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Atomistic Insight into the Formation of Metal-Graphene One-Dimensional Contacts

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Motivated by the need to control the resistance of metal-graphene interfaces, we have simulated the structural and transport properties of edge contacts upon their formation. Our first-principles calculations reveal that the contacts evolve in a nontrivial way depending on the type of metal and the chemical contamination of the graphene edge. In particular, our results indicate that the origin of the low experimental resistance of chromium-graphene edge contacts is related to their weaker variation upon contamination and defect formation. In summary, by analyzing the distance dependence of the graphene-metal interaction and the relation between the reactivity and forces at the graphene edge, we shed new light on the mechanisms responsible for the diverse performance of experimentally fabricated graphene edge contacts.

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I. INTRODUCTION

The idea of using graphene in electronic components, such as interconnects, comes naturally due to the material’s small volume and excellent electrical and thermal properties [1,2]. However, many of the envisioned applications involve the formation of contacts between graphene and metallic electrodes [3–7], which will necessarily alter the electrical characteristics of graphene near the contact region [8–10]. Accordingly, huge efforts are being made to design metal-graphene (M-Gr) interfaces with minimal contact resistance [11–16].

Achieving a good contact performance has been proven difficult to achieve with conventional on-top contact schemes, in which the large overlap between the graphene π-system and the metal often yields a large contact resistance [17–21]. An alternative approach is to use edge contacts, where the graphene edge is directly connected to the metal, forming a one-dimensional (1D) metal-graphene interface [22]. Within this new contact scheme, the spatial overlap between the graphene and the metal is reduced to a minimum and, ideally, a covalent bond is formed between the metal atoms and the graphene. This leads to substantially lower resistance in edge contacts as compared to on-top contacts [22–28].

Despite their outstanding potential, understanding and controlling the transport properties of edge contacts remains a challenging task, mainly due to the difficulty of obtaining statistically reproducible M-Gr 1D contact geometries. Indeed, considerable device-to-device resistance variations have been observed [15,22,26], which can be ascribed to the use of different fabrication strategies. For instance, before electrode deposition the graphene is often shaped by reactive ion or plasma etching. This technique is efficient in exposing graphene edges and removing residues [15,22,25,29,30], but might also significantly alter the structural and chemical conformation of the contact.

A few first-principle calculations have already discussed the influence of chemical contamination by foreign atomic species in graphene edge contacts, but assuming chemically abrupt M-Gr interfaces at equilibrium bonding conditions [22,31]. Yet, the final properties of the contacts are most likely governed by electrochemical processes involving both graphene (clean or contaminated) and metal. Additionally, the chemical reactivity during the contact formation will also depend on the metal of choice. This complex atomic-scale scenario is not accessible experimentally, which indicates an urgent need for a systematic theoretical study.

In this work, we investigate the formation of M-Gr edge contacts using different contaminants and metal electrodes. Special attention is paid to the reactivity of the metal and the graphene edges, a relevant question so far neglected in the study of this type of contact. Based on our first-principles calculations, we elucidate the complex dependence of the graphene edge-contacts resistance on both the termination chemistry of graphene and the metal work function [32,33].
II. METHODS

The electronic structures and geometries of 1D metal-graphene contacts upon their formation are calculated using density-functional theory (DFT), as implemented in the SIESTA code [34]. To describe the model graphene edge contact schematically depicted in Fig. 1(a), we use a supercell made up of a metallic slab containing five layers of Ni(111), Cr(110), TiO(001), or Au(111), and a graphene sheet containing 40 C rows. The graphenic subsystem is passivated with H at the edge furthest from the contact region, and unpassivated or functionalized with F, F2, or O at the edge facing the metal surface. This choice of contaminants is based on stability calculations of different graphene edge functionalizations, performed using the VASP [35–38] code (for details see Ref. [39]).

In order to determine the relevant contact configurations, we first calculate the total system energy as a function of the separation distance between both subsystems (metal, and pristine or contaminated graphene sheet) at frozen geometries. This allows us to approximately estimate the minimum-energy and maximum-force separations, from where all the atoms of the graphenic subsystem (C, H, and contaminant atoms) as well as the three topmost metal layers are allowed to relax (for more details about metal-graphene distances see Ref. [39]). The optimized interface geometries obtained in this way are referred to as minimum-energy and maximum-force contact conformations throughout the paper. It is worth mentioning that more realistic interface structures might be obtained using modern stochastic methods [40,41]. However, this would pose a challenging task that falls outside the scope of the present work. Thus, herein we focus on the optimized structures obtained by standard DFT, which might indeed serve as excellent starting points for future studies based on stochastic approaches.

Based on the relaxed contact geometries, we compute the elastic transmission with TRANSIESTA [42,43], which combines DFT with nonequilibrium Green’s function methods. The scattering region is constructed by connecting the metal and graphene electrodes at the left-hand and right-hand sides of the contact region, aiming to prevent spurious effects related to the graphene channel length. Due to the asymmetric arrangement of the two electrodes, a buffer region of 16 C atom rows is introduced. Regarding the termination of the graphene edge, it is found to have a minor influence on the resistance of the uncontaminated contacts (see Ref. [39]). Although the scenario is likely to change in the case of contaminated graphene edges, to allow direct comparisons between different contacts to the same metal, here we have decided to consider only zigzag-edged graphene both for uncontaminated and chemically modified cases. Moreover, since the metal contacts used experimentally are not crystalline and thus difficult to simulate, we have chosen one single crystalline orientation for each of the metals considered. This approach allows us to focus on trends, making our results for different systems fairly comparable.

For all the calculations, a Monkhorst-Pack k-point grid with $23 \times 3 \times 1$ k-points is used for the Brillouin zone sampling and the mesh cutoff is set to 400 Ry. A double-$\zeta$ polarized basis set is utilized with 0.02 Ry energy shift to control the cutoff radii of the basis orbitals. The exchange-correlation energy is approximated by the Perdew-Burke-Ernzerhof (PBE) [44] flavor of the generalized gradient approximation (GGA). Structures are relaxed until forces are smaller than 0.01 eV/Å. The transmission functions are sampled over $767 \times 5 \times 1$ k-points.

Based on the first-principles transport results, the effect of gate voltage is modeled by a rigid shift of the Fermi level. The thermally averaged resistance as a function of gate voltage ($V_G$) is then obtained from the calculated transmission curve $T(E)$ as follows [26]:

$$\frac{1}{R_C} = \frac{G_0}{L} \int T(E) \frac{e^{(E-(E_F+eV_G))/k_BT}}{1 + e^{(E-(E_F+eV_G))/k_BT}^2} \frac{dE}{k_BT},$$

FIG. 1. (a) Side (left) and top (right) view of a model M-Gr edge contact. (b) Calculated contact resistance at 300 K as a function of gate voltage ($V_G$) for different graphene edge contacts with different metals, namely Ni (orange), Cr (brown), Ti (brown), and Au (cyan). The intrinsic resistance of free-standing graphene is included as reference (black curve). A gate voltage of $V_G = \pm 0.2$ V corresponds to a doping level of approximately $4 \times 10^{12}$ cm$^{-2}$. 
where $L$ is the transverse length of the contact along the periodic $x$ direction, $G_0$ is the quantum of conductance, and $T$ is the temperature.

III. RESULTS AND DISCUSSION

A. Conductance properties of clean contacts

To examine whether the wide range of resistance values reported for different metals can be attributed to their intrinsic different nature [22,26], we first study clean M-Gr contacts at equilibrium. Figure 1(b) shows the calculated resistance of uncontaminated edge contacts to a representative set of metal surfaces [31], i.e., Cr(110), Ti(0001), Ni(111), and Au(111). Unexpectedly, the maximum resistance is obtained when contacting graphene to Ti. In fact, the performance is even worse than with the more inert Au electrode. The origin of such poor electronic transmission across the Ti-Gr interface is the mismatch between the Fermi level states of the metal and those at the graphene edge, as was reported to occur for epitaxial graphene on Ti [45]. In any case, despite small variations, all four metals exhibit contact resistances of the same order of magnitude, within roughly a factor of two from the ideal contact resistance of graphene. Our calculated resistances are in qualitative agreement with those reported in Ref. [46], despite some numerical discrepancies which can be attributed to methodological issues (in particular, to the short channel length employed in their studies). Remarkably, this indicates that the experimental orders-of-magnitude disparity in the resistance of edge contacts can not be solely due to the use of different metal electrodes. For this reason, in the following we turn our attention to the influence of graphene’s chemical termination.

B. Interface reactivity of contaminated contacts

The plasma employed to etch graphene is usually generated from a mixture of gases containing fluorine (F) or oxygen (O) [22,25,26,47] and, thus, we consider such elements as possible edge contaminants. According to our energetics considerations (see details in Ref. [39]), mono- and difluorination [48], and mono-oxidation are found to be the most favorable chemical edge modifications. For each of these contaminant groups, we mimic the experimental fabrication process by investigating the contact formation upon approach of the metal to the graphenic system (see the details in the Methods section). Particular focus is placed on the maximum total attractive force formation, as this corresponds to the separation at which metal and graphene will be most reactive.

Figure 2 shows the forces in the $z$ direction (perpendicular to the metal surface) on the atoms of the clean or contaminated graphene sheet, at a distance of maximum total attractive force. The following representative systems are studied: (i) three Ni-based contacts, composed of Ni(111) and clean, O-contaminated, or F-contaminated graphene (Ni-Gr, Ni-O-Gr, and Ni-F-Gr, respectively); (ii) a Cr-based contact composed of Cr(110) and F-contaminated graphene (Cr-F-Gr). Blue (red) shades indicate forces on the individual atoms pointing towards (away from) the metal surface. To allow direct comparisons, the forces for the contaminated graphene contacts are rescaled with respect to those for clean graphene. The arrows in Fig. 2 indicate the magnitude and direction of the forces on the atoms closest to the metal, i.e., on the first C row for the clean Ni-graphene contact, and on the F or O atoms for the contaminated contacts. In all cases, the net forces on the edges of the graphenic subsystems are pointing towards the metal surface, i.e., graphene feels an attractive force exerted by the metal. We see how the magnitude of the force strongly depends both on the chemical termination of graphene as well as on the metal of choice. For the Ni electrode we find that the attraction is maximum for the clean metal-graphene interface, the attractive force always being reduced upon contamination. However, the degree to which the force is quenched depends on the type of contaminant: for the oxidized case the forces remain quite large [Fig. 2(b)], while in the fluorinated case they become extremely small [Fig. 2(c)]. Interestingly, if one substitutes Ni by Cr, the force on the fluorinated graphene edge increases significantly [Fig. 2(d)], while it remains almost constant upon contamination with O (see Fig. S7 in [39]).

In addition, we investigate to which degree the interface evolution depends on the specific contacting site between the metal and the graphenic system. With this aim, we
consider a Ti(0001) electrode and a difluorinated graphene sheet, which are brought into contact at two different relative positions, as shown in Fig. 3. Both systems are fully relaxed from their maximum-force conformation (upper figures) and, interestingly, the equilibrium structures are remarkably different (lower figures). For the initial contact configuration of Fig. 3(a), only one of the F atoms gets stripped off from the graphene edge and bonds to the metal. For the contact geometry in Fig. 3(b) instead, both F atoms are removed from the edge and adsorbed onto the metal substrate. In the latter case, the graphene eventually forms a clean contact to Ti and the contact resistance is thus significantly reduced.

C. Resistance characteristics of 1D contacts

The results presented in the previous section clearly indicate that an appropriate interpretation of experimental results requires more realistic considerations of the complex interplay between the interface structure and its reactivity. Accordingly, we carry out a systematic analysis of the structural and chemical changes of the 1D contacts upon their formation, for all the possible combinations of metal electrodes (Au, Ni, Cr, and Ti) and graphene terminations (clean, and F-, F₂-, or O-contaminated) considered in this work. We find that, when transition metals are used, the final contacts always fall in one of the three conformations schematically depicted in Fig. 4(a). Thus, the F₂ modification of the graphene edge, despite being energetically more favorable for isolated graphene [48], is not likely to occur under the influence of transition-metal electrodes in edge-contacted topology. When Au is used, instead, the graphene edges (clean or contaminated) are not modified by the interaction with the metal. The reason behind this is the low reactivity of Au and the subsequent weak interplay between the metal and the graphene edge, which might also explain the bad performance of experimentally fabricated Au-based 1D contacts [22].

Having determined the structural and chemical conformation of the 1D contacts, we now focus on their conductance properties. The calculated zero gate voltage resistances of the different interfaces studied here are compiled in Fig. 4(b). From our results, it is clear that, regardless of the metal, the presence of contaminants reduces the efficiency of electron transmission across the contacts. However, the degree to which the contamination reduces the conductance and, thus, deteriorates the contact performance depends on the chemical species at the graphene edge. Oxygen contamination does not significantly alter the contact resistance for Ni, Cr, and Ti. In contrast, when graphene is terminated with fluorine, the resistance is always drastically enlarged (by at least an order of magnitude). There are two intertwined factors that play a role in this increase of contact resistance upon F contamination. The first is that F can only form one covalent bond and, therefore, the contact between the metal electrode and the graphene edge is quite weak. The second is
the larger separation distance between the metal and the F-
contaminated graphene edge as compared to the clean one
(see Table S1 in [39]). As expected from its inert charac-
ter, these two effects are magnified for Au, giving rise to a
contact resistance more than an order of magnitude larger
than for other metals.

The contacts employed in our simulations, made up
of perfect crystalline metals and graphene with straight
edges, are highly idealized structures. Experimental con-
tacts will deviate in different ways from these theoretical
systems, e.g., structural defects are expected and chemical
contamination is most likely not uniform. Figure 5(a) dis-
plays a schematic model of a corrugated metal-graphene
junction, which contains an irregular graphene edge with
inhomogeneous fluorine and oxygen contamination. In this
model, we consider the following representative distances
from the metal to the graphene edge (clean or contam-
inated): \( d^1 \) (distance of minimum energy), \( d^2 \) (distance
of maximum force), and \( d^3 \) (distance of maximum force
plus 1 Å). Taking this model as a reference, and using
Ni and Cr as representative metals, transport calculations
for each distance and different graphene edge terminations
(clean, oxidized, mono- or difluorinated) are carried out
separately. The realistic system can be viewed as a com-
bination of the different simulated scenarios and, thus, we
can extract a range of values for the resistance of the model
contact in Fig. 5(a) from the calculations reported in the
following.

Figures 5(b) and 5(c) show the zero-gate-voltage resist-
ance for clean, oxidized, and mono- or difluorinated
graphene sheets at different distances from the Ni and Cr
metal electrodes (additional results in [39]). As expected,
the resistance rises as the graphenic subsystem gets further
from the metal. However, the degree at which the qual-
ity of the contact is deteriorated depends on the type of
metal. For Ni contacts, the resistance smoothly increases as
the graphene sheet is moved further away. The increase is
more pronounced for contaminated interfaces, particularly
for fluorinated ones. At \( d^3 \), the resistance of Ni-F-Gr is
more than two orders of magnitude larger than at the equi-
librium distance.

Interestingly, the situation is quite different for Cr(110).
When connected to clean graphene, we find that the con-
tact resistance does not depend much on the Cr-Gr sep-
oration distance. This behavior can be ascribed to the
relatively short bond distance, both at minimum-energy
and maximum-force conformations. Furthermore, when
the graphene edge is contaminated, either with F or O,
increasing the distance to the Cr electrode yields a lower
relative variation of the resistance as compared to Ni [31].
In other words, the quality of Cr-based contacts is less
damaged upon contamination. We can thus conclude that
Cr is the best candidate to form 1D graphene contacts, in
agreement with the experimental observations reported by
Wang et al. [22].

D. Spin polarization at contaminated interfaces

Finally, the spin-polarization character of the electron
transport across 1D M-Gr contacts is addressed, using Cr-
F-Gr and Ni-F-Gr as illustrative systems. Figure 6 shows
the spin-resolved density of states projected onto the inter-
face atoms of a Cr-F-Gr contact, at the representative
separation distances described above. At the distance of
minimum energy (\( d^1 \)), the coupling to the metal destroys
the spin-polarized attributes of the graphene zigzag edge,
and the corresponding projected density of states (PDOS)
exhibits spin-unpolarized behavior. As the graphenic sub-
system is detached from the Cr surface, the interaction with
the metal is weakened and a net spin polarization rapidly
builds up. Indeed, at \( d^3 \) the PDOS on the first C row is com-
parable to that for isolated hydrogen-passivated zigzag-
edged graphene. In the case of Ni-F-Gr, instead the elec-
tronic properties exhibit strong spin-polarization character
at any separation distance, which can be attributed to the
intrinsic spin-polarized nature of Ni (see Fig. S10 in [39]).
FIG. 6. Spin-up (↑) and spin-down (↓) calculated density of states projected onto the interface atoms of the Cr-F-Gr contact, at distances \(d^1\) (top panel), \(d^2\) (middle panel), and \(d^3\) (bottom panel). In the bottom panel the corresponding PDOS of a graphene zigzag edge is shown for comparison. For visualization purposes, the PDOS of F has been multiplied by 4.

The spin-polarization properties studied above will eventually manifest in the conductance across the contacts. In fact, as shown in Table I, the Ni-based contact exhibits a large spin-filtering efficiency at any separation distance. In contrast, when Cr is used as electrode material, the spin polarization of the conductance predicted at large separation distances is completely quenched when the graphene is bound to the metal. Hence, the spin-polarization properties of the contact might be greatly modified both by contamination and by the use of different metallic electrodes.

IV. CONCLUSIONS

In summary, we investigate the role of atomic-scale chemical reactivity on the formation and resistance performance of metal-graphene 1D contacts. We demonstrate that the final interface configuration depends on a delicate interplay between the type of metal, the contamination of the graphene edge, and the contacting site. Moreover, structural defects and chemical inhomogeneities are found to induce significant changes in the conductance properties and even the appearance of spin-filtering effects. Interestingly, we find that Cr-based contacts exhibit the lowest dependence on contamination and structural irregularities, which explains why 1D contacts fabricated using Cr electrodes have the lowest interface resistance.

All in all, our results allow us to conclude that the use of different metals and etching agents is crucial to produce 1D contacts with resistance values suitable for their use in real applications. Of course, the actual resistance of a 1D M-Gr contact depends on the density and distribution of defects and contaminants, which in turn is conditioned by the exact fabrication process. This kind of information is very difficult to extract experimentally and, thus, a quantitative assessment of measured resistances is out of the scope of this work. Nevertheless, our atomistic simulations are very valuable for understanding the origin of the device-to-device variations and, more importantly, might be helpful for guiding the future design of metal-graphene contacts.

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