

## Thin Films Division

### Room A124-125 - Session TF+PS-TuA

#### Epitaxial Thin Films

**Moderator:** Robert Grubbs, Micron Technology

**2:40pm TF+PS-TuA2 Van der Waals Layer Promoted Heteroepitaxy in Sputter-deposited Transition-metal Carbide and Sulfide Thin Films, Koichi Tanaka<sup>1</sup>, P. Arias, M.E. Liao, Y. Wang, H. Zaid, A. Aleman, University of California, Los Angeles; K. Hojo, Nagoya University, Japan; A. Deshpande, M.S. Goorsky, S. Kodambaka, University of California, Los Angeles**

Over the past decade, two-dimensional (2D) layered materials such as graphene, MoS<sub>2</sub>, etc., have attracted considerable attention for a variety of applications, primarily in nanoelectronics and optoelectronics. An exciting and relatively little explored application of these van der Waals (vdW) layered materials is their use as templates for crystal growth. In the recent years, vdW layers present at the substrate-film interface have been shown to promote 'remote epitaxy', by relaying the epitaxial registry between the film and the substrate.

Here, we demonstrate that the crystallinity of sputter-deposited thin films can be significantly improved using vdW layered materials as buffer layers on growth substrates. Using 2D hexagonal boron nitride (hBN,  $a = 0.250$  nm and  $c = 0.667$  nm) as the buffer layer, we grow hexagonal-MoS<sub>2</sub> ( $a = 0.315$  nm and  $c = 1.23$  nm), trigonal-structured Ta<sub>2</sub>C ( $a = 0.310$  nm and  $c = 0.494$  nm), and NaCl-structured TaC ( $a = 0.446$  nm) of desired thickness on Al<sub>2</sub>O<sub>3</sub>(0001) substrates via ultra-high vacuum direct current magnetron sputtering of Mo and TaC targets respectively, in Ar/C<sub>2</sub>H<sub>4</sub> and Ar/H<sub>2</sub>S gas mixtures. hBN layers are deposited in the same system via pyrolytic cracking of borazine (~600 L) onto the substrates at prior to the growth of the thin films. The as-deposited films are characterized using a combination of *in situ* using Auger electron spectroscopy and low-energy electron diffraction and *ex situ* X-ray diffraction (XRD), X-ray photoelectron and Raman spectroscopies, and transmission electron microscopy (TEM) based techniques.

We find notable differences in the layers deposited on hBN-covered Al<sub>2</sub>O<sub>3</sub>(0001) compared to those grown on bare substrates: significantly stronger 0002 (or 111 in case of TaC) reflection intensities and observation of Laue oscillations in  $\omega$ -2 $\theta$  XRD scans and higher intensity of MoS<sub>2</sub> characteristic peaks in Raman spectrum. Furthermore, we show that inserting hBN layers at regular intervals results in highly-0002-oriented growth and suppression of polycrystallinity in thicker Ta<sub>2</sub>C films. Our results indicate that hBN layers enhance the crystallinity, irrespective of the crystal structure, of sputter-deposited thin films.

**3:00pm TF+PS-TuA3 Molecular Beam Epitaxy Applied to Tensile-Strained Quantum Dots for Quantum Optics and Band-Structure Engineering, Paul Simmonds, Boise State University** **INVITED**

Since the early 1990s, solid-state self-assembled quantum dots (QDs) have been the subject of intensive research for devices and technologies ranging from high-stability lasers, to intermediate band solar cells. Driven by compressive strain, semiconductor QDs form spontaneously on the (001) surfaces of both III-V and group IV materials during growth by molecular beam epitaxy (MBE). But several years ago, I became interested in the question of why QD self-assembly seemed to be limited to materials with this specific combination of compressive strain, and a (001) surface orientation. For example, why could we not grow QDs under *tensile* rather than compressive strain or on non-(001) surfaces, especially since QDs with these characteristics are predicted to be highly desirable for certain applications. The low fine-structure splitting of (111) QDs should make them ideal entangled photon sources; tensile-strained QDs would have dramatically reduced semiconductor band gaps, with implications for infrared optoelectronics and nanoscale band structure engineering.

The first step towards answering this question was to understand how the competition between plastic and elastic strain relief mechanisms made it enormously challenging to synthesize non-(001) or tensile-strained QDs without the formation of crystallographic defects. The outcome of this analysis was the discovery of a robust new approach to QD self-assembly based on MBE that overcomes these difficulties, and enables the reliable, controllable growth of defect-free, tensile-strained QDs on (111) and (110) surfaces.

I will describe the model upon which tensile-strained QD self-assembly is founded, and then discuss the application of this novel growth mode to several different material systems. I will present data confirming that the (111)-oriented QDs we can now grow do indeed show promise as entangled photon sources. I will highlight the possibilities for band structure engineering that are now available with tensile-strained QDs, using the example of transforming germanium into a direct band gap semiconductor.

In summary, I hope to demonstrate that tensile-strained self-assembly represents a powerful new tool for heterogeneous materials integration, and nanomaterial development.

**5:00pm TF+PS-TuA9 Low-temperature Homoepitaxial Growth of N-type Superlattices for Ultrastable, Ultrafast X-Ray and Charged Particle Detectors, April Jewell, Jet Propulsion Laboratory, California Institute of Technology; M.E. Hoenk, Jet Propulsion Laboratory; Q. Looker, M.O. Sanchez, B.D. Tierney, Sandia National Laboratories; A.G. Carver, Jet Propulsion Laboratory; S. Nikzad, Jet Propulsion Laboratory, California Institute of Technology**

We present a low-temperature process for the homoepitaxial growth of antimony superlattices in silicon. The all low temperature superlattice doping process is compatible as a post-fabrication step for device passivation. We have used low-temperature molecular beam epitaxy (MBE) to embed atomically thin (2D), highly concentrated layers of dopant atoms within nanometers of the surface. This process allows for dopant densities on the order of  $10^{13}$ - $10^{14}$  cm<sup>-2</sup> ( $10^{20}$ - $10^{21}$  cm<sup>-3</sup>); higher than can be achieved with three-dimensional (3D) doping techniques. This effort builds on our prior work with n-type delta doping; we have optimized our growth processes to achieve delta layers with sharp dopant profiles. By transitioning from a standard effusion cell to a valved cracker cell for antimony evaporation, we have achieved carrier densities approaching  $10^{21}$  cm<sup>-3</sup> with peak distribution at  $\sim 10$  Å FWHM for single delta layers. We will discuss details related to growth optimization, and show results from *in situ* monitoring by electron diffraction. We will also report on elemental and electrical characterization of our films.

The performance of our low-temperature 2D-doping processes has been validated by applying both p-type and n-type superlattice-doping to fully depleted photodiodes. The superlattice-doped devices show significantly higher responsivity than the equivalent ion-implanted devices. Additionally, when exposed to pulsed X-rays the superlattice-doped devices exhibit fast response and recovery times required for use in pulsed power experiments.

**5:20pm TF+PS-TuA10 Epitaxial Growth of Ultrathin Molybdenum Nitrides on Ru(0001) and Ag(100), Asim Khaniya, M. Sajid, A. Kara, W.E. Kaden, University of Central Florida**

Molybdenum-nitrides are known to possess interesting mechanical, electronic, and catalytic properties. For example, (i) hexagonal  $\delta$ -MoN exhibits mechanical elasticity and hardness values comparable to cubic BN and diamond, (ii) both hexagonal and cubic phases of molybdenum nitrides are known to be superconducting, and (iii) mixed-phase structures have been shown to outperform commercial hydrotreatment catalysts for selective nitrogen removal from heterocyclic organic feedstocks. To better understand these properties, many groups have worked to create improved recipes to grow different phase-pure crystallographic phases of the material. To-date, the most successful procedures have leveraged epitaxy to improve long-range bulk order, but have lacked the well-defined, planar terminations suitable for controlled surface-science investigations. To establish such samples, our group has opted to use low energy nitrogen ions in tandem with molybdenum physical vapor deposition to grow and characterize molybdenum-nitride films on Ru(0001) and Ag(100) supports, which have been chosen to template the growth of hexagonal and cubic phases of the nitride. At the time of this abstract submission, we have succeeded in the growth of a  $\delta$ -MoN-like film that appears to grow layer-by-layer and in registry with the Ru(0001) support, and are now in the early stages of repeating the process to create  $\gamma$ -Mo<sub>2</sub>N on Ag via an analogous process. This talk will focus on the interesting aspects of these materials (particularly those relevant to catalysis), our approach to film preparation, and a thorough analysis of the physical properties of the resultant films and growth modes via: XPS, LEED, He<sup>+</sup> Ion Scattering Spectroscopy, STM, and DFT.

# Tuesday Afternoon, October 22, 2019

5:40pm **TF+PS-TuA11 Using Time and Temperature of the Purge Step to Control Crystallinity, Phase Assemblage, and Epitaxy in Atomic Layer Deposited (ALD) Thin Films**, *Mark Losego, B.D. Piercy, R.J. Petrie*, Georgia Institute of Technology

The purge step between precursor and co-reactant doses in an atomic layer deposition (ALD) process is often viewed as a process liability. The goal for most manufacturing processes is to make this purge step as short as possible without disrupting the quintessential self-limited growth of ALD. In our lab, we have instead viewed this purge step as a potential opportunity to influence the crystallinity and phase assemblage of our materials. In actuality, each of these purge steps are an opportunity to allow surface diffusion to rapidly reform the film's microstructure before the next layer is deposited. Throughout the literature are interesting, but often conflicting reports of how ALD films crystallize with temperature and thickness. In our recent work, we have asked some simple questions, like how does the onset of such crystallinity change with purge time? We have found, for example, that the onset of anatase formation in the  $\text{TiCl}_4\text{-H}_2\text{O}$  ALD system can be reduced by more than 40 °C by simply extending the purge time between each cycle. While potentially time intensive, these results have implications for depositing crystalline materials on temperature-sensitive substrates, like polymers. We also find that often an initial seeding of the crystallinity can lead to accelerated growth of crystalline phases with subsequent cycles. In a second paradigm to be discussed, we have introduced a high-temperature pulsed heating source to an ALD system to intentionally crystallize materials and drive epitaxial growth. As proof-of-concept, we have studied epitaxial growth of ZnO on c-plane sapphire using a diethylzinc (DEZ) / water chemistry. DEZ is known to decompose above about 180°C, and the DEZ-H<sub>2</sub>O system cannot be grown epitaxially on c-sapphire with traditional thermal ALD approaches. Here, we show that heating pulses up to 900°C can be used to drive epitaxy. Interestingly, we find that a template layer of only 20 pulsed heating ALD cycles is sufficient to template ZnO epitaxy with subsequent low temperature ALD growth (180 °C) to film thicknesses of up to 100 nm.

6:00pm **TF+PS-TuA12 The Role of Template Layers in Heteroepitaxial ALD Growth of Crystalline  $\text{La}_2\text{O}_3$  on  $\text{GaN}(0001)$** , *Pei-Yu Chen, T. Hadamek*, University of Texas at Austin; *S. Kwon*, University of Texas at Dallas; *F. Al-Quaiti, A. Posadas*, University of Texas at Austin; *M.J. Kim*, University of Texas at Dallas; *A.A. Demkov, J.G. Ekerdt*, University of Texas at Austin

The high switching frequency, operating temperatures and voltages make GaN the material of choice for higher power applications and instrumental to reducing power consumption. In many of these applications, there is a need for a high quality gate dielectric. Lanthanum sesquioxide,  $\text{La}_2\text{O}_3$ , is one of the promising gate insulator candidates. In this work, we compare  $\text{La}_2\text{O}_3$  thin films grown by atomic layer deposition (ALD) and molecular beam epitaxy (MBE), and explore the formation of ALD- $\text{La}_2\text{O}_3$  films on  $\text{GaN}(0001)$ . An island growth mode (Volmer-Weber growth) was observed when  $\text{La}_2\text{O}_3$  films were deposited directly on  $\text{GaN}(0001)$  at 250 °C by ALD using tris(*N,N'*-diisopropylformamidinato)-lanthanum as the precursor and  $\text{H}_2\text{O}$  as the co-reactant. Only with use of a thin template layer, 2 nm-thick hexagonal  $\text{La}_2\text{O}_3$  grown by MBE or 3 nm-thick cubic  $\text{Er}_2\text{O}_3$  grown by ALD, can a 2-dimensional ALD- $\text{La}_2\text{O}_3$  thin film be formed. The 2-dimensional ALD- $\text{La}_2\text{O}_3$  growth on templated- $\text{GaN}(0001)$  was confirmed by RHEED and AFM. The macrostructure and microstructure of ALD- $\text{La}_2\text{O}_3$  films were verified with XRD, STEM, and atomic structure modeling. The ALD- $\text{La}_2\text{O}_3$  film retains a cubic structure on ALD- $\text{Er}_2\text{O}_3$  templated- $\text{GaN}(0001)$  while it transforms from the cubic phase to mixture of cubic and hexagonal phases on MBE- $\text{La}_2\text{O}_3$  templated- $\text{GaN}(0001)$  when the film is thicker than 15 nm. Hexagonal  $\text{La}_2\text{O}_3$  is more thermodynamically stable than cubic bixbyite  $\text{La}_2\text{O}_3$ ; the stabilization of cubic ALD- $\text{La}_2\text{O}_3$  on ALD- $\text{Er}_2\text{O}_3$  templated- $\text{GaN}(0001)$  can be attributed to the use of the cubic ALD- $\text{Er}_2\text{O}_3$  template and relatively low growth temperature. Analogies are presented for the  $\text{In}_2\text{O}_3$  system, which has similar cubic bixbyite and hexagonal structures as  $\text{La}_2\text{O}_3$ , except the phases are reversed in  $\text{In}_2\text{O}_3$ . We calculate the surface energy of hexagonal  $\text{In}_2\text{O}_3$  and compare the result with reported cubic  $\text{In}_2\text{O}_3$  values to explore the relative contribution of bulk and surface energies in stabilizing the structure of thin crystalline films. Stabilization of thin cubic ALD- $\text{La}_2\text{O}_3$  on hexagonal MBE- $\text{La}_2\text{O}_3$  templated- $\text{GaN}(0001)$  is attributed to likely surface energy differences between cubic and hexagonal  $\text{La}_2\text{O}_3$ .

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