Clustering and Morphology Evolution of Gold on Nanostructured Surfaces of Silicon Carbide: Implications for Catalysis and Sensing

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ABSTRACT: A fundamental understanding of the behavior of gold (Au) nanostructures deposited on functional surfaces is imperative to discover and leverage interface-related phenomena that can boost the efficiency of existing electronic devices in sensors, catalysis, and spintronics. In the present work, Au layers with nominal thickness of 2 nm were sputter-deposited on graphenized SiC substrates represented by buffer layer (BuL)/4H-SiC and monolayer epitaxial graphene (MLG)/4H-SiC. Morphometric analysis by means of scanning electron microscopy shows that Au on BuL self-assembles in nearly round-shaped plasmonically active islands, while on MLG, a fractal growth of considerably larger and ramified islands is observed. To correlate the experimentally established differences in surface morphology on the two types of graphenized substrates with energetics and kinetics of Au nanostructure growth, the deposit–substrate interaction strength was studied using density functional theory (DFT) calculations, molecular dynamics simulations, and optical measurements. The theoretical considerations involve participation of Au clusters with different sizes and energetics at the initial stages of the metal nanostructure formation. The results indicate that gold exhibits a considerably stronger interaction with BuL than with MLG, which can be considered as a key aspect for explaining the experimentally observed morphological differences. From the statistical analysis of Raman spectra, indications of Au intercalation of MLG are discussed. The current research shows that, due to its unique surface chemistry, buffer layer has peculiar affinity to gold when compared to other atomically flat surfaces, which is beneficial for boosting high-performance catalytic and sensing technologies based on low-dimensional materials.

KEYWORDS: gold, epitaxial graphene, SiC, buffer layer, adsorption, growth

1. INTRODUCTION

Monolayer epitaxial graphene on the Si face of 4H-SiC (MLG/SiC) is an important ingredient of today’s advanced technologies in quantum metrology, electronics, and sensors.1,3,4 Due to several fundamental and practical advantages over other graphene-family materials,11 MLG/SiC has gained wide-ranging recognition as a multifunctional monolithic architecture, which can perfectly satisfy application-specific requirements while maintaining the key properties of true graphene crystal. An expected step in the development of epitaxial graphene devices is the integration of MLG/SiC with noble metals (NMs) to complement or expand the functionalities of the existing MLG/SiC host system. For instance, interfacing gold (Au) with epitaxial graphene has a huge potential to be exploited as (i) ohmic contact, due to a low Au-MLG/SiC contact resistivity of $<1 \times 10^{-6}$ $\Omega$·cm$^{-2}$,12 (ii) a catalyst for electrochemical hydrogenation of MLG,13 (iii) an active sensing component on MLG for detection of biomolecules and gases,14–16 and (iv) a plasmonically active phase enabling SERS (surface-enhanced Raman scattering) applications.17,18 In this regard, atomistic-level interfacial processes governing the growth energetics and kinetics of Au and other NM nanostructures and layers on epitaxial graphene have attracted considerable attention during the last years.19–23 Indeed, critical knowledge of the mutual interplay between NMs and MLG/SiC is required to accelerate the development of tunable NM-MLG/SiC hybrids and related devices.

In addition to MLG growth, another unique opportunity provided by SiC is the growth of the so-called buffer layer (BuL), formed at the early stages of the annealing process used for MLG synthesis. The BuL is a C-rich reconstructed SiC surface, which is partly bonded to the substrate and can be decoupled from the substrate by elemental intercalation and transformed to quasi-free standing (QFS) monolayer graphene. Some of the technological implications of buffer layer have been described in our previous works,24,25 where BuL in place of MLG enables homogeneous deposition of metals. The
homogeneous deposition of Pt is only possible on buffer layers and not on MLG (nor other atomically flat surfaces such as 4H-SiC or sapphire). In ref 24, we describe that buffer layer allows the formation of atomically thin Pt, which works as a chemical sensor with sub-part-per-billion sensitivities and fast operation. In this regard, buffer layer, which has been thought as a technological barrier for epitaxial graphene technology due to high intrinsic doping, is now considered to have its own technological relevance for catalysis and biosensing. Therefore, its high affinity to transition metals needs to be studied comprehensively in comparison with MLG surfaces. However, the major focus of existing studies of the Au-BuL system is the controllable Au intercalation beneath BuL to form high-quality QFS graphene with tunable properties. For instance, homogeneous deposition of Au on BuL followed by annealing-induced Au intercalation enables ambipolar charge transport of quasi-freestanding graphene, which may find applications in sensors and spin-related transport. Au as intercalant species is also anticipated to initiate new physical phenomena at the interface, including tuning the energy scale of van Hove singularity and Fermi velocity increase, doping control, and giant Rashba-type spin-orbit splitting. Although substantial progress has been made towards clarifying Au intercalation conditions and identification of Au intercalants’ morphology, the resulting interplay between graphene and the confined gold atoms after intercalation is still an open issue. There are hints that the morphology of intercalated Au atoms determines the properties of the resulting graphene–Au intercalated systems.

Based on the analysis of the existing scientific background and applications, it can be stated that the interaction between gold and graphenized SiC is decisive for the application-specific properties of the Au-MLG and Au-BuL systems. Fine tuning this interaction can enable manipulating Au morphology from dispersed three-dimensional (3D) nanoscale islands to continuous two-dimensional (2D) layers. Thus, a holistic understanding of Au adsorption and kinetics on the graphenized Si face of on-axis 4H-SiC is an essential precondition to realize a full control on the structural morphology of deposited and/or intercalated Au. Furthermore, because of the complex structure of the MLG/BuL/SiC system that offers several options for Au accommodation, it is of fundamental importance to distinguish between the processes occurring on the buffer layer and those on the graphene monolayer. Despite the pronounced interest in the investigation of Au behavior on graphenized SiC, such distinction is still not fully explored and remains a challenging task for researchers.

In this work, we shed light on fundamental aspects of Au nanostructures grown by direct current (d.c.) magnetron sputtering on buffer layer and monolayer epitaxial graphene synthesized on the Si face of on-axis 4H-SiC by the well-recognized method of high-temperature thermal decomposition in argon atmosphere. As far as the authors are aware, there are no studies reported on selective gold growth on BuL with MLG patches by d.c. magnetron sputtering. Considering the high technological importance of the magnetron sputtering for the formation of homogeneous metallic coatings on large-area substrates, it is sensible to investigate the morphology of vapor-deposited gold layers on both BuL and MLG and to link the structural uniqueness of gold on each type of surfaces to the specific gold–template interaction, which is a key determining factor in processes of nucleation, diffusion, and clustering. By performing complementary experimental work and ab initio calculations, we first demonstrate fundamentally important material–property relationships for the Au-graphenized SiC system. We are focusing on the investigation of the mesoscopic morphology of gold on graphenized SiC rather than on the investigation of detailed morphology features of the Au nanoclusters. We show that the BuL acts as a strongly interacting substrate for gold deposition and the presence of sp2-bonded C atoms limits the diffusion length of Au, which results in the formation of high-density isolated round islands. Concomitantly, we observe growth of larger and ramified irregularly shaped Au islands on MLG, which is explained considering the weak interaction between gold and epitaxial graphene and strong metal–metal attractive forces. The obtained results provide yet-unknown in-depth atomic and thermodynamic insights into the nature of the interaction between gold and BuL or MLG, thereby expanding the existing knowledge on gold self-assembling on atomically thin, flat, and corrugated substrates as well as promoting further development of the mentioned system for biosensing and catalysis.

2. METHODS

2.1. Experimental Procedures. To distinguish between the behavior of Au on buffer layer and graphene monolayer, we prepared corresponding templates by high-temperature (~1700 to 1900 °C) thermal decomposition of the Si face of nominally on-axis 4H-SiC (0001) substrates in Ar atmosphere. The graphenization proceeds via Si sublimation in an inductively heated graphite enclosure under strictly controlled temperature–gas pressure–time conditions, enabling to achieve a complete coverage of the 4H-SiC substrate with the C-rich 6√3 × 6√3 R30° surface pattern (BuL) due to characteristic reconstruction of the (0001) face, also known as zero graphene layer. While heating, BuL transforms into monolayer epitaxial graphene (MLG) above a subsequent buffer layer. Therefore, we prepared two types of graphenized surfaces: buffer layer and monolayer graphene over a buffer layer, respectively. BuL formation was realized at a lower temperature in comparison to the MLG growth temperature, and the BuL templates intentionally contain MLG graphene inclusions that give the opportunity to compare the different surface properties of BuL and MLG on the same SiC substrate. Both studied templates have atomic thicknesses and two-dimensional structures providing nonstandard substrates for further nanoscale metal deposition and related phenomena. Prior to the Au deposition process, we performed detailed Raman and optical reflectance mappings of the substrates to evaluate the quality of the carbon layers (thickness, homogeneity, defects, etc.). The Raman fingerprint of BuL, which includes structured bands within the frequency range of 1355–1600 cm−1, was observed at every set point of the investigated area, suggesting a homogeneous BuL coverage (see Figure S1, Supporting Information), typically with few submicrometer patches of MLG. The formation of MLG was then evidenced by the observation of the characteristic G peak (around 1602.78 ± 3.55 cm−1) and 2D peak (around 2738.73 ± 6.42 cm−1) during Raman mapping (Figure S2, Supporting Information). This observation is consistent with the topmost graphene layer being n-doped and under biaxial compressive strain. Optical reflectance mapping (Figure S3, Supporting Information) confirmed that most of the SiC surface (~81%) is covered by monolayer graphene, while the remaining ~19% are small inclusions of bilayer epitaxial graphene (BLG) regions. Notably, no defect-related D phonon mode was observed in the Raman spectra. This indicates a high crystalline quality of the epitaxial graphene, required to avoid defect-mediated Au nucleation that might mask eventual van der Waals epitaxy of Au on the basal plane of graphene.

Au layers with a nominal thickness of 2 nm were deposited on BuL and MLG by direct current (d.c.) magnetron sputtering of an Au target at room temperature and UHV conditions (base pressure <
laser with 1 mW power was used as an excitation source. The spectral focused on the sample surface. A 532 nm diode-pumped solid-state (NA = 0.95) resulting in an objective lens has a magnification on the phonon modes of graphene and BuL were explored by micro-

10^-8 Pa). The total pressure of Ar (purity 99.999%) in the working chamber was kept at 6.7 Pa (50 mTorr), while the deposition rate was maintained at 0.1 Å/s. Such mild deposition conditions were used to ensure that the epitaxial graphene remains undamaged after magnetron sputtering of Au. From our previous work, with the growth of noble metals on weakly interacting substrates, a 2 nm thickness results in a discontinuous metal layer that allows us to study film morphology while leaving a part of graphene exposed for various analyses. Concurrently, differences in morphology observed during the stage of discontinuous layer translate consistently into differences in morphology of continuous layers (i.e., more pronounced 3D morphology of islands in discontinuous layer typically means larger roughness and a continuous film is formed). Scheme 1 summarizes the steps used in the preparation of the gold-decorated BuL and MLG samples, respectively. To explore whether the observed gold morphologies are related with features of the sputtering process (energetic bombardment, plasma, etc.) or they are process-independent and can only be ascribed to the film/substrate system, extremely thin gold nanofilms were also deposited using thermal evaporation and additionally studied by atomic force microscopy (AFM).

The surface morphology and microstructure of the as-deposited Au films were examined by scanning electron microscopy (SEM) (Leo 1550 Gemini SEM instrument) at an operating voltage ranging from 10 to 20 kV and a standard aperture value of 30 mm. The quality of the Au-coated graphenized surfaces and the effect of the Au deposits on the phonon modes of graphene and BuL were explored by micro-Raman spectroscopy mapping. Raman spectra were recorded with a micro-Raman setup based on a monochromator (Jobin-Yvon, model HR-460) equipped with a CCD (couple-charged device) camera. The objective lens has a magnification of 100x and numerical aperture (NA = 0.95) resulting in an ~0.85 μm diameter of the laser spot focused on the sample surface. A 532 nm diode-pumped solid-state laser with 1 mW power was used as an excitation source. The spectral resolution of the system is ~5.5 cm^-1. Absorbance measurements were performed at room temperature using a Shimadzu UV-2100 spectrometer.

2.2. Theoretical Details. To elucidate the specifics of Au growth kinetics and energetics, we studied theoretically Au behavior on two different interfaces, namely, BuL/4H-SiC and MLG/BuL/4H-SiC. We employed the √3 × √3 R30°-reconstructed 4H-SiC (0001) substrate (Figure S4), which is consistent with the experimentally observed features of the MLG/BuL/4H-SiC system. According to this model, a 2 × 2 honeycomb array of carbon atoms is accommodated on top of the reconstructed surface of 4H-SiC (0001). To satisfy charge neutrality conditions, the dangling bonds of the bottom carbon layer (or carbon-terminated face) of a 4H-SiC slab were passivated by forming C–H bonds. The buffer layer is partly covalently bonded to the Si surface atoms of the SiC and is known to be corrugated. It does not exhibit graphene-like properties, while the first graphene layer above the BuL is an n-doped honeycomb array of C atoms with a cone-like electronic band structure.

To study the Au atom adsorption and migration on the graphenized SiC surfaces, we used the following scheme: (1) positioning the Au atom at the specific site (top site) in the beginning of the migration path → (2) geometrical optimization of the Cartesian coordinates with constraints to Au movements in x and y directions at the starting point of the diffusion path → (3) physical movement of the Au atom to the next point along the migration path followed by relaxation only along the z direction (Figure S5, Supporting Information). Despite the fact that thermal vibrations may modify diffusion behavior, the static potential energy landscapes calculated from energy minimization can provide a qualitative meaningful picture of the adsorption and migration dynamics in weakly interacting film/substrate systems, which is consistent with experimental results. The calculated adsorption energy difference at specific sites determines how long an adsorbed Au atom will remain trapped at the surface of interest: the larger the adsorption energy difference, the shorter the diffusion path. We also studied the Au nucleation process on the considered interfaces by using larger slabs: 4 × 4 BuL/4H-SiC and 4 × 4 MLG/BuL/4H-SiC, respectively (see Figure S6, Supporting Information).

All ab initio calculations were carried out by using the SIESTA code within the van der Waals density functional (vdW-DF) method. The construction of pseudopotentials for H, Si, C, and Au was performed within the Troullier–Martins scheme by using the ATOM code. A vacuum layer of 20 Å was added above the graphenized surface along the slab-normal direction to avoid undesired interactions with neighboring unit cells. Double-ζ polarized (DZP) basis set with an energy shift of 200 meV was chosen. Although DZP basis set usually allows obtaining reasonable results, some extra calculations using larger basis sets have been carried out to validate the results of the model (see Section S1, Supporting Information). The Cartesian coordinates of the considered structures were relaxed until the force on each atom reached less than 0.02 eV/Å. A Monkhorst–Pack k-point 3 × 3 × 1 mesh was used to sample the Brillouin zone during the optimization process. The mesh cutoff was set at 400 Ry. All obtained results are strictly valid only at T = 0 K and provide deeper insights into the nature of the low-energy properties of Au–template systems. A better understanding of ground-state properties will benefit further experimental work and future applications, since fundamentally crucial material–property relationships can be extrapolated to temperatures beyond 0 K.
3. RESULTS AND DISCUSSION

3.1. Morphology of Au Deposited on Buffer Layer and Monolayer Epitaxial Graphene. As a first step toward understanding energetic and kinetic aspects of Au growth on graphenized surfaces of 4H-SiC, we studied experimentally the morphology of as-deposited layers on both buffer layer and monolayer epitaxial graphene using SEM. In the case of BuL/SiC substrates (Figure 1a–c), Au self-assembles into largely isolated and nearly circular nanoscale islands with no evident interconnections between them. Analysis of the SEM images shows that the island density is \(9.6 \times 10^{11}\) cm\(^{-2}\), while the substrate area coverage is 43.7%. Moreover, the island size (i.e., projected area) follows a bell-shaped distribution with a mean size of 45.6 nm\(^2\) (Figure 1d). We also note that some parts of the BuL substrate contain MLG overlaid regions (see the outlined regions in Figure 1a–c), which show completely different Au morphology, i.e., Au islands are considerably larger in size and exhibit dendritic shapes.

The overgrowth of monolayer graphene layers on BuL allows to study the Au growth simultaneously on both surfaces and to elucidate the difference in Au growth morphology on BuL and MLG. The differences in growth morphology on the two types of graphenized surfaces are further confirmed by the SEM data from the Au/MLG/SiC sample (Figure 2), which reveal the formation of islands with an approximate mean size of 1623.7 nm\(^2\) and ramified island shapes (note that the island size in Figure 2 was estimated only based on the analysis of the isolated islands, while most of the MLG surface is covered with large and dendritic Au structures).

A similar picture was observed for 4 Å gold deposited by thermal evaporation on BuL with MLG patches (see Figure 3). In this case, Au completely covers the BuL forming a smooth
coverage. On the other hand, a pronounced Au agglomeration takes place on the MLG patches (see regions indicated by the arrows in Figure 3). The morphological similarities of Au layers obtained by different deposition methods illustrate a common trend in Au growth on graphenized SiC surfaces. More details on the specific structural features of thermally evaporated gold layers on graphenized surfaces of SiC studied by using scanning tunneling microscopy (STM) and synchrotron facilities can be found in our recently published work.25

The analysis of the SEM image of sputter-deposited Au on MLG (Figure 2) shows that the substrate coverage is 36.6%, this being smaller than that for the Au/BuL/SiC system (43.7%). Given that on both surfaces the same amount of Au was deposited, we conclude that Au exhibits more pronounced 3D growth morphology on MLG compared to BuL. From a thermodynamic point of view, the observed results could be understood by considering the difference between the surface energies of the two substrates. Since the surface energy is strongly dependent on the structural parameters (including geometric configuration of the adsorption sites) and electronic properties of the substrate, it positively correlates with the reactivity of the surface sites and, consequently, with the adsorption energy of external adsorbates: the larger the surface energy, the higher the adsorption energy. In our case, the surface energy of MLG (0.1 J/m²) is much smaller compared to that of BuL, thereby providing lower wettability and weaker nanoscale adsorption of Au during early growth stages. It should be noted, however, that the surface energy of BuL is not known exactly and is assumed to be close to that of the nongraphenized SiC substrate (1.4 J/m²). Because of the substantial difference in the surface energetics of the two material systems, it seems that the morphology of Au deposited on MLG is insensitive to the underlying buffer layer i.e., the contribution of Au-BuL interaction to the total Au-MLG/BuL interaction is negligibly small, which can be probably related to prevailing van der Waals forces on MLG. By using ImageJ software with the FracLac plugin, we performed additional analysis of the dendritic microstructure of Au islands on MLG and found that the fractal dimension is approximately equal to 1.84. Due to the thermalized nature of the deposition fluxes used in our experiments (sputtering at 50 mTorr) that cause limited atomic diffusion at island edges, the diffusion-limited aggregation (LDA) is believed to be a key mechanism underlying the formation of dendritic structures exhibiting fractal dimensions. According to this mechanism, the surface diffusion of the as-deposited gold atom can be considered as the random walk on the weakly interacting substrate containing a preformed stable nucleus. Once this gold atom reaches the nucleus, its further movement is constrained. Newly arriving gold species repeat this process, thereby forming the fractal aggregates.

Finally, we noticed that the observed morphological properties of gold correlate with its optical properties (Figure 4). More specifically, dendritic gold nanostructures on MLG show a higher absorbance compared to gold on BuL, without any pronounced feature. Meanwhile, isolated Au nanoscale islands on BuL exhibit an absorption band at 634 nm related to localized surface plasmon resonance. This feature can be used for design of highly selective sensor arrays based on buffer layer for optical detection of biomolecules.

To understand the way by which morphology differences are related with the fundamental processes of adatom adsorption and diffusion, we study the deposit/substrate interactions using Raman spectroscopy and DFT calculations, as presented in the next two sections.

Figure 3. (Left panel) AFM image and (right panel) phase image of 4 Å-thick Au deposited onto the BuL/4H-SiC surface by the thermal evaporation technique, respectively. Field of view is 1 µm × 1 µm, respectively.

Figure 4. Absorption spectra of 2 nm gold deposited on the buffer layer (blue curve) and monolayer (orange curve), respectively, as measured with UV–visible spectroscopy.

3.2. Probing Gold–Substrate Interaction by Raman Spectroscopy. Figure 5 displays typical Raman spectra taken from Au/MLG/BuL/4H-SiC, Au/BuL/4H-SiC, and the corresponding reference samples (MLG/BuL/4H-SiC and BuL/4H-SiC). As seen from this figure (cf. also Figure S1 in the Supporting Information), the Raman spectrum of the bare
buffer layer exhibits two broad phonon bands within the frequency range peaking at 1380 to 1597 cm$^{-1}$, which match qualitatively the theoretically predicted vibrational density of states (vDOS) of the buffer layer on SiC.\textsuperscript{56}

Raman mapping confirms the formation of BuL within the whole investigated area, which is evidenced by the observation of the abovementioned two bands (Figure S1, Supporting Information). Since the Raman signature of the buffer layer is not related to specific discrete Raman modes (unlike the graphene case), but is instead density-of-states-like, we discuss only qualitatively the effect of Au on the phonon properties of the BuL. After gold deposition, the Raman spectra of the buffer layer undergo a substantial change (Figure 5 and Figure S7a, Supporting Information). More precisely, the low-frequency band is replaced by two bands peaking at about 1372 and 1492 cm$^{-1}$ (the latter is denoted by an asterisk (*) in Figure 5). The new features with respect to the spectrum of pristine BuL are observed over the whole mapped area (see also Figure S7b,c).

These changes indicate structural modifications of the BuL induced by interaction that entails a modification of the vibrational density of states and, consequently, of the experimental Raman spectra of the buffer layer. All measured spectra (121 in total) on the Raman map are similar, indicating a homogenous deposition of Au islands over the surface of the BuL (cf. the mapping results in Figure S7b,c, Supporting Information).

The Raman spectrum of the Au-covered graphene monolayer expressed in the G and 2D peaks is remarkably like that of the reference sample (apart from minor peak shifts and broadening to be discussed below), but the buffer layer contribution experiences a significant change after the Au deposition, if compared to the BuL spectrum of the MLG/4H-SiC reference. We notice that the BuL contributions for the BuL sample and for the MLG/4H-SiC sample have quite different structures. However, after Au deposition, the buffer layer contributions for the samples with and without graphene have very similar structures comprising three main bands already discussed in the previous paragraph regarding the BuL after Au deposition [cf. Figure 5a,c]. The reason for this similarity is most probably due to an interaction between Au atoms and BuL; in the case of the MLG/BuL/4H-SiC sample, it is due to a possible gold intercalation underneath the topmost graphene layer. To justify this assumption, we simulated eventual vertical Au diffusion through the hollow sites of MLG (note: an empirically measured Au atomic radius of 0.135 nm\textsuperscript{57} is very close to the geometrical graphene pore size of 0.246 nm,\textsuperscript{58} which indicates a nonzero probability of Au penetration beneath graphene). The simulation details are shown in Section S2 (Supporting Information). By analyzing Figure 6, we noticed the presence of a high barrier (6.84 eV) at $z = 28.53$ Å. This point of the potential energy profile can be regarded as a transition state for which we observed a pore size

![Figure 5. Typical Raman spectra of (a) Au/MLG/4H-SiC, (b) MLG/4H-SiC (reference), (c) Au/BuL/4H-SiC, and (d) the buffer layer reference (cm$^{-1}$). The reference spectra are recorded with higher excitation power (17 mW vs 1 mW for the Au-deposited samples). In addition, the Au islands are the screening part of the Raman signal, which together with the lower excitation power stipulates the lower signal-to-noise ratio in (a) and (c). The vertical scaling is chosen for the convenient view only. The main features in the spectra are denoted with their peak positions (cm$^{-1}$) for G and 2D, the peak positions are for the spectra displayed; the statistical distribution is given in the Supporting Information). The appearance of the relatively pronounced background in the Raman spectra after Au deposition is ascribed to luminescence phenomena. If we consider the Raman luminescence background maximum at 3100 cm$^{-1}$, with a 532 nm excitation wavelength, it gives a PL signal at $\sim$637 nm, which is in very good agreement with the localized surface plasmon resonance obtained from absorbance.](https://dx.doi.org/10.1021/acsanm.0c02867)

![Figure 6. Free-energy profiles (color coded with Hirshfeld charge on Au) corresponding to intercalation of Au underneath the topmost graphene layer in MLG/BuL/4H-SiC. The most stable adsorption site has energy of 0 eV, and less stable sites have positive energies. The optimized geometrical structures corresponding to different intercalation stages are illustrated inside the figure.](https://dx.doi.org/10.1021/acsanm.0c02867)
expansion and a final destruction of the hexagonal ring of graphene (see the Figure S8, Supporting Information). Our calculations show that the intercalation barrier is causally related to the penalty for escaping the hollow site above graphene in order to occupy the hollow site from the other side of graphene. Since the sputtered Au target atoms in our experiment have energies in the order of few electron volts with energy tails of several tens of electron volts, the kinetic energies of selected Au atoms can be compatible with the energies needed for Au intercalation. Thus, some fraction of Au atoms can be accommodated in the space between MLG and BuL (the middle structure inside Figure 6), thereby influencing the phonon dispersion of BuL in the same manner as in the cases of the graphene-free sample.

Another important argument based on our computations is that the MLG/BuL/4H-SiC system with intercalated Au atoms has more negative Gibbs free energy with respect to the MLG/BuL/4H-SiC system with just the adsorbed Au atom. Since the sputtered Au target atoms in our experiment have energies in the order of few electron volts with energy tails of several tens of electron volts, the kinetic energies of selected Au atoms can be compatible with the energies needed for Au intercalation. Thus, some fraction of Au atoms can be accommodated in the space between MLG and BuL (the middle structure inside Figure 6), thereby influencing the phonon dispersion of BuL in the same manner as in the cases of the graphene-free sample.

It should be also noticed that the MLG after Au deposition preserves its characteristic Raman modes, which appear slightly shifted (downshift to $\sim 1597.75 \pm 1.52 \text{ cm}^{-1}$ for the G peak and upshift to $2742.22 \pm 4.72 \text{ cm}^{-1}$ for the 2D peak) relative to those of pristine MLG [Figure 5a,b, see also Figure S9a, Supporting Information]. More significantly, no defect-related Raman peaks were observed, this pointing to the robustness of epitaxial graphene upon Au deposition by sputtering. Further information can be acquired by inspecting the 2D vs G peak position dependence (Figure S9b). For Au-covered MLG, the points scatter primarily along a line (called the strain line) with a slope of 2.4, partly overlapping the same line for the pristine MLG sample. However, the data points shift to the left with respect to the strain line and the main group of data points for pristine MLG, which can be interpreted because of reduction of the electron density in MLG due to a screening effect. The nature of such a screening effect could be related to Au intercalation that weakens MLG-substrate interaction, decreasing the charge transfer from the buffer layer. Furthermore, the color-coded Raman maps (Figure S9c–e, Supporting Information) exhibit inhomogeneous broadening of G and 2D peaks for the MLG after Au deposition, which can be attributed to spatial charge and/or strain fluctuations at the interface.

In overall, the Raman data indicate that there is a significantly weaker interaction between Au and MLG compared to that between Au and BuL. This aspect is further explored in sections 3.3 and 3.4, where the correlation between deposit-substrate interaction strength and the experimentally

Figure 7. Free-energy profiles (color coded with Au adsorption height) corresponding to migration of Au along the diffusion path over (a) buffer layer and (b) monolayer epitaxial graphene surfaces. The most stable adsorption site has energy of 0 eV, and less stable sites have positive energies. Fluctuations of the effective charge on a Au adatom at each stage of the diffusion path for Au movement on (c) BuL and (d) MLG. The charge population analysis has been performed by using Hirshfeld and Voronoi schemes, respectively.
observed layer morphology is established using DFT calculations.

3.3. Potential Energy Landscapes for Gold Migration on Graphenized Surfaces of SiC. To quantitatively explain the experimental observations and to understand the kinetics of Au on BuL and MLG, we investigated the relative potential energy profiles (i.e., landscapes) for in-plane migration of a single Au adatom above both considered surfaces along predefined migration paths with a 0.3 Å translation step (see Figure S5, Supporting Information). A Au adatom is allowed to relax only along the z Cartesian vector, while the atomic movements along the x and y directions were constrained. The relative energy profiles color coded by Au adsorption height for the Au on BuL and MLG are plotted in Figure 7a,b, while the corresponding optimized structures related to different migration steps are shown in Figures S10 and S11 (Supporting Information), respectively.

We start by describing the Au migration on BuL. According to the data in Figure 7a, there is a significant energy difference of 350 meV between the minimum-energy adsorption site (T1) and maximum-energy adsorption on-top site position T2 (above sp3-bonded carbon). The center of the distorted hexagonal ring with evident protrusions of sp2-bonded C atoms is also an energetically unfavorable adsorption position. Thus, on the BuL, Au atoms prefer to occupy the top site position above the protruded sp3-bonded carbon atom of the BuL (T1 position). The situation is different for the MLG surface (Figure 7b) on which the Au adatom favors the hollow site position H (center of the hexagonal ring). The analysis of the relative energy profile for the Au atom on the MLG surface suggests negligibly small diffusion barriers (energy differences of 10−20 meV) among the adsorption sites (Figure 7b). Molecular dynamics calculations performed at room temperature for both considered systems additionally corroborate the abovementioned findings (see Section S3, Supplementary Information). Indeed, root mean square displacement (RMSD) in the case of Au/MLG is steadily increasing with time, which can be interpreted as a consequence of the weak interaction with graphene (Figure S12, Supporting Information). Meanwhile, the RMSD curve for Au/BuL contains a pronounced maximum and minimum, suggesting that the gold atom fluctuates around a stable adsorption site.

In addition to diffusion, we also studied the Au-position-dependent charge transfer on both surfaces (Figure 7c,d), which can be related to Au sharing electrons between different number of carbon atoms at each diffusion stage. More specifically, the largest charge transfer occurs at the hollow site for Au on MLG, which is energetically the most favorable adsorption site. Meanwhile, in the case of Au adsorption on the preferred T1 site at the BuL surface, the charge transfer is minimal. According to Voronoi and Hirshfeld population analyses, the Au adatom donates electrons to both BuL and MLG, which is evidenced by the positive charge of Au at all positions along the diffusion path. The obtained results suggest that it is easier for electrons to be transferred from the Au adatom to MLG than to BuL at each diffusion stage (see also Figure S13, Supporting Information). Since a carbon-rich buffer layer has partial sp3 hybridization, some of the carbon atoms cannot accept electrons due to the complete saturation of the covalent bonding, causing an overall reduced charge transfer from Au.

From the comparison of the relative energy profiles for gold on BuL and MLG, it can be determined that the probability for trapping gold adsorbates by the BuL is much higher than that by the MLG. As a direct consequence, the large energetic difference between T1 and T2 sites for the BuL surface will decrease the diffusion length of the Au adatom, thereby causing the formation of a larger number of early-stage nuclei compared to the MLG surface. On the contrary, the small energetic difference in the case of MLG results in an increased diffusion length of Au, which yields smaller and larger Au islands. As these islands attain relatively large size (compared to their counterparts on BuL), their shape becomes heavily dependent on the atomic diffusion at the island edges.60 Considering that Au is deposited at a pressure−distance product of 82.5 Pa·cm, at which gas-phase collisions are frequent and vapor flux becomes largely thermalized,61,62 we attribute the dendritic shape of the Au nanostructures on MLG to limited edge diffusion that prevails during their formation and growth.63 From a fundamental point of view, the early-stage growth morphology of vapor-deposited films is governed by the atomic-scale processes, which are operating during island nucleation, growth, and coalescence. Since in weakly interacting film/substrate systems63,64 the nominal thickness of 2 nm is about an order of magnitude larger than the critical thickness at which island density saturates and film morphology is governed by nucleation, it is highly likely that island coalescence is the dominant structure-forming process. Hence, the results in Figure 1 indicate faster island reshaping during Au coalescence on BuL/SiC, while in the Au/MLG/SiC system, Au cannot completely coalesce to form rounded islands and become irregularly shaped. Unexpectedly, the growth landscape of gold on MLG resembles more Au growth on graphite than on monolayer graphene.65 Indeed, as was shown by Liu et al.,65 5 Å Au on graphene/SiO2/Si is composed of many well-dispersed, compact Au nanoparticles, while using a graphite substrate led to a formation of ramified islands. Due to the reduced adatom mobility on graphene/SiO2/Si in comparison to graphite, much smaller Au island density is expected on graphene than on graphite.65 The discrepancy between our experimental data and literature strongly suggests that the SiC substrate significantly weakens the interaction between the graphene layer and gold nanostructures, thereby favoring smaller density and larger size of Au atoms.

3.4. Adsorption and Nucleation of Auₙ = 1−7 Clusters on BuL and MLG Surfaces. The details of adsorption and nucleation energy calculations are set in Sections S4−S6 (Supporting Information). The summary for single Au adatoms is that Au preferentially occupies the top site of BuL above the sp3-bonded carbon atom (Figure S14a, Supporting Information) and the hollow site of MLG (Figure S15a, Supporting Information), respectively. Au on BuL has adsorption energy of 0.820 eV, being larger than that of the Au/MLG system (0.144 eV). Such a distinct difference provides clear evidence for chemisorption of Au on the BuL surface and weak Au physisorption, mainly governed by van der Waals forces, on the MLG surface.66 For multiple gold atoms, an important finding is that the Au clusters on BuL are more stable than those on MLG, which is confirmed by the larger Eads (note that this energy includes two components: metal−template interaction and cohesive energies) values for all considered cases.

Finally, as can be seen from Table S5 (Supporting Information), the nucleation energy of Au on BuL and MLG is negative for all considered clusters. The obtained results...
clearly demonstrate that the metal clustering is preferred on the monolayer epitaxial graphene compared to the buffer layer, which is supported by the more negative values of $E_{\text{vac}}$. From the experimental point of view, such a difference will correlate with a difference in the nucleation rates. To be more specific, the existing Au clusters on MLG can capture easier newly incoming Au atoms to form larger clusters than those on BuL. To sum up, Au on BuL has much larger adsorption energy, higher energetic difference between minimum-energy and maximum-energy adsorption sites, and lower diffusivity than Au on MLG. As a direct consequence of that, Au atoms or small clusters on BuL prefer to stick to the most favorable adsorption sites, thereby resulting in the growth of a high density of Au islands. Furthermore, the restricted Au adsorption on top site positions (above the C atom that is covalently bonded to Si) of BuL prohibits, to some extent, the Au adatoms’ diffusion on the BuL, thereby also causing the formation of a large amount of nucleation centers followed by self-assembling of a discontinuous island-like film (see Figure 1a,b). In contrary, Au species adsorbed on MLG are mobile and have enough time for self-assembling in large clusters.

4. CONCLUSIONS

Combining experimental investigations and density functional theory calculations, we explored surface kinetics of magnetron-sputtered gold on carbon-rich buffer layer and monolayer epitaxial graphene on 4H-SiC. We demonstrated the selectivity of Au growth morphology on buffer layer with MLG patches. The difference in surface energetics of both graphenized surfaces is anticipated to play a major role in determining not only growth evolution of gold nanostructures but also in controlling the wettability and permeability of the topmost graphene layer. Gold on buffer layer exhibits dispersed nanoscale morphology of rounded islands, while a dendritic-shaped landscape appeared on MLG. Monolayer graphene is suggested to be nontransparent to the interaction between the underlying buffer layer and gold nanostructures; thus, the formation of large ramified and fractal-like Au islands on MLG/BuL/4H-SiC is mainly driven by weak van der Waals interaction between Au and graphene. These are prerequisites of easy self-organization of gold and prevention of defect generation. As was confirmed by the DFT results, the morphological distinctiveness of gold on both substrates is intrinsically linked to the adsorption capacity of the surface and substrate-influenced Au diffusivity. The unique corrugated topology of the buffer layer related to the mixed sp$^2$–sp$^3$ chemical bonding is identified to be decisive for the high Au adsorption and limited diffusivity. The current results shed light on the atomistic level interplay between gold clusters and graphenized surfaces of 4H-SiC and can be exploited to conceptualize novel applications in catalysis, sensors, and spintronics. We found that during the early nucleation stages, gold atoms (from 2 to 7) supported by nanostructured surfaces of SiC choose planar configuration rather than the three-dimensional one. Furthermore, we identified an odd-numbered Au$_n$ cluster as a minimal structural unit, acting as a growth nucleus. Raman results complemented by ab initio predictions provide reasoned arguments in favor of in situ gold intercalation beneath monolayer graphene, thereby opening a way for realization of, e.g., spatially confined 2D metal growth. Our finding also suggests that coexisting buffer layer–monolayer graphene regions providing unique adsorption capacity with respect to gold can be beneficial for designing

novel robust artificial materials with patterned nanoscale wettability and selective electronic performance. The strong affinity of transition metals on buffer layer can be used as a chemical sensor that can detect catalytic amount of organic and inorganic gases, such as benzene and NO$_x$. Furthermore, the observation of localized surface plasmon resonance in gold-decorated buffer layer could open the possibility for optical sensing of biomolecules. Further studies may provide a new platform for catalytic processes and biodetection.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsanm.0c02867.

- Raman data for bare buffer layer; Raman data for bare monolayer epitaxial graphene; reflectance map of bare monolayer epitaxial graphene; the 2 × 2 supercell of buffer layer and monolayer epitaxial graphene; the migration path; the 4 × 4 supercell of buffer layer and monolayer epitaxial graphene; molecular dynamics results; Raman data for Au-decorated buffer layer; intercalation of gold; Raman data for Au-decorated monolayer epitaxial graphene; optimized structures of 3D Au islands; optimized structures of 2D Au islands; adsorption of gold clusters; and parameters describing gold–template interaction (PDF)

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Author Contributions
I.S. performed all theoretical calculations and prepared the original draft of the paper. K.S. and N.P. were responsible for magnetron sputtering of Au, morphology characterization, and contribution to writing. I.G.I conducted reflectance mapping analysis and the Raman study. T.I. fabricated the bulk graphene/silicon carbide interface for monolithic wafer-scale electronics. Nat. Commun. 2012, 3, 957.

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