Atomic-scale structural evolution of Ge(100) surfaces etched by H and D

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The atomic-scale structural evolution of Ge(100) surfaces etched by H(g) and D(g) at $T_s=400$ K is studied using scanning tunneling microscopy (STM) and field emission-scanning electron microscopy (FE-SEM). The STM investigation reveals that etching of the Ge(100) by H(g) and D(g) proceeds initially via the production of single atom vacancies (SV), dimer vacancies (DV), and subsequently, line defects along the Ge dimer rows. It is also observed that D(g) etches the Ge(100) surface eight times faster than H(g) does. After extensive exposures of the surface to H(g), the FE-SEM images show square etch pits with V-groove shapes, indicating that H(g) etching of the Ge(100) surface proceeds anisotropically. © 2004 American Institute of Physics.

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The interaction of hydrogen with semiconductor (100) surfaces has been of great technological interest in the field of semiconductor processing. It is well known that etching of a Si(100) surface by atomic H(g) depends on the surface temperature $T_s$. Previous experiments have suggested that H etching of a Si(100) surface proceeds initially on the dihydride (1×1) surface: H(g) + SiH$_2$(a) → SiH$_3$(a), followed by H(g) + SiH$_3$(a) → SiH$_4$(g). H(g) breaks the Si–Si bonds of SiH$_2$(a), producing gas-phase SiH$_3$(g) via successive H(g) additions.

It has been generally believed, however, that atomic H(g) does not etch the surface of Ge(100), with even high H(g) exposures yielding only a stable monohydride (2×1) phase. Recently, a scanning tunneling microscopy (STM) study has shown that this 2×1:H phase is largely maintained even after extensive H(g) exposure, due to the instability of the surface dihydride GeH$_2$(a). To our knowledge, there have been no reported reports to date on H etching of Ge(100) surfaces, though studies have been published for hydrogen etching on Ge/Si(100) surfaces.

In this letter, we present an atomic-scale STM study of structural changes of Ge(100) surfaces etched by atomic H(g) and D(g) at $T_s=400$ K. The variation of the surface morphology after extensive H(g) exposure was also investigated using field emission-scanning electron microscopy (FE-SEM). The etching rates of Ge(100) surfaces by H(g) and D(g) are compared.

The studies presented here were performed in an ultra-high vacuum chamber described in detail elsewhere.

We note that hydrogen etching at $T_s<400$ K occurs randomly and forms large, irregular bright features ascribed to the etching intermediate, GeH$_3$(a), while for $T_s>400$ K the etching rate rapidly drops due to the difficulty of GeH$_3$(a) formation.

Figure 1(a) shows the STM image of an initially etched Ge(100)-2×1:D surface obtained after dosing 3000 L of D$_2$(g). Some bright ball-like features (<1%) are observed and distributed randomly over the surface. A prior study has indicated that these ball-like features are due to unpaired dangling bonds on dimers that are occupied by only a single D atom (D–Ge–Ge). A significant number of such unpaired dangling bonds still persist even after prolonged D(g) exposure. A few atomic vacancies due to initial etching as well as antiphase boundaries are observed near local GeD$_2$(a) rows. Figures 1(b)–1(d) show high resolution STM images of a single atom vacancy [SV, circle in Fig. 1(a)], single atom vacancy row [SVR, square in Fig. 1(a)], and dimer vacancy [DV, triangle in Fig. 1(a)] appearing after initial D etching. It is known that adjacent GeH$_3$(a) species are precursors to breaking of the Ge–Ge bonds. The surface trihydride species, GeH$_3$(a), produced by H insertion into GeH$_3$(a), can be hydrogenated by H(g) and removed as GeH$_4$(g), as shown by mass spectrometry. Figure 1(e) shows a STM image of a Ge(100) surface exposed to 7×10$^5$ L of D$_2$(g). In contrast to previous studies, the STM image shows that significant etching by D(g) occurs on the Ge(100) surface. The STM images recorded with increasing D(g), and also of H(g), exposures reveal linear line defects formed along the Ge dimer rows. These STM results indicate that the GeD$_2$(a) or GeH$_2$(a) species on the Ge(100) surface are quite stable as etching precursors. However, even after extensive exposure of the Ge surface to D(g) or H(g), the large-scale 1×1 or 3×1 phases composed of GeD$_2$(a) or GeH$_2$(a) were not observed, unlike in the H/Si(100) case, due to repulsion driven H$_2$(g) or D$_2$(g) desorption, as reported previously.

Figure 1(f) shows the etching mechanism in which Ge dimers on a Ge(100) surface form GeH$_2$(a) or GeD$_2$(a) pairs and how these dihydrides or dideuteride species desorb, generating SVs and subsequently DVs via successive H(g) or D(g) additions. However, recombinative H$_2$(g) or D$_2$(g) desorption, due to steric repulsion between the H or D atoms of two adjacent GeH$_2$(a) or GeD$_2$(a) species, may reduce the

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amount of GeH$_2$(a) or GeD$_2$(a) species on the surface by decomposition into GeH(a) or GeD(a) species. This indicates that the GeH$_2$(a) or GeD$_2$(a) species either quickly return to GeH(a) or GeD(a) by collision-induced H$_2$ or D$_2$ dissociation [2GeH$_2$(a)→H−Ge−Ge−H+H$_2$(g)]$^4$ or etch the Ge surface by the adsorption of incoming H(g) or D(g) [GeH$_2$(a)+H(g)→GeH$_3$(a), followed by GeH$_2$(a)+H(g)→GeH$_4$(g)]. This DV then destabilizes adjacent dimers due to tensile stress. The incoming H(g) or D(g) attacks them and a pit grows as a string of DVs parallel to the dimer row by successive removal of neighboring dimers. In addition, row features, expected by [GeH$_3$(a)→GeH$_2$(a)+H(a),GeH$_2$(a)→GeH(a)+H(a)] at $T_s$=400 K, are also observed [the circle in Fig. 2(a)].$^7$

Figures 2(a) and 2(b) show STM images of Ge(100) surfaces exposed to $7 \times 10^5$ L of H$_2$(g) and D$_2$(g), respectively. The areas of the etch pits produced by H(g) and D(g) etching occupy ~10% and ~65% of the whole surface area, respectively. The STM images in Figs. 2(a) and 2(b) reveal linear line defects formed along the Ge dimer rows. In these STM images, the etched surface area rapidly increases with increasing flux of H$_2$(g) or D$_2$(g). Figure 2(c) shows the linear dependence of the etched monolayer (ML) of the Ge(100) surface on H$_2$(g) and D$_2$(g) exposure. The linear dependence is consistent with an etching reaction involving a direct attack of the Ge hydride or Ge deuteride species by H(g) or D(g) through the Eley–Rideal mechanism.$^5$ It can be seen in Fig. 2(c) that the rate of etching of the Ge(100) surface by D(g) is eight times faster than that by H(g), indicating that the formation of GeH$_2$(a) as an etching precursor is more difficult than the formation of GeD$_2$(a). This is due to an isotope effect, as previously reported.$^8$ Specifically, the collision induced dissociation of GeH$_2$(a) pairs occurs more easily than that of GeD$_2$(a) due to the larger vibrational frequency of Ge−H compared with that of Ge−D.$^9$ Hence the efficiency of dry etching of a Ge(100) surface by D(g) is greater than that by H(g).

We examined the surface morphology after extensive exposures of the Ge(100) surface to H(g) using FE-SEM. Figure 3(a) shows a SEM image of a Ge(100) surface recorded after a H$_2$(g) dosage of $\sim 1 \times 10^6$ L at $T_s$=400 K. The SEM image reveals one of the square etch pits with a V-groove shape of size $\sim 2 \times 2 \, \mu m^2$ found randomly distributed on the substrate. Rectangular etch pits are also observed. Similar etch pits are observed in wet and dry etching processes.$^{10}$ The shapes of the etch pits were expected to be limited by the planes with the lowest etching rate. It is known that (111) planes develop during anisotropic etching because of the lower etching rate of the (111) plane compared to other planes.$^{10}$

Figure 3(b) shows a cross-sectional view of the etch pit with a V-groove shape seen in Fig. 3(a). The angle between the two faces of the etch pit is about 55$^\circ$ and the etch pit is bounded by a set of four (111) planes, as shown in the sche-
The repulsion resulting from the insertion of a H atom is relaxed by rotation about the remaining bond. The ability to stabilize such etch intermediates helps to promote the etching reaction on the Ge(100) surface. The results show that the H(g) etching of the Ge(100) surface is highly anisotropic, resulting in predominantly square etch pits with V-groove shapes.

In summary, the present STM studies show that etching of a Ge(100) surface by H(g) and D(g) at $T_s = 400$ K proceeds initially by producing SVs and DVs. Etch pits of V-groove shape produced by anisotropic H(g) etching of the Ge(100) surface were observed. Compared to H(g), D(g) was found to etch the Ge(100) surface with greater etching efficiency.

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FIG. 2. Filled-state STM images (70×70 nm²) of Ge(100) surfaces exposed to (a) H₂(g) of $7\times10^4$ L, and (b) D₂(g) of $7\times10^4$ L at $T_s = 400$ K. The circle in (a) shows a regrowth feature; (c) shows a plot of an etched Ge monolayer (ML) vs exposure (L) with H₂(g) and D₂(g).

FIG. 3. FE-SEM images of (a) a typical etch pit of V-groove shape formed on a Ge(100) surface exposed to H₂(g) of $\sim 1\times10^5$ L at $T_s = 400$ K, (b) the cross sectional view of (a), and (c) a schematic diagram of (b) showing that the etch pits are bounded by a set of four (111) planes.

It is well established that the (111) face is generally the most stable because the Ge atoms on the (111) face are bonded to the surface by three Ge–Ge bonds. Insertion of a H atom into one of these bonds will greatly strain the remaining two bonds. However, on the (100) face, each Ge atom has two highly strained bonds to the surface. The repulsion resulting from the insertion of a H atom is relaxed by rotation about the remaining bond. The ability to stabilize such etch intermediates helps to promote the etching reaction on the Ge(100) surface. The results show that the H(g) etching of the Ge(100) surface is highly anisotropic, resulting in predominantly square etch pits with V-groove shapes.

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