Memristors Using Solution-Based IGZO Nanoparticles

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ABSTRACT: Solution-based indium–gallium–zinc oxide (IGZO) nanoparticles deposited by spin coating have been investigated as a resistive switching layer in metal–insulator–metal structures for nonvolatile memory applications. Optimized devices show a bipolar resistive switching behavior, low programing voltages of ±1 V, on/off ratios higher than 10, high endurance, and a retention time of up to 10⁴ s. The better performing devices were achieved with annealing temperatures of 200 °C and using asymmetric electrode materials of titanium and silver. The physics behind the improved switching properties of the devices is discussed in terms of the oxygen deficiency of IGZO. Temperature analysis of the conductance states revealed a nonmetallic filamentary conduction. The presented devices are potential candidates for the integration of memory functionality into low-cost System-on-Panel technology.

1. INTRODUCTION

There are fundamental limits to the downscaling of materials involved in the current complementary metal-oxide-semiconductor technologies. Resistive random-access memories (RRAMs) can be considered as a highlight as they consist of two terminal devices with metal–insulator–metal (MIM) structure, allowing large-scale-integration circuits.¹

They can be described, in simplified form, as a nonlinear passive resistor (memristor). RRAMs combine the best features of current memories such as the fast speed of SRAM, high density of DRAM, and nonvolatile properties of flash memory. These advantages support the idea that the future of memory devices will be based on RRAM devices.²–⁵ This memory technology will revolutionize all consumer electronic products from computing technology to healthcare and the Internet of Things.

An ideal System-on-Panel (SoP) technology utilizes electronic circuits, which are deposited with the same tools and materials as those used for the display components. This explains the choice to investigate amorphous indium–gallium–zinc oxide (a-IGZO) for RRAMs, because it can be commercially applied in the thin-film transistors (TFTs) of driver circuits for active-matrix displays.⁶,⁷ The highly competitive markets for such displays, as mentioned above, demand the lowest production cost possible, which motivates a solution-based deposition route. Here, the main bottleneck is to find the optimal compromise between electrical performance/reliability and production cost effectiveness.

Developments in IGZO-based RRAM have been reported elsewhere, focusing on studies of bipolar behavior in fully transparent devices,⁷,⁸ unipolar switching behavior,⁹ electrode influence,¹⁰ filament nature,¹¹ and flexible devices.¹² A single synaptic device based on an a-IGZO memristor has been reported,¹³ demonstrating the versatility of these devices. The most common industrially relevant deposition technique for IGZO films is radio frequency magnetron sputtering, and several ways to improve the performance of the corresponding memory devices have been reported. An interlayer of Ti at the interface improves the device yield and resistive switching properties due to the formation of TiOₓ.¹⁵ Microwave irradiation can be used to obtain a smoother surface, which results in more stable resistive switching properties.¹⁶ By engineering the concentration of oxygen vacancy defects, stable forming-free resistive switching characteristics are obtained.¹⁷ SiO₂ has been used as a series resistance material to produce a thinner conductive filament.¹⁸ These reports show the excellent resistive memory properties that can be obtained with sputter-deposited IGZO. This justifies the investigation of low-cost fabrication routes such as deposition from solution, because such methods typically lead to more defective materials with poorer electrical properties. However, to the best of our knowledge, only two reports on solution-based IGZO non-volatile memories exist so far: Inspired by low-cost IGZO...
solution-based TFTs, Kim et al. developed a-IGZO solution-based bipolar RRAM. However, the devices operate at high current level and the reported on/off ratio did not exceed 1 order of magnitude. Later, Hu et al. reported an a-IGZO-based RRAM device fabricated by a photochemical solution deposition with improved unipolar memory performance. This work intends to further explore the potential of solution-based synthesis of oxide-based RRAM. The here presented method comprises the synthesis of nanoparticles followed by thin-film deposition from a dispersion. An important reason to choose such a dual-stage process is the greater production flexibility compared to that of direct deposition from solution. To the best of our knowledge this is the first report on a RRAM device based on IGZO nanoparticles. Furthermore, a maximum process temperature as low as 200 °C is introduced because this enables the integration on flexible substrates.

2. RESULTS AND DISCUSSION

IGZO nanoparticles (IGZOnp) used as the memristive switching layer were produced by hydrothermal synthesis. The film was deposited on substrates using a common Ti bottom contact. The annealing temperature of 350 °C was optimized by Santos et al. and applied to two substrates, the remaining two substrates were annealed at 200 °C for comparison.

For each annealing temperature, two types of MIM structures were fabricated. One structure is symmetric using Ti top electrodes (Ti/IGZOnp/Ti). The other structure is asymmetric by applying Ag as top contact (Ag/IGZOnp/Ti). Figure 1 depicts a typical substrate with the 6 by 6 matrix of MIM structures used in this study. A cross-section view of the device by scanning electron microscopy (SEM) in Figure 1b shows the 20 nm thick IGZO nanoparticle layer. Furthermore, Figure 1c,d shows the surface morphology characterized by atomic force microscopy (AFM) of the IGZOnp thin-film deposited on Ti on Corning glass. A smaller nanoparticle size is observed for the lower temperature. For 200 °C, the mean particle radius was approximately 3 nm whereas for 350 °C, it was ~4 nm. It was also observed that for lower temperatures, there is a decrease of surface roughness.

As is often reported for resistive switching systems, the MIM structures undergo an irreversible electroforming process before they can act as a memory device. The original conduction state is referred to as the pristine state. Figure 2 compares the current–voltage characteristic (I–V) of a typical pristine state as well as off-state and on-state I–V curves of all four devices. In the pristine state, samples annealed at 350 °C show higher conductance than that of the ones annealed at 200 °C. When Ti is used for both electrodes, the I–V curve is symmetric for both polarities. In contrast, MIM structures using Ag as top contact electrode and Ti as bottom contact show a rectifying behavior in the pristine state. To electroform the memory cell, particular care has to be taken to prevent the immediate breakdown, thus current compliance (CC) is required. For the Ti/IGZOnp/Ti devices, the forming step occurs by a positive voltage applied to the top
electrode. However, it should be noted that structures with the same type of metal electrode (Ti) at both contacts do not have a preferential voltage polarity to electroform, thus applying positive or negative voltages led to identical resistive switching behavior.

For the MIM structure using the silver top electrode, electroforming occurs only when a negative bias is applied to the silver contact. Forming under positive bias leads to unstable switching behavior after a few programming cycles. Furthermore, electroforming is established at lower voltages compared to that for the symmetric MIM structures using Ti on both top and bottom electrodes. Finally, the low conductance state (off-state) has a higher conductance than the pristine state.

Cycles of endurance and retention tests were carried out for all four devices to evaluate the impact of different electrodes and annealing temperature on memory performance. Reading voltages were ±0.1 V, corresponding to set polarization. Figure 3 shows endurance tests over 100 cycles in sweep voltage mode carried out for all four test devices. The Ti/IGZO/Ti device annealed at 350 °C shows a set voltage at 1.3 V in the first cycle, with a tendency to shift to higher voltages as the number of cycles increases. Noncontrollable intermediate switching events to off-state appear during the endurance cycles under the reset process between −0.8 and −1.5 V (see Figure 3a). For the Ti/IGZO/Ti device annealed at 200 °C, set voltages between 0.8 and 1.3 V and reset voltages of −1 to −1.5 V are observed (see Figure 3b). Here, the on/off current ratio and the memory performance gradually deteriorate after the 40th cycle.

Figure 3c,d shows endurance cycle tests carried out on Ag/IGZO/Ti devices annealed at 350 and 200 °C, respectively. The hysteresis shows a clockwise direction that is consistent with the negative bias forming loop. For the first device annealed at 350 °C, the set process is performed at −1.4 V and reset occurs at 2 V. For the device annealed at 200 °C, the set voltage is even lower, between −1 and −0.8 V, and the reset is obtained by applying a positive voltage of 1 V. The operating on/off ratio window is more than 1 order of magnitude.

Concerning the data retention presented in Figure 3, the comparison of all four devices shows that all devices except Ti/IGZO/Ti fabricated at 200 °C annealing temperature have a retention time of at least 10⁴ s when keeping the switching margin equal to or even higher than 10.

Table 1 compares the main electrical characteristics for both Ti/IGZO/Ti and Ag/IGZO/Ti memory devices annealed at different temperatures. The yield was defined here as the number of devices out of 36 that show at least 10 stable and consecutive endurance cycles.

Figure 2. Typical I–V characteristics showing the forming process and memory states. (a, b) Ti/IGZO/Ti annealed at 350 and 200 °C, respectively. (c, d) Ag/IGZO/Ti annealed at 350 and 200 °C, respectively.

Figure 3. I–V characteristic over 100 cycles and retention time characteristics at ±0.1 V at room temperature for devices: (a) Ti/IGZO/Ti annealed at 350 °C, (b) Ti/IGZO/Ti annealed at 200 °C, (c) Ag/IGZO/Ti annealed at 350 °C, and (d) Ag/IGZO/Ti annealed at 200 °C.
This temperature dependence rules out metallic-like conductance, have shown a metallic-like temperature dependence of contact. If some authors tentatively explain it as caused by the formation of a thin oxide layer between the switching layer and the bottom electrode when the samples are annealed at the higher temperatures, the IGZOnp films annealed at 350 °C hinder the Fermi energy to enter the conduction band and cause the observed nonmetallic temperature dependence of conductance.

The temperature dependence for the off-state shows an anomalous behavior. For the off-state below 270 K, a very large positive temperature coefficient (PTC) is observed. The increase of the current upon lowering the temperature of the device is more than 2 orders of magnitude in a temperature range of 60 K ($\alpha = 0.02$ K$^{-1}$). This PTC is anomalously large when compared with typical values for metals ($\alpha = 4 \times 10^{-3}$ K$^{-1}$ for Ag). This behavior has been observed on other RRAM devices and explained as due to filamentary conduction. For temperatures above 270 K, the temperature dependence shows a semiconductor-like behavior. The transition point between the two behaviors occurs close to the temperature of the phase transition of water (273 K), which is unavoidably present due to the fabrication process.

A stochastic electrical behavior is observed for the symmetric Ti-based device structure. By replacing the Ti top electrode with Ag, the memristor shows stable switching properties with higher switching yield during endurance tests. In the following, the role of the titanium at the interface to the IGZO shall be discussed. Figure 5 serves as an accompanying graphical illustration. Titanium is referred to as an oxygen-getter material due to its high affinity for forming titanium oxide. In the devices presented here, this leads to the removal of oxygen from the IGZO layer. Because this process is accelerated at higher temperatures, the IGZOnp films annealed at 350 °C are more conductive than the ones annealed at 200 °C. In Figure 5, this is represented by the more intense red color toward the titanium bottom electrode. Although the annealing is done in contact with ambient air, the electrical conducance of the devices is dominated by the oxygen removal at the titanium bottom electrode and not by an eventual oxidation at the IGZOnp film surface. The presence of an anatase phase could be confirmed by grazing incidence X-ray diffraction (XRD) in the samples annealed at 350 °C, whereas the same could not be detected in the samples annealed at 200 °C (see Figure S3). This confirms the increased oxidation of the titanium bottom electrode when the samples are annealed at the higher temperature.

### Table 1. Performance Comparison between Ti/IGZOnp/Ti and Ag/IGZOnp/Ti for 350 and 200 °C Annealing Temperatures

| Structure     | $T$ [°C] | Yield [%] | CC$^a$ [mA] | Forming [V] | Set$^b$ [V] | Reset$^b$ [V] | Read [V] | On/off ratio | Retention [s] | Endurance |
|---------------|---------|----------|-------------|-------------|------------|-------------|----------|-------------|--------------|-----------|
| Ti/IGZOnp/Ti  | 350     | 20       | 1           | +3.5        | +1.3       | −0.8        | +0.1     | >10         | 10$^6$       | 80/100    |
|               | 200     | 14       | 1           | +4.7        | +2.3       | −1.8        | +0.1     | <10         | 10$^4$       | 40/100    |
| Ag/IGZOnp/Ti  | 350     | 64       | 0.5         | −1.9        | −1.4       | +2          | −0.1     | >10         | 10$^6$       | 100       |
|               | 200     | 68       | 0.01        | −1.5        | −1         | +1          | −0.1     | >10         | 10$^4$       | 100       |

$^a$CC during forming. $^b$Set and reset values are corresponding to the first cycle.
The electron beam deposition of the titanium top electrode leads to the formation of a thin layer of oxidized titanium at the interface. Hence, a similar—although probably less severe—oxygen-getter effect occurs at the top contact of the Ti/IGZO/Ag devices, which also leads to an increase of conductivity close to the top surface of the IGZO layer. In Figure 5, this is shown by the red color below the titanium top electrodes. When silver is used instead, the conductivity at the top interface is lower, because silver typically does not oxidize in contact with IGZO (based on the Gibbs free energies of formation). Moreover, silver is reported to form a Schottky barrier with IGZO, which implies a high resistivity in the IGZO/titanium interface. Hence, a similar leads to the formation of a thin layer of oxidized titanium at the interface. Therefore, under negative forming bias, a conductive filament based on oxygen vacancies growing from the bottom contact toward the top contact results in more stable switching events, as compared to the titanium top contact. Furthermore, the conductance in the pristine state under negative bias allows for the use of a lower CC during electroforming (see Table 1). In this way, the risk of breakdown due to Joule heating effects is minimized.

3. CONCLUSIONS

The solution-based Ti/IGZO/Ag/Ti and Ag/IGZO/Ag/Ti memory devices were fabricated at different annealing temperatures of 200 and 350 °C. The lowest switching voltages were observed for Ag/IGZO/Ag/Ti devices annealed at 200 °C with set and reset voltages of +1 and −1 V, respectively, and retention times of up to 10^4 s. The Ag/IGZO/Ag/Ti devices showed higher yields compared to those of the symmetric Ti-based devices. Temperature-dependent measurements reveal a nonmetallic conductive filament at the on-state.

It could be shown that the oxygen-getter effect due to the oxidation of titanium increases the conductivity of IGZO. If both IGZO interfaces are oxygen deficient, stochastic switching properties are observed. The solution is a nonsymmetric device architecture using a nonoxidizing top electrode such as silver. The resulting lower conductivity is favorable to establish a well-defined filamentary path by applying lower current compliance during electroforming.

In summary, we demonstrated an RRAM device processed from a dispersion of solution-based IGZO in comparison to the previously reported works on solution-based IGZO, we have achieved lower operation power. Also, self-compliant operation is a major feature that simplifies the circuit design. This RRAM device has the potential to be integrated into SoP technology.

4. EXPERIMENTAL SECTION

4.1. Device Fabrication. IGZO nanoparticles were produced following a recipe previously developed by Santos et al. with gallium nitrate hydrate (Ga(NO3)3·H2O, Sigma-Aldrich, 99.9%), indium acetate (In(CH3COO)3, Sigma-Aldrich, 99.99%), and zinc acetate (Zn(CH3COO)2·2H2O, Sigma-Aldrich, 99%) precursors with a molar ratio of 3:6:2, respectively. The mixture was dissolved in 2-methoxyethanol (6 mL, Sigma-Aldrich, 99.5%) with ethanolamine (0.2 mL, Fluka, 98%) as a cation reducer, and the solution was left stirring at 50 °C for 1 h. The final solution was transferred to the polytetrafluoroethylene cup of a 23 mL stainless-steel autoclave (4745 general purpose vessel, Parr), and placed in an oven (L3/11/B170, Nabetherm) at 180 °C for 24 h. The resulting product was collected by centrifugation at 4000 rpm for 5 min (F140, Focus instruments) and dispersed in ethanol (20 mL).

IGZO ink was ensured by mixing ethylene glycol (Carlo Erba, 95.5%) with the previous dispersion in 0.4:1.6 proportion, respectively, and consequently, the mixture was left stirring for 24 h. Then, it was sonicated for 2 min and filtered with a 0.45 μm porous diameter filter (Sartorius CA). The deposition process was performed by spin coating 4 layers at 2000 rpm for 35 s and drying at 100 °C for 1 min between each layer. The annealing process was done in air after the fourth deposition at 350 and 200 °C for 1 h.

MIM structures were fabricated on glass substrates (1737, Corning). First, a 60 nm Ti thin-film was deposited by e-beam evaporation (homemade apparatus) followed by spin coating deposition of the IGZO ink, creating an IGZO thin-film layer of approximately 20 nm, see Figure 1b. The top electrode was deposited with the same procedure as the bottom electrode but using a shadow mask, where titanium (60 nm) or silver (60 nm) were deposited on different substrates. A device area of 19.6 μm² was defined by top electrode deposition through a shadow mask.

4.2. Materials and Device Characterization. Structural and morphological characterization of the IGZO nanoparticles was performed by scanning electron microscopy (Carl Zeiss AURIGA). The nanoparticles were deposited on a silicon substrate for the X-ray diffraction (XRD) measurements and scanned in the 20–70° 2θ range with a step size of 0.033° in a PANalytical’s X’Pert PRO MPD diffractometer with Cu Kα radiation. In addition, the devices were measured by grazing incidence X-ray diffraction with the same diffractometer at an incident angle of θ = 1.1° and a step size of 0.1°. The IGZO thin-film surface roughness was analyzed by atomic force microscopy in a commercial microscope, Asylum Research MFP-3D. Attenuated total reflectance Fourier-transformed infrared spectroscopy was measured with a Thermo Nicolet 6700 Spectrometer, using a single bunched diamond crystal (4500–525 cm⁻¹).

IGZO/Pd memristor electrical characterization was performed using a semiconductor characterization system (Keithley...
4200SCS). Positive bias refers to the top electrode in the sandwich structure.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.7b01167.

Confirmation of the presence of organic residues due to the solution process by attenuated total reflection Fourier-transformed infrared spectroscopy; Space-charge-limited conduction mechanism of the Ti/IGZOOnp/Ti device; X-ray diffractograms of the synthesized IGZO nanoparticles and the Ag/IGZOOnp/Ti devices after different annealing temperatures (PDF).

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#### Author Contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

#### Notes

The authors declare no competing financial interest.

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