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Annealing of isolated amorphous zones in silicon

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In situ transmission electron microscopy has been used to observe the production and annealing of individual amorphous zones in silicon resulting from impacts of 200-keV Xe ions at room temperature. As has been observed previously, the total amorphous volume fraction decreases over a temperature range from room temperature to approximately 500 °C. When individual amorphous zones were monitored, however, there appeared to be no correlation of the annealing temperature with initial size: zones with similar starting sizes disappeared (crystallized) at temperatures anywhere from 70 °C to more than 400 °C. Frame-by-frame analysis of video recordings revealed that the recovery of individual zones is a two-step process that occurred in a stepwise manner with changes taking place over seconds, separated by longer periods of stability. © 2003 American Institute of Physics. [DOI: 10.1063/1.1562336]

Ion implantation is a major technology for semiconductor device manufacture, but it unavoidably leads to radiation damage that must be minimized, controlled, or removed. Heterogeneous nucleation of the amorphous phase may result when an ion impact causes a sufficiently high local-defect density to render a small region amorphous.1–4 For large energy-deposition densities, this may occur for a single ion impact. As ion fluence further increases, these zones accumulate and overlap to form a continuous amorphous layer.

Thermal annealing can convert such amorphous layers to crystal by advancement of the interface with good crystal into the layer. Motion of the interface, so as to produce additional amorphization or recrystallization, can also be controlled by varying the ion flux and substrate temperature.5 This suggests that recrystallization involves the generation and movement of defects at the amorphous–crystalline (a/c) interface. Possible defects include dangling bonds,6 charged kinks on terraces,6 the so-called “bond defect,”7 and divacancies.2

Recrystallization of incompletely amorphized layers has been observed to occur thermally and by electron irradiation.8 The annealing behavior has been studied by Rutherford backscattering,9 which determines the fractional lattice disorder, and by transmission electron microscopy (TEM),1 in which amorphous zones were observed to recrystallize at lower temperatures than the continuous amorphous layer, although details of the dynamics of this zone annealing were not recorded. The zones were also observed to recrystallize over a wide range of temperature.

The thermal recrystallization rate of the planar a/c interface exhibits Arrhenius-type behavior with a single activation energy of 2.7 eV for recrystallization rates varying over ten orders of magnitude and temperatures from 500 to 1400 K.3 However, recent molecular dynamics (MD) simulations yielded two ranges of activation energies at low and high temperatures, respectively.10,11 The difference between these results was attributed to a higher concentration of defects near the interface in the MD simulation. Similarly, MD simulations of the recrystallization of small amorphous zones yielded a range of activation energies that increase with zone size.12 In these latter simulations, annealing was observed to result in periods during which no crystallization occurred, separated by periods of rapid partial recrystallization. It should be noted that MD simulations of annealing are carried out over very short time scales (<100 ps).

The present work was undertaken to examine the kinetics of thermal annealing of isolated amorphous zones in Si using a TEM in which in situ ion irradiation can be carried out. Using a heating stage, the entire “life-cycle” of individual amorphous zones can be followed, from creation to recrystallization, and recorded on film and videotape.

Specimens were prepared by diamond core drilling 3-mm disks from single-side-polished, 400-μm-thick, p-type (boron, 80 to 160 Ω cm) {110} Czechralski silicon wafers. Disks were mechanically ground to a thickness of 200 μm before being dimpled and electrochemically jet-thinned to perforation in a non-acid etch. In situ ion irradiations and anneals (using a Gatan double-tilt heating stage) were carried out in a Hitachi H-9000 TEM, operating at 100 keV, located at the IVEM/Accelerator Facility at Argonne National Laboratory.13 The ion beam is oriented 30° from the microscope axis. The electron energy was chosen so as to minimize electron-beam annealing of amorphous zones in silicon.8 Under our viewing conditions, no effects of the electron beam on the amorphous zones were observed over a time scale of 30 min. During isochronal annealing experi-
The projected areas of more than 400 zones were below 40 °C and realigned for down-zone, structure-factor was maintained for 10 min. The specimen was then cooled to temperature as rapidly as possible during which the temperature rose from 302 to 308 °C. Shrinkage of this zone took place in two distinct stages labeled "A" and "B." Each stage lasts on the order of 1 s, and the zone changed little during the period between the two stages. In general, during a slow temperature ramp, zones measured and converted to effective radii by assuming a spherical shape. The number of amorphous zones remaining following each annealing step is illustrated in Fig. 1. This behavior is in good agreement with results obtained previously. Also plotted in Fig. 1 is a dashed line indicating the theoretical annealing behavior of these zones if the motion of the a/c boundary followed the same kinetics as in planar geometry.

Two important points are observable on this plot: first, an activation energy of 2.7 eV would result in all zones surviving to above 500 °C, and second, the temperature range over which the zones would recrystallize due to a singly activated process would be much smaller than is actually observed. Clearly, the recrystallization behavior of small amorphous zones is very different from that of the planar a/c interface and does not involve a single, well-defined activation energy.

The averaged information shown in Fig. 1 fails to reveal the erratic annealing of individual zones. This is illustrated in Fig. 2, where the annealing behavior of three amorphous zones of almost identical starting sizes (effective radius of just over 2 nm) is plotted. This wide range of recrystallization behavior is typical. Analysis of more than 400 zones on each annealing step reveals little consistency in their behavior and little or no correlation between their size and recrystallization temperature. On each annealing step, zones of every size disappear. The behavior is thus neither consistent with a singly activated process nor with the findings of MD simulations in which the activation energy for recrystallization was seen to increase with zone size.

The dynamical shrinkage of individual zones was monitored by slowly ramping the temperature while recording the image on videotape. Figure 3 shows the behavior of a single isolated amorphous zone (initial effective radius 1.7 nm) as a function of time during a brief period of just over a minute, during which the temperature rose from 302 to 308 °C. Shrinkage of this zone took place in two distinct stages labeled “A” and “B.” Each stage lasts on the order of 1 s, and the zone changed little during the period between the two stages. In general, during a slow temperature ramp, zones...
The precise atomic arrangement near the interface will vary from zone to zone and from point to point around the surface of each zone. The energy required to create the trigger defect responsible for recrystallization will thus vary from zone to zone (as possibly will the migration energy).

The time for the rapid recrystallization steps (as seen in Fig. 3) is on the order of 1 s at 300 °C. Annealing of a nanometer-sized amorphous volume involves several hundred atomic rearrangements. At a temperature of 300 °C, a singly activated process with a rate constant of $10^{13}$ s$^{-1}$ that requires between 100 and 1000 atomic jumps to go to completion, would have an activation energy in the range 1.14 to 1.25 eV. A migration energy of 1.2 eV has been measured for the defect responsible for both thermal and ion-beam-induced recrystallization (implying a formation energy of 1.5 eV). Although Jackson suggests that this defect is the dangling bond, it should be noted that this is generally assumed to have a formation energy of 2.25 eV (half the cohesive energy), which implies a migration energy of 0.43 eV.

We have shown that recrystallization of nanometer-sized amorphous zones is not a singly activated process, but one that takes place over a wide range of temperatures. It involves at least two thermally activated processes: a triggering event and the recrystallization process itself. Triggering events occur over a temperature range of more than 300 °C. The defect responsible for recrystallization has not yet been identified, but the dangling bond, the kink site, and the bond defect have all been suggested as possible candidates. The irregularity of the interface of any nanometer-sized zone and the variation in local atomic arrangement at the interface may be responsible for a wide range of energies for the creation of the defect responsible for recrystallization. This creation is the rate-limiting triggering event. The subsequent rapid recrystallization appears to have an associated energy of about 1.2 eV.

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