Computer Simulation of Precipitation Process in Si/Ge Amorphous Multi-Layer Films: Effects of Cu Addition

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We have simulated the precipitation process in an amorphous Si/Ge multi-layer film, with and without Cu addition, by a molecular dynamics method. Four specimens were prepared for this study: Si/Ge layers, Si/(Ge + Cu) layers, (Si + Cu)/(Ge + Cu) layers and Si/Cu/Ge/Cu layers. After the multi-layered films became amorphous, we tracked the movement of individual atoms at 1000 K, the annealing temperature. When Cu was present in the Ge layer or both the Si and Ge layers, the precipitation of nano-clusters was less than that in Cu-free Si/Ge layers. We think that the Cu atoms block the precipitation and make the Si and Ge become more stable in the amorphous state. If Cu atoms are not present in a layer, however, like the Si layer in Si/(Ge + Cu) and Si/Cu/Ge/Cu specimens, the precipitation of nano-clusters in the Cu-free layer is enhanced. Therefore we conclude that precipitation of nano-clusters in Si/Ge layers can be controlled by how Cu atoms are added to the amorphous Si/Ge system, and that this will improve the thermoelectric performance. [doi:10.2320/matertrans.M2013191]

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1. Introduction

People in Japan have become even more interested in renewable and conservation energy techniques after their experiences in the summer months following the massive earthquake and tsunami that struck northeastern Japan in March 2011 and led to the Fukushima nuclear accident, when the government implemented various energy saving measures (superlattice thin film, there was no sign of the MIC effect. In fact, high thermoelectric power (>1 mV K⁻¹) was observed in the superlattice thin film only when nanocrystals (≤10 nm) were present; however, limitations of the specimen preparation system and the use of B as the dopant made it difficult to prepare specimens with a precisely controlled nanostructure.

In this paper, we report our research using computer simulation on the effect of Cu dopant in Si/Ge amorphous multi-layer films. We used a molecular dynamics calculation method to obtain a meta-stable structure of the specimen. We used Cu-free and Cu-added multi-layer films of Si and Ge with an amorphous structure, and then we annealed the films to generate nano-clusters. We analyzed the behavior of the precipitation of nano-clusters by extracting the atoms with the same coordination number as the bulk in the obtained structure.

2. Simulation Method

In this study, we prepared four types of specimens: multi-layers of Si/Ge, Si/(Ge + Cu), (Si + Cu)/(Ge + Cu) and Si/Cu/Ge/Cu, where “+Cu” means that Cu atoms were added into Si and Ge layers as substitutional atoms and “/” indicates layers. The first, second and third multi-layer films contained the same numbers of unit cell, ten in the (100) and (010) directions and six in the (001) direction, where the lattice constant of the unit cell was set to be the average of the lattice constants of Si and Ge. The dimensions of the unit cell were 5.54 nm in the (100) and (010)
directions and 3.33 nm in the (001) stacking direction of the multi-layers. On the other hand, the fourth specimen had ten unit cells in the (100) and (010) directions and eight unit cells in the (001) direction. Due to the different number of unit cells in the (001) direction, this specimen was 4.43 nm long in the stacking direction of the multi-layers. The first specimen contained only Si and Ge, described as Si/Ge. In the second specimen, we added Cu to only the Ge layer at the ratio of Ge:Cu = 6:4, described as Si/(Ge + Cu). In the third specimen, we added 20 at% of Cu to both the Si and Ge layers, described as (Si + Cu)/(Ge + Cu). The fourth specimen, we formed a thin Cu layer at each Si and Ge interface, described as Si/Cu/Ge/Cu. Side views of the four types of specimens from the [010] direction are shown in Fig. 1.

To calculate the interaction between these atoms, we used the extended Tersoff potential.5 This is one of the families of potentials developed by Tersoff,6,7 and it is based on the concept of bond order: the strength of a bond between two atoms is not constant, but depends on the local environment. Moreover, this potential has been proven to be reliable, and it is widely used recently. The periodic boundary conditions were imposed in all the directions, (100), (010) and (001). The Verlet algorithm was used for the calculation of atomic movement,8 while the discrete time Δt was set as 0.3 fs. The book-keeping and cell-index methods were used to reduce computing time.9 First, all four specimens were melted at a temperature 5000 K and quenched at 4.2 K. The time durations of them were both 9 fs. After the multi-layered films became amorphous, we tracked the movement of individual atoms at 1000 K, the annealing temperature, for 7200 fs. For the atomic structures before and after the annealing process, the distributions of the four-coordination atoms were calculated.

3. Results and Discussion

In this study, we used Si/Ge as the reference system. We compared the four specimens to each other to determine how Cu affects the precipitation of nano-clusters in Si/Ge amorphous multi-layer films. After that, we counted the numbers of four-coordination atoms in every 0.2 nm interval of the layers from bottom to top along the (001) direction. Then, by checking which layer had more four-coordination atoms, we could identify which parts of the system had more nano-cluster precipitates. Takiguchi’s previous experimental results10 showed that the size of the precipitated nano-clusters in the Si/Ge amorphous multi-layered films would be around 6 nm in diameter. This size is too large compared to the present calculation. However, it is possible to deduce important information about the precipitation process in the multi-layer films.

The results were visualized to observe the precipitation of nano-clusters. Figure 2 shows the images of all atoms (left) and the four-coordination atoms (right), in amorphous state, for the specimens of (a) Si/Ge, (b) Si/(Ge + Cu) with Ge:Cu = 6:4, (c) (Si + Cu)/(Ge + Cu), with 20 at% Cu in both the layers and (d) Si/Cu/Ge/Cu. Si, Ge and Cu atoms are represented as green, red and yellow circles, respectively.

The multi-layer films became amorphous, we tracked the movement of individual atoms at 1000 K. The annealing temperature, for the same specimens as Fig. 2. It should be noted that the structures shown in Fig. 3 are at 1000 K.

In these figures, we can clearly see how the presence of Cu affects the precipitation of the four-coordination atoms before and after the annealing process. Comparing the results for Si/Ge shown in Figs. 2(a) and 3(a) with those for Si/(Ge + Cu) in Figs. 2(b) and 3(b), we can deduce the effects of Cu addition to the Ge layer in the Si/Ge multi-layer system. The density of the precipitated four-coordination atoms in the Ge layer in Si/(Ge + Cu) is lower than that in Si/Ge. However, there are slightly increases in the density of the precipitated four-coordination atoms in the Si layer of Si/(Ge + Cu). Comparing the results for Si/Ge with those for (Si + Cu)/(Ge + Cu) shown in Figs. 2(c) and 3(c), we can deduce the effects of Cu addition to both the layers of Si and Ge. The density of the four-coordination atoms in (Si + Cu)/(Ge + Cu) is lower than their density in Si/Ge for both the Si and Ge layers.
In Si/Cu/Ge/Cu, we added a thin Cu layer between each Si and Ge interface. As shown in Fig. 2(d) for the amorphous state specimen and Fig. 3(d) for the specimen after the annealing process, the densities of the four-coordination atoms in both the Si and Ge layers are much higher than in (Si + Cu)/(Ge + Cu). Just like Si/(Ge + Cu), it seems that Cu has enhanced the precipitation of four-coordination atoms in the Cu-free layers.

Figure 4 shows the distributions of the four-coordination atoms for the amorphous state and after the heat treatment process, for the four specimens, Si/Ge, Si/(Ge + Cu), (Si + Cu)/(Ge + Cu) and Si/Cu/Ge/Cu multi-layer films. Comparing the results of (Si + Cu)/(Ge + Cu) to Si/Ge, (Si + Cu)/(Ge + Cu) gives very low numbers of four-coordination atoms, and no significant change in Si and Ge layers is seen even after the heat treatment. It should be noted that the structures after heat treatment are at 1000 K.

Furthermore, we can see clearly the differences between the results of Si/(Ge + Cu) and Si/Ge, in which when the Cu dopants are present only in the Ge layer, the precipitation of nano-clusters decreases only in the Ge layer. This can be attributed to Cu atoms stabilizing Si and Ge in the amorphous state. In other words, Cu blocks crystallization during the heat treatment process. By placing a Cu layer at the interface between Si and Ge layers in Si/Cu/Ge/Cu, we get a result that is just like the result for Si/(Ge + Cu). In Fig. 4(d), at the Si and Ge layers distinct from the Cu layer, the number of four-coordination atoms after the annealing process increases significantly. Unlike in (Si + Cu)/(Ge + Cu) and Ge layers in Si/(Ge + Cu), these Cu layers do not block the precipitation of the four-coordination atoms in both the Si and Ge layers. Moreover, it seems these Cu layers enhance the precipitation process. Interestingly, this result is very similar to the result for the Si layer in Si/(Ge + Cu), in which
Cu dopants are present only in the Ge layer, not in the Si layer.

An actual experiment has been reported studying the role of gold (Au) in the thermoelectric properties of amorphous Ge/Au and Si/Au thin films. In the experiment, four specimens containing two different layers were prepared. Two of them had Ge and Au layers, with different thicknesses of the Au layer for each of the two specimens. The other two specimens had Si and Au layers, with different thicknesses of the Au layer for each of the two specimens. The results showed that Au enhanced the crystallization of Si and Ge. By increasing the Au composition, the crystallization was enhanced in both Si and Ge amorphous layers, and the grain size of the thin films, i.e., the diameter of the precipitated nano-cluster was also increased. In other words, by controlling the structure of the Si/Ge/Au multi-layer, the thermoelectric performance will be much improved.

In our experiment, we considered that Cu will behave just like Au because both elements have similar physical and chemical characteristics according to the similarity of their outermost electron shells. In Si/Cu/Ge/Cu, we can see that both Si and Ge show increased precipitation of the four-coordination atoms. Thus, by placing a thin layer of Cu at the interfaces of Si/Ge multi-layers, we can enhance the precipitation of the four-coordination atoms and can improve the thermoelectric performance. Unfortunately, we do not have details about the distribution sizes of the nano-clusters for the present calculation or the actual Au experiment. We plan to make a detailed analysis for both computer simulations and actual experiments.

### 4. Conclusion

We have simulated the precipitation process in Cu-free and Cu-added amorphous Si/Ge multilayer films by a molecular dynamics method. Four specimens were prepared: Si/Ge, Si/(Ge + Cu), (Si + Cu)/(Ge + Cu) and Si/Cu/Ge/Cu. From the analysis of the distribution of the four-coordination atoms before and after the heat-treatment, we clarified the following points. (1) In the Si/(Ge + Cu) system, Ge did not diffuse into the Si layer during annealing. (2) In the (Si + Cu)/(Ge + Cu) system, very low numbers of four-coordination atoms were found, and no significant change was seen in both Si and Ge layers even after the heat treatment. We attributed this to the fact that Cu stabilized the Si and Ge in the amorphous state. (3) In Si/Cu/Ge/Cu system, the thin layer of Cu at the interfaces of Si/Ge multi-layers enhanced the precipitation of the four-coordination atoms. (4) Moreover, the Si/(Ge + Cu) system showed the same behavior as the Si/Cu/Ge/Cu system regarding the precipitation process at the Si layer. Therefore we believe that precipitation of nano-clusters in Si/Ge can be controlled by changing the amount of Cu impurity and the way that we place Cu in the system. Hence, this technique will improve thermoelectric performance in Si layer.

### REFERENCES

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