Genesis and evolution of extended defects: The role of evolving interface instabilities in cubic SiC

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I. INTRODUCTION

The growth of high-quality substrates for microelectronic applications is one of the key elements that drive society toward a more sustainable green economy. The development of new high-power and high-current devices for clean energy production, energy harvesting, and energy conversion is a crucial asset for limiting the use of carbon-based fuels. Decreasing the size and weight of power devices would be very desirable, as well as reducing their cooling requirements while maintaining their high performances at elevated temperatures.

Emerging wide bandgap semiconductor devices, like those built with SiC have the potential to revolutionize the power electronics industry through faster switching speeds, lower losses, and higher blocking voltages, which are superior to standard silicon-based devices. The current epitaxial technology enables more controllable and less defective large area substrate growth for the hexagonal polymorph of SiC (4H-SiC) with respect to the cubic counterpart (3C-SiC). However, the cubic polymorph exhibits superior physical properties in comparison to its hexagonal counterpart, such as a narrower bandgap (2.3 eV), possibility to be grown on a silicon substrate, a reduced density of states at the SiC/SiO₂ interface, and a higher channel mobility, characteristics that are ideal for its incorporation in metal oxide semiconductor field effect transistors. The most critical issue that hinders the use of 3C-SiC for electronic devices is the high number of defects in bulk and epilayers, respectively. Their origin and evolution are not understood in the literature to date. In this manuscript, we combine ab initio calibrated Kinetic Monte Carlo calculations with transmission electron microscopy characterization to evaluate the evolution of extended defects in 3C-SiC. Our study pinpoints the most critical issue that hinders the use of 3C-SiC for electronic devices is the high number of defects in bulk and epilayers, respectively. Their origin and evolution are not understood in the literature to date. In this manuscript, we combine ab initio calibrated Kinetic Monte Carlo calculations with transmission electron microscopy characterization to evaluate the evolution of extended defects in 3C-SiC. Our study pinpoints the antiphase boundaries is particularly important to achieve good quality crystals, which can then be incorporated in electronic devices.
temperature and 200 °C, which leads to a large reduction of device-on-resistance at realistic junction temperatures for power device operation.

The growth and fabrication of high-quality 3C-SiC epilayers and bulk wafers is becoming very important. The most crucial issue that hinders the use of 3C-SiC is the high number of defects in both bulk and epilayers. Therefore, an intense research effort has been dedicated to this problem.

With this respect, as in the synthesis process of any material, the interplay between surface instabilities that occur during the crystal growth, bulk defects, and crystal boundaries are key issues for the understanding of the kinetics involved in the growth process. Indeed, morphological and micro-structural characterizations of grown crystals often indicate a clear correlation between the defective structures and the evolution of the interfaces during the synthesis stage. Silicon carbide crystals are a test bed for these studies due to the extreme polymorphism caused by the small energetic cost of stacking disorder. The different sequences of SiC bi-layers in the three symmetric positions (conventionally named ABO) of the hexagonal close packing creates more than 250 different polytypes (polypeptide instability). A direct consequence of the huge polymorphic character of SiC is the facile generation of extended defects in a plethora of configurations. The Stacking Faults (SFs) class is probably the most common and most studied one, due to the tight relationship with the SiC polytypism. In addition, crystallography studies often focus on antiphase boundaries (APBs), micropipes, dislocations, and other kinds of missing crystal homogeneity.

Knowledge on defects in 3C-SiC crystals is usually obtained by post-growth experimental characterizations. However, the genesis and the evolution of crystal imperfections is, in the majority of cases, a moving interface/surface. As a consequence, due to the lack of reliable in situ dynamic analyses of the evolving systems, it is difficult to achieve the correct and complete description of the atomic mechanisms governing this complex phenomenology. The integration of structural analysis and process simulations could overcome these difficulties. However, it is easy to understand that the simulation tools should achieve the following unconventional performances: (a) a structural evolution occurring on macroscopic time scales (from seconds to hours); and (b) a simulation mechanism that can be correctly modeled only at the atomic scale, as a result of the cooperative dynamics of (at least) billions of atoms.

Several theoretical approaches have been suggested to study the SiC crystal growth, which can be categorized as off-lattice approaches (ab initio, molecular dynamics, and related structure optimization procedures) or on-lattice (e.g., Kinetic Monte Carlo (KMC)). Off-lattice codes with continuous particle positions can be used for the research of defects but are unable to reach the time scales of the epitaxial process. The latter can be hindered either by the full simulation with lattice vibrations or by the evaluation of the relevant barriers from the potential energy surface on the fly.

Stochastic simulations within the KMC approach could satisfy the two requirements of atomic accuracy and large-scale simulations, allowing for a multi-scale description of the epitaxial growth process. On-lattice models (including Lattice Gas and Solid-on-Solid approaches) are characterized by evolving Monte Carlo particles that lay on the perfect lattice sites but fail in correctly describing the generation and evolution of extended defects.

In particular, previous KLMC methods formulated for silicon carbide materials suffer from fundamental limitations: they either allow approximate defect simulation but do not simulate separately the evolution of the two system atoms, i.e., Si and C, in the compound, or they simulate Si and C individually, but the defect formation and evolution cannot be estimated.

A Kinetic Monte Carlo (KMC) formulation on augmented or super-lattices offers great flexibility to simulate the structural evolution of defective systems, overcoming all previously noted limits. We simulate the complex kinetics scenario during growths of cubic SiC by means of a recently developed stochastic Kinetic Monte Carlo superLattice (KMLC) code, which aims at simulating with an atomistic resolution the growth process of compound materials characterized by sp³ bond symmetry. The peculiar characteristic of this method is the possibility to study the generation and evolution of extended and point defects during the growth process. In particular, in the class of extended defects, stacking faults and APBs are of primary interest due to their impact on the material quality and their negative consequences when building devices and applications. The code treats carbon and silicon atoms as independent particles. In order to increase the predictive power of the KMCsL simulations, we make use of ab initio calculations at a density functional theory level to extract the energetics needed to set rate constants within the Transition State Theory for the active Monte Carlo particles.

We combine ab initio calculations of several surface configurations and the temperature dependence of the solid/vapor SiC-phase equilibrium to specifically calibrate the event rates for 3C-SiC. The analysis of several equivalent replicas of the KMCsL evolution starting from the same initial defective seed reproducing a 3C-SiC crystal with two different (anti)phase domains discloses a variety of kinetic behaviors, which finds a clear counterpart in real growth processes, revealing the atomic mechanisms responsible for extended defect generation and evolution.

II. EXPERIMENTAL

The growth process was performed using a Chemical Vapor Deposition (CVD) reactor, on (001) “on axis” oriented Si substrates, using silane (SiH₄), propane (C₃H₈), and H₂ as silicon precursor, carbon precursor, and gas carrier, respectively. A multi-step growth process was carried out: it comprises “etching,” “carbonization,” “growth,” and “cooling down” steps. Each of these steps consists of further sub steps. Etching was performed under an H₂ flux at a temperature of 1100 °C. During carbonization, the temperature was set to 1120 °C, and propane was added to the carrier gas. The growth was performed at 1300 °C with both Si and C precursors. The growth rates were maintained at 1 μm/h. Finally, the temperature was decreased until reaching room temperature, and the chamber was flushed with Ar in order to avoid etching of the surface due to the presence of hydrogen.

Focused samples were characterized by means of High Angle Annular Dark Field Scanning Transmission Electron Microscopy (HAADF-STEM) and conductive atomic force microscopy (CAFM). HAADF-STEM was carried out on a Cs-corrected JEOL ARM200F probe, equipped with a cold field emission gun and working at 200 kV. We operated with three detectors, having a low, medium, and very high scattering angle, respectively. The dark field detector inner semi-angle (80 mrad) allowed the signal to be roughly proportional to the atomic number Z. Under these conditions the intensities observed in the scanning transmission electron microscopy micrographs could be...
directly interpreted. The microscope has a nominal resolution of 0.68 Å.

Nanoscale resolution current mapping and morphological images of the 3C-SiC layers grown on the Si (001) substrate were obtained by CAFM. These analyses were carried out using the TUNA module of the DI-3100 microscope with the Nanoscope V controller. The current maps were acquired with a Pt coated Si tip scanned on the sample surface, by applying a DC bias between this conductive tip and a macroscopic Ni2Si Ohmic contact fabricated onto 3C-SiC. In this configuration, local inhomogeneities in the conductivity of 3C-SiC associated with electrically active defects reaching the sample surface can be directly visualized.

III. COMPUTATIONAL

Crystal growth was simulated by means of a KMCsL approach, designed to study at an atomic resolution the growth kinetics of elements, alloys, and compounds characterized by the sp3 bond symmetry. Formalization and implementation details of the KMCsL code are discussed in Ref. 52. Deposition and evaporation are the active Monte Carlo events, driving the stochastic evolution. In the application of the KMCsL to the SiC material presented here, the silicon and carbon atoms undergo independent kinetics. We point out and demonstrate in the following that a dense superlattice, where the original lattice of the ideal crystal is a sublattice of the superlattice itself, can correctly accommodate a large class of SiC defective configurations. Indeed, the code is able to simulate the evolution of both point-like and extended defects, like stacking faults of different symmetries, APBs, and grain boundaries. Moreover, the KMCsL can also simulate the morphology evolution during the growth (e.g., the epitaxial growth or etching of flat, structured, or patterned substrates, as well as nanoparticles of various shapes). In the case of surfaces, periodic boundary conditions are applied in the planes that are orthogonal to the growth direction.

In order to be reliably applied to a given material, configuration-dependent parameters in KMCsL have to be calibrated using a more fundamental approach, or by means of an ad hoc experiments performed in controlled conditions. Following a sequential multiscale approach, we used ab initio energetics at a density functional theory level to set temperature-dependent input frequencies for the local transitions involving atoms bonded X ∈ {C, Si} to the evolving SiC surface/interfaces (i.e., the subset of under-coordinated atoms at a given time t of the simulation). As such transitions imply bond breaking, we assume that the related KMCsL evaporation frequencies follow Arrhenius-type functions of the generalized binding energies $E\{\Sigma_{\text{loc}}(X,t)\}$ for the X detachment,

$$\nu(X,t) = \nu_0(T) \times \exp \left(-\frac{E\{\Sigma_{\text{loc}}(X,t)\}}{kT}\right).$$

We note that the frequency in Eq. (1) depends on the local atomic configurations $\Sigma_{\text{loc}}(X,t)$ around atom X before the transition, which we classify in the current KMCsL approximation by means of the number (one/two/three) of sp3 X-Si and X-C bonds in the first-neighbor shell (see Sec. S1 of the supplementary material for the setting of the evaporation energetics and frequencies). Arrhenius pre-factors $\nu_0(T)$ have been set with partial pressures at equilibrium in the gas phase during silicon carbide sublimation. We notice that in the absence of deposition, the method is also calibrated to the temperature-dependent sublimation kinetics of the SiC material in equilibrium with its gas components.

Deposition frequencies depend on the experimental controlled growth conditions implemented in the CVD or physical vapor deposition chambers. In our modeling approach, these frequencies are again configuration dependent. They reproduce the average rate of
the Si or C atoms attached/released at the solid surface independently from the particular reactions’ mechanisms involving atomic or molecular components in the vapor phase. We have calibrated these parameters in order to reproduce experimental time-dependent SiC profiles in CVD growth processes on structured silicon substrates with inverted pyramid patterns grown with the same method described in Sec. II. The simulated evolution starts from a SiC seed, reproducing the growth step only (for the calibration of the KMCsL deposition frequencies see Sec. S1 of the supplementary material). In our case, KMCsL deposition parameters guarantee, as in the experimental case, controlled simulated epitaxial growth with a low generation of point defects (vacancies or anti-sites). The generation flux rate of point defects is below $2.0 \times 10^{10} \text{cm}^{-2} \text{s}^{-1}$, while bulk annihilation events are not considered here, since we focus our analysis on the evolution of extended defects.

We simulated the epitaxial growth of 3C-SiC substrates exposing the (001) surface (the $z$ axis of the Cartesian system lies along the [001] direction). A cubic simulation box with a side of 740.8 Å was set. Considering the superlattice description, the box contained $\sim 8 \times 10^9$ sites. We started the KMCsL run from a 3C-SiC slab with a thickness of 137.8 Å. The initial active KMC particles lying at the (001) surface consisted of 51 200 under-coordinated atoms.

The simulation results starting from an ideal 3C-SiC slab (i.e., no defects are considered in the bulk of the initial substrate) with the surface aligned to a (001) plane are discussed in Sec. S2 of the supplementary material. Here we notice that in many replicas of equivalent simulated growths (from the stochastic point of view), no generation of extended defects has been observed, demonstrating that this growth direction is rather robust against extended defect formation. Contrarily, in the absence of a step-flow growth mode, stacking faults form due to the polytype instability implemented in the KMCsL model in the case of growth over [111] facets.

In order to evaluate the conduction properties of stacking faults and APBs, we performed quantum transport calculations within the nonequilibrium Green’s function formalism, considering electronic Hamiltonians obtained from the density functional theory (e.g., see Ref. 66). We used the SIESTA code along with the local density approximation for the exchange-correlation functional. Electronic wave functions were constructed on a double-$\zeta$ polarized basis set for both Si and C, whereas the ionic cores were described with norm-conserving Troullier-Martins pseudopotentials. All atoms were allowed to fully relax until forces were less than 0.04 eV/Å. The

![FIG. 2](image-url)

FIG. 2. Evolution of an APD with four APBs along the (110) planes during the 3C-SiC substrate growth along the [001] $z$ direction. Epitaxially grown 3C-SiC substrate: (a) atoms at the antiphase boundaries in the bulk; (b) under-coordinated atoms of the input KMC substrate, highlighting the initial $xy$ surface exposed to the deposited gases; (c) under-coordinated atoms taken after the start of the KMC deposition, highlighting the formation of the surface depletion at the APB; and (d) under-coordinated atoms taken at the end of the KMC evolution, highlighting the final $xy$ surface. The dashed and continuous segments highlight, respectively, the (111) Si-face and C-face; the red arrow depicts the APB drift during an epitaxial growth due to the different growth velocity of the exposed (111) Si-face and C-face.

![FIG. 3](image-url)

FIG. 3. (left) Scanning transmission electron microscopy image of an APB within a 3C-SiC (001) substrate. (right) Zoom of the surface depletion induced by the APB and lying at the (001) surface. It is worth noting that during the experimental growth along the [001] direction, the APB moves along the [110] direction, reaching a final tilted configuration with respect to the (001) surface. The KMC evolution reported in Fig. 2 agrees quite well with the experimental analysis of this figure.
geometrical configuration of the supercell structures allowed for a direct contact between the metallic electrodes and the defects (see Fig. S2 of the supplementary material).

IV. RESULTS
The ideal conditions reproducing a flat substrate are difficult to achieve experimentally, since the initial substrate itself usually presents non-homogeneous regions and preexisting defects. AntiPhase Domains (APDs) and related APBs are rather common defects when growing 3C-SiC crystals, and they can form as possible boundaries of 3D structures merging in Volmer-Weber or Stranski-Krastanov growth modes. A growing [e.g., (001)] surface with a preexisting extended defect (as the APB) exposes a defective region (linear "cut"), which can cause a disturbance on the growth kinetics. This growth kinetics unravels the eventual genesis, evolution, and interaction of extended defects (of various nature) starting from a defective seed layer.

We have considered a suitable model for the study of the kinetics of APBs within our KMCsL approach. At an atomic level, we introduce the APB as a planar distribution of anti-sites along a {110} crystal plane. At a mesoscopic level, we include four APBs, which reproduce a space distribution of APDs (considering the periodic boundary conditions imposed in the simulation box). This symmetric choice minimizes the possible artifacts caused in other possible configurations by periodic conditions and opposite phase domains. Green (silicon) and gray (carbon) atoms in Fig. 1(a) outline the four APBs and the different polarities of the substrate: considering the periodic boundary conditions on the xy plane, the substrate looks like an outline of APDs, favoring defect interactions in a reasonable time and space scale. Anti-sites along a [110] crystal plane are highlighted in Figs. 1(b) and 1(c). Figure 1(c) represents the orthogonal view of a zoom of the APB showed in Fig. 1(b). Dashed and continuous segments of Fig. 1(c) highlight, respectively, the [111] Si-face and the C-face with the different
polarity. Note that no under-coordinated defects are present in the bulk of this structure, since the defective region is composed of planes with homoatomic Si–Si and C–C bonds.

The analysis reported in Secs. IV A–IV E discusses several representative features occurring randomly in the many-replicas of the KMC simulation (in total we have investigated \(25^2\) replicas of simulated evolution starting from the same initial state). We note that all figures in the paper refer to independent KMC runs where we modified only the input seed of the random number generator.

A. Antiphase boundary kinetics

Figure 2 reports the typical KMC evolution of a domain with four APBs along the \{110\} planes during the 3C-SiC substrate growth along the [001] z direction. Figure 2(a) shows only atoms at the APBs in the whole simulation box, where the initial 3C-SiC (001) substrate grows in the z direction. Initial symmetric APBs are visible at surface portions in Fig. 2(b). Contrarily to the ideal case, the presence of the APBs in the underlying bulk crystal significantly alters the morphology of the growing surface. Figures 2(b)–2(d) reveal under-coordinated atoms, highlighting the “running” (001) surface. We observe that the bulk antiphase boundaries generate surface local depletion, with the two facets oriented very close to two \{111\} planes that originate from the (110) lines in the growing surface. For example, the APB on the (110) plane realizes a surface depletion with \(111\) and \(111\) surfaces. Such depletion is formed at the start of the KMC deposition, as highlighted by Fig. 2(c).

The APBs show a clear drift with different direction of motion for the boundaries of different symmetry, as observed in Fig. 2(a). The depletion depth, as well as the drift velocity of the antiphase boundaries depend on the deposition conditions. The drift mechanism [represented by the red arrows of Figs. 1(c) and 2(d)] relies on the different velocity of the \{111\} facets forming the groove. Indeed due to the phase change induced by the APB, these facets are of opposite polarity (i.e., C-face and Si-face). Dashed and continuous segments in Fig. 2(d) highlight, respectively, the \{111\} Si-face and the C-face forming the surface depletion at the APB. \{111\} Si-face and C-face are also emphasized in Fig. 1(c).

In our deposition condition, C-faces have different growth velocity with respect to Si-faces (see Sec. S1 of the supplementary material). As a consequence of this non-symmetric APB kinetics, the APD gradually switches from a square shape to an elongated one as shown by the final KMC snapshot of Fig. 2(d).

In Fig. 3, we show a scanning transmission electron microscopy image of an APB within a 3C-SiC (001) substrate. It is worth noting that during the experimental growth along the [001] direction, the APB moves along the [110] direction, reaching a final tilted configuration with respect to the (001) surface. Figure 3 also provides a zoom of the surface depletion induced by the APB and lying at the (001) surface. The KMC evolution reported in Fig. 2 agrees quite well with the experimental analysis of Fig. 3.
that are not in the ideal crystal configuration. Figure 4 represents an orthogonal view of an SF sequence, the presence of \{111\} planes on the surface depletion allow for ABC ABC …, which lies on the crystal symmetric positions usually named A, B, or C. The cubic 3C-SiC on the surface depletion causes (due to polytype instability) the generation of triple SFs [see Figs. 5(a) and 5(b)]. Once the SF(3)s are generated by the APB, the two extended defect-types (SF(3)s and APB) follow independent kinetics: the APB continues its drift along the [110] direction, while the SF(3)s grow on the (111) planes [see snapshots of Figs. 5(b)–5(d)]. SF(3)s hold an elongated shape as shown in Fig. 9(e).

Figure 6 shows a scanning transmission electron microscopy image of a stacking fault generated by an APB during the epitaxial growth of a 3C-SiC (001) substrate. It evolves on \{111\} planes independently from the APB kinetics. The surface depletion is also visible at the terminal (001).

Figure 7 shows the calculated quantum transport properties of a triple stacking fault and an antiphase boundary, and compares them with the values expected for non-defected 3C-SiC. Both defects show an increased conductivity with respect to the ideal crystal due to an enhanced density of states at the defect area (see Fig. S3 of the supplementary material). Additionally, the antiphase boundary introduces triple SF (also known by the term "micro-twin," ABC ACB ABC) separates two crystal regions in a perfectly epitaxial order, whereas the only atoms that are not in the correct crystal positions are those inside the extended defect. The faulted stacking sequence of the Si-C dimers along \{111\} directions with respect to the ideal periodic sequence, ABC ABC ABC …, is indicated by the light purple (silicon) and light blue (carbon) atoms. The orange line is a guide for the eye, highlighting the maintained epitaxial order in the two semi-spaces separated by the defect.

Figure 5(a) shows the generation of a couple of triple stacking faults from the surface depletion induced by an APB. Figures 5(b)–5(d) represent a sequence of snapshots for the undercoordinated atoms taken at different KMC times. The triple SFs are visible in this representation due to the presence of tree-fold coordinated Monte Carlo particles at their boundary (i.e., at the associated partial dislocation). The zoom in Fig. 5(c) shows the three-bilayers structure of the two generated SF(3)s.

The local asymmetry of the APB and the presence of \{111\} facets on the surface depletion causes (due to polytype instability) the generation of triple SFs [see Figs. 5(a) and 5(b)]. Once the SF(3)s are generated by the APB, the two extended defect-types (SF(3)s and APB) follow independent kinetics: the APB continues its drift along the [110] direction, while the SF(3)s grow on the (111) planes [see snapshots of Figs. 5(b)–5(d)]. SF(3)s hold an elongated shape as shown in Fig. 9(e).

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states within the 3C-SiC bandgap that narrow the conduction gap. The large transmission coefficients of these defects as compared to the bulk indicates that both should be extremely conductive when contacted by the metallic electrodes. On the contrary, their effect on the transport properties should be restrictive when not in direct contact with the electrodes and perpendicular to the transport direction. In that case, they should act as potential barriers hindering the conduction of current.

Figures 8(a) and 8(b) show a representative morphological image and the corresponding current map collected on the 3C-SiC surface by applying a DC bias of 0.5 V to the tip. The boundary between two neighboring domains can be observed in the morphological map [see Fig. 8(a)]. On the other hand, a large density of conductive features, mostly uncorrelated to the topography, is revealed by the current map in Fig. 8(b) and reflects the conductivity of APBs and SFs. The structural properties of these extended defects connecting the surface to the bulk of 3C-SiC layers will be extensively discussed in this paper. This preliminary electrical investigation clearly indicates how the presence of such a high density of conductive paths can have strong implications on the performances of vertical or quasi-vertical devices based on heteroepitaxial 3C-SiC on Si. As an example, they can account for the typically low values of the turn-on voltage and the high leakage current observed in metal/3C-SiC Schottky barrier diodes.

C. SF\(_{(3)}\)-APB interference: Growth termination and polarity changes of triple SFs

After its generation, an SF\(_{(3)}\) can continue its extension during the epitaxial process at the 3C-SiC (001) surface, eventually increasing its size longitudinally (i.e., moving the partial dislocations that bind the SF\(_{(3)}\) on the slipping plane). In the case of the triple SF, which perturbs minimally the crystal order of the embedding crystal, the defect’s evolution is essentially driven by atomistic transitions occurring at the growing surface. The analysis of several equivalent replicas of the simulated kinetics seems to indicate that the persistence and enlargement of the SF\(_{(3)}\) is ruled by a critical nucleation phenomenon: thanks to random fluctuations only defects that reach a critical lateral size manage to survive and extend during the simulated growth. However, the high kinetic (meta)stability of large SF\(_{(3)}\)s is significantly hampered if...
another extended crystal imperfection interferes with the defect evolution.

In particular, we investigated this defect-defect interaction, thanks to the presence of multiple APBs in the same KMC framework. As a possible result of this extended defects interaction, we observed both SF(3) growth termination and survival (followed by a polarity change) due to a different APB coming from the orthogonal direction to the original APB that generated the SF(3).

Figure 9 illustrates the whole process of the SF(3) growth suppression (Multimedia view). A couple of triple SFs are generated at the surface depletion induced by an antiphase boundary [Fig. 9(a)]. Then they come in contact with another APB orthogonal to the APB that generated the two SF(3)s [Fig. 9(b)]. The red arrow of Fig. 9(a) highlights the APB that generated the SF(3)s, while the blue arrow points to the coming APB, which suppresses the SF(3)s. The SF(3)s growth is then suppressed (in this case) by the APB, as shown by snapshots in Figs. 9(c) and 9(d), taken at subsequent times (orange arrows highlight the contact points between the APB and the SF(3)s). Finally, driven by the APB drift along the [110] direction, the SF(3)s close, and the crystal surface is healed [see Fig. 9(e)]. A movie of the KMC evolution described in Fig. 9 is reported in the supplementary material. Figure 9(e) also shows small SF(3)s that are generated and immediately closed. It is worth recalling now that SF(3)s and the associated partial dislocations cannot be self-terminated, whereas either the generation or the termination of SF(3)s is always connected with the presence of a domain boundary. The generation and termination processes are stochastic in nature, due to the isoenergetic configurations of 3C-SiC, and depend on the local epitaxial growth at an atomic level.

KMC results for the growth termination of triple SFs at defective domain boundaries agree with experimental observations of the epitaxial growth of 3C-SiC antiphase domains. Figure 10 provides a scanning transmission electron microscopy image showing the growth termination of various SF(3)s at a defective antiphase boundary during the epitaxial growth of a 3C-SiC (001) substrate. Both SF(3)s and APB defects show independent kinetics during the epitaxial growth. Once they come in contact, the SF(3)s disappear. A posteriori experimental characterization is not able to reveal the mechanism beyond their interaction, which, conversely, can be easily deduced from the KMC simulation.

Another kind of behavior can be observed from the KMC simulations: an SF(3) can continue its extension during epitaxial growth, surviving from the interaction with an APD boundary by changing its polarity. Figure 11 illustrates the polarity change of a triple SF.
following its interaction with a moving APB during the 3C-SiC epitaxial growth along the [001] z direction. We show under-coordinated atoms taken at different KMC times. First, the SF(3) interacts with the APB [see Fig. 11(a)]. The SF(3) survives and follows independent kinetics with respect to the APB [Fig. 11(b)]. The latter moves toward the [110] direction, while the triple SF instead turns away from the APB along the (111) planes. To highlight the SF polarity change, we show in Figs. 11(c) and 11(d) the atoms that are not epitaxially ordered and associated, respectively, to the triple SF of Figs. 11(a) and 11(b). The zoom in Figs. 11(b) and 11(d) highlights the triple stacking fault and the region where the polarity change takes place, respectively.

The original plane of the SF(3) does not change since a polarity change involves only a local reversal of the Si–C doublet. So the original plane of the SF, in the reference system of the initial substrate, remains the (111) plane. For similar reasons, the stacking sequence does not change once the SF passes from a crystal domain to another crossing an APB. The basal plane dislocations at the boundary of the stacking faults, which disappear by crossing in the case of SFs growth termination [see Figs. 9(c)–9(e)], react in the case of polarity change and modify their atomic configuration. This modification of the bonding network can be described by a direction change of the dislocation vectors within the SF(3) habit plane and an inversion Si–C ⇒ C–Si of the defects’ pairs (due to the polarity change) along the dislocations’ direction.

Figure 12 shows a scanning transmission electron microscopy image that reveals the interaction between a triple stacking fault with a grain boundary. It is apparent that an SF(3) belonging to the crystal on the left side encounters the APB and emerges in the crystal on the right side. Although it is impossible to detect the polarity change of the SF, we note its survival after the interaction with an APB, and its subsequent independent kinetics with respect to the APB.

**D. Generation and bulk growth of SF(1) tetrahedra**

The similar energetics of different SiC polytypes can accommodate a large family of extended defects, characterized by various wrong stacking sequences with respect to the cubic polymorph. As a result, very complex behavior can also be obtained. Three dimensional arrangements of SFs called SF tetrahedra (SF(T)) can be generated. They are constituted by 4 intrinsic single SFs (SF(1), i.e., SF with a single wrong sequence) limited by stair rod dislocations forming a tetrahedron. An SF(T) can originate on the [111]-oriented surface depletion induced by an APB, in a similar way to SF(3)’s reported in Secs. IV B and IV C. Figure 13 shows the generation of a single stacking fault from an APB and the bulk growth of the SF tetrahedron. Initially, the epitaxial growth produces two consecutive triple SFs at the APB [Fig. 13(a)]. Then, the same APB in Fig. 13(b) generates a single SF [aligned to the previous triple SFs, i.e., lying on the (111) plane]. Differently from the triple SF, the single SF has a non-negligible Burgers vector and separates the crystal into two regions that are not in perfect epitaxial arrangement. As a consequence, an SF with finite size, transmits this atomic misalignment along its border, which in turn becomes a mobile interface in the crystal that rearranges itself. Hence, the SF(T) expands, while the apical part of the defect grows, following the surface motion.

The atomistic KMC code is able to reproduce the bulk expansion of this portion related to finite size single SFs. Snapshots of Figs. 13(c)–13(e) show the growth of the SF tetrahedron in the bulk 3C-SiC crystal, while the triple SF mainly evolves only at the surface of the growing crystal. In the same KMC framework, we can unravel the opposite behaviors of the extended defects and triple SFs. The first can expand in the bulk with independent kinetics from the superficial epitaxial growth, while the second, once generated at the {111} planes

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**FIG. 14.** (a) Lateral view along the [110] direction of the extended defect shown in Fig. 13(e): atoms that are not epitaxially ordered, representing the single and triple SFs; (b) orthogonal view of a single stacking fault SF(1): the faulted stacking sequence of the Si–C dimers along (111) directions with respect to the ideal ABC ABC ... one is indicated; the orange line is a guide for the eye, highlighting the non-maintained epitaxial order in the two semi-spaces separated by the defect.
of the APB depletion, maintains a fixed shape. It is interesting to note that the SF(T) adapts the edges on [110] directions. It is known that the edges of the SF(T) are stair-rod dislocations,2 which suggests that partial dislocations tend to be straight and with screw character.

Figure 14(a) shows a lateral view along the [110] direction of the snapshot of Fig. 13(e). It indicates atoms that are not epitaxially ordered, representing the single and triple SFs. Figure 14(b) represents a cartoon with the orthogonal view of a single stacking fault SF(1) with a stacking sequence of ABC AC ABC. The faulted stacking sequence of the Si–C dimers along (111) directions with respect to the ideal ABC ABC ABC … one is indicated. An SF(1) does not maintain any epitaxial order in the two semi-spaces separated by the defect as highlighted by the orange line crossing the extended defect. Please note that a possible microscopy analysis in cross section mode of this defective volume will appear as a triple SF sequence and two split SF(1)s, which form two edges of the SF(T). Therefore, the KMCsL simulation can unravel the kinetic mechanism underlying some experimental evidence obtained by means of transmission electron microscopy.

E. Multiple sequence of triple SFs

As shown in Figs. 5, 9, and 13, triple stacking faults can be generated by the APB in sequence (two consecutive triple SFs are present in these cases). Figure 15 demonstrates the generation of a multiple sequence of triple SFs from an APB during 3C-SiC epitaxial growth along the [001] z direction. Under-coordinated atoms taken at different KMC times have been selected. In particular, Fig. 15(a) shows the formation of the first triple SF. The APB continues to generate adjacent triple SFs as illustrated in the snapshots of Figs. 15(b)–15(e). This continuous sequence of triple SFs represents a local nanocrystal domain with sequence ABC ACB ACB … (that is an extended twin configuration). A further consequence of such a triple SF sequence is a local rectangular depletion at the growing (001) surface.

Figure 16(a) shows a lateral view along the [110] direction of the snapshot of Fig. 15(e). We select atoms that are not epitaxially ordered representing the multiple sequence of triple SFs. We also show in Fig. 16(b) a cartoon representing the orthogonal view of a multiple sequence of triple stacking faults SF(3)s. The faulted stacking sequence of the Si–C dimers along <111> directions with respect to the ideal ABC ABC ABC … one is indicated. The orange line is a guide for the eye, highlighting the epitaxial order in the two semi-spaces where just one of three bilayers of ABC represented by the dark purple (silicon) and dark blue (carbon) atoms follows the stacking sequence of the 3C-SiC crystal.

V. CONCLUSIONS

This work has been motivated by the quest for a deeper control of material quality in one of the most promising semiconductors for future power electronic devices, i.e., the cubic polymorph of SiC. Indeed, the high number of defects characterizing the material obstructs its high potential in, e.g., 3C-SiC power MOSFETs, where the 3C-SiC superior physical properties could increase the performance with respect to current state-of-the-art power devices.

In spite of recent intense research focusing on the improvement of the 3C-SiC material, only incremental results have been achieved so far. We have indicated that these difficulties could also derive from the lack of complete understanding of the mechanisms ruling the generation, evolution, and interaction of extended defects in SiC. Indeed, the kinetics of defective SiC systems has been subjected to partial and often qualitative theoretical analyses, whereas experimental investigations provide an accurate description of the material status only after the synthesis process. Likely, they do not have access to the microstructural evolution during the process itself.

The results discussed demonstrate that the combination of advanced simulation methodologies and experimental characterization techniques closes this knowledge gap. We would like to summarize the main scientific achievements of the research presented here. First, in terms of theoretical and computational research, we have demonstrated that long-time kinetics of complex systems (all the examples discussed refer to growths lasting several minutes) can be predicted and
analyzed with atomic resolution. The accuracy of the method relies on an *ab initio* calibration and can be directly evaluated by the experiments, since it virtually reproduces equivalent growth conditions. As a relevant general outcome for materials science research, our investigations suggest that the processing of a defective system does not provide univocal upshots. Indeed, a non-pre-definite branching of the evolutive features in the 3C-SiC growths has been theoretically and experimentally evidenced. In particular, the stochastic simulation analysis, which was initiated exactly with the same defective APD system and with the same growing conditions, demonstrates without any doubts that this branching is “intrinsic” and not dependent on the possible experimental variance of the initial state.

For specialists in the growth processes of SiC materials, our results identify a critical source of other extended defects in the anti-phase boundary; therefore, the control of APBs and their eventual reduction is particularly important for achieving a good quality of the grown crystals. We have shown, by means of combined conductivity measurements with nanoscale resolution and quantum transport calculations, that the association of APBs and SFs strongly alters the surface conductive paths and can have a detrimental impact on the performances of vertical or quasi-vertical devices based on heteroepitaxial 3C-SiC on Si.

Finally, since the atomic mechanisms at the basis of extended defect evolution are accessible by the method, several particular aspects have been analyzed and discussed in Sec. IV. We have shown how the interaction of different types of defects often derives from the rearrangement of the atomic configuration of the defect portion emerging at the growing surface, whose morphology is in turn affected by the defect proximity. We would like to advise the reader that surface mediated interactions are an important, but partial aspect of the defect-defect correlation modes. Indeed, defect induced instabilities could give rise to “pure bulk” kinetics, which can be accessed by the *ab initio* calibrated KMCsL approach and will be investigated in the future.

**SUPPLEMENTARY MATERIAL**

See the supplementary material for the following: calibration details of the KMCsL code (Sec. S1) and KMC results for the epitaxial growth of an ideal 3C-SiC (001) substrate (Sec. S2); and details for the quantum transport calculations (Sec. S3). The movie SF-APB-generation-termination.mp4 shows the generation and termination of a couple of triple stacking faults from an APB during the 3C-SiC epitaxial growth along the [001] z direction.

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