Crystal growth of Ge$_2$Sb$_2$Te$_5$ at high temperatures

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Abstract

Phase-change materials (PCMs) have important applications in optical and electronic storage devices. Ge$_2$Sb$_2$Te$_5$ (GST) is a prototypical phase-change material (PCM) employed in state-of-the-art storage-class memories. In this work, we investigate crystallization of GST at temperatures 600–800 K by ab initio molecular dynamics. We consider large models containing 900 atoms, which enable us to investigate finite-size effects by comparison with smaller models. We use the metadynamics method to accelerate the formation of a large nucleus and then study the growth of the nucleus by unbiased simulations. The calculated crystal growth speed and its temperature-dependent behavior are in line with recent experimental work.

Introduction

Phase-change materials (PCMs) are employed in storage devices (rewritable optical discs, non-volatile electronic memories), which exploit their ability to undergo fast and reversible transitions between the crystalline and amorphous state.[1–3] Long-term information storage is possible, thanks to the optical and electronic contrast between the two phases, as well as the high stability of the amorphous state near room temperature. Phase-change storage-class memories have recently entered the market,[4,5] where subnanosecond crystallization processes are challenging, due to the small time- and length-scales accessible and the stochastic nature of nucleation. Considerable experimental and theoretical work has been devoted to the study of the strong temperature dependence of the dynamics of PCMs.[12–18] This is a key property that ensures fast crystallization at elevated temperatures and high stability of the amorphous phase at low temperatures. Such property has been linked to the high fragility of the supercooled liquid state, namely the fact that the viscosity behaves in a non-Arrhenius fashion.[12] The atomic processes involved in the high-temperature crystallization of PCMs have also been addressed by several computational studies based on density-functional–theory (DFT) molecular dynamics (the so-called ab initio molecular dynamics——AIMD) or accurate neural-network potentials fitted to DFT data.[19–33]

Upon fast crystallization of the glass, GST forms a metastable rocksalt phase,[34] in which Te atoms occupy one sublattice, while Ge, Sb, and vacancies occupy the second one in a random fashion. It has been shown experimentally[35–37] and theoretically[38–40] that the degree of disorder in rocksalt GST and similar compounds on the pseudobinary line can be tuned by thermal annealing or even pressure, so as to induce Anderson insulator–metal transitions induced by disorder.

GST is a nucleation-dominated PCM, implying that crystallization of an amorphous mark occurs via formation and subsequent growth of crystalline nuclei. AIMD simulations of these processes are challenging, due to the small time- and length-scales accessible and the stochastic nature of nucleation. In our previous work,[26] we considered models of GST containing about 500 atoms and performed two sets of simulations. In one set, we generated amorphous models inside a crystalline matrix (obtained by keeping two crystalline layers fixed during melting and quenching), and studied crystal growth at the planar crystal–amorphous interface. In the second one, we employed the metadynamics method[41] (a powerful and versatile enhanced-sampling technique) to accelerate the formation of postcritical nuclei and then monitored the growth of these nuclei during unbiased AIMD simulations. Here, we carry

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out large-scale AIMD simulations of crystallization of 900-atom models of GST at high temperatures ranging between 600 and 800 K. Similarly to our previous work, the crystallization kinetics is determined from a crystalline nucleus, which is created by means of metadynamics. The goal of this work is twofold: (a) address finite-size effects by comparing these simulations with previous work employing smaller models; (b) determine how temperature affects crystal growth and the size of the critical nucleus.

Methods

We perform large-scale AIMD simulations based on the “second-generation” Car–Parrinello scheme,[42,43] which is implemented in the Quickstep code of the CP2K package.[44] The generalized gradient approximation to the exchange-correlation potential[45] and the scalar-relativistic Goedecker pseudopotentials[46] are employed. A triple-zeta plus polarization Gaussian-type basis set is used to expand the Kohn–Sham orbitals, whereas plane waves with a cutoff of 300 Ry are used to expand the charge density. The Brillouin zone is sampled at the Γ point of the supercell and the periodic boundary conditions are imposed. The time step is 2 fs. The temperature is controlled by a stochastic Langevin thermostat. The simulations are conducted in the canonical ensemble (NVT) with a fixed density of 0.03173 Å lying between the experimental amorphous value (0.0300 Å) and the crystalline one (0.03283 Å). Fixing the density during the simulation is relevant to phase-change memory cells, where the presence of a crystalline matrix constrains the volume available during recrystallization of the amorphous mark. A purely amorphous model containing 900 atoms in a cubic 3.11 nm × 3.11 nm × 3.11 nm supercell is obtained by a melt-quenching procedure. In this procedure, the model is equilibrated at the melting temperature.

Results

In standard AIMD runs, the generation of a crystalline nucleus is computationally extremely heavy, since nucleation is a stochastic process. The first computational studies of crystallization kinetics of GST used very small cells containing <200 atoms,[19,20] which facilitated crystallization. Subsequent studies on larger systems were either focused on models containing a crystalline seed, which was fixed during the melt-quench procedure,[21] or were based on very long simulation times of the order of a few nanoseconds to enable the formation of large nuclei.[24,27] In Ref. 26, we employed metadynamics to create a postcritical crystalline nucleus inside a 460-atom model at $T = 600$ K. In this study, we follow a similar procedure and run a metadynamics simulation prior to the unbiased run of crystallization. Since we are only interested in accelerating the nucleation and do not aim at the estimation of free energies, we sample the phase space in a coarse and efficient manner using large Gaussians ($\omega = 0.5$ meV/atom, $\sigma_1 = 0.01$, $\sigma_2 = 3$ meV/atom, $\tau = 0.1$ ps) and employing the multiple walkers scheme[48] with seven independent walkers. In the multiple walker scheme, the sampling speed is increased by simultaneously running independent simulations that add Gaussians to the same shared bias potential. The metadynamics run is conducted at ~600 K, which is also one of the target temperatures of the unbiased simulations. The most important step for the success of a metadynamics simulation is the choice of appropriate CVs. In the case of nucleation, they should be able to discern atoms in a crystalline-like environment. One of the CVs in our metadynamics simulation is based on the local parameter $q_{4,\text{dot}}$, which is computed for each atom and measures the average correlation of the bonding arrangement between the atom and its neighboring atoms. It takes values close to one for atoms in a crystalline-like environment and close to zero for atoms in a disordered, amorphous-like environment. The top left panel of Fig. 1 shows the distributions of $q_{4,\text{dot}}$ in the amorphous and the crystalline phases: since the two distributions display almost no overlap, this quantity indeed allows to discriminate the two phases and is therefore well suited as a CV in nucleation studies. By setting a threshold value of 0.45, we define an atom to be crystalline-like if its $q_{4,\text{dot}}$ value exceeds this threshold. This definition is also used for the analysis of the crystal growth. The CV in our metadynamics simulation is obtained by averaging $q_{4,\text{dot}}$ inside a spherical region around a preselected atom. The size of the spherical region is chosen to be large enough to encompass a postcritical nucleus (which, in our previous work, was estimated to contain <150 atoms at 600 K), but as small as possible to minimize the finite size effects due to the periodic boundary conditions. As second CV, we use the potential energy of the system $U$.

As a result of the metadynamics simulations, some of the walkers sample the relevant region of the CV space where a crystalline nucleus starts to form. The bottom left panel of Fig. 1 shows the time evolution of the first CV for one such walker. On the right panel of Fig. 1, the formation of a crystalline nucleus...
inside the amorphous phase is visualized at a few selected snapshots corresponding to the times shown as black circles in the left bottom panel. Although size and shape of the nucleus can vary significantly due to the fluctuations of the CVs, we obtain a single, compact crystalline cluster at final stages of this walker.

Subsequently, we performed a long, unbiased simulation at \( \sim 600 \) K starting from a snapshot with a sizable crystalline nucleus containing about \( N_c \sim 175 \) crystalline-like atom. The crystallization process completed in \( \sim 810 \) ps and the recrystallized structure was in a metastable cubic phase. Three representative snapshots of the growing crystalline nucleus (solid balls) within the amorphous phase (transparent bonds) are depicted in Fig. 2. The crystalline nucleus is initially quasi-spherical; however, it becomes elongated after some transient period.

**Figure 1.** (Left) Distribution of \( q_{4\text{tot}} \) in the amorphous and crystalline phases (top) and evolution of the averaged \( q_{4\text{tot}} \) employed in the metadynamics simulation as first CV (see also main text) for a selected walker. The insets in the top panel show the typical atomic structures of amorphous and cubic rocksalt GST. (Right) Formation of a crystalline nucleus of GST (solid spheres and sticks) inside the amorphous phase (small, transparent sticks and spheres). Te atoms are green, Sb atoms are yellow, and Ge atoms are blue. The snapshots are taken at the corresponding time steps (black circles) of the biased simulations shown in the bottom left plot of the CV.

**Figure 2.** Unbiased simulations at \( \sim 600 \) K. Rapid crystal growth around the crystalline nucleus generated by metadynamics are observed. The snapshots are taken at \( t = 32, 421, 990 \) ps of the unbiased simulations.
The presence of a geometrically complex amorphous–crystal interface makes the computation of the crystal growth velocity \( v_c \) difficult. Considering a general crystalline volume \( V_c \) with a bounding surface \( S_{ac} \), which divides the crystalline and amorphous regions, we define the \( v_g \) as the average flux through the surface:

\[
v_g = \frac{1}{S_{ac}} \times \frac{dV_c}{dt}.
\]

Here we use the Voronoi tessellation of the supercell in order to define \( V_c \) and \( S_{ac} \). We obtain \( V_c \) by summing up the volumes of the Voronoi polyhedra of each crystalline-like atom. \( S_{ac} \) is the total area of the faces that are shared by Voronoi polyhedra of amorphous-like and crystalline-like atoms. We use the voro++ code\cite{50} to compute the Voronoi polyhedra in our models. In Fig. 3, we compare the 900-atom model with a 460-atom model from our previous work\cite{26} by plotting \( V_c, S_{ac}, \) and \( v_g \) as a function of time. This allows us to assess the finite size effects on the estimation of \( v_g \). The time derivative \( dV_c/dt \) is calculated by smoothing \( V_c \) with a Savitzky–Golay filter with a window size 2 ps. The computed velocities are only meaningful in the range in which \( V_c \) is increasing. The start and end of this range are indicated by thin, vertical lines in Fig. 3. The crystallization of the larger model containing 900-atom requires considerably longer simulation time as compared with the 460-atom models, because of the larger amorphous volume and the lower crystal growth velocity. After a closer inspection of the \( v_g \) curve for the 900-atom model, we can divide the crystallization of the 900-atom model into three distinct regimes: two regimes with a plateau of \( v_g \) values around \( \sim 0.5 \) and \( \sim 0.8 \) m/s and a third one with a peak value around \( \sim 1.4 \) m/s. In the first two regimes, the amorphous–crystalline interface area \( S_{ac} \) is increasing and the growth velocity is dominated by the derivative \( dV_c/dt \). In the first regime, the crystalline nucleus is small enough so that it is not connected with its periodic images or, at most, elongated along one direction. It corresponds to the crystalline nucleus shown in the left snapshot in Fig. 2. In this regime, the amount of amorphous volume is maximal and the finite size effects are minimal. Therefore, this region provides the best estimate for \( v_g \). In the second regime, the crystalline nucleus is connected with its periodic image, as in the middle snapshot in Fig. 2. The crystalline volume \( V_c \) of the nucleus becomes roughly comparable to the volume of the smaller 460-atom model of Ref. 26. The slope \( dV_c/dt \) of the 900-atom model in this regime is similar to that of the 460-atom at the initial stage, so that the values of \( v_g \) fall into a similar range close to \( \sim 1 \) m/s. In the final regime, the surface effects dominate, since \( v_g < S_{ac}^{-1} \) and there is an abrupt decrease in \( S_{ac} \) due to the vanishing amorphous region in the last stages of the crystallization process. Overall, we conclude that the finite size effects in the 460-atom models lead to a non-negligible overestimation of \( v_g \) and its values are roughly doubled by going from 900 to 460 atoms.

Next we address the crystal growth velocities at higher temperatures. In principle, independent amorphous models containing a postcritical crystalline nucleus at the target temperature should be considered to accurately estimate \( v_g \). Since such simulations would be computationally too demanding, we consider a simpler approach that allows us to approximately assess the crystallization kinetics at high temperatures. It has to be noticed, however, that the higher the temperature, the stronger the finite-size effects (for a fixed simulation box), since the size of the critical nucleus typically increases with temperature. The latter property stems from the fact that the interfacial energy is weakly dependent on temperature, whereas the chemical potential difference between the two phases (which, according to classical nucleation, is inversely proportional to the radius of the critical nucleus\cite{51}) decreases roughly linearly with temperature. Concretely, we perform a new set of simulations at \( \sim 700 \) and \( \sim 800 \) K starting from a snapshot at \( t = 222 \) ps of the crystallizing model at \( \sim 600 \) K. The choice of the starting configuration at a later stage is appropriate, because the size of the critical nucleus is larger at higher temperatures, as mentioned above. In our case, the crystalline nucleus already contains 245 atoms in this starting configuration. The simulation temperature is raised directly from \( \sim 600 \) K to the target temperatures. In Fig. 4, we show the evolution of the crystalline volumes \( V_c \) for the three sets of simulations (at \( \sim 600, \sim 700, \) and \( \sim 800 \) K, respectively) and the corresponding crystal growth velocities. At \( \sim 700 \) K, the crystalline cluster starts to grow after a short stabilization period. This indicates that its size is large enough to be postcritical. By contrast, the nucleus is not sufficiently large to grow at \( \sim 800 \) K. Instead, the cluster shrinks and dissolves quickly at this

![Figure 3. Evolution of the crystalline volume \( V_c \) (top), amorphous–crystal interface area \( S_{ac} \) (middle) and the corresponding crystal growth velocity (bottom) for 900- and 460-atom models of GST at \( \sim 600 \) K. Vertical lines indicate the start and end points of the time window in which \( V_c \) is increasing.](https://www.cambridge.org/core/='')
temperature. Therefore, we only plot \( v_g \) for the simulations at \( \sim 600 \) and \( \sim 700 \) K in Fig. 4. The initial crystallization regimes with minimal finite size effects are indicated with shaded rectangular regions. From this, we extract a crystal growth velocity \( v_g \sim 1.5 \text{ m/s} \) at \( \sim 700 \) K, which is approximately three times larger than the one at \( \sim 600 \) K. This trend is in line with kinetic measurements based on ultrafast differential scanning calorimetry,\(^1\) where the maximum growth rate is reached at temperatures close to 700 K.

**Conclusions**

In this work, we have studied the crystallization kinetics of a large model of GST containing 900 atoms starting from a crystalline nucleus of about 175 atoms. The crystalline nucleus was successfully generated from a metadynamics simulation with an appropriate set of CVs. Comparing the crystal growth velocities for this model with the ones obtained for a smaller 460-atom model in our previous work, we have assessed the finite size effects and found a reduced crystal growth velocity of \( \sim 0.5 \text{ m/s} \) in the larger model, which is roughly half of the value obtained for the 460-atom model. The crystal growth velocity at a higher temperature of \( \sim 700 \) K is estimated to be \( v_g \sim 1.5 \text{ m/s} \) from a single run with an initial configuration containing a crystalline cluster of 245 atoms. At even higher temperature of \( \sim 800 \) K, the latter crystalline nucleus is found to be unstable.

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