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Toward epitaxial ternary oxide multilayer device stacks by atomic layer deposition

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The authors demonstrate multilayer epitaxial films by atomic layer deposition and postdeposition annealing. Their example features two ABO3 type perovskite oxide films with different materials properties—a conductor (LaNiO3) and an insulator (SrTiO3)—that can be integrated epitaxially once the geometric interaction between the two oxides’ lattices is understood. Once preliminary epitaxial materials had been developed, the pilot multilayer device fabricated was an epitaxial metal-insulator-metal structure with layers of ∼5 nm thickness. This work shows the potential for advanced device types based on epitaxial atomic layer deposited films, assuming that care is taken in the selection of processes and starting substrate. Published by the AVS. 

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I. INTRODUCTION

Atomic layer deposition (ALD) has already delivered unparalleled thickness, conformity, and composition control in thin films—enabling the continued scaling of MOSFET and DRAM devices to feature sizes previously thought impossible.1 Additionally, ALD is now exploited as a feature-defining tool utilizing the self-aligned layer principle to set the tolerance of device dimension into the nanoscale.2 Despite its inventor Suntola originally coining the ALD technique “Atomic Layer Epitaxy,”3 most ALD films are amorphous or polycrystalline. Further device-enabling materials property control is potentially available from films that are single-crystal and registered to the substrate. The advantages this class of materials offer are improved materials properties and better-defined interfaces for improved device properties. For example, the reduction or elimination of the grain boundary network in ultra-thin high-κ layers, and preservation of atomically sharp interfaces, would be expected to reduce leakage current.

Several groups have reported epitaxial ternary and quaternary films by ALD, focusing on materials with exotic device properties like ferromagnetic,4 multiferroic,5 and superconducting6 layers. A group from the University of Texas at Austin has made huge inroads into perovskite integration onto Si and Ge substrate platforms.7 These examples in the literature show that ternary and quaternary epitaxial layers can be produced by ALD, an established manufacturing technique which is already vital for advanced electronic device fabrication.

For the device integration of epitaxial materials, it is necessary to form heterostructures from multilayer stacks of thin films, in order to optimize the materials through the stack and across each interface. Such heterostructures have been fabricated by pulsed laser deposition and molecular beam epitaxy in the literature,8,9 but metal-insulator-metal (MIM)-type structures are uncommon due to the paucity of options for an epitaxial oxide conductor. In this letter, we describe the use of conductive (LaNiO3) and insulating (SrTiO3) perovskite oxide thin films to demonstrate the possibility of fabricating epitaxial heterostructures by ALD and annealing steps.

II. EXPERIMENT

Depositions were performed at 275 °C in an ASM F-120 hot-wall flow type reactor with N2 as the process gas and a working pressure of ∼10 mbar. Substrates were SrTiO3(100) (Pi Kem Ltd.) prepared using a published methodology.10 One-side-polished SrTiO3(100) was heated in air for 1 h at 1000 °C, cooled, and agitated in de-ionized water for 30 s. This process was performed twice in order to provide atomically flat starting surfaces for deposition. The LaNiO3 process used La(2,2,6,6-tetramethyl-3,5-heptanedionato)3, Ni(2,2,6,6-tetramethyl-3,5-heptanedionato)2 (both from Volatec Oy), and O3 as precursors.11 The SrTiO3 process used Sr(1,2,4-tri-tert-butylcyclopentadienyl)2 (Air Liquide), Ti(O(Me)3 (Sigma Aldrich), and deionized H2O.12 The deposition recipes were organized in subcycles of 10 cycles of each binary oxide, e.g., [10(Ni(thd)2/O3)+ 10(La(thd)3/O3)]n, with n supercycles then performed to result in the desired film thickness. This approach has been found to promote stoichiometry in the resulting ternary oxide after the layers intermix during heat treatment. Films were measured using ellipsometry (Film Sense FS-1) and grazing-incidence x-ray diffraction (GIXRD, Rigaku SmartLab). Crystallization temperatures of both oxides were found by variable-temperature XRD in an Anton Paar HTK1200N oven fitted to a PANalytical X’Pert Pro MPD diffractometer. Cross-sectional

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samples for scanning tunneling electron microscope (STEM) analysis were prepared by a Focused Ion Beam instrument. High-angle annular dark field (HAADF)-STEM and energy-dispersive x-ray spectroscopy (EDS) images for the samples were taken using a probe aberration corrected STEM (FEI Titan Cubed 60-300) operating at 300 keV with a convergence semi-angle of 26.7 mrad for the incident electron probe. Stoichiometry of LaNiO₃ was measured by time-of-flight elastic recoil detection analysis (TOF-ERDA) measurements, after which deposition parameters could be optimized. The resistivity was assessed by averaging 5 four-point probe measurements made across each film using a Keithley 2400-SourceMeter and Alessi C4S four-point probe head. This average sheet resistance was then converted into resistivity with the film thickness.

III. RESULTS AND DISCUSSION

A. Epitaxial LaNiO₃/SrTiO₃(100)

LaNiO₃ is considered a so-called “bad metal” — a correlated oxide material that exhibits low resistivity if the oxygen stoichiometry can be preserved during deposition, and the Ni valence state can be maintained as close to 3+ as possible. The LaNiO₃ material can display insulating (for majority of valence state can be maintained as close to 3+ as possible) and metallic conducting (for majority Ni³⁺ properties). In the epitaxial MIM structure presented later, LaNiO₃ acts as the conductor because epitaxial metals are extremely difficult to produce, especially in stacks where heat treatment is employed to crystallize layers.

LaNiO₃ films were amorphous as-deposited and were crystallized by annealing in air at 750 °C. Compositional measurements by TOF-ERDA confirmed the ABO₃ stoichiometry of the deposited oxide. Initial attempts to form epitaxial films from layers with a thickness in the order of 30 nm were unsuccessful as polycrystalline films resulted. It was found that epitaxial film formation only worked for layers below a thickness limit of ~10 nm. Figure 1 is an HAADF-STEM image of such a film with a thickness of ~8.5 nm.

The primitive cell of LaNiO₃ has rhombohedral symmetry with the space group R3c, but the material forms a pseudocubic arrangement on SrTiO₃(100) due to a very close match in the lattice parameters (0.391 nm for pseudo-LaNiO₃, 0.383 nm for SrTiO₃). The substrate/film lattice continuity can be seen clearly in Fig. 1 in addition to what appears to be antiphase boundaries, defects in the lattice that are presumed to relieve the strain of the pseudocubic epitaxial relationship. To make thicker epitaxial LaNiO₃ films, a seed layer methodology was necessary where a thin seed layer less than this ~10 nm thickness limit for epitaxy was deposited and crystallized first, before additional growth was added and crystallized by a second anneal. Using this method with a 5 nm seed, films of (5 + 25) nm were grown and compared with layers that were grown as a 30 nm continuous deposition. Figure 2(a) is a θ/2θ XRD measurement for a 30 nm LaNiO₃ film, which can be compared with a measurement on a seed-layered (5 + 25) nm film in Fig. 2(b).

To confirm the epitaxial relationship between the substrate and film in the [5 + 25] nm sample, a reciprocal space map (RSM) was recorded around the SrTiO₃(103) reflection, as presented in Fig. 3. The LaNiO₃(103) (pseudocubic) reflection is noted in the image, confirming the epitaxial arrangement and demonstrating the close in-plane match between the substrate and film (x axis in the RSM). The [5 + 25] nm film had a low resistivity of ~300(±50) μΩ cm which compares well with the data obtained from the seed-layered film.

![Diagram](image-url)

**Fig. 1.** HAADF-STEM image of an ~8.5 nm epitaxial LaNiO₃ film on SrTiO₃(100) after annealing. The top black area is a carbon film deposited prior to FIB sectioning. Scale bar is 5 nm.

**Fig. 2.** XRD summary for LaNiO₃ films grown on SrTiO₃(100) and annealed. (a) θ/2θ measurement on a 30 nm film showing polycrystalline reflections; notably the (110) reflection ~33°. (b) θ/2θ measurement on a [5 + 25] nm film where a 5 nm seed layer is annealed before subsequent deposition. The sharp, unlabeled peaks in the two diffractograms are from Cu Kβ and W Lx1 signals. The data are logarithmic scale in the y axis, x axis units are degrees 2θ.
with values for this material in the literature.\textsuperscript{16,17} Thinner films such as those depicted in Sec. III B would be expected to have higher resistivity. The \( \sim 8.5 \text{ nm} \) \LaNiO\textsubscript{3} film shown in Fig. 1 was found to have resistivity closer to \( \sim 1500 \mu \Omega \text{ cm} \).

B. Multilayer epitaxial \LaNiO\textsubscript{3}/\SrTiO\textsubscript{3}/\LaNiO\textsubscript{3}/\SrTiO\textsubscript{3}(100)

Multilayer epitaxial samples were formed by keeping the layer thicknesses \( \sim 5 \text{ nm} \) and alternating \LaNiO\textsubscript{3} and \SrTiO\textsubscript{3} deposition, with each layer having been annealed at 750 °C in air after growth to form a crystalline layer. The \SrTiO\textsubscript{3} layer was found to develop polycrystallinity if the layer thickness was greater than \( \sim 10 \text{ nm} \) which is similar to the epitaxy thickness limit seen for the \LaNiO\textsubscript{3} films grown on \SrTiO\textsubscript{3}(100) substrates. XRD performed at a range of temperatures showed the crystallization temperature of both of these oxides to be <600 °C; so the thermal budget for fabrication could potentially be kept lower than that used here. Figure 4 is an HAADF-STEM image of a typical FIB section of the sample, demonstrating that three concurrent epitaxial layers had formed in a heterostructure on the \SrTiO\textsubscript{3} substrate.

Three FIB sections were analyzed, and common to each was the breakdown of the top \LaNiO\textsubscript{3} layer in the stack, as can be seen in Fig. 4 (denoted \LaNiO\textsubscript{3}/\La\textsubscript{2}O\textsubscript{3} in the figure). STEM-EDS measurements showed a Ni deficiency in this top layer but whether there was an issue with the deposition process itself or this breakdown happened during the annealing step is currently not certain. Careful refabrications of the stack depicted in Fig. 4 have so far all resulted in some polycrystallinity in the topmost layer. Figure 5 presents GIXRD measurements made on a multilayer sample after each layer had been deposited and annealed. The GIXRD measurement setup is usually not considered useful for epitaxial films but it was found to be a vital and sensitive tool to check for failure of epitaxy for each individual layer during fabrication. If all layers in the sample crystallize epitaxially, the GIXRD should show only background signal. The small peak at \( \sim 30^\circ \) in Fig. 5 shows the formation of randomly orientated crystallites after the third layer had been annealed. Scans made over a wider angular range were best fitted to hexagonal \La\textsubscript{2}O\textsubscript{3} XRD pattern. The information in Figs. 4 and 5 is congruent and demonstrates the power of GIXRD to monitor the quality of epitaxial ultra-thin films.

Figure 4 demonstrates that it is possible to produce epitaxial device stacks within a modest thermal budget (275 °C deposition and 600–750 °C heat treatment), and it now remains to improve the quality of the samples produced. As the specimens discussed here involved vacuum breaks after each deposition and an air anneal, it is likely that eliminating
these sources of contamination would improve the interface and film quality. Additionally, it would be interesting to see if such layer stacks could be deposited on a substrate like Ge as has been demonstrated recently by the Texas group in the case of BaTiO₃.¹⁸

IV. SUMMARY AND CONCLUSIONS

The plausibility of fabricating multilayer epitaxial films by ALD in simple stack device configurations was presented. A pilot MIM-type device was fabricated utilizing ALD processes for LaNiO₃ and SrTiO₃, combined with postdeposition heat treatment. These results pave the way for the fabrication of advanced ALD-based material devices and are only a preliminary indication of what is likely to be possible with proper fabrication equipment and process knowledge.

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¹²Q. Lei et al., npj Quantum Mater. 2, 10 (2017).