Search for encapsulation of platinum, silver, and gold at the surface of graphite

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Using scanning tunneling microscopy, we show that Pt clusters can be encapsulated beneath the surface of graphite, whereas Ag and Au cannot. This is in complete agreement with independent predictions from density functional theory, which show that surface intercalation of single metal atoms is favorable for Pt, but unfavorable for Ag and Au. This supports the validity of using single-metal-atom energetics for predicting encapsulation of metal nanoparticles at the graphite surface. We also demonstrate that the optimal temperature for encapsulation scales with the cohesive energy of the metal.

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I. INTRODUCTION

Many applications of solid metals—as catalysts, magnets, sensors, heat sinks, or electrodes, to name a few—are most efficient and cost effective when the surface-to-volume ratio of the metal is high. Recently, we have reported that high surface-to-volume ratios in metals can be achieved by encapsulating metal nanoclusters at the surface of a layered material, graphite. Effectively, the layered material resists deformation, thereby forcing the metal cluster between the layers to adopt a much flatter profile (higher aspect ratio) than it would otherwise [1,2]. Meanwhile the valuable properties of the metal may remain accessible or even be enhanced, e.g., in catalysis [3,4], magnetism [5], photonics [6], or other applications, despite being covered by a graphene layer(s).

The question then naturally emerges, which metals can be encapsulated at the surface of graphite? We have found that transition metals Cu, Ru, and Fe, as well as the rare earth Dy, can be embedded, provided two conditions are met in synthesizing the metal-plus-graphite surface system [7–10]. First, defects must be created on the clean graphite surface, to provide entry portals for the metal [11]. We achieve this by ion sputtering with argon. Second, metal must be deposited on the defect-rich graphite surface at elevated temperature, rather than being deposited at low temperature and heated. We believe the elevated temperature is necessary to prevent the portals from becoming blocked by metal atoms [11], which have a high affinity for undercoordinated carbon atoms such as those at steps and defects [12,13]. Dissolution of metal clusters at portals engenders an expectation that the temperature required for encapsulation should scale, at least roughly, with the cohesive energy of the metal [9].

The observation of surface encapsulation of these metals is somewhat surprising, given that there is no precedent for the transition metals (Cu, Ru, Fe) to form bulk graphite intercalation compounds [14]. In the case we have studied where precedent does exist—for the rare earth Dy [14]—features of Dy clusters enclosed at the graphite surface depart significantly from those known for Dy enclosed in the graphite bulk. One main difference is that Dy clusters at the surface consist of three metal layers, whereas a Dy layer in the intercalation compound does not exceed a single layer in height [7]. The transition metals can form even taller surface-encapsulated clusters, up to 200 metal layers for Cu [8]. Clearly, the forces and processes that lead to surface encapsulation are different than those that would lead to bulk intercalation.

We have, in fact, conjectured that the formation of encapsulated clusters may be driven, in part or in whole, by surface kinetics rather than thermodynamics (depending on the metal). This point is made most clearly in the Cu/graphite system, where density functional theory (DFT) showed that there is no driving force for encapsulation of Cu multilayers, despite the experimental observation of very large embedded Cu nanoclusters. However, DFT showed that there is a large driving force for encapsulation of single Cu atoms [8]. This would lead to a high population of Cu atoms in the gallery beneath the topmost graphene monolayer (GML). These Cu atoms could then nucleate and become trapped in the gallery.

In this paper we extend the investigations to three more transition metals: Pt, Ag, and Au. Pt nanoclusters are, of course, of general interest due to the broad catalytic properties of Pt [15]. Ag and Au nanoclusters generate interest, not only...
because of their catalytic properties [16,17], but also because of their useful photonic properties [18,19]. We show that Pt nanoclusters can be embedded in the graphite surface, but Au and Ag cannot. This is completely supported by our DFT calculations, which indicate that single atoms of Pt are stable beneath the topmost GML, whereas Au and Ag atoms are not.

This paper is organized as follows. Section II provides a summary of experimental methods and computational methodology, benchmarking, and energy definitions. Sections III and IV present experimental data and DFT results, respectively, for all three metals on/in graphite. Section V is a discussion, and Sec. VI concludes the paper.

II. EXPERIMENTAL AND COMPUTATIONAL METHODS

A. Experimental methods

Experiments were performed in an Omicron ultrahigh-vacuum (UHV) chamber with base pressure of $2 \times 10^{-11}$ mbar. In short, the metals were deposited via physical vapor deposition from an e-beam evaporator onto commercially available highly oriented pyrolytic graphite (HOPG, ZYB grade). The graphite surface was either pristine (cleaved using Scotch tape; referred to as p-graphite hereafter) or defect rich (bombarded with argon ions; referred to as i-graphite hereafter). Metals were deposited onto either type of surface while the graphite sample was held at different temperatures, $T_{\text{dep}}$. Major characterization techniques included scanning tunneling microscopy (STM; images were acquired in constant-current mode) and x-ray photoelectron spectroscopy (XPS), both performed with the sample at room temperature. More experimental details are available in the Supplemental Material [20].

B. Computational methods

DFT methodology. We perform first-principles DFT total energy calculations for the Ag-, Au-, and Pt-graphite systems using the plane-wave VASP code [21]. The projector-augmented-wave (PAW) method [22] is used for the electron-core interactions, and the optB88-vdW functional, where the exchange functional is optimized for the correlation part [23], is used to approximately account for dispersion interactions. The pseudopotentials were generated and released in 2013 by the VASP group. Spin-polarization effects and dipole corrections have been taken into account in all DFT calculations. The $\Gamma$-centered $k$ mesh will be specified for each type of system to be calculated.

In calculations for various metal-plus-graphite systems, we always use a graphite slab consisting of 4 GMLs as the substrate with a 2-nm vacuum thickness along the direction perpendicular to the slab surface, and the lateral size of the supercell is taken to be $6 \times 6$ in units of the lateral lattice constant of graphite, $a_C$. During relaxation for energy minimization, the bottom-most GML of the substrate is fixed. The $k$ mesh is taken to be $7 \times 7 \times 1$, and the energy cutoff is 600 eV. The force-convergence criterion is 0.1 eV/Å. All these parameters have been carefully tested for energy convergence.

Benchmarking. We have carried out benchmark calculations for the single-phase systems graphite, Pt, Ag, and Au. For graphite, from our optB88-vdW calculation, we obtain a lattice constant of $a_C = 0.2465$ nm (cf. experimental value $a_C = 0.246$ nm [24]). For more information, see Ref. [25]. For the three metals, in these benchmark calculations, we consistently use a $k$ mesh of $61 \times 61 \times 61$ for the primitive unit cell and an energy cutoff of 600 eV. For fcc Ag, we calculate $a_{Ag} = 0.4132$ nm (cf. experimental value $a_{Ag} = 0.4069$ nm extrapolated to 0 K [26]), and cohesive energy of 2.821 eV (cf. experimental value 2.95 eV [27]). For fcc Au, we obtain a lattice constant of $a_{Au} = 0.4161$ nm (cf. experimental value $a_{Au} = 0.4065$ nm extrapolated to 0 K [26]), and cohesive energy of 3.404 eV (cf. experimental value 3.81 eV [27]). For fcc Pt, we obtain $a_{Pt} = 0.3980$ nm (cf. experimental value $a_{Pt} = 0.3916$ nm extrapolated to 0 K [28]), and cohesive energy of 5.817 eV (cf. experimental value 5.84 eV [27]). In short, the agreement between theory and experiment is very good.

Energy definitions. To determine whether it is favorable for a metal cluster with $n$ atoms to be adsorbed on (top of) a graphite surface or intercalated underneath the top GML, we define the chemical potential $\mu_M (M = \text{Pt, Ag, or Au})$ of the metal cluster as

$$\mu_M = \frac{E_{\text{tot}} - E_{\text{graphite}} - E_M}{n},$$

where $E_{\text{tot}}$ is the total energy of the metal-plus-graphite system, $E_{\text{graphite}}$ is the energy of the graphite substrate, and $E_M$ is the energy of one metal atom in the gas phase. For one adatom ($n = 1$) adsorbed on the substrate, Eq. (1) is reduced to the conventional expression for the adsorption energy $E_{\text{ads}} = E_{\text{tot}} - E_{\text{graphite}} - E_M$.

III. EXPERIMENTAL RESULTS

A. Characteristics of graphite surfaces: General comments

Detailed characterization and descriptions of $p$- and $i$-graphite surfaces are available elsewhere [7,8]. In short, $p$-graphite has flat and smooth terraces. Triangular arrays of C atoms (three out of six atoms in each hexagon) can be resolved, with lattice spacing of 0.247 ± 0.003 nm. This agrees well with the literature value of graphite lattice spacing of 0.246 nm [24]. After subjecting $p$-graphite to ion bombardment, surface defects appear as bright but rather diffuse protrusions with height $\leq 0.35$ nm. STM reveals that defects come in a variety of shapes and sizes. Some are similar to single-atom vacancies with threefold symmetry [29–31] [Figs. 1(a) and 1(b)]. Other types presumably include multiatom vacancies and interstitial C atoms that are displaced within the graphite gallery (the space between graphene layers) as a result of ion bombardment [31–33] [Figs. 1(c) and 1(g)]. Evidence of electronic perturbation is often present around defects, manifest as a well-known $\sqrt{3} \times \sqrt{3} \text{R}30^\circ$ superlattice [34,35] [Figs. 1(e) and 1(f)]. Examples of these features are also shown in Refs. [8,10].

B. Temperature dependence: General comments

We have demonstrated that bare (unencapsulated) metal clusters predominate at low $T_{\text{dep}}$, then diminish in population
while encapsulated clusters emerge at higher temperature [7–10]. To guide any exploration of surface intercalation by different metals, it is important to gauge the optimal value of $T_{\text{dep}}$, which we define as the point where there is a large population of encapsulated clusters with minimal bare (unencapsulated) clusters adsorbed on the graphite surface. The optimal $T_{\text{dep}}$ values for the three transition metals that we have reported to date—Cu, Ru, and Fe—scale quite linearly with the metal’s cohesive energy [27], as shown in Fig. 2(a). As explained in Sec. I, we conjecture that this correlation reflects irreversible heterogeneous nucleation of metals at entry portals at low $T_{\text{dep}}$, which blocks the portals.

By fitting the data to a straight line as shown in Fig. 2(a), and using 5.84 eV/atom as the cohesive energy of Pt [27], the optimal $T_{\text{dep}}$ for Pt is predicted to be about 1080 K. Our experimental strategy is to start at a lower temperature, where bare metal clusters can be observed and characterized, then increase $T_{\text{dep}}$ and place a fine grid of 50–100 K near the value of $T_{\text{dep}}$ where optimal encapsulation is predicted.

**C. Pt growth**

*Temperature dependence.* Pt was deposited on graphite at different $T_{\text{dep}}$ as shown in Fig. 3. Focusing on Pt on i-graphite first, at 300 K, Pt forms round and small clusters (0.65 ± 0.10 nm tall, averaged over $X = 60$ measurements). At 800 K, much less Pt is observed on the surface than at 300 K. Some Pt forms tall (bright) clusters on the surface, with heights of 1.14 ± 0.24 nm ($X = 57$). The amount of tall clusters continues to decrease at 900 K. At 950 K, however, in addition to the tall clusters, a unique feature appears that has a rounded profile and also a round footprint. Examples are marked by arrows in Figs. 3(c)–3(e). Starting at 1050 K, the number density of the rounded features becomes much smaller, and the tall clusters primarily decorate graphite step edges with only a few on graphite terraces. Finally, at 1100 K, the rounded features are scarce, rendering graphite terraces quite clean. At this high $T_{\text{dep}}$, tall clusters are only found along graphite steps [Fig. 3(f)]; round features are not observed along steps. In the following, we will present evidence that
the tall features are bare clusters, and the rounded features are
encapsulated islands. The trends in dimensions and densities of
these two types of features, with increasing \( T_{\text{dep}} \), are shown
in Fig. 4, and are discussed more fully below.

As a control experiment, Pt was also deposited on
\( p \)-graphite at 1000 K as shown in Fig. 3(g). Graphite terraces
are clean without noticeable features, which contrasts Pt on
\( i \)-graphite at 1000 K where rounded features are observed as
in Fig. 3(d). This experiment demonstrates that the defects
introduced by ion sputtering are essential to produce the
rounded features.

Identification of rounded features as encapsulated Pt clusters.
To further investigate the rounded features on \( i \)-graphite
at 950–1100 K, we examine high-resolution STM images
of rounded features shown in Fig. 5. Zooming in directly
on these rounded features, we resolve a hexagonal lattice
that has spacing of \( 0.247 \pm 0.003 \) nm \((X = 26)\). The spacing
corresponds to that of the graphite lattice. Furthermore, the
graphite lattice is resolved on both graphite substrate and
edges of the rounded features, demonstrating the continuity
of the graphite lattice, as shown in Figs. 5(a’) and 5(b’). The
robust observation of a continuous graphite overlayer serves
to identify these rounded features as encapsulated Pt clusters.

Atop all of the rounded features with resolved graphite
lattice, triangular arrays of C atoms are observed, where only
three out of six C atoms are visible in a honeycomb. As dis-
cussed elsewhere, this threefold symmetry can signal a single
graphene sheet interacting strongly with a metal substrate,
known to occur for Ru(0001) [36–39]; or multiple graphene
sheets stacked in a way that breaks the sixfold symmetry, e.g.,
the conventional AB stacking of graphite [24]. At present the
appropriate interpretation for Pt is not clear. We note that
precedent exists for both interpretations: multiple graphene
sheets have been observed atop metal clusters for two other

FIG. 3. STM images after seven separate depositions of Pt on \( i \)-graphite at (a) 300 K, (b) 800 K, (c) 950 K, (d) 1000 K, (e) 1050 K, and (f) 1100 K, and on \( p \)-graphite at (g) 1000 K. Examples of rounded features are marked by arrows. Tip bias (\( V_{\text{tip}} \)) and tunneling current (\( I \)) are (a) 4.0 V, 0.28 nA; (b) 4.5 V, 0.27 nA; (c) 1.2 V, 0.27 nA; (d) 1.6 V, 0.27 nA; (e) 1.3 V, 0.26 nA; (f) 1.1 V, 0.26 nA; (g) 2.7 V, 0.27 nA.
metals, Cu [8] and Fe [10], whereas a single graphene sheet has been identified atop Ru [9].

In general, there are noticeable defectlike features on top of and beside encapsulated Pt clusters. These features are especially visible in Fig. 5. More examples of atomically resolved defectlike features are shown in Figs. 1(e) and 1(f). These features are similar, whether they are on encapsulated clusters or beside them on the graphite substrate. Many of these...
features have a somewhat compact, cuprous appearance, which distinguishes them from the diffuse appearance of preexisting defects that result from ion bombardment alone. They are usually slightly taller than preexisting defects [Fig. 1(b)]. Thus, it is reasonable to assign features like the brightest ones in Figs. 1(e) and 1(f) as a few Pt atoms anchored at defect sites. Elsewhere, we have made similar arguments to identify Cu-decorated defects [8].

Multiple moiré patterns have been reported on epitaxial graphene grown on Pt(111) [40,41]. However, no moiré is observed on encapsulated Pt clusters in the present study. The absence of moiré is potentially due to defects that interfere with the undulation of the C overlayer. Another possibility is that multiple graphene layers exist atop the Pt island, which would also dampen or extinguish the moiré, as they do on Ru(0001) [36].

In keeping with literature reports of low reactivity between Pt and C [42,43], there is no known form of Pt carbide. Similar to Ru, the solubility of solid C in solid Pt is low, at only 1.76 at. % at 1270 K. Thus, it is highly likely that the encapsulated Pt clusters are metallic in nature, as are other encapsulated transition metals [8–10].

Encapsulated Pt cluster densities and dimensions. Having identified the rounded features as encapsulated metallic Pt clusters, we comment on their relative density, which is shown in Fig. 4(c). At 950 K, the density of bare islands is much higher than that of encapsulated Pt clusters. At slightly higher temperatures of 1000 and 1050 K, the relationship is reversed, with encapsulated clusters outnumbering bare clusters. At 1000 K the absolute density of encapsulated clusters is about twice that at 1050 K. Together, these observations lead us to conclude that the optimal $T_{\text{dep}}$ for Pt, under our experimental conditions, is 1000 K, only 80 K below the value predicted from cohesive energies (Sec. IIIB).

We also measure the cluster dimensions as a function of $T_{\text{dep}}$ as shown in Figs. 4(a) and 4(b). The average encapsulated cluster height increases with deposition temperature, especially from 1000 to 1100 K. Cluster area increases smoothly throughout the temperature range. Thus, an increase in cluster dimensions accompanies a decrease in island density as $T_{\text{dep}}$ increases.

D. Ag growth

Temperature dependence for Ag: Predictions. Following the same line of reasoning as presented in Sec. III B for Pt, the predicted optimal $T_{\text{dep}}$ for Ag is about 740 K, based on its cohesive energy of 2.95 eV/atom. This is shown by the open circle labeled Ag in Fig. 2(b), where the experimental value for Pt has been added to the dataset used to make the prediction. To also help guide in the search, we measured Ag 3d peak intensity (which should at least roughly correlate with Ag coverage) and binding energy as a function of $T_{\text{dep}}$, with results shown in Figs. 6(a) and 6(c). The peak intensity is measurable through 850 K, but reaches zero at 900 K, indicating that 850 K is the practical upper limit. The binding energy of the Ag 3d peak is constant at 368.5 eV, consistent with metallic Ag. We thus searched for Ag encapsulation in the range 700–850 K.

STM results, 700–850 K. Starting our examination at 700 K, Figs. 7(a)–7(d) show representative STM images. As observed for Cu [8], bare features are easily removed from the graphite surface during scanning. For example, the feature present in Fig. 7(a) is gone in several subsequent images, one of which is shown in Fig. 7(b). Locations of surrounding features are circled as reference points to guide the eye. After removal of bare Ag, the STM tip usually becomes unstable, which can be remedied by pulsing the tip between a large voltage difference, for example, −10 to +10 V. There are also other bare Ag features [Fig. 7(c) and 7(d)] but they are all removed by the STM tip during scanning, even though they can be quite large in size, as shown by the profiles in Figs. 7(e) and 7(f).

At 800 K, Ag deposited on $i$-graphite generally results in graphite step edges decorated by bare clusters and relatively empty graphite terraces as seen in Fig. 7(g). Small bare clusters on terraces are about 1.0–1.2 nm tall, as shown in Figs. 7(h) and 7(k). Some of the bare clusters are unstable under tunneling, so they can be removed by the STM tip, resulting in minor streaking in scanning as pointed by an.
arrow in Fig. 7(g). This behavior is similar to Cu deposited on $i$-graphite and $p$-graphite at 300 K, where bare Cu clusters are unstable during scanning and can be picked up by the STM tip [8].

Rarely, there is an area that has a significantly higher coverage of bare Ag than elsewhere on graphite as shown in Fig. 7(j). An arrow points out a feature with a rough top and hexagonal footprint. In the process of zooming in on this feature, tunneling becomes unstable, and the hexagonal feature along with surrounding lumps are removed by the tip. In this case, the removal of bare Ag likely results from reducing the bias voltage during zooming in, which leads to a smaller tip-sample distance and causes the tip to interact more strongly with the Ag. This indicates that the hexagonal feature is a bare Ag island that is unstable under tunneling, similar to the large clusters in Figs. 7(a)–7(d).

After Ag deposition at 850 K, the graphite terraces are almost cluster free, even cleaner than at 800 K. Terraces show defects resembling the clean ones in Figs. 1(a) and 1(b), indicating that the defects no longer trap Ag. Very rarely, a larger feature is observed on a terrace, but it exhibits the streaking characteristic of a bare cluster.

Overall, the above observations of instability during tunneling suggest that the clusters observed after Ag deposition at 700–850 K are bare clusters. In contrast, encapsulated metal clusters are always very stable under tunneling, based on data for Cu, Fe, Ru, and Pt. For Ag, we find no evidence of stable, encapsulated metal clusters.

E. Au growth

Temperature dependence for Au: Predictions. Following the same line of reasoning as presented in Sec. III B for Pt or Sec. III D for Ag, the predicted optimal $T_{\text{dep}}$ for Au is about 830 K, based on its cohesive energy of 3.81 eV/atom [see Fig. 2(b)]. However, from XPS, Au coverage approaches zero at about 700 K [Figs. 6(b) and 6(c)]. Also similar to Ag, the binding energy of the Au $4f$ peak is constant at 84.1 eV, consistent with metallic Au. This suggests that it may be impractical to reach conditions where Au becomes encapsulated,
under our experimental conditions. We nonetheless searched for Au encapsulation in the range 300–700 K.

STM results, 300–700 K. As shown in Fig. 8(a), Au deposited on i-graphite at 300 K forms bare clusters that are 1–3 nm tall. Similar to Ag and Cu on i-graphite, bare Au clusters can be perturbed by the STM tip during scanning, causing frequent streakiness and image instabilities. To reduce or prevent streaking, bias voltage was usually kept high (>4.0 V) during scanning.

Additional depositions of Au on i-graphite above 300 K, in the range $T_{\text{dep}}=500–700$ K, result in decreasing coverages of Au on the graphite surface, evident in Figs. 8(b)–8(d) and the downward trend in Fig. 6(c). There is no evidence of encapsulated Au clusters. In order to broaden our parameter space somewhat, we performed two other tests: (i) increasing defect density in graphite by increasing ion emission current during ion bombardment, prior to Au deposition; and (ii) depositing Au at 300 K, then annealing to 700 K. In both cases the only result was bare Au clusters on terraces, with STM images similar to those in Figs. 8(a)–8(d). There was no evidence of encapsulated Au clusters.

In summary, Ag and Au do not form encapsulated clusters at the graphite surface, whereas Pt does. We next examine whether this is consistent with theoretical predictions.

IV. DFT CALCULATIONS

Pt ± graphite. We first examine the adsorption energies of one Pt adatom after relaxation from the seven high-symmetry positions on graphite: TH, TT, HT, BM, MB, MM, and CC, as shown in Fig. 9(a). As shown in Table I, the most favorable adsorption (top) site is at BM with adsorption energy $-1.813$ eV. The adsorption of a Pt adatom at MB, CC, and MM sites is unstable, and the adatom will move to a local minimum close to the initial site after full relaxation. Thus, the minimum energy path for diffusion of the adatom should be along TT to BM to TH, and the lower limit of the diffusion barrier is about $0.161$ eV.

| Site  | $\mu_{\text{Pt}}$ (eV) | $\Delta\mu_{\text{Pt}}$ (eV) | $m$ (Bohr magneton) | Notes                      |
|-------|------------------------|----------------------------|---------------------|-----------------------------|
| tBM   | $-1.813$               | $0.000$                    | $0.000$             | Local equilibrium           |
| tCC   | $-1.094$               | $+0.719$                   | $0.039$             | Moves to tBM                |
| tHT   | $-1.669$               | $+0.144$                   | $0.000$             | Local equilibrium           |
| tMM   | $-1.652$               | $+0.161$                   | $0.000$             | Local equilibrium           |
| tTH   | $-2.045$               | $-0.232$                   | $0.000$             | Local equilibrium           |
| tTT   | $-2.037$               | $-0.224$                   | $0.000$             | Local equilibrium           |

TABLE I. DFT results for chemical potential $\mu_{\text{Pt}}$ (in units of eV) and magnetic moment $m$ (in units of Bohr magneton per cell) for adsorption and intercalation of one Pt atom after full relaxation at different initial sites (“t” denotes adsorption on top of graphite, and “i” denotes intercalation beneath the top GML). Sites BM, CC, etc., are defined in Fig. 9(a) and its caption. $\mu_{\text{Pt}}$ is calculated from Eq. (1). $\Delta\mu_{\text{Pt}}$ is the energy (or chemical potential) difference relative to the lowest-energy adsorption site, tBM.

$^{a}$il1 is a local minimum at midpoint between iBM and iTT.
$^{b}$il2 is a local minimum (lowest) close to midpoint between iMB and iTT [see Fig. 9(b)].
$^{c}$il3 is a local minimum close to iCC, with a small shift of top GML.
TABLE II. DFT results for chemical potential $\mu_{Ag}$ (in units of eV) and magnetic moment $m$ (in units of Bohr magneton per cell) for adsorption and intercalation of one Ag atom after full relaxation at different initial sites (“t” denotes adsorption on top of graphite, and “i” denotes intercalation beneath the top GML). Sites BM, CC, etc., are defined in Fig. 9(a) and its caption. $\Delta \mu_{Ag}$ is calculated from Eq. (1). $\Delta \mu_{Ag}$ is the energy (or chemical potential) difference relative to the lowest-energy adsorption site, tTH.

| Initial site | $\mu_{Ag}$ | $\Delta \mu_{Ag}$ | $m$ | Notes |
|-------------|-------------|------------------|-----|-------|
| tBM         | -0.281      | +0.003           | 0.993 | Local equilibrium |
| tCC         | -0.280      | +0.004           | 0.996 | Local equilibrium |
| tHT         | -0.272      | +0.012           | 1.001 | Local equilibrium |
| tMB         | -0.278      | +0.005           | 1.001 | Local equilibrium |
| tMM         | -0.283      |                 | 0.990 | Local equilibrium |
| tTH         | -0.282      | +0.001           | 1.003 | Local equilibrium |
| iBM         | 0.578       | +0.861           | 0.000 | Local equilibrium |
| iCC         | 0.542       | +0.826           | 0.000 | Local equilibrium |
| iHT         | 0.515       | +0.798           | 0.000 | Local equilibrium |

0.14 eV [the true value would require a nudged elastic band (NEB) calculation].

We also look at the energies of one Pt atom intercalated beneath the top GML. Any intercalated (i) site is more favorable than any top (t) site (Table I), and the most favorable i site is close to the midpoint between tMB and iTT, which is shown in Fig. 9(b). We designate it iL2, and its chemical potential of $-2.045$ eV is $0.232$ eV lower than the adsorption energy at tBM. Therefore, for one Pt atom, intercalation beneath the top GML is always more favorable than adsorption on top of the graphite surface. We also note that both the adsorption system and the intercalated system have zero or nearly zero magnetism (Table I).

$Ag + graphite$. Let us next examine the adsorption energies of one Ag adatom after relaxation from the same seven initial positions shown in Fig. 9(a). Although the lowest adsorption energy of $-2.083$ eV is at the tTH site, all local energy minima for adsorption are very close. The adsorption energies at iTT and tHT are $-0.282$ and $-0.272$ eV, respectively. Turning to the energies of one Ag atom intercalated underneath the top GML, all i sites are less favorable than t sites (Table II). The most favorable i site is at iTT with chemical potential of $+0.515$ eV, which is $0.789$ eV higher than the adsorption energy at tTH. Therefore, for one Ag atom, intercalation beneath the top GML is always less favorable than adsorption on top of the graphite surface. As an aside, the adsorption system is magnetic and becomes nonmagnetic after intercalation (Table II). (This feature is very similar to Cu adsorption and intercalation at another layered material, MoS$_2$ [44].)

$Au + graphite$. Finally, we look at the adsorption energies of one Au adatom after relaxation from the same initial positions. The lowest adsorption energy $-0.492$ eV is at the tTH site, which is the same as that for Ag. For one Au atom intercalated beneath the top GML, the only local stable site is iTT (again the same as for Ag). It has $\mu_M = +0.735$ eV, which is $1.227$ eV higher than at tTH. Therefore, for one Au atom, intercalation beneath the top GML is always less favorable than adsorption on top of the graphite surface. Also, the adsorption system is magnetic while the encapsulated system is nonmagnetic (Table III). (Again, this feature is very similar to Cu adsorption and intercalation on MoS$_2$ [44].)

In summary, the DFT results for single metal atoms predict that intercalation is favorable for Pt but unfavorable for Ag and Au, in agreement with the experimental results for surface encapsulation.

V. DISCUSSION

The main result of this paper is the experimental observation and theoretical prediction that Pt metal clusters can become encapsulated at the surface of graphite, whereas Ag and Au cannot. These transition metals, unlike others, have been found to not undergo encapsulation. Counterexamples include not only Pt, but also Cu, Ru, and Fe, as reported elsewhere [8–10].

This result supports the validity of using single-metal-atom energetics to predict encapsulation. As noted in Sec. I, we have postulated that the key to encapsulation is having a high population of single atoms beneath the top GML (a high-density two-dimensional gas), so that there is a high probability of nucleation and growth, leading to irreversible trapping. For each of the other transition metals, Cu, Ru, and Fe, we have reported that single-metal-atom energetics favor encapsulation, even in the case of Cu, where the stability of embedded Cu clusters or embedded Cu slabs predicts the opposite [8]. In the present study, encapsulation of Ag and Au single atoms is strongly disfavored—by 0.8 and 1.2 eV—relative to adsorption on top of graphite, whereas encapsulation of Pt is favored by about 0.2 eV. The exact reason why
Ag and Au are so different (even from Cu), in terms of their interaction with graphite, is a topic of current investigation in our group. The validity of using single-metal-atom (plus graphite) calculations to predict encapsulation is an important result, because calculations involving single metal atoms are generally simpler and less expensive than calculations involving metal clusters or metal slabs (plus graphite).

Elsewhere, we have argued that encapsulated metal islands tend to fall into two groups: Those with small sizes and high number densities (Ru, Dy) and those with large sizes and low densities (Cu, Fe) [10]. We speculated that this might reflect a division between clusters that diffuse and ripen (Cu, Fe) vs those that are fixed in place and do not ripen (Ru, Dy). Pt appears to be intermediate in its properties. The Pt island heights around 1 nm, and widths around 10 nm, resemble those of Ru and Dy [10]. However, the rather low maximum density of 36 \( \mu \text{m}^{-2} \) resembles that of Cu and Fe [10]. Perhaps in this case the low density has a different origin than diffusion-mediated coarsening. For Pt, the difference in \( \mu_M \) between the most favored adsorption site and the most favored interstitial site is only 0.2 eV. This is rather small when compared with the other three transition metals, for which the difference in \( \mu_M \) ranges from 0.5 eV for Cu to 1.2 eV for Fe. Hence the density of the single-Pt-atom gas beneath the graphite surface is relatively low, leading to a lower probability for nucleation and growth. We thus propose that the density of Pt islands is low because the probability of nucleation and growth is low.

The shapes of the Pt islands are also distinctive. All other encapsulated metal islands investigated until now have flat tops, though for Cu round-top islands can be observed in addition to flat-top islands, depending on island size [8]. In other systems the islands with flat tops also show faceted footprints, which are absent for Pt. The factors that favor round vs flat tops, and correspondingly round vs faceted footprints, are not presently understood.

VI. CONCLUSIONS

Using STM, we have shown that Pt clusters can be embedded beneath the surface of graphite, whereas Ag and Au cannot. This is in complete agreement with independent predictions from density functional theory, which show that surface intercalation of single metal atoms is favorable for Pt, but unfavorable for Ag and Au. This agreement between theory and experiment supports the validity of using single-metal-atom energetics for predicting encapsulation of metal nanoparticles at the graphite surface. This is appealing, because calculations involving single metal atoms are generally simpler and less expensive than calculations involving metal clusters or metal slabs. It also supports a model in which the density of a two-dimensional gas of metal atoms beneath the graphite layer is key, leading to nucleation, growth, and trapping of metal nanoclusters.

We have also shown that the optimal temperature of metal deposition scales well with the cohesive energy of the metal, and can be used as a predictive tool. This is important in designing searches for metal encapsulation at graphitic surfaces. Presumably, this scaling reflects the necessity of metal cluster dissolution at defect sites, which maintains open portals during metal deposition.

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