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Accessibility
Amorphous-crystal interface in silicon: A tight-binding simulation

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The structural features of the interface between the crystalline and amorphous phases of Si solid are studied in simulations based on a combination of empirical interatomic potentials and a nonorthogonal tight-binding model. The tight-binding Hamiltonian was created and tested for the types of structures and distortions anticipated to occur at this interface. The simulations indicate the presence of a number of interesting features near the interface. The features that may lead to crystallization upon heating include (110) chains with some defects, most prominently dimers similar to those on the Si(001) 2×1 reconstructed free surface. Within the amorphous region order is lost over very short distances. By examining six different samples with two interfaces each, we find the energy of the amorphous-crystal interface to be 0.49 ± 0.05 J/m².

I. INTRODUCTION

The crystalline and amorphous phases of silicon are considered prototypical examples of a tetrahedrally coordinated network in ordered and disordered forms. Each phase has been intensively studied experimentally and theoretically, and both are used in a very broad spectrum of electronic applications. Currently all Si integrated circuits employ several ion implantation steps in their fabrication. Regions that receive a sufficiently high implantation dose become amorphous within approximately 100 nm of the free surface; the crystal structure is restored by an interface-mediated crystallization process called solid-phase epitaxial growth (SPEG). While much is known about the structure of the crystal and amorphous phases individually, considerably less direct information is available about the structure of the interface between them. In light of the importance of SPEG, and of the intrinsic interest of interfacial phenomena, a detailed atomistic study of the amorphous-crystal interface in Si is appropriate. The inaccessibility of the interface atomic structure by experimental probes leaves as the only alternative realistic simulations of this system. In this paper we discuss such simulations and the insight they provide into the atomic structure and dynamics at the amorphous-crystal interface in Si.

There are two major obstacles in simulating this interface: first, a relatively large number of atoms must be included in the simulation to ensure that the character of the two phases is represented accurately; second, extensive exploration of configuration space is required to ensure that the system is not locked in some small (and potentially not representative) region of the accessible configuration space. Ideally one would like to simulate this system by means of unbiased, parameter-free quantum-mechanical calculations [such as density functional theory in the local density approximation (DFT/LDA)], but both the size of systems that can be handled and the extent of configuration space that can be explored through such calculations are severely restricted. Past efforts have either used hand-built models,¹,² computer relaxed geometrical models,³ or molecular-dynamics (MD) simulations based on empirical interatomic potentials.⁴,⁵

Here we combine the use of the Stillinger-Weber empirical interatomic potential and a recently developed semi-empirical quantum-mechanical technique, based on a nonorthogonal tight-binding (TB) Hamiltonian that was parametrized to represent accurately a wide range of bulk and surface structures of Si.⁶ The use of the empirical potential was motivated by the fact that it affords fast but less accurate calculations for parts of the simulation where maintaining high accuracy is not important; specifically it is used to bring the system from a high-temperature, liquid-crystal interface far from equilibrium, to a low-temperature amorphous-crystal interface near equilibrium. Once the system is close to equilibrium, we switch to the TB model, which can handle reasonably large systems and is sufficiently fast to allow exploration of configuration space, while maintaining the basic quantum-mechanical treatment of electronic degrees of freedom. As such, it is superior to empirical interactions that are considerably more restricted in their ability to describe large structural distortions and the breaking and formation of covalent bonds. The results of the tight-binding studies can also be used as starting points for more elaborate parameter-free quantum-mechanical calculations, although we anticipate that the essential features will remain unchanged.

II. METHODOLOGY

We use constant-temperature, constant-stress MD to prepare the amorphous-crystal interface samples starting with a liquid-crystal interface as described below. Because of the large time scale necessary to create reasonably equilibrated amorphous samples, the use of the tight-binding Hamiltonian to compute the interatomic forces while the system is very
far from its equilibrium state is impractical, and indeed not beneficial. Instead we use the Stillinger-Weber interatomic potential\(^7\) to bring the system reasonably close to equilibrium and only then turn on the tight-binding Hamiltonian. The equations of motion are integrated with a Gear predictor-corrector algorithm\(^8\) with a time step of 1 fs. The temperature is kept constant using a velocity rescaling algorithm where the atomic velocities are uniformly scaled to give the desired temperature once every 500 time steps. Zero stress is maintained with an extended system Parrinello-Rahman approach.\(^9\)

The simulation cell includes 320 atoms in a \([220]\times[2\overline{2}0]\times[005]\) box with periodic boundary conditions in all three directions. The interface lies in the (001) plane, with the [220] and [2\overline{2}0] vectors forming its sides (in the following the [001] direction is referred to as the z axis). To create the interface, we maintain a portion of the simulation cell in the crystalline phase by keeping it below the melting point, while melting and then quenching the remainder of the cell. The crystalline region includes 128 atoms (8 monolayers) that are kept at 100 K, and the remaining 192 atoms (12 monolayers) form the amorphous region. The amorphous region is produced by cooling a molten region from 5000 to 1000 K, using the method of Luedtke and Landman.\(^10\) A total of six samples, cooled to 100 K and equilibrated with the Stillinger-Weber potential, were relaxed using a conjugate-gradient algorithm with the tight-binding Hamiltonian to calculate forces and stresses. The relaxed samples were used in the analysis of interface features. A typical sample is shown in Fig. 1.

### III. ANALYSIS

#### Structural analysis

Standard measures for characterizing the structure of the bulk phases are the radial pair correlation \(g(r)\) and bond-angle distribution \(p(\theta)\) functions. These are shown in Fig. 2 averaged over all six samples. For the amorphous regions (those that were thermally cycled), the functions were computed from samples where the atoms in the crystal regions (those that were kept cold) were removed, but using the original periodic boundary conditions. For the crystal regions the atoms in the amorphous regions were removed. Because of the missing neighbors at the edges of each region the normalization for the curves is nonstandard, although the overall shape is not affected. The pair correlation functions exhibit the expected features: averages over atoms in the crystalline regions show distinct order at all ranges allowed by the size of the simulation cell; averages over atoms in the amorphous regions show distinct first- and second-neighbor peaks, but no order at longer range. In particular, they do not have a third-neighbor peak, a feature also seen in DFT/LDA simulations\(^11\) and in experiment.\(^12\) In the following, the position of the minimum between the first two peaks of the pair correlation function \((r=2.7 \text{ Å})\) is used as the criterion for defining the neighbors of an atom in the amorphous regions. The mean bond angles are 108.9\(\pm\)6° and 108.4\(\pm\)14° in the crystalline and amorphous regions, close to the ideal tetrahedral angle of 109.5°.

Coordination statistics and ring statistics based on the same nearest-neighbor criterion are listed in Table I. The coordination of the atoms in the crystalline region is nearly perfect; in the amorphous region there is a significant number of defects, with overcoordinated atoms predominating. There is also a significant number of minimal rings (computed using shortest path analysis\(^13\)) with size other than six, including a few eight-membered rings. In agreement with the results of DFT/LDA MD simulations by Stich et al.,\(^11\) we also observed more 5-membered than 7-membered rings. The total ring statistics indicate more even-membered rings than the random bond switching model of Wooten et al.,\(^14,15\) and fewer odd-membered rings.

The characterization of the interface is somewhat more demanding. In order to identify the interface region and to characterize its features we define three different quantities. The first of these is the rms deviation of the bond angles from the ideal tetrahedral angle \(\Delta\theta\). The bond angle devia-
FIG. 2. Measures of order in the bulk of the crystalline and amorphous regions (as defined in the text): Pair correlation functions $g(r)$ and bond angle distribution functions $p(\theta)$.

A second quantity we define to characterize the interface is the sum of the vectors pointing from an atom to its nearest neighbors. This vector quantifies the asymmetry of the atomic environment. For example, if an atom is missing one of its neighbors while retaining $sp^3$ bonding, this vector will point away from the missing atom. We refer to this vector as the "tetrahedral" vector $\mathbf{v}$. Because of the difficulty of plotting vector quantities, the magnitude of $\mathbf{v}$ versus the $z$ position of each atom is plotted in Fig. 3(b), averaged over all samples and smoothed as described earlier. The differences between the crystalline and amorphous regions are again small but distinct. The extent of the interface using $\mathbf{v}$, is very similar to that indicated by $\Delta \theta$. In the interface region, both $\Delta \theta$ and $\mathbf{v}$ vary monotonically between the values in the amorphous and the crystal regions. The definition of the vector sum becomes more useful when its values and directions at individual interface atoms are considered; these indicate the direction and amount by which a given atom (or one of its neighbors) should move in order to create an environment closer to the crystalline state.

A third local quantity we employed to characterize the interface region is the volume of the Voronoi polyhedron associated with each atom $\Omega_v$. Plotted in Fig. 3(c), averaged over all samples and smoothed as described earlier. This quantity gives a local measure of the density, as well as an estimate of the free volume around each atom. $\Omega_v$ is about $19.0 \, \text{Å}^3$ in the crystal, which corresponds to a $3.5\%$ compression of the unstrained bulk crystal volume. In the amorphous phase $\Omega_v$ ranges from $20.0 \, \text{Å}^3$ to $20.5 \, \text{Å}^3$, i.e., the amorphous phase is a few percent less dense than the crystal. To determine the variation of the strain with position and direction, we calculated the mean nearest-neighbor spacing projected along the in-plane axes, $x$ and $y$, and the perpendicular axis, $z$. In the crystalline region, the $x$ and $y$ spacings are $7\%$ smaller than the spacing along the perpendicular direction, indicating that the crystal is under biaxial compression. In the amorphous region the $x$ and $y$ spacings are $3\%$ larger than in the perpendicular direction, indicating that the amorphous is under biaxial tension. Because the two adjacent phases are in opposite strain states, it is impossible to resolve the sign or magnitude of the interface stress.
For a better understanding of the structure of the amorphous-crystal interface we created slices of the samples parallel to the interface. Perspective views of these slices reveal some interesting characteristics: Fig. 4 is an example where the prominent features of the crystalline portion are chains of atoms along the [110] direction, with very few defects. The atoms that are not in ideal positions form dimers, where pairs of atoms on adjacent [110] chains have come close together to form a bond, a feature that was also seen in the hand-built model of Saito and Ohdomari. One example of this defect is seen on the left side of the image in Fig. 4 (between the two vertical [110] chains). This feature is very similar to the well-known Si(001) $2 \times 1$ free surface reconstruction, although in the present case the atoms participating in the dimer have four bonds (each with two more neighbors on the crystalline side and one more neighbor on the amorphous side). On the amorphous side of the interface, some atoms are beginning to assume positions compatible with the crystal lattice. They line up in chains along [110] directions (top of image in Fig. 4), as would be expected for the next layer in the crystal. The remaining atoms are arranged in more disordered configurations.

**IV. INTERFACE ENERGIES**

One important quantity that characterizes the interface is the interfacial tension $\sigma_{ac}$, which is, for a single-component system, the excess free energy per unit area. This excess is responsible for the barrier to nucleation of the crystal in the middle of the amorphous phase; typically the interfacial tension is determined experimentally by interpreting nucleation rate measurements under conditions where heterogeneous nucleation is believed to be insignificant. Because it is difficult to ensure that this condition has been achieved, experimental values for the interfacial tension, such as those estimated by Tsao and Peercy or Yang, represent a lower limit on the true value of the interfacial tension. At sufficiently low temperatures the entropic contribution to the interfacial tension can be neglected and $\sigma_{ac}$ can be approximated by the excess interfacial energy per unit area $\epsilon_{ac}$, which is easier to determine theoretically. Mathematically, $\epsilon_{ac}$ is defined as the excess energy of a system with an interface over the weighted sum of the energies of the two constituent phases,

$$\epsilon_{ac} = \frac{(E - N_c \epsilon_c - N_a \epsilon_a)}{A}.$$

$E$ is the total cohesive energy of the sample with the interface, $\epsilon_c$ and $\epsilon_a$ are the cohesive energies per atom of the reference crystal and amorphous states, $N_c$ and $N_a$ are the number of atoms in the crystalline and amorphous phases, respectively, and $A$ is the total area of the interface. An analogous equation to Eq. (1) for $\sigma_{ac}$ can be obtained by replacing $\epsilon_c$ and $\epsilon_a$ by the corresponding free energies of these phases per atom, $g_c$ and $g_a$. When the system is in equilibrium, the assignment of atoms to the individual phases (i.e., the determination of $N_c$ and $N_a$) is unnecessary because, by definition, $g_c$ and $g_a$ are equal in equilibrium. For the silicon amorphous-crystal interface, even when $\sigma_{ac}$ can be approximated by $\epsilon_{ac}$, the determination of $N_c$ and $N_a$ is necessary because the two phases are not in equilibrium with each other. Hence we must determine which atoms should be considered “crystalline” and which “amorphous.”

To do that we visualize slices of our samples parallel to the interface and label as crystalline any atoms that are bonded to two atoms that were kept frozen or two other atoms that are labeled as crystalline by this procedure, provided that the two atoms would share a common neighbor in the perfect crystal. This ensures that all the atoms that are considered part of the crystal are in a nearly ideal crystal environment on at least one side, and all are members of sixfold rings that are contained in the crystal. Because the calculated interface energy is sensitive to the number of crystal atoms we need to employ a more rigorous definition of the bond between atoms than the one used earlier, which relied simply on distance (atoms closer than 2.7 Å were considered bonded). To this end, we consider atoms bonded only if the tight-binding charge density half way between them is above a threshold value that is obtained by using representative $s$ and $p$ orbitals attached to each atom. Typically, between 10 and 20 pairs of neighboring atoms (out of about 650 pairs in each sample) have charge densities that fall below this threshold and are not considered to be bonded to each other, even though their distance is shorter than 2.7 Å.

A second complication in using our tight-binding Hamiltonian to compute the interface energy is the precise value of $\epsilon_c$ and $\epsilon_a$. The reference crystal state is an uncompressed diamond lattice (the compression energy is negligible), trivial to generate and its cohesive energy $\epsilon_c$ is easy to compute. To compute an appropriate reference amorphous state from which $\epsilon_a$ can be estimated, we take each interface sample and apply the same procedure we used to create the amorphous portion, but this time keeping a 4.75 Å slab centered in the middle of the amorphous portion frozen. In this way, we make the entire sample amorphous. Each bulk amorphous sample is then relaxed with the tight-binding
Hamiltonian, and used as the reference state when computing the interface energy for the corresponding interface sample. The resulting bulk amorphous samples have cohesive energies $e_c$ between 4.519 and 4.536 eV/atom, corresponding to an excess energy for the amorphous phase $\Delta e_{ac}$ of 0.17 to 0.19 eV/atom. These values are a factor of two higher than an experimental value for $\Delta e_{ac}$ of 0.097 eV/atom, as extrapolated to 0 K from Donovan's measurement at 960 K using the specific heat listed in that work.

The resulting interface energies $\sigma_{ac}$ range from 0.39 to 0.54 J/m$^2$ for the six different samples, with a mean of 0.49 J/m$^2$ and a standard deviation of 0.05 J/m$^2$. The scatter is due to several factors. The total energy of the two interfaces in each sample is a small number (about 15 eV) computed by subtracting large numbers (total energies for the interface and reference states, each of order 1500 eV). Scatter of 0.3% in the total energy of the interface samples or reference amorphous samples (which is inevitable due to their disordered nature and small size of the systems) causes a scatter of 30% in the computed interface energy. Partitioning the atoms into crystalline and amorphous parts also involves an error of about two or three atoms per interface, arising from both the threshold charge density value for considering two neighboring atoms bonded and from errors made in the manual counting process. There is also a potentially larger source of error in the arbitrary definition of what is required for an atom to be considered "crystalline." Some other criteria we considered, using the values of different measures of order to distinguish between "crystalline" and "amorphous" atoms, gave values for $N_c$ that differed by as many as tens of atoms from the topological criteria described previously.

The only previous attempt to compute the interface energy through simulation we are aware of is Spaepen's work using a Keating potential to evaluate the energy of each atom. Using a combination of interatomic potentials and a specially optimized nonorthogonal tight-binding Hamiltonian we have created amorphous-crystal interfaces in silicon by performing melt and quench numerical experiments. The interfaces are about 7 Å thick. All measures of order we considered interpolated smoothly between the crystalline and amorphous phases. Slices of the sample along the interface reveal features analogous to dimers on the Si(001) surface and short crystal-like chains in the amorphous layer adjacent to the crystal. By comparing the energies of samples with and without interfaces we compute an interface energy of about 0.49 J/m$^2$, in good agreement with experimental evidence and other theoretical work.

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