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2018-09-06


http://hdl.handle.net/10138/246369
https://doi.org/10.1149/2.0261809jss

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Atomic Layer Deposition and Properties of HfO2-Al2O3 Nanolaminates

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Nanocrystalline HfO2-Al2O3 mixture films and nanolaminates were grown by atomic layer deposition at 350 °C from metal chloride precursors and water. Formation of metastable HfO2 polymorphs versus monoclinic phase was affected by the relative amount and thickness of constituent oxide layers. The films exhibited saturative magnetization and charge polarization in externally applied fields at room temperature. The films also demonstrated resistive switching behavior with considerable window between low and high resistance states.

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Manuscript submitted June 29, 2018; revised manuscript received August 21, 2018. Published September 6, 2018.

Composites and multilayers of chemically distinct metal oxides are of continuous interest as materials tailoring useful electronic, mechanical and structural properties of their constituents which increase their suitability to many applications. Hereewith hafnium-aluminum-oxide thin films have been studied as insulating dielectrics potentially suited to application as gate dielectric materials in MOSFET structures with Si,1–3 Si0.7Ge0.3,4 In0.53Ga0.47As,5,6 or InP7,8 as well as memory capacitor dielectrics in three-dimensional DRAM structures.9 For flash memory cell structures, different charge trapping materials HfO2, Hf-Al-O and Al2O3 have been compared in metal-oxide-semiconductor capacitors with fixed Al2O3 tunneling and blocking layers and Pd-electrode.12 Al2O3 nanolaminates, as well as mixtures, have also been investigated for ferroelectric-like behavior26 have already been described in some papers to date. At the same time, magnetization behavior in the same hystereses as characteristic of ferroelectric materials. Corresponding polarization, since, due to its noncentrosymmetric lattice, it could be responsible for the ability of the material to nonlinearly and saturatively polarize in external electric field. Such films can demonstrate remnant polarization and coercivity, forming polarization loops with marked hystereses as characteristic of ferroelectric materials. Corresponding behavior in HfO2 films doped with Al has been reported earlier.7,24 Stabilization of metastable (incl. orthomorphic) HfO2 films grown by atomic layer deposition using various dopants including Al2O3,25 and their ferroelectric-like behavior26 have already been described in some papers to date. At the same time, magnetization behavior in the same films has not yet been studied.

In general, layered transition metal oxides have gained attention as materials able to exhibit both magnetoelectric and resistive switching behavior.27 Resistive switching, ferromagnetic and ferroelectric polarization have simultaneously been registered and investigated in rather complex stacked compounds, such as Bi4Ti3O12, Pr2Fe2O7, MnO0.05O2/Bi2O3, Gd0.04O12/Ti2O3, W0.05O12/Pt(111)/Ti/SiO2/Si(100).28 Defects, in particular oxygen vacancies, have presumably been responsible for the switching behavior, concurrent electrical modulation and the following electrical field coupling with magnetization of the same material. Resistive switching possibly affecting electrical control of ferromagnetism in Ag/HfO2/Nb/SrTiO3/Ag stacks has been reported.29 Coexistence of electric field controlled ferromagnetism and resistive switching in transition metal oxide films has been recorded at room temperature earlier in TiO2 based stacks.30 Switching currents were also measured in atomic layer deposited, presumably ferroelectric, Hf3(Sr0.75Tl0.25)O3 thin films.31 Therefore, simultaneous evaluation of magnetization behavior, electrical polarization and switching resistivity effects in the same solid films appear justified. Recently, a study on ZrO2-Al2O3 doped films and nanolaminates was published, describing the electric and magnetic behavior of these materials.32 Nonlinear saturative magnetization was achieved in the laminated structure, referring to ferromagnetic-like behavior. At the same time, electric charge polarization in ZrO2-Al2O3 stacked layers was markedly affected by the interfacial polarization, possibly with weak ferroelectric component contributing to the polarization-field curves measured. The electronic leakage currents in these laminates and doped films were considerable, although allowing definitive switching between high and low resistance states. The switching was positively influenced by the inclusion of Al2O3 dopant layers.32

Apart from scarce studies on HfO2-Al2O3 multilayers produced by physical methods such as sputtering, HfO2-Al2O3 films have mainly and most feasibly been processed by atomic layer deposition (ALD). HfO2-Al2O3 films and multilayer films have been synthesized by ALD from Al(CH3)3, Hf[N(CH3)(C2H5)]4, and H2O;7,12,24,33 Al(CH3)3, Hf[N(CH3)2]2 and H2O;3,4 Al(CH3)3, Hf[N(CH3)2]2 and O2 plasma;34 Al(CH3)3, Hf[N(CH3)2]2, and O2 plasma;35 Al(CH3)3, Hf[N(CH3)2]2, and O2 plasma;36 Al(CH3)3, HfCl4, and H2O;6,15 At least one of the metal precursors in those experiments were aluminum alkyl or hafnium alkylamid compounds, i.e. precursors which knowingly provide growth of thin films uniformly over arbitrarily shaped substrates. On the other hand, the growth experiments are then limited to temperatures 200–250 °C or below, because these precursors are prone to thermal decomposition. For instance, Al2O3-HfO2 nanolaminates were grown by ALD from Al(CH3)3, Hf[N(CH3)2]2 and H2O at 225 °C,3,4 and from Al(CH3)3, Hf[N(CH3)2]2, and O2 plasma at 150 °C with post-deposition annealing at 350 and 850 °C, respectively, in order to improve the film structure and reduce the defect densities.

In the present study, HfO2-Al2O3 nanolaminates were grown by ALD in order to investigate the phase composition of HfO2 layers grown to variable thicknesses between the intermediate Al2O3 constraint layers. The experiments were carried out using thermally stable
metal chloride precursors, i.e. AlCl3 and HfCl4. The deposition temperature, 350 °C, was chosen higher than that most commonly used, 300 °C, in chloride-based ALD process, to enhance ordering and provide well-defined crystal growth in the as-deposited HfO2 films.35 At the same time, the temperature chosen was sufficiently low to enable reasonably high nucleation density followed by development of continuous solid material layers beginning from the earliest stages of the growth process, since even higher deposition temperatures might reduce the nucleation density, increase the grain size and cause significant structural defects and voids.36 After the growth experiments, the films were electrically evaluated in their as-deposited states without further heat-treatments. The films were also characterized in terms of electrical charge and magnetic polarization in external fields.

**Experimental**

The films were grown in a commercial flow-type hot-wall reactor F120 (ASM Microchemistry, Ltd.)37 from HfCl4 (99.9%, Strem), AlCl3 (99%, Acros Organics) and H2O. The growth temperature was held at 350 °C. HfCl4 and AlCl3 were evaporated at 170 and 95 °C, respectively from open boats inside the reactor, and transported to the substrates by the N2 flow. The cycle times used were 0.5–0.5–0.5–0.5 s, denoting the sequence metal precursor pulse–purge–water pulse–purge. The substrates were rectangular pieces cut out of undoped Si(100) with maximum edge length of 5 cm, covered with a 1.5–2.0 nm thick wet-chemically-grown SiO2. In addition, also conducting electrode substrates were used for the deposition of HfO2, based on (100) silicon with a resistivity of 0.014–0.020 Ω cm, i.e., boron-doped to concentrations up to 5 × 1018–1 × 1019/cm3, and coated with 10 nm thick chemical vapor deposited titanium nitride layer. The films were grown to thicknesses ranging from ca. 5 to 100 nm, i.e. also relatively thinner than those subjected to electrical measurements, in order to make the structural and compositional measurements more convenient. The Al to Hf content was varied by changing the ratio of the subsequent Al2O3 and HfO2 deposition cycles. The number of subsequent Al2O3 and HfO2 deposition cycles was varied separately in order to change the thicknesses of constituent metal oxide layers. In this way, films of different structures were deposited, ranging from HfO2 mixed with low amounts of Al2O3 to Al2O3–HfO2 nanolaminates. The growth cycle sequences applied for the different samples are given in Table I.

<table>
<thead>
<tr>
<th>HfO2:Al2O3 cycle ratio</th>
<th>growth cycle sequences</th>
<th>d, nm</th>
<th>Hf:Al</th>
</tr>
</thead>
<tbody>
<tr>
<td>(15:4)</td>
<td>31 × [15 × HfO2 + 4 × Al2O3] + 15 × HfO2</td>
<td>41</td>
<td>1.8</td>
</tr>
<tr>
<td>(15:3)</td>
<td>33 × [15 × HfO2 + 3 × Al2O3] + 15 × HfO2</td>
<td>34</td>
<td>2.7</td>
</tr>
<tr>
<td>(15:2)</td>
<td>34 × [15 × HfO2 + 2 × Al2O3] + 15 × HfO2</td>
<td>32</td>
<td>2.8</td>
</tr>
<tr>
<td>(50:5)</td>
<td>11 × [50 × HfO2 + 5 × Al2O3] + 50 × HfO2</td>
<td>37</td>
<td>4.4</td>
</tr>
<tr>
<td>(15:1)</td>
<td>35 × [15 × HfO2 + 1 × Al2O3] + 15 × HfO2</td>
<td>33</td>
<td>3.3</td>
</tr>
<tr>
<td>(150:10)</td>
<td>4 × [150 × HfO2 + 10 × Al2O3] + 150 × HfO2</td>
<td>44</td>
<td>3.5</td>
</tr>
<tr>
<td>(170:10)</td>
<td>4 × [170 × HfO2 + 10 × Al2O3] + 170 × HfO2</td>
<td>48</td>
<td>6.4</td>
</tr>
<tr>
<td>(100:5)</td>
<td>5 × [100 × HfO2 + 5 × Al2O3] + 100 × HfO2</td>
<td>41</td>
<td>4.2</td>
</tr>
<tr>
<td>(120:6)</td>
<td>5 × [120 × HfO2 + 6 × Al2O3] + 120 × HfO2</td>
<td>24</td>
<td>4.5</td>
</tr>
<tr>
<td>(200:10)</td>
<td>3 × [200 × HfO2 + 10 × Al2O3] + 200 × HfO2</td>
<td>48</td>
<td>5.4</td>
</tr>
<tr>
<td>(150:5)</td>
<td>3 × [150 × HfO2 + 5 × Al2O3] + 150 × HfO2</td>
<td>38</td>
<td>5.7</td>
</tr>
<tr>
<td>(100:2)</td>
<td>6 × [100 × HfO2 + 2 × Al2O3] + 100 × HfO2</td>
<td>24</td>
<td>2.3</td>
</tr>
<tr>
<td>(30:5)</td>
<td>2 × [300 × HfO2 + 5 × Al2O3] + 300 × HfO2</td>
<td>19</td>
<td>1.9</td>
</tr>
<tr>
<td>(100:1)</td>
<td>6 × [100 × HfO2 + 1 × Al2O3] + 100 × HfO2</td>
<td>43</td>
<td>-</td>
</tr>
<tr>
<td>HfO2</td>
<td>700 × HfO2, reference film</td>
<td>-</td>
<td>42</td>
</tr>
</tbody>
</table>

The PANalytical X’pert Pro MPD diffractometer was used to measure grazing incidence X-ray diffraction patterns with the incident beam angle of 1°. Energy dispersive X-ray spectrometry (EDX) was applied for the measurements of the hafnium to aluminum atomation ratio, and also for the estimation of the film thicknesses, using a Hitachi S-4800 scanning electron microscope (SEM) equipped with an Oxford INCA 350 EDX spectrometer. The EDX spectra were measured at 5 keV. The beam current and spectrometer gain were determined from a calibration measurement under the same beam conditions. The film thicknesses and ratios of the different elements were calculated from the k ratios of Hf Mα, and Al Kα X-ray lines measured with the calibrated beam. These calculations were done with a GMRFILM program.38 The thicknesses were calculated using a film density of 9 g/cm³. The selected nanolaminates on Si substrate were investigated by scanning electron microscopy (SEM) using FEI Helios Nanolab 600 DualBeam microscope equipped with focused ion beam (FIB) module and Omniprobe model 100.7 in-situ nanomanipulator. High-resolution transmission electron microscopy (HRTEM) studies for the characterization of the cross-sections of the nanolaminate structures were carried out using FEI Titan Themis 200 instrument with a field emission gun operated at 200 kV.

Magnetic measurements on selected samples were performed using vibrating sample magnetometer (VSM) option of the Physical Property Measurement System (PPMS) 14 T (Quantum Design), with the magnetic field parallel to the film surface. For electrical measurements, Al/Ti/HfO2–Al2O3/TiN/Si/Al capacitor stacks were constructed with the effective capacitor area of 0.204 mm². Double-layer 110 nm/50 nm thick Al/Ti dot electrodes were evaporated through a shadow mask on the top HfO2 layers by electron beam evaporation, the Ti layer being in direct contact to the top HfO2. Backside ohmic contact was provided by evaporating 100–120 nm thick Al layer on HF-etched Si. Electrical measurements were done by means of an Agilent DSO-X 3104 digital oscilloscope with a built-in wave generator. The standard Sawyer-Tower experiment was carried out by applying a periodic triangular-shaped stimulus and recording the voltage loops data from the oscilloscope. Charge values were obtained from the sensed voltage across a stabilized capacitance.

**Results and Discussion**

**Film structure.**—The HfO2-based films grown in the present study, except those mixed with Al2O3 in large amounts, tended to crystallize in the as-deposited state, as expected. The multilayered films consisted of distinct Al2O3 and HfO2 layers, as revealed by HRTEM images (Figure 1). One could also see that the very first HfO2 layers, closest to the Si substrate, occurred more disordered, while the crystal growth inside the hafnium oxide layers became more prominent toward the surface of the nanolaminate film.

The HfO2 films grown without mixing with Al2O3, i.e. the reference HfO2 films, were polycrystalline, as recognized by X-ray diffraction studies (Fig. 2), consisting of the stable monoclinic phase and metastable phases. The metastable phases of HfO2 to be considered are cubic, tetragonal and orthorhombic polymorphs. Almost all the reflection peaks in the GI-XRD pattern of the reference hafnium oxide can be attributed to the monoclinic HfO2. There is only one strong
peak, apparent at 30.3°, which does belong to some metastable phase and not to the monoclinic one. It is, however, to be mentioned, that many reflections originating from the different phases locate close to each other. Thus, it is rather complicated to unambiguously separate the monoclinic phase from the metastable ones at diffraction angles higher than 33–35°. It is also complicated to unambiguously distinguish between the orthorhombic, cubic, and tetragonal phases because even the strongest characteristic peak at 30.3° may shift due to internal stresses and be too wide, as characteristic of nanocrystalline materials, for the precise phase determination.

Also the nanolaminate films containing HfO2 layers grown to relatively large nominal thicknesses (300 cycles, nominally ca. 15 nm) between very thin (e.g. 5 cycles) intermediate Al2O3 layers, can be regarded as multiphase materials, but with strengthened contribution from the metastable polymorphs, decided mainly on the basis of the relative intensity of the peak at 30.3°.

It is to be noted that the aluminum to hafnium cation ratios as measured and revealed in the Table I, tend to be higher than those expected after considering the nominal ratios of Al2O3 and HfO2 growth cycles. This can, partially, become explained by difficulties in accurate determination of the content of metals in the films. Under the measurement conditions used here, it is more complicated for hafnium than for aluminum due to the proximity of the Hf Mα and Si Kα lines which may make the Hf content appear higher than it actually is. On the other hand, the relative content of aluminum may also increase through surface reactions between adsorbing AlCl3 and underlying HfO2, resulting in conversion of some HfO2 to the thermodynamically more stable Al2O3, similarly to the process considered earlier in the case of ZrO2-Al2O3 nanolaminates grown from ZrCl4, AlCl3 and H2O.39

Nanolaminate films built up on thin HfO2 films separated by amorphous intermediate layers, such as Al2O3 in this study, demonstrate mixed crystal structures with strong contribution from both metastable and stable HfO2 polymorphs. We have earlier observed that in HfO2-Ta2O5 nanolaminates, where 2–20 nm thick HfO2 intermediate layers were also grown in HfCl4-based ALD process, the monoclinic phase started to form in addition to the cubic/tetragonal phases in quite early stages of growth, i.e. after the HfO2 layer thickness exceeded 5 nm.40

In the HfO2-Al2O3 nanolaminates with somewhat thinner HfO2 layers of ca. 7 nm (Figure 3), separated by ca. 1 nm Al2O3 layers,
a metastable HfO2 polymorph with reflections designated to the orthorhombic lattice appeared dominating over the monoclinic phase. One could also see that post-deposition annealing as well as the growth of the films on nanocrystalline 10 nm thick TiN layer did not induce qualitative changes in the phase composition of the oxide films. These films were thus notably resistant against phase changes upon annealing.

In the HfO2-Al2O3 nanolaminates with HfO2 and Al2O3 layers both grown to even lower thicknesses, nominally 5–6 nm and 0.23–0.30 nm, respectively, the metastable phases were dominant at least in the as-deposited state (Figure 4). Also notable was the amorphous background in the GIXRD patterns. Quite aggressive annealing at 800 °C in air ambient for half an hour reduced the amorphous background significantly and increased the degree of crystallinity of the films, decided on the basis of the intensity and number of the diffraction peaks (Figure 4). Also the contribution from the monoclinic polymorph became apparent upon annealing.

**Electrical and magnetic performance.**—Selected films with the largest contribution from the metastable HfO2 phases were subjected to electrical measurements. The relative permittivity of the films tended to increase with the HfO2:Al2O3 cycle ratio. The relative permittivity, measured at 100 kHz, was increased from 14 to 22 (Table II) with the increase of Hf:Al cation ratio from 4.2 to 6.4. In the film containing aluminum below the EDX detection level, i.e. in the sample grown with the HfO2:Al2O3 cycle ratio of 1:1, the permittivity was measured somewhat lower again, not exceeding 18. In the latter film, the contribution from monoclinic phase of HfO2 was also increased compared to the samples with lower Hf:Al ratio and Al content measurable by EDX. One can see, that in the present series of films, the permittivity was somewhat correlated to the phase composition, increasing with the content of metastable phases expressed by the increasing intensity of the corresponding peaks in the XRD patterns, as was described above.

Somewhat surprisingly, the permittivity dispersion in the film grown with the HfO2:Al2O3 cycle ratio of 170:10 was exceptionally strong (not shown), allowing one to measure relative permittivities above 60 in the frequency range of 1–10 kHz, probably due to the presence of strong dipolar and interfacial polarization mechanisms. The apparently high permittivity and dispersion might arise from the strongly polycrystalline and mixed phase nature of this film. For the rest of the samples, the dispersion at low frequencies was essentially weaker, demonstrating only 10% increase in the permittivity toward measurement frequencies down to 1 kHz. These low-frequency permittivities are quite close to those reported in our earlier studies on HfO2 films grown by ALD in the temperature range of 300–400 °C from HCl and H2O. Certain increase in the permittivities compared to the earlier published results is likely connected to the increased amount of metastable hafnium oxide phases in the films grown in the present study. Cubic and tetragonal phases of HfO2 are known to...
possess higher permittivities than the stable monoclinic polymorph. Orientationally averaged static dielectric constants of 29, 70, and 16–18 have been obtained for the cubic, tetragonal, and monoclinic HfO₂ phases, respectively.²¹

Figure 5 demonstrates current-voltage behavior measured in resistive switching regime from the HfO₂-Al₂O₃ nanolaminates grown using the HfO₂-Al₂O₃ cycle ratio of 200:10. One can see that the films demonstrate current-voltage behavior similar to those studied as potential memristor materials. At both positive and negative bias voltages marked differences between the high and low resistance states were achieved, with a window ranging over almost an order of magnitude in the conduction current. The forming voltages, necessary to achieve the first highly conductive state in the material, varied between 1 and 2 V, which is still reasonably low voltage range considering the thickness of the films, exceeding 20 nm.

Figure 6 depicts the charge polarization–voltage loops measured by means of the Sawyer-Tower circuit from selected samples, in most of which the metastable HfO₂ phase or phases were dominating in accord with the GIXRD results. In these loops, one can recognize features characteristic of ferroelectric materials—charge being stored and released upon the application and removal of an electric field. The initial increase of the charge is most pronounced in the films grown with the cycle ratio of 200:10.

Table II. HfO₂:Al₂O₃ cycle ratio, Hf:Al cation ratio and permittivity, measured for selected representative films, in the ascending order for Hf content. Also indicated are the estimated values for the half-width of charge-field loops, designated as coercive field, and the charge density at zero field value, assigned as remnant polarization. The latter values are calculated for the charge-voltage loops, measured in the medium voltage range of −1 to 1 V, approximately, as depicted in Fig. 6.

<table>
<thead>
<tr>
<th>HfO₂:Al₂O₃ cycle ratio</th>
<th>Hf:Al cation ratio</th>
<th>Relative permittivity</th>
<th>Coercive field, MV/cm</th>
<th>Remnant polarization, mC/cm²</th>
</tr>
</thead>
<tbody>
<tr>
<td>(100:5)</td>
<td>4.2</td>
<td>14.0</td>
<td>0.23</td>
<td>8.3</td>
</tr>
<tr>
<td>(120:6)</td>
<td>4.5</td>
<td>15.8</td>
<td>0.18</td>
<td>9.2</td>
</tr>
<tr>
<td>(200:10)</td>
<td>5.4</td>
<td>22.0</td>
<td>0.20</td>
<td>2.3</td>
</tr>
<tr>
<td>(170:10)</td>
<td>6.4</td>
<td>22.0</td>
<td>0.25</td>
<td>12.5</td>
</tr>
<tr>
<td>(100:1)</td>
<td>Al not measurable</td>
<td>18.0</td>
<td>0.28</td>
<td>3.7</td>
</tr>
</tbody>
</table>

The coercive field was somewhat larger at low measurement temperatures, reaching 400 oe at 2 K in the film with a Hf:Al ratio of 5.4, i.e. in the nanolaminates grown with the HfO₂-Al₂O₃ pulse ratio of 200:10 (Table I) and possessing a structure of dominantly metastable HfO₂ polymorph (Fig. 3, upper panel). Within the composition and structure range of HfO₂-Al₂O₃ films explored in this study, the saturative behavior of both charge and magnetic polarization in external electric and magnetic fields was most clearly recognized in the film grown with the cycle ratio of 200:10.

Nonlinear saturative magnetization characteristic of ferromagnetic materials has been achieved and observed in undoped HfO₂ films in some works earlier, such as in 10–200 nm thick HfO₂ films grown by pulsed laser deposition⁴²–⁴⁴ and sputtering.⁴⁵ Magnetization in crystallized HfO₂ films has thereby been explained by presence of oxygen vacancies,⁴²–⁴⁵ and the magnetization indeed could be suppressed by annealing in oxygen environment, i.e. upon improvement of the stoichiometry of the dioxide.⁴²–⁴⁵ Magnetization in HfO₂ films has been similar to that in soft ferromagnetic materials with narrow hysteresis, which was observed also in the present study.

The residual impurities in the films can affect their physical properties, in general. In terms of electrical performance, the existence of measurable content of residual chlorine may, quite likely, increase the conductivity of the films and, thus, also the role of interfacial polarization contributing to the polarization charge and the remnant polarization as recorded at zero sample voltage. At the same time, the existence of residuals can assist in the stabilization of artifically defective structure, up to the partial formation of metastable hafnium oxide phases, and, in this way, give rise to magnetization of the solid films.

In ALD processes using metal halide and water precursors, the amounts of residual impurities decrease rapidly and nonlinearly with the increase in the growth temperature in the range of 200–600 °C. In regard with the ALD process based on the HfCl₄ and H₂O precursors, ion beam analysis has shown residue contents in the ranges of 0.1–0.6 and 0.5–2.3 at.% for chlorine and hydrogen respectively, in films deposited at 300 °C. In the present study, the amount of residual chlorine in the films grown at 350 °C remained, quite unexpectedly, lower than the limit for its reliable detection by EDX.

Summary

HfO₂-Al₂O₃ films and nanolaminates were grown by atomic layer deposition from hafnium tetrachloride, aluminum trichloride and water on silicon and titanium nitride substrates. The main goal of the work was to study the feasibility of the stabilization of the metastable polymorphs in HfO₂ by alternate layering of crystalline HfO₂ and amorphous Al₂O₃. The HfO₂-Al₂O₃ ALD cycle ratio was varied between 200:10 and 15:1 in order to grow of multilayers consisting of distinct binary oxides or mixtures and, by this way, tune the phase composition of the resulting films. The contribution of the metastable cubic and, possibly, orthorhombic HfO₂ polymorphs increased with the relative content and thickness of aluminum oxide layers, while the intensity of the crystal growth in the HfO₂ layers increased with their thickness.
The films exhibited polarization charge-electric field loops, affected by interfacial polarization. Two distinctive resistance states were observed in the current-voltage loops of the nanolaminate films as characteristic of resistive switching memory materials. The films could also be saturatively magnetized and they exhibited soft ferromagnetic behavior with weak coercive field in the magnetization-field hystereses. The magnetoelectric properties were recognized at room temperature and most clearly in the well-defined nanolaminate film.
grown with the HfO2:Al2O3 cycle ratio of 200:10. In this nanolaminate, the metastable polymorphs of HfO2 dominated the phase composition.

Acknowledgments

The study was partially supported by the Finnish Centre of Excellence in Atomic Layer Deposition (284623), European Regional Development Fund project “Emerging orders in quantum and nanomaterials” (TK134), Spanish Ministry of Economy and Competitiveness (TEC2014-52152-C3-3-R) with support of Feder fund, Estonian Research Agency (IUT2-24, IUT23-7, PRG4), and Estonian Academy of Sciences (SLTFYPROP).

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References

7. C. Mahata, Y.-C. Byun, C.-H. An, S. Choi, Y. An, and H. Kim, “Comparative study of atomic-layer-deposited stacked (HfO2/Al2O3) and nanolaminated (Hf/AlOx) di-electrics on In0.53Ga0.47As,” ECS Appl. Mater. Interfaces, 5, 4195 (2013).
8. E. Cianci, A. Molle, A. Lamperti, C. Wiemer, S. Spiga, and M. Fanciulli, “Phase stabilization of Al2O3/In0.53Ga0.47As grown on In0.53Ga0.47As substrates (x=0.00, 0.15, 0.53) via trimethylaluminum-based atomic layer deposition,” ECS Appl. Mater. Interfaces, 6, 3455 (2014).


