

# Matching different symmetries with an atomically sharp interface: Epitaxial Ba<sub>2</sub>SiO<sub>4</sub> on Si(001)

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In this paper, we present a comprehensive investigation of the epitaxial growth of Ba<sub>2</sub>SiO<sub>4</sub> on Si(001), a system in which neither crystal symmetry nor lattice constants match in a simple manner. In addition, it has the potential to become the first crystalline high-*k* gate dielectric. We combined x-ray photoelectron spectroscopy, low-energy electron diffraction, and aberration-corrected scanning transmission electron microscopy (STEM) in order to optimize the epitaxial growth by molecular beam epitaxy. Our focus was on the formation of a high quality crystalline interface. The films were grown by a co-deposition method that requires no diffusion of Si from the substrate. An annealing temperature of 400 °C turned out to be sufficient to form chemically homogeneous films. However, crystalline films require an annealing step to 670–690 °C for the formation of the epitaxial interface necessary for breaking Si-O bonds. STEM confirms that the interface is atomically sharp and that a single layer of the silicate is changed to a (2 × 3) structure at the interface from the (2 × 1.5) bulk structure. Based on our experimental results, we propose a geometrical model for the epitaxial interface. The growth of films with an understoichiometric Si flux leads to the formation of a near-surface Ba silicide that does not restrict the epitaxial silicate growth.

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## I. INTRODUCTION

Ultrathin oxide films in combination with semiconductors can be viewed as prototype systems on which formation of interfaces, conditions of growth, and defect densities have been studied in detail [1–3]. They are of ever-increasing scientific and technological interest since they provide many possibilities of functionalization and application, such as the use as a gate insulator for field effect transistors [4], wave guides [5], barriers for tunnel junctions, and spin filters [6]. The interfaces of these materials can exhibit properties that are completely different from those of the corresponding bulk. For example, ferromagnetism or superconductivity can emerge at interfaces of materials that are nonmagnetic or nonconducting, respectively, in the bulk [6,7].

The interface between silicon surfaces and dielectric layers is of particular technological interest for the continued scaling of complementary metal-oxide semiconductor transistors. In order to achieve the best possible scaling using a planar transistor structure, the formation of an atomically sharp interface with an extremely low defect concentration, surpassing even the SiO<sub>2</sub>/Si interface, is required. The most promising pathway in this direction is the growth of crystalline gate oxides [8–10]. McKee *et al.* have demonstrated that epitaxial BaTiO<sub>3</sub> can be grown with very small trapped interface charge density ( $D_{it} < 10^{10} \text{ eV}^{-1} \text{ cm}^{-2}$ ) using a layer sequencing technique to manipulate the interface [11]. Nonetheless, a crystalline gate oxide that fulfills all of the requirements placed upon

high-*k* gate dielectrics [4] is still missing to date. In the case of BaTiO<sub>3</sub>, the band gap is too small for practical application. Epitaxial Ba<sub>0.7</sub>Sr<sub>0.3</sub>O layers, lattice matched to Si(001), are thermodynamically not sufficiently stable in contact with silicon and turn into silicates at temperatures above 400 °C [12].

Epitaxial Ba<sub>2</sub>SiO<sub>4</sub> on Si(001) is a candidate that could potentially fulfill all requirements for a crystalline gate oxide. Interestingly, this material was shown to grow as crystalline layers on Si(001) [13,14], although neither symmetry (tetragonal vs cubic) nor lattice constants match in a simple manner between the two systems. Ba<sub>2</sub>SiO<sub>4</sub> grows with its longest crystal axis perpendicular to the Si(001) surface. The lattice constants in the other two crystal directions are approximately equivalent to 2 and 1.5 times the Si lattice constant in the [110] direction, respectively [lattice constants of orthorhombic Ba<sub>2</sub>SiO<sub>4</sub>:  $a, b = 5.81, c = 10.21 \text{ Å}$  [15],  $S(001)$ :  $a = 3.84 \text{ Å}$ ].

Furthermore, barium silicates are thermodynamically and kinetically very stable. In particular, the reaction of the oxides SiO<sub>2</sub> and BaO to silicates is exothermic [16]. Generally, silicates are spontaneously generated from the oxides as shown in several examples in the past [13,17–19], and these silicate layers are stable up to their desorption temperature above 750 °C. The thermal stability is, thus, comparable to SiO<sub>2</sub>. Moreover, Ba<sub>2</sub>SiO<sub>4</sub> also fulfills other requirements necessary for high-*k* materials. Epitaxial films have shown a dielectric constant of  $22.8 \pm 0.2$ , band offsets to *p*-Si(001) of over 2 eV, and an acceptable leakage current of 3 mA/cm<sup>2</sup> at –1 V [14]. The only problem is that these films still featured a high density of interface traps of  $D_{it} = (1 \pm 0.5) \times 10^{13} \text{ eV}^{-1} \text{ cm}^{-2}$ . This interface property is closely related to the formation of

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the film itself for which diffusion of Si from the Si(001) substrate into a deposited BaO layer was used, which leads to an atomically rough interface, only works in the presence of concentration gradients of Si and, thus, may introduce local disorder and/or local variations of composition.

In order to avoid the diffusion problem, we used a co-deposition growth method in the present paper in which Ba and Si were evaporated simultaneously in an oxygen atmosphere. In the following, we show that precise control of the oxygen background pressure and the individual fluxes is a prerequisite for high quality  $\text{Ba}_2\text{SiO}_4$  thin films with an atomically sharp interface. Furthermore, we analyze the formation and structure of the epitaxial interface in order to understand how the lattice matching is achieved and develop an appropriate model.

## II. EXPERIMENTAL METHODS

$\text{Ba}_2\text{SiO}_4$  layers with thicknesses between 1 and 16 nm [4 and 63 monolayers (MLs)] were prepared under UHV conditions (base pressure  $8 \times 10^{-11}$  torr) on Si(001) samples (size  $6 \times 14 \text{ mm}^2$ ,  $p$  type, B doped, and resistivity 10–20  $\Omega \text{ cm}$ ) by molecular beam epitaxy.

The Si samples were cleaned *ex situ* with purified water ( $\rho \geq 18 \text{ M}\Omega \text{ cm}$ ), petroleum ether, acetone, and isopropanol in an ultrasonic bath for 15 min per step. Then, they were dipped in hydrogen fluoride solution (1%) until the oxide was completely removed (i.e., for approximately 30 s) as indicated by a hydrophobic surface, followed by a controlled reoxidation in  $\text{H}_2\text{O}_2$  solution (35%, 40 s). After each step, the samples were rinsed in purified water. In vacuum, the samples were degassed for more than 12 h at temperatures up to 600 °C and, subsequently, trained by heating to temperatures between 700 and 800 °C for a few minutes per step. Finally, the oxide film was removed by flash annealing to 1050 °C.

$\text{Ba}_2\text{SiO}_4$  films were grown by evaporation of Ba from a crucible and of Si from a rod, both heated by electron bombardment in an oxygen background pressure. Fluxes were controlled by a quartz crystal microbalance at the sample position, which was also used to calibrate relative thicknesses. The films were grown at room temperature (RT) in order to minimize oxidation of the Si surface and Si diffusion from the substrate. At the beginning of the growth, 0.5-ML Ba were first deposited before the Si flux was added. Subsequently, oxygen was introduced into the chamber, and the oxygen partial pressure was carefully increased to its final value within 30 s after the addition of the Si flux. The deposition rate was 1.1 ML/min. The films were annealed using direct current heating at temperatures between 400 and 700 °C for 30–40 min. The temperatures were determined with a pyrometer (minimum 300 °C) with emission coefficients adjusted to Si. In particular,  $\varepsilon = 0.574$  was used for  $T = 680$  °C. The optimum annealing temperature and oxygen background pressure will be discussed below. Annealing was always carried out in vacuum without oxygen. A new Si sample was used for each film.

The BaO film was grown similar to the  $\text{Ba}_2\text{SiO}_4$  films but without the Si flux. The  $\text{SiO}_2$  film was grown by thermal oxidation, and its thickness was measured by ellipsometry.

The  $\text{Ba}_2\text{SiO}_4$  films were investigated using a high-resolution low-energy electron-diffraction system (SPA-

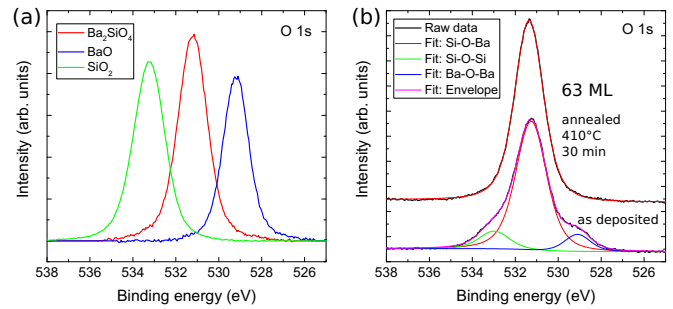


FIG. 1. Identification of chemical bond formation from the chemical shift of the O 1s signal in XPS: (a) Amorphous layers of BaO (blue line, 16-nm thick),  $\text{SiO}_2$  (green, 18-nm thick), and  $\text{Ba}_2\text{SiO}_4$  (red, 16-nm thick, annealed at 400 °C) on Si(001) are compared. The intensity of the  $\text{SiO}_2$  peak was not calibrated with respect to the others. (b) O 1s signal of a 16-nm- (63-ML-) thick  $\text{Ba}_2\text{SiO}_4$  film as deposited at room temperature (bottom curve) and after annealing at 410 °C (top curve). The deconvolution into the individual components is shown by colored lines as indicated in the inset.

LEED), x-ray photoelectron spectroscopy (XPS), and aberration-corrected scanning transmission electron microscopy (STEM). The SPA-LEED and XPS investigations were performed *in situ*. The XPS spectra were recorded with a SPECS Phoibos 100 spectrometer with an MCD-5 detector using an Al  $K\alpha$  source. The energy scale of the XPS system was calibrated using the Si 2p peak of a clean Si substrate (99.15-eV binding energy). For the evaluation of the XPS data, a Shirley background was first subtracted, and the O 1s peaks were then fitted with a LA(1.5,1.5,100)-type peak in CASAXPS [20]. This line shape is a convolution of a Lorentzian and a Gaussian and with the given parameters symmetric. It was carefully chosen since it perfectly describes the O 1s peaks of 16-nm-thick BaO and  $\text{Ba}_2\text{SiO}_4$  films, both of which feature single O 1s peaks.

In order to compare integrated XPS intensities, the measuring conditions have to be identical for both measurements, i.e., the instrument has to be in the exact same state and the sample has to be in the exact same position. We ensured the first part by keeping the instrument running when the measurements were performed on the same day, i.e., when comparing the same sample before and after annealing. When the measurements were performed on different days [Fig. 2(a)], the instrument was turned off between the measurements, however, all potentiometers were locked in place, and only on/off switches were used so that the applied voltages and currents remained the same for all measurements. Also, only measurements from one measurement period were compared with no bake-out of the system in between. Furthermore, the manipulator is equipped with micrometer screws that allow the sample to be reliably put in the same position for every measurement. The identical measuring conditions were ultimately confirmed by the background levels being the same for the compared measurements.

For the *ex situ* STEM investigations, the samples were capped with 1 ML of Si followed by 300 nm of Au. Two cross-sectional TEM lamellae in [110] and  $[1\bar{1}0]$  Si zone axes have been prepared using a focused ion-beam facility

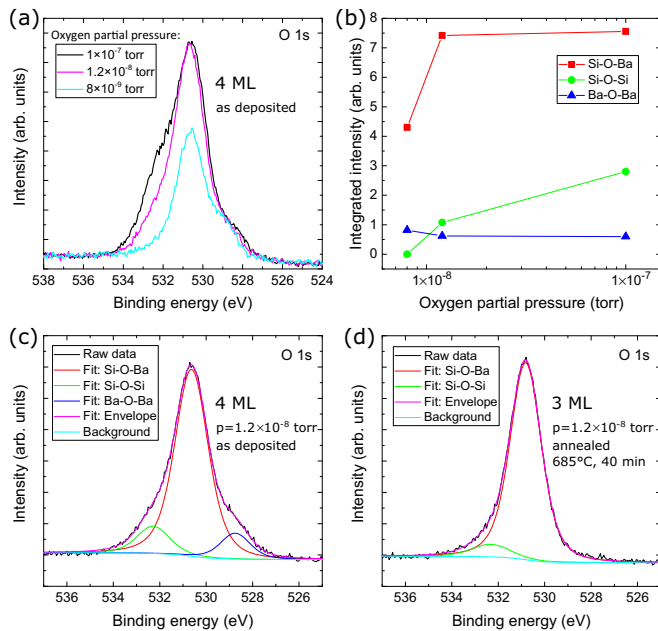


FIG. 2. Determination of the optimal oxygen background pressure: (a) XPS O 1s spectra for (approximately) 4-ML-thick Ba-silicate layers grown with different oxygen background pressures. The spectra were taken directly after growth at room temperature. The Si/Ba ratio was kept at the optimal value as determined in the context with Fig. 1. (b) Integrated intensities of contributions from different chemical bonding environments obtained from fits of the curves in (a). An example of this fit is shown in (c) for the optimal oxygen partial pressure of  $1.2 \times 10^{-8}$  torr. (d) O 1s peak after annealing the layer in (c) to 685 °C for 40 min.

with a final low-energy milling step in order to remove amorphized surface layers arising from previous milling steps. The two zone axes were used to obtain structural images of the epitaxial layer in two projections rotated by 90°. Scanning transmission electron microscopy was performed using a FEI Titan 80/300 STEM instrument equipped with an aberration corrector for the probe-forming lens [21]. The microscope was operated at 200 kV, and high-angle annular dark-field (HAADF) Z-contrast STEM images were acquired using a Fischione 3000 detector at a camera length of 135 mm. The final absolute thickness calibration of the Ba<sub>2</sub>SiO<sub>4</sub> films was performed using these STEM images.

### III. RESULTS AND DISCUSSION

#### A. Stoichiometry and chemical homogeneity

The exact ratio between the Si and the Ba fluxes was carefully adjusted for stoichiometric growth conditions since this ratio turned out to be non-self-limiting. The oxygen concentration, on the other hand, is self-limiting in the bulk as long as the oxygen partial pressure is high enough for a complete oxidation of the films. Under this condition, the oxygen content of the films is only determined by the Si and Ba fluxes as confirmed from the O 1s signal in XPS: Neglecting interface contributions, this XPS signal does not change at fixed Ba and Si fluxes even if the oxygen pressure is increased by one order of magnitude.

As it turned out, the line shape of the O 1s peak in XPS is ideal to identify any deviation from the Ba<sub>2</sub>SiO<sub>4</sub> stoichiometry and, thus, to calibrate the system. The relative chemical shift between the covalent bonds in SiO<sub>2</sub> and the ionic bonds in BaO is approximately 4 eV for O 1s as demonstrated in Fig. 1(a) where we compare the O 1s signals of BaO and SiO<sub>2</sub> with that of Ba<sub>2</sub>SiO<sub>4</sub>. All films are, at least, 16-nm thick so that the interface contribution to the signals is negligible. Conveniently, every O atom in the particular silicate compound we are interested in, Ba<sub>2</sub>SiO<sub>4</sub>, is bound to exactly one Si atom, i.e., no Si-O-Si bonding states are present [22], in contrast to other barium silicate compounds, such as BaSiO<sub>3</sub> [23] or BaSi<sub>2</sub>O<sub>5</sub> [24]. This results in a single O 1s peak for Ba<sub>2</sub>SiO<sub>4</sub>, which is located roughly in the middle of the SiO<sub>2</sub> and BaO peaks as also shown in the figure. This chemical oxidation state will be referred to as a Si-O-Ba bonding state in the following. The energy difference of about 2 eV between the peaks can easily be resolved by a standard XPS. Therefore, excess concentrations of Si or Ba lead to clearly visible shoulders at higher or lower binding energies, respectively. It is important to see these peaks as a representation of chemical bonding states and not of compounds since a shoulder at higher binding energy, in particular, can be either due to local SiO<sub>2</sub> or due to a barium silicate compound with a higher Si concentration than Ba<sub>2</sub>SiO<sub>4</sub>.

We first adjusted the Si and Ba fluxes to the required ratio of 1:2 using a quartz crystal microbalance located at a sample position by weighting the changes in frequency over time with the atomic masses. In order to confirm that we have obtained the correct Si:Ba atomic ratio in the deposited films, a 63-ML- (16-nm-) thick Ba<sub>2</sub>SiO<sub>4</sub> film was grown. Since the contribution of the interface to the XPS signal is negligible at this thickness, the peak shape is only determined by the Si to Ba ratio. The O 1s peaks after deposition at RT and after annealing at 410 °C for 30 min are shown in Fig. 1(b). Directly after deposition, the O 1s peak has two shoulders with approximately the same area. This demonstrates that the reaction at RT is incomplete so that local deviations from stoichiometry lead to various bonding characters and to only about an 80% preference for the Si-O-Ba bonding state. Only after annealing to 410 °C, which does not lead to any desorption, a single peak at the Si-O-Ba position is formed, i.e., the reaction is complete. As expected, this peak has approximately the same integrated intensity as the RT peak including the shoulders. This symmetric peak demonstrates that the correct deposition ratio for stoichiometric deposition was chosen. Furthermore, it shows that temperatures around 400 °C are sufficient for getting a *chemically* homogeneous film. This temperature, however, is too low to crystallize amorphous layers. In fact, all films starting with an amorphous interface layer turned out to remain amorphous after an annealing step to 400 °C.

#### B. Interface properties and oxygen pressure

In contrast to the bulk material, the interface between the silicate and the Si(001) surface is highly sensitive to the background oxygen pressure. On one hand, this is due to the oxidation of the Si(001) surface, which leads to the formation of amorphous SiO<sub>x</sub>. On the other hand, the Si wafer acts as a reservoir for Si so that the formation of Si-rich silicate

compounds is possible, irrespective of the Si flux. Both effects limit the crystalline quality of the deposited film. In order to achieve high quality epitaxial films with an abrupt interface, it is, therefore, absolutely essential to fine-tune the oxygen partial pressure to a value just above the saturation point for complete oxidation.

For calibration of the oxygen partial pressure, we again use the line shape of the O 1s peak in XPS, but this time, we investigate ultrathin films so that the interface is visible. For this purpose, we grew 4-ML-thick films at a rate of 1.1 ML/min in different oxygen partial pressures using the stoichiometric Si:Ba ratio determined above. The corresponding O 1s peaks and the integrated intensities of their individual components are shown in Figs. 2(a) and 2(b), respectively. Since the spectra were taken directly after growth at RT, the reaction is still incomplete, and, thus, all peaks feature a component at the BaO position. Nevertheless, the conclusion is still clear:  $1.2 \times 10^{-8}$  torr is the optimum oxygen partial pressure for this growth rate. For positive deviations from this optimum pressure, the concentration of O atoms in Si-O-Si bonding states is increased significantly, whereas the concentration of O atoms in Si-O-Ba bonding states stays nearly constant, indicating an oxidation of the silicon surface. At negative deviations, the main peak at the Si-O-Ba position is drastically reduced, indicating an oxygen deficiency for silicate formation.

The O 1s peaks for a 4-ML film grown at optimum conditions before and after annealing at 685 °C for 40 min are shown in Figs. 2(c) and 2(d), respectively, together with the deconvolution into the individual components. During the annealing process, about 1 ML of the film was desorbed as derived from the intensity of the Ba 3d and O 1s peaks in XPS so that the 4-ML film was reduced to only 3 ML after annealing. Even after annealing, the O 1s peak of the now 3-ML thick film still shows a small concentration of O atoms in Si-O-Si bonding states, unlike the peak of the 63-ML film. This concentration remains unchanged when the film is grown with a small Ba excess. Therefore, this signal is most likely due to interface bonds that are required between the film and the substrate and is not due to defects. Assuming that the O atoms in Si-O-Si bonding states sit in the monolayer closest to the interface and that this layer has the same oxygen concentration as the overlying layers, the Si-O-Si signal corresponds to a concentration of 1/4 ML. In order to correct for the damping of the signal by the overlying layers, we used an inelastic mean-free path of 2.072 nm calculated with the TPP-2M equation using the National Institute of Standards and Technology Electron Inelastic-Mean-Free-Path Database [25].

The important difference to all previous steps, however, is the appearance of a crystalline film after annealing. When using stoichiometric Ba and Si fluxes, crystallinity was only achieved when the oxygen background pressure during growth was close to the saturation point. Although lower oxygen pressures will not allow to obtain stoichiometric conditions, higher pressures lead to oxidation of the interface [see Fig. 2(b)] that prevents crystalline growth.

This behavior is in strong contrast to conditions of large Ba excess concentrations. Islam *et al.* [14] obtained crystalline Ba<sub>2</sub>SiO<sub>4</sub> thin films by Si diffusion from the substrate into

a BaO film (BaO contains twice as much Ba per O atom as Ba<sub>2</sub>SiO<sub>4</sub>) even though the oxygen background pressure during the growth of the BaO film was one order of magnitude above the saturation point determined in this paper at a comparable growth rate, i.e., a SiO<sub>x</sub> must have formed at the interface.

If Si is supplied by diffusion from the substrate, it is the ratio of Ba:O in the system that determines the silicate formation, which is always thermodynamically favorable to keep either SiO<sub>2</sub> or BaO in the system [16]. Under conditions of large Ba excess, Ba<sub>2</sub>SiO<sub>4</sub> is formed predominantly since it is the Ba-silicate compound with the highest Ba concentration. Therefore, as long as there is sufficient excess Ba in the system, a SiO<sub>x</sub> layer at the interface does not prevent the formation of crystalline Ba<sub>2</sub>SiO<sub>4</sub> since it can be incorporated into the Ba<sub>2</sub>SiO<sub>4</sub> film during reactive growth, thus, scavenging interfacial SiO<sub>x</sub> in favor of silicate formation. However, when we start with a stoichiometric Ba<sub>2</sub>SiO<sub>4</sub> layer and force the system to incorporate additional SiO<sub>x</sub> at the interface, the formation of silicate compounds with higher Si concentrations than Ba<sub>2</sub>SiO<sub>4</sub> will occur for thermodynamic reasons [16]. These compounds again hinder the epitaxial growth of Ba<sub>2</sub>SiO<sub>4</sub> due to their crystal structures that do not match with Si(001). We tested this hypothesis by growing films with a *reduced* Si flux, i.e., a BaO surplus. Indeed, when the Si flux was reduced by 50%, crystallinity could still be achieved even with an oxygen partial pressure 50 times higher than the saturation pressure from above.

### C. Formation of the epitaxial interface

As mentioned, the key to the growth of high quality crystalline films under stoichiometric Ba:Si ratios is the correct adjustment of the oxygen partial pressure. However, an annealing step at high temperatures close to desorption is still required to crystallize the films. The minimum temperature at which crystallinity was achieved is 640 °C, but the quality was still rather poor, and most films even remained amorphous at this temperature. An annealing temperature of 670–690 °C produced the best and, most importantly, reproducible results. At these temperatures, there was already a slight evaporation of the film as mentioned above, which drastically increased at temperatures above 690 °C. This reduction in thickness was most evident for very thin films, whereas for more than 40-ML-thick films, the material loss during annealing was negligible.

The LEED image of the same 3-ML film whose O 1s peak is shown in Fig. 2(d) is presented in Fig. 3(a). The LEED pattern originates from the superposition of two rectangular domains rotated by 90°. Their reciprocal lattice constants are very close to one-half and two-thirds, respectively, of the reciprocal lattice constant of the unreconstructed Si(001) surface. Thus, they correspond to the two shorter real-space lattice constants *a* and *b* of the Ba<sub>2</sub>SiO<sub>4</sub> crystal structure, whereas the *c* axis is oriented normal to the surface (*a* = 7.51, *b* = 5.81, *c* = 10.21 Å [15]). A schematic of this structure in reciprocal space is shown in Fig. 3(b). The same structure was found in Ref. [14].

As seen from the line profile in the horizontal direction through the center of Fig. 3(a), shown in Fig. 3(c), there

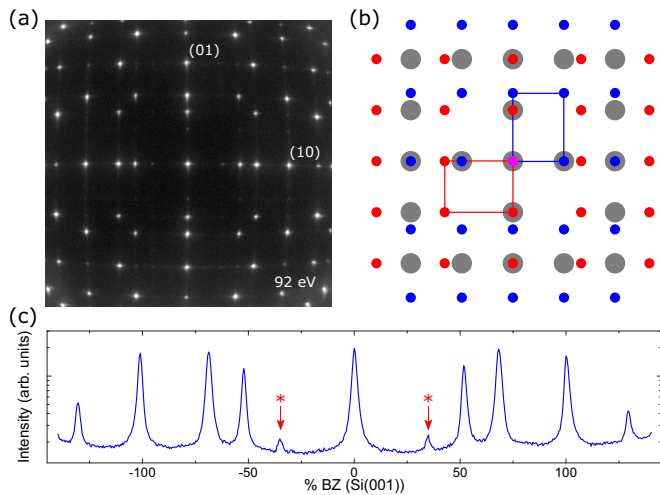


FIG. 3. LEED pattern of epitaxial  $\text{Ba}_2\text{SiO}_4$ : (a) LEED image of a 3-ML film after annealing to  $685^\circ\text{C}$  taken at 92 eV. The film was grown under optimal conditions as determined from Figs. 1 and 2. (b) Schematic of this pattern with two orthogonal rectangular unit cells (red and blue dots), demonstrating epitaxial growth with axes  $a$  and  $b$  of the silicate  $\parallel \text{Si}[1\bar{1}0]$ . Gray balls indicate the positions of the diffraction spots of the  $\text{Si}(001)$  substrate prior to silicate deposition. (c) Horizontal line scan through the center of (a). Spots marked by \* are not indicated in (b).

are some less intense spots, which correspond to a  $(2 \times 3)$  structure. These can be explained by a modulation with twice the lattice constant in the  $b$  direction close to the interface (see below). Since the value of  $2b$  is close to three times the silicon lattice constant, it is conceivable that the epitaxial growth leads to a doubling of the lattice constant in this direction close to the interface.

The HAADF  $z$ -contrast STEM images in Fig. 4 confirm that the interface between the crystalline  $\text{Ba}_2\text{SiO}_4$ , grown under the optimized conditions, and  $\text{Si}(001)$  is atomically sharp. Whereas the HAADF-STEM technique clearly shows Ba in the epitaxial layer, the Si and O atoms are not visible within the silicate layer due to their low atomic number compared to Ba. Projections along  $\text{Si}[110]$  are shown. As expected from Fig. 3, two orientations of  $\text{Ba}_2\text{SiO}_4$  can be seen in this projection, which are depicted in Figs. 4(a) and 4(b). As can be seen there, the bulk crystal structure of  $\text{Ba}_2\text{SiO}_4$  is maintained up to the penultimate layer at the interface, i.e., only one layer at the interface is geometrically modified, neglecting relaxations, in order to form the bonds to the  $\text{Si}(001)$  substrate. The Ba atoms that are shifted from their bulk positions in  $\text{Ba}_2\text{SiO}_4$  are depicted in dark green in the overlaid crystal structures. This rearrangement of the Ba atoms at the interface results in a period doubling at the interface in the  $b$  direction as compared to the  $\text{Ba}_2\text{SiO}_4$  bulk structure in this layer as visible in Fig. 4(a). Furthermore, there are only three Ba atoms instead of four within this period. This confirms that the  $(2 \times 3)$  structure observed in LEED (see Fig. 3) is, indeed, due to the epitaxial interface. In the perpendicular direction [see Fig. 4(b)], the size of the unit cell at the interface is the same as in the bulk, again in agreement with the LEED data, but clear shifts of Ba atoms at the interface can be seen.

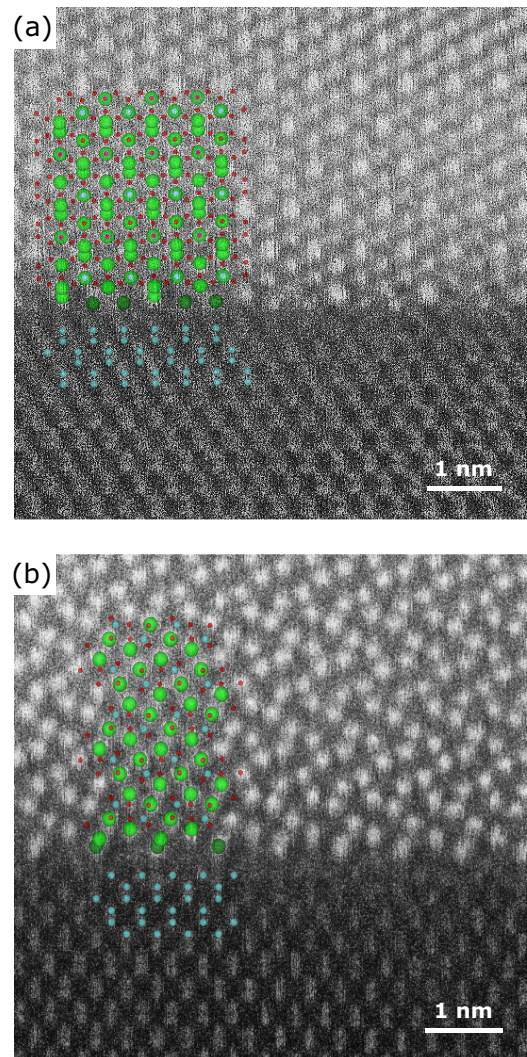


FIG. 4. HAADF  $z$ -contrast STEM images: (a)  $b$  and  $c$  projections and (b)  $a$  and  $c$  projections of the  $\text{Ba}_2\text{SiO}_4$  crystal structure. The bright spots are Ba atoms. The bottoms of both images show the  $\text{Si}(110)$  plane. The images were taken from a 44-ML  $\text{Ba}_2\text{SiO}_4$  film, which was prepared by first growing a crystalline 3-ML film under optimized conditions. Then, an additional 41 ML were deposited at RT, and the film was annealed at  $680^\circ\text{C}$  for 40 min. The crystal structures overlaid on the images were drawn with VESTA [26] based on Refs. [22,27]. Green: Ba; blue: Si; red: O. The Ba atoms at the interface that deviate from the  $\text{Ba}_2\text{SiO}_4$  bulk structure are depicted in dark green.

Based on the information from XPS, LEED, and STEM and the chemical boundary conditions that Ba atoms are divalent and that the Ba-O bonds are ionic, we have designed a plausible model of the interface, which is depicted in Fig. 5. The model has six O atoms per unit cell in Si-O-Ba bonding states in the interface layer. The  $(2 \times 1)$  reconstruction of the  $\text{Si}(001)$  surface is, for the most part, still present [see Fig. 5(b)]. The model only deviates from the  $\text{Si}(001)$   $(2 \times 1)$  reconstruction at two Si atoms per interface unit cell. Here, the bonds are elongated by an O bridge. Thus, these Si atoms can get closer to the Ba atom in the second silicate layer so that ionic bonds can be formed. The concentration of O

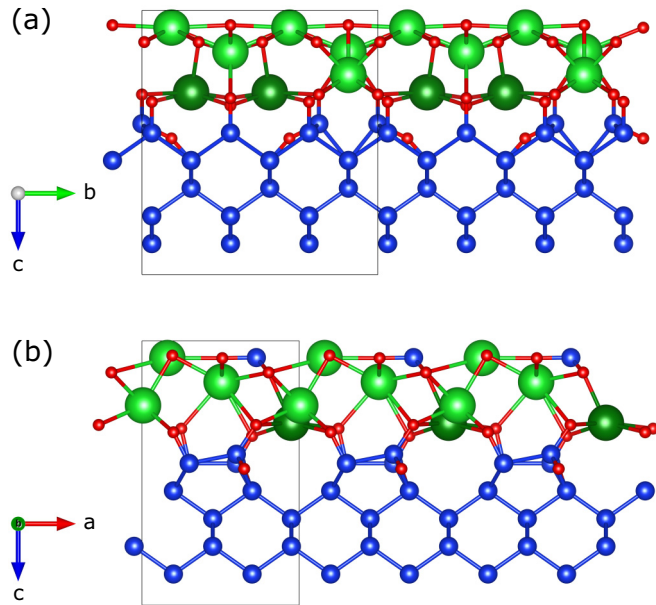


FIG. 5. Proposed interface model: (a)  $b$  and  $c$  projections and (b)  $a$  and  $c$  projections with respect to the  $\text{Ba}_2\text{SiO}_4$  crystal structure. Green: Ba; blue: Si; red: O. The structure was drawn with VESTA [26]. In this model, the  $\text{Ba}_2\text{SiO}_4$  bulk structure [22] was slightly stretched so that the  $a$  and  $b$  axes fit exactly 1:2 and 2:3, respectively, to the lattice constant in the [110] direction of the Si lattice [27], i.e., small lattice mismatches were ignored. The Ba atoms at the interface that deviate from the  $\text{Ba}_2\text{SiO}_4$  bulk structure are depicted in dark green.

atoms in Si-O-Si bonding states, two per interface unit cell, is equivalent to  $1/4$  ML of the  $\text{Ba}_2\text{SiO}_4$  bulk structure and, thus, in agreement with the XPS results shown in Fig. 2(d). The pseudo- $(2 \times 1)$ -reconstruction results in a 1:1 match in the  $a$  direction between  $\text{Ba}_2\text{SiO}_4$  and the Si surface.

We now deal with the question why the high-temperature annealing step is necessary for the formation of crystalline layers. Even though no long-range diffusion is necessary to form the silicate since conditions very close to the perfect stoichiometry were chosen, the formation of crystalline layers still requires the same high-temperature step as under conditions of silicate formation by Si diffusion [14]. It is clear from the LEED images that the crystallization has to start at the interface since the film is perfectly aligned with the substrate and there is no rotational disorder. Moreover, when we deposited an additional 4 ML on a crystalline 3-ML film, the film could already be crystallized by annealing to  $400^\circ\text{C}$ . The crystalline quality was comparable to that of the initial film. We tested this method up to a thickness of 63 ML and always obtained crystalline layers at  $400^\circ\text{C}$ , although with slightly decreasing crystal quality for the thickest films as evident from the spot profile in LEED, which could be improved after annealing to higher temperatures. Hence, the high annealing temperature is only needed to form the epitaxial interface.

An adjustment of the stoichiometry at the interface cannot be the reason for the requirement of the high temperature since Si is already sufficiently mobile at lower temperatures. As an example, we investigated the reaction of a 1-nm-thick BaO film with Si by diffusion from the Si bulk. Annealing at  $400^\circ\text{C}$

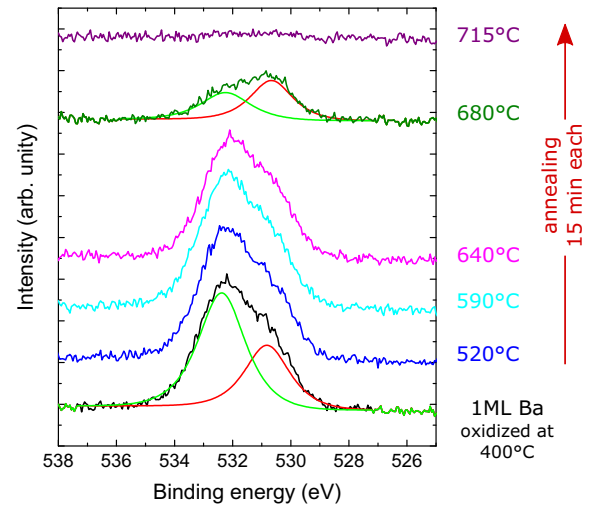


FIG. 6. Effects of high-temperature annealing on the interface: A 1-nm-thick Ba film on Si(001) was oxidized in an oxygen partial pressure of  $5 \times 10^{-8}$  torr for 10 min at  $400^\circ\text{C}$  and, subsequently, annealed at increasing temperatures for 15 min per step. The O  $1s$  spectra after oxidation and after each annealing step are shown. For the spectra after oxidation and after annealing to  $680^\circ\text{C}$ , the deconvolution into the individual components (Si-O-Si, green and Si-O-Ba, red) is also shown. The spectra were shifted for better visibility.

for 30 min was sufficient to turn this film completely into a silicate.

The high temperature is also not needed to break the original  $(2 \times 1)$  reconstruction of the Si(001) surface since it is possible to produce crystalline films with an oxygen partial pressure during growth that is more than one order of magnitude above the saturation point if the additional oxygen concentration at the interface is compensated by lowering the Si flux and thereby indirectly lowering the oxygen content of the bulk (see Sec. III B). In this case, the  $(2 \times 1)$  reconstruction at the interface is definitely destroyed due to the oxidation of the Si(001) surface. However, the high-temperature step is still needed to crystallize these films.

In order to better understand the effect of the high-temperature annealing step on the interface, we deposited one monolayer of Ba on a Si(001) surface and oxidized it at  $400^\circ\text{C}$ . Then, we annealed it at increasing temperatures for 15 min per step. The corresponding O  $1s$  peaks are shown in Fig. 6. Directly after oxidation at  $400^\circ\text{C}$  the O  $1s$  peak consists of two components at the Si-O-Si and Si-O-Ba locations with a ratio of roughly 2:1. The annealing steps at  $520^\circ\text{C}$  and  $590^\circ\text{C}$  have no effect on the peak. At  $640^\circ\text{C}$ , the Si-O-Si component is already slightly reduced. At  $680^\circ\text{C}$ , the Si-O-Si component is reduced significantly, whereas the Si-O-Ba component is only slightly reduced. Finally, at  $715^\circ\text{C}$ , the oxygen is completely desorbed.

The temperature range at which the Si-O-Si peak is reduced lines up perfectly with the temperatures needed for crystallization of the  $\text{Ba}_2\text{SiO}_4$  films. In particular, the temperature at which the reduction of the Si-O-Si peak sets in is exactly the lowest temperature at which crystallization was observed. These findings indicate that the high temperatures

required for the formation of the epitaxial interface are needed to break interfacial Si-O bonds for O atoms with two bonds to Si, possibly for O atoms with two bonds to the substrate. Under our conditions, the transient formation of this type of bonds could not be avoided, not even by starting with an amorphous monolayer of Ba.

If we assume the proposed interface model in Fig. 5 to be correct, the high temperatures can be explained by the need to (largely) restore the  $(2 \times 1)$  reconstruction of the Si(001) surface that was destroyed by oxidation during the silicate growth and by the need to reduce the concentration of Si-O-Si to 1/4 th of a monolayer.

#### D. Robustness of crystalline growth: Ba excess concentrations

Finally, we tested the robustness of the interface when modifying the composition of ultrathin layers and its consequence on crystalline silicate growth. Some 4-ML films (given as the necessary Ba amount) were grown, again at RT, with a reduced Si flux of  $x = 0.37, 0.57, 0.81$ , and 1 times the flux needed for the exact stoichiometry of  $\text{Ba}_2\text{SiO}_4$ . As expected, the O 1s peaks of the Si deficient layers (not shown here) show a larger BaO fraction than the stoichiometric film that increases with decreasing  $x$ . After annealing at  $685^\circ\text{C}$  for 40 min, all films turned completely into silicate due to Si diffusion from the substrate. Although the thickness of all films was reduced during the annealing, the reduction in thickness increased with increasing Si deficiency. A possible reason is the lower initial amount of O atoms per Ba atom and desorption of Ba during annealing (there was no oxygen background pressure during annealing). The thicknesses of the films after annealing were 1.6, 2.3, 2.5, and 3 ML for  $x = 0.37, 0.57, 0.81$ , and 1, respectively, as derived from the Ba 3d and O 1s peaks in XPS.

The LEED images for  $x = 0.37$  and  $0.81$  as well as the line scans through the (00) and (10) spots after annealing for all films are shown in Fig. 7. As expected from the discussion in Sec. III B, the varying concentrations of Si relative to Ba do not suppress the formation of crystalline silicate layers. The interesting part, however, is the appearance of a faint but quite sharp  $(4 \times 2)$  structure for  $x < 1$  that we assign to the formation of a near-surface Ba silicide, following Ref. [28]. The high intensity of the  $(4 \times 2)$  peaks for  $x = 0.37$  is most likely due to the smaller film thickness. The absence of the  $(4 \times 2)$  structure for  $x = 1$ , however, seems to be real since the presence of the  $(2 \times 3)$  structure (spots marked with \*'s) associated with the epitaxial interface means that the interface is still visible in LEED at this thickness. Moreover, neither a  $(4 \times 2)$  structure nor Ba diffusion into the substrate was observed in STEM, see Fig. 4. Therefore, the appearance of the  $(4 \times 2)$  structure seems to be coupled with a surplus of Ba or BaO in the insulator film. Although most of the Ba seems to desorb during the annealing process, there is also the possibility of diffusion of Ba to the Si interface and the subsequent formation of silicide [28–30].

Crystalline silicate formation turns out to be quite robust with respect to these variations of the interface. Although a sharp  $(2 \times 1.5)$  structure appears under all conditions in LEED, the disappearance of the  $(2 \times 3)$  structure for  $x = 0.57$  and  $0.37$  indicates that the silicide distorts the period doubling.

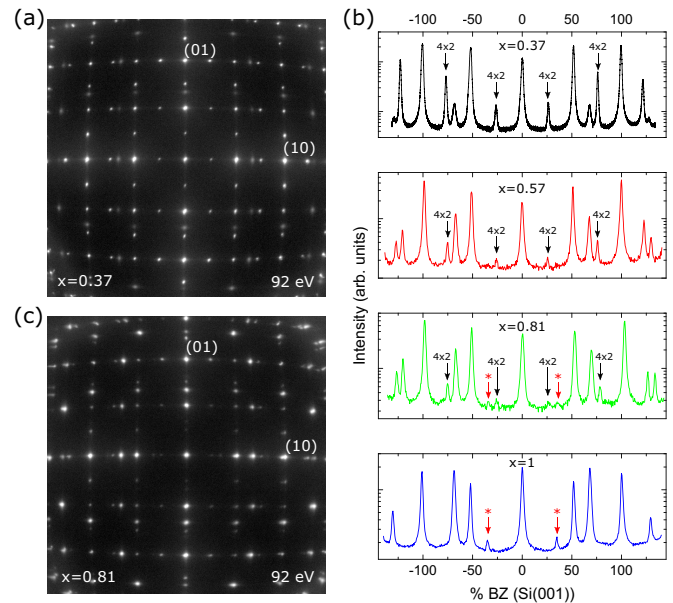


FIG. 7. Effects of an understoichiometric Si flux on the interface: 4-ML films were grown with a reduced Si flux and, subsequently, annealed to  $685^\circ\text{C}$ .  $x$  denotes the normalized Si flux with  $x = 1$  corresponding to the flux needed for the exact stoichiometry of  $\text{Ba}_2\text{SiO}_4$ . The LEED images for  $x = 0.37$  and  $0.81$  are shown in (a) and (c), respectively. The horizontal line scans through the center of the LEED patterns for  $x = 0.37, 0.57, 0.81$ , and  $1$  are shown in (b).

This does not necessarily mean that the local structure is changed. Only long-range order is distorted.

#### IV. CONCLUSIONS

We have demonstrated that epitaxial  $\text{Ba}_2\text{SiO}_4$  thin films can be grown on Si(001) with an atomically sharp interface and investigated the formation of this interface. In order to match  $\text{Ba}_2\text{SiO}_4$  with its  $(2 \times 1.5)$  structure to Si(001), a  $(2 \times 3)$  structure is needed at the interface. However, the system is flexible enough so that this change in crystal structure from the  $\text{Ba}_2\text{SiO}_4$  bulk structure is limited to only one layer at the interface. Considering the  $\text{Ba}_2\text{SiO}_4$  crystal structure, the cause for this flexibility is most likely the undirected Ba-O bonds that hold the  $\text{SiO}_4$  tetrahedra together. The formation of the epitaxial interface neither requires a special surface preparation nor requires an intentional lattice matching. Only well-optimized growth conditions are needed. The stability of this system is further demonstrated by the fact that the crystalline growth is not prevented by silicide formation at the interface.

The key to the successful growth of crystalline films in this paper was the precise determination of the saturation point for the oxygen background pressure based on the line shape of the O 1s peak in XPS. This method might also be helpful for the growth of other oxide species on silicon since it is, according to our experience, more precise than methods based on the frequency change in a quartz crystal microbalance.

As mentioned in the Introduction,  $\text{Ba}_2\text{SiO}_4$  has many attractive properties for a high- $k$  dielectric and may serve as a crystalline high- $k$  gate insulator. For this purpose, the

precise definition of the epitaxial interface is a necessary and important step. Furthermore, the proposed interface model can serve as a starting point for theoretical investigations. This paper may also provide a basis for controlled manipulation of the epitaxial interface.

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