BN sheet. At each point, the adsorption energies and the lateral forces were computed under zero load, which necessitates a geometry optimization with only the \( z \)-coordinates of the atomic positions free to move. We used \( 3 \times 3, 4 \times 4, \) and \( 6 \times 6 \) h-BN simulation cells for the Au_{11}, Au_{13}, and Au_{19} clusters with Monkhorse-Pack meshes of \( 6 \times 6 \times 1, 4 \times 4 \times 1, 6 \times 6 \times 1, \) and \( 6 \times 6 \times 1 \) kpoints, respectively. We used a larger kpoint set for the two largest cluster sizes for a more accurate calculation of potentially high magnetic moments. The plane-wave kinetic energy and the augmentation charge thresholds used for all cluster calculations were 40 and 400 Ry, respectively. All cluster calculations were conducted within spin-polarized DFT.

The morphology of Au clusters has been the subject of many theoretical studies.\(^{35–37} \) For the current treatment, the rhomboid, icosahedral truncated, and pyramid geometries were selected for the Au_{11}, Au_{13}, and Au_{19}, respectively. All clusters were initially geometry optimized in a large cubic unit cell at the \( \Gamma \) point. Following this initial optimization, the clusters were placed on the h-BN network at a suitable high-symmetry location, and the combined system was geometry optimized one more time. For details on the cluster geometries and their magnetic properties before and after their adsorption on the h-BN layer, we refer the reader to the Supporting Information.

In spite of the fact that variations in the energy surface are more relevant to the problem of tribology instead of the absolute values of the adhesion energies, we calculate the adhesion energies using the formula

\[
E_{\text{adh}} = E_{\text{sub+ads}} - E_{\text{sub}} - E_{\text{ads}}
\]

(5)

where \( E_{\text{sub+ads}} \) is the total energy of the substrate and the adsorbate, \( E_{\text{sub}} \) is the total geometry optimized energy of the substrate by itself, and \( E_{\text{ads}} \) is the total energy of the geometry optimized adsorbate. In the case of the bulk interface, the h-BN layer is considered to be the adsorbate while in the case of the Au clusters, the term adsorbate refers to the cluster in question. The isolated adsorbate energies as well as those of the substrate were calculated by using the same vDw-DF2-C09 protocol of the main calculations for the sake of consistency.

3. RESULTS AND DISCUSSION

3.1. h-BN/Au(111) Interface.

In most periodic DFT-based bulk interface studies, a strain is applied to one or both of the materials in question to fit both materials into a computationally accessible shared simulation cell. This strain may be tensile or compressive as needed. In the case of the h-BN/Au(111) interface, the choice of a \( 2 \times 2 \) h-BN cell on top of a \( 3 \times 1 \) Au(111) surface cell, shown in Figure 1, turns out to be a near perfect match, using our calculated bond lengths of \( d_{\text{Au-Au}} = 2.90 \) Å and \( d_{\text{N-N}} = 1.45 \) Å. As a result, no strain was applied to either material for our bulk calculations. If used instead of the computational ones, the experimentally determined bond lengths (2.87 Å for Au\(^{38} \) and 1.44 Å for h-BN\(^{15} \)) necessitate a 0.2% tensile strain on the h-BN component. This strain is still minimal and is not expected to affect the results significantly. The interface constructed as described has a high degree of coincidence because of its small repeating unit in addition to being strain-free. This model therefore presents an opportunity for exploring the upper limits of friction between the two components.

The PES calculated at zero load for this interface (Figure 1) presents an exceptionally mild corrugation of \( \sim 0.3 \) meV/BN pair. As a comparison, Table 1 lists a set of corrugations calculated for a set of interfaces using DFT, taken from various sources in the literature. The h-BN/Au(111) interface studied here falls in the low corrugation regime among the values presented in this table, much smaller than the perfectly matching, symmetric interfaces displayed. Its corrugation is, on the other hand, much closer to the asymmetric interfaces FG/MoS\(_2\)^{7} and graphene/MoS\(_2\)^{8} The corrugation of the graphene/MoS\(_2\) interface is the lowest of all the values presented here; however, the degree of incommensurability is larger than our case. Furthermore, the “per atom” \( \Delta E \) values
maximum adhesion energies and corrugations for perfect match the underlying Cu substrate.

The friction coefficients exhibit a rapidly decreasing trend with increasing vertical load. For the smallest load considered here, namely $\sigma_N = 0.01$ GPa, the friction coefficient calculated by using the derivative of the PES is 0.12 for sliding along the $x$-direction. As the load increases, $\mu$ decreases all the way down to 0.005 at $\sigma_N = 1.4$ GPa. This behavior is commonly observed in atomically perfect interfaces. A careful analysis of the individual lateral components of the Hellmann–Feynman forces yields insight into this behavior. For this analysis, we concentrate on the range of separations between $d = 2.8$ Å and $d = 3.1$ Å. As depicted in Figure S2, the projection of the atomic forces onto the plane of the h-BN sheet varies greatly both in direction and in magnitude depending on relative location of the two materials. We therefore collect all individual atomic forces at all locations along each path and consider the trends. In this range of separations, overall, the lateral forces grow for motion along the $x$ direction (Path 1 in Figure 1) as a function of increasing load as shown in Figure 3. However, the increase in the average lateral forces in the direction opposite to motion turns out to be slower than the increase in the vertical load, thereby causing the emergence of a decreasing trend in $F_L/F_N$ (lower panel of Figure 3). In addition to the slower increase of $F_L$, it is also shifted; in other words, $F_L$ is not zero at zero load. Along the $y$ direction (Path 2), the forces appear to be largely independent of separation, once again resulting in a decrease in $F_L/F_N$. Separations larger than $d = 3.1$ Å resulted in atomic forces mostly along the direction of motion rather than in the opposite direction and are therefore excluded from this analysis. Exceptions to this behavior can be seen in other two-dimensional interfaces. For instance, the Ti$_{n+1}$C$_n$ MXenes investigated computationally by

Table 1. Potential Energy Surface Corrugation (meV/Atom) Values Calculated for Interfaces in Several Computational Studies Are Compared with the Value Obtained in the Current Work

<table>
<thead>
<tr>
<th>Interface</th>
<th>$\Delta E$ (meV/BN unit)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>graphene/Cu(111)</td>
<td>$\approx 12$</td>
<td>ref 18</td>
</tr>
<tr>
<td>graphene/graphene</td>
<td>$\approx 10$</td>
<td>ref 39</td>
</tr>
<tr>
<td>h-BN/h-BN</td>
<td>$\approx 10$</td>
<td>ref 40</td>
</tr>
<tr>
<td>MoS$_2$/MoS$_2$</td>
<td>$\approx 3.3$</td>
<td>ref 41</td>
</tr>
<tr>
<td>HG/HG</td>
<td>0.96</td>
<td>ref 7</td>
</tr>
<tr>
<td>FG/FG</td>
<td>0.31</td>
<td>ref 7</td>
</tr>
<tr>
<td>FG/MoS$_2$</td>
<td>0.14</td>
<td>ref 42</td>
</tr>
<tr>
<td>graphene/MoS$_2$</td>
<td>0.046</td>
<td>ref 8</td>
</tr>
<tr>
<td>h-BN/Au(111)</td>
<td>0.15</td>
<td>current work</td>
</tr>
</tbody>
</table>

*HG and FG stand for hydrogenated graphene and fluorographene, respectively.

Table 2. Maximum Value of the Adhesion Energy ($E_{\text{adh}}$ in meV/BN unit) and the Corrugation ($\Delta E_x$ and $\Delta E_y$ in meV/BN unit) of the h-BN/Au Interface as a Function of Interlayer Separation $z$ (Å)

<table>
<thead>
<tr>
<th>$z$ (Å)</th>
<th>$E_{\text{adh}}$ (meV/BN)</th>
<th>$\Delta E_x$ (meV/BN)</th>
<th>$\Delta E_y$ (meV/BN)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.8</td>
<td>120</td>
<td>1.83</td>
<td>1.45</td>
</tr>
<tr>
<td>2.9</td>
<td>145</td>
<td>1.47</td>
<td>1.30</td>
</tr>
<tr>
<td>3.0</td>
<td>160</td>
<td>1.22</td>
<td>1.09</td>
</tr>
<tr>
<td>3.1</td>
<td>168</td>
<td>0.93</td>
<td>0.84</td>
</tr>
<tr>
<td>3.2</td>
<td>170</td>
<td>0.73</td>
<td>0.70</td>
</tr>
<tr>
<td>3.3</td>
<td>168</td>
<td>0.58</td>
<td>0.65</td>
</tr>
<tr>
<td>3.4</td>
<td>164</td>
<td>0.48</td>
<td>0.45</td>
</tr>
<tr>
<td>3.5</td>
<td>157</td>
<td>0.40</td>
<td>0.39</td>
</tr>
<tr>
<td>3.6</td>
<td>150</td>
<td>0.33</td>
<td>0.25</td>
</tr>
</tbody>
</table>
Zhang show the opposite trend, where $\mu$ increases almost linearly with $\sigma_N$.44

Several studies have linked the tribological behavior of interfaces to the interfacial charge density profile.45−47 The majority of these studies focus on symmetric interfaces where both materials donate an equal amount of charge to the interface, whose magnitude determines the adhesion energy and friction forces. The h-BN/Au(111) interface is, however,
between a covalently bonded material with polar bonds and a metal. The behavior of the charge transfer upon the formation of the interface is therefore expected to be somewhat different. In Figure 4, we present the charge density difference and its plane-average (along $z$) for $d = 2.8$ Å and $d = 3.1$ Å. The interface appears to be highly polarized with an electron excess region near the Au(111) surface and an electron depletion region near the h-BN layer. Interestingly, the net electron charge is supplied by both materials. As a result, although the interface is polarized, its overall charge remains negative. It is also worth noting that although the amount of positive and negative charges are individually smaller for the $d = 3.1$ Å separation, the total integrated negative charge in the interface (defined as the distance between the top Au layer and the h-BN layer) increases from $1.0 \times 10^{-5}$ to $1.6 \times 10^{-5}$ e/Å$^3$ going from $d = 2.8$ Å to $d = 3.1$ Å. This is consistent with the increase in the adhesion energy for the larger separation (see Table 2).

3.2. Au Clusters on h-BN. The bulk model of the interface provides a rather idealized view of its tribological properties. In actual experiments, the metal component is either in the form of a sharp AFM tip$^{10}$ or islands manipulated by an AFM tip. Through these experimental studies, the importance of the size and edge effects have been convincingly established. To address these issues and to understand the transition between a high-adhesion cluster geometry and a low-friction bulk interface, we analyze the frictional properties of the h-BN/Au$_n$ interface ($n = 1, 4, 13, 19$) interface by means of comparing the lateral forces (Figure 5). Before transferring the clusters to the h-BN surface, a geometry optimization was performed. Upon interaction with h-BN, the Au$_4$ and Au$_{19}$ clusters maintain their geometry while the Au$_{13}$ cluster was seen to undergo a substantial structural rearrangement. For details of the cluster geometries before and after adsorption, see section 2 of the Supporting Information along with Figure S3.

The optimized geometries and optimized distances of the clusters at the beginning of the sliding path are displayed in Figure S4. All four clusters are seen to rest at a vertical distance much smaller than the equilibrium h-BN/Au(111) separation in bulk form. The interaction is especially strong for Au$_4$, which assumes a tilted configuration even though the initial configuration before the geometry optimization is flat. This final tilted configuration was reproduced for different coverages and starting conditions and was therefore deemed to be robust. All cluster calculations were performed at zero load. The energy corrugation and the average friction force values are presented in Figure 5. Similarly to the bulk interface calculations, the forces are computed by using both the numerical derivatives of the energy and the Hellmann–Feynman forces. Once again, the agreement is very good (Figure S5).

Considered on a per atom basis, the single Au atom has the largest corrugation. However, in the case of the tilted Au$_4$ cluster, there is a single atom in immediate contact with the surface. Similarly, in the case of the Au$_{13}$ and Au$_{19}$ clusters, four and ten atoms are in direct contact with the surface, respectively. Viewed from this perspective, Au$_4$ appears to display the largest $\Delta E$/atom value while the corrugation is the smallest for Au$_{19}$. The decreasing lateral forces with increasing cluster size may possibly indicate a trend of decreasing friction force with increasing contact area.

Beyond these general observations, some details of the cluster results are also noteworthy. The maximum and the minimum of the Au$_4$ energy profile shown in Figure 5 correspond to locations that are on top of a hollow site and directly on top of a B atom, respectively. As mentioned above, similarly to Au$_4$, the Au$_{19}$ cluster also interacts with the surface mostly through a single atom namely the atom that is closest to the surface. As a result of this, the maximum and the minimum of the energy landscape appear at identical locations, namely on top of the B atom (configuration 1 in Figure 5) and the hollow site (configuration 3 in Figure 5). Because of the larger contact area of the Au$_{13}$ cluster, the minima and maxima are not as well-defined as the smaller clusters. As explained in the Supporting Information, upon adsorption on the h-BN sheet, the Au$_{13}$ cluster undergoes a sizable distortion. In its new distorted cuboctahedron geometry, the bottommost four atoms form a slightly distorted square. Configuration 1, displayed in Figure 5, corresponds to a geometry in which the geometric center of these four atoms are directly above a N atom. On the other hand, for configuration 3, this center is on top of a hollow site. In contrast, the Au$_{13}$ cluster largely maintains its truncated pyramidal shape upon adsorption. In configuration 1, all ten of the bottommost atoms are almost perfectly coincident with hexagon centers of the h-BN layer, whereas in configuration 3 the Au atoms are largely on top of B or N atoms.

As a means of providing a detailed account of atomic contributions to the lateral forces on the clusters, we break down the lateral forces as a function of vertical distance from the h-BN sheet. As shown in Figure S6, Au$_4$, Au$_{13}$, and Au$_{19}$ can all be broken down into three atomic layers each. The bottom layer is defined to be the collection of those atoms (one, four, and ten atom in the case of Au$_4$, Au$_{13}$, and Au$_{19}$).
clusters, respectively) that are closest to the h-BN sheet. The total lateral forces, summed over each layer, when plotted against displacement along the sliding direction, reveal that the interaction of $A_{13}u$ with h-BN is predominantly due to the bottommost layer (i.e., a single atom). This also applies to a large extent to $A_{13}u$ and $A_{19}u_5$, however, there is a sizable contribution, at some points of equal magnitude, of the mid and top layers to lateral forces. Along most of the sliding path for both clusters, the forces on the mid and top layers are of similar magnitude and opposite direction and therefore cancel each other.

Another manifestation of the strong interaction and of the edge effects is the induction of ripples in the h-BN substrate as the clusters move along the sliding path illustrated in Figure 5. As the cluster moves, portions of h-BN directly underneath are depressed while adjacent portions are elevated (Figure S7). This effect is seen clearly in the animated sequences created for all four clusters as they slide, which is presented as a part of the Supporting Information. The amplitudes of the ripples, defined as the difference between the highest and the lowest point of the h-BN sheet ($\Delta z_n$), are shown in Figure S7. According to these results, while the general trend is similar across all clusters, $A_{13}u$ and $A_{19}u_5$ clusters both induce distortions that are about 4 times larger than those of the two other clusters. It is also interesting to note that for all four clusters the extrema of the ripple magnitudes coincide with those of the lateral forces. This emphasizes once again the importance of the edge effects, especially for small clusters.

4. CONCLUSIONS

The interface between the h-BN and Au(111), explored here, to our knowledge for the first time in the nanotribological context, displays a wealth of interesting behavior. This interface offers a unique opportunity to explore the electronic origins of ultralow friction forces rather than structural ones since it presents a high level of coincidence between the two materials.

At zero vertical load, the interface between the bulk materials presents an exceedingly small corrugation of about 0.3 meV/atom. As a direct result of this corrugation, the lateral friction forces opposing motion were found to be small. The friction coefficients were found to vary inversely with vertical load between 0.12 and 0.005 for the range of loads studied here. These values can be considered low as defined in the literature.

In contrast to the bulk interface, the calculated corrugation for the sliding of $A_{n}$ ($n = 1, 4, \text{ and } 13$) clusters on h-BN is much larger. The largest contribution to the lateral forces are due to those atoms of the cluster that are in direct contact with the h-BN layer. For the $A_{13}u$ and $A_{19}u_5$ clusters, this amounts to one and four atoms, respectively. In addition to the larger lateral forces, secondary effects such as structural deformation of the clusters upon adsorption and ripples formed in the h-BN substrate during sliding provide further evidence to the importance of the edge and size effects on the nanotribological properties.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.9b06767.

Adhesion energy of the hBN/Au(111) surface along two perpendicular directions of sliding, atomic forces in vector representation, geometry comparisons of isolated and adsorbed clusters, in-depth analysis of the forces on the clusters as they slide (PDF)

Four animated GIF sequences of the four clusters as they slide along the h-BN surface (ZIP)

AUTHOR INFORMATION

Corresponding Author

*E-mail: ustunel@metu.edu.tr.

ORCID

Daniele Toffoli: 0000-0002-8225-6119
Oguz Gulseren: 0000-0002-7632-0954
Hande Ustunel: 0000-0003-0307-9036

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

This work is financially supported byTUUBITAK, Scientific and Technological Research Council of Turkey (Grant 115F493). The computational resources were provided by the National Center for High Performance Computing (UHeM). The authors thank Dr. Mehmet Zeyyad Baykara for his valuable comments on the manuscript.

REFERENCES

instability originated from geometrically corrugated graphene on a strongly interacting substrate. 2D Mater. 2017, 4, 025079.
(28) Broyden, C. G. The convergence of a class of double-rank minimization algorithms 1. general considerations 1970, 6, 76–90.
Supporting Information for: Nanotribological Properties of the h-BN/Au(111) Interface: a DFT Study

Merve Baksi,† Daniele Toffoli,‡ Oguz Gulseren,¶ and Hande Ustunel∗,†

†Department of Physics, Middle East Technical University, Dumlupinar Blv 1, 06800, Ankara, Turkey
‡Dipartimento di Scienze Chimiche e Farmaceutiche, Università degli Studi di Trieste, Via L. Giorgieri 1, I-34127, Trieste, Italy
¶Department of Physics, Bilkent University, 06800, Ankara, Turkey

E-mail: ustunel@metu.edu.tr
1 Bulk h-BN/Au(111) interface

Figure S1: Adhesion energies per BN unit of h-BN on Au(111) as a function of sliding distance along x for Path 1 (left panels) and along y for Path 2 (right panels) as defined in the main text. These adhesion energies are calculated at a set of fixed h-BN—Au(111) distances.

2 Au clusters

The Au$_4$, Au$_{13}$ and Au$_{19}$ clusters used in the current work were first geometry optimized in cubic simulation cells of sides 21.7 Å, 30 Å and 25 Å, respectively. The optimized geometries are displayed in Figure S3. The rhombohedral Au$_4$ maintains its geometry upon adsorption
Figure S2: The xy-plane projections of the forces on the h-BN atoms in the bulk interface at d=2.8 Å are shown in vector form (upper panels).

on h-BN. It is also found to be magnetically unpolarized both before and after adsorption. On the contrary, the Au$_{13}$ geometry is significantly distorted post adsorption, as evidenced by the bond lengths shown in Figure S3. In accordance with this distortion, the magnetic moment goes from very close to 5 $\mu_B$ in the isolated state to zero as an adsorbate. The Au$_{19}$ cluster has a zero magnetic moment and largely maintains its truncated pyramidal shape both before and after adsorption. The single Au atom has a magnetic moment of 1.0 $\mu_B$ both in isolation and also when in contact with the substrate. Throughout the sliding motion reported in the main manuscript, the distorted zero magnetic moment configuration is preserved. Therefore, the corrugations reported for all clusters are consistent.

Figure S3: Au$_4$ (left panels), Au$_{13}$ (middle panels) and Au$_{19}$ (right panels) clusters geometrically optimized in isolation and in contact with the underlying h-BN substrate.
Figure S4: Initial geometrically optimized Au$_1$, Au$_4$, Au$_{13}$ and Au$_{19}$ clusters on the h-BN substrate. The equilibrium vertical distances are also given.

Figure S5: Comparison of the total "exact" Hellman-Feynman forces on the clusters against the numerical derivative of the PES along the sliding direction.

Figure S6: A "layer-by-layer" decomposition of the total lateral force on Au$_4$, Au$_{13}$ and Au$_{19}$.
Figure S7: The extent of the ripples caused by the clusters on the h-BN sheet throughout the sliding path. The definition of the parameters $\Delta z$ that quantifies the extent of the ripples (top) and $\Delta z$ as function of displacement, $x$ along the sliding path for all three clusters (bottom).