

PCSI

Room Ballroom South - Session PCSI-SuA

New Developments in Oxide Materials & Growth

Moderator: Alex Demkov, The University of Texas

4:00pm PCSI-SuA-1 Crystal-Chemical Origins of the Ultrahigh Conductivity of Metallic Delafossites, *Chris Leighton*, University of Minnesota **INVITED**

The delafossites are a class of complex oxides with general formula ABO_2 that have been available synthetically since 1971 [1]. Some of these delafossites are metallic, where conductive triangular sheets of A^{+} ions are interspersed with insulating $B^{3+}O_6$ edge-sharing octahedral layers, generating a remarkably simple electronic structure at the Fermi level [1,2]. Only in the last 10-20 years, however, was it understood that despite their highly anisotropic complex-oxidic nature, metallic delafossites (particularly $PdCoO_2$ and $PtCoO_2$) are the *most conductive oxides known*, for reasons that remain poorly understood [1]. In particular, their room-temperature resistivity is lower than Au and their low-temperature resistivity falls as low as 8 nW cm, implying mean-free-paths of ~ 20 nm [1]. These extraordinary values have led to a slew of recent advances [1]. To reach such low-temperature values, it is widely accepted that these materials must be ultrapure and ultraperfect, although the methods for their bulk growth (which produce only small crystals) are not typically capable of such [1].

In this presentation, we first report a new approach to $PdCoO_2$ crystal growth, using a novel chemical vapor transport method to achieve order-of-magnitude gains in size and mass, the highest structural qualities yet reported, and record residual resistivity ratios ($440 < RRR < 760$) [3]. Nevertheless, the first detailed mass spectrometry measurements on these materials reveals that they are *not* ultrapure, typically harboring 100s-of-part-per-million impurity levels [3]. Through detailed crystal-chemical analyses, we resolve this apparent dichotomy, showing that the vast majority of impurities are forced to reside in the insulating Co-O octahedral layers, leaving the conductive Pd sheets highly pure (~ 1 ppm impurity concentrations). These purities are shown to be in quantitative agreement with measured residual resistivities [3]. We thus conclude that a previously unconsidered "sublattice purification" mechanism is essential to the ultrahigh low-temperature conductivity and mean-free-path of metallic delafossites [3], opening up many exciting device possibilities.

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[1] Mackenzie, *Rep. Prog. Phys.* **80**, 032501 (2017).

[2] Zhang, Saha, Tutt, Chaturvedi, Voigt, Moore, Garcia-Barriocanal, Birol and Leighton, *Phys. Rev. Mater.* **6**, 115004 (2022).

[3] Zhang, Tutt, Evans, Sharma, Haugstad, Kaiser, Ramberger, Bayliff, Tao, Manno, Garcia-Barriocanal, Chaturvedi, Fernandes, Birol, Seyfried Jr., Leighton, arXiv:2308.14257

4:40pm PCSI-SuA-9 Superconductivity and Magnetism in Infinite-Layer Nickelate Heterostructures, *Jennifer Fowlie*, SLAC National Lab **INVITED**

Nickel and copper are nominally very similar in chemistry so the search for superconductivity in nickelates is a story as old as the quest to understand the high temperature superconductivity of the cuprates.

In this talk, I will introduce the recent discovery of superconductivity in infinite-layer nickelates [1] and the ever-growing family of nickelate superconductors. I will touch on some of the materials challenges involved before summarizing the key physics we have learned so far including results from x-ray scattering [2] that identify a Mott-Hubbard-like character to the infinite-layer nickelate electronic structure as well as a significant rare earth 5d influence at the Fermi level. In particular I will focus on muon spin rotation [3] that reveals local magnetism in these materials that 1) onsets at rather high temperature, 2) is independent of the rare earth 4f electrons, 3) appears to be robust to doping 4) is antiferromagnetic and possibly short-range-ordered in nature and 5) coexists with superconductivity at low temperatures.

Finally, I will come back to the comparison between nickelates and cuprates and discuss how the disparities in the magnetic properties may be understood.

[1] D. Li et al, *Nature* **572**, 624 (2019).

[2] H. Lu et al, *Science* **373**, 213 (2021).

[3] J. Fowlie et al, *Nat. Phys.* **18**, 1043 (2022).

Sunday Afternoon, January 14, 2024

5:20pm PCSI-SuA-17 The Redox Chemistry of Oxide Molecular Beam Epitaxy, *Oliver Bierwagen*, Paul-Drude-Institut für Festkörperelektronik Leibniz-Institut im Forschungsverbund Berlin, Germany

The materials class of crystalline oxides provides a plethora of functional (dielectric, semiconducting, superconducting, ferroelectric, or ferromagnetic) properties. For harnessing this potential, molecular beam epitaxy (MBE) has proven an established method that realizes high quality oxide thin films. In its mostly used variety, the corresponding cation is evaporated from an effusion cell in vacuum onto the heated substrate where it gets oxidized by molecular oxygen, ozone, or an oxygen plasma.

This contribution will review the peculiar redox chemistry of oxide MBE, related to the existence of suboxides with significantly higher vapor pressure than their cation elements, not only taking place at the growth surface but also in effusion cells. I will discuss the implications of this chemistry on growth rate, film composition, and flux emanating from the effusion cells with the example of the plasma-assisted MBE growth of the semiconducting oxides Ga_2O_3 , In_2O_3 , GeO_2 , SnO_2 and its suboxide SnO . Three major reactions describe the suboxide-related chemistry:

(1.) The reaction of the metal with its oxide towards the suboxide, which can be utilized to etch oxide films in-situ [1,6], but also provides efficient suboxide sources [2].

(2.) The reaction of the cation element with oxygen, resulting in unintentional suboxide fluxes from elemental sources, leading to unexpectedly high cation incorporation in the grown films [3]. On the substrate, the same reaction provides the *p*-type oxide SnO [4] as well as an efficient way of removing a cation layer in-situ. Generally, the suboxide formation on the growth front is the first step during oxide growth, its desorption limits the growth rate.

(3.) The reaction of the suboxide with activated oxygen in the source leads to source passivation [9]. On the growth front, it is the second step that completes oxide growth. A kinetic growth model that involves both these steps, describes the observed, peculiar growth rate dependence on metal-to-oxygen flux ratio and substrate temperature for SnO_2 , Ga_2O_3 , In_2O_3 [5], and GeO_2 [6]. A simpler, single- and zero-step growth kinetics can be realized when the suboxide instead of an elemental cation flux is provided to the growth front, as demonstrated for SnO_2 [2], Ga_2O_3 [7], and SnO [8] films grown by suboxide MBE (S-MBE).

5:25pm PCSI-SuA-18 Optical Phonon Modes in $LaInO_3$: Lattice Dynamics and Complete Polarization Analysis of Raman-Active Modes, *Hans Tornatzky*, Paul-Drude Institute for Solid State Electronics, Germany; *Z. Galazka*, Institut für Kristallzüchtung, Germany; *R. Gillen*, Friedrich-Alexander-University Erlangen-Nürnberg (FAU), Germany; *O. Brand*, *M. Ramsteiner*, *M. Wagner*, Paul-Drude Institute for Solid State Electronics, Germany

$LaInO_3$ is part of the family of ABO_3 perovskites, and is considered promising for next generation devices, such as for power electronics, due to its band gap of about 4.5 eV. A detailed knowledge of phonon modes in $LaInO_3$ is important as they determine a number of material properties, such as the mechanical and elastic properties, thermal transport and carrier dynamics, phonon-assisted optical excitations, and many more. However, little is known about the vibrational properties of this material. In this study, we investigate the lattice dynamics by polarization- and angle-resolved Raman spectroscopy and density functional theory (DFT). We experimentally observe all but one of the Raman active modes and compare them to our simulated values from DFT. Furthermore, we present the DFT-derived phonon dispersion relation along the high symmetry directions in reciprocal space and depict the oscillation patterns for selected phonons at the Γ point. Finally, we determine the relative Raman tensor elements of the observed modes from the angular dependence of their corresponding scattering efficiencies (cf. Fig. 2).

5:30pm PCSI-SuA-19 Non-Trivial Electronic States in the $EuO/KTaO_3$ Interface Revealed by Quantum Oscillations in High Magnetic Fields, *K. Rubi*, Los Alamos National Laboratory; *M. Dumen*, *S. Chakraverty*, Institute of Nano Science and Technology, India; *S. Zeng*, *A. Ariando*, National University of Singapore; *M. Chan*, *N. Harrison*, Los Alamos National Laboratory

The coexistence of electric-field controlled superconductivity and spin-orbit interaction in two-dimensional electron gas (2DEG) based on complex oxides (e.g., $SrTiO_3$ and $KTaO_3$) hold great promise for advancement in spintronics and quantum computing. However, a comprehensive understanding of the electronic bands that give rise to the multifunctional character of these 2DEGs remains elusive. To address this, we recently

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investigated quantum oscillations in the magnetoresistance of the KTaO_3 -2DEG in high magnetic fields (60 T).

KTaO_3 is a $5d$ transition metal oxide, exhibiting a lighter effective mass of electrons and a stronger spin-orbit interaction at its conducting surface/interface than its counterpart SrTiO_3 [1-2]. A high-mobility spin-polarized 2DEG with the superconducting feature is discovered at the EuO/KTaO_3 interface [3]. In this talk, I will present novel insights into the electronic states of the EuO/KTaO_3 interface investigated through Shubnikov-de Haas (SdH) oscillations (Fig.1a). Remarkably, we observed a progressive increase in cyclotron mass and oscillation frequency with the magnetic field (Fig. 1b and c), indicating the presence of non-trivial electronic bands [4]. Besides providing experimental evidence for topological-like electronic states in KTaO_3 -2DEG, these findings shed light on the recent predictions of

[1] K. Rubi *et al.*, *npj Quantum Materials* **5**, 1 (2020); King, P. D. C. *et al.* *Phys. Rev. Lett.* **108**, 117602 (2012).

[2] K. Rubi *et al.*, *Phy. Rev. Research* **3**, 033234 (2021).

[3] H. Zhang *et al.*, *Phys. Rev. Lett.* **121**, 116803 (2018); Liu *et al.*, *Science* **371**, 716 (2021).

[4] K. Rubi *et al.*, *arXiv:2307.04854* (2023).

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Room Ballroom South - Session PCSI-SuE

Probing Exotic Order Parameters with Photoemission Spectroscopy

Moderator: Chris Leighton, University of Minnesota

7:30pm PCSI-SuE-1 Searching for the Excitonic Insulator State in Quantum Materials, *Edoardo Baldini*, The University of Texas at Austin **INVITED**

The excitonic insulator is an electronically driven phase of matter that emerges upon the spontaneous formation and Bose condensation of excitons. Detecting this exotic order in candidate materials is a subject of paramount importance, as the size of the excitonic gap in the band structure establishes the potential of this collective state for superfluid energy transport. However, the identification of this phase in real solids is hindered by the coexistence of a structural order parameter with the same symmetry as the excitonic order. Only a few materials are currently believed to host a dominant excitonic phase, Ta_2NiSe_5 being the most promising. In this talk, I will describe how advanced protocols based on time- and angle-resolved photoemission spectroscopy can shed light on primary order parameter of a candidate excitonic insulator [1]. Finally, I will discuss the opportunities offered by the development of novel momentum microscopy tools to extend these studies to the realm of two-dimensional material flakes that may host similar physics.

[1] E. Baldini et al., Proc. Natl. Acad. Sci. 120, e2221688120 (2023)

8:10pm PCSI-SuE-9 Comparative Study on Non-Linear and Linear Least Square Analyses Applied to X-Ray Induced Auger Electron Spectroscopy Transitions, *A. Gagliardi*, CNRS, ILV, France; *N. Fairley*, Casa Software Ltd, UK; *Solene Bechu*, CNRS, ILV, France

With the exception of the modified Auger parameter, X-ray induced Auger electron (X-AES) transitions aren't exploited to their full potential. Indeed, they can provide as much information (oxidation degree, chemical environment, atomic composition) as the classic photopeaks used in XPS, but their shapes' complexities limit their decompositions.

We offer here to explore the decomposition of Ga $L_{3/2}M_{4,5}M_{4,5}$ and In $M_{4,5}N_{4,5}N_{4,5}$ X-AES lines by comparing two approaches: the non-linear [1] and the linear [2] least square analyses. By combining non-linear and linear fitting procedures, PCA, and vectorial method [3], those two analyses have been implemented on the materials $\text{Cu}(\text{In}_x\text{Ga}_{1-x})\text{Se}_2$ and InSb , to unveil their surface oxidation when exposed to different atmospheres. The growth of oxide phases (Ga_2O_3 and In_2O_3 , determined by PCA, vectorial method and by comparison with reference spectra) was monitored on the X-AES lines with non-linear and linear approaches, showing a very good coherence between both, as illustrated in Fig 1 for the In $M_{4,5}N_{4,5}N_{4,5}$ X-AES transition of InSb . We will provide keys to perform non-linear and linear least squares analysis on X-AES lines, to explore new approaches for chemical determination.

[1] J.J. Moré, Numer. Anal. **630**, 105 (1978).

[2] G.H. Golub and C. Reinsch, Linear Algebr. **420**, 403 (1971).

[3] S. Béchu et al., Appl. Surf. Sci. **447**, 528 (2018).

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8:15pm PCSI-SuE-10 Probing Electrons and Light in Nanomaterials Using the Photoelectric Effect, *Taisuke Ohta*, *A. Boehm*, *S. Gennaro*, *C. Doiron*, *A. Kim*, *K. Thuermer*, *J. Sugar*, *C. Spataru*, Sandia National Laboratories; *J. Fonseca Vega*, *J. Robinson*, Naval Research Laboratory; *T. Beechem*, Purdue University; *M. Sinclair*, *I. Brener*, *R. Sarma*, Sandia National Laboratories

The photoelectric effect is sensitive to both the occupied electronic density of states and the electromagnetic field distribution. Thus, capturing the energy, yield, and spatial origin of photoelectrons from the sample enables us to examine local electronic properties and light-matter interactions concurrently. In this talk, we will describe two case studies using photoelectron emission microscopy (PEEM), revealing the spatial variations of Schottky barrier height between WS_2 and Au, and the local electromagnetic near-field profiles of Si metasurfaces. We will discuss the impact of crystallographic facets of Au grains as well as how the attractive interaction of Au with WS_2 can modify the crystallographic alignment among WS_2 layers. For near-field imaging, we will demonstrate the sensitivity of photoemission yield to the light absorptivity in visible to near infrared range, and evaluate the field profiles around Si meta atoms at the sub-photon wavelength scale on and off resonance excitation. Altogether we will discuss the potential of photoelectron imaging to examine the

intertwined light-matter coupled phenomena abundant in two-dimensional and quantum materials.

The work was supported by Sandia's LDRD program and in part by the US Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering). The work performed at the U.S. Naval Research Laboratory (NRL) was supported through Base Programs funded by the Office of Naval Research and through the NeuroPipe ARAP funded by the Office of the Secretary of Defense. Samples were fabricated, in part, at the Center for Integrated Nanotechnologies, an Office of Science User Facility operated for the US Department of Energy, Office of Science. Sandia National Laboratories is a multimission laboratory managed and operated by National Technology and Engineering Solutions of Sandia, LLC., a wholly owned subsidiary of Honeywell International, Inc., for the U.S. Department of Energy's National Nuclear Security Administration under contract DE-NA0003525. This paper describes objective technical results and analysis. Any subjective views or opinions that might be expressed in the paper do not necessarily represent the views of the U.S. Department of Energy or the United States Government.

8:20pm PCSI-SuE-11 Layer-by-Layer Engineering and Deciphering of Topological Orders in Magnetic Topological Insulators, *W. Lee*, University of Chicago; *S. Fernandez-Mulligan*, Yale University; *H. Tan*, Weizmann Institute of Science, Israel; *C. Yan*, University of Chicago; *Y. Guan*, *S. Lee*, *R. Mei*, *C. Liu*, Pennsylvania State University; *B. Yan*, Weizmann Institute of Science, Israel; *Z. Mao*, Pennsylvania State University; *Shuolong Yang*, University of Chicago **INVITED**

The advent of intrinsic magnetic topological insulators enables us to envisage various low-dimensional topological orders, such as the quantum anomalous Hall insulators and the axion insulators, at realistic cryogenic temperatures. These materials are represented by MnBi_2Te_3 and its derived superlattices $\text{MnBi}_{2n}\text{Te}_{3n+1}$. However, it has been controversial whether these materials exhibit the key ingredient for magnetic topological phases: an energy gap due to the time-reversal symmetry breaking. Moreover, the construction of high-quality magnetic topological insulators at the ultrathin limit has met significant challenges. In this talk, I will present a new technique, layer-encoded frequency-domain photoemission spectroscopy, which allows us to decipher the layer origins of various electronic states. By encoding layer indices with intralayer phonon frequencies, we measure the strengths of coupling with layer-specific phonons. This experiment reveals that the topological surface states on antiferromagnetic MnBi_4Te_7 are partially relocated to the nonmagnetic layers, reconciling the mystery of vanishing broken-symmetry gaps [1]. Moreover, I will present our recent progress on the "carpet-growth" of Bi_2Te_3 ultrathin films and $\text{MnBi}_2\text{Te}_4/\text{Bi}_2\text{Te}_3$ heterostructures using molecular beam epitaxy. These thin films extend coherently across a millimeter spatial scale without disruptions by substrate step edges. Angle-resolved photoemission spectroscopy studies yield unprecedentedly sharp electronic structures in agreement with first-principles calculations layer-by-layer, and suggest opportunities to realize the quantum spin Hall effect and quantum anomalous Hall effect at near-ambient temperatures [2].

[1] W. Lee et al., Nature Physics **19**, 950-955 (2023).

[2] W. Lee et al., Submitted (2023).

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Room Ballroom South - Session PCSI-MoM1

Semiconductor Heterostructures (Growth, Nanostructures & Interfaces) I

Moderator: Kirstin Alberi, National Renewable Energy Laboratory

8:30am PCSI-MoM1-1 Mechanisms and Applications for Remote Epitaxy of Heusler Compounds, Jason Kawasaki, University of Wisconsin - Madison

INVITED

Remote epitaxy on monolayer graphene is promising for synthesis of highly lattice mismatched materials, exfoliation of free-standing membranes, and re-use of expensive substrates. However, due to contaminants at the transferred graphene/substrate interface, other mechanisms such as pinhole-seeded lateral epitaxy often dominate rather than the intrinsic growth via remote interactions [1]. I will describe our understanding of the synthesis science of remote epitaxy, focusing on III-V semiconductors and Heusler compounds [1,2]. I will also show how exfoliated free-standing membranes of rare earth Heusler compounds can be used to tune (flexo)magnetism and novel superconductivity [3,4]

[1] S. Manzo, et. al., Nature Commun., 13, 4014 (2022). <https://doi.org/10.1038/s41467-022-31610-y>

[2] D. Du et. al., Nano Lett. 22, 21, 8647 (2022). <https://doi.org/10.1021/acs.nanolett.2c03187>

[3] D. Du, et. al., Nature Commun., 12, 2494 (2021). <https://doi.org/10.1038/s41467-021-22784-y>

[4] D. Du, et. al. APL, 122, 170501 (2023). <https://doi.org/10.1063/5.0146553>

9:10am PCSI-MoM1-9 UPGRADED: High-Mobility Two-Dimensional Electron Gas with Quantized States in Polar-Discontinuity Doped $\text{LaInO}_3/\text{BaSnO}_3$ Heterostructure Grown by Molecular Beam Epitaxy, G. Hoffmann, Paul-Drude-Institut für Festkörperelektronik Leibniz-Institut im Forschungsverbund Berlin, Germany; A. Hartl, Paul Scherrer Institut, Switzerland; M. Zupancic, Leibniz-Institut für Kristallzüchtung, Germany; A. Riaz, University College London, UK; V. Strocov, Paul Scherrer Institut, Switzerland; M. Albrecht, Leibniz-Institut für Kristallzüchtung, Germany; A. Regoutz, University College London, UK; Oliver Bierwagen, Paul-Drude-Institut für Festkörperelektronik Leibniz-Institut im Forschungsverbund Berlin, Germany

Transistor applications of semiconducting oxides require, both high room-temperature electron mobilities (μ_{RT}) and high charge carrier densities (CCDs), ideally realized with a two-dimensional electron gas (2DEG). So far, prototype oxide 2DEG systems have either high μ_{RT} but limited CCD such as modulation-doped $(\text{Al}_x\text{Ga}_{1-x})_2\text{O}_3/\text{Ga}_2\text{O}_3$, or a high CCD but low μ_{RT} such as the polar-discontinuity doped $\text{LaAlO}_3/\text{SrTiO}_3$ interface. Interfacing the more suitable, wide-bandgap, nonpolar semiconductor BaSnO_3 (BSO), having high bulk μ_{RT} (up to $320 \text{ cm}^2/\text{Vs}$), with polar LaInO_3 (LIO) is predicted to create and confine a 2DEG with CCD up to $2 \times 10^{14} \text{ cm}^{-2}$ for the SnO_2/LaO interface termination.

We demonstrate the adsorption-controlled growth of the LIO[1] on BSO heterostructure by molecular beam epitaxy using a shutter sequence to control the SnO_2/LaO interface termination. The films were analyzed by reflection high-energy electron diffraction (RHEED) [Figs. 1(a) and 1(b)], x-ray diffraction, atomic force microscopy (AFM) [Fig. 1(c)]. The interface structure is investigated by cross-sectional transmission-electron microscopy. The formation of the quantized 2DEG at their interface is confirmed by capacitance-voltage (CV) [Fig. 1(d)] and angular-resolved photo-electron spectroscopy (ARPES) [Fig. 1(e)]. Van der Pauw-Hall measurements confirm $\text{CCD} > 10^{13} \text{ cm}^{-2}$ and $\mu_{\text{RT}} > 100 \text{ cm}^2/\text{Vs}$.

[1] G. Hoffmann et al., Phys. Rev. Mater. 7, 084606 (2023).

9:30am PCSI-MoM1-13 Enabling Direct-Write Fabrication of Low Dimensional Micro- and Nanostructures on Supported and Suspended Substrates, Irma Kuljanishvili, Saint Louis University

Low-dimensional nanomaterials, such as one-dimensional (1D) or two-dimensional (2D) systems, assembled in vertical or lateral arrangements, often lead to enhanced properties and new functionalities. Nanotubes, nanowires (NWs), and 2D layered structures (graphene and graphene like materials) are emerging as key building blocks for the next generation devices and emerging technologies. Practical implementation of such nanomaterials necessitates their successful incorporation with well-established processes for fabricating electronic and/or mechanical devices.

While preparing layered architectures usually involves multi-step fabrication processes but relies on mask-assisted fabrication techniques. Here, we present a methodology for the controlled and selective preparation of nanostructures such as 1D NWs on 2D materials-substrates in various controlled geometric assemblies by employing direct write patterning (DWP) of custom ink precursors on supported or suspended architectures for subsequent chemical vapor deposition (CVD) synthesis. Our two-step fabrication approach enables simple and flexible routes to produce various architectures in a precisely controlled fashion. Location-specific materials synthesis provides access to as-grown interfaces and rapid testing of materials' quality, crystallinity, chemical composition, etc.

9:35am PCSI-MoM1-14 Silicon (111) - Aluminum (111) - Amorphous Alumina: Asymmetric Quantum Well and Band Alignment, Hanran Jin, University of Texas at Austin, China; A. Demkov, University of Texas at Austin

Thin single crystal films of Al can be epitaxially grown on Si(111) [1]. When taken out of ultra-high vacuum, Al oxidizes and forms a quantum well structure Si-Al- Al_2O_3 . Theoretical calculations can shed light on finding the band alignment across the hetero-structure and its electronic and optical properties. However, one needs to build a structural model. The structure of Al on Si(111) has been experimentally determined via the electron microscopy by McSkimming et al. [1]. The Al layer is found to be (111) oriented, with a linear ratio of 3 Si to 4 Al atoms along the interface. Using this information, we construct a periodic model of Si-Al interface and perform density functional theory (DFT) calculations to find the relaxed structure, layer-projected density of states, and planar averaged potential across the interface. From there, we find the band alignment using the valence band offset from the bulk averaged potential. When aluminum metal is exposed to air, a several nanometers thick layer of alumina rapidly forms [2]. This oxide layer is amorphous. We simulate the structure of amorphous alumina using ab-initio molecular dynamics with the melt-and-quench technique [3]. This amorphous alumina can then be directly combined with the previous system to form an asymmetrical quantum well, where the well states and band alignment can be directly obtained from the DFT calculations.

[1] McSkimming, B. M., Alexander, A., Samuels, M. H., Arey, B., Arslan, I., & Richardson, C. J. (2016). Metamorphic growth of relaxed single crystalline aluminum on silicon (111). J. Vac. Sci. & Technol. A 35, 021401 (2017).

[2] Jurgens, L. P., Sloof, W. G., Tichelaar, F. D., & Mittemeijer, E. J. Growth Kinetics and mechanisms of aluminum-oxide films formed by thermal oxidation of aluminum. Journal of Applied Physics, 92, 1649 (2002).

[3] Gutiérrez, G., and Johansson, B. Molecular dynamics study of structural properties of amorphous Al_2O_3 . Physical Review B 65, 104202(2002).

9:40am PCSI-MoM1-15 Silicene Ribbons: Synthesis, Electronic and Geometric Structure at the Atomic Scale, A. Costine, University of Virginia; Z. Gai, Orak Ridge National Laboratory; Petra Reinke, University of Virginia

Silicene is a 2D material which poses significant challenges in synthesis and device integration but offers unique electronic properties which are not yet fully understood. Challenges remain in silicene synthesis on non-metallic substrates and the control of buckling. The degree of in-plane buckling is tied to the emergence of a Dirac point and intriguing quantum phases have been predicted.[1-4] In our previous work we discovered a new pathway to silicene synthesis using h-MoSi₂ (0001) surfaces as templates,[5] and we present here the formation of silicene nanoribbons on the same surface, and measure their geometric and electronic structure with STM and STS at 77 K. Our work promises a new pathway to create silicene layers and nanoribbons with molecular beam epitaxy directly on silicon wafers.

The h-MoSi₂ (0001) crystallites are synthesized on Si(100) by deposition of a thin Mo film with electron beam evaporation, and subsequent annealing to form well-defined crystallites with 15 - 120 nm in diameter. Nanoribbons cover the entire surface for all "flat top" silicide crystallites with (0001) surfaces. Several surface reconstructions present intermediate superstructures, and are precursors for the nanoribbons. The STM images in Fig. 2 illustrate the geometric structure of the nanoribbons which are 1.8 nm in width and separated by a distinct groove of roughly a single atomic row in width. Models of the nanoribbon's geometric structure will be discussed. The atoms located at the ribbon edge appear brighter which is likely due to density of states (DOS) modulation by edge states. Characteristic defects are seen in the ribbons and express identical STM. The average DOS (dI/dV) measured with STS is indicative of a Dirac type electronic signature with a V-type dip around E_F , and the position dependent DOS across the ribbons allows to identify the Dirac type regions.

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We will discuss all aspects of electronic and geometric structure of the nanoribbons including electronic confinement, and localized Dirac type signatures. In addition, point defects, and long range deformations induced by strain are addressed and inform our structural models.

[1] C. Grazianetti et al.. ACS Nano **11**, 3376-3382 (2017). [2] C.-C. Liu et al.. Physical Review Letters **107**, 076802 (2011). [3] A. Molle et al. Chemical Society Reviews **47**, 6370-6387 (2018). [4] H. J. W. Zandvliet. Nano Today **9**, 691-694 (2014). [5] C. Volders et al. Nano Letters **17**, 299-307 (2017).

9:45am **PCSI-MoM1-16 Spontaneous Growth of Silver on Si(001) Tuned by Substrate Temperature**, *Xiaohang Huang, K. Huang*, Guangdong Technion - Israel Institute of Technology, China

The development of semiconductor technology features miniaturization of device that approaches the physical limit, i. e., nanostructure of a few atoms and/or molecules. It has been aimed that one could use nanostructures to fabricate functional devices at will. For the system of silver on Si(001), there is a rich collection of nanostructures as revealed in our recent works [1, 2], serving as the candidates to test this aim.

In this work, we examined the spontaneous growth of silver on Si(001) held at 109 to 298 K by scanning tunneling microscopy. As shown in Figure 1, the dimension of the formed silver depends on the substrate temperature during the deposition. That is, silver is formed predominantly as zero-dimensional (0D) objects at less than 120 K, one-dimensional (1D) objects at 120-200 K, and two-dimensional (2D) objects at 200-270 K. These observations are linked to the anisotropic migration of the key intermediate of silver tetramer, as supported by density functional simulations; the barriers are calculated as 0.24 eV along Si-rows, and 0.68 eV across Si-rows [2]. A schematic of the growth dynamics is sketched in Figure 2.

[1] K. Huang, X. Huang, J. Nogami, Phys. Chem. Chem. Phys. **7**, 23 (2021).

[2] X. Huang, A. Hoffman, K. Huang, J. Phys. Chem. C. **18**, 126 (2022)

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Room Ballroom South - Session PCSI-MoM2

2D Materials and Graphene I

Moderator: Kunal Mukherjee, Stanford University

11:00am **PCSI-MoM2-31 Interplay of Valley Polarized Dark Trion and Dark Exciton-Polaron in Monolayer WSe₂**, *Xiao-Xiao Zhang*, University of Florida

INVITED

The interactions between charges and excitons involve complex many-body interactions at high densities. The exciton-polaron model has been adopted to understand the Fermi sea screening of charged excitons in monolayer transition metal dichalcogenides (TMD). The results provide good agreement with absorption measurements, which are dominated by dilute bright exciton responses. The Fermi-polaron model treats the quasiparticle responses of a single mobile impurity in a surrounding Fermi sea. In comparison, the exciton density in monolayer TMD can be tuned by laser fluence and be comparable to or exceed the charge density, where the analogy to a single mobile impurity no longer applies. The modification to Fermi sea screening at high exciton densities, however, is still not well understood. Apart from the bright excitons previously studied in reflection contrast measurements, different spin and momentum dark exciton species have been established, which are also expected to have many-body interactions with charges. The coupling between these different species of exciton-polarons has not yet been experimentally investigated.

Here we investigate the Fermi sea dressing of spin-forbidden dark excitons in monolayer WSe₂. With a Zeeman field, the valley-polarized dark excitons show distinct p-doping dependence in photoluminescence when the carriers reach a critical density (see Fig. 1). This density can be interpreted as the onset of strongly modified Fermi sea interactions and shifts with increasing exciton density. Through valley-selective excitation and dynamics measurements, we also infer an intervalley coupling between the dark

trions and exciton-polarons mediated by the many-body interactions. Our results reveal the evolution of Fermi sea screening with increasing exciton density and the impacts of polaron-polaron interactions, which lay the foundation for understanding electronic correlations and many-body interactions in 2D systems.

11:40am **PCSI-MoM2-39 Evidence of Single Photon Emitters from 1L WSe₂ under Electrostatically Induced Strain**, *Frances Camille Wu, S. Wu, B. Fang, X. Li, J. Inconvia, E. Yu*, The University of Texas at Austin

Strain engineering is a powerful tool that strongly influences electronic structure and exciton states in 2D transition metal dichalcogenides (2D TMDs). Among 2D TMDs, monolayer WSe₂ has gained attention as a host for quantum emitters due to its lowest lying dark exciton state that hybridizes with mid-gap defect states under tensile strain, giving rise to bright single photon emitters. The creation of a hybridized state consisting of dark excitons and mid-gap defect states results in the radiative recombination of dark excitons that is otherwise forbidden due to spin and momentum conservation. At cryogenic temperatures, the energetic alignment and coupling of the two abovementioned states results in localized defect emission which possess significant characteristics of single photon emitters. Strain-tunable devices are crucial for investigating the nature of TMD-based single photon emitters which can be beneficial for quantum information processing and secure communications.

In this study, we demonstrate strain modulation of monolayer WSe₂ suspended over a hole-patterned substrate via electrostatic deflection and characterize the resulting photoluminescence. This approach enables the creation of strain-tunable WSe₂ devices that can be operated at cryogenic temperatures, wherein strain fields are generated by applying a bias voltage to the suspended monolayer WSe₂ membrane. We observe a significant monolayer deflection of ~50 nm at 15V gate bias and a ~20 meV redshift of dark exciton peak as the applied is increased to 20V, corresponding to a 0.2% increase in tensile strain of WSe₂. Sharp localized emitters, typically associated with single photon emitters, were observed at 4K which showed less dependence on strain as the applied bias increased. Thus, we attribute these localized emitters to the presence of localized defects which are only weakly influenced by strained lattice environment. We also observe luminescence lifetimes of ~3 ns and saturation of luminescence intensity with incident power, both of which are also characteristic of single photon emitters. These realizations are critical for understanding the origin of single photon emitters based on strained monolayer WSe₂.

11:45am **PCSI-MoM2-40 Comprehensive Study of Interface Chemistry and Electrical Property of Metal Contacts on TMDs**, *S. Kim, Joy Roy, X. Wang, R. Wallace*, University of Texas at Dallas

Transition metal dichalcogenides (TMDs) have been introduced due to their exceptional electronic, optical mechanical, and magnetic properties, even in atomically thin thickness, for advanced electronic, optoelectronic, and spintronic devices [1]. However, the limitations in tuning the Schottky barrier height with metal contacts, based on work function greatly hinder efficient carrier injection and electronic performance of TMD-based devices [2]. This study examines contact interfaces and their relationship to electrical contact characteristics. The research encompasses interface chemistry, band alignment, and electronic contact properties of Ni, Ag, Bi, Co and Sn.

An ultrahigh vacuum (UHV) cluster system was employed to investigate the contact properties where in-situ X-ray Photoelectron Spectroscopy (XPS) showed the contact bonding features of metal/TMD interfaces. Ni and Co contacts exhibited stronger bonds with TMD surfaces, resembling covalent-like interfaces, with notable interface reaction products resulting from annealing. However, Bi and Sn showed no robust chemical bonding features under the XPS analysis and van der Waals contact interface was formed due to a weak interaction between metal and TMD. The subsequent ex-situ atomic force microscopy (AFM) measurement supported these contact interface properties. The subsequent electrical characterization using XPS and scanning tunneling microscopy (STM) suggests that the roles of surface defect impact the metal contacts as well. In conclusion, comprehensive research and investigation of metal materials and their contact interface properties with TMDs have shed light on the potential and advantages of metal contact studies.

This work was supported in part by NEWLIMITS, a center in nCORE, a Semiconductor Research Corporation (SRC) program sponsored by NIST through award number 70NANB17H041, and by the National Science

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11:50am **PCSI-MoM2-41 Transport Anisotropy in One-dimensional Graphene Superlattice in the High Kronig-Penney Potential Limit**, *Tianlin Li, H. Chen, K. Wang, Y. Hao, L. Zhang*, University of Nebraska - Lincoln; *X. Watanabe, T. Taniguchi*, National Institute for Materials Science, Japan; *X. Hong*, University of Nebraska - Lincoln

One-dimensional (1D) graphene superlattice (GSL) has drawn considerable research interest as it is promising for realizing the electron lensing effect [1]. Despite the intensive theoretical studies, 1D GSL has only been realized experimentally via periodic dielectric gates in previous studies [2, 3], which yields a moderate Kronig-Penney (KP) potential profile that is not viable to achieve electron supercollimation.

In this work, we demonstrate 1D GSL in the high KP potential limit exploiting nanoscale domains patterning in a ferroelectric bottom gate [4]. We work with 50 nm (001) $\text{PbZr}_{0.2}\text{Ti}_{0.8}\text{O}_3$ (PZT) films deposited on 10 nm $\text{La}_{0.67}\text{Sr}_{0.33}\text{MnO}_3$ buffered SrTiO_3 substrates. Monolayer graphene field-effect transistors with top h-BN global gates are fabricated on PZT prepatterned with periodic polarization up (P_{up}) and down (P_{down}) stripe domains, with the period L varying from 200 to 300 nm (Fig. 1a). The polarization shifts the Fermi level of graphene, leading to a KP potential V_0 of about 0.9 eV at 2 K. We fabricate 1D GSL samples in two configurations, with current along the SL vector \hat{s} and perpendicular to \hat{s} . For the former samples, additional Dirac points (DP) emerge in the sheet resistance (R_{xx}) vs. top-gated induced electron doping δn (Fig. 1b), from which emanates multiple Landau fan branches in the magnetic field (Fig. 1c). This feature is absent in the latter configuration (R_{yy}), which can be attributed to the SL modulated band crossings along \hat{s} . The carrier density between consecutive DP positions (Δn_{DP}) scales with the SL period as $\Delta n_{\text{DP}} \propto L^\theta$, with $\theta = -1.18 \pm 0.06$ (Fig. 1d), which closely resembles the inversely proportional relation predicted for the high KP potential limit. Figure 1e shows the simulated 1D GSL band structure for our ferroelectric doping scheme, with dimensionless KP potential $u = V_0 L / \hbar v_F = 90\pi$, which reveals a highly flattened band that can potentially host electron lensing effect.

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11:55am **PCSI-MoM2-42 Terahertz Emission Spectroscopy Revealing Nanoscale Vectorial Photocurrents in Symmetry-Broken Optoelectronic Metasurfaces**, *J. Pettine, P. Padmanabhan*, Los Alamos National Laboratory; *L. Gingras, R. Holzwarth*, Menlo Systems, Germany; *R. Prasankumar, A. Taylor, S. Lin, Hou-Tong Chen*, Los Alamos National Laboratory

Terahertz (THz) emission spectroscopy has emerged in the past several decades as a versatile method for directly tracking the ultrafast evolution of physical properties, quasiparticle distributions, and order parameters within bulk materials and nanoscale interfaces. Ultrafast optically-induced THz radiation is widely utilized to analyze nonlinear polarization, magnetization, and various transient free charge currents, where the underlying broken symmetries (surface and bulk) enable THz emission by defining a system directionality in space and/or time [1]. The broken spatial or temporal symmetries responsible for second-order nonlinearities and bias-free photocurrent generation in materials are typically either intrinsic to the lattice and thus constrained to specific light-matter interaction geometries, or otherwise dependent upon applied static fields that are difficult to texture on small length scales.

In this talk we show that asymmetric gold nanoantennas on graphene exhibit strong light-driven directional responses [2]. The local photocurrent directionality is determined by the orientation of individual nanoantennas, which can be patterned into an arbitrary direction profile for global spatially-varying and optically-controlled photocurrents. We use THz emission spectroscopy, complementary to the direct photocurrent readout,

to validate such a concept. Our experimental results clearly demonstrate that these vectorial optoelectronic metasurfaces serve as efficient and versatile sources of ultrafast THz radiation, including broadband THz vector beams. Electrostatic gating and multiphysics modeling reveal a local photothermoelectric driving mechanism and elucidate previously unexplored dynamics occurring at the intersection of femtosecond excitation and nanoscale localization.

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12:00pm **PCSI-MoM2-43 Excitons, Electrons, and Holes in Monolayer Semiconductors: Insights from Spectroscopy in (Really) High Magnetic Fields**, *Scott Crooker*, National High Magnetic Field Lab

Historically, magnetic fields have played an essential role in revealing the properties of semiconductors, and the many-body physics that can emerge when they are doped with mobile carriers. However, for atomically-thin ‘transition metal dichalcogenide’ (TMD) semiconductors such as MoS_2 and WSe_2 , the relevant field scale is substantial (of order 100 tesla!) due to heavy carrier masses, huge exciton binding energies, and typically large Fermi energies. Fortunately, modern pulsed magnets can achieve this scale. This talk will discuss a few recent optical studies that probe the physics of -- and many-body correlations between -- excitons, electrons, and holes in TMD monolayers. These experiments used dual-gated TMD monolayers, assembled via van der Waals stacking directly atop single-mode optical fibers to enable polarized absorption spectroscopy at low temperatures in 60-100T fields.

*In charge-neutral monolayers, spectroscopy up to $\sim 90\text{T}$ reveals the diamagnetic shifts of the neutral exciton’s $1s$ ground state *and* its excited $2s$, $3s, \dots ns$ Rydberg states, revealing exciton masses, radii, binding energies, dielectric properties, and free-particle bandgaps – essential ingredients for the rational design of optoelectronic van der Waals structures. [1]

*In hole-doped monolayer WSe_2 , high-field spectroscopy of both neutral and charged exciton transitions revealed the (often-hypothesized) spontaneous valley polarization of mobile holes, due to exchange interactions, occurring at $\sim 40\text{T}$. [2]

*In electron-doped WSe_2 monolayers, the ordering of the conduction bands in the K and K' valleys allows studies of not only neutral excitons ($X0$) and charged excitons (X^- trions) at low carrier density, but also many-body states that can emerge at higher doping. We investigate the so-called X'^- resonance that emerges at high electron density, known since 2013 but never understood. The data suggest that X'^- is, in fact, a six-particle “hexicton” state that arises when the photoexcited electron-hole pair couples simultaneously to two Fermi seas having quantum-mechanically distinguishable spin/valley quantum numbers. This state also appears in WS_2 and may appear in MoS_2 , and appears in MoSe_2 at the B-exciton resonance [3,4]

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PCSI

Room Ballroom South - Session PCSI-MoA1

Materials for Catalysis, Energy Storage, and Energy Harvesting

Moderator: Edward Yu, The University of Texas at Austin

2:00pm PCSI-MoA1-1 Interface Control of III-Nitride Semiconductors: From High Efficiency Artificial Photosynthesis to Ferroelectric Switching, *Zetian Mi*, University of Michigan, Ann Arbor INVITED

In this talk, I will discuss the recent advances of nanoscale III-nitride semiconductors and their applications in artificial photosynthesis and ferroelectric devices. Artificial photosynthesis, the chemical transformation of sunlight, CO₂, and H₂O into clean chemicals and fuels, has been extensively studied but faces fundamental challenges of efficiency, stability, and selectivity. Recent studies of III-nitride semiconductors, e.g., GaN, InN, and their alloys, have shown that their surfaces can be transformed to be oxynitride during harsh photocatalysis conditions, leading to significantly improved efficiency and stability. With the integration of various co-catalysts, we have demonstrated high efficiency, long-term stable solar water splitting and hydrogen production. The recent advances of converting CO₂ to liquid fuels, reduction of N₂ to ammonia, and methane oxidation to methanol will also be discussed.

Another recent exciting development is the discovery of ferroelectricity in III-nitride semiconductors. The incorporation of rare-earth elements such as scandium (Sc) can transform conventional III-nitride semiconductors to be ferroelectric. I will present recent advances of ferroelectric Sc-III-nitride heterostructures and nanostructures, including epitaxy, properties, and emerging device applications. Molecular beam epitaxy and properties of ScAlN and ScGaN with a wide range of Sc compositions will be discussed. The realization of ultrathin ferroelectric nitride heterostructures and the underlying physics and interface properties will be discussed, together with their applications in quantum photonics and electronics.

2:40pm PCSI-MoA1-9 UPGRADED: Wafer-Scale Si-Based Metal-Insulator-Semiconductor Photoanodes for Water Oxidation Fabricated Using Thin Film Reactions and Electrodeposition, *Shang-Hsuan Wu, S. Lee, Y. Choi, E. Yu*, The University of Texas at Austin

The environmentally friendly generation of hydrogen (H₂) is anticipated to have a pivotal role in shifting from fossil-based to greener and more sustainable energy systems. Photoelectrochemical (PEC) water splitting is a promising technology for converting solar energy into clean and storable chemical energy and providing carbon-free production of hydrogen for other key applications, e.g., ammonia production. In PEC cells, semiconductors play a key role in absorbing photons from the light source to create mobile charge carriers. Si-based photoelectrodes have drawn much attention due to their moderate bandgap, high charge mobility, long carrier diffusion length, cost-effectiveness, and scalability in manufacturing. To improve the stability of Si-based PEC cells in operation, metal-insulator-semiconductor (MIS) structures have been widely employed. In MIS photoelectrodes, the insulator thickness plays a key role in such MIS photoelectrodes since it influences both efficiency and long-term stability. Photo-generated charges are typically extracted from the semiconductor to the metal catalyst via tunneling through the insulator, mandating the use of extremely thin insulators. However, optimal stability generally motivates the use of thicker insulators.

In this work, we employ a simple and highly scalable method to fabricate high-performance, extremely stable Si-based MIS photoanodes and demonstrate its application to the fabrication of wafer-scale photoanodes. Localized conduction paths formed via an Al/SiO₂ thin-film reaction enable low-resistance charge extraction even through thick insulating layers, and this approach has been shown in our previous work to yield photoanodes with excellent stability. In addition, we demonstrate a two-step Ni/NiFe electrodeposition process to create efficient OER catalysts. The Ni/NiFe catalyst allows for a high Schottky barrier between Si and Ni, lowering the photoanode onset potential, while the NiFe surface layer improves catalytic performance. An unassisted solar-driven water splitting system integrated with wafer-scale photoanode and monocrystalline Si solar cells is demonstrated under both AM 1.5G sunlight simulator and outdoor illumination, with solar-to-hydrogen efficiency of 6.9% achieved with a full-wafer photoanode and minimal optimization.

3:00pm PCSI-MoA1-13 UPGRADED: Field-Assisted Oxidation of a Fe Single Nanoparticle, Nanoscale Observations by Operando Atom Probe, *Sten V Lambeets*, Pacific Northwest National Laboratory; *N. Cardwell, I. Onyango*, Washington State University; *T. Visart de Bocarmé*, Université Libre de Bruxelles, Belgium; *J. McEwen*, Washington State University; *D. Perea*, Pacific Northwest National Laboratory

Mechanisms governing surface chemical reactions involved in heterogeneous catalysis fundamentally depends on the synergistic interactions between the reactants and the different surface structures present at the surface. Recently, special attention has been raised regarding the influence of intense electric fields on these mechanisms [1]. An increasing number of analytical surface science techniques are achieving their conversion to their respective in-situ/operando version to study surface reactions at the "applied" conditions. Amongst them, Atom Probe Microscopy (APM) techniques are particularly interesting for their inherent use of intense electric fields and their capability to image matter at the nanoscale. In this work, we will present a nanoscale study of the field-assisted oxidation of a single Fe nanoparticle using Field Ion Microscopy (FIM) and Operando Atom Probe (OAP).

APM techniques are capable of imaging the apex of sharp needles, mimicking model nanoparticles, with nanometric lateral resolution. FIM is used to image apices with atomic resolution and to identify the crystal orientation. The resulting FIM image corresponds to a stereographical projection of the apex and allows the identification of the crystal orientation. OAP relies on the thermally assisted field evaporation of positively charged ions from a needle shaped specimen [2]. Once the FIM characterization is complete the sample is maintained at 300K with an applied electric field of ~20V/nm, before starting OAP analysis and introducing 1.1×10⁻⁷mbar of pure O₂. As soon as the O₂ is introduced, Fe₂O⁺ ion species formation are observed starting from open facets structures, such as Fe{244} and {112}, towards the central Fe(011) and {024} (Fig.1). OAP results allow us to reconstruct the full movie of the surface oxidation in real-time and show how intense electric fields (>10V/nm) play a central role in surface chemistry.

PCSI

Room Ballroom South - Session PCSI-MoA2

Topological Materials & Interfaces I

Moderator: Jun Sung Kim, Pohang University of Science and Technology (POSTECH), Republic of Korea

4:30pm PCSI-MoA2-31 Crystalline Materials with Anisotropic Conduction Polarities, *Joshua Goldberger*, The Ohio State University INVITED

It is conventionally thought that a single material will exhibit a single kind of conduction polarity, either n-type or p-type, uniformly along all directions of the crystal. Then, in all modern electronic devices, functionality is achieved by integrating together these p-type or n-type materials together. Here we will describe our recent work in the synthesis, properties, and applications of metals and semiconducting materials that exhibit either n-type or p-type conduction behavior depending on the crystallographic direction, a phenomenon we refer to as "goniopolarity". We will establish the origin of this exotic behavior and the band structure design principles for identifying new goniopolar materials.^[1] This has led to a large expansion in the number of compounds that we have experimentally demonstrated to exhibit this effect, such as NaSn₂As₂, NaSnAs, WSi₂ and PdSe₂.^[2-5] Finally, we will show that the unique charge separation in goniopolar materials can overcome limitations of energy-harvesting technologies including thermoelectrics and photocatalysis.

5:10pm PCSI-MoA2-39 Weyl Semimetals and the Interface: Surface State Transport Probed via Weak Antilocalization in Ultrathin TaAs Films, *Ian Leahy, A. Rice, C. Jiang, G. Paul, K. Alberi, J. Nelson*, National Renewable Energy Laboratory

Topological semimetals hold promise for their use in low-powered electronics and spintronic devices [1-4] but these applications await targeted growth on conventional semiconducting substrates and the exploration of their properties in the ultrathin limit. Weak antilocalization (WAL) has been used extensively in the study of surface states in topological insulators and shows promise for the study of surface states in Weyl semimetals (WSMs). WAL is a quantum interference effect that results in an increase in a system's conductivity owing to the suppression of back-scattering from self-intersecting carrier paths. This quantum interference requires carriers maintain phase coherence over multiple scattering events.

The length over which carriers maintain coherence is defined as the decoherence length. In an applied field, the WAL is destroyed when the magnetic length approaches the decoherence length, offering a natural insight into the localizing disorder length scales. Here we report on insights from WAL into the surface state and interface properties of the recently synthesized, single-crystal-like ultrathin films of Weyl semimetal TaAs(001) grown on GaAs(001) substrates [5-7].

Figure 1 shows atomic force microscopy and magnetoconductance for representative TaAs ultrathin films. AFM on a 10 nm thick TaAs film on GaAs reveals oriented, rod-like growth along $\bar{1}10$ direction. At low temperatures, the magnetoconductance exhibits clear signatures of WAL. Intriguingly, we find that the number of apparent WAL conduction channels depends on the orientation of the applied current relative to the film topography as well as the number of GaAs/TaAs interfaces. We hypothesize that this unique anisotropic WAL stems from a topological and trivial state with different decoherence lengths localized at each interface.

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5:15pm **PCSI-MoA2-40 Topological Hall Effect in Dirac Semimetal**, *Saurav Islam*, E. Steinebronn, Pennsylvania State University; *B. Neupane*, University of North Texas; *K. Yang*, Pennsylvania State University; *Y. Wang*, University of North Texas; *C. Liu*, Pennsylvania State University; *S. Ghosh*, University of Minnesota; *K. Mkhoyan*, University of Minho, Portugal; *J. Chamorro*, T. McQueen, Johns Hopkins University; *N. Samarth*, Pennsylvania State University

Magnetic skyrmions are chiral spin textures whose non-trivial real space topology is often created by an interfacial anisotropic Dzyaloshinskii-Moriya exchange interaction (DMI) that originates from spin-orbit coupling and broken inversion symmetry [1]. They have been observed in a wide variety of bulk single crystals such as MnSi [2] and thin films such as $\text{Fe}_{1-x}\text{Co}_x\text{Si}$ [3]. More recently, magnetic skyrmions have been probed at ferromagnet/topological insulator interfaces [4] and in magnetic Weyl semimetals [5]. This motivates similar explorations of skyrmion formation in Dirac semimetals (DSMs). We investigate the formation of skyrmions at the interface of a canonical DSM (Cd_3As_2) and a ferromagnetic semiconductor ($\text{In}_{1-x}\text{Mn}_x\text{As}$) with perpendicular magnetic anisotropy. Our calculations indicate nonzero spin susceptibility in such heterostructures due to Rashba spin-orbit coupling from broken inversion symmetry, implying the DM interaction necessary for skyrmions. To experimentally test this idea, we grew $\text{Cd}_3\text{As}_2/\text{In}_{1-x}\text{Mn}_x\text{As}$ bilayers (Fig. 1a) and mapped out the behavior of the Hall effect as a function of temperature, magnetic field, and gate voltage in electrostatically top gated devices. Below $T = 6$ K, we observe an emergent gate-tunable topological Hall effect (THE) indicated by an excess Hall resistance (Fig. 1b). This signature is most pronounced at the charge neutrality point, suggesting the formation of a Dirac-electron mediated chiral spin texture at the DSM/ferromagnet interface. Our study provides a new platform to study the interplay between the topological states in DSMs and the chiral spin textures associated with the THE. Supported by the NSF Graduate Research Fellowship Program (Grant No. DGE1255832).

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5:20pm **PCSI-MoA2-41 Helical Dislocations in 2D Materials and the Connection to Transport in Topological Insulators**, *T. Rakib*, *M. Choi*, *E. Ertekin*, University of Illinois at Urbana-Champaign; *P. Pochet*, Université Grenoble-Alpes, France; *Harley Johnson*, University of Illinois at Urbana-Champaign

Layered two-dimensional materials host a variety of crystalline defects, including dislocations either in-plane or out-of-plane with respect to the 2D layered structure. Recently, twisted multilayer 2D material structures have been of interest due to the presence of flat bands and other emergent properties associated with moiré superlattices.[1] Periodic regions of crystalline commensurability making up these superlattices are now understood to be separated by interlayer dislocations, with Burgers vectors and line directions in the plane of the 2D material, and having either edge or screw character.[2] Using density functional theory and quantum Monte Carlo-fitted total energy tight-binding calculations, we show that out-of-plane relaxation of the structures makes possible unique helical dislocations in bilayer graphene, and that the presence of these helical dislocation lines coincides precisely with the so-called magic-angle condition at which unconventional superconductivity is observed.[3] We then illustrate a different dislocation structure, with line direction oriented out-of-plane, but which also has a helical structure. Such a screw dislocation, which adopts a double-helix dislocation core configuration in bilayer structures, is expected to create conditions for exotic transport properties in certain classes of layered topological insulator materials. We present initial results demonstrating this possibility in BiTe and BiSe compounds. In these examples, we present relaxed dislocation core structures computed using first-principles methods, and show that the observed configurations match both experimental observations and the theoretical conditions that are expected to lead to quantum conduction in these otherwise topologically insulating materials.

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5:25pm **PCSI-MoA2-42 Layer-dependent Optical Conductivity of MBE-grown ZrTe_2** , *E. Houser*, *Frank Peiris*, Kenyon College; *A. Richardella*, *M. Stanley*, *N. Samarth*, Pennsylvania State University

Besides providing an interesting platform to interrogate fundamental physics questions, two-dimensional transition metal dichalcogenides (TMDCs) are well suited to advance the development of optoelectronic technologies. In this work, we investigated the growth and the optical properties of ZrTe_2 , a candidate topological Dirac semimetal, grown using molecular beam epitaxy. During the growth of 12 unit cells (u.c.) of ZrTe_2 on a sapphire substrate, we obtained in-situ spectroscopic ellipsometry after the deposition of each u.c. Additionally, we obtained temperature dependent ellipsometry data on the sample between 20 °C and 350 °C. After the deposition of the ZrTe_2 layers, a Te capping layer was deposited in order to protect the TMDC film. Post-growth X-ray reflectivity measurements indicated that the total thickness of ZrTe_2 and the thickness of Te to be 5.95 nm and 19 nm, respectively.

A standard inversion technique was used to model the ellipsometry spectra by specifying a three layer model (i.e., sapphire substrate, ZrTe_2 layer and the Te capping layer) to fit the final ellipsometry spectra. The thicknesses obtained from X-ray reflectivity allowed us to obtain the precise dielectric function of the final ZrTe_2 layer (i.e., 12 u.c.), which was converted to the optical conductivity. Subsequently, we fit the remaining ellipsometry spectra obtained for 11 u.c. through 1 u.c. ZrTe_2 layers. Clearly, the optical conductivity shows a noticeable change with the thickness of the ZrTe_2 layers, where the real part increases with the thickness of ZrTe_2 , as shown in Fig. 1. The layer-dependent conductivity was further analyzed by incorporating a Drude oscillator to account for free electrons, and two Kramers-Kronig-consistent oscillators to represent the band-to-band transitions. Interestingly, we find that the Drude contribution reduces as the thickness of ZrTe_2 gets smaller, suggesting that its metallic character diminishes as the thickness reduces.

Monday Afternoon, January 15, 2024

5:30pm **PCSI-MoA2-43 Surface Dependent Doping Efficiency in Te: Cd₃As₂ Thin Films**, *Anthony Rice, I. Leahy, K. Alberi*, National Renewable Energy Laboratory

Cd₃As₂ is a prototypical Dirac semi-metal, a class of materials with gapless topologically protected electronic states. In this system, these topological electronic states are close to the intrinsic Fermi level and are well isolated from non-trivial bands. Additionally, this system is air stable and compatible with molecular beam epitaxy, including lattice matching to III-Sb and II-Te layers, and similar elements to conventional semiconductors. These materials could play a role in a large number of applications, including transistors, spintronics, photodetectors, and thermoelectrics. To do this, however, significant progress must be made on achieving tunability of these materials. In particular, routes to altering its typical n-type carrier concentration must be developed.

Previous attempts to dope Cd₃As₂(112) with group VI elements, including Te and Se, were successful, allowing for increases of n_{3d} from $5e17\text{ cm}^{-3}$ up to slightly over $3e18\text{ cm}^{-3}$ [1]. Concentration vs mobility relationships appeared similar to doping in convention semiconductors, with mobility decreasing with increased ionized impurities. Attempts to increase doping beyond this level by using increased group VI fluxes resulted in lower measured Hall concentrations and even larger decreases in mobility, a sign that compensating defects are forming. When similar doping is attempted on the (001) surface, however, doping beyond $1e19\text{ cm}^{-3}$ is possible. Furthermore, an order of magnitude smaller Te fluxes are required to achieve similar doping levels. Finally, smaller unintentionally doped carrier concentrations are achievable on this surface. This work highlights the role of surface kinetics in defect incorporation in topological semi-metals.

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5:35pm **PCSI-MoA2-44 Investigating the Structural and Electronic Properties of FeSn on LaAlO₃ (111) Grown by Molecular Beam Epitaxy**, *T. Erickson, Sneha Upadhyay, A. Shrestha, A. Abbas, H. Hall, D. Ingram, S. Kaya, A. Smith*, Ohio University

The Kagome lattice of 3-d transition metals produces exciting electronic excitations through correlated topological phases due to a mixture of the unique geometry and spin-orbit coupling [1,2]. Research into these materials continues to provide insight into electronic properties of Dirac-bands in transition metal materials. FeSn with its Fe₃Sn Kagome layers separated by honeycomb Sn-2 layers provides ample opportunity to study these phenomena. Recent scanning tunneling microscopy (STM) studies into FeSn confirm the expected antiferromagnetic spin order consistent with bulk measurements, demonstrating a ferromagnetic alignment within Kagome layers and antiferromagnetic coupling between separate layers [3,4]. Currently, these findings are for bulk FeSn samples transported to and cleaved in ultra-high vacuum chambers. Here, we perform direct in-situ UHV-STM analysis of FeSn samples as-grown by molecular beam epitaxy. We grew our FeSn on LaAlO₃ substrates at temperatures ranging from 450 to 550 °C and Fe:Sn flux ratios of 0.64:1 to 1.52:1. LaAlO₃ and FeSn have a lattice match with a difference of only 1%. We also compare the results samples by means of RHEED, XRD, RBS, and AFM. In all cases, smooth streaky RHEED patterns are observed, and from the streak spacing we calculate the in-plane lattice constants which are then complemented by the lattice constants calculated from the XRD spectra. For the case of the 1.52:1 flux ratio, using RHEED we find an $a = 5.240 \pm 0.017\text{ \AA}$ as compared to the expected value for the FeSn lattice parameter $a = 5.297\text{ \AA}$ [2], and using XRD we find $c = 4.436 \pm 0.042\text{ \AA}$ as compared to the expected c for FeSn = 4.481 \AA [2]. In this presentation, we will discuss the lattice parameters as functions of the incident flux ratios as well as the phases and phase purity of the resultant samples. Additionally, AFM and RBS results are used to describe the smoothness and stoichiometry respectively.

This work is supported by the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering under Award No. DE-FG02-06ER46317.

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[2] Yin, JX., Zhang, S.S., Li, H. et al. Nature 562, 91–95 (2018).

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[4] Lee, SH., Kim, Y., Cho, B. et al. Commun Phys 5, 235 (2022).

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5:40pm **PCSI-MoA2-45 Ultra-quantum Limit Magnetotransport in the Topological Pentatellurides**, *Johanna Palmstrom, C. Ribeiro, C. Mizzi, L. Winter, S. Thomas*, Los Alamos National Laboratory; *J. Liu, L. Jauregui*, University of California Irvine; *J. Mutch, Q. Jiang, J. Ayres-Sims, J. Chu*, University of Washington; *E. Peterson, J. Zhu*, Los Alamos National Laboratory

With low carrier concentrations and high mobilities, the pentatelluride material family (HfTe₅ and ZrTe₅) typically have a quantum limit of a few Tesla, making them an ideal platform to study ultra-quantum limit phenomena and magnetic field-induced effects in a three dimensional (3D) Dirac fermion system. In the quantum limit, the magnetic field is strong enough to confine all the electrons to their lowest Landau levels, resulting in a quantized in-plane dispersion, lower effective dimensionality, and a system that is more unstable to electronic correlations. Previous experiments in the pentatellurides have revealed many exotic high-field phenomena including an interaction driven instability [1] and a field induced Lifshitz transition [2]. These materials sit right at the cusp of a strong-to-weak 3D topological phase transition, resulting in a band structure and electronic properties that are extremely sensitive to external tuning parameters such as magnetic field and strain [3]. Consequently, while these systems are promising for the extrinsic control of topological properties, there are many controversies surrounding their intrinsic behavior as the electronic properties depend on the sample growth and preparation conditions [4].

In this work we report on the ultra-quantum limit electronic properties and magnetic field-temperature phase diagram of flux grown, bulk HfTe₅ as revealed by magnetotransport measurements in pulsed magnetic fields up to 65 T (Fig. 1). These samples show a purely insulating resistance vs temperature behavior in zero field. We find a strong and non-monotonic angle dependence of the magnetoresistance and several high field features in the ultra-quantum limit. The interpretation of these features will be discussed during the presentation.

[1] L. Yanwen et al. Nat. Commun. 7, 12516 (2016)

[2] S. Galeski et al. Nat. Commun. 13, 7418 (2022)

[3] J. Mutch et al. Sci. Adv. 5, eaav9771 (2019)

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PCSI

Room Ballroom South - Session PCSI-MoE

Topological Materials & Interfaces II

Moderator: Joshua Goldberger, The Ohio State University

7:30pm PCSI-MoE-1 Large Magnetotransport Responses and Spintronic Functionalities of Topological van der Waals Ferromagnets, *Jun Sung Kim*, Pohang University of Science and Technology (POSTECH), Republic of Korea
INVITED

Topological van der Waals (vdW) ferromagnets has emerged as a promising material platform for investigating novel magnetotransport responses and spintronic functionalities. Their unique topological electronic structures in combination of magnetism, spin-orbit interaction, and orbital-driven topological band degeneracy gives rise to large magnetotransport responses and magnetic tunability. In addition, their unique vdW structure allows for the isolation of atomically thin layers as well as the creation of atomically sharp and clean interfaces in heterostructures. In this talk, I will discuss our recent demonstrations of these magnetotransport and spintronic properties on various topological vdW ferromagnets and their heterostructures, highlighting large anomalous Hall effect [1], large angular magnetoresistance [2], highly-tunable spin-valve operations [3], and highly efficient magnetic switching [4]. These findings demonstrate that topological vdW ferromagnets have great potential for realizing novel spin-dependent electronic functionalities, which may be suitable for all-vdW-material-based spintronic applications.

[1] K. Kim, et al. Nat. Mater. 17, 794 (2018)

[2] J. Seo et al. Nature 599, 576–581 (2021).

[3] K.-H. Min, et al. Nat. Mater. 21, 1144 (2022)

[4] G. S. Choi et al. submitted

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8:10pm PCSI-MoE-9 Tuning the Curie Temperature of a 2D Magnet/Topological Insulator Heterostructure to Above Room Temperature by Epitaxial Growth, *Wenyi Zhou, A. Bishop*, The Ohio State University; *X. Zhang*, Cornell University; *K. Robinson, I. Lyalin, Z. Li, R. Bailey-Crandell*, The Ohio State University; *T. Cham*, Cornell University; *S. Cheng*, The Ohio State University; *Y. Luo*, University of Southern California; *D. Ralph, D. Muller*, Cornell University; *R. Kawakami*, The Ohio State University

Heterostructures of two-dimensional (2D) van der Waals (vdW) magnets and topological insulators (TI) are of substantial interest as candidate materials for efficient spin-torque switching, quantum anomalous Hall effect, and chiral spin textures. However, since many of the vdW magnets have Curie temperatures below room temperature, we want to understand how materials can be modified to stabilize their magnetic ordering to higher temperatures. In this work, we utilize molecular beam epitaxy to systematically tune the Curie temperature (T_C) in thin film $\text{Fe}_3\text{GeTe}_2/\text{Bi}_2\text{Te}_3$ from bulk-like values (~ 220 K) to above room temperature by increasing the growth temperature from 300 °C to 375 °C (Figure 1). For samples grown at 375 °C, cross-sectional scanning transmission electron microscopy (STEM) reveals the spontaneous formation of different $\text{Fe}_m\text{Ge}_n\text{Te}_2$ compositions (e.g. $\text{Fe}_3\text{Ge}_2\text{Te}_2$ and $\text{Fe}_2\text{Ge}_6\text{Te}_2$) as well as intercalation in the vdW gaps, which are possible origins of the enhanced Curie temperature. This observation paves the way for developing various $\text{Fe}_m\text{Ge}_n\text{Te}_2/\text{TI}$ heterostructures with novel properties.

8:15pm PCSI-MoE-10 Kagome Antiferromagnetic Mn_3GaN grown on $\text{MgO}(001)$ using Molecular Beam Epitaxy, *A. Abbas, A. Smith, Ashok Shrestha, S. Upadhyay, T. Erickson*, Ohio University; *K. Sun*, University of Michigan; *D. Ingram*, Ohio University

Antiperovskite materials are intermetallic compounds with perovskite crystal structure (space group $\text{Pm}\bar{3}\text{m}$) but with anion and cation positions interchanged in the unit cell [1]. Similar to oxide-perovskite structure, antiperovskite materials have a variety of physical properties including antiferromagnetism, superconductivity and giant magnetoresistance [2]. There have been very few studies of antiperovskite structure Mn_3GaN in general although it was seen in molecular beam epitaxial growth as a second-phase precipitate when growing MnGaN [3]. Here we discuss the molecular beam epitaxial growth and surface study of Mn_3GaN . In our work, Mn_3GaN is deposited at 250 ± 10 °C onto magnesium oxide (001) substrates with a Mn: Ga: N flux ratio of 3:1:1. The sample surface is continuously monitored throughout the growth using reflection high energy electron diffraction. During the growth, the RHEED pattern was observed to

be highly streaky, indicating an atomically smooth surface. The calculated *in-plane* lattice constant based on RHEED is 3.89 ± 0.06 Å. This value is close to the theoretical lattice constant a of Mn_3GaN (3.898 Å) [3]. X-ray diffraction confirms the majority 002 peak, and the value calculated is 3.84 ± 0.06 Å which also agrees well with the theoretical value (3.898 Å) [3] and with the experimental reported c value (3.881 Å) [2]. Since we did not observe significant second-phase peaks, the phase purity of the sample is quite high. Furthermore, cross-sectional STEM was done to understand the interface and the surface of the film. The plan is to also present *in-situ* scanning tunneling microscopy results for the surfaces of these MBE-grown Mn_3GaN layers.

This work is supported by the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering under Award No. DE-FG02-06ER46317.

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8:20pm PCSI-MoE-11 Investigation of Smooth Epitaxial Growth of Mn_3Sn Films on C-Plane GaN Using Molecular Beam Epitaxy, *Sneha Upadhyay, H. Hall, C. D'Mello*, Ohio University; *J. Hernandez*, Universidad Autonoma de Puebla, Mexico; *T. Erickson*, Ohio University; *K. Sun*, The University of Michigan, Ann Arbor; *G. Coccoletzi*, Universidad Autonoma de Puebla, Mexico; *N. Takeuchi*, Universidad Nacional Autónoma de México; *A. Smith*, Ohio University

Recently, Chen *et al.* studied the all-antiferromagnetic tunnel junction consisting of $\text{Mn}_3\text{Sn} / \text{MgO} / \text{Mn}_3\text{Sn}$ (011 $\bar{1}$), where they observed a tunnel magnetoresistance (TMR) effect at a ratio of 2% at room temperature.¹ Furthermore, Bangar *et al.* reported the epitaxial growth of *c*-plane Mn_3Sn on the Al_2O_3 substrate using a Ru seed layer. They demonstrated a technique of engineering intrinsic spin Hall conductivity in Mn_3Sn by adjusting the Mn composition slightly for functional spintronic devices.² These works indicate great potential for kagome antiferromagnetic material, and it is essential to investigate the growth of Mn_3Sn on various substrates. In our previous work, we demonstrated the deposition of Mn_3Sn (0001) on Al_2O_3 (0001) at 524 ± 5 °C, which resulted in a 3D island growth. We observed dome-like structures, which may be related to the significant lattice mismatch with sapphire (19%).³ Subsequently, we began to explore new substrates, and recently, we tried the growth on the MBE-grown N-polar GaN (000 $\bar{1}$). The growth was monitored *in-situ* using reflection high energy electron diffraction and measured *ex-situ* using X-ray diffraction, Rutherford backscattering, and atomic force microscopy. The sample grew at 524 ± 5 °C for 71 mins, resulting in an epitaxially smooth growth of Mn_3Sn on GaN (000 $\bar{1}$). The *in-plane* lattice constants indicate a strain of -2.13 %, while the XRD indicates a 0001 orientation with a strain of -0.53% and an 11 $\bar{2}$ 0 orientation with a strain of + 2.73%. Furthermore, the effect of varying growth temperature and Mn: Sn flux ratio on film orientation and crystallinity will be discussed in detail. We are also planning to begin scanning tunneling microscope experiments.

The authors acknowledge support from the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering under Award No. DE-FG02-06ER46317. We acknowledge the financial support of the Nanoscale & Quantum Phenomena Institute.

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8:25pm PCSI-MoE-12 Symmetry Constraints on Topological Invariants, *Jing Zhang*, Imperial College London, UK

Classification of topologically trivial/non-trivial crystalline insulators are based on the homology of Berry connection on the Bloch (vector) bundle

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with the BZ as the base manifold. Specifically, the trivial phase correspond to zero Berry phase defined in terms of Wilson loop operator integrated along closed path in the BZ. (Eq.1). The transformation properties of eigenstate in the Berry connection are well defined for a given k but it is generally a function of k . This makes the analysis of symmetry properties of Berry phase difficult (path integrals in BZ do not form a representation). There are symmetry analysis such as symmetry indicator method but they lack the group theoretical justification like Wigner-Eckart theorem. In this contribution, it is shown that ϕ_B can be evaluated using the band representation basis (EBRs) under the tight binding model and Stokes theorem. (Eq.2). The transformation properties of these EBRs contains no explicit k dependence and RHS of Eq.2 form a representation of the space group (Symmetry operation takes the closed path to others in the BZ belonging to a closed set. These set of Berry phases along different paths within the set form the representation). As a closed path in the BZ is frequently not contained in the representation domain, full group method is used.

The BZ are 2-torus or 3-torus. It is not simply connected and one needs to consider inequivalent un-contractable closed path, as in homotopy analysis involving the fundamental group. The symmetry operation naturally permute closed path between such set. Symmetry analysis shows the ϕ_B is generally not forbidden by symmetry of layer/space group. However, presence of some symmetry (e.g. inversion) may leads to specific selection rules that forces the Berry phase to be zero.

For a set of physically connected bands with symmetry at high symmetry points identical to direct sum of the EBRs, they have the same transformation properties as the set of EBRs and may be represented as such with appropriate interactions. The same symmetry analysis then may be applied. For close path containing the Γ point, what forms a representation of the group is not necessarily restricted to the whole close path, but half of a close path given that Γ point is invariant. Graphene is used as an example to illustrate both the trivial (sp^2 bands) and non-trivial (p_z band with spin).

The general conclusions are that not all occupied EBRs are symmetry forbidden from having non-zero Berry phase and occurrence of trivial phase are the exceptions. The symmetry indicator method at identifying trivial phase may include non-trivial phases.

8:30pm **PCSI-MoE-13 UPGRADED: Epitaxial Kagome Thin Films as a Platform for Topological Flat Bands and Dirac Cones**, *S. Cheng, M. Nrisimhamurty*, Ohio State University; *T. Zhou*, University at Buffalo; *N. Bagues, W. Zhou, A. Bishop, I. Lyalin*, Ohio State University; *C. Jozwiak, A. Bostwick, E. Rotenberg*, Advanced Light Source, Lawrence Berkeley National Laboratory; *D. McComb*, Ohio State University; *I. Zutic*, University at Buffalo; **Roland Kawakami**, Ohio State University

Metals consisting of kagome lattices have interesting band structures consisting of topological flat bands and Dirac cones. Systems with flat bands are ideal for studying strongly correlated electronic states and related phenomena due to the smaller bandwidth W compared to the Coulomb repulsion U . Kagome metals such as CoSn have been recognized as promising candidates due to the proximity between the flat bands and the Fermi level. A key next step will be to realize epitaxial kagome thin films with flat bands to enable tuning of the flat bands across the Fermi level via electrostatic gating or strain. Here we report the band structures of epitaxial CoSn thin films grown directly on insulating substrates [1]. Flat bands are observed using synchrotron-based angle-resolved photoemission spectroscopy (ARPES). The band structure is consistent with density functional theory (DFT) calculations, and the transport properties are quantitatively explained by the band structure and semiclassical transport theory. We are also developing kagome metals that have the Dirac cones near the Fermi level, which are interesting for investigating the intrinsic anomalous Hall effect and to potentially realize the quantum anomalous Hall effect at elevated temperatures.

[1] Cheng *et al.*, Nano Letters, 23(15), 7107-7113 (2023).

PCSI

Room Ballroom South - Session PCSI-TuM1

Magnetic Materials (2D, Monolayers, & Heterostructures)

Moderator: Xiao-Xiao Zhang, University of Florida

8:30am PCSI-TuM1-1 Efficient Control of 2D Magnets, **Cheng Gong**, University of Maryland, College Park **INVITED**

The emergent two-dimensional (2D) layered magnets provide ideal platforms to enable the atomically thin magneto-optical and magnetoelectric devices. Though many have envisioned that 2D magnets should allow efficient control of magnetism by a variety of external stimuli, true breakthroughs are still lacking, with limited proof-of-concept demonstrations reported thus far. There appear to be fundamental obstacles for efficient control, e.g., through electrical and optical means. In this talk I will analyze the challenges and present our theoretical and experimental progress on efficient electrical and optical control of 2D magnets. Specifically, the results show that the voltage of a few volts can effectively change the magnetic anisotropy of 2D magnets and the laser shining of tens of $\mu\text{W}/\mu\text{m}^2$ can effectively affect the domain behaviors of 2D magnets. These efficient controls of 2D magnets potentially open up new avenues towards low-power spintronics and photonics.

9:10am PCSI-TuM1-9 Surface-Bulk Difference in van der Waals Magnets, **Liuyan Zhao**, University of Michigan, Ann Arbor **INVITED**

The successful isolation of monolayer to few-layer magnetic atomic crystals from van der Waals (vdW) magnets have opened a new pathway of researching two-dimensional (2D) magnetism [1,2,3]. Over the past half a decade, the vdW and 2D magnet library has been greatly expanded, and new magnetic phenomena have been discovered in the 2D limit. Yet, one key question has been brought up: what is the distinction amongst bulk, surface and 2D magnetism for a vdW magnet? This question is well motivated by the observations of 2D behaviors in 3D vdW magnets, as well as the contrasts between 2D layers and 3D bulk, for systems such as CrI_3 , CrSBr , NiPS_3 , etc.

In this talk, we will show the surface-bulk difference in two archetype vdW magnets, CrI_3 [4] and CrSBr [5]. In CrI_3 , it has been thought that the 3D bulk hosts the ferromagnetic (FM) state below $T_c = 61\text{K}$ whereas the 2D films realizes the layered antiferromagnetic (AFM) order below $T_N = 45\text{K}$. We will show from our optical magneto-Raman spectroscopy measurements that even in a 3D bulk CrI_3 , we capture clear signatures of layered AFM, in addition to the known bulk FM. We attribute the layered AFM signature here to the surface magnetism, which is the same as that of the 2D layers but distinct from that deep in the 3D bulk (Figure 1a). In CrSBr , it has the same layered AFM order in both 3D bulk and 2D layers, but surprisingly with a higher critical temperature in the 2D case. We will show with our nonlinear optical measurements that multiple characteristic temperature scales appear in the 3D bulk CrSBr , including a surface (T_{surface}) and a bulk (T_{bulk}) onset temperature between which the surface one is unexpectedly higher than the bulk one (Figure 1b). Our results on these two systems demonstrate that the surface of vdW magnets can well be distinct from their bulk.

[1] Cheng *et al* Nature 546, 265 (2017)

[2] Huang *et al* Nature 546, 270 (2017)

[3] Wang *et al* ACS Nano, 16, 6960 (2022)

[4] Li *et al* Phys. Rev. X, 10, 011075 (2020)

[5] Guo *et al* arXiv 2309.01047 (2023)

9:50am PCSI-TuM1-17 Surface Investigation of Hexagonal Non-Collinear $\text{D}_{019}\text{-Mn}_3\text{Ga}$ Thin Film on $\text{GaN}(0001)$ Substrate, **Ashok Shrestha**, A. Abbas, D. Ingram, A. Smith, Ohio University

In recent years, Mn_3Ga has garnered significant attention due to its exotic physical properties and potential applications in spintronic devices [1,2]. One of the most intriguing, yet less explored, phases is the hexagonal antiferromagnetic phase of Mn_3Ga (D_{019}), which exhibits anomalous Hall effect and topological Hall effect in distinct temperature ranges [2]. In this presentation, we will delve into the growth and surface studies of a thin film of $\text{D}_{019}\text{-Mn}_3\text{Ga}$ on a Ga polar- GaN (0001) substrate.

The experiments are carried out in an ultra-high vacuum chamber equipped with a molecular beam epitaxy system and a room-temperature scanning tunneling microscope. Initially, the GaN epilayer is deposited on a GaN (0001) substrate at 700°C under gallium-rich conditions, followed by the growth of $\text{D}_{019}\text{-Mn}_3\text{Ga}$ at 250°C under manganese-rich conditions. Reflection high-energy electron diffraction is used during growth to monitor

the sample, and the *in-plane* lattice constant is evaluated. Various *in-situ* techniques confirm that the grown sample exhibits epitaxial growth. Furthermore, scanning tunneling microscopy image shows the hexagonal atomic arrangements with an average *in-plane* atomic spacing of $5.37 \pm 0.05 \text{ \AA}$. However, the atomic spacing varies in the local region. The 1×1 surface structure of hexagonal $\text{D}_{019}\text{-Mn}_3\text{Ga}$ ($a = 5.40 \text{ \AA}$ [2]) is shown in Fig. 1. Moreover, multiple flat terraces and steps with height of 2.20 \AA are observed. The measured step height corresponds to the $c/2$ value of $\text{D}_{019}\text{-Mn}_3\text{Ga}$ ($c = 4.39 \text{ \AA}$ [2]). The *ex-situ*- X-ray diffraction clearly shows the Mn_3Ga 0002 peak, and the calculated *d*-spacing matched well with the step heights measured by scanning tunneling microscope. These measurements are consistent with the theoretically reported *c*-value of $\text{D}_{019}\text{-Mn}_3\text{Ga}$. The concentration of manganese and gallium in the sample is confirmed to be $3.2:1.0$ by Rutherford backscattering. Various *in-situ* and *ex-situ* measurements confirm the $\text{D}_{019}\text{-Mn}_3\text{Ga}$ growth. Further work is planned to investigate the non-collinear antiferromagnetism using spin polarized scanning tunneling microscope.

This work is supported by the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering under Award No. DE-FG02-06ER46317.

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[2] Z. H. Liu, Scientific Reports **7**, 515 (2017).

9:55am PCSI-TuM1-18 Enhancement of Microwave to Optical Spin-Based Quantum Transduction via a Magnon Mode, **Tharnier O. Puel**, Department of Physics and Astronomy, University of Iowa; A. T. Turflinger, S. P. Horvath, J. D. Thompson, Department of Electrical Engineering, Princeton University; M. E. Flatté, Department of Physics and Astronomy, University of Iowa, Department of Applied Physics, Eindhoven University of Technology, Eindhoven, The Netherlands

The highly localized 4f electrons of rare-earth-doped materials provide a simple atom-like level structure with a spin-photon interface, telecom-wavelength optical transitions, potential for long spin and optical coherence times, and the ability to realize high-density doping. Proposals for microwave to optical quantum transduction using rare-earth ions [PRL113,203601(2014)] rely on spin-flip transitions from microwaves that couple to optical inter-4f transitions. An example is the Er^{3+} ion's transition $|J=15/2\rangle$ to $|J=13/2\rangle$ at telecom wavelength. The oscillator strengths (g_b) of the microwave excitations of the Er^{3+} are particularly weak leading to poor transduction efficiencies. We describe an approach to dramatically enhance the microwave coupling without diminishing the optical oscillator strength (g_a) for Er^{3+} ions. The microwave excitation is coupled to a magnon (g_m) of a magnetic material, e.g., yttrium iron garnet (YIG). The Er^{3+} ions are embedded in an insulator and live close to the interface with the magnet. The iron lattice of the YIG will strongly couple to the Er^{3+} . We predict that the microwave-magnon coupling allows higher transduction rates that dramatically exceeds the previous set up.

10:00am PCSI-TuM1-19 Magnetic Modulation and Large Magnetoresistance in Cr_5Te_8 , M. Vaninger, S. Kelley, University of Missouri; F. Ye, Oak Ridge National Laboratory; X. Zhang, Nanjing University, China; T. Heitmann, University of Missouri; A. Mazza, Los Alamos National Laboratory; Y. Hor, A. Sarikhan, Missouri S&T; G. Bian, Paul Miceli, University of Missouri

Because of the ability to manipulate their structure and properties, metallic 2D van der Waals materials that exhibit ferromagnetism (FM) are of considerable potential interest for spintronics applications. Cr_5Te_8 is such a system whose structure consists of layers of CrTe_2 having additional Cr intercalated between the layers. CrTe_2 itself is known to be a strong ferromagnet up to room temperature [1]. Cr_5Te_8 is FM below $T_{c1}=155\text{K}$ with perpendicular magnetic anisotropy and it exhibits a large (10%) negative magneto-resistance effect above T_{c1} over a narrow temperature range [2].

We have performed neutron diffraction measurements to explore the magnetic behavior in a temperature range above T_{c1} and as a function of applied magnetic field. A modulated antiferromagnetic phase is observed, which has a wavevector perpendicular to the van der Waals layers and a period that is triple the unit cell length. The modulated spin structure is canted with a significant component in the van der Waals layers. The modulation is robust with field applied *in-plane* but it is quickly destroyed with a field applied perpendicular to the layers. Our magnetic phase diagram shows that the transition from FM to the modulated phase at T_{c1} is strongly first-order with a true FM transition occurring at a higher temperature, $T_c=180\text{K}$. We show that the large magnetoresistance observed in transport arises from the *in-plane* components of the magnetic moments. Since the spin modulation is controlled at relatively low magnetic

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field and the intercalated Cr can be tuned, 2D systems such as these have potential for spintronic applications.

Support: NSF-DMR; the University of Missouri Research Reactor. Spallation Neutron Source at Oak Ridge National Lab is supported by the US Department of Energy.

[1] Room-temperature intrinsic ferromagnetism in epitaxial CrTe₂ ultrathin films X. Zhang *et al.*, *Nature Communications* **12**:2492 (2021) [<https://www.nature.com/articles/s41467-021-22777-x>]

[2] Self-Intercalation Tunable Interlayer Exchange Coupling in a Synthetic Van der Waals Antiferromagnet X. Zhang *et al.*, *Advanced Functional Materials* **2202977** (2022) [<https://onlinelibrary.wiley.com/doi/abs/10.1002/adfm.202202977>]

PCSI

Room Ballroom South - Session PCSI-TuM2

Organic and Hybrid Semiconductor Materials & Interfaces

Moderator: Wanyi Nie, Los Alamos National Laboratory

11:00am PCSI-TuM2-31 Development of Surface Chemistry on-Top of Organic Semiconductor Thin Films to Improve Optoelectronic Devices, *Jacob W. Ciszek*, Loyola University Chicago

Organic devices (OLED, OFETs, OSCs) are commonly configured such that a metal contact must be deposited on top of the organic semiconductor. These interfaces often suffer from numerous flaws including poor adhesion, metal penetration, poor surface wetting (discontinuous films), and significant charge injection barriers. We feel that the ability to selectively change the surface of the organic layer via chemistry would allow for improved interactions with the deposited metal and, in-turn, improve device performance. Vast literature shows monolayers (deposited on top of gold, silicon, etc.) can address many of these issues when the system is reversed (organic on metal), but the methodology for installing them is inapplicable to organic thin-films. This points to an obvious need for controllable surface chemistry on top of the organic semiconductor.

We have developed click-like "Diels-Alder chemistry that allows prototypical OFET films (tetracene, pentacene, rubrene) to be appended with a variety of small molecules to form an interfacial layer only ~1 nm thick. The reacted surface is highly tailorable with dozens of combinations explored for a variety of application. This basic approach is presented in detail. This work then summarizes many of the expected and unexpected deviations which occur due to the very unusual nature of the organic surface, namely its high anisotropy, recessed reaction loci, and weakly bound molecular units. Concluding the fundamental components, we highlight some of our recent work in generating analogous chemistry on OLED electron transport layers, specifically TPBi.

This work concludes by highlighting many interfacial components the chemistry aims to improve, focusing primarily on the morphological and mechanical structure of applied metal top contact. Specifically, we examine how the various functional groups improve surface wetting of the metal, film continuity, adhesion in flexible contacts, metal penetration, and sheet resistance.

11:05am PCSI-TuM2-32 Characterizing Nanopattern Formation of Polymer Thin Films on Silicon Substrates with Ion Beam Sputtering, *Jocelyn Zhang*, Boston University, Del Norte High School; *G. Pettis*, Oregon State University, Boston University; *B. Jiang*, Boston University, Turkey; *N. Baker*, Boston University; *E. Guney*, Sabanci University, Turkey; *G. Ince*, Sabanci University IICEC, Turkey; *K. Ludwig, Jr.*, Boston University

Surface nano-patterns formed by ion beam sputtering (IBS) have been reported by many research groups with most focus on semiconductor and metal materials [1]. However, limited study has been conducted on polymer nanopatterning introduced by IBS [2]. This study aims to understand polymer film wrinkling in relation to ion beams and humidity conditions, which could potentially introduce effective methods of tuning chemical and physical characteristics of polymer film surfaces. Poly(4-Vinylpyridine) (4-VP) and Polystyrene (PS) polymer thin films on silicon substrates were sputtered with Ar⁺ ions under ultra-high vacuum (UHV) and then placed in either humid or dry conditions. The results show that wrinkling patterns are formed in 4-VP film after sputtering at 2 keV and consecutive humidification, but no wrinkling patterns are observed in PS under the same condition. The wrinkling amplitude of 4-VP films increases over time under humidity. Areas of lower ion flux have less order and higher amplitude wrinkling in 4-VP. Increasing film thickness increases wrinkle

wavelength and decreases the order in the wrinkles. It is also found that the surface contact angle with water increased on both 4-VP and PS after ion beam sputtering, presumably due to increased roughness.

11:10am PCSI-TuM2-33 Functionalizing Organic Semiconductors with Dipole Monolayers, *Matthew Williams*, Loyola University Chicago

This work seeks to improve OFETs performance via the addition of a chemisorbed monolayer on top of the organic material to address various issues such as contact resistance and charge trapping. The monolayer is made possible via site-specific Diels-Alder chemistry which only reacts with the semiconductor. By installing specific functional groups to the surface, the aforementioned issues can be eliminated, resulting in improved charge injection and/or charge transport in organic material.

We focused on the addition of monolayers which contain significant electronic charge separation (or molecular dipole) within their structure. This feature can act as a potential step, shift the mean energy in gaps, and/or align energy levels between materials. We first utilize the chemistry to selectively append dipole-containing molecules to trap states at grain boundaries for polycrystalline OFETs in order to shift the mean energy within the grain boundary and improve device performance. Grain boundaries are unavoidable flaws inherent to the materials, and the ability to address trap states would be a powerful way to address these flaws post-fabrication. This performance improvement is exemplified in conductance measurements with two orders of magnitude increases, improved threshold voltages, and doubling of mobility.

Additionally, we can optimize the surface potential of thin films by systematically varying similar dipole-containing monolayers. A linear relationship between dipole strength and surface potential shift is seen, as predicted by the Helmholtz equation. The achieved potential adjustments are attributed to the monolayer and result in a high degree of tunability of surface potential. With the capability to shift potential up to ~800mV, reducing electron charge injection barriers is hypothetically feasible. The challenges arising in conductance measurements are discussed.

PCSI

Room Ballroom South - Session PCSI-TuE

Point Defects for Quantum Information Applications

Moderator: Roland Kawakami, The Ohio State University

7:00pm **PCSI-TuE-1 Rare Earth Doped Oxide Thin Films on Silicon for Chip Scale Quantum Emitters and Memories**, *Supratik Guha, D. Awschalom*, University of Chicago, Argonne National Laboratory; *C. Ji, G. Grant, S. Seth, I. Masiulionis*, University of Chicago; *A. Dibas, J. Zjang*, Argonne National Laboratory; *S. Chattaraj*, University of Chicago; *M. Singh*, University of Chicago, memQ; *J. Wen*, Argonne National Laboratory **INVITED**

Quantum memories are an enabling technology for long distance repeater based quantum communications via optical fibers. Embedded within a host dielectric, the Er ion, with its 1.5 μm 4f-4f optical transition and its expected long spin coherence times, presents a convenient solid-state spin-optical interface that is telecom wavelength compatible for such quantum memory applications. Furthermore, it is desirable that such memories be scalable and compatible with silicon electronics for large scale deployment. Consequently, we have been exploring the properties of Er doped (few to ~ 100 ppm) dielectric oxide thin films grown on silicon substrates through detailed microstructural, growth and optical studies. Those oxides are also judiciously chosen to have low nuclear spin noise in the host to foster long Er electron spin coherence for memory applications. In these studies, using Er doped TiO_2 , Y_2O_3 , and CeO_2 as epitaxial and polycrystalline thin film hosts, and careful correlations of electron microscopy and X-ray diffraction based microstructural studies with optical properties, we find that while extended defect densities do not appear to have a significant effect upon the inhomogeneous linewidths, the Er doping levels, proximity of surfaces, the substrate interface, and film thickness have strong effects upon the optical properties including spectral diffusion and optical lifetime besides inhomogeneous linewidth, all critical for memory applications. We will discuss these results and the models of interaction that arise from these results. For the case of epitaxial CeO_2 on Si(111) we measure a narrow homogeneous linewidth of 440 kHz with an optical coherence time of 0.72 μs at 3.6 K when studying the Z_1 - Y_1 optical transition near 1530 nm at ~ 3.5 K, along with an inhomogeneous linewidth of 10 GHz, an optical excited state lifetime of 3.5 ms. Using Er doped TiO_2 films on silicon grown via both molecular beam deposition as well as atomic layer deposition (where we had to develop mechanisms of ppm level doping of Er), we further show that such structures can be processed into good quality factor Si nanophotonic cavity devices and demonstrate a large Purcell enhancement (~ 300) of their optical lifetime leading to higher emission rates. These results indicate the significant promise of Er doped thin films as silicon compatible qubit devices for optical quantum memory and emitter applications. We will discuss these results with a focus on the materials science engineering aspects of this work.

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7:40pm **PCSI-TuE-9 Erbium sites in Silicon for Quantum Information Processing**, *Sven Rogge*, University of New South Wales, Australia **INVITED**

Rare-earth ions incorporated in several solid-state hosts were shown to exhibit low homogeneous broadening and long spin coherence at cryogenic temperatures making them a promising candidate for quantum applications, such as optical quantum memories, optical-microwave transductions, and quantum communication. However, long electron spin coherence has not been demonstrated in Si, a leading material platform for electronic and photonic applications. Here, we present the first demonstration of Er sites in semiconductor (Si) with a millisecond electron spin coherence time, optical homogeneous linewidths below 100 kHz, spin and optical inhomogeneous broadening approaching 100 kHz and 100 MHz, correspondingly. Er properties were measured using photoluminescence excitation spectroscopy within a nuclear spin-free silicon crystal ($< 0.01\%$ ^{29}Si) doped at 10^{16} cm^{-3} Er level. Er homogeneous linewidth and spin coherence were addressed using optical comb-based spectral hole burning and optically detected magnetic resonance. To enhance Er emission collection efficiency, samples were directly positioned on top of dedicatedly fabricated superconducting single photon detectors and resonantly excited using fiber optics. Measurements in naturally abundant Si revealed that the Er electron spin coupling to ^{29}Si nuclear spins significantly shortens Er spin coherence times. Long spin coherence time and narrow optical linewidth show that Er in ^{28}Si is an excellent candidate for future quantum information and communication applications.

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PCSI

Room Ballroom South - Session PCSI-WeM1

Ferroelectric & Neuromorphic Computing Materials

Moderator: Alec Talin, Sandia National Laboratories

8:30am PCSI-WeM1-1 Emergent Phenomena at Ferroelectric/van der Waals Heterointerfaces, *Xia Hong*, University of Nebraska - Lincoln INVITED

The heterointerfaces between ferroelectrics and two-dimensional (2D) van der Waals materials present a versatile platform for achieving novel interfacial coupling, nonvolatile field effect control, and nanoscale programmable functionalities. In this talk, I will discuss a range of emergent phenomena in ferroelectric/vdW heterostructures mediated by interfacial coupling of charge, lattice, and polar symmetry. By combining polarization doping with nanoscale domain patterning in a ferroelectric polymer P(VDF-TrFE) top-gate, we create directional conducting paths in an insulating 2D channel, which reveals highly anisotropic conductivity in monolayer (1L) to 4-layer 1T'-ReS₂ between the directions along and perpendicular to the Re-chain [1]. The interface-epitaxy between P(VDF-TrFE) and ReS₂ leads to large scale P(VDF-TrFE) thin films composed of highly ordered, close-packed, 10 and 35 nm wide crystalline nanowires [2]. We observe enhanced polar alignment, piezoelectricity, and Curie temperature in thin CuInP₂S₆ (CIPS) flakes prepared on ferroelectric oxide PbZr_{0.2}Ti_{0.8}O₃ (PZT), which can be attributed to the interfacial strain imposed by PZT [3]. An unconventional filtering effect of second harmonic generation signal is enabled by the polar coupling of 1L MoS₂ with either the polar domain or the chiral dipole rotation at the domain wall surface in PZT thin films or free-standing membranes [4,5]. Our study showcases the rich research opportunities offered by integrating ferroelectrics with 2D materials.

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9:10am PCSI-WeM1-9 Impact of High-Power Impulse Magnetron Sputtering Pulse Width on the Nucleation, Crystallization, Microstructure, and Ferroelectric Properties of Hafnium Oxide Thin Films, *Samantha Jaszewski*, Sandia National Laboratories

The impact of the high-power impulse magnetron sputtering (HiPIMS) pulse width on the crystallization, microstructure, and ferroelectric properties of undoped HfO₂ films is reported. HfO₂ films were sputtered from a Hf target in an Ar/O₂ atmosphere, varying the instantaneous power density by changing the HiPIMS pulse width with fixed time averaged power and pulse frequency. The pulse width is shown to affect the ion-to-neutral ratio in the depositing species with the shortest pulse durations leading to the highest ion fraction, as shown in Figure 1. *In-situ* X-ray diffraction measurements during crystallization demonstrate that the HiPIMS pulse width impacts nucleation and phase formation, with an intermediate pulse width of 110 μs stabilizing the ferroelectric phase over the widest temperature range. Although the pulse width impacts the grain size with the lowest pulse width resulting in the largest grain size (Figure 2), grain size does not strongly correlate with phase content or ferroelectric behavior in these films. These results suggest that precise control over the energetics of the depositing species may be beneficial for stabilizing the ferroelectric phase in this material.

9:15am PCSI-WeM1-10 Fabrication and Gamma Radiation Effects on Endurance of Ferroelectric Hafnium Zirconium Oxide Capacitors, *M. David Henry*, Sandia National Laboratories; *M. Lenox*, University of Virginia; *A. Hillsman*, North Carolina State University; *S. Jaszewski*, *G. Esteves*, Sandia National Laboratories, USA; *J. Jones*, North Carolina State University; *J. Ihlefeld*, University of Virginia

Ferroelectric hafnium zirconium oxide (HZO) is attracting significant interest in the semiconductor microelectronics industry with attributes including coercive voltages compatible with CMOS, retention of memory states after power down and reasonable polarizations achieved with films 8 to 15 nm thick. An immediate application of the HZO capacitors include non-volatile memory (NVM) with insertions in the back end of line (BEOL) fabrication. Although devices such as ferroelectric capacitors are most applicable for FeRAM integrations, subtle details in their fabrication including the electrodes used and thickness can have impact in the device performance metrics.

This work investigates electrode configurations, including W and TiN, ferroelectric thickness and anneals utilized in BEOL processes for effects on endurance and polarization. Insertion of thin linear dielectrics, 1 nm of alumina on the bottom electrode, is also investigated to determine properties impactful to FeRAM circuit design. To determine the stability of the film, device polarization and endurance was measured after 5 MRad of Co⁶⁰ gamma cell irradiation over differing voltage rails and cycling frequencies. This work extends the knowledge base of ferroelectric HZO with radiation effects for non volatile memory applications in CMOS.

9:20am PCSI-WeM1-11 Design of Memristive Devices Towards Neuromorphic Computing, *Aiping Chen*, Los Alamos National Laboratory INVITED

Current digital computing based on Von Neumann architecture suffers from several key bottleneck including von Neumann bottleneck, Moore's law, and the breakdown of Dennard scaling. Developing new computing platforms provide solutions towards Beyond Moore's computing. Recently, emergent devices such as memristive switching devices have been used to emulate some brain functions including synaptic behavior and neuronal behavior and therefore they have been proposed for developing low-power neuromorphic computing. Oxide-based memristive devices with excellent scalability have the potential to revolutionize not only the field of information storage but also neuromorphic computing.

In this talk, I will first discuss some basics of the brain, brain-inspired neuromorphic computing and artificial intelligence. In the second part of my talk, I will then discuss the roles of defects and interfaces on switching behavior in different types of memristive devices and their impacts on neuromorphic computing. Material systems have profound effects on switching behavior. For example, ferroelectric and non-ferroelectric systems show completely different switching behavior [1-2]. Defects also dominate the switching behavior. Figure 1 compared switching behavior in a variety of materials with different type of defects. Among different types of switching, filament-type switching and interface-type switching are two most distinct switching modes. I will focus on a specific interface-type switching we observed in Au/Nb:SrTiO₃ system [3]. It shows the switching is controlled by protons in the environment. We also explored the applications of such systems for neuromorphic computing applications [4].

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10:05am PCSI-WeM1-20 Neuromorphic Memristors with TiO₂ and a-IGZO Bilayer Structure, *Jae-Yun Lee*, College of Electrical and Computer Engineering, Chungbuk National University, South Korea; *H. Zhao*, *X. Wang*, *S. Shi*, College of Electrical and Computer Engineering, Chungbuk National University, South Korea, China; *B. Lee*, *S. Kim*, College of Electrical and Computer Engineering, Chungbuk National University, South Korea

In recent years, ReRAM devices have gained significant attention in neuromorphic applications and hardware-based artificial intelligence [1-2]. Specifically, the resistive memory devices exhibit ultrafast read and write speeds, high retention time [3], low voltage operation and low power consumption, emerging an attractive research target from the perspective of the modern low-cost portable devices [4].

Our proposed device performance and physical properties of the fabricated ReRAM devices were assessed at various annealing temperatures. The analysis from XPS results confirms that the device operation was mostly driven by the density of oxygen vacancies in the TiO₂ and a-IGZO bilayer structure. The optimal density of oxygen vacancies in the a-IGZO causes the drift of O²⁻ ions to and from the TiO₂ layer that induced a significant variation in the resistivity of the device, providing switching behavior.

10:10am PCSI-WeM1-21 Origin of Large Electro-Optic Response in Ferroelectrics, *Alex Demkov*, *I. Kim*, *T. Paoletta*, *S. Apte*, The University of Texas at Austin

Integrated silicon photonics experiences a revolution [1]. The key element of this technology is an optical modulator (OM) playing a role similar to that of a usual transistor. OMs based on a phase shifter using a linear electro-optic (EO) effect are an attractive option for building ultra-compact, fast and low power OMs [2]. Linear EO effect can be only observed in non-centrosymmetric materials, such as ferroelectrics, which started a search for ferroelectrics that can be integrated with Si and maintain a strong EO effect in thin films [3]. Ab initio calculations became an indispensable tool in this search [4].

We will discuss our recent progress in understanding the microscopic mechanism behind the EO response in three ferroelectrics successfully

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integrated on Si:BaTiO_3 (BTO), LiNbO_3 (LN) and $\text{Sr}_x\text{Ba}_{1-x}\text{Nb}_2\text{O}_6$ (SBN). There are three parts to the EO effect in a ferroelectric, they are ionic, piezo and electronic contributions [5,6]. In different materials, different components of the EO tensor are dominated by different contributions. This has implications for the device design, depending on the temperature and frequency range. For example, optical quantum computing occurs at cryotemperatures (when optical phonons are frozen out), and thus has rely on the electronic and piezo contributions. On the other hand, at high RF frequencies, only the ionic and electronic contributions survive. On the fundamental level, our results support the notion that P4mm BTO is a dynamic average of lower symmetry Cm structures (Fig. 1). We also discover that in SBN, surprisingly the major contribution to the EO effect comes from high frequency optical phonons (Fig. 2). And in LN, ferroelectricity and the EO response are essentially decoupled.

The work is supported by the AFOSR under Award No FA9550-18-1-0053.

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PCSI

Room Ballroom South - Session PCSI-WeM2

Spin Transport and Spintronics

Moderator: Ezekiel Johnston-Halperin, The Ohio State University

11:00am PCSI-WeM2-31 Direct Visualization of Electronic Transport in a Quantum Anomalous Hall Insulator, *Katja Nowack*, Cornell University INVITED

A quantum anomalous Hall (QAH) insulator is characterized by quantized Hall and vanishing longitudinal resistances at zero magnetic field that are protected against local perturbations and independent of sample details. This insensitivity makes the microscopic details of the local current distribution inaccessible to global transport measurements. Accordingly, the current distributions that give rise to the transport quantization are unknown. Here, I will discuss how we use magnetic imaging to directly visualize the transport current in the QAH regime [1]. As we tune through the QAH plateau by electrostatic gating, we clearly identify a regime in which the sample transports current primarily in the bulk rather than along the edges. Furthermore, we observe a local response of the equilibrium magnetization to electrostatic gating, whose spatial structure is strongly correlated with the observed current density. Combined, these measurements are consistent with the current flowing through incompressible regions whose spatial structure can change throughout the QAH regime.

At sufficiently high currents in the QAH regime and generally outside the QAH regime, we observe a weak response of the magnetization to the applied current. We show that this response can be explained by current-induced heating of the electrons. Effectively this allows us to image local dissipation in the QAH regime. As an example, I will show images of hot-spots localized in the corners of the electrical contacts through which the transport current enters our devices.

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11:40am PCSI-WeM2-39 Magneto-Optical Detection of the Orbital Hall Effect in Chromium, *Igor Lyalin, R. Kawakami*, The Ohio State University
The Hall effect was discovered by a PhD student Edwin Hall and his advisor Henry Rowland in 1879. Since then, the family of Hall effects has grown considerably. The anomalous Hall effect, integer and fractional quantum Hall effects, spin Hall effect, quantum anomalous Hall effect are fundamental physics phenomena of great importance. The orbital Hall effect (OHE) with giant orbital Hall conductivities has recently been theoretically predicted [1-4], however its direct observation is a challenge. Here, we report the magneto-optical detection of current-induced orbital accumulation at the surface of a light 3d transition metal, Cr. The orbital polarization is in-plane, transverse to the current direction, and scales linearly with current density, fully consistent with the orbital Hall effect. Comparing the thickness-dependent magneto-optical measurements with ab initio calculations, we estimate an orbital diffusion length in Cr of 6.6 ± 0.6 nm. Along with Choi et al. study of the OHE in Ti [5], our work [6] provides strong evidence for the OHE. The detection of the orbital Hall effect in light metals can have important consequences for future spintronics applications that could utilize orbital currents rather than spin currents.

11:45am PCSI-WeM2-40 Temperature Dependent Study of $\text{Na}_x\text{Si}_{136}$ Type II Si Clathrate Spin Dynamics, *Joseph Briggs, Y. Liu, S. Saiter, A. Faricy, C. Burns, C. Taylor, M. Singh, R. Collins, C. Koh*, Colorado School of Mines
We report the temperature dependence of relaxation time (T_1) and phase memory (T_M) of Na dopants in type II Si clathrate films utilizing electron paramagnetic resonance (EPR). There is a rich history directed at understanding defect properties in conventional diamond Si motivated by its dominant position in the microelectronics industry. Type II Si clathrates represent an alternative crystal structure to diamond Si. This cage-like inclusion compound is made up of a Si lattice with interstitial "guests" situated inside the cages. Our recent advances have allowed the synthesis of Na guest, type II Si clathrate films with low enough Na concentration for the Na to be considered a dopant and the spin dynamics of isolated Na donors to be investigated.[1]

EPR gives insight into the electron spin dynamics of the Na donors and their placement and interactions within the Si cages. The naturally occurring Na isotope, ^{23}Na , has nuclear spin 3/2 with the EPR spectrum exhibiting four hyperfine lines associated with the interaction of the electron and nuclear spins. Hyperfine features associated with Na atoms in neighboring cages, clustered Na, and interactions with ^{29}Si isotopes on the cage, are also observed.[2] Pulsed-EPR spectra exhibit clear spin echo signals with T_1 times in the hundreds of microseconds at temperatures near 7 K, and T_M times above a microsecond. The effects of various parameters (i.e. temperature, magnetic field center, Na concentration) on the relaxation time and phase memory are reported. The relaxation time exhibits thermally activated behavior from 6-14K suggesting an Orbach relaxation pathway. Strong similarities and important differences between the spin dynamics of Na in clathrate and P in diamond Si will be discussed along with Na's potential to function as a qubit in quantum applications. This work was supported by National Science Foundation award #2114569.

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11:50am PCSI-WeM2-41 Spin-orbit coupling in InGaAs random and digital alloy quantum wells, *Jason Dong*, University of California at Santa Barbara; *Y. Gul*, University College London, UK; *A. Engel, C. Dempsey*, University of California at Santa Barbara; *T. van Schijndel*, University of California Santa Barbara; *M. Pepper*, University College London, UK; *C. Palmström*, University of California at Santa Barbara
InGaAs two dimensional electron gases (2DEGs) have high spin-orbit coupling, making them potentially useful for spintronics [1] and topological quantum computing applications [2,3]. With increasing In concentration, InGaAs quantum wells will have lower effective masses, higher spin-orbit coupling, and higher g-factors than GaAs quantum wells [4]. Digital alloying, or growing the ternary as a superlattice, is an alternative to growing ternary III-V as a random alloy. However, the effect of digital alloying on the spin-orbit coupling in semiconductor quantum wells is not understood. Digital alloy quantum wells can potentially enhance the Rashba spin-orbit coupling by forming asymmetric interfaces with the barrier layers. Here, we use molecular beam epitaxy and magnetotransport to the role of random and digital alloying of the spin-orbit coupling of InGaAs quantum wells.

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We report the growth of high electron mobility $\text{In}_{0.81}\text{Ga}_{0.19}\text{As}$ quantum wells grown as a random and a digital alloy. From low temperature magnetotransport (2 K), the electron mobility of the random alloy quantum well is in excess of $450,000 \text{ cm}^2/\text{Vs}$ and the electron mobility of the digital alloy quantum well is in excess of $540,000 \text{ cm}^2/\text{Vs}$. The spin-orbit coupling of the quantum wells is extracted from fits to the weak localization in the magnetotransport data and will be presented. We will also discuss the role of interfaces on the differences in the spin-orbit coupling observed in the random and digital alloy quantum wells.

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11:55am **PCSI-WeM2-42 Screw Dislocations-Based Spin Valves**, *Finley Haines, E. Renteria, M. Debasu, F. Cavallo*, University of New Mexico

We fabricated and characterized a vertical spin valve (VSV) based on single-crystalline Si nanomembranes (NMs) engineered with 2D arrays of screw dislocations (SDs) throughout their thickness. The device includes a bottom soft ferromagnetic contact (NiFe), Si NMs, and a top hard ferromagnetic contact (Co). Based on previously reported theoretical calculations, we expect that the operation of the VSV relies on the coherent transport of spin-polarization through SDs [1]. The first step in the fabrication of the VSV is patterning a 220 nm-thick Si NM into a 2D array of pixels with lateral sizes in the range of 200-400 nm. At this stage of the process, the NM is bonded to a SiO_2 -coated bulk Si substrate. Si pixels are released in place by selective etching of the SiO_2 layer. An adhesive stamp removes the pixels from the original substrate and transfers them onto a second array of patterned pixels at a controlled twist angle, Ψ . The twisted NM pairs are then annealed at 1000-1200 °C in N_2 atmosphere to grow the SDs. Annealed NMs are finally transferred to a bulk substrate coated with NiFe. A dielectric barrier and a Co/Au top contact are fabricated using conventional top-down processes. The coercivity of the ferromagnetic films used for the contacts is extracted from measured magnetization curves on a Quantum Design Magnetic Properties Measurement System 3 (QD-MPMS3). Magneto-transport measurements characterize the resistance of the VSV at different magnitudes of magnetic induction (B). We observe a change in resistance at B corresponding to the measured coercivity of NiFe. The estimated magnetoresistance ratio, $MR(\%) = (R_{AP} - R_P) / R_P$ is -0.38 % at 300 K. No change in resistance was measured for VSVs based on Si NM that did not include SDs, suggesting that the line defects are responsible for the probed MR at 300K.

PCSI

Room Ballroom South - Session PCSI-WeA1

Characterization of Interfaces and Devices

Moderator: Paul M. Koenraad, Eindhoven University of Technology, Netherlands

2:00pm **PCSI-WeA1-1 Imaging the Properties of Atoms and Fields at the Picometer Scale inside Materials and Devices**, *David Muller*, Cornell University **INVITED**

Electron microscopes use electrons with wavelengths of a few picometers, and are potentially capable of imaging individual atoms in solids at a resolution ultimately set by the intrinsic size of an atom. Even with the rapid advances in aberration-corrector technology, both residual aberrations in the electron lenses and multiple scattering of the incident beam inside the sample, the best resolution possible was an order of magnitude worse than this limit. However, with recent advances in detector technology [1] and ptychographic algorithms to unscramble multiple scattering, the resolution of the electron microscope is now limited only by the dose to the sample, and thermal vibrations of the atoms themselves [2]. At high doses, these approaches have allowed us to image the detailed vibrational envelopes of individual atom columns as well as locating individual interstitial atoms that would be hidden by scattering of the probe with conventional imaging modes. The three-dimensional nature of the reconstruction means surface relaxations can be distinguished from the bulk structure, and interface roughness and step edges inside devices can be resolved – including gate-all-around transistors and Josephson junctions. Even the location of all atoms in thin amorphous films now seems within reach. These approaches have also allowed us to image the internal structures of both magnetic and ferroelectric vortices, skyrmions and merons, including their singular points that are critical for accurately describing the topological properties of these field textures.

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2:45pm **PCSI-WeA1-10 Cryogenic Growth and *in-Situ* Fabrication of Superconducting Tantalum Devices**, *Teun van Schijndel*, UC Santa Barbara; *A. McFadden*, NIST-Boulder; *A. Engel*, *J. Dong*, *S. Chatterjee*, UC Santa Barbara; *R. Simmonds*, NIST-Boulder; *C. Palmstrøm*, UC Santa Barbara

Superconducting devices are crucial in various fields of quantum information technology, including superconducting qubits and topological quantum computing. The vast majority of either of these qubit technologies use Aluminum as the superconducting component. Al is generally grown at low temperatures to achieve smooth thin films. This allows for easy integration of Al-based devices with material systems such as sapphire, Si, Ge, or III-V materials due to minimal interfacial reactions. Also, Al is often used for Josephson Junctions (JJ) that require *in-situ* oxidation to form AlO_x barriers. While Al is the most common superconductor, other superconductors show promising results as well. In particular, Tantalum-based superconducting qubits on sapphire show low loss and long coherence times.^{1,2} One of the contributing factors to the enhancement of qubit performance is their higher chemical resistance during device fabrication. However, only a few substrates can be used to stabilize the required α -Ta phase with a BCC lattice structure. Due to its resilience to high temperatures, low-loss sapphire can withstand the growth of Ta at elevated temperatures necessary for the realization of desirable superconducting properties. Growth of Ta on Silicon or III-V substrates remains a challenge.

In this work, we explore the MBE growth and *in-situ* fabrication of superconducting tantalum films. The growth at ultralow temperatures below 10K shows the stabilization of the required superconducting phase of Tantalum (α -Ta). Moreover, Figure 1 shows that α -Ta can be stabilized at low temperatures regardless of the substrate choice. In each case, a superconducting transition temperature of above 4K is observed. Furthermore, this deposition technique can be combined with *in-situ* shadow masks, which allows for patterns with at least 1 μ m precision. This can be used to realize Ta/Ta₂O₅/Ta JJ's by using the native oxide, which is something that has never been shown before, or by depositing other

dielectrics *in-situ* such as Silicon or Germanium. Our work demonstrates the growth of high-quality superconducting devices, which enables the exploration of different superconductors and dielectric combinations for use in quantum information technology.

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2:50pm **PCSI-WeA1-11 Multi-Technique Characterization of GaN-Based Devices: A Powerful Tool to Probe the in-Depth Chemistry**, *Kirène Gaffar*, CNRS, Université Paris-Sud, France; *S. Béchu*, *G. Patriarache*, *M. Bouttemy*, CNRS, France

Gallium Nitride (GaN) technology has proven to be a contender for power electronic applications and has shown its suitability for GaN-based devices such as High Electron Mobility Transistors (HEMT) for high-frequency applications. However, the miniaturization of the device dimensions, such as gate length, requires a thorough mastering of device fabrication process with the help of suitable analytical techniques. In particular, systematic electrical characterization has shown that the interface states have a significant impact on the electrical performance and long-term reliability of GaN HEMT devices. This work ambitions to develop a robust methodology to perform advanced chemical characterization of GaN transistors and better understand how the chemistry of the constitutive layers and interfaces properties impact the electrical response. In addition to STEM/EDX analysis on cross sections, conventionally used to access quantitative information and epitaxy quality, an innovative methodology combining X-ray photoemission spectroscopy (XPS) and Auger Electron Spectroscopy (AES) is developed. Indeed, these techniques not only provide access to elemental compositions but also to key information on chemical environments, especially modifications induced by each technological step during device fabrication. In particular, (nano)-Auger spectroscopy, with its high spatial resolution (12 nm), is a very promising tool to access buried interfaces directly on cross-section, bringing complementary information to STEM/EDX such as oxidation states or contaminant presence. If the added value of this multi-technique approach is obvious, the direct implementation and interpretation of XPS and Auger analyses is not straightforward. In fact, access to the ultimate composition of any GaN-based structure is conditioned by the fitting procedure to ensure a reliable nitrogen content determination. We propose here a reliable methodology to decompose gallium L₂M_{4.5}M_{4.5} transition and nitrogen N 1s photopeak overlap for precise quantification. We will show how these preliminary results obtained by XPS and STEM/EDX analysis are crucial for an accurate interpretation of Auger spectrum acquired on the same materials with the nano-probe for chemical state identification and to refine the quantification. The correlation between structural observations, chemical information and electrical performances measured on a HEMT transistor will be illustrated on a concrete case.

2:55pm **PCSI-WeA1-12 Mo-SiN_x Granular Metal High-pass Filters**, *Laura Biedermann*, *M. McGarry*, *S. Gilbert*, *W. Bachman*, *M. Meyerson*, *L. Yates*, *P. Sharma*, *J. Flicker*, *P. Kotula*, *M. Siegal*, Sandia National Laboratories

Granular metals (GMs) comprise a 3D network of metal nanoparticles embedded in a dielectric matrix. Over the past ~50 years, GM investigations have spanned fundamental physics to unique applications, including Au-SiO₂ GMs used as insulating contacts in vidicons, video cameras used in NASA's Apollo and Voyager missions [1]. As a controlled platform for electron transport studies, GMs exhibit tunneling transport (e.g. variable-range hopping, Poole-Frenkel conduction in Ni-SiO₂ GMs) and frequency-dependent conductivity $\sigma(\omega)$ in Pt-SiO₂ and Pd-ZrO₂ GMs [2–4]. These prior GM investigations focused almost exclusively on metal-oxide GMs. Our goal—to develop nanosecond-responsive high-pass filters for electrical grid applications—has advanced development of Mo-SiN_x GMs that exploit these conductivity mechanisms.

High-dielectric strength SiN_x is an attractive matrix for GMs, enabling Mo-SiN_x and Co-SiN_x GMs [5]. However, initial Mo-SiN_x GMs showed weak $\sigma(\omega)$; thermally-excited resistive transport through defective SiN_x overwhelmed the desired transport mechanisms. Fortunately, sputtering Mo-SiN_x in a partial N₂ environment ameliorates these SiN_x matrix defects. X-ray photoemission spectroscopy (XPS) analysis shows deleterious MoSi₂ is further reduced by annealing in H₂/N₂ forming gas (Fig. 1a). Improvements in SiN_x insulator quality resulted in the desired many decades reduction in σ_{DC} (Fig. 1b). This evaluation of nanostructure and chemical structure has enabled optimization of high-frequency and high electric (*E*) field transport

(Fig. 1c, d), key properties of high-pass filters for electric grid applications [6].

This work was supported by the Laboratory Directed Research and Development program at Sandia National Laboratories (SNL). SNL is managed and operated by NTESS under DOE NNSA contract DE-NA0003525. SAND2023-09502A.

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3:00pm **PCSI-WeA1-13 Restructuring Cracks in Rutile TiO₂ with Radiolysis-Driven Rolling of Octahedral Units**, *Silu Guo, H. Yun, S. Nair, B. Jalan, K. Mkhoyan*, University of Minnesota, USA

When energetic electrons interact with crystals in transmission electron microscope (TEM), a combination of "knock-on" and radiolysis effects takes place [1]. Radiolysis, in particular, is known to either amorphize or crystallize materials and limit the accuracy of the measurements [2]. However, the precise atomistic mechanisms of these transformations are still under debate. Here, we use scanning TEM (STEM) imaging and electron energy loss spectroscopy (EELS) to study the bond-breakage, atomic movements and crystallization mechanisms in rutile-TiO₂ driven by radiolysis [3].

Thin-film rutile IrO₂ was grown on top of rutile TiO₂, introducing nanometer width cracks due to an anisotropic epitaxial strain (Figure 1a). To assess the impact of electron beam exposure on the structural healing process of the atomically sharp cracks in rutile-TiO₂, high angle annular dark-field STEM (HAADF-STEM) time-lapse images of the cracks were obtained (Figure 1b). With the accumulation of electron doses, the crack undergoes a self-healing restructuring process. Based on these observations and quantitative EELS analysis, we propose a "2-step rolling" model for the TiO₆ octahedral building blocks located at the crack's edge of rutile-TiO₂ as a possible mechanism for radiolysis-driven atomic migration (Figure 1c and d). With radiolytic bond breakage (Figure 1c), the TiO₆ octahedral units from the edge of the crack can roll and occupy new sites and, in the process, move materials from both sides of the crack into the gap (Figure 1d) [3]

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3:05pm **PCSI-WeA1-14 UPGRADED: Growth and Angle-Resolved Photoemission of Strain- and Thickness- Tuned Epitaxial α -Sn Thin Films**, *Aaron Engel, H. Inbar*, University of California, Santa Barbara; *P. Corbae, C. Dempsey, S. Nishihaya, Y. Chang*, University of California, Santa Barbara; *A. Fedorov*, Advanced Light Source, Lawrence Berkeley National Laboratory; *M. Hashimoto, D. Lu*, SLAC National Accelerator Laboratory; *C. Palmstrøm*, University of California, Santa Barbara

α -Sn, the diamond structure allotrope of Sn, is a zero-gap semiconductor with band inversion. Calculations suggest that epitaxial tensile strain induces a 3D topological insulator (TI) phase, while epitaxial compressive strain induces a 3D Dirac semimetal (DSM) phase [1,2]. When this DSM phase is confined in a thin film, it has been suggested to form a quasi-3D TI phase [3]. Transitions to other phases instead, such as 2D TI, have been suggested as well [4].

We first explore the topological phase of ultrathin unintentionally doped α -Sn thin films. Using spin- and angle-resolved photoemission spectroscopy (ARPES), we study compressively strained α -Sn films on InSb(001). We find clear evidence of the confinement-induced quasi-3D TI phase in compressively strained α -Sn. We also find that the spin-polarization of the topological surface states differs markedly from reports in the literature where the films are intentionally doped with Te [5], indicating this intentional doping (a frequently used procedure) could have a significant effect on the electronic structure of α -Sn.

With the previous behavior benchmarked, we then alloy the α -Sn films with Ge to decrease the bulk lattice constant and switch from compressive to tensile strain when grown on InSb(001). Morphology changes as a function of Ge alloying were studied with *in-situ* scanning tunneling microscopy, and strain was confirmed through X-ray diffraction. The presence of a topological phase transition induced by tensile strain away from the expected 3D TI phase is found in ARPES (Fig. 1). Our results pave the way for a better understanding of the effect of strain and confinement on α -Sn's band structure.

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3:25pm **PCSI-WeA1-18 Characterization of Buffer Layers for Remote Plasma-Enhanced Chemical Vapor Deposition of Germanium-Tin Epitaxial Layers**, *Stefan Zollner, C. Armenta*, New Mexico State University; *B. Rogers*, Vanderbilt University; *G. Grzybowski, B. Claffin*, Air Force Research Laboratory

Germanium-tin alloys are of interest for infrared light detectors to increase capabilities in image and data capture and transmission, because they can have a direct band gap with more than about 7% tin. Remote plasma-enhanced chemical vapor deposition (RPECVD) is attractive for growth of Ge-Sn alloys because it enables low-temperature epitaxy on Si using common precursors GeH₄ and SnCl₄. The growth of such epilayers can be optimized with an initial high-temperature buffer layer. This work focuses on the characterization of this buffer layer using atomic force microscopy, ellipsometry, thin-film powder x-ray diffraction, and x-ray photoelectron spectroscopy (XPS) for different growth conditions.

Thin Ge and Ge-Sn buffer layers with 10-20 nm thickness were deposited on Si (100) substrates for one minute at temperatures from 360°C to 500°C with varying SnCl₄ precursor flows mixed with GeH₄ and helium. Ellipsometry spectra for all films show critical point structures in the E1, E1+ Δ 1, and E2 region of Ge, indicating that all layers are crystalline. A layer grown at 360°C without SnCl₄ can be described reasonably well as an 11 nm thick layer of crystalline germanium with 2 nm of roughness. Adding SnCl₄ to the gas flow significantly reduces the height of the ϵ_2 maximum at E2, indicating that the layer is rough. In addition, a new broad peak appears near 1.3 eV, which is attributed to plasmonic effects arising from metallic β -tin inclusions. The plasmon peak disappears in the layers grown at 490°C with the same SnCl₄ flow. We conclude that depositing the buffer layer with SnCl₄ at low temperatures leads to β -Sn precipitates, where plasmon oscillations can be excited, but are not present for high-temperature growth.

The tin contents in the layers were also estimated by x-ray photoelectron spectroscopy. While XPS measures the total amount of tin in the layers, the presence of substitutional tin in Ge_{1-x}Sn_x alloy buffers is best determined with x-ray diffraction. The (002) diffraction peak is forbidden in pure Ge, because the contributions from the two Ge atoms in the primitive unit cell cancel. It is absent in our buffers grown without SnCl₄ or at high temperature. The (004) XRD peak position in these layers is also very similar to pure Ge. The Ge_{1-x}Sn_x (002) peak does appear in buffers grown at temperatures lower than 460°C. From the position of the (004) XRD peak, we can estimate the tin content to be below 7%, ignoring the effects of residual stress. This tin content determined from XRD shifts is much lower than the total tin content of about 20% estimated by XPS.

3:30pm **PCSI-WeA1-19 Near Zero-Field Magnetoresistance and Defects in GaN pn Junctions**, *M. Elko, A. Higgins, D. Hassenmayer, Patrick Lenahan*, Pennsylvania State University; *M. Flatte, D. Fehr*, University of Iowa; *T. Larsen, M. Craven*, NexGen Power Systems

We report on observation of near zero-field magnetoresistance (NZFMR) in GaN devices, in this case, pn junction diodes. NZFMR is a new technique with great potential in electronic materials physics. [1,2] The NZFMR response is due to recombination centers within the diode depletion regions. A representative NZFMR amplitude versus magnetic field plot is shown in figure 1. Figure 2(a) illustrates the anticipated depletion region recombination current versus bias predicted by standard first order expressions. [3] Figure 2(b) shows the measured NZFMR amplitude versus junction bias. The agreement between the calculated response of figure

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2(a) and the experimental observations of figure 2(b) should be considered reasonably convincing, considering multiple approximations involved. The NZFMR pattern peaked near the built-in voltage is expected from recombination within the depletion region. [3,4] The NZFMR phenomena are somewhat similar to low field magnetoresistance phenomena observed in some organic semiconductors. The NZFMR response can be understood within the framework of the stochastic quantum Liouville expression. [1,2] Preliminary analysis of traces represented by figure 3, based upon this framework, indicates that the NZFMR response is consistent with nitrogen vacancies.

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PCSI

Room Ballroom South - Session PCSI-WeA2

Semiconductor Heterostructures (Growth Nanostructures & Interfaces) II

Moderator: Jason Kawasaki, University of Wisconsin - Madison

4:30pm PCSI-WeA2-31 Heteroepitaxy of PbSe-SnSe Semiconductors on GaAs for Infrared Optoelectronics, Kunal Mukherjee, Stanford University

INVITED

The IV-VI semiconductor alloy of PbSe-SnSe has rich materials physics and several device applications[1]. This narrow band gap semiconductor system spans a 3D-bonded rocksalt structure on the PbSe-rich side and a closely related Van der Waals bonded 2D/layered-orthorhombic structure on the SnSe-side, and has long been studied for its rare electronic, photonic, and thermal properties. We present results from epitaxial integration of IV-VI semiconductor films with III-V templates using molecular beam epitaxy to understand how we may harness their properties for emerging applications infrared optoelectronics.

With attractively low temperatures for epitaxy below 300 °C, we describe the nucleation and growth of IV-VI materials on III-V substrates and the formation of extended crystal defects that arise due to integration[2]. We find bright band-edge photoluminescence in the mid-infrared (3–4 μm) from PbSe and PbSnSe epitaxial films on ~8% mismatched GaAs at room temperature, despite a threading dislocation density exceeding 10⁹/cm². We present measurements of carrier recombination in PbSe and show preliminary results from junction devices of mid-infrared light emitting diodes and photodetectors that highlight the promise of this materials platform.

The IV-VI on III-V template also enables us to probe deeper into the structural phase boundaries and miscibility gaps in PbSnSe, with an aim to harness the high contrast in optical properties across the transition between 3D/rocksalt to 2D/layered bonding. We show that MBE synthesis can stabilize the layered phase deep in the bulk miscibility gap. Close to a composition of Pb_{0.5}Sn_{0.5}Se, we find evidence for a displacive or martensitic transformation (without composition change) between the rocksalt and layered phases in our thin films. These results, in agreement with recent reports in high temperature quenched samples[3], point the way to phase change devices.

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5:10pm PCSI-WeA2-39 Investigation of Localized Electric Fields of InAs/GaAs Quantum Dot Interfaces, T. Kang, Jong Su Kim, Department of Physics, Yeungnam University; S. Lee, Division of Convergence Technology, Korea Research Institute of Standards and Science

In InAs/GaAs QDs, the strain-induced lattice deformation and strain induced dislocation could generate localized electric fields (LEF) due to the piezo-electric fields [1]. Therefore the electric field distribution could be more complex at the interface of QDs than for the 2D superlattice (SL). The strain-induced complex electric fields can significantly modify the quantum confinement states. Direct observation of strain-related effect, such as the LEFs caused by strain-induced polarizations and defects, is therefore very important. Photoreflectance spectroscopy (PR) is useful for investigating LEFs of semiconductors [2]. In the case of GaAs, the Franz-Keldysh oscillations (FKOs) that appear above the band gap of GaAs and contain information about the LEFs in GaAs. To investigate InAs/GaAs quantum confinement states PR experimental results were reported [3]. However, experimental observation of LEFs attributed to strain between InAs QD and GaAs has not been the focus of many studies.

In this work, we investigated LEFs between InAs QD and GaAs by PR. FIG. 1 shows the InAs/GaAs QDs sample used in this work. FIG. 2 shows the PR spectra of the InAs/GaAs QD samples obtained at low temperature. The low-temperature PR spectrum of the InAs/GaAs QD shows clear FKO transitions above the GaAs band gap energy. This work suggests that the interface electric fields attributed to strain originate from the strain-induced polarization near the InAs QD interface in GaAs metric. We suggested that the FKOs originated from the LEFs predominately caused by the strain-induced polarization at GaAs interface near the InAs QDs. The InAs/GaAs QDs have a broad range of interface electric fields from ~10⁴ V/cm to ~2x10⁵ V/cm.

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5:15pm PCSI-WeA2-40 X-STM Study of Interlayer Effects on InAs Quantum Dots in InP, Edoardo Guido Banfi, Eindhoven University of Technology, Netherlands; E. Sala, Sheffield University, UK; R. Gajjala, Eindhoven University of Technology, Netherlands; J. Heffernan, Sheffield University, UK; P. Koenraad, Eindhoven University of Technology, Netherlands

Quantum dots (QDs) have been studied in the past two decades in order to optimize their performances in a wide range of applications, from QD lasers, photovoltaics, single photon emitter, memories, and last but not least, quantum communication and information technologies [1-6]. In order to further follow this line of optimization it is relevant to achieve more size and shape uniform QDs. This work targets the optimization process of droplet epitaxy (DE) InAs QDs in InP by the use of interlayers.

In this work we performed, with atomic scale resolution, cross-sectional scanning tunneling microscopy (X-STM) on InAs DE QDs in InP, we also characterized the QDs with atomic force microscopy (AFM) and performed finite element simulations (FES). We measured two samples with different compositions in the interlayer (IL), the region below the QDs. The first sample has an In_{0.53}Ga_{0.47}As IL and consists of two separate QDs layer with different amount of In (referred to as QDL1 and QDL2). The two regions are separated by 100nm of InP in order to reduce the chances of seeding of the QDs formation. The second sample has an IL composed by In_{0.719}Ga_{0.281}As_{0.608}P_{0.392} (Fig. 1) lattice matched to the InP of the host. Our study performed on these samples, highlighted how some previously identified effects (trenches and etch-pit formation [7]) were successfully avoided in these samples. We studied size, shape and composition of all the QDs measured. AFM and X-STM proved that these dots have a rhombic base elongated preferentially along the {110}. We performed FES and we can conclude that in the second sample the QDs purity is 95±5% InAs, while in the other sample the purity is around 90±5%. Results confirmed are not only by the relaxation and lattice constant but also from our X-STM measurements. These results prove that introducing P in the IL region can enhance the purity of the QDs and allow for a stress-free structure.

5:20pm PCSI-WeA2-41 UPGRADED: Atomic Scale Analysis of N Dopants in InAs, T. Verstijnen, D. Tjeertes, E. Banfi, Eindhoven University of Technology, Netherlands; Q. Zhuang, Lancaster University, UK; Paul Koenraad, Eindhoven University of Technology, Netherlands

The band gap of most III-V semiconductors is strongly reduced with the introduction of only a few percent of N, even if the III-N alloy has a much

bigger band gap. N impurities in InAs introduce an impurity state around 1 eV above the conduction band minimum, much deeper in the band than in other III-V materials. Topographic scanning tunneling spectroscopy measurements (STS) and areal spectroscopy measurements performed on N atoms up to two layers below the (110) surface of InAs show a reduction of the resonance energy of the N atom with increasing depth. This is attributed to tip induced band bending, pulling the N states up at positive bias and acting most strongly on surface N atoms. An example of STM images on InAs:N is shown in figure 1. STS measurements obtained on undoped InAs and N-doped InAs show a band gap reduction of <0.1 eV. Spatial imaging of features corresponding to N dopants up to two layers below the surface are also compared to density functional theory simulations and show excellent correspondence. Spectroscopy maps of N atoms up to two layers below the surface provide a high resolution spatial and spectroscopic view of the N atoms. Here the characteristic shape of the N atoms in different layers below the surface is observed as an enhancement of the dI/dV signal compared to the InAs background. At energies above the enhancement a reduction of the dI/dV is observed, which has the same shape and size as the enhancement. This shows that the redistribution of density of states caused by the N impurities is mainly energetic in nature.

5:40pm **PCSI-WeA2-45 Direct Wafer Bonding of GaN on AlN Through the Optimization of Chemical Mechanical Polishing**, *Kaicheng Pan, K. Huynh, M. Li, Y. Ge, T. Fisher, Y. Hu, M. Goorsky, UCLA*

Optimization of surface preparation and interfacial characterization of direct wafer bonded GaN to AlN are presented in this study. In particular, the Ga face of the GaN wafer was bonded to the N face of the AlN wafer. The as received Ga-face of GaN substrates showed <1 nm roughness while the N-face AlN had a starting roughness of ~3 nm RMS. The N-face of the AlN was successfully polished to <1 nm RMS roughness suitable for direct wafer bonding using chemistry based on our previous work with GaN CMP [1]. For bonding, 2" GaN (Unipress) and 2" AlN (Hexatech) were bonded using standard cleaning and immersion in a $(\text{NH}_4)_2\text{S}$ solution. The samples were rinsed, dried and pressed Ga- to N-face (AlN) under moderate pressure (~50 kPa) and room temperature bonding was initiated. A significant fraction of the surfaces bonded, except for a couple of triangular regions associated with growth sector boundaries in the GaN. Subsequent annealing up to 800 °C was performed to strengthen the bond and to test the structure for high temperature stability. Similar coefficient of thermal expansion between GaN and AlN at high temperatures allows for high temperature annealing without debonding or cracking. The GaN substrate was then grinded and also subject to CMP to < 1µm for transmission electron microscopy and time-domain thermal reflectance measurements of the bonded interface.

High resolution transmission electron microscopy shown in Figure 1 reveals complete crystallinity across the interface. However, only a ~1.5 nm interfacial region is observed, which is suspected to be caused by reconfiguration of the interface after a total anneal of 350 °C 22 hours, 600 °C 1 hour, and 800 °C 1 hour. No thicker amorphous or oxide interfacial layer commonly found in other bonding methods (surface activated bonding, plasma treatment, or other interfacial layers) [2-5] are observed in this study. Preliminary thermal boundary conductance measurements via time domain thermal reflectance have been measured and will be reported.

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5:45pm **PCSI-WeA2-46 Strategies for Analyzing Non-Common-Atom Heterovalent Interfaces: The Case of CdTe-on-InSb**, *Esperanza Luna, A. Trampert*, Paul-Drude-Institut für Festkörperelektronik Leibniz-Institut im Forschungsverbund Berlin, Germany; *J. Lu, T. Aoki, Y. Zhang, M. McCartney, D. Smith*, Arizona State University

Semiconductor heterostructures are intrinsic to a wide range of modern-day electronic devices. Knowledge of chemical interfacial profiles in these complex structures is critical to the task of optimizing the device performance. Here, we report on an innovative methodology that enables reliable interface structure analysis of non-common-atom heterovalent interfaces on all relevant length scales from hundred-nm to atomic resolution.

Non-common-atom (NCA) heterovalent interfaces offer potential benefits arising from the valence mismatch but also contain challenges due to the

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large charge imbalances. Whether a sharp polar interface is formed or, on the contrary, there are mixtures of chemical bonds across the interface leading to a nonpolar graded interface is under intense discussion. Furthermore, very little has so far been done to exploit the opportunities offered by NCA heterovalent interfaces, in part due to challenges determining the structure and properties of these types of interfaces, for example, by using scanning transmission electron microscopy (STEM) techniques. This work presents a comprehensive analysis of the composition profile across the case study NCA heterovalent CdTe/InSb interface, carried out using a combination of (S)TEM imaging and spectroscopic techniques. Techniques such as high-angle annular-dark-field and large-angle bright-field STEM, as well as electron energy-loss spectroscopy, give results from the interface region on the atomic scale. These measurements, however, are inherently difficult to interpret because of the close atomic numbers of the constituent elements. In contrast, use of the 002 dark-field TEM imaging mode emphasizes the interface location by comparing differences in structure factors between the two materials. Based on the 002 dark-field TEM, a methodology was developed for reliable determination of the composition profile across the interface and systematic quantification of the interface width. Since the identities of both cations and anions change across the heterointerface, the respective contributions of different elements need to be inserted separately. The intermixing at each sub-lattice is thus independently and completely determined. Comparisons of experimental and simulated CdTe-on-InSb profiles reveal that the interface is structurally abrupt to within about 1.5 nm defined by the variation between 10 and 90%. The present investigation opens new routes to the systematic investigation of heterovalent interfaces, formed by the combination of other valence-mismatched material system.

5:50pm **PCSI-WeA2-47 Multi-Material Deposition for Spatial Atomic Layer Deposition Process**, *A. Varga, M. Carnoy, M. Plakhotnyuk, I. Kundrata*, ATLANT 3D, Denmark; *J. Bachmann*, Friedrich-Alexander-University Erlangen-Nürnberg (FAU), Germany; *M. Baraket*, ATLANT 3D, Denmark; *Simone Santucci*, ATLANT 3D Nanosystems, Denmark

Spatial Atomic Layer Deposition (sALD) offers a unique opportunity for localized deposition due to its physical separation and isolation of precursor and co-reagent dosing.^[1] While simple in theory, due to well-developed examples of sALD, in practice miniaturization of sALD requires substantial effort into the creation of suitable micro-nozzles.^[1] Uniquely, ATLANT 3D has developed proprietary sALD micronozzles, called microreactor Direct Atomic Layer Processing - μ DALP™.

The μ DALP™ process undergoes the same cyclic ALD process but is only done in a spatially localized area.^[2] The microreactor or micronozzle confines the flows of gases used for ALD within a defined μ m-scale area on the substrate, to deposit the desired material.

Since sALD and the μ DALP™ process are based on physical separation, it is theoretically compatible with any ALD material process however requires development as ALD processes are highly tool dependent.^[3] As such, the material capabilities can match traditional ALD and exceed other patterning techniques, such as lithography, which can be costly and time-consuming, especially for rapid prototyping required for innovation.^[4,5]

Using a small amount of precursor multiple film materials and thicknesses can be deposited onto a single wafer within only a few hours, compared to days for a traditional ALD process (Fig 1.). Films deposited with ATLANT 3D technology have been shown to produce high-quality, crystalline, atomically precise thin films used to fabricate temperature (Fig 2.) and capacitive sensors with sensitivities that meet or exceed those of devices made using conventional vapor phase deposition techniques. Low-cost rapid prototyping facilitated by ATLANT 3D technology of such devices enables design innovation and optimization not possible with other thin film deposition techniques.

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PCSI

Room Ballroom South - Session PCSI-ThM1

Wide Bandgap Materials

Moderator: Christopher Palmström, University of California, Santa Barbara

8:30am PCSI-ThM1-1 Surface Transfer - Modulation Doping at a Diamond-Dielectric Interface, **Robert Nemanich**, Arizona State University **INVITED**

Great strides in diamond wafer technology and diamond epitaxy have inspired new concepts for diamond electronics particularly for power conversion and RF applications. However, the high activation energy of substitutional p- and n-type dopants in diamond has limited the development of field effect transistors (FET). An alternative approach of charge transfer doping at a diamond-dielectric interface, which results in the formation of a hole accumulation layer, is not limited by thermal activation [1]. However, the hole transport shows a mobility that is much lower than predicted. It is widely accepted that the low mobility is due to scattering from the near interface negative charges transferred into the dielectric layer.

Following the concept of modulation doping at heterostructure interfaces [2], we have proposed and demonstrated (Fig. 1) a dielectric layer configuration that results in a nearly ten-fold mobility increase for the accumulated holes at the diamond interface [3]. In this approach MoO_3 is used as the charge transfer dielectric, and Al_2O_3 is employed as the modulation doping spacer layer. The charge transfer is driven by the energy difference between the diamond valence band and the charge transfer states in the MoO_3 . The thickness of the spacer layer also affects the hole accumulation layer charge density.

In this study photoemission spectroscopy is employed to measure the band alignment and band bending throughout the multi-layer structure. The relative distribution of the charge near the interface is deduced from the band diagram. Another study developed a modulation doping approach using acceptor molecules (NO_2) to enable the charge transfer from the diamond valence band. They achieved results that were similar to the multi dielectric layer approach.

These experiments and the model of Surface Transfer - Modulation Doping demonstrates a new approach to FET channel doping for diamond field effect transistors. We also discuss options for improving modulation doping in diamond FET's.

This research was supported by a grant from MIT-Lincoln Laboratories and the NSF through Grant Nos. DMR-1710551 and DMR-2003567.

9:10am PCSI-ThM1-9 Operation-Induced Short-Term Degradation Mechanisms of 275-Nm-Band AlGaN-Based Deep-Ultraviolet Light-Emitting Diodes Fabricated on a Sapphire Substrate, **Shigefusa Chichibu**, Tohoku University, Japan; **K. Okuno**, **M. Oya**, **Y. Saito**, **H. Ishiguro**, Toyoda Gosei Co. Ltd., Japan; **T. Takeuchi**, Meijo University, Japan; **K. Shima**, Tohoku University, Japan **INVITED**

The short-term degradation mechanisms of 275-nm-band AlGaN multiple quantum well (QW) deep-ultraviolet light-emitting diodes fabricated on a (0001) sapphire substrate were investigated under hard operation conditions with the current density of 66 A/cm^2 and the junction temperature of $105 \text{ }^\circ\text{C}$. The optical output power (P_o) decreased by about 20 % within the operating time (t_{op}) less than 102 h and then gradually decreased to about 60 % by 484 h, as shown in Fig. 1. For elucidating the cause for the initial degradation ($t_{\text{op}} < 102 \text{ h}$), complementary electrical, time-resolved photoluminescence (TRPL), and impurity characterizations were carried out making a connection with the energy band profiles.

The initial degradation was accompanied by the increases in both the forward current (I_f) below the turn-on voltage (V_b) and reverse leakage current (I_R). These results are consistent with those reported previously [1-5]. Because the weak-excitation room-temperature PL lifetime for the near-band-edge emission using the QW-selective TRPL showed only slight change by the operation at least until 1002 h, the initial degradation is attributed essentially to the decrease in carrier injection efficiency. From the correlation between the energy band profiles and H concentration profiles before and after the operation, the output power reduction is ascribed to be due to de-passivation of initially H-passivated preexisting nonradiative recombination centers (NRCs) in a Mg-doped p-type $\text{Al}_{0.85}\text{Ga}_{0.15}\text{N}$ electron blocking layer (EBL) caused by certain breaking of H bonds and the electric field induced drift of H^+ . According to our database on the species of vacancy-type defects acting as NRCs in AlN [6] and GaN [7], vacancy clusters comprised of a cation vacancy (V_{III}) and nitrogen vacancies (V_{N}), such as

$V_{\text{III}}(V_{\text{N}})_{2-4}$, are the most suspicious origins of the NRCs in the Mg-doped p-type AlGaN layers [8].

This work was supported by MOE program for implementation of innovative infection-control and digital technologies with low CO_2 emissions and MEXT Crossover Alliance, Japan.

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9:50am PCSI-ThM1-17 Impact of Interfacial Defects and Lattice Strain on NbN_x Films for Integration with Wide Bandgap Semiconductors, **Annaliese Drechsler**, University of Maryland College Park; **P. Shea**, Northrop Grumman; **A. Christou**, University of Maryland College Park

Niobium nitride (NbN) films have garnered significant attention due to their high critical temperature (T_c) and their usage in infrared-sensitive superconducting nanowire single photon detectors (SNSPDs). Devices fabricated from NbN have demonstrated single photon detection to mid-wave infrared wavelengths, which unlocks possibilities for novel applications such as long-range laser detection and ranging (LiDAR), interferometry of planetary emissions, quantum key decryption, and optical communications. To expand beyond a laboratory, however, these devices must be fabricated into focal plane arrays (FPAs), requiring integration with semiconducting device materials. In this work, we report progress on achieving a device structure comprised of a NbN SNSPD monolithically integrated with a wide bandgap semiconductor-based amplifier. This investigation is motivated by recent reports of monolithic integration of NbN with aluminum nitride (AlN) to provide a superconducting load for an amplifier [1].

NbN films for SNSPDs must be thin, typically $\ll 100$ nanometers. As a result, the film quality and defectivity, and ultimately SNSPD performance, are highly correlated to the interface between the NbN film and underlying lattice, lattice-mismatch strain, and deposition parameters of the NbN processing. In this talk, we investigate the impacts of the semiconducting interface on the NbN films utilized for SNSPD fabrication through XRD of grown films, AFM surface studies, cathodo-luminescence (CL), and TEM analysis. To optimize this interface, similar materials 6H-SiC (3.57% lattice mismatch) and wurtzite GaN (6.6% lattice mismatch) chosen to minimize intrinsic defect sources. XRD analysis of grown films indicates that growth on these substrates is possible with long range crystallinity, suggesting the presence of epitaxial growth for high quality films. Optimization of the stress of the film due to the lattice mismatch with the substrate is also investigated by modifying the growth temperature, pressure, and power to reduce lattice strain-induced defects. The presence of threading and other dislocations stemming from interface defects analyzed through CL and TEM will also be discussed.

9:55am PCSI-ThM1-18 Impact of Unintentional Boron Supply on Sapphire Nitridation Process for GaN Growth by RF-MBE, **Tohru Honda**, **K. Yajima**, **T. Yayama**, **T. Onuma**, **T. Yamaguchi**, Kogakuin University, Japan

GaN, InN and their related alloys grown by radio-frequency excited plasma-assisted molecular beam epitaxy (RF-MBE) [1] have investigated for the application to light-emitting devices operating in visible and infrared spectral regions [2]. Although the plasma assisted nitrogen source is a useful for III-nitrides by MBE, unintentional boron (B) supply, which is coming from the nitrogen cell, during the growth was reported [3]. The nitridation process was used for N-polar GaN growth on a sapphire substrate [4]. Although some nitridation models were reported [4], the boron incorporation during the nitridation is still unclear. Thus, the impact on unintentionally supplied boron incorporated during the nitridation is discussed.

Chemically cleaned (0001)sapphire substrates were used for the study. Thermal cleaning at $850 \text{ }^\circ\text{C}$ for 15 min. was performed before the nitridation. The nitrogen plasma (200 W, N_2 flow of 0.6 ccm) was used for the nitridation, whose temperature was fixed at $500 \text{ }^\circ\text{C}$. The samples were taken out the growth chamber to air, subsequently, XPS and Auger electron spectroscopy (AES) spectra were observed. Boron 1s peaks (B-N and B) [5] were observed from the sapphire surface with the nitridation. This means that boron atoms or compounds were deposited on the sapphire. N 1s peaks related Al-N and B-N were also observed. To confirm the deposition

of boron related layers, AES was also observed for the samples. We observed B (KVV) signals from those. These indicate that the boron related layers were deposited on the sapphire substrates during the nitridation.

Generally, it was reported [6] that the suitable nitridation time led to the high quality GaN growth. We believed AlN coverage on the sapphire substrate by the nitridation was a key for the GaN growth with high crystalline quality. XPS results indicate that the amount of AlN is increased as a function of the irradiation time and it's saturated. On the other hand, the boron deposition is monotonically increased. The boron on the surface obstructs the GaN growth although the AlN enhanced it with high crystalline quality.

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10:00am **PCSI-ThM1-19 Photoluminescence Maps of Surface Defects in β -Ga₂O₃**, **Matthew McCluskey**, Washington State University; *J. Huso*, Klar Scientific; *C. Remple*, *J. McCloy*, Washington State University; *S. Rebollo*, *S. Krishnamoorthy*, *J. Speck*, University of California at Santa Barbara

Monoclinic gallium oxide (β -Ga₂O₃) is an ultrawide bandgap semiconductor with potential applications in power electronics [1]. Photoluminescence (PL) spectroscopy is an important method to characterize dopants and defects in this material. Common features in the PL spectrum include the intrinsic UV band, blue and green bands that involve donor-acceptor pairs, and red emission due to Cr³⁺ impurities.

PL mapping with excitation wavelengths ranging from 266 to 532 nm reveals the spatial distribution of these features with micron resolution. In Czochralski-grown β -Ga₂O₃, the Cr³⁺ emission intensity shows striations that are attributed due to inhomogeneities during growth [2]. In addition to defects in the bulk, PL microscopy has revealed several specific defects on the surface. Some of these localized centers are very bright UV emitters [3]. Homoepitaxial layers show defects that are observed via the shifts in the PL band, likely due to the strain field around a dislocation core. Damage due to high-intensity laser pulses results in significant changes in the intensity and energy of the UV band. *In situ* PL spectroscopy performed with a pulsed 266 nm laser shows characteristic emission peaks attributed to Ga atoms ablated from the surface.

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10:05am **PCSI-ThM1-20 UPGRADED: Epitaxial Growth and Properties of Wide Bandgap P-Type NiGa₂O₄ on β -Ga₂O₃ for High Voltage P-N Heterojunctions with Superior Performance at Elevated Temperatures**, **Kingsley Egbo**, *B. Tellekamp*, *W. Callahan*, *A. Zakutayev*, National Renewable Energy Laboratory

Gallium oxide (β -Ga₂O₃) is a promising wide bandgap oxide semiconductor material with properties well-suited for high-power electronics, and recent results show superior high-voltage performance compared to the commercial state-of-the-art [1],[2]. Due to the difficulty in the *p*-type doping of Ga₂O₃, unipolar devices based on Ga₂O₃ are prevalent. Several studies have explored bipolar devices using polycrystalline *p*-type oxides such as Nickel oxide and Tin (II) oxide grown on Ga₂O₃ to form heterojunctions[3][4]. However, resulting interface defects and grain boundaries decrease the electrical performance of these devices which directly affects the power device performances, such as breakdown characteristics, on-resistance, and mobility. Hence, the development of high-quality heteroepitaxy of a *p*-type layer with low structural defects on *n*-type Ga₂O₃ is essential to improve device performance in Ga₂O₃-based bipolar devices. For operation at high temperatures, thermodynamically stable interfaces are also critical. Recent observations show that NiGa₂O₄ forms as a thermodynamical reaction product between Ga₂O₃ and NiO at the *p*-*n* heterojunction interface during high-temperature operation. Hence the possibility of developing a *p*-type NiGa₂O₄ on Ga₂O₃ can circumvent this interface reaction and lead to the development of thermodynamically

stable high-temperature devices. In this work, we demonstrate the epitaxial growth of wide bandgap *p*-type NiGa₂O₄ thin films on Ga₂O₃ and the device performance of vertical *p*-*n* heterojunction diodes processed using these heterostructures. Undoped NiGa₂O₄ thin films were grown on three different orientations of β -Ga₂O₃ wafers and on reference Al₂O₃ substrates by pulsed laser deposition. Structural characterizations of the NiGa₂O₄ thin films show that 002-oriented NiGa₂O₄ grows epitaxially on β -Ga₂O₃ (100) while NiGa₂O₄(220) was stabilized on β -Ga₂O₃ (010) orientation. But thin films of NiGa₂O₄ grown on Ga₂O₃(001) were polycrystalline. The reflection high energy diffraction (RHEED) patterns during growth were streaky indicating relatively flat surfaces. A bandgap of ~3.95 eV is obtained for NiGa₂O₄ thin films from spectroscopic ellipsometry. The fabricated NiGa₂O₄/ β -Ga₂O₃ vertical *p*-*n* heterojunction devices demonstrated good specific on-resistance, excellent temperature-dependent reverse leakage current, and lower on-voltage compared to widely used NiO-Ga₂O₃ heterojunctions. These performances demonstrate that NiGa₂O₄/ β -Ga₂O₃ *p*-*n* heterojunction diodes can be promising for high-power devices with low-on-state power dissipation capable of operating in extreme environments

10:25am **PCSI-ThM1-24 Quantum Oscillations in GaN/AlN 2D Hole Gas and Extraction of Light Hole Effective Mass**, **Chuan Chang**, *J. Dill*, *Z. Zhang*, Cornell University; *S. Crooker*, *O. Valenzuela*, *R. McDonald*, Los Alamos National Laboratory; *D. Jena*, *G. Xing*, Cornell University

Gallium Nitride (GaN) has been a leading contender in commercial high-frequency and high power applications due to its internal polarization field and a wide bandgap of 3.4 eV [1]. However, while *n*-channel high-electron mobility transistors (HEMT) based on GaN's polarization-induced two-dimensional electron gas (2DEG) progress towards higher performance, its *p*-type counterpart has been lagging due to the low mobility of the polarization-induced two-dimensional hole gas (2DHG), hindering the development of GaN-based CMOS and an extraction of hole effective mass by Shubnikov de-Haas (SdH) oscillations and cyclotron resonance. In the absence of reliable and uniform experimental data, researchers have had to rely on theoretical calculations [3]. In this talk, we report the first observation of SdH oscillations in any *p*-type GaN platform and subsequent extraction of hole effective mass. Here, a technique pioneered by Chaudhuri et. al. [2] is used to form a high-density (~5×10¹³ cm⁻² at 300 K) 2DHG at the heterointerface between GaN and an AlN substrate via the large internal electric fields induced by spontaneous and piezoelectric polarization. Magnetoresistance measurements up to 63 T is performed at the National High Magnetic Field Laboratory Pulsed Field Facility, showing

Shubnikov de-Haas (SdH) oscillations with an onset at around B=25 T (Fig. 1). Fig. 2a shows R_{xx} with a polynomial background subtracted plotted against B-1 and Fig. 2b shows its power spectrum. A strong peak is located at f =168 T in the power spectrum at all temperatures corresponding to a density of 8.2×10¹² cm⁻². In lower magnetic fields (< 9T), R_{xx}(B) and R_{xy}(B) are fitted to a classical two-band model (Fig. 3), revealing the coexistence of two carrier populations – low-mobility (~230 cm²/Vs) heavy holes with a density of 4.2×10¹³ cm⁻² and high-mobility (~1400 cm²/Vs) light holes with a density of 7×10¹² cm⁻² in agreement with the density extracted from SdH frequency. Attributing the oscillations to the light holes, we extract their effective mass from the temperature dependence of the amplitudes, yielding a value of 0.48 ± 0.02 m₀.

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PCSI

Room Ballroom South - Session PCSI-ThM2

2D Materials and Graphene II

Moderator: Scott Crooker, Los Alamos National Laboratory

11:00am **PCSI-ThM2-31 UPGRADED: Reduced Metal Contact Resistances for Moire MoS₂ Interfaces**, *John Robertson*, Cambridge University, UK
We show how rotational Moire interfaces for electrical contacts between metals and monolayer MoS₂ can create weakly bonded physisorptive interface sites with weaker Fermi-level pinning. This creates smaller n-type Schottky barrier heights, giving the lowest contact resistances for In and a noble metal Ag, as seen experimentally, but previously unexplained. Analogous sites are found for p-type interfaces on WSe₂.

Scaling of semiconductor devices requires lower contact resistances by reducing Schottky barrier heights (SBH) for metals on transition metal dichalcogenide (TMD) contacts. Duan [1] achieved unpinned Fermi levels and physisorbed interfaces by using metal films mechanically transferred onto unmanufacturable exfoliated MoS₂. This effectively increases the interfacial bond length and makes physisorbed interfaces [2]. We suggest longer bonds can be formed by making Moire interfaces between contacts and TMDs.

The TMD lattice allows an alternative way to create longer interfacial bonds using Moire interfaces. These apply a rotational twist between MoS₂ layer and the metal contact layer [2]. There are three types of interface, on-top site (T), a hollow site (H), and Moire sites (M), Fig. 1. We calculate the interfacial binding energy of each metal interface, and find the most stable configuration, as a function of metal work function ϕ . The data shows two zones of physi- or chemi-sorptive interfaces. We then calculate SBHs for the various bonding sites for each contact metal and show these as a function of ϕ . The previous scattered pattern of SBHs sorts into two trends; most SBHs have a slope with ϕ of 0.24 for T or H interfaces. But Moire sites have a clear depinning trend for E_F, with the slope increasing to 0.37 (Fig. 2). This gives a small n-SBH to the MoS₂ conduction band, and lower contact resistances for In and Ag, as seen experimentally. Similar results are found for p-contacts on WSe₂.

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11:20am **PCSI-ThM2-35 UPGRADED: A Generalized and Modular Approach to Tunnel-Junction Spectroscopy for Quantum Systems**, *M. Kavand*, *Z. Phillips*, *M. Hamilton*, *E. Perez-Hoyos*, The Ohio State University; *D. Freedman*, Massachusetts Institute of Technology; *M. Flatté*, University of Iowa; *J. Gupta*, *Ezekiel Johnston-Halperin*, The Ohio State University

We present a generalized and modular scheme for tunneling spectroscopy of 0D quantum systems based on the exfoliation and stacking of 2D heterostructures. In this scheme, layers of graphene/graphite (gr) and hexagonal boronitride (hBN) are assembled into a gr/hBN/hBN/gr tunnel junction. The differential conductance (dI/dV) of this structure is sensitive to both direct tunneling through the insulating hBN and resonant tunneling through any impurity states within the bandgap. As a proof of principle, we demonstrate the ability to resolve a variety of structural defects in hBN as well as the direct observation of the HOMO and LUMO states of vanadyl phthalocyanine (VOPc) encapsulated at the interior hBN/hBN interface of the heterostructure (Fig. 1). The VOPc tunneling spectra directly correlate with scanning tunneling microscopy (STM) of witness samples and are consistent with density functional theory (DFT) of VOPc. This technique is extensible to a wide variety of 0D systems encapsulated at the hBN/hBN interface, including electrically (or redox) active molecular systems, adatoms, and point defects in 2D materials. This generality and flexibility provides an exciting opportunity for both electronic/structural characterization of these quantum states as well as potential applications in quantum information.

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