





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# Electroforming-free threshold switching of NbO<sub>x</sub>-based selector devices by controlling conducting phases in the NbO<sub>x</sub> layer for the application to crossbar array architectures

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Bipolar threshold switching characteristics, featuring volatile transition between the high-resistance state (HRS) at lower voltage than threshold voltage ( $V_{th}$ ) and the low-resistance state (LRS) at higher voltage irrespective of the voltage polarity, are investigated in the Nb(O)/NbO<sub>x</sub>/Nb(O) devices with respect to deposition and post-annealing conditions of NbO<sub>x</sub> layers. The device with NbO<sub>x</sub> deposited by reactive sputtering with 12% of O<sub>2</sub> gas mixed in Ar shows threshold switching behaviors after electroforming operation at around +4 V of forming voltage ( $V_f$ ). On the other hand, electroforming-free threshold switching is achieved from the device with NbO<sub>x</sub> deposited in the reduced fraction of 7% of O<sub>2</sub> gas and subsequently annealed at 250 °C in vacuum, thanks to the increase of the amount of conducting phases within the NbO<sub>x</sub> layer. Threshold switching is thought to be driven by the formation of a temporally percolated filament composed of conducting NbO and NbO<sub>2</sub> phases in the NbO<sub>x</sub> layer, which were formed as a result of the interaction with Nb electrodes such as oxygen ion migration either by annealing or electrical biasing. The presence of a substantial amount of oxygen in the Nb electrodes up to ~40 at%, named Nb(O) herein, would alleviate excessive migration of oxygen and consequent overgrowth of the filament during operation, thus enabling reliable threshold switching. These results demonstrate a viable route to realize electroforming-free threshold switching in the Nb(O)/NbO<sub>x</sub>/Nb(O) devices by controlling the contents of conducting phases in the NbO<sub>x</sub> layer for the application to selector devices in high-density crossbar memory and synapse array architectures.

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## 1. Introduction

In order to fabricate high-density two-terminal resistive random access memory (RRAM) and artificial synapses, crossbar array architectures have been employed for the benefit of their  $4F^2$  cell sizes ( $F$  is the minimal feature size).<sup>1</sup> In the crossbar array architectures, it is essential to minimize unintended leakage current through neighboring unselected cells, called sneak path leakage current, to avoid erroneous operation as well as to reduce operation power.<sup>2–4</sup> Various approaches have been pursued to minimize sneak path leakage current by integrating additional selector devices such as transistors,<sup>5</sup> diodes,<sup>6</sup> tunnel barriers,<sup>7</sup> mixed ionic electronic conduction (MIEC) devices<sup>8,9</sup> having highly non-linear current–voltage characteristics, and threshold switching selectors having abrupt switching between

insulating and conducting states.<sup>10–16</sup> Among these, the use of two-terminal selectors has advantages of high scalability due to small feature size and the same two-terminal structure with memory or a synapse cell that enables it to be directly integrated by stacking devices.

As two-terminal selectors, the threshold switching selectors feature to have an abrupt increase of current when the applied voltage exceeds a threshold voltage ( $V_{th}$ ), resulting in switching operation. They have a reversible and volatile change of resistance between a high-resistance state (HRS) at a lower voltage than  $V_{th}$  and a low-resistance state (LRS) at a higher voltage than  $V_{th}$ , which is analogous to resistance change in non-volatile RRAM cells except that the resistance change is volatile in nature upon removing applied voltage due to temporary formation of the conducting filament.<sup>16–20</sup> The analogous nature of threshold switching selectors and RRAM cells enables them to be readily integrated and operated by tuning the conditions to obtain volatile unstable filament formation in threshold switching selectors and non-volatile stable filament formation in RRAM cells.

One route to realize threshold switching selectors is the use of diffusive memristors operated by temporal and diffusive

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dynamics of filament evolution consisting of active metal species such as Ag and Cu in electrolyte oxide layers.<sup>17,21,22</sup> The filament is formed by electrochemical metallization for temporary switching from HRS to LRS only while the voltage higher than  $V_{th}$  is applied, and it is spontaneously dissolved after removing the applied voltage. Another route is to form the temporary filament composed of oxygen vacancies, corresponding to valence change memory (VCM)-type RRAM.<sup>23–25</sup> In this device, oxygen ions migrate from the resistive switching oxide layer to oxygen-reactive electrode, then oxygen vacancies in the oxide layer coalesce and form a conducting filament. In contrast to VCM-type RRAM, the filament is unstable and suddenly dissociated when the applied voltage is removed, thereby leading to volatile threshold switching.

In this study, the VCM-type threshold switching characteristics particularly with electroforming-free (also called “forming-free”) operations were demonstrated with a simple metal-insulator-metal (MIM) structure of Nb/NbO<sub>x</sub>/Nb, where niobium (Nb) and niobium oxide (NbO<sub>x</sub>) play the role of oxygen-reactive electrode and resistive switching layer, respectively. To date, NbO<sub>x</sub> has been examined as promising selector material with the structures such as W/NbO<sub>x</sub>/TiN,<sup>14</sup> Pt/NbO<sub>x</sub>/Pt,<sup>15</sup> Pt/NbO<sub>2</sub>/Nb<sub>2</sub>O<sub>5</sub>/Pt,<sup>12</sup> and Ti/NbO<sub>x</sub>/Ti.<sup>26</sup> However, most of the devices generally require the forming operation to obtain consequent threshold switching characteristics. Distinct from those previous reports, in this study, Nb was used as oxygen-reactive electrodes for the threshold switching of NbO<sub>x</sub> switching layer, particularly for the forming-free threshold switching. In addition, it is expected to have reliable interfaces and to simplify fabrication procedures for reduced processing budget and increased production yield by using Nb electrodes for NbO<sub>x</sub> switching layer. Electroforming-free threshold switching could be achieved by controlling deposition conditions and employing the post-deposition annealing process of NbO<sub>x</sub> layer to obtain proper contents of conducting NbO and NbO<sub>2</sub> phases mixed with insulating Nb<sub>2</sub>O<sub>5</sub> phase.

## 2. Experimental

The threshold switching devices were prepared with the structure of Nb(O)/NbO<sub>x</sub>/Nb(O) as follows. A 70 nm-thick Pt with a 30 nm-thick Ti adhesion layer was deposited on SiO<sub>2</sub>/Si substrate by e-beam evaporation under the base pressure lower than 10<sup>−6</sup> Torr. Then, a 90 nm-thick Nb(O) bottom electrode (BE) was deposited by direct-current (dc) magnetron sputtering with a 99.995% pure Nb target in a diameter of 2 inches at room temperature in an Ar ambient. The base pressure was in the range of 10<sup>−6</sup> Torr and a working pressure was 10 mTorr. Its deposition rate was 9 nm min<sup>−1</sup>. Although the electrode was deposited in an Ar ambient, it was deposited as the mixture mostly of Nb and NbO phases with oxygen content of almost 40 at%, as discussed layer. Therefore, the electrodes are presented herein as Nb(O). A 90 nm-thick NbO<sub>x</sub> resistive switching layer was subsequently deposited by dc magnetron reactive sputtering using the same Nb target in 7% or 12% O<sub>2</sub> gas mixed in Ar, which is named hereafter 7%-device and 12%-device with these switching layers, respectively. The deposition rates of 7% and 12% NbO<sub>x</sub> layers were 14.3 and

16.7 nm min<sup>−1</sup>, respectively. The other conditions for the deposition were maintained to be same for each other. A Nb(O) top electrode (TE) with a diameter of 100 μm was finally deposited using a shadow mask. In addition, the device with NbO<sub>x</sub> deposited in 7% O<sub>2</sub> gas and subsequently annealed at 250 °C in the deposition chamber at a pressure in the range of 10<sup>−4</sup> Torr for an hour right after deposition of Nb(O) TE was prepared as named 7% annealed-device. The threshold switching characteristics were examined using Agilent 4156B and Keithley 4200A-SCS semiconductor parameter analyzers. The voltage was applied to Nb(O) TE and the Nb(O)/Pt BE was kept grounded. The microstructures of NbO<sub>x</sub> layers and device structures were analyzed by using scanning electron microscope under acceleration voltage of 10 kV (SEM, Quanta 200FEG) and transmission electron microscope under acceleration voltage of 200 kV (TEM, ARP200P). TEM specimens were prepared by focused ion beam method with Ga ions (FIB, Helios NanoLab 450/FEI). The chemical bonding states and compositions of the layers were analyzed by using X-ray photoelectron spectroscopy (XPS, K-alpha) and Auger electron spectroscopy under acceleration voltage of 3 kV (AES, PHI-700).

## 3. Results and discussion

Fig. 1 shows the current-voltage ( $I$ - $V$ ) sweep curves of three Nb(O)/NbO<sub>x</sub>/Nb(O) devices with respect to preparation conditions. Fig. 1(a)–(c) show the  $I$ - $V$  sweep curves of 12%-, 7%-, and 7%-annealed devices, respectively. The voltage sweep was performed by increasing applied voltage amplitude in a linear staircase manner and the current was measured with an integration time of 0.64 ms. First, the 12%-device required forming operation by sweeping voltage 0 → +5 → 0 V as shown at the inset of Fig. 1(a). The compliance current for the forming operation was set to be 0.02 A to prevent excessive filament growth. At forming operation, the current was abruptly increased from ~10<sup>−3</sup> to 0.02 A at the forming voltage ( $V_f$ ) of around +4 V during forward sweep of 0 → +5 V and decayed back to ~10<sup>−3</sup> A at +1 V during backward sweep of +5 → 0 V. In general, the forming operation is required to make either volatile or nonvolatile conducting path through insulating layers to alternately form and rupture filament at subsequent voltage application steps for threshold switching or memory operations, respectively. Even at forming operation, the  $I$ - $V$  curve shows volatile threshold switching, implying that the filament formed at the forming operation is temporarily sustainable as volatile filament. As will be further discussed with Fig. 3, the resistivity of the NbO<sub>x</sub> layer at HRS (off-state) was substantially decreased by up to four orders of magnitude of the increased current level after forming operation. It implies that the layer became to have partially formed filament or abundant conducting phases or defects that provide the conducting path by the forming operation. As shown in Fig. 1(a), after forming operation, the device shows threshold switching behaviors upon alternatively repeating positive voltage sweeps 0 → +2 → 0 V and following negative voltage sweeps 0 → −2 → 0 V at the compliance current of 0.1 A. During repeated positive and negative voltage sweeps by total forty times, the threshold switching characteristics remained stable at both polarities



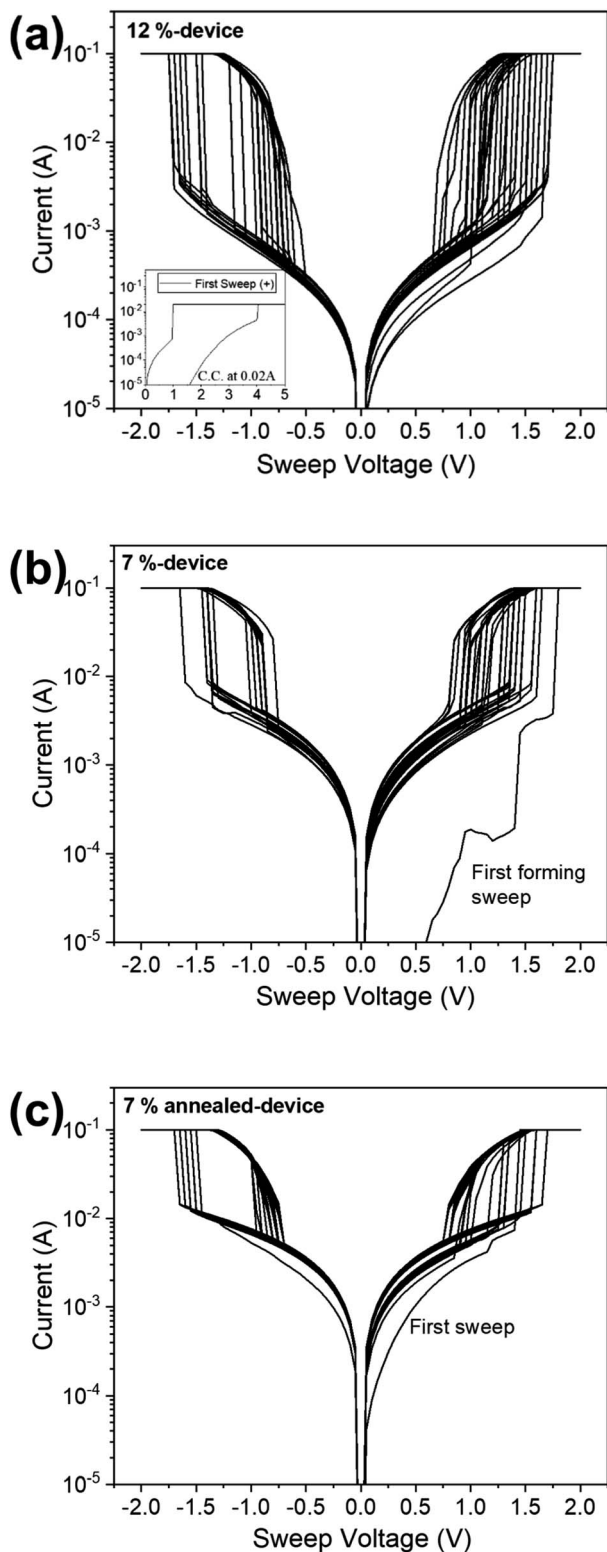


Fig. 1  $I$ - $V$  sweep curves of (a) 12%-device, (b) 7%-device, and (c) 7% annealed-device.

with  $V_{th}$  of around  $\pm 1.2$  V for an abrupt increase of current and the hold voltage ( $V_{hold}$ ) of around  $\pm 0.9$  V at which the current descended back abruptly. The HRS/LRS ratio is measured to be in the range of 30  $\sim$  100 at the  $V_{th}$ .

The 7%-device with reduced  $O_2$  gas mixture required also forming operation at the  $V_f$  of around +1.8 V during the first voltage sweep of  $0 \rightarrow +2 \rightarrow 0$  V at the compliance current of 0.1 A as shown in Fig. 1(b). Its compliance current for the forming operation was set to be higher than that of the 12%-device because of the higher conductivity of the  $NbO_x$  layer in the 7%-device. The  $V_f$  of about +1.8 V in the 7%-device is much smaller than around +4 V in the 12%-device because the  $NbO_x$  layer in the 7%-device is initially more conducting with a higher content of conducting phases, as confirmed from XPS analysis (Fig. 11), to readily form the filament. After the first sweep, the device shows threshold switching with  $V_{th}$  of around  $\pm 1.2$  V and  $V_{hold}$  of around  $\pm 0.9$  V during repeated  $0 \rightarrow \pm 2 \rightarrow 0$  V sweeps. It is noted that the HRS/LRS ratio ranges from 10 to 20, which is smaller than that of the 12%-device due to higher conductivity of the  $NbO_x$  layer at the HRS in the 7%-device.

Compared to these devices, the  $I$ - $V$  curves of the 7% annealed-device in Fig. 1(c) show notable feature of forming-free threshold switching. The threshold switching is observed even at the first sweep curve of  $0 \rightarrow +2 \rightarrow 0$  V as to be almost identical to those in the following sweeps. Except for the forming-free operation, other properties such as the values of  $V_{th}$ ,  $V_{hold}$ , and the HRS/LRS ratio are nearly the same with those of the 7%-device shown in Fig. 1(b). The forming-free operation in the 7% annealed-device comes from the interaction between Nb(O) electrodes and  $NbO_x$  switching layer such as oxygen migration during the post-annealing at 250  $^{\circ}$ C. The interaction such as oxygen migration depletes oxygen from the  $NbO_x$  layer to increase the content of conducting NbO and  $NbO_2$  phases, as confirmed with XPS analysis in Fig. 11. As schematically illustrated in Fig. 2, the post-annealing drives oxygen atoms to the Nb(O) electrode and increases the concentration of oxygen vacancies in the  $NbO_x$  layer (Fig. 2(a) and (b)), and these vacancies form the volatile filament as being aligned under the electric field (Fig. 2(c)). It verifies that the post-annealing to migrate oxygen atoms to electrodes and subsequent increase of oxygen vacancy concentration (content of conducting phases) could be a simple and viable route of forming operation alternative to electric biasing. The mechanism of threshold switching is further discussed later with the results of structure, composition, and bonding state analyses.

Fig. 3(a)–(c) show the  $I$ - $V$  curves of randomly chosen seven different cells at the first two voltage sweeps of the 12%-, 7%-, and 7% annealed-devices, respectively, to examine the forming characteristics. In the 12%-device, as shown in Fig. 3(a), the forming took place at around  $\pm 4$  V of the first sweep. The HRS current before forming was about  $10^{-7}$  A at +1 V, which was the lowest among those of the three devices. Then, it was increased to the range of  $10^{-4}$  A at following sweep, and  $V_{th}$  was around +1 V. The measured seven devices showed a narrow distribution of switching parameters such as the currents at HRS and LRS, the values of  $V_f$  and  $V_{th}$ . The  $I$ - $V$  curves of the 7%-device are found to be rather dispersed at the first sweep for the forming operation with deviated  $V_f$  values and multiple current changes. Then, threshold switching behaviors are stabilized at the second sweep. The  $I$ - $V$  sweeps of the 7% annealed-device show the forming-free and uniform threshold switching



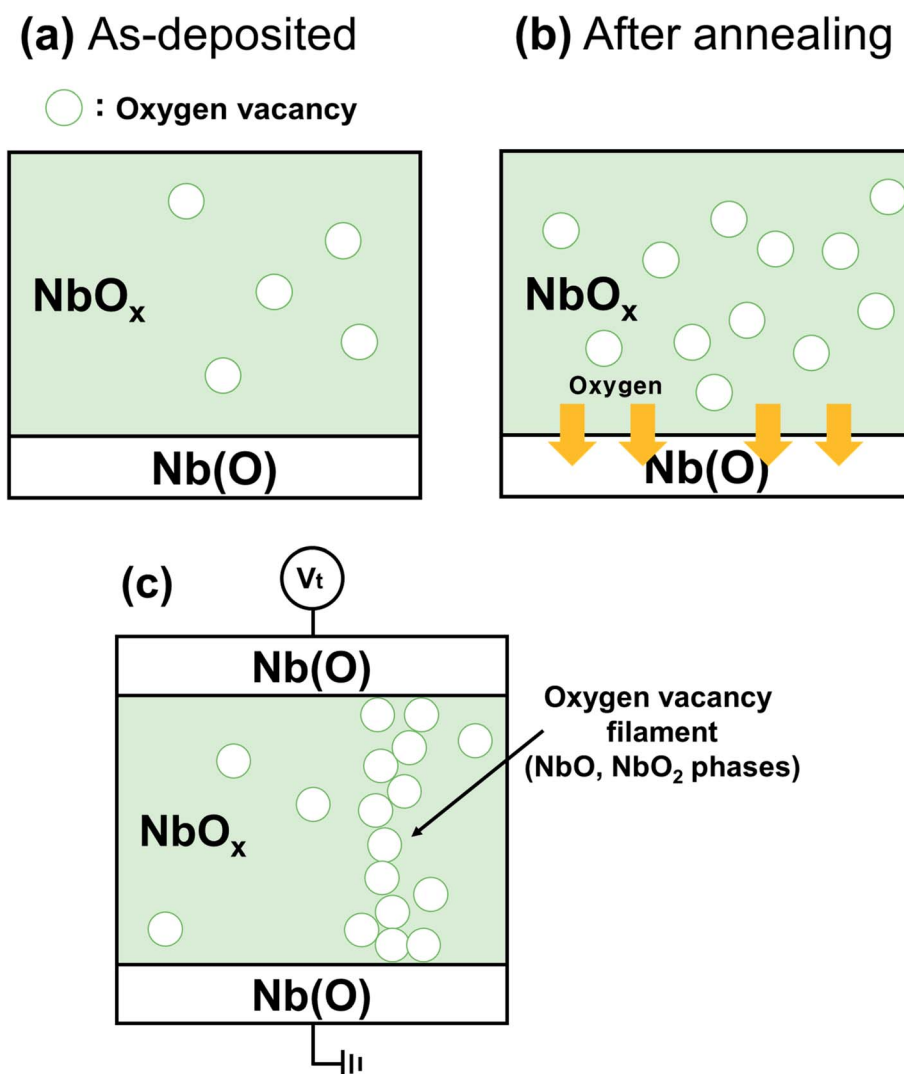


Fig. 2 Schematic illustration of forming-free threshold switching mechanism in the 7% annealed-device associated with the increased oxygen vacancy formation by annealing and the field-induced alignment of vacancies for filament formation: (a) as-deposited state, (b) after annealing to increase oxygen vacancy concentration, *i.e.*, increase of NbO and NbO<sub>2</sub> phases, and (c) alignment of vacancies for temporary formation of conducting filament.

characteristics, where the curves at the first and the second sweeps are almost same with a little higher  $V_{th}$  of +1.5 ~ +2.0 V at the first sweep than about +1 V at the second sweep.

Fig. 4 compares current levels at HRS and LRS of the 12%-, 7%-, and 7% annealed-devices when the voltage sweeps for the threshold switching operations were repeated eighty times to assess endurance properties. The currents values right before and after threshold switching at  $V_{th}$  are presented for those of HRS and LRS, respectively. As shown in Fig. 4(a), the 12%-device has a lowest HRS current of  $\sim 5 \times 10^{-4}$  A due to its lowest conductivity of the NbO<sub>x</sub> layer, resulting in the highest HRS/LRS ratio of  $\sim 200$ . However, the device experienced a hard break-down as being stuck to LRS after repeating forty times of voltage sweeps possibly due to the overgrowth of filament. The gradual increase of current manifests continued growth of filament during the repetition of sweeps. On the other hand, both the

7%- and 7%-annealed devices exhibit the reliable threshold switching up to eighty times of operations, although there is a trade-off with the smaller HRS/LRS ratio of around 10 ~ 30 due to their higher HRS currents in the range of  $10^{-3}$  A.

The uniformity of threshold switching characteristics was evaluated with cycle-to-cycle and cell-to-cell variations of  $V_{th}$  and  $V_{hold}$  values. Fig. 5 presents cycle-to-cycle distribution of  $V_{th}$  and  $V_{hold}$  values extracted from a hundred switching cycles as cumulative probability functions for the 12%-, 7%-, and 7% annealed-devices. It is found that all three devices have  $V_{th}$  and  $V_{hold}$  values within narrow ranges. The mean value ( $\mu$ ) and standard deviation ( $\sigma$ ) of  $V_{th}$  of the 12%-device are 1.52 V and 0.31 V, respectively, corresponding to  $\sigma/\mu \sim 0.2$ , *i.e.*,  $\sim 20\%$  deviation from mean value, as shown in Fig. 5(a). Also,  $\mu$  and  $\sigma$  of  $V_{hold}$  are 1.23 V and 0.15 V, respectively, corresponding to  $\sim 12\%$  deviation. As shown in Fig. 5(b), the 7%-device has



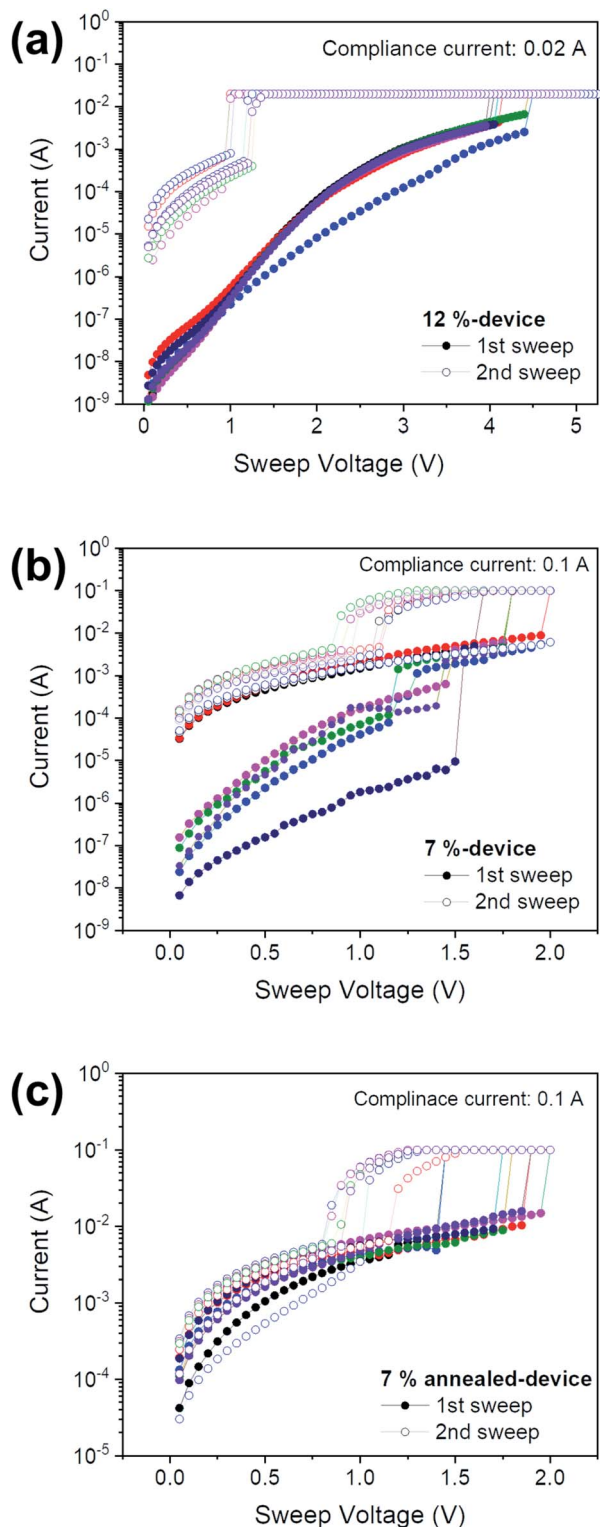


Fig. 3  $I$ - $V$  curves of randomly chosen seven different cells at the first two voltage sweeps of (a) 12%-, (b) 7%-, and (c) 7% annealed-devices.

similar distributions that  $\mu$  and  $\sigma$  of  $V_{th}$  are 1.24 V and 0.15 V, respectively, rendering  $\sim 12\%$  deviation, and  $\mu$  and  $\sigma$  of  $V_{hold}$  are 0.87 V and 0.06 V, respectively, with  $\sim 7\%$  deviation. In addition, the 7% annealed-device has similar values that  $\mu$  and  $\sigma$  of  $V_{th}$  are

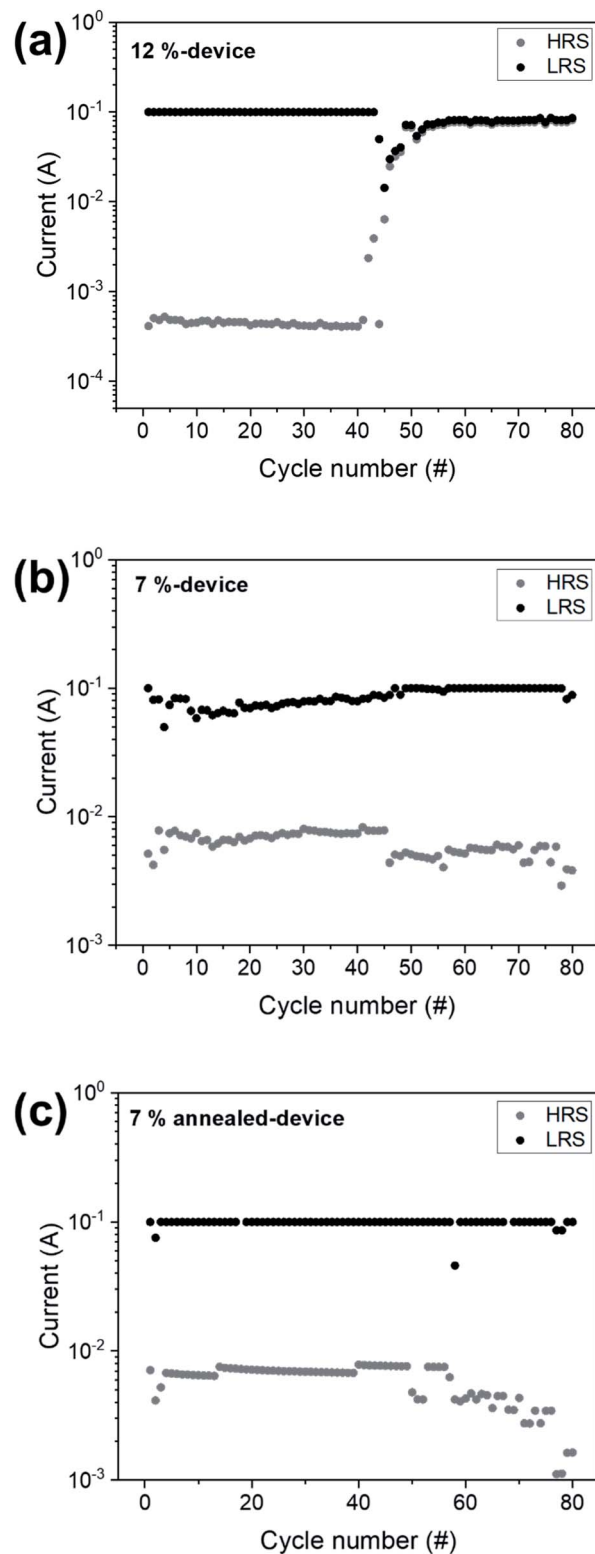


Fig. 4 Current levels at HRS and LRS of (a) 12%-, (b) 7%-, and (c) 7% annealed-devices as repeating threshold switching operations eighty times.

1.42 V and 0.11 V, respectively, rendering  $\sim 8\%$  deviation, and  $\mu$  and  $\sigma$  of  $V_{hold}$  are 0.81 V and 0.07 V, respectively, with  $\sim 9\%$  deviation (Fig. 5(c)).

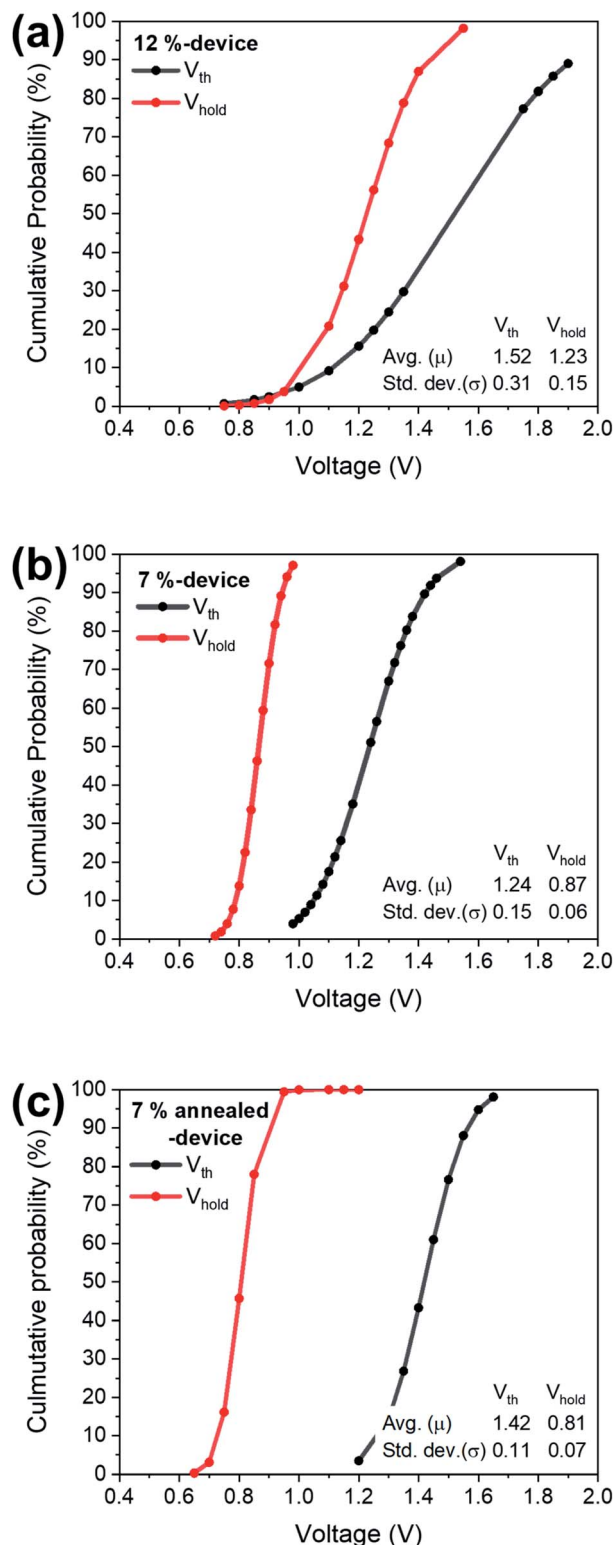


Fig. 5 Cycle-to-cycle distribution of  $V_{th}$  and  $V_{hold}$  values extracted from a hundred cycles by cumulative probability functions for (a) 12%-, (b) 7%-, and (c) 7% annealed-devices.

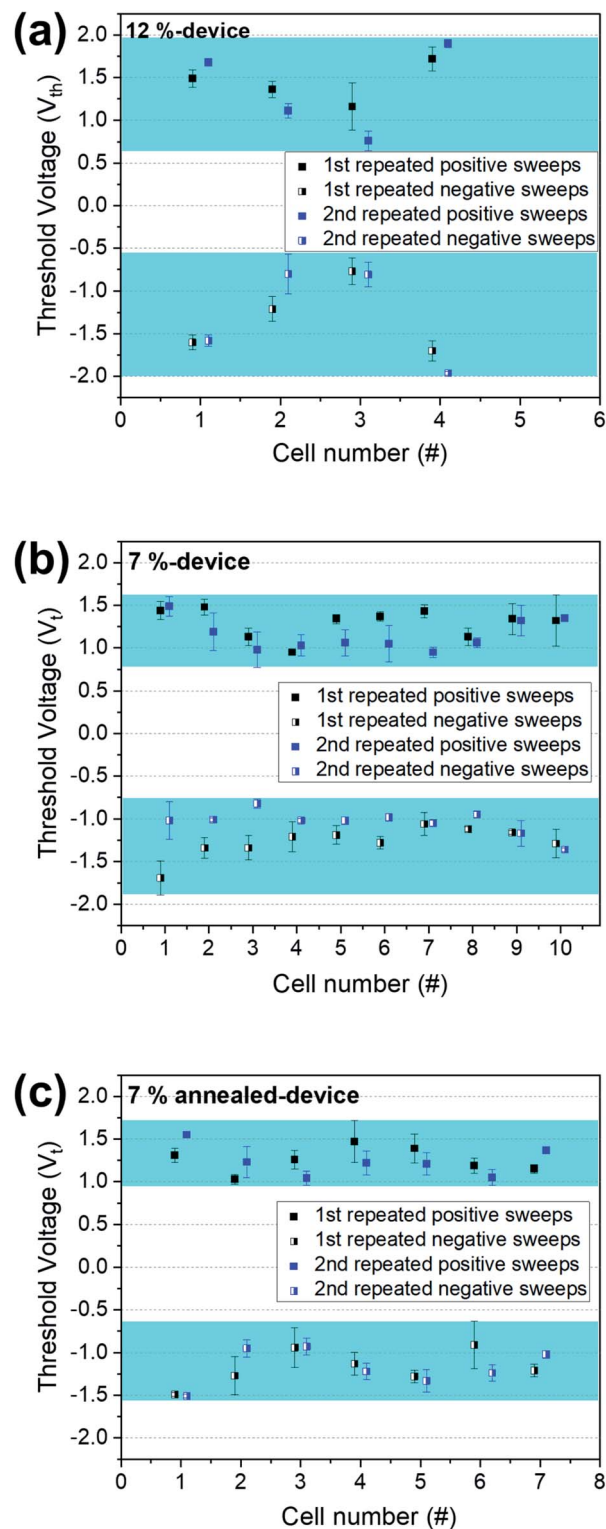


Fig. 6 Cell-to-cell variation of  $V_{th}$  values obtained at positive and negative voltage sweeps of randomly selected cells of (a) 12%-, (b) 7%-, and (c) 7% annealed-devices.

Fig. 6 shows the cell-to-cell variation of  $V_{th}$  values obtained at positive and negative voltage sweeps of randomly selected cells of the 12%-, 7%-, and 7% annealed-devices. The sets of positive

voltage sweeps repeated five time and following negative voltage sweeps five times were repeated consecutively. The  $V_{th}$  values at both the positive and negative sweeps are found to be within



narrow ranges marked as cyan color bands. These results indicate good threshold switching uniformity with a narrow distribution of switching parameters, irrespective of the  $\text{NbO}_x$  preparation conditions. Considering a half-bias operation scheme in crossbar array architectures, where the selected cell is full-biased while unselected cells are half-biased,<sup>27</sup> it is important to achieve the conditions for on-state (LRS) at “ $\pm V$ ” and off-state (HRS) at “ $\pm V/2$ ” simultaneously. Although the measured  $V_{\text{th}}$  values are little scattered from cell to cell, it is found that the values of  $\pm V_{\text{th}}/2$  for the off-state are mostly out of the  $\pm V_{\text{th}}$  ranges for the on-state thanks to the narrow  $V_{\text{th}}$  distribution, satisfying the requirement of half-bias scheme.

The comparison of threshold switching characteristics of three devices demonstrates that all devices have threshold switching behaviors for the application to selector devices while the switching parameters such as  $V_{\text{f}}$ ,  $V_{\text{th}}$ ,  $V_{\text{hold}}$ , HRS/LRS ratio, and endurance properties are determined by the preparation conditions of the  $\text{NbO}_x$  layers. It is also notable that the

threshold switching characteristics remained stable and even further improved by post-annealing. This thermal stability of threshold switching is beneficial for the integrated device fabrication requiring the processing steps at elevated temperatures.

Fig. 7 shows the threshold switching of the 7% annealed-device in response to voltage pulse application. During +1 V pulse application with a width of 100  $\mu\text{s}$ , the current remained to be low with the value of about  $8 \times 10^{-4}$  A which coincides with the off-state (HRS) current in  $I$ - $V$  sweep results in Fig. 1. Then, applying +2 V pulse increases the current to 0.02 A set to be the compliance current, which is the on-state (LRS) current in the

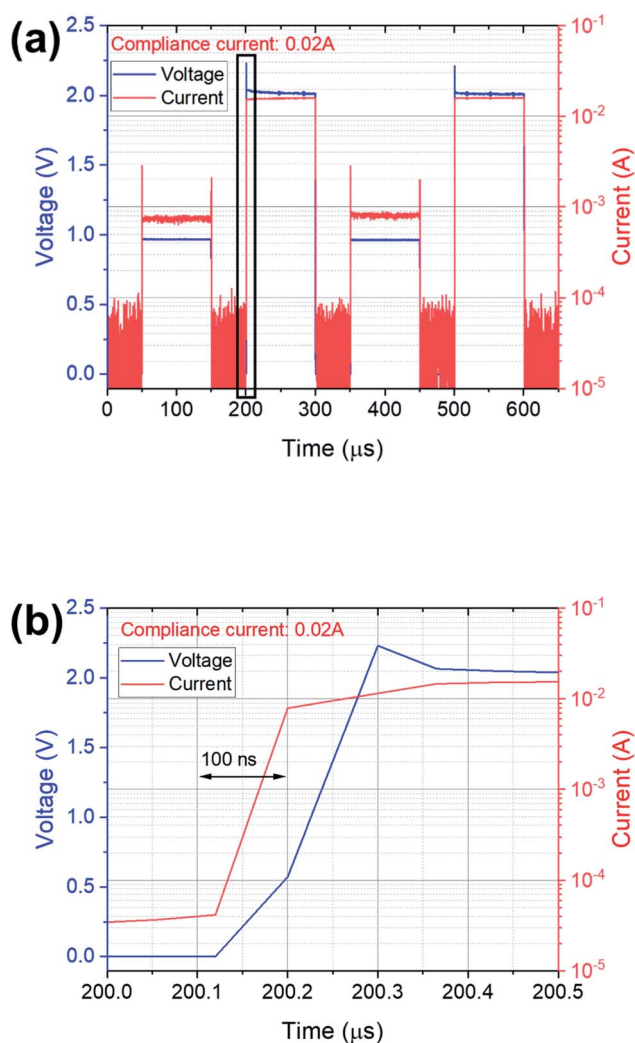


Fig. 7 (a) Pulse measurements of the 7% annealed-device by applying +1 V for HRS and +2 V for LRS with a pulse width of 100  $\mu\text{s}$  and (b) magnified graph at the onset of the increase of current upon applying +2 V pulse marked with black square in (a).

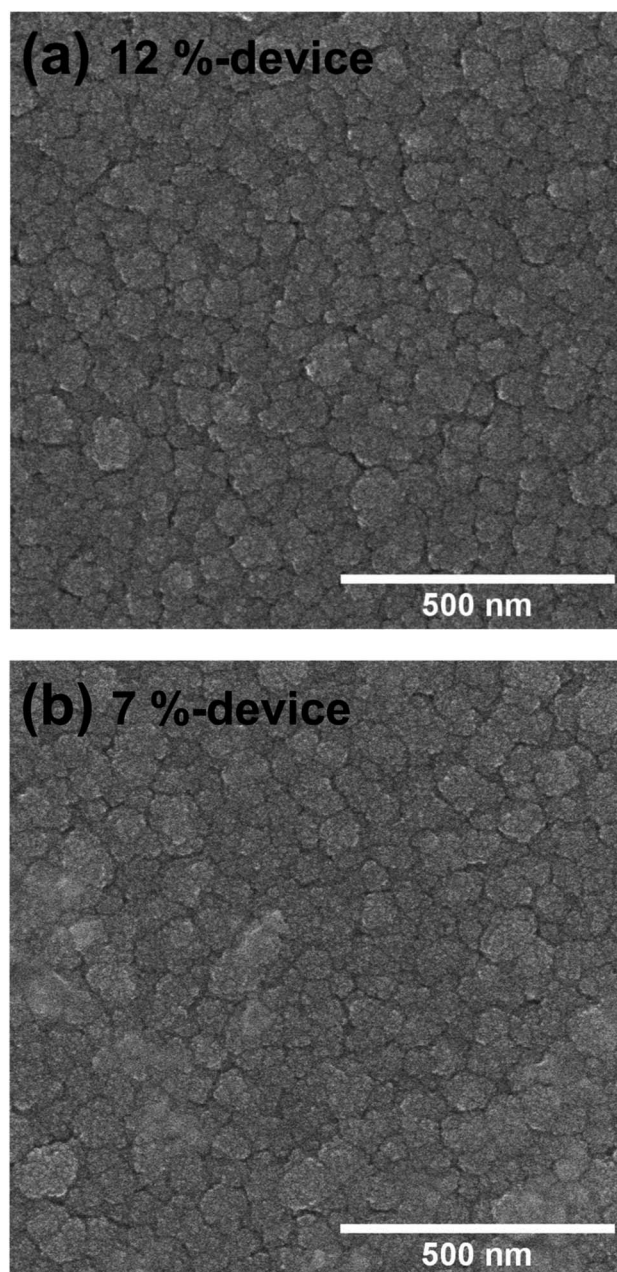


Fig. 8 Plan-view SEM micrographs of the surface of  $\text{NbO}_x$  layers deposited in (a) 12% and (b) 7%  $\text{O}_2$  gas.

threshold switching. Fig. 7(b) is the magnified graph at the onset of the increase of current upon applying +2 V pulse marked with black square in Fig. 7(a). The abrupt increase of current to LRS occurred clearly within the response time  $< 100$  ns, which demonstrates a fast switching behavior. From the hypothesis of threshold switching mechanism associated with alignment of oxygen vacancy as a VCM-type switching, it implies that the local alignment of oxygen vacancies to form temporary filament within  $\text{NbO}_x$  layer takes place very quickly within tens of ns.

Microstructures of the  $\text{Nb(O)}/\text{NbO}_x/\text{Nb(O)}$  devices were also investigated using SEM and TEM analyses. Fig. 8(a) and (b) are plan-view SEM micrographs of the surface of  $\text{NbO}_x$  layers deposited in 12% and 7%  $\text{O}_2$  gas, respectively. Even though the threshold switching behaviors and the current levels of the devices with these layers are distinguishable from each other, their microstructures are almost identical to have granular structures with the size in the range of  $50 \sim 100$  nm. However, it should be noted that the layers are not crystalline even with granular structures as confirmed from selected area diffraction patterns (SADP) analysis in Fig. 9.

Fig. 9(a)–(d) are bright-field (BF) and high-resolution (HR) cross-sectional TEM micrographs, and high angle annular dark field (HAADF) micrograph in STEM (scanning TEM) mode, and SADP of the 12%-device, respectively. Similar to SEM micrograph, the  $\text{NbO}_x$  layer is found to have columnar structure with undulating contrast within the layer, implying the presence of

porous regions inside the layer.<sup>28</sup> These porous regions would be facilitated path for the filament formation under the electric field. The HR-TEM micrograph in Fig. 9(b) and HAADF micrograph in Fig. 9(c) confirm that the  $\text{NbO}_x$  layer is amorphous with locally porous regions as shown to be with undulating contrast. The  $\text{Nb(O)}$  electrodes have also the columnar structure as generally observed from polycrystalline metal layers. Its crystalline feature is also found in HAADF micrograph with white spots representing diffracted grains in Fig. 9(c), implying that their size is tens of nanometers scattered inside granular structures. The SADP analysis in Fig. 9(d) discloses also that  $\text{Nb(O)}$  electrode is polycrystalline. However, it is not a pure Nb, but includes  $\text{NbO}$  crystallites,<sup>29</sup> due to the incorporation of oxygen atoms inside the layer as also confirmed from following AES analysis results in Fig. 10. The  $\text{NbO}_x$  layer and  $\text{Nb(O)}$  electrodes in the 7%-device shown in Fig. 9(e) and (f) have almost the same microstructures and phases as those of the 12%-device, and it indicates that the difference in  $\text{O}_2$  content from 7 to 12% in the sputtering conditions does not alter the microstructures and phases of the  $\text{NbO}_x$  layers significantly although the threshold switching characteristics are substantially distinguishable.

The compositional analyses were carried out in the 12%- and 7%-devices through AES depth profiling as shown in Fig. 10(a) and (b), respectively. In the  $\text{NbO}_x$  layers of both devices, the approximate ratios of  $\text{Nb} : \text{O} = 1 : 1.9$  were observed regardless

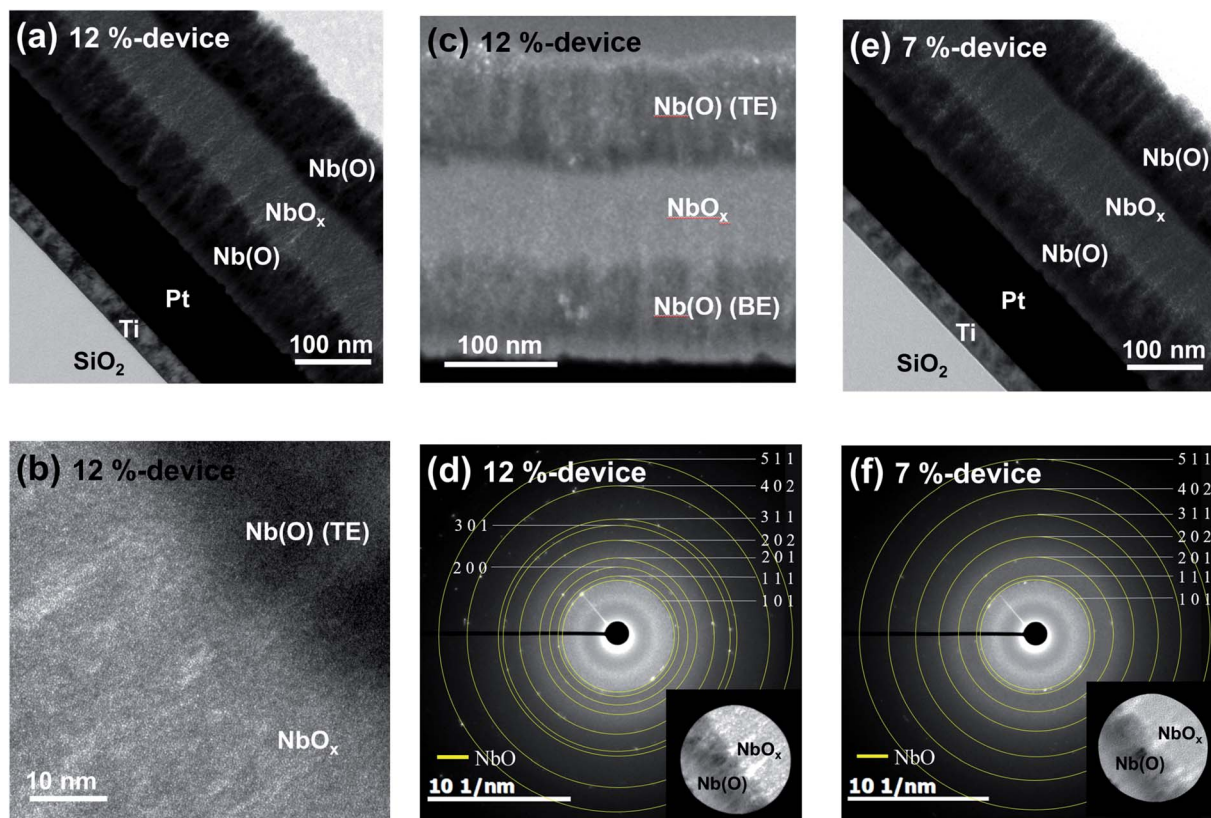


Fig. 9 (a) Bright-field and (b) high-resolution cross-sectional TEM micrographs, (c) high angle annular dark field micrograph in STEM mode, and (d) SADP of 12%-device, (e) bright-field cross-sectional TEM micrograph and (f) SADP of 7%-device.



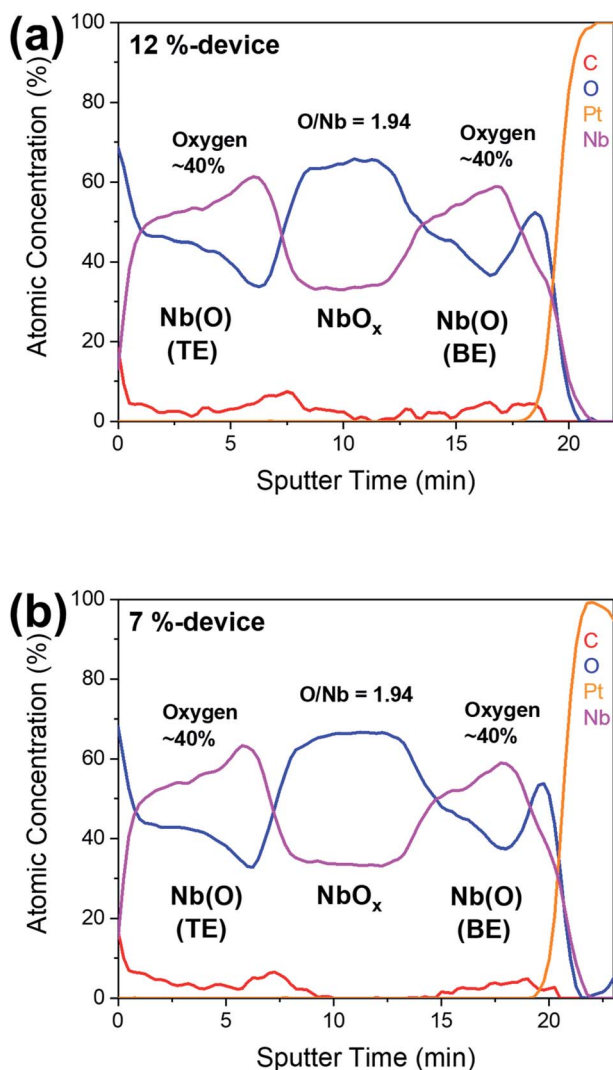


Fig. 10 AES depth profiles of (a) 12%- and (b) 7%-devices.

of the preparation conditions of  $\text{NbO}_x$  layers. It implies that  $\text{NbO}_x$  layers are the mixture of insulating  $\text{Nb}_2\text{O}_5$  and conducting  $\text{NbO}_2$  and  $\text{NbO}$  phases.<sup>30</sup> In addition,  $\text{Nb(O)}$  TE and BE are found to have substantial oxygen content up to almost 40 at%, resulting from the incorporation of oxygen atoms through their grain boundaries and also during their sputtering deposition.<sup>31</sup> It is also notable that oxygen content in  $\text{Nb(O)}$  TE near top  $\text{Nb(O)}/\text{NbO}_x$  interface is depleted while it is accumulated near bottom  $\text{Nb(O)}/\text{NbO}_x$  interface. The decent gradient of oxygen content in the  $\text{Nb(O)}$  TE with its depletion near top interface is explained with that oxygen atoms were incorporated from top surface as the device was exposed to air. The accumulation of oxygen in the  $\text{Nb(O)}$  BE near bottom interface could result from the penetration of oxygen atoms during  $\text{NbO}_x$  deposition in  $\text{O}_2$  mixed gas due to high reactivity of Nb with oxygen. Indeed, the resistivity of partially oxidized  $\text{Nb(O)}$  electrode was measured to be around  $9.4 \times 10^{-4} \Omega \text{ cm}$  using a 4-point probe (CMT 2000N, AIT), which is much higher than the reported value of  $2 \times 10^{-6} \Omega \text{ cm}$  of pure Nb layer.<sup>32</sup>

Fig. 11 shows Nb 3d spectra presenting chemical bonding states of Nb cations in  $\text{NbO}_x$  layers analyzed by using XPS.<sup>33,34</sup> For XPS analysis, the measurement spot size was  $400 \mu\text{m}$  in diameter. The measurement was performed at the middle positions in thickness of  $\text{Nb(O)}$  TE and  $\text{NbO}_x$  layers in the devices after etching with Ar gas. The peaks were calibrated with C 1s peak at 284.5 eV. The spectra were analyzed by fitting with Shirley background and Gaussian peak shape. Similar to previous studies reporting the coexistence of various Nb valence states such as  $\text{Nb}^{5+}$ ,  $\text{Nb}^{4+}$ , and  $\text{Nb}^{2+}$  in sputter-deposited  $\text{NbO}_x$  layers,<sup>11,13,26,30</sup> the Nb 3d spectra exhibit that the  $\text{NbO}_x$  layers are composed of mixed  $\text{Nb}_2\text{O}_5$ ,  $\text{NbO}_2$ , and  $\text{NbO}$  phases with varied ratio with respect to the  $\text{NbO}_x$  preparation conditions. The  $3d_{3/2}$  and  $3d_{5/2}$  peaks are observed near at 210.3 and 207.5 eV for  $\text{Nb}^{5+}$  cations in  $\text{Nb}_2\text{O}_5$ , and 208.4 and at 205.6 eV for  $\text{Nb}^{4+}$  in  $\text{NbO}_2$ , and 206.7 and 203.9 eV for  $\text{Nb}^{2+}$  in  $\text{NbO}$ .<sup>11,26,33–35</sup> Thus, the peaks with the energies for those phases were indexed accordingly with red, blue, and green lines, respectively. The  $\text{Nb(O)}$  TE deposited in pure Ar ambient consists of most of Nb and  $\text{NbO}$  with very little  $\text{NbO}_2$ , and  $\text{Nb}_2\text{O}_5$ , where the peaks at 205.1 and 202.3 eV for  $3d_{3/2}$  and  $3d_{5/2}$  peaks of Nb are shown with purple line.<sup>36,37</sup> The fraction of each phase in these layers was extracted from the relative area of each peak in the spectra. In the  $\text{NbO}_x$

layer of the 12%-device, the fraction of  $\text{Nb}_2\text{O}_5$ , i.e.,  $\frac{\text{Nb}^{5+}}{\text{Nb}^{4+} + \text{Nb}^{2+}}$ , is higher than those of other phases. Also, it is the highest in the 12%-device among three devices. The highest fraction of insulating  $\text{Nb}_2\text{O}_5$  phase led to the lowest HRS current in the pristine state. Comparing the spectra of  $\text{NbO}_x$  layers in the 7%- and 7%-annealed-devices, the fraction of  $\text{Nb}_2\text{O}_5$  phase was decreased while the more conducting  $\text{NbO}$  phase was enriched in the 7% annealed-device. It is explained by the interaction between  $\text{NbO}_x$  and  $\text{Nb(O)}$  electrodes that the migration of oxygen atoms from  $\text{NbO}_x$  layer to  $\text{Nb(O)}$  electrodes during post-annealing led to the increased fraction of  $\text{NbO}$  phase. The increased fraction

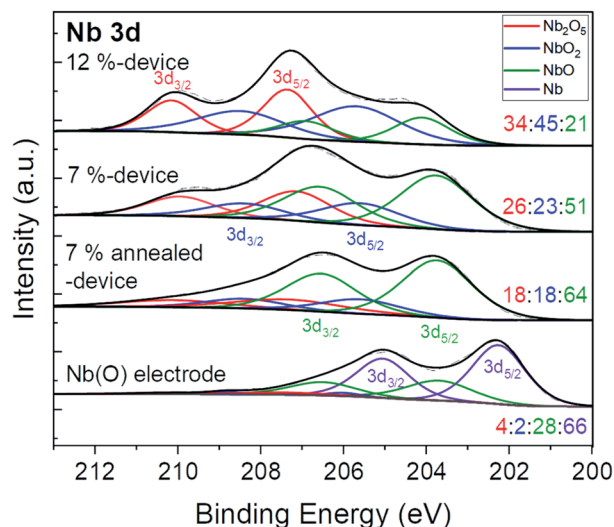


Fig. 11 Nb 3d spectra of the  $\text{Nb(O)}$  electrode and the  $\text{NbO}_x$  layers in the 12%-, 7%-, and 7%-annealed devices.



of NbO phase coincides well with the results of forming-free threshold switching in the 7% annealed-device.

As illustrated in Fig. 2, the threshold switching mechanism in the Nb(O)/NbO<sub>x</sub>/Nb(O) devices is associated with the percolation of conducting phases as conducting filament that depends on the oxygen vacancy concentration with respect to the preparation conditions of NbO<sub>x</sub> layer. From the XPS results of chemical bonding states analyses of the NbO<sub>x</sub> layers, the NbO<sub>x</sub> layers consist of the mixture of insulating Nb<sub>2</sub>O<sub>5</sub> and fairly conducting NbO<sub>2</sub> and NbO phases.<sup>30</sup> The resistivities of Nb<sub>2</sub>O<sub>5</sub>, NbO<sub>2</sub>, NbO, and Nb are known to be around 10<sup>11</sup>, 10<sup>1</sup> ~ 10<sup>4</sup>, 2 × 10<sup>-5</sup>, and 1.5 × 10<sup>-5</sup>, respectively.<sup>38</sup> The 12%- and 7%-devices require the forming operation for the threshold switching. At forming operation, the content of conducting phases such as NbO<sub>2</sub> and NbO is increased as Nb(O) electrodes absorb oxygen atoms and further deplete oxygen from NbO<sub>x</sub> layer by applied electric field.<sup>10,39</sup> The conducting phases are percolated to form conducting filament upon applying the voltage higher than  $V_{th}$ , which would be partially dissolved readily as reducing the voltage to be lower than  $V_{hold}$  and the devices return to be HRS. The HRS current after dissolution of filament is determined by the resistivity of NbO<sub>x</sub> layer. Thus, the 12%-device with higher content of insulating Nb<sub>2</sub>O<sub>5</sub> phase than other devices has the lowest HRS current. On the other hand, because the 7% annealed-device has the increased content of conducting phases settled in the NbO<sub>x</sub> layer by post-annealing, the forming-free threshold switching could be obtained readily by the alignment of conducting phases upon applying the voltage.

Since the threshold switching is strongly related with the compositions and phases of NbO<sub>x</sub> layers, the interaction with electrodes plays an important role in threshold switching behaviors.<sup>13,40</sup> Because the reaction of  $Nb + 2Nb_2O_5 \rightarrow 5NbO_2$  is thermodynamically favorable with Gibbs free energy of formation  $\Delta G_f^\circ = -164.3 \text{ kJ mol}^{-1}$ ,<sup>41</sup> the reduction of Nb<sub>2</sub>O<sub>5</sub> to enrich oxygen-depleted conducting phases would be facilitated in the device with Nb(O) electrode by the post-annealing or application of electrical bias. Even with the reduction of Nb<sub>2</sub>O<sub>5</sub>, because the Nb(O) electrodes have substantial oxygen content of almost 40 at%, it restricts excessive migration of oxygen atoms from NbO<sub>x</sub> layer. Thus, the overgrowth of conducting filament could be effectively prevented and reliable threshold switching could be obtained. These results indicate that the precise control of oxygen amount in the NbO<sub>x</sub> switching layer by deposition and annealing processes and the proper interaction with Nb(O) electrodes are crucial for reliable threshold switching characteristics of the Nb(O)/NbO<sub>x</sub>/Nb(O) devices.

## 4. Conclusion

The bipolar threshold switching characteristics of the Nb(O)/NbO<sub>x</sub>/Nb(O) devices with respect to the preparation conditions of the NbO<sub>x</sub> layers are demonstrated. The 12%-device with NbO<sub>x</sub> switching layer deposited at 12% of O<sub>2</sub> gas mixed in Ar shows threshold switching behavior at  $V_{th} < \pm 2 \text{ V}$  after forming operation at  $V_f$  of about +4 V. The 7%-device with NbO<sub>x</sub> layer deposited at 7% of O<sub>2</sub> gas is also found to require forming operation but with the lower  $V_f$  of around +2 V, and subsequent

threshold switching at  $V_{th} < \pm 2 \text{ V}$ . Notably, the 7% annealed-device with post-annealing process exhibits the forming-free threshold switching behaviors as a result of increased contents of conducting NbO and NbO<sub>2</sub> phases by post-annealing. The LRS/HRS ratios are in the range from 10 to 20 in the 7%-device and from 30 to 100 in the 12%-device, respectively. The cycle-to-cycle and cell-to-cell distributions of  $V_{th}$  and  $V_f$  values are found to be fairly uniform with maximum deviation < 20% from randomly chosen devices. The voltage pulse measurements disclose the switching time < 100 ns. From the composition and chemical bonding state analyses, the NbO<sub>x</sub> layers have the approximate ratio of Nb : O = 1 : 1.9 with the mixture of Nb<sub>2</sub>O<sub>5</sub>, NbO<sub>2</sub>, and NbO phases with different relative fractions. The fraction of insulating Nb<sub>2</sub>O<sub>5</sub> phase is about 34% in the 12%-device, which is the highest among three devices, so the forming operation with the highest  $V_f$  is required. On the other hand, the fraction of Nb<sub>2</sub>O<sub>5</sub> is about 18% in the 7% annealed-device, which is the lowest and then renders the forming-free threshold switching operation. The threshold switching mechanism is associated with the percolation of conducting phases, which is determined by the NbO<sub>x</sub> layer preparation conditions and the interaction with Nb(O) electrodes. The Nb(O) electrodes have substantial oxygen content of almost 40 at%, which prevents excessive redistribution of oxygen atoms, then enables the reliable threshold switching characteristics. The results of threshold switching in these devices demonstrate that the precise control of oxygen amount in the NbO<sub>x</sub> switching layer by appropriate interaction with the Nb(O) electrodes through post-deposition annealing is an effective route to realize the forming-free threshold switching characteristics for the application to selector devices in high-density crossbar array architectures.

## Author contributions

Kitae Park: conceptualization, data curation, formal analysis, investigation, writing – original draft. Jiyeon Ryu: data curation, formal analysis. Dwipak Prasad Sahu: writing – review & editing. Hyun-Mi Kim: data curation, formal analysis. Tae-Sik Yoon: funding acquisition, project administration, supervision, visualization, writing – review & editing.

## Conflicts of interest

There are no conflicts to declare.

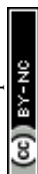
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