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HRTEM Study on Resistive Switching ZrO$_2$ Thin Films and Their Micro-Fabricated Thin Films

Ying Li, Gaoyang Zhao, Zhibo Kou, Long Jin and Yajing Wang

Abstract

For the next-generation nonvolatile memory material, the most promising candidate is resistive random access memory (RRAM) which is nonvolatile memory with high density, high speed, and low power consumption. Resistive switching (RS) behavior had been reported in various films including transition metal oxides, perovskite, and chalcogenide. For further application, it is still a challenge to fabricate nanostructures of RS material. Micro-fabrication method involves traditional lithography, chemical etching, electron beam direct writing, nano-imprint, and so on. However, the procedure and the cost of these methods are relatively complex and high for semiconductors process. In this chapter, we demonstrate a method for fabricating sub-micro ZrO$_2$ lattice by using sol-gel method combined with laser interference lithography and micro-analysis with high-resolution transmission electron microscopy (HRTEM).

Researchers used all kinds of techniques to investigate the mechanism such as conductive atomic force microscopy, HRTEM, and scanning electrical microscopy. Despite the extensive research, much of the underlying mechanism is still unclear and controversial. This task can be accomplished only with advanced measurement and technique such as in HRTEM, local conductive atomic force microscopy, and so on. HRTEM sample preparation method for array dots is also discussed. In our research, the bipolar RS behavior can be observed successfully in this structure. And HRTEM observation was used to study the interface between the layers. RRAM unit consists of a conductive atomic force microscope tip as an anode, a ZrO$_2$ lattice dot as RS material, and a copper electrode as a cathode.

Keywords: Resistive switching, HRTEM, Sample preparation
1. Introduction

The development of information technology has brought rapid changes to people’s life. Resistive random access memory (RRAM), because of its fast response, high stability, low power consumption, with the traditional advantages of the semiconductor technology, has good compatibility, and according to scientists, is expected to become one of the next generations of memory [1-5]. In this chapter, by using shadow lithography and laser interference lithography, two kinds of methods to micro-fabricate the RS oxide film, it is suggested that the oxide film is suitable for micromachining process than using atomic force microscopy (AFM) in situ electrical measurement technique for fine-patterned films after the oxide film was studied by means of in situ micro area electrical performance. In this chapter, we used sol-gel method and chemical modification method for the preparation of a variety of ZrO₂ sol, and optimizing the good comprehensive properties of oxide RS material, through the analysis of the HRTEM studies of the oxide film [6-10].

2. Experiment

2.1. ZrO₂ thin films preparation

Photosensitive ZrO₂ films were fabricated with the metal alkoxide zirconium tetran-butoxide [Zr(OC₄H₉)₄], benzoylaceton (BzAcH), and ethanol (EtOH) in the molar ratio of Zr:BzAcH:EtOH at 1:1:40. After being mixed and refluxed for 2 to 3 hours, the photosensitive ZrO₂ solution was obtained. Then using dip-coating process, ZrO₂ organic films were formed on copper substrate. The copper substrates were fabricated by magnetic sputtering.

\[
(C_4H_9O)_4Zr + H_2 \rightarrow (C_4H_9O)_2Zr + C_4H_9OH
\]

2.2. ZrO₂ lattice preparation

ZrO₂ lattice was fabricated by double exposures of two-beam laser interference through 90° rotation of the sample in its own plane between twice exposures with 350 nm Krypton ion laser. After laser exposure, the exposed part of ZrO₂ film is not soluble in leaching solution while the un-exposed part is still soluble. Therefore, ZrO₂ lattice can be obtained [11-14].

The UV–visible absorption spectra were measured and these spectra of the ZrO₂ films were dip-coated on quartz glass substrate. It was known that an absorption peak of BzAcH at 308 nm was due to the π-to-π* transition. These peaks shift to 338 nm in the spectra of the ZrO₂ gel films. This red shift indicates the chelation between Zr⁺⁺ and BzAcH, and the reaction re-
results in the partial replacement of the alkoxy group by β-diketone ligand [see Eq. (1)]. Fig. 1 also shows the AFM image of ZrO$_2$ films irradiated by a UV Krypton ion laser. The intensity of absorption peaks at around 338 nm decreases step by step with increasing irradiation time from 0 second to 400 seconds. These decreasing spectra showed that the chelate rings were dissociated by UV irradiation.

Figure 1. Observation of ZrO$_2$ lattice dot by AFM measurement systems.

2.3. I-V measurement with ZrO$_2$ thin films

Before the local conductive AFM (LC-AFM) measurement of the ZrO$_2$ lattice dots, a traditional I-V probe station measurement was carried out with the normal ZrO$_2$ thin film which
is made of the photosensitive ZrO$_2$ solution. This RS structure is formed with platinum (top electrode), ZrO$_2$ photosensitive film (RS layer), and copper (bottom electrode). The platinum top electrode is made by sputtering with shadow mask. The pattern of the shadow mask is periodic lattice dot with a diameter of 2 mm.

2.4. I-V measurement with ZrO$_2$ lattice

This LC-AFM measurement system consists an atomic force microscope (AFM, Seiko Instruments Inc, SPI3800N/SPA-400) with a resolution of 0.1 nm in x–y directions and 0.01 nm in the z direction and a Keithly 2400 I-V source meter (Keithly, USA.). The silicon tips (PPP-CONT, Nanosensors) were coated by sputtering deposition with a 25 nm thick double layer of chromium and platinum/Iridium (Pt/Ir) with an initial 20 nm radius. In our LC-AFM measurement study, RRAM unit consists of a conductive atomic force microscope tip as an anode, a ZrO$_2$ lattice dot as RS material, and a copper electrode as a cathode. In order to identify the RS properties of ZrO$_2$ array dot, the Pt/Ir conductive atomic force microscope tip is tightly positioned at the center of one of the dots. During the LC-AFM measurement, Pt/Ir conductive probe and the base of the LC-AFM are connected to Keithly 2400 meter to apply the excitation signal and record voltage and current response of the RS array dots. When the voltages from Keithly 2400 meter are applied to the Pt/Ir conductive AFM tip and the bottom electrode of the sample, the current flow occurs through the cantilever and goes back to the Keithly 2400 meter. Then a local I-V curve can be obtained.

In order to solve the influence from huge contact resistance between the Pt/Ir conductive atomic force microscope tip and a single ZrO$_2$ array dot, a platinum layer was spread on the whole ZrO$_2$ lattice by sputtering. In this way, the contact can be improved.

2.5. HRTEM sample preparation

The HRTEM observation was carried out by HRTEM (JEOL 3010, Japan). The diagram of sample preparation process is shown in Fig. 2. In Fig. 2(a), we cut some tiny part from the platinum-covered sample which can achieve I-V curves by LC-AFM systems for HRTEM observations.

Two tiny parts were pasted by AB glue face-to-face. AB glue should be used as less as possible because the more AB glue used the worse ion milling would be. And this unit was milled by hand in x direction as smooth as possible. When the thickness in z direction decreases down to less than 1 µm, this unit can be pasted to nickel (Ni) grid. This will be ready for Argon ions milling. A cross-section sample of Pt/ZrO$_2$ lattice dot/Cu unit was milled by the low-angle Argon milling machine (Gatan, USA). The milling angle was less than 8°. Microstructural analysis was done on the basis of HRTEM observations. It is well known that the preparation of good specimens for high-resolution transmission microscopy is a very important process. The milling angle of Argon ions was less than 8° for the reasons of the damage from Argon ions. The transparent area to electrons can be widened if the Argon ions milling angle was small. However, it cannot mill if the milling angle was less than 3°. The ideal milling angle of Argon ions was between 4° and 8°.
3. Discussion and results

3.1. Normal probe station I-V curves

For the electrical measurements, probes contact top electrode (Cu) and bottom electrode SnO$_2$:Sb (ATO), respectively. The bottom electrode was grounded. In Fig. 3, it shows the I-V characteristics of the Cu/ZrO$_2$/ATO structure after fifty times of I-V measurements cycles. The current compliance (CC) is applied to 10 mA during I-V sweep. When the voltage goes to 0.5 V, the current increases rapidly which means the OFF state switches to the ON state. During the I-V sweep at negative side, when the voltage goes to -1.4 V, the current decreases rapidly which means the ON state switches to the OFF state. This behavior can be repeated more than fifty times.

Figure 2. Illustration of HRTEM samples preparation process.

3. Discussion and results

3.1. Normal probe station I-V curves

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Figure 3. I-V curve of the Cu/ZrO$_2$/ATO structure after fifty cycles.

Bipolar RS characterization was stable for ZrO$_2$ thin film in the structure unit. Normally, the voltage sweeps start from 0 V and go to the positive side (1 V) and goes back to 0 V. Resistance switches from high resistance state ($R_{H}$) to low resistance state ($R_{L}$) and it is called SET process. And then the voltage sweeps from 0 V to the negative side (-2 V) and goes back to 0 V. Resistance switches from $R_{L}$ to $R_{H}$ and it is called RESET process. When the voltage swept at -0.5 V, $R_{L}$ was 70 Ω and $R_{H}$ was 100 KΩ. This behavior can be repeated hundreds of times. The CC is applied with 10 mA. The highest ratio of $R_{H}$:$R_{L}$ can reach to $10^4$.

During the RESET operation, the interesting thing we found is that $R_{H}$ varied when we changed the RESET voltage. $R_{H}$ decreased when the RESET voltage value decreased. For I-V curve in Fig. 3, voltage sweep started from 0 V to the -2 V and then from -2 V to 0 V during the RESET process. When the voltage swept at -0.5 V, $R_{L}$ was 100 Ω and $R_{H}$ was 250 KΩ. This behavior can be repeated more than several times. The CC is applied with 10 mA. The highest ratio of $R_{H}$:$R_{L}$ can reach to $10^4$.

3.2. Local I-V curves by LC-AFM measurement systems

During the LC-AFM measurement, the parameters of I Gain and P Gain should be adjusted. The value of P Gain should be set as one-third of I Gain value. The Pt/Ir conductive coating could be switched off when the value of Force Reference is too small. This value should be set from -1 nm to -4 nm. The sweep voltage can be set from -25 V to 25 V. The height of array dot A ($\Delta$Z) is about 61 nm and dot B ($\Delta$Z) is about 58 nm. The diameter of dot A is about 1.2 µm and that of dot B is about 1.1 µm. Local I-V curves are measured while the LC-AFM image was scanning. The CC is applied to 100 µA during I-V sweep. When the voltage goes to
18 V, the current increases rapidly which means the OFF state switches to the ON state. During the I-V sweep at negative side, when the voltage goes to -5.0 V, the current decreases rapidly which means the ON state switches to the OFF state. In order to make sure that the array dot is not shifted during the LC-AFM measurement, an AFM image scan is very necessary after the local I-V measurement is finished.

A platinum film of 30 nm thickness was sputtered on ZrO$_2$ lattice dot. Fig. 4 is a local I-V curve which is also measured while the LC-AFM image was scanning. The CC is applied to 100 µA during I-V sweep. When the voltage goes to 0.8 V, the current increases rapidly which means the OFF state switches to the ON state. During the I-V sweep at negative side, when the voltage goes to -2.2 V, the current decreases rapidly which means the ON state switches to the OFF state.

![Image of I-V curve](image_url)

**Figure 4.** A local I-V curve which is measured while the LC-AFM image was scanning.

### 3.3. HRTEM observations

We took one unit of samples for TEM observations in which the ZrO$_2$ thin film in the structure was annealed at 300°C. Microstructural analysis has been done on the basis of TEM observations. Fig. 5 shows the interface image of Cu/ZrO$_2$/ATO structure in which the ZrO$_2$ thin film was annealed at 300°C. The bottom electrode (BE) is ATO film and the thickness is about 400 nm. The ZrO$_2$ work layer is about 200 nm thick and it is amorphous according to the selected area diffraction pattern. In Fig. 5, it is shown that copper ions are penetrated into the ZrO$_2$ layer. It can be proved by the Fast Fourier Transform patterns. According to the filament formation mechanism, resistance reduction in the devices is due to the existing
copper to form conducting copper-rich pathways. An opposite bias takes the existing copper back to the copper electrode to its high resistance state [15-19].

Figure 5. HRTEM image of Cu/ZrO$_2$/ATO structure.

Fig. 6(a) shows the interface image of Pt/ZrO$_2$ lattice dot/Cu sandwiched structure. Two lattice dots were seen in this figure, which are labeled as A and B. These two dots were not similar in size. This phenomenon was also seen in the rest of the HRTEM sample observations. The diameter of dot A is around 800 nm while that of dot B is around 700 nm. This is not agreeing with the LC-AFM image results. Through LC-AFM image observation, it is shown that the ZrO$_2$ array dots are even and in the same diameter range of about 1 µm. This is because when the HRTEM sample was milling manually in the first stage, it was difficult to keep the milling plane vertical to the ZrO$_2$ array plane. This is shown in Fig. 7(a) as the ideal case. It was always tilted a little bit which was perhaps due to the milling stage, sandpaper, or movement by hands. This is shown in Fig. 7(b) which is the practical case. For the conventional TEM specimen preparation techniques such as polished milling, the milling direction is controlled only by hands. Therefore, in the future HRTEM observation, an automatic monitor for sample milling system is necessary.

This HRTEM image is accepted for interface analysis. In inset of Fig. 6(a), it shows the selected area diffraction pattern of platinum, ZrO$_2$, and copper which are corresponding to polycrystalline, amorphous, and polycrystalline behaviors, respectively. Fig. 6(a) shows the HRTEM image of copper layer and Fig. 6(c) shows the interface between ZrO$_2$ array dot and platinum layer. In HREM images, copper ions filaments were not found in the amorphous ZrO$_2$ array dot. Therefore, it is possible that the Redox process could have happened in the interface between copper layer and ZrO$_2$ array dot.
Figure 6. HRTEM interface image of Pt/ZrO$_2$ lattice dot/Cu structure.
4. Conclusion

We fabricated ZrO$_2$ thin films after annealing at 300°C through sol-gel deposition. Reproducible I-V curves can be obtained with these samples at room temperature (300 K). The ZrO$_2$ thin film annealed at 300°C in Cu/ZrO$_2$/ATO device can also be operated in RS sweeps cycles. In summary, we fabricated TiO$_2$ thin films by sol-gel deposition. Successful Cu/ZrO$_2$/ATO structure device in which ZrO$_2$ thin film was calcined at 300°C can be obtained. The bipolar RS behavior was observed and the ratio of $R_{off}/R_{on}$ can be reached to $10^4$. We also investigated the switching device of Pt/ZrO$_2$ lattice dot/Cu sandwiched structure by HRTEM.

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