Understanding the growth mechanism of graphene on Ge/Si(001) surfaces

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The practical difficulties to use graphene in microelectronics and optoelectronics is that the available methods to grow graphene are not easily integrated in the mainstream technologies. A growth method that could overcome at least some of these problems is chemical vapour deposition (CVD) of graphene directly on semiconducting (Si or Ge) substrates. Here we report on the comparison of the CVD and molecular beam epitaxy (MBE) growth of graphene on the technologically relevant Ge(001)/Si(001) substrate from ethene (C2H4) precursor and describe the physical properties of the films as well as we discuss the surface reaction and diffusion processes that may be responsible for the observed behavior. Using nano angle resolved photoemission (nanoARPES) complemented by transport studies and Raman spectroscopy as well as density functional theory (DFT) calculations, we report the direct observation of massless Dirac particles in monolayer graphene, providing a comprehensive mapping of their low-hole doped Dirac electron bands. The micrometric graphene flakes are oriented along two predominant directions rotated by 30° with respect to each other. The growth mode is attributed to the mechanism when small graphene “molecules” nucleate on the Ge(001) surface and it is found that hydrogen plays a significant role in this process.

Graphene is widely supposed to be a material on the way to enable new development of many modern technologies1,2. This includes microelectronics and optoelectronics, which require graphene of the highest quality, inexpensively produced and processed on large areas3–5. Such graphene can be grown on Cu or on Ni3,6–8, but then it must be transferred to the target wafer. The transfer process is problematic; one of the reasons is contamination with residual metal atoms9,10. The other approach is the direct growth of graphene on the insulating target, like hexagonal BN (h-BN)11–14, which can then be used for the immediate device fabrication. Therefore, researchers examine the prospects of growing graphene on microelectronics-friendly substrates, such as Si or Ge15–25. Direct growth of graphene on Si is hindered by high reactivity of Si against C16. In contrast to Si, Ge does not form stable carbide. However, much higher growth temperatures are needed for CVD graphene on Ge than on insulating h-BN, indicating that the formation of C-Ge bonds plays a major role in the former process21,26.

Graphene films grown on Ge may be used in two ways. First, if large area graphene (like 300–450 mm in diameter) is obtained, the film may be transferred to the target wafer at any point of the fabrication process (i.e., during front end of line or back end of line steps) without the hazard that the production tools become contaminated. In practice this means that the growth must be performed on Ge(001)/Si(001) wafers, because Ge wafers of this size are not commercially available and it would most probably be prohibitively expensive to develop and introduce a Ge wafer growth technology solely for this purpose. The second way to use graphene films grown on Ge may be open when graphene can be grown directly on device-size Ge(001) islands pre-deposited on the Si(001) substrate. For example, the width of the active region of a graphene base transistor may be smaller than a micron, and its whole area may be of the order of μm2 27,28.

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Graphene growth on the technologically relevant Ge(001) and Ge(001)/Si(001) wafers has been reported for a CH₄-based CVD and for a molecular beam epitaxy (MBE) (atomic carbon), respectively. In both cases, the structural and electrical quality of graphene significantly improves at temperatures so high (900–930 °C) that the Ge substrate begins to melt. Besides causing practical problems associated with strong surface roughening, that takes place at such temperatures, this again highlights the importance of the C-Ge interaction. In the present manuscript, we report on the observation that illustrates the influence of this interaction on the growth process – the appearance of two orientational domains. These two domains are revealed by the presence of two sets of Dirac cones in the electronic structure of the graphene film as deduced from the nanoARPES data. The interaction of the C₂H₄ precursor with the Ge(001) surface is then analyzed in the framework of ab initio density functional theory (DFT) calculations and conclusions for the microscopic mechanisms of the growth process are drawn. These mechanisms are then compared to those expected for the growth of graphene on Ge(001) from atomic carbon.

**Results**

**Experiment.** Morphology of the CVD- and MBE-grown graphene on Ge/Si(001) was studied by means of scanning electron microscopy (SEM) (Fig. 1). The graphene flakes in the CVD and MBE prepared films appear similar in SEM, at least in the growth stage depicted in the figure, but one interesting difference between them is that the MBE prepared flakes have long straight edges, while the CVD synthesized flakes are of much more rounded shape. The flakes are separated by a material that is far less conducting. In the SEM images these areas appear as bright spots, and the ARPES spatial maps show that the density of states at the K-point around the Fermi surface forms a circle centred at \( \Gamma \) and similar to the one observed for highly oriented pyrolytic graphite (HOPG).

Figure 1. Distribution of graphene flakes in the CVD and MBE synthesised samples on Ge(001). Conductive graphene is dark in SEM images for (a) CVD and (b) MBE grown graphene.

The electronic structure in the vicinity of \( E_F \) the Fermi level \( (E_F) \) of the CVD-produced graphene flakes on Ge(001)/Si(001) substrate was studied in the micro and nanoARPES experiments with 120 μm and 100 nm spot size resolution, respectively.28–31 As it is well-known for a free-standing graphene, the charge neutrality point (Dirac point, \( E_D \)) coincides with \( E_F \). In the vicinity of \( E_D \), the energy of graphene electronic states depends linearly on the wave-vector \( k \) (Fig. 2a). These energy bands form the so-called Dirac cones at the six equivalent \( K \) points in the corners of the hexagonal Brillouin zone (BZ) of graphene (Fig. 2b). In case of the n- or p-doped graphene, the Fermi level is located either above or below the Dirac point, respectively. In-plane long-range disorder in the graphene lattice blurs the energy states imaged by taking the constant energy cuts in the 3D ARPES data, \( I(\mathbf{E}_D, k_x, k_y) \). Here, \( \mathbf{E}_D \) is the binding energy and \( k_x, k_y \) are two in-plane components of the wave vector of electron. When a neutral graphene has two domains rotated with respect to each other by 30° and the same area of each of them is illuminated by the photon beam, the Fermi surface in the ARPES experiment consists of 12 equivalent spots centred at the 12 equivalent \( K \) points around the BZ centre, \( \Gamma \). When a neutral graphene is fully disordered or polycrystalline, the Fermi surface forms a circle centred at \( \Gamma \) and similar to the one observed for highly oriented pyrolytic graphite (HOPG).

Figure 2c shows a constant energy cut taken at the Fermi level and extracted from the 3D photoemission intensity data set, \( I(E, \theta, \phi) \) measured for graphene/Ge(001)/Si(001) at \( h\nu = 100 \text{ eV} \) (spot size 120 μm) as a function of two emission angles, polar \( \theta \) (detector angle) and azimuth \( \phi \) (in-plane sample rotation), and then converted into \( I(\mathbf{E}_D, k_x, k_y) \). One can clearly see that the photoemission at the Fermi level has a circle-like shape centred at \( \Gamma \) and having equidistant intensity maxima. The radius of the circle in the reciprocal space is 1.7 Å⁻¹ and the angular distance between the maxima is 30°. This indicates that the observed photoemission intensity originates from graphene flakes with orientation distribution that has probability maxima in the directions corresponding to graphene hexagons being parallel and/or perpendicular to the surface dimer rows of the Ge(001) surface (see discussion below).

A detailed map of photoemission intensity measured with circularly polarized light of \( h\nu = 30 \text{ eV} \) along the \( \Gamma - K \) direction for one of the preferential domains is given in Fig. 2d. Due to the fact that circularly polarized light is used, the photoemission bands along both \( K - \Gamma \) and \( K - M \) directions are detected. One can also see that the photoemission intensity between two branches does not vanish; this is due to the partially random orientation of graphene flakes. However, a clear Dirac cone is visible; this is valid for all intensity spots in Fig. 2c.

These data allow us to estimate the energy distance between the Fermi level and the Dirac point, and hence the doping level of graphene: \( E_F \) is located by 0.185 eV below \( E_D \) which corresponds to the hole concentration
of $2.3 \times 10^{12} \text{ cm}^{-2}$. The linear fit of the Dirac cone in the ARPES intensity map in Fig. 2d gives the Fermi velocity of $(1.36 \pm 0.3) \times 10^6 \text{ m/s}$. This can be compared with the graphene/Ge/SiC(0001) system, where $p$-doping of graphene was obtained via intercalation of Ge atoms in graphene/SiC at $720 \text{ °C}$ and the hole doping of $4.1 \times 10^{12} \text{ cm}^{-2}$ was measured\cite{32}. The measured radius of the Fermi surface around K is $0.027 \text{ Å}^{-1}$.

Graphene flakes distribution was probed in the real-space-resolved nanoARPES experiments. Figure 3a shows the respective photoelectron spectroscopy (PES) survey spectrum for our sample before ARPES mapping where C 1s and Ge 3d core levels are marked. The photoemission intensity at the K-point for one of the preferential graphene domains and in the vicinity of the Fermi level (the energy and wave-vector windows are marked as a rectangular in inset of Fig. 3a and in Fig. 3b) was acquired as a function of space coordinate (see Fig. 3c). The photoemission intensity map indicates that the graphene film consists of flakes with size in the $\mu\text{m}$ range, which is also consistent with the SEM measurements.

Results of a electrical measurements in the field-effect transistor (FET) configuration in 4-point STM (Fig. 4a) are consistent with the observation of $p$-type doping detected in nanoARPES experiments (cf. Fig. 2d): the Fermi level crosses the Dirac point (and the graphene channel conductance reaches its minimum) when the gate is positively biased. This direction of the shift corresponds to $p$-doped channel. The same sign of doping was detected for CVD as well as for MBE prepared samples (Fig. 4b,c). The gate bias $V_{FB}$ at which the channel conductance reaches its minimum is a measure of the Fermi level position (with respect to the Dirac point), which in turn is determined by the doping level; therefore $V_{FB}$ measures the doping level. Comparison of the $V_{FB}$ value for CVD and MBE prepared films indicates that former films are more strongly $p$-doped. The measurements in the FET configuration allow to estimate the carrier mobility of the graphene channel. For this purpose, one assumes that the film...
The area controlled by the gate tip corresponds to a square and one measures the sample-tip distance \( d \) by approaching the tip towards the sample until mechanical contact is observed. The measured carrier mobilities were lower than for high-quality graphene and amounted to \( 1000 \pm 500 \text{ cm}^2/\text{V} \cdot \text{s} \) at room temperature for CVD and MBE prepared graphene/Ge(001). Figure S1 in the Supplementary materials reveals the fit according to ref. 33.

For epitaxial graphene on SiC(0001) with 2 monolayer of Ge intercalated hole mobilities up to \( 1800 \text{ cm}^2/\text{V} \cdot \text{s} \) are reported34. This shows that rather the defect density in CVD- and MBE-grown graphene in our case is significantly higher.

Figure 5 summarizes the results of our Raman analysis of the MBE- and CVD-grown graphene layers on Ge(001) described above (see Fig. S2 in the Supplementary materials for the comparison of the Raman spectra of graphene grown by different methods). In both cases, the phonon frequencies of the 2D and G modes differ from their reference values measured on undoped and unstrained graphene films. Following the analysis of Lee et al.35, we can disentangle the effects of strain and doping in our graphene films on Ge(001) (see Fig. 5b). Since the 2D-mode position for the MBE-grown graphene is upshifted compared to the reference value, the strain in the MBE-grown graphene is compressive, assuming that the strain is biaxial; the average strain can be estimated to approximately \( -0.3\% \). In contrast, the strain in the CVD-grown graphene films is tensile and can be evaluated to about \( 0.4\% \). If we would instead assume uniaxial strain for both samples, the strain would amount to \( -0.8\% \) and \( 1.4\% \) for the MBE- and CVD-grown graphene, respectively. However, such large uniaxial strain would lead to a splitting of the double-resonant 2D mode by more than 20 cm\(^{-1}\)36. This is in contrast to our experimental data, where a symmetric, Lorentzian-shaped 2D mode is observed during all measurements. Thus, strain in our samples is likely to be biaxial.

In accordance with the electrical 4-point STM measurements, the doping level of the CVD graphene film is higher than that of the MBE-grown film as deduced from the Raman spectra measurements. A rough estimate yields the average doping below \( 1 \times 10^{12} \text{ cm}^{-2} \) for the MBE graphene and around \( 3 \times 10^{13} \text{ cm}^{-2} \) for the CVD sample. The presence of data points with G-mode positions on the left side of the zero-doping line can be partially explained with the anomalous phonon softening of the G mode for Fermi-level shifts close to the Dirac point35,37,38. Additionally, we have to consider other effects than strain and doping to affect the 2D- and G-mode positions. In particular, the background subtraction, which is necessary due to strong luminescence from the substrate, introduces an additional uncertainty to the strain/doping analysis that is in the order of 0.2% and \( 5 \times 10^{12} \text{ cm}^{-2} \).
Theory. Although carbon and Ge form no stable carbide phase (GeC), the chemical interaction between the Ge(001) surface and C atoms is strong. This is particularly pronounced when atomic carbon is deposited. In this case, there is a significant probability that a mobile C atom becomes immobilized by substituting a Ge atom in a Ge surface dimer (Ge₂), whereby a mobile Ge adatom is released. The process is reversible: a Ge adatom may eject the C atom from a C-Ge surface dimer. The mobile and highly reactive C interstitially produced in this process can diffuse under the surface. The mobile species controlling the MBE growth mode of graphene on Ge(001) are C atoms, short chains consisting of few C atoms, and Ge adatoms; the latter are attracted to the edge of growing graphene. Carbon dimers (C₂) are trapped by Ge dimer vacancies (DV) to become graphene seeds; a similar role is attributed to longer C chains (like C₈) trapped between dimer rows. In contrast to that, results of our calculations suggest that nucleation of CVD graphene on a flat Ge(001) surface takes place predominantly on the DV defects (see Supplementary Materials for details and further discussion).

The vast majority of C₂H₄ precursor molecules do not give all H atoms away to the substrate. Calculations indicate that the major unpolymerized diffusing species may be, apart from adsorbed H, C₂H₃ molecules, with small admixture of C₂H and C₂ (see Supplementary Materials for details and further discussion). This is because most of C₂H₂ (acetylene) produced during C₂H₄ decomposition desorb before they decompose further. The surface reactions do not lead to significant production of Ge adatoms. Moreover, although there are Ge adatoms on the surface (due to high temperatures used in these experiments), they are repelled from the edge of a hydrogen-terminated graphene. This is in strong contrast to the MBE process: adsorption of Ge adatoms at edge sites of hydrogen-free graphene is energetically preferred to adsorption at surface steps.

The C atoms on a clean graphene edge tend to form chemical bonds with the substrate (Fig. 6a), immobilizing the molecules consisting of several carbon rings. On the other hand, when the molecule edge is completely saturated with hydrogen (what our calculations predict to happen during our ethylene CVD synthesis), the graphene-substrate bond formation is suppressed, apart from the possibility that some of the edge C atoms may become attached to the substrate by a weak sp³-like bond (Fig. 6b). These bonds are however not strong enough to prevent the molecule from diffusing on the surface during the growth process, so that the orientation transfer from the substrate to CVD graphene may occur only during nucleation on immobile seeds, like on the DV defects.

Discussion. Formation mechanism of two domains in graphene growth. In the MBE process, graphene may nucleate on C dimers, (C₂)ₓDV, trapped inside Ge dimer vacancy defects (Fig. 7a). DV defects are abundant in STM
images of Ge(001) even at room-temperature. At 930 °C, C2 dimers reside about 50 times longer in the trapped
than in the diffusing state. [Assuming thermal equilibrium DV concentration at 900 °C, the DV formation energy
of 0.6 eV , and the same attempt rates for C2 diffusion and C2 escape from DV . In order to escape from the trap,
a C2 must overcome a barrier of 2.3 eV , higher by 1.0 eV than the barrier for C2 diffusion on the perfect
surface. The total energy of C2 is by 1.0 eV lower in a dimer vacancy at the most favorable adsorption site on the perfect
surface.] (C2)DV is produced also by atomic C in two kick-out events23. [C(diff)
→
CGe
+ Ge(bulk)
− 0.13 eV , and
C(diff)
+ CGe
→ (C2)DV + Ge(bulk) + 0.31 eV , where C(diff) is the mobile atomic carbon diffusing under the sur-
face and Ge(bulk) is a Ge atom having the same energy as in the Ge bulk. The total energy gain of 0.2 eV is in this
case much smaller than for trapping on a DV (1.0 eV), because a new DV site must be produced, by subsequent
ejection of two Ge atoms. The difference between these two energies (0.8 eV) should thus be equal to the DV
formation energy (0.6 eV). The discrepancy (0.2 eV) comes from inaccuracies in the calculation of atomic relax-
ations, caused by the presence of nearly degenerate metastable configurations of defects and by finite size effects.
This inaccuracy amounts to at least 0.1 eV per defect.] Dimer vacancies collect carbon; e. g., when the second C2
is trapped by a DV already occupied by C2, the energy decreases by 0.9 eV, which is similar to the energy gained
by trapping of the first C2. Trapping of carbon by dimer vacancies provides a means to align the orientation of
graphene with that of the substrate (Fig. 7c,d). Variations in the orientation of MBE graphene may be enhanced

Figure 5. Raman characterization of graphene films. (a) Collection of 130 spectra from a CVD film after
background subtraction; the spectrum closest to the average is shown in black. Energies of D, G, and 2D
modes in an unstrained and undoped graphene are indicated. (b) 2D-position vs. G-position plot illustrating
the difference in the doping and strain levels in MBE (blue squares) and CVD (orange circles) graphene films
on Ge(001). The black straight lines indicate the 2D (G) mode dependence for biaxial strained undoped and
unstrained p-doped, free-standing graphene; the distances between the grey lines amount to 0.1% and to
1 × 10¹⁵ cm⁻², respectively. The average strain is compressive in the MBE film and tensile in the CVD film. The
MBE graphene film is separated from Ge(001) by a graphene-like interfacial layer23.

Figure 6. Small graphene molecule (29 atoms) on Ge(001)-p(2 × 2). (a) MBE: the edge C atoms tend to make
bonds with the substrate. (b) CVD: the edge C atoms are terminated with H, and some of these atoms (here: one
atom) makes a weak bond to the substrate.
by nucleation on longer C strings (like $C_8$, Fig. 7b), which prefer to reside between dimer rows and may initiate the growth of graphene in directions somewhat misaligned with dimer rows.

The contribution of Ge DV to graphene nucleation during CVD is less direct, although – as it will be argued – important. During the first moments after the exposure of Ge(001) to ethylene, DVs have no influence: at the growth conditions, they do not trap $C_2H$, $C_2H_2$, or $C_2H_3$. Rings such as $C_6H_4$ or $C_{10}H_6$ can be produced on the flat surface by polymerization of the direct products of $C_2H_4$ decomposition. These arenes stand vertically on the surface and are strongly bonded to the Ge(001) dimers by Ge-C bonds. But if a polyarene has grown large enough to lean towards the horizontal position as the result of van der Waals attraction, the Ge-C bonds become weakened and are now vulnerable to attack by H atoms. The latter leave the Ge(001) dimers and stick to C. We estimate that, at the high temperatures needed for the CVD growth, even when the H originates only from $C_2H_4$ decomposition and when the partial pressure of $C_2H_4$ is only $2 \times 10^{-2}$ mbar (as in our experiments), the whole edge of such a polyarene becomes saturated with H (Fig. 6b) and the molecule can move freely. It is therefore not obvious how the crystallographic information is transferred from Ge(001) to graphene grown from $C_2H_4$, as we observe in experiment (Fig. 2).

In order to preserve its orientational correlation with the substrate, a “graphene molecule” must – until it grows to a size precluding its diffusion on the surface – retain the Ge-C bonds that prevented its small seed from evaporating. This is where the DV defects play an important role. It turns out that DV sites catalyze polymerization of $C_2H$ molecules. First, the process of $C_4H_2$ formation is significantly more favorable inside a DV (Fig. 8a) than on the perfect surface: the energy gain increases from 1.6 eV to 2.1 eV. $C_4H_2$ (DV) traps $C_4H_4$; the resulting $C_8H_6$ molecule is likely to remain inside the dimer vacancy (Fig. 8b). The Ge-C bonds are stronger inside a DV than on the perfect surface, and the local geometry of the defect makes them less exposed to H attack. A graphene flake having orientation correlated to that of the orientation of the DV defect may grow in this way. The probability that the nucleus becomes detached before its orientation is fixed by van der Waals forces increases however with the background concentration of H.

**Crystallographic and electrical quality of graphene.** ARPES measurements strongly indicate that the orientational domains are so small that many of them fit into the area sampled by the X-ray beam spot, i.e. into space of about $100 \times 100$ nm$^2$. The presence of two orientational domains, or more precisely, of grain boundaries between these domains, is thus a factor limiting the electrical quality of graphene grown on Ge(001)/Si(001). According to the mechanism proposed above, the domain size is associated with the abundance of Ge dimer vacancies (on which graphene nucleates), and with the presence of surface steps on the substrate surface (which are
a/(\tan(\alpha) \cdot 2) \approx 15 \text{nm}, where \( a_0 \) is the surface lattice constant of Ge(001). This is in accordance with the observation that both orientations occur with the same frequency on the area covered by the X-ray spot. The quality of graphene increases with the substrate temperature because the concentration of dimer vacancies \( N_{\text{DV}} \) (thermally activated with \( E_{\text{form}}^{\text{DV}} \approx 0.6 \text{eV} \)) increases with temperature slower than is the decrease of the fraction \( p_{\text{occ}}(\text{DV}, x) \) of DVs occupied with hydrocarbons \( x \). For example at thermal equilibrium, \( p_{\text{occ}}(\text{DV}, \text{C}_2\text{H}_4) \) is thermally activated with \( E_{\text{form}}(\text{C}_2\text{H}_4) \approx 1.7 \text{eV} \). In other words, the concentration \( s - p_{\text{occ}}N_{\text{DV}} \) of the growing nuclei decreases with temperature. Small domain size is responsible also for the curved shape of CVD graphene edges (Fig. 1). The sharp form of MBE graphene edges (Fig. 1) is probably an indication that the graphene-like interfacial layer underlying most of the MBE graphene \(^{35} \) contains much smaller concentration of efficient nucleation seeds, so that the domains may grow much larger, in principle reaching even into the \( \mu \text{m} \) range.

**Doping and strain.** Doping of the MBE film may originate solely from molecules from the ambient air that are adsorbed on top of the graphene. Higher doping of the CVD film may be caused by Ge atoms \(^{35} \), since the CVD graphene remains in direct contact with Ge(001).

The compressive strain in the MBE film (−0.3%) may possibly be caused by thermal expansion coefficient difference between graphene (−3 \( \times \) 10\(^{-6} \text{K}^{-1} \)) and Ge (6 \( \times \) 10\(^{-6} \text{K}^{-1} \)). Cooling from 930°C to 25°C would produce strain of about −0.8%, which by its absolute value is higher than the −0.3% estimated under the assumption of biaxial strain (Fig. 5b). It follows that the graphene film forming on the interfacial graphene-like layer grows under tensile strain, possibly induced by the strain in the interfacial layer. Indeed, the strain in the CVD film, which remains in the direct contact with Ge(001), is tensile, indicating that the strain in the growing graphene is also tensile and higher than that the +0.4% measured at room temperature. This may be caused by coalescence of the small graphene domains.

**Conclusion**

Here we report on the successful growth of graphene on the CMOS (complementary metal-oxide-semiconductor) technology compatible Ge(001)/Si(001) substrates from C\(_2\)H\(_4\) precursor in a cold-wall CVD chamber. Experimental results are interpreted on basis of \textit{ab initio} DFT calculations. Presented mechanisms of the graphene synthesis are deduced from experimental and theoretical data.

The major experimental result is that the graphene film on Ge(001)/Si(001) consists of two orientational domains rotated by 30° with respect to each other, and that the grain size is comparable to step-step separation of the substrate. Poor crystallographic quality (high Raman D mode peak) and poor electrical quality (low carrier mobility) of the CVD graphene produced in this process are attributed to the small domain size. Arguments linking the grain size with the size of surface terraces separated by Ge(001) monatomic steps are presented. It was argued that the dimer vacancies omnipresent on the surface act as seeds on which graphene can nucleate and that the orientation of a domain is determined by the orientation of Ge dimers on the surface terrace in which the domain nucleated. Small graphene nuclei released from the dimer vacancy sites by hydrogen atoms are volatile; this contributes to the reduction of graphene growth rate by increased surface concentration of hydrocarbon and to the increase of the average domain size. For completeness and illustration, the CVD growth mode and mechanism was compared to those for the growth from atomic carbon in MBE chamber.

**Methods**

**Samples preparation.** Graphene was grown in a multi chamber ultrahigh vacuum system (UHV) with a base pressure in the lower 10\(^{-10} \text{mbar} \) range. The substrate was a Ge(001) film (2 \( \mu \text{m} \)) grown by CVD on a Si(001) wafer and then overgrown \textit{in situ} by epitaxial Ge buffer (about 4 nm). The substrate temperature during graphene growth was close to but below the melting point of bulk Ge (it was around 930°C). Growth time was varied between 90 min and 200 min in both cases. The substrate was heated from the backside by a graphite filament and the temperature was calibrated by pyrometer. The graphene CVD growth was performed by using ethene (C\(_2\)H\(_4\)) as a precursor in a UHV-CVD chamber (Riber). The pressure was in the range of 10\(^{-2} \text{mbar} \) during deposition.
and the flow of ethylene was 5 sccm. In addition, graphene samples were grown by a physical method in a MBE chamber (DCA), as described in ref. 23: carbon atoms were evaporated via e-gun, and the pressure was in the lower $10^{-7}$ mbar range.

**Sample characterization.** Quality of graphene was evaluated by μ-Raman spectroscopy using Renishaw In-viaFlex spectrometer and green laser light (514 nm) with a spot diameter of about 500 nm. Transport measurements were performed with a 4-tip scanning tunnelling microscope (4-tip STM, Omicron) with a high resolution scanning electron microscope (SEM) used as a tip-navigation tool.

**ARPES experiments.** Photoemission experiments were performed at the nano-ARPES microscope of the ANTARES beamline of the SOLEIL synchrotron, equipped with two Fresnel zone plates (FZP; i.e. for low and high photon energy) for focalization of the synchrotron radiation beam and an order selection aperture (OSA) to eliminate higher diffraction orders. Samples are mounted on a nano-positioning stage placed at the coincident focus of the Scienta R4000 analyser and the FZP focal point, which is responsible for the sample rastering during the image acquisition. The spatial resolution is determined by the FZP resolution and the mechanical stability of the sample stage. Experiments were performed at photon energies $h\nu = 100$ eV with linearly polarized light for Fermi surface mapping and at $h\nu = 30$ eV with circularly polarized light for the energy scans around the K point of graphene. The circular polarization in these experiments was selected in order to avoid the “dark corridor” problem along the $\Gamma - K$ direction for graphene. The sample was kept at room temperature during ARPES experiments. The base pressure during all experiments was better than $5 \times 10^{-11}$ mbar.

**DFT calculations.** Pseudopotential density functional theory (DFT) total energies and atomic and electronic structures were calculated by Quantum Espresso. Exchange and correlation energy was approximated by the PBE gradient-corrected functional. Periodically repeated slab consisting of eight Ge(001) planes separated by 10 Å of the empty space was used. The calculations were done for square Ge(001)-p(2×2) surface cells, each containing eight dimers on one side and passivated with H atoms on the other side. Further calculation details can be found elsewhere.

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Author Contributions
J.D. performed DFT modeling. G. Lippert, G. Lupina, T. Schaffus, M.K. and D.V. prepared graphene on Ge substrates by CVD and MBE methods. G. Lippert, J.A., I.C., M.K., D.V. and M.-C.A. performed nano-ARPES measurements. J.B. and C.T. carried out 4-tip STM characterizations. F.H. and J.M. performed Raman experiments. J.D., G. Lippert, J.A., I.C., Y.S.D., F.H., G. Lupina, J.M., T. Schaffus, T. Schroeder, M.K., C.T., D.V. and M.-C.A. analyzed the data and contribute in the preparation of the manuscript.

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