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To cite this version:
A. Curcella, R. Bernard, Y. Borensztein, M. Lazzeri, G. Prévot. The mechanism for the stabilization and surfactant properties of epitaxial silicene. Nanoscale, Royal Society of Chemistry, 2018, 10 (5), pp.2291-2300. 10.1039/C7NR06833F . hal-01701778

HAL Id: hal-01701778
https://hal.sorbonne-universite.fr/hal-01701778
Submitted on 6 Feb 2018

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Mechanism for the stabilization and surfactant properties of epitaxial silicene

A. Curcella, Y. Borenszttein, M. Lazzeri, G. Prévot

Using real-time in-situ scanning tunneling microscopy and density functional theory simulations, we have studied the growth of Si films on Ag(111) beyond the silicene monolayer, evidencing the existence of metastable phases and an original growth mechanism. Above monolayer Si coverage, a first \( \sqrt{3} \times \sqrt{3} \) structure forms, which is identified as an Ag-free Si bilayer with additional Si adatoms. With further deposition, this structure is replaced by a distinct bilayer structure covered by Si trimers and Ag atoms. The formation of these bilayers follows a counterintuitive dynamics: they are partially inserted within the Ag substrate and form by expelling, from the underlying substrate, atoms that reinsert below the adjacent silicene layer. The growth is therefore characterized by an unexpected "surfactant competition" between Ag and silicene: while silicene is a metastable surfactant for the Ag(111) surface, Ag plays the role of a surfactant for thicker diamond-like Si islands. In spite of being thermodynamically unfavoured, the silicene monolayer is, thus, a remarkably stable structure because of the high kinetic-barrier for the growth of thicker layers.

Introduction

Two-dimensional atomic layers, like graphene, silicene, germanene or phosphorene, metal dichalcogenides or trichalcogenides, or boron nitrides, show an increasing interest due to their specific and promising properties.\(^1,2\) Whereas graphene, phosphorene and chalcogenides can be obtained by exfoliation of the bulk material, this is not the case for silicene, since no graphite analogue exists for silicon. However, since the first theoretical predictions of the existence of metastable free-standing silicene,\(^3-5\) many efforts have been devoted to the synthesis of epitaxial silicene.\(^6-11\) Epitaxial Si monolayers have been first synthesized on Ag substrates: Si dimers have been observed on Ag(100),\(^12\) Si nanoribbons on Ag(110),\(^13\) with a pentagonal organization recently determined,\(^14,15\) whereas formation of a silicene layer was evidenced on Ag(111).\(^16\) On this surface, various ordered structures have been found to coexist, with a relative proportion depending on the preparation conditions.\(^17-19\) They all display a buckled hexagonal organization, with a buckling related to the matching between the Si and Ag lattices.\(^20\)

The silicene structures that form on Ag(111) are however metastable. During growth at temperatures higher than 540 K, the silicene layer dewets before its completion, leading to the formation of large three dimensional (3D) silicon islands separated by bare Ag regions.\(^21\) A similar result is observed after annealing, at 600 K, of a silicene layer grown at lower temperature.\(^21,22\) However, if growth is performed at temperatures lower than 540 K, a pseudo layer-by-layer growth mode (Frank van der Merwe)\(^23-27\) is observed, leading to the formation of thin silicon films terminated by a \( \sqrt{3} \times \sqrt{3} R30^\circ \) reconstruction. This reconstruction is observed after completion of the first monolayer, and corresponds thus to a thicker layer.\(^25,26,28\) On the basis of X ray diffraction and Raman spectroscopy, such films have been described as a new class of material, silicate, the equivalent of graphite for silicon atoms, also called multilayer silicene.\(^29-31\) Following these observations, various atomic models have been proposed, based on density functional theory (DFT) calculations, as alternatives to the classical diamond structure of Si, for bilayers or multilayer films.\(^32-40\) However, these results have been questioned by further independent experimental studies, concluding that the multilayer Si films structure is similar to the diamond-like structure of bulk Si, and that Ag atoms segregates at the surface during growth.\(^41-45\)

The exact termination of the surface remains also debated. Reconstructions involving only silicon atoms have been suggested on the basis of experimental and simulation results.\(^46-49\) However, the presence of Ag atoms on top of the surface has been confirmed by spectroscopic and microscopic measurements,\(^50-52\) proposing that the observed \( \sqrt{3} \times \sqrt{3} R30^\circ \) reconstruction is similar to the well known Ag/Si(111) reconstruction.\(^53,54\) Open questions remain on the nature of the bulk and surface structure of the silicon films that can be grown at moderate temperature (470 K-500 K). In particular, is it possible to grow metastable multilayer silicene structures? On the contrary, how does the surface evolve from a single silicene layer to a thick Si film covered with Ag? Most important, silicene, which is not stable with respect to bulk diamond, is commonly observed, in standard growth conditions, even in the presence of bulk nucleation centers. Why is silicene so unusually stable? Silicene/Ag(111) is a paradigmatic case of a 2D material which cannot be obtained by exfoliation from bulk material. Understanding these specific points is, thus, a step towards a more general goal: the synthesis of new phases, potentially thermodynamically unfavoured, keeping a weak interaction with the substrate.

To answer these questions, we have used real-time in-situ STM and DFT calculations to study the growth of Si films on Ag(111) in the 470 - 500 K temperature range. We have evidenced the existence of metastable phases and an original growth mechanism, which is schematized in Figure 1. In spite of being thermodynamically unfavoured, the silicene monolayer is a relatively stable structure with respect to atom addition. Above one monolayer Si coverage (Figure 1a), a first \( \sqrt{3} \times \sqrt{3} R30^\circ \) structure forms which is identified as an Ag-free bilayer with additional Si atoms (Figure 1b). With further Si deposition, this structure is replaced by a slightly thicker one, covered by Si trimers and Ag atoms (Figure 1c). The formation of these Si bilayers follows a counterintuitive dynamics: the bottom layer is located below the original silicene monolayer and forms by dislodging the atoms of the outermost Ag layer. These Ag atoms either wet the bilayer, or
intercalate between the substrate and the remaining silicene monolayer. The growth is thus associated with a high kinetic barrier which further enhances the stability of the silicene monolayer which behaves as a surfactant.

Results

The mechanism of growth of the silicene monolayer on Ag(111) has been recently described.55–57 At 500 K, Si atoms first insert in the Ag surface plane and act as nucleation centers for the growth of large silicene domains with mainly (4×4) and (√13×√13) reconstructions (these periodicities refer to the Ag surface, and correspond to (3×3) and (√7×√7) reconstructions, respectively, when referred to the pristine silicene plane). Ag atoms are expelled and form new terraces that are further covered with silicon. For coverage larger than nearly one monolayer (1 ML = 1.56×10¹⁵ at/cm²), the growth is characterized by the development of (4/√3×4/√3) reconstructed islands, commonly referred to as (√5×√5) (with respect to the pristine silicene plane). Such islands cover relatively smaller areas with respect to the silicene monolayer. Sone et al.26 have actually shown that two kinds of islands are observed: (√5×√5)−α and (√5×√5)−β, the latter ones progressively increasing at the expense of the former ones with further Si deposition.

STM measurements: (√3×√3)−α and (√3×√3)−β islands.

In Figure 2 (present measurements), the (√3×√3)−α and β islands are visible, surrounded by different phases of silicene (1 ML) with (4×4) (lower right) and (√13×√13) (upper left) reconstructions. The measured size of the unit cell is 0.66±0.03 nm. The α and β islands do not display the same surface appearance. First, their contrast is inverted: while the (√3×√3) triangular pattern of the α islands corresponds to outward protrusions, the β one corresponds to intrusions. Second, their corrugations are different, being ~0.06 nm for the α islands and ~0.02 nm for the β ones (Figure 2b). This corrugation difference suggests an hypothetically more metallic character of the β islands. Finally, contrary to previous observations,26 the apparent heights of the islands are markedly different: while the α islands are observed at 0.21 ± 0.01 nm above the surrounding silicene monolayer (Figure 2), the β ones are at 0.31 ± 0.01 nm. Thus, without making assumptions on the exact structure of the α and β islands, Figure 2 clearly indicates that their atomic and electronic structures are different.
Evolution of the islands with increasing Si deposition

Figure 3 displays the evolution of the same region during Si growth above the silicene monolayer at 500 K. The set of images correspond to detailed views of a 700 x 700 nm$^2$ area that is also periodically fully scanned with a lower resolution. The regions labeled S$_1$ (magenta), S (lavender), S$_1$ (seafoam/light-green), and S$_2$ (light-yellow) correspond to silicene monolayers lying on different Ag terraces. All these regions are staggered in height by multiples of 0.235 nm which is the height of a Ag(111) step. Within each of these regions, the monolayer displays domains with different reconstructions, mainly (4x4) and ($\sqrt{3}$$\times$$\sqrt{3}$). The darker dotted lines are defects present at the boundaries between the domains. After completion of the first Si layer, islands with ($\sqrt{3}$$\times$$\sqrt{3}$) - $\alpha$ reconstruction are growing. They correspond, for example, to the small green islands visible in Figure 3a (labeled "$\alpha$"). For larger amounts of silicon, they are progressively replaced by the ($\sqrt{3}$$\times$$\sqrt{3}$) - $\beta$ islands (green areas in Figure 3c, labeled "$\beta$", then brown and red areas in Figure 3c-e). This evolution can be seen more easily in the zoom reported in Figure 4.

In Figure 4, the different domains of silicene monolayer (lavender area) are clearly visible. Two islands at a higher level can also be seen. In Figure 4a, the bottom-left one partly consists of ($\sqrt{3}$$\times$$\sqrt{3}$) - $\alpha$ (turquoise), where the reconstruction is visible, and partly of ($\sqrt{3}$$\times$$\sqrt{3}$) - $\beta$ (green), which is not resolved as in Figure 2a because of the different tunneling conditions. For a larger amount of deposited Si (Figure 4b), the $\alpha$ domain of the lower island is completely replaced by the $\beta$ structure. Upon further Si evaporation, the $\beta$ domains continue growing laterally while the $\alpha$ ones progressively disappear, as it is visible in Figure 3b and c. Eventually, only silicene areas and $\beta$ islands are present. $\beta$ islands also grow vertically in a layer-by-layer way. In Figure 3, $\beta$ (green), $\gamma$ (brown), and $\delta$ (red) correspond to areas of increasing heights with the same apparent topographical contrast. The interlayer spacing, 0.30 $\pm$ 0.01 nm, is similar to what observed by others$^{25,42,52}$ and could correspond to the distance between two silicene layers as well as to the height of a (111) step of bulk Si.

Intercalation of Ag layers under the silicene single layer

In addition to this classical layer-by-layer growth of the $\beta$ islands, the evolution of the silicene regions displays a much unexpected behavior. Surprisingly, the comparison of Figures 3a-f shows that the height of the silicene monolayer increases during Si evaporation. For example, the region on the left of the cross is 0.235 nm higher on Figure 3e-f (S$_1$, seafoam) than in Figure 3a-d (S, lavender). Other examples are the S$_1$ region (magenta, on the left part of Figures 3a,b,c), which progressively disappears at the expense of the S one (lavender), and the S$_2$ region visible in the top-right position in Figures 3c,d, which laterally grows at the expense of the S$_1$ (seafoam) one.

In order to better explain these evolutions, Figure 5 reports STM scanning at a finer scale. It must be noticed that the amounts of deposited Si in Figure 5 do not exactly correspond to the ones in Figure 3, as Si was continuously deposited during the scanning of the STM. On the left side of Figure 5a, two parts of a $\beta$ island (coloured in green) are present. They are identified by their surface appearance and by their height ($h_{Si}=0.32 \pm 0.01$ nm) with respect to the silicene layer (S) as it is shown by the profile along the red line, Figure 5d. With increasing amount of deposited Si, several modifications can be observed in Figure 5b, then in Figure 5c: (i) the $\beta$ island is
spreading out on the surface and covers a larger part of the silicene layer (left part of the figures, in green); (ii) the majority of the silicene layer ($S_1$ in seafoam colour in Figure 5b,c) is now $0.235 \pm 0.01$ nm above the initial silicene layer ($S$ in lavender colour in Figure 5a), as shown by the height profiles along the black line; (iii) a new level of silicene layer is appearing at the upper right part of Figure 5c ($S_2$, yellow), $0.235 \pm 0.01$ nm above the $S_1$ silicene layer; (iv) a new flat island coloured in brown ($\gamma$) is developing upon the $\beta$ island coloured in green in Figure 5c, corresponding to a height increase $h_{Si} = 0.30 \pm 0.01$ nm.

The most striking feature is that the height increase of the silicene regions, when changing from $S$ to $S_1$ and from $S_1$ to $S_2$, coincides with $h_{Ag} = 0.235$ nm, which is the height of a single Ag(111) step. It is also very remarkable that the different domains and their orientations observed on the initial silicene layer ($S$ region, lavender in Figure 5a) are generally preserved in the corresponding higher silicene terrace ($S_1$, seafoam in Figure 5b). This is clearly seen in the bottom of Figure 5b where the growth front separates upper ($S_1$) and lower terraces ($S$) with the same (4x4) reconstruction. These
observations indicate that the growth (from Si to Ag) corresponds to the insertion of an additional Ag layer (located at \( h_{\text{Ag}} \) above the initial (111) Ag surface) below the silicene layer and not to a second silicene layer which would grow above the first one at \( h_{\text{Si}} = \frac{4}{3} h_{\text{Ag}} \). This is the signature of an unexpected intercalation of an Ag layer below the silicene layer, which, in turn, preserves its structure during the process.

**DFT Calculations: structures and simulation of STM topography**

The structure and energetics of various Si/Ag multilayers were calculated within DFT. Shown results are obtained with the generalized gradient approximation (GGA),\(^{38}\) which gives the best agreement with X ray diffraction measurements for the structure of \((4 \times 4)\) silicene.\(^{20}\) Results were tested against the inclusion of dispersive corrections as in refs. 59-60. Unless otherwise stated, we consider silicon-based super-cells with \((3 \times 3)\) periodicity lying on a \((4 \times 4)\) Ag(111) surface super-cell. In order to determine the nature of the \((\sqrt{3} \times \sqrt{3})\) and \((3 \times 3)\) – \(\beta\) reconstructions, we tested various possibilities. A first set corresponds to pure silicon structures lying on Ag(111), labelled “BL-AB”, “ML+Si”, “BL+Si”, and “BL-AA”. The “BL-AB” structure is a silicon bilayer with AB stacking; “ML+Si” corresponds to a Si monolayer in which three Si atoms are added to form the trigonal dumbbell (TDS) structure proposed in ref. 34; “BL+Si” corresponds to an AB bilayer in which three Si atoms are added, in analogy to the TDS one. In both the ML+Si and BL+Si structures the outermost three Si atoms are added on the top of three Si atoms inward buckled forming a \((\sqrt{3} \times \sqrt{3})\) periodicity. This configuration is inspired by the very stable \((7 \times 7)\) reconstruction of the Si(111) surface. “BL-AA” is a silicon bilayer with stacking AA as proposed by Pfugradt et al.\(^{38}\) (the BL-AA structure has a true periodicity of \((2 \sqrt{3} \times 2 \sqrt{3})\), and lies on a \((\sqrt{12} \times \sqrt{12})\) Ag super-cell). A second set of structures are silicon layers covered by Si and Ag atoms arranged in analogy to the reconstruction observed when the Si(111) surface is covered by Ag: \(\text{Ag/Si}(111)\) \((\sqrt{3} \times \sqrt{3})\). In this configuration, a silicon monolayer covered by half a silicon layer and nine Ag atoms per unit cell will be labeled as “ML+Si/Ag”. “BL+Si/Ag” labels an analogous structure obtained from a Si AB bilayer. We recall that the Ag/Si(111) \((\sqrt{3} \times \sqrt{3})\) reconstruction is observed in two different configurations which pose, or not, a mirror symmetry. An inequivalent triangle structure (IET, non-symmetric) is observed at temperatures smaller than 150 K, while a honeycomb chained triangle (HCT, symmetric) appears at \(T > 150\) K.\(^{61}\) In the DFT calculations, starting from an appropriate initial guess, it is possible to obtain symmetric and non-symmetric structures for the Ag/Si(111) \((\sqrt{3} \times \sqrt{3})\) and for both the ML+Si/Ag and BL+Si/Ag structures. In the following, we will only consider the symmetric ones, assuming that they are stabilized by temperature. Details of the simulated structures are given in the ESI.

Figure 6 shows the simulated STM images for some of these structures. These images were used to estimate the apparent height \(h\), as seen by an STM probe, of the different structures with respect to the \((4 \times 4)\) silicon monolayer, and their corrugation \(\Delta h\). \(h\) is the average height probed by an ideal spherical tip with radius 1 nm touching the constant density surface. \(\Delta h\) is the difference between the maximum and the minimum height. These values are given in Table 1. Although these numbers cannot be considered as an absolute determination of the corresponding measured quantities, the relative heights of the different theoretical structures should be more comparable with measurements.

**Energetics**

To confront the stabilities, we consider the grand canonical surface energy:

\[
\gamma(\mu_{\text{Si}}, \mu_{\text{Ag}}) = \frac{1}{A} \left[ E_{\text{Slab}}(N_{\text{Si}}, N_{\text{Ag}}) - \mu_{\text{Si}} N_{\text{Si}} - \mu_{\text{Ag}} N_{\text{Ag}} \right] \tag{1}
\]

where \(E_{\text{Slab}}\) is the energy of a slab containing \(N_{\text{Si}}\) Si atoms and \(N_{\text{Ag}}\) Ag atoms. The bare Ag surface on the bottom of the slab was kept fixed during relaxation. \(A\) is the surface area of the slab, \(\mu_{\text{Si}}\) and \(\mu_{\text{Ag}}\) are the chemical potentials of Si and Ag, \(E/\Delta\) is the surface energy of the bottom bare Ag surface, which is calculated independently. Figure 7 compares the surface energies of the different structures calculated for \(\mu_{\text{Ag}} = E^{\text{bulk}}_{\text{Ag}}\), the Ag cohesive energy in the fcc Ag lattice. The energies are plotted as a function of \(\Delta\mu_{\text{Si}} = \mu_{\text{Si}} - E^{\text{bulk}}_{\text{Si}}\), where \(E^{\text{bulk}}_{\text{Si}}\) is the Si cohesive energy in the diamond Si lattice.

<table>
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<th>(h) (nm)</th>
<th>(\Delta h) (nm)</th>
<th>(\Delta h) (nm)</th>
<th>(\alpha)</th>
<th>(\beta)</th>
<th>(\mu_{\text{Ag}})</th>
<th>(\mu_{\text{Si}})</th>
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<td>0.09</td>
<td>0.02</td>
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Table 1: Characteristics of the simulated STM images of the structures described in the text. \(h\) is the apparent height with respect to the silicene monolayer and \(\Delta h\) is the corrugation (see the text). The quantities in boldface are from measurements; \(\alpha\) and \(\beta\) refer to the two \((\sqrt{3} \times \sqrt{3})\) phases; ML to the monolayer \((4 \times 4)\) reconstruction. For the structures containing a Si bilayer, \(h\) has been reduced by 0.235 nm (the Ag-Ag interlayer distance) assuming that the atoms of the outermost Ag layer are expelled during the growth (this choice is clarified in the discussion). The values in parenthesis are obtained without applying this correction.
To further elucidate the growth dynamics, we also calculated the adsorption energy of one single Si atom on various sites on the silicene/Ag(111) structure. Results are reported in the Electronic Supplementary Information.

**Discussion**

The present findings allow a clear attribution of the \((\sqrt{3} \times \sqrt{3}) - \alpha\) and \((\sqrt{3} \times \sqrt{3}) - \beta\) phases. There are already several experimental evidences that the \((\sqrt{3} \times \sqrt{3}) - \beta\) phase consists of both Ag and Si atoms lying on a silicon layer in a structure analogous to the \((\sqrt{3} \times \sqrt{3})\) reconstruction observed on Ag/Si(111).\(^{34,45,50-52}\) This corresponds to the present ML+Si/Ag and BL+Si/Ag structures and, indeed, the corresponding simulated STM images (Figure 6) display intrusions with \((\sqrt{3} \times \sqrt{3})\) periodicity as those experimentally observed for the \((\sqrt{3} \times \sqrt{3}) - \beta\) (Figure 2). Note that the simulated STM images of the ML+Si/Ag and BL+Si/Ag structures are quite different from those of the \((\sqrt{3} \times \sqrt{3})\) Ag/Si(111) reported in literature,\(^{62}\) despite the fact that the atomic structures of the three systems are very similar. This issue is discussed in the Supporting Information.

Concerning the \(\alpha\) phase, the present STM measurements reveal that it is a structure clearly distinct from the \(\beta\) one (Figure 2). Assuming thus that the \((\sqrt{3} \times \sqrt{3}) - \alpha\) is an Ag-free structure, we can compare it to the BL-AB, BL-AA, ML+Si, BL+Si structures. The BL-AB and BL-AA ones are unlikely to correspond to the \(\alpha\) phase: according to our calculation (which confirms what already remarked in ref. 38) the BL-AB structure does not reconstruct with the \((\sqrt{3} \times \sqrt{3})\) periodicity of the \(\alpha\) phase, and the BL-AA structure (which actually presents the \((\sqrt{3} \times \sqrt{3})\) reconstruction) has a very high energy, compared with the one of the bilayer (BL-AB, Figure 7), probably because of the AA stacking. On the other hand, both ML+Si and BL+Si are good candidates for the \(\alpha\) phase: they are significantly more stable than the BL-AB structure (Figure 7) and reconstruct with \((\sqrt{3} \times \sqrt{3})\) periodicity, displaying STM images which nicely resemble those of the \(\alpha\) phase (Figure 6).

In the previous two paragraphs, we concluded that the \(\alpha\) phase corresponds to either ML+Si or BL+Si structures, and that the \(\beta\) phase corresponds to either ML+Si/Ag or BL+Si/Ag.
ones. We now consider the energetics reported in Figure 7. Keeping in mind that we are not at thermodynamical equilibrium, we argue that a reasonable $\Delta \mu_{Si}$ range to describe the present experimental conditions is $0 < \Delta \mu_{Si} < 0.13$ eV. Indeed, we expect silicon to be metastable with respect to the formation of silicon bulk since, for deposition at 600 K, the silicon layer dewets to form large Si bulk islands. $\Delta \mu_{Si}=0.13$ eV corresponds to the point where the surface becomes fully covered with silicon. When the $\alpha$ islands start to grow, the monolayer is nearly (but not entirely) complete and the chemical potential must thus be close to the value $\Delta \mu_{Si}=0.13$ eV.

We remark that, in a large range around $\Delta \mu_{Si}=0.13$ eV, the ML+Si and ML+Si/Ag structures are less stable than the monolayer (ML). On the other hand, the BL+Si and BL+Si/Ag ones are more stable than the clean silicon bilayer (BL-AB). As a consequence, the ML+Si and ML+Si/Ag terminations are not expected to form in the presence of the monolayer. On the contrary, the BL+Si and BL+Si/Ag ones should form when the second silicone layer starts to grow. The fact that the ML+Si and ML+Si/Ag terminations are less stable than the monolayer, while BL+Si and BL+Si/Ag are more stable than the bilayer is not entirely surprising. Indeed, while silicon (ML) is a relatively stable configuration with an hybridation different from that of diamond, the bilayer (BL) is more similar to diamond and needs to be stabilized by non-ideal reconstructions as it is the case for bulk-silicon surfaces. As a consequence, the $\alpha$ and $\beta$ phases are attributed to the BL+Si and BL+Si/Ag structures, respectively.

One of the most interesting observation, among the present ones, is that the formation of the $(\sqrt{3} \times \sqrt{3}) - \beta$ phase is accompanied by an increase in height of the adjacent silicone regions, which testifies the intercalation of Ag atoms under these monolayers (Figures 3-5). The emergence of these additional Ag atoms is, at first sight, very puzzling and raises the question of their origin. Knowing that the $\beta$ phase is a bilayer (BL+Si/Ag), the most natural answer is that the bottom Si-layer of the bilayer is located below the original silicone monolayer and forms by dislodging the atoms of the outermost Ag layer (Figure 1). The expelled Ag atoms partially intercalate under neighboring silicone regions, increasing the height of the terraces, and partially go on the top of silicon (to form the $(\sqrt{3} \times \sqrt{3}) - \beta$ reconstruction). We argue that this dynamics has already started during the formation of the $(\sqrt{3} \times \sqrt{3}) - \alpha$ phase. This is not evident on the sole basis of Figures 3-5. However, we have already concluded that the $\alpha$ phase is a bilayer (BL+Si). If we also consider that the measured STM height of the $\alpha$ phase is smaller than that of the $\beta$ one (0.21 vs. 0.30 nm) we can deduce that the BL+Si structure has also formed by expelling the outermost Ag layer (otherwise the height would be much higher). Note also that the attribution of $(\sqrt{3} \times \sqrt{3}) - \alpha$ and $(\sqrt{3} \times \sqrt{3}) - \beta$ to bilayers (BL+Si and BL+Si/Ag) that are inserted below the outermost Ag layer is compatible with the calculated STM heights reported in Table I.

The presently described dynamics is compatible with the energetics for the adsorption of one Si atom on silicone/Ag(111). According to the calculations described in the Supporting Information, the substitution of a Si atom with one Ag atom of the outermost Ag layer (which is below the silicone) is energetically favoured with respect to the adsorption of one Si atom on the top of silicone. This dynamics contrasts with a more intuitive and classical picture in which a Si atom would stick on the top of the surface and act as a nucleation centre for the growth of the second layer. Exchange between Si and Ag atoms already occur in the initial stage of growth: the insertion of a Si atom within the clean Ag(111) surface is a relatively fast process associated with an experimentally-determined small energy barrier of 0.43 eV.66 However, it is safe to assume that the insertion of a Si atom below an existing silicone layer should be associated with a high activation barrier. Thus, if we also assume fast diffusion of the Si atoms adsorbed on the silicone layer, the adsorbed Si atoms should be allowed to reach a particularly favoured location (possibly a step or a defect) to insert below the silicone layer. This picture could explain why the bilayer starts growing only when the monolayer has completely covered the Ag surface, a fact observed experimentally.

Indications that the Si adatom diffusion is fast are given by DFT calculations on free-standing silicone,65 which provide an energy barrier of 0.75 eV for the diffusion of a dumbbell unit created by addition of a Si adatom. This should result in a high jump rate at $T = 500$ K, of the order of $10^7$ s$^{-1}$, considering a usual attempt frequency of $10^{13}$ s$^{-1}$. Note also that the reorganization of the silicone layer during growth (which involves Si-Si bond breaking) has been predicted64 and experimentally observed.66 Thus, energy barriers for Si diffusion and Si-Si bond reorganization are low enough to enable 2D growth at 500 K.

Concerning the Ag kinetics, the expelled Ag atoms are expected to easily wet Si islands due to their fast diffusion. Indeed, an activation barrier of 0.25 eV has been measured for Ag atoms on the Si(111) (7×7) surface.67 The diffusion of Ag atoms on the Ag induced $(\sqrt{3} \times \sqrt{3})$ reconstruction of Si is also fast, with a barrier of 0.33 eV.66 Lower barriers should be obtained for Ag atomic jumps on the silicone surface, in analogy with graphene. Finally, further growth of the Si islands proceeds by exchange between incoming Si atoms and surface Ag atoms forming the $(\sqrt{3} \times \sqrt{3})$ reconstruction. Such effect has been already evidenced during Si homoepitaxy on Si(111)-$(\sqrt{3} \times \sqrt{3})$ Ag at 570 K.67 Silicon diffusion through the Ag layer has also been evidenced during Co deposition on Si(111)-$(\sqrt{3} \times \sqrt{3})$ Ag where cobalt silicide formation on the $(\sqrt{3} \times \sqrt{3})$ reconstruction is observed above 400 K.68
Methods

Sample preparation and Si evaporation

All experiments were performed in set-ups working under ultra-high vacuum (base pressure 10^{-10} mbar). The Ag(111) sample was prepared by repeated cycles of Ar` sputtering and annealing at 800 K. Si was evaporated from a thermally heated crucible using a commercial Omicron Nanotechnology e-beam evaporator. The evaporation rate was around 0.004 ML/min.

STM experiments

STM experiments were carried out using an Omicron Nanotechnology VT STM. STM images were directly acquired during Si evaporation. In order to avoid a continuous shadowing of the surface by the STM tip, various areas were scanned successively. Temperature measurement is performed with a thermocouple located on the sample heating stage, previously calibrated with another thermocouple soldered on the surface of a testing sample. The uncertainty on the absolute value of the temperature is around 10 to 20 K at 500 K.

DFT calculations

We used the Quantum ESPRESSO package, and the projector augmented plane wave method of ref. 70. The energy cut-off of plane waves was 30 Ry and that of the charge 240 Ry. The Ag (111) surface is described by a four layers slab. Atomic positions were always relaxed by energy minimization (only the Ag-slab bottom-layer positions were fixed) fixing the in-plane waves to the theoretical Ag bulk one (0.2935 nm). The Brillouin zone of the Ag-slab bottom-layer positions was fixed the local density of states at \( \alpha \) above the Fermi energy and, then, the height of the surface at constant density. Results were systematically tested against the inclusion of dipole corrections.\(^7\)

Conclusions

To conclude, we have studied silicene growth dynamics (beyond the monolayer) on the Ag(111) surface by means of real-time in-situ STM measurements and DFT calculations, providing a consistent scenario explaining all the experimental and numerical observations. The silicene monolayer is a relatively stable structure with respect to atom addition, always maintaining its honeycomb structure. The \((\sqrt{3}\times\sqrt{3})-\alpha\) and \((\sqrt{3}\times\sqrt{3})-\beta\) reconstructions observed experimentally are non-ideal terminations of silicon bilayers. The bilayers form by expelling the underneath atoms of the outermost Ag layer. An ideal bilayer, i.e. a stacking of two silicene planes, is never observed and the thicker layers are stabilized in two alternative ways. In the early stages of the growth, a Ag-free structure (\( \alpha \) phase) bearing analogies with that of the \((7\times7)\) reconstruction of the bulk Si(111) surface can be observed. After this initial stage, the bilayer is stabilized in a structure covered by Si trimers and Ag atoms (\( \beta \) phase), analogous to the structure of the Ag/Si(111) \((\sqrt{3}\times\sqrt{3})\) reconstruction. The next stages of the growth follow a standard layer-by-layer growth, in which the outermost Si-Ag structure does not change. During this growth, the silicon interlayer distance is very similar to that of the diamond structure, making hard to probe the existence of a “multilayer silicene” structure bearing distinct properties from those of a diamond with an important stacking disorder. The present scenario indicates a surfactant competition between Si and Ag, driven by the high kinetic barrier to transform a silicene monolayer into thick Si islands. While silicene is a metastable surfactant for the Ag(111) surface, Ag plays the role of a surfactant for diamond-like Si islands. If we now consider the experimental results discussed in the introduction: for temperatures below 540 K, the atomic mobility is low enough to inhibit the complete dewetting of the silicene layer. Above 540 K, Si thick islands act as nucleation centers for the dewetting of silicene, leading to thick 3D islands separated by bare Ag regions.\(^21\)

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by French state funds managed by the ANR within the Investissements d’Avenir programme under reference ANR-11-IDEX-0004-02, and more specifically within the framework of the Cluster of Excellence MATISSE led by Sorbonne Universités.

Notes and references


