Phase transition between \((2 \times 1)\) and \(c(8 \times 8)\) reconstructions observed on the \(\text{Si}(001)\) surface around 600°C

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The \(\text{Si}(001)\) surface subjected to different treatments in ultrahigh vacuum molecular beam epitaxy chamber for \(\text{SiO}_2\) film decomposition has been \textit{in situ} investigated by reflected high energy electron diffraction (RHEED) and high resolution scanning tunnelling microscopy (STM). A transition between \((2 \times 1)\) and \((4 \times 4)\) RHEED patterns was observed. The \((4 \times 4)\) pattern arose at \(T \leq 600°C\) during sample posttreatment cooling. The reconstruction was observed to be reversible. The \(c(8 \times 8)\) structure was revealed by STM at room temperature on the same samples. The \((4 \times 4)\) patterns have been evidenced to be a manifestation of the \(c(8 \times 8)\) surface structure in RHEED. The phase transition appearance has been found to depend on thermal treatment conditions and sample cooling rate.

Development of a procedure of atomically clean \(\text{Si}(001)\) surface preparation at lowered temperatures and/or by short thermal treatments is a keystone of creation of a CMOS compatible process of nanoelectronic VLSI fabrication [1, 2]. One of the ways of solving this problem is formation of a thin protective \(\text{SiO}_2\) film on a \(\text{Si}\) surface or surface passivation by hydrogen atoms during wet chemical etching with posterior silicon dioxide decomposition or hydrogen desorption from the surface in ultrahigh vacuum (UHV) ambient [3, 4]. In this connection, an issue of surface structure after these treatments becomes a task of primary importance taking into account a possible effect of \(\text{Si}\) surface atomic-scale roughness on formation of nanostructured elements (e. g., self-assembled Ge quantum dot nucleation on wetting layer in Ge/\(\text{Si}(001)\) heterostructures [2, 5, 6]).

This letter presents data of mutually complimentary investigation of clean \(\text{Si}(001)\) surfaces prepared by different methods in an UHV molecular beam epitaxy (MBE) chamber after different processes of wet chemical etching which has been carried out by means of high resolution scanning tunnelling microscopy (STM) and \textit{in situ} reflected high energy electron diffraction (RHEED).

The experiments were made using an integrated ultra-high-vacuum (UHV) system [1] based on the Riber EVA 32 molecular beam epitaxy chamber equipped with the Staib Instruments RH20 diffractometer of reflected high energy electrons and coupled through a transfer line with the GPI 300 UHV scanning tunnelling microscope [7, 8]. Samples were \(8 \times 8 \times 0.4\) mm cut from the specially treated KDB-12 commercial B-doped CZ \(\text{Si}(100)\) wafers (\(p\)-type, \(\rho = 12 \text{ Ω cm}\)). Initially, the specimens were chemically etched in the RCA etchant or a mixture of HNO\(_3\) and HF [3]. The obtained samples had different thicknesses of protective \(\text{SiO}_2\) films. A part of the samples, after etching in RCA, were immersed in a dilute HF solution to form a surface passivated by hydrogen atoms (\(\text{Si:H}\)). Then, the samples (except for \(\text{Si:H}\)) were annealed at 600°C for not less than 6 hours at the residual gas pressure of less than \(5 \times 10^{-9}\) Torr; the \(\text{Si:H}\) samples were treated at the temperature of \(\sim 300°C\) and the pressure of less than \(5 \times 10^{-11}\) Torr.

To obtain a clean \(\text{Si}(001)\) surface two standard for MBE methods of surface preparation were used: a short high-temperature annealing and decomposition of the \(\text{SiO}_2\) film by a weak flux of \(\text{Si}\) atoms. The short annealings were performed at 925, 800 or 650°C. A pressure in the MBE chamber did not exceed \(5 \times 10^{-9}\) Torr. Surface cleaning was monitored by RHEED: a characteristic \(\text{Si}(001)-(2 \times 1)\) pattern indicated that the surface was clean. Preparation of a clean surface by silicon dioxide decomposition in \(\text{Si}\) flux was conducted at the temperature of 800°C and \(\text{Si}\) deposition rate of less than 0.1 Å/s. The deposited \(\text{Si}\) layer thickness was greater than 30 Å. The samples were heated from the rear side by Ta radiators. The temperature was monitored with the IMPAC IS12-Si pyrometer which measured the sample temperature through chamber windows. Two modes of cooling was applied: quenching at the rate of \(\sim 0.4\) K/s and

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Fig. 1. A complimentary pair of STM images of the Si(001) surface after two cycles of annealing at 925°C for ∼3 min. with quenching (50 × 60 nm): (a) empty state mode, $U_s = +1.9$ V, $I_t = 80$ pA; (b) filled state mode, $U_s = -1.5$ V, $I_t = 80$ pA; inserts present the corresponding Fourier transforms ($2.9 \times 2.9$ and $3.7 \times 3.7$ nm$^{-1}$, respectively).

slow cooling at the rate of ∼0.17 K/s The atmosphere composition in the MBE camber was monitored using the SRS RGA-200 residual gas analyser. The STM images were obtained in the constant tunnelling current ($I_t$) mode at room temperature. The pressure in the STM chamber was less than $10^{-10}$ Torr. The STM tip was zero-biased while the sample was positively or negatively biased ($U_s$) when scanned in empty or filled states imaging mode. The STM tips were ex situ made of the tungsten wire and cleaned by ion bombardment [9] in a special UHV chamber connected to the STM. The STM images were processed afterwards using the WSxM software [10].

Application of high-temperature annealings for silicon dioxide film removal can lead to appearance of a reconstruction on the Si(001) surface during wafer cooling which is different from the (2 × 1) one. This reconstruction has previously been found to be $c(8 \times 8)$ [11]. This structure consists of ordered Si dimer pairs and di-vacancies which form “rectangles” gathered in rows running along ⟨110⟩ axes [11]. Surface coverage by this structure somewhat varies in different samples cooled at the same rate. Decrease of the coverage is observed with lowering of the sample cooling rate. Figs. 1 and 2 show STM images of the Si(001) surface obtained from samples cooled at different rates. It is seen that the same structure forms on the surfaces but with different coverages.

Fig. 2. STM images of the Si(001) surface after annealing at at 925°C for ∼3 min. and slow cooling: (a) 70 × 70 nm, $U_s = +1.6$ V, $I_t = 200$ pA; (b) 104 × 109 nm, $U_s = +2.0$ V, $I_t = 100$ pA.
Surface deoxidization has been explored during processing by means of RHEED; it has been established that according to electron diffraction patterns the (2 × 1) structure forms on the surface during the high-temperature treatment at 925°C. Then, on sample cooling, a phase transition goes on which is characterized by gradual appearance of a diffraction pattern corresponding to the (2 × 1) pattern change on screen is apparent to the eye in the sample temperature interval from ∼ 600 to ∼ 550°C. The phase transition is reversible: recurring heating returns the (2 × 1) pattern at the same temperature (∼ 600°C); the (4 × 4) pattern arise again on repeated cooling. Fig. 1 presents STM images of a sample after two cycles of annealing and quenching. The c(8 × 8) structure is seen on the images obtained in the empty-state (Fig. 1a) and filled-state (Fig. 1b) modes; RHEED (4 × 4) patterns corresponding to this surface reconstruction are presented in Fig. 3. Inserts of Fig 1 show the Fourier transforms which are evidently different for different scanning modes: the empty-state image Fourier transform corresponds with a periodicity of 8 translations along the [1 1 0] directions. Fig. 5 represents RHEED patterns obtained from one of these samples with reflexes exactly matching to the reconstruction observed by STM. The origin of this reconstruction has not been understood thus far and requires further study.

The (2 × 1) RHEED pattern was observed for the passivated samples after annealing at 800°C for 5 minutes and quenching.

Earlier, we have already presented a model of the c(8 × 8) structure formation based on the assumption that it consists of two layers. The uppermost layer is composed by Si ad-atoms which, at high temperatures, migrate along the underlying (2 × 1) reconstructed layer. Si ad-atoms stay on the surface after the SiO2 decomposition reaction: SiO2 + Si → 2SiO↑. This is additionally confirmed by the observation of the (2 × 1) reconstruction after cleaning at 800°C of the surface passivated by hydrogen because in this case Si atoms are not spent for hydrogen removal. RHEED data may be interpreted as follows: at high temperatures, we observe a pattern produced by diffraction on the underlying layer, which corresponds to (2 × 1). As the sample is cooled atoms of the uppermost layer lose the mobility and form the c(8 × 8) structure which is exhibited in diffraction patterns as (4 × 4). Possible models of atom ordering in such structure will be considered in a different article.
A reason of the discrepancy of STM and RHEED results may be understood from the fact that the c(8 × 8) structure consists of “rectangles” whose location is strictly predetermined by the dimer rows of the underlying layer \([11]\). The dimers located on short sides of the “rectangles” lie on top of the lower dimer rows and are somewhat higher than the rest dimers forming the c(8 × 8) structure. Fig. \([11]\) demonstrates an STM image of the surface and its Fourier transform which correspond to the (4 × 4) reconstruction. Depending on a magnitude of the negative bias applied to the specimen, an image can be obtained in which the “rectangle” looks as two parallel bars positioned exactly on places of the extreme dimers \([11]\). This usually takes place at small values of the negative bias. In this case the entire visible structure corresponds to the (4 × 4) reconstruction. Thus, a cause of the discrepancy of STM and RHEED data is likely that the high energy electrons diffract on the topmost dimers, situated on sides of the “rectangles”, rather than on different dimers of the surface structure who are somewhat lower but usually manifested in STM images except for some special mode of scanning at low negative bias when only the highest dimers contribute to the tunnelling current.

In summary, it has been established that at high-temperature processes of the silicon dioxide film removal from the Si(001) surface a reversible phase transition from \((2 \times 1)\) to \(c(8 \times 8)\) structure takes place at the temperature of \(\sim 600°C\) on sample cooling. The \((2 \times 1)\) structure restores on heating at the same temperature. The low-temperature structure is exhibited as \((4 \times 4)\) one in RHEED patterns. STM images show this structure to cover different fractions of the surface area. The coverage decreases as the sample cooling rate is reduced. At small coverages, RHEED patterns correspond to the \((2 \times 1)\) reconstruction. STM images also demonstrate the \((2 \times 1)\) structure in areas free of the \(c(8 \times 8)\) one.

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