I. INTRODUCTION

One-dimensional systems have been extensively studied over the last decades due to their intriguing physical properties and potential applications in nanometer-scale devices [1–7]. A bottom-up approach based on self-assembly on nanotemplates represents an attractive method for fabricating one-dimensional structures. The regular arrangement of the atomic steps of vicinal semiconductor surfaces makes them perfect templates for the growth of one-dimensional systems, e.g., nanowires. Here, we report on the atomic structure of the vicinal Si(111) surface with 10° miscut investigated by a joint frequency-modulation scanning force microscopy (FM-SFM) and ab initio approach. This popular stepped surface contains 7 × 7-reconstructed terraces oriented along the Si(111) direction, separated by a stepped region. Recently, the atomic structure of this triple step based on scanning tunneling microscopy (STM) images has been subject of debate. Unlike STM, SFM atomic resolution capability arises from chemical bonding of the tip apex with the surface atoms. Thus, for surfaces with a corrugated density of states such as semiconductors, SFM provides complementary information to STM and partially removes the dependency of the topography on the electronic structure. Our FM-SFM images with unprecedented spatial resolution on steps coincide with the model based on a (7 7 10) orientation of the surface and reveal structural details of this surface. Two different FM-SFM contrasts together with density functional theory calculations explain the presence of defects, buckling, and filling asymmetries on the surface. Our results evidence the important role of charge transfers between adatoms, restatoms, and dimers in the stabilization of the structure of the vicinal surface.

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II. METHODS

Both sample preparation and experiments were carried out in an ultrahigh vacuum (UHV) chamber with a base pressure of less than 3 × 10⁻⁸ Pa. Stripes of low n-doped Si(111) (phosphorus, \( \rho = 1–10 \) Ωcm, Virginia Semiconductor) and an inclination angle of 10 ± 0.5° towards the [112] direction were used. The silicon stripes were cleaned in a diluted
aqueous HF solution prior to loading into the UHV chamber. The sample was resistively heated by direct current with the current direction parallel to the steps on the vicinal Si(111). The surface was prepared by several short flashes to 1420 K, the last flash was proceeded by a fast ramp down to 1200 K, followed by a slower cooldown. The silicon sample was then transferred to a variable-temperature scanning force microscope (Omicron NanoTechnology GmbH, Germany) equipped with Nanosensors cantilevers (Switzerland) and a Nanonis phase-locked loop electronics (SPECS, Switzerland). All measurements were performed in the dynamic frequency modulation (FM) mode. Topographical imaging was carried out at constant frequency shift using sputter-cleaned silicon cantilevers with a force constant of 30-50 N m⁻¹, and a free resonance frequency of 270–300 kHz. Some of the cantilevers had a platinum-iridium-coated silicon tip. The long-range electrostatic interaction was minimized by applying a voltage that compensated the contact potential difference between the tip and the sample. For characterizing the FM-SFM images, the normalized frequency shift \( \gamma = \Delta f / k A^{1/2} f_0^{-1} \) has been used. In order to preserve the tip, the images were obtained scanning almost parallel to the [110] direction where the corrugation of the surface is the lowest. Since all measurements were performed at room temperature, the images (particularly in the y axis, the slow scan direction) are influenced by experimental drift. For the discussion of the structure, this drift has been compensated in Figs. 2 and 4. These images have been additionally slightly rotated for clarity.

The electronic structure calculations were performed within density functional theory (DFT) and the Purdue, Burke, and Ernzerhof (PBE) generalized gradient approximation [16], as implemented in the Vienna ab initio simulation package (VASP) [17]. The Brillouin zone (BZ) integrations in the electronic structure calculations were done using uniform meshes, equivalent to 224 points for the \((1 \times 1)\) surface unit cell. The starting structures were prepared based on the experimental observation of a 16-fold lateral periodicity [2], the model introduced by Teys et al. [2], and variations thereof. Due to the mismatch between the sevenfold periodicity of the terraces and the twofold periodicity of the dimer rows, the smallest possible unit cell size parallel to the step edges that does not contain any obvious defects has a 14-fold periodicity. Thus all calculations were performed within a \((16 \times 14)\) surface unit cell. The surface has been modeled by a slab containing four Si bilayers along the (111) direction, resulting in 2302 atoms per unit cell. The bottom Si bilayer was frozen at the equilibrium DFT lattice constant with the dangling bonds terminated by hydrogen. To avoid a spurious interaction between periodic images along the surface normal, a vacuum distance of 50 Å between the surface and the bottom layer of its periodic image has been employed. This vacuum distance has been used in conjunction with a dipole correction by introducing a step discontinuity inside the vacuum region which cancels out the surface dipole [18]. Forces were relaxed below a threshold of 0.01 eV Å⁻¹.

III. RESULTS AND DISCUSSION

Figure 1(a) shows an FM-SFM image of the silicon surface after preparation. The surface consists of large terraces, two of them can be distinguished in Fig. 1(a). Each of these large terraces contains periodically spaced steps with the height of three interplanar (111) distances and Si(111)-7 × 7 reconstructed areas. The separation of the two large terraces in Fig. 1(a) corresponds roughly to the height of two periodic steps, as shown in the profiles of Fig. 1(c). Such large terraces appear when the miscut angle of the crystal does not exactly correspond to 10°. In general, we avoid scanning with the fast axis parallel to the step edges by rotating the scan direction by 5°. Still, sometimes the tip apex becomes unstable, as it can be seen in Fig. 1(a), where a jump occurred in the middle of
the scan over the large step edge between the large terraces. In this image, silicon atoms are resolved in the upper and the lower large terraces. A magnification of the small region marked in Fig. 1(a) is displayed in Fig. 1(b). A unit cell of the $7 \times 7$-reconstructed part is indicated by two triangles. Large protrusions are observed on the step edges of the flat $7 \times 7$-reconstructed terraces [an example is indicated with a white arrow in Fig. 1(b)].

All SFM images in this paper are oriented like Fig. 1. This orientation implies that considering the vicinal Si(111) surface as an infinite stair, the upper part of the image corresponds to the lower steps, and the lower part of the image to the higher ones. This vicinal surface is inclined towards the $[\bar{1}\bar{1}\bar{2}]$ direction. The $[\bar{1}\bar{1}\bar{2}]$ direction is parallel to the surface of the $7 \times 7$-reconstructed part. We do not directly show such direction in the images but a projection of it. We have decided to mark it in gray on the images axes as a guide to the eye.

In Fig. 2, a close look into the atomic structure of this vicinal surface is shown. Figures 2(a) and 2(b) display two different FM-SFM images obtained with the same cantilever. Figure 2(a) shows the typically observed contrast, where the adatoms of the $7 \times 7$-reconstructed surface and the adatoms of the steps are the most protruding features. This contrast is also the most presented SFM contrast for the flat surface in the literature, and has been explained by a chemically reactive tip [19,20]. This image coincides almost one to one with the empty-states STM image [21] described in Ref. [2]. Due to changes of the tip apex during scanning, in addition, the image shown in Fig. 2(b) was obtained. In this other contrast, the restatoms of the $7 \times 7$ become prominent, therefore we named it here, the restatom contrast. This kind of contrast has previously been reported on the flat surface, and it has been explained by electrostatic interactions between tip and surface [22,23]. Thus the tip change may involve the loss of one or more reactive atoms of the tip apex. Alternatively, the tip change could be dominated by a change in local contact potential difference leading to a local effective bias, and therefore, a change in the contrast. Such a bias dependence of the Si(111)-$7 \times 7$ images is known from the literature [24]. The different contrasts highlight different features of the surface, providing supplementary information about its configuration.

In order to discuss the atomic arrangement of the steps in more detail, we performed DFT calculations for the Si(7 7 10) surface within the full ($16 \times 14$) surface unit cell. Although the experimentally determined 16-fold periodicity perpendicular to the step edges [2] strongly constrains the structural possibilities in the modeling, there are still several options how the $7 \times 7$ terraces and their partial counterparts are interfaced to the step edges and extra rows. Figure 3(a) shows the top and side views of the lowest energy structure obtained from our calculations. The calculations reveal several interesting structural details in addition to the experimental data.

In the following, a detailed analysis of the atomic structure of the triple step will be presented. For this purpose, the SFM images in Fig. 2(a) (normal contrast) and Fig. 2(b) (restatom contrast) will be described and compared to the resulting calculated structure and the STM data reported by Teys et al. [2,10]. We will start at the upper $7 \times 7$ terrace and move down along the $[\bar{1}\bar{1}\bar{2}]$ direction, layer by layer. For the discussion, a schematic representation of the vicinal surface has been included in Fig. 3(b). In the model, the terraces will
FIG. 3. (a) Top and side views of the structural model obtained from DFT calculations performed in a (16 × 14) supercell. RA1, RA2, and RB denote the restatoms of the surface. (b) Schematic representation of the top view of the vicinal Si(111) surface. The blue rectangle indicates the (16 × 14) supercell shown in (a). At the right side, an experimental SFM profile over the surface is included [partially shown in Fig. 4(a)]. An analog profile to this SFM profile has been indicated by a black dashed line in the scheme. An adapted side view of the structural model has been plotted next to the experimental profile for clarity. The adatoms and restatoms of the 7 × 7 terrace, A2, and R rows are plotted as black circles. Every atom of the parallel dimers (D∥3) is plotted as a red circle. In D⊥3, the upper atoms are plotted as dark blue circles whereas the lower atoms are plotted as light blue circles. Each atom of the ZR is plotted as a grey circle. A Si(111)-7 × 7 unit cell is marked with two triangles.

be denoted with Greek letters and the layers with the Latin letters. The blue rectangle indicates the (16 × 14) supercell shown in Fig. 3(a). In Fig. 2(a), a (16 × 14) supercell has been indicated as well. To enhance the features on the triple step, we have additionally applied a line subtraction fit to Fig. 2 and the resulting images are displayed in Fig. 4. For clarity, we have also included a table in Ref. [25], where the most important features of the distinct atomic species are summarized.

The terraces are formed by one cell of reconstructed 7 × 7 silicon (as indicated by two triangles in the figures), and an additional row of atoms, R row, located at the bottom of the triple step. This additional row makes the width of the terrace slightly larger than one 7 × 7 unit cell. The R row resembles a partial 7 × 7 unfaulted half cell. Each adatom and restatom of the 7 × 7 reconstruction, and each R row atom has one dangling bond (DB) pointing upwards, i.e., perpendicular to the terrace surface [plotted as a black circle in the model of Fig. 3(b)]. As previously mentioned, in the normal contrast, Figs. 2(a) and 4(a), the higher topographic protrusions imaged are the adatoms and the R atoms, with a weak contribution of the restatoms, as in empty-states STM images. In the restatom contrast shown in Figs. 2(b) and 4(b), the adatoms are still topographically higher but they are imaged as smaller cloudy protrusions, whereas the restatoms become more noticeable. This contrast is similar to the filled-states STM image, but with the significant difference that the R row has almost disappeared in the SFM image.

On the edge of the terraces, the last row of silicon adatoms, denoted as A3 (the index indicates the layer of the triple step), is accompanied by a row of parallel dimers (D∥3). Every atom of the D∥3 dimer has a DB perpendicular to the step surface, pointing slightly upwards and tilted apart from each other [red circle in Fig. 3(b)]. In general, the D∥3 atoms are seen as protrusions at the side of the A3 atoms. In the normal contrast, sometimes we are able to distinguish both atoms of the dimer [see Fig. 4(a), inside the dashed-line ovals]. In the restatom contrast, Figs. 2(b) and 4(b), the D∥3 dimers display a similar contrast to the restatoms.

The lower layer (with index 2) is formed by a row of adatoms (A2). Each A2 adatom has a DB pointing upwards, i.e., perpendicular to the terrace surface (black circle in Fig. 3(b)). The A2 atoms are located either opposite to the A3 atoms, as indicated with white dashed lines in Figs. 2(a) and 2(b), and depicted at step α of Fig. 3(b); or opposite to the gaps between the A3 atoms as indicated with white solid lines in Figs. 2(a) and 2(b), and depicted at step β of Fig. 3(b). Often defects are observed in the A2 row, in particular at the corner vacancies of the 7 × 7 surface, these are denoted as RD (row defect). At these sites there are no D∥3 dimers. Such RDs appear due to the mismatch between the twofold periodicity of the dimer rows and the sevenfold periodicity of the terraces. RDs are therefore, not true defects but part of the reconstruction. At the RD position, the adatom shifts towards the step, as indicated in Figs. 2(a) and 3(b). This shift uncovers the restatom in the second layer close to RD denoted as F. The restatom F is indicated with a white arrow in Fig. 4(a). In general, the appearance of the adatoms of the terrace, the A2 row, and R row is similar: in the normal contrast as prominent protrusions and in the restatom contrast as cloudy protrusions.

Below the row of A2, there are rows ascribed to perpendicular and parallel dimers (D⊥3 at the step, and D∥3 at the first layer), as indicated in Figs. 2 and 3. In the D⊥3 dimer, the
FIG. 4. A line subtraction fit was applied to the atomically resolved FM-SFM images of Fig. 2 (slightly larger area) for enhancing the features of the triple step structure. (a) Normal contrast. (b) Restatom contrast. For comparison with Fig. 2, the same unit cells are marked with two triangles. The white dashed line indicates partially the experimental SFM profile shown in Fig. 3(b). Black dashed lines are guidelines of the triple step structure. Dashed-line ovals indicate $D_{\perp}$ dimers where the signal of the two atoms is split. The solid-line rectangle indicates a location where only the upper atoms of the $D^+_{\parallel}$ dimers that are located opposite to the gaps between the $A_2$ atoms are imaged. The dashed-line rectangle indicates a location where all upper atoms of $D^+_{\perp}$ are imaged. Black dashed-line arrows indicate adsorbates at step edges of the second layer ($A_2$). The white arrow indicates a restatom in the second layer close to RD, denoted as F, that becomes visible due to the shift of RD towards the step. The black V marks a defective 7×7 half cell. The white double arrow indicates a defective triple step: the distance between the two 7×7 flat terraces is larger than the one usually expected for the triple step. Images size 14×20 nm².

two atoms are asymmetric: the upper atom features one DB perpendicular to the surface pointing slightly upwards [dark blue circle in Fig. 3(b)], whereas the lower atom is saturated (no DB) [light blue circle in Fig. 3(b)]. Consequently, we expect to predominantly image the upper atom. In Figs. 2 and 4, for both contrasts, we mainly image $D^+_{\perp}$ that are located opposite to the gaps between the $A_2$ atoms, an example is indicated with a solid-line rectangle in Fig. 4(a). In our FM-SFM images, only in the lower part of Fig. 4(a) (inside the dashed-line rectangle) all upper atoms of $D^+_{\parallel}$ are imaged. Still, in the images, many dimers of the $D^+_{\perp}$ rows seem to be missing. Either these dimers are absent, or the interaction of the dimers with the tip is not strong enough to image them. The low interaction with the tip may be due to their position close to the more protruding $A_2$ atoms. In the $D^+_{\parallel}$ dimer, as in the $D^+_{\perp}$ dimer, each atom has a DB perpendicular to the step surface tilted apart from each other [red circle in Fig. 3(b)]. Also in the $D^+_{\parallel}$ row some dimers are missing in the SFM images. Again here, either the dimers are absent or the bond formation with the tip is sterically hindered.

The environment of the $D^+_{\perp}$ and $D^+_{\parallel}$ dimers with the step oriented at the [100] direction is similar to the one of the dimers on the Si(100)-(2×1) surface [26]. In the Si(100)-(2×1) dimers, charge transfer from the lower DB to the protruding DB causes buckling [27,28]: the lower DB is empty ($\delta^+$) and the upper one is fully filled ($\delta^-$), similar to the case of a zwitterion. In our surface, however, because the $D^+_{\parallel}$ has only one DB, the partner-DB is at one of the atoms of the $D^+_{\perp}$ dimer. In Fig. 5, the charge transfer between $D^+_{\perp}$ and $D^+_{\parallel}$ dimers is sketched in similarity to the one observed on the Si(100)-(2×1) surface. Likewise, Fig. 5 displays the charge transfer between both atoms of the $D^+_{\parallel}$ dimer, evidencing the interconnection of the two adjacent $D^+_{\parallel}$ and the corresponding $D^+_{\perp}$ dimer [10]. $D^+_{\perp}$ dimers located between two $A_2$ adatoms prefer a more upright orientation, whereas the ones directly opposite to an $A_2$ adatom assume a flatter orientation. The $D^+_{\perp}$-$D^+_{\parallel}$ buckling in turn induces a corresponding tilt in the directly adjacent $D^+_{\parallel}$ row. Interestingly, due to the
mismatch between the twofold periodicity of the dimer rows and the sevenfold periodicity of the terraces, there can be no defect-free configuration within the buckling orientation of the D_2-D_1 dimers. The row defects (RDs) accommodate this mismatch, making the two D_2 dimers between the A_2 adatoms adjacent to the RD to be both preferentially oriented in the upright configuration. However, these two D_2 dimers are directly adjacent to each other, locally interrupting the twofold order of the buckled dimer row. Likewise, in the D_1 dimers, there is charge transfer between both atoms in analogy to the D_1 dimers, thus the D_1 dimers prefer a similar tilt angle than the D_2 dimers.

Finally, in the lowest layer at the bottom of the triple step, another extra row of Si atoms is observed. This row, denoted as ZR, is at the level of the R row and shows a weak contribution to the image in Fig. 2. In Fig. 4, this contribution is enhanced by the line subtraction processing, although this slightly distort the position of the row. The configuration of this row is zigzag-like as reported from STM analysis [2,10], similar to the zigzag chains observed on the Si(111)-(2 × 1) surface [29,30]. Each atom of the ZR has a DB almost parallel to the terrace slightly pointing upwards [grey circle in Fig. 3(b)]. The zigzag structure causes the DBs of consecutive atoms to point towards opposite directions, away from the ZR row. This zigzag chain is also buckled since the DBs are asymmetrically filled: the DBs of the atoms facing the R row are double filled whereas the ones of the atoms facing the D_1 dimers are empty. This asymmetry has been observed in the STM images, with a shift of the bright positions between empty and filled states images. In our SFM images a similar structure of ZR is observed for both contrasts, being better resolved in the restatom contrast owing to the lack of signal from the R row. The ZR row itself is preferentially tilted slightly either towards or away from the step edge, with the tilt away from the edge being the lower-energy configuration. However, even a mixed configuration within the ZR row represents a local energy minimum.

The calculations uncover as well the presence of additional restatoms. Between ZR and R, analogous to the usual restatoms of the 7 × 7 reconstruction, RA_3, there are restatoms denoted as RB. RB are the deepest located atoms within the reconstruction with dangling bonds. These RB restatoms are visible in both SFM contrasts as shoulders of the ZR row. In Fig. 2, RB atoms observed opposite to the gaps between the R atoms are indicated with a black dashed-line arrow. The ZR row is located at the position where the adatoms next to the RB restatoms would be if there was another 7 × 7 half cell. In addition, other restatoms are found between RDs, A_2 adatoms, and D_1 dimers, denoted as RA_2. The area close to the corner holes at the upper part of the triple step features an absence of restatoms, due to the row defects RD passing through three potential restatom sites. In contrast, the corner holes at the bottom of the triple step (at the R row) exhibit three restatoms as the close proximity of the ZR row leaves no space for row defects within the R row analogous to the ones in the A_2 row (RD). RB and RA_2 restatoms are indicated in Figs. 3 and 5 [orange and green circles in Fig. 3(b), respectively].

Apart from the atomic defects reported in the description of the triple step, other structural irregularities are observed. There are several types of adsorbates at step edges, especially at the 7 × 7 terrace step edge, as mentioned above. Their contrast depends on the tip, e.g., in Fig. 2 they are better resolved than in Fig. 1. Some of these adsorbates are indicated with white arrows in Fig. 2. Such protrusions can be observed at the step edge of the second layer (A_2) as well, as indicated with black arrows in Fig. 4(a). They look very similar to silicon atoms, therefore we tentatively ascribe them to additional silicon clusters produced during the preparation of the sample, while further work is needed to fully identify them [15]. Besides the Si clusters, we observe parts of the surface that are not properly reconstructed. First, on the 7 × 7 terrace in the middle of Fig. 4(b), we have marked with a V one not fully 7 × 7-reconstructed half cell. Second, in the lower part of the image, the distance between the two 7 × 7 flat terraces, marked with a double arrow in Fig. 4(b), is larger than the one usually given by the triple step.

IV. CONCLUSIONS

Summarizing, we have investigated with a joint FM-SFM and ab initio approach the structure of the popular vicinal Si(111) surface inclined towards the [112] direction. We observe two different SFM contrasts. First, the normal atomic contrast, which is expected from the bonding of the tip apex with the dangling bonds of the atoms of the surface. Second, the restatom contrast, which is explained taking into account additional short-range electrostatic interactions. Our atomically resolved images of unprecedented resolution
disclose the detailed structure of the triple step, and show the presence of several atomic defects, such as missing dimers and other structural irregularities. The calculations reproduce the features of the SFM image of the surface and reveal, besides the presence of different restatoms, a number of structural details on this surface: the buckling of the dimer rows at the step edges, the filling asymmetry and preferential orientation of the zigzag row, the degrees of freedom in the orientation of both dimers, and the possibility of orientational defects. All these features originate from the (partial) charge transfer between the dangling bond states.

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