In situ transmission electron microscopy analysis of conductive filament during solid electrolyte resistance switching

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An in situ transmission electron microscopy (TEM) analysis of a solid electrolyte, Cu–GeS, during resistance switching is reported. Real-time observations of the filament formation and disappearance process were performed in the TEM instrument and the conductive-filament-formation model was confirmed experimentally. Narrow conductive filaments were formed corresponding to resistance switching from high- to low-resistance states. When the resistance changed to high-resistance state, the filament disappeared. It was also confirmed by use of selected area diffraclometry and energy-dispersive x-ray spectroscopy that the conductive filament was made of nanocrystals composed mainly of Cu. © 2011 American Institute of Physics. [doi:10.1063/1.3593494]

Solid electrolyte resistance random access memory (ReRAM) has great potential as nonvolatile memory.1,2 It is scalable to a nanometer size because of its simple structure and its conduction paths at the nanometer scale.3 Many previous studies reported resistance switching in solid electrolytes such as AgGeS2,4–7 CuGeS5,6 Cu2S,8 Ag2S,9,10 Ta2O5,11,12 Cu-SiO2,12 and bilayer-types.13,14 The mechanism of resistance switching is attributed to the formation and disappearance of the conductive filament in the solid electrolyte. When a bias voltage is applied, the ions generated at the anode are thought to migrate toward the cathode where they undergo reduction and become metal atoms.8 Contrarily, an opposite bias voltage dissolves the metal composing filaments into the solid electrolyte.11 Therefore, an analysis of the conductive filament must provide important information for understanding this switching mechanism. Recently, conductive filament, which is formed in not only cation-type but also in anion-type electrolytes, was observed by scanning electron microscopy (SEM).5,15,16 However, no detailed experimental results to confirm the existence of the filament during the switching process have been reported. Therefore, in situ transmission electron microscopy (TEM) with simultaneous electrical measurements has attracted a great deal of attention.10,17–21

In this letter, we use this in situ TEM method to reveal the switching mechanism of a solid electrolyte. Real-time observations of the filament formation and disappearance process were performed with the TEM instrument. We also clarified the structure and composition of the filament by the use of selected area diffraction microscopy (SAD) and energy-dispersive x-ray spectroscopy (EDX).

For the in situ TEM experiments,21 commercially available Pt–Ir tips for scanning tunneling microscopy were further sharpened by ion milling. One Pt–Ir tip was used as the substrate and the other was used as a counter electrode. Cu–GeS thin films were deposited at room temperature by rf sputtering on Pt–Ir substrate. Since the substrate and the counter electrode of Pt–Ir were different in shape, the structure of Pt–Ir/Cu–GeS/Pt–Ir was asymmetric. The atomic composition of the sample layer was analyzed by means of EDX. The proportion of Cu:Ge:S was 4:4:2. The film thickness was between 8 and 60 nm, and no remarkable difference depending on thickness was recognized. The system was composed of a custom-made TEM holder with a PCI-controlled operating system.21–23 The TEM instrument used mainly was a JEM-2010 microscope (200 kV, Cc=0.5 mm), having a vacuum of about 10−5 Pa. The conduction properties were measured between the Pt–Ir counter electrode and the Cu–GeS/Pt–Ir. The best location of measuring I-V characteristics was selected in the fixed Cu–GeS/Pt–Ir sample by moving the Pt–Ir counter electrode. The TEM images were recorded using a CCD video camera.

Prior to the current-to-voltage (I-V) measurement, TEM observation of Cu–GeS was carried out. It was confirmed that the Cu–GeS layer was smoothly deposited on the Pt–Ir tips and that Cu–GeS was in an amorphous phase including the nanocrystals. This was also recognized in SAD patterns obtained from a wide area. The structure changes were not observed even during high resolution TEM observations (beam current density at the sample; about 170 fA/nm2). The in situ TEM experiments described below were performed with lower beam current. Thus, the influence of the electron beam should be negligible.

The I-V curve measured in the TEM instrument is shown in Fig. 1, in which the voltage is the potential of the substrate relative to the counter electrode. The voltage was swept from 0 to 7 V, from 7 to −2 V, and back to 0 V. The I-V curve shows a typical resistance switching characteristics with hysteresis. Figure 2 shows a set of TEM images corresponding to Fig. 1. The resistance changed from high resistance states (HRS) to low resistance states (LRS) occurred at around 1 V. Corresponding to this change, a small deposit appeared as shown in Fig. 2(b).

The growth of the deposit was almost stopped after current increase [Figs. 2(c)–2(g)] even though the current level and the resistance change up and down according to the voltage scan. In this process, even if the size of the deposit is almost constant, some changes were occurred around the deposit. It was confirmed that the diffraction spots of the SAD twinned continuously during the voltage scan. The current crossing at about 2 V and sudden current increase at about
-0.5 V may be caused by over accumulation of Cu to the needle side in the Cu–GeS film. The accumulation was released by applying slight negative voltage of -0.5 V.

The deposit suddenly shrank [Fig. 2(h)] at about -2 V, and completely disappeared [Fig. 2(i)]. The sample resistance turned back to HRS. The size of the deposit and the current value corresponded to each other. From these results shown in Figs. 1 and 2, it was concluded that the deposit observed in Fig. 2 constitutes the conducting path.

As shown in Fig. 3, the crystal structure of the deposit was studied by observing real-time SAD patterns during voltage application, which were from the wide area where the Pt–Ir substrate and the counter electrode were situated.

To minimize the influence of regions other than those near the deposit, a corner of the Cu–GeS layer was selected to be investigated [Fig. 3(a)]. The SAD pattern before applying voltage [Fig. 3(b)] was composed of a faint background and Debye-ring patterns. This corresponds to characteristics found in the TEM image described above, i.e., amorphous with nanocrystals. The clear spots in this pattern were caused by Pt–Ir. Using these spots as references, it was concluded that the faint rings corresponded to the 111, 220, and 311 reflections of the Ge. By applying 1 V to the substrate, a deposit appeared as seen in Fig. 2. Its width was about 40 nm at about 1 V. At this voltage, sharp spots appeared in the SAD pattern [Fig. 3(c)]. At this voltage, sharp spots appeared in the SAD pattern in addition to Fig. 3(b). They twinkled like stars in the night sky. This SAD pattern indicates that well crystallized nanocrystals were generated by the voltage application, and their orientation frequently changed during the growth of the deposit.

To analyze this process in detail, 1152 frames of video images totaling 35 s were superposed [Fig. 3(d)]. Relatively sharp spots that formed rings were recognized. These rings have d-values similar to those of the 111, 200, 220, and 311 reflections of the Cu. These results clearly show that main material forming the deposit was the nanocrystals of Cu or diluted Cu alloy with either Ge and/or S.

The EDX spectra of the area before and after positive voltage (1 V) application are shown in Fig. 4. The sample was the 8-nm-thick Cu–GeS where the deposit appeared. The beam size for the EDX measurement was about 10 nm. Therefore, in the obtained spectra [Figs. 4(a) and 4(b)], the Pt peaks of the electrodes were superposed. The intensity of the Cu peak greatly increased during the voltage application. By assuming the thin foil approximation, the estimated composition of the deposit for Cu:Ge:S was 7:2:1 [Fig. 4(b)], while it was 4:2:1 without the deposit [Fig. 4(a)]. Although this approximation gives rough estimation to discuss in de-
tail, it can be summarized by saying that the deposit was an agglomerate of crystals with a relatively large amount of Cu.

The conductive filament is assumed to be formed in the observed deposit. During filament formation, Cu ions dissolving in the GeS may move to the cathode and become metal atoms. The polarity dependence in this work may be attributed to the asymmetry of the electric field caused by the shape difference between the substrate and counter electrode as clearly seen in Fig. 2. We investigated various thicknesses of the Cu–GeS layers, and similar phenomena were observed in all cases.

In summary, we confirmed the resistance switching of Cu–GeS films by in situ TEM and confirmed the conductive-filament-formation model experimentally by means of real time observation. The formation and disappearance of the conductive filament were clearly observed in the switching operation. From SAD and EDX results, it was determined that the conductive filament consisted of nanocrystals composed mainly of Cu.

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FIG. 4. (Color online) (a) EDX spectrum before voltage application. The relative concentration was estimated as Cu:Ge:S=4:4:2. (b) EDX spectrum during 1 V application. The Cu peak was strong from the deposit. The relative concentration of the deposit was estimated as Cu:Ge:S=7:2:1.