The reliability of organic depth profiling was tested recently in two VAMAS interlaboratory studies. Results from these studies highlight the rapid developments that have recently been made. The application of this technique to medical devices and biological materials will be reviewed and the remaining challenges described.

The emerging new field of “live-cell nanoscopy” has revolutionized the way biologists explore the living cell to molecular resolution. Whereas far-field fluorescence nanoscopy enables to study the nanoscale localization and dynamics of biomolecules in cells, recent developments in atomic force microscopy (AFM) techniques offer unprecedented opportunities for imaging the supramolecular organization of cell surfaces, and for probing the functional properties and interactions of their molecular machineries. In the past few years, AFM-based nanoscopy has enabled key breakthroughs in cell biology, including deciphering the nanoscale architecture of cell surfaces and their remodelling upon changing the cells functional state, understanding cellular mechanics and its functional implications, quantifying cell adhesion forces contributing to processes like tissue development and bacterial infection, unravelling the molecular elasticity of cellular proteins such as sensors and adhesion molecules, and elucidating how cells reassemble membrane receptors into nanodomains and modulate their functional state. In this talk, I will provide a survey of the recent work we have done using the AFM multifunctional toolbox, emphasizing its potential for understanding cell surface properties and interactions on the nanoscale.

References
Nat. Methods, 8 (2011), 123-127.

The biointerface and related surface science ideas have had profound impact on biomaterials science since at least the 1960’s. In the 21st century there is much discussion of tissue engineering, clearly a “3-D” phenomenon. This talk will review some history of the biointerface, illustrate modern trends, and show how biointerface ideas can be applied to tissue engineering and 3-D scaffolds. Techniques such as electron spectroscopy for chemical analysis (ESCA) and secondary ion mass spectrometry (SIMS) will be featured. SIMS has proven to be powerful for cell identification in culture dishes and for the analysis of decellularized extracellular matrix scaffolds. SIMS also provides much information on synthetic scaffold materials. There are now profound challenges in interpretation of complex spectra and in gleaning useful, biomedically relevant information from complex data sets. Finally, this talk will discuss healing, biointegration and regeneration, particularly in the context of new scaffolds made by a sphere-templating process.

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References
Nat. Methods, 8 (2011), 123-127.
8.20am AS-MoM1 Characterization of Nano-structures from Analysis of the XPS Background: Automation and 3D-imaging. S. Tougaard, University of Southern Denmark

The XPS peak intensity and its background of inelastically scattered electrons varies strongly with the depth distribution of atoms. This phenomenon is the basis for a widely used method for nano-structure analysis. With the rapid increase in the application of XPS, there is a growing need for automated XPS analysis. To meet this demand, a modified simpler and less accurate algorithm, which however is suitable for automation was developed. For each XPS peak, this algorithm determines the approximate depth of the atom’s. The validity of the algorithm was found to significantly improve the 3D images of a thermally patterned 3D XPS imaging.

9:00am AS-MoM4 Corrections for Backscattering Effects in Quantitative Auger Analyses. A. Jablonski, Polish Academy of Sciences, Poland. C.J. Powell, National Institute of Standards and Technology

The backscattering factor (BF) has long been recognized as an important matrix-correction factor for quantitative AES. The BF definition (“fractional increase in the Auger current due to backscattered electrons”) [1] has been shown to be unsatisfactory since the “fractional increase” can be negative at low primary energies and more grazing-incidence of the primary electrons [2]. ISO/TC 201 is considering the definition of a new term, the backscattering correction factor (BCF), as a “factor equal to the ratio of total Auger-electron current arising from ionizations in the sample caused by both the primary electrons and the backscattered electrons to the Auger-electron current arising directly from the primary electrons.” NIST recently released a new BCF Database [3] for AES. This database provides BCFs from Monte Carlo simulations for two models, a simplified model based on the previous BF definition which is relatively rapid and an advanced model based on the proposed BCF definition which is more accurate but slower. BCFs can be determined for a solid of arbitrary composition, a user-specified instrumental configuration, and primary energies up to 30 keV.

Examples will be given of the dependence of the BCF for Pd Mn2N3 Auger electrons on various instrumental conditions (primary energy, primary-beam angle of incidence ($\theta_0$), and analyzer acceptance angle) [4]. BCFs calculated from the advanced model of electron transport in the surface region of the Pd sample varied slightly at various analyzer acceptance angles for $\theta_0 = 0^\circ$ but not for $\theta_0 = 80^\circ$ where there were BCF differences varying between 19% at a primary energy of 1 keV and 6% at a primary energy of 5 keV. These BCF differences are due partly to variations of the density of inner-shell ionizations within the information depth for the detected Auger electrons. The latter variations are responsible for differences larger than 10% between BCFs from the widely used simplified BCF model and those from the more accurate advanced model for primary energies less than about 5 keV for $\theta_0 = 80^\circ$. These and other BCF differences indicate that the simplified model can provide only approximate BCF values. In addition, the simplified model does not provide any BCF dependence on Auger-electron emission angle or analyzer acceptance angle. Comparisons will also be made with measured BCFs for elemental solids.


Monday Morning, October 31, 2011
align extracted electrical behaviors and chemical information from the sample. For a better understanding of a realistic performance (or failure) of a device during operation, we have extended our measurements to a CdS-based photoresistor, where we have mapped the position of the CdS peak over an area of 5 mm x 5 mm while the device is under operation without and under photoillumination. This method allows us to detect defects and malfunctioning sites/domains. We will present our methodology with experimental results and an electrical circuit model.


XPS has long been a primary technique for the analysis of material surfaces, providing quantitative analysis of elemental atomic concentrations and chemical bonding states. When combined with ion sputtering, it has provided a similar level of detail in depth profiling of metals, semiconductors, and inorganic compounds; however, the damage produced by ion bombardment precluded the depth profiling of organic and polymeric specimens. More recently it has been shown that sputter etching by carbon cluster ions significantly reduces the apparent damage to such materials, presumably the result of a combination of a reduced damage layer thickness and efficient removal of the damaged material. In this work we have used a coronene (C28H16) ion source to sputter etch a series of organic compounds and polymers, including trichloro, poly(aniline), poly(glycidyl acrylate), poly(ethylene glycol), poly(ethyl methacrylate), poly(acrylonitrile), poly(ethylene terephthalate), poly(vinyl chloride), and polytetrafluoroethylene. Specimens were sputtered to a steady-state condition and analyzed by XPS. Elemental composition and chemical bonding were compared to the original surface and the theoretical values. These data will be used to refine our understanding of cluster ion bombardment effects in polymers, and to interpret data from the analysis of future materials and devices.

11:00am AS-MoM9 Using A C60 Ion Source For Routine Surface Chemical Analyses, W. Stickle, M.D. Johnson, D.K. Bilich, HP ADL Corvallis, C. Knutson, W. Wang, W. Cowell, Oregon State University

C60® ion sources are becoming one of the routinely used tools in the surface chemical analysis laboratory. While finding early acceptance and utility in the TOF-SIMS community because of its direct applicability to TOF-SIMS systems, the C60 ion source is now a standard option for photoelectron spectroscopy tools. Routine use of this ion source requires characterization and understanding of the surface that is being created during the ion milling process. Such characterizations include the shape and roughness of the resulting surface, changes in chemistry of the surface created by the ion milling process and changes in the mechanical characteristics of the surface that might influence the interpretation of the surface analytical results. Full characterization requires the preparation of standards and also validation of the measurements. The characterization of the application of the C60 ion source for a variety of common polymers such as PMMA, SU8 and polycarbonate are discussed. Further, anticipating the examination of mixed polymer systems, the application of the C60 source for the characterization of inorganic materials such as titania, hafnia and alumina multilayered structures as well as metal-metal oxide layered structures is presented where sputter induced chemistry from the C60® source is observed.


Many technologically important devices require nanometre level chemical characterisation over areas of several square mm, or even cm, for applications such as coating continuity and integrity or contamination monitoring.

The traditional approach when employing XPS for this application has been to acquire a number of small area spectra at different points. The advent of quantitative parallel XPS imaging introduced the possibility of faster acquisition times and higher spatial resolution than the traditional small XPS probe approach.

Here we present an alternative approach whereby multiple features are simultaneously analysed over a large area using XPS imaging. Acquisition conditions and post acquisition data treatment to optimise this approach on a range of samples will be discussed

This approach provides both a faster and more easily interpreted graphical method for multiple feature analysis.

We present examples of this approach on a variety of samples.
Fusion of lipid vesicles and cells is a natural process which takes place in eukaryotic cells. It is a vital process, since it enables cells to exchange material with the outside, both via vesicle content release and through delivery of e.g. membrane proteins to the outer cell membrane. Exocytosis is still not fully understood and although attempts have been done to deliver membrane constituents to supported lipid bilayers, improvement in e.g. efficiency remains. In order to gain a deeper understanding of the mechanics of membrane fusion as well as improve the delivery of functional membrane constituents including complete cell membrane fragments, to supported lipid bilayers, we have in this work developed two novel and powerful techniques.

To mimic exocytosis, we use giant unilamellar vesicles (GUVs) as a model of the cell membrane, cholesterol-DNA[1-3] as a mimic of the SNARE-proteins and small unilamellar lipid vesicles filled with easily oxidized catechol to represent the cellular vesicles. We probe the fusion process using sum-frequency generation (SFG) spectroscopy, detecting the released catechol. By building on this advanced but yet controllable model system of exocytosis, we believe that a wide range of studies can be made in order to decipher the process of exocytosis. Future applications are delivery of e.g. membrane proteins to GUVs, as well as for vesicular drug delivery to cells.

In order to deliver membrane constituents to SLBs, we use a controlled bulk flow through a microfluidic channel to move the front edge of a supported lipid bilayer and fuse it with vesicles adsorbed in front of it. The membrane constituents of the vesicles are efficiently incorporated into the supported lipid bilayer and can be manipulated in 2D (accumulated and separated) by again using the bulk flow. We show that this method is perfectly compatible with cell membrane fragments derived directly from 3T3 fibroblast cells[4]. The method enables studies of e.g. receptor-ligand interactions as well as membrane protein separation in a native environment.

Simonsson et al., Submitted


Biomineralization proteins act as nature's crystal engineers and adsorb onto crystal surfaces with high binding affinity and precision using specific substrate-surface binding motifs. Owing to the importance of the underlying physiological processes and a general interest in biomineralization mechanisms, the binding of regulatory proteins has attracted significant interest. We have studied statherin, which regulates the growth of hydroxyapatite (HAP) in bone and tooth enamel and prevents the buildup of excess HAP by inhibiting spontaneous calcium phosphate growth. A detailed understanding of the underlying molecular recognition mechanisms would help bioengineers and scientists to devise new biomimetic approaches for clinical applications and biomimeral nanofabrication. Sum frequency generation (SFG) spectroscopy can probe protein orientation and secondary structure at the solid-liquid interface and we have recently shown that it can address specific protein regions with atomic resolution when combined with isotopic labeling and solid state NMR (ssNMR) data.(1) We have combined both techniques with near edge X-ray absorption fine structure (NEXAFS) spectroscopy to characterize the structure of the binding domain of statherin, SN-15, on HAP. Protein adsorption was verified using XPS and ToF-SIMS. NEXAFS N K-edge spectra clearly show that hydrogen bonding is important for the binding of both peptides. SFG confirmed an α-helical secondary structure of SN-15 on HAP with the helix axis parallel to the surface. Deuteration was used to specifically probe the orientations of all hydrophobic side chains (leucine, isoleucine, phenylalanine) with SFG in situ. The labeled 12O' from the surface normal (pointing towards the surface) and isoleucine was tilted 5° from the surface normal. We also used fluorine labels to probe individual phenylalanine rings with NEXAFS spectroscopy. Measurements of ring orientations in combination with ssNMR surface distance and rotamer dynamics data allowed us to develop a clear picture of the side chain structure.


10:00am BI-MoM6 Structure and Function of von Willebrand Factor on Synthetic Surfaces and Collagen. E. Hillemeijer, O. Yakovenko, R. Pensala, W. Thomas, D.G. Castner, University of Washington

von Willebrand Factor (VWF) is a soluble clotting protein responsible for binding platelets through the glycoprotein 1b platelet receptor. VWF can become activated and bind platelets when bound to collagen in an injured blood vessel or under increased shear. VWF can also bind platelets when adsorbed to synthetic surfaces, specifically biomaterials. There is evidence that surface characteristics influence VWF adsorption. Previous AFM studies of VWF adsorbed to hydrophilic (mica) and hydrophobic (octadecyltrichlorosilane modified glass) surfaces showed differences in adsorbed topography. However, studies were not performed to relate adsorption differences to VWF function.

Previous studies in our lab have shown differences in function and structure of the platelet binding domain of VWF (A1 domain) when adsorbed to three surfaces: polystyrene (PS), tissue culture polystyrene (TCPs), and glass, with A1 most active when adsorbed onto PS. A1 function was tested by measuring platelet binding. Flow A1 was investigated using time of flight secondary ion mass spectrometry (ToF-SIMS) and binding of conformation-dependent antibodies in ELISA assays. In the work presented here, we have used surface analysis techniques to obtain greater detail about structural differences of the VWF A1 domain adsorbed onto synthetic surfaces. We used near-edge x-ray adsorption fine structure (NEXAFS) to examine the amide backbone, corresponding to the π* feature of the nitrogen edge NEXAFS spectrum. Differences were observed in the angle dependence of the spectra when A1 was adsorbed onto PS, TCPs, and glass, indicating significant structural differences in the protein when adsorbed onto different surfaces. Sum Frequency Generation (SFG) was also used to probe the structure of the amide backbone using amide I spectra.
In addition to examining VWF on synthetic surfaces, we have used ToF-SIMS to obtain structural information about A1 bound to collagen, as well as determining effect on cell-surface interactions. Principal of ToF-SIMS data showed differences between A1 bound to collagen and A1 adsorbed directly onto polystyrene. This suggests that A1 adopts different conformations on the natural versus synthetic substrates, potentially leading to different mechanisms of activation.

In this work, we use surface analysis tools to increase our understanding of VWF behavior, both on synthetic surfaces and in complex, layered protein systems. Increasing our fundamental knowledge of VWF can improve our understanding of VWF interactions with biomaterial surfaces, as well as thrombosis during injury.


10:40am Bi-MoM8 ToF-SIMS Study of Orientation of FNIII10 Fibronectin Fragment on Self-Assembled Monolayers. L. Árnadóttir, L.J. Gambale, University of Washington

Fibronectin (FN) is an extra cellular matrix protein that is involved in many cell processes such as adhesion, migration and growth. The orientation and conformation of FN adsorbed onto surfaces can therefore have a critical effect on cell-surface interactions. Experimental orientation studies of model systems also provide an important benchmark for molecular simulations and are of great value for further development of force fields used in many theoretical studies of protein adsorption. In this study the adsorbed orientation of the 9-10 fragment of FNIII10 was studied on three different model surfaces (self-assembled monolayers (SAM) of C1, alkanethiols on Au, -CH2, -NH2, and COOH terminated SAMs.) The FNIII10 fragment includes the two binding sites for the FN protein, a RGD site on segment 10 (seg10) and a synergy site on segment 9 (seg9). This fragment also has a significant asymmetry in its amino acid distribution with His found exclusively on seg9, Lys only on seg10 and three times more Tyr on seg10 than seg9. Taking advantage of this asymmetry, we use time-of-flight secondary ion mass spectroscopy (ToF-SIMS) to study the different orientation on different surfaces and X-ray photoelectron spectroscopy (XPS) to determine the difference in surface coverage. While CH3 and NH2 have similar full coverage the highest COOH coverage is about half the maximal coverage observed for the two other surfaces. A ToF-SIMS comparison of the FNIII10 on COOH and NH2 at similar coverages shows significantly more His on the COOH and more Tyr on the NH2 surfaces suggesting an opposite orientation of the fragment on these two surfaces. Results indicate that on the COOH surface the fragment is oriented with seg10 down while on the NH2 surface seg9 is closer to the surface. Comparing similar coverages of the FNIII10 on NH2 and CH3, the later has less Tyr signal (from seg10) suggesting that the seg9 is closer to the surface compared to COOH.

11:00am Bi-MoM9 Single Protein Manipulation with STM. S. Khan, K. Clark, C. Hennekens, E. Rauh, S.-W. Hsu, Ohio University

Scanning tunneling microscope (STM) is not only an instrument to image atomic landscape of material surfaces but also is a tool to manipulate individual atoms and molecules. If STM manipulation and spectroscopy can be applied to individual biomolecules, it will be advantageous for multiple research areas. Here we use a low temperature STM in an ultrahigh environment to image individual protein molecules with molecular resolution on Ag (111) surface at 4.6 K. A-b type amyloid precursor proteins molecules and various single chain proteins were deposited on atomically clean Ag (111) surface. STM images and tunneling spectroscopy enables direct sequencing of amino acid groups in these molecules. Moreover, using a STM manipulation technique of manipulating an attractive tip-molecule interaction, individual molecules were relocated on the surface; thereby their mechanical integrity can be tested. Protein folding is a major issue in biological processes. Here, we will demonstrate that folding of carbon back-bone in these proteins is possible using an STM manipulation procedure. This experiment presents a novel avenue of biological research where sequencing and manipulation may be performed on one molecule at a time. We acknowledge the financial support provided by US-DOE-DFG02-02ER46012.

11:20am Bi-MoM10 DNA Origami from Inkjet Synthesis Produced Strands. I. Saaem, A. Marchi, J. Tian, T. LaBean, Duke University

High-throughput synthesis of quality mixed oligonucleotides (oligos) allows for exhaustive studies of DNA nanostructured material formation. In situ DNA synthesis was achieved with a custom piezoelectric inkjet system using novel inkjet chemistry, on功能alyzed cyclic olefin copolymer (COC) as a chip substrate. When amplified off of the chip via strand displacement amplification (SDA), the oligos were eluted and used directly for constructing multiple DNA origami nanostructures in a one-pot assembly. By performing SDA, each chip may be reused multiple times to produce pools of staple strands for repeated experiments. Two separate DNA motifs were produced from a double-stranded scaffold strand. By redesigning specific staple strands, these structures can be connected to provide a larger nanostructured template.


We found that solid-state electron transfer (Etp) studies are possible across non-covalent linked protein interfaces, between two solid electrodes, using the light-induced proton pumping protein, Bacteriorhodopsin (bR), the electron transfer (ET) protein, Azurin (Az), and Bovine Serum Albumin (BSA). Clear differences between these proteins, where for bR and Az we can show that they preserve their structure in the solid state measurement configuration, were observed, with small tunneling decay constants for all three proteins, suggesting that incoherent transport is the dominant mechanism. Putting our data in perspective by comparing them to all known protein Etp data in the literature, we concluded that, in general, proteins behave more akin to molecular wires than to insulators. An important part of these studies was the modification of the proteins by, e.g., removing or disconnecting the retinal in bR and removing the Cu redox center in Az.

We now report that, notwithstanding the above-noted earlier conclusion on hopping, which was based on the Etp efficiency, Az shows 9.360K higher Etp than its denatured (ET protein). We now also show that changing the back configuration of Cu to Arhenius-like thermally activated Etp, which becomes temperature independent below ~200K, a behavior that apo-Az shares, qualitatively with bR, its variants and with BSA.

This difference between bR and Az leads to the truly remarkable situation that above room temperature the non-ET protein bR shows significantly better Etp than ET protein Az.

Electronic Materials and Processing Division

Room: 210 - Session EM-MoM

Moderator: S. King, Intel Corporation

Dielectrics for Novel Devices and Process Integration

8:20am EM-MoM1 Surface Cleaning and Monolayer Seeding for ALD of High-k Studied by In Situ STM, STS, and XPS. W. Mellett, T. Kent, J. Shen, A.C. Kummel, University of California San Diego

Air exposed III-V surfaces nearly always defects which prevent full modulation of the Fermi level thereby impeding their use in practical semiconductor devices such as MOSFETs. For a high speed device, the air induced defects need to be removed to reduce the trapping states while maintaining an atomically flat surface to minimize interface scattering thereby maintaining a high carrier mobility. For silicon, the only commercial atomic layer deposition (ALD) high-k fabrication process is a replacement gate process to avoid processing induced damage. Surface channel III-V MOS devices can be fabricated with ALD high-K gate-first processes; while ALD is known to greatly reduce surface contaminants, contamination removal is not complete and the order of the surface after ALD cleanup is unknown. Using in-situ scanning tunneling microscopy (STM) and scanning tunneling spectroscopy (STS) the surface morphology of a multipstep process was explored for gate-last unpinning of air exposed InGaAs and InP surface via cleaning with atomic H and ALD nucleating/passivating with trimethyl aluminum (TMA). STM of atomic H cleaned surfaces show the residual temperature and a post deposition anneal are critical to forming surfaces that resemble the decapped InGaAs or a highly ordered InP surfaces. For InGaAs(100), 300K H dosing can produce large multilayer etch features which cannot be annealed out; however by dosing at elevated temperatures these features can be avoided. After H cleaning at 380°C, the surface contains dark features consistent with monolayer etch pits, and these features are reduced by a factor -50 with a post deposition anneal. The H cleaned and annealed surface can be unpinned by a half cycle dose of TMA followed by annealing because it generates an ordered dimethyl aluminum layer providing monolayer nucleation density, and an atomically flat surface, critical for aggressive EOT scaling. For InP, a low dose of H at similar temperatures and post deposition anneal generates a mixed surface reconstruction; however, with a higher cleaned surfaces show the residual temperature and a post deposition anneal around 440°C, the single surface reconstruction is observed. A similar atomic H cleaning and TMA dosing procedure has been demonstrated to produced an ordered passivation layer on air exposed InP(100). It has been shown that for InGaAs and InP an oxide free surface can be achieved with atomic H dosing and annealing, however there is a dependence of surface roughness and defect densities on dosing and annealing conditions. The combination of atomic H cleaning and TMA
PECVD and e-beam evaporation. A combination of electron beam and with UV o-zone and a dilute 10 H2O:1 HCL dip. Samples were heated to spacer pillars. Oxidation and oxide removal of exposed InGaAs was done 420 °C and treated with thermally cracked hydrogen (≈ V:III = 4.0 and 5.6 indicating a group In-rich surface regrowths with V:III ratios of 4.0 and 5.6. AFM showed roughened RHEED images during regrowth showed 4x2 surface reconstructions for Contact resistances were extracted by transmission line measurements at 500 °C with V:III beam equivalent pressures of 4.0, 5.6, and 8.0. After regrowth, shorts over the dummy pillar were removed, and samples were metalized with in-situ deposited e-beam evaporated Ti/Pd/Au and mesa isolated. Contact resistances were extracted by line transmission measurements (TLM).

RHEED images during regrowth showed 4x2 surface reconstructions for regrowth with V:III ratios of 4.0 and 5.6 showing a 4x2 In-rich surface reconstruction. SEM of regrowths at V:III ratios of 4.0 and 5.6 showed no faceting and fill-in to the dummy pillar edge. AFM showed roughened surfaces possibly due to high Si incorporation and lattice mismatch between InGaAs and InAs. Regardless of the V:II ratio during growth, 25 nm thick InGaAs channels showed contact resistances of 190 μΩ while 15 nm thick InGaAs showed contact resistances of 105 μΩ. Metal-semiconductor contact resistances were 2.1 μΩ. Local electrode atom probe shows that the regrowth carries some of the Ga along with creating a varying InGaAs alloy concentration throughout the regrowth.

The continuous need for films with lower dielectric constant, higher strength, and greater etch resistance drives the need to explore new materials. In this paper we present a study of boron nitride and other boron-based materials for multiple applications in semiconductor devices discussing the benefits and integration challenges. As critical dimensions shrink and RC delay increases, the dielectric constant of the interconnect is a continuous area of focus. The current silicon carbide-nitride (SiCN) Cu barrier film has a dielectric constant greater than 5.0 and relatively poor step coverage. Significant advances have been made in recent years to develop a low k, conformal and manufacturable BN thin film for Cu barrier applications. The BN film was shown to have improved leakage, mechanical properties and insensitivity to UV cure relative to SiCN.

Back-end of line patterning has increased in complexity with the introduction of ultra-low k (ULk) dielectric materials. Dual hardmask (HM) patterning scheme eliminates the ULk damage caused by photoresist strip process. The TIN HM has faced challenges in extending below 20nm due to post etch residue and high stress leading to line bending. A boron-based HM material was developed to address these integration challenges. The new material has a low and tunable stress eliminating the line bending concerns. Boron content was optimized for the best selectivity to ULk without impacting the film stress. Significant defectivity and queue time improvement was observed with the boron-based HM due to volatility of the HM byproducts. 9% RC reduction relative to the conventional tri-layer patterning scheme was measured using 45nm 2-metal level electrical test structures. Double or quadruple patterning technique is required for critical dimensions reduction due to the lack of manufacturable EUV lithography. Spacer-based double patterning (SADP) is one of the most adopted process flows to generate one-dimension regular array structures. Its implementation is impacted by the poor step coverage of the conventional PECVD SiN spacer leading to metal line cuts after final polishing step. A low temperature BN film with superior step coverage, minimum pattern loading, good uniformity and low cost was developed for 20nm node and beyond. Its benefit for SADP was verified using a 20nm half pitch logic structure where a 200nm long serpentine yield was improved by 80%.

Evaluation of the boron-based thin films for Cu barrier and patterning applications has shown their potential to replace the conventional materials used today in the logic and memory process flow and thus enabling scaling below 20nm.

Contact resistances were extracted by transmission line measurements (TLM).

InGaAs, respectively. 300 nm of SiO2 and 20 nm of Cr were deposited by PECVD and e-beam evaporation. A combination of electron beam and with UV o-zone and a dilute 10 H2O:1 HCL dip. Samples were heated to spacer pillars. Oxidation and oxide removal of exposed InGaAs was done 420 °C and treated with thermally cracked hydrogen (≈ V:III = 4.0 and 5.6 indicating a group In-rich surface regrowths with V:III ratios of 4.0 and 5.6. AFM showed roughened RHEED images during regrowth showed 4x2 surface reconstructions for Contact resistances were extracted by transmission line measurements (TLM).

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becoming in nano generation in forces to Logic. Logic demands new scheme technology for realizing lower power operation and managing the total power consumption. On the other hands, Memory, especially new volatile memory demands new cell technology for shrinking cell size and realizing high speed programming, low voltage operation and good reliability including endurance. From above viewpoint, we will show the excellent performance of both Logic-in-Memory Architecture [5-7] using MTJ, and MTJ based Vertical structured cell, as follows. First, it is shown that by Logic-in-Memory Architecture using MTJ, a compact LSI with a standby-power-free and immediate-power-up capability can be realized. Next, it is shown that by Vertical structured cell using MTJ, ultra high density non-volatile memory can be realized with utilizing both a capability of Vertical structure MOSFET such as large drive current, excellent gate controllability and compact footprint, and a capability of MTJ such as unlimited endurance and manufacturability integrated in backend metal line of Silicon CMOS technology. Finally, we discuss the impact of spintronic devices for future Nano Si-LSI.


Energy Frontiers Focus Topic
Room: 103 - Session EN+PS-MoM

Plasmas for Photovoltaics & Energy Applications
Moderator: S. Agarwal, Colorado School of Mines


Plasmas for photovoltaics and energy applications are inevitably and thus easily eroded by the reaction catalysts produced for the coal gasification.

9:00am EN+PS-MoM3 Synthetic Fuel Processing through Plasma-Assisted CO2 Conversion, S. Wezel, S. Ponduri, F. Brehmer, M. Creatore, M.C.M. van de Sanden, R. Engeln, Eindhoven University of Technology, Netherlands

Continuously increasing green house gas emissions and forthcoming (fossil) fuel depletion has stimulated research in novel fuel processing, cleaner combustion as well as CO2 capturing and conversion. Conventional fuel processing usually aims at producing syngas (CO/H2) mixtures that may be further converted into value added hydrocarbons and oxygenates (CxHyOz). Photocatalytic CO2 hydrogenation is now considered as alternative approach which would simultaneously lead to a global green carbon cycle. It could reduce atmospheric CO2 concentrations, while at the same time provide fuels on a renewable basis that can directly be supplied to our present energy infrastructure. Since the efficiency of such an artificial photosynthesis is low, we propose the plasma-assisted hydrogenation of CO2 into hydrocarbons.

This contribution focuses on the efficiency of CO2 depletion and selectivity of CH4 production in a low-temperature plasma expansion. The plasma is created from mixtures of argon and hydrogen while CO2 is injected into the expansion part where the dissociation mechanism might be radical- and/or ion-driven. Results on measurements of the (steady state) gas composition obtained by mass spectrometry and mid-infrared tuneable diode laser absorption spectroscopy will be reported. Especially under argon rich conditions, where the chemistry is mainly driven by a combined charge exchange with the Ar ions and dissociative recombination, a CO yield of 50% was achieved. CH4 formation was particularly detected at high hydrogen admixtures. C2Hy hydrocarbons were in most cases absent while H2O and CO turned out to be the main stable products. The results suggest an inherent syngas step during the plasma-assisted conversion approach, particularly a successive hydrogenation of CO.


Currently, in the photovoltaic (PV) industry, wet chemistry based etching is used for saw damage removal and surface texturing. It is known that plasma based dry etching leads to an improved light trapping on multi-crystalline silicon material and, therefore, it has the potential to increase the solar cell efficiency. However, up to now plasma based texturing has not been implemented in the PV industry, because of the very low etch rate (<1 µm/min) and the high cost of ownership. In this study, different front surface textures obtained by means of the Linear Microwave Plasma (LMP, commercialized by Roth&Rau) technique and the high rate expanding Thermal Plasma (ETP, commercialized by OTB-Solar) technique, are investigated to reduce the overall reflection losses of mono-crystalline silicon solar cells and, therefore, has the potential to increase the solar cell efficiency. The different front surface textures obtained by means of the Linear Microwave Plasma (LMP, commercialized by Roth&Rau) technique and the high rate expanding Thermal Plasma (ETP, commercialized by OTB-Solar) technique, are investigated to reduce the overall reflection losses of mono-crystalline silicon solar cells and, therefore, has the potential to increase the solar cell efficiency. The different front surface textures obtained by means of the Linear Microwave Plasma (LMP, commercialized by Roth&Rau) technique and the high rate expanding Thermal Plasma (ETP, commercialized by OTB-Solar) technique, are investigated to reduce the overall reflection losses of mono-crystalline silicon solar cells and, therefore, has the potential to increase the solar cell efficiency.
9:40am EN+PS-MoM5 RF-PECVD Processes Excited by Asymmetric Voltage Waveforms, P.-A. Delattre, S. Poudouque, Laboratoire de Physique des Plasmas, France, E.Y. Johnson, Laboratoire de Physique of Interfaces and Thin Films, France, J.-P. Booth, Laboratoire de Physique des Plasmas, France

Voltage Waveform Tailoring (VWT) is a promising new technique for Radio-Frequency (RF) process plasma excitation. It is known that asymmetric waveforms resembling peaks (short positive, and long negative voltage) in valleys (long positive, short negative voltage) can produce higher voltage self-bias, even in a symmetrical reactor [1], known as the Electrical Asymmetry Effect (EAE). We have implemented a system to provide such voltage waveforms on the RF electrode of our Capacitively Coupled Plasma (CCP) reactor. For a peak to peak voltage (Vpp) of 300 V, we can control the self-bias from -190 V to 15 V, without changing any other process parameter. A new differential RF probe gives us the real-time current and voltage diagnostics of the plasma in the discharge process. For a specific waveform composed of a 15 MHz fundamental and three harmonics, instantaneous power changes from +1 kW to –1kW in 10 ns. Using a hairpin resonator probe in hydrogen at 13 Pa, we have measured an electron density of 28E16 cm⁻³ with a standard sine waveform, 2E19 cm⁻³ with a valleys waveform and 2E10 cm⁻³ with a peaks waveform (all with Vpp= 300V). With a view towards photovoltaic applications, using a gas mixture of 4% of SiH₄ in H₂ at 65 Pa, we have achieved a deposition rate of high-quality amorphous silicon of 1 Å/s for sine, 2.7 Å/s for valleys, and 3.8 Å/s for peaks voltage waveforms.


10:00am EN+PS-MoM6 Spontaneous and High Rate Synthesis of Nanocrystalline Silicon by Expanding Thermal Plasma, I. Doğan, N.J. Kramer, M.A. Verheijen, Eindhoven University of Technology, Netherlands, K. Dohnalova, T. Gregorkiewicz, University of Amsterdam, Netherlands, M.C.M. van de Sanden, Eindhoven University of Technology, Netherlands

Silicon nanocrystals (Si-NCs) draw attention since they exhibit size dependent luminescence, improved charge storing capacity and increased surface reactivity. For instance, the size dependent optical properties of Si-NCs show great promise for increasing the efficiency of solar cells. Si-NCs could be used as spectrum down converters by converting the excess energy of a photon into a carrier which can then generate multiple electron-hole pairs. For such applications, the main issues on Si-NC synthesis are size control and surface engineering for improved optical properties, and high throughput. Among these points, the amount of throughput is highly critical for large scale applications however, it is not possible to achieve with current production techniques. Our research goal is to show that it is possible to fulfill this demand with a novel route by using the remote expanding thermal plasma (ETP) technique. Synthesis of Si-NCs in a remote Ar/SiH₄ plasma occurs by means of a reaction sequence of ion-SiH₄ charge exchange and subsequent addition of SiH₄ molecules. This realizes spontaneous and nearly complete conversion of SiH₄ into Si-NCs with very high throughputs of about 25mg/min, the fastest deposition rate reported in the literature so far. Moreover, ETP has the freedom of tuning the size of the Si-NCs by controlling the flow rates of SiH₄ and Ar, i.e. the residence time in the plasma. Synthesized Si-NCs have a bimodal distribution of small (4-7nm) and large (50-80nm) sizes as confirmed by TEM, which is a consequence of the plasma expansion and reactor geometry. Bimodality has been analyzed with Raman spectroscopy by studying the asymmetry and shift in the transverse optical vibration mode of bulk-Si at 521cm⁻¹. Photoluminescence spectroscopy confirms the presence of monomodal size distribution of small Si-NCs (4-7nm) leading to luminescence in the region 600-900nm. Observation of luminescence illustrates the quality of the nanocrystal surface passivated by the rapid native oxidation. Preliminary results show separation of the bimodal distribution can be achieved by means of geometrical isolation of the regions, in which small and large particles are formed.

10:40am EN+PS-MoM8 SiHₓ and SiFₓ, Dissociation in Matrix Distributed ECR Sources, and Potential for High Deposition Rate of Thin Film Silicon Alloys, S. Kasouti, Total S.A., France, P. Builkin, P. Roca i Cabarrocas, LPICM, France

Deposition at heights and on ever larger areas are important objectives for the reduction of thin film silicon modules costs. High deposition rates have been achieved so far using different plasma sources but uniformity over large areas is still problematic. Matrix distributed electron cyclotron resonance (MDECRL) systems consist of individual ECR plasma sources, which can be arranged in arrays with virtually no size limitations. Deposition rate of silicon alloys exceeding 10 nm/s, has been demonstrated, but little is known so far about the precursors’ dissociation and species fluxes onto the surface.

We study here the dissociation of Silicon film precursors, using optical emission spectroscopy and quadrupole mass spectrometry, both in the plasma itself and in the plasma plume. We correlate the one hand to the electron density and temperature measured using Langmuir probes and microwave interferometry, and to film growth rate and structure on the other. It is found that more than 90% of SiH₄ is used across a wide range of microwave powers, with a small dependence on gas residence time and pressure. Deposition rate depends primarily on the total flow of SiH₄, and values above 3 nm/s are easily obtained. Transition from amorphous to microcrystalline silicon growth, on the other hand, occurs at high powers, and correlates to the electronic temperature and the high H/Si ratio observed in the gas phase. Direct dissociation of SiH₄ by electron impact, followed by deposition from atomic Si and subsequent crystallization by hydrogen is therefore a satisfactory explanation.

On the contrary, for deposition from SiF₄ , consumption is found to be lower than 60%, and independent of the gas residence time and microwave power. The deposition rate is also much lower, on the order of 0.3 nm/s. SiF₄ consumption increases with the total hydrogen flow rate added to the mixture. The total concentration of atomic Si in the gas phase varies linearly with the product of SiF and H species, estimated by actinometry, suggesting fluorine abstraction by hydrogen as a possible mechanism. Deposition rate is also proportional to the Si content in the plasma phase and increases with the hydrogen flow rate added to the plasma. We propose that film growth from MDECR plasma mixtures occurs through the deposition of atomic silicon, obtained from hydrogen abstraction of fluorine in the gas phase, direct electron impact dissociation of SiF₄ playing a smaller role.

This illustrates the importance of gas phase reaction for achieving high deposition rates, even in the case of high density low pressure plasma sources.

11:00am EN+PS-MoM9 The Effects of Showerhead Hole Structure on the Deposition of uc-Si:H Thin Films by VHF PECVD, S.-S. Wi, T.-G. Kim, H.-J. Lee, Pusan National University, Republic of Korea, D. Kim, D. Hwang, W.S. Chang, LG Electronics, Republic of Korea

We presents the characteristics of hydrogenated microcrystalline silicon thin films deposited from SiH₄/H₂ in 40 MHz plasma enhanced chemical vapor deposition (PECVD)equipped with multi-hole-array showerhead. The effects of hole array structure are analyzed in terms of their diameter and depth. Cross dependences between the hole structure and process parameters, such as SiH₄ concentration, rf power, pressure, substrate temperature and total gas flow rate, are also investigated. The results show that deposition rate is not a strong function of hole structure compared with other process parameters. However, it is found that uniformity can be controlled by varying the surface density of hole array. With decreasing total flow rate, faster increase in deposition rate is found at the multi-hole array compared with flat electrode. This may be attributed to the high electron density and longer residence time within the holes. It is demonstrated that the multi hole array electrode can be used as an effective control variable for optimization of Si thin film solar cell PECVD process.

11:20am EN+PS-MoM10 Plasma-Enhanced CVD and ALD Prepared Nanolayers for High-Efficiency Solar Cell Manufacturing, W.M.M. Kessels, Eindhoven University of Technology, the Netherlands

INVITED Photovoltas has become a very innovative field of research and manufacturing due to the continuous improvement in the solar cell cost-performance ratio and its tremendous growth opportunities (past average annual growth rate of 40%). Several innovations with respect to the improvement of the cell efficiency lie in the field of thin film technology, not only for thin-film solar cells but also for crystalline silicon solar cells which are currently still dominating the market (87% market share in 2010). One particular trend is the application of ultrathin films or “functional nanolayers” for solar cell interface engineering. In crystalline silicon technology these nanolayers have as a main application the reduction of charge carrier recombination at interfaces through “surface passivation”. Plasma-based deposition processes such as plasma-enhanced chemical vapor deposition (CVD) are key for the preparation of such films and recently the interest also shifted to (plasma-enhanced) atomic layer deposition (ALD) processes due to their precise growth control and their excellent uniformity and conformity [1]. Moreover, ultrathin films of < 10 nm have been found feasible in terms of reaching very well passivated surfaces. In this contribution, the application of ultrathin films of various materials such as a-Si:H, a-SiNₓ:H, SiOₓ, and in particular Al₂O₃ will be addressed. The preparation methods will be described as well as the relevant surface reaction mechanisms during the film synthesis. Passivation and solar cell results will be presented with a main emphasis on the key mechanisms underlying the good passivation performance of the ultrathin
films. Also the market feasibility of new ALD technologies, e.g., in terms of high throughput processing, will be addressed.


Energy Frontiers Focus Topic
Room: 104 - Session EN-MoM

Industrial Physics Forum on Energy I
Moderator: R.A. Sears, Massachusetts Institute of Technology, B. Clark, Schlumberger

8:20am EN-MoM1 Energy Security and Energy Policy. W.W. Hogan, Harvard University INVITED
Energy security is broader and different than energy independence. Different definitions of energy security produce different policy prescriptions. A consistent framework for energy security must address basic principles.

9:00am EN-MoM3 Technology Innovation and China's Skyrocketing Demand for Energy. E. Steinfeld, Massachusetts Institute of Technology INVITED

9:40am EN-MoM5 Making Energy Sustainable – Scientific Challenges in Determining the Pathways to the Future. E.D. Williams, BP plc, UK INVITED
The scale and cost of the energy challenge are immense. Everyone wants secure, reliable, and affordable energy, but climate change, demand growth and, increasingly, resource scarcity, are transforming the energy landscape and it will continue to evolve.

There are many possible technical pathways to a low-carbon energy future, and each presents unresolved technical challenges that will influence the time, money and global-scale asset and infrastructure deployment that will take place over the next decades. BP’s energy portfolio demonstrates many of the research challenges in this arena, and some examples from Carbon Capture and Storage and Biofuels will be discussed here specifically. Ultimately, research, technology, policies and partnerships will determine the pace of change.

10:04am EN-MoM8 Synthetic Biology for Energy and the Environment. A.A.N. Patrinos, Synthetic Genomics (SGI) INVITED
Synthetic biology is one of the major “tools” that are converting biology from a concept-driven scientific revolution to a tool-driven scientific revolution. This paradigm shift will enable significant applications of the new biology to major challenges in medicine, energy, and the environment.

11:00am EN-MoM10 Manufacturing Innovations for a Sustainable Energy Future. R. Castro, O. Nalamasu, Applied Materials, Inc. INVITED
Nanomanufacturing technology, the cost-effective and practical manufacturing solutions based on equipment and process solution platforms have been translating the promise of nanotechnology to reality in advancing transforming the electronics and display industries. Technology, Scale and Innovation would continue to be fundamental to meet the global inflections associated with Electronics and Display industries and more importantly, advances in nanomanufacturing technology are critical to solving the energy and environment challenges. In this presentation, I will detail the challenges and opportunities in building a sustainable energy future based on nanomanufacturing innovations.

Electron Transport in Low Dimensional Materials Focus Topic
Room: 209 - Session ET+EM+SS-MoM

Quantum Transport: From 0- to 2-Dimensions
Moderator: A.-P. Li, Oak Ridge National Laboratory, K. Varga, Vanderbilt University

8:20am ET+EM+SS-MoM1 Charge and Spin Transports at Surfaces of Strong Spin-Orbit-Coupling Materials. S. Hasegawa, T. Hirahara, University of Tokyo, Japan INVITED
Transports of charge as well as spin at crystal surfaces are now intensively studied by various kinds of experiments. Surface electronic states are generally decoupled from the bulk states and therefore intrinsically low-dimensional. Furthermore, space-inversion symmetry is broken down at crystal surfaces; one side of the surface is empty vacuum while other side is full of electrons in the crystal. These effects provide rich physics of transport, especially on surfaces of strong spin-orbit-coupling (SOC) materials. The surface-state bands are known to be spin-split of such strong SOC crystals such as Bi and Bi alloys, which is called by Rashba effect [1-4]. Similar effect is observed on a special kind of materials called topological insulators such as BiSb, BiSe, and BiTe alloys. Some of them have spin-split Dirac-cone type surface-state bands. This implies that spin-polarized current will flow at the surfaces of such materials.

In my presentation, by using samples of pure Bi [1-4], BiSb [5], BiSe [6,7], and BiTe, I will show that the surface-state bands are really spin-split and the Dirac-cone conductivity is directly measured by microscopic four-point probe method. An on-going project to detect the spin-polarization of surface current by using magnetic tips in a four-tip STM will be also introduced.


9:00am ET+EM+SS-MoM3 Electron Transport in Ferroelectric Domains and Walls. A. Baddorf, Oak Ridge National Laboratory INVITED
Ferroelectric tunneling, where electron transport is controlled by the polarization state, has recently been realized in a number of experiments. Polarization-controlled transport effects have been detected in tunnel junctions, thin films, single crystals and at domain walls. Yet, little analysis of data has undertaken to determine the transport mechanisms and their interaction with ferroelectric fields and domain boundaries involved in switching. We present seminal experimental observations of transport in thin films of Pb(Zr0.2Ti0.8)O3 (PZT) and BiFeO3 (BFO). Earlier we have shown that both materials exhibit pronounced polarization-controlled electroresistance [1]. Temperature and voltage dependence of currents are not well fit by any one standard model. Instead a transition between surface and bulk limiting effects is observed. Upon close inspection, I-V curves exhibit a reproducible region of negative differential conductance associated with ferroelectric switching. Although this anomaly may originate from extrinsic processes, e.g. due to oxygen vacancies or charge injection, we have carried out a series of control experiments on PZT films that unequivocally connect variation of conductance with the size of the polarization domain in the plane of the surface. The I-V anomaly therefore originates from significant conductivity of the domain wall and a relatively slow expansion of the domain following polarization switching. However, our results do not imply simply that transport is through domain walls, but further that nanoscale domains formed by switching have fundamentally different conduction behavior [2]. We suggest that domains formed by tip-applied bias have curved walls and are consequently charged, modifying adjacent material much as charge accumulation modifies a semiconductor. Engineering the ferroelectric domain size produces a tunable conductance reminiscent of analogue memristors, providing a quasi-continuous spectrum of non-volatile resistive states, even though the PZT polarization itself is bistable. Ferroelectric memristive behavior, which based on our measurements is likely to be universal for ferroelectric semiconductors, is a striking departure from the conventional picture of discrete electron transport states in ferroelectrics.

Research was conducted at the Center for Nanophase Materials Sciences and sponsored by the Division of Scientific User Facilities, U.S. Department of Energy.

Monday Morning, October 31, 2011
Quantum transport is at the heart of nanoscience and carries a fundamental law of nature — quantum mechanics — with applied electrical engineering and emerging materials technologies. Ultimately, nanoscale electronic devices will contain networks of wires whose cross sections will be so small as to represent one-dimensional conductors with novel transport properties. We have fabricated exceptionally long and uniform YSi$_2$ nanowires via self-assembly of YSi$_2$ on Si(001). The wire widths are quantized in odd multiples of the Si substrate lattice constant. The thinnest wires represent one of the closest realizations of the isolated Peierls chain, exhibiting van Hove type singularities in the one-dimensional density of states and charge order fluctuations below 150 K. Conduction through individual nanowires follows an inverse Arrhenius behavior, indicative of thermally-assisted tunneling of small polarons between defect centers. Quantitative analysis of individual wire resistances, probe resistances, and negative differential resistances of nanowire networks indicates significant electronic interwire coupling below 150 K. The long-range coupling mechanism involves the dielectric polarization of the substrate, which induces current blockades in neighboring conduction channels.

This work is sponsored by the NIH/NHGRI and was partially conducted at the Center for Nanophase Materials Sciences, which is sponsored at Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy.

We propose a Hubbard model to describe the tunneling effect of electrons in a double quantum dot system connected in the parallel circuit configuration to electrodes. The change in the interdot coupling is shown to dramatically influence the Coulomb blockade properties. For magnetic double dots, the interdot coupling can be tuned by the external magnetic field, leading to a giant Coulomb blockade magnetoresistance. Possible detection of this effect in organic systems is discussed.

This research was conducted at the Center for Nanophase Materials Sciences, which is sponsored at Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy.

**Graphene and Related Materials Focus Topic**

**Room: 208 - Session GR-MoM**

**Graphene Growth**

**Moderator:** D.K. Gaskill, U.S. Naval Research Laboratory

**Tuesday, October 31, 2011 9:40 am**

**ET+EM+SS-MoM5 Electronic Instabilities, Fluctuations, and Transport in Epitaxial Nanowires, H.H. Wettiner**, University of Tennessee at Knoxville and Oak Ridge National Laboratory

**INVITED**

Quantum transport is at the heart of nanoscience and carries a fundamental law of nature — quantum mechanics — with applied electrical engineering and emerging materials technologies. Ultimately, nanoscale electronic devices will contain networks of wires whose cross sections will be so small as to represent one-dimensional conductors with novel transport properties. We have fabricated exceptionally long and uniform YSi$_2$ nanowires via self-assembly of YSi$_2$ on Si(001). The wire widths are quantized in odd multiples of the Si substrate lattice constant. The thinnest wires represent one of the closest realizations of the isolated Peierls chain, exhibiting van Hove type singularities in the one-dimensional density of states and charge order fluctuations below 150 K. Conduction through individual nanowires follows an inverse Arrhenius behavior, indicative of thermally-assisted tunneling of small polarons between defect centers. Quantitative analysis of individual wire resistances, probe resistances, and negative differential resistances of nanowire networks indicates significant electronic interwire coupling below 150 K. The long-range coupling mechanism involves the dielectric polarization of the substrate, which induces current blockades in neighboring conduction channels.

This work is sponsored by the NIH/NHGRI and was partially conducted at the Center for Nanophase Materials Sciences, which is sponsored at Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy.

**Monday Morning, October 31, 2011 10**
Epitaxial graphene (EG) has largely garnered enormous interest, due to its high free-carrier mobility and compatibility with semiconductor processing. Furthermore, EG RF field effect transistors have been demonstrated. Current RF device work has been on the Si face (0001) semi-insulating 6H-SiC substrates as EG on this face mainly consists of mono- and bilayer graphene. In contrast, the C-face consists of up to a dozen or more graphene layers and has a rougher morphology. Even so, there is significant interest in obtaining few layer, smooth EG on the C-face of SiC due to its superior mobility (for similar charge density) as compared to growth on the Si-face. However, the growth mechanism of this material is not well understood. Recently, it was shown that C-face EG grown in an argon ambient slows the growth rate and, under certain conditions, results in localized growth of the graphene on the C-face. These localized areas, referred to herein as graphene covered basins (GCBs), create the possibility of investigating the initial stages and mechanism of graphene growth on the C-face of SiC.

Previously, we had used electron channeling contrast imaging (ECCI) to investigate GCB morphology as a function of GCB size and growth conditions for EG growth on C-face SiC. Threading screw dislocations (TSDs) in the SiC substrate were found to be nucleation sites for GCBs. The TSDs were easily identified at the centers of small EG GCBs (~20 μm diameter). This work shows the evidence that the TSDs fade then disappear with increasing CGB size, suggesting that as the GCBs grow or coalesce to larger diameters the TSDs become buried. Concurrently, Raman mapping experiments determined graphene thickness and quality at GCB genesis; the maximum graphene thickness for TSDs in the SiC by ECCI was also determined. Initial findings with Raman mapping confirm ECCI results showing that the graphene is thicker in the middle of the GCB. The small GCBs (~20 μm), which exhibit a strong TSD signal, are comprised of roughly 3-4 monolayers of graphene in the center, with decreased thicknesses near the edge. In addition, the Raman 2D spectral linewidth for these GCBs correlated with thickness. This was experimentally shown that the group velocity of the bands can be tuned anisotropically by external potentials of a few tens nanometer scale. [1] A three-hold anisotropy of the conical band edge is complex on this polar surface of SiC.

3 J.K. Hite et al., Nano Lett. 11, 1190, 2011.

9:40am GR-MoM5 Graphene Band Engineering on One-Dimensionally-Modulated SiC Substrate, K. Nakatsuiji, T. Yoshimura, University of Tokyo, Japan, K. Morita, S. Tanaka, Kyushu University, Japan, F. Komori, University of Tokyo, Japan

Engineering of the gapless graphene Dirac bands has been studied for the graphene on the 4-degree-off substrate are trigonally warped, and almost agree with the previous reports. On the other hand, the intensity maps for the graphene on the 8-degree-off substrate are significantly elongate in the [1-100] direction. The group velocity of the pi* band in the G-K direction parallel to [1-100] is more than 20% lower than the velocity in the other G-K directions. The velocity reduces in the direction parallel to the substrate slope in contrast to the theoretical prediction.[1]

References

10:00am GR-MoM6 Graphene Growth on Au(111), J. Wofford, University of California at Berkeley and Lawrence Berkeley National Laboratory, E. Starodub, N.C. Bartelt, K. McCarty, Sandia National Laboratories, O. Dubon, University of California at Berkeley and Lawrence Berkeley National Laboratory

Studies on the growth of graphene on metal surfaces indicate that the strength of the interaction between the two materials plays a significant role in determining the evolution and final properties of the resulting film. A number of relatively strongly interacting graphene-metal systems, such as – Ru and –Ir, have been studied comprehensively, but the graphene-Au system remains the only comparatively weakly interacting combination to have been so scrutinized [1]. Comparisons between graphene growth on Cu and Au provide an opportunity to systematically understand graphene growth with both weakly interacting substrates. For example, both Cu and Au have low C solubility, but the mismatch between the lattice of graphene aligned with that of the metal’s (111) surface is substantially larger for Au than for Cu. To examine what effect these differences have, we used low-energy electron microscopy (LEEM) to observe graphene growth on Au(111) in UHV by direct deposition of C from a heated graphite rod. Low-energy electron and electron diffraction (LEED) analysis of the pre-growth Au (111) surface showed the characteristic “herringbone” reconstruction peculiar to Au. Graphene islands nucleate rapidly upon exposure to the C flux, suggesting a relatively low equilibrium C adatom concentration on the otherwise bare Au surface. The graphene islands nucleate simultaneously across the surface with a slight preference for nucleation along Au step edges rather than on terraces. The nucleation density is substantially higher than previously observed on other metals, causing inter-island impingement to begin at sub-micron sizes. Similar to Cu, no island growth due to C precipitation from the bulk of the Au occurs during sample cooling. We find that the graphene lattice prefers strongly to be aligned with the Au lattice with a small minority of the domains rotated by 30 degrees. The prevalence of the aligned graphene orientation is particularly surprising due to the substantial lattice mismatch involved and calculations predicting a 30 degree relative rotation is preferred [2,3]. We draw comparisons between the observed rotational structure and those predicted by first principle calculations.

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Epitaxial graphene (EG) enables wafer-scale production needed to realize graphene-based technologies. Since monolayer graphene is all surface, any impurities or adsorbed atoms can alter the graphene by doping, acting as scattering sites which decrease the mobility, and creating additional resistances that degrade device performance. Also, impurities and resist residues can increase ohmic contact resistance and impact the ability to deposit conformal gate dielectrics. Achieving pristine epitaxial graphene surfaces allows ultimate control over future interface formations and thus improves device performance. In this work, an in-situ H2 anneal is explored to attain the pristine epitaxial graphene required to fully realize the advantages of this material for future electronic applications.
EG was grown on semi-insulating, on-axis, (0001) 6H-SiC substrates in an Aixtron VP508 CVD reactor from 1575 to 1650°C for 60 to 180 min. Initial EG samples were grown at 1000°C for 30 min. Ar/H₂ anneal. Chemical analysis, using x-ray photoelectron spectroscopy (XPS), of the as-grown graphene surfaces showed the presence of both O and excess C impurities. These impurities can dope the EG and reduce the mobility, suggesting the need for methods to obtain pristine surfaces.

To address this need, several post-growth treatments were studied. Unlike exfoliated graphene, uncontaminated EG surfaces could not be obtained using a 400°C Ar/H₂ anneal, and wet chemical cleans reduced the surface impurities. A pristine post-growth EG surface was only accomplished with a 1300°C UHV anneal for 30 min. In-situ UHV anneals are not practical for manufacturing graphene devices, so we explored a new technique incorporating a H₂ anneal during cool down from growth temperature. First, EG samples were cooled in Ar from growth temperature to 100°C. Next, Ar was evacuated and H₂ introduced at constant pressure. Samples were annealed at 1000°C for 30 min., cooled in H₂ to 700°C, and then the chamber was evacuated. XPS analysis showed that the EG samples have no O impurities and reduced excess C impurities compared to Ar cooled samples. Also, the signal from the interfacial layer is reduced, and both the graphene and SiC peaks are shifted to lower binding energies, suggesting a reduction in strain between the EG and the SiC substrate. These samples also had a more inert surface, showing only a small amount of adsorbed O (< 2%) after 3 days in atmosphere. Further, van der Pauw Hall measurements revealed a decrease in mobility over Ar cooled samples with no change in the sheet concentration magnitude.

1. Ishigami, et. al. Nano Letters 7(6) 1643 (2007)


Previous studies have shown that monolayer graphene films can be grown on Cu substrates by the catalytic decomposition of various carbon containing molecules. These films are typically grown on poly-crystalline Cu foils, which have a tendency to recrystallize into a {100} texture during the growth procedure. Since graphene crystalizes in a hexagonal lattice and the {100} surface of a face centered cubic lattice has a square symmetry, this is expected to result in multi-domain graphene growth. Because the Cu(111) surface has hexagonal symmetry and a lattice mismatch of 3.7% with graphene, growth on this surface termination has potential for producing films with a lower defect density.

There have been relatively few studies of graphene growth on single crystal Cu substrates, owing to the fact that hydrocarbon source pressures in the range of 100 millitorr are needed, which is incompatible with most UHV systems. In this study, graphene films were grown on Cu(111) substrates by first preparing the clean surface in UHV by sputtering with inert gas ions followed by annealing. The sample was then transferred to a conventional tube furnace where the graphene film was grown by annealing in forming gas, which reduces the surface oxide, followed by annealing in ethylene to produce the graphene film.

The growth morphology of the graphene overlayer was characterized using low energy electron microscopy (LEEM) and low energy electron diffraction (LEED). The LEEM shows a mesa-like surface texture with relatively flat terraces covered with graphene and valleys between the terraces with low carbon coverage. Micro-Raman spectroscopy of the terrace regions produces a signal characteristic of a well-ordered graphene overlayer, whereas the Raman signal in the valleys is much weaker and shows a distinct D-peak. The typical lateral dimension of the terraces was ~10 μm. Micro-LEED performed on the terrace sites shows a sharp Moiré pattern. Conventional LEED, which probes a lateral area of ~1 mm, shows a ring structure with increased intensity in the six high symmetry directions of the Cu(111) substrate lattice. This indicates that for this growth procedure there is a quasi-epitaxial relationship between the graphene overlayer and the Cu(111) substrate with rotational disorder of the graphene from mesa to mesa. Synchronous-scanning angle-resolved ultraviolet photoelectron spectroscopy (ARUPS) measurements have been performed to probe the electronic band structure of the graphene overlayer. A linear dispersion has been measured in the K direction with the Dirac point located near the Fermi level.

11:20am GR-MoM10 Graphene Growth on Cu(111). S. Nie, Sandia National Laboratories, J. Wofford, University of California at Berkeley and Lawrence Berkeley National Laboratory, O. Dubon, University of California at Berkeley and Lawrence Berkeley National Laboratory, K. McCarty, Sandia National Laboratories

Large area graphene growth on copper foils has attracted considerable interest because of the low cost and high graphene quality. However, relatively little is understood concerning the effect of substrate crystallographic orientation on the morphological evolution of graphene islands. Complication arises by the fact that the foil surface texture depends on how the foil is manufactured. In previous work we have examined graphene growth on the commonly occurring (100) surface of cold-rolled Cu foil. However, growth on the Cu(111) surface, another often observed orientation in textured Cu foil was not studied.

In this work we investigate graphene growth on single crystal Cu(111) in situ using a carbon evaporator in a low-energy electron microscope (LEEM). We find that graphene nucleates at defects and at terraces. A considerable fraction of the islands is misaligned in plane with the substrate, generating rotational boundaries upon inter-island impingement. Islands are dendritic with distinct lobes, similar to those reported on graphene/Cu(100).1 However, instead of each lobe being a graphene sheet with a different orientation, all lobes in an individual island form a single crystal. We propose that these dendritic shapes are the result of diffusion-limited growth. Consistent with this, we find that new island nucleation caused by an increase in the carbon flux occurs equidistant from existing islands. Furthermore, we show that the growth velocity of each lobe is accurately predicted by simulations assuming diffusion limited growth. This diffusion limited growth is in stark contrast with the large carbon attachment barriers seen on Ru and Ir.2 Unlike graphene growth on Ruthenium and Ir(111),3 large densities of carbon adatoms are not detected before island nucleation. Temperature plays an important role in the crystallographic alignment of the graphene film. At high growth temperature (~900 °C), graphene islands are found closely aligned with Cu, while at low temperature (~800 °C), increased disorder in the orientation of graphene with respect to Cu(111) is observed in the low-energy diffraction patterns.

Work at Sandia and LBNL was supported by the Office of Basic Energy Sciences, Division of Materials Sciences and Engineering, U.S. Department of Energy under Contracts No. DE-AC04-94AL85000 and No. DE-AC02-05CH11231 respectively. JMW acknowledges support from the National Science Foundation Graduate Research Fellowship Program.


Graphene has attracted recent interest due to its unique electronic properties. Epitaxial graphene (EG) grown on different planes of 6H-SiC can have different morphological and electrical properties. For example, EG grown on the (0001) plane produces single or bilayer regions, step bunched heights of 5–10nm and lower carrier mobility compared to EG on the (000-1) plane that produces multilayer EG with ridge heights up to 80nm. Step bunching introduces conduction anisotropy and poses a challenge to lithography at ≤100nm. To mitigate step heights and improve carrier mobility, we studied the formation of EG on non-/semi-polar surfaces, specifically the (10-10), (11-20), and (03-312). Here, we show the morphology of EG on these non-traditional planes is improved by an order of magnitude compared to vicinal plane growths.

EG samples (16x16mm²) were synthesized on non-polar (10-10), (11-20), and (03-312) substrates. Growth was carried out in an Aixtron VP508 chemical vapor deposition (CVD) reactor. Prior to EG growth, substrates underwent an in situ H₂ etch at 1520°C for 50min., producing a controlled nucleating surface with average RMS of 0.1nm. Subsequent EG growth was conducted under a flowing Ar ambient of 20 standard liters per minute at 100mbar. Micro-Raman spectroscopy contour mapping was used to confirm the presence, thickness and strain variation of EG. Atomic force microscopy (tapping mode) and scanning electron microscope was used to extract surface morphology variations.

Initial results show that 15min. of EG growth on (10-10) non-polar surface has horizontal wrinkles, indicative of multiple layers of graphene, peak-valley heights ≤5nm, and an RMS of ~0.73nm from a 10x10µm² AFM scan. Growth on (11-20) non-polar plane showed the presence of wrinkles (~3nm in height) and an average RMS of 1.1nm. Both (10-10) and (11-20) plane EG do not show the step bunch and terrace morphology typically associated with polar surfaces. While uniform terraces on the (0001) face can be up to
several µm wide, the (10-10) and (11-20) plane exhibits smaller terrace widths ~75 and ~115nm, respectively. Finally, EG grown on high order (03-312) plane was observed to have wrinkles as well as vertical ridges that were preferentially oriented parallel to each other and an average RMS of 1.17nm. The distance between the successive ridge peaks was ~500nm. Compared to the RMS values of 2.3 and 3.7nm for EG on (0001) and (000 ) planes, respectively, all EG films grown on non-semi-polar orientations were markedly smoother. This work shows that the morphology of EG can potentially be controlled using different SiC orientations.


**In Situ Spectroscopy and Microscopy Focus Topic**

**Room: 106 - Session IS+AS+SS-MoM**

**In Situ Studies of Catalysis and Gas-Solid Reactions**
**Moderator:** G. Rijnber, University of Twente, the Netherlands

8:20am IS+AS+SS-MoM1 In Situ X-ray Studies of Model and Real Catalysts: Bridging the Complexity Gap. A.L. Frenkel, Yeshiva University

**INVITED**

In the last decade, there was a surge in advanced characterization methods to study catalytic materials at work. Most notable innovations in synchrotron-based techniques include the coupling of in situ operando x-ray absorption and scattering methods to vibrational spectroscopies, empowered by improved time and energy resolutions. For example, in situ XAFS-XRD combination enables complementary studies of short and long range order in the same system, a great tool when multiple spatial dimensions evolve in a certain process, such as: modifications of both the catalyst and the support during catalytic reaction, the nucleation and growth of a nano-catalyst, oxidation/reduction of a bulk oxide. Combining in situ XAS or XRD with infrared or Raman spectroscopy is critical for understanding how the structural and electronic properties of a catalyst relate to its reactivity.

Although these are important new improvements in the way we currently study, and understand, processes in nanomaterials, they are done by methods that are not sensitive to local fluctuations in size, shape, structure of nanomaterials, that are present even in well-defined, model catalysts. Thus, in addition to the ensemble averaging that these and other commonly used methods provide, local information, such as one provided by electron microscopy, is needed. In this talk, I will focus on the new efforts in combining the local and average information by coupling the in situ x-ray absorption spectroscopy to in situ environmental transmission electron microscopy (E-TEM) for in situ investigations. Such experiments, done in two separate facilities (NSLS and CFN) at Brookhaven National Laboratory, revealed anomalous, mesoscopic phenomena in the electronic, structural and thermal properties of supported Pt nanoparticles. These systems have long been excellent model systems in catalysis research, yet, at a closer look, as our in situ experiments demonstrated, they turned out to be much more unstable and complex than previously perceived. These clusters exhibited unique physical properties, such as negative thermal expansion in the Debye temperature, broad amorphous-to-crystalline transition zone, large surface strain, as well as charge exchange with support and adsorbates.

I will review recent works showing how such complex behaviors can be, in the case of Pt on g-alumina and carbon supports, theoretically understood by separately studying the effects of their size, shape, support and adsorbates.


9:40am IS+AS+SS-MoM5 New Assignment for Ag(III) from In Situ XPS of Highly Oxidized Silver Films. T.C. Kaspar, T. Drobay, S.A. Chambers, Pacific Northwest National Laboratory, P.S. Bagus, University of North Texas

For decades, it has been a goal to elucidate the mechanisms behind the unique chemistry of both oxygen-exposed silver metal and silver oxides. Silver compounds in bulk, thin film, and nanoparticle form are widely investigated for applications including industrially-relevant catalysis,
Ambient pressure x-ray photoelectron spectroscopy (APXPS) is a method that makes possible x-ray photoelectron spectroscopy (XPS) measurements at pressures of around 0.1 to 10 mbar, the use of a retractable ambient pressure cell. For ambient pressure experiments and science, the instrument is capable of performing XPS measurements on the same sample in both types of environment. This is made possible by a unique design, which is based on the use of a retractable ambient pressure cell. For ambient pressure measurements at pressures of around 0.1 to 10 mbar the cell is docked to the electron energy analyser. Once the sample is loaded the cell is locked, and the only leak to the vacuum is through the nozzle of the analyser’s lens system. Hence, even during ambient pressure measurements the vacuum remains intact in the analysis chamber. For UHV measurements the cell together with the nozzle is retracted into a separate chamber, and UHV XPS measurements can be performed normally. This entails also another attractive feature of the instrument, namely, that the high pressure cell can be replaced by dedicated cells for other sample environments.

In this contribution the design and concept of the APXPS instrument at MAX-lab will be discussed and first results shown. Also plans for an upgraded and dedicated new beamline at MAX-lab will be presented.

In order to shed some light on the state of Pd during complete methane oxidation, we have performed in-situ X-ray Diffraction (XRD) over a Pd(100) surface in a realistic reaction environment combined with DFT calculations. Our study demonstrates that significant roughening of the surface occur during the reaction, which increases the active surface area and thus affects the overall reactivity. Nevertheless, our study strongly suggests that the Pd metal is the most active phase for the full oxidation of methane.

Under realistic or close-to-realistic conditions, while conventional XPS is limited to vacuum conditions of 10^{-6} mbar or better, APXPS thus contributes to closing the “pressure gap” of surface science, which has inhibited the understanding of processes and chemical reactions, for which the chemical potential of the gas atmosphere plays a decisive role. It also renders possible experiments on samples with a large vapour pressure, such as liquids or solid samples with a high degassing rate. Today, there exist a number of APXPS instruments around the world, including a small number of systems that perform well at elevated pressures, but none of them is specifically designed to also allow studies under ultrahigh vacuum conditions. This complicates the connection to results from ultrahigh vacuum studies.

A new instrument for APXPS has just been installed at beamline 1511 of the Swedish Synchrotron Radiation Facility MAX-lab. This instrument, which has been delivered by SPECS GmbH, Berlin, Germany, and which makes use of a PHOIBOS 150 NAP analyser, has been developed with the particular aim of building a strong link between ultrahigh vacuum and ambient pressure experiments and science. The instrument is capable of performing XPS measurements on the same sample in both types of environment.

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11:20am IS+AS+SS-MoM10 In Situ XPS and STM Studies of GeH₂ Interactions with the Si(100) Surface. S. McDonnell, J.F. Veyan, Y.J. Chabal, R.M. Wallace. We present a study of the reactions between Ge₂H₆ and Si(100) surfaces. Ge₂H₆ is a potential precursor that could allow atomic layer epitaxy (ALE) on Ge(100) and Si(100) surface [1,2] which will be a vital component for atomically precise manufacturing (APM). We investigate the effects of various growth conditions such as substrate temperature, dosing pressure and post deposition annealing. We study the formation of seed layers for ALE along with the reactions on both the atomically clean and the hydrogen-passivated surfaces, were we see evidence of Ge₂H₆ reacting with the dangling bonds.

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Ambient pressure x-ray photoelectron spectroscopy (APXPS) is a technique, which dates back to the 1970s and 1980s, but which only during the past ten years has developed a very significant impact, driven forward especially by groups at the Advanced Light Source and BESSY. APXPS makes possible x-ray photoelectron spectroscopy (XPS) measurements under realistic or close-to-realistic conditions, while conventional XPS is limited to vacuum conditions of 10^{-6} mbar or better. APXPS thus contributes to closing the “pressure gap” of surface science, which has inhibited the understanding of processes and chemical reactions, for which the chemical potential of the gas atmosphere plays a decisive role. It also renders possible experiments on samples with a large vapour pressure, such as liquids or solid samples with a high degassing rate. Today, there exist a number of APXPS instruments around the world, including a small number of systems that perform well at elevated pressures, but none of them is specifically designed to also allow studies under ultrahigh vacuum conditions. This complicates the connection to results from ultrahigh vacuum studies.

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Marine Biofouling Focus Topic
Room: 105 - Session MB-MoM

Interfacial Aspects of Marine Biofouling
Moderator: D. Barlow, Naval Research Laboratory

8:20am MB-MoM1 Fouling in the Face of a "Little" Surface Roughness, R. Lamb, A. Wu, K. Cho, H. Zhang, The University of Melbourne, Australia

INVITED
Nanoe- and engineered superhydrophobic surfaces have been investigated for potential fouling resistance properties. Integrating hydrophobic materials with nanoscale roughness generates surfaces with superhydrophobicity that have water contact angles (θ) in excess of 160° and low hysteresis (≤ 10°).

Small angle x-ray scattering (SAXS) was used to investigate the presence of air incursions at the superhydrophobic aqueous interfaces with varying nano/microscale architecture. This technique, sensitive to local changes in electron density, looks at the nanoscale wetting of the rough interface. Three superhydrophobic coatings differing in their chemical compositions and architecture were analyzed using SAXS and tested against major fouling species (Amphora sp., Ulva rigida, Polysiphonia sphaerocarpa, Bugula neritina, Amphibalanus amphitrite) in settlement assays.

Varying extents of attachment-inhibiting properties were observed across the tested coatings and appeared to correlate with the resistance to nanowetting rather than macroscopic contact angle measurements.

9:00am MB-MoM3 Chemistry Depending Surface Conditioning and its Implication for Colonization by Microorganisms, I. Thomé, Karlsruhe Inst. of Tech. (KIT), Germany, M.E. Pettitt, University of Birmingham, UK, S. Kirchen, T. Schwarz, S. Heissler, Karlsruhe Inst. of Tech. (KIT), Germany, M.E. Cullow, J.A. Cullow, Univ. of Birmingham, UK, G. Swain, Florida Inst. of Tech., M. Grunze, A. Rosenhahn, Karlsruhe Inst. of Tech. (KIT), Univ. of Heidelberg, Germany

Biofouling is a ubiquitously occurring phenomenon in tidal zones worldwide [1]. To prevent unwanted effects caused by biofouling, suitable non-toxic coatings for these environments are required. Changing the surface chemistry and the composition of a coating changes not only its properties but also the formation and composition of a conditioning layer. We use self-assembled monolayers (SAMs) on gold as highly controlled surface chemistries which allow to fine tune the physicochemical surface properties. In order to correlate colonization with surface conditioning, we varied the surface chemistry and thus their wetting properties. In agreement with previous work, chemical termination of the surface affects not only the settlement kinetics of spores of the macrofouler Ulva linza [2] but also the settlement of other species. As different SAMs have different affinity towards macromolecules, settlement is controlled by both, surface chemistry itself and an adsorbed conditioning layer. To disentangle both effects, formation of conditioning layers depending on the surface chemistry was investigated in greater detail by spectral ellipsometry and IRRAS. Organism settlement is significantly changed if pristine chemistries are compared to conditioned surfaces.


9:40am MB-MoM5 Probing Molecular Details of Marine Bioadhesion with In Situ Infrared Spectroscopy, A.J. McQuillan, University of Otago, Dunedin, New Zealand

INVITED
The critical step which triggers biofouling at interfaces is the initial adhesion of an invading species to a solid substrate. Of less importance in what occurs thereafter is growth of the species under the influence of nutrients and with the protection of its more sheltered environment. Understanding the factors determining the propensity of species to adhere to substrates is the key to developing new strategies aiming to more effectively inhibit the development of biofouling in many contexts.

Much of the thinking about biofilm formation and the adhesion of biological species to surfaces has been extrapolated from macroscopic observations about the aggregation of colloids. This has been largely based on the interplay between attractive dispersion forces and repulsive electrostatic forces and has given rise to adhesion descriptions in terms of reversible and irreversible stages. A major advance has been the use of in situ atomic force microscopy (AFM) to measure forces during adhesion of microbes to substrates and evaluate their environmental influences.

Nevertheless, this approach is unable to provide in situ molecular details of the chemical components which are suspected to play major roles in adhesion processes.

Vibrational spectroscopy is powerful to reveal the identity and environmental details of molecules in wet interface environments. Vibrational sum frequency spectroscopy and infrared spectroscopy are increasingly thus employed. However, attenuated total reflection infrared (ATR-IR) spectroscopy has the advantages of relative simplicity and greater general familiarity in spite of its use for wet surface situations having only recently been recognised. The ATR-IR approach employs total internal reflection at a high index refraction crystal such as ZnSe or diamond resulting in an evanescent wave sampling a few micrometers of material. In this talk I will outline the principles of the ATR-IR method and how we have adapted them for studies of initial settling of live marine organisms settling onto surfaces under controlled temperature and environment conditions. Observations from recently published work on the settling of Perna canalicula mussel larvae and Undaria pinnatifida kelp spores will be presented and prospects for breakthroughs studies of the settling and propagation of the fleshy seaweed diatom Didymosphenia geminata, invasive to New Zealand, will also be discussed.


Marine organisms attach themselves to a wide variety of submerged surfaces. The barnacle is one of the most pervasive and persistent species to do so, securing itself by forming a thin film of permanent proteinaceous adhesive. For hard-shelled acorn barnacles like Balanus amphitrite, this process involves the recurring sequential release of two major secretions at the adhesive interface. We have shown that as the barnacle grows lateral came of these cement precursor solutions, CPS1, is released on a fairly continuous basis, while the other, CPS2, is released cyclically. By utilizing the differences in secretion patterns, we have begun to deconvolute the contributions of these cement precursors to adhesion. Barnacles were resettled on CaF2 substrates and release of the components at the interfaces was distinguished by optical and fluorescence microscopy. Shear detachment measurements of resettled barnacles showed that the release of CPS2 into the interface corresponded with a roughly twofold increase in adhesion versus CPS1 alone. AFM and FTIR also showed distinct differences in morphology, protein conformation, and chemical functionality for the CPS mixture versus CPS1. Possible ways in which the two components contribute to barnacle adhesion will be discussed based on these results.

11:00am MB-MoM9 Micro to Nanostructured Stimuli-Responsive Surfaces for Study and Control of Bioadhesion, G.P. Lopez, Duke University

This contribution will present recent results on the development and study of bioadhesion on stimuli responsive surfaces that are patterned on lateral length scales of the order of 10 microns and below. These length scales are commensurate with the sizes of the smallest creatures known to be problematic in marine biofouling. Model natural responsive surfaces include patterned polymer brushes and model marine organisms include marine bacteria grown in culture. Our previous studies have demonstrated that stimuli responsive materials can be used to control the adhesion of model marine organisms and this presentation will provide our latest advancements in this line of study, as regards to both molecular and cellular bioadhesive phenomena. We described how to prepare nanopatterned stimuli responsive polymer brushes over areal scales necessary for biofouling studies will be presented, along with characterization of their structure and dynamic behavior.

11:20am MB-MoM10 Surface Topographic Features to Control Biofouling, L. Xiao, University of Heidelberg, Germany, M. Röhrig, Karlsruhe Institute of Technology, Germany, S.E. Thompson, M.E. Cullow, J.A. Cullow, University of Birmingham, UK, A. Rosenhahn, M. Grunze, University of Heidelberg, Germany

Marine biofouling is the undesirable accumulation of microorganisms, plants and animals on artificial surfaces immersed in the sea [1]. The increased hydrodynamic drag caused by fouling leads to higher operating costs of vessels. Studying the interaction between marine organisms and surfaces enhances the development of environmentally compatible approaches to control fouling [2]. Surface microtopography has been found to influence the settlement of cells and larvae [3]. We have studied the influence of surface topographic features on the biofouling process. Honeycomb gradient structures, inspired by the pattern found on the skin of
the pilot whale [4], were obtained by a hot embossing process, and the effect on the density of spores of the green alga Ulva that attached in laboratory assays was quantified. Spore settlement density was higher on the microstructured gradients than the smooth background. The highest density of spores was found when the size of the microstructures was similar to or larger than the size of a spore. With decreasing size of the honeycombs, spore settlement decreased to a level similar to that on the smooth background. In line with the results from the Brennan group [5], spore settlement correlated with Wenzel roughness.


11:40 am MB-MoM11 Biofouling: It’s a Rough Business. A. Wu, R. Lamb, A. McDonald, The University of Melbourne, Australia

The effect of minute changes in nanoengineered superhydrophobic surfaces on the attachment behaviour of several fouling species (Amphora sp., Ulva rigida, Bugula neritina) was investigated.


11:40 am MB-MoM11 Biofouling: It’s a Rough Business. A. Wu, R. Lamb, A. McDonald, The University of Melbourne, Australia

The effect of minute changes in nanoengineered superhydrophobic surfaces on the attachment behaviour of several fouling species (Amphora sp., Ulva rigida, Bugula neritina) was investigated.

Superhydrophobic surfaces were fabricated from latex-templated silica sol-gels. Nanoscale features of the surfaces were varied using two building blocks; silica nanoparticles ranging from 7 – 40 nm and PMMA templating latex ranging from 400 – 800 nm. The combination of these building blocks affords various roughness changes at the nanoscale. In maintaining surface chemistry identical, all fabricated surfaces exhibited superhydrophobic characteristics.

Attachment assays of each surface were conducted and a large variation in attachment-inhibition behaviour was observed. Correlating this behaviour against measured AFM roughness suggests that an intricate relationship exists between surface roughness and attachment behaviour. Data indicate that surface fabricated using small nanoparticles (7 nm) coupled with large templating latexes and high RMS roughness (> 130 nm) exhibited significant attachment inhibiting behaviour. This set of criteria also conforms to the definition of a fractal surface, where self-similarity is present different length scales.

Coupled with previous work that linked pseudo-fractal dimension with nanowetting, this work strongly suggests that resistance to nanowetting has a key influence on attachment of marine organisms.


Nanomanufacturing Science and Technology Focus Topic Room: 207 - Session NM+MS+NS+TF-MoM

ALD for Nanomanufacturing Moderator: B. Lu, AIXTRON Inc.


The demonstrated benefits provided by Atomic Layer Deposition (ALD) in producing films of exceptional uniformity, and conformality, has set the stage for its use in large area, batch processing, and Roll-to-Roll applications. In this work we discuss the use of Computational Fluid Dynamics (CFD) as a means of gaining insight into the system performance of such industrial instruments, but also as a technique for refining system design. Additionally we describe the basic underpinnings of design for ALD systems operated under atmospheric conditions, (for Roll-to-Roll use), along with the design factors which must be considered for zone separated ALD methods. Finally we will present film results taken from a zone-separated ALD system, and discuss the salient aspects of the deposition process.

9:40am NM+MS+NS+TF-MoM5 Improved MOS Characteristics of CeO2/La2O3 and MgO/La2O3 Gate Stacks Prepared by ALD. T. Suzuki, M. Kouda, Tokyo Institute of Technology and AIST, Japan, K. Kakushima, P. Ahnmet, H. Iwai, Tokyo Institute of Technology, Japan, T. Yasuda, AIST, Japan

La2O3 is one of the candidate materials for the next-generation high-k gate stacks because it can achieve sub-1 nm EOT by forming direct-contact La silicate with Si. There have been many ALD studies for La2O3 however, the performance of the MOSFETs incorporating ALD-La2O3 needs much improvement. Our previous studies using EB evaporation showed that capping the La2O3 dielectrics with an ultrathin layer of CeO2 or metallic Mg (~1 nm) significantly improves the channel mobility. In this work we report fabrication of CeO2/La2O3 and MgO/La2O3 gate stacks by ALD/CVD for the first time, and demonstrate that these stacks show improved electrical properties (k value, channel mobility, etc.) as compared to single-layer ALD-La2O3.

The experiments were carried out using a multi-chamber ALD/CVD system which was capable of in-situ metallization and RTA. The CeO2/La2O3 and MgO/La2O3 gate stacks were formed on H-terminated Si(100) using Ce[OCEt2Me]4, La(PrCp3), and Mg(0.8nm)/La2O3(4nm) stack capacitors were approximately 16, which was significantly larger than those for La silicate without any capping layer (k~10–12). The k-value improvement by the CeO2 capping is presumably due to the higher k value of CeO2 (~23), whereas the improvement by the MgO capping is ascribed to suppression of excessive La-silicate formation.

We have also found that the CeO2/La2O3 gate stack leads to excellent mobility characteristics. The mobility for the MOSFET with 1.43 nm EOT was 214 cm2/Vs at an effective field of 1.0 MV/cm, which was 85% of the Si universal mobility. The mobility improvement by the CeO2 capping is attributed to the reduced fixed-charge density, since Vth approached to the ideal values by the CeO2 capping. On the other hand, the MgO capping induced a negative shift in Vth and consistently degraded the mobility. These effects of ALD-MgO capping are qualitatively different from those observed for EB-evaporated Mg [2]. The mechanisms causing such a difference between EB evaporation and ALD are now under investigation. This work was carried out in Leading Research Project for Development of Innovative Energy Conservation Technologies supported by NEDO.


10:00am NM+MS+NS+TF-MoM6 Highly Uniform and Conformal Thin Film Deposition of Metal Layers with Chemical Vapor Deposition. M. Toivola, J. Kostamo, T. Malinen, T. Pili, T. Lehto, C. Dezelaar, Picouson Oy, Finland

Ultra-thin, nanometer-scale metal or metallic films are a crucial component in e.g. several applications of modern MEMS/NEMS (Micro/NanoElectroMechanical Systems) and other advanced IC technologies. Sensors, optoelectronic devices and catalysts are just a few examples. In many of these applications the component sizes keep diminishing and at the same time, the level of system integration increases (for example the so-called “System-in-a-Package” multifunctional chip devices), it creates a drive from “conventional” 2D device architecture to 3D component integration. Through Silicon Vias (TSV) are a central structure in these 3D-stacked devices and there’s a need to produce highly uniform and conformal thin films of metals or otherwise conducting materials on the insides of the vias. Due to the often very high aspect ratio (AR) of the TSV structures, Atomic Layer Deposition (ALD) is one of the only methods with which reliably uniform and conformal material layers can be deposited on the via walls.

Industrially upscaleable ALD processes were developed for several metals and metallic compounds, i.e. Pt, Ir, Ru, Cu, Ag, Au, TiN and TiAlCN. Deposition of metals can be done with thermal ALD and plasma-enhanced (PEALD) the main benefits of the PEALD technology is that we use reductive processes instead of oxygen, lower deposition temperatures which decreases the thermal stress on the substrates, and a wider variety of precursor chemicals.
Inducively coupled remote plasma source system was further developed to reduce any possibility of plasma damage, which can often happen in the more conventional static mode. As a result of our plasma design, the plasma system utilizes highly reactive radicals. Protective flows and separating metal precursor inlets shield the plasma source from getting short-circuited by films from precursor back-diffusion. E.g. N\textsubscript{2}, H\textsubscript{2}, H\textsubscript{2}/Ar, O\textsubscript{2} and mixed gas plasmas can be generated with the system.

Structural design solutions were optimized for ALD reators. Top flow delivery of the precursor gases ensures even distribution of reactive molecules inside the reactor vessel. This is beneficial especially in the case of non-optimal processes with precursor decomposition or etching or poisoning of reactive sites by reaction by-products. Less impurity and thickness gradient scan be achieved with the top flow, compared to the side-flow (cross-flow) design since all the area reacts at the same time leaving less reactive sites left for reaction with the by-products. Therefore, it is possible to get more than 10\% reaction efficiency in ALD reators working with the top-flow design, and also a forced flow for through-portous samples is possible. With modified stopped flow design, extended reaction time inside the chamber can be reached while still keeping the protective flows from the inlets on to prevent any back-diffusion of precursor and subsequent particle formation in the inlet lines.

Upscalable structure was specifically designed to bridge the gap between R&D and production. Smaller ALD tools can be used for process and chemist precursor development at for universities and research centers, whereas the larger, ALD tools can be fully automatized, upscalled and clustered into full scale high volume throughput industrial production unit capable of coating even several thousands of wafers per hour.

**10:40am NM=NS+NS+TF-MoM8** Atomic Layer Deposition for Continuous Roll-to-Roll Processing - S.M. George, P.R. P. Ryan Fitzpatrick, University of Colorado at Boulder

Atomic layer deposition (ALD) is currently being developed for continuous roll-to-roll processing. This development is significant because roll-to-roll processing would allow ALD to address many applications in a cost effective manner. This talk overviews the approaches and progress to date. The original idea of ALD with moving substrates and constant precursor flows was presented in a patent by Santulo and Antson in 1977. This scheme involved rotating the substrate between alternating precursor source regions and vacuum pumping regions. One current approach under development is based on moving the substrate close to a gas source head. The ALD precursors continuously flow through slits in the gas source head that are separated and isolated by inert gas purging. A second version of this design involves using a gas bearing to set the gap spacing between the gas source head and substrate. Another ongoing approach is based on moving the substrate through separate regions of precursor pressure and inert gas purging. Limited conductance between the regions prevents the gas phase reaction of the ALD precursors. The talk examines the issues and prospects for achieving ALD processing.

**11:20am NM=NS+NS+TF-MoM10** High Rate Continuous Roll-to-Roll Atomic Layer Deposition - D. Dickey, Lotus Applied Technology

Atomic Layer Deposition (ALD) is a unique thin film deposition process, capable of producing coatings with unmatched quality and performance. Its unique attributes include high conformality and outstanding thickness precision, enabling the deposition of dense, continuous pinhole-free films, even when extremely thin, and even on highly imperfect substrate surfaces. These attributes have made the process attractive for applications in electronics, semiconductors, and high performance gas diffusion barriers to encapsulate and protect environmentally sensitive devices such as OLED displays and lighting, and CIGS photovoltaic modules. Until recently, ALD films have generally been deposited using conventional static processing, in which the individual precursors are sequentially introduced into and purged from a common vessel containing the substrate. However, this sequence, typically requires at least several seconds and results in the growth of approximately 0.1nm thickness. As a result, the time required to deposit films of reasonable thickness can be quite long. Furthermore, the static nature of the process makes roll-to-roll processing impractical. In this presentation, we discuss the development of a new ALD process based on substrate translation, which can still be called an ALD process, that can be used to deposit films on a substrate without the need for a substrate rotation or translation. This ALD process consists of a series of cycles, each beginning with the transport of the flexible substrate back and forth between the precursor zones. Because a substrate is not required for introducing, saturating, and removing precursors for each cycle, the deposition speed is dramatically increased. In addition, this configuration naturally provides the unique feature of film deposition only on the substrate itself, as it is the only surface which is exposed to both precursors. In turn, this allows the use of steady-state plasma as the oxygen source, enabling a new technique of precursor isolation; “precursor separation by radical deactivation”, in which the oxygen gas precursor actually mixes with the metal precursor, but is only reactive in the region of the plasma source. Together, this technology set has allowed the deposition of high quality ALD films on polymer substrates, including ultra-barrier films, at substrate speeds in excess of one meter per second.

**8:20am NS+EM-MoM1** ZnO Nanowire Logic Inverter with the Difference of Two Gate Electrode - J.K. Kim, Y.T. Lee, R. Ha, H.J. Choi, S.I. Im, Yonsei University, Republic of Korea

Recently, zinc oxide nanowires (ZnO NWs) have attracted much attention for high mobility and sensing properties. These advantageous give us strong possibility to use nanostructures as non-electronic device application; such as field effect transistors (FETs), diodes, and logic circuit devices.[1] In this work, fabricated the logic circuit inverter using difference of gate electrodes which have the different work function.[2]

In order to fabricate the inverter devices, grown ZnO NWs were dispersed to the SiO\textsubscript{2}/Si substrate by using a drop-and-dry method. The Ni/Ti source electrode was patterned by photolithography and liftoff process. To make 30nm-thick Al\textsubscript{2}O\textsubscript{3} gate insulator layer, we used Atomic Layer Deposition (ALD) system. And then, Pd and Ni/Ti top gate electrodes were deposited and these two devices were connected by wire bonding technique. The threshold voltage of the Pd gate ZnO NWs FET shows much positive value (~0 V) than that of the other FET (~ -1 V) with Ni/Ti top gate, and these transistors are able to be used as a driver and a load, respectively. The linear mobility of the driver shows about 119 cm\textsuperscript{2}/Vs at VDD = 0.6 V and the inverter device has high gain value ~15 at VDD = 5 V. Furthermore, the dynamic property of the logic inverter was measured under the 5 V square input voltages.


The geometry of semiconductor nanowires (NWs) allows for both vertical and coaxial heterostructures, while only vertical heterostructures can be formed using planar structures. This is especially important for III-nitride NWs because crystallographic directions in which heterostructures are formed largely determine the magnitude of internal electric fields due to polarization. Here we describe a method to control the relative vertical and coaxial growth rates in catalyst-free GaN/AlN NW heterostructures grown on Si(111) substrates by plasma-assisted molecular beam epitaxy*. A growth phase diagram is established relating NW density to substrate parameters for a model gas source head.

**References**


**9:40am NS+EM-MoM4** Nanowires and Nanoparticles I: Assembly and Devices - M. Hines, Cornell University

**Moderator:** M. Hines, Cornell University
period GaN/AlN (2 nm / 2 nm) superlattices along either the vertical or coaxial NW axis, which exhibit atomically sharp compositional profiles. A coaxial, NW-GaN resonant tunneling diode structure is presented. Large areas of nanowires are processed for electrical measurements without removing them from the Si(111) substrate. Preliminary electrical measurements are provided for both room temperature and low temperature conditions. This work is supported by the ONR under grant N00014-09-1-1153.


9:00am NS+JM-MoM3 INVITED III-V Nanowire MOSFETS, L. E. Wernersson, Lund University, Sweden

In this talk, we will review some of the efforts made in Lund to realize high-performance III-V nanowire transistors using vertical nanowires grown by MOVPE. We will show how bottom-up technologies can be combined with top-down processing to realize nanowire-based RF-devices on Si 2° wafers. We use CV techniques to characterize the properties of the high-k material in vertical nanowire capacitors and compare the data to the 1/nm characteristics of scaled transistors to evaluate the influence of the high-k material on the transistor performance. We also show that the transistor channel may be reduced down to a diameter of 15 nm without degradation of the transport properties. Finally, we explore the use of novel materials in the transistor structures as we developed GaSb/InAs heterostructures with excellent Esaki diode characteristics to be used for TFET implementations.

9:40am NS+JM-MoM5 Optimizing Quantum Efficiency in Quantum Dot Display, S.J. Lim, J. Kwon, Y. Oh, Seoul National University, Republic of Korea; B.L. Choi, K. Cho, Samsung Advanced Institute of Technology, Republic of Korea; Y. Kuk, Seoul National University, Republic of Korea

In our previous study, we were able to fabricate full-color, 4-inch display made of colloidal quantum dot (QD). Despite such a demonstration of QD light emitting device which is one of candidates for next-generation display, understanding the interface characteristics between QD layer and electron (or hole) accumulation layer is still lacking and further study for improvement of quantum efficiency is essential. Here, we report on a study of scanning tunneling microscopy (STM), spectroscopy (STS) and cathode luminescence induced by tunneling current, performed on individually manipulated QDs using STM. We control the distance between two QDs using STM to reveal the mechanism of interaction between QDs. STS measurement showed shift of energy levels as manipulating the distance between two QDs. This result suggests that there exists the optimal distance between QDs for efficient light emission. Besides by making contacts between separated QDs and organic molecules, we simulated contacts between QD layer and electron (or hole) accumulation layer. From these experiments, we could understand excitonic behavior and carrier hopping from QD to QD or surrounding materials. Our findings thus suggest optimal configuration for QD application in display.

10:00am NS+JM-MoM6 Polarization Engineered 1-Dimensional Electron Gas, D.N. Nath, P.S. Park, M. Esposito, Ohio State University; D. Brown, S. Keller, U.K. Mishra, University of California Santa Barbara; S. Rajan, Ohio State University

One-dimensional structures (nanowire) based devices are of great interest due to their promise in high-performance electronics and other future device applications. However, synthesis and patterning of arrays of nanowires is a challenge in all material systems since both bottom-up and top-down approaches have their own merits and demerits. Here we report on the demonstration of pure 1-dimensional arrays of electrons with current density up to 130 mA/mm and carrier confinement greater than 100 meV using lateral polarization engineering in N-polar vicinal GaN/GaN heterostructures. The width of the atomic characteristic of vicinal surfaces defines the dimensions of the nanowires which are found to exhibit sharp and clear signatures of 1-dimensionality at room temperature making them promising for novel device applications. We report on devices fabricated on MOCVD grown N-polar AlGaN/GaN HEMT structures on vicinal sapphire substrate (4° miscut towards a-plane) with anisotropy in current and channel pinch-off voltages. Channels parallel to the miscut direction pinched off at higher negative gate biases than those perpendicular to the step and carried more charge as measured by direction-dependent C-V profiling. An electrostatic model which predicts a saw-tooth energy band profile in the lateral direction has been proposed to explain the charge anisotropy. Each atomic terrace characteristic of the surface morphology of vicinal GaN with its corresponding saw-tooth energy profile is proposed to exhibit quasi-1D confinement. We will discuss the heterostructure/polarization engineering of structures demonstrating pure 1-D transport in direction parallel to steps.

Gated structures were fabricated to investigate the physics of the system as the Fermi occupation function is varied by varying gate bias. To confirm that the carriers are indeed 1-dimensional, we used direction-dependent small-signal capacitance voltage measurements to probe the density of state function and hence dimensionality of electrons as a function of gate bias. We developed a 2-band model consisting of one 1-D and one 2-D subband to describe the behavior of these wires at room temperatures. The variation of capacitance as well as charge density for a pure 1-D and a pure 2-D system as a function of applied gate bias as predicted by our 2-band model based on density of states matches very well with the data measured experimentally for 1-D and 2DEG respectively. This confirms that the channels created are indeed 1-dimensional in nature. Since 1-D channels are atomic terrace defined, they are promising for eliminating the disadvantages of both bottom-up and top-down approaches.


INVITED

Integrating functionalized nanowires directly onto Si CMOS chips has the potential to combine highly selective and sensitive chemical and/or biological sensing capabilities with electronic signal processing in a single ultra compact, low power platform. Conventional integrated circuit manufacturing methods place considerable limits on the range of and number of different materials and molecules that can be incorporated onto Si chips, making it difficult to realize this goal. This talk provide an overview a new deterministic assembly approach that uses electric field forces to direct many different types of bioprobe-terminated nanowires to specific regions of the chip and to provide accurate registration between each individual nanowire and a specific transistor on the chip. This is achieved by synchronizing sequential injections of nanowires carrying different bioprobe molecules with a programmatically controlled electric field profile that directs nanowire assembly. Subsequent back-end lithographic and metal deposition processes are then used to electrically and mechanically connect all of the nanowire devices to the Si chip at the same time. Using this technique, individual nanowire device integration yields exceeding 90% have been demonstrated with a less than 1% mismatch across three populations of DNA-coated nanowires for arrays with densities of 106 cm-2. The nanowire-bound DNA retained its ability to selectively bind complementary target strands following assembly and device fabrication, demonstrating that this process is compatible with these end-manufacturing steps. The uniformity in the electrical properties of device arrays that were fabricated using this hybrid integration strategy will also be discussed.

11:20am NS+JM-MoM10 Solid-State Dewetting of Direct Nanoimprinted Metallic Thin Films, R. Clearfield, North Carolina State University; J.D. Fowlkes, Oak Ridge National Laboratory; P.D. Rack, University of Tennessee Knoxville; A.V. Melchko, North Carolina State University

Heat applied to thin films below a critical thickness will generally cause transformation of the film into isolated particles. This process is known as dewetting. Solid state dewetting occurs below the melting temperature of the film and is governed by diffusive mass transport. Currently two mechanisms of dewetting are distinguished: hole nucleation and growth, and spinodal dewetting. Nanowire dewetting mechanisms via film surface undulations that have characteristic wavelengths related to the thickness of the film. Lithographic patterning of thin films has been utilized to direct the dewetting instability development toward designed nanostructured geometry of nanoparticle arrays. Tailoring the geometry of thin film edge have been shown to affect both heterogeneous nucleation and spinodal dewetting regimes. Nanoimprint lithography, conventionally used for definition of the edges of thin films, is a fabrication method where a stamp is pressed into a thin normally monomer or polymer film at elevated temperatures. Nanoimprinting can also be conducted in direct mode where the stamp is pressed into a metallic film. Surface undulations characteristic for spinodal dewetting will be used to direct the stamp design. Such imprinting allows setting initial conditions, programming instability, in the thin metallic film that is linked to the spinodal surface instability. In this work we are presenting the results of the investigation into behavior of thin
films in which a 3D structure has been imprinted. We present observations on the effect of direct nonlinear lithography on nanoscale Au and Ni films using periodic arrays of cylinders. Our focus is on the spatial distribution of the particles produced from dewetting of the nanoimprinted films. Particles in patterned regions are characterized in terms of their spacing, periodicity and size, and shape. The geometry of the dewetted patterns is compared to that of the 3D features created after direct nanoimprinting of the films. Analysis of spatial correlation of the final dewetted patterns to stamp patterns is presented.

**Plasma Science and Technology Division Room: 201 - Session PS-MoM**

**Advanced FEOL / Gate Etching I**

**Moderator:** A. Agarwal, Applied Materials, Inc.


Plasma etch processes are the only technological solution to address the critical dimension control at the nanometer range imposed by the continuous downsizing of CMOS device dimensions in microelectronics. However, the current processes are reaching their limits of controlling etch selectivity, defects, critical dimensions and uniformity. Lately, pulsed plasma etching has been used to overcome some of these limitations. In this paper, we will show how plasma pulsing can be used to improve the passivation layer, the reactive etch layer, the uniformity and the passivation layer. To investigate the passivation layer at various trench depths and CDs we developed a new XPS technique for quasi in-situ chemical topography analysis.

The experiments are carried out in a 300 mm AdvantEdge™ tool from Applied Materials Inc. The two generators to sustain the plasma (ICP) and to polarize the wafer (CCP), operated at 13.56 MHz, have been modified with the Pulsync™ system to allow pulsing at a wide range of parameters. However, only synchronous pulsing is studied here. Moreover, a Theta300 angle resolved XPS system from Thermo VG is connected under vacuum, allowing quasi in-situ analysis of an etched sample. Several additional diagnostic methods including time resolved UV-absorption, mass spectrometry and ellipsometry are used to study the impact of pulsing the plasma on the surface and the gas phase compared to a continuous wave process.

We demonstrate that plasma pulsing can minimize the mask faceting and consumption which is due to a change in the ion energy distribution and the presence of larger molecular ion species. The duty cycle controls the formation of the passivation layer that governs in turn the form of the etched trenches. The passivation layer can partly be explained by less dissociated oxygen molecules in low duty cycle pulsing. Nevertheless, XPS analyses show that the chemical composition of the passivation layer changes little with height and trench CD while its thickness is directly linked to the aspect ratio of the analyzed part of the sidewall. Less thickness variation with respect to the aspect ratio is observed in pulsed mode, which explains an improved local uniformity in profile evolution. Furthermore, the time compensated etch rate with respect to the actual ON time of the plasma is increasing at low duty cycles due to a change of etch regime from more radical flux limited to ion flux limited. Nevertheless, the overall process etch rate is decreasing in pulsed mode which can be partly counteracted by increasing the source and the bias power.

**8:40am PS-MoM2 Effect of Si Damage on Shallow Source-Drain (SSD) Recess Structures.** J. Ghaa, S. Sriraman, Lam Research Corporation.

Continued scaling in the semiconductor industry provides new challenges for critical Front-end-of-the-line (FEOL) process etch applications in front-end logic devices. One such application that is utilized in the PMOS transistor is the Strained Source Drain recess (SSD) structure that embeds an epitaxial strained SiGe thin film that significantly improves hole mobility in the channel region. Scale down of critical dimensions (CD) in current and future CMOS devices puts ever increasing emphasis in reducing post-etch Si surface damage in a source-drain (SD) recess structures. For a typical SSD application, the roughness of Si surface obtained after SD etch governs both the epitaxial growth of SiGe as well as the roughness of the SiGe layer, and ultimately determines the device performance. This paper will discuss the factors that contribute to the Si surface roughness arising from a representative SD process etch step and its impact on the subsequent SiGe/Al contact and device performance. Typically, this may consist of an anisotropic etch (halogen/oxygen based chemistry) followed by an isotropic etch (halogen/halogen based chemistry). Surface roughness of the etched silicon is quantified and spatially resolved through atomic force microscopy and surface haze measurements, and contributions of the anisotropic and isotropic etch steps to surface roughness are inferred. The effects of halogen ratio and relative halogen atom reactivity in the isotropic etch chemistry on surface roughness and the vertical-to-lateral (V/L) etch ratio in the SSD recess feature will be discussed and a surface reaction model proposed to characterize roughness evolution.


INVITED

The etching processes for integrated circuits fabrication becomes extremely challenging when the devices dimensions are downsizing and the etch process has to stop on the ultra thin layers (less than 2nm) of materials. Perfect control of the passivation layers and mask erosion are required to precisely control the patterns profiles, which necessitates using flows and energies of plasma species in ranges which can not be attainable with current plasma technologies. In addition, limiting the plasma induced damage to thin layers requires very low ion energies which are not accessible in current plasma technologies. One promising solution is to pulse the plasma in conventional industrial reactors at very low duty cycle. Using this solution, the ratio of radicals and ions as well as their energy can be fine tuned in ranges that are unreachable otherwise, and very low energy ions are produced which strongly minimizes the damage to the thin exposed to the plasma.

Experiments are performed on a 300mm diameter Inductively Coupled Plasma chamber commercialized by Applied Materials (DPSII G5) equipped with the pulsync system to enable plasma pulsing in an industrial reactor.

In this paper, we will show how plasma pulsing can be used to improve uniformity and pattern profiles by better controlling the passivation layer formed on the patterns sidewalls during the etching. In addition, we will demonstrate that the reduction of the average ion energy as well as the low use of molecular ions decreases mask erosion, as well as damage to ultrathin etch stop layers. Morphological characterization, surface characterization and plasma diagnostics will be correlated to explain the mechanisms responsible for the process improvement.


Plasma etching of high-k materials, including HfO2, has attracted much attention due to the necessity to integrate these materials in MOSFET transistor. After the metal gate patterning process, the high-k dielectric film must be removed from the source and drain regions of the transistor. It is today well established that HfO2 can be etched selectively towards SiO2 and Si, which allows good plasma etching. High-k dielectric materials are sensitive to plasma induced damages (Si amorphisation and recession) in the source/drain transistor regions in such processes. Since pulsed plasmas have shown a capability to minimize Si-recess in typical silicon gate oxide etching processes, we have investigated their potential for HfO2 etching.

The experiments are performed in a 300mm DPS tool from Applied Materials. The chamber is modified to allow plasma diagnostic like UV-broad band absorption spectroscopy. The reactor is also connected to an Angle-Resolved X Ray Photoelectron Spectroscopy (ARXPS) spectrometer and a calibrated ultrahigh vacuum chamber. The etching rate of HfO2 sample (either 40 or 3.5nm-thick HfO2 films with 8 Å SiO2 interlayer deposited on Si substrates) and of SiO2 samples (10 nm thick) are measured in real time by in situ multi-wavelength ellipsométrie.

We have evaluated the capability of pulsed plasmas to minimize the plasma induced damage of silicon during the etching of HfO2 high-k gate dielectric. XPS analyses show that the perturbation of the bulk Si lattice is less pronounced with pulsed conditions, which is attributed to a lower ion bombarding energy. However, the formation of a BCl polymer selectively on the silicon (which is responsible for the HfO2 to Si etching selectivity) is reduced when the plasma is pulsed leading to a loss of selectivity. UV absorption spectroscopy indicates that the fragmentation of BCl molecules, which leads to the formation of B-rich polymer precursors, decreases dramatically when the plasma is pulsed at low duty cycle. Therefore, as the duty cycle is reduced, the fluxes of B radicals to the wafer are reduced while the amount of Cl in the polymer increases. It follows that the polymer
In particular, Fins for FinFET technology represent the tightest pitch (approximately 40nm for the technology nodes in development at this time) and yield well to Sidewall Image Transfer. We will present the issues surrounding mandated definition and spacer film properties for Fin definition. The Gate level poses competing requirements of overlay control between the simpler patterns and the need to attain multiple Critical Dimensions (CDs) and pitches. We will examine this and the challenges of etch into advanced gate stacks for the 14nm node. At the Interconnect levels, we will consider the challenges of etching not only trenches but also self-aligned via patterns at sub-40nm half pitch into ultra low-k (ULK) dielectrics.


To continue scaling CMOS devices at the traditional pace following Moore’s law, high selectivity of etch processes towards multiple materials is approaching non-dimensional spaces. The spacer etch process is a very critical element in the CMOS device process flow as it ensures and enables the electrical isolation of source/drain and gate regions. Extremely high precision is needed to form a silicon layer that is free from imperfections such as exposed Si, SiO2, or other surfaces (SiGe or SiC for example). This process is even more challenging for non-planar devices (such as FinFETs and Trigates), where the plasma process needs to be able to form the spacer on the gate sidewall, but not the fin sidewall. At the same time the exposed SiO2 and Si surfaces (if applicable) have to withstand the extended processing necessary to form the spacer.

Multiple etch gas chemistries have been evaluated and their impact on etch rates and selectivities for spacer applications have been evaluated. Surface analysis techniques such as XPS and FTIR have been applied in conjunction with OES analysis of the plasma to study the etch mechanisms leading to the observed etch rates. Fundamental differences in etch mechanism were found for different etch gas chemistries. We observed that during conventional spacer processes, very little difference in plasma polymer deposition onto the respective substrates could be noted. A successful SiN spacer process was rather facilitated by a SiO2 chemistry that was able to achieve a conversion of Si to SiO2. This also means that the etch rates of the SiN are limited by the simultaneous oxidation of the same. A potential solution to overcome this limitation would be to control the etch rate by polymer thickness, similar to high selectivity SiO2 etching. An evaluation of this approach has yielded similar results as the general etch mechanism proposed by Schaepkens et al. [1] A SiN spacer chemistry was also evaluated that enables a different etch mechanism that cannot be described by the general model. The impact of the described mechanisms on actual CMOS devices will be discussed in detail.

oxidation is just limited onto the higher radio frequency bias within the above condition. As for the dependence on the trench aspect ratio R, the oxidation at the trench bottom was attained 100% of the trench top up to R=3, while at the trench bottom with R more than 4 it was only 60%. The oxidation at the bottom of deep trench with the aspect ratio R= 6 was improved at the higher radio frequency bias. It is concluded that the oxidation is due to the negative oxygen ions in downstream.

Surface Science Division
Room: 109 - Session SS1-MoM

Water Films & Environmental Interfaces
Moderator: H. Fairbrother, Johns Hopkins University

8:20am SS1-MoM1 Surface Science of Acetonitrile on Model Interstellar Ices and Grains, A. Abdulgalil, M. Collings, M. McCoustra, Heriot-Watt University, UK.

Physical and chemical processes occurring on the surfaces of icy dust grains in the cold, dense interstellar medium have a crucial role to play in the evolution of the modern Universe. The formation of simple hydrides, e.g. water, on grain surfaces explains the infrared astronomical observations of icy grain mantles and provides a reservoir of coolant molecules during the early stages of star formation that helps maintain the current rate of star formation and favours the formation of small, long-lived stars. Energetic processing of icy mixtures by short wavelength radiation and cosmic rays turns the icy grain mantles into chemical nanofactories producing increasingly complex organic molecules. Nitriles (RCN) represent a potential intermediate on the pathway to forming amino acids in these icy mantles. However, the formation of nitriles in and on icy grain mantles is not understood. Nor is the surface science of these simple species on models of the interstellar grain population. Acetonitrile (CHCN) is the simplest organic nitrile and is amongst the list of chemical species observed in the interstellar medium. This paper will report our recent investigations of the interaction of CHCN with model silica and ice surfaces representative of bare and icy interstellar dust grains. We will report on the nature and strength of the interaction of CHCN with these substrates and on the thermal and non-thermal (electron-induced) desorption of CHCN from these surfaces as part of our on-going programme in solid state and surface astrophysics.

8:40am SS1-MoM2 Energetic Xenon Sputtering and Embedding at Ice Surfaces, D.R. Killelea, K.D. Gibson, H. Yuan, S.J. Sibener, University of Chicago

***PLEASE NOTE YOU MUST IDENTIFY A DIFFERENT PRESENTER FOR THIS ABSTRACT. YOU MAY PRESENT ONE PAPER ONLY (ORAL OR POSTER) AT THE CONFERENCE. YOU ARE LISTED AS PRESENTER FOR ABSTRACT #278***Energetic collisions between gas-phase atoms or molecules and ice surfaces are listed as presenter for abstract #278 energetic xenon sputtering and embedding at ice surfaces. The probe depth of the experiment. The result is a depth profile of the incident species. Moreover, metastable absorption states can be observed in XPS. Experiments presented were performed at Beamline U41 at the BESSY II synchrotron facility. By tuning the energy of the incoming photons, the kinetic energy of the photoelectrons is varied, allowing measurements from different depths of solution. As a strong diprotic acid, an aqueous solution of sulfuric acid will have $\text{HSO}_3^-$ and $\text{SO}_4^{2-}$ present, and at high concentrations, (greater than 16M) undissociated $\text{H}_2\text{SO}_4$ is present. In XPS, all of these components are distinguishable using their 52p binding energy shifts. A series of sulfuric acid aqueous solutions with concentrations up to 16M is measured and using the peak areas in the XP spectra, the dissociation at the surface relative to the bulk can be determined for different solutions.

Additionally, the effect of solution temperature on the acid dissociation at the surface is explored.

9:00am SS1-MoM3 In Situ Studies of Sulfuric Acid Aqueous Solutions by X-ray Photoelectron Spectroscopy, A. Margarella, T. Lewis, University of California, Irvine, M. Faubel, Berliner Institut fur Dynamik und Selbstorganisation, Germany, B. Winter, Hemholtz-Zentrum Berlin fur Materialien und Energie, Germany, J.C. Hemminger, University of California, Irvine

Sulfuric acid is amongst the most widely used acids in the chemical industry, as well as having an important presence in atmospheric aerosols. Using a micro-liquid jet, the chemistry at the liquid-vapor interface of aqueous solutions is explored in situ by x-ray photoelectron spectroscopy (XPS). Experiments presented were prefereed at Beamline U41 at the BESSY II synchrotron facility. By tuning the energy of the incoming photons, the kinetic energy of the photoelectrons is varied, allowing measurements from different depths of solution. As a strong diprotic acid, an aqueous solution of sulfuric acid will have $\text{HSO}_3^-$ and $\text{SO}_4^{2-}$ present, and at high concentrations, (greater than 16M) undissociated $\text{H}_2\text{SO}_4$ is present. In XPS, all of these components are distinguishable using their 52p binding energy shifts. A series of sulfuric acid aqueous solutions with concentrations up to 16M is measured and using the peak areas in the XP spectra, the dissociation at the surface relative to the bulk can be determined for different solutions.

Additionally, the effect of solution temperature on the acid dissociation at the surface is explored.


The structure of water at interfaces is crucial for processes ranging from photocatalysis to protein folding. We have investigated the structure of thin water films adsorbed on TiO$_2$(110) using reflection-absorption infrared spectroscopy (RAIRS), temperature programmed desorption and ab-initio molecular dynamics simulations. Infrared spectra were obtained for light with the plane of incidence parallel and perpendicular to the [001] azimuth of TiO$_2$(110) for water coverages ≤ 4 monolayers (ML). The spectra indicate strong anisotropy in the water structure along the two major azimuths. For both 1 and 2 ML coverages, the water films form “strings” of water molecules parallel to the bridging oxygen rows on the (110) surface. Within the strings, each molecule participates in 4 bonds, such that the films have no dangling hydrogen bonds. The vibrational densities of states predicted by the ab-initio simulations for 1 and 2 monolayer coverages agree well with the observations. Despite extensive prior research, the structure of water films on TiO$_2$(110) has remained controversial. Our results provide crucial, molecular-level information about the structure of water films on this benchmark transition metal oxide.


Amorphous solid water (ASW), a metastable phase of water occurring when water is deposited on a substrate cooled below 140K, is of fundamental interest for an array of applications including but not limited to desorption from liquid/ice interfaces and melting of diversified processes. When deposited over an immiscible substance, such as CCl$_4$, the underlayer desorption is dictated by the morphology of the ASW overlayer. This desorption process culminates in the rapid release of the underlayer commensurate with the development of crystallization induced cracks through the ASW. This episodic release is termed the “molecular volcano”.

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Thus, the desorption event reports on changes in the structure of the ASW overlayer. Infrared spectroscopy combined with programmed desorption elucidates the complex interplay between dewetting and etching processes which dominate the eruption event. The effects of the ASW deposition temperature and overlayer thickness on the CC4 desorption process will be presented and discussed.

This work was supported by the U.S. Department of Energy (DOE), Office of Basic Energy Sciences, Division of Chemical Sciences, Geosciences, and Biosciences. The research was performed using EMSL, a national scientific user facility sponsored by DOE’s Office of Biological and Environmental Research and located at Pacific Northwest National Laboratory, which is operated by Battelle, operated for the U.S. DOE under Contract DE-AC05-76RL01830.

Surface Science Division
Room: 110 - Session SS2-MoM

Surface Chemical Dynamics
Moderator: L. Bartels, University of California, Riverside

8:20am SS2-MoM1 STM Dynamics Studies of Tip-Induced Reactions of Anthracene Derivatives on TiO2(110). D.V. Potapenko, R.M. Osgood, Columbia University

In photocatalysis, reactions are driven by the charges that are generated in the bulk of the catalyst and then are transferred to the adsorbed molecules. Injection of electrons from the STM tip into the adsorbed molecules is a useful tool for surface dynamics studies as the energy the electrons as well as the adsorption state of the molecules could be precisely controlled. We use this technique to study tip-induced reaction of anthracene and its derivatives on the TiO2(110) surface. These molecules self-assemble on the rutile(110) surface at room temperature driven by electrostatic interaction. Applying electric pulses from the STM tip to individual molecules causes desorption and in some cases dissociation as indicated by the changes in the STM images. We have observed dissociative electron capture of 2-chloroanthracene molecules that leaves behind a surface chlorine atom. The energy threshold for dissociation found as ~ 3.1 V. We report on the dynamics of this tip-induced chemistry and propose a mechanism for the observed phenomena.


New possibilities for the study of chemical reactions on surfaces using X-ray free-electron lasers (Linac Coherent Light Source, or LCLS, at SLAC Nat. Accelerator Laboratory) will be presented. We induced the hot electron mediated excitation of CO on Ru(0001) with synchronized excitation by a femtosecond optical laser pulse. We have followed the ultrafast evolution of the bond distortions, weakening and breaking, using x-ray emission spectroscopy resonantly tuned to the oxygen core level with ultrashort x-ray pulses delivered from LCLS. We can directly study the time evolution of the molecular orbitals in an atom-specific way on a subpicosecond timescale. Different adsorption configurations explored have been characterized by comparing the measured time dependent energy shifts of the molecular orbitals in the valence band with theoretical results.

This research was carried out on the SAX Instrument on the Linac Coherent Light Source (LCLS) at the SLAC National Accelerator Laboratory. The SAXR Instrument is funded by a consortium whose membership include the LCLS, Stanford University through the Stanford Institute for Materials Energy Sciences (SIMES), Lawrence Berkeley National Laboratory (LBNL), University of Hamburg through the BMBF priority program FSP 301, and the Center for Free Electron Laser Science (CFEL). The LCLS is funded by the U.S. Department of Energy's Office of Basic Energy Sciences.

9:00am SS2-MoM3 The Oxidation of Benzyl Alcohol on Gold and its Implications Towards a Gold-Mediated Conversion of Hydrocarbons. J.C. Rodriguez-Reyes, C.M. Friend, R.J. Marks, Harvard University

In order to understand alcohol conversion on gold, we have explored the mechanism of the oxidation of benzyl alcohol (CH3–CH2–OH) on a Au(111) surface - a test reaction commonly used to determine the efficiency and selectivity of novel gold-based catalysts. Using adsorbed atomic oxygen as the active oxidizing species, we find that a high selectivity towards the corresponding aldehyde (CH3–CH2–CHO) is achieved at low oxygen coverages. However, in excess oxygen the selectivity decreases dramatically and benzaldehyde is readily transformed to adsorbed benzoate, which is eliminated as benzoic acid (C6H5–COOH) and CO2. In addition, we show that the production of the ester produced by self-coupling (benzylic benzoate) does not require the separate presence of acid, as might be expected from acid-alcohol condensation reactions. These surface transformations are extremely facile and exhibit sub-picosecond rates of the rate-limiting steps. Lastly, by comparing the mechanism of benzyl alcohol oxidation to that of toluene oxidation, we extract information from a model hydrocarbon-to-alcohol reaction (toluene to benzyl alcohol, C6H5–CH3 to C6H5–CH2–OH). Our results suggest that due to the relatively large kinetic requirements for toluene conversion, the alcohol cannot be selectively produced on a gold surface; indeed, preliminary TPRS data indicates that the gold-mediated conversion of toluene yields benzoic acid and combustion gases.

9:20am SS2-MoM4 Determination of the Structure and Vibrational Dynamics of Methyl-Terminated Si(111) Using Helium Atom Scattering. R.D. Brown, S.J. Sibener, University of Chicago

Methyl-terminated Si(111) interfaces are of great interest in the fields of solar energy, solar fuels, and electronics due to their superior electronic and chemical stability and device performance. Recent advances in synthetic techniques allow for the fabrication of high-quality, passivated organic-semiconductor hybrid interfaces on single-crystalline silicon substrates. This termination maintains the bulk electronic behavior of the underlying silicon, but eliminates surface trap states which hinder the performance of any device. We directly examined the structural and dynamical characteristics of the CH3-Si(111) and CD3-Si(111) interfaces using helium atom scattering. Helium atom scattering is a uniquely surface-sensitive technique which probes both atomic structure and dynamics. These results are the first measurements of this nature for an organic-semiconductor hybrid interface. Helium atom diffraction confirmed a high quality (1x1) methyl termination of the Si(111) substrate. We observed unusual dynamical characteristics for these interfaces through Debye-Waller attenuation measurements of helium atom diffraction peaks. These measurements characterized the thermal motion of the interface, and elucidated the surface Debye temperature and gas-surface interaction well depth. The Debye Waller measurements yielded temperature dependences of the surface mean-square displacements displaying polarization dependence, and were similar to the thermal motion of the vibrational modes. The effective surface Debye temperatures were much higher than expected, and correspond to the local vibrational modes of the C-Si bond. These two unusual results indicate that termination with the simplest monolayer organic film initiates a transition away from phonon dominated vibrational dynamics to dynamics dominated by local molecular vibrational modes. We also performed a series of high precision inelastic time-of-flight measurements to characterize the phonon band structure of these interfaces. Our measurements have clearly resolved multiple single phonon surface modes on CH3-Si(111), and are the first observations of this nature on an organic-semiconductor hybrid interface.

9:40am SS2-MoM5 STM Insights into Single-Molecule Dynamics. K. Morgenstern, Leibniz University of Hannover, Germany

INVITED

The development of molecular switches on the single molecule level is a major challenge on the path towards incorporating molecules as building units into nanoelectronic circuits. With a scanning tunneling microscope (STM) it is possible to induced chemical reactions on a single molecule basis by electrons tunneling inelastically from the STM tip into a molecule. The method is based on high-resolution imaging at low-temperature (5K) that allows us to identify different groups within the molecule. Chemical reactions are induced by injecting selectively electrons into specific parts of the molecule. The success of the manipulation is visualized in the recorded tunneling current during the manipulation and in STM images taken afterwards. We investigate isomerization of individual molecules adsorbed on metal surfaces. For chlorobenzene and azobenzene derivatives, the effects of different substitutional groups and different substrates are explored.
We are able to quantify the state-resolved reactivity of methane in this clusters. At low $E_i$ and $T_s$, coverage dependent adsorption probabilities, bending combination vibration of CH$_4$ on Ni(111) as a function of reactivity of methane prepared in select vibrational states reveal how both fundamental and practical impact. Experiments that measure the increase in impact energy, $E_i$, and adsorption temperature, $T_s$, which is dissociative chemisorption. The result permits a detailed comparison of the mechanism on a metal surface remains scarce. We will present recent mechanisms.”

Methane dissociation on transition metal surfaces is the rate-limiting step in methane dissociation via a precursor-mediated mechanism, but evidence for Evib activation via a precursor-mediated chemisorption via a direct mechanism, but evidence for Evib activation via a precursor-mediated mechanism on a metal surface remains scarce. We will present recent experimental results from our lab that demonstrate the ability of Evib to promote methane dissociation via both direct and precursor mechanisms on Ir(111). These results build on prior studies of trapping-mediated reactivity on Ir(111) by exploring how Evib in the incident methane molecule promotes reactivity, and they are consistent with the timescales and pathways of vibrational energy flow and damping on these surfaces.


10:40am SS2-MoM8 State-resolved Reactivity of Methane ($v_2 + v_4$) on Ni(111). N. Chen, Y. Huang, A. Uts, Tufts University Methane dissociation on transition metal surfaces is the rate-limiting step in the steam reforming reaction, which is the principal route for converting CH$_4$ to H$_2$. Understanding the dynamics of energy flow during this process has both fundamental and practical impact. Experiments that measure the reactivity of methane prepared in select vibrational states reveal how specific nuclear motions promote methane dissociation. State-resolved measurements for methane reactivity on Ni and Pt surfaces show that molecules prepared in vibrationally excited states are more reactive than those without laser excitation, and the efficiencies of vibrational energy in different excited states can differ significantly. Here, we present state-resolved reactivity measurements of the $v_2 + v_4$ bending combination vibration of CH$_4$ on Ni(111) as a function of translational energy. This state is a member of the pentad of vibrational states that can play an important role in the thermal activation of methane. We are able to quantify the state-resolved reactivity of methane on Ni(111) for particular vibrational state over a wide range of translational energies. We compare the efficiency for $v_2 + v_4$ bend with that of the $v_3$ stretching and $v_4$ bending states to gain insight into the ability of $v_3$ excitation to promote dissociative chemisorption. The result permits a detailed comparison of the role of stretch and bend excitation of methane dissociative chemisorption on Ni(111).

11:00am SS2-MoM9 Molecular Beam Scattering of CO and CO$_2$ on CuO, Nanoclusters Supported on Silica Fabricated by Electron Beam Lithography. M. Komarneni, U. Burghaus, North Dakota State University Adsorption dynamics of CO and CO$_2$ on 12 nm CuO, nanoclusters supported on silica are studied by molecular beam scattering. Samples are fabricated by electron beam lithography. Scanning electron microscopy, auger electron spectroscopy, and x-ray photoelectron spectroscopy are used to characterize the sample. Cu clusters which are nearly metallic, partially oxidized, and fully oxidized are prepared by varying the sample treatment. The initial adsorption probability, $S_0$, of both CO and CO$_2$ decreased with an increase in impact energy, $E_i$, and adsorption temperature, $T_s$, which is consistent with non-activated molecular adsorption. $S_0$ of CO approaches the values obtained for copper single crystals for $E_i$ of 0.4 eV. This is a result of the so-called capture zone effect, where CO molecules are trapped on the support and subsequently diffuse to the metallic clusters. $S_0$ of CO$_2$ is larger than for single crystals. The differences in $S_0$ for metallic and oxidic clusters can be explained by deviations in the mass matching. In contrast with CO, $S_0$ of CO on metallic clusters is slightly larger than that of oxidic clusters. At low $E_i$ and $T_s$, coverage dependent adsorption probabilities, S($\Theta$), of CO and CO$_2$ on metallic clusters obeyed precursor-mediated Kistliuk-like dynamics. At large $E_i$, Langmuirian-like dynamics are seen for CO which is less pronounced for CO$_2$ on metallic clusters.

11:20am SS2-MoM10 Precursor-Mediated Reactivity of Vibrationally Hot Molecules. D. DelSesto, E. Peterson, E. Dombrowski, A. Uts, Tufts University Methane’s dissociative chemisorption is highly activated on a range of transition metal surfaces. Internal-state-averaged and state-resolved beam-surface scattering experiments show that vibrational energy (Evib) can be highly effective in promoting dissociative chemisorption via a direct mechanism, but evidence for Evib activation via a precursor-mediated mechanism on a metal surface remains scarce. We will present recent experimental results from our lab that demonstrate the ability of Evib to promote methane dissociation via both direct and precursor mechanisms on Ir(111). These results build on prior studies of trapping-mediated reactivity on Ir(111) by exploring how Evib in the incident methane molecule promotes reactivity, and they are consistent with the timescales and pathways of vibrational energy flow and damping on these surfaces.

11:40am SS2-MoM11 Three-Dimensional Spatial Distribution of Desorbing N$_2$ and N$_2$O from Pd(211). M. Sakurai, T. Kondo, J. Nakamura, University of Tsukuba, Japan Spatial distributions of N$_2$ and N$_2$O desorbing from Pd(211), Pd(S)-[3(111)](100), in temperature programmed desorption (TPD) were measured. After the N$_2$O desorbing from Pd(211) was introduced, N$_2$O molecules on the surface are found to desorb as N$_2$ by the decomposition with sharp concentration to 25$^\circ$ ($\Theta$ = -25$^\circ$) (100) side from the normal direction of (211) plane at 108 K in the parallel plane to [211] and [-111] directions at N$_2$O coverage of 1.0 ML. This indicates that N$_2$ desorbs without thermal equilibration by N$_2$O decomposition on Pd(211). From the angle-scan differential desorption structure of N$_2$O just before the desorption is assigned as the bridge-structure at step-site, where O and terminal N bonded with Pd at ontop (100) and 360 hollow(111), respectively along the [-111] directions. On the other hand, the spatial distribution of desorbing N$_2$O obeyed cosine distribution directing the normal direction to the (211) plane, suggesting the existence of pre-desorption state on the surface.

Thin Film Division
Room: 107 - Session TF-MoM

Thin Films: Growth and Characterization I
Moderator: J.M. Fitz-Gerald, University of Virginia


There has been growing interest in thin bi-metallic multilayer films for the usage under extreme radiation conditions because of their radiation healing properties. Recent discovery and research indicate that materials can be hardened against radiation damage by building nanolayered structures with an optimized layer thickness to increase point defect recombination relative to the density of defects that can self-heal. In this study, we investigate whether the internal interfaces can be manipulated at the nanoscale to enhance dynamic recombination of radiation-produced defects, or self-healing, so as to dramatically reduce radiation damage without compromising other properties using Ti/Al multilayer films. Ti/Al multilayer films were fabricated on Si (100) and epi polished MgO (100) substrates using DC magnetron sputtering and Molecular beam epitaxy (MBE). The growth parameters for each method, for sputtering- pressure, power and substrate temperature deposition rate; and for MBE- deposition rate and substrate temperature were optimized to achieve high-quality thin films. The films were characterized using x-ray diffraction (XRD), x-ray reflectivity (XRR), Rutherford backscattering spectrometry (RBS), and x-ray photoelectron spectroscopy (XPS) measurements. The films show mostly polycrystalline structure with no elemental interdiffusion at the interfaces. Detailed structural and compositional analysis was also performed using high resolution TEM/STEM and atom probe tomography (APT). The films were irradiated using 1-8 MeV Au ions to understand the radiation effects. The damage peak, stopping range and ion distribution were simulated using binary collision approximation based Monte Carlo method (SRIM software program). Au ion energies, estimated from the simulation, were used to position the damage peak at the interfaces and away from the surfaces to obtain the complete picture at the interfaces and in the bulk of the films. The surface and crystal lattice damage, amorphization, and defect density were studied by RBS, HAADF-STEM and APT, and compared with those results from the pristine samples. The relationships between film properties and radiation healing characteristics will be presented and discussed.

8:40am TF-MoM2 Flux and Surfactant-Assisted Physical Vapor Deposition: New Approaches for Improving Complex Oxide Thin Film Growth. J.-P. Maria, E.A. Paisley, B.E. Gaddy, North Carolina State University, M.D. Biegalski, Oak Ridge National Laboratory, D.L. Irwin, A.R. Rice, R. Collazo, Z. Sitar, North Carolina State University INVITED Epitaxial integration of complex oxides with wide band gap polar semiconductors such as GaN (0002) presents the possibility for high-power, high-frequency, and high temperature GaN electronics by virtue of 2-D charge carrier materials at polar interfaces and possible access to non-linear dielectric properties. This work describes synthesis and characterization of such thin film heterostructures by MBE, with specific attention given to controlling film growth. Our previous work has shown growth of high-quality MgO (111) and CaO (111) films on GaN with measured valence band offsets of 2 and 2 eV (MgO) and 2.5 eV (CaO). However, the terminal 3D rocksalt growth surface is determined by the tendency for the high-energy (111) polar orientation to form (100)-oriented low-energy facets, and the interfacial symmetry between cubic MgO and hexagonal GaN. Therefore, realizing smooth oxide films on GaN requires the ability to overcome the rocksalt (001) faceting tendency through a surfactant growth method. In this presentation we will discuss a newly-developed surfactant approach to MBE growth of rocksalt oxides utilizing water vapor to hydroxylate CaO and MgO (111) surfaces in situ, which changes the equilibrium habit from cubic to octahedral.
eliminating the (100) facets and tendency. RHEED oscillations and AFM images of these films show 2D growth, suggesting that altering the surface chemistry during growth plays a critical role in controlling the surface orientation. We demonstrate through electrical property measurements the impact of smooth (111) rocksalt surfaces as leakage current densities for thin CaO films are reduced by two orders of magnitude when films of equivalent thickness are grown using a 2-D vs. a 3-D mode. Finally, we will present a set of temperature-ab-initio thermodynamic calculations of CaO surface energies with and without H-containing terminations that validate the surface-chemical mechanism of facet stabilization.


The state-of-the-art approach in the encapsulation of high-end devices such as solar cells (polycrystalline Si) and organic light emitting diodes is based on an organic/inorganic multilayer system. Although this approach allows increasing the lifetime of the encapsulated device, the optimization of a multilayer is rather empirical as the mechanisms behind the improvement of the barrier performance are not yet unraveled. In particular, the role of the organic interlayer is rather controversial since it does not act as a moisture vapor barrier, yet its application appears to be fundamental in the multilayer solution.

In this contribution, the role of the organic interlayer is investigated by selecting a system in which the barrier layer, a 100 nm-thick SiO2 film, is plasma-deposited while the organic interlayer, a 200 nm-thick organosilicon film, is synthesized by means of initiated-chemical vapor deposition, i.e. via thermal desorption of an initiator molecule promoting the polymerization of 3,5-trimethyl-1,3,5-trivinyl-cyclooctasiloxane (Vd3) at the substrate.

The implementation of in situ (real time) spectroscopic ellipsometry allows following the different growth stages in the Vd3 polymerization process. In particular, when applied to the polymer bulk growth, the determination of the growth rate allows monitoring the transition from a kinetic-limited (with activation energy of 65 ± 4 kJ/mol) to a mass transfer-limited regime. Furthermore, the deposition process is found to be monomer adsorption-limited, characterized by an adsorption energy of 39 ± 4 kJ/mol. When close to the deposition process is initiated, the open porosity in the micro/meso transition region (pore radius ≤ 2nm). The microstructure characterization by means of the above-mentioned studies explicitly points out the role of the i-CVD organic interlayer in the multi-layer barrier structures, i.e. the filling of the open micro/meso porosity of the inorganic barrier layer, therefore, improving the intrinsic barrier quality of the underlying SiO2 barrier layer, characterized by a residual open porosity in the micro/meso transition region (pore radius ≤ 2nm). The microstructure characterization by means of the above-mentioned studies implicitly points out the role of the i-CVD organic interlayer in the multi-layer barrier structures, i.e. the filling of the open micro/meso porosity of the inorganic barrier layer, therefore, improving the intrinsic barrier quality of the underlying SiO2 barrier layer, characterized by a residual open porosity in the micro/meso transition region (pore radius ≤ 2nm).

9:40am TF-MoM5 Processing and Characterization of Iron and Fluorine Co-Doped Ba0.6Sr0.4TiO3 Thin Films, F. Stemme, H. Gesswein, A. Azucena, Karlsruhe Institute of Technology (KIT), Germany, M. Sazegar, Darmstadt University of Technology, Germany, J.R. Binder, M. Bruns, Karlsruhe Institute of Technology (KIT), Germany

Barium strontium titanate (BST) is a very promising material for tunable microwave applications like phase-shifters and tunable filters. Due to the influence of the a-gelation conditions and processes on thin film properties and their dielectric performance were largely investigated. But very few researchers have tried to change the sputtered BST thin film properties by using different dopants at the same time. Such iron and fluorsine co-doped thin films can be achieved by RF magnetron sputtering, with a co-sputter target and a two step annealing process after deposition. The first annealing provides the crystallization of the films. The second annealing process the fluorsine co-dopant is introduced into the BST thin films by a diffusion controlled process.

The present contribution focuses on the processing and characterization of the iron doped BST thin films with various amounts of fluorsine co-dopant. The characterization of the thin films by X-ray photoelectron spectroscopy (XPS) provides chemical binding states and film composition. XPS and time of flight secondary ion mass spectrometry (ToF SIMS) sputter depth profiling prove the chemical homogeneity and the film thickness. Grazing incident X-ray diffraction (XRD) and Raman spectroscopy validate the crystallinity and the identification of chemical phases. Furthermore film morphology is determined by scanning electron microscopy (SEM) and atomic force microscopy (AFM). Dielectric measurements, to investigate the influence of the donor and acceptor co-doping on the dielectric performance, were carried out in metal insulator metal (MIM) structures with ground signal ground probes.

10:00am TF-MoM6 Advances in the Growth of Epitaxial Oxides for Neuromorphic Computing Applications, J.D. Greeneel, W.L. Calley, W.A. Doolittle, Georgia Institute of Technology

Present-day computing devices are approaching performance limits due to excess heat and parasitic capacitance degrading performance. One proposed solution, termed ‘neuromorphic computing,’ is to perform computations similarly to the brain.

A key enabling device for neuromorphic computing is the lithium niobate (LiNbO2) memristor[1]. Like synapses in the brain, LiNbO2 memristors can express excitatory or inhibitory behavior. This is due to the ability to produce both n- and p-type LiNbO2 memristors, a unique property for an oxide. As shown in Figure 1, when a voltage is applied across an n-type memristor (left), resistance increases over time, whereas a p-type memristor exhibits decreasing resistance (right). N-type memristors are grown with excess oxygen vacancies while p-type memristors are grown with excess lithium vacancies. N- and p-type LiNbO2 membranes exhibit resistances of 4.3x10-4 Ω-cm (n-type) and 3.9x10-4 Ω-cm (p-type), respectively.

The epitaxial growth of LiNbO2 is the enabling technology for the memristors described above. Growths are conducted using an oxy-chloride MBE system in which metal-halide sources interact with lithium and oxygen [2]. Lithium acts as a getter for the chlorine from the metal-halide, and excess LiCl is deposited from the heated growth surface. The bare metal then oxidizes under the application of oxygen plasma to form a high-k oxide. This growth chemistry has been used to produce dielectric/ferroelectric lithium niobate (LiNbO3), semiconducting lithium niobite (LiNbO2), and lithium cobalt oxide (LiCoO2).

Insulating LiNbO3 (Figure 2) and semiconductor LiNbO2 (Figure 3) are grown using NbC5, and the phase of the material is controlled by the niobium to oxygen ratio delivered to the growth surface. Both single crystal LiNbO3 and LiNbO2 have been epitaxially grown on Al2O3 and SiC. LiNbO3 is a promising high-speed waveform generator and high-k oxide for its semiconducting sub-oxide, LiNbO2. LiNbO2 has a layered, lithium-intercalated structure [3] (Figure 4). When lithium is removed, holes are introduced which increases the conductivity of p-type material and decreases the conductivity of n-type material, thus producing the memristance effect described herein.

LiCoO2 is grown on Al2O3 using CoC12 as the precursor (Figure 5). Like LiNbO2, LiCoO2 has a layered structure that facilitates lithium movement. The lattice spacing of LiCoO2 is closely matched to LiNbO2 (2.81 Å and 2.91 Å respectively), thus making it an attractive material for heterostructures of lithium-bearing semiconductor materials.

10:40am TF-MoM8 New Method to Produce High-Quality Epitaxial Ge on Si Using SiO2-Lined Etch Pits and Epitaxial Lateral Overgrowth for III-V Multijunction Solar Cells, D. Lennes, S.M. Hay, University of New Mexico

Integrating a high-quality layer of epitaxial Ge on Si has been a longstanding engineering challenge, despite its technological importance. The applications of Ge-on-Si include ‘virtual substrates’ for III-V multijunction solar cells, high-mobility-effect transistors, and optical interconnects monolithically integrated with Si-based circuitry. The primary difficulties in achieving Ge films of sufficient quality stem from the lattice mismatch that leads to a large density (~10^9 cm^-2) of threading dislocations (TDs) and the thermal expansion coefficient mismatch between Ge and Si that leads to microcracks or delamination of Ge film upon cooling from growth to room temperature. Herein, we present a new method to reduce the TD density, using a minimal number of standard microfabrication steps. The method begins with growing a 500-nm-thick epitaxial Ge layer on Si. A post-growth anneal step leads to a TD density of approximately 5x10^7 cm^-2, as measured by electron channeling contrast imaging (EMCD) and etch pit density (EPD) measurements. The close agreement between EPD measurements and TEM shows that the EPD measurements reliably decorate all TDs. Etch pits are created around the dislocation cores in the Ge film. A 15-nm-thick layer of SiO2 is subsequently deposited on the etch-pit-decorated Ge film. A thin layer of polyimine methacrylate is then spin-coated onto the sample, which fills the etch pits and planarizes the Ge surface. Next, a reactive ion etching step is used to remove the polymer and SiO2 from the planar regions of the sample surface surrounding the etch pits. An O2 plasma is then used to selectively remove the remaining polymer, so that SiO2 remains only within the etch pits. Lastly, a second layer of Ge is selectively grown on the exposed Ge surface and laterally over the SiO2-lined etch pits until a fully coalesced Ge film is created. A final polishing step produces an atomically flat continuous Ge film. ensuing EPD measurements reveal that the density of twin defects and TDs in the
upper Ge layer is approximately $1.7 \times 10^8$ cm$^{-2}$, such that the overall defect density is reduced by a factor greater than 30 compared to that in the initial Ge layer. Both theoretical and experimental results suggest that the defect density in GaAs films on Ge/Si must be less than $2 \times 10^7$ cm$^{-2}$ to have a minor carrier lifetime comparable to GaAs films grown on Ge and GaAs substrates. Therefore, our new method of using SiO$_2$-lined etch pits to block the propagation of TDs in Ge may finally lead to device quality III-V materials integrated on Si substrates.

11:00am TF-MoM9 Three Dimensional Reciprocal Space Measurements by X-ray Diffraction using Linear and Area Detectors: Application to Texture and Defect Determination in Oriented Thin Films and Nanoprecipitates, S. Gaudet, S. Lambert-Milot, P. Desjardins, Ecole Polytechnique de Montréal, Canada, A. Vasher, F. Desnoyers, Hohenheim Université, Berlin, J.L. Jordan-Sweet, C. Laviole, IBM T.J. Watson Research Center, New York, USA

Very thin films grown or deposited on oriented substrates are often composed of highly oriented phases. Traditional x-ray diffraction approaches, scanning only limited portions of the reciprocal space, can fail to characterize some phenomenon occurring in these thin films. We developed an approach for the fast and efficient measurement of complete volumes of the reciprocal space by x-ray diffraction using linear and area detectors. In this presentation, we show how it allowed a detailed understanding of the solid-state thin film reaction to form contacts on transistors and of the growth of magnetic nanoprecipitates by metalorganic vapor-phase epitaxy. We first explain the procedures and scan strategies required for transforming raw scattering data into three-dimensional maps of the reciprocal space, and we present a complete open source software package for advanced data processing, analysis, and visualization. Case studies are chosen to highlight the overall capabilities of the technique, which are then introduced. First, thermal diffuse scattering from a monocry stalline Si substrate is characterized. The presence of lines linking diffraction peaks in reciprocal space reflects the interaction of the x-ray beam with the phonon spectrum of the monocrystal. Second, a detailed investigation of texture in multiphase thin layers permits to reveal the unambiguous presence of fiber, axiotaxial, and epitaxial components in highly oriented films. The visualization of the entire reciprocal space allowed us to identify unexpected metastable phases that could not be deduced or observed from measurements carried out in the Bragg-Brentano geometry. Finally, we present an investigation of defects in two thin film systems: planar defects in nickel silicide layers formed by solid-state reactions and microtwins in a GaP matrix containing coherent Mn$^2+$ precipitates. In both systems, the systematic acquisition of significant volumes of the reciprocal space allows for the observation of behaviors that could otherwise remain undetected when restricting the analysis to typical measurement scans.

11:20am TF-MoM10 Surface Characterization of Zr/Ti/Nb Tri-layered Films Deposited by Magnetron Sputtering on Si(111) and Stainless Steel Substrates, D.A. Tallarico, Federal University of Sao Carlos, Brazil, E.L. Gross, Brazilian Nanostructure Light Laboratory, Brazil, P.J. Pauvin-Filho, Federal University of Sao Carlos, Brazil, A. Calbaygues, Ecole Nationale Superieure de Chimie de Paris, France. P.A.P. Nascente, Federal University of Sao Carlos, Brazil

Titanium is a metal commonly used in medical implants, due to its interesting properties, such as high mechanical strength, good corrosion resistance in extreme conditions, and excellent thermal stability. Ti-6Al-4V is one of the main bi-metallic alloys employed as implants, but the release of Al and V ions is associated to health problems and adverse tissue reactions. A new class of Ti alloys employs Zr for solid-solution hardening and Nb as phase stabilizer. Metals such as Ti, Nb, and Zr, known as valve metals, usually have their surfaces covered by a thin oxide film spontaneously formed in air. This oxide film constitutes a barrier between the metal and the medium. The Ti-Nb-Zr alloys have mechanical and corrosion resistance characteristics which make them suitable for use as implants. Tri-layered films of Ti-Nb-Zr were deposited on both Si(111) and stainless steel substrates using a DC magnetron sputtering equipment, under an argon atmosphere. The films were deposited in the following manner: a 100 nm thick layer of Nb was deposited on a Si(111) substrate, then a 200 nm layer of Ti was deposited on top of Nb/Si, and lastly a 50 nm layer of Zr was deposited on top of Ti/Nb/Si. A similar Ti/Nb/Zr film was also grown on stainless steel substrate. The structure, morphology, and chemical composition of the films were analyzed by means of X-ray diffraction (XRD), atomic force microscopy (AFM), X-ray photoelectron spectroscopy (XPS), and time-of-flight secondary ion mass spectrometry (ToF-SIMS). The XPS results for the Zr/Ti/Nb layers deposited on Si(111) presented a predominant presence of the surface Ti-Nb-O and a small quantity of Ti-O. For the layers deposited on stainless steel, only Zr-O and a small amount of niobium oxide were detected. The ToF-SIMS results indicated the formation a three-layered film on Si(111), with each metal in a distinct layer and a well-defined interface between the layers, while the deposition on the stainless steel substrate caused slight intermixing at the Nb/Ti and Ti/Zr interfaces. AFM images showed that the Zr/Ti/Nb tri-layer films present a rougher surface than those grown on Si(111), but the roughness, with the film deposited on stainless steel having the roughest surface.


Hybrid organic-inorganic nanolaminates combine the functionality of an inorganic material with the flexibility and mechanical integrity provided by the organic polymer layer. They are integral components in various approaches serving as advanced dielectrics, flexible barrier coatings, and optical components. This work focuses on the low temperature synthesis of alumina/silicone nanolaminates by plasma-enhanced chemical vapor deposition (PECVD) in a single chamber for dielectric applications.

Self-limiting synthesis of alumina was accomplished via pulsed PECVD at the synthesis temperature of ~ 100 °C using trimethyl aluminum (TMA) and oxygen as precursors. The deposition kinetics and film quality were evaluated as a function of precursor exposure, plasma power, substrate temperature, and pulse parameters. Precise control of nanolaminate construction was confirmed through field emission scanning electron microscopy (FE-SEM) and transmission electron microscopy (TEM). The dielectric performance of these structures was examined by using capacitance-voltage and current-voltage measurements. The effective dielectric constant could be controlled by changing the alumina content in the nanolaminates, and modeling these structures as capacitors in series accurately described the observed variations in $\varepsilon$.

Vacuum Technology Division

Room: 111 - Session VT-MoM

Vacuum Measurement, Calibration & Primary Standards, Gas Flow and Permeation

Moderator: R. Garcia, SAES Getters

Over the past decade, NIST has designed and built several high-stability transfer standard packages (TSPs) that have proven to be ideally suited for inter-laboratory comparisons in the atmospheric pressure and vacuum pressure range [1]. In the mid 1990’s the development and use of micro electro mechanical systems (MEMS) enabled pressure sensor technology to make significant advances in both precision and accuracy. Resonant silicon gauges (RSGs) are MEMS sensors that are manufactured by microfabricating silicon to produce silicon diaphragms nominally a few millimeters square by a fraction of a millimeter thick [2]. NIST has found that these gauges are very stable, rugged, and ideally suited as core technology for high-stability precision pressure and vacuum standard. The RSG sensors with full-scale ranges of 10 kPa and 130 kPa have shown excellent stability [3]. However, one drawback of the RSGs is that they lack the sensitivity and resolution of capacitance diaphragm gauges (CDGs) with full-scale ranges of 133 Pa. The downside of only using these CDGs is the relatively poor calibration stability when compared to 10 kPa full scale range RSGs. The NIST solution has been to combine the high-resolution of the 133 Pa CDGs, with the high-stability of the 10 kPa RSGs into one transfer standard package. The RSG gauges are then used to determine the calibration shift in use. A recently completed set of transportable NIST traceable vacuum standards with a range of 1 Pa to 10,000 Pa will be highlighted. These TSPs consist of pairs of 10 kPa RSGs and pairs of 133 Pa CDGs enclosed in temperature controlled enclosures that further enhance gauge performance. These standards will be used for an
The spinning rotor gauge (SRG) has long been the primary transfer standard used by metrology laboratories for providing traceability of measurements from field vacuum devices to primary vacuum standards maintained at national measurement institutes. The useful range of the SRG, in its current commercial form, is somewhat limited by several factors including the relatively small change in momentum of the rotating sphere caused by the adsorption/desorption of gas molecules as the gas pressure (and thus number of molecules adsorbing/desorbing) decreases. Current technology limits the low pressure end of this range to about 2E-5 Pa with a K=2 uncertainty of approximately 1-3%.

Efforts are currently underway at the Oak Ridge National Laboratory to increase the range of the commercial SRG by approximately a decade by increasing the sensitivity of the rotor's momentum change to the low number of gas molecules available for adsorption/desorption at the lower pressures. Efforts are also underway to improve the uncertainty of the existing commercial SRG through better measurement and characterization of the thermal expansion of the spinning rotor itself. This paper will report on the progress of these efforts to date, some of the challenges discovered during the development process and plans to address those challenges.
A pinhole orifice with a known conductance can be used as a secondary flow standard. We are interested in using pinhole orifices to produce nitrogen gas flows into vacuum in the range of $10^{-11}$ mol/s to $10^{-4}$ mol/s ($10^{-7}$ to $10^{-2}$ cm$^3$/s; STP) for vacuum gauge calibrations because a flowmeter based upon an appropriate set of orifices is easy to operate and automate. Commercially available laser-drilled pinhole orifices with diameters from 1 μm to 50 μm can have molecular-flow conductances, $C_0$, ranging from about 0.1 μL/s to 230 μL/s for N$_2$ at 23 °C, and can be used to produce gas flows in the range of interest by applying an upstream pressure in the range of 10 Pa to 100 kPa (0.1 to 760 torr). Accurate measurements of the orifice conductance, $C$, as a function of pressure are required to use the pinhole orifice as a basis of a flowmeter. The NIST bellows flowmeter is a primary gas flow standard that was used to directly measure the conductance of a pinhole orifice to better than 0.2 % over the entire pressure range of interest. We present results of the conductance measurements for nitrogen and other gases. One might expect that the differences among the gases would be mainly due to their different thermal velocities, and that those differences would disappear when the normalized reduced flow rate ($C/C_0$) is plotted as a function of inverse Knudsen number. However, this was not the case at higher pressures. For example, the reduced flow rate for nitrogen was smaller than for argon at the same inverse Knudsen number. Following a suggestion by Jitschen (Vacuum 76 (2004) 89-100), the effect of the heat capacity ratio on $C/C_0$ was investigated.
transferring surface-adsorbed biomolecules into the gas phase [1]. Using Neutral cluster-induced desorption and ionization is a very soft method for detecting the dipole moment of the cluster’s constituents which allows both for solvation and in great detail. From the experimental data, combined with polymer theory, we develop a better understanding of the relationship between the supramolecular organization and dynamics of biomolecular hydrogels, their physico-chemical properties and their biological function. To illustrate this concept, I will present some of our recent work on the “sweet” jelly-like matrix that forms around the mammalian egg during ovulation (the so-called cumulus cell-oocyte complex matrix) and that is crucial for fertility, and on the proteoglycan-meshwork that contributes to the load-bearing and lubricating properties cartilage

In order to interrogate biomolecular hydrogels directly on the supramolecular level, we have developed an unconventional approach that draws on knowledge from several scientific disciplines. Exploiting surface science tools, such as supported lipid bilayers, we tailor-make model systems by directed self-assembly of purified components on solid supports. With a toolbox of surface-sensitive analytical techniques, including quartz crystal microbalance, ellipsometry, atomic force microscopy and microinterferometry, these model systems can be investigated quantitatively and in great detail. From the experimental data, combined with polymer theory, we develop a better understanding of the relationship between the supramolecular organization and dynamics of biomolecular hydrogels, their physico-chemical properties and their biological function. To illustrate this concept, I will present some of our recent work on the “sweet” jelly-like matrix that forms around the mammalian egg during ovulation (the so-called cumulus cell-oocyte complex matrix) and that is crucial for fertility, and on the proteoglycan-meshwork that contributes to the load-bearing and lubricating properties cartilage.

Desorption Efficiency

Biomolecules – Influence of Surface Load and Sample Morphology on Desorption Efficiency. M. Baur, B.-J. Lee, HS Esslingen, Germany, C.R. Gebhardt, Bruker Daltonik, Germany, H. Schroder, K.-L. Kompa, MPI for Quantum Optics, Germany, M. Durr, HS Esslingen, Germany

Neutral cluster-induced desorption and ionization is a very soft method for transferring surface-adsorbed biomolecules into the gas phase [1]. Using neutral cluster-induced desorption and ionization, the method makes use of the dipole moment of the cluster’s constituents which allows both for solvation and charge transfer processes in the cluster [2]. Thus the cluster provides not only the energy for the desorption process but also serves as a transient matrix. As a consequence, desorption and ionization of oligopeptides and proteins is observed at low energies of the impacting clusters and without any fragmentation of the biomolecules.

Here we show that cluster-induced desorption and ionization of biomolecules can be efficiently applied for a wide range of surface concentrations and configurations, i.e. from µm-thick films down to surfaces prepared with submonolayer surface concentration of biomolecules. Highest signal intensity in the respective mass spectra was observed from thick films, indicating an efficient desorption mechanism from bulk-like material. In the submonolayer regime, the ion signal of the desorbed biomolecules was found to depend nonlinearly on surface concentration of the wet-chemically applied biomolecules. The behavior is traced back to the formation of multilayered islands of biomolecules on the surface, as observed by means of SEM and AFM, and a dominant contribution to the ion signal from these islands even at low coverage. With the current set-up and preparation scheme, the lower detection limit was shown to be 10⁻¹⁷ mol.


Monday Afternoon, October 31, 2011

Applied Surface Science Division
Room: 102 - Session AS-MoA

Quantitative Surface Chemical Analysis and Technique Development - Part II
Moderator: M.S. Wagner, The Procter & Gamble Company
electrical effects may be convoluted with geometric factors and influence mass resolution in a number of unpredictable ways.

For data acquired in "raw" mode (full spectrum at every pixel), it is sometimes possible to correct for these real-world difficulties after data acquisition. Two approaches are employed. The first involves subdividing the analysis area into a regular grid of smaller regions and extracting mass spectra from each region. The extracted spectra are individually calibrated by an automated process, and all or an optimized portion of the spectra are summed to produce a new spectrum with higher mass resolution than the original total spectrum. Interestingly, the spectral calibration information can be used as a diagnostic tool for instrument alignment and tuning.

The second approach is effective for improving mass resolution in spectra of rough surfaces, such as fabrics. Unlike the first approach, the analysis area is not subdivided into a regular pattern. Rather, spectra are obtained from regions of similar height, identified by any of four methods ranging from manual selection of regions-of-interest to automated pixel selection using principal component analysis and multivariate curve resolution. The automated methods have the advantage of simultaneously optimizing the mass resolution and the spectral counts without having to take a trial-and-error approach.

With these methods, mass resolution improvements of 20% - 50% are typical for smooth surfaces, and much larger improvements can be achieved for rough surfaces.

4:20pm AS-MoA8 ToF-SIMS Analysis of Iron Oxide Particle Oxidation by Isotopic and Multivariate Analysis, J. Ohlhausen, E. Coker, A. Ambrosini, J. Miller, Sandia National Laboratories

A procedure for quantitative ToF-SIMS analysis of the re-oxidation of iron oxide particles in a ceramic matrix is discussed. Iron oxide is reacted with yttria stabilized zirconia (YSZ) to create a composite that facilitates the high temperature decomposition of CO2 and H2O. In the two step process, Fe2O3 is partially reduced to FeO by heating to high temperatures (~1300 °C) under inert atmosphere. It is then re-oxidized at < 1200 °C under CO2 or H2O yielding CO or H2 respectively. The reactivity of this two step solar-thermochemical process is being investigated by varying the concentration of iron in YSZ up to and past its solid solubility point, thus affecting the size of iron oxide particles in the matrix, and hence their rate and extent of re-oxidation. For the SIMS experiment, the YSZ sample containing natural abundance iron oxide was mixed with an organic binder, isostatically pressed into a disc and calcined in air at 1450 °C. This disc (~10mm diameter, 2mm thickness) was thermally reduced at 1400 °C and then re-oxidized at 1100 °C in the presence of CO2. The ratio of 18O to 16O shows the extent of oxygen exchange for each iron oxide particle.

For ToF-SIMS analysis, samples are prepared by cross-sectioning and polishing by conventional metallographic preparation techniques followed by ion milling with Cs+ in the ToF-SIMS. ToF-SIMS data are acquired from the cross section only after surface contaminants are removed and a "bulk" condition exists on the exposed surface. Data are acquired in a fashion that maximizes the ability to correct for detector saturation, thus providing quantitative oxygen isotopic results with little error. Data analysis methods uses a combination of multivariate analysis for particle identification and conventional analysis for quantitative isotopic ratioing. Details of analysis procedures will be discussed along with results for a range of iron oxide particle sizes.

Sandia National Laboratories is a multi program laboratory managed and operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Corporation, for the U.S. Department of Energy's National Nuclear Security Administration under contract DE-AC04-94AL85000.

4:40pm AS-MoA9 Informatics for SIMS: Identifying Molecules in Complex Mass Spectra, J.S. Gilmore, F.M. Green, M.P. Seah, J.L.S. Lee, National Physical Laboratory, UK

High-throughput screening using mass spectrometry for proteomics has driven the need to move from manual methods for protein identification to automated methods. Metabolomics has similar needs owing to the complex chemical mixture studied. A combination of three important developments has allowed major progress in the automated interpretation of spectra to identify chemical and biological constituent substances. These are (i) the explosion in the amount of publicly available chemical information (PubMed, for example indexes over 71 million substances) (ii) advances in mass spectrometry search engines and fragmentation tools and (iii) rapid growth in high performance mass spectrometers (mass accuracy < 1 ppm and mass resolution > 100,000). These recent developments in informatics are the endeavours of a very much larger community than the surface analysis community. We can utilise this rich resource.

We show this in three parts. Firstly, we analyze the popular PubChem database in terms of the population of substances with mass when resolved with typical mass spectrometer mass accuracies. In general, in ToF-SIMS the mass accuracy is ~ 30 ppm for an unknown substance. For a typical molecule (the modal mass in PubChem is 385 u) there are ~ 30,000 substances within this mass tolerance. In high performance mass spectrometers (~ 1 ppm mass accuracy) this range reduces to ~ 1000 substances which may be further reduced to around 50 substances using isotope pattern matching. Clearly, the mass accuracy in organic SIMS needs to improve significantly to benefit from chemical databases in the same manner as the metabolomics community. Secondly, we have previously shown that G-SIMS simulated spectra so that the most structurally significant peaks are dominant and we now show a new development called the g-ogram. This gives a visually simple chromatographic method to interpret spectra and allows separation of, for example, substrate, polymer and molecule peaks based on the fragmentation energy. Thirdly, we show how the G-SIMS spectra are a bridge to the informatics methods used by the metabolomics community providing identification automatically linked to public chemical databases. Present challenges and future opportunities will be discussed.

References
[1] F.M. Green, J.S. Gilmore & M.P. Seah, Analytical Chemistry 2011, dx.doi.org/10.1021/ac200676a

5:00pm AS-MoA10 keV Ion Impact Effect on the IonCCD™ Surface and Mass Spectra Peak Shape in Non-Scanning Sector-Field Instrument. O. Hadjar, G. Kibelka, S. Kaisan, C. Cameron, K. Kuhn, OI Analytical

Particle-surface interactions are very important processes making physics practically impossible to apply without putting those interactions into the equation. For particle detection applications, the detection event is triggered by total or partial particle energy deposition upon impact on the detector. Mass spectrometry common ion detectors are Channeltrons and MCPs, which inherently destroy the particle upon measurement. The IonCCD, a product from the rapidly emerging technology will be characterized against keV ion impact when used in a dispersive mass analyzer.

The IonCCD is used as focal plane array in a sector field instrument of Mattauch-Herzog geometry (MH-MS). When miniaturized, MH-MS is best suited for low mass range applications (< 100 u). Different from the two first detector families that most often operate in particle counting mode (time resolved detection) the IonCCD operates in an integration mode (charge integrator). In this case, dispersed ions neutralize on the electrode pixels for a well-defined time known as the integration time. While the potential energy of the detected ions is used for detection, the kinetic energy leads to ion-surface interaction, an artifact amplified at extreme low mass detection. This latter can be eliminated by floating the IonCCD or operating it in higher magnetic fields.

The artifact manifesting in the mass spectra as distortion (negative peak) due to keV ion impact induced secondary electron emission was modeled and investigated experimentally using electronic stopping power fingerprints. We demonstrate that the artifact increases linearly with ion impact velocity and is dependent in an oscillatory fashion on ion nuclear charge. Both findings are in agreement with the electronic stopping of keV ions with the TiN surface of the IonCCD. 3D simion modeling suggests efficient peak artifact suppression by operating the IonCCD in higher B-field (~ 4000 G) and less elegantly by IonCCD-magnet face-retarding field. Same model was used to enhance the performance of the instrument, confirming the dynamic mass range (Mmax/Mmin) increase from 16 to 70.

The potential IonCCD damage upon keV ion impact through the nuclear stopping effect was investigated by means of atomic Force Microscopy and X-ray Photoelectron Spectroscopy. While AFM confirmed the expected increase in surface roughness, XPS showed no stoichiometry change due to implantation or preferential ion sputtering. The discoloration observed after extensive use was related to carbon layer formation in the roughened irradiated pixel area. Nuclear stopping effects do not seem to affect the detector performance at practical doses.

5:20pm AS-MoA11 First use of ToF-SIMS for Screening Assays: Enzymes Active on Wood. R.E. Goacher, E.A. Edwards, C.A. Mims, E.R. Master, University of Toronto, Canada

Proteomic and metagenomic studies are rapidly increasing the number of proteins available for enzymatic screening. However, current high-throughput enzyme assays have limited applicability for an important class of biochemical substrates – complex solid materials. The present work aims...
to utilize the strengths of Time-of-Flight Secondary Ion Mass Spectrometry for the direct measurement of enzyme activity on solid substrates. Particularly, ToF-SIMS is applied to the detection of wood-modifying enzymes.

Proof-of-principle ToF-SIMS enzyme assays were performed by immersing extracted wood fibers in solutions of commercial cellulase and laccase enzymes (utilizing water/buffer and denatured enzymes for controls). The laccase enzyme was also tested with and without several small molecule mediators. Principle Component Analysis (PCA) clearly distinguished cellulase tests from controls through the loss of polysaccharide peaks and relative enrichment of lignin peaks. Additionally, PCA differentiated laccase test samples (with mediator) from controls through a shift in lignin-characteristic peaks. The active laccase was indicated by a relative decrease in guaiacyl-lignin and syringyl-lignin peak intensities and increase in generic aromatic peaks, resulting from the cleavage of hydroxyl and methoxy groups from lignin benzoic units.

These proof-of-principle assays demonstrate that ToF-SIMS is capable of providing yes/no screening information for enzyme activity on complex solid substrates, such as wood.

We compared the $I_{DS-VDS}$ characteristics of devices before and after hybridization. A decrease of drain current was noticed, revealing the binding of negatively charged DNAs. More importantly, the decreases in $I_{DS-VDS}$ demonstrated a clear dependence of charge layer distance by an observation of the current change ratio, defined by $\Delta I/\Delta V$, (gate bias: -1 V): 16.1% for fully complementary target DNA (0.1 μm), 14.9% for 21-bp complementary DNA, 10.9% for 13-bp complementary DNA, 6.89 % for 9-bp complementary DNA, and 0.75% for mismatch DNA, which was comparable to insensitivity. The current change decreases with the distance between the target DNA and sensing surface, although the amount of charges for the target DNA is constant. These results demonstrate that the detection sensitivity is significantly dependent on the charge layer distance. Theoretical analysis for varying charge layer distances of target DNAs will be also studied and compared with the experimental results.

4:00pm BI-MoA7 Effect of Analyte Flow Rate on the Sensitivity of Microcantilever Biosensors, R.P. Desikan, C.W. Van Neste, T.G. Thundat, University of Alberta, Canada

The past decade has witnessed the use of microcantilevers as mechanical transducers of molecular recognition and for the development of miniaturized and sensitive biochip platforms. Microcantilever based biosensors can be based on either mass adsorption or surface stress variation. The potential to operate a microcantilever sensor in liquid medium with extreme sensitivity makes it an ideal choice for the development of biological sensors. Selective detection is obtained by immobilizing receptor molecules on one side of the cantilever. Microcantilever based biological sensors predominantly operate in a liquid environment; this is done in order to retain the functionality of the biomolecules immobilized on the surface. Typically reference cantilevers serve to preserve and monitor the effect of non-specific interactions on the flow rate from specific biomolecular interactions. We have additionally observed that the interaction of analytes on the functionalized surface of the cantilever is influenced by the varying the flow rate of the solution used. The surface stress observed due to the adsorption of analyte molecules on to the receptors on cantilever surface in static condition is much higher compared to a dynamic condition where analytes are allowed to flow across the cantilever surface with the help of a flow control system. Here we address the effect of flow rate on the biomolecular adsorption kinetics of the system and how it affects the sensitivity of Microcantilever based biological sensors.

4:20pm BI-MoA8 Fabrication of Nanowire FETs for pH Sensing, C. D’Emic, S. Zafar, A. Afzali, B. Fletcher, T. Ning, M.A. Guillorn, D.-G. Park, IBM T.J. Watson Research Center

Sensors for measuring pH are very important for understanding reactions of biological species such as proteins, enzymes and cells. While traditional sensors based upon such techniques as infrared spectroscopy, fluorescence and others have low sensitivity and slow response time, more recent nanowire field effect transistor sensors offer improved sensitivity and response time due to smaller size and increased surface areas. [1, 2]

We have fabricated nanowire FET sensors using conventional CMOS semiconductor processes. The nanowires were patterned by electron beam lithography and reactive ion etched into 30 nm thick silicon dioxide (SOI) substrates. The gate sensing surface is comprised of a hafnium oxide/silicon dioxide stack covering the nanowire, while the source/drain regions are comprised of boron activated SOI with nickel-platinum silicide contacts. The resulting 16 nm wide nanowire devices show high sensitivity for pH measurements. The FET drain current increases by a factor of eight per unit change in pH, while the subthreshold slope is ~77 mV/decade. The sensors operate at a reduced sensing voltage of 0.5 V, making them promising candidates for low power, bio-medical applications.


4:40pm BI-MoA9 La$^{3+}$ doped TiO$_2$ Nano-engineered Platforms for Biosensor, R.R. Pandey, Centre for Cellular and Molecular Biology, India, K.K. Saini, National Physical Laboratory, India, M. Dhaival, Centre for Cellular and Molecular Biology, India

The use of nanotechnology tools has opened new opportunities to explore analytical applications of the nano-engineered materials which attracted great attention due to their unique morphology, extraordinary physical and chemical properties towards development of biosensors to facilitate the improvement of the selectivity and sensitivity of the current methods. In this work, the importance of semiconductor La$^{3+}$ doped TiO$_2$ a metal-oxide-based nanostructure platform is highlighted for biosensors platforms. La$^{3+}$ doped TiO$_2$ nanostructure having nanometer-scale inner-core cavity which were exposed to the outer surface with different oxidation states having possibilities for redox-activity can make them attractive for sensing uses. Therefore, the use of La$^{3+}$ doped TiO$_2$ for the development of electrochemical sensors will be discussed. These platforms has been characterized by XRD, XPS, FTIR, SEM, cyclic voltametry to determine structural and surface chemical information, respectively, for biosensor applications. Cholesterol oxidase immobilized onto La$^{3+}$ doped TiO$_2$-based nanostructured surfaces exhibited a pair of well-defined and quasireversible voltammetric peaks in CV measurements. We will also discuss the potential prospect of these surfaces as low cost stable platforms for biomedical diagnosis.

5:00pm BI-MoA10 Spray Deposition of Functional Antibody Films, J. Figueroa, S. Magana, D. Gomez, D.V. Lim, R. Schlaf, University of South Florida

Antibody films for the use in biosensors and assays are usually deposited via wet-chemical attachment methods. The presented experiments demonstrate that pneumatic spray deposition of antibody thin films from aqueous solution yields films of similar sensitivity and durability without special surface treatments and attachment chemistries. The experiments were performed using a commercially available low flow nebulizer in combination with a syringe pump and a substrate rotation stage to homogenize the coating.

In the experiments E. coli O157:H7 antibody was deposited on cleaned microcopy glass slides without any other pre-treatment. Standard wet-chemically prepared silanized glass slides using the avidin-biotin attachment scheme were also prepared for direct comparison of sensitivity and longevity of the spray vs. substrates. After incubation with GIP-labeled E. coli O157:H7 cells (ATCC 35150) ranging from 10$^3$-10$^6$ CFU/ml the slides were rinsed and AF647-labeled antibody detector was added and incubated. After rinsing and drying the slides were interrogated with a 635 nm laser and visualized using a CCD camera. Slides were also visualized by epifluorescence microscopy to examine antibody patterns and determine E. coli capture efficiencies.

The results of the experiments demonstrate that there is little difference between spray and standard protocol wet-chemically prepared substrates. This indicates that antibody films can be prepared via physiosorption without complex attachment chemistries, and that antibodies can directly attach to glass slides, while retaining their functionality.


Genome-wide analysis of single cells is important in life science research and modern medicine in applications ranging from cancer diagnosis to understanding tissue development. Microfluidic devices have been explored as a promising platform for single cell studies, providing superior handling of minute sample and reagent volumes in engineered microstructures. Isolation of nucleic acids from biological samples is an essential step of every type of genomic analysis. While numerous isolation methods have been explored, it remains rather challenging to isolate and analyze genomic DNA from small cell populations and individual cells. Traditional microfluidic devices utilize solid phase extraction (SPE), a method based on binding of DNA to chemically functionalized solid phase matrices for separation of nucleic acids from cell lysates. The binding affinity is sensitive to factors such as pH, temperature, and buffer composition which must be controlled, often dynamically, to minimize DNA losses. Even when the extraction process is optimized, it is difficult to ensure that all of the DNA fragments are adsorbed on the solid phase matrix and that the whole genome is represented in the purified extracts. An appreciable fraction of genomic DNA is often lost during the purification process when the cell debris is washed away. Additional DNA losses can be caused by incomplete elution of the-art microfluidic devices for DNA from cell debris and cell lysates exhibit rather modest extraction efficiencies of 60-85%. This is sufficient for genetic analysis of cell populations because multiple copies of every gene are present in the extract, which statistically guarantees complete genome coverage, but such losses are hardly acceptable when single-copy genes in a single cell need to be investigated. This work describes a valveless two-channel microfluidic nanofluidic device for efficient isolation and fluorescent analysis of DNA contents of single cells. Long strands of human chromosomal DNA released from the cell by chemical lysis loop around PDMS micropillars and are physically retained while the remaining cellular contents are washed away under hydrodynamic flow. DNA fragmentation is minimized by operating at low flow rates. Hydrodynamic entrainment of DNA in non-functionalized obstacle array allows separation of very large genomic DNA from cell debris and components such as proteins and membrane fragments as well as from much smaller mitochondrial DNA and RNA. The purified DNA was subsequently released from the device by enzymatic fragmentation with restriction endonucleases under continuous flow and collected for fragment-size analysis and evaluation of the extraction efficiency. Fluorospectrometric measurements indicate that the microdevice extracts
Electronic Materials and Processing Division
Room: 209 - Session EM1-MoA

Group III-Nitrides and Hybrid Devices
Moderator: K. Kavanagh, Simon Fraser University, L. Porter, Carnegie Mellon University

2:00pm EM1-MoA1 Electrically-Monitored Gate-Recess for Normally-Off AlGaN/GaN High-Electron Mobility Transistors. H. Kim, M. Schuette, W. Lu, The Ohio State University

GaN-based power devices have been intensively investigated for high power switching applications as well as high power microwave applications. Particularly, high breakdown voltage and high saturation velocity of GaN-based heterostructures facilitate reduction in on-state loss and switching loss compared to currently dominant Si-based power devices. Moreover, normally-off GaN-based power field-effect transistors (FETs) offer their inherent safety, reduced power consumption, and diverse circuit functionality with normally-on ones. Recently, we developed a zero-bias and laser energy. The concentration profile of In_xGa_1-xN/GaN MQW obtained from our Cl-based gate-recess for target Gm,MAX of 300 mS/mm. In addition, our recess approach has been implemented, which provides additional process control templates and/or sapphire substrates. In this study, the separation times $S_1$ and $S_2$ affect the growth of InN alloys at higher growth temperatures. In order to reduce gas phase fluctuations from 0 to 10%. Both heterostructures were grown on chemomechanically polished (0001) GaN substrates. Calculations using temperature-dependent photoluminescence spectra revealed a four-fold behavior on N-polar vs. Ga-polar GaN. Furthermore, based on isocantention surface analysis and proximity histograms the upper surfaces of InGaN QWs appear to be more flat than the lower surfaces. These results indicate a surface roughening of the InGaN layer. A detailed comparison of the two structures will be presented and the ability of 3-D atom probe tomography for such an analysis and the impact of the results on next generation LED technologies will be discussed.


2:40pm EM1-MoA3 Local Surface Electrical Characterization of Polar GaN Surfaces. J.D. Ferguson, M.A. Reshchikov, A.A. Baski, Virginia Commonwealth University

While GaN is a widely-used material in optoelectronic devices, localized surface-related electrical properties are not well-understood. These properties affect the operational performance and lifetimes of GaN-based devices. Here, several atomic force microscopy (AFM) techniques were used to characterize the Ga-polar, +c [0001], and N-polar, -c [0001], surfaces of free-standing bulk GaN. Samples were prepared by either a chemical-mechanical polish (CMP) or mechanical polish (MP) of HVPE-grown GaN. AFM data showed that the Ga-polar surfaces (MP and CMP) were uniformly flat with rms roughness of less than 1 nm over a 5x5 micron image. In contrast, the N-polar surfaces were significantly rougher (~5 nm rms) with scratch-like features (100 nm wide, microns long), where the CMP treatment resulted in the presence of surface protrusions (~100 nm dia.) in proximity of the scratches. We then examined the local electrical properties using conducting AFM (C-AFM) to map surface conductivity and to obtain I-V spectra. C-AFM images at forward-bias (<6V) showed small contrast variations for all samples except the N-polar CMP surface. In that case, we observed less conducting behavior on the protrusions as compared to the surrounding surface. Local I-V data also revealed a higher forward voltage turn-on voltage for the N-polar vs. Ga-polar samples. To investigate the local surface charging behavior, we used a two-step technique. First, a metallized AFM tip was used to locally charge the surface by applying a DC voltage, and then the resulting change in surface potential was monitored as a function of time with scanning Kelvin probe microscopy (SKPM). These surface charging data showed a smaller change in surface potential for the N- vs. Ga-polar samples, which appears to be consistent with the lower onset of conduction for the N-polar orientation. We measured the photo-induced changes in surface potential under UV light exposure (100W Hg lamp), otherwise known as the surface photovoltage effect (SPV). The N-polar samples had a smaller SPV compared to Ga-polar, which indicates a smaller amount of band bending at the surface. Additionally, N-polar GaN restored to dark-state conditions at a much faster rate, regardless of CMP or MP treatment. In summary, we observed differences in morphology and electrical behavior for the two polar, c-plane GaN surfaces, as well as differences in behavior due to CMP and MP treatments. These data suggest a less pronounced surface charging behavior on N-polar vs. Ga-polar GaN.

3:00pm EM1-MoA4 The Effect of Ammonia - TMI Pulse Separation on the Structural Properties of InN Epiayers. R. Atlay, Georgia State University, M. Buegler, Technische Universität Berlin, Germany, S. Gamage, I. Senevirathna, A.G.U. Perera, Georgia State University, J.S. Tweedie, R. Collazo, North Carolina State University, N. Dietz, Georgia State University

Over the last decade, significant research efforts have been devoted to understand and improve the properties of InN epiayers. However, even today, there is a significant lack of understanding of the InN surface and growth chemistry and its affect on the physical bulk properties. Crucial challenges in the growth of InN epiayers are the vast different partial pressures between indium and nitrogen and associated fragments of the utilized precursors trimethylindium (TMI) and ammonia (NH3). In this work, high-pressure chemical vapor deposition (HPCVD) is employed and explored in order to control and suppress the dissociation of InN alloys at higher growth temperatures. In order to reduce gas phase reactions and to control the surface chemistry, a pulsed precursor injection approach has been implemented, which provides additional process control parameters for the optimization of the physical epiayers properties.

In the pulsed precursor injection approach, the precursor separation times between the metal organic (MO) sources (TMI and TMG) and ammonia (S1), and ammonia and MO (S2) are two critical process parameters.

This contribution will present results on how the precursor separation times between the TMI and ammonia (S1) and ammonia and TMI (S2) affect the structural and optical properties of InN epiayers grown on GaN/sapphire templates and/or sapphire substrates. In this study, the separation times S1and S2
and S2 have been varied from 1000 to 2500 ms and 250 to 500 ms, respectively. The results show that a longer S1 separation of around 1700 ms and short S2 separation of around 360 ms provide the best structural properties of InN epilayers. The InN epilayers have been analyzed by Raman scattering, X-ray diffraction, IR reflection, and transmission spectroscopy. The Raman studies showed that the lowest FWHM of E2(high) vibrational mode with about 8 cm−1 for a S2 separation time of 358 ms. XRD 2θ−ω scans on these InN epilayers and the analysis of the InN (0002) Bragg reflex confirm the improved structural properties for the optimized S2 separation.

3:40pm EM1-MoA6 Neutron Scattering Provides a New Model for Optimal Morphologies in Organic Photovoltaics: Rivers and Streams, M. Dudman, W. Yin, University of Tennessee, J. Ankner, K. Xiao, Oak Ridge National Laboratory

Organic Photovoltaics (OPVs) have attracted increasing interest as a lightweight, low-cost and easy to process replacement for inorganic solar cells. Furthermore, the morphology of the OPV active layer is critical to its performance, a bicontinuous, interconnected, phase-separated morphology of pure electron donor and acceptor phases is currently believed to be optimal. In this work, we use neutron scattering to investigate the morphology of a model OPV conjugated polymer bulk heterojunction, poly[3-hexylthiophene] (P3HT) and surface-functionalized fullerene 1-(3-methoxyxycarbonyl) propy(1-phenyl [6,6]) C61 (PCBM). These results show that P3HT and PCBM form a homogeneous structure containing crystals of P3HT and an amorphous P3HT/PCBM matrix, up to ca. 20 vol% PCBM. At 50 vol% PCBM, the samples exhibit a complex structure containing at least P3HT crystals, PCBM crystals, and a homogeneous mixture of the two. The 20 vol% PCBM samples exhibit behavior consistent with the onset of phase separation after 6 hours of thermal annealing at 150 °C, but appears to be miscible at shorter annealing times. This suggests that the miscibility limit of PCBM in P3HT is near 20%. Moreover, for the 50 vol% PCBM sample, the interface roughens under thermal annealing possibly owing to the growth of PCBM crystals. These observations suggest a different morphology than is currently presented in the literature for optimal bulk heterojunctions. We propose a novel ‘rivers and streams’ morphology to describe this system, which is consistent with these scattering results and previously reported photovoltaic functionality of P3HT/PCBM bulk heterojunctions.

4:20pm EM1-MoA8 Novel Boron Carbide-Based Semiconducting Polymers for Enhanced Electronic Properties, F. Pasquale, J. Kelber, University of North Texas

We report the controlled modification of valence band electronic structure of semiconducting boron carbides with sharply narrowed band gap, by combination of 1,4 dianinobenzene (DAB) with orthocarboranes to form a novel P3HT/PCBM type novel semiconducting film. Semiconducting boron carbide films (B3C3H4) and related materials formed from crosslinking of boron carbide isoosahedra are of rapidly increasing interest in neutron detection, nanoelectronics, spintronics, and even conventional CMOS ULSI applications. The ability to modify the material's electronic structure in a controlled manner is of obvious interest in all these areas. We report x-ray and ultraviolet photoelectron spectra (XPS, UPS) demonstrating that electron-induced cross-linking of boron carbide and DAB results in a novel semiconductor with a significantly enhanced valence band density of states near the Fermi level, resulting in a shift, as determined by UPS, in the valence band maximum from ~ 4.3 eV to ~ 1.7 eV below the Fermi level. Importantly, this effect is relatively insensitive to total film thickness and for DAB/orthocarborane atomic ratios (determined by XPS) ranging from 1:4 to 1:1. Films were formed by condensation and cross-linking of alternating layers of orthocarborane and DAB precursors under ultrahigh vacuum (UHV) conditions, allowing precise control of relative DAB and boron carbide concentrations. This procedure also yielded films of systematically varied DAB/carborane concentration and film thicknesses between 20 Å and 100 Å average thickness, as determined by attenuation of CuKα photoemission intensity from the substrate. The films exhibited shifts in the valence band spectra of ~ 0.2 eV to lower binding energies upon cross-linking, indicating a surface photovoltaic effect, and the formation of a true p-type semiconducting film. The results demonstrate formation of a new semiconducting material based on boron carbide isoosahedra but modified by the addition of an organic species. These data further indicate the potential for the development of a broad range of novel boron carbide-based polymers using carboranes and other organic additives, with film formation by a variety of methods, including photon or electron bombardment, or plasma-enhanced chemical vapor deposition.


Electronic functionalization of complex fibrous systems is of interest for developing new hybrid electronic systems geared toward integrating biological detection and energy harvesting devices in textile materials. Reliable methods to evaluate the electrical properties of these textiles are necessary for future device design and performance improvement. This work investigates conformal coating of textile substrates produced by atomic layer deposition (ALD) on natural and synthetic fibers structures, resulting in novel hybrid-based electronic structures. A modified 4-probe test method is introduced to evaluate the effective conductivity of these coatings. An applied normal force orthogonal to the current and field direction improves the fiber/fiber contact, resulting in consistent evaluation of the effective conductivity of the coatings across fiber systems and is a universal method of evaluating the mechanical behavior of these hybrid fiber structures. Optimization of the coatings has resulted in conductivity values as high as 40 S cm−1 for zinc oxide coatings (~75 nm) on polypropylene and cotton fiber, as well as 1150 S cm−1 for ALD tungsten (~50 nm) on quartz fiber matrices. Device application of these coated fiber matrices are benefited by their ‘all-fiber’ structure, with characteristic high porosity and surface area. For example, a textile-based flow-through metal-insulator-metal capacitors fabricated from tungsten-coated quartz fibers is shown as an application in liquid chemical sensing. The mechanisms related to electronic transport in a surface-coated textile fabric and implications on device fabrication and improvement will be discussed.

5:00pm EM1-MoA10 Quantum Dot Transfer using Patterned Self-Assembled Monolayers, S. Miller, A.J. Muscat, University of Arizona

Nano particles such as light emitting quantum dots offer many exciting new possibilities for device manufacturing. Among the many advantages of quantum dot manufacturing, the ability to make quantum dot based solar cells and LEDs, has lead to the potential to make arrays of light detectors and nanometer scale patterned LEDs. Using two such devices together results in an optical switch, which may be employed in computer processing as a fast memory readout device. In addition ROM memories can be created using direct patterning of light emitting quantum dots on a disk. Because quantum dots of different size emit different wavelengths it is possible to code data in several wavelengths allowing the data from each wavelength to be written in the size limited area of focused light, dramatically increasing data storage. Many of the potential devices which can be made using nano particles require precise control over nano particle placement and layer thickness. We have found that it is possible to control quantum dot placement using a patterned self-assembled monolayer (SAM). Due to the hydrophobic properties of octadecyltrichlorosilane (OTS) it is possible to form an OTS SAM on which quantum dots will not deposit. This SAM layer can then be patterned using one of several methods. One approach is to use direct UV light exposure in air through a shadow mask, which will remove the OTS SAM in the open areas of the mask, allowing for quantum dot deposition to occur only in the open areas. OTS SAMs can also be formed on pre-made chemical patterns. Ultra high resolution patterns can be formed using nano-lithography including conductive atomic force microscopy, STM, and EBL. Arrays of conductive atomic force microscopic tips can be used to pattern repeated patterns such as those required for detectors. Quantum dots or an alternate SAM can then be deposited directly into the open pattern areas. However, because of the time required for nano-lithography, it is desirable to use the patterned OTS SAM as a nano particle master stamp, where the quantum dots are transferred from the patterned OTS surface to a surface which chemically binds them. A self-assembled monolayer such as APTMS can be used to bind the quantum dots and remove them from the master patterned stamp, allowing the patterned OTS SAM to be repeatedly filled. Quantum dots can be patterned by quantum dot deposition demonstrated using CdTe quantum dots with TGA ligands; however, there are a variety of nano particles with which this technique will work. Therefore by creating a patterned SAM and utilizing selective deposition and appropriate transfer chemistry it is possible to open a new realm of potential device manufacturing.

5:20pm EM1-MoA11 Solution Processed Quantum Dots for Infrared Imaging, J. Lewis, E.J.D. Klein, C. Gregory, G. Cunningham, D. Temple, RTI International

While InGaAs-based focal plane arrays (FPAs) provide excellent detectivity and low noise for short wave infrared (SWIR) imaging applications, wider scale adoption of systems capable of working in this spectral range are limited by high costs, limited spectral response, and costly integration with Si readout circuits (ROICs). RTI has demonstrated a novel photodiode technology based on C3H5I as SWIR (0.7-1.7 μm) detectors with an operating temperature of 75°C and actively dimmed InGaAs FPAs. The detector response is linear from 0.9 to 1.7 μm, with a peak response at 1.6 μm. The dark current is <100 fA and the FWHM of the photoresponse is 150 MHz. A number of characteristics such as high performance, high reliability, and low cost make the InGaAs photodiodes an attractive option for a wide range of applications. The InGaAs photodiodes are further integrated into a 320x256 pixel InGaAs FPA using an ultra-thin liquid phase epitaxy (LPE) InGaAs substrate. The LPE InGaAs substrate is thermalized to 75°C and a backside-on bonding process is used to transfer the FPA to the substrate. The FPA is thinned down to a thickness of 100 μm and the bond between the InGaAs photodiode and the LPE substrate is confirmed using Raman spectroscopy. The LPE substrate is removed and the FPA is mounted and wire bonded to a ceramic package. The ceramic package is then attached to an InGaAs backside-on process to form the final FPA. The InGaAs FPA is characterized and the results are compared to other SWIR FPAs. The InGaAs FPA demonstrates excellent performance and is a cost-effective solution for SWIR imaging applications.
2:00pm  EM2-MoA1 Oxides for Spintronics, K.L. Wang, P. Khalili, F. Xu, University of California Los Angeles

Mainstream CMOS technology in today’s electronics continues to scale down in its feature size. However, power dissipation per unit area and variability post two major issues and challenges for the continuing scaling. Spintronics, an emerging technology that exploits the intrinsic spin of the carriers, could potentially offer power-efficient, low variability and low scalability. In the talk, we will address the importance of functional oxides such as MgO in field controlled spin FET devices and magnetic tunnel junctions.

Toward the realization of spin logic devices, electric-field manipulation of ferromagnetism offers a potential for achieving low power dissipation. The control of collection of spins is critical in accomplishing room-temperature spin-polarized field effect transistors for dilute magnetic semiconductors such as Mnixer or MnGe. We found that by using high-quality MgO as gate oxide, the ferromagnetism of the MnGe thin film can be modulated up to 300 K, which opens the possibility to build room-temperature spin FETs. In addition, MgO was also shown to be effective for unpinning Schottky barrier height and improving the spin injection. Using a epitaxially grown Fe/MgO/Ge tunnel junction, we have obtained single crystalline and atomically smooth Fe/MgO on Ge. This high quality Fe/MgO/Ge junction not only passivates the Ge-Si surface states to favor electronic transport, but also leads to an enhanced spin injection efficiency due to the symmetry induced spin filtering property of the MgO. By using this junction, we show electrical spin injection to bulk Ge.

We also studied the effect of MgO tunnel barrier thickness on the spin-transfer torque-induced switching of CoFeB-MgO-CoFeB magnetic tunnel junction (MTJ) devices used for nonvolatile memory. We studied the effect of MgO thickness on the resistance-area product (RA) and tunneling magnetoresistance (TMR) of the structures using both film-level current-in-plane tunneling (CIP) and device-level electrical transport measurements. The TMR showed a large distribution for RA values lower than 4 Ω-μm (corresponding to an MgO thickness ~0.85 nm), while it increased to ~150% for larger RA > 6 Ω-μm. The results allow for optimization of RA and MgO thickness for low write energy and high-density of charge limited transient currents) to form conducting filaments (leakage current), to study the effect of MgO layer composition and thickness on device performance.

2:40pm  EM2-MoA3 Charge Trap Memories and 3D Approaches. G. Molas, CEA Leti Minace Campus, France

Charge Trap Memories and 3D Approaches

The standard planar Floating gate Flash memory has been scaled down over 20 years. However, many critical limitations are appearing (charge loss through the top or bottom dielectric, cell to cell coupling interference, Random Telegraph Noise, reduction of the number of stored electrons, process induced variability...), making difficult further scaling of the memory device.

In this context, charge-trapping memories, based on the TANOS (TaAl2O5-Un-doped Si-N2O5-SiO2-Si) gate stack, are foreseen as the backbone of future NAND technologies, allowing to reach the 20nm era with planar device structures and to overcome the 1X node when coupled to novel 3D vertical memory architectures. Nevertheless, to face this challenging Flash memory evolution, several process innovations are still required, and an in-depth physical understanding of the gate stack materials property, is needed.

This paper discusses the potentialities and limitations of charge trap memories, and proposes some paths of improvements to fulfill the stringent requirements of future memory generations.

First the engineering of the memory gate stack is investigated. In particular, engineered tunnel dielectrics, alternative charge trapping layers and improved control dielectric stacks are proposed, and their impact on the memory performances and reliability is debated. Experimental results are analyzed by means of models and simulations.

Then in a second part, the integration of charge trap memories in 3D architectures is studied. The various approaches investigated in the literatures are reported, and an original method to process stacked 6nm crystalline nanowires with gate all around SONOS configuration is proposed.

5:40pm  EM2-MoA6 A Survey of Cross Point Phase Change Memory Technologies, D. Kau, Intel Corporation

This survey reviews the current advances in phase change memory and the integrated selector. Based on memory cell configuration in array, there are 3 basic array types, including 2-terminal cross point array [1-6], 3-terminal NOR array [7, 8], NAND string [9]. Among all the configurations, stackable thin-film cross point memory delivers the densest array, therefore the most compact size. Combining its attributes in cost, performance and reliability, cross point phase change technologies stimulate potential opportunities in computing memory hierarchy.

Energy Frontiers Focus Topic
Room: 103 - Session EN+EM+NS-MoA

Nanostructured Materials for Third Generation Solar Cells
Moderator: W.A. Tisdale, Massachusetts Institute of Technology

2:00pm EN+EM+NS-MoA1 Fabrication of Two-dimensional Array of Sub-10nm GaAs Nanodisk using Bio-template Neutral Beam Etching Process. S.-H. Lin, X.-Y. Wang, C.-H. Huang, Y. Ohno, M. Igarashi, Tohoku University, Japan, A. Murayama, Hokkaido University, Japan, S. Samukawa, Tohoku University, Japan

Recently, 3-dimensional (3D) or 2-dimensional (2D) quantum dot superlattices are widely investigated to develop the new generation devices, such as quantum dot solar cell. Quantum dot superlattice consists of quantum dots with lower band gap energy in the matrix with higher band gap energy. For the device application, the structure requires high QD density, periodic QD array and uniform dot size and inter-dot spacing. Molecular beam epitaxy (MBE) and Metal-organic chemical vapor deposition (MOCVD) are two attractive methods among various bottom-up fabrication methods to fabricate quantum dots. However, there is limitation of bottom-up process to control the size, spacing and density of quantum dots. To breakthrough these problems, we have proposed the ultimate top-down process by combination of bio-template and damage-free neutral beam etching (NBE) for fabricating defect-free 2D array of quantum dots.

In this study, we developed a series of novel process for fabricating uniform size and high density 2D array of GaAs nanodisk with uniform inter-dot-spacing. Firstly, the hydrogen-radical treatment was used to remove the native oxide on GaAs surface. To utilize two-dimensional array of ferritin (protein including 7-nm-diameter iron core) as an etching mask, the formation of a hydrophilic 1-nm thick GaAs neutral beam oxidation film is key point. It is found that protein shell can be removed with oxygen-radical treatment at a low temperature of 280°C without thermal damage to GaAs. After protein shell removal, the iron oxide cores inside the ferritins remained on the surface. Then, using the iron core as an etching mask, neutral beam could etch the defect-free nanodisc structure of GaAs. Finally, we developed that iron oxide core was removed by wet etching with diluted hydrogen chloride and completed a fabrication process without inflicting any damage to GaAs. The result shows the quantum dot superlattice structure with a two-dimensional array of GaAs quantum dots with a diameter of ~7-nm, a height of ~10-nm, and a quantum dot density of more than 7 x 10^10 cm^-2 was successfully demonstrated without any damage to GaAs.

This work is supported by the Core Research of Evolutional Science and Technology (CREST) of Japan Science and Technology Agency (JST).

2:40pm EN+EM+NS-MoA3 Solution Processed Quantum Dots for Low Cost Photovoltaics. E.J.D. Klem, J. Lewis, C. Gregory, G. Cunningham, D. Temple, RTI International INVITED

For solar energy to be a significant component of our energy supply new technologies are needed that enable the fabrication of low cost, high efficiency solar cells. Research into solar energy devices which incorporate technologies are needed that enable the fabrication of low cost, high performance devices. The band gap of Ge nanowires blue shifts with H passivation and red shifts with CH3 termination. This relationship. Germanium is an ideal system to study because of its large Bohr exciton radius, low rate of oxidation, and chemical similarity to silicon. In this work we controllably synthesize germanium nanowires with diameters from 5 to 20 nanometers using gold catalyst particles with a narrow size distribution. To obtain epitaxial Ge nanowires with uniform diameters and lengths, a two-step growth process is employed that includes a brief, high-temperature nucleation (390°C) above the Au-Ge eutectic point, followed by elongation at various process conditions, generally below the eutectic point (280 – 340°C, 1x10^ -1 – 1x10^-2 Torr). To limit catalyst diffusion and formation of the V3 x V3 Au/Si reconstruction on the Si(111) surface, a key problem for controllable sub-20 nm growth, the substrate is passivated with CH3 groups prior to nanowire synthesis by introducing small quantities of methylgermane. Nanowire surface chemistry is probed with in-situ transmission infrared (IR) spectroscopy both during and after growth. We introduced different adsorbates (-H and -CH3) and monitored their influence on the band gap as a function of surface coverage. We show that the band gap of Ge nanowires blue shifts with H passivation and red shifts with CH3 termination.

4:00pm EN+EM+NS-MoA6 Role of Quantized and Mid-Gap States in “Dark” Charge Transport and Photoconductivity in Semiconductor Nanocrystal Films. P. Nagpal, Los Alamos National Laboratory

Colloidal semiconductor nanocrystals (NCs) have attracted significant interest for applications in solution-processable devices such as light-emitting diodes and solar cells. However, poor understanding of charge transport in NC assemblies, specifically the relation between electrical conductance in dark and under light illumination, hinders their technological applicability. Here, we simultaneously address the issues of “dark” transport and photoconductivity in films of PbS NCs by incorporating them into optical field-effect transistors (OFETs), in which the channel conductance is controlled by both gate voltage and incident radiation. Spectrally resolved photoresponses of OFETs reveal a weakly conductive mid-gap band (MGB) which is responsible for charge transport in dark. The mechanism for conductance, however, changes under illumination when it becomes dominated by band-edge quantized states. In this case, the MGB still plays an important role as its occupancy (tuned by the gate voltage) controls the dynamics of band-edge charges. Our study has broad implications for NC-based electronics and optoelectronics, and specifically suggests that design guidelines for NC devices might be different depending on whether they are intended for operation in dark (diodes and transistors) or under illumination (photodetectors and solar cells).

4:20pm EN+EM+NS-MoA7 Using Surface Chemistry to Modulate the Bandgap of Ge Nanowires. S. Sivaram, M.A. Filler, Georgia Institute of Technology

Small-diameter semiconductor nanowires are highly attractive building blocks for next generation photovoltaic devices because they exhibit highly tunable optoelectronic properties as a result of quantum confinement. Bottom up approaches, such as the vapor-liquid-solid (VLS) growth mechanism, are controllable down to ~20 nanometers but significant challenges exist at smaller lengthscales where property tuning is maximized. Furthermore, it is expected that the optoelectronic properties of quantum-confined nanowires will be heavily dependent on surface chemistry, yet there are no experimental studies that fundamentally probe this relationship. Germanium is an ideal system to study because of its large Bohr exciton radius, low rate of oxidation, and chemical similarity to silicon. In this work we controllably synthesize germanium nanowires with diameters from 5 to 20 nanometers using gold catalyst particles with a narrow size distribution. To obtain epitaxial Ge nanowires with uniform diameters and lengths, a two-step growth process is employed that includes a brief, high-temperature nucleation (390°C) above the Au-Ge eutectic point, followed by elongation at various process conditions, generally below the eutectic point (280 – 340°C, 1x10^ -1 – 1x10^-2 Torr). To limit catalyst diffusion and formation of the V3 x V3 Au/Si reconstruction on the Si(111) surface, a key problem for controllable sub-20 nm growth, the substrate is passivated with CH3 groups prior to nanowire synthesis by introducing small quantities of methylgermane. Nanowire surface chemistry is probed with in-situ transmission infrared (IR) spectroscopy both during and after growth. We introduced different adsorbates (-H and -CH3) and monitored their influence on the band gap as a function of surface coverage. We show that the band gap of Ge nanowires blue shifts with H passivation and red shifts with CH3 termination.

4:40pm EN+EM+NS-MoA8 Production of Multi-milligram Yields of Ternary II-VI Semiconductor Nanocrystals Under Non-coordinating Amine Activated Synthesis. M. Plaisant, P.H. Holloway, University of Florida

Thermolysis of Zn and Cd precursors under a non-coordinating one-pot wet-chemical synthesis has successfully produced a heterogeneous ternary CdZnSnSe alloy with an extensive spectral red shift of 200nm beyond the visible range. Large yields of twenty milligram of the CdZnSnSe alloy are minimally required for inclusion of the material in the active layer of an inorganic/organic photovoltaic device. The facile synthesis of such large yields of ternary semiconductor nanocrystal alloys has not commonly been reported in the literature. Herein we discuss first the thermolytic synthesis of the material beginning from the ZnSe core through the inclusion of the Cd precursor. We then discuss the production of the ternary II-VI semiconductor CdZnSnSe material through a process of non-coordinating

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synthesis with amine-activation to produce the high multi-milligram yields required for device inclusion.


Intermediate band solar cells and impurity photovoltaics have been proposed to achieve high efficiency solar energy conversion by introducing electronic states within the bandgap of the host material. The intermediate electronic states provide enhanced photocurrent through the addition of sub-bandgap optical transitions while maintaining a voltage that follows the host material. Approaches to realize these solar cells include the incorporation of dopants/impurities, dilute alloys, and nanostructures such as quantum dots. Self-assembled quantum dots such as InAs/GaAs possess excellent optical properties and have been applied to numerous optoelectronic devices including demonstration of the intermediate band solar cell concept. The highly radiative transitions in these type-I materials result in short radiative carrier lifetimes – a feature that is desirable for light emitters, but can be problematic for photodetectors and solar cells. Semiconductor nanostructures with type-II band alignment result in spatially separated electron and hole wavefunctions, resulting in reduced oscillator strength and corresponding reduction in optical absorption, spontaneous radiative recombination rate, and increased radiative carrier lifetime. The tradeoff between carrier lifetime and optical absorption may be effectively used to provide a closer match to the generation-recombination rates desired for intermediate band solar energy conversion. In this work, ZnTe/ZnSe type-II quantum dots are proposed for intermediate band solar energy conversion. The theory of intermediate band solar energy conversion in this material system will be presented along with initial experimental results on the epitaxial growth, structural properties, and optoelectronic response of the materials. The ZnTe/ZnSe quantum dots were grown by molecular beam epitaxy, where three-dimensional island formation is observed via strained layer growth in the Stranski-Krastanow growth mode. Low temperature photoluminescence spectra reveal optical transitions from ZnSe (2.8eV), Te$_{se}$ isoelectronic centers (2.6eV), and broad emission in the range of 2.0-2.4eV attributed to type-II quantum dots.

Energy Frontiers Focus Topic
Room: 104 - Session EN-MoA

Industrial Physics Forum on Energy II
Moderator: J.W. Rogers, Idaho National Laboratory, J.N. Hollenhorst, Agilent

2:00pm EN-MoA1 The Role of Nuclear Energy in a Sustainable Energy Scenario, H.F. McFarlane, Idaho National Laboratory INVITED

Only three primary energy sources power the planet for humankind’s benefit. Though seldom identified as such, the most familiar is nuclear fusion, which provides the solar flux to warm the earth, power the renewable wind and water cycles, and drive photosynthesis for plant growth. Ancient carbon bonds, formed over millions of years and stored in fuels and their resource constraints, along with the increasing world energy consumption, have spurred great interest in renewable energy from sources such as wind and solar. However, the power from these intermittent sources is constantly varied, making quite challenging for its use and dispatch through the aging electrical grid. One effective way to smooth out the intermittency is to employ electrical energy storage (EES). As such EES has been widely considered as a key enabler of the future grid or smart grid that is expected to integrate a significant amount of renewable, while providing electricity or “fuel” to hybrid and electrical vehicles. Among the potential technologies are electrochemical energy storage technologies or batteries that are capable of storing a large quantity of electricity and releasing it according to demands. There remain significant challenges however for the current technologies to meet the performance and cost matrices for broad market penetration. This paper offers an overview on varied technologies, in particular batteries, and discusses the status, challenges and research needs.

Graphene and Related Materials Focus Topic
Room: 208 - Session GR+TF+ET-MoA

Graphene: Electronic Properties and Charge Transport
Moderator: L. Colombo, Texas Instruments Incorporated

2:00pm GR+TF+ET-MoA1 Electronic and Magnetic Properties of a Graphene Line Defect, D. Guoyleke, Naval Research Laboratory INVITED

Although graphene exhibits excellent electron and thermal transport properties, it does not have an intrinsic band gap, required to use graphene as a replacement material for silicon and other semiconductors in conventional electronics. The band structure of graphene, however, offers opportunities to develop non-traditional applications. One such avenue is to exploit the valley degeneracy in graphene. In this presentation, I will present a two-dimensional valley filter based on scattering of electrons and holes off a recently observed extended line defect. The transmission probability depends strongly on the valley and the angle of incidence of the incident quasiparticles. Quasiparticles arriving at the line defect at a high angle of incidence lead to a valley polarization of the transmitted beam that is near 100%.

I will also discuss results showing that the extended line defect gives rise to ferromagnetically coupled local moments. This ferromagnetism can be understood from a symmetry analysis of the boundary-localized eigenstates. The symmetry requires that the principal moments couple ferromagnetically both along and across the line defect, leading to approximately 2/3 more spin-up electrons and than spin-down electrons per repeat unit along the line defect.

This work was supported by the Office of Naval Research, directly and through the Naval Research Laboratory.
2:40pm GR+TF+ET-MoA3 Hydrogenation Induced Graphene-Metal Contact: Observation of States at Fermi Level, S. Rajasekaran, Stanford University, S. Kayu, T. Amo, A. Stanford Synchrotron Light Source, F. Yang, D. Stacchiola, Brookhaven National Laboratory, H. Ogasawara, A. Nilsson, Stanford Synchrotron Light Source

Graphene has received tremendous interest due to its unique electronic structure. Manipulating its electronic structure has received considerable interest. Hydrogenating graphene to open a band gap has been proposed and certain groups have demonstrated hydrogenation induced band opening for graphene on metal substrates.

We employed carbon specific soft x-ray spectroscopy (X-ray photoelectron (XPS), X-ray absorption (XAS) and X-ray emission spectroscopy (XES)) and scanning tunneling microscopy (STM) to investigate how hydrogenation changes the geometric and electronic structure of graphene on Pt(111). Graphene growth on Pt(111) is accompanied with Moiré structure due to periodic rippling in the graphene overlayer due to lattice mismatch and weak interaction between graphene and Pt. Hydrogenation leads to complete disappearance of long range order, although STM indicates that ripple periodicity survives even after hydrogenation. We show that hydrogenation of the unit cell of Moiré nano-patterns is accompanied by pinning of the graphene layer to underneath metal substrate. Structural changes involved in the process of hydrogenation induce covalent graphene-metal interaction. Angle resolved XES and XAS make it possible to probe symmetry resolved states in $\sigma$ and $\pi$ bond geometry. XES-XAS indicates that the density of states (DOS) of graphene is very similar to that of graphite. Hydrogenation induces significant changes in the electronic DOS, most of which reflect the formation of C-H $\sigma$ bonds. For disordered hydrogenated graphene, contrary to band opening, we observe states at the Fermi level after hydrogenation which we reason to arise due to graphene metal interaction and localized C-H $\sigma$ bonds. This effect is proposed to be a way to tailor its electronic properties as a possible method to form better graphene-metal contact.

3:00pm GR+TF+ET-MoA4 First-principles Studies of Atomic and Electronic Structure of Graphene on Sn/Ni(111) Surface Alloy, L. Adamska, R.G. Addou, A. Dahal, M. Batzill, I.I. Oleynik, University of South Florida

Graphene-substrate interactions can be exploited to modify the electronic structure of free-standing graphene. Although most of the efforts were directed towards investigation of graphene on pure metallic substrates, less is known about properties of graphene grown on surface alloys. Sn/Ni(111) surface alloy is of particular interest because this structure has the surface lattice constant of Ni(111), i.e. almost a perfect match with graphene’s lattice constant, and its surface is highly corrugated due to the substantially large atomic radius of Sn compared to Ni. Here we present results of first-principles density functional theory investigations of structural and electronic properties of graphene on (3x3)Sn(3x3)R30 Sn/Ni alloy substrate. It was found that the presence of Sn atoms (1/3 monolayer) results in substantial weakening of graphene-substrate interactions, as is evidenced by large graphene-substrate separation. Nevertheless, the electronic structure of graphene is substantially affected by the underlying substrate as is seen on simulated STM images. The theoretical predictions for the atomic and electronic structures of graphene on Sn/Ni(111) substrate are compared with experimental results.

3:40pm GR+TF+ET-MoA6 Charge Transport through Graphene: the Role of Metal Contacts, M.Y. Chou, M.E. Kindermann, S. Barraza-Lopez, Georgia Institute of Technology

Graphene is a flat form of carbon only one-atom thick. Formed by two interlaced triangular sublattices, it has two atoms on its unit cell. Under normal conditions the charge carriers in graphene can be described with an effective single-particle picture. They display an isotropic and linear (i.e., conical) electronic dispersion around the charge neutrality level. This dispersion is described by a Dirac-like equation with a pseudo-spin (related to the sublattice degree of freedom) instead of the standard electron spin. The pseudo-spin is responsible in part for the strongly suppressed back-scattering in graphene that results in coherent quantum charge transport on lengths up to the micrometer scale, making graphene a relevant material for electronic applications. Yet for applications and for electrical measurements metal contacts are patterned onto graphene. Then the two following questions naturally arise: (i) How does graphene interact with these metal contacts? (ii) How this interaction affects the conductance of a metal/graphene/metal junction in comparison with the hypothetical conductance of pristine (and contactless, infinite) graphene?

In this talk I will discuss quantitatively –within a single-particle description– the quantum transport of charges through graphene, with focus on the signatures that metal/graphene interfaces imprint on the conductance features. The metals considered are normal (i.e., not superconducting), and spin unpolarized. A crucial observation is that not all metal form covalent bonds to graphene. The conductance is obtained as a function of the separation between contacts, the width of the junctions, as well as the thickness of the metal layers when necessary. From these calculations we are able to extract the basic physics involved in transport for all normal metals [1]. Our studies aim towards a comprehensive modeling of graphene devices at the quantitative level.[1] S. Barraza-Lopez, M. Vanevic, M. Kindermann, and M.-Y. Chou. “Effects of metallic contacts on electron transport through graphene.” Phys. Rev. Lett. 104, 076807 (2010); S. Barraza-Lopez, M. Kindermann and M.-Y. Chou. "Charge transmission through short two-terminal graphene junctions with normal bonding metal contacts." (In preparation.)

4:00pm GR+TF+ET-MoA7 Layer Number Determination and Thickness-dependent Properties of Graphene Grown on SiC, W. Zhu, C. Dimitrakopoulos, M. Freitag, Ph. Avouris, IBM T.J. Watson Research Center

The electronic properties of few-layer graphene grown on the face of silicon carbide (SiC) are found to be strongly dependent on the number of layers. The carrier mobility is larger in thicker graphene because substrate-related scattering is reduced in the higher layers. The carrier density dependence of the mobility is qualitatively different in thin and thick graphene, with the transition occurring at about 2 layers. The mobility increases with carrier density in thick graphene, similar to multi-layer graphene exfoliated from natural graphite, suggesting that the individual layers are still electrically coupled. The Hall coefficient peak value is reduced in thick graphene due to the increased density of states. A reliable and rapid characterization tool for the layer number is therefore highly desirable. To date, AFM height determination and Raman scattering are typically used since the optical contrast of graphene on SiC is weak. However, both methods suffer from low throughput. We show that the scanning electron microscopy (SEM) contrast can give similar results with much higher throughput.

4:20pm GR+TF+ET-MoA8 Graphene: Scratching the Surface. M. Fuhrer, University of Maryland at College Park

Graphene is of interest for its unique electronic structure: electrons in graphene obey the Dirac equation for massless particles, complete with a two-component spinor degree of freedom that mimics the spin of a relativistic particle. But graphene is also composed entirely of surface atoms, orienting the techniques of surface science to alter its properties. I will discuss experiments which combine ultra-high vacuum (UHV) surface science with electronic transport measurements to understand graphene and the adsorbed species on its surface. Surface science techniques can be used to controllably modify graphene’s properties: potassium atoms can be deposited to form charged impurity scatterers; ice can be deposited to modify the dielectric environment of graphene and tune the electronic charge density, electron-electron interaction and band gap; ion irradiation can be used to create atomic vacancies which act as Kondo impurities. Graphene’s transport properties are extraordinarily sensitive to surface adsorbates, and can be used to detect e.g. correlations in the positions of potassium atoms at concentrations below 1/1000th of a monolayer, and phase transitions in few-monolayer water.

5:00pm GR+TF+ET-MoA10 Metallic and Insulating Adsorbates on Graphene, K.M. McCready, R.K. Kawakami, University of California, Riverside

While several experiments have separately investigated the doping of graphene by metallic and insulating adsorbates, the transition from metallic to insulating behavior of the adsorbates has not yet been explored. We directly compare the effect of metallic titanium (Ti) and insulating titanium dioxide (TiO$_2$) on the transport properties of single layer graphene. The deposition of Ti results in substantial n-type doping and a reduction of graphene mobility by charged impurity scattering. Subsequent exposure to oxygen largely reduces the doping and scattering by converting Ti into TiO$_2$. In addition, we observe evidence for short-range scattering by TiO$_2$ impurities.


Graphene and carbon nanotubes are two of the most promising materials for future applications due to their unique properties. Devices combining the two materials are expected to be particularly advantageous. The interface of carbon nanotubes and various metal electrodes has been previously studied, both experimentally (Chen et al. Nano Lett. 2005; Zhang et al. Nano Lett. 2007) and theoretically (Shan et al. PRB 2004, Zhu et al. APL 2006, He et al. APL 2009). These studies focus on the search for materials with a low p-type Schottky barrier. Hybrid graphene-carbon nanotube structures have been previously demonstrated (Tung et al. Nano Lett. 2009). We propose...
the use of graphene as a electrode material for carbon nanotube based FET devices. To this end the carbon nanotube - graphene contact is investigated with different deposition techniques. Simulations with the help of the Schottky barrier height (SBH) and transport properties. Total energy and electronic structure calculations are carried out with a plane-wave basis set and the transport characteristics are calculated with a localized atomic orbitals basis within the non-equilibrium Green's function framework. We consider only the side-contact geometry, nanotubes laying atop graphene. The SBH for (5,0), (8,0) and (10,0) nanotubes on graphene is calculated. The transport characteristics of a combined graphene - nanotube device are considered. In the case of small diameter nanotubes (~0.6nm) a SBH of ~0.09 eV is found when graphene contacts are used, much lower than the typical reported values (0.3 eV and higher).

In Situ Spectroscopy and Microscopy Focus Topic
Room: 106 - Session IS+AS+SS-MoA

In Situ Characterization of Solids: Film Growth, Defects, and Interfaces
Moderator: M. Salmeron, Lawrence Berkeley National Laboratory

2:00pm IS+AS+SS-MoA1 A New Approach to Defect Evolution Studies – Combined In Situ Experiments and Electron Tomography, J.M. Robertson, J. Kacher, G. Liu, University of Illinois at Urbana-Champaign

Electron micrographs are two-dimensional images capturing specific instances in the evolution of the microstructure and composition as well as the electronic and magnetic state. As these yields no insight as to how the state evolved, post mortem knowledge is used to determine the most likely pathway. This challenge can be addressed by conducting experiments in situ in the transmission electron microscope, which allows direct observation and in some cases quantification of the reactions and interactions responsible for the evolved structure. The information, however, remains two-dimensional and with increasing use of this technique it is becoming apparent that the three-dimensional aspects of defects are underreported and requires new techniques to detect studies and despite its potential it remains a static snapshot. In this talk, I will illustrate how time-resolved deformation studies have improved our understanding of the behavior of dislocations and how this information has informed the development of new models. I will also demonstrate how three-dimensional images yield a better understanding of complex dislocation interactions and configurations. Finally, I will address the challenges faced in combining these two techniques such that three-dimensional snapshots of the evolving microstructure can be acquired periodically.

2:40pm IS+AS+SS-MoA3 Real-time Oxide Growth Characterization using Atomic Force Microscopy, G. Bijlenders, University of Twente, the Netherlands

Complex oxides have attracted great interest since they exhibit a rich spectrum of physical properties such as ferromagnetism, antiferromagnetism, colossal magnetoresistance, ferroelectricity, dielectricity, and superconductivity. Novel heteroepitaxial devices based on these complex oxides, like spin-polarized ferromagnetic tunnel junctions, superconducting devices and piezoelectric devices, have great potential and are currently under investigation in many groups. The nature of the above-mentioned physical properties in complex oxides is determined by very small characteristic length scales, comparable to the unit cell lattice parameters of complex oxide. Because of these small characteristic length scales, growth control on an atomic level as well as understanding of the different mechanisms affecting the growth mode is essential for the fabrication of epitaxial heterostructures.

Two independent processes, i.e., nucleation and growth of islands, play an important role during vapor-phase epitaxial growth on an atomically flat surface. Here, nucleation causes the formation of surface steps and subsequent growth causes the lateral movement of these steps. Both processes are determined by kinetics, since they take place far from thermodynamic equilibrium. These kinetic processes affect the final surface morphology and are, therefore, extensively studied. I will demonstrate the applicability of high-pressure RHEED as well as Scanning Force Microscopy (SFM) to monitor the growth of complex oxides during Pulsed Laser Deposition (PLD). Because of recent developments, SFM is nowadays also used to study dynamic processes, such as thin film growth and surface reaction mechanisms.

We have realized a system, in which SFM can be performed during Pulsed Laser Deposition (PLD). Deposition and force microscopy are performed in one vacuum chamber and via a fast transfer (in the order of seconds) the surface of a sample can be scanned. In our system we take advantage of the pulsed deposition process, because microscopy measurements can be carried out between the pulses. This provides real-time morphology information on the microscopic scale during growth. The transfer mechanism allows switching between microscopy and deposition with a repetition rate of ~500 nm which gives new opportunities to study growth processes. Furthermore, it can provide information if RHEED is not possible, for example during amorphous and polycrystalline growth.

In this contribution, I will highlight recent advances in oxide thin film growth as well as the latest equipment developments.


Auger Electron Spectroscopy (AES) analysis is a surface sensitive technique for thin film analysis, able to detect nearly all elements [1]. Not only can AES help determine the species present at the surface, but AES can also yield information about the chemical bonding [1]. However, this analysis tool has historically been an ex situ technique with a few noted exceptions [2]. Herein we demonstrate the capabilities and usefulness of an Auger probe, the Staib In situ Auger Probe (SIAP) that has a sufficient position accuracy (tested up to 82 nm) so as to not shadow beam fluxes allowing use during growth. The probe leverages an existing RHEED gun as an e-beam source for Auger electron excitation.

The configuration and operation of the SIAP has been described in detail [3]. The tool is installed on an MBE system configured for Terfenol growth, a miscible alloy of TbFe2 and DyFe2. Initial growths performed without Auger monitoring exhibited substantial oxidation even after Ar etched to remove surface contamination. Figure 1. The SIAP was then employed to determine the sources of oxygen contamination. Three sources of O were identified. The growth chamber had a high enough partial pressure of oxygen to oxidize the highly reactive rare earth elements. After 30 hours in the chamber, uncapped Tb showed a substantial increase in the ratio of O to Tb, figure 2. The Tb source material also delivered O and the rate of O delivery increased with cell temperature, figure 3. A final oxygen source was identified, a substantial Tb/SiO2 interface shown in the SIMS data in figure 4. Figure 5 shows the Si wafer with C and O present at the surface. After growth is initiated the C is not detected after 40 Å of deposition, however, the O can be detected until 200 Å are deposited, indicating intermixing with the SiO2 interface. Further tests were conducted starting with a layer of Dy and depositing part of a monolayer of Tb in 2% increments. Figure 6 shows a clear distinction between bare Dy and 2, 4, 6, 8, and 10% monolayer coverage of Tb on a Dy layer, demonstrating the SIAP’s sensitivity is at least 2% of a monolayer for these heavy elements.

The SIAP is complementary to existing RHEED systems. While RHEED gives information about the crystal structure of the growing film, the SIAP provides chemical information. This is especially useful when working with films without line compositions. The SIAP coupled with a future closed loop control system may enhance growth of films with multiple oxidation states or other similar phase/chemical transitions. Finally this technique could give information about transitions between multilayered films grown via MBE.

4:00pm IS+AS+SS-MoA7 Quantum Size Effect Driven Structure Modifications of Bi-films on Ni(111). T.R.J. Bollmann, R. van Gastel, H. Zandvliet, B. Poelsema, University of Twente, The Netherlands

We have investigated the initial growth of Bi/Ni(111) using Low Energy Electron Microscopy (LEEM) and Selected Area Low Energy Electron Diffraction (μLEED). Bismuth represents an interesting material since 1) it has a tendency for allotropism, 2) it forms several ordered alloys with Ni, and 3) with Bi being a neighbor of Pb in the periodic system, one may find evidence for quantum size effects in ultrathin Bi layers. Indeed we obtain ample evidence for Bi/Ni(111) as being a very rich system, even at a fixed substrate temperature of 474 K.

We find first that the deposition of Bi leads to the formation of a surface alloy with a (\sqrt{3}x\sqrt{3})-R30° structure at a Bi-coverage of 1/3. Continued Bi deposition leads to the formation of an incommensurate wetting layer with a complex periodicity, finally ending in a (7x7) structure. From the variation of the step position at the buried interface, nicely accessible with LEEM, we conclude that the dealloying of the Bi takes place during the growth process. For thicker Bi layers we find that the (7x7) structure becomes the dominant structure, while the (7x7) structure remains stable for thinner Bi layers. The substrate temperature of 474 K is the result of a compromise between the need to have a fixed substrate temperature and the need to have a Bi coverage which is low enough to not have a Bi NP layer present instead of the (7x7) structure. In this contribution, I will highlight recent advances in oxide thin film growth as well as the latest equipment developments.
Bi deposition elongated, 3-4 layers high nanowires emerge, with a p(5x2) structure and a width of about 80 nm, oriented along <110> and <100> azimuths. Further deposition of Bi-leads to different (sometimes coexisting) structures: (3x3)-patches with a thickness of three atomic layers and patches with a matrix structure ($m_1=3$, $m_2=1$, $m_3=1$, $m_4=2$) and a thickness of five atomic layers. This accurate height assignment is uniquely enabled by the analysis of LEEM-IV data.

The results are fully consistent with quantum size effect driven thin film morphology: the different film structures and their thicknesses nicely fit with integer numbers of nodes in their specific Fermi wave function, even for the thin layers that were thick at a lower temperature of 422 K. Tensor LEED calculations of the interlayer spacing of the different structures concur with this assignment.

The influence of the structure and morphology on electronic properties of various materials is well known. The interaction between electronic and crystal structure should be reciprocal. The Bi/Ni(111) system provides a nice and we think first illustration: electronic properties, in particular quantum size effects, actually drive the structure of the thin bismuth films.

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**Marine Biofouling Focus Topic**

**Room:** 105 - Session MB+B1+PS-MoA

**Marine Antifouling Coatings**

**Moderator:** A. Rosenhahn, Karlsruhe Institute of Technology, Germany

2:00pm MB+B1+PS-MoA1 Advances in Sustainable Technologies for the Prevention of Marine Biofouling. R. Deshmukh, University of Texas at Arlington, P. Sheth, University of North Texas Health Science Center, R.B. Timmons, University of Texas at Arlington, J.A. Schetz, University of North Texas Health Science Center

The cost associated with preventing the attachment of marine organisms to underwater surfaces (biofouling) is billions of dollars annually impacting numerous sectors including shipping, aquaculture, offshore drilling, and offshore tidal and wind power. Invariably the most effective biofouling control methods are ones that are biocidal in nature whereby a coating is impregnated with a toxin that leaches out over time. However, persistence and bioaccumulation of heavy metal-based and organic biocides, creates environmental and health problems that have resulted in their regulation. Vigorous hull cleaning, aimed at periodically removing biofouling organisms, disperses fine biocide-impregnated coating particles and this significantly increases biocide persistence in sediments where they are ingested by a variety of marine organisms and bioaccumulate.

Coatings that peel off under conditions of high shear, so called foul-release coatings, are another approach but such coatings are easily damaged, not amenable to grooming and ineffective against microfouling. Further, the most effective foul-release coatings are silicon-based and it is becoming increasingly apparent that leaching of toxic silicon oils from the coating matrix plays a role in their superior antifouling performance. Other approaches have been to develop durable coatings with little antifouling activity but that can withstand repeated intense grooming. One of our long term objectives is to identify coating materials with surface properties discouraging the settlement of biofouling organisms but also durable enough to withstand cleaning.

Here we report on the discovery of a transparent, conformal, pin-hole free, and plasma-polymerizable polymer coating made from an inexpensive monomer material that drastically reduces (>90%) algae (Ulva) sporing settlement and biomass accumulation without any observable toxicity towards Ulva or a non-target brine shrimp species (Artemia). Further the unpolymerized monomer has no antifouling effect indicating that potential leaching of monomer from coated surfaces does not account for the observed antifouling activity. Rather the antifouling effect is believed to rely on the presence of a specific chemical moiety because chemical modification of that moiety abolishes antifouling activity. Testing of other marine antifouling species such as microfouling bacteria (Cellulophaga) and macrofouling mussels (Mytilus) suggests that the antifouling properties of the polymer surface are selective for algae. Potential unique...
Polyelectrolyte Brushes in Water

3:00pm MB+BI+PS-MoA4 Development of a Comparative Protocol for Anti-Fouling Surfaces Based on Polymer Brushes. A. Serrano, S. Zürcher, S. Tosatti, SuSoS AG, Switzerland, N.D. Spencer, ETH Zurich, Switzerland

Marine surfaces are known to accumulate fouling material through the starting point of adhesion and settlement of proteins and cells. This effect can be manipulated through the modification and control of the substrate properties via surface functionalization. This approach has led to successful anti-fouling coatings based on biocidal agents containing copper or zinc compounds.[1,2] The environmental toxicity of these latter materials, however, has increased the demand for less adverse coatings. The use of ultra-thin films consisting of polymer brushes has been considered a promising alternative and many studies have been published in this field.[3-5] None, however, has focused on developing a protocol that allows a reliable comparison between the efficiency of different well-known anti-fouling polymers. This is one of the aims of this work and has been achieved by using a common, azide-terminated monolayer to which different non-fouling polymers, such as PEG, PEOXA, PVP and PVA, have been covalently bound. The different materials were compared by characterizing the structure-property relationship of the formed polymeric brushes. Also investigated was the role of the solvent used in the anti-fouling polymer solution as a key element to better control the surface homogeneity. A thorough analysis of the influence of this parameter on the conformation of the final polymer brush was based on ellipsometry, XPS and imaging ToF-SIMS. Finally, the anti-fouling surfaces were subjected to a comparative biological study by exposure to complex protein solutions and Ulva zoospores, in order to validate the developed protocol.

References:

3:40pm MB+BI+PS-MoA6 Surface Modification of Polymers via Self-Stratification: Decoupling of Bulk and Surface Properties. D.C. Webster, North Dakota State University, INVITED

Since materials interact with their surroundings via their surfaces, controlling the surface properties of a material are of critical importance. For materials to be used to mitigate biofouling, having the correct surface properties is a means to achieve anti-fouling properties to the material and one that is unsuitable for the application. In many cases, the material properties of compositions which yield useful surface properties are not suitable for the bulk properties of the material. Thus, being able to decouple the surface and bulk properties is of interest in many areas. Combining polydimethylsiloxane (PDMS) with other polymer systems generally results in materials covered with PDMS due to its low surface energy and incompatibility with other polymers. However, if the PDMS is not chemically bound into the system it can be easily removed from the surface. Thus, forming a copolymer of the PDMS with the other polymer is required for a durable system. Since polyurethanes are known as tough polymers due to extensive internal hydrogen bonding, combining PDMS with a polyurethane could lead to a material which is tough, but has a low surface energy surface provided by the PDMS. Thus, we have found that incorporating a reactive PDMS into a crosslinked polyurethane system can result in a material which has a low surface energy which is stable when immersed in water. High throughput screening has been used to aid in the identification and optimization of PDMS molecular weight, composition of end groups, and the amount of PDMS in the coating. A unique coating was also identified having discrete domains of PDMS on the surface. Coatings having good fouling-release properties have been prepared and tested in ocean immersion testing.

4:20pm MB+BI+PS-MoA8 Antifouling Behavior on the Surface of Polyelectrolyte Brushes in Water. M. Kobayashi, M. Terada, Jst, Erato, Japan, A. Takahara, IMCE, Kyushu University, Japan

Nature utilizes super-hydrophilic surfaces under wetted state by water to achieve oleophobicity and self-cleaning behavior. For example, fish can maintain a clean body surface by surrounding a thin layer of mucus containing calcium phosphate and protein, which protects oil attachment, marine fouling, and adhesion of marine organisms. In this study, high-density hydrophilic polymer brushes were prepared on Si-wafer by surface-initiated controlled radical polymerization of methacrylate monomers with ionic functional groups.[1] For example, poly[3-sulfopropyl methacrylate potassium salt] (PSPMK) and poly[2-[methacryloyloxy]ethyl phosphorylcholine] (PMPC) brushes with 50 - 100 nm thickness repelled both of air bubble and hexadecane droplet in water.[2] Even when the silicone oil was spread on the polyelectrolyte brush surfaces in air atmosphere, once the oil-sitting brush substrates were immersed in water, the oil quickly rolled up and detached from the brush surfaces due to the low adhesion force between water and oil caused by the low surface energy of polyelectrolyte brushes to water. Similar oil detachment behavior was observed on the hydrophilic poly(sodium methacrylate) (PMANa) and poly[3-[dimethyl(2-[methacryloyloxy]ethyl)ammonium] propanesulfonate (MAPS) brushes,[3] whereas the oil still remained attached on the hydrophobic poly(2-perfluorooctylstyryl acrylate) (PFA-CS) brush even though the PFA-CS brush showed relatively oleophobic properties under air atmosphere. These hydrophilic brush surfaces would contribute to the self-cleaning, antifouging, and antifouling properties without any surfactants.


4:40pm MB+BI+PS-MoA9 Non-fouling Polymer Chemical Gradients for the Investigation of Marine Bioadhesion. O. Sterner, ETH Zurich, Switzerland, S. Zürcher, SuSoS AG and ETH Zurich, Switzerland, S. Tosatti, SuSoS AG and ETH Zurich, Switzerland, N.D. Spencer, ETH Zurich, Switzerland

The accumulation of marine organisms on submerged man-made structures (referred to as marine biofouling) has great economical and environmental impact [1]. Numerous strategies to prevent or lower the extent of marine biofouling have been developed, ranging from biocidal coatings to coatings that either prevent adhesion, lower the strength of adhesion or combinations thereof [2-4]. Surface gradients offer a high-throughput approach to investigate the potency of such coatings, and have the additional advantage of reducing the error in experiments by replacing a set of single samples, including positive and negative controls, with a single substrate [5]. In this project, polymeric ultrathin coatings have been prepared using a versatile surface functionalization system based on a self-assembled monolayer of poly(allyl amine) grafted with photo sensitive perfluorophenyl azide functional groups. Gradients have been prepared using a straightforward approach to control the extent of azide to nitrile conversion over the surface, forming a polymer density gradient. Gradients of poly(ethylene glycol), poly(2-ethoxy-2-oxazoline) and poly(vinyl pyrrolidone) have been prepared and investigated for non-fouling action against zoospores from green macrofouling algae Ulva and two strains of marine bacteria. The gradients reveal a drastic reduction in bacterial adhesion at low polymer densities for all polymers investigated. Gradients have been characterised with variable angle spectroscopic ellipsometry (VASE) and the properties of the polymer coatings have been evaluated with time-of-flight secondary ion mass spectroscopy (ToF-SIMS) and XPS.

References:

5:00pm MB+BI+PS-MoA10 Development of Poly(silyl urethanes) with Tethered Quaternary Ammonium Biocides as Antifouling Marine Coatings. P.N. Coneski, N.K. Weise, J.H. Wynne, Naval Research Laboratory

Due to the significant economic burden posed by high operational and maintenance costs of biofouled ships, the design of environmentally benign, antifouling marine coatings has been a significant interest for many researchers since the ban on traditional coatings was enacted. Current approaches for developing new marine coating materials have primarily focused on the preparation of low surface energy materials, such as modified polydimethylsiloxane polymerized from photo-initiated polymers. These materials have shown great promise for reducing the adhesion strength of various fouling organisms, thus allowing hydrodynamic forces to release the fouling as the ship moves through water. Unfortunately, no existing material has been shown to eliminate adhesion of all of the numerous...
different fouling organisms, including zoospores, microalgae and diatomaceous species. As such, the development of marine coatings with multiple mechanisms of fouling prevention may be an important avenue of antifouling materials research. Polysilyl urethanes coatings have been developed via the reaction of various quaternary ammonium modified orthosilicates with polyisocyanates. Material properties including glass transition temperature, surface energy, and thermal stability have been investigated as a function of orthosilicate and polyisocyanate composition as well as inclusion of low energy soft segments. The low surface energy of these materials should provide excellent fouling release properties, as has been seen with other PDMS-like materials, while the inclusion of bound quaternary ammonium biocides should further reduce the adhesion and propagation of fouling organisms at the material interface. Finally, the hydrolysable silyl ether crosslinking may prove to enhance the antifouling capabilities of these materials by allowing any fouled portions of the coating to slough away over time, generating a new active biocidal interface.


We investigate a novel device that features a reduction in frictional resistance and antifouling hull surface of seagoing ships which are activated electrochemically by a series of porous Pt-TiO2/Ti electrodes. This device includes a series of anodic and cathodic porous Pt-TiO2/Ti electrodes insulatedly mounted on the hull surfaces of which are electrically connected to a direct current (DC) power supply. The above-mentioned porous Pt-TiO2/Ti electrodes are fabricated by two steps; porous TiO2/Ti plates are firstly prepared on pure titanium plates via a micro-arc oxidation technique and Pt nanoparticles are thereafter deposited on the porous TiO2/Ti samples using magnetron sputtering. The DC power supply provides an adequate DC bias to the Pt-TiO2/Ti electrodes where hydrogen and oxygen microbubbles are electrochemically nucleated and formed herein in seawater. As a result, the microbubble-induced superhydrophobic surfaces are created and the reduction in frictional resistance and antifouling hull surface are obtained. Our preliminary tests reveal that no attachment organisms are found on the Pt-TiO2/Ti electrodes after 15 days of field seawater tests at an applied potential of 1.2 V and an energy consumed rate of about 4 W/m2. On the other hand, organisms, mainly Crassostrea gigas and barnacles, attached and grew on the Pt-TiO2/Ti electrode that was not applied a DC potential. That is to say that attachment of organisms can be prevented. Furthermore, the Pt-TiO2/Ti electrodes yields a gas production rate of 800 cm3/min m2 by electrolysis of seawater at an energy consumed rate of about 18 W/m2. Hence, the microbubble-induced superhydrophobic surfaces can be realized with this gas production rate for drag reduction of the ship hull.

Nanomanufacturing Science and Technology Focus Topic
Room: 207 - Session NM+MS-MoA

Challenges Facing Nanomanufacturing (All Invited Session)
Moderator: S. Rosenthal, Vanderbilt University, S. Butler, Texas Instruments Incorporated

2:20pm NM+MS-MoA2 Sustainable Nanomanufacturing. M. Roco, National Science Foundation

INVITED Nanomanufacturing has been defined as an approach to design, produce, control, modify, manipulate, and assemble nanometer-scale elements or features for the purpose of realizing a product or system that exploits properties seen at the nanoscale. Nanomanufacturing R&D has as its goal enabling the mass production of reliable and economical nanoscale materials, structures, devices, and systems. The current relatively rudimentary capabilities for systematic control and manufacture at the nanoscale are envisioned to evolve faster after 2011 as we develop new models and instrumentation and enter production of nanosystems for revolutionary new products and processes. We have estimated the global market of final products that incorporate nanotechnology increases by about 25 percent per year reaching $1 trillion by 2015. This estimation made in 2000 [1] holds in 2011, after passing two thirds of the interval.

Three challenges of nanomanufacturing will be discussed: supporting innovation (beyond scaling), realizing efficiency (beyond new functions) and sustainability (of nanoscale processes and of global development). The research trends and application opportunities in nanomanufacturing will be presented by considering four generations of products by 2020 [2]. Most of what has already made it into the marketplace is in the form of “First Generation” products (passive nanostructures with steady behavior) and more recently “Second Generation” (active nanostructures, such as advanced transistors, amplifiers, targeted drugs and chemicals, sensors, actuators, and adaptive structures), while embryonic “Third Generation” (nanosystems, such as bio-assembling; networking at the nanoscale, nanoscale robotics and multiscale architectures) products are in the pipeline. Concepts for the “Fourth Generation” products, including molecular nanosystems, are only in research. Convergence with modern biology, digital revolution, cognitive sciences and other areas is expected to accelerate nanotechnology manufacturing. The role of NNI Signature Initiative on Sustainable Nanomanufacturing will be discussed.


Vanadium dioxide (VO₂) films and nanostructures in contact with gold (Au) or silver nanostructures can form the building blocks of active metamaterials that can be modulated in response to various stimuli such as the presence of chemical agents, changes in temperature, or irradiation. The semiconductor-to-metal phase transition (SMT) of VO₂, which can be induced thermally (T = 68°C), optically, or electrically — leads to a change in the dielectric function of the film. Since the localized surface plasmon resonance (LSPR) of the metal nanoantenna is sensitive to the local dielectric environment, the SMT allows the optical response of the metal:VO₂ nanocomposite to be tuned. Thus these Au:VO₂ nanocomposites are unique probes of strong-correlation physics because, during the phase transition, the electron-electron interactions in VO₂ that drive the SMT are coupled with the plasmonic excitation of the Au nanostructure.

We fabricated arrays of Au nanoparticles (NPs), 180 nm in diameter and 20 nm high on indium-tin-oxide coated glass by electron-beam lithography. Subsequently, the nanoparticle arrays were coated with a 60 nm VO₂ film by pulsed laser ablation of vanadium metal targets in 10 mTorr oxygen (O₂) background gas, then annealed for 45 minutes at 450°C in 250 mTorr of O₂. Using Peltier hot and cold plates and thermocouples, the VO₂ film could be cooled to just above the SMT and rapidly cycled between the frozen and conducting states. The LSPR wavelength of the NPs was 1000 nm in the semiconducting state and approximately 840 nm in the metallic state, thus overlapping the VO₂ electronic transitions from the occupied vanadium 3d₅ band to the empty 3dₓᵧ band at approximately 885 nm. The film undergoes the SMT, the split 3d₅ bands merge and, with the 3dₓᵧ band, form the metallic VO₂ conduction band. Since the Au NPs are sensitive to changes in both the real and imaginary parts of the VO₂ local dielectric function, they serve as a direct probe of the SMT. The results show a 30% decrease in plasmon dephasing time during the transition due to an increase in carrier-carrier scattering in the VO₂. Both Maxwell-Garnett and Bruggeman effective-medium theories predict the decrease in dephasing time during the SMT; however, a linear theory is a more accurate model for the hysteresis in the LSPR wavelength and linewidth during the SMT.

The LSPR wavelength of the NPs was 1000 nm in the semiconducting state and approximately 840 nm in the metallic state, thus overlapping the VO₂ electronic transitions from the occupied vanadium 3d₅ band to the empty 3dₓᵧ band at approximately 885 nm. The film undergoes the SMT, the split 3d₅ bands merge and, with the 3dₓᵧ band, form the metallic VO₂ conduction band. Since the Au NPs are sensitive to changes in both the real and imaginary parts of the VO₂ local dielectric function, they serve as a direct probe of the SMT. The results show a 30% decrease in plasmon dephasing time during the transition due to an increase in carrier-carrier scattering in the VO₂. Both Maxwell-Garnett and Bruggeman effective-medium theories predict the decrease in dephasing time during the SMT; however, a linear theory is a more accurate model for the hysteresis in the LSPR wavelength.

NS-MoA2 Metamaterial Nanosensors based on the Metal-Insulator transition in VO₂. K. Appavoo, R.F. Haglund Jr., Vanderbilt University
The use of solid-solid phase transitions to modulate the plasmonic response of metal nanostructures is a promising approach to nanophotonic technologies, including sensors based on signal modulation in confined nanoscale volumes [1]. Consideration of phase-transforming materials has typically focused on composition, whereas relatively little attention has been paid to the question of size dependence in determining stable phases. However, size effects play a crucial role in determining the coupling with mechanical, optical, chemical or thermal input required to effect the phase transformation [2]. With rapid progress in nanofabrication techniques, size-dependent properties become relevant and systematic studies to assess both the role of nucleation in forming a new state and of the nanoscale dynamical effects are needed.

Here, we describe an example that shows how, by systematically varying the gap between the arms of split-ring plasmonic resonators, the in-arm coupling resonance in a split-ring metamaterial can be used to monitor the metal-insulator transition in discrete volumes of the strongly correlated VO₂. Moreover, this “plasmonic hysteresis” technique also provides a means to correlate the electronic phase-transition with its structural counterpart which was previously measured using SERS technique [3]. If the number of intrinsic nucleation sites is directly proportional to the interrogated volume (a reasonable assumption) [4], we have effectively shown that well-crafted plasmonic metamaterials with well-understood modes can be helpful to probe size-dependent properties. Full field 3D finite-difference time-domain simulations show that the physical origins of these non-isotropic electronic oscillations leads to concentration of the electromagnetic energy for focused interrogation and high sensitivity.

As an additional example, we briefly describe an investigation into the use of similar nanostructures as chemical sensors based on coupling of atomic vibrations to the VO₂ interface and molecular absorption moieties. In this case, the detection method involves the change in optical transition of a metamaterial array incorporating VO₂ when the heat of decomposition is sufficient to initiate the metal insulator transition.

2:40pm NS-MoA3 Enhanced Photoluminescence from Gd₂O₃:Eu³⁺ Based Core/Multi-shell Nanoparticles. J. Choi, M.R. Davidson, P.H. Holloway, University of Florida
Core-shell and core-multi-shell nanoparticles with luminescent Gd₂O₃:Eu³⁺ were successfully synthesized by a high boiling-point alcohol (polyol) and solution precipitation methods, respectively. The hetero-structured nanoparticles with Eu doped Gd₂O₃ exhibited intense metal to semiconductor plasmonic (PL) from Eu³⁺ after calcination at 600 °C for 2h in air. Photoluminescence excitation (PLE) data showed that while a small fraction of the emission originated from direct excitation of Eu³⁺, most of the excitation resulted from adsorption in the Oxygen to Europium charge-transfer band (CTB) between 225 and 275 nm. Gd₂O₃:Eu³⁺/Y₂O₃ core/shell nanoparticles exhibited PL intensities up to 40% larger than from bare Gd₂O₃:Eu³⁺ nanoparticles and SiO₂/Gd₂O₃:Eu³⁺/Y₂O₃ core-multi-shell samples showed quantum yield (QY) up to 72% larger than that of SiO₂/Gd₂O₃:Eu³⁺ core/single-shell nanoparticles. The increased PL and QY were attributed to reduced non-radiative recombination based on longer luminescence decay time. Potential applications of the nanoparticles as scintillation radiation detectors will be discussed.

3:00pm NS-MoA4 Au₃SiO₄:Yb⁺:Er⁺:Y₂O₃ Coreシェル Optical Nanooantenna: Experiment & Simulation, V. Jankovic, J.P. Chang, University of California Los Angeles
The conversion of electromagnetic (EM) energy from free propagating radiation to localized electromagnetic fields in an optical nanoantenna in the EM and microwave domains is accomplished with the use of antennas. Optical antennas are analogous to their RF and microwave counterparts, but there are crucial differences in their physical properties and scaling behavior because metal is a highly dispersive material with finite conductivity at optical frequencies. Optical antennas are not driven by galvanic transmission lines like RF antennas, instead, localized oscillators such as atomic emitters are brought close to the field point of the antennas, and electronic oscillations are driven capacitatively.

In this work, Au nanoparticles of different shapes (spheres, rods and stars) were used as antenna elements, Er³⁺ ions in a Y₂O₃ host matrix were used as atomic emitter antenna driving elements while the capacitative gap between the antenna element and the atomic emitter was controlled by deposition of an ultra-thin SiO₂ inner layer between the Au nanoparticle and the Yb:Er:Y₂O₃ outer shell. A 4-5nm silica spacer layer was deposited through a controlled TEO₂ hydrolysis reaction and was shown to be effective in preventing quenching yet enabling energy coupling between the Au nanorod and the RE-ion doped oxides. Spatially and compositionally controlled Yb:Er:Y₂O₃ outer shells were deposited using both wet chemistry methods and radical enhanced atomic layer deposition (RE-ALD).

Upconversion (UC) spectral, power dependence and radiative lifetime measurements with 532nm, 750nm 980 nm and 1064nm laser excitation were used to assess the coupling of the Au optical antenna to the emitter ions as a function of antenna shape, spacer layer thickness and spectral and spatial mode overlap efficiency. Preliminary optical characterization showed a 2X earlier onset of upconversion with 980nm excitation for Yb:Er:Y₂O₃ coupled to an Au nanorod antenna compared to pure (uncoupled) Yb:Er:Y₂O₃ nanoparticles. Power dependence measurements with 980nm excitation showed a >5 slope indicating a multi-photon absorption-induced luminescence process. Further work found a >2 slope for the uncoupled erbium, indicating a two photon absorption (expected for erbium with 980nm excitation). These optical antenna core/shell particles have potential application in bio-imaging and light trapping for solar and sensor applications.

Optical cavities can tightly confine light in the vicinity of optical emitters, enhancing the interaction of light and matter. The modes or optical states of the cavity can be precisely designed and engineered, and in recent years there has been remarkable progress in demonstrations of ‘cavity quantum electrodynamics (cQED)’ in solid state platforms. Such progress has been primarily for cavities fabricated in dielectric materials, with a steady improvement in cavity quality factor Q in the range of 10⁶ – 10⁹ realized for cavities with coupled emitters [1],[2]. These high-Q coupled emitter systems have demonstrated heralded single photon emission [3], ultra-low threshold lasing [4] and strong light-matter coupling [5],[6].
Metal-based optical cavities would have inherently lower Q’s (and greater loss) than dielectrics; however, metal cavities utilizing surface plasmon polaritons (SPPs) can have sufficiently small mode volume to produce a substantial Q/V, the quantity relevant for high Purcell factors, a measure of the light-matter interaction. This talk will focus on such plasmonic cavities, with optical modes formed within the gap of the two metal layers which defined the cavity [7]. Initial structures comprised silver (Ag) nanowires (NW), 70 nm in diameter and 1 - 3 µm in length, placed into close proximity to a Ag thin film substrate, with the NW axis parallel to the substrate surface. Optically active material was interposed between the nanowire and the Ag substrate: this comprised one to two monolayers of PbS colloidal quantum dots, clad on top and bottom by thin dielectric layers of varying composition and thickness. The fluorescence spectrum of PbS quantum dots within the gap was strongly modified by the cavity mode, with peak position in quantitative agreement with numerical calculations, and demonstrating Q values of ~ 60.

Such plasmonic cavities allow the easy incorporation of a variety of light-emitting active areas, and we have also explored the incorporation of various organic, dye-containing layers within the gap-mode plasmonic cavities. In addition these structures lend themselves to relatively simple modifications of geometry, allowing effective tuning of cavity modes, and also control of modes through the use of photonic crystal geometries, fabricated into metal.

The high Q/V possible for these cavities, and the range of organic and nanocrystalline emitters they can accommodate make these important building blocks for the exploration of light-matter interaction in the solid state.

References
In this talk, we will present two approaches to microplasma-based electrochemistry that we have developed for nanoparticle synthesis. In one approach, microplasmas are formed at the surface of a liquid electrolyte and operated similar to an electrochemical cell with the plasma as the cathode and a solid metal immersed in the electrolyte as the anode [1]. Metal cations in solution such as Ag⁺ are electrochemically reduced by the plasma to solid metal, resulting in the formation of metal nanoparticles without any chemical reducing agent. Alternatively, thin films of metal cations dispersed on a polymeric surface are electrochemically reduced by a rastered microplasma [2]. This configuration allows macroscale patterns of metal nanoparticles to be produced without the need for lithography. Recently, we have extended our patterning method to the reduction of metallopolymers which are novel molecular structures that can be used as a template for metal ion and metal particle formation [3]. This strategy has enabled patterns of metal nanoparticles to be prepared which are beyond lithographic limits. We will discuss our experimental techniques in detail, as well as the properties of the nanoparticles as assessed by UV-Visible absorbance spectroscopy, X-ray diffraction, and transmission electron microscopy.

References:
3. S. W. Lee et al., in preparation.

Low temperature plasma (LTP) treatment of surfaces is a promising path toward sterilization of bacteria [1]. Past works have shown plasma-induced degradation of bacteria [2], but little knowledge exists regarding the plasma-induced chemical modifications in biomolecules that result in inactivation since various plasma species, e.g. ions, reactive radicals, and UV/VUV photons may aid in inactivation. Lipo polysaccharides (LPS) are a main component of the outer membrane of gram-negative bacteria and are difficult to remove from surfaces by conventional methods [3]. LPS is made up of a polysaccharide chain and lipid A and lipid A elicits an immune response in animals [1]. Previous studies have found that adding H2O to an Ar plasma leads to a reduction in the filament currents originating from broad absorption bands of lipid A, namely C-Hx stretching, C=O, and amide bands [4]. This study aims to distinguish the roles of physical sputtering, chemical attack by H-atoms, and plasma-generated VUV. LPS-coated silicon chips were exposed to LTP (Ar, H2, and Ar/H2 mixtures) to explore the effects of plasma species on the optical density during plasma exposure. The real-time data showed that Ar plasmas create a dense film on the sample's surface that decreases in density with H2addition. The films were etched fastest in Ar discharges mixed with ~10% H2 and were slowest in pure H2. Since previous work [4] found that adding H2O to an Ar discharge enhanced sterilization, these results may indicate that chemical modification rather than rapid erosion may be more important for inactivation. After LTP treatment, samples were characterized by vacuum-transfer to x-ray photoelectron spectroscopy (XPS) to measure the chemical modifications taking place in the LPS layer. With XPS, we measured a decrease in the pulse width of about 1 μs. Plasma was characterized by optical emission spectroscopy (OES). The OES gives information about the presence of the relative number densities of the radicals as OH, O, and O2 and from these data the electron temperature and density were evaluated.

On surface, in high vacuum environment, the Au NPs were characterized by Transmission Electron Microscopy (TEM), Secondary Ion Mass Spectrometry, Transmission Electron Microscopy (SIMS - ToF) and X-rays Photoelectron Spectroscopy (XPS). In solution, as prepared, the Au NPs were characterized by UV-visible spectroscopy and Attenuated Total Reflectance Fourier Transform Infrared (ATR-FTIR) spectroscopy.

The size of the synthesized Au NPs depends on the initial pH of the solution, low-pH solutions having the particle size around 2.5 nm in diameter and in a solution with a higher pH value than 6, the NPs diameter was found to be 2 – 5 nm. The negative SIMS-ToF measurements reveal the presence of the Au, AuCl and AuN⁺ ions in the solution with a pH 3 and AuO⁺ in a solution with pH 11. In a solution before plasma processing, with a low pH value the negative ions AuCl⁺ and AuCl⁻ were detected, which suggest an initial agglomeration of Au atoms in solution. The binding functional groups on the Au NPs are confirmed by the XPS analysis.

The correlation among information of the Au NPs morphology, the binding atoms on Au surface, plasma electron temperature and density and the formation of the Au complexes, during the Au NPs synthesis will be presented and discussed.

References:
3. S. W. Lee et al., in preparation.
intensity of the C=C=C-H peak, which indicates that the aliphatic chains in lipid A were removed. The N/C ratio increases approximately equally in all dissections, which suggests that gas chemistry does have a large impact on amides. Complementary studies with lipid A will be presented as well as results of a UUV optical filter approach used to probe UUV-induced LPS modifications in real time by in-situ ellipsometry while protecting the material against ion bombardment.


4:20pm PS+Bi-MoA8 Development of Plasma Treated Mn Induced Nano-arrayed Structures in Sol-gel Derived TiO2 Matrix for Biosensing Applications, R.R. Pandey, Centre for Cellular and Molecular Biology, India, K.K. Salmi, National Physical Laboratory, India, M. Dhaval, Centre for Cellular and Molecular Biology, India

We describe Mn doped TiO2 nanomaterial-based biosensors modified by high pressure RF plasmas for biosensor applications which activates nanotechnology matrix and greatly enhanced enzyme loading capacity for development of high sensitivity biosensors. As a test model, effects of plasma treatment on a novel potentiometric urea biosensor for selective and quantitative recognition of urea by immobilizing urease onto Ti/urease-imprinted Mn incorporated TiO2 film has been studied and monitoring the potentiometric response caused by the immobilized urease/urea reaction system was carried out. These platforms has been characterized by XRD, XPS, FTIR, SEM, cyclic voltammetry to determine the changes in structure, surface chemistry and electron transfer characteristics of platforms after plasma treatments and have been correlated with improved response of biosensor.

4:40pm PS+Bi-MoA9 Growth Promotion of Bread Yeast using Atmospheric Pressure Dielectric Barrier Discharges, S. Kitazaki, K. Koga, M. Shiratani, Kyushu University, Japan, N. Hayashi, Saga University, Japan

Nonthermal atmospheric pressure plasmas have been employed for biomedical processing applications, because they provide high density radicals at a low gas temperature [1]. Recently, nonthermal atmospheric pressure plasmas as well as low pressure plasmas have been employed for growth promotion of plant cells [2,3]. In this study, we have developed a scalable atmospheric dielectric barrier discharge (DBD) device for biomedical processing in a large area and have applied the device to growth promotion of bread yeast. The device consisted of 20 electrodes of a stainless rod of 1 mm in outer diameter and 60 mm in length covered with a ceramic tube of 2 mm in outer diameter. The electrodes were arranged parallel with each other at a distance of 0.2 mm. The dry yeast was set at 1 mm under the electrodes. The discharge voltage and frequency were 10 kV and 10 kHz, respectively. The plasma treatment was carried out in the air. The treatment duration $t_{\text{w}}$ was 50, 100 and 150 s. After the treatment, yeast was suspended in 0.5 ml yeast extract peptone dextrose (YPD) medium and monitored the absorbance increases from $t=0$ hrs. At $t=10$ hrs, the maximum absorbance for $t_{\text{w}}=150$ s is 6.6 times as high as that for the control. From $t=10$ to 15 hrs, the gradient of the absorbance becomes gradual. After $t=15$ hrs, the absorbance increases exponentially with $t$. The plasma treatment reduces the lag phase of yeast growth and enhances the growth rate. The growth promotion tends to be enhanced with increasing $t_{\text{w}}$ from 50 to 150 s. The growth promotion, therefore, depends on the dose of radicals produced by discharge plasmas.


5:00pm PS+Bi-MoA10 Plasma Deactivation of Pyrogenic Biomolecules: Vacuum Ultraviolet Photon and Radical Beam Effects on Lipid A. T.-Y. Chung*, N. Ning, J.-W. Chu, D.B. Graves, University of California, Berkeley, E. Bartis, J. Seog, G.S. Oehrlein, University of Maryland, College Park

Conventional medical instrument sterilization methods are generally ineffective in completely removing harmful biological residues [1]. Biomolecules such as proteins and other pyrogens from bacterial residues are poorly resistant to heat and temperature and are not easily removed by conventional procedures [2, 3]. For example, the presence of lipopolysaccharide (LPS) in host tissue or blood circulation could lead to a generalized sepsis syndrome including fever, hypotension, and respiratory dysfunction and may lead to multiple organ failure and death [4]. Low temperature plasma is a promising technique for sterilization/deactivation of surgical instruments or medical devices, but its effectiveness against such targets is incompletely understood [5]. In this study using a vacuum beam system, we chose lipid A, the major immune-stimulating region of LPS, as a model biomolecule to study. Lipid A consists of a β-1,6-linked D-glucosamine (GlcN) disaccharide carrying two phosphoryl groups. This structure is attached to multiple acyl chains by ester or amide linkage [6]. After vacuum ultraviolet (UV) photon exposure, loss of CH₂/CH₃, C=O ester, and P-O absorption peaks were observed by ex-situ transmission Fourier transform infrared (FTIR) spectroscopy, but the C=O amide absorption peak was only mildly affected. Monitoring photolysis products from lipid A films by in-situ mass spectrometry, we observed cracking patterns similar to those of alkanes/alkenes with a carbon number ~11-13. This result suggests that UV photons remove phosphate groups and break ester linkages leading to desorption of acyl chains. Endotoxicity of lipid A is known to be primarily determined by the number and length of acyl chains as well as the phosphorylation state and the disaccharide backbone [6]. The present results therefore indicate that plasma-generated UUV reduces the endotoxicity of lipid A, in support of the hypothesis of Rossi et al. [7]. We report the effects of UV and radical (H, O) exposures on endotoxicity based upon chemical structural change in Lipid A. Synergism of various beams is compared with plasma exposures and corresponding molecular dynamic (MD) simulations.

electrolysis and measured hydrogen generation by mass spectrometry. In this talk, we will present our overall methodology and discuss these results in detail.


Plasma Science and Technology Division
Room: 201 - Session PS+SE-MoA

Advanced FEOL / Gate Etching II
Moderator: A. Kadavanchik, Mattson Technology


In this study, the deterioration of line edge roughness during plasma etching process was analyzed to find possible control parameters. According to the aggressive design rule shrinkage of memory devices, the physical width of line patterns has become around 20nm. Controlling line edge and width roughness now becomes one of the biggest challenges in patterning process. Though a number of studies on this matter have done, the wiggling of line patterns is still making the migration slower. It is well known that the line edge and width roughness are mainly caused by the projection of poor photoresist patterns. However, the recent line patterns of around 20nm design rules experience severe wiggling added from the plasma etching process. This study focused on this etch-induced deformation by measuring line edge roughness from the prior status. The final line edge roughness was assumed as the sum of the wiggling projected from that of mask pattern and the deformation during plasma process. The deformation part was explained with the equations of the mechanical beam theory assuming several stress sources originated from the plasma. This newly introduced approach could suggest the direction of process modification for more robust profile against deformation. Moreover, combining with the material properties of commonly used materials, this approach could estimate the extent of lateral deformation so that the ultimate size of line patterns could be expected in the aspect of line edge roughness.

2:20pm PS+SE-MoA2 Dependence of ArF Photore sist Polymer Structure on Line-Edge-Roughness Formation during Plasma Etching Processes. T. Uesugi, A. Wada, Tohoku University, Japan, S. Maeda, K. Katou, A. Yaishida, S. Sakuma, Mitsubishi Rayon, Japan, S. Samukawa, Tohoku University, Japan

ArF excimer laser (193nm) lithography technique is widely used in the fabrication of sub-50nm devices. During plasma etching processes, however, the activated species radiated from plasma, such as ions, radicals, and photons, cause damages to ArF photore sist, resulting in low etching resistance and formation of line-edge roughness (LER). To solve these issues, we investigated the interaction between irradiated species from plasma and polymer structure of ArF photore sist. In our previous study, we found that improvement of stability of lactone group in side chain of ArF photore sist realized decrease in etching rate and reduction of the surface roughness of ArF photore sist.

In this study, to further improve the etching resistance and the surface roughness of ArF photore sist, we proposed a new polymer structure of ArF photore sist. Our newly developed ArF photore sist structure has acrylate group in main chain polymer structure, while usual ArF photore sist has methacrylate group in main chain polymer structure. We prepared silicon wafers coated by ArF photore sistis with methacrylate group and acrylate group and etched them using chlorine plasma. As a result, these two types of photore sistis had almost same etching rates. This result suggests that main chain structure of ArF photore sist does not affect its etching rate. On the other hand, the surface roughness of acrylate type photore sist is drastically reduced in comparison with that of methacrylate type photore sist. It is considered to be due to stronger bonding energy of acrylate group than methacrylate group. From this result, it is concluded that the acrylate type ArF photore sist structure is very effective to suppress the roughness formation in ArF photore sist.

2:40pm PS+SE-MoA3 193nm Photore sist Pre-Treatments Before Plasma Transfer to Improve LWR Transfer and CD Control. E. Pargone, CNRS-LTM, France, L. Azarnouche, ST Microelectronics, France, M. Foucheir, K. Mengueli, O. Joubert, CNRS-LTM, France

Linewidth roughness (LWR) is today one of the main parameters that limits our ability to shrink the transistor gate dimension down to 20nm. Indeed, LWR needs to be controlled down to 2nm to ensure good electrical performance of the future CMOS device, while state of the art patterning techniques only allows 4-3 nm gate LWR at best. The major issue in decreasing the gate LWR comes from the fact that the significant LWR of the resist pattern printed after 193nm lithography (about 6nm measured by CDAFM) is transferred into the gate stack materials during the subsequent plasma etching processes. One way to minimize the final gate LWR is to apply various resist post-treatments obtained right after lithography (before any plasma pattern transfer step).

In the present study, we have used CD-SEM and CD-AFM techniques to investigate the impact of different types of resist pre-treatments (combining plasma exposure (HBr, Ar, H2 plasmas), vacuum ultra violet (VUV) light exposure, and annealing) on the photore sist LWR and profile. Many characterization techniques (FTIR and Raman spectrometries, ellipsometry, chromatography, DMA, TGA) have also been used to characterize the physico-chemical modifications of photore sist films responsible for the resist smoothing. We have also investigated the benefits of those resist pre-treatments on both LWR and CD control after pattern transfer in different stacks of materials.

We will show that all treatments generate resist chemical modifications that lead to a decrease in resist LWR while the etch resistance is not always improved. All treatments have in common the cleavage of the side groups (lactone group for plasma treatment and protecting group for annealing treatment) and a decrease of the glass transition temperature that seems to have a direct impact on the LWR decrease. But some other mechanisms compete according to the treatment used and its duration: main chain scission and crosslinking, leading to some different etch resistance improvement. Consequently even if some specific treatments (VUV light exposure, annealing) can improve the resist LWR before transfer, the subsequent plasma etching steps can degrade it and also induce a loss of CD control. In this talk, we will show that combining plasma exposure and annealing treatment the photore sist LWR could be decreased down to 2.6nm and that this LWR could be transferred into polysilicon gate without LWR and CD degradation.

3:00pm PS+SE-MoA4 Plasma Smoothing of Extreme Ultraviolet Photore sist: LWR Reduction at 30nm Half Pitch. E. Altamirano-Sanchez, A. Pret Vaglio, R. Gronheid, D. Marc, W. Boullart, IMEC, Belgium

Over the past years a tremendous amount of effort has been put on Extreme Ultraviolet lithography (EUVL) for printing the 16nm node. In 2010 imec’s EUV alpha demo tool printed for the first time a 16nm node SRAM cell using state-of-the-art EUV photore sist (PR). In order to meet the line width roughness (LWR) requirements for the 16nm node, we have investigated plasma smoothing techniques on 30nm half pitch lines after exposure.

In this contribution we will report our findings on EUV PR plasma treatment (FT) using H2, Ar and HBr for LWR reduction; and subsequent, in-situ PR encapsulation for preserving the improved LWR during subsequent pattern transfer. Currently we are investigating the vacuum UV role during the H2 PT using MgF2 windows. In parallel, we are characterizing a PR resist encapsulation carried out in-situ (in the etching chamber). The encapsulating layer is characterized through various analytical techniques, such as: XPS, Ellipsometry, mass metrology and TEM. These characterizations will provide understanding of how the H2 plasma improves the LWR and of how the PR encapsulation preserves the PR pattern allowing a straight HM patterning profile.

This study was carried out on 300 nm silicon wafers with the following patterning stack, from top to bottom: 50 nm PR/20 nm under layer (UL)/15 nm SiOC/40 nm amorphous carbon layer (ACL). The dry etching was carried out in an ICP like reactor from Lam research (Kioy C reactor TCPTM).

Power spectral density (PSD) analyses have shown that Ar and HBr plasmas do not improve the LWR of EUV PR as they do on 193i PR. Using the correct conditions in the ICP reactor, H2 plasmas can improve LWR by ~30% without CD bias; in other words no PR reflow was detected.
After the H2 plasma treatment, the pattern was transferred into the UL using a CHF2/CF4/O2 gas mixture. This chemistry provides on the one hand a high PR passivation but on the other hand induces a PR pattern degradation which is translated into higher LWR. A novel alternative for maintaining the improved LWR after H2 PT was to deposit in-situ (in the ICP reactor) a silicon containing layer that encapsulates the PR and preserves the improved LWR when the pattern is transferred into the UL, the SiO/C and the ACL.


As the feature size in CMOS technology continues to shrink, control over line edge roughness (LER) and line width roughness (LWR) is approaching atomic scale for the 14 nm node and beyond. When the line/space patterns are decreased, deformation of the organic underlayer material occurs during plasma processing, which adversely impacts pattern transfer into substrate material to fail. We previously reported that vacuum ultraviolet (VUV) “curing” (modification) of the organic underlayer material by plasma discharges is a promising approach to extend the process window for obtaining high fidelity pattern structures [1]. To understand the details of this approach in more detail, we exposed VUV light at various wavelengths corresponding to the absorption energies of various underlayer materials by synchrotron radiation on “unopened” and “opened” samples comprising stacks inclusive of the underlayer material. We found that exposing underlayers to certain absorption maxima of the patterning material maximized the curing effect. Specifically, for the underlayer material trademarked as NFC, absorption maxima at 155 nm had relatively large effects, reducing LER by as much as 37%. We also found that post-treatments on “unopened” (post lithography) samples comprising varying underlayer materials effectively reduced the pattern deformation, though much less effective than on previously “opened” samples comprising the same. Similarly, chemical and physical effects of the plasma on underlayer material deformation were investigated. To investigate the changes based on chemical modification by plasma, we generally found that the non-selective pattern transfer plasmas are better than highly selective processes to minimize the deformation. In addition, we also noticed that the ion energy of the discharge plays an important role in the deformation and found that the extent of pattern deformation decreased for lower energies. In addition, many commercially available organic underlayer materials were tested with respect to their composition and hardness. We found that the relative hydrogen content of the underlayer material seemed to correlate with the deformation behavior, while little effect was seen for hardness. These initial findings show that a close interlock between patterning materials, lithography and plasma processes has to be executed to minimize effects such as LER for future technology nodes.


4:00pm PS+SE-MoA7 Single Digit Nano Plasma Etching. D.L. Olynick, Lawrence Berkeley National Laboratory

One of our themes at the Molecular Foundry at LBNL is “Single-Digit Nano fabrication” (SDN) which describes our efforts to pattern materials with resolution, precision, and control at the sub-10 nm scale. At this scale, we enable research and applications in areas such as nanoelectronics, nanomagnetics, nanofluidics and plasmonics. For instance, the Molecular Foundry’s work on graphene nanomeshes using SDN shows a band gap opening with sub-bands.1 However, the patterning and plasma pattern transfer for SDN present significant challenges and the question arises, “What are the limits?”

In this talk, I will survey nanoscale etching work from 30-3 nm to demonstrate the challenges and opportunities for plasma nanopatterning. High resolution patterns are made using a variety of materials and techniques including atomic layer deposition, directed self-assembly and electron beam and nanoimprint lithographies. Recent work with cryogenic etching and simulation in the SDN regime will be highlighted.


This work was performed at the Molecular Foundry, Lawrence Berkeley National Laboratory, and was supported in part by the U.S. Department of Energy under Contract No. DE-AC02-05CH11231

4:40pm PS+SE-MoA9 Ar and He Plasma Pretreatments of Organic Masking Materials for Performance Improvements during Plasma Pattern Transfer. D. Mezler, P. Wellenbock, N. Kumar, G.S. Chedin, University of Maryland, S. Engelmann, R.L. Bruce, N.C.M. Fuller, IBM T.J. Watson Research Center

Plasma based pretreatments of organic masking materials have been shown to offer significant potential for reduction of surface, line edge and line width roughness during the subsequent pattern transfer process. Since one of the underlying mechanisms of roughness formation has been reported to be a synergistic effect of energetic ion bombardment, ultraviolet (UV) / vacuum ultraviolet (VUV) plasma radiation and increased temperature, a possible mechanism of organic mask curing may be the elimination of the above synergism by sequential exposures to VUV plasma radiation followed by ion bombardment dominated plasma etching. To examine this question, and establish the impact of pre-treatments on roughness introduction, etch resistance and pattern transfer fidelity during the pattern transfer process, we have studied pretreatments of 193nm PR and other organic masking materials in Ar and He discharges with pronounced UV/VUV emission. The impact of pretreatments and subsequent pattern transfer processes on the organic materials were monitored in real-time by in-situ ellipsometry. Multilayer modeling of these data allows determination of optical material density and film thickness of various layers, i.e. the surface layer densified by ion bombardment, the UV/VUV modified layer in the material bulk, and the rough surface layer. Additionally, post plasma characterization by Fourier transform infrared spectroscopy (FTIR) and atomic force microscopy (AFM) provides information on bulk material modifications and surface roughness improvements, respectively. UV/VUV spectroscopy combined with Langmuir probe characterization provides information on plasma parametric dependencies of photon and ion fluxes, respectively. Pretreatments in a He discharge reduced roughness introduction in a C6F5/Ar pattern transfer process by ~50% (RMS=4nm) while increasing film etch resistance by ~30% leading to only a slight increase in the total material removal when compared to the uncured material (pattern transfer only). Time resolved studies of this process allow detailed investigation of distinct stages and underlying mechanisms of materials modification, such as radiation induced changes in the materials bulk, ion crust formation at the film surface, and surface roughness development. Correlating these observations with post plasma characterization by AFM and FTIR allows a mechanistic understanding of plasma based pretreatments or organic materials and their impact on the subsequent pattern transfer. The dependence of improvements in masking performance on Ar and He plasma based pretreatments and a description of the underlying mechanisms will be presented.

5:00pm PS+SE-MoA10 Sub-32nm Node Mask Patterning for Deep Silicon Trench Ech. J. Yarmush, H. Haga, Y. Chiba, K. Kumar, P. Biolsi, TEL Technology Center America, LLC, J. An, H. Hichri, B. Divahouri, X. Li, N. M. Microsystems, R. Wise, IBM Research

In the last several semiconductor device generations, one of the complexities in fabricating ever smaller feature sizes and increased density, has been the stringent requirements placed on photolithographic processes and mask scheme formation. The use of Immersion Lithography, reduced resist layer thicknesses and planarity requirements have driven the need for complex multilayer-multimaterial stacks that can be utilized for subsequent plasma Etching masking.

One requirement by manufacturers of Sub-32nm DRAM technology, utilizing deep silicon memory cells, is an extremely high aspect ratio mask that enables the anisotropic etch profile of the Silicon Trench. To meet both the needs of the photolithographic processes and the high aspect ratio mask requirements of the Silicon Trench etch, a complex Photo Resist, Silicon Anti-reflective coating, Optical Dispersive Layer, CVD Oxide layer is used. This deposited mask stack also sits on top of a Silicon-on-Insulator layer that must also be etched through plasma process. In this paper, we describe the unique requirements of etching each film stack in order to meet the overall physical requirements of this high aspect ratio mask patterning etch. It also describes the process capabilities of a commercially available Capacitively Coupled Plasma reactor that enables it to meet these advanced complex film stack requirements.

This work was performed by the Research and Development team at TEL Technology Center America in joint development with IBM Semiconductor Research & Development Center.


The production of atomically perfect surfaces by simple solutions is both intrinsically fascinating and technologically important. For over half a century scientists have known that many aqueous bases — so-called...
“anisotropic etchants” — selectively attack all silicon faces except Si\{111\}. As a result, a macroscopic silicon sphere placed into one of these solutions spontaneously transforms into a polyhedron. Twenty years ago, the surface science community was rocked when researchers at Bell Labs showed that, in some cases, the etched surfaces are not just smooth, they are atomically flat and passivated by a single monolayer of H atoms. This type of highly precise but inexpensive chemical machining is used in diverse applications ranging from the production of ink-jet nozzles to the fabrication of ultraminiaturized transistors to the cleaning and polishing of silicon wafers. However, the chemical reactions that govern this behavior remain a source of controversy. We resolve this controversy and give the first quantitative, atomic-scale understanding of anisotropic etching across all silicon surface — not just Si\{111\}.

The reactivity of a wide variety of Si\{100\} surface sites towards a prototypical anisotropic etchant, ammonium fluoride, is quantitatively determined from measurements of the atomic-scale morphology and chemical composition of etched surfaces. These measurements enable predictions of the effects of chemical strain, steric hindrance, and chemical structure to be separately determined. The high selectivity of the etchant is explained by the strain energy released during the chemical reaction; steric hindrance plays an important, but distinct, role. This pattern of reactivity is consistent with previously postulated mechanisms of aqueous silicon etching, which predict insertion reactions across rigid, essentially immobile Si-Si backbones. Instead, we propose that cleavage of the backbond occurs during the formation of a surface silanone which is driven by simultaneous interadsorbate strain release. On Si\{100\} surfaces, this hypothesis quantitatively explains the characteristic alternating-row etch morphology on both flat and vicinal surfaces, the observed site-specific reactivity, the unusual reaction kinetics, and the hydrogen termination of the etched surface, without invoking an unreasonably strained reaction intermediate. This mechanism also explains the atomic-scale reactivity and relative etch rates of the three principal faces of silicon, thereby giving the first atomic-scale understanding of anisotropic silicon etching.

**Surface Science Division**

**Room: 109 - Session SS1-MoA**

**Selectivity and Reactivity of Chemisorbed Species**

**Moderator:** A.J. Gellman, Carnegie Mellon University

**2:00pm SS1-MoA1 Observation and Modeling of Chiral Modifier-Substrate Complexes on Pt(111).** B. Hammer, Aarhus University, Denmark, V. Demers-Carpentier, P.H. McBreen, Université Laval, Quebec, Canada INVITED

We present a combined density functional theory (DFT) and scanning tunneling microscopy (STM) study of the complex formation between the chiral modifier naphthyl ethylene (NEA) and the model substrate trifluorocyclophenone (TFAP) on a Pt\{111\} surface. The two molecules interact via a hydrogen bond between the amine group on the NEA and the carbonyl on the TFAP. Many different realization of the hydrogen bond exist depending on the relative position and orientation of the two molecules. However, only very few are observed experimentally. These are all among the most stable structures found in DFT. The chemisorbed TFAP is prochiral and hydrogenation of the carbonyl would lead to a chiral product. The prochirality of TFAP when forming a complex with NEA is regiospecific, i.e., it depends on the position at which the TFAP is chemisorbed relative to the NEA. The origin of this regiospecificity is discussed.

**2:40pm SS1-MoA3 Enantiospecific Decomposition of Tartaric Acid on Spherically Curved Copper Single Crystals.** B. Holcslaw, P. Kronasticky, A. De Alwis, A. Reinicker, V. Pushkarev, A.J. Gellman, Carnegie Mellon University

The kinetics and mechanisms of many catalytic surface reactions depend on the atomic level structure of surfaces. Detailed study and understanding of the influence of surface structure on a given reaction is experimentally arduous. The space of possible surface orientations spans a two-dimensional continuum. A detailed study of structure sensitive surface chemistry requires preparation and study of many different single crystal substrates, each with a different orientation. A new high-throughput methodology has been developed based on the use of Surface Structure Spread Single Crystals (S4C), spherically curved single crystals that expose a continuous distribution of different crystallographic planes across their surfaces. These S4C surfaces are being studied using surface analysis tools capable of spatially-resolved measurements that can sample the continuous space of surface orientations.

Six copper S4C surfaces have been produced in-house: three low Miller index centered surfaces, (100), (110), and (111); and three high Miller index surfaces, (211), (311), and (411). Each surface has been chosen and spherically curved so that the complete set of surfaces spans the entire stereographic triangle. The surfaces have been characterized using laser profilometry and oriented using x-ray diffraction and low energy electron diffraction. A benefit of the S4C surfaces is that they are naturally chiral, forming regions of either R- or S- chirality. The surface chemistry of a chiral molecule, such as tartaric acid, on a naturally chiral surface can be enantiospecific. Tartaric acid decomposition from copper surfaces shows very high enantioloselectivity due to its highly nonlinear surface explosion kinetics. The nature and magnitude of the enantioloselectivity of tartaric acid decomposition across the stereographic triangle is not currently well-understood. Copper S4C surfaces provide an opportunity to study tartaric acid decomposition on virtually all possible copper surface structures. This research is the first attempt to map surface enantioselectivity for any compound across the entire stereographic triangle in high detail.

**3:00pm SS1-MoA4 Reaction Pathways of Alcohols with Transition Metal Oxides: A Comparison between WO3 and MoO3.** Z.J. Li, Y.K. Kim, R.J. Rousseau, B.D. Kay, Z. Dohnalek, Pacific Northwest National Laboratory

The reactivity of C1-C4 aliphatic alcohols over cyclic (WO3), and (MoO3), (0 ranges from 3 to 6) clusters were studied experimentally and theoretically using temperature-programmed desorption, infrared reflection-absorption spectroscopy, and density functional theory. Three reaction channels, dehydration, dehydrogenation, and condensation, have been identified on (WO3)n clusters while only dehydration and dehydrogenation have been observed on (MoO3)n. The desorption temperature of reaction products decreases with increasing cluster size and the dehydration reaction channel on (MoO3)n is attributed to the lower reactivity of alcohols with (MoO3)n as compared to (WO3)n and consequently a negligible concentration of the Mo(VI) centers coordinated with two alkox groups required for this reaction are formed. DFT calculations provide a detailed explanation for the reactivity and relative selectivity among the reaction channels and (VII) and Mo(VI) metals centers.

This work was supported by the U.S. Department of Energy Office of Basic Energy Sciences, Division of Chemical Sciences, Biosciences and Geosciences, and was performed at EML, a national scientific user sponsored facility of the Department of Energy’s Office of Biological and Environmental Research located at Pacific Northwest National Laboratory (PNNL). PNNL is operated for the U.S. DOE by Battelle Memorial Institute under contract no. DE-AC06-76RLO 1830. Computational resources were provided at EMSL and the National Energy Research Scientific Computing Center at Lawrence Berkeley National Laboratory.

**3:40pm SS1-MoA6 Molecular Dynamics Simulations of Oligomer Film Stabilization through Ion-Bond Deposition.** T. Kemper, University of Florida, D. Lee, Lawrence Livermore National Laboratory, S.R. Phillpot, S.B. Sinnott, University of Florida

Ion-bond deposition is used to stabilize conducting oligomer films for use in organic photovoltaic devices. The goal is to prevent structural changes caused by chemical attack, by charge, or by desorption, that would degrade device performance. The second-generation reactive empirical bond-order (REBO) potential has been successfully applied to the irradiation and modification of crystalline, polymer and nanostructures, such as carbon nanotubes. In this work the atomic-level processes involved in selective modification of oligomers for optoelectronic applications through thermal energy particle deposition are explored to identify the mechanisms by which different polyatomic and radicals assist in the stabilization of oligomer films. In particular, oligomer films of polythiophene are bombarded with incident H, CH3, C2H and thiophene in the hyperthermal regime that involve incident energies of 4-50 eV. The results of classical molecular dynamics simulations with the REBO potential are compared to experimental findings, as well as linearly scaled density-functional theory molecular dynamics results. This work was supported by the NSF (CHE-0809376).

**4:00pm SS1-MoA7 Structure-Reactivity Relationships in the Electron Induced Reactions of Surface Bound Organometallics.** H. Fairbrother, S. Rosenberg, J. Wvuk, Johns Hopkins University, C. Hagen, W. vanDorp, T. Landheer, Delft University of Technology, Netherlands

Electron beam induced deposition (EBID) is a direct-write lithographic technique where volatile or metallic precursors are decomposed by a focused electron beam in a low vacuum environment to create metallic nanostructures. As a tool for fabrication, EBID offers an attractive and unique combination of capabilities including high spatial resolution and the flexibility to deposit free-standing three-dimensional structures without the need for resist layers. However, a major limitation of EBID is that
Weidner, CaF$_2$ as well as RF glow discharge deposited films of allylamine. The SDS positively charged and hydrophilic surfaces. The surfaces studied included spectroscopy and surface plasmon resonance (SPR) sensing to investigate with sodium dodecyl sulfate (SDS) being one of the most widely used Surfactants are important compounds used in many industrial applications, National ESCA and Surface Analysis Center for Biomedical Problems Ge(100)-2×1 is highly sensitive to subsequent exposure to water vapor. The experimental results. We show that phenyl isocyanate pre-adsorbed on calculations are also used to corroborate and help understand these isocyanates to form a diphenyl urea compound, similar to what is expected the molecular environment, fits of the SFG data were used to quantify the orientation and alignment of the SDS layers across the wide range of SDS concentration. Since SFG is sensitive to both orientational order and the amount of material adsorbed we used SPR to determine the SDS concentration from 0.067 to 20 mM. Since the water molecules above the positively charged CaF$_2$ surfaces by the anionic charged head group of SDS to grow laterally as well as across is shown to occur in nanobonding. Wet densities typically found on Si(100) with miscuts < 0.025° and particulates received wafers or post-processing. A surface step density ≥ 500 step/µm across atomic terraces direction) and very low particulate density > 0.1 µm-2 particulates results in 3-dimensional isolated bonding. For this to occur, the surfaces need to exhibit wide flat atomic terraces (width >10 nm), low atomic step density (< 500 steps/µm across atomic terraces direction) and very low particulate density. For this to occur, the surfaces need to exhibit wide flat atomic terraces (width >10 nm), low atomic step density (< 500 steps/µm across atomic terraces direction) and very low particulate density. For this to occur, the surfaces need to exhibit wide flat atomic terraces (width >10 nm), low atomic step density (< 500 step/µm across atomic terraces direction) and very low particulate density (< 1/100 µm2). This contrasts with the typical density of surface steps (~ 500 steps/µm a.a.t.d.) and particulate density ~0.1 - 1 µm$^2$ as received wafers or post-processing. A surface step density ≤ 50 steps/µm a.a.t.d., typically found on Si(100), and a particulate density ≤ 0.025 µm$^2$ and particulates densities ≥ 0.1 µm-2 particulates results in 3-dimensional isolated bonding points of contacts as opposed to more uniform, 2-dimensional interphases that grow laterally as well as across is shown to occur in nanobonding. Wet chemical processing and SEZ spin technology are compared and combined to smmoth substrates via the H-A chemistry [1,2] via Tapping Mode Atomic Force Microscopy, before and after nanobonding. Our results show nanobonding can result in bonding strength larger than 10 MPa/cm$^2$ as measured by mechanical bond pull tests. Wafers fracture within the bulk of both Si and SiC substrates rather than interfacial delamination.

1] US Patent 6,613,677, issued 9/2/03 "Long range ordered semiconductor interface surface and oxides," 6,613,677, Herbots, N; Atluri, V. P.; Bradley J.D.; Swati, Banerjee; Hurst, Q.B.; Xiang, J.
Molecular Ordering and Electrochemical Interfaces
Moderator: S.L. Tait, Indiana University

2:00pm SS2-MoA1 Self-assembled Chains of 4,4'-azopyridine on Cu(100) Stabilized by Metal-Organic Coordination Interactions. H. Lim, S.L. Tait, Indiana University

Understanding self-assembled molecular architectures at surfaces is essential to control and tune low-dimensional nanometer-scale organic structures. 4,4'-azopyridine (APY) has been vapor deposited on the clean Cu(100) surface in ultra-high vacuum and investigated using scanning tunneling microscopy and X-ray photoelectron spectroscopy. APY is chosen as a model building block for metal-organic frameworks, and for its conformational switching by the photoactive azo bridge. At low coverage, APY self-assemblies into highly-ordered one-dimensional chains oriented along the low symmetry directions of the surface, many of which grow from the Cu step edges and some of which form tee junctions with other chains. Careful analysis of high resolution STM images points to pyridyl-Cu interactions being responsible for the high chain stability at room temperature. The related molecule, azobenzene, forms 1D chains on Au(111), but these grow side-to-side by hydrogen bonding at the azo bridge nitrogens and are only observed at cryogenic temperatures [1]. At higher coverages, APY chains align in a parallel fashion to form 2D islands, which increase in size with annealing.


2:20pm SS2-MoA2 Molecular Self-assembly of Terephthalic Acid and Sodium Chloride on the Cu(100) Surface. D. Skomski, S. Abb, S.L. Tait, Indiana University

To expand the catalogue of available interactions for the efficient self-assembly of highly-ordered nanoscale structures, we have investigated the formation of new supramolecular networks of terephthalic acid (TPA) and sodium chloride (NaCl) on the copper (100) surface. The mixture of this organic species with salt is prepared by vapor deposition in an ultra-high vacuum system and represents a model system for ionic self-assembly in two-dimensions. Several structures have been observed by means of scanning tunneling microscopy molecular resolution imaging. Chemical shifts in the sodium 1s photoelectron peak have been observed by X-ray photoelectron spectroscopy upon addition of TPA to the surface, confirming a direct interaction. Our research indicates that TPA and sodium produce new structures due to favorable ionic interactions between sodium cations and the negatively charged carboxylate groups in TPA’s deprotonated form. Interactions with the metal surface and organic molecules appear to be sufficient to break the ionic bonds of the NaCl lattice. The formation of new TPA structures integrates the interplay between adsorbate-substrate and ionic interactions and opens new possibilities for ionic self-assembly at surfaces with highly ordered structure and specific chemical function.

3:00pm SS2-MoA4 Formation of Closed Shell Quantum Dots as a Driving Force for Molecular Ordering. L. Bartels, J. Wyrick, Z. Cheng, D. Sun, D. Kim, University of California, Riverside; T.L. Einstein, University of Maryland

Anthrquinone self-assembles on Cu(111) into a giant honeycomb network with exactly three molecules on each side. Here we propose that the exceptional degree of order achieved in this system can be explained as a consequence of the confinement of substrate electrons in the pores, with the pores being ordered so that the confined electrons can adopt a chiral-like two-dimensional quasi-atom configuration with two filled shells. Formation of identical pores in a related adsorption system (at different overall periodicity due to the different molecule size) corroborates this concept. A combination of photoemission spectroscopy with density functional theory computations (including van der Waals interactions) of adsorbate-substrate interactions allows quantum mechanical modeling of the spectra of the resultant quasi atoms and their energetics.

The resultant pores have about 4 nm in diameter. In this study we explore how the behavior of adsorbates inside them differs from that on extended terraces. CO molecules and adlayers exhibit properties under such nanoscale confinement that markedly depart from those of extended adlayers: a) the confinement stabilizes dislocation lines (anti-phase domain boundaries) in the adlayer that affect roughly 1/3 of the adsorbed molecules; b) confinement prevents the formation of dense islands of adsorbed molecules, depending on coverage either causing dispersion of vacancies in the adlayer or preventing the growth of molecular islands; c) at a coverage of just a few molecules on the facet, we observe that a molecular shell structure is formed, resembling in its underlying mathematics the atomic model. Confined structures are an ideal test bed for measurement of the coverage dependence of molecular diffusion and in this study we find a reduction of the diffusion barrier at a slope of 57%/ML.

4:20pm SS2-MoA8 Influence of Solvent on the Chiral Resolution of Organic Molecules on Au(111): EC-STM Study of Biphenyl Dicarboxylic Acid on Au(111) in an Aqueous Environment. B.L. Kim, J.A. Hanson, M.W. Turner, J.L. Reeder, Boise State University

Adsortion induced chiral resolution of organic molecules is important due to its potential applications in stereo-selective catalysis. We studied the adsorption induced chiral resolution using a model achiral molecule of 4,4’-biphenyl dicarboxylic acid (BPDA) on Au (111) in 0.1 M perchloric acid (HClO4) by scanning tunneling microscopy (STM). The BPDA molecules are known to have chiral resolution on Au(111) in an ultrahigh vacuum (UHV). However, our experimental data show that the molecules form island structures with distinctive preferred orientations at the length scale of the molecular size, whereas they have no orientation order at the length scale bigger than the molecular size. We calculated angle dependent binding energy between the substrate and a BPDA molecule, the intermolecular interactions between the BPDA molecules and their interactions with water molecules. The calculation suggests that the absence of chiral resolution in the aqueous environment may originate from the increase of effective rotation energy barrier of the BPDA molecules due to its hydrogen bonding with the surrounding water molecules. The strength hydrogen binding between BPDA molecules is sufficient to overcome the energy barrier for chiral resolution through rotational motion in UHV, but not in an aqueous environment.
Diffusion of water (H₂S) and hydrogen sulfide (H₂S) on metal surfaces can be characterized using a variety of techniques. We present novel setups for both an improved electrochemical AFM [1] and an advanced electrochemical surface force apparatus [2]. These tools allow for force measurements under electrochemical potential control between a reference electrode and the metal surface of interest. We describe the results of the first surface force measurements on metal surfaces under electrochemical control between a (111) oriented gold film and 5×10⁻⁵ mbar of ethanethiol vapor at room temperature.

Additional degree of freedom, such as rotation and tilting of the adsorbed molecule, can significantly affect the physical and mechanical properties of surfaces and play a critical role in all of these systems. We present a systematic study of the potential-dependent interaction forces and how these forces contribute to deviations from DLVO theory, particularly at distances less than two Debye lengths, or 2-3 times the rms roughness, whichever is greater.

fabrication of Al₂O₃ dielectric thin films using atomic layer deposition (ALD) from trimethylaluminum and both H₂O and D₂O precursors as a function of substrate temperature and precursor exposure. ALD enables precise control of film growth at the atomic scale, while comparison of H₂O vs. D₂O as the oxidation precursor enables isotopic defect loading in the Al₂O₃ and replacement of the –OH defect with the –OD defect. We have developed the D₂O based ALD process, and characterized both D₂O and H₂O based ALD Al₂O₃ thin films by SIMS, XPS and spectroscopic ellipsometry. Our resulting Al₂O₃ films are physically identical with the exception of –OH and –OD defects, making this system an ideal platform for the study of TLS defect related dielectric loss in JQHs. To this end we have fabricated MIM capacitor test structures and superconducting resonators using Re and Al electrodes and performed electrical measurements. Correlation between these data and low temperature superconducting microwave resonator performance will be discussed.

3:00pm  TF-MoA4 Supported Core-Shell Pd/Pt Nanoparticles Synthesized by Atomic Layer Deposition. M.J. Weber, A.J.M. Mackus, Eindhoven University of Technology, Netherlands, M.A. Verheijen, C. van der Marel, Philips Innovation Services, Netherlands, W.M.M. Kessels, Eindhoven University of Technology, Netherlands

Recently, atomic layer deposition (ALD) has become the subject of great interest as a new way to synthesize supported metallic nanocatalysts (1, 2). In comparison to their single-metal counterparts, bimetallic nanoparticles (NPs) can present a higher selectivity and an enhanced catalytic activity. In this contribution, we present an innovative ALD process enabling the synthesis of supported bimetallic core-shell NPs. Although ALD was primarily developed to deposit conformal thin films, metals have the tendency to form nanoclusters on the substrate during the initial cycles of the process. In this work, Platinum-Palladium core-shell NPs were successfully deposited on Al₂O₃ using ALD of Pt and Pd. The selective growth property of noble metal ALD (3) has been exploited to cover Pt nanoclusters with a Pd shell. An ALD process using Me₂CPPMe₂ and O₂ has been used to deposit Pt nanoclusters, and ALD with Pd(Hfacac) and H₂ enabled to selectively cover them with a Pd shell. High Angle Annular Dark Field (HAADF) TEM images confirm the fact that bimetallic core-shell NPs of 3-4 nm were synthesized (on Al₂O₃ covered TEM windows). Information on the chemical state of the Pd and Pt was obtained from XPS analysis. Average values of the Pd shell thickness as obtained by a model analysis of the XPS data were found to be in agreement with the TEM data. This new process is expected to be also applicable to other Platinum group metal cores, for example, with tailoring the Au NPs by changing the ALD process parameters in order to obtain different nanoparticles sizes and compositions is also addressed. It is expected that the fuel cells industry, but also advanced sensors technologies, can benefit considerably from an enhancement of catalytic activity and selectivity of nanocatalysts prepared by ALD.

2: Christensen et al., Small 2009, 5, No. 6, 750-757
3: Mackus et al., J. Appl. Phys. 2010, 107 (11), 116102-1/3


Titanium dioxide (TiO₂) has been regarded as one of the most promising photocatalysts for environmental protection due to its high photocatalytic activity, high chemical stability, low toxicity, and low cost. Anatase and rutile structures of TiO₂ are two main phases with band gap energies of 3.3 and 3.1 eV, respectively. The structures of both anatase and rutile consist of chains of TiO₆ octahedra. However, two crystal structures are obviously different. It is well-known that the difference in crystal structures causes different mass density (3.894 g/cm³ for anatase and 4.250 g/cm³ for rutile) and electronic band gap. The anatase phase has a more negative conduction band edge than that of the rutile phase. It results in superior photocatalytic activity of anatase phase than that of the rutile phase. Therefore, the structure of TiO₂ films and high surface area are heavily important factors in photocatalytic efficiency.

In this study, TiO₂ film was deposited by remote plasma atomic layer deposition (RPALD) and its phase formation temperature, impurity concentration, and chemical states of the were characterized by various analyses such as Auger electron spectroscopy (AES), X-ray diffractionmetry (XRD), X-ray photoelectron spectroscopy (XPS), and transmission electron microscopy (TEM). The TiO₂ film was deposited on the flat Si substrate and vertically oriented Si nanowire by using remote plasma atomic layer deposition (RPALD) method. RPALD method has advantage to deposit thin film with uniform thickness on the 3-dimensional structure like vertically oriented Si nanowire. In addition, the RPALD method is possible to reduce damage by high energetic ion and to enhance the chemical reactivity between metal-organic precursor and reactant gas for deposition of high quality film. The vertically oriented Si nanowire was fabricated by Ar-anneciling of Si substrate with Al₂O₃ nanocrystal at 1000 °C. We have compared the PL properties of these TiO₂ films deposited with these two different specific surface areas. We have found that TiO₂ film deposited vertically aligned Si nanowire has higher PL intensity than that of TiO₂ deposited flat Si substrate due to large area density of vertically oriented Si nanowire.

4:00pm  TF-MoA7 Fast Atomic Layer Deposition for High Throughput and Low Temperature Applications. P. Poedt, A. Hilbert, M. Smeets, R. Knaapen, TNO, Netherlands, F. Roozeboom, M. TNO & Eindhoven University of Technology, Netherlands, A. van Asten, TNO, Netherlands

Atomic Layer Deposition is a deposition technique capable of producing ultrathin conformal films with control of the thickness and composition of the films at the atomic level. The major drawback of ALD is its low deposition rate (~1 nm/min). Recently, fast ALD concepts were developed based on the spatial separation of the half-reactions, instead of temporal, combined with gas-bearing technology. With this technique, deposition rates for Al₂O₃ of more than 1 nm/s have been reported. This has led to the development of the high-throughput, industrial scale ALD tools for surface passivation of crystalline silicon solar cells.

A new field of applications for fast ALD are flexible electronics, including system-in-foil, flexible displays, OLEDs and solar cells. Flexible electronics are slowly but surely evolving from lab-scale to industrial production. This opens up new possibilities for fast ALD as high-throughput production tool for functional layers such as transparent oxide (semi)conductors (e.g. ZnO) and moisture barriers (e.g. Al₂O₃). One important prerequisite is that these applications require low temperature processes (below 100°C), as they are often temperature sensitive. We present two approaches for low temperature Fast ALD: low temperature fast ALD and atmospheric plasma enhanced fast ALD.

A low temperature thermal fast ALD process for alumina from tri-methyl aluminum and water has been developed. It was observed that the kinetics of the water half-reaction is significantly different at high temperatures (>200°C). Multilayer adsorption of water molecules at low temperatures seems to hinder the self-limiting nature of the ALD process at temperatures below 75°C. Nevertheless, if very low temperatures are not required, low temperature thermal fast ALD is a very suitable technique. Potentially lower deposition temperatures could be achieved by atmospheric plasma enhanced fast ALD. We have integrated an atmospheric plasma source in our reactor in which an He/O₂ plasma is created to act as the oxidant half reaction. However, the chemistry of atmospheric plasmas is different than that of conventionally used low pressure plasmas and has a lower reactivity that can limit the throughput.

Another important aspect of fast ALD for flexible electronics is the processing of flexible substrates, either sheet-to-sheet or roll-to-roll. Approaches for roll-to-roll fast ALD, their challenges and possible solutions will be briefly discussed.

4 J. van den Brand et al., Microelec. Rel. 48 (2008) 1123
We have obtained thin film thickness uniformity of ± 3 % with the cycle time of 2.0 s. With 60 x 120 cm² substrate size these numbers will be doubled to 24–48 wafers/hour. These values fulfill the throughput requirements of CIGS production lines (20-60 modules/h). Even higher throughputs can be obtained using batch type tools for CIGS buffer layers. TFS NX 300 is a fully automated cassette to cassette ALD manufacturing system for Al₂O₃ surface passivation. It consists of 4 processing tubes each with the capacity for a 500 wafer batch. For this application we have modified thermal ALD-process for Al₂O₃ which has a higher growth rate than the conventional thermal (TMAl₂H₂O) ALD process and the film quality is equal to plasma assisted ALD film quality giving throughput of >3000 wafers/h.

The Beneq TFS 600 is a vacuum-line integrated ALD system for OLED moisture barrier coating. Reaction chamber is designed for batch processing up to 35 substrates measuring 500 mm x 400 mm. For this application we have been developing low-cost multilayer barrier structure giving WVTR <10⁻⁶ g/m²/d.

4-40pm TF-MoA9 Conductive Coatings on Nonwoven Fiber Mats by Atomic Layer Deposition. W. Sweet, J.S. Jar, G.N. Parsons, North Carolina State University

Conductive fiber mats offer unique possibilities in the development of many sensing and protective electronic systems. In this work, atomic layer deposition (ALD) is applied to produce conformal coatings of conductive ZnO on complex fiber systems such as nonwoven polypropylene and nylon. The conductivity of these materials were evaluated using a modified 4-probe method specific for fabric structures and correlated with the overall mass gain of the fabric samples after ALD processing. This analysis provided a detailed evolution of the conductive films on the fabric mats. For example, nonwoven polypropylene coated with ZnO showed a significant initial mass increase, eventually becoming linear with increasing ALD cycles. The corresponding conductivity of these films remained low (< 3 S cm⁻¹) after 500 ALD ZnO cycles. Inserting a thin insulating Al₂O₃ layer prior to the ZnO growth, linear ZnO mass gain was achieved after low ALD cycles. Transmission electron microscopy shows that the Al₂O₃ barrier reduces the penetration of the ZnO vapor phase reactants into the polypropylene. As a result, a higher effective conductivity (> 25 S cm⁻¹) was achieved after 200 ALD ZnO cycles that corresponded well with measurements from simultaneous ZnO growth on a planar silicon oxide surface. For comparison, ZnO deposition on nonwoven nylon-6 exhibits uniform growth without Al₂O₃ pretreatment and the conductivity again changes significantly with the Al₂O₃ pretreatment. Implications of the deposition temperature, film thickness, and the use of aluminum doping, on the chemical, mechanical, and electrical properties of ZnO films deposited on polypropylene and nylon nonwovens will be discussed.

5:00pm TF-MoA10 ALD IrOx Thin Film to Improve Microelectrode Array Performance in Stem Cell Applications. T. Ryynänen, J. Lekkala, Tampere University of Technology, Finland, L. Yli-Outinen, S. Narkilahti, University of Tampere, Finland, J. Häimaläinen, M. Leskelä, University of Helsinki, Finland

We promote atomic layer deposition (ALD) and especially ALD deposited iridium oxide (IrOx) thin film [1] as a novel alternative to improve electrical characteristics of microelectrode arrays (MEAs) used in stem cell and other tissue engineering applications. The additional microelectrode coating on microelectrode base material is traditionally used to decrease impedance and noise levels, and to increase charge transfer capacity of microelectrodes in MEAs. However, most of the common microelectrode coatings suffer from certain drawbacks. For example 1) electrochemical activation used to decrease the impedance of sputtered IrOx by one decade, which is considered to be a standard and widely accepted technique, which would make ALD IrOx thin film a strong competitor for the other commonly used microelectrode coatings. In addition to offering an economical, easily controllable, and highly reproducible fabrication process for a thin film improving the electrical characteristics of MEAs, ALD technology may in the future offer also other interesting coating solutions for cell culturing platforms, for example controlling cell growth via functionalized coatings.


Vacuum Technology Division
Room: 111 - Session VT-MoA

5:00pm VT-MoA1 Low Uncertainty Measurements of Trace Water Vapor Based on Cavity Ring-Down Spectroscopy. T. Hodges, National Institute of Standards and Technology INVITED

I will discuss how cavity ring-down spectroscopy (CRDS) can be applied to accurately measure the concentration of residual water vapor which is present in a vacuum system or process gas stream. In CRDS, a monochromatic laser beam is injected into an evacuated or sample-gas-containing optical resonator and the transient decay of light exiting the cavity is monitored to quantify the optical losses. For water detection, the laser wavelength is tuned to probe characteristic rotation-vibration absorption features of the water molecule. The sample absorption coefficient is determined from observations of the ring-down cavity decay time and laser frequency, both of which can be precisely measured. Also, because CRDS uses a resonant optical cavity, extremely long effective optical pathlengths (up to tens of km) can readily be achieved in the laboratory. These properties make CRDS a high-spectral resolution, species-selective method, with relatively small combined uncertainty, and high sensitivity. I will show that when CRDS measurements are combined with first-principles spectroscopic models, this technique can yield concentration measurements with sub-percent-level relative uncertainty for absolute concentrations as low as 10⁻⁰⁰ cm⁻³. I will present examples of CRDS-based trace water detection in vacuum and ultra-high purity gas systems, and I will talk about recent CRDS measurements of the vapor pressure of ice over the temperature range 0 deg C to -100 deg C.

2:40pm VT-MoA3 Comparison of Cavity Ring-Down Spectroscopy, Oscillating Quartz Crystal and Electrical Impedance Technologies for Trace Water Vapor Detection below 100 ppb. M.W. Raynor, J. Feng, Matheson

Control of trace water vapor in high purity process gases at low ppbv levels is critical to the performance of many micro-electronic and photonic devices [1]. Consequently a variety of measurement technologies, have been developed to detect water below 100 ppb. However, the performance characteristics of each technology can vary and this is not always well understood by users. In this presentation three different approaches are considered: Oscillating quartz crystal microbalance (QCM), Al₂O₃ based electrical impedance sensor and laser induced cavity ring-down spectroscopy (CRDS). QCM technology, developed in the early 1960’s, is still widely applied today. It is based on adsorption of water vapor on the hygroscopic coating of the QCM, which causes an increase in the mass of the crystal, and in turn, increases its oscillation frequency. CRDS is a laser absorption technique based on the light decay in a high finesse optical cavity. The high resolution laser ~1-2 MHz (~10⁻⁰⁰ cm⁻³), high reflectivity mirrors (~ 0.99998) results in a long effective path-length which enables high selectivity and sensitivity for H₂O detection. Impedance-based sensors for trace water vapor detection have typically suffered from drift and equilibration issues. However, recently an impedance-based Al₂O₃ sensor chip with integrated heater for cycling the temperature within 60°C to 200°C has been developed. Water vapor is measured dynamically as impedance changes during wet-up of the sensor resulting in rapid response. In this work, we present and discuss data showing the performance of the above parameters from the default 3000 cycles of Ir(acac)₃ and ozone, the impedance level of the ALD IrOx coated microelectrodes can be decreased even further than 10⁻⁰⁰ cm⁻³. This improved performance could be reached, which would make ALD IrOx thin film a strong competitor for the other commonly used microelectrode coatings. In addition to offering an economical, easily controllable, and highly reproducible fabrication process for a thin film improving the electrical characteristics of MEAs, CRDS technology may in the future offer also other interesting coating solutions for cell culturing platforms, for example controlling cell growth via functionalized coatings.

Hydrogen getters are used in many industries including aerospace, defense, and electronics (e.g., MEMS packaging) to control levels of hydrogen in sealed atmospheres and vacuum systems. In this research, we explore not only the getter's ability to operate the sensor remotely, but also its capability to provide fast and sensitive data at an ultrahigh vacuum level (UHV). The ability to operate the sensor remotely, without the need for connecting the electronics unit away from the gauge head, has also made ART MS sensors more suitable for higher parts-per-billion measurement, this technology is now increasingly sought for higher parts-per-million applications as well. Taking advantage of its exceptional dynamic range, with a variety of flexible configurations, CRDS-based instruments address these various applications with a self-verifying measurement solution that is fast and sensitive, yet extremely robust and simple to operate.

With over 1000 measurement points worldwide, CRDS has gained widespread acceptance and growing use in a series of challenging, real-world industrial applications, well beyond trace moisture in simple matrices. We will demonstrate the strong capability of CRDS for a diverse group of analytes over a large dynamic range and under widely varying application conditions. In addition, starting with ultrahigh-purity, sub or low parts-per-billion measurement, this technology is now in practice at a number of organizations where it is making a significant impact. Maximum advantage of its exceptional dynamic range, with a variety of flexible configurations, CRDS-based instruments address these various applications with a self-verifying measurement solution that is fast and sensitive, yet extremely robust and simple to operate.

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The use of hydrogen getters is widespread, and in many industries, including aerospace, defense, and electronics (e.g., MEMS packaging), they are used to control levels of hydrogen in sealed atmospheres and vacuum systems. In this research, we explore not only the getter's ability to operate the sensor remotely, but also its capability to provide fast and sensitive data at an ultrahigh vacuum level (UHV). The ability to operate the sensor remotely, without the need for connecting the electronics unit away from the gauge head, has also made ART MS sensors more suitable for higher parts-per-billion measurement, this technology is now increasingly sought for higher parts-per-million applications as well. Taking advantage of its exceptional dynamic range, with a variety of flexible configurations, CRDS-based instruments address these various applications with a self-verifying measurement solution that is fast and sensitive, yet extremely robust and simple to operate.
Numerical analysis of performances has to be founded on a careful modeling of the local gas flow features, such as pump leakage and development of the rarefied gas flow along the curved channels. Here, gas flows are in general considered three-dimensional, because of the spiral groove curvature, and driven by pressure gradients and the applied rotation speed as well as inertial forces (centripetal and Coriolis effects), which play the most important role.

Following the assumptions made for a Holweck model by Sharipov et al., we propose a lower-order model for steady flows in spiral molecular drag stages, based on the solution of the Boltzmann Equation (BE) with a BGK closure, in general curvilinear coordinates (properly fitted to the geometrical design of the channel), where the inertial effects explicitly appear in the governing equation. The order of the 3D original problem is reduced in the physical space (2D), by introducing assumption of “locally” known flow development of the distribution function along the spiral channel. Thus, 2D-BE calculations of the flow rates and stresses will be performed in a finite number of sections, suitably positioned along the spiral channel, from the outlet up to the inlet, in order to recover the pressure and torque distribution. In particular, the 2D Boltzmann equation is linearized in the most significant parameters (local rotation speed and pressure gradients along the pump radial direction), and solved in the reference section. The local values of pressure and torque are obtained consistently by enforcing the mass flow conservation constraint.

A Discrete Velocity Method (DVM) is used to solve the Boltzmann Equation, with an explicit pseudo-time dependent technique to relax the flow up to its stationary solution. In order to decrease the computational time employed, the solver is designed to work on parallel architectures (MPI).

The performance prediction of the model will be assessed using test cases from the literature and compared to the available experimental data, on both Holweck and Siegbahn geometries. A further verification test will be carried out, to test prediction capabilities in the continuum regime by direct comparison with results obtained by a Navier-Stokes solver, with slip-boundary conditions.
Recent Applications of GCIB Depth Profiling with XPS and TOF-SIMS. Takuya Miyayama, N. Sanada, ULVAC-PHI Inc., Japan. J.S. Hammond, Physical Electronics

The development of new electronic devices incorporating organic materials, such as Organic Light Emitting Diodes (OLEDS) and Organic Photovoltaics (OPVs) is rapidly increasing. To control quality, performance and lifetimes of these devices, it is necessary to characterize the layer structures and the dopant distributions in the thin organic materials. Conventional surface analysis techniques such as XPS and TOF-SIMS, combined with mono-atomic ion beam sputtering, have been widely used for chemical profiling of inorganic films. However, this approach has not been successful for the depth profiling of organic materials due to the loss of chemical information during the sputtering process. Recent cluster ion beam developments utilizing C60 and Coronen ions have also had limited success for the XPS and TOF-SIMS depth profiling of OLEDS and OPVs due to similar modification of chemical and molecular information as a function of sputter depth.

The chemical depth profiling of organic layers with thicknesses greater than one micron has been problematic utilizing XPS and TOF-SIMS with C60 cluster sources. The implementation of new cluster ion sources that could extend chemical depth profiling of organics to more than several microns should also expand the applications of the XPS and TOF-SIMS techniques.

Recent studies have shown the successful use of a GCIB (gas cluster ion beam) source on XPS (X-ray photoelectron spectroscopy) instruments to quantify the chemical depth profile of polyimide films without sputter cooling. [1] Based on these earlier experiments, additional GCIB depth profiling applications of organic and biomaterials with XPS and TOF-SIMS will be discussed. Examples will be presented for the characterization of ultra-thin organic electronic layers in OLED’s and OPVs. The successful organic depth profiling to depths of several tens of microns will also be discussed.


9:00am AS-TuM4 Advances in Organic Depth Profiling for Polymer Devices. J.L.S. Lee, I.S. Gilmore, National Physical Laboratory, UK, A. Ligiardello, University of Catania, Italy

Knowledge of the distribution of organics within an organic matrix is important to the innovation and manufacture of many advanced technologies including polymer electronics and photovolataics, medical devices, ink-jet printing technologies and drug delivery systems. Organic depth profiling using sputtering with cluster ions and imaging by SIMS or XPS have revolutionised the analytical capability for these important systems, providing uniquely detailed 3D chemical information. However, despite recent progress, organic depth profiling is not yet routinely applied to industry problems. The principal reason for this is that organic sputtering only works for a limited set of materials [1] and unfortunately it fails completely for many industrially important materials, such as conjugated polymers in the organics electronics industry. Consequently, an important recent development is the use of nitric oxide (NO) sputtering [2] as a radical scavenger to reduce ion induced cross-linking during depth profiling.

In this study, we use model polymer layers, including polystyrene and industrially relevant conjugated polymers, to demonstrate the benefits of using NO sputtering for polymers that do not sputter under normal conditions and evaluate the basic metrology. Understanding the mechanisms for damage, cross-linking, radical gas production and detection is vital in developing this technique to work with industrial materials. Using C60 as a sputtering ion beam, it is found that NO sputtering combined with sample cooling [3] significantly reduce the disappearance cross-section for characteristic fragments. For polystyrene model systems, the steady state intensity of C6H5 can be increased from typically < 0.1% of the initial intensity at normal experimental conditions to 50% when NO flooding is used in conjunction with cooling. We also demonstrate successful depth profiles on 1 µm thick polymer material, showing high and constant sputtering yields with ~ 30 nm depth resolution to the interface. Our results show a dramatic improvement for the depth profiling of difficult type II polymers. NO flooding may be used along with other developments e.g. large cluster ions [4] for sputtering and sample rotation [3], and has the potential to provide a step change in analytical capability for industrial samples.


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due to the fact that each image slice of the surface is displayed as a 2-D image taken from a 3-D surface. This results in a type of inverted topography of the 3D structure within the ToF-SIMS image volume. To correct for this, the National ESCA and Surface Analysis Center for Biomedical Problems QSEC/KYOTO, Japan has developed a Matlab (Mathworks, Natick MA) toolbox to correct the z-axis of cell depth profiles and display the data properly.

Since the shape and topography of a cell can be complex, we have created a set of features of known chemistry and controlled geometry in order to test and validate the zcorrectorgui is accurately correcting the z-axis. For this, a well-defined model system was transferred to a 3D analysis of an OLED display device using knowledge about the optimum analysis conditions gained on the model system. The resulting depth profile data was then imported into the zcorrectorgui and processed. In this presentation we will highlight the results from this study and show example data obtained from a real cell depth profile processed with the zcorrectorgui.


Organic multilayer systems are of increasing importance in many technological fields. The entry of the OLED technology into commercially available multi-color displays is one example. Optimization of this technology in terms of lifetime and efficiency requires a detailed knowledge of the layer and interface composition.

From the analytical point of view, these OLED multilayer structures are quite challenging. A detailed analysis requires the identification of complex high mass organic molecules in thin layers of only several nanometer thickness with a lateral resolution in the micrometer range. In particular the identification of the molecular composition is challenging for SIMS as molecular information does usually not survive high dose sputtering conditions.

In the past different projectiles such as low energy Cs+ [1] and keV C60 cluster [2-4] have been explored for their depth profiling capabilities on organic layers. For some materials molecular information survives but for the majority of organic molecules depth profiling using these projectiles fails. Recently the application of large Ar clusters for the non-destructive removal of organic material has been discussed in the SIMS community. The GCIB (gas cluster ion beam) technique was developed by the group of Isao Yamada [5] at the University of Kyoto and has already demonstrated some potential in this field of application [6].

We used a TOF.SIMS 5 instrument equipped with a Bi cluster ion gun for the analysis and an Ar GCIB as well as an C60 cluster ion gun providing both analysis and sputter gun capabilities. In this contribution we will focus on the variation of the sputter projectile, the energy dependence of the resulting depth resolution and the survival of specific molecular ion signal under high dose sputtering conditions. For this purpose a well-defined multilayer model sample similar to a working OLED device was used. The knowledge about the optimum analysis conditions gained on the model system was transferred to a 3D analysis of an OLED display device using an Ar GCIB for sputtering.


The stratum corneum (SC) is the outermost layer of epidermis that acts as a barrier to keep foreign objects out of the body and to keep water in. It is composed of multi-layered sheets of dead cells containing keratin that continuously fall off and are regenerated from live cells in the underlying layers. The total thickness of the SC layer is less that 40 μm. In order to study the barrier properties of the SC and how this function fails in certain skin disorders, it is necessary to visualize the distribution of different biomolecules within the multi-layered SC structure. It is also valuable to compare the penetration of various foreign chemicals into the SC layer in mice with and without the skin disorder. In this initial study, we applied TOF-SIMS imaging using a bismuth cluster ion beam to characterize the multi-layer structure of mouse skin. Samples were prepared by quick freezing of mouse tails followed by cross-sectioning by cryostat. TOF-SIMS imaging provided the molecular specificity to clearly visualize dead cell layers and living layers of the epidermis. By using peaks characteristic of specific molecules, it was possible to image the distribution of amino acids, cholesterol, and lipids within the SC. The results suggest that SC might contain several chemically distinct layers. Skin samples were also depth profiled using GCIB sputtering. Imaging of cross-sections and depth profiling from the skin surface will be compared for obtaining molecular profiles within the SC structure.

Biofabrication and Novel Devices Focus Topic
Room: 105 - Session BN+NM-TuM

Biofabrication Applications
Moderator: G.F. Payne, University of Maryland, College Park


Stimuli-responsive polysaccharides, such as chitosan and alginate, are useful biomaterials that can be induced to undergo a reversible sol-gel transition to generate biologically-relevant scaffolds. The recent discovery that their gelation can be triggered by imposing an electrical signal opens many avenues for the creation of biologically functional hybrid structures and the localization onto and within microfabricated devices for biofabrication and biosensing applications. Here we report two different mechanisms for creating polysaccharide hydrogels in microfluidics by electrical signal. The cathodic electrodeposition of the cationic chitosan hydrogel was achieved by electrochemically generated OH- ions at the cathode surface, creating a localized pH gradient at the sol-gel interface. The anodic electrodeposition of calcium alginate hydrogel was achieved by electrical-signal-mediated release of Ca2+ ions as a result of electrochemically generated H+ ions at the anode surface reacting with suspended CaCO3 particles in alginate solution. Localization of the
As microbial communities occupy a confined space over time, concentrations of extracellular signaling molecules accumulate, providing stimuli for unique and varied cellular responses as well as protection from competing microbial communities. Referred to as “quorum sensing” for it’s often reported and coincident dependence on high population density, extracellular signaling provides a new basis for control over molecular and cellular processes as well as population behavior, perhaps in a manner more consistent with that of native machinery. Among behaviors guided by QS are the establishment and persistence of bacterial infections.

Our laboratory has uncovered many of the molecular features of the QS autolinker-2 (AI-2) system using traditional methods that probe bacterial physiology and by exploiting newer principles of biofabrication. That is, we employed electrodeposition methods to assemble complex biological subsystems onto specific sites on microfabricated devices and within microfluidic channels via electrical signals. We have also used genetic engineering techniques to create a signal activated fusion tags that covalently link proteins to the device/bio interface. We have designed and synthesized "biological nanofactories" that provide small signal molecule processing at the surface of targeted and captured cells - enabling programmable control of cell function.

Using these methods, we have discovered attributes of the natural switching mechanism that can be exploited for developing next generation antimicrobials. That is, we decomposed elements of the QS “switch” via mutation and a mathematical model of the regulatory elements and coupled this understanding with devices designed to appropriately interrogate these molecular features. Finally, we have developed alkyl analogs of AI-2 that elucidate structural detail and have potential for affecting behavior in diversity and modes of action. As such, quorum sensing represents a “guide” for learning how signals can be translated into altered phenotype.

Overall, we believe that in vitro microfabricated devices with cell cultures can be an in vitro system that combines microfabrication and cell cultures and is guided by a computer model of the body. We called this in vitro system a micro cell culture analog (microCCA) or a “Body-on-a-Chip”. A microCCA device contains mammalian cells cultured in interconnected micro-chambers to represent key body organs linked through the circulatory system and is a physical representation of a physiologically based pharmacokinetic model. MicroCCAs can reveal toxic effects that result from interactions between organisms, as well as provide specific, inexpensive, accurate, rapid throughput toxicological studies that do not require animals. The advantages of operating on a microscale include the ability to mimic physiological relationships more accurately as the natural length scale is order of 10 to 100 microns.

We have used a microCCA to test potential combination therapies (Tegafur and uracil) for colon cancer. Tegafur is a produg for 5-FU and uracil an inhibitor of DPD, an enzyme which deactivates 5-FU. Simple microwell plate can be a probe for this system, but the microCCA predicts the types of responses observed experimentally. A "pumpling" system that would be easy to utilize has been demonstrated with Tegafur also. We have coupled these body modules with a micro model of the GI tract to examine the response to oral exposure of drugs, chemicals, or nanoparticles. Overall, we believe that in vitro, microfabricated devices with cell cultures provide a viable alternative to animal models to predict toxicity and efficacy in response to pharmaceuticals.

References


Transformation, the process by which a bacterium takes up and incorporates extracellular DNA, is one of the primary enabling technologies in the biotechnology field. This allows a researcher to program bacteria, equipping them with a complement of genes to accomplish a task, such as producing a molecule of interest or acting as a sensor. We describe the simultaneous transformation and localization of Escherichia coli bacteria in response to an electric signal within a microfluidic device. We demonstrate that these transformed bacteria can act as fluorescent sensors of isopropyl β-D-1-thiogalactopyranoside (IPTG), a chemical stimulus, or low dissolved oxygen levels, an environmental stimulus.

This method focuses on bacterial transformation with the added benefit of simultaneous entrapment within an alginate hydrogel at a desired electrode address. This offers the ability to create microfluidic cell-based sensors in a single, simple step. To transform and deposit bacteria, the device was filled with a mixture of electrocompetent cells, 200ng plasmid, 0.5% alginate, and 0.125% CaCO3 and subjected to a 30V/cm DC electric field for 3 minutes on ice. The cells were allowed to recover at 37°C for an hour, cultured for 16 hours, and induced with a chemical signal, IPTG, for 4 hours. This resulted in increased expression of DsRed, a red fluorescent protein. Dissolved oxygen is an important parameter for many cell culture experiments. To create a dissolved oxygen sensor, E. coli were transformed with a plasmid that causes production of green fluorescent protein (GFP) in response to decreased dissolved oxygen concentration in the surrounding medium. Following the above transformation and culturing method, the cells were induced with media that had been deoxygenated in a vacuum chamber, resulting in an increase in GFP expression.

This method is versatile in terms of creating microfluidic cell-based sensors. We envision many exciting applications of this work, including the development of dynamically reconfigurable microfluidic biosensors and high-throughput screening methods for plasmid libraries generated by protein engineering and directed evolution experiments.

Electronic Materials and Processing Division Room: 210 - Session EM+TF-TuM
High-k Dielectrics for MOSFETs Part 1
Moderator: R.M. Wallace, University at Albany, State of New York, USA
8:20am EM+TF-TuM2 In Situ TMA Pre-Treatment Study of GaAs and \(\text{In}_{x}\text{Ga}_{1-x}\text{As}\) Surfaces. B. Brennan, D.M. Zhermak, H. Dong, R.V. Galatage, J. Kim, E.M. Vogel, R.M. Wallace, University at Albany, State of New York, USA
One of the major issues preventing the integration of high mobility III-V semiconductors into next generation CMOS devices is the formation of high levels of interfacial defects at the high-k/BiF interface. These can have the effect of pinning the Fermi level and preventing optimal operation of the devices. Engineering the interface between these materials therefore becomes of critical importance to try and reduce the defect density. Identification of the individual defects however is not a trivial matter with correlation between electrical data and physical measurement techniques rarely seen. This study aims to investigate the effect of in-situ chemical treatments prior to Al2O3 deposition on (NH4)2S treated GaAs and InGaAs surfaces, in terms of both physical characterization by X-ray photoelectron spectroscopy (XPS) and electrical measurements from MOS capacitors.

The reduction of interfacial oxides through a “clean up” effect by a ligand exchange mechanism with the tri-methyl aluminum (TMA) precursor for atomic layer deposition (ALD) of Al2O3 is well known, however little work has been carried out to optimize this process and determine whether variations in the effect are seen as a result of changes in the number of TMA cycles or pulse time prior to oxide deposition. Variations in the presence of arsenic surface features, (i.e. As-As bonding or surface dimers) come under particular focus. The effect of post deposition annealing is also investigated specifically in terms of the potential role hydrogen could play in passivating defects at the interface. [3]

9:00am EM+TF-TuM4 Remote Phonon and Surface Roughness Limited Universal Electron Mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\)/GaSb Surfaces. E.M. Vogel, A.M. Sonnet, R.V. Galatage, University at Albany, State of New York, USA
The results show that room temperature mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\) on GaSb(001) surface after sulfur passivation procedures and HCl etching designed to removed the native oxides. Monochromatic XPS is used to examine the surfaces following the various surface treatments and then without breaking vacuum, after each individual ALD pulse of tetrakis(dimethyl-amino-hafnium (TDMA-Hf) and deionized water (DIW) precursors (i.e. single TDMA-Hf pulse/XPS scan; single DIW/XPS scan; etc.) for two full cycles and finally after 1 nm of HfO2 deposition to determine whether there is any “clean up” effect of the native oxides due to the ALD process. The various surface preparation techniques are compared to determine which is more effective at minimizing native oxides. The behavior of the sulfides and the effect of HCl surface cleaning procedure upon H2O deposition are discussed as well as a comparison to previous results from half cycle Al2O3 deposition on GaSb.[2]. This work is supported by the Semiconductor Research Corporation (SRC) MRF Focus Center, the Nanoelectronics Research Initiative and the National Institute of Standards and Technology through the Midwest Institute for Nanoelectronics Discovery (MIND), and the NSF (ECCS-0925844).


9:40am EM+TF-TuM6 Structural Characterization of Ultra-thin High-k Gate Oxide Films through a Multi-technique Approach. E.J. Bersch, J.D. Latoe, L.B. Wells, University at Albany, State of New York, USA
The results show that room temperature mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\) surface channel MOSFETs with HfO2 gate dielectric with several substrate impurity concentrations (~1×1016 cm-3 to ~1×1018 cm-3) and various surface preparations (HF surface clean, (NH4)2S surface clean and PECVD a-Si interlayer with a HfO2 gate dielectric) have been studied. The peak electron mobility is observed to be strongly dependent on the surface preparation, but the high field mobility is observed to be almost independent of the surface preparation. A detailed analysis of the effective mobility as a function of electric field, substrate doping, and temperature was used to determine the various mobility components (surface roughness, phonon, and Coulombic scattering limited mobility components). For the substrates with high doping concentration, the electron mobility at low vertical electric field is dominated by Coulomb scattering from the substrate dopants, whereas, for lower substrate doping the Coulombic scattering is dominated by the disorder induced gap states. Low temperature measurements were used to determine the surface roughness scattering and phonon components. The results show that room temperature mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\) surface channel MOSFETs with HfO2 gate dielectric at high electric field is limited primarily by remote phonons whereas the Al2O3 gate dielectric is limited by surface roughness scattering.

9:40am EM+TF-TuM6 Structural Characterization of Ultra-thin High-k Gate Oxide Films through a Multi-technique Approach. E.J. Bersch, J.D. Latoe, L.B. Wells, University at Albany, State of New York, USA
The results show that room temperature mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\) surface channel MOSFETs with HfO2 gate dielectric with several substrate impurity concentrations (~1×1016 cm-3 to ~1×1018 cm-3) and various surface preparations (HF surface clean, (NH4)2S surface clean and PECVD a-Si interlayer with a HfO2 gate dielectric) have been studied. The peak electron mobility is observed to be strongly dependent on the surface preparation, but the high field mobility is observed to be almost independent of the surface preparation. A detailed analysis of the effective mobility as a function of electric field, substrate doping, and temperature was used to determine the various mobility components (surface roughness, phonon, and Coulombic scattering limited mobility components). For the substrates with high doping concentration, the electron mobility at low vertical electric field is dominated by Coulomb scattering from the substrate dopants, whereas, for lower substrate doping the Coulombic scattering is dominated by the disorder induced gap states. Low temperature measurements were used to determine the surface roughness scattering and phonon components. The results show that room temperature mobility of \(\text{In}_{0.53}\text{Ga}_{0.47}\text{As}\) surface channel MOSFETs with HfO2 gate dielectric at high electric field is limited primarily by remote phonons whereas the Al2O3 gate dielectric is limited by surface roughness scattering.
efforts to increase the dielectric constant (k) of HfO2-based gate oxides are ongoing. Recent work has shown that by tailoring annealing procedures, k values for HfO2 films of greater than 30 have been obtained. These higher k values for HfO2 occur for the metastable tetragonal and cubic crystalline phases, while the thermodynamically preferred monoclinic phase has a lower k value (~20). To evaluate the crystalline structure of ultra-thin (<100 Å) HfO2 films which have undergone various annealing treatments, we used several techniques, including grazing incidence X-ray diffraction (GIXRD), X-ray photoelectron spectroscopy (XPS and UPS, respectively) and spectroscopic ellipsometry (SE). GIXRD measurements showed that ~60 Å HfO2 films grown with a sequence of depositions and anneals (so-called DADA process) were monoclinic, while those which were post deposition annealed (PDA) were in a mixture of monoclinic and either tetragonal or orthorhombic phases. Pole figure measurements of the dipoles showed that the DADA film had a monoclinic (~111) fiber texture, while the PDA film was randomly oriented. For HfO2 films with thicknesses of ~25 Å, GIXRD measurements showed that DADA films were tetragonal or orthorhombic, while PDA films were also tetragonal or orthorhombic, but also possibly with a monoclinic component. XPS and UPS measurements of the valence bands of HfO2 films were found to be useful in distinguishing between crystalline and non-crystalline films, but were not useful in distinguishing between crystalline phases. SE has been shown to be useful in indentifying crystallinity in HfO2 through a feature that appears in the HfO2 extinction coefficient curve at the absorption edge. We have observed this absorption edge feature for films that were crystalline and strongly monoclinic, but not for films that were only weakly crystalline or mostly non-monoclinic, in keeping with previous work.

References:

10:40am EM+TF-TuM9 Research Advances on III-V and Ge MOS/MOSFETs Beyond Si CMOS. T.D. Lin, M.R. Huang, Y.C. Chang, H.C. Tsao, National Tsing Hua University, Republic of China, and L. Gao, P. Li, National Synchrotron Radiation Research Center, Taiwan, Republic of China, and J. Kwo, National Tsing Hua Univ. and National Taiwan Univ., Taiwan, Republic of China, and M. Hong, National Tsing Hua University, Taiwan, Republic of China

Invited
Metal-oxide-semiconductor (MOS) and MOS field-effect-transistors (MOSFETs) of high k dielectrics on high carrier mobility channels of InGaAs and Ge have been fiercely studied, as they are now strongly considered for technologies beyond Si complementary MOS (CMOS) integrated circuits (ICs). The post Si CMOS research is now facing unprecedented challenges in materials and physics, as key material/electrical/processing issues have to be met/solved in order to realize the new advanced devices; these include equivalent oxide thickness (EOT) < 1 nm, internal field density of state < 1011 eV·cm-2, temperature thermal stability for self-aligned process, low parasitic, and integration with Si. Using in-situ ultra high vacuum (UHV) and in-situ/ex-situ atomic layer deposited (ALD) high k’s of Ga2O3(Gd2O3), Al2O3, and HfO2 on InGaAs and Ge, this research group has made advances in achieving an EOT of 0.5 nm, Dit of low 1011 eV·cm-2 with a flat distribution versus energy within the semiconductor bandgap), and high-temperature stability of the MOS structures (rapid thermal annealing to 800-900°C and 500-600°C for the high k’s/InGaAs and /Ge, respectively). Atomic manipulation to perfect the high k’s/InGaAs and /Ge interfaces is the key for the above achievements. Probing of the interfaces and gaining insightful understanding of the electronic properties was made possible using our uniquely designed experiments of in-situ synchrotron radiation photonics. High-performance self-aligned inversion-channel high k’s/InGaAs MOSFETs in achieving record-high drain currents and transconductances, and record-low sub-threshold swings, and high k’s/Ge MOSFETs without employing interfacial passivation layers will also be discussed.


In this contribution, we investigate the influence of the SiO2 thickness on the density and polarity of built-in charges in SiO2/Al2O3 stacks deposited on Si(100). Such charges lead to the development of a space-charge region (SCR) in the Si at the dielectric interface, having consequences such as flat band voltage shifts in MOS devices and electric-field induced passivation in optoelectronic devices like solar cells. We have employed the nonlinear optical technique of second-harmonic generation (SHG) to probe the Si(100) SCR electric field through the effect of electric-field-induced SHG (EFISH). Using this non-invasive and contactless technique we found previously that the built-in charge density at the SiO2/Al2O3 interface is independent of the Al2O3 thickness down to ~2 nm. Here we report on the influence of the interfacial SiO2 layer, present between the Si(100) substrate and the atomic layer deposited (ALD) Al2O3 film, addressing the origin, density, and polarity of the charges. For this reason, we have synthesized SiO2/Al2O3 stacks with intentionally grown SiO2 interlayers having a thickness in the range ~1.4-1.5 nm using various deposition methods (e.g. thermal oxidation, PECVD, ALD). Spectroscopic SHG measurements were carried out with a femtosecond pulsed Ti:sapphire laser tunable in the 1.33-1.75 eV photon energy range. From the obtained spectra we found that the charge density is highly influenced when increasing the SiO2 thickness, dropping from ~1013 to 1011 cm-2 with the polarity switching from negative to positive. These measurements were confirmed by C-V measurements and surface voltage measurements employing corona charging of the stacks. On the basis of the obtained data, we derived charge trapping in Si/SiO2/Al2O3 stacks will be addressed.

Tuesday Morning, November 1, 2011

Ultrafast Charge and Energy Transfer in Nanomaterials Moderator: J.B. Baxter, Drexel University 8:00am EM+NS+TuM1 Controlled Deposition of Nanocrystall Quantum Dots on Silicon Surfaces: Demonstration and Application of Forster Resonant Energy Transfer. Moderator: E. Seitz, H.T. Nguyen, Y.N. Gurtstein, A.V. Malko. University of Texas at Dallas, Y.J. Chabal, The University of Texas at Dallas

Studying Forster resonant energy transfer (FRET) at semiconductor surfaces has been a challenge because of difficulties in grafting reliably nanocrystal quantum dots (NQDs) onto electronically passivated substrates.
Poor control has often resulted in formation of aggregates (3D growth), inhomogeneity, and poor adhesion. In this study, combining IR absorption spectroscopy (IRAS) and X-ray photoelectron spectroscopy (XPS), photoluminescence, atomic force microscopy (AFM) and electrical measurements, we have prepared self-assembled monolayers (SAMs) on both oxidized and oxide-free silicon surfaces with appropriate functionality to obtain dense monolayer of NQDs and to study FRET. SAMs that are directly attached to the silicon via Si-C bonds display a high interface quality as well as a low density of interfacial defects. This makes it possible to prepare systems with tunable thicknesses necessary for FRET investigation. The time evolution of the fluorescence intensity is in good agreement with the predicted thickness dependence. We are currently developing 3D structures to enhance energy collection for a given surface area. Such hybrid colloidal NQD/Silicon optoelectronic structures could potentially be attractive for both photovoltaic as well as light emitting applications.

8:20am EN+NS-TuM2 Orbital-dependent Charge Transfer Dynamics in Potential Molecular Wires, H. Hamoudi, Technische Universität München, Germany, S. Nebpl, Technische Universität München, Germany, P. Kao, Penn State University, U. G. E. Perera, S.-W. Hla, Ohio University

Continued progress in technologically important fields such as molecular and organic electronics as well as organic photovoltaics depends on reliable information about the charge transport (CT) through individual molecular groups, above all so-called molecular wires, since these represent important building blocks of a variety of devices. In this context, femtosecond CT dynamics in a series of self-assembled monolayers with oligo(phenylethenylene) and oligoacetylene backbone, which can be used for building prototypes of potential molecular wires, was addressed by resonant Auger spectroscopy using the core hole clock method. The length of the molecular backbone was varied to monitor the respective dependence of the CT time. The CT pathway was unambiguously defined by resonant excitation of the nitrile tailgroup attached to the backbone. Due to the conjugation of the electronic systems of this group and the adjacent terminal phenyl rings of the backbone, a splitting of the degenerated unoccupied molecular orbital (MO) of nitrile occurred, resulting in two different MOs which could be selectively addressed by X-rays and used as the starting points for CT. The characteristic CT times were found to depend strongly on the character of the MO which mediates the CT process. This demonstrates that the efficiency and rate of CT in molecular wires can be controlled by resonant injection of the charge carriers into specific MOs.

8:40am EN+NS-TuM3 Photophysics of Semiconductor Nanostructures in Relation to Problems of Solar Energy Conversion, V. I. Klimov, Los Alamos National Laboratory

This presentation provides a brief overview of research activities in the Center for Advanced Solar Photophysics with focus on spectroscopic properties of semiconductor nanocrystals studied from the perspective of solar cell assembly. One process, which can be used for photocurrent of solar cells, is carrier multiplication (CM) or multielectron generation. Our recent activities in this area include the development of reliable methods for efficient screening of CM performance using photon counting with superconducting nanowire detectors, the studies of the impact of “extraneous” processes on CM measurements, and the evaluation of the efficiency of the solar cell assembly, composition, dimensions, and shape of CM yields. As part of our effort on controlling excited-state dynamics, we study hot-electron transfer in nanocrystals. We find that the efficiency of this process can approach 10% even with incidental impurity-like acceptors, suggesting that even higher probabilities are possible with engineered acceptors designed for testing the ideas of hot-electron extraction. We also apply spectroscopic tools for probing the physics of charge transport in nanocrystal assemblies using exploratory devices such as optical fiber transistors (OFETs). The OFET studies help to understand the nature of conducting states in dark and under illumination and to rationalize many previously unexplained observations including a weak sensitivity of conductance to particles’ polydispersity and a significant difference in a photovoltage compared to a nominal band-gap energy. These studies illustrate how key insights into the performance of nanoscale materials are gained through close integration of spectroscopic, materials and device efforts across the Center.

9:20am EN+NS-TuM5 Hot Electron Transfer from Semiconductor Nanocrystals, W.A. Tisdale, Massachusetts Institute of Technology

In conventional semiconductor solar cells, absorption of photons with energies greater than the semiconductor band gap generate “hot” charge carriers that quickly “cool” before all of their energy can be captured – a process that limits device efficiency. Semiconductor nanocrystals (or quantum dots) have been touted as promising materials for photovoltaics because discretization of their electronic energy levels can slow down this cooling process, which might enable the extraction of photogenerated charge carriers before their excess energy is converted to heat.

In this talk, I will demonstrate hot electron transfer from PbSe nanocrystals to delocalized conduction band states of TiO2 and the concomitant excitation of coherent surface vibrational modes associated with this ultrafast process. In order to make these measurements, we developed the use of optical second harmonic generation (SHG) for femtosecond time-resolved studies of interfacial charge separation. I will discuss the information we obtain from this technique as well as the effect of temperature, nanocrystal size, and surface chemistry, and how these observations inform our understanding of electronic coupling at interfaces between confined states and bulk materials.

10:40am EN+NS-TuM9 Single Molecule Study of Charge Transfer in 6T-TBrPP-Co Molecular Complex, F. Zhang, U.G.E. Perera, S.-W. Hla, Ohio University

When two molecules having tendency to donate or accept electronic charge are put together, charge transfer between the molecules can take place. By a suitable selection of donor and acceptor molecules, it is possible to engineer an entire class of materials having metallic, semiconducting, insulating, or even superconducting properties [1, 2]. Here, we present a low temperature scanning tunneling microscopy and spectroscopy study of single molecule level charge transfer process between α-sexithiophene(6T) and TBrPP-Co molecules on a Cu(111) surface. We form molecular clusters composed of both molecular species on Cu(111). The charge transfer between the molecules is directly evident in the tunneling spectroscopy data, which reveals the shift of 6T HOMO towards the surface Fermi level indicating donation of charge from 6T to TBrPP-Co. This work is supported by the US-DOE-DE-FG02-02ER46012 grant.

Reference:

11:00am EN+NS-TuM10 NEGF Quantum Simulation of Nanotip Thermionic Emitters for Direct Energy Conversion, T.D. Musho, D.G. Walker, Vanderbilt University

Wide band-gap diamond nanotip field emission devices have been experimentally shown to have superior performance and lifetime. However, theoretical studies of the electronic emission from these devices using standard Fowler-Nordheim (FN) theory does not fully capture the physics as a result of the fitting parameters inherent to the FN approximation. The following research numerically models wide band-gap nanotip field emission devices from a quantum point of view using a novel nonequilibrium Green’s function (NEGF) approach previously applied to modeling the transport in solid-state electronic devices. In this research the IV characteristics of a single square tip diamond emitter are investigated under several bias conditions. Those bias conditions include both field emission in response to a potential bias and thermionic emission in response to a temperature bias. The NEGF model calculates the ballistic transport using a self-consistent Schrödinger-Poisson solver, calculating the transmission at discrete energy levels which is then used by the Landauer formalism to determine the total current. Ultimately, this model allows the inherent quantum mechanical transport to be captured without any fitting parameters. Findings from this research have confirmed non-linearities in the FN curve and have demonstrated the experimental transport trends. Additionally, thermionic emission trends suggest that select geometric parameters are target for enhanced emission.

11:20am EN+NS-TuM11 Mechanisms of Heterogeneous Charge Transfer at the Quantum Dot-Organic Interface, A. Morris-Cohen, M. Frederick, L. Cas, E.A. Weiss, Northwestern University

We examine the rates and mechanisms of electron transfer between colloidal semiconductor quantum dots (QDs) and viologen derivatives using ultraviolet transient absorption spectroscopy. Viologen accept electrons from photoexcited QDs on the femtosecond-to-single picosecond timescale. Modifications of the chemistry by which the ligands link to the QD surface, and the density of ligands on the QDs, facilitates control of charge separation and recombination rates, and determination of the operative mechanisms of charge transfer.
Silicon-Based Materials and Technology for Efficient Energy Conversion - A Focus on Novel Materials and Devices

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8:00am  EN-TuM1  Materials for Low Risk Nuclear Reactors, T.R. Allen, University of Wisconsin, Madison

Advanced reactor concepts have been proposed for many reasons that could be attributed to “risk reduction.” Risk reduction could mean improved safety margins, more certain economic performance, greater resistance to proliferation, or reduction of risk of exposure during the long-term storage of waste. For many of the envisioned advanced reactor concepts that have been proposed to reduce some form of risk, the ultimate deployment hinges on overcoming challenges in fuels and materials performance. The limits of performance unique to nuclear systems are typically associated with either radiation damage from high-energy particles or due to high-temperature corrosion or stress corrosion cracking. This presentation will provide an overview of the unique operating conditions in proposed reactor concepts and special challenges associated with structural materials operation. Then an overview of two approaches to improve material performance will be presented, specifically use of nanoparticles to improve high temperature strength in radiation fields and the use of tailored systems of grain boundaries to improve corrosion and stress corrosion cracking resistance.

8:40am  EN-TuM3  Battery 500 - the Li-Air Battery Opportunity, S.A. Swanson, IBM Almaden Research Center

In 2009, IBM started a project to develop rechargeable Li-air batteries for electric cars with a range of 500 miles per charge. This type of high density energy storage technology could become a game changer for the widespread adoption of electric vehicles but it also presents enormous technical challenges. Published work on Li-air batteries has only reported small fractions of the theoretical limit with limited rechargeability. Many aspects of the technology, including the lithium-oxygen electrochemistry, appear to have been poorly understood. This presentation will give an overview of Li-air battery technology. We will describe our ongoing research including our investigation of decomposition of carbonate based solvents during cell discharge using Differential Electrochemical Mass Spectrometry, the characterization and identification of the cathode electrodeposits, and enhanced cell capacities achieved using alternative aprotic solvents.

9:20am  EN-TuM5  Advanced Thermoelectric Technology for Waste Heat Recovery, G.P. Meisner, General Motors Research & Development

In today’s internal combustion engine based vehicles, more than two-thirds of the fuel energy is lost as waste heat. At General Motors Global Research & Development, we aim to demonstrate a viable thermoelectric (TE) generator system to recover that waste heat by converting it into useful electricity using advanced TE technology and thereby reduce vehicular fuel consumption. Essential to the long term success of TE technology in the automobile industry, and for waste heat recovery applications in general, is new materials research, specifically fundamental physics and materials research aimed at discovering and understanding new high performance TE materials, and the development of those materials into robust and high performance TE devices. Our work, which is generously supported by the U.S. Department of Energy’s Vehicle Technologies Program, focuses on (1) bulk TE materials (e.g., filled skutterudites) and their temperature dependent physical, mechanical, and TE properties, and (2) TE device and module development, TE generator design, and prototype TE generator fabrication, assessment, and validation. We have made significant progress on constructing working prototype automotive TE generator and testing it on a GM production vehicle.

10:40am  EN-TuM9  Improving Solar Energy Conversion with Nanoscale Materials, S.F. Bent, Stanford University

With the intensifying global need for alternative energy, there is strong interest in new approaches to materials for sustainable energy devices. A variety of different energy technologies must work in concert to produce, store, and consume the 20 TW of energy that humans will soon demand. Underlying the diverse set of energy conversion devices are similar physical and chemical phenomena, many of which can be controlled with nanoscale materials. This talk will describe research on nanoscale materials for photovoltaics and solar fuel production. Synthetic strategies including atomic layer deposition are used to generate nanoscale materials with a high level of control over composition, structure, and thickness. The materials are then tested in energy conversion devices. The prospects and challenges for such materials to contribute to higher energy conversion efficiencies will be discussed.

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8:00am  ET+EM+NS+GR-TuM  Electron Transport Study of Graphene on SiC Using Scanning Tunneling Potentiometry, K. Clark, S. Qin, Oak Ridge National Laboratory, G. He, Carnegie Mellon University, G. Gu, The University of Tennessee, R.M. Feenstra, Carnegie Mellon University, A.-P. Li, Oak Ridge National Laboratory

The unique electronic and transport properties of graphene have helped this material emerge as a perspective graphene based electronic system. Single layers of graphene formed on SiC look to be a promising system for the realization of graphene electronics. To utilize the full potential of graphene on SiC a complete understanding of the physical and electronic properties of this system is needed. This study uses Scanning Tunneling Microscope (STM) images along with scanning tunneling spectroscopy to characterize the surface, STM images clearly show the distinction between 1 monolayer (ML) and 2ML regions. The 1ML to 2ML transition is further confirmed by point spectroscopy measurements and spectroscopic mapping across the boundary. Defects, grain boundaries, step edges and other potential scattering centers are thought to play a major role in the electronic properties, especially in transport, along the graphene sheets. Using a low temperature four-probe scanning tunneling microscope, potentiometry measurements are performed on epitaxial graphene grown on 4H-SiC. Potentiometry maps spanning the transition from 1ML to 2ML graphene layers show a contrast change indicating a potential change at this interface. Preliminary results of the transport along this potentially revolutionary new electronic system will be presented. This research was conducted at the Center for Nanophase Materials Sciences, which is sponsored at Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy.

8:20am  ET+EM+NS+GR-TuM2  Engineering the Electronic States of CVD Grown Few Layer Graphene by Twisting and Lattice Distortion, M.H. Pan, Oak Ridge National Laboratory, X.T. Jia, S. Bhavaripudi, Massachusetts Institute of Technology, F. Meunier, Rensselaer Polytechnic Institute, J. Kong, Massachusetts Institute of Technology

Few layer graphene (FLG) can have advantages over single layer graphene because it has a larger current-carrying capacity and the electronic properties are sensitive to more engineerable system parameters. In particular, Hass et al. have demonstrated that orientational disorder is normally present in carbon-face SiC epitaxial FLG samples. [1] Recently both theoretical and experimental studies suggest that strain can be used to engineer graphene electronic states through the creation of a pseudo-magnetic field. [2] Here we present both scanning tunneling microscopic/spectroscopic (STM/S) studies of chemical vapor deposition grown few layer graphene samples. There indeed exists a twisting between the stacked graphene layers, confirmed by both high-resolution STM images and low temperature spectroscopic measurements. Our results show that, by stretching graphene along three symmetry directions, a strain-induced pseudo magnetic field can lead to the formation of different Charge Density Wave (CDW) states at the top layer of graphene.


8:00am  ET+EM+NS+GR-TuM3  Unique One- and Two-Dimensional Phenomena Observed in Carbon Nanotubes and Graphene, S. Cronin, University of Southern California

Our ability to fabricate nearly defect-free, suspended carbon nanotubes (CNTs) has enabled us to observe several phenomena never seen before in CNTs, including breakdown of the Born-Oppenheimer approximation[1], mode selective electron-phonon coupling[2], leading to negative differential resistance (NDR) and non-equilibrium phonon populations, and a Mott
tunneling microscopy is used to probe the electrical transport through...0937060 (subaward CIF-146) and DOE.

Acknowledgement:...coupling. And the reversible remnant polarization of the ferroelectric gate observed at room temperature, owing to the enhanced gate capacitance...We measure the membrane conductance as a function of both contribution, enabling experimental determination of the surface...the backgate voltage is varied, and the conductance in this regime does not depend on membranes...Because they are so thin, SiNMs display interesting transport phenomena...The typical membrane thickness is less than a few hundred nanometers. Because they are so thin, SiNMs display interesting transport phenomena influenced by surface effects. Here, we demonstrate a novel method to probe surface transport via conductance measurements on SiNMs. When contacts are placed on the front surface, a current flows between the source and the drain via the membrane body as well as its surface. By utilizing an underlying back gate (the Si handle substrate), the conductance through the membrane can be continuously tuned and made smaller than the surface contribution, enabling experimental determination of the surface conductance. We measure the membrane conductance as a function of both the membrane thickness and the backgate voltage in ultra-high vacuum. In contrast to H-terminated Si surfaces, clean reconstructed Si(001)(2×1) surfaces show a constant-conductance regime when the back-gate voltage is varied, and the conductance in this regime does not depend on membranes thickness. We demonstrate that the constant conductance (on the order of 10⁸ Siemens) stems from an additional conduction channel through the dimer-reconstructed surface π* band. By comparing the experimental results to numerical simulations, the surface band mobility is determined to be in the range 10-50 cm²/Vs.

Research supported by NSF [UW MRSEC, award DMR-0520527, as well as awards 0937060 (subaward CIF-146) and ECCS-0547415] and DOE.


Complex nanostructures such as branched semiconductor nanotetrapods are promising building blocks for next-generation nanoelectronics. Here we report on the electrical transport properties of individual CdS tetrapods in a field-effect transistor (FET) configuration with a ferroelectric Ba,Sr,TiO₃ film as high-k, switchable gate dielectric. A cryogenic four-probe scanning tunneling microscopy is used to probe the electrical transport through individual nanotetrapods at different temperatures. A p-type field effect is observed at room temperature, owing to the enhanced gate capacitance coupling. And the reversible remanant polarization of the ferroelectric gate dielectric leads to a well-defined nonvolatile memory effect. The field effect is shown to originate from the channel tuning in the arm/core/arm junctions of nanotetrapods. At low temperature (8.5 K), the nanotetrapod devices exhibit a ferroelectric-modulated single-electron transistor behavior. The results illustrate how the characteristics of a ferroelectric such as switchable polarization and high dielectric constant can be exploited to control the functionality of individual 3-dimensional nanoarchitectures.

Acknowledgement: The research at the Center for Nanophase Materials Sciences is sponsored at Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy. The research in Beijing is supported by MOST and CAS of China.


Strong electron-electron correlations occur in nanoscale systems in a variety of contexts – when electrons form a crystal at low density, for example, or in correlations between quantum dots. Nanoscale systems introduce in addition an unprecedented level of control over the physical parameters determining such correlations. As electron transport is one of the primary probes of nanosystems, the understanding of correlations on transport is a key issue. I shall discuss an example in which we used quantum Monte Carlo (QMC) techniques to calculate the conductance.

Consider a system of four quantum dots designed to study the competition between three types of interactions: Heisenberg, Kondo, and Ising. We find that the competition produces a rich phase diagram containing two sharp features: a quantum phase transition (QPT) between charge-ordered and charge-liquid phases, and a dramatic resonance in the charge liquid visible in the conductance. The conductance is calculated using a world-line QMC method: extrapolation of the imaginary time QMC data to zero frequency yields the linear conductance, which is then compared to numerical renormalization group results in order to assess its accuracy. The QPT is of the Kosterlitz-Thouless type with a discontinuous jump in the conductance at the transition. We connect the sharp resonance phenomenon with the degeneracy of three levels in the isolated quadruple dot and argue that this leads to an emergent symmetry. I shall end by discussing the sensitivity to parameter variation and possible experimental realizations in laterally gated quantum dots as well as carbon nanotubes.

This work was done in collaboration with Dong E. Liu and Shailesh Chandrasekharan (Duke University).

11:20am ET+EM+NS+GR-TuM11 Resistivity Increase due to Electron Scattering at Surfaces and Grain Boundaries in Metal Thin Films and Nanowires. J.S. Chawla, D. Gall, Rensselaer Polytechnic Institute

The effect of surface and grain boundary scattering on the resistivity of Cu thin films and nanowires is quantified using (i) in situ transport measurements on single-crystal, atomically smooth Cu(001) layers, (ii) textured Cu(111) layers and patterned Cu wires with independently varying grain size, thickness and line width, and (iii) in situ grown interfaces including Cu-Ta, Cu-MgO, Cu-SiO₂ and Cu-oxygen. In addition, the electron surface scattering is also measured in situ for single-crystal Ag(001) and TiN(001) layers. These findings are important for the development of future generation narrow low-resistivity Cu interconnects and TiN metal gates.

Cu(001), Ag(001), and TiN(001) layers with a minimum continuous thickness of 4, 5 and 1.8 nm, respectively, are grown by ultra-high vacuum magnetron sputter deposition on MgO(001) substrates and are found to be atomically smooth single crystals by a combination of x-ray diffraction θ-2θ scans, o-trocking curves, pole figures, reciprocal space mapping, Rutherford backscattering, x-ray reflectometry, transmission electron microscopy, and in-situ scanning tunneling microscopy. Polycrystalline Cu layers with a 111-texture are deposited on thermally grown SiO₂, with and without Ta barrier layer. Subsequent in-situ annealing at 550°C followed by sputter etching in Ar plasma yields Cu layers with independently variable thickness and grain size. Cu nanowires, 50 to 150 mm long, 70 to 350 nm wide, and 45 nm thick, are patterned using electron beam lithography and sputter etching.

In-situ electron transport measurements at room temperature in vacuum and at 77 K in liquid nitrogen for single-crystal Cu and Ag layers is consistent with the Fermi-Sondheimer (FS) model and indicates specular scattering at the metal-vacuum boundary with an average specularity parameter p = 0.6 and 0.4, respectively. In contrast, layers measured ex-situ show complete diffuse surface scattering due to sub-monolayer oxidation. Electron transport measurements for polycrystalline Cu/Ta layers and wires show a ~10% and ~11% decrease in resistivity, respectively, when increasing the average lateral grain size by factor 2. In-situ deposition of 0.3 to 8 nm thick Ta barrier layers on Cu(001) leads to a resistance increase that indicates a transition from p = 0.8 to p = 0, independent of the Ta thickness. In-situ exposure of Cu(001) layers to O2between 10-3 and 105 Pa-s results in a sequential increase, decrease and increase of electrical resistance which is attributed to specular surface scattering for clean Cu(001) and for surfaces with a complete adsorbed monolayer, but diffuse scattering at partial coverage and after chemical oxidation.

* NSTD Student Award Finalist
Graphene and Related Materials Focus Topic
Room: 208 - Session GR+EM-TuM

Graphene: Optical Properties, Optoelectronics and Photonics
Moderator: P.E. Sheehan, U.S. Naval Research Laboratory

The richness of optical and electronic properties of graphene attracts enormous interest. So far, the main focus has been on fundamental physics and electronic devices. However, we believe its true potential to be in photonics, plasmonics and optoelectronics, where the combination of its unique optical and electronic properties can be fully exploited, the absence of a bandgap can be beneficial, and the linear dispersion of the Dirac electrons enables ultra-wide-band tunability [1]. The rise of graphene in photonics and optoelectronics is shown by several recent results, ranging from solar cells and light emitters to touch screens, photodetectors and ultrafast lasers. Despite being a single atom thick, graphene can be optically visualized [2]. Its transmittance can be expressed in terms of the fine structure constant [3]. The linear dispersion of the Dirac electrons enables ultra-wide-band tunability [1]. 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A plasmonic nanostructure can act like an optical antenna, concentrating light into a deep sub-wavelength volume and enabling manipulation of light-electron interactions at the nanometer scale. Achieving efficient coupling from such antennas to functional electrical devices has been challenging. Alas, the effective region of field enhancement is so small. We achieve direct electrical read out of the wavelength and polarization dependence of the plasmon resonance in individual gold nanogap antennas by positioning a graphene nanoconstriction within the gap as a localized photodetector. The polarization sensitivities can be as large as 99%, while the plasmon-induced photocurrent enhancement is 2-100. The plasmon peak frequency, polarization sensitivity, and photocurrent enhancement all vary between devices, indicating the degree to which the plasmon resonance is sensitive to nanometer-scale irregularities.

11:00am GR+EM-TuM10 Large Area Graphene Growth for Optoelectronic Applications. C. Edwards, C.L. Berrie, J. Liu, J. Wu, University of Kansas
Graphene shows great promise for numerous applications within the field of optoelectronics due to its high charge mobility, high optical transmittance, chemical inertness, and flexibility. We are focused on developing large-area graphene sheets that sustain or enhance these characteristics currently present in small-area growth. Chemical vapor deposition of graphene onto various copper substrates has been investigated to understand the role of the substrate in graphene epitaxy and its deposition mechanism. With this understanding it will be possible to use the substrate structure to control the density of surface defects, which is high in current methods for large area fabrication. The effect of nanopatterning and doping graphene by chemical vapor deposition has been investigated, and observed improvements in light transmittance and electrical conductivity suggest the potential to favorably modify graphene optical and electrical properties for these applications.

Since becoming experimentally available by mechanical exfoliation, graphene has been used in various devices such as field effect transistors (FETs), Schottky based solar cells and sensing applications. Although graphene based devices with modest characteristics have been reported, in some of the device geometries a lower graphene sheet resistance with different Fermi level values is still desired. To achieve these ends, graphene’s physical properties have been adjusted by n- or p- chemical doping using AuCl4-, Br-, N2, and organic solutions. However, these techniques have several drawbacks which prevent their use in devices, namely: environmental instabilities, aging effects and a reduction in optical transparency. Here, we describe our use of a hydrophobic organic complex dopant with strong electronegativity, tight bonding, environmental stability and high optical transmittance which is spin cast onto CVD-prepared graphene films. We observe a typical 75% reduction in sheet resistance upon chemical modification of the graphene. Resistance vs. temperature / magnetic field and Hall measurements imply that the modified graphene sheets are doped, and time-dependent resistance measurements show excellent stability. Using a Horiba Micro Raman instrument we confirm the doping of graphene sheets from the shifts in G and 2D peak positions and intensity ratios. We show transmittance and SEM characteristics of the graphene sheets before and after doping. The presented results may serve as a guide for modification of graphene’s properties as desired for various applications.

In Situ Spectroscopy and Microscopy Focus Topic

In Situ Studies of Organic and Soft Materials and Liquid-Solid Interfaces

Moderator: A.I. Frenkel, Yeshiva University
8:00am IS+AS+SS-TuM1 Solid-Vacuum, Solid-Gas, and Solid-Liquid Interfaces: Structure and Dynamics under Environmentally Relevant Conditions. M. Salmeron, C. Escudero, Lawrence Berkeley National Laboratory
Surfaces play a fundamental role in many of today's frontier topics, such as clean and renewable energies, efficient and highly selective chemical processes (green catalysis), high capacity rechargeable batteries and fuel cells, and also environmental problems. To advance our understanding it is imperative to develop new fundamental approaches to the study of the interface of solid materials with gases, liquids and solids, because it is in these environments that crucial processes occur that need to be understood to enable game-changing discoveries.

One way to control the structure of interfaces and their properties is through the design of materials of nanoscale dimensions, with specific shape, size and composition. It is equally imperative to develop and use techniques for in-situ and in-surface level structural and spectroscopic characterization of the interfaces. New advances in instrumentation are fulfilling this need. I will illustrate this with examples from research carried out in my laboratory, which include scanning tunneling microscopy (STM), photoelectron and x-ray absorption spectroscopies (XPS) under ambient conditions, for studies of catalyst models, thin films, single crystals and nanoparticles, for applications in catalysis and electrochemistry. The results obtained so far demonstrate that the information obtained with these new techniques is unique and could not have been obtained or extrapolated from other more traditional surface sensitive techniques.

8:40am IS+AS+SS-TuM3 Imaging Tagged Proteins in Whole Eukaryotic Cells in Liquid with Scanning Transmission Electron Microscopy. N. De Jonge, D.B. Peckys, Vanderbilt University School of Medicine
We have recently introduced a novel electron microscopy technique for the imaging of whole cells in aqueous media using scanning transmission electron microscopy (STEM) [1, 2]. Eukaryotic cells in liquid were placed in a microfluidic chamber with a thickness of 5 - 10 μm contained between two ultra-thin electron-transparent windows. On account of the atomic number (Z) contrast of the STEM, nanoparticles of a high-Z material (e.g., gold) were detected within the background signal produced by a micron-thick layer of a low-Z liquid (e.g. water, or cellular material). Nanoparticles specifically attached to proteins can be used to study protein distributions in whole cells in liquid, similar as proteins tagged with fluorescent labels can be used to study protein distributions in cells with fluorescence microscopy.

COS7 fibroblast cells were labeled with gold nanoparticles conjugated with epidermal growth factor (EGF). Intact fixed cells in liquid were imaged with STEM with a spatial resolution of 4 nm and a pixel dwell time of 20 microseconds [1]. In test experiments we demonstrated a maximal spatial resolution of 1.5 nm on gold nanoparticles placed above a water layer of a thickness of 3 micrometer, consistent with theoretical predictions, and with Monte Carlo simulations of the STEM imaging [3]. The use of quantum dots (QDs), which are fluorescent nanoparticles, allowed STEM images to be correlated with fluorescence images [4]. Eukaryotic cells were grown directly on microchips for the microfluidic chamber, fixed, and imaged with fluorescence microscopy. The intact cells were then imaged in liquid with STEM. The STEM images showed individual QDs, and their locations were correlated with the cellular regions, as imaged with fluorescence microscopy. We have also demonstrated the imaging of nanoparticle uptake in live cells [5], and the ultrastructure of pristine yeast cells was studied [6]. Liquid STEM presents an innovative approach for the imaging of whole cells, with significantly improved spatial resolution and imaging speed over existing methods.

References

URL: http://www.mc.vanderbilt.edu/labs/dejongelab/

Tuesday Morning, November 1, 2011
9:20am IS+AS+SS-TuM5 Imaging Live Cells in Liquid with Scanning Transmission Electron Microscopy. D.B. Peckys, N. De Jonge, Vanderbilt University School of Medicine

We have applied a novel electron microscopy technique, referred to as liquid scanning transmission electron microscopy (liquid STEM) [1, 2] for the imaging of live eukaryotic cells. In two separate experiments, we studied a) nano particle (NP) uptake in COS-7 cells [3], a green monkey kidney fibroblast cell line, and b) the ultrastructure of Schizosaccharomyces pombe cells [4], also known as fission yeast. The cells were confirmed to be alive at the onset of the liquid STEM imaging using specific fluorescent, live indicating dyes and correlated fluorescence microscopy. For the STEM imaging in liquid the cells were placed (in liquid) in a microfluidic chamber. The chamber had two ultra-thin electron-transparent windows allowing the passage of electrons and photons. The dimensions of the COS-7 cells required a thicker liquid filled space in the microfluidic chamber compared to the experiments with the fission yeast cells, and contrast was mainly obtained on the gold NP's. However, the thinner S. pombe cells allowed a thinner liquid layer, and images were recorded of the cellular ultrastructure.

Despite the fact, that the cells were not anymore alive after the STEM imaging, we consider the first STEM images taken from a cell or a specific cellular region, to represent the unperturbed and therefore physiological state. We derived this assumption after evaluation of the STEM images for signs of radiation damage at the achieved resolution. Our STEM results were found to be consistent with known data about intracellular NP trafficking and storage in mammalian cells and data about the dimensions and distribution of organelles in fission yeast.

In conclusion, we have demonstrated the feasibility of STEM imaging live eukaryotic cells. The advantages of this approach are a) a several-fold higher resolution than live cell imaging with conventional light microscopy, b) a much faster (hours versus days) sample preparation than needed for conventional transmission electron microscopy (TEM) imaging of cells, c) absence of artifact introduction associated with conventional TEM sample preparation, and d) no need for introducing any kind of labels in order to achieve a similar range of resolution as possible with the new nanoscopic imaging techniques.

References

9:40am IS+AS+SS-TuM6 Microscopic Imaging of Biological Samples using Coherent Soft X-rays from Free-Electron Laser and Synchrotron Sources. T. Gorniak, T. Senkbeil, M. Beckers, C. Christopher, University of Heidelberg, Germany, K. Giewekemeyer, University of Göttingen, Germany, M. Granze, University of Heidelberg, Germany, T. Salidtt, University of Göttingen, Germany, A. Rosenhahn, University of Heidelberg, Germany

Coherent X-ray microscopy of hydrated biological samples – especially in the so-called water window of 284-540 eV – is of tremendous interest for life sciences due to the high contrast of organic matter with respect to the aqueous background. Especially free-electron lasers can provide highly intense and coherent pulses, which allow single pulse imaging to overcome resolution limits set by radiation damage. We present the first holographic microscopy images of dehydrated biological material acquired in the water window with higher harmonic radiation provided by the free-electron laser FELIX. In order to increase the photon flux we used high efficiency zone plates instead of pinholes to create the divergent light cone for holography. The results pave the way to the vision of holographic imaging of hydrated biological samples with single FEL pulses. We supplement single pulse imaging experiments by ptychographic imaging with synchrotron radiation at BESSY II. This method uses coherent diffraction imaging at different sample positions while maintaining a fixed spatial overlap between the fields of view. By introducing this spatial redundancy to the data an additional constraint for the iterative reconstruction algorithm is achieved. This enhances the convergence of phase retrieval drastically. The spatial resolution of below 50 nm and the imaging properties were characterized using lithographic and biological test samples. We also show results on resonant imaging with chemical contrast caused by both, absorption and phase shifts, in the vicinity of core level absorption edges.

10:40am IS+AS+SS-TuM9 Surface Chemistry of Amino Acids at Near Ambient Pressure of Water Vapor. A. Shavrovskiy, Lawrence Berkeley National Laboratory, T. Eralp, The University of Reading, UK, F. Aksoy, Nagde University, Turkey, M.E. Grass, Z. Liu, H. Bluhm, Lawrence Berkeley National Laboratory, G. Held, The University of Reading, UK

The co-adsorption of water with organic molecules under near-ambient pressure and temperature conditions opens up new reaction pathways on model catalyst surfaces that are not accessible in conventional ultra-high vacuum surface-science experiments. The surface chemistry of glycine and alanine at the water-exposed Cu{110} and Pt{111} interface was studied both in situ and in UHV using ambient-pressure photoemission and X-ray absorption spectroscopy techniques [1,2]. At water pressures above 10-5 Torr a significant pressure-dependent decrease in the temperature for dissociative desorption was observed for both amino acids on Cu{110}[3]. On Pt{111}, on the other hand, desorption temperature does not depend significantly on the presence of water vapor. The most likely reaction mechanism of decomposition involves dehydrogenation induced by O and/or OH surface species resulting from the dissociative adsorption of water on Cu{110}, but not on Pt{111}.

The linear relationship between the inverse decomposition temperature on Cu{110} and the logarithm of water pressure enables determination of the activation energy for the surface reaction, between 213 and 232 kJ/mol, and a prediction of the decomposition temperature at the solid-liquid interface by extrapolating towards the equilibrium vapour pressure. Such experiments near the equilibrium vapour pressure provide important information about elementary surface processes at the solid-liquid interface, which can neither be retrieved under ultra-high vacuum conditions nor from interfaces immersed in a solution.


11:00am IS+AS+SS-TuM10 STM Tip Catalyzed Adsorption of Thiol Molecules and Functional Group-Selective Adsorption of Bi-Functional Molecule Using This Catalysis. Y.H. Min, S. Kim, S.J. Jung, Y.-S. Youn, Korea Advanced Institute of Science and Technology, Republic of Korea, D.H. Kim, Daegu University, Republic of Korea, E.H. Park, Korea Advanced Institute of Science and Technology, Republic of Korea

In this study, in contrast with cases in which Scanning Tunneling Microscopy (STM) tip-induced reactions were instigated by the tunneling current, local electric field, or the mechanical force between a tip and a surface, we found that the tungsten oxide (WO₃) covered tungsten (W) tip of a STM acted as a chemical catalyst for the S-H dissociative adsorption of phenylthiol and 1-oxanethiol on a Ge(100) surface. By varying the distance between the tip and the surface, the degree of the tip-catalyzed adsorption could be controlled. We have found that the thiol head-group is the critical functional group for this catalysis and the catalytic material is the WO₃ layer of the tip. After removing the WO₃ layer by field emission treatment, the catalytic activity of the tip has been lost.

3-mercapto isobutyric acid is bi-functional molecule which has two functional groups, carboxylic acid group and thiol group, at each end. 3-Mercapto Isobutyric Acid adsors at Ge(100) surface only through carboxylic acid group at room temperature and this adsorption was enhanced by the tunneling electrons between a STM tip and the surface. Using this enhancement, it is possible to make thiol group-terminated surface where we desire. On the other hand, surprisingly, the WO₃ covered W tip of STM was found to act as a chemical catalyst to catalyze the adsorption of 3-mercapto isobutyric acid through thiol group at Ge(100) surface. Using this catalysis, it is possible to make carboxylic acid group-terminated surface where we want. This functional group-selective adsorption of bi-functional molecule using the catalysis may be used in positive lithographic methods to produce semiconductor substrate which is terminated by desired functional groups.

11:20am IS+AS+SS-TuM11 CO2 Capture in Aqueous Monoethanolamine Solutions: Role of the Solution Interface Investigated with X-ray Photoelectron Spectroscopy, T. W. Negishi, University of California, Irvine, Irvine, CA. Winter, Helmholtz-Zentrum Berlin für Materialien und Energie, Germany, M. Faubel, Max-Planck-Institut für Dynamik und Selbstorganisation, Germany, J.C. Hemminger, University of California, Irvine

Aqueous monoethanolamine (MEA) solutions are commonly used to capture CO2 emitted into the atmosphere. An important challenge in industry is likely that interactions between MEA and CO2 at the aqueous solution surface are important to this process, yet surprisingly very few studies have explicitly addressed the role of the solution-gas interface. In the present study, interfacial chemistry of CO2 capture is studied by surface sensitive photoelectron spectroscopy measurements from a liquid microjet of CO2-reacted MEA solutions with carbon loadings of 0.1 to 0.9 mol/mol. These experiments determine the spatial distribution of MEA and reaction products into the solution as a function of CO2 loading. Results show that neutral MEA exhibits a propensity for the solution surface, whereas protonated MEA and reaction products prefer bulk solvation, suggesting enhanced reactivity at the solution interface, especially at high CO2 loading. These observations indicate that a detailed understanding of the chemistry of CO2 at the liquid-vapor interface and interface to bulk transport of the products will be important in understanding CO2 capture.


Over the past several years significant advances have been made concerning our understanding of the growth of crystalline small molecule organic thin films consisting of a single component. A major challenge in electronics, photonics and photovoltaics is to develop and improve methods to integrate both p-type and n-type small molecule organic semiconductors into the same device microstructure. Thus, developing an understanding of the molecular scale events that lead to heterojunction formation is essential in these systems consisting of multiple components. Here we report on our examinations of the nucleation, growth, and dynamics of adsorption of a n-type organic semiconductor, N,N-ditridecylerpylene-3,4,9,10-tetracarboxylic diimide (PTCDI-C13), on SiO2 surfaces modified by self-assembled monolayers (SAMs) and on a pre-deposited monolayer of pentacene (a p-type semiconductor) using supersonic molecular beam techniques, in situ synchrotron x-ray scattering and ex situ atomic force microscopy. From real-time x-ray scattering we find that PTCDI-C13 exhibits prolonged layer-by-layer growth for approximately the first 10 monolayers (MLs) of deposition on all three SAMs examined. Concerning the kinetics of growth we find that the adsorption probability of PTCDI-C13 on itself is similar to that observed on two SAMs that possess aromatic endgroups, but it differs significantly to that observed on a relatively short, methyl-terminated SAM. These differences could reflect mechanisms such as direct molecular insertion of PTCDI-C13 into either the existing PTCDI-C13 film, or the longer chain SAMs with aromatic endgroups. Concerning growth in the submonolayer regime, we find that nucleation is homogenous, and that the absolute density of islands depends on the nature of the surface, while the relative change of the island density with increasing growth rate is essentially independent of the underlying SAM. From the latter we find that a critical island size of a single molecule of PTCDI-C13 can describe all the data. Finally, we will discuss our most recent results concerning the growth of heterostructures composed of a few to several monolayer stacks of PTCDI-C13 and pentacene. In this work we find that PTCDI-C13 grows in a smooth layer-by-layer fashion on pentacene, but the opposite is not true—pentacene grows in a purely 3D mode when deposited on PTCDI-C13. We will discuss implications of this observation concerning the growth of organic heterostructures for applications in electronics, photonics and photovoltaics.

**Nanomanufacturing Science and Technology Focus Topic**

Room: 207 - Session NM+MN+MS+TF-TuM

**Lithography Strategies for Nanomanufacturing**

Moderator: T.S. Mayer, Penn State University

8:00am NM+MN+MS+TF-TuM1 A SANE Approach to Programmable Soft Lithography. T.W. Odom, Northwestern University

The prototyping of nanoscale features has rarely been separated from the scaling of them. In order to create arbitrary patterns, serial techniques such as e-beam lithography or focused ion beam milling must start from scratch every time; also, the patterns cannot be generated over large areas. In contrast, parallel fabrication techniques such as imprint lithography and soft lithography can scale patterns, but they are limited to transferring the same pattern on the mold. The development of new tools that can combine the strengths of serial approaches (prototyping patterns, high resolution) with those of parallel ones (high throughput, large patterned areas) is critical for next-generation applications based on nanostructures.

This talk will describe an all-moldable nanofabrication platform that can generate—from a single master—large-area nanoscale patterns with programmable densities, fill factors, and lattice symmetries. Solvent-assisted nanoscale embossing (SANE) can increase the spacing of patterns up to 100% as well as decrease them down to 50% in a single step by stretching or heating a thermoplastic substrate. In addition, SANE can reduce critical feature sizes as small as 45% compared to those on a master by controlled swelling of patterned molds with different solvents. SANE can also produce different and reconfigurable lattice symmetries, which enables new opportunities to manipulate the electronic, photonic, and magnetic properties of nanomaterials.


This presentation will introduce the use of confined surface-initiated ring-opening metathesis polymerization (SI-ROMP) of perfluoroalkyl or alkyl norbornene monomers from solid substrates to synthesize surface-bound polymer structures with tunable physical and chemical properties that accurately replicate those exhibited by Nature’s engineered, microscopically rough, and highly functional surfaces. This approach not only allows mimicking of highly evolved and functional surface architectures but also provides versatility in that it introduces a wide variety of chemical compositions available in materials chemistry, including partially fluorinated polymers with ultralow critical surface tensions. Optical microscopy and scanning electron microscopy confirm growth of the polymer structures and the precise replication of the microscale and nanoscale features exhibited by the target natural surface with the added freedom to expand beyond the intrinsic feature size of the organic mask. Angle dependent measurements show that the surface architectures exhibit both hydrophobic and oleophobic behavior, and in some cases, superhydrophobic properties. This approach is not limited to natural surfaces and could be applied in a straightforward manner to a variety of synthetic surfaces that have microscale features.

9:00am NM+MN+MS+TF-TuM4 Si Mold Etching with Hard Mask for Bit-Patterned Media. M. Kurihara, Hitachi, Ltd., Tokyo, M. Satake, T. Tsuchiya, T. Nishida, Central Research Laboratory, Hitachi, Ltd., Tokyo, Y. Tada, H. Yoshida, Hitachi Research Laboratory, Hitachi, Ltd., Japan, N. Negishi, Central Research Laboratory, Hitachi, Ltd., Japan

Bit-patterned media (BPM) is one of the promising candidates for hard disk media with areal density greater than 1.0Tb/in2. Nanoimprint lithography (NIL) for BPM has also been investigated as a patterning technique to reduce the production cost. The critical issue in BPM mold fabrication is the etching selectivity between silicon and the organic mask pattern due to the following two reasons. One is a significant decrease of pattern thickness to meet the photo-lithography requirements. This decrease remained when applying the self-assembly polymer process to fine patterning. The other is the micro-loading effect that causes the etching rate drop with pattern size shrinking.

In this work, we have developed a hard mask process to compensate for the low etching selectivity. The micro-loading effect caused by the SiO2 hard mask was evaluated with a SiO2 hard mask of 20-nm thickness. This SiO2 hard mask was patterned from 30 to 50 nm by EB lithography and etched with CHF3 gas chemistry. The coefficients of micro-loading in silicon etching were evaluated based on the relationship between hole depth and etching time with a hole diameter of 30, 40, and 50 nm respectively. With increase of the sidewall taper angle, the micro-loading effect could be improved by about 60%. We also confirmed that there was a hole with a depth of 87 nm with a diameter of 30 nm. Extrapolating this micro-loading effect, it is expected that a hole with a depth greater than 80 nm with a diameter of 10 nm will be achieved. This result will satisfy the pattern aspect ratio of 2, which is required in NIL. We will also demonstrate the Si mold etching with a hard mask by applying the self-assembly polymer in which the areal density is greater than 1.0Tb/in2.
Our research program aims to integrate self-assembling block copolymers into current manufacturing practice. The fundamental concepts of the approach are that 1) the most advanced production-oriented exposure tools (e.g. 193 nm, EUV, or electron beam lithography) and resist materials are used to create patterns of differing chemical functionality on the substrate, and 2) use of block copolymers can be directed to assemble in the presence of the chemical pattern into predictable and desirable morphologies, thereby augmenting and enhancing the lithographic process. In comparing the pattern in resist to the pattern of domains induced to morphologies, thereby augmenting and enhancing the lithographic process. In order to measure the backscattered profile, we use a pattern of lines with varying doses. By measuring the thickness of the resist, the contribution from both the lines and the annulus can be determined for different doses and different shapes by using the contrast curve. By using the contrast curve to assign doses to specific resist thicknesses, the specific resist and developer effects are removed from the data. The resultant backscattered electron profile can be used as input into the FEM model to create more accurate resist dose assignments for proximity effect corrected patterns for all resists.

The use of planar Si CMOS device technology may continue beyond the 22 nm node. However, the requirements for the gate dielectric and junction depth needed to maintain control of short channel effects might prove to be unobtainable in devices scaled to meet the integration density requirements of the 14 nm node and beyond. Consequently, an additional method for improving the electrostatics of the device is required. This realization has driven a steady increase in research on non-planar multi-gate CMOS devices over the past 5 years. Raising the Si channel out of the plane of the substrate creates the opportunity to form the gate electrode above multiple sides of the channel. This geometry results in a superior situation from an electrostatics standpoint compared to a planar device where the gate electrode is present only on the top surface of the channel.

In this talk, we will discuss the challenges of fabricating three non-planar multi-gate devices from Si on insulator (SOI) substrates: (1) the FinFET, where the gate controls two sides of a thin Si mesa or fin (2) the Trigate where the gate controls three sides of a thin Si mesa and (3) a gate-all-around nanowire transistor where the gate electrode surrounds all sides of a suspended Si channel. We will present experiments results from advanced prototypes of these devices fabricated at dimensions and densities relevant to 14 and 10 nm node technology. An emphasis will be given to the unique role lithography and patterning play in determining the electrical behavior of these devices. These results offer insight into what may lie ahead for Si CMOS scaling and how it will impact the demands placed on patterning and metrology.
9:00am NS-TuM4 Semiconductor Nanowires: From Materials Physics to Devices. L. Samuelson, Lund University, Sweden. INVITED

The field of self-assembled growth of semiconductor nanowires has emerged as a powerful way to form advanced materials and devices on the 10-nm scale. In this talk I will review the state of the art of the field, exemplified by our recent progress in growing ideal and defect-free III-V and III-nitride nanowires, on either III-V or silicon as substrates. I will discuss the hot materials physics aspects of structural control and the influence of zincblende vs wurzite crystal structures on materials properties. I will then review recent progress in realizing various electronic as well as opto-electronic devices, such as nanowire field-effect transistors, tunnel devices, solar-cells as well as light-emitting diodes.

9:40am NS-TuM6 Self-Catalyzed Growth of InP$_x$Sb$_{1-x}$ Nanowires on InP(111)B C. Ngo, M. Pozuelo, M. Mecklenburg, H. Zhou, B.C. Regan, R.F. Hicks, S. Kodambaka, University of California Los Angeles

Group III–V semiconductors possess high carrier mobilities and small band gaps, making them applicable to nanoelectronics and optoelectronics.[1] Properties of these materials can in principle be controllably tuned by the fabrication of low-dimensional structures, such as nanowires. Nanowires are most commonly grown via the vapor-liquid-solid (VLS) process using Au as the catalyst. A variant of this approach is the self-catalyzed VLS process, where one of the elements of the growing material promotes the one-dimensional growth on the substrate to yield nanowires. Recent studies show that InP and InP$_x$Sb$_{1-x}$ nanowires can be grown via metalorganic chemical vapor deposition (MOCVD) using indium and indium-antimonide droplets as catalysts.[2] In this talk, we present results from studies focused on understanding the influence of metallicorganic precursor flow rates on the compositional and structural evolution of InP$_x$Sb$_{1-x}$ alloy nanowires.

All of our samples are grown via MOCVD using trimethylindium (TMIN), tertiarybutylphosphine (TBP), and trimethylantimonide (TMSb) as precursors, with liquid indium and indium-antimonide as substrates. The as-grown structures are characterized using scanning transmission electron microscopy, selected area electron diffraction, energy dispersive x-ray spectroscopy, and scanning TEM to determine their morphology, crystallinity, and composition.

We demonstrate the successful growth of InP$_x$Sb$_{1-x}$ alloy nanostructures of desired Sb content that is tunable with substrate temperature and TMSb flow rate. Interestingly, doubling the precursor flow rates at a given temperature leads to crystalline growth of pure wurzite-structured nanowires at the catalyst-wire interface. We attribute this phenomenon to the precipitation of excess Sb present in the indium droplets during InP$_x$Sb$_{1-x}$ alloy growth. By taking advantage of the differences in growth kinetics of InP and InSb, we demonstrate the formation of compositionally-abrupt interfaces in InP/InSb axial nanowire heterostructures. [1] M. Pozuelo, S.V. Prikhodko, R. Grantab, H. Zhou, L. Gao, S.D. Sitzman, V. Gambin, V.B. Shenoy, R.F. Hicks, and S. Kodambaka, “Zincblende to Wurtzite Transition (During the Self-Catalyzed Growth of InP Nanowires).” J. Crystal Growth 312, 2305 (2010).

phase-boundary is dramatically reduced by hydrogen adsorption and drives a transition from \( \langle 111 \rangle \) to \( \langle 110 \rangle \) oriented growth. We propose a simple nucleation model that explains this observation. This knowledge is then applied to rationally fabricate nanowire superstructures through the judicious incorporation of small quantities of Ge during growth. This work demonstrates the important role that surface chemistry plays in the growth of semiconductor nanowires, and the extensive use of hydride chemistries for most group IV and III-V semiconductor nanowire syntheses suggests significant implications for many materials systems.

11:00am NS-TuM10 Inhomogeneous Longitudinal and Radial Dopant Distribution Measurements in Si Nanowires Using Scanning Auger, J.S. Hammond, D.F. Paul, Physical Electronics, U. Given, Northwestern University

The incorporation of electrically active dopants into nanowires (NW) is essential to the development of semiconductor NWs based electronic devices. The ability to engineer the electrical properties of nanowires grown by the vapor liquid solid (VLS) process is currently limited by our incomplete understanding of the doping mechanism. Recently, several studies have shown evidence of inhomogeneous radial dopant distributions in Si NWs and the resulting effects on their electrical properties [1-2]. However, the longitudinal dopant profile has not been addressed to the same extent. Studies employing both indirect and direct measurement techniques (such as scanning photocurrent microscopy, Kelvin probe force microscopy and atom probe tomography) have addressed variations in longitudinal dopant profiles in Si NWs and related them to radial dopant variations induced by the growth process [3-4]. There have not yet been direct measurements of dopant concentrations along VLS grown nanowires.

We have measured the longitudinal and radial doping profiles of phosphorus-doped Si NWs using scanning Auger. We have found order of magnitude enhancements in the dopant concentration toward the NW's base as expected from previous indirect measurements. Importantly, the physical dopant profile is not identical to the active dopant profile, as shown by comparison with scanning photocurrent microscopy and Kelvin probe force microscopy measurements. The resolution and sensitivity of scanning Auger as an analytical tool for dopant concentration measurements will be compared to the other techniques to indicate unique capabilities that can advance our understanding of nanowire doping.


11:20am NS-TuM11 Crystal Structure Engineering of Ge Nanowires, I.R. Musin, M.A. Filler, Georgia Institute of Technology

Semiconductor nanowire engineering provides a promising route to achieve next generation energy conversion, photonic, and electronic materials. In order to enable the appropriate function for a particular application, control of nanowire crystal structure (e.g. lattice, orientation, faceting) is critical. Unfortunately, this remains a challenging task with bottom-up nanowire structure syntheses. To this end, we rationally control Ge nanowire crystal structure for the first time via the addition of bifunctional alkylgermananes, which adjust the interface energetics near the three-phase line. More specifically, Ge nanowires are grown using the vapor-liquid-solid (VLS) technique with germane combined with methyl-, ethyl- or tertbutyl-germane. Scanning electron microscopy (SEM) and high resolution transmission electron microscopy (TEM) reveal that nanowires transition to a new growth direction upon addition of an alkylgermane, but remain single crystalline throughout. For the case of methyl-germane, nanowires transition from the \( \langle 111 \rangle \) to \( \langle 110 \rangle \) crystal growth direction. A significant reduction in tapering is also observed in all cases. Infrared spectroscopy (IR) shows that nanowire sidewalls are alkyl terminated and X-ray photoelectron spectroscopy (XPS) indicates this termination reduces the rate of oxidation. The impact of alkylgermane identity and surface coverage on crystal growth direction will be discussed in detail. The control of interface chemistry demonstrated by this work provides an important new handle for controlling nanowire structure and properties. Furthermore, the ability to effectively passivate nanowire sidewalls during growth is expected to enable more robust doping profiles by only permitting precursor incorporation through the catalyst tip.

8:00am PS+MN+TF-TuM1 Scallop Free TSV Etching Method for 3D LSIs, Y. Morikawa, T. Maruyama, T. Sakuishi, S. Toyoda, K. Sune, ULVAC, Inc., Japan

INVITED Thru silicon via (TSV) etching process for deep and high-aspect ratio structure has been studied thoroughly for applications such as MEMS and CMOS devices. Recently, TSV used in 3D-LSI devices for logic devices may be a few microns in diameter and about 50 um deep. On the other hand, TSVs used in stacked memory devices, the via diameter and depth would be several tens of microns. Therefore, etching of dielectrics and electronic materials is very important for realizing these applications. In this study, a large size etching in a high-pressure process was focused by using very high frequency capacitive coupled plasma (VHF -- CCP) with an ultra self-confined system. This plasma system is simple parallel plate CCP about 100Pa or more process. High-pressure process was carried out on the plasma confined, because mean free path is very short. And, ion energy distribution (IED) is also controllable by high-pressure process with VHF bias. The bimodal IED changes under high-pressure. The peak of high-energy side is reduced, and a charge exchange peak appears. It is considered that the charge exchange is important to anisotropic Si etching of large size TSV with VHF bias.

And next, the high-density and small size of TSV below 10um diameter is indispensable to the utilization and improvement in high performance of 3D-LSI. We have developed a new etching system for TSV etching process for small size and high aspect ratio via. This system is a planar type metal neutral loop discharge (NLD) plasma. For high rate silicon etching, it is very important to understand not only the high density of the ICP plasma generation but also the high density of fluorine atoms. In this study, a novel RF antenna ‘Multi Stacked rf Antenna’ has been developed for highly accurate and high rate etching process. This antenna consists of multistage neutral loop discharge (NLD) plasma. Finally, the diameter of 2um was attained by the anisotropic etching.
etching of 5 um/min, and the aspect ratio is above 10 using the planer NLD etcher. VHF CCP and planer NLD etching processes are non-cycle etch methods, and these processes were demonstrated above smooth sidewall TSV formation.

8:40am PS+MN+TF-TuM3 Deep Silicon Etching of 0.8 µm to Hundreds of Microns Wide Trenches with the STIGer Process. T. Tillocher, W. Kafrouni, GREMI, France, J. Ladroue, STMicroelectronics - GREMI, France, P. Lefauxeches, GREMI, France, M. Bobbischel, STMicroelectronics, France, P. Rautson, R. Dussart, GREMI, France

The STIGer process is designed to achieve high aspect ratio features in silicon. Like the Bosch process, passivation steps (SiF4/O2 plasmas) and etching steps are cycled to get vertical structures. The etching steps can be purely isotropic (SF6 plasmas) or anisotropic (SF6/O2 plasmas). It is required to cool the silicon substrate with liquid nitrogen to form a SOxFy passivation layer. It desorbs and disappears when the substrate is heated back to temperatures above the liquid nitrogen. Thus, thermal optimization has been carried out in order to optimize the STIGer process. Next to the STIGer process, the formation of microstructures nor the chamber walls after each process run. Then, the robustness of the STIGer process is enhanced in comparison with standard cryo-etching: the profiles are less sensitive to temperature or flow rate variations. But, like in Bosch etching, a scalloping is present on the sidewalls.

Submicron trenches having critical aperture of about 0.8 µm can be etched with high aspect ratios (> 40). In these cases, the average etch rate is around 1.8 µm/min. These features exhibit both undercut and a special defect, which is called “extended scalloping”. This defect is composed of anisotropic cavities developed on the feature sidewalls, just below the mask. It originates from ions scattered at the feature entrance that hit the top profile and remove locally the passivation layer. This defect is observed only for high aspect ratios (typically above 10). Thus, we will also investigate the role of trench critical dimension (from 0.8 µm to 100 µm). A mechanism explaining the formation of the extended scalloping will be proposed.

We have studied the influence of both the duty cycle (etch/etch+passivation)) and the chamber pressure on the profiles and the extended scalloping. Basically, when the duty cycle increases, etching dominates passivation, which leads to higher defects. Pressure is a way to tune the slope of the sidewalls. Actually, decreasing the chamber pressure helps to shift from positively tapered features to more vertical profiles, and even negative slopes, hence with dovetailed shape.

This will be correlated with plasma analysis by means of mass spectrometry and optical emission spectroscopy. Actually, it is relevant to investigate how changes in the plasma chemistry can modify the trench profiles.

These trends have been used to optimize two methods that can help to reduce the extended scalloping. The first consists in adding a low oxygen flow in the etch cycle; favouring a low additional passivation. The second technique consists in gradually increasing the SF6 flow from a low value to the nominal value. Consequently, the process starts with a low etch rate and a more efficient passivation, which helps to limit the extended scalloping.


With the current advent of 3D integration for advanced interconnect and packaging applications, there has been a renewed focus on deep silicon etch technology to satisfy the need for Through Silicon Via (TSV) patterning. The most common etch method used to fabricate said devices is a time-multiplexed (BoschTM) process, based on years of maturity in the MEMS field.[i] However, issues such as scalloping, mask undercut and limited etch rates are becoming more pronounced as feature sizes scale to meet the ITRS roadmap requirements. This has prompted efforts to attempt to either develop a more conventional etch process[ii][iii][iv][v] or to modify the Bosch process to circumvent these issues.[vi][vii][viii] In this work, we explore a novel polymerizing feedgas chemistry for the deposition step of the Bosch process to improve mask undercut while simultaneously increasing TSV etch rate. Initial results indicate a 5x larger deposition rate as compared to C4F8 (under nominal conditions) and under optimized conditions, enables a 50% decrease in undercut along with 10% increase in TSV etch rate. Optical emission spectra also differ substantially between the two feed gases, indicating different dissociation pathways and radical densities. Further results and a detailed characterization of the deposition properties of the novel chemistry will also be discussed leading to a proposed mechanism for the profile improvements as compared to C4F8.[v] B. Wu, A. Kumar and S. Narayu, J. of Applied Physics 108, 051101 (2010) [vii] I. Sakai, N. Sakurai and T. Ohiwa, J. Vac. Sci. Technol. A 29(2), Mar/Apr 2011 [iii] N. Ranganathan et al, Proceedings of the Electronics Components and Technology Conference, 2005 [iv] H. Rhee et al, J. Vac. Sci. Technol. B 27(1), Jan/Feb 2009 [v] S.-B. Jo et al, J. Vac. Sci. Technol. A 23(4), Jul/Aug 2005

9:20am PS+MN+TF-TuM5 Wafer Scale Hermetic Packaging of MEMS. C.S. Gudeman, IMT

The explosion of MEMS in automotive and cell phone markets has been enabled by low cost wafer level packaging (WLP) technology that provides a robust and hermetic enclosure for an otherwise delicate device. The more obvious advantage of WLP is greatly improved reliability, because the device is protected from organic and particulate contaminants while in the hands of the end user. However, other advantages related to package design and manufacture are enabled by WLP during the manufacturing process, which often produces the highest levels of stress that a MEMS device experiences. These processes include wafer grinding, wafer dicing, and chip solder re-flow attachment to circuit boards and other chips. Firstly in this talk, wafer level packaging technologies will be outlined, focusing on the truly hermetic methods -- alloy, glass frit, Au-Au thermo-compression, anodic, and fusion bonding. Secondly the integration of Through Silicon Via (TSV) with WLP will be discussed. Finally the performance of these technologies will be compared from a manufacturing perspective, including yield and thermal budget.


INVITED

With advances in non-volatile memory, the major challenge confronting plasma etch is the introduction of new materials while simultaneously shrinking critical dimensions. This talk will address key development aspects including profile control, feature level uniformity, and plasma microdamage. Plasma microdamage is not the traditional, charge/voltage/current based impact to, for example, gate oxides. Rather, it is the changes to the atomic scale morphology in the sidewall or landing film of a plasma etch process. This results in a disruption of local stoichiometry, film defects, and other issues that impact device performance. The etch process and hardware changes to address this will be presented.

11:40am PS+MN+TF-TuM12 Mechanisms of Selective Etching for Magnetic Materials: Ni, Co and Ta Etching by Carbon Monoxide/Methyl Alcohol Based Plasmas. K. Karahashi, T. Ito, S. Hamaguchi, Osaka University, Japan

Dry etching of magnetic thin films is a crucial step in micro fabrication of magnetic random access memories (MRAMs) and write/read heads for magnetic data storages. Argon (Ar) ion milling seems to be almost the only etching technique available in the current manufacturing processes. However Ar ion milling is incapable of achieving anisotropic and selective etching of magnetic films (Ni, Co etc.) over hardmasks (Ta etc.) and therefore highly selective reactive ion etching (RIE) of magnetic thin films is a highly sought-after technology. RIE processes based on CO/NH3 or CH3OH is a candidate for selective etching of magnetic thin films. In this study, we have examined etching processes of Ni, Co and Ta thin films by energetic CO+ or O+ ions, which are considered to be major etchants of CO/NH3 or CH3OH plasmas. We have determined the etching yields and analyzed surface reactions, using a mass-selected ion beam system. The ion beam system is designed to perform metastable gas reactions onto a sample surface in ultra-high vacuum conditions. The reaction chamber, where the sample is placed, is equipped with an X-ray photoelectron spectroscopy (XPS) for in-situ chemical analyses of irradiated surfaces. The ion beam energy used in this study is in the range of 150-1600 eV. The etching yields are determined from measured depth profiles of irradiated surfaces and ion fluxes. The etching yields of Ni and Co by CO+ ions are higher than that by O+ ions but lower than the yields of possible physical sputtering, which are estimated from interpolation of sputtering yield data of inert atom ions (He+, Ne+, Ar+, Kr+ etc.). From XPS analysis for O2+ irradiated Ni and Co surfaces, oxidation is found to occur under O2 irradiation, which suggests that the oxide layer hinders sputtering by ion bombardment. It is found that little oxidation occurs on Ni or Co surfaces under CO+ irradiation and etching by CO+ ion bombardment proceeds. On a Ta surface, on the other hand, in the both cases of O2+ and CO+ irradiations, oxidation occurs and its etching yield is far smaller than the yield of its possible physical sputtering. Therefore we have found that high selectivity of Ni and Co etching against hard masks (Ta, TaN) arises from the prevention of sputtering by mask oxidation. Etching characteristics by OH+ irradiation were also studied in a similar manner. This work was...
supported by the Semiconductor Technology Academic Research Center (STARC).

**Plasma Science and Technology Division**

*Room: 201 - Session PS-TuM*

**Advanced BEOL / Interconnect Etching I**

**Moderator:** A. Balakrishna, Applied Materials, Inc.

8:00am **PS-TuM1** Narrow Pitch Dual Damascene Patterning using EUV Lithography in Association with a Spin-On Trilayer Resist System, F. Lazzarino, V. Truffert, B. Vereecke, S. Demaynyc, IMEC, Belgium

Extreme ultraviolet lithography (EUVL) is one of the leading candidates for the 22-nm node device manufacturing. However, a major issue is the necessity to use thin photoresist (between 55-nm and 80-nm after development) because of resolution requirement and limited depth of focus. In addition, its low etch resistance does not allow high aspect ratio pattern transfer. In this context, a new hardmask strategy called spin-on trilayer resist system has been considered.

In this work, we mainly focus on the etch patterning capability of narrow pitch dual damascene structures by using EUVL combined with a spin-on trilayer resist system. The latter consists of three layers. The photoresist on top is used to pattern a thin spin-on glass layer which is then used to pattern a thick spin-on carbon layer (SOC). The SOC has two functions. It is used for its good gap-filling capability to avoid patterning over nonplanar surfaces but it also acts as a hardmask to pattern the dielectric stack (150-nm of oxide on top of 15-nm of SiCO and 5-nm of SiCN). Regarding the dual damascene architecture, two different approaches have been considered: the via-first and the trench-first. Despite few challenges such as the well-known fencing issue, the via-first approach has been chosen as it is less sensitive to misalignment. In this scheme, two lithography and etch steps are needed, first to form the via then to pattern the trench and etch the barrier layer. In this study, we compared the via opening by using the standard PECVD carbon layer and by using the SOC layer. As expected, the PECVD carbon layer has a better process window compared to the SOC layer. The selectivity is greater and allows many chemistry variations to fine tune for instance the profile. To get similar process window with the trilayer resist system, we introduced C4F8 and CO to substitute C4F6 and O2 in the original chemistry. This modification clearly improved the process by having a better control on the passivation layer formation. Regarding the trench opening, we observed a significant line wiggling of the SOC hardmask for 50-nm half-pitch structures and beyond. We characterized this instability thanks to stress measurements and we kept it under control by changing three different process parameters: the bottom electrode temperature, the baking conditions after coating and the film thickness. Each of them has an impact but the best result came from combining all three together.

To conclude, we demonstrated that narrow pitch dual damascene structures can be obtained by using EUVL in association with a spin-on trilayer resist system. The structures formed in this way shows good electrical characteristics.


Cu resistivity increase at smaller geometries and the use of lower k dielectrics forced several changes on BEOL integration. Susceptibility of ULK material to strip damage drove the adoption of TiN hard mask integration starting with a few players at 65nm node to full adoption at 32nm for advance logic.

TiN hard mask demonstrated good selectivity during dielectric etch but concerns arise as we move to smaller pitch: 1- TiFx residues may compromise metallization of small features and 2- Residual compressive stress in the TiN may induce line wiggling. This last concern is the focus of this work.

We develop a simple analytical model based on energy balance to predict wiggling onset based on geometrical and mechanical properties of the materials. We showed that the wiggling onset aspect ratio reduces with the pitch. We also showed that even when at the end of the process the no-wiggling condition is satisfied, wiggling might have occurred during etch causing feature distortions.

8:40am **PS-TuM3** Surface Reaction Control for BEOL Application. M. Fukasawa, T. Tsutsumi, Sony Corporation, Japan

Increasingly there are more challenges of controlling the plasma processes for BEOL integration, which include the Cu/low-k interconnects, contact hole etching, etc. The presentation will give an overview of the surface reaction control during plasma processes. The main focus is damage reduction and suppression of process fluctuations.

Reducing the damage to low-k dielectrics caused by plasma exposure is one of the key issues. Ox-based plasma has been widely used for ashing the photoresist on low-k SiOCH. H2-based plasma is employed for the etching of organic low-k film as well as the ashing of photoresist. The thickness of damage generation is classified by ions, radicals, and UV/VUV radiation. It was found that not only the ions but also the synergy of radicals and UV/VUV radiation cause a significant amount of damage in the SiOCH, measured by using the pallet for plasma evaluation (PAPE). Hence, precise control of incident ions, radicals, and UV/VUV radiation is required for controlling the surface reactions.

In addition to the precise control of incident species, the optimization of subsequent processes (wet clean, dry clean, etc.) is very important to obtain sufficient electrical yields and reliabilities. The remaining damaged layer after wet treatment degrades the Cu and SiOCH surface and corrupts the interface between the Cu and barrier metal. For instance, the desorbed H2O from the SiOCH damaged layer causes the oxidation of barrier metal, which results in a shorter EM lifetime. The surface modification during SiCN and SiN etching by CHF3-based plasma and its impact on the electrical yields will also be discussed.

The suppression of fluctuation is also required in the advanced interconnects. Although statistical prediction is one of the approaches to realize stable processes, there is a limitation of prediction accuracy for purely statistical predictions due to the lack of physical models. Thus, the combination of statistical and physical models for highly accurate prediction has become an emerging trend in mass production. We will demonstrate a novel statistical and etch rate prediction model by considering the fluctuation caused by the plasma-wall interactions.

The developed prediction model is one of the approaches to realize stable processes. Changes in the spatial distribution of reactive species in the etching chamber are, however, very difficult to detect in current mass production tools. Thus, greater progress of the in-situ monitoring tools and prediction methods based on the physical model (simulation) are strongly required in the near future.

2. M. Fukasawa et al., JAP 48, 08HCC01 (2009).


Trench First metal Hard Mask (TFmHM) integration scheme for BEOL has gained traction over recent years because it can mitigate many challenges that are inherent with Via First Trench Last (VFTL) scheme. This integration scheme was more recently shown to enable Self-Aligned Via (SAV) patterning. The SAV patterning implies a pretty drastic change of the via process as, on top of the usual via requirements, the via patterning process needs to be selective to the metal HM. Key process parameters including temperature, gas chemistry, power and pressure were investigated. The required selectivity of the materials and tight CD control capability necessitate temperature and pressure adjustments. Each of them has an impact but the best result came from combining all three together.

The simultaneous control of via, trench and chamfer profiles (i.e. Critical Dimensions, depth, taper profile, etc), implies the need for better control of the metal HM selectivity during both SAV and trench patterning and the need for flexible adjustment of the ion energy and control of the flux of ions and active neutrals. Low-k material damage control is always pertinent in the RIE process as dimensions get smaller. As the direct result of such tight process guidelines, the hardware challenges arise and new dimension of process controls are needed.

In this paper, the RIE efforts on process controls of the via and trench profiles, the metal HM selectivity, associated hardware solutions and future process flow options under TFmHM scheme will be discussed.

This work was performed by the Research and Development team at TEL Technology Center America in joint development with IBM Research Alliance Teams in Albany, NY 12203. This work has also been supported by the independent Bulk CMOS and SOI technology development projects at the IBM Microelectronics Div. Semiconductor Research & Development Center, Hopewell Junction, NY 12533.
Matsuura, during plasma etch. We also determined that hydrogen addition to the fluorocarbon (FC) layers generated on oxide and organic mask materials Ar/C4F6/O2 in terms of oxide-to-organic mask etch selectivity and line- Ar/C5HF7/O2 chemistry was shown to have significant advantages over

originated from the different thickness and composition of steady-state evaluated for high aspect ratio dielectric etch. Plasma etching with a high etch selectivity hydrogen-containing fluorocarbon gas, C5HF7, was Corpo rating...
presence of cobalt in the framework affects the reducibility of the cobalt species. The data also suggests mixed phases of cobalt metal and cobalt monocarbide in the reduced and post FTS samples. The as prepared samples show only the Co3O4 phase, while the post FT and post TPR show both CoO and Co metal. The good agreement between XAS and XRD measurements demonstrates the use of these techniques to quantitatively determine the cobalt phase composition.

8:20am SS1-TuM2 Measurement of the C-Pt Bond Energy for Adsorbed Methyl on Pt(111) by Methyl Iodide’s Dissociative Adsorption Energy. E.M. Karp, T.L. Silbaugh, C.T. Campbell, University of Washington

Knowing the bond energies of molecular fragments adsorbed to surfaces is an important ingredient in the fundamental understanding needed for rational design of heterogeneous catalysts. This study utilizes the ultrahigh vacuum technique of single crystal adsorption calorimetry to study the energetics of adsorbed methyl (CH₃ad) on Pt(111), using measurements of the dissociative adsorption energy of methyl iodide (CH₃I) on Pt(111). At temperatures above 250 K, adsorbed CH₃I on Pt(111) is known to undergo C-I bond scission to form CH₃ad and adsorbed iodine (Iad). At 270 K, the rate of C-I scission is fast relative to the 100 ms time of calorimetric measurement, so that it provides the enthalpy for the net, two-step dissociative adsorption reaction:

CH₃I(ad) → CH₃(ad) + I(ad) (1)

The integral heat of Reaction (1) is -224 kJ/mol at a coverage of 0.05 ML. Through the use of a simple thermodynamic cycle and other known adsorption energies, this provides a heat of formation of CH₃ad of -63 kJ/mol and a Pt—CH₃ bond energy of 209 kJ/mol.

Mass spectrometer measurements taken during experiments at 300 K and 320 K indicate that methane is produced. Previous studies indicate that at these temperatures, adsorbed methyl both decomposes to form adsorbed methylene (CH₂ad) and hydrogen (H₂ad) and reacts with H₂ to form methane according to the reactions:

CH₃ad → CH₂ad + H₂ (2),
CH₂ad + H₂ → CH₄ (3)

where CH₄ is gas phase methane. The microcalorimetric measurements performed at 300 K and 320 K, along with information on the rates of Reactions (2) and (3) available in the literature, allowed for the decoupling of Reactions (1)-(3) and an analysis of the energetics of all three reactions as a function of surface coverage, and the heat of formation of CH₂ad,

8:40am SS1-TuM3 A DFT Study of Methanol Reaction Pathways on the Au₅₀/TiO₂(110) Surface. S. Hong, University of Central Florida, D.A. Chen, University of South Carolina, T.S. Rahman, University of Central Florida

We have performed density functional theory calculations to understand the reaction pathway selection for methanol decomposition on clean and Au₅₀-doped TiO₂(110) surface. We find that when methanol adsorbs on the clean and reduced TiO₂(110) surface, it decomposes spontaneously into methoxy, which adsorbs on the O-vac site and desorbs as methylene leaving behind O-br on the TiO₂(110) surface. On the stoichiometric TiO₂(110) surface, we find that while methanol decomposition has a small activation energy of 0.2 eV, methoxy decomposition (i.e. formaldehyde formation) is not possible because of high activation energy of 3 eV. Nevertheless, once the surface is oxidized formaldehyde formation becomes spontaneous, through C-H bond scission by surface oxygen species. These results confirm the experimental results of methyl formation on the reduced and formaldehyde formation on the oxidized TiO₂(110) surface. Turning to the case of partially reduced Au₅₀/TiO₂(110) surface, our DFT calculations show that the adsorption of methanol and its intermediate methoxy does not occur on gold sites but on TiO₂ sites Methoxy, in fact, forms at the Au-titania interface as a result of O-H bond scission by lattice oxygen (O-br), and its optimized titled structure makes C-H bond scission -- through H abstraction by lattice oxygen near the interface site -- much easier than that for a vertical, non-tilted structure of methoxy in a non-interfacial site. Moreover, depending on the reduction level of the TiO₂(110) surface, the adsorption site preference for methanol and methoxy species on the Au₅₀/TiO₂(110) surface flips between O-vac and interfacial Ti-cus > stoichiometric Ti-cus site. This hierarchy implies that the major product will switch from methanol on the fully-reduced surface, to formaldehyde on the partially-reduced or oxidized Au₅₀/TiO₂(110) surface. This conclusion is in full agreement with experiment.

This work is supported in part by US-DOE under Grant No. DEFG02-07ER15842.

9:00am SS1-TuM4 Turning Aluminum into a Noble-metal like Catalyst for Low Temperature Molecular Hydrogen Activation. J. Chopra, University of Texas at Dallas, S. Chaudhuri, Washington State University, J.F. Veyan, Y.J. Chabal, University of Texas at Dallas

There has been an ongoing quest to find cheaper hydrogen activation routes based on aluminum. Despite theoretical suggestions[2] it has been difficult to obtain unambiguous experimental evidence for such catalytic activity towards hydrogen activation.

We demonstrate here that aluminum doped with very small amounts of Ti can activate molecular hydrogen at temperatures as low as 90K. The method is based on the ability to introduce a high flux of molecular hydrogen seeded with a guest molecule to probe the catalytic activity and H₂ dissociation. Once dissociated, hydrogen forms a complex with adsorbed CO (CO-H), characterized by a substantially and uniquely blue-shifted CO internal frequency. This complex is metastable, and is removed at a temperature (115K). We find that CO does not adsorb even weakly on H-covered Au, and use this as a clear demonstration that dissociated hydrogen diffuses away from the catalytic site onto Al sites (i.e. spills over). We use this new method to determine the dependence of the catalytic activity of aluminum surfaces on Ti coverage. Finally we show that the complex with activated hydrogen leads to further reactions at remarkably low temperatures (115K), such as formation of formyl (HCO), formaldehyde (HCHO) or methanol. These results provide the first direct evidence that Ti-doped Al can perform the quintessential first step of molecular hydrogen activation under nearly barrier-less conditions, thereby challenging the monopoly of noble metals in hydrogen activation.3

References:

9:20am SS1-TuM5 On the Role of Hydrogen in Heterogeneously Catalyzed Reactions, M. Mavrikakis, University of Wisconsin Madison

INVIITED

Hydrogen is a frequent participant in several heterogeneously catalyzed reactions, including Fischer-Tropsch Synthesis (FTS) of fuels, ammonia synthesis, oxygen reduction reaction (ORR), NO reduction, preferential oxidation of CO in the presence of CO (PROX), etc. Having analyzed the detailed aspects of the reaction mechanism for a number of these reactions on various transition metal and alloy surfaces using first-principles methods, some common principles governing the role of hydrogen in a wide range of catalytic transformations begin to emerge. In this presentation, we will discuss these common mechanistic principles by examples, including FTS[1], NO reduction, ORR[2], PROX[3], through an analysis of the energetics of alternative elementary reaction steps and the resulting potential energy diagrams. Connections to observations from experimental studies provide an invaluable perspective for the evaluation of our theoretical assessments.

References

10:40am SS1-TuM9 Prediction of Surface Ensembles in Au-based Bimetallic Alloys using Combined DFT and Monte Carlo Simulations. J.A. Stephens, H.C. Ham, G.S. Hwang, University of Texas at Austin

Bimetallic materials have shown great promise for the development of superior catalysts. The recent surge of new interest in catalysis by gold has led researchers to investigate the effects of adding gold to other metals. While mechanisms underlying the alloying effect are still not understood in detail, recent evidence suggests that the enhanced reactivity of bimetallic catalysts can be attributed to a combination of metal-metal interactions.
Technology Laboratory

the CO$_2$ at the oxygen vacancy defect is in a tilted configuration and the
On the reduced surfaces (x = 0.1 – 0.4), acetaldehyde interacts more strongly
with its C-C bond perpendicular to the surface plane and the acyl hydrogen
adsorption of CO$_2$ takes place at the oxygen vacancy defect. At higher
dispersion corrected density functional theory (DFT). At low coverage the
coverage the CO$_2$ starts to adsorb on the Ti rows. The DFT results show that the
CO$_2$ at the oxygen vacancy defect is in a tilted configuration and the
molecular axis is perpendicular to the bridging oxygen row. On the Ti row,
the DFT results confirm that the CO$_2$ adsorbs in a flat configuration with its
axis parallel to the bridging oxygen row. Electron injection from the STM
tip into the CO$_2$ at the oxygen vacancy defect induces the dissociation of
CO$_2$. The oxygen vacancy defect is found to be healed by the O atom
released during the CO$_2$ dissociation process. Statistical analysis shows that
the dissociation of CO$_2$ is a one-electron driven process with a threshold
voltage of 1.4 eV above the conduction band minimum of TiO$_2$.
The formation of a transient negative ion by the injected electron is considered to be
the key process in the CO$_2$ dissociation.

11:00am SS1-TuM10 Adsorption and Electron-induced Dissociation of CO$_2$ on TiO$_2$(110). J. Lee, X. Deng, D. Sorensen, National Energy Technology Laboratory

Adsorption and electron-induced reaction of CO$_2$ on the TiO$_2$(110) surface has been investigated using scanning tunneling microscopy (STM) and
dispersion corrected density functional theory (DFT). At low coverage the
adsorption of CO$_2$ takes place at the oxygen vacancy defect. At higher
coverage the CO$_2$ starts to adsorb on the Ti rows. The DFT results show that the
CO$_2$ at the oxygen vacancy defect is in a tilted configuration and the
molecular axis is perpendicular to the bridging oxygen row. Electron injection from the STM
tip into the CO$_2$ at the oxygen vacancy defect induces the dissociation of
CO$_2$. The oxygen vacancy defect is found to be healed by the O atom
released during the CO$_2$ dissociation process. Statistical analysis shows that
the dissociation of CO$_2$ is a one-electron driven process with a threshold
voltage of 1.4 eV above the conduction band minimum of TiO$_2$.
The formation of a transient negative ion by the injected electron is considered to be
the key process in the CO$_2$ dissociation.

11:20am SS1-TuM11 Adsorption and Dissociation of Propionate-3-diol Molecules on Reduced TiO$_2$(110) Surface- A Scanning Tunneling Microscopy Study. D. Acharya, X. Lin, B.D. Kay, Z. Dohnálek, Z. Zhang, Pacific Northwest National Laboratory

Rutile TiO$_2$(110) surface is one of the most studied model systems for the
fundamental investigations of a variety of processes on metal oxide
surfaces, including heterogeneous catalysis, greenhouse gas reforming,
solar cells, photodecomposition of organic pollutants, and photoinduced
water splitting. It is widely accepted that the surface chemistry of the oxide
surfaces are mainly influenced by the defects sites, particularly oxygen
vacancies. Here we study the adsorption of propionate-3-diol molecules on
partially reduced rutile TiO$_2$(110) surface using variable temperature
scanning tunneling microscopy (STM). STM images obtained before and after
the deposit of propionate-3-diol molecules at room temperature shows
that the molecule preferentially binds at the bridging oxygen vacancies via
bond scission of one of the OH groups. The hindered rotation of the 3-
hydroxypropoxide species was seen at room temperature, while the
diffusion and dissociation of the species were observed at elevated
temperatures. The experiment was carried out at the Environmental
Molecular Science Laboratory, a National Scientific user facility supported by the U.S. Department of Energy, Office of Biological and Environmental
Research at Pacific Northwest National Laboratory.

11:40am SS1-TuM12 Oxygen-Vacancy Assisted Formation of Enolate Species on Reduced CeO$_2$(111) Surfaces. F.C. Calaza, Y. Xu, D.R. Mullins, S.H. Overbury, Oak Ridge National Laboratory

Enolate species are key intermediates proposed in a number of important
organic reactions heterogeneously catalyzed by metals and metal oxides, but
enolate has been difficult to identify on active catalytic surfaces due to
difficulties of isolating it in the keto-enol equilibrium.

Reflection absorption infrared spectroscopy (RAIRS) was coupled with
density functional theory (DFT) to study the adsorption of acetaldehyde, a simple C$_2$H$_4$ aldehyde, on CeO$_2$(111) surfaces of different extent of oxidation(where x = 0 – 0.5). It is found experimentally that the molecule
adsorbs weakly on the fully oxidized surface (x=0) at low temperatures and desorbs without further reaction near 215 K. The molecule bonds to c.c.s.
Ce$^{IV}$ cations through the oxygen lone pair electrons in the carbonyl group
with its C-C bond perpendicular to the surface plane and the acyl hydrogen
tilted slightly towards one of the lattice oxygen anions of the first layer.

On the reduced surfaces (x=0.1 – 0.4), acetaldehyde interacts more strongly
with the surface upon adsorption at low temperatures by losing its carboxyl
bond character and adsorbing as 1,1-dioxymethylene and forming dimers and
polymers. Heating the surface to 400 K leads to desorption of some amount of
these strongly adsorbed species as acetaldehyde and the appearance of
hydroxypropoxide species, and yet a different organic species.

The identities and structures of the different intermediates on the CeO$_2$ and
CeO$_2-x$ surfaces have been determined by their characteristic signatures in
RAIRS and by DFT calculations. Our observations for the CeO$_2-x$ surfaces are consistent with the vacancy-promoted dehydrogenation in the original
methyl position of acetaldehyde and the formation of enolate (CH$_3$CHO-
Ce$^{IV}$).

Experiments with isotopically labeled acetaldehyde have verified the
vibrational assignments for the enolate species and are in excellent
agreement with DFT results. The assignment of the enolate species is
furthermore consistent with C 1s XPS and C k-edge NEXAFS results.

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Biosciences, Office of Basic Energy Sciences, US Department of
Energy. Use of the National Synchrotron Light Source, Brookhaven National Laboratory, was supported by the US Department of Energy, Office of Science, Office of Basic Energy Sciences.

Surface Science Division

Room: 110 - Session SS2-TuM

Self Assembled Monolayers and Networks

Moderator: J.E. Reutt-Robey, University of Maryland

8:20am SS2-TuM2 In Situ UHV Growth and XPS/NEXAFS Characterization of Aromatic Self-Assembled Monolayers on Gold Substrates. A. Nefedor, Karlsruhe Institute of Technology, Germany, H. Mazik, University of Bielefeld, Germany, M. Naboka, Karlsruhe Institute of Technology, Germany, A. Turchanin, A. Götzlühäuser, University of Bielefeld, Germany, C. Will, Karlsruhe Institute of Technology, Germany

Aromatic self-assembled monolayers (SAMs) on gold show high potential for applications in nanobiotechnology [1]. Although the SAMs are typically
pre pared by immersing gold substrates in a solvent, their growth by vapor
deposition in vacuum has various advantages both for the technological
applications and fundamental studies [2, 3]. Thus, it is possible to
characterize in detail the monolayer growth by various UHV compatible
surface science techniques such as, e.g., electron spectroscopy. Here, we
employ a new establishment of the HESG beamline at BESSY II to study the
in situ growth of 4’-nitro-1,1’-biphenyl-4-thiol (NBPT) SAMs on gold/mica
substrates by X-ray photoelectron spectroscopy (XPS) and near edge X-ray
absorption fine structure spectroscopy (NEXAFS). XPS Cls, N1s, O1s and
S2p spectra as well as NEXAFS C K-, N K- and O K-absorption edge spectra
were analyzed. These results demonstrate the formation of NBPT SAMs by
vapor deposition in UHV with a similar packing density and molecular
orientation as usually obtained in solvents. Moreover, we analyze the
electron-radiation-induced modification of NBPT SAMs, that is widely
employed in chemical nanolithography [4]. By varying the dose of electron
irradiation we study two effects: (i) mechanisms of the electron-induced
cross-linking [5] and (ii) conversion of the nitro groups into amino groups
[6]. We analyze these transformations and compare the experimental
NEXAFS data with calculations made by the StoBe software package.


8:40am SS2-TuM3 Structure, Bonding and Electric Properties of Self-Assembled 2D Organic Nanostuctures at Surfaces: Negatively Charged TCNQ Networks and Other Systems. S.L. Zalt, Indiana University

INVITED

Interfaces between organic materials and inorganic supports are critical for the
design and function of new organic-based technologies (e.g., OLEDs, organic
photovoltaics, and molecular electronics) as well as novel routes to
chemical sensors and catalysts. There are vast opportunities for designing
structure-function relationships in these systems due to the immense
library of organic compounds and metal-organic chemistries available. Molecular
self-assembly at surfaces by covalent, metal-organic, ionic, and weaker
interactions are active fields of research, but much remains to be determined with regard to the complex interplay of intermolecular and adsorbate-substrate interactions and how these impact structure and function. TCNQ undergoes a charge transfer from a Cu surface to adopt a bond conformation to the surface that enables stable adsorption and the self-assembly of highly ordered 2D structures via surface-mediated attractive interactions. These structures are stable at room temperature. Addition of Mn triggers a structural transformation to a highly-ordered porous network with Mn centers in a high spin state. These studies have allowed new insights into organic/metal interfaces by collaborative work involving high-resolution scanning tunneling microscopy, photoelectron spectroscopy methods, synchrotron measurements, other UHV surface experiments, and density functional theory calculations. Recent progress on other 2D and multilayer organic systems in our group lends further evidence to the significance of directed intermolecular interactions in such systems, the problems these interactions may pose, and solutions for balancing them at a desired level. We are making progress towards tailored chemical function by rational design of molecular architectures at surfaces and tuning such function through supramolecular design strategies.

9:20am SS2-TuM5 Ultrafast Self-Assembly of 1-Adamantanethiol and \( p \)-Terphenylthiol on Au(111)-Surface. V.V. Korolkov, S.A. Allen, C.J. Roberts, S.B. Tendler. The University of Birmingham, UK

Study of organic thiols adsorption on noble metals continues to be a subject of many research papers within the last three decades. Such steady interest arises mainly from the possibility to design metal surfaces with pre-defined chemical and physical properties that makes them suitable for a wide range of theoretical and applied studies.

Here we propose and investigate an idea that thiols with rigid or spherical hydrocarbon moieties should self-assemble on Au-surface much quicker than those with long-chain moieties. To support this idea we have investigated adsorption dynamics and monolayer structure for 1-adamantanethiol (AdSH) and \( [1,1':4',1''\text{-terphenyl}]-4\text{-thiol} \) (TPT) molecules self-assembled on Au(111)-surface at 393K. We have demonstrated that both thiols form defect free and uniform SAMs almost instantly at elevated temperature. It takes \( \sim 1 \text{sec} \) for AdSH to self-assemble on gold. In case of TPT \( \sim 90 \text{sec} \) of the monolayer forms within the first \( \sim 10 \text{sec} \) of adsorption, and then it takes \( \sim 40 \text{sec} \) to develop into a well-ordered structure. Both SAMs on Au were characterized with ambient STM up to a single-molecule level, X-ray photoelectron spectroscopy and spectroscopic ellipsometry. We were able to obtain near atomic resolved STM images for AdSH SAM on Au. They clearly showed that all AdSH molecules adsorbed in the same conformation with the molecular tilt and the twist angles being \( 114^\circ \) and \( 0^\circ \) correspondingly. We have also demonstrated that most AdSH molecules are arranged in a head-to-tail orientation, with some molecules having a head-to-head orientation without forming a disulfide bond.

Molecular resolved STM images of TPT SAM/Au clearly demonstrated that within first seconds of adsorption the Au-surface is equally covered with two different phases (\( \alpha \)- and \( \beta \)-phase) which dominate on the Au-surface from \( \sim 10 \text{sec} \) of exposure and onwards. A closer inspection of the \( \beta \)-phase on a single-molecule level allowed us to propose an upright conformation for TPT molecule on Au with the tilt angle of \( 0^\circ \). We also estimated the tilt angle for the \( \alpha \)-phase to be \( 10-20^\circ \). For both systems STM and XPS analysis suggest a high chemical and structural quality of the monolayers.

Overall we have proposed a simple and straightforward protocol for ultrafast fabrication of AdSH and TPT SAMs on Au-surface, which we believe can be readily extended for other similar molecular systems.

9:40am SS2-TuM6 Role of van der Waals Interaction in the Binding of 1,4-diaminobenzene to the Au(111) Surface. D. Le, M. Aminpour. University of Central Florida, A. Kijeva, University of Wroc\’law, Poland, T.S. Dresselhaus, University of Central Florida

The physisorption of 1,4 diaminobenzene (BDA) molecule on Au(111) surface is studied within the generalized gradient approximation of the density functional theory with the PBE [1], vdW-DF [2], and vdW-DF2 [3] exchange correlation functionals. The binding energy of an isolated BDA adsorbed on Au(111) surface calculated from vdW-DF (vdW-DF2) is 0.78 eV (0.84 eV) in better agreement with the experimental value (1.0 eV) [4] than that from PBE (0.77 eV), while the alignment of the molecule along the surface obtained from PBE (20°) is closer to the experimental value of 24° [4] than that of about 5° derived from vdW-DF (or vdW-DF2). On the other hand, when interactions between the BDA molecules is included in the calculations, as would be the case if the molecules were to self-assemble, say in the form of a linear line structure on Au(111), inclusion of vdW interactions gives excellent agreement with experimental observations. In particular vdW-DF and vdW-DF2 predict the BDA tilt angles to be 23° and 21°, respectively. The binding energy of a BDA on Au(111) in this case is 0.70 eV and 0.71 eV, respectively. As is to be expected, PBE does not account for intermolecular interactions and does not give good agreement with the data. We suggest the presence of such alignment of molecules along the surface in the experiments, and that they are governed by hydrogen bonding between N and H atoms of neighboring BDA molecules. We compare our results also with unpublished STM data for the system.


*Work supported by DOE Grant DE-FG02-07ER15842.

10:40am SS2-TuM9 Conformational Chirality, Chiral Switching and Chiral Induction in Self-Assembled Molecular Structures. T.R. Linderoth. University of Aarhus, Denmark

INVITED

Chiral self-assembled structures formed from organic molecules have been subject to intense investigation, motivated both by applications such as enantiospecific heterogeneous catalysis as well as by fundamental interest e.g. in relation to the origin of biomolecular homochirality. Chirality on surfaces may arise both for intrinsically chiral molecules and for prochiral molecules that become chiral due to reduced symmetry upon adsorption. However, chiral effects originating from conformational degrees of freedom have received relatively little attention.

Here we use a combination of organic synthesis and UHV-STM experiments to address how rational design of molecular building blocks allows transfer of chirality from the molecular to the supra-molecular level. We investigate a class of custom-designed molecules based on a linear oligo-phenylene-ethynylene backbone and characterize their adsorption structures on the Au(111) surface. Most of these compounds are prochiral and display conformational chirality in the sense that they can adopt in different chiral conformations distinguished by the positions of two tert-butyl side-groups. A novel chiral switching mechanism, involving a conformational change where the terminal groups rotate around the molecular axis, is directly revealed from time-resolved STM. We demonstrate that it is possible through control of the terminal group functionalization to steer the molecular backbones into surface assemblies that are either mirror symmetric or display pronounced organizational chirality in the form of a characteristic windmill motif. We furthermore achieve control over the absolute chirality of windmill assemblies by synthesizing an intrinsically chiral variant where the tert-butyl side pendant is replaced by a chiral (S)-sec-butyl group. This intrinsically chiral compound is finally used in co-deposition experiments as an induction seed to control the chirality of assemblies formed from the original prochiral compound.

Chiral switching by spontaneous conformational change in adsorbed organic molecules


Steering organizational and conformational surface chirality by controlling molecular chemical functionality


Controlling chiral organization of molecular rods on Au(111) by molecular design

repulsion. In recent years charge transfer through PNA molecules has been a focus of research due to potential applications in self-assembled molecular circuits. This makes it interesting to investigate the electronic structure of PNA interfaces to electrode materials. A widely used strategy to ‘connect’ PNA molecules to metallic electrodes is through thiol-Au bonds using a terminal cysteine appended to PNA oligomers. This motivated the here presented research where the electronic structure of self-assembled PNA monolayers on Au substrates was investigated. Cys-appended PNA 7-mers of thiolamine (Cys-7T) were incubated on Au substrates in a nitrogen glove box attached to a photoemission spectrometer. Ultraviolet and x-ray photoemission spectroscopy (UPS and XPS) measurements on the resulting SAMs revealed the hole injection barrier at the interface and the interface dipole. Electronic structure calculations based on molecular dynamics sampling of the PNA structure yielded the band gap and the electronic density of states for PNA. Combined with the UPS data, the theoretical calculation enabled the estimate of the electron injection barrier at the interface, as well as the assignment of individual UP spectral features to specific molecular orbitals. Control measurements on Cys-appended, abasic PNA backbone 7-mers allowed the identification of the emissions related to the PNA backbone in the UP spectra. The orbital line-up at the interface between the Au substrate and the Cys-PNA indicates a significant interface dipole resulting in the alignment of the Au Fermi level near the center of the PNA HOMO-LUMO gap. This alignment causes large charge injection barriers for both holes and electrons, and thus impedes charge transfer from Au into the Cys-PNA SAM.

11:40am SS2-TuM12 Immobilization of Single-Stranded DNA Probe on InAs Surfaces for Biosensor Application, E.K. Cho, University of Wisconsin, A. Brown, Duke University, T.F. Kuech, University of Wisconsin

We study the immobilization of single-stranded DNA (ssDNA) probe on InAs surfaces and its characterization for diagnostic application. In contrast to other semiconductors, the Fermi level in InAs is typically pinned above the conduction band minimum, resulting two-dimensional electron gas (2DEG) located immediately below the surface. The InAs based system forms the basis of a DNA sensing platform because dimensional electron gas (2DEG) located immediately below the surface.

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Thin Film Division
Room: 107 - Session TF+EN-TuM

ALD for Energy
Moderator: W.M.M. Kessels, Eindhoven University of Technology, the Netherlands


Platinum nanoparticles are used as the catalyst on the cathodes of proton exchange membrane (PEM) fuel cells. Controlling the dispersion and size of the Pt nanoparticles is important for efficient and cost-effective fuel cells. When Pt atomic layer deposition (ALD) is performed on oxide or carbon substrates, nucleation difficulties and the high surface energy of Pt lead to Pt nanoparticles rather than continuous films. This research explored strategies to control the dispersion and size of Pt nanoparticles using Pt ALD together with various surface treatments. Pt ALD was performed using Pt hexafluoroacetylacetonate Pt(hfac); and formalin as the reactants. Titanium oxide (TiO2) and tungsten oxide (WO3) were explored as the substrates. We used in situ transmission Fourier transform infrared (FTIR) spectroscopy to monitor the surface species during Pt ALD. Surface poisoning by hfac species was observed during the nucleation of Pt ALD on TiO2, in agreement with previous studies of Pt ALD [1]. Trimethylaluminum (TMA) was able to remove the hfac species from TiO2, and promote more facile nucleation of Pt ALD, also as expected by our earlier work on Pd ALD [2]. We then used hfac adsorption prior to Pt ALD to block surface sites, delay Pt ALD nucleation and decrease Pt nanoparticle dispersion. In addition, we used TMA exposures after Pt(hfac); exposures to facilitate Pt ALD nucleation and increase Pt nanoparticle dispersion. The Pt nanoparticles were detected by the rising absorbance baseline of the FTIR spectrum, and transmission electron microscopy images of Pt nanoparticles on TiO2 and WO3, particles. The size of the Pt nanoparticles was dependent on the number of ALD reaction cycles.


Transition metals, such as Pt, are important as catalysts in fuel cells. The cost of Pt requires that this metal be utilized as efficiently as possible. Previous studies have demonstrated that flat, continuous Pt surfaces have an activity for the fuel cell oxygen reduction reaction (ORR) per surface Pt atom that is 5-10 times larger than the activity of 3 nm Pt nanoparticles on carbon-supported electrocatalysts. Consequently, very thin continuous Pt layers are needed to achieve high activity per mass of Pt and favorable economics.

Pt has a high surface energy and does not readily wet low surface energy oxide and carbon material surfaces. Instead, Pt nanoclusters are formed during the nucleation of Pt ALD on these low surface energy supports. A continuous Pt film is possible only after the coalescence of the nanoclusters when the film thickness is >5 nm. One possible route to obtain a continuous and ultrathin Pt film is to deposit on an adhesion layer that has a higher surface energy than Pt. In this case, the Pt will wet the adhesion layer because the deposited Pt film will lower the surface energy. One material that has a higher surface energy than nearly all other metals, including Pt, is tungsten (W).

W ALD using WF6 and SiH4 is known to nucleate and grow rapidly on Al2O3 ALD surfaces [1]. The surface chemistry of W ALD is able to form chemical bonds to the underlying Al2O3 substrate that overcome the surface energy differences. In this study, Pt ALD is performed using MeCpPMe3 and H2 plasma on W ALD adhesion layers grown on Al2O3 ALD films. X-ray photoelectron (XPS) and x-ray reflectivity (XRR) studies have demonstrated that Pt ALD nucleates nearly immediately on the W ALD adhesion layers. The XPS and XRR data are consistent with a layer-by-layer growth model. Pt ALD films can be deposited that are 1 nm thick, conformal and continuous. Initial electrochemical measurements on these
thin films have yielded a large proportion of the anticipated ORR activity benefit.


9:00am TF-EN-TuM4 Engineering Li,Al,Si,O Ion Conducting Thin Films by Atomic Layer Deposition for Lithium-Ion Battery Applications, Y.-C. Perng, J. Cho, D. Membrano, N. Cirigliano, B. Dunn, J.P. Chang, University of California Los Angeles

Lithium (Li)-ion batteries have drawn much attention for their outstanding performance in portable electronics applications. However, formation of the solid electrolyte interphase (SEI) layer on the surface of electrodes during the charge-discharge cycling can reduce battery capacity and the long-term reliability of current battery technology. The use of solid electrolyte layers can effectively suppress formation of the SEI. Another application for thin solid electrolytes is in microbatteries, especially those based on the engineering of electrodes into 3D architectures involving high aspect ratio pillars. To realize this potential, an ultra-thin and highly conformal electrolyte layer is needed to coat the 3D electrode array. The ionic conductor lithium aluminosilicate (LiAlSiO4) synthesized by atomic layer deposition (ALD), is a promising candidate for these battery applications. The material exhibits high ionic conductivity along its c-axis because of channels formed by the alternating tetrahedra of aluminum-oxygen (Al-O) and silicon-oxygen (Si-O).

The self-limiting characteristic of ALD allows for precise control of thickness and composition of complex oxides and results in a highly conformal and pinhole-free coating suitable in 3D micro-battery applications or electrolyte surface coatings. The metal precursors used in this work are tetraethyl orthosilicate (TEOS), trimethylaluminum (TMA) and lithium t-butoxide (LTB). These precursors, along with water vapor as the oxidant, were used to deposit SiO2, Al2O3 and Li2O, with deposition rates in the range of 0.8-2 Å/cycle, respectively. The deposition rate of stoichiometric LiAlSiO4 was ~5 Å/cycle at a temperature of 290°C. The concentration of each metal element in Li1-xAlxSi1-O (LASO) thin films is found to correlate closely to ALD cycles and the associated incubation times. Complex impedance measurements show that the ionic conductivities found to correlate closely to ALD cycles and the associated incubation times.

The results are relevant to important energy research fields where they need finely tuned nanostructures to meet intricate requirements.

Second, scaleup of ALD equipment, processes, and control is required for cost-effective ALD manufacturing. Significant efforts are already underway to develop these technologies, notably including roll-to-roll and other high throughput approaches, as well as atmospheric pressure ALD to circumvent the cost and complexity of vacuum-based equipment.

* Supported by the US Department of Energy, Office of Basic Energy Sciences as part of an Energy Frontier Research Center, and by the Laboratory for Physical Sciences.


Dye-sensitized solar cells (DSCs) are a promising alternative to Si-based photovoltaic cells because of their high efficiency, low cost, and simple processing. To improve the efficiency, the light harvesting properties of the DSC photoanode structure should be improved.

This work shows the ability to improve DSC efficiency by the incorporation of quartz fiber mats functionalized by ALD with a conformal nanoscale TiO2 coating for the photoanode. The TiO2 atomic layer deposition allows precise nanoscale thickness control throughout the quartz fiber mat. In addition, thermal stability of the quartz fiber also enables the annealing of TiO2 films at high temperatures in order to obtain anatase crystallinity, which is the preferred TiO2 phase in DSCs, due to faster electron transfer kinetics. As a result, we obtained very stable anatase TiO2 on quartz fibers by annealing at as high as 1050 °C. The micro-sized randomly oriented structure of coated quartz fibers caused high light scattering effect inside the photoanode, so that it increased the photon adsorption. At the same time, TiO2-coated quartz fibers were successfully sensitized by more dye molecules compared to bare quartz fibers.

The use of the dye-sensitized quartz fiber mats in a DSC framework showcased devices with overall efficiencies exceeding 7% in our laboratory, compared to ~ 6% for similar devices without the coated fiber scattering layer. We will present X-ray diffraction, morphological changes, and optical properties of TiO2-coated quartz fibers, as well as IV and IPCE data for the DSCs. The effects of various integration schemes for introducing quartz fibers into the DSC photoanode will be discussed. Moreover, it is believed that the combination of quartz fiber and ALD is very attractive especially to energy research fields where they need finely tuned nanostructures to meet intricate requirements.

9:20am TF-EN-TuM5 ALD: Enabling Designer Nanostructures for Energy Applications, G.W. Rubloff, University of Maryland INVITED

Nanostructures will dominate next-generation energy technologies. Progress in nanofabrication increasingly allows design flexibility control of structural geometry and material combinations to achieve high performance multifunctional 3-D nanostructures for energy harvesting and storage. Such designs derive advantages from high aspect ratio pillars. To realize this potential, an ultra-thin and highly conformal structural electrode layer is needed to coat the 3D electrode array. The ionic conductor lithium aluminosilicate (LiAlSiO4) synthesized by atomic layer deposition (ALD), is a promising candidate for these battery applications. The material exhibits high ionic conductivity along its c-axis because of channels formed by the alternating tetrahedra of aluminum-oxygen (Al-O) and silicon-oxygen (Si-O).

The self-limiting characteristic of ALD allows for precise control of thickness and composition of complex oxides and results in a highly conformal and pinhole-free coating suitable in 3D micro-battery applications or electrolyte surface coatings. The metal precursors used in this work are tetraethyl orthosilicate (TEOS), trimethylaluminum (TMA) and lithium t-butoxide (LTB). These precursors, along with water vapor as the oxidant, were used to deposit SiO2, Al2O3 and Li2O, with deposition rates in the range of 0.8-2 Å/cycle, respectively. The deposition rate of stoichiometric LiAlSiO4 was ~5 Å/cycle at a temperature of 290°C. The concentration of each metal element in Li1-xAlxSi1-O (LASO) thin films is found to correlate closely to ALD cycles and the associated incubation times. Complex impedance measurements show that the ionic conductivities found to correlate closely to ALD cycles and the associated incubation times.

The results are relevant to important energy research fields where they need finely tuned nanostructures to meet intricate requirements.

Atomic layer deposition (ALD) plays a pivotal role in this paradigm because of its intrinsic attributes: thickness control at the atomic scale; exceptional conformality to apply this control in the most demanding nano geometries; and a growing portfolio of ALD materials choices from new precursors and processes. As a result, ALD is becoming common, even pervasive, in nanofabrication-based energy research, spanning applications from heterogeneous nanowires to passivating electrochemical layers and high-efficiency catalytic nanostructures.

Two primary challenges will determine whether ALD’s potential is realized in next-generation energy technology.

First, integration of ALD with other processes into suitable process sequences determines how well nanostructure designs can be fabricated and tailored for the energy application. For example, in high aspect ratio trenches, pores, or aperiodic porous materials, ALD films can be highly conformal at the top; a more conformal or tapered “bottom-up” to thinner layers deeper in the structure, while electrodeposited films can be grown “bottom-up” from a working electrode at the bottom, together offering more design flexibility in vertical profiles than does either process alone.

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energy generation from radiation, sensors, and electromagnetic wave detectors.

Thin Film Division

Room: 104 - Session TF+SE-TuM

Glancing Angle Deposition (GLAD) I

Moderator: T. Karabacak, University of Arkansas at Little Rock

8:20am TF+SE-TuM2 Progress in Glancing Angle Deposition Technology for Practical Applications. M. Suzuki, Kyoto University, Japan

Recently, advanced top-down processes for the fabrication of nanostructures have been developed; however, these processes are long predicated by the processing window of the technology in question. By employing a GLAD technique to fabricate thin film, we can improve upon these limitations and achieve enhanced characteristics for the new materials. In this presentation, we will introduce the technique and discuss the importance of learning from the work of the previous researchers and moving forward.


Organic photovoltaics (OPVs) represent one possible route to widespread adoption of solar energy production. The most promising OPV technology to date has been the bulk heterojunction, a photoactive mixture which breaks the gap between exciton diffusion lengths (~15 nm) and typical device length scales (~200 nm). Here we introduce the recent developments in OPV technology that have led to improvements in device performance, including the use of novel materials and nanostructured electrodes.

9:20am TF+SE-TuM5 Enhanced Photoconductivity Response of Glancing Angle Deposited Indium Sulfide Nanorod Arrays for Photodetector Device Applications. H. Is, M.F. Canizoigulo, T. Karabacak, University of Arkansas at Little Rock

Photoconductivity (PC) can be improved by altering the structure of the photodetectors. This can be achieved by using advanced top-down processes that employ GLAD to fabricate thin films. In this study, we fabricated photodetectors with indium sulfide nanorod arrays, which exhibited enhanced PC response due to their low dark conductivity and superior capability of absorbing light. We will discuss the results of this study and the potential applications of this technology.

10:40am TF+SE-TuM9 Catalytic Nanomotor Control: Design Techniques Using Dynamic Shadowing Growth. J.G. Gibbs, Y.P. Zhao, University of Georgia

Catalytic nanomotors can be controlled by changing the location of the catalyst. In this study, we investigated the use of dynamic shadowing growth (DSG) to manipulate the position of the catalyst and control the motion of the nanomotor. The results of this study suggest that DSG can be used to control the movement of catalytic nanomotors with high precision.

Here we present a summary of organic solar cell research utilizing GLAD at the University of Alberta. GLAD’s flexibility lends itself to structuring both the transparent conductor layer as well as the donor and acceptor photoactive layers. We have demonstrated GLAD structuring of indium tin oxide (ITO), copper (II) phthalocyanine (CuPc), and fullerene (C60). ITO nanopillars fabricated by GLAD have been incorporated as three-dimensional high surface area electrodes in organic photovoltaic devices. The nanostructured electrodes demonstrated 30% improved performance compared to planar ITO anodes, due to increased optical absorption and high surface area. Typical OPV donor and acceptor materials, CuPc and C60, have also been successfully structured by the GLAD technique [4,5]. Advanced motion control algorithms, designed to form thinner columns, were used to optimize the material nanostructures and produce highly desirable ordered bulk heterojunctions when coupled with complementary polymers. In the case for GLAD C60 devices, the short-circuit current was double that of bulk heterojunction devices. The GLAD technique has proven to be an incredibly useful method for fabricating and tuning electrode and bulk heterojunction morphologies in OPVs.

efficiency and IPCE of the Cr:TiO$_2$ films is reduced when compared to grain boundaries and the surface of the nanorods. The photocatalytic produced mixed results regarding the effect of the Cr dopant on intrinsicTiO$_2$ due to changes in the photocatalyst’s surface and charge rotation rate, deposition rate, and final film thickness. By qualitatively fingerprints with the CEFR technique. We varied: base pressure during comparing fingerprints before and after development for various values of pressure. An optimal base pressure of 0.1 mTorr significantly relaxes the vacuum requirements of an evaporation chamber to be used for this purpose, making the technique very accessible to forensic scientists and law vacuum requirements of an evaporation chamber to be used for this purpose, making the technique very accessible to forensic scientists and law challenges encountered in the fabrication of vacuum chambers, photon absorbers and RF shielded bellows will be described. The status of the project will also be summarized.

9:00am VT-TuM4 The Large Cryopump System for the Heating Neutral Beam Injection of ITER, S. Hanke, M. Scannapiego, X. Luo, X. Day, Karlsruhe Institute of Technology (KIT), Germany, F. Fellin, P. Zaccaria, Consorzio RFX, Italy The ITER Neutral Beam Injection system (NBI) is one of the heating systems to achieve the required plasma temperatures to start the fusion process. Thereby, the NBI system is basing on the principle of provision and acceleration of deuterium and protium ions and the re-neutralization of the high energy ions to be injected into the plasma through the confining magnetic fields. Each heating NBI is designed to insert 16 MW of heating power to the plasma and presents major technical and physical challenges. In order to solve these and to demonstrate the achievement of the required parameters, a robust R&D program is under way. A central milestone for this development is the establishment of a full scale test facility, which will be built on site of Consorzio RFX, Padova. Part of this test facility is MITICA (Megavolt ITER Injector and Concept Advancement), a hardware bed for the entire neutral beam injection system, Karlsruhe Institute of Technology (KIT), which is the lead party in design and R&D of the ITER cryopumps since more than a decade, is supporting this project with the development of a customized cryopump design which ensures that the requested density profiles for optimum beam performance can be produced. The main operational task which has to be provided by the cryogenic pump at a speed of ~5000 m$^3$/s is to handle very high gas loads of protium and deuterium. As basic pumping concept, cryosorption concept was adopted and the cryopump is operated with supercritical helium at 4.5 K for the adsorbing and gaseous helium at 80 K for the shielding circuit. As demonstrated in other NBI applications, cryosorption provides a wide and robust operational window at acceptable cryogenic loads to the cryoplant. The design was driven by two competing requirements: The high thermal heat loads ask for a closed pump, whereas the need for a high pumping speed asks for an open structure. To reconcile both objectives in an optimized geometry, modeling simulations were performed using the Test Particle Monte Carlo code MOVAK3D. To properly describe the density distribution in the NBI vessel with large thermal and pressure gradients, the time-of-flight cell code ProVac3D was developed.

Additional to the design activities for an optimized cryopump, a considerable effort has been spent to investigate the thermal hydraulic properties of the cryopump during the different operational and failure modes.

9:20am VT-TuM5 Design and Construction of the Vacuum System for SuperKEKB, Y. Suetsugu, K. Shibata, H. Hisamatsu, M. Shirai, T. Ishibashi, K. Kanazawa, KEC, Japan The upgrade project of KEKB B-factory, that is SuperKEKB, has started last year. The SuperKEKB is a two-ring electron-positron collider with 7.0 GeV electrons and 4 GeV positrons aiming a goal luminosity of 8.0×10$^{35}$ cm$^{-2}$s$^{-1}$, which is approximately 80 times higher than that of KEKB. In order to realize the unprecedented luminosity, the stored beam currents are increased to 2.4 A and 3.6 A for electrons and positrons, respectively. The vertical beam sizes at the collision point are also squeezed to approximately 60 and 50 nm with the beam emittances of 5/13 and 3/9 nm/pm (horizontal/vertical) for electrons and positrons, respectively. The vacuum system is accordingly improved to achieve the challenging goal. Beam pipes with antechambers are adopted for the reduction in the beam impedance of beam channel and also in the irradiation power density of vacuum chambers will be presented, including operational experience with the newly available Watanabe Bent-Belt Beam (BBB) gauge, and the incorporation of a bakable cryopump into the pumping configuration.

8:00am VT-TuM1 Continued Work toward XHV for the Jefferson Lab Polarized Electron Source, M.L. Stutzman, P.A. Adderley, Thomas Jefferson National Accelerator Facility The Jefferson Lab DC, high voltage polarized electron source utilizes deep-UHV pressures to limit photocathode damage due to ionization and acceleration of residual gasses into the photocathode, as well as preserve the surface chemistry necessary for electron emission. Continued efforts toward improving and quantifying pressures below 5×10$^{-10}$ Torr for electron source trajectories change as a function of arm length. Simulations based on the method of regularized Stokeslets are also described and correctly capture the trends observed in the experiments.
synchrotron radiation from intense stored beams. Various vacuum components adaptable to the antechamber scheme with low beam implosion and high thermal strength had been developed. The bellows chambers, for example, have a comb-type RF-shield, and the main vacuum pump is NEG strips inserted into one of antechambers. Special attention is paid for the mitigation of the electron cloud issues in the positron ring to avoid unwanted increase in the beam emittance. In order to reduce secondary electron emission, the inner surface of beam pipes is coated with TiN, where mound layer is prepared in dipole magnets, and the newly developed clearing electrodes are introduced in wiggler magnets. The antechamber scheme is also effective to suppress the photocurrent effect. The beam pipes in drift spaces, furthermore, are winded by solenoid coils. The design of vacuum system has been mostly completed, and the mass production of beam pipes has started. Copper beam pipes with clearing electrodes for wiggler magnets had been already delivered. Aluminum beam pipes for arc section of positron ring are under manufacturing, together with bellows chambers, gate valves, NEG pumps, and so on. The installation to the ring will start next year after TIN coating, expecting the start of commissioning from 2014. The overall vacuum system design and some key issues for SuperKEKB together with the present status will be reported here.

Since the introduction, extensive studies have been carried out indicating vacuum pumps with better performance in smaller packages. This is driving pump manufacturers to redesign their pumps and/or to consider new ways to combine pumping technologies more efficiently. In response to this trend cryopump manufacturers are revising their product range in order to combine pumping technologies more efficiently. In response to this trend it has been found it may be advantageous in UHV-XHV systems to use Non Evaporable Getter (NEG) pumps as the main pumping element and complement it with a small ion pump to remove inert gases and methane. A novel design of such a combination, called NEXTorr®[1], was first introduced in 2010 at this conference.

The model used the values obtained by Grant, Cummings, and Blackburn for the diffusivity, recombination, and permeation of hydrogen in stainless steel. If one assumes a plausible value for the initial hydrogen concentration, it gives outgassing rates in rough agreement with the measurements.

11:00am VT-TuM10 Modelling and Simulation of the ITER Cryopumping Systems. C. Day, Karlsruhe Institute of Technology, Germany

A cryopump is probably the most versatile and flexible vacuum pump. In large R&D applications where cryopumps are available and high pumping speeds at high throughputs are required, it is often advantageous to exploit a directly cryogen-supplied cryopump. The nuclear fusion project ITER is a perfect example for such a project, which triggered the development of customized cryopumps. To name just two advantages of this approach, a cryopump can be designed to perfectly fit the available space, and can be installed in-situ without any conductance losses, if regeneration frequency allows for that.

Karlsruhe Institute for Technology (KIT) is developing tailor-made cryosorption pumps for fusion applications over the last 20 years. This has been associated with an extensive design supporting R&D programme which has provided a broad parametric database and stimulated the development of modeling and design tools.

This paper will delineate the essential steps one has to consider when designing a cryosorption pump. The design process of a customized cryopump starts with the proper identification of the set of requirements, which defines the requested integral pumping speed at the given pump location and space. The tools needed for individual cryopump design are described and typical examples are given. This includes the calculation of capture coefficients and distributed pumping on the cold surfaces by means of Test Particle Monte Carlo methods.

Cryopump examples are taken from the area of large cryopumps for ITER, such as the torus cryopumps (~80 m³/s) and the cryopumps for high energy neutral beam injection development (~5000 m³/s). Although being cryogenic pumps, these applications are characterised by relatively moderate vacua due to the high gas throughputs during pumping. This also leads to the fact that transitional flow conditions prevail inside the pump, which results in additional challenges with regard to modelling and operation. Both pump types are currently in the stage of build-to-print design finalization and prototypes will be manufactured to validate this design in dedicated testbeds at KIT and Padova, Italy.

11:40am VT-TuM12 Design and Construction of the Ultrahigh Vacuum System for the 3 GeV TPS Accelerator. G.Y. Hsiao, H.P. Hsueh, C.L. Chen, J.R. Chen, National Synchrotron Radiation Research Center, Taiwan, Republic of China

The ultrahigh vacuum system for the electron storage ring of the 3 GeV Taiwan Photon Source (TPS) accelerator has been started the construction since 2010. The critical vacuum components with lower impendence design including the bellows with spring-finger rf-contact, the metal gate valves with comb-finger rf-contact, pulsed magnet kicker ceramic chambers, beam position monitors, crotch absorbers, and the precise machined sector bending chambers and BPM chambers, have been manufactured. The large aluminum alloy (Al-) bending chambers for the arc-cells have been precisely machined with oil-free machine tools in the clean room, and undergoing the ozonate water cleaning after with the precisely in-house welding. Mass production of the vacuum equipments including the ion gauges, ion pumps, NEG pumps, and gate valves, has been contracted out and partially delivering following the schedule of the cell assembling. Each cell, contains two short AI-straight chambers and two AI-bending chambers, has been started the assembling and on-site welding on the pre-aligned girders in clean room forming an one-piece vacuum vessel about 14 m in length following by the vacuum baking to the ultra-high vacuum. The progress of prototyping development and the status of construction for the TPS ultrahigh vacuum system will be described in this paper.
1:00pm EW-TuL4 Optimized XPS Depth Profiling of Organic Materials using Polyatomic Ion Sources, D. Surman, Kratos Analytical Inc., C. Blomfield, A. Roberts, S. Page, Kratos Analytical Ltd., UK
Over the last several years multiple methods have developed for the depth profile analysis of organic materials using a variety of different polyatomic species as the sputtering ion. It has also become clear that different classes of polymers require quite different sputtering parameters to be effectively profiled with no loss of chemical information. In this presentation we discuss a variety of approaches that have developed in order to effectively sputter profile a wide range of materials. These approaches utilize differences in ion beam energy, impact angle and sample temperature in order to achieve an effective sputter profile.

1:20pm EW-TuL5 Advances in XPS Chemical Imaging and Depth Profiling, J.S. Hammond, D.G. Watson, P.E. Larson, S.N. Raman, Physical Electronics
Optimized scanning x-ray microprobe technology has been shown to provide superior sensitivity with minimal data artifacts for micro-area XPS. To improve the chemical state sensitivity, computation methods have been developed to provide a 10X improvement in count rate for chemical state spectroscopy with reduced x-ray damage. Utilizing the unscanned analyzer mode of operation, a user selectable number of data channels are now available for optimized chemical state imaging and chemical state depth profiling. Examples of these new computational methods will be presented for XPS chemical state imaging of patterned semiconductor and polymer samples as well as organic XPS depth profiling.

Tuesday Lunch, November 1, 2011
Exhibitor Technology Spotlight
Room: West Exhibit Hall - Session EW-TuL

Exhibitor Technology Spotlight
Moderator: Langley

Surface structure and chemistry are crucial to the successful production and operation of innumerable devices, materials and coatings. X-ray photoelectron spectroscopy, with its high specificity and chemical state sensitivity, is an ideal tool for the evaluation of material composition. XPS depth profiling allows the identification of chemical variations in materials from the surface to bulk, and facilitates characterisation of complex layer structures. Recent advances in ion source design have seen the introduction of noble gas cluster ion beams for depth profiling applications, which allow materials that are unstable under monatomic ion bombardment to be analysed.

Solutions to structural and chemical problems using the full range of state-of-the-art, fully integrated X-ray photoelectron spectrometers from Thermo Fisher Scientific are presented. These include the characterisation of thin film polymer coatings, and determination of the structure of multilayer stacks. The effectiveness of the analyses, and the automated data refinement processes using new features of the award-winning Avantage data system, are shown for each of these examples.

The application potential of graphene is currently being extensively explored by the materials science community. Its utility as a transparent conductive electrode for the microelectronics industry is already being exploited. More recent progress has demonstrated how the unique combination of electronic, chemical, and structural properties of graphene will have a significant impact on the development of next-generation thin film transistors. Additional applications of graphene to molecular sensors are under way. In all stages of development there is a critical requirement for materials characterization and analysis: from the initial research stages through to testing of the finished devices. Because most materials need to be analyzed for compositional homogeneity both across the surface and through the thickness of a given sample, a complementary approach involving several techniques is often required.
In this presentation we will discuss how a multi-technique approach using Raman spectroscopy and XPS can address the problems associated with the analysis of ultra thin film materials. Raman microscopy is a vibrational technique that is very sensitive to small changes in a molecule’s geometric structure and its environment. This sensitivity allows Raman to be used as a probe for a number of properties important to a specific graphene sample. These properties include, but are not limited to layer thickness, the presence or absence of defects, and local strain. XPS enables complete characterization of thin graphene films with respect to chemical modification, in addition to the chemical interaction between the film and the substrate. The combined XPS/Raman measurement approach will be applied to graphene produced by both exfoliation and CVD methods, providing a full comparison of the chemical and structural information offered by each technique.

Tuesday Lunch, November 1, 2011

1:00pm EW-TuL4 Optimized XPS Depth Profiling of Organic Materials using Polyatomic Ion Sources, D. Surman, Kratos Analytical Inc., C. Blomfield, A. Roberts, S. Page, Kratos Analytical Ltd., UK
Over the last several years multiple methods have developed for the depth profile analysis of organic materials using a variety of different polyatomic species as the sputtering ion. It has also become clear that different classes of polymers require quite different sputtering parameters to be effectively profiled with no loss of chemical information. In this presentation we discuss a variety of approaches that have developed in order to effectively sputter profile a wide range of materials. These approaches utilize differences in ion beam energy, impact angle and sample temperature in order to achieve an effective sputter profile.

Tuesday Lunch, November 1, 2011
2:00pm AS-TuA1 Integration of an External Cavity Quantum Cascade Laser Into a Scattering-Type Infrared Scanning Near-Field Optical Microscope, A.S. Lea, M.S. Taubman, M.C. Phillips, Pacific Northwest National Laboratory, M. Raschke, University of Colorado, Boulder

Molecular nanostructures, polymer and supramolecular assemblies, proteins, biomembranes, correlated systems, and many other natural and synthetic materials gain their unique functionalities from intra- and intermolecular interaction and electron correlations on mesoscopic length scales of 10’s of nm. Gaining a molecular level understanding of the materials structure and function has remained a major experimental challenge. This is due to the lack of experimental techniques that can routinely provide a chemically specific spectroscopic identification with simultaneous nanometer spatial resolution on the relevant length scale associated with the size and interactions of the molecular building blocks: within the 10 – 100 nm range. We have developed an instrument for spectroscopic infrared vibrational near-field nano-imaging. The instrument is based on a monopolar semiconductor laser devices that can be fabricated to cover significant regions of the mid IR spectrum, specifically 3.5 to 20 microns. Moreover, these devices work extremely well in the molecular fingerprint region (8 to 12 microns), which will be of particular use in this instrument as this is where many fundamental vibrational bands are found, where MCT IR detectors work best, and the region is relatively free from water interference. The rapid scanning capabilities of External Cavity QCXs (ECQCLs) in the 100s of Hz, will allow IR spectra to be taken point by point across the sample, allowing rapid spectral data coverage. This is to be compared with the Hz-rate scanning of the OPO chain. The second significant advantage of OCL incorporation will be its ease of use (permitting wide-spread usage), low cost and ruggedness. This presentation will cover the integration of an ECQCL into the existing scattering-type IR scanning near-field optical microscope and demonstrate its ability to provide spatially resolved IR spectroscopic signatures on a sub-100nm scale. Ultimately, we anticipate this instrument will be able to provide chemical binding information of molecular adsorbates on nanostructured materials.

2:20pm AS-TuA2 An Investigation Into the Aging of Paintings using Surface Analysis Techniques, T.S. Nunney, Thermo Fisher Scientific, UK, J.J. Boon, AMOLF, Netherlands, E.S.B. Ferreira, Swiss Institute for Art Research (SIK-ISEA), Switzerland

The understanding of the processes involved in the aging of artworks is of considerable importance in maintaining these pieces for future generations. Delamination of paint layers, chemical reactions of constituents in and between paint layers, interactions with the environment and the influence of the environment on the works can all have an effect on their appearance. By understanding these processes, conservators can attempt to retain the appearance as the artist intended. Surface analysis has been used to investigate a cross-section sample taken from “Autumn in Oschwand” (1906) by Cuno Amiet (1865-1961). The painting has become lighter in certain areas, and the cause of this was investigated. EDS analysis indicated that a higher proportion of lead had accumulated near the surface of the outer paint layer. XPS imaging of the sample in conjunction with multivariate statistical analysis of the data allows the chemical state of the individual components to be distinguished, from lead carbonate (or “lead white” pigment) to the lead soaps that are causing the change in the appearance of the painting, and the copper, chromium and arsenic pigments that form the paint layers. These analyses were compared with spectra from reference pigments to identify the original paints used. Further investigation using Raman and FTIR microscopy was used to complete the analysis.


Insect cuticle is a matrix of structural proteins and long chain polymers of chitin. The chemical composition of the cuticle is dictated by the biomechanics of the insect. Regions of the exoskeleton where extra stiffness is required - this matrix become mineralized and form scalaris. Elastic properties of the cuticle are due to the pliable structural proteins. While engineers have set out to copy these biomimeralization processes and design materials that mimic the extraordinary structural capabilities of these exoskeletons – a complete understanding of the structure of this chitin/protein/mineral matrix does not currently exist. In this study, we set out to spatially resolve the chemistry, at the cuticle surface, of an African Flower Scarab (Eudicella grallti) by near edge x-ray adsorption fine structure (NEXAFS) imaging. The NEXAFS images are produced by a new parallel process magnetic field electron yield optics detector, and a full field incident soft X-ray beam on the sample. The rapid parallel process magnetic field electron yield optics detector (LARIAT: Large Area Imaging Analytical Tool) produces a series of two-dimensional NEXAFS spatial images as the incident x-ray energy is scanned above a K or L absorption edge. A spatially resolved view of cuticle surface chemistry was created by mapping spectral features within the carbon (270-370 eV), nitrogen (380-430 eV), and oxygen (520-580 eV) K-edges. Distributions of protein and chitin rich regions around the beetle were defined by changes in intensity of π∗(284.5 eV) and C-Η* (290 eV) within the C K-edge spectra. Regions of high mineralization were observed around the joints of the beetle’s head and were assigned by tracking the intensities of the calcium (330-360 eV) and iron (700-740 eV) L-edges. These images also showcase the strengths of NEXAFS imaging, which unlike other photoelectron spectroscopy modalities, allowed us to collect high quality spectra over a large field of view (12 mm x 18 mm) at a range of x-ray incidence angles, with little beam damage to the tissue.

3:00pm AS-TuA4 Surface Cleaning of Organic and Inorganic Materials with Argon Cluster Ion Beams, A.E. Wright, P. Mack, O. Greenwood, Thermo Fisher Scientific, UK

Surface analysis of many materials can be impeded by the presence of contaminant substances at the surface. Signals from these contaminants can dominate a photoelectron spectrum, leaving little contribution from the underlying material of interest. Spectroscopic analysis of contaminated samples can be a considerable challenge in surface science, so methods of cleaning samples are of some importance. One common method of removing contamination in vacuo is argon ion sputtering, which can be effective at removing surface materials. Damage to the underlying material can be substantial with argon ions, however, particularly at high impact energies. Ion impacts can disrupt polymer structures and reduce inorganic oxides, so that the surface after cleaning does not closely resemble the intact material.

The recent development of argon cluster ion beam sources promises significant improvements in depth profiling of soft materials. Cluster beams can offer exceptionally gentle sputtering, and so may be ideally suited to removal of contaminants and surface modifications from a variety of samples. In this presentation, we evaluate the utility of argon cluster ions for the surface cleaning of various materials, in combination with the Thermo Scientific ESCALAB250Xi surface analysis instrument for characterisation of the samples. The results of gas cluster profiling on polymeric and inorganic materials will be presented, with spectroscopic and imaging analyses showing the benefits of this method.


The sputtering of solids with ion beams followed by analysis with photoelectron spectroscopy has been widely used as it offers a powerful strategy for the in-depth characterization of complex inorganic materials. The combination of imaging techniques with depth profiling to create three-dimensional information is an obvious and exciting extension of these experiments. Recent introduction of cluster ion beams, such as coronene,
into XPS instrumentation offers capabilities in quantifying the chemical and molecular gradients in the near surface region of soft materials, such as polymers and biomaterials. XPS data can be used as a function of sputter depth into organic materials while maintaining molecular integrity. There are only a handful of studies combining XPS imaging and depth sputtering. One of the biggest problems when dealing with 3D imaging data sets is visualizing the lateral distribution of chemical moieties as a function of depth. Multispectral images are acquired at each sputtered depth, four-dimensional datasets can result with a full spectrum acquired at each voxel of the space. These datasets represent a huge amount of data which can only be interpreted with assistance of Multivariate Analysis.

In this research we report on application of various types of MSA methods such as two-step PCA and two-step MCR-ALS, PARAFAC, MFA, Tucker and 3D MCR-ALS to multispectral XPS imaging data acquired as a function of sputtering depth for different heterogeneous biomaterials and polymers. The component images extracted from MSA, which show the spatial distribution of the various chemical components, will be visualized in 3D individually or together representing an overall chemistry of individual layers.


Intermixing between thin film layers can alter mechanical and thermal transport properties, phase stability and growth textures. Quantification of the degree of intermixing is crucial to identify the mechanisms of intermixing and their scaling effect on properties as listed previously. Atom probe tomography has received considerable attention for this type of characterization because of its ability to identify and provide reconstructions of atoms with near atomic spatial 3D resolution. In general, these atom probe reconstruction algorithms assume a constant evaporation field across the surface of the specimen. In reality, chemical inhomogeneity (i.e. discrete interfaces) modulates the evaporation field at the specimen surface. This introduces reconstruction artifacts and degrades the spatial resolution of the atom probe tomography technique. Multilayer thin films provide ideal specimen geometries to measure and quantify these artifacts since thin films can be deposited with near atomic layer precision and can exhibit the planar surfaces with various degrees of intermixing across the interfaces. A series of Fe/Ni and Ti/Nb multilayers with bilayer repeat distances of 4 nm have been sputter-deposited onto n-doped Si [001] substrates. The multilayers were anular focus ion beam milled into the interfaces oriented with the bilayer chemical modulations parallel and distances of 4 nm have been sputter-deposited onto n-doped Si [001] interfaces. A series of Fe/Ni and Ti/Nb multilayers with bilayer repeat distances of 4 nm have been sputter-deposited onto n-doped Si [001] substrates. The multilayers were annular focus ion beam milled into the required needle-shaped geometry for the atom probe analysis with the film substrates. The multilayers were annular focus ion beam milled into the interfaces oriented with the bilayer chemical modulations parallel and distances of 4 nm have been sputter-deposited onto n-doped Si [001] interfaces. A series of Fe/Ni and Ti/Nb multilayers with bilayer repeat distances of 4 nm have been sputter-deposited onto n-doped Si [001] substrates.

4:40pm AS-TuA9 Atom Probe Tomography and Spectroscopic Analysis of Wide Bandgap Nanostructures, N. Dawahre, G. Shen, W. Baughman, S. Balci, S. Wilbert, N. Harris, L. Butler, S. Kim, P. Kung, University of Alabama

Wide bandgap ZnO based semiconductors are materials of great importance in regulating the levels of the phosphatidylinositoltriphosphate phosphatase activity as an interfacial enzyme at the plasma membrane-cyttoplasm boundary. Acting as an antagonist to phosphoinositide-3-kinase (PI3K) in cell signaling, it is deleted in many human cancers. Despite its importance in regulating the levels of the phosphatidylinositoltriphosphate (PI(3,4,5)P3), there is little understanding of how PI3K binds to membranes, is activated and then acts as a phosphatase. The interaction of the protein with membranes is highly dynamic and is at least partially controlled by the in-plane fluidity of the bilayer. PI3K function requires multiple, lipid-specific interactions with the target membrane. These interactions regulate enzyme activity as well as lateral and subcellular distribution of the enzyme. From studies of the membrane association of PI3K under well-defined conditions in model systems, we report recent insights in the structural and functional basis for PI3K membrane binding and regulation.

We subsequently discuss the sample preparation techniques employed and the influence of various APT measurement parameters on the quality of the data. Atom probe tomography (APT), which combines a field ion microscope and a time-of-flight mass spectrometer, is an analytical technique which is unmatched in identifying composition at the atomic scale and in 3D. However, proper interpretation of the APT data required thorough analysis of the mass spectra. Data analysis was also carried out in correlation with the nanowire synthesis conditions (e.g. carrier gas and dopant) as well as with other characterization techniques aimed at assessing the nanowire optical and electrical properties. These included high resolution transmission electron microscopy along with energy dispersive spectroscopy mapping, confocal Raman spectroscopy and imaging, confocal photoluminescence and imaging, as well as terahertz time domain spectroscopy.

5:00pm AS-TuA10 Probing 3D-Semiconductor Structures, Vandervoort, IMEC, Belgium

The trends in advanced semiconductor devices and technologies call for the ability to probe compositional and impurity distributions with a depth resolution approaching near-atomic resolution. At the same time, quantitative interpretation is of utmost importance in particular in multilayer structures and at interfaces such as local silicide composition and phase, interfacial interactions high k metal gate, etc.. At the same time one must admit that the analysis of nanoscale devices and their analogues in sample structures is an ongoing research challenge. We therefore conducted an experimental study of Finfets, nanowires necessitates techniques which provide 3D-resolution. In this paper we will discuss recent approaches in extracting local 2D, 3D information on dopant distributions, carrier distribution, defects in thin dielectrics, phase and composition analysis, detection of small voids in ultra narrow interconnect lines (> 15 nm) as well as in (large, > 50 micron) Cu-interconnects (TSV’s). For nanoscale 1D and 2D characterization we presently rely on concepts such as EXLE-SIMS for ultra high depth resolution dopant profiling, SSRM for 2D- carrier analysis, C-AFM for dielectrics, STM (+ELNES, EDX, HAADF) for quantitative composition analysis. 2D- and 3D-characterization represents a serious challenge and one must rely on concepts like the tomographic Atom Probe to extract 3D-composition analysis on the nm-scale or Tomographic TEM. We will show examples for both cases and address issues of Atom Probe in particular as the latter, although very appealing, does contain many artifacts as well. The latter are linked to sample preparation, laser-tip interaction, and reconstruction artifacts as well as issues inherently linked to instrument performance (mass resolution, sensitivity...). In this paper we will discuss recent approaches in extracting local 2D, 3D information on dopant distributions, carrier distribution, defects in thin dielectrics, phase and composition analysis, detection of small voids in ultra narrow interconnect lines (> 15 nm) as well as in (large, > 50 micron) Cu-interconnects (TSV’s).

For back-end applications, the detection voids in narrow Cu-lines (15-30 nm) is a big challenge. An approach used during the deposition of Cu TSV’s requires sophisticated ion milling approaches (based on machine FIB, sicle and view) as an effective milling times and surface topography evolution (curing) can hide the required information.

Biomaterial Interfaces Division
Room: 105 - Session BI-TuA

Protein-Membrane Interactions
Moderator: L.J. Gamble, University of Washington

4:00pm BI-TuA7 Membrane Binding, Structure and Regulation of the PTEN Phosphatase, M. Lische, Carnegie Mellon University and National Institute of Standards and Technology

PTEN Phosphatase and tensin homologue deleted on chromosome 10 (PTEN) is an important regulatory protein and tumor suppressor that performs its phosphatase activity as an interfacial enzyme at the plasma membrane-cyttoplasm boundary. Acting as an antagonist to phosphoinositide-3-kinase (PI3K) in cell signaling, it is deleted in many human cancers. Despite its importance in regulating the levels of the phosphatidylinositoltriphosphate (PI(3,4,5)P3), there is little understanding of how PTEN binds to membranes, is activated and then acts as a phosphatase. The interaction of the protein with membranes is highly dynamic and is at least partially controlled by the in-plane fluidity of the bilayer. PTEN function requires multiple, lipid-specific interactions with the target membrane. These interactions regulate enzyme activity as well as lateral and subcellular distribution of the enzyme. From studies of the membrane association of PTEN under well-defined conditions in model systems, we report recent insights in the structural and functional basis for PTEN membrane binding and regulation.
interactions remain unknown. Antigens as well as the viral lipid membrane. However, the mechanistic antigen designs induce polyreactive antibodies that recognize MPER interactions and antigen localization at the membrane interface, with the ultimate vision of developing immunogens. Recent imaging studies have shown that induction of antibodies that avidly bind the gp41-MPER antigen is not sufficient for neutralization. Rather, it is required that antigen designs induce polyreactive antibodies that recognize MPER antigens as well as the viral lipid membrane. However, the mechanistic details of how membrane properties influence NAb-lipid and NAb-antigen interactions remain unknown. Methods: To understand how membrane properties contribute to 2F5/4E10 membrane interactions, we engineered biomimetic supported lipid bilayers (SLBs) and have developed a surface plasmon resonance (SPR) spectroscopy based assay that monitors antibody binding to thiol monolayers, which mimic salient surface chemical properties of lipid membranes. Our results showed that 2FS and 4E10 bound preferentially on charged and hydrophobic thiol surfaces. This supports the theory that NAbs interact with lipid head groups before embedding into hydrophobic tail regions. We have also engineered supported lipid bilayers (SLBs) whose compositions mimic both the host cell membrane and the HIV-1 envelope. These SLBs have planar surfaces that facilitate quantitative surface-characterization techniques such as high-resolution scanning-probe imaging, detection of fluorescence recovery after photobleaching, and neutron reflection measurements. Using these characterization techniques we have begun to i) visualize domains of lateral membrane organization; ii) determine SLB domain diffusivity; iii) determine differences in adhesion force (surface energy) of domains; and iv) correlate these membrane properties with NAb-binding and NAb/antigen localization. Our research is significant in that it provides a biologically relevant system to screen interactions of lipid-reactive antibodies with a broad range of diagnostic tools. Because current 2FS/4E10 immunogens have not yet elicited antibodies with the required membrane reactivity it is important to reveal the role of guiding immunogens designs. Recent imaging information will elucidate how membrane properties can enhance antigen recognition and thus enable the design of next generation HIV-1 immunogens.

5:00pm Bi-TuA10 Interactions between the Norovirus and Glycosphingolipids Studied with Cell Membrane Mimics. M. Bally, Chalmers University of Technology, Sweden, G. Larson, University of Gothenburg, Sweden, F. Höök, Chalmers University of Technology, Sweden

The determining initial step of viral infection is mediated by highly specific recognition events between the viral shell and ligands on the host cell surface. Detailed understanding of virus-membrane interactions is therefore of central importance to the development of new antiviral therapies, new vaccines and high-performance diagnostics platforms. In this context, assays based on virus-membrane mimicking consideration platforms are particularly valuable to offer the possibility to study interactions between controlled ligand mixtures with surface-sensitive techniques, while presenting the ligand in a more native configuration. Potentially relevant characteristics such as membrane fluidity, ligand mobility and their ability to organize into microdomains can be preserved [1].

In this work we investigate the interactions between norovirus capsids and phospholipid bilayers containing glycosphingolipids (GSL). The norovirus is well known as the major causative agent of acute viral gastroenteritis and its human target cells and the precise mechanism for viral entry are still poorly understood.

In a first example, we identify galsactosylceramide - a major glycosphingolipid in the small intestine- as a ligand for Norovirus-like particles (VLP) from the Dijon strain. Quartz Crystal Microbalance and Atomic Force Microscopy studies on GaCer-containing supported lipid bilayers reveal that a clustered arrangement of the glycosphingolipids plays a crucial role in promoting a firm attachment of the pathogen to the lipidic membrane, most likely via the establishment of multiple contacts between the particle and the membrane [2].

We further investigated the interaction between individual fluorescent GSL liposomes and surface immobilized VLPs with single virus particle sensitivity. Besides representing the ultimate sensitivity for diagnostics purposes, our method makes it possible to study weak interactions. Kinetics analysis of vesicle residence times over large time scales reveals a highly heterogeneous behavior and yields information on multivalency, on the presence of domains and the role of cell-membrane curvature.

As exemplified here, simplified membrane models have a unique potential in providing fundamental understanding on the contribution of individual components to complex biological processes [3].

References:

5:20pm Bi-TuA11 Binding of C-reactive Protein to Lipoprotein Nanoparticle Mimics: A Gel Electrophoresis Study. M.S. Wang, S.M. Reed, University of Colorado Denver

C-reactive protein (CRP) is an acute phase serum protein involved in inflammation that recognizes pathogens, and activates complement. Because CRP levels can increase by over 1000-fold from basal levels within 72 hr in response to inflammation, it has been used as a biomarker to predict the risk for cardiovascular disease (CVD). In addition, CRP has been shown to bind oxidized low density lipoprotein (oxLDL) through the phosphatidylcholine (PC) headgroup that is exposed on the surface of oxLDL and has been found to co-localize with atherosclerotic lesions. It is suggested that CRP was deposited in these lesions when engulfed by macrophage while it is still bound to oxLDL. Because LDL and the other lipoprotein particles (LPPs) have diameters (d) in the nanometer range, they can be considered as biological nanoparticles. In fact, LPPs are classified into four major categories according to their size and density: high density lipoprotein (d=8-13nm), LDL (d=20-30nm), intermediate density lipoprotein (d=10-40nm), and very low density lipoprotein (d>40nm). Moreover, the physiological functions of LPPs are greatly influenced by the size. For example, the presence of sdLDLs (d<25.5nm) are associated with increased risk of coronary artery disease (CAD) and diabetes; while increased levels of HDL are considered to be atheroprotective. Therefore, LPP profiling has emerged as a tool to more accurately assess the development of metabolite risk factors such as CAD. While the PC moieties on oxLDL are likely ligands for CRP recognition, it is still unclear why CRP binds only oxLDL although PC is expressed on native LDL and on all cell membrane. In this work, we correlated the effects of particle size (i.e. membrane curvature) to CRP binding using LPP mimics. To this end, we engineered LPP mimics using lipid-coated gold nanoparticles (PC-AuNPs) to explore the influence of LPP size on CRP binding. Binding analysis of CRP to the mimics was performed using gel electrophoresis (GE). The migration of CRP and PC-AuNP was directly visualized after electrophoresis, and the presence of CRP was confirmed using Western blots. The overlapping bands from the gel and Western blot confirmed that CRP bound to the PC-AuNPs and co-migrated during GE. Together, we demonstrated that 1) PC-AuNPs are size-separable, 2) the lipid layer around the PC-AuNP remained intact during GE, and 3) CRP binds to a relevant LDL-sized PC-AuNP mimic. This is, to our knowledge the first use of GE to separate lipid-coated nanoparticle and to evaluate non-covalent binding of protein-nanoparticle interactions.

Biofabrication and Novel Devices Focus Topic Room: 105 - Session BN-TuA

Biofabrication Methods and Devices
Moderator: L. Gamble, University of Washington

2:00pm BN-TuA1 Microengineered Hydrogels for Stem Cell Bioengineering and Tissue Regeneration. A. Khademhosseini, Brigham and Women’s Hospital, Harvard Medical School, MIT, and Harvard University

INVITED Micro- and nanoscale technologies are emerging as powerful tools for controlling the interaction between cells and their surroundings for biomedical studies, tissue engineering, and cell-based screening. In addition, hydrogel biomaterials have been increasingly used in various tissue engineering applications since they provide cells with a hydrated 3D microenvironment that mimics the native extracellular matrix. In our lab we have developed various approaches to merge microscale techniques with hydrogel biomaterials for directing stem cell differentiation and generating complex 3D tissues. In this talk, I will outline our work in controlling the cell-microenvironment interactions to direct the differentiation of stem cells. In addition, I will describe the fabrication and the use of microscale hydrogels for tissue engineering by using a ‘bottom-up’ and a ‘top-down’ approach. Top-down approaches for fabricating complex engineered tissues involve the use of miniaturization
The features of the scaffolds lead to the development of a geometrically defined neuronal network when applied as platforms in a primary-neuron culture. Cell morphology and expression of the neuronal markers were characterized by fluorescent microscopy.

Electronic Materials and Processing Division
Room: 210 - Session EM-TuA

High-k Dielectrics for MOSFETs Part 2
Moderator: A.C. Kummel, University of California San Diego

2:00pm EM-TuA1 High Mobility Channel Materials and Novel Devices for Scaling of Nanoelectronics beyond the Si Roadmap, M. Heyns, IMEC, Belgium

The introduction of high-k dielectrics and metal gates in advanced CMOS has opened the door to Ge and III-V compounds as potential replacements for Si to further increase the device performance. Using MOCVD the selective area growth of low-defect InP and InGaAs layers in submicron trenches on Si was demonstrated. These virtual Ge-III/V substrates can be processed in a standard CMOS line. Short channel Ge pMOS devices with high drive currents were fabricated. Strain engineering using GeSn source/drain allows to boost the performance of these devices so that they can outperform their strained Si counterparts. One of the key problems in developing III/V devices is the near midgap Fermi level pinning associated with the high density of defect states present at the high-k/III-V interface. The origin of these states is still under debate but there are clear indications that there exists a strong relationship with native antiposite point defects. Various sulfide and other treatments were investigated to passivate the surface. The measured distribution of interface states and border traps on typical III/V MOS structures has some special consequences on the electrostatic operation of different transistor designs. Since inversion mode devices do not seem to be the appropriate choice for III/V based logic applications, other device types have been explored. The Implant-Free Quantum Well (II-QW) device enables VLSI-compatible processing by self-aligned source/drain definition. Strained Implant Free Quantum Well Ge-based pFETs show excellent short channel control and record drive currents. The concept was also used to demonstrate high mobility n-channel InGaAs devices. For III/V pMOS devices GaSb is at present the material of choice. Very encouraging results have been obtained on direct heteroeptaxy of GaSb epilayers on InP(001) combined with in-situ deposition of an AlOx high-k gate dielectric. The introduction of these advanced materials also allows the development of new device concepts that can fully exploit the properties of these new materials. Tunnel-FETs, where the III/V material may be either introduced only in the source or in the complete device, can provide superior performance at lower power consumption by virtue of their improved subthreshold behavior, allowing to reduce the supply voltages. Vertical surround gate devices can be produced from III/V nanowires directly grown on silicon, allowing the introduction of a wide range of III/V materials and functionalities on Si. This illustrates some of the possibilities that are created by the combination of new materials and devices to allow scaling of nanoelectronics beyond the Si roadmap.

2:40pm EM-TuA3 Local Profile of the Dielectric Constant Near the Oxygen Vacancy in the GeO2 Films, J. Nakamura, M. Tamura, The University of Electro-Communications (UEC-Tokyo), Japan

Ge-based metal-oxide-semiconductor (MOS) devices are focused as complementary-MOS devices for the next-generation in the post-Si technology. However, the dielectric properties of GeO2 gate ultrathin films have not been clarified yet in detail. Our purpose is to clarify the spatial variation of the local dielectric constant for the GeO2 thin films using first-principles ground-state calculations in external electric fields [1,2]. In particular, we reveal the local profile of the dielectric constant near the oxygen vacancy in the film, focusing on the crystal phase dependence. We have adopted quartz (0001) and rutile (001) films with/without oxygen vacancies, in which Ge atoms at the topmost surfaces are terminated by H atoms. We have evaluated the optical and the static dielectric constants that are attributed to the electronic polarization and both the lattice and electronic polarizations, respectively.

From the local profile of the dielectric constants for the ideal films, it has been clarified that the dielectric constants change gradually from the surface and approach an atomic constant at the center of the film. Such features have also been confirmed for the Si and SiO2 films [1,2]. The dielectric constant for the defective model of the quartz film becomes larger locally “at” the oxygen vacancy site compared with that for the ideal model, but at “adjointing” oxygen sites to the vacancy for the rutile film. Such features
stem from the difference in the fashion of the chemical bonding between Ge and O atoms: The dielectric constant for the defective quartz model becomes large at the vacancy site where the vacant Ge-Ge bond is formed. For the rutile, on the other hand, the Ge-O bondings surrounding the vacancy site are softened because of their less ionic character, which results in the larger displacement in external electric fields, leading to the larger lattice polarization around the vacancy.


In this work, the effect of dielectric post deposition anneal (PDA) passivated InP is thermally stable up to ~460° C. However, little work has been performed to understand the thermal stability of the high-k/InP interface. In this work, the effect of dielectric post deposition anneal (PDA) passivated InP is thermally stable up to ~460° C. However, little work has been performed to understand the thermal stability of the high-k/InP interface.

3:00pm EM-TuA4 Two Step Passivation and ALD Monolayer Nucleation on Ge(100). T. Kaufman-Osborn, J.S. Lee, K. Kianjait, W. Melitz, A.C. Kummel, University of California San Diego, A. Delabie, S. Sioncke, M. Caymax, G. Pourtois, IMEC, Belgium

Germanium is a promising channel material for next generation MOSFET. The best method to passivate Ge(100) is to form a layer of GeO2 free of Ge suboxides, using high pressure O2 or O. However, there are three challenges: (1) it is difficult to keep a stoichiometric GeO2 monolayer (ML) at elevated temperatures, (2) the thermal oxidation process creates a rough interface degrading mobility at high field, and (3) scaling the passivation layer to only 1 ML is a challenge. This study presents a process to form a ½ ML of Ge-H and ½ ML of Ge-OH bonds without disrupting the Ge(100) surface. In-situ scanning tunneling microscopy (STM), in-situ scanning tunneling spectroscopy (STS), and in-situ X-ray photoelectron spectroscopy (XPS) were employed to determine the atomic and electronic structure of the passivation monolayer.

Using a differentially-pumped H2O dosing system, an ordered, full monolayer of H2O chemisorption sites on Ge(100) was formed with a low density of unreacted dangling bonds at 300K. STS data showed that the Ge-H and Ge-OH sites removed the bandgap states from the Ge(100) dangling bonds. Annealing the surface between 20°C and 250°C gradually decreased the coverage of H2O sites. However, even at 300K, the H2O surface is highly reactive to metal alkyls (TMA) since it contains a half monolayer of Ge–OH which catalyzes the breaking of Al-CH3 bonds thereby inducing the formation of Al-O bonds, and the Ge sites block ALD ligand chemisorptions. STM experiments showed that the H2O chemisorbed Ge surface provides a half monolayer of nucleation centers with approximately 0.5 nm spacing for TMA dissociative chemisorption at 300K. High resolution XPS experiments indicated that the highly reactive Ge–OH bonds were converted to thermally stable Al-O bonds. Furthermore, passivating the surface with H2O prior to TMA dosing doubles the aluminum coverage compared to the TMA only dosed Ge(100) surface. The higher nucleation density from the two step functionalization process, TMA + H2O, should be favorable for pinhole reduction. DFT calculations are consistent with the data showing TMA reaction with either –Ge-H and –Ge-OH at the interface. The reaction of TMA with H2O provides a low activation barrier and higher exothermicity (-41.4 kcal/mol) compared to TMA reaction on the –Ge-H site (-10.8 kcal/mol). The calculation is consistent with the key to full monolayer nucleation, the formation of a full monolayer of Ge-OH chemisorption sites, which is being studied with HOOH dosing.

4:00pm EM-TuA7 Bilayer High-k Gate Stacks on Ge and InGaAs. P.C. McIntyre, Stanford University

In this work, the effect of dielectric post deposition anneal (PDA) passivated InP is thermally stable up to ~460° C. However, little work has been performed to understand the thermal stability of the high-k/InP interface. In this work, the effect of dielectric post deposition anneal (PDA) passivated InP is thermally stable up to ~460° C. However, little work has been performed to understand the thermal stability of the high-k/InP interface.


Due to high defect density between III-V semiconductors and high-k dielectrics, buried channel structures with InP barrier layers are being considered for CMOS applications1,2. It has been observed that sulfur passivated InP is thermally stable up to ~460° C3. However, little work has been performed to understand the thermal stability of the high-k/InP interface.

In this work, the effect of dielectric post deposition anneal (PDA) on InP MOS capacitors with high-k dielectrics was studied. Temperatures above 450° C result in an increase of the interface trap density.

MOS capacitors were fabricated on both n-type and p-type InP substrates with HfO2 and Al2O3 dielectric. Trimethyl aluminum (TMA) and water by (TDMA-Hf precursor) and water by in-situ monochromatic X-ray photoelectron spectroscopy (XPS). The "clean up" effect is examined and compared to the results for Al2O3 on InP. The significant reduction of low activation barrier and higher exothermicity compared to TMA reaction on the –Ge-H site (-10.8 kcal/mol). The calculation is consistent with the key to full monolayer nucleation, the formation of a full monolayer of Ge-OH chemisorption sites, which is being studied with HOOH dosing.

5:00pm EM-TuA10 ALD Half Cycle Study of HIO3 on InP by In Situ XPS. H. Dong, D.M. Zhernokletov, B. Brennan, J. Kim, R.M. Wallace, University of Texas at Dallas

InP attracts significant attention as a high mobility channel material for Metal-Oxide-Semiconductor Field Effect Transistors (MOSFETs) with many excellent electrical measurements reported on InP based devices. It is well known that one of the electrical limitations of surface channel III-V MOSFETs comes from the interface defects generated from surface/interface oxidation between high-k dielectrics and III-V materials, which cause Fermi levels to shift[1]. InP is also considered as a potential buffer layer material for quantum well FETs, which is in direct contact with a gate dielectric[2]. To examine the interfacial chemistry for high-k/InP, we present a half cycle Atomic Layer Deposition (ALD) study on native oxide and wet chemically treated InP (100) samples with Hf (TDA-Hf precursor) and water by in-situ monochromatic X-ray photoelectron spectroscopy (XPS). The "clean up" effect is examined and compared to the results for Al2O3 on InP. The significant reduction of low activation barrier and higher exothermicity compared to TMA reaction on the –Ge-H site (-10.8 kcal/mol). The calculation is consistent with the key to full monolayer nucleation, the formation of a full monolayer of Ge-OH chemisorption sites, which is being studied with HOOH dosing.

5:00pm EM-TuA11 Nonvolatile Memristive Nano-Crossbar Switches in Pt/Ta2O5/Cu Solid Electrolytes. P.R. Shrestha, K.P. Cheung, National Institute of Standards and Technology (NIST), H. Baumgart, Old Dominion University

Memristive devices operate by changing resistance from high (Roff) to low (Ron) values in response to an applied voltage. Despite a tremendous amount of work in the scientific literature, the actual underlying switching mechanism has yet to be fully explained. Few studies, most of which lack measurement details, have reported on the transient current response and high speed switching characteristics of memristive devices. The papers suggest the presence of the variety of active current and thermal dissolution of the low resistance filament while switching OFF. The switching ON has been attributed to the movement of the metal ions towards the cathode and being neutralized by the electrons to form the metal filament. Transient current measurements for switching ON have not yet been analyzed in detail in the literature. Another crucial problem in measurements of these devices is the difficulty in designing a nanoscale crossbar switch, and such two terminal "memristive" devices are being considered for next-generation non-volatile memory due to the inherent simplicity, scalability and low cost (1). Additionally, these devices show potential to replace static random access memory (SRAM) as high performance switches for reconfigurable devices.

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These results to X-ray photoelectron spectroscopy (XPS) analysis will be presented. This work is sponsored by SRC FCRP MARCO Materials Structures and Devices Center and the National Science Foundation.


devices is due to the preferred values of \( R_{\text{on}} < 1 \Omega \) and \( R_{\text{off}} > 1 \Omega \). Thus the key obstacle preventing fundamental understanding has been the lack of reliable and accurate measurement of the transient response while switching ON (from \( R_{\text{off}} \) to \( R_{\text{on}} \)). In this work, we developed a new measurement capability that enables reliable and accurate investigation of the transient switching response, monitoring change from low \( R_{\text{on}} \) to high \( R_{\text{on}} \).

In order to address the need to reliably and accurately monitor the device I-V transient response extremely fast, we have designed an amplifier with low-gain and high bandwidth (1.7 GHz) to accommodate much faster (and realistic) “programmable” voltage pulses. For switching purposes, this amplifier allows for fast transient current monitoring during programming with proficiency but not limited to follow 2 nsec of pulse rise time. The high gain bandwidth of the amplifier allows us to monitor change from low current (\( R_{\text{off}} \)) to high current (\( R_{\text{on}} \)) accurately.


5:40p.m. EM-TuA12 Metrology for Interfaces and Mass Transport in C-MOS Related Nanofilms; A. Herrera-Gomez, A. Sanchez-Martinez, O. Ceballos-Sanchez, M.O. Vaquez-Lepe, CINVESTAV-UNidad Queretaro, Mexico. P. Lysaght, SEMATECH

Interface layers play a fundamental role in determining the electrical properties of CMOS devices because their thicknesses are of a magnitude comparable to that of the dielectric layers currently employed. The main (top) techniques traditionally used for characterizing the chemical depth profile of MOS structures have been XPS-Sputter and Back-Side TOF-SIMS. However, they lack the appropriate resolution to characterize the thickness and composition of nano or sub-nano layers. Due to the lack of appropriate metrology methods, the structure of interface layers is usually assessed indirectly through their effect on the device’s capacitance. Another important issue in the processing of MOS devices is the diffusion control of chemical species. To quantify to what extent, observe displacements on the order of 2 or 3 nm of low concentration elements with those techniques is close to impossible. The semiconductor industry will greatly benefit from a metrology method capable of: (a) characterizing the thickness and composition of the various layers constituting a MOS device, including the interface layers; (b) assessing the effect of process driven diffusion of various critical chemical species present in the film. There is a growing consensus that X-ray Photoelectron Spectroscopy (XPS), specifically Angle-Resolved XPS (ARXPS), has the appropriate chemical and depth resolution for assessing the chemical depth profile of films between 0 and 8 nm. The precise methodology for applying the technique, however, varies widely among different groups. In many cases the analysis algorithms of ARXPS data are highly susceptible to noise. Because of this, ARXPS is frequently regarded as qualitative techniques. In this talk we briefly describe a robust ARXPS analysis methodology that minimizes the sensitivity to noise. This methodology has been successfully applied to characterize various systems. One example that will be addressed regards the failure mechanism for the degradation of the electrical performance of TiN/HfO2/InGaAs devices during thermal processing. We investigated the change on the structure caused by annealing, such as the possible formation of As, Ga or In oxides, accumulation of these elements in the dielectric, formation of metallic arsenic, and/or any other change on the structure that could be correlated to device degradation. ARXPS experiments were performed on those samples and analyzed using the robust methodology. One important finding was that indium diffuses through the dielectric all the way into the metallic layer upon annealing.

Energy Frontiers Focus Topic Room: 103 - Session EN+NS-TuA

Nanostructured Materials for Thermophotovoltaics, Thermoelectrics & Plasmonics

Moderator: P. Nagpal, Los Alamos National Laboratory

2:00p.m. EN+NS-TuA1  Thermoelectric and Plasmonic Crystals as a Route to Photovoltaics?; D.J. Norris, ETH Zurich, Switzerland INVITED

Patterned metallic films allow the generation and manipulation of specific electromagnetic waves known as surface plasmons that propagate along a metal interface. Because these waves allow the concentration of light in nanoscale volumes, they have implications for fundamental phenomena as well as applications such as imaging, sensing, and solar cells. Thus, the field of plasmonics has arisen to study and utilize surface plasmons. While light is typically used to create these waves, in some cases it would be more convenient to have a non-optical source for surface plasmons. One route is thermal excitation (i.e., heat). Here, we will explore hot plasmonic structures for obtaining new optical behavior. For example, we will examine films patterned with series of concentric grooves (a bull’s eye pattern). We show that, when heated, these films can emit light that is amazingly narrow, both in terms of its spectrum and its angular divergence. Thus, a simple metallic foil can generate a highly directional beam of monochromatic light by a thermal process. This effect has implications for creating efficient thermophotovoltaic devices, which convert heat into electricity. During these studies, we also developed a simple, high-throughput method for obtaining ultrasmall patterned metal films. Previously, roughness in such films has hindered the field of plasmonics. Therefore, our approach solves a critical problem and allows many high-quality plasmonic structures to be available for the first time.

2:40p.m. EN+NS-TuA3 Nanocrystal Assemblies: A Modular Approach to Materials Design; D.V. Talapin, University of Chicago INVITED

Colloidal nanocrystals can be the advent of new classes of crystalline inorganic semiconductors with the size-tunable electronic structure and inexpensive solution-based device fabrication. Single- and multicomponent nanocrystal assemblies, also known as superlattices, provide a powerful general platform for designing two- and three-dimensional solids with tailored electronic, magnetic, and optical properties. Unlike atomic and molecular crystals where atoms, lattice geometry, and interatomic distances are fixed entities, the nanocrystal arrays represent ensembles of “designer atoms” with potential for tuning their electronic structure and transport properties. Generally speaking, nanocrystal assemblies can be considered as a novel type of condensed matter, whose behavior depends both on the properties of the individual building blocks and on the particle interaction exchange interactions.

The ability to assemble precisely engineered nanoscale building blocks into complex structures is opening the door to materials where components and functionalities can be added and removed in a finely controlled manner. I will show how self-assembly of nanocrystals can lead to a palette of unprecedented phases including superlattices isostuctural with the Archimedean tilings and dodecaogonal quasicrystals. Efficient charge transport is crucial for performance of nanocrystal-based electronic and optoelectronic devices. The insulating nature of surface ligands traditionally used for nanocrystal synthesis results in the poor electronic coupling between individual nanocrystals. To facilitate charge transport we introduced strong organic ligands for colloidal nanocrystals. These ligands, namely metal chalcogenide complexes, can be applied to a broad range of inorganic nanomaterials. I will demonstrate the power of this approach on several examples of prospective electronic, thermoelectric and photovoltaic materials.

4:00p.m. EN+NS-TuA7 Photonic and Plasmonic Crystals for Thermophotovoltaics and Energy Conversion; R. Biswas, Iowa State University & Ames Laboratory - US DOE INVITED

Photonic and plasmonic crystals that have various energy-related applications will be discussed. Metallic plasmonic crystals consisting of an array of nano-holes or nano-pillars on common substrates, with pitch at infrared length scales, have very sharp absorption properties at infrared wavelengths. Such arrays have diverse applications to sensors and thermophotonic applications. The absorption, thermal emission, and angular characteristics from these plasmonic arrays will be described with rigorous scattering matrix simulations and compared to measurements. I will also survey the rich physics underlying plasmonic nano-arrays at optical length scales, and a few energy related applications.

4:40p.m. EN+NS-TuA9 Molecular and Hybrid Solution Processible Thermoelectric Materials; R.A. Segalman, S. Yee, University of California, Berkeley, N. Coutts, J. Urban, Lawrence Berkeley National Laboratory INVITED

Thermoelectric materials for energy generation have several advantages over conventional power cycles including lack of moving parts, silent operation, miniaturizability, and CO2 free conversion of heat to electricity. Excellent thermoelectric efficiency requires a combination of high thermopower (S, V/K), high electrical conductivity (σ, S/cm), and low thermal conductivity (κ, W/mK). To date the best materials available have been inorganic compounds with relatively low earth abundance and highly complex, vacuum processing routes (and hence greater expense), such as Bi2Te3. Molecular materials and hybrid organic-inorganic bring the promise of inexpensive, solution processible, mechanically durable devices. While highly conductive polymers are now common place, they generally demonstrate low thermoelectric performance. Our work on such polymers suggests that nanostructuring of organics allows them to act as thermionic filters between inorganic junctions which can lead to enhanced thermoelectric properties. We have taken inspiration from this fundamental
understanding to design material systems in which we combine a high electrical conductivity, low thermal conductivity polymer with a material that contributes high thermopower. Additionally, the work functions of the two materials are well-aligned which introduces the possibility of thermionic filtering at the interface and an additional boost to the power factor. The combination of these effects results in a new hybrid, solution processible material with a thermoelectric figure of merit approaching those of Bi$_2$Te$_3$. In this talk, I will discuss both the use of thermoelectric measurements to gain insight into molecular junctions and how this insight translates to design principles for polymer and hybrid thermoelectrics.

5:20pm EN+NS-TuA11 Semiconductor Nanowire Networks as Thermoelectric Platforms, A.J. Lohn*, University of California Santa Cruz, E. Coleman, G.S. Tompa, Structured Materials Industries, Inc., N.P. Kobayashi, University of California Santa Cruz

Current energy production mechanisms for electrical power and transportation are plagued by inefficiencies which results in most of the energy source being lost as heat. In most cases that heat is found in the form of low-grade heat with temperatures below approximately 200 degrees C. Unfortunately, typical methods such as the Rankine cycle for converting heat to electricity suffer from poor efficiency for low-grade heat. Direct thermoelectric conversion is currently struggling to match the efficiency of the Rankine cycle at high temperatures but offers advantages in terms of reduced maintenance and form-factor which enable energy scavenging in places such as the exhaust line of a vehicle where larger systems could not be implemented.

Dominated by recent progress in nanostructured materials, the unitless thermoelectric figure of merit ZT has increased to well beyond 1 such that efficiencies are reaching a range which makes them cost effective. Typically thermoelectric materials include elements such as lead or tellurium which are toxic and rare therefore alternative materials are being sought. Recent progress in silicon nanowire thermoelectric has shown a reduction in thermal conductivity, and therefore an increase in ZT of two orders of magnitude, making them viable candidates in the thermoelectric marketplace. Decreased cost and toxicity of silicon as compared to conventional thermoelectric materials make it an attractive candidate but to date nearly all studies on thermoelectricity of nanowires have focused on nanowires in isolation. Our platform based on interconnected 3-dimensional nanowire networks grown directly on metallic substrates provides large area thermoelectric modules capable of scavenging low-grade heat for low cost. The materials properties comprising ZT: thermal conductivity, electrical conductivity and Seebeck coefficient will be discussed for undoped, p-type and n-type silicon nanowire networks with particular emphasis on electrical conductivity and Seebeck coefficient within the temperature range of low-grade heat.

5:40pm EN+NS-TuA12 Diamond as an Electrode Material for the Direct Conversion of Thermal to Electrical Energy through Thermionic Emission, W.F. Paxton, J.L. Davidson, W.P. Kang, Vanderbilt University

Introduction

Thermionic emission conversion is a candidate technology for the efficient conversion of thermal energy directly to electrical energy. In a thermionic converter, thermally excited electrons are emitted from the surface of a heated cathode into a vacuum gap. These electrons are then collected by a cooler anode and driven through an external load back to the cathode. The thermionic emission current density of a heated cathode can be described by the Richardson Equation (Eq. 1).

\[ J = A T^2 e^{\frac{-\Phi}{kT}} \]

where: J: Thermionic emission current density (A/cm$^2$); A: Richardson constant (A/cm$^2$·°K); T: Temperature (K); Φ: Material’s work function (eV); and k: Boltzmann constant (eV/°K)

It can be seen from Eq. 1 that a material with a lower work function, Φ, can achieve higher current densities at lower temperatures than a material with a higher work function, which implies lower work function values translate into better energy converters. Prior attempts to construct efficient thermionic converters were limited by available materials with work function values ranging from 3.5eV to 5eV requiring extreme cathode temperatures in order to achieve useable output power values. In this study, the thermionic emission properties of nitrogen-incorporated diamond films are examined as a potential electrode material to enhance the efficiency of such a device.

Experimental

Polycrystalline diamond films were synthesized on molybdenum substrate via Microwave Plasma-Enhanced Chemical Vapor Deposition (MPCVD). The nitrogen and hydrogen partial pressures were 0.3% and 2%, respectively, the power density was 500 W/cm$^2$, and the nominal pressure was 50Torr. Scanning electron micrographs of the samples demonstrated uniform film coverage and a thickness of ~100μm. Thermionic emission characterization was performed in a vacuum environment with a base pressure of 1 x 10$^{-7}$ Torr. The diamond films were resistively heated and the temperature was constantly observed by a dual color pyrometer. Electron emission current was collected with an electrically isolated anode biased at a constant voltage of 100V positioned 0.5cm above the heated cathode.

Results and Discussion

Observation of the electron emission current above the noise level began at 600°C and increased exponentially with temperature up to 800°C. Analysis of this data demonstrated agreement with the Richardson equation with a correlation coefficient of 0.99. From this data, the nitrogen-incorporated diamond samples were determined to have a work function value less than 2eV which is considerably lower than previously mentioned materials. These results exhibit diamond’s potential as an interesting cathode material for a thermionic energy converter.

Energy Frontiers Focus Topic

Room: 108 - Session EN+TF-TuA

Thin Films for Solar Cells

Moderator: C.A. Wolden, Colorado School of Mines

2:00pm EN+TF-TuA Thin Film Silicon Approaches to Future Generations of PV Materials, P.C. Taylor, Colorado School of Mines

INVITED

Polycrystalline silicon (poly-Si) is considered to be a promising candidate for thin film PV, coupling the high quality crystalline Si technology with large area and low-cost manufacturing. Our initial studies [1] on poly-Si layers have shown grains extending through the whole thickness (1 μm) upon solid phase crystallization (SPC) of high growth rate plasma deposited amorphous silicon (a-Si:H) films. Furthermore, larger grains are promoted by an increase in the a-Si:H microstructure parameter R* [2], which represents the order (low R*)/disorder (high R*) in the matrix according to the Si-H bond distribution in mono-di-vacancies (~low stretching mode-LSM) and nano-sized voids (~high stretching mode-HSM), and it is quantified by the integrated IR absorption band ratio $I_{LSM}/I_{LS}$ of poly-Si. The SPC of a-Si:H follows the steps of incubation, nucleation and grain growth. With the purpose of providing insight on the crystallization process, this contribution addresses a detailed crystallization kinetic study of plasma deposited a-Si:H films by means of in-situ X-ray diffraction (XRD), a-Si:H films having R* in the range of 0.03-0.6, with an hydrogen content of 5-14 at. %, were annealed at 600 °C.
The medium range order (MRO) of the a-Si:H layers, quantified by the XRD line-width, and representing the most ordered regions in the matrix (up to 3 Å) from the mono-vacancies), is found to affect the incubation time \( t_{\text{inc}} \), in agreement with [3]: low \( R^* \) and high MRO promote a faster nucleation \( t_{\text{nuc}} \) in the range of 50-100 min, since the most ordered regions act as nucleation centers; as the structural disorder increases, the MRO decreases and the incubation step is delayed up to 450 min. However, for \( R^* > 0.3 \) and a hydrogen content above \( 9\% \), the incubation time unexpectedly decreases. Therefore, the MRO evaluations during the annealing step are studied. High \( R^* \) layers, characterized by hydrogen mainly bonded to nano-sized voids, are more prone to hydrogen out-diffusion upon annealing, as inferred by the quantitative decrease of the HSM mode with respect to the LSM mode. The hydrogen evolution is then followed by the rearrangement of the a-Si:H into more ordered regions, as witnessed by the increase of the MRO upon annealing, prompting a decrease in incubation time. In conclusion, next to the established role of the MRO, the nano-sized voids play also a role in the crystallization kinetics, as they affect the overall microstructure and medium range order upon annealing.


3:00pm EN+TF-TuA4 Deposition of Microcrystalline Silicon Thin Films by Radio Frequency PECVD using Voltage Waveform Tailoring S. Pouliquen, P.-A. Delatour, E.V. Johnson, J.-P. Booth, Ecole Polytechnique (Paris), France

This paper concerns the feasibility of depositing thin films of hydrogenated microcrystalline silicon (µc-Si:H) for photovoltaic applications by radio-frequency discharge using voltage waveform tailoring. Contrary to typically used sinusoidal waveforms, the films were deposited using asymmetric voltage waveforms, resembling “peaks” and “valleys”, composed of a fundamental frequency of 15 MHz and three harmonics. Such waveforms lead to an electrical asymmetry effect in the plasma, and a different ion bombardment energy on each electrode. For these experiments, we used a gas mixture of silane diluted in hydrogen (from 1-4%), a total flow rate less than 100 sccm, and a range of pressure from 100 up to 1000 mTorr. We show that the chemical, optical and structural properties - as measured by FTIR, spectroscopic ellipsometry, Raman scattering and profilometry - are strongly determined by the shape of the voltage waveform, and therefore the ion bombardment energy. We observe this effect on the Si-Si bond configuration (microcrystalline vs amorphous peaks in Raman scattering between 480 and 520 cm\(^{-1}\)) as well as on the Si-H bond configuration (FTIR peaks at 2000-2100 cm\(^{-1}\)). We examine the strength of the effect for varying process parameters, including pressure, total gas flow, voltage amplitude, and \( \text{SiH}_4/\text{H}_2 \) dilution ratio. For the range of parameters used, a growth rate for highly microcrystalline silicon of up to 3 Å/s was achieved without exceeding a pressure of 100 mTorr.

4:00pm EN+TF-TuA7 Amorphous and Nanocrystalline Silicon Thin Film Photovoltaic Technology on Flexible Substrates. B. Yan, A. Banerjee, J. Yang, S. Guha, United Solar Ovonic LLC

INVITED
Because of the rapid increase of energy demand and growing concern of environmental impact, renewable energy from photovoltaic (PV) has gained a great deal of attention in the last decade. Various PV technologies have been developed. However, solar panels using conventional crystalline silicon have dominated the market. Thin film silicon is one of the so-called second generation PV technologies. Nowadays, majority of thin film silicon PV products are made with hydrogenated amorphous silicon (a-Si:H) and amorphous silicon germanium (a-SiGe:H) alloys. The advantages of a-Si:H based technology are low cost, capability of large scale manufacturing, abounding of raw materials, and no environmental concerns. One disadvantage of a-Si:H PV technology is its low efficiency compared to solar panels made of crystal silicon and compound crystal thin film semiconductors. To resolve the low efficiency issue, significant effort has been made by the researchers. In order to use the solar spectrum effectively, multi-junction structures are normally used by incorporating a-SiGe:H in the bottom cell. In recent years, hydrogenated nano-crystalline silicon (nc-Si:H) has been used as a potential replacement of a-SiGe:H bottom cell in multi-junction structures. The pros of nc-Si:H are its stability under sunlight, high photocurrent capability, and no Ge-containing gases required in the process; the cons are thin intrinsic layer that needs high rate deposition and technical challenges for large-area deposition. United Solar has been heavily involved in a-Si:H and nc-Si:H based thin film silicon PV technology. We have made significant progress in efficiency improvements of a-Si:H and nc-Si:H multi-junction solar cells and modules. We have achieved (i) a 15.4% initial active-area (~0.25 cm\(^2\)) solar cell efficiency, (ii) an NREL measured stable total area (~0.25cm\(^2\)) efficiency of 12.5%, and (iii) NREL measured initial and stable module (~400 cm\(^2\)) efficiencies of 12.0% and 11.4%, which all set new record efficiencies achieved in a-Si:H, a-SiGe:H, and nc-Si:H multi-junction cell structures. Based on these achievements, we have started working on the development of roll-to-roll manufacturing technology for a-Si:H and nc-Si:H multi-junction structures on flexible substrates. We expect to launch 12% stable aperture area a-Si:H and nc-Si:H product in 2012. In this presentation, we will review the progress made by the community and challenges a-Si:H and nc-Si:H PV technology face.


INVITED
Hydrogenated amorphous and microcrystalline silicon based thin film solar cells have long shown great potential for various photovoltaic applications due to its better electricity generation under low light and high temperature, shorter energy payback time and greener manufacturing processes, as well as unique applications for BIPV. However, this technology hasn’t fully realized its market potential, as low module efficiencies at mass-production level being one of the major limiting factors. In this paper we report over 9% stabilized module efficiencies in volume production achieved on 5.7m \( \times \) 2 substrates from a-Si:H/nc-Si:H tandem junction solar modules, which reflects significant efficiency improvement by process tuning of silicon thin films, back contact, and device structure, etc. Several application case studies are also reported which demonstrate excellent field performance of ENN silicon thin film solar modules.


InGaN alloys has great potential in the field of photovoltaics due to their excellent light absorption and tunable bandgap (0.7-3.4eV) which spans the visible spectrum. However, the growth of this material remains a challenge due to thermal decomposition, indium surface segregation, and phase separation. To combat these issues, low substrate temperatures and high growth rates must be implemented in combination with in situ surface monitoring via RHEED. Metal modulated epitaxy (MME) has been shown to achieve single-phase InGaN alloys throughout the miscibility gap with sub-nm RMS roughness.

In this study, MME is applied to the growth of n-GaN/p-InGaN/p-GaN solar cells. Unintentionally doped InGaN layers 50/200nm thick with 16% In composition are grown on a 3µm n-type GaN template. The topmost layer consists of p-type GaN with varying hole concentrations. Transient RHEED signals are monitored closely during InGaN growth to prevent indium surface segregation and ensure a smooth film.

The absorption characteristics are determined as shown in Figure 1, and the material is characterized via AFM, XRD and PL prior to being fabricated into device structures. The InGaN layers are found to be approximately 80% strained as determined by XRD analysis of the (20-25) reflection (Figure 2). During device testing, the solar cells exhibit low turn-on voltages below the expected value, as well as low external quantum efficiency. These inconsistencies indicate carrier loss due to electrical shorts and defects. The scanning TEM images in Figure 3 show striations in the InGaN layer similar to those in a superlattice structure but with a period that does not correspond to the shutter cycles. Even though the x-ray diffraction indicates predominantly relaxed epitaxy, TEM images are unable to resolve individual threading dislocations. It is not clear at present if this is indicative of extremely large dislocation density or a new method of relaxation related to the above mentioned striations. Finally, post-growth annealing processes are explored as possible ways to improve device performance.

The work at Georgia Tech was supported by Air Force Office of Scientific Research, under a basic science grant managed by Kari Reinhardt.

Sandia National Laboratories is a multi-program laboratory managed and operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Corporation, for the U.S. Department of Energy’s National Nuclear Security Administration under contract DE-AC04-94AL85000.
relaxation times measured in epitaxial graphene fabricated with similar is expected due to the lack of a tunnelling barrier contact. The comparable spin show a coupling between the magnetic contacts and graphene channel. This deposition on the graphene. Spin precession was observed and the spin measurement devices were fabricated on epitaxially grown graphene on Graphene is an ideal candidate for the transport channel in future spintronic T.J. Watson Research Center, A.A. Sandin, D.B. Dougherty, J.E. Rowe, North Carolina State University Graphene may be an ideal material for spin field effect transistors because of its high charge carrier mobility and long spin relaxation times due to small spin-orbit coupling. However, efficient spin injection into graphene requires overcoming conductivity mismatch through the use of tunnel barriers and/or spin filters. It is possible that organic films can serve as tunnel barriers/spin filters with highly tailorable properties. In particular, metal phthalocyanines have recently been shown to exhibit spin dependent interfacial coupling on magnetic electrodes. A study of the coupling and morphology of such molecules on graphene is a crucial first step to understand potential spin enhanced interfaces. We deposit monolayer iron phthalocyanine (FePc) on both single layer and bilayer epitaxial graphene on the Si-terminated polar face of SiC, named SiC(0001). Scanning tunneling microscopy reveals an adsorbed molecular lattice periodicity of 1.8 nm, close to that of the graphene/SiC buffer layer corrugation periodicity. This lattice spacing is larger than that of FePc adsorbed on a graphite surface that shows a smaller spacing of ~1.4 nm. This implies a stronger interaction of the FePc with epitaxial graphene than expected and is possibly due to the modification of graphene by the SiC substrate. Tunneling spectroscopy has been used to study the occupied and unoccupied electronic states of the adsorbed monolayer FePc. Broad unoccupied states indicate significant electronic coupling between the molecules and the graphene and suggest a promising future for molecular strategies for spin injection. *Supported by the NSF Center for Chemical Innovation: Center for Molecular Spintronics under CHE-0943975.


Graphene: Magnetic Properties and Spin-Dependent Phenomena
Moderator: A.C. Ferrari, University of Cambridge, UK

We deposit monolayer iron phthalocyanine (FePc) on both single layer and bilayer epitaxial graphene on the Si-terminated polar face of SiC, named SiC(0001). Scanning tunneling microscopy reveals an adsorbed molecular lattice periodicity of 1.8 nm, close to that of the graphene/SiC buffer layer corrugation periodicity. This lattice spacing is larger than that of FePc adsorbed on a graphite surface that shows a smaller spacing of ~1.4 nm. This implies a stronger interaction of the FePc with epitaxial graphene than expected and is possibly due to the modification of graphene by the SiC substrate. Tunneling spectroscopy has been used to study the occupied and unoccupied electronic states of the adsorbed monolayer FePc. Broad unoccupied states indicate significant electronic coupling between the molecules and the graphene and suggest a promising future for molecular strategies for spin injection. *Supported by the NSF Center for Chemical Innovation: Center for Molecular Spintronics under CHE-0943975.


Graphene Energy Gap
We have performed scanning tunneling spectroscopy measurements on a gated bilayer graphene device. In graphene bilayer, a potential asymmetry between the layers induces an energy gap in the electron spectrum. The formation of the energy gap is investigated as a function of carrier density and magnetic field. We found that in zero magnetic field, the reliable determination of the gap can be complicated because of disorder scattering. However, in the quantum Hall regime, the energy gap can be quantitatively determined by measuring the layer-polarized low index Landau levels. Our scanning tunneling spectroscopy measurements reveal that the microscopic nature of the bilayer gap is very different from what was expected due to the lack of a tunnelling barrier contact. The comparable spin relaxation times measured in epitaxial graphene fabricated with similar contacts and the multi-layer flake is believed to be caused by a large contact induced relaxation due to the contacts coupling with the graphene channel. The strong coupling effectively removes the spin from the channel.
observed in previous macroscopic measurements or expected from current theoretical models. The potential asymmetry varies spatially in both magnitude and sign on a nanometer length scale, showing strong correlation with the disorder potential. This random pattern of alternating dipole fields is qualitatively consistent with the reduced disorder-induced density fluctuations in the top layer.

4:40pm GR+MI-TuA9 Atomic, Electronic, and Magnetic Properties of Metal-Graphene Interfaces. I.I. Oleynik, L. Adamska, Y. Lin, University of South Florida, A. Ross, Saint Anselm College, M. Batzill, University of South Florida
Metal/graphene interfaces play an important role in both surface science studies of the epitaxial growth of graphene on metallic substrates, as well as in metal/graphene contacts in graphene nanoelectronic devices. We present results of first-principles density functional theory (DFT) investigations of structural, electronic, and magnetic properties for graphene/Ni(111) and graphene/Ni(111) interfaces relevant to experimental studies of graphene growth on metallic substrates. The favored interface geometries and binding sites for different interface configurations were identified. Additional adlayers of Ni and Cu were either adsorbed on top of the graphene/metal interface, or placed between the graphene and substrate to model processes of metal intercalation. It was also found that the interaction between graphene/Ni(111) and the top Cu adlayer is much weaker compared to that for a Ni adlayer. The atomic, electronic, and magnetic properties of these interfaces, including induced magnetic moments in graphene/Ni(111), Ni/graphene/Ni(111) systems, are also discussed.

5:00pm GR+MI-TuA10 Spin-Derived Scattering from Gated Potential Obstacles in Graphene Systems. M. Asmar, S. Ulloa, Ohio University
We study the scattering of Dirac fermions in a sheet of graphene from potential obstacles created by external gates in the presence of both intrinsic and extrinsic spin-orbit (SO) interactions [1]. Obtaining analytical solutions in a real-space representation for the eigenvectors allows us to calculate the phase shifts generated by a finite-size obstacle in the presence of SO interactions [2]. From the phase shifts extracted from these solutions we can calculate the differential, total and transport cross sections. The knowledge of these quantities allows us to obtain the spin-flip and momentum relaxation times. The dependence of both relaxation times on the strength of the SO interaction was analyzed showing comparable relaxation times for relatively large values of energy, while displaying a big difference for small values of energy. The relaxation times of the injected electrons exhibit a number of resonances in energy associated with the structure of the scattering obstacle. In the presence of SO, new resonances appear at energies that depend on the strength of the SO interaction, and as such contain spectroscopic information on the system. It has been shown that the main scattering mechanism in graphene is due to strong defects [3]. Therefore, the analysis performed in our work can help understand the role of SO interactions in the scattering processes in these and related experiments.


We investigate the magnetic field and temperature-dependent transport properties of CVD-grown graphene subjected to different strains. The graphene is transferred to kaptop substrates to which a bending-force can be applied. In zero magnetic field, the prefactor to the logarithmic-in-temperature conductivity correction decreases by an approximate factor of 3 for strains as high as 0.6 %. There is also a concomitant decrease in diffusivity by a factor of 6. At 5 K we observe negative magnetoresistance for fields up to 0.5 Tesla followed by positive magnetoresistance at higher fields. We attribute the low field negative magnetoresistance to weak-localization and find that it is well described by the theory. The strains resulting from the applied bending force inhibit the intervalley scattering more than an order of magnitude and decrease the phase coherence length, thereby leading to a suppression of weak-localization.

5:40pm GR+MI-TuA12 Simulation of Electron-Ion Dynamics in Pristine and Functionalized Graphene in External Fields. S. Buhin, K. Varga, Vanderbilt University
In the framework of real-time real-space time-dependent density functional theory (TD-DFT) we have studied coupled electron-ion dynamics in small fragments of graphene, graphene, and fluorinated graphene subjected to short (a few femtoseconds) intense laser pulses or irradiated by energetic ions. The goal of this study is to investigate the possibility of defect creation in graphene and desorption of hydrogen/fluorine from graphene surface. We will present the results of our simulations, discuss the mechanisms that take place, and identify the parameters of the laser or energetic ions necessary for those processes to occur.

5:40pm GR+MI-TuA7 Studies on Ozone Based Atomic Layer Deposition of High-k Dielectrics on Graphene. S. Jandhyala, G. Mordi, B. Lee, J. Kim, University of Texas at Dallas, P.-R. Cha, Kookmin University, Korea
Graphene, being a two dimensional material, is one of the most promising alternative channel materials for post-Si generation [1-3]. However, being just one atom thick and having an inert surface, it poses a huge challenge to develop a top-gate dielectric process for graphene-based devices. Several
techniques are currently being explored for depositing dielectrics including physical-vapor deposition (PVD), chemical vapor deposition (CVD) and atomic layer deposition (ALD) after chemical functionalization of graphene (using NO$_2$ or O$_3$) or after depositing nucleation layers (such as Al, PtCA, PVA) on graphene [3].

Here, we will present a novel technique developed by our group for depositing ALD high-k dielectrics such as Al$_2$O$_3$ on graphene through ozone functionalization [4]. Physisorption of ozone has been claimed to be the plausible mechanism for functionalizing the graphene surface [5]. Based on Langmuir adsorption equation, the amount of ozone adsorbed on graphene can be increased by increasing the partial pressure of ozone. By utilizing this, we have been able to precisely control the dielectric thicknesses and successfully scale dielectrics on graphene down to a thickness of ~3 nm. We employed both AFM on HOPG/graphene and in-situ electrical characterization of graphene-FETs in order to understand the adhesion mechanisms of ozone with graphene, enabling the deposition of ALD dielectrics.

In situ electrical characterization, we used package-level devices with back-gated graphene devices to detect molecules adsorbed on graphene surface. The observed charge scattering mechanisms and effect on mobility due to the interaction of ozone with graphene as a function of temperature and amount of ozone will be presented. In-situ studies regarding the role of TMA (Tri-methyl Aluminum) will also be discussed based on experiments in actual ALD chambers.

Acknowledgement
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References


Thermal atomic layer deposition (ALD) is a viable approach to attain high-quality graphene dielectric films needed for graphene devices, but the hydrophobic nature of the graphene surface inhibits direct application of thermal ALD oxides. Several methods have been explored to render the surface more susceptible to ALD[1-3], but these techniques often result in graphene mobility degradation and/or shifts in the Dirac voltage due to charge in the gate stack. In this work, we investigated a simple dry chemical process using XeF$_2$ to functionalize the graphene surface prior to ALD which results in conformal ultrathin high-k oxides without degradation of the underlying graphene electrical properties. Epitaxial graphene samples were grown on semi-insulating, on-axis (0001) 6H-SiC substrates using an Aixtron VP508 SiC reactor at 1650°C for 120 min. Fluorination of the graphene was performed in a XeF$_2$ Xe$_2$ etcher operating in pulse mode. Optimum fluorine exposure conditions consisted of six, 20s pulses with constant XeF$_2$ and carrier gas partial pressures of 1 and 35 torr, respectively. X-ray photoelectron spectroscopy (XPS) was used to chemically analyze the functionalized surface prior to oxide deposition. ALD Al$_2$O$_3$ and HfO$_2$ films (~15 nm) were deposited at temperatures between 150 - 225°C using TMA or TEMAHf and deionized (DI) water precursors. Growth was initiated with 20 DI water pulses. Oxide coverage was characterized with atomic force microscopy and scanning electron microscopy, while graphene mobility changes were observed with Van der Pauw Hall measurements. Capacitance-voltage (C-V) measurements were conducted on Ta/Au/C-V dots to extract the dielectric constant and electrical quality of the oxide. Initial results show that 15 nm conformal, uniform Al$_2$O$_3$ and HfO$_2$ films are obtained with an optimized XeF$_2$ surface treatment prior to ALD. XPS showed that the optimum XeF$_2$ treatment resulted in ~6% fluorine on the surface and the presence of only C-F bonds which provide ALD reaction sites needed for uniform oxide deposition. Graphene mobilities were maintained, and occasionally increased, implying little impact of the XeF$_2$ treatment on the underlying graphene properties. Raman spectroscopy reveals no change in the D/G ratio after XeF$_2$ and oxide deposition, verifying that the graphene lattice quality is maintained. The viability of the fluorination method for achieving ultrathin films (~10 nm) will be presented along with electrical C-V data to show the electronic quality of the ALD oxides.

A. Robinson, et.al., ACSNano 4(5) 2667 (2010)


Graphene grown by chemical vapor deposition (CVD) provides a promising pathway for large area fabrication of graphene field effect transistor (FET). However, the performance of CVD graphene FETs reported to-date is lower than that of FETs fabricated using exfoliated graphene. CVD graphene FETs often exhibit strong hysteresis accompanied with low mobility, large positive Dirac point (V$_{Dirac}$) and large intrinsic carrier concentration. CVD graphene is exposed to a number of aqueous solutions and deionized water when it is transferred to a device substrate. We find that the large V$_{Dirac}$ shift and strong hysteresis observed in CVD graphene FET are largely due to water trapped in the graphene/substrate interface during the transfer process.

In this study, CVD graphene grown on copper is transferred to SiO$_2$ substrates with the following three interfacial conditions: i) normal hydrophilic SiO$_2$, ii) SiO$_2$ with 20nm of Al$_2$O$_3$, and iii) a hydrophobic surface prepared by coating hexamethyldisilazane (HMDS). Device performance, including mobility, V$_{Dirac}$ and intrinsic carrier concentration are compared in ambient as well as in vacuum. Gate hysteresis is analyzed by measurement of time-resolved channel resistance at various back-gate bias voltages. We find that the gate hysteresis is partially reduced by transferring the graphene onto a substrate coated with HMDS. Vacuum pump down and low temperature (80 °C) annealing can remove the remaining gate hysteresis and V$_{Dirac}$ shift. The resulting hole mobility is 5.420cm$^2$/Vs, which is high compared to most of the CVD graphene mobility values reported in the literature.

As a control experiment, the CVD graphene FET fabricated on untreated SiO$_2$ shows a smaller mobility, a larger V$_{Dirac}$ and a stronger hysteresis compared to the HMDS coated sample. However, the hysteresis is reduced but remains significant. We believe the remaining hysteresis is due to adsorbrates trapped at the substrate/graphene interface. A graphene FET prepared on a substrate with an Al$_2$O$_3$ interface shows less hysteresis than the sample fabricated on an untreated SiO$_2$ surface but more than that of the HMDS coated substrate. In order to study the influence of water trapped between the graphene and the substrate, water is intentionally replaced by isopropanol at the end of the transfer process before drying. In samples prepared using this method, hysteresis and V$_{Dirac}$ point shift are both reduced. These results indicate that efforts to prevent trapping of water molecules at the graphene/substrate interface during the transfer process will improve the performance of CVD graphene FETs.

This work was supported by the NRI SWAN center, ONR, NSF and Sandia’s LDRD program.

5:00pm GR-TuA10 Improved Performance of Top-Gated Graphene-on-Diamond Devices, J.F. Sumanth, Argonne National Laboratory, J. Yu, G. Liu, A. Balandain, University of California, Riverside

Since the discovery of graphene and realization of its exceptional electronic properties in suspended form, there have been many efforts in fabricating FET-type devices based on single and bilayer graphene on SiO$_2$ substrate. However, performance of these devices is found to be inferior to the expected intrinsic properties of graphene. It has been observed that apart from carrier mobility in graphene, which is sensitive to trapped charges, and surface impurities at the graphene-oxide interface, breakdown current density in graphene depends sensitively on the heat dissipation property of the underlying supporting substrate. Although graphene has extremely high intrinsic thermal conductivity, it is reported that in graphene devices, more than 70% of the heat dissipates through the 300 nm SiO$_2$ on silicon directly below the active graphene device, while the remaining ~2% flows via the graphene that extends beyond the device and metallic contacts. Such a distribution of heat in to the substrate cause undesirable effects on the overall performance of the device. We show for the first time that by the use of thin CVD-grown ultrananocrystalline diamond thin films on silicon in graphene-on-diamond configuration, the heat dissipation can be improved substantially leading to the higher breakdown current density of more than 50% as compared to conventional graphene-on-oxide substrates. We also describe the fabrication of the top-gate graphene-on-diamond devices and discuss their performance. The obtained devices had the carrier mobility ~ 2354 cm$^2$/Vs for holes and ~1293 cm$^2$/Vs for electrons. The obtained results are promising for developing high-performance graphene-on-diamond devices and interconnects for future electronics.

Use of the Center for Nanoscale Materials was supported by the U. S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Contract No. DE-AC02-06CH11357. The work in Balandain group at UCR was supported, in part, by DARPA – SRC Center on Functional Engineered Nano Architectonics (FENA).
electron diffraction, which was possible due to the low Debye-Waller factor. After obtaining a clean reconstructed surface, the sample was cooled in 1 – 7 E-7 Torr of acetylene. Transmission electron diffraction showed very sharp, nearly continuous rings, indicating large domains and no preferential azimuthal rotation between planes. X-ray diffraction showed an increased layer separation of 0.345 nm compared to graphite. Energy loss spectroscopy showed bulk-like plasmons or interband transitions, indicative of multilayer graphene. Raman spectra showed 2D/G peak intensity ratios of 0.5 to 1, comparable to literature values for transition structures. Water nanomanipulation under graphene could help elucidate water’s complex bonding structure and charge very efficiently, and with enhanced materials contrast when ion beam interaction are readily seen due to the surface sensitivity of the technique. At sufficiently high current densities the sharply focused He ion beam may very efficiently decompose or cross-link residual hydrocarbons that are present in the instruments vacuum, more so than an electron beam may very efficiently decompose or cross-link residual hydrocarbons.
sputtering rates. On the one hand, this has proven extremely useful in the field of nanopatterning for sensors, plasmonics or other device fabrication applications at the sub-10nm level when operating at high doses. On the other hand, it is undesirable when the instrument is used for materials characterization.

In this presentation we will discuss the basic considerations that went into the design of a Near-UHV (NUHV) Orion Plus® He Ion Microscope. We will detail how the improved vacuum level is anticipated to alter these processes that are directly relevant to the imaging performance of the instrument such as beam interaction in the surface region and the emission of secondary electrons. First applications that the instrument was used for will be highlighted and its impact in the areas of surface physics, notably catalysis, corrosion, and other research areas that require increased imaging sensitivity, both laterally and in depth, will be discussed.

References

3:00pm HI+AS-TuA4 Sub-10 nm Scanning Helium Ion Beam Lithography, K. van Langen, E.W.J.M. van der Drift, Delft University of Technology, Netherlands, E. van Veldhoven, D.J. Maas, TNO, Netherlands, P.F.A. Alkemade, Delft University of Technology, Netherlands
Since the launch of the novel sub-nanometer helium ion microscope by Zeiss / Alis in 2006 nanofabrication with this tool has gained a lot of interest [1]. Key advantage in this matter is the directional interaction of the helium ion with matter with negligible backscattering. In ion milling it enables very steep structuring when compared to the Ga+ ion beam [2]. In a similar comparison helium ion beam-induced deposition in a precursor gas ambient yields tall and smooth nanostructures [3], partially also because the sputtering by helium ions is at least an order of magnitude lower than by gallium ions.

The present contribution deals with scanning helium ion beam lithography (SHIBL). Thus far two initial SHIBL studies on hydrogensulphosxacn (HSQ) resist were reported [4,5]. In the present work performance of SHIBL is compared with state-of-the-art electron beam lithography (EBL). As resist materials we explored HSQ, polymethylmethacrylate (PMMA), and the inorganic resist of aluminumoxide. The latter material choice is motivated by the need for enhanced mask selectivity in pattern transfer in the sub-10-nm area.

The results for HSQ and PMMA can be summarized as:
- smallest feature size of 5 nm, equivalent to the best EBL performance [6]
- clear pattern densities up to 10 nm full-pitch, which is better than in EBL
- sensitivity 1-2 orders of magnitude better than in EBL.

As for the inorganic resist, 5-nm features have been realized.

In a semi-quantitative and comparative approach the results will be explained and future prospects will be outlined.


4:00pm HI+AS-TuA7 Contrast Performance in Helium Ion Microscopy, D.C. Bell, Harvard University
In order to achieve ultra high resolution imaging with secondary electron imaging, it is critical that the electric potential of the specimen surface is well controlled. For electrically conductive samples this can be achieved by simply grounding the specimen. However, imaging of electrically insulating specimens can provide challenging or impossible to image due to uncompensated charge resulting from the electron or ion beam interaction with the specimen surface. The main reason for the uncompensated charge is that the insulating specimen has insufficient conductivity through mobility of either electrons or holes to quickly restore the neutrality of the scanned area. The buildup charge causes significant deflection and distortion of the ion or electron beam. Which is more appropriate, to use charge compensation with high kV helium Ions or employ a low kV SEM image to obtain the required surface information? This paper will present a systematic examination of the surface information provided by both techniques, including SEM charge compensation mechanisms.

One key advantage of the Helium Ion Microscope technology in the case of imaging highly charging specimens is the electron flood gun can be utilized to neutralize the positive charge buildup and facilitate high-resolution imaging. A flood gun is used to charge the surface to a positive potential (using electrons as the neutralizing particles). When utilizing the electron flood gun, the electron beam and He ion beam are synchronized and adjusted with respect to one another, so that the low energy electrons are not interfering with the secondary electron imaging.

Some of our research from the past year has been surprising and may provide a foundation for a change in analysis techniques of different materials. The nature of the Helium ion beam interactions with the sample shows enhanced edge contrast which is especially useful for critical dimension measurements; one particularly interesting development is the imaging of non-conducting materials showing a contrast due to three apparent mechanisms simultaneously - atomic number, channeling contrast and a possible enhanced edge contrast. The advantages of Helium ion microscopy is still being investigated and still are proving some exciting results.

5:00pm HI+AS-TuA10 Helium Ion Beam Induced Deposition Examined using a 3D Monte Carlo Simulation, D.A. Smith, P.D. Rack, University of Tennessee Knoxville, P.F.A. Alkemade, H. Miro, Delft University of Technology, The Netherlands
The growth of nanostructures has traditionally been dominated by electron beam induced deposition (EBID) or gallium ion beam induced deposition (Ga-IBID). While EBID provides smooth sidewalls and good resolution for nanopillar growth, the cross-section for dissociation is low and etching is difficult as sputtering is negligible. Ga-IBID is a relatively faster method of producing nanostructures, however it suffers from lower resolution, alters deposited materials, and leaves an etching residue. A new tool in this field has been recently explored: the helium ion beam microscope. This tool has been modified to perform IBID using high energy helium ions. It has been found that He-IBID combines the high resolution of EBID with the speed of Ga-IBID. Moreover, there is less implantation damage and minimal sputtering compared to Ga beams.

To examine this process, a 3-dimensional Monte Carlo simulator has been designed based ion-solid-precursor interactions. This simulation system, named EnvisION, can provide useful knowledge of how user-controlled parameters can be optimized for highly efficient growth of nanostructures using this tool. In this work, an in-depth explanation of the simulation will be presented including an example of its use examining the growth efficiencies of nanopillars grown on a silicon substrate using the (CH3)3P(CpCH3) precursor via He-IBID. Furthermore we compare how the morphology changes with dwell times, refresh time, precursor coverage and surface diffusion in order to span the range of growth regimes from mass-transport limited to reaction-rate limited deposition. The simulated morphologies predicted using the EnvisION simulator are compared to experimentally grown pillars to validate the simulation.

5:20pm HI+AS-TuA11 TEM Specimen Preparation with Light Ions, L. Giannuzzi, L.A. Giannuzzi & Associates LLC
Much research with light energetic ions such as He+ and Ne+ from gas field ionization sources has focused on imaging and nano-machining. It is a natural progression to question to the viability of TEM specimen preparation using these light ions. Of vital importance for TEM specimen preparation quality is the understanding of surface ion implantation and amorphization damage. Theoretical calculations using SRIM indicate that there may be a damage trade off between vacancy formation, ion range, and dose. That is, the range of light ions is much greater than conventional heavy ions (e.g., Ga+), and can indeed penetrate directly through a TEM specimen. While this may indicate the possibility of light ions damaging the entire TEM specimen thickness, light ions produce far less vacancies per ion compared to heavy ions for the same dose. However, since the sputter yield of light ions is smaller than heavy ions, a larger dose of light ions may be necessary to achieve sufficient material sputtering. This theory will be supplemented with experimental results.
Manufacturing
Berkeley, R. Mu, Fisk University

Moderator:
Manufacturable Nanoscale Devices and Processes
Nanomanufacturing Science and Technology Focus
D.J. Maas, E. van Veldhoven, E.W.J.M. van der Drift, Delft University of Technology, Netherlands

Although Helium Ion Microscopy (HIM) was introduced only a few years ago [1], many new application fields are emerging. Key issue is the directional interaction of the primary helium ion beam with the sample at and just below its surface with negligible backscattering. The sub-nanometer sized probe of the 10-35 keV ion beam generates Secondary Electrons (SEs) that have a typical energy between 0 and 20 eV. Taking all together the SE signal stems from an area that is very well localized around the point of incidence of the primary beam. This makes the HIM well-suited for both high-resolution imaging as well as high resolution nanofabrication [2]. We explore the possibilities to use the HIM simultaneously for imaging and nano patterning of EUVL masks.

The HIM is a high-resolution surface imaging tool. In practice, the optimum dose for imaging is a balance between maximizing SN ratio, while minimizing sample damage. Imaging work at TNO van Leeuwenhoek Laboratory (VLL) [3] focuses on sensitive materials such as e.g. DUV and EUV resists and EUV masks, which are difficult to image in a SEM due to their charging behavior. An electron flood gun in the HIM offers effective charge cancellation, which enables high-resolution imaging of insulation structures and for pin pointing defects on a EUV reticle. In this presentation we will show images of particles down to 5 nm in reticles.

Furthermore, to explore the possibilities of the helium ion microscope as a nanofabrication tool, the HIM at the TNO VLL is equipped with a pattern generator and a gas injection system (GIS). This presentation will show our latest nano structuring results made with Helium Ion Beam Induced Processes: deposition and etching. It is expected that the unique capabilities of the HIM in combination with the GIS are suited for EUV mask repair. These capabilities offer the possibility of circuit repair in the latest and smallest semiconductor technology nodes (beyond 22 nm). In both cases, sub-surface damage due to scattered He ions is a matter of concern and topic of investigation. At this moment we are capable of etching 13 nm lines with 25 nm spacing on a EUV dummy mask with approximately 80 nm of TaN absorber. Furthermore we demonstrate Pt deposited lines of 13 nm width at a 16 nm spacing.

References

Invited: N. B. Ramananathan, AIXTRON Inc.
Atomic Layer Deposition enables conformal coating of high-quality thin film on complex nano-scale structures. It has been the preferred choice of deposition technology for high-k and metal films in high-aspect ratio capacitor structures for modern 30nm DRAM devices. This talk will review the latest nano-structuring results made with Atomic Layer Deposition (ALD), as well as practical manufacturing targets (such as throughput, cost of ownership) in order to provide a production-worthy ALD solution. Applications in new high-k oxide (La/Sr/Ba oxides), metals, and PCRAM materials (such as GST) will be discussed.

4:00pm NM+NS+MS-TuaA7 The Metal-Oxide-Metal Vacuum Drift Memristor - A CMOS Compatible, High Speed, Non-Volatile Switch for Universal Memory and Storage, R.S. Williams, J.P. Strachan, Hewlett-Packard Labs

The existence of the fourth passive circuit element was proposed by Prof. Leon Chua of UC Berkeley in 1971 from fundamental symmetry arguments to augment the familiar resistance, inductance and capacitance equations. Although he showed that such a ‘memristor’ had many interesting and useful circuit properties, until 2008 no one knew if such a circuit element existed or not. In fact, researchers had been making and studying memristors for decades without knowing it - examples are resistive RAM devices, STTRAM devices and phase change memory devices. At HP, we have focused primarily on metal-oxide-metal bipolar resistive switches. Memristance arises naturally in these systems via coupling of electronic and ionic transport in thin semiconducting metal-oxide films under an external bias voltage. Simple analytical models show that memristance becomes much more important as the thickness of the active device region decreases, and thus memristors are mainly nanoscale structures. Memristor theory serves as the foundation for understanding a wide range of hysteretic current-voltage behavior, including both unipolar and bipolar switching, observed over the past 50 years. We have built nanoscale titanium dioxide and tantalum pentoxide memristors in our laboratory and have demonstrated both their fundamental electrical properties and several potential uses. They can be incorporated into electronic circuits using conventional fabrication techniques and materials available in standard CMOS fabrication facilities. I will discuss recent results on such topics as device switching speed, endurance, measurements required to parameterize a physics-based SPICE model, and 3D stacking of memristive crossbars.

4:40pm NM+NS+MS-TuaA9 Large Scale Graphene: Progress and Challenges, R.S. Ruoff, The University of Texas at Austin

Graphene-based materials hold promise due to their electronic and thermal transport properties, mechanical properties, high specific surface area, and that they can act as an atom thick layer, barrier, or membrane. Here, I focus on growth of large area graphene on metal substrates and the structure and thermal and mechanical properties of such graphene. A history of experimental work on graphene (from its discovery in 1969 until 2010) is provided at: http://bucky-central.me.utexas.edu/.

Support of our work by The WM Keck Foundation, DARPA, ONR, SWAN NRI, NSF, ARO, AEC, and Graphene Energy, Inc., is appreciated.
Omniprobe, Inc. successful application of high K dielectrics to DRAMs, it is essential to constraint at the same time, high K gate dielectrics should be adapted. For a tuner method was realized. SiGe channel. To conclude, the channel SiGe SEG process has magnitudes, which consequently results in the degradation of transistor DRAM process. Therefore, so called 'gate-first approach' has been examined, i.e. LaO, or MgO capping layers for NMOS [1,2], and AlO XeF2.

As the DRAM technology evolved towards the sub 2x era, the need for high Yoo, S. Ahn, I. Jeon, B. Kim, S. Nam, S. Kim, K. Jung, J. Lee, S. Jung, T. Lee, K. Huh, S. Yamada, Samsung Electronics Co., Ltd, Republic of Korea As the DRAM technology evolved towards the sub 2x era, the need for high performance transistor grows higher for the DRAM peripheral transistors. The novel technologies such as embedded SiGe, high K gate oxide, or 3-dimensional transistor technologies are indispensable in a near future. Especially, to scale the gate oxide further and to meet the gate oxide leakage treatment [1,2], a variant of noncontact atomic force microscopy, with simultaneous steps, kinks, impurities, and domain boundaries as active sites. For example, in the case of a catalytically active surface, this would allow electron densities, quantify force interactions, and identify chemical species. In this talk, we will show with the example of an oxygen/copper(100) surface that such of this information can be derived from combining the new method of three-dimensional atomic force microscopy (3D-AFM) [1,2], a variant of noncontact atomic force microscopy, with simultaneous scanning tunneling microscopy. The surface oxide layer of Cu(100) features domain boundaries and a distinct structure of the Cu and O sublattices that is ideally suited for such model investigations. By combining experimental results with theoretical simulations, we will show how 3D data sets enable the site-specific quantification of force interactions and tunneling currents, how different chemical species can be imaged using different tips, different tunneling conditions, and different interaction mechanisms, and how

5:20p GM+-NS+MS-TuA1 Laser-Assisted Electron-Beam Induced Deposition and Etching N.A. Roberts, University of Tennessee and Omniprobe, Inc., J.D. Fowlel, Oak Ridge National Laboratory, P.D. Rack, University of Tennessee and Oak Ridge National Laboratory, G.A. Magel, H.M. Marchman, C.D. Hartfield, T.M. Moore, Omniprobe, Inc., etc. Focused electron-beam induced deposition (EBID) and etching (EBIE) are direct-write nanofabrication techniques that allow localized deposition or etching of materials without the need for resists. These deposition and etching processes are controlled by sputtering-electron beam dissociation of a precursor gas. In both cases, by-product species are created, and if the unwanted byproduct is not desorbed from the surface it will be incorporated into the deposit or reduce the etch rate for deposition and etching, respectively. Substrate heating has been used in experiments to enhance desorption by reducing the residence time of the by-product. The substrate heating has the same impact on the residence time of the precursor gas and thereby reduced the growth or etch rate of the process. Ex situ analysis of deposits have also been investigated to remove impurities with some success, but these treatments results in void formation and shape changes. In situ laser processing at short pulse widths is ideal for electron-beam induced processing because desorption of the by-products can be achieved by local heating of the sample, but the narrow pulse width results in a short heating time and cooling time. Thus the by-products can be effectively desorbed and adequate recovery time for fresh reactant to re-adSORB.

Laser-assisted EBID and EBIE processes are made possible through the use of the OptoProbe™, which is an optical imaging and processing system that can be attached to an SEM and used in conjunction with an appropriate gas injection system. The design of this port-mounted optical accessory enables simultaneous optical imaging and delivery of laser irradiation to a sample within the SEM, without interfering with normal SEM/FIB imaging and processing modes. The optical system is mounted on a 3-dimensional nanomanipulator so that precision alignment and focusing is easily achieved. For this work, the OptoProbe™ has been optimized to deliver a high-irradiance near-infrared laser spot to provide localized time-dependent sample heating for enhancing focused electron-beam induced deposition and etching. In this presentation, we will discuss recent experimental results as well as modeling of laser-assisted EBID of Au and EBIE of SiO2 using XeF2.

5:40p GM+-NS+MS-TuA2 Channel SiGe Selective Epitaxy Process for DRAM High K Peripheral Transistors. J. Yeo, H. Hwang, S. Lee, W. Yoo, S. Ahn, I. Jeon, B. Kim, S. Nam, S. Kim, K. Jung, J. Lee, S. Jung, T. Lee, K. Huh, S. Yamada, Samsung Electronics Co., Ltd, Republic of Korea As the DRAM technology evolved towards the sub 2x era, the need for high performance transistor grows higher for the DRAM peripheral transistors. The novel technologies such as embedded SiGe, high K gate oxide, or 3-dimensional transistor technologies are indispensable in a near future. Especially, to scale the gate oxide further and to meet the gate oxide leakage treatment [1,2], a variant of noncontact atomic force microscopy, with simultaneous steps, kinks, impurities, and domain boundaries as active sites. For example, in the case of a catalytically active surface, this would allow electron densities, quantify force interactions, and identify chemical species. In this talk, we will show with the example of an oxygen/copper(100) surface that such of this information can be derived from combining the new method of three-dimensional atomic force microscopy (3D-AFM) [1,2], a variant of noncontact atomic force microscopy, with simultaneous scanning tunneling microscopy. The surface oxide layer of Cu(100) features domain boundaries and a distinct structure of the Cu and O sublattices that is ideally suited for such model investigations. By combining experimental results with theoretical simulations, we will show how 3D data sets enable the site-specific quantification of force interactions and tunneling currents, how different chemical species can be imaged using different tips, different tunneling conditions, and different interaction mechanisms, and how
structure-induced stress fields and their influence on the local chemical activity and topographical deformation can be studied.


4:00pm NS+AS-TuA7 The Role of Surface States in Inelastic Electron Tunneling Into Metal Surfaces, P. Maksymovych, M.H. Pan, Q. Li, Oak Ridge National Laboratory

A small fraction of electrons tunneling across a vacuum junction will undergo inelastic scattering, exciting surface phonons, molecular vibrations, magnons and plasmons in the contact leads. Although the study of surface phonons is a most straightforward inelastic electron tunneling spectroscopy (IETS) experiment, relatively few systematic studies have been done to date. One of the most surprising observations was that of atomic resolution in the IETS of Cu(111). In this talk we will present results from a new two-color method that completely eliminates this interference. We will show results for two cases: (1) where the tip is retracted from the surface far enough to prohibit tunneling, and (2) where the tip is within tunneling range of the surface. A delay-modulation technique isolates the two-color photo-emission from concurrent one-color two-photon photoemission and the conventional tunneling current, and also enables subpicosecond time-resolved detection of the photoexcited surface electrons. Advantages of the two-color approach are highlighted by comparison with the one-color case where optical interference causes current modulations that are orders of magnitude larger than the desired signal. The two-color approach represents an important step toward the ultimate goal of simultaneous subnanometer and subpicosecond measurement of surface electron dynamics.


5:20pm NS+AS-TuA11 High Precision local electrical Probing: A New Low Temperature 4-Tip STM with Gemini UHV-SEM Navigation, B. Guenther, A. Bettac, M. Maier, M. Oertel, Omicron NanoTechnology, Germany, F. Matthes, C.M. Schneider, Forschungszentrum Juelich, Germany, A. Feliz, Omicron NanoTechnology, Germany

A major challenge in the development of novel devices in nano- and molecular electronics is their interconnection with larger scale electrical circuits required to control and characterize their functional properties. Local electrical probing with STM tips can significantly improve efficiency in analyzing individual nano-electronic devices without the need of a full electrical integration. Among a very few commercial approaches, the Omicron UHV NANOPROBE has been established as a suitable instrument for local electrical probing in UHV on nano-structures down to structure sizes in the 10 nm range. The major technical requirements for such sophisticated instrumentation are:

- Rapid and simultaneous SEM navigation of four local STM probes on nano-structures
- Localization of nanostructures by high resolution SEM (UHV Gemini)
- Individual probe fine positioning by atomic scale STM imaging
- STM based probe approach for “soft-landing” of sharp and fragile probes and controlled electrical contact for transport measurements
- Preparation techniques towards sharp and clean and STM tips
- Suitable low noise signal re-routing for transport measurements with third party electronics

Although the UHV NANOPROBE has been successfully used for various applications, today’s scientific requirements motivated the development of the next generation probing system. We will present the newly developed LT NANOPROBE which takes experimental capabilities one step further and opens up new research opportunities in nano-electronics, spintronics, and molecular electronics. Besides SEM/STM probe fine navigation and imaging, the excellent STM performance level of the LT NANOPROBE expands applications to tunneling spectroscopy and even the creation or modification of nano-structures by an ultimately precise STM probe. The R&D project has been driven by the following major milestones:

- Operation at temperatures of T<5 K for STM imaging and STM based probing
- Operation at temperatures of T<5 K for UHV imaging and STM based probing
- SEM navigation at base temperature T<5 K
- Simultaneous operation of STM and SEM at base temperature
- Thermal equilibrium of sample and STM at base temperature
- Performance and stability level of each individual STM Probe suitable for STM spectroscopy and atom manipulation

First evaluation measurements with the system installed at the Forschungszentrum Jülich will be presented: STM on Au(111) with pm...
stability, STS revealing the superconducting gap of a Nb tip with approx. 3meV gap size, and transport measurements on nanowires at T<5K.


Plasma Science and Technology Division
Room: 202 - Session PSI-TuA

Advanced BEOL / Interconnect Etching II
Moderator: S. Sriraman, Lam Research Corp.

Feature profile control for HARC etching is a very important issue. Possible profile defects such as bowing, necking, variations in depth and shape, among others, become critical for the next technological steps with ever shrinking dimensions of the features. Using numerical simulations as a tool, we studied HARC etching of SiO2 by the fluorocarbon-argon-oxygen plasma in a capacitive-type plasma system. The first step in our simulations involved estimation of neutral and ion fluxes, and their angle-energy distributions for most important species involved into processing in the system chosen for comparison. Then the feature profile simulator FPS-3D [1] was run to estimate various chemical and physical effects on HARC systems chosen for comparison. Then the feature profile simulator FPS-3D [1] was run to estimate various chemical and physical effects on HARC profile evolution during processing. The difficult part in those simulations was taking into account deposition of polymer films and etching through those films. The final results come as a competition between counteracting effects of etching and deposition. Important effects of ions for etching at the bottom of the contact holes, and of polymer films for protecting sidewalls from further etching, were investigated. Possible bowing and necking effects were considered in detail. Calculations were carried out for different sizes of the features, different gas compositions, and different RF powers leading to different ion energies. Published data for the underlined reaction mechanisms were analyzed and used for tuning the chemical reactions database in the FPS-3D code. Comparison of simulations with different experiments and the corresponding analysis of HARC etch processing are presented. The authors are thankful to Masanobu Honda and Akira Tanabe for providing experimental data.


Etching of high aspect ratio (HAR) features in dielectric substrates is a challenging process, growing increasingly difficult with each new device generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals. [1] Oxygen atoms act to suppress excessive fluorocarbon generation. A common approach uses a multi-frequency capacitively-coupled plasma (CCP) reactor to achieve high bias powers and high density of fluorocarbon and oxygen feed gases. Here, F atoms are the primary etchant, assisted by high energy ions to penetrate the fluorocarbon polymer layer formed by CFx radicals.
Because the strip plasma tends to remove the passivation from the sidewall, and because the damage rate increases in the final part of the strip process, better results can be achieved if the protective films is periodically re-applied. A cyclic process which alternates between passivation and strip conditions minimizes the strip-induced damage at the via sidewall. To facilitate an efficient hybrid strip process, the etch reactor design can be optimized for rapid switching between the two different plasma conditions. This requires minimization of both the gas exchange time and the RF coupling stabilization time.

99 Tuesday Afternoon, November 1, 2011
4:00pm PS1-TuA7 Plasma-induced Damage Reduction in Porous SiOCH Dielectrics by Replacement of H2 and N2 by CH2F2 and Ar in Fluorocarbon Based Plasmas. L. Souriau, F. Lazzarino, L. Carbonell, I. Ciofi, P. Verdonck, J.F. de Marneffe, M. Baklanov, IMEC, Belgium

With the scaling of devices and related interconnects, the integration of dielectric materials with a k-value lower than 2.5 is greatly required. In particular, hybrid porous SiOCH low-k materials are considered as promising candidates. However, such low-k dielectrics generally have a poor tolerance to process-induced damage due to their porosity. It is therefore important to develop less damaging etching chemistries as well as more resistant low-k materials. During the discharge, ion bombardment, active radicals and VUV radiations are responsible for the degradation of the low-k. Mainly, CH3 groups are removed from the film resulting in its hydrolyzation and consequent moisture uptake which has a severe impact on the final k-value after patterning.

In order to limit the damage during plasma etch, fluorocarbon-based chemistries allow to thin a protective etch-by-product film to be deposited on the surface of the low-k film during plasma etch and therefore effectively prevents possible damage by penetration of active radicals. We have evaluated the etching of a SiOCH dielectric with a k value of 2.0 using CF4 in combination with either CH4 or CHF3 and using H2, N2 or Ar as tuning gas. The plasma damage has been evaluated by measuring the loss of –CH3 groups by –CH2 groups –O–H uptake by FTIR and by measuring the k-value variation by forming MIS planar capacitor. Starting from a CF4/H2/N2 plasma, it has been found that the damage is reduced by substituting H2 by N2. H is able to diffuse deep into the low-k film and probably remove –CH3 groups by forming CH4. Switching to a CF4/CH3/N2 plasma further allows to decrease the damage, mostly probably through the formation of a thicker passivation layer on the low-k. Plasma damage can be further reduced by replacing N2 by Ar, leading to a CF4/CH3/Ar plasma. Possibly, N2 is causing damage by extracting C from the low-k film by forming HCN by-product. However the replacement of N2 by Ar resulted in a reduction of the etch rate which could be recovered by finally replacing CH4 by CHF3 while maintaining a low damage level.

Finally, 90nm half pitch trenches have been patterned into the SiOCH dielectric film using a dual hard mask approach (spin-on glass and spin-on carbon films). The influence of the hard mask on the plasma composition requires the optimization of the etch conditions in order to obtain high-quality profiles. Integrated k-value after complete processing will be discussed.

In conclusion, we have shown that H2 and N2 are responsible for the low-k damage during etch in fluorocarbon based plasmas. Switching to a CF4/CHF3/Ar plasma allows to reduce damage while maintaining good patterning capability.


Feature scale etch/deposition properties in plasma processing of microelectronic devices are determined by the energies and fluxes of radicals and ions to the wafer. These fluxes are ultimately controlled by the etching species, ion and neutral energies of the ions impacting the wafer. These values were used to evaluate the etching of a SiO2 wafer in Ar/CF4/O2 capacitive coupled plasma tool. Starting from a CF4/H2/N2 plasma, it has been found that the damage is decreased by substituting H2 by N2. H is able to diffuse deep into the low-k film and probably remove –CH3 groups by forming CH4. Switching to a CF4/CH3/N2 plasma further allows to decrease the damage, mostly probably through the formation of a thicker passivation layer on the low-k. Plasma damage can be further reduced by replacing N2 by Ar, leading to a CF4/CH3/Ar plasma. Possibly, N2 is causing damage by extracting C from the low-k film by forming HCN by-product. However the replacement of N2 by Ar resulted in a reduction of the etch rate which could be recovered by finally replacing CH4 by CHF3 while maintaining a low damage level.

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In conclusion, we have shown that H2 and N2 are responsible for the low-k damage during etch in fluorocarbon based plasmas. Switching to a CF4/CHF3/Ar plasma allows to reduce damage while maintaining good patterning capability.
Patterned structure etch testing was performed and optimized using a DOE approach. CF3I was found to be suitable to obtain nicely etched features. In many cases, better etched structures and higher selectivity were obtained using CF3I compared to the standard gases.

5:20pm PS1-TuA11 Mechanism of Highly Selective SiO2 Etching over Si,N4, Si and Photoresist Using Hydro-Fluorocarbon Gases. Y. Miyawaki, Y. Kondo, K. Asano, M. Sekine, K. Ishikawa, T. Hayashi, K. Takeda, H. Kondo, M. Hori, Nagoya University, Japan

Highly selective etch of dielectric films is one of the key technologies in integrated circuits fabrication. We achieved extreme high SiO2 selectivity against Si,N4, Si and ArF photoresist using hydro-fluorocarbon gas, CH,F2, and O2, Ar gas mixture [1] in a dual frequency (60 MHz / 2 MHz) capacitive coupled plasma (CCP). The selectivity over SiN4 seemed to be infinite because selective fluorocarbon deposition occurred on SiN4. The surface roughness that leads to line edge roughness was reduced by the selective deposition. The selective etch mechanism among plasma etch using fluorocarbon gases with and without hydrogen in gas molecule was investigated by the diagnostics of gas phase species and etched surfaces. As comparing CxFy, CxFyO and CHFxY gases, it was found that the hydrogen containing species with larger molecular weight play an important role for forming thick and carbon-rich polymer film especially on Si,N4, as well as on Si, photore sist. For the CHFxY gas plasma, the density of F radical that degrades the selectivity is low compared to the CxFy species. We speculated that the thick polymer formation on Si3N4, Si and photore sist. Here, we need some sacrifice layer to initiate that selective deposition. The photore sist surface suffered roughness formation in the beginning, then the roughness was cured by the further deposition with ion bombardment [2]. This protective and roughness curing deposition on photore sist should occur for CxFy gas plasma in some condition (e.g. higher gas mixing ratio of CF,Ar) however, it also forms a thick polymer on SiO2 to stop etching. The temperature dependence for the etching and deposition properties [3] suggested that the difference of sticking coefficient for species with and without hydrogen was not so important. It is also speculated that the thick polymer formation on SiN4, Si and photore sist while less reducing the SiO2 etch rate is due to the relative large flux of specific hydrogen containing species in C,FyH2 plasma.

Acknowledgement: Tokyo Electron Ltd., ZEON Corp., JSR Corp.


Plasma Science and Technology Division
Room: 201 - Session PS2-TuA

Plasma Diagnostics, Sensors and Control I
Moderator: V. Nagorny, Mattson Technology, Inc.

2:00pm PS2-TuA1 Prediction of Ion Sheath Shape and Ion Trajectory during Plasma Etching Processing using On-Wafer Monitoring Technique. R. Araki, K. Miwa, T. Kubota, Tokoh University, Japan, T. Iwashiki, K. Ono, Mizuhpo Information & Research Institute, Inc., Japan, S. Samukawa, Tokoh University, Japan

Precise plasma processes are indispensable for the fabrication of ULSI and MEMS devices. Some MEMS devices have larger scaled 3D structures comparable to the ion sheath thickness on the surface in plasma processing. In such cases, because of distortion of sheath shape due to the MEMS structure, ions trajectory are distorted to the surface and it causes etched shape anomaly. In order to solve these problems, we are developing a system to measure sheath thickness and to predict sheath shape, ion trajectory, and etched shape by fusion of a new on-wafer monitoring data and computer simulation. Our newly developed on-wafer sheath sensor can measure the surface potential and ion saturation current at wafer surface. Based on these results, the sheath thickness, shape, and ion trajectory were calculated by using our developed simulation. In this study, we could measure the sheath thickness, and then calculate the sheath shape deformation around a structure having large step in the case of using SF6 inductively coupled plasma. We found that the sheath thickness was about 1 mm and ion trajectory was bent near the large steps on the wafer surface. This result was completely corresponding to the actual etching pattern profile near a structure having large step. It is suggested that our proposed fusion system of on-wafer monitoring data and computer simulation is very effective to predict real etched shape during plasma etching processes.


Plasma density measurements are an essential tool in understanding and controlling processing plasmas across a wide range of applications. Charge collection probes (Langmuir probes) are of limited utility in depositing plasmas, high pressure applications or in processes that require the use of reactive gases, as these environments result in unreliable data acquisition. Plasma frequency probes are an attractive alternative to Langmuir probes in such applications since they do not suffer significant performance degradation in these environments. This work presents frequency probes measurements of plasma density over a range of 1010 to 1012 cm-3 in a variety of process plasmas, plasma chemistries (N2, CH4, NH3, O2 and SF6). In addition to electron density measurements frequency probes are also useful for measuring plasma potential, electron temperature, and electron energy distribution functions in the gas chemistries mentioned above.

2:40pm PS2-TuA3 Effects of Wire Thickness, Neutral Pressure and Gas Composition on the Inflection Point Technique. B. Dechawatapanisal, N. Hershkovitz, J.P. Sheehan, CS. Tip, University of Wisconsin-Madison

The inflection point technique in the limit of zero emission determines the plasma potential by fitting a straight line to the graph of the emission current versus the inflection point of the emissive probe I-V traces. The plasma potential is determined by extrapolating the line to the limit of zero emission. The effects of wire thickness, gas composition, neutral pressure and position on the technique were investigated. Experiments were performed in a multi-dipole filament discharge. Wire thicknesses of 0.013, 0.025, 0.05 and 0.01mm were studied. Experiments were done in Argon, Xenon and Helium plasmas at neutral pressures ranging from 0.5torr to 3Mtorr. Measurements were performed from the bulk of the plasma to the sheath edge near a 10cm diameter negatively biased plate.

This work is supported by U.S. DOE under the Grant and Contract Nos. DE-FG02-97ER55437 and No. DE-FG02-03ER55728, DE SC0001939 and by the National Science Foundation Grants under the Grant and Contract Nos. CBET-0903832, and No. CBET-0903783.

3:00pm PS2-TuA4 A New Diagnostic Tool System of Radio-Frequency Plasmas by Employing Floating-Emissive Probe. Y. Taniuchi, M. Usumi, Tokai University, Japan, M. Yanagisawa, Landmark Technology Corporation, Japan, H. Shindo, Tokai University, Japan

A new diagnostic tool to measure plasma parameters as well as Electron Energy Distribution Function (EEDF) by a floating-emissive probe has been proposed[1], and a diagnostic system has been newly developed and applied to radio-frequency (RF) plasmas. It is generally difficult for a conventional probe method to measure EEDF in RF plasmas, because of the plasma potential fluctuation, particularly in the capacitive mode. The present probe method to measure EEDF in RF plasmas, because of the plasma potential fluctuation, particularly in the capacitive mode. The present method has an advantage that there is no need of an external compensation circuit and all measurements can be made in the floating condition. The method is based on measurement of the functional relationship between the floating potential change ΔVf and the heating voltage VF of emissive probe. If the plasma electrons are in Maxwellian, the equation can be obtained for the value of ΔVf as a practical and useful formula.[1] It is important to know that the value of ΔVf contains information of electron energy distribution. In the experiment, the data of Vf and ΔVf was measured in a 13.56 MHZ RF plasma produced by single-loop antenna[2], as a function of Vf. In the conditions of high RF power, the plasma mode was ICP and the measured values of ΔVf were in agreement with the theoretical value, stating that the plasma electron was in Maxwellian. The electron temperatures thus obtained were very consistent with those measured by Langmuir probe. The electron density was also obtained from the value of ΔVf near the plasma space potential and they were consistent with Langmuir probe data. Consequently, by using a new diagnostic system one can obtain the electron temperature and density, the plasma space potential as well as floating potential, not only as EEDF in the floating condition of the probe. It should be stressed that this is the first success of floating probe to be able to measure all plasma parameters. One can also expect that the present method is applied to plasmas which are produced in insulated vessels.

References:
Accurate measurements of the plasma potential is a critical challenge especially for complex plasmas such as magnetized and flowing. We compare various emissive probe techniques for measurements of the plasma potential. The measurements were conducted under a low-pressure noble gas discharge of the Hall thruster. The thruster was operated with xenon gas in subkilowatt power range and the discharge voltage range of 200-400 V. The probe was placed at the channel exit where, the electron temperature is in the range of 10 to 60 eV and the plasma potential is in the range of 50 to 250 V. The floating point method is expected to give a value $-\epsilon e$ below the plasma potential. The experimental results are consistent with these expectations. Sperically, it is shown that the floating potential of the emissive probe is $-\epsilon e$ below the plasma potential. It is observed that the separation technique varies wildly and does not give a good measure of the plasma potential.

This work was supported by US Department of Energy grants No. DE-AC02-09CH11466, No. DE-FG02-97ER54437, and No. 3001346357 and the Fusion Energy Sciences Fellowship Program administered by Oak Ridge Institute for Science and Education under a contract between the U.S. Department of Energy and the Oak Ridge Associated Universities.

**Advanced Surface Engineering Division**

**Room: 104 - Session SE+TF-TuA**

**Glancing Angle Deposition (GLAD) II**

**Moderator:** N.A. Beckers, University of Alberta

**1:50pm SE+TF-TuA1 Evolution of Crystal Orientation during Oblique Angle Deposition. G.-C. Wang, T.-M. Lu, Rensselaer Polytechnic Institute**

The oblique angle deposition (OAD) has attracted attention due to the shadowing effect that results in interesting morphology of isolated nanostructures. These artificial nanostructures have rich textures depending on the growth conditions such as the angle of incidence, substrate temperature, deposition rate, and substrate rotation speed or mode. These OAD films often have biaxial textures even on amorphous substrates because the in-plane symmetry is broken under the off-normal incidence of the deposited atoms on the substrate. The most frequently used characterization technique for biaxial texture films is x-ray pole figure analysis. However, the texture obtained by x-ray is an average texture from the entire thickness of the film due to the x-ray’s few micron penetration depth. As the texture of a film often changes during growth, information on the basic mechanisms that control the final texture is often lost. In this talk we will show how we can use our newly developed reflection high energy electron diffraction (RHEED) surface pole figure technique to probe the surface texture evolution of the growth front from the initial stage (a few nm thick) to the later stage of thick films [1] either in situ or ex situ. Examples of biaxial texture evolution of CaF$_2$ [2], Mg, W [3] and Mo nanostructures as a function of thickness, incident angle, and rotation schemes, will be presented. The surface sensitive RHEED pole figure technique also reveals that biaxial texture can be formed at thickness as small as 20 nm. In some cases, surface texture at the growth front is seen to be quite different from the bulk texture measured by x-ray pole figures. We also show that these biaxial textured films can be used as buffer layers to grow near-single crystal semiconductor films which may find important applications in energy conversion devices.

Work was supported by NSF-0506738.


deposition (VLS-GLAD), of fabricating crystalline semiconductor nanowire arrays with controlled geometry. VLS-GLAD is a physical vapor deposition based semiconductor fabrication technique which relies on selective deposition of source atoms onto metal catalyst nanosized lands placed on a crystal wafer. In this technique, collimated obliquely incident flux of source atoms selectively deposit on catalyst islands by using "shadowing effect". Geometrical showing effect combined with VLS growth mechanism leads to the growth of crystalline semiconductor nanowire arrays. In this work, we show the morphological and structural properties of tilted single crystal semiconductor nanowire arrays fabricated by utilizing a conventional thermal vaporization system for VLS-GLAD.

3:00pm SE+TF-TuA4 Lithographic Processing of Nanostructured Thin Films Grown Using Oblique Angle Deposition Method. P. Shah, University of Dayton Research Institute, A. Sarangan, University of Dayton

It is known that exposing structured thin films (STF) grown using oblique angle deposition (OAD) to liquids such as DI water or any solvents permanently deforms the physical structure of the thin films and alters their properties. This is a severe limitation of STFs because the films cannot be patterned into useful devices using conventional wet lithographic processes. In this work, we overcome this challenge and propose to demonstrate conventional i-line lithography technique for patterning STF's grown using OAD. The ability to selectively fabricate STF in chosen areas of the active devices will be beneficial for numerous applications. It is shown that the structure of these thin films is preserved after lithographic processing. Processing limits in terms of dimensions of the devices or patterns that may possibly be fabricated are discussed.

4:00pm SE+TF-TuA7 Automated Measurement Technique for Growth Scaling in Glancing-Angle Deposited Films. J.M. Siewert, J.M. LaForge, M.T. Taschuk, M.J. Brett, University of Alberta, Canada

With growing applications of glancing-angle deposition (GLAD) thin films, there is increasing need to understand and engineer GLAD growth mechanisms. Devices that make use of GLAD films, such as RH sensors, optical filters, and solar cells require precise knowledge and control of GLAD nanostructure. Typically, GLAD vertical post diameters are described by a power law, \( d = w_0 p^h \), where \( d \) is column diameter, \( h \) is height, and \( w_0 \) and \( p \) are material dependent constants which describe column diameter and broadening, respectively [1, 2]. Based on theoretical growth models, \( p \) is expected to range from 5/16 to 1/2. While many GLAD materials have been characterized for \( p \), reliable trends have not yet been obtained [2,3,4,5].

Most groups measure \( p \) from cross-sectional SEM images, recording diameters of clearly distinct posts as a function of height. This technique is labour-intensive and large scatter in the data has been observed [4]. One alternative, focused ion-beam cross sections of GLAD films [5], produces very precise measurements but is impractical for characterizing the ensemble properties of a GLAD film. In both cases, the limited number of measurements restricts what can be learned about GLAD films. To eliminate this limitation, we have been developing an automated characterization method for GLAD posts that enables effective measurement of thousands of posts. Using this technique, we have measured \( p \) and \( w_0 \) as a function of pitch for 1500 nm TiO₂, 81° deposition angle, vertical post films. We examine post growth for “extinct” and full height posts, discovering markedly different growth scaling behaviors. Characterization of phi-sweep tilted posts structures is underway. We will present the optimized technique, current experimental results, and comparisons of automated measurements with existing measurement methodologies.


2:00pm SS+EM-TuA1 The Effect of Structure on Interfacial Energy. F. Sfeilacei, EPFL, Switzerland

In this talk I will highlight recent result on solid-liquid interfaces where the solid has a nanostructure with a characteristic length scale that is only a few liquid molecule wide. The talk will start with the presentation of a new scanning probe technique able to measure interfacial energy (work of adhesion to be precise) with atomic/molecular resolution. It will then show how surfaces with alternating stripe-like domains a few nanometer thick have a structural component such as to their work of adhesion that can account for as much as 20% of the total energy. Finally novel self assembly approaches to achieve such surfaces will be discussed.

2:40pm SS+EM-TuA3 Interfacial Electronic Structure of Dipolar Organic Semiconductors. O.L.A. Monti, M.P. Steele, N. Ilyas, L.L. Kelly, D.A. Racke, University of Arizona

We present an investigation of the evolution of the electronic structure at the interface of the dipolar organic semiconductor vanadyl naphthalocyanine with both highly oriented pyrolytic graphite and Au (111). Using angle-resolved two-photon photoemission and other photoelectron spectroscopies we observe both excitonic as well as strictly interfacial states in both ground and excited state manifolds, with large differences between the two surfaces. Simple electrostatic considerations provide a chemisorption model that is capable of quantitatively describing long- and short-range interface-mediated intermolecular coupling, significantly altering the molecular electronic structure. Additional insights are available from full-scale first-principles calculations at these interfaces. As a consequence, we show that electrostatic multipoles can significantly influence molecular and interfacial electronic structure, with direct and observable impact on interfacial charge-transfer dynamics. Interfacial electronic fields may therefore be used to manipulate in a concrete fashion processes of critical importance to solar energy conversion such as photoinduced interfacial electron transfer.


There is considerable interest in the functionalization of metal surfaces by molecules with large dipole moments. For this purpose, p-benzoquinonemonoxygen-type zwitterions represent ideal candidates. These zwitterions can be anchored on gold surfaces where they form homogenous thin films with the dipole preferentially oriented along the surface normal. Some zwitterions will selectively adsorb on patterned gold substrates from
solution while selective deposition of the zwitterions onto specific ferroceneic domains has been demonstrated. This represents an attractive approach to pattern molecular deposition on transparent planar substrates through electrostatic dipolar interactions or orientation dipole controlled surface chemistry. Taking advantage of the high solubility of two zwitterions in both organic solvents and water, we studied the influence of the solvent on the functionalization of surface.\(^{[2]}\) The goal is to control both packing and selective deposition on a variety of substrates through zwitterion solvent combination.

Acknowledgement
This research was supported by the CNRS and the Ministère de la Recherche et des Nouvelles Technologies, the ANR (07- BLAN-0274-04), the National Science Foundation (grants CHE-0909580 and DMR-0851703), and the Nebraska Center for Materials and Nanoscience at the University of Nebraska-Lincoln.

4:00pm SS+EM-TuA7 Directed Organization of C\(_{60}\) Kagome Lattice by TiOPc-Monolayer Template. J.E. Reutt-Robey, T. Wei, University of Maryland

Abruptly-ordered semiconductor interfaces between titanyl phthalocyanine (TiOPc) and C\(_{60}\) were prepared by physical vapor deposition and characterized by UHV-STM. Molecular TiOPc is a highly anisotropic molecule with a 3.5 dipole moment. Ordered TiOPc monolayer films of the honeycomb phase thus represent a regular 2-d dipolar lattice, which was investigated as an electrostatic template for the growth of the highly polarizable C\(_{60}\) films. A single-layer grown layer-by-layer revealed the directed formation of a Kagome lattice. Atomically detailed structural models were obtained for the 0-6 nm C\(_{60}\) thickness (up to 5 ML) range over which the ordering influence of the TiOPc dipolar substrate persists. Unusually low-density C\(_{60}\) molecular packing arrangements result from the ellipsoidal shape, curved surfaces and high polarizability of C\(_{60}\). While Kagome lattices have been frequently observed in colloidal and magnetic systems, this represents the first electrostatically-induced Kagome lattice involving a molecular film.

This work has been supported by the National Science Foundation under Surface Analytical Chemistry grant CHE0750203 and under the University of Maryland MRSEC DMR-05-20471

4:20pm SS+EM-TuA8 Solvation-Assisted Growth of ZnPc and 4NO\(_2\)-ZnPc Monolayer Films on Au(111). L. Tskipuri, Q. Shao, J.E. Reutt-Robey, University of Maryland, College Park

The UHV spray-jet molecular beam deposition technique marries the chemical versatility and efficiency of solution-film processing with the atomic-level control and analytical tools of UHV surface science. Here we show how the solvent used in aerosol deposition can be used to fine tune atomic-level control and analytical tools of UHV surface science. Here we investigate how the solvent used in aerosol deposition can be used to fine tune atomic-level control and analytical tools of UHV surface science. Here we investigate how the solvent used in aerosol deposition can be used to fine tune atomic-level control and analytical tools of UHV surface science. Here we investigate how the solvent used in aerosol deposition can be used to fine tune atomic-level control and analytical tools of UHV surface science. Here we investigate how the solvent used in aerosol deposition can be used to fine tune atomic-level control and analytical tools of UHV surface science. 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Surface Science Division
Room: 109 - Session SS-TuA

Catalysis on Metals and Alloys
Moderator: G. Fisher, University of Michigan

2:00pm SS-TuA1 La1-xCexMnO3. Perovskite Structural Fates and Performance for Preferential CO Oxidation Reaction. S.S. Maluf, Federal University of Sao Carlos, Brazil, B.E. Hayden, University of Southampton, UK, C.R.M. Afsanno, Federal University of Sao Carlos, Brazil, E.M. Assaf, University of Sao Paulo, Brazil, P.A.P. Nascente, Federal University of Sao Carlos, Brazil

The perovskite structure is characterized by a large capacity to stabilize unusual valence states of different metal ions and can also accommodate variable amounts of different lattice defects. Several lanthanum transition metal based perovskites, represented as La$_x$A$_{1-x}$MnO$_3$, have been known as very good oxidation catalysts in a variety of reactions. Particularly, LaMnO$_3$ perovskites have shown to be the most active. The catalytic activity of these compositions in different reactions can be further enhanced by the substitution of lanthanum by cerium. The catalysts were prepared by co-precipitation method from aqueous solutions of La, Mn, and Ce nitrates with Na$_2$CO$_3$/NaOH solution (pH 10). The prepared samples were La$_x$Ce$_{1-x}$MnO$_3$ (x = 0, 0.05, and 0.10) and were characterized by EDS-SEM, surface area BET method, X-ray diffraction (XRD), and temperature programmed reduction (TPR). The catalytic activity was tested over the temperature range of 130-230°C in a fixed-bed tubular glass micro-reactor, with 200mg of catalyst. The reaction products were analyzed in-line by gas chromatography. The amounts of the constituent metals obtained by EDS were in agreement with expected values; the presence of cerium did not change the surface area of the samples (26-27m$^2$ g$^{-1}$). The X-ray diffraction lines corresponded to cubic LaMnO$_3$ structure; the replacement of La did not change the diffraction peaks. Peaks of cerium oxide were not observed, suggesting the incorporation of Ce ions into the LaMnO$_3$ lattice. The sample with the highest content of cerium presented an increase on thermal stabilization of Mn$^{4+}$ ions, and Mn$^{3+}$ species were reduced to lower temperatures. The SEM results indicated that the presence of cerium decreased slightly the grain size (from 56 to 43 nm) and the range of distribution sizes is homogeneous and the most grains are spherical. All samples presented the same catalytic behavior, starting their activities at 130°C, and their activities increased with the temperature. At 150°C, the sample with 5% of Ce presented a slight advantage compared to other samples, and this was more significant at 230°C, indicating that the promoter effect of cerium can be observed for higher reaction temperatures. The results suggested that the replacement of La by Ce caused an increase of cation/anion vacancies and also in the Mn$^{4+}$/Mn$^{3+}$ ratio change, and consequently the catalytic behavior for CO oxidation changed, facilitating the CO adsorption.

Acknowledgements

The authors thank FAPESP for financial assistance, Ilika Technologies and University of Southampton for the XRD analyses.

2:20pm SS-TuA2 Faceted Metal Surfaces: Surface Chemistry and Growth of Metallic Nanoclusters. W. Chen, Q. Shen, R.A. Bartynski, Rutgers University

In this work, surface faceting is used to prepare a nanoscale model catalyst for surface reactions and a nanoscale template for growth of metallic nanoclusters. Faceting occurs when an initially planar surface converts to a "hill and valley" structure exposing new crystal faces of nanometer scale dimensions. Nanoscale three-sided pyramids exposing (311), (31-1) and (110) faces with tunable facet sizes are formed on an initially planar Ir(210) surface upon annealing in O$_2$ and a clean faceted Ir(210) surface can routinely be prepared in situ by heating in H$_2$. Surface reaction of CO has been carried out on clean planar Ir(210) and clean faceted Ir(210) with varying facet size (5-14nm). Both planar and faceted Ir(210) favor reduction of NO by CO with high selectivity to N$_2$, which is accompanied by simultaneous oxidation of CO. The reaction is not only structure sensitive on faceted Ir(210) versus planar Ir(210) but also exhibits size effects on faceted Ir(210) for average facet size ranging from 5nm to 14nm without change in facet structure. Strong interaction between NO and CO at high NO exposure and IML CO pre-coverages result in “explosive” evolution of N$_2$ and CO$_2$ on planar Ir(210). Annealing Ru(11-20) in NO$_2$ leads to formation of ridge-like faceted surface exposing (10-11), (10-1-1), (01-1-1) and (01-1-1) faces. Faceted O/Ru(11-20) facilitates growth of nanoscale metallic clusters at room temperature which preferentially nucleate within valleys of the faceted surface.

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2:40pm SS-TuA3 Modeling the Complexities of Heterogeneous Catalysts. D.W. Goodman, Matt Lundwall, Texas A & M University

INVITED

In the past several decades, surface science methodologies have contributed significantly to our understanding of reaction mechanisms of heterogeneous catalysts. From fundamental studies on metal single crystals to investigations of supported metal clusters, the relative complexities of model catalyst systems have continued to evolve in complexity and likeness to industrial catalysts. Studies on metal single crystals have aided researchers in understanding the effect of surface structure on catalyst reactivity and selectivity for a number of important reactions. More recently model systems consisting of metal clusters deposited on planar oxide surfaces under ultra high vacuum conditions have facilitated the study of metal oxidation effects. These model systems are useful for carrying out kinetic investigations, yet are amenable to surface spectroscopic techniques, thus enabling investigations under realistic pressures and at working temperatures. This talk will present selected investigations of catalytic reactions on single crystal and model supported catalysts with emphasis on recent work addressing the nature of the active site in structure sensitive reactions.

4:00pm SS-TuA7 Nanocatalysis: New Developments in Catalytic Performance of Size- and Shape-Controlled Metal Nanoparticles. B. Roldán Cuenya, University of Central Florida

Tailoring the chemical reactivity of nanomaterials at the atomic level is one of the most important challenges in catalysis research. In order to achieve this elusive goal, fundamental understanding of the geometric and electronic structure of these complex systems at the atomic level must be obtained. The study of metal nanoclusters, the relative complexities of model catalyst systems have continued to evolve in complexity and likeness to industrial catalysts. Studies on metal single crystals have aided researchers in understanding the effect of surface structure on catalyst reactivity and selectivity for a number of important reactions. More recently model systems consisting of metal clusters deposited on planar oxide surfaces under ultra high vacuum conditions have facilitated the study of metal oxidation effects. These model systems are useful for carrying out kinetic investigations, yet are amenable to surface spectroscopic techniques, thus enabling investigations under realistic pressures and at working temperatures. This talk will present selected investigations of catalytic reactions on single crystal and model supported catalysts with emphasis on recent work addressing the nature of the active site in structure sensitive reactions.

4:20pm SS-TuA8 Oxidative-Coupling Reactions via Nucleophilic Attack on Gold Surface. B. Xu*, C.M. Friend, R.J. Madix, Harvard University

Metallic gold owes its centuries-old mystique and intrinsic value to its chemical inertness toward bulk compound formation. In the past decade, however, it has been discovered that the surface of gold is far from inert, and that gold can catalyze important chemical transformations – particularly with the assistance of molecular oxygen or other oxidizing agents. These processes are of particular significance because they may occur at remarkably low temperatures and pressures, suggesting the possibility of energy efficient and environmentally benign reaction conditions using metallic gold catalysts. Recently, there has been a focus on selective
oxidation of alcohols to aldehydes, oxidative self-coupling of alcohols to form esters and, most recently, acylation via amine-formaldehyde coupling. Our work under high vacuum in UHV or on Au(111) surface has unequivocally proved that surface adsorbed atomic oxygen is critical in facilitating a range of nucleophilic reaction–reactions among alcohols, aldehydes and amines. A general reaction mechanism for this class of coupling–reactions is established: surface alkoxy or amide is formed via the deprotonation of the corresponding alcohols or amines by surface atomic oxygen, which can subsequently nucleophilically attack the aldehydes (formed in situ from alkoxy or introduced directly) and produce the corresponding ester or amide. The product distribution of our low-pressure experiments match remarkably well with gas phase reactions carried out in the ambient condition as well as liquid phase reactions, proves the generality of the mechanism.

The key features of Cu-Au-Pd alloys relevant to their application as National Energy Technology Laboratory (NETL) catalysts have been studied by scanning tunneling microscopy, temperature programmed desorption and density functional theory calculations. Methanol reaction on 0.25 ML Au clusters produces formaldehyde as the major product at 535 K with methanol desorption observed at the same temperature; water and hydrogen evolution are detected below room temperature, as well as methyl radical evolution at 575 K. On the titania surface itself, methanol reaction also produces methyl radical around 600 K. Formaldehyde production reaches its maximum value between 0.25 and 2 ML, but decreases dramatically as the coverage is increased to 5 ML. This behavior suggests that formaldehyde is produced at the Au-titania interface since formaldehyde production does not occur at pure Au sites. STM experiments confirm that the 0.25 ML Au coverage has the greatest number of Au-titania interface sites at the perimeter of the clusters. For higher coverages of 2 and 5 ML, cluster coalescence diminishes the number of Au-titania sites, and this is consistent with the decreased formaldehyde yield at higher coverages. When the titania surface is reoxidized with O2 prior to Au deposition and exposure to methanol, lattice oxygen is incorporated into the water that is evolved at low temperature. We propose that the role of the titania support is to facilitate the formation of the reactive methoxy intermediate via abstraction of the hydroxyl hydrogen in methanol by lattice oxygen. Density functional theory calculations also indicate that methoxy is the intermediate immediately at the Au-titania interface after O2 bond scission is induced by lattice oxygen.

The changes induced in the structure and chemical state of size-selected Pt nanoparticles deposited on rutile TiO2(110) have been studied by scanning tunneling microscopy, temperature programmed desorption and density functional theory calculations. Methanol reaction on 0.25 ML Au clusters produces formaldehyde as the major product at 535 K with methanol desorption observed at the same temperature; water and hydrogen evolution are detected below room temperature, as well as methyl radical evolution at 575 K. On the titania surface itself, methanol reaction also produces methyl radical around 600 K. Formaldehyde production reaches its maximum value between 0.25 and 2 ML, but decreases dramatically as the coverage is increased to 5 ML. This behavior suggests that formaldehyde is produced at the Au-titania interface since formaldehyde production does not occur at pure Au sites. STM experiments confirm that the 0.25 ML Au coverage has the greatest number of Au-titania interface sites at the perimeter of the clusters. For higher coverages of 2 and 5 ML, cluster coalescence diminishes the number of Au-titania sites, and this is consistent with the decreased formaldehyde yield at higher coverages. When the titania surface is reoxidized with O2 prior to Au deposition and exposure to methanol, lattice oxygen is incorporated into the water that is evolved at low temperature. We propose that the role of the titania support is to facilitate the formation of the reactive methoxy intermediate via abstraction of the hydroxyl hydrogen in methanol by lattice oxygen. Density functional theory calculations also indicate that methoxy is the intermediate immediately at the Au-titania interface after O2 bond scission is induced by lattice oxygen.

This study hints on how the surface arrangements of atoms and the influence of ensembles on the reaction kinetics and energetics can offer insight to properly tailor CO-tolerant Au-based bimetallic catalysts for fuel cell applications.

Thin Film Division
Room: 107 - Session TF-TuA

ALD: Fundamental Reactions and Film Properties
Moderator: S.M. George, University of Colorado, Boulder

2:00pm TF-TuA11 Indium Oxide Atomic Layer Deposition Facilitated by the Synergy between Oxygen and Water, J.A. Libera, J.N. Hryn, J.W. Elam, Argonne National Laboratory

This study describes how In2O3 films can be prepared by ALD using alternating exposures to cyclopentadienyl indium (In Cp) and combinations of H2O and O2, even though H2O and O2 are ineffective when used individually. Nanostructured photovoltaics would benefit enormously from the capability to deposit conformal indium-tin oxide (ITO) films inside of high aspect ratio structures. For instance, our previous In2O3 ALD method using In Cp and O2 enabled ITO growth inside of anodic alumina membranes producing higher photocurrents through radial charge collection in dye sensitized solar cells. However, this process yielded poor thickness conformity due to the In2O3-catalyzed thermal decomposition of O2, and this shortcoming motivated our search for an O2-free process. We were surprised to discover that when H2O and O2 were used together, either as a simultaneous exposure (SE) or in the sequence O2-H2O2-WO (WO) or O2-WO (OW), very uniform, highly conducting In2O3 films were deposited at 1.0-1.6 Å/cycle over large areas (12"x18") at temperatures as low as 100°C. In-situ quartz crystal microbalance, mass spectrometry, and Fourier transform
infrared measurements revealed that the H₂O and O₂ work synergistically to facilitate the In₂O₃ ALD. Each molecule performs a necessary but distinct role in the growth mechanism. Next, we conducted a thorough study of this process for all three growth modes (SE, WO, OW) over the temperature range 100-250°C using spectroscopic ellipsometry, ultraviolet-visible transmission, X-ray diffraction, scanning electron microscopy, and Hall probe measurements. These measurements identified a remarkable correlation between the film structure and electrical properties around an amorphous-to-crystalline phase transition near the deposition temperature of 140°C.

2:20pm TF-TuA2 Engineering AlN Thin Films by Atomic Layer Deposition on Wide Bandgap Semiconductors as Gate Dielectric. Y.-C. Perng, J.P. Chang, University of California Los Angeles

Wide bandgap semiconductors, such as SiC and GaN, are known as based materials in electronic devices operating under high temperatures and high electric fields. Atomic layer nitride (AlN) is a promising dielectric material for SiC and GaN due to its material properties, such as wide bandgaps, similar atomic arrangement and small lattice mismatch (1.3% and 2.6%). Although various deposition methods, such as chemical vapor deposition (MOCVD) and molecular beam epitaxy (MBE), have been used, synthesizing high quality AlN thin films for those applications is still an issue, especially in the view of controllability in atomic scales. Atomic layer deposition (ALD) is used to synthesize AlN thin layer on SiC, GaN and Si to achieve atomic controllability and assess the possibility to extend this deposition method to nitride growth and how the substrate properties affect the interface quality and corresponding electrical properties of the synthesized film.

The AlN thin films were synthesized at 400–600°C via a chamber with the base pressure as 10⁻¹⁰ torr, using trimethylaluminum (TMA) and blue grade ammonia (NH₃) as precursors. The surface structure during the deposition was monitored via in-situ reflection high electron energy diffractometry (RHEED) and the growth rate was 0.5–2 Å/cycle as a function of the deposition temperature. The surface composition, including the impurities, was found to be correlated to the deposition temperature, verified by in-situ x-ray photoelectron spectroscopy (XPS). The surface morphology of the films was studied by transmission electron microscopy (TEM) and atomic force microscopy (AFM) and found to be conformal because the rms value was found to be less than 10 Å on SiC, which is the same as that of the substrate surface roughness. The capacitance/conductance-variation voltages were measured to determine the dielectric constant of the AlN film and interpret the interface states density, which were 8 and on the order of 10¹⁵ cm⁻²eV⁻¹, respectively, comparable to that reported for MBE synthesized AlN.

2:40pm TF-TuA3 Paul Holloway Award Lecture - Gas-Surface Interactions during Atomic Layer Deposition. S. Agarwal, Colorado School of Mines INVITED

Atomic layer deposition (ALD) is an ideal technique for depositing amorphous and polycrystalline films that require atomic layer control over the film thickness and chemical composition, and can provide conformal coverage with sharp interfaces on high-aspect-ratio nanostructures. ALD has been used to deposit oxide, nitride, elemental and compound semiconductor material films for various applications. The main challenges in ALD are control over the film’s microstructure and residual impurities, which in turn affect their structural, electrical, and transport properties. To overcome these problems, an understanding of atomic-scale events that occur during growth is required. In this presentation, we will discuss the mechanisms of the surface reactions of radicals and molecular precursors supplied during growth using a suite of in situ diagnostic tools. We will specifically discuss the reaction mechanisms during plasma- and ozone-assisted ALD of TiO₂, Al₂O₃, and SiO₂ and how such reactions influence the film properties.

4:00pm TF-TuA7 Mechanical Properties of ALD Thin Films. H. Baumgart, Old Dominion University INVITED

Atomic Layer Deposition (ALD) is recognized as the preferred method to deposit technologically important thin films of novel high-k dielectric metal oxides or semi-insulating metal oxides for growth rates of 0.2 Å/cycle with Angstrom accuracy. Large bodies of extensive studies exist on the electrical characterization of ALD films, however, there exists a lack of systematic studies regarding the mechanical properties of ALD grown thin films. Elevated temperatures cause phase changes in many ALD metal oxide films, which affect the mechanical properties and surface morphologies. Little is known about the impact of those phase changes on the nanomechanical properties of ALD HfO₂. Phase change of ALD HfO₂ impacts the mechanical and electrical properties of high-k dielectric gate insulators depending on whether a Gate First or Gate Last process integration has been adopted. Nanoinindentation is the most appropriate testing mechanism that accurately investigates the mechanical properties of extremely thin film specimens as microcrystalline films. Nanoinindentation testing was conducted to investigate the impact of the different phase changes of HfO₂ on the mechanical properties. We have deposited ALD HfO₂ at low temperature and measured the mechanical properties of the various phase changes of HfO₂ following various thermal annealing cycles. After crystallization by annealing in a rapid thermal annealing (RTA) chamber, the modulus was found to decrease from 370±20GPa to 240±20GPa as the HfO₂ films transition from amorphous to polycrystalline structure past the phase change transition temperature of 600°C. Similarly, the hardness measurements reveal a high value of 18±1GPa for amorphous HfO₂ films and a decrease to 15±1GPa following the transition temperature to polycrystalline HfO₂ films.

Piezoelectric films such as ALD ZnO are finding applications in microelectromechanical systems (MEMS), piezoelectric transducers and various micro-electronic devices. ZnO piezoelectricity involves internal generation of electrical charge resulting from an applied mechanical force deforming the static structure of the ZnO crystals, studying the mechanical properties of novel ALD thin films of ZnO is important for those technical applications. Obtaining a better understanding of the structural and mechanical properties of novel ALD ZnO films is essential to improve key performance parameters of MEMS micro-devices.

An overview of the measured mechanical properties of selected ALD thin films will be presented. The mechanical properties of ALD thin films differ significantly from published values of bulk material.

5:00pm TF-TuA10 In Situ Studies of Oxide ALD for Crystalline Oxide Growth on Silicon. B.G. Willis, H. Wang, University of Connecticut, C. Zhang, Chinese Academy of Sciences, China, X. Jiang, University of Connecticut INVITED

Thin films of crystalline oxides integrated with silicon are of interest for novel electronic devices that utilize a rich variety of functional properties of oxide materials. These useful properties include piezoelectricity and ferroelectricity among others. Growth of crystalline oxides on silicon requires careful control of a nanometer thin transition region between the film and substrate to avoid deleterious side reactions such as oxidation of the substrate. Submonolayer surface reconstructions of alkaline earth metals such as Sr are critical for controlling the interface for successful growth. In this paper, we show how atomic layer deposition (ALD) can be used to precisely control the interface reactions for enthalpic growth. X-ray photoelectron spectroscopy and in-situ spectroscopic ellipsometry are used to monitor the growth and interface evolution of oxides deposited by ALD. Ellipsometry provides real-time feedback on sequential adsorption cycles including precursor adsorption and oxidation. X-ray photoelectron spectroscopy provides detailed insight into the evolution of the interface structure during annealing and oxidation. Using this high degree of...
control enables the successful deposition of critical Sr/Si reconstructions for heteroepitaxy. The results show that ALD crystalline oxide heteroepitaxy is promising.

5:40pm **TF-TuA12 Nucleation and Interface Formation of Al2O3 on HF-treated InGaAs(100) by Atomic Layer Deposition, A.J. Muscat, B. Granados, F.L. Lie, University of Arizona**

III-V compound semiconductors are alternative channel materials for high-speed, low-power digital logic devices because their electron transport and breakdown properties are superior to that of Si. However, robust surface passivation processes and thermodynamically stable interfaces are barriers to its wider adoption. Additionally, the scaling down of integrated circuits has prompted the use of alternative high-k dielectric films to replace SiO2 as the gate in metal-oxide-semiconductor field effect transistors (MOSFETs). Recent work demonstrated oxide removal and passivation of III-V surfaces by depositing high-k dielectrics using atomic layer deposition (ALD).

In this study, the ALD of Al2O3 was investigated on liquid and vapor HF-etched In0.53Ga0.47As(100) samples. Both half and complete ALD cycles of trimethylaluminum (TMA) and H2O at 170°C were used to better understand nucleation and film growth. Aqueous HF etching was performed by a 49% HF dip 1 min and 15 s water rinse. In situ gas phase HF/H2O etching was run at 29°C and 100 Torr with an HF to water partial pressure ratio of 1.23.

The initial 8.0±1.4 Å-thick native oxide contained 21% In, 27% Ga, and 52% As oxides and was reduced to a 4.5±1.5 Å-thick oxide containing 91±7% As by aqueous HF. In contrast, the gas phase HF produced 7 Å-thick mixed oxide and fluoride overlay containing 30% In, 40% Ga, and 30% As.

Large reductions of substrate oxides were observed after the first TMA pulse on both liquid and gas phase HF-treated samples. The intensity of the O 1s XPS peak was constant but the peak shifted by 1 eV to higher binding energy (BE) due to the conversion of the oxide to Al2O3. On the gas phase HF-treated samples removal of In, Ga and As atoms in the fluoride-rich overlay layer was also observed after the first TMA pulse. The intensity of the F 1s peak was reduced and the peak shifted by 2 eV to higher BE, indicating the etching of fluoride as well as the conversion of the bonding from substrate fluorphides to Al-F.

Subsequent H2O and TMA pulses up to three cycles of TMA/H2O revealed a systematic peak shift of the overlay atom signals. F 1s, O 1s and Al 2p peaks shifted 0.9 eV, 0.4 eV and 0.2 eV, respectively, towards lower BE after a H2O pulse and shifted back to their original positions after a TMA pulse. The systematic shifts could be attributed to the change in surface termination after every half-cycle reaction, methyl termination after the TMA pulse and hydroxyl termination after a water pulse. Understanding surface reactions involved in the nucleation phase and early cycles of ALD is important in achieving control of the III-V dielectric interface.

**Vacuum Technology Division**

**Room: 111 - Session VT+MN+NS+SS+AS-TuA**

**Surface Science for Future Electronic Materials and Accelerator Applications**

Moderator: M. Wüest, INFICON Ltd, Liechtenstein

2:00pm **VT+MN+NS+SS+AS-TuA1 New UHV Low Temperature Scanning Probe Microscopy Facility for the Study of Future Electronic Materials, J.A. Stroscio, National Institute of Standards and Technology, INVITED**

Since the beginning of the last century new frontiers in physics have emerged when advances in instrumentation achieved lower experimental operating temperatures. Notable examples include the discovery of superconductivity and the integer and fractional quantum Hall effects. New experimental techniques are continually adapted in order to meet new experimental challenges. A case in point is scanning tunneling microscopy (STM) which has seen a wealth of new measurements emerge as cryogenic STM instruments have been developed in the last two decades. In this talk I describe the design, development and performance of a scanning probe microscopy facility operating at a base temperature of 10 mK in magnetic fields up to 15 T [1]. The STM system can be connected to, or disconnected from, a network of interconnected auxiliary UHV chambers used for sample and probe tip preparation. Results from current measurements on graphene and topological insulators will be described.

3:00pm **VT+MN+NS+SS+AS-TuA4 Surface Issues for Solid Niobium SRF Accelerator Cavities, M. Kelley, College of William and Mary**

The world-wide physics community looks forward to a slate of accelerator protocols of unprecedented magnitude and diversity. Certainly its sheer size makes the International Linear Collider the most visible to the public eye, with 16,000 solid niobium cavities performing at historically high gradient, and built (and operated) for historically low unit cost. Net performance makes superconducting radiofrequency (SRF) technology the approach of choice.

Solid niobium is the material most widely used for construction of SRF cavities because it has the highest critical transition temperature (Tc = 9.2 K) of the pure metals, sufficiently high critical magnetic field (Hc2 > 2 k Oe) for SRF accelerator applications, and metallurgical properties adequate for fabrication and service load. Studies of the SRF performance of niobium cavities began to be reported more than 30 years ago and continue now with the application of improved experimental techniques. Niobium metal superconductivity is a nanoscale, near-surface phenomenon because of the shallow RF penetration. Considerable evidence indicates that cavity interior surface chemistry and topography strongly impact SRF accelerator performance. This talk will motivate investigation of how they are affected by post-fabrication treatments. Current status and prospects are discussed with respect to accelerator needs and opportunities.
Examples of Surface Related Reactivity on Nb Samples and SRF Cavities for Particle Accelerators at JLab

This contribution will review some examples of surface related R&D on small and flat niobium (Nb) samples and single cell Nb superconducting radio frequency (SRF) cavities done at Jefferson Lab in the past few years. Most of the surface measurements were performed via the experimental systems available in the surface science lab that was set up at JLab to study the various problems on the Nb surfaces in the SRF field.

The first topic is about a new Nb surface polished technique called buffered electro-polishing (BEP) that was developed at JLab. This technique can produce the smoothest surface finish ever reported in the literature. It was also demonstrated that under a suitable condition, a Nb removal rate higher than 10 μm/min could be realized. Efforts have been made to try to understand the polishing mechanism through experiments with a well defined experimental geometry on small flat Nb samples. A unique versatile vacuum polishing system was constructed to perform BEP on Nb single cell cavities. Small flat samples, Nb dumbbells and Nb single cell cavities were also studied and treated at CEA Saclay in France and Peking University in China and the cavities were RF tested at JLab. Experimental results will be analyzed and summarized. It is showed that BEP is a very promising candidate for the next generation surface polishing technique for Nb SRF cavities.

A second topic will deal with a new Nb surface cleaning technique employed gas cluster ion beam (GCIB) treatment. This is a result of collaboration with Epion Corporation, Fermi Lab, and Argonne Lab. Beams of Ar, O2, N2, and NF3 clusters with accelerating voltages up to 35 kV were employed in this technique to bombard Nb surfaces. The treated surfaces of Nb flat samples were examined by several surface experimental systems such as SEM, EDX, AFM, SIMS, and 3-D profilometer. The experiments revealed that GCIB technique could not only modify surface morphology of Nb, but also change the surface oxide layer structure of Nb and reduce the amount of field emission sites on the surface dramatically. Computer simulation via atomistic molecular dynamics and a phenomenological surface dynamics was employed to help understand the experimental results. A system was set up at Epion Corporation to do treatments on Nb single cell cavities and then RF-tested at JLab. The experimental results will be summarized and the perspective of this technique for real applications is discussed.

Finally, I will show two typical examples of surface studies of Nb using a high resolution transmission electron microscope and a home-made scanning field emission microscope respectively.

Early Stages of Nb Growth on Cu for SRF Accelerator Applications, C. Clavero, The College of William and Mary, N.P. Guisinger, Argonne National Laboratory, R.A. Lukaszew, The College of William and Mary

Among the large range of possible applications for superconducting Nb thin films, coatings for superconducting radio-frequency (SRF) cavities in linear accelerators have greatly aroused the interest of researchers in the last years [1]. Superconducting thin films and multilayer coatings are expected to increase further the maximum field gradients that SRF cavities can withstand, pushing them above 100 MeV/m [2]. In this regard, Nb coated Cu cavities have been proposed as a prototypical system for this purpose since they combine the better thermal stability of Cu due to its much higher thermal conductivity and the superconducting properties of Nb thin films [3]. Nevertheless, it is well known that structural dislocations and localized surface resistive defects on the thin films have a dramatically negative influence on their superconducting properties and resonator quality. Indeed, the quality of the films is strongly conditioned by the growth mode and the single atomic layer coverage at the very early stages of growth, and thus specific attention must be devoted to this range. Here we present a complete study on the early stages of growth of Nb on Cu(111). Different growth and annealing temperatures ranging from room temperature (RT) to 600 °C were used in order to investigate the characteristic growth mode of Nb in the sub-monoatomic coverage range. Scanning tunneling microscopy (STM) and scanning tunneling spectroscopy (STS) were used to investigate morphology and chemical composition of the surfaces with atomic resolution. Growth of sub-monolayer coverages at RT leads to amorphous Nb islands with 1 and 2 AL heights. Annealing at 350 °C gives rise to crystallization of the islands pseudomorphically with the substrate, i.e. Nb(111). Further annealing at 600 °C promotes interdiffusion of Nb atoms into the Cu substrate and alloying of the islands. Growth of higher coverages above 1 AL at 350 °C reveals preferential Volmer-Weber growth mode.

References:

The currently proven superconducting radio frequency (SRF) technology used in linear accelerators is based on bulk niobium cavities. Since this has a high cost and these cavities are approaching the maximum field gradients that they can withstand [1], development of a suitable, reliable, cost effective alternative to bulk niobium SRF cavities is needed. Attempts have been made to replace bulk niobium cavities with niobium-coated copper cavities since the thermal conductivity of a suitable base material such as copper is better than bulk niobium [2]. Coating niobium on SRF cavities is a promising but also challenging path, since there are several difficulties associated with various thin film deposition techniques and a lack of systematic studies pertinent to niobium thin film nucleation and growth leading to surfaces of greatest benefit.

Our systematic studies show that the transport properties, in particular the residual resistance ratio (RRR), are improved when niobium is epitaxially grown on crystalline ceramic substrates such as MgO and Al2O3, compared to niobium grown on (001) copper templates. Since grain boundaries are typically one of the main obstacles to superconducting transport, we show how the increased number of crystallographic domains that can occur during epitaxial niobium growth onto copper surfaces leading to higher density of grain boundaries can explain our results. We will discuss a route to improved transport properties while maintaining thermal efficiency by using alternative seed-layers grown on copper templates that can limit increased grain boundary density. We will show our correlated studies of microstructure and surfaces of Nb films on MgO and Al2O3 and the resulting transport/magnetic properties (four point probe and SQUID magnetometry) illustrating possible mechanisms to improve SRF cavity performance of such niobium films.

This work is funded by HDTRA1-10-1-0072 from the Defense Threat Reduction Agency as well as a subcontract from Thomas Jefferson National Accelerator Facility under contract DE-AC05-06OR23177 from the Department of Energy as supplemented by ARRA funds.

References:

5:00pm VT+MN+NS+SS+AS-TuA10 Development via Energetic Condensation of Niobium Thin Films Tailored for Superconducting RFApplications, A.-M. Valente-Feliciano, Jefferson Lab

For the past three decades, bulk niobium has been the material of choice for SRF cavities applications. In the recent years, RF cavities performances have approached the theoretical limit for bulk niobium. For further improvement of RF cavity performance for future accelerator projects, an interesting alternative has been recently proposed by Alex Gurevich with the Superconductor-Insulator-Superconductor multilayer approach, using the benefit of the higher critical field $H_c$ of higher-$T_c$ superconductors without being limited with their lower $H_c$.

JLab is pursuing this approach with the development of multilayer structures based on NbTiN via magnetron sputtering and High Power Impulse Magnetron Sputtering (HIPIMS). Insulators such as, AlN, Al2O3 and MgO are being investigated as candidates for the insulator layers. This paper present the the characteristics of NbTiN and insulator layers produced and results on NbTiN-based multilayer structures on bulk Nb and thick Nb films.


The performance of particle accelerators may be significantly limited due to buildup of the electron cloud at the opposing ends of the vacuum chambers. The electron buildup intensity is strongly affected by secondary electron emission from interior surfaces of the chambers. Application of coatings with reduced secondary electron yield (SEY) onto vacuum chamber interior surfaces is one of the
most economical EC suppression techniques. As a part of the International Linear Collider (ILC) R&D program, the Cornell Electron Storage Ring (CESR) has been successfully reconfigured as a Test Accelerator (CesrTA) to study EC buildup and suppression techniques. During the CesrTA program, various passive SEY-reduction coatings (TiN, amorphous-carbon and diamond-like carbon thin films) have been applied to diagnostic vacuum chambers in CESR in order to evaluate the efficacy of the EC suppression and the vacuum performance of these coatings in an accelerator environment. These chambers are equipped with both vacuum instrumentation (ion gauges and residual gas analyzers), as well as EC diagnostics (retarding field analyzers and RF-shielded pickups). In this paper, we present the results of studies of the vacuum conditioning and EC mitigation performance of these coatings.

5:40pm VT+MN+NS+SS+AS-TuA12 Electron Cloud Mitigation for the Large Hadron Collider (LHC), V. Baglin, G. Bregliozzi, P. Chiggiato, P. Costa Pinto, J.M. Jimenez, G. Lanza, M. Taborelli, C. Yin Vallgren, CERN, Switzerland

One of the main issues for the vacuum system of the Large Hadron Collider (LHC) is the build-up of electron clouds generated by electron multipacting in presence of beams. The occurrence of spatially distributed negative charges can lead to beam instabilities and emittance blow-up, pressure rises with a consequent background growth in the experimental areas, and increased thermal load in the cryogenic sections. The development of electron clouds depends on beam intensity and structure, magnetic field, and, in particular, the secondary electron emission of the beam pipe walls.

With respect to this latter point, electron clouds can be eradicated whenever the maximum secondary electron yield becomes lower than a critical threshold. In the LHC the problem has already been tackled at the design phase by introducing TiZrV non-evaporable getter thin film coatings as the baseline for most of the room temperature sectors of the ring. After activation by in situ heating, this material provides maximum secondary electron yield lower than 1.1. In addition, during operation, dedicated scrubbing runs are carried out by generating intentionally electron clouds and electron impingement onto the non-coated vacuum chambers, in a way to reduce their secondary electron yield. Recently magnetron sputtered carbon coatings have been also studied because they can reach exceptionally low secondary electron emission without any heating; their application in the LHC injectors and future LHC components is under investigation.

The effect of electron clouds in the pressure variations during the first months of LHC operation will be presented, together with the effects ascribed to the mitigation techniques.
There are only a few studies reporting density-functional-theory (DFT) being able to calculate binding energy shifts based on molecular structure. Spectra present a big problem for accurate identification of unknown peaks in calibrating spectra and also sensitivity of BE to secondary environments. However, in many cases the reference materials available in the databases. Induced on an atom by its surrounding ligands and by comparing to them to spectroscopy (XPS) measures the binding energy (BE) changes that are caused particular materials behavior is crucial for the rational development of functional motifs that knowledge of geometries and chemistry. Halevi, K. Artyushkova treatment of core electrons. Their high kinetic energy can require to include the specimen surface. Its electron optical design is an Einzel lens of asymmetrical electrode structure and operates with a positive potential of up to 50kV applied to its central electrode. The lens operation is extremely stable with no detectable electrical instabilities. To aid the development of the electrostatic lens we use simulation software to calculate the lens parameters. At the objective lens WD of 10mm and object at infinity, low values of spherical aberration (75mm) and chromatic aberration (19mm) were computed. These values are comparable to the best magnetic lenses of similar external dimensions and working distance. The column is mounted on a small analytical chamber for test purposes. We have demonstrated a 3nm SEM spatial resolution at beam energies of 10keV with 250pA of current. At a beam energy of 3keV with 150pA of current, we achieved and 7nm SEM spatial resolution. An Auger signal was detected using a new form of hyperbolic field parallel energy analyser. A Gold Au (64eV) line scan demonstrated an edge resolution of 4.1nm using a primary probe of 10keV energy and 400pA of current. For conventional energy analyzers, a probe current of 1nA is more appropriate and the spatial resolutions have been measured as 5nm and 7nm for respective beam energies of 10keV and 7keV.

AS-TuP3 Core Level Shift from Experiment and First-Principles-Theory – A Comparison, B. Kiefer, New Mexico State University, B. Halevi, K. Artyushkova, University of New Mexico The knowledge of geometries and chemistry of functional motifs that causes particular materials behavior is crucial for the rational development of novel materials and materials based technologies. X-ray photoelectron spectroscopy (XPS) measures the binding energy (BE) changes that are induced on an atom by its surrounding ligands and by comparing to them to databases. However, in many cases the reference materials available in the databases are limited to limited to common/model compounds, while actual materials designed may have more complicated chemistries. Furthermore, the big spread in the reported values of BE due to various ways to charge calibrate spectra and also sensitivity of BE to secondary environments present a big problem for accurate identification of unknown peaks in spectra. Being able to calculate binding energy shifts based on molecular structure can be very important tool for identifying structural motifs in materials. There are only a few studies reporting density-functional-theory (DFT) calculations of BE energies due to challenges that are associated with the treatment of core electrons. Their high kinetic energy can require to include relativistic effects especially for heavy elements and absolute values of binding energies cannot be expected to be accurate. Furthermore the differences in reference state complicate the direct comparison between experiment and theory.

A possible solution to these challenges is to compare BE to a reference system that is accessible both in experiment and theory. We will report on BE shift that have been calculated and compared to experimentally obtained values for metallic, bimetallic and covalent systems. For metallic Pd (111) systems, diverse layers have been found to surface binding energy, while sub-surface layers exhibit a metallic-like. For bimetallic PdZn system, Pd(3d) and Zn(2p) shift upward with respect to fcc-Pd(3d) and for hcp-Zn(2p), respectively. Magnitudes of shifts are very similar to that obtained experimentally. For covalent system, such as metal-less and metal-containing porphyrins, shift for N 1s BE in Co 2p environment compared to N 1s without metal is also very close to that obtained experimentally.

AS-TuP4 Evaluation of CuO Depth Profiling Conditions for XPS Organic Films Analysis, S. Alnabalsi, S.R. Bryan, J. Moulder, Physical Electronics Type-II organic materials can be successfully depth profiled by XPS using C60 ion beam sputtering at room temperature. Success is defined by achieving steady state composition as a function of depth. However, depth profiling of Type-I polymers have been marginally successful due to rapid sample damage under C60 bombardment. The use of high C60 beam energy at a glancing angle has been shown to extend the maximum depth of successful depth profiling, yet it eventually fails due to sample roughening, concomitant carbon build up, and a strong reduction in sputter rate. Zalar (azimuthal) rotation has been shown to minimize sputter induced roughness formation, improve depth resolution in multi-layer film structures, and allow depth profiles to be extended to even greater depths. Cooling of the sample to below its glass transition temperature is expected to reduce the mobility and reactivity of free radicals that are formed during the sputtering process. The purpose of this study is to evaluate the relative importance of using Zalar rotation and sample cooling to characterize standard organic thin film structures and to quantify the benefit to using them simultaneously.

AS-TuP5 Development and Application of Novel Electron Energy Analyzers for Chemical Analysis of Surfaces, D. Cubric, Shimadzu Research Laboratory (Europe) Ltd., UK, N. Kholine, Institute for Analytical Instrumentation, RAS, Russian Federation Various instruments have been developed previously for analyzing the spectrum of energies of beams of charged particles emanating form surfaces and described in numerous publications [1]. Among all those instruments the concentric hemispherical analyser (CHA) and the cylindrical mirror analysers (CMA) have been the most often used instruments both commercially and for various devices developed by different laboratories. At first sight, geometry of the two analyser types is very different. However, topologically their geometries are very similar. This paper presents a line of thought where geometries of both analysers are just particular examples of a more general geometry that in turn provides numerous analyser configurations, that we now call spheroid energy analysers (SEA), often having excellent electron optical characteristics [2]. The SEA instrumental concept enables electron optical exploration of various new analyser embodiments, particularly well suited for chemical analysis of surfaces and surface nano-features. Progress in this instrumentation development and example spectra will be presented.


AS-TuP6 Extreme Brightness: Reaching the Ultimate Limits of the Electron Beam, J.D. Jarvis, J.L. Kohler, B. Ivanov, N. De Jonge, B.K. Choi, A.B. Hmelo, C.A. Brau, Shimadzu Research Laboratory (Europe) Ltd. We report recent results from field emission microscopy studies of multivall carbon nanotubes (MCWNT) and from energy spectrum measurements of beams from diamond field emitters (DFE). As expected, resonant tunneling through adsorbed species on the emitter surface is an important and sometimes dominant effect. For diamond emitters our observations include order-of-magnitude emission enhancement without spectral broadening, complex spectral structure, and sensitivity of that structure to the applied electric field. For carbon nanotubes we have observed electron beams from individual adsorbates which are estimated to approach the maximum beam brightness allowed by Pauli exclusion. The development of CNTs as field emitters for the purposes of microscopy, lithography, radiation generation, and display production has been a protracted endeavor. Working in their favor, CNTs have excellent emission stability, high activation energy for thermal migration, excellent current throughput capacity, and a high degree of chemical inertness. Not
surprisingly, these properties extend to other covalent-carbon structures such as CVD diamond field emitters. DFEs have shown great promise as high-pressure electron beam sources in recent years. One of the more exciting revelations regarding CNTs and DFEs has been the realization that resonant tunneling through adsorbed species can be exploited for the generation of multi-microamp-level beams with perfect transverse coherence.

For resonant tunneling through a single molecule the electron source size is on the order of the electron wave length. Estimates of the transverse momentum spread of the resulting electron beams suggest that the transverse emittance is Heisenberg limited. Such beams have been produced using single atom tungsten, or single atom noble-metal field emitters, however the relatively weak binding of the metal atoms has limited the total current that can be extracted to the sub-microamp regime. In this paper we will discuss our efforts to generate, stabilize, and characterize, multi-microamp beams from individual molecules on CNTs.


Disposable elastic gloves are ubiquitous in scientific laboratories and are also widely used in many industries during handling of critical surfaces. Disposable gloves are typically made from nitrile, latex, neoprene, or other polymers and are used to protect users protected from various aqueous solvents, biological fluids, organic solvents, and other potentially harmful chemicals. A second major application of disposable gloves is to protect manufactured products and analytical samples from contamination caused by the transfer of skin cells, oils, salts, or other residues resulting from contact with bare hands. However, disposable gloves can also be a potential source of contamination. In addition to the primary polymer structure, many types of common laboratory gloves also contain a variety of inorganic materials in the glove formulation. Mold-release agents that allow the gloves to be easily stripped from the glove formers during fabrication may be present on glove surfaces. Furthermore, many types of disposable gloves have polymeric surface coatings that provide improved donning properties. Contamination resulting from surface residues on gloves can adversely affect materials used in industries where surface cleanliness is essential for optimum product performance and can also interfere with the analysis of samples depending on the specificity and sensitivity of the analytical technique. X-ray photoelectron spectroscopy (XPS) is a quantitative and quantitative surface sensitive technique that can be used to evaluate the surface composition of disposable gloves and to determine if contamination transfer occurs from gloves in a specific process. In this study, XPS was used to characterize the surface compositions of a variety of common laboratory gloves. The transfer of surface components from gloves to other material surfaces and changes in the surface composition of gloves following exposure to several common laboratory solvents were also investigated.

AS-TuP10 Large Area Cross Sectional Microstructural Characterization of FIB-TOF-SIMS Depth Profile Crater Walls, F.S. Smentkowski, D. Ellis, GE-GRC

Focused Ion Beam (FIB) techniques are widely used in order to generate cross sections of samples which can then be analyzed using a variety of analytical instrumentation in order to obtain microstructural information. In order to rapidly generate FIB cross sections, small areas (typically 5 to 10 microns) are milled. Time of Flight Secondary Ion Mass Spectrometry (ToF-SIMS) is a powerful surface analytical technique that is able to measure low concentrations of sub surface species. ToF-SIMS depth profile measurements are collected by eroding a large area (50 to 1,000 microns in size) of a sample using an ion beam for a certain amount of time, pausing the erosion, analyzing a smaller (central) region of the eroded area, and repeating the cycle until the desired depth has been reached. Ion images are often recorded at each depth and a full mass spectrum is saved at every vol, element allowing for 3-D analysis. Often times, both ToF-SIMS depth profile analysis and high resolution microstructural SEM characterization are required on the same sample. Ideally, these complimentary analyses would be performed in the same region of a sample. In this presentation, we will demonstrate that FIB techniques can be used to clean up the large area craters produced by ToF-SIMS depth profile analysis thereby enabling microstructural characterization over larger areas of samples.

Advantages of this approach are: larger FIB cross sections can be prepared since a significant amount of material was removed during the erosion cycles in the ToF-SIMS analysis, combining the chemical information provided from the ToF-SIMS analysis with the large area microstructural characterization provided by FIB-SEM analysis, and having both the ToF-SIMS and the large area cross sectional analysis performed in the same region of the sample. The ToF-SIMS depth profile craters have a slope of about 45 deg (the ion column is positioned at an angle of 45 deg from the sample normal), and hence an advantage of performing the FIB clean up at this angle is the entrainment in the layer thickness that will be available for microstructural analysis.

AS-TuP11 Microphase Separation of Various Diblock Copolymers Investigated by TOF-SIMS Depth Profiling, Y. Lee, Korea Institute of Science and Technology, Republic of Korea, J. Lee, Korea University, W.C. Lim, Korea Institute of Science and Technology, K. Shin, Sogang University, Korea, K.-J. Kim, Korea University

Diblock copolymers, composed of two distinct homopolymers covalently bonded together at one end, exhibit a microphase separation from a disordered state to an ordered state on cooling or heating. The microphase separation of diblock copolymers has been investigated by many different research groups for many years, because of the increasing use of diblock copolymers as compatibilizers, dispersants, impact modifiers, nanocarriers, and templates. Investigation of the phase behaviour of a family of diblock copolymers between styrene and a homologous series of methacrylates or acrylates has revealed that significantly different phase behaviours are seen for these materials, depending on the temperature, molecular weight, and alkyl side chain length. In this work, we characterized the morphology from various diblock copolymers of poly(styrene-b-alkyl acrylate)(PS-PAA), where P5 block was perdeuterated, near the copolymer/air and copolymer/substrate interfaces and in the bulk using time-of-flight Secondary Ion Mass Spectrometry (TOF-SIMS). TOF-SIMS Depth profiling was obtained for the lamellar morphology of PS-PAA which is found to orient parallel to the surface of the substrate. This preferential orientation resulted in a periodic variation in the composition of each block that continued through the entire copolymer film. Temperature- and chain length-dependent annealing studies on PS-PAA thin films on the silicon substrates were performed to investigate the order-to-disorder transition (ODT) properties of diblock copolymers.


Stainless steel coupons approximately 0.5” in diameter and 0.125” thick were passivated with five different surface treatments and an untreated coupon was left as a control. These surface treatments are being explored for use in tritium storage containers. These coupons were made to allow surface analysis of the surface treatments using well-known surface analysis techniques. Depth profiles using Auger electron spectroscopy (AES) and X-ray photoelectron spectroscopy (XPS) were performed on these coupons to characterize the surface and near surface regions. Scanning electron microscope (SEM) images were collected as well. All of the surface treatments studied here appear to change the surface morphology dramatically, as evidenced by lack of tool marks on the treated samples. In terms of the passivation treatment, Vendors A-D appeared to have oxide layers that were very similar in thickness to each other (0.7-0.9 nm thick) as well as the untreated sample (the untreated sample oxide layers appeared to be somewhat larger). Vendor E’s silicon coating appears to be on the order of 200 nm thick.


Discharged firearm cartridges produce unique microscopic particles referred to as gunshot residue (GSR). GSR is primarily composed of the products of combustion of the cartridge primer materials and typically contains varying amounts of Pb, Sb, and Ba, plus other elements. GSR particles can range in size from tens of nanometers to hundreds of micrometers. Because of the high temperatures (~1,500-3,600 °C) and pressures (~14,000-65,000 psi) that result within 1 millisecond of discharging a firearm cartridge, highly complicated chemical interactions are likely to occur that will affect the chemical composition of the GSR particles. Computer-controlled scanning electron microscopy (CCSEM) is the method preferred by the forensic community for the automated analysis of GSR. With CCSEM, large populations of potential GSR particles in the size range of 1–100 microns) are rapidly scanned for the characteristic chemical compositions of Pb, Sb, and Ba using energy dispersive X-ray spectroscopy (EDS). Determining the presence of these three elements fused together in a single particle having the correct morphology is all that is normally required for the positive identification of GSR. CCSEM, however, does generally not provide information regarding the population of particles much less than 1 µm. In addition, little is presently known regarding the surface chemistry of GSR. Because of its nanometer-scale sampling depth and the ability to provide
detailed chemical state information, X-ray photoelectron spectroscopy (XPS) can provide important information regarding the surface chemistry of QSSDs. In this work, we describe the fine particle synthesis and development of high resolution electron microscopy methods and complements the microscopy data with surface chemistry information obtained from XPS.

AS-TuP14 Analysis of Graphene and Other Graphitic Materials using XPS and AES, H.M. Meyer III, Oak Ridge National Laboratory

Synthesis and characterization of advanced carbon materials, including graphene, has accelerated over the past several years at the Oak Ridge National Laboratory. Projects include, among others, the development of nanostructured carbon materials for batteries, fuel cells and as supports for revolutionary catalysts. The past year has also brought to ORNL new projects related to the growth, characterization and application of graphene. This poster will highlight x-ray photoelectron spectroscopy and scanning Auger microanalysis of a variety of important carbon materials from several of these research projects, including nano-sized carbon powders for battery applications, graphite carbon powders used in fuel cells, graphene flakes synthesized for supporting catalyst materials and CVD grown graphene. In particular, an approach for curve fitting highly graphitic materials will demonstrate internally consistent results, giving insight into the sp2 vs. sp3 type carbon atoms present in the material.

Research sponsored by the Assistant Secretary for Energy Efficiency and Renewable Energy, Office of FreedomCAR and Vehicle Technologies, as part of the High Temperature Materials Laboratory User Program, Oak Ridge National Laboratory, managed by UT-Battelle, LLC, for the U.S. Department of Energy under contract number DE-AC05-00OR22725.


Ceria-zirconia mixed oxide (CZO) has been incorporated into automotive catalysts as a support material for precious metals (Pt, Rh, Pd) due to its highly desirable redox properties. However, at high operating temperatures, certain interactions between CZO and precious metal particles may adversely affect the performance of the catalytic converter. Thus, it is important to investigate the CZO/precious-metal interactions under redox conditions. Here, the preparation of Pd/CZO thin films as model planar catalyst systems will be described together with a systematic study of an important adverse interaction, the encapsulation of Pd particles by CZO, using a combination of x-ray diffraction (XRD) and high resolution transmission electron microscopy (HRTEM).

Ceria-zirconia (Ce0.2Zr0.8O2-δ) thin films with thickness ranging from 10-200 nm were grown onto (111) yttria-stabilized zirconia (YSZ) by oxygen plasma assisted molecular beam epitaxy (OPA-MBE). The epitaxial layer growth of CZO on YSZ(111) was confirmed by in-situ reflection high energy electron diffraction (RHEED). Atomic force microscopy (AFM) images revealed the smooth surface of the films with low roughness values (3-8 Å). Rutherford backscattering spectrometry (RBS) data along channeling and random geometries showed the minimum yield ( Yam) of 13% for Ce with no interdiffusion of metal atoms at the film/substrate interface. The single crystal nature of the film with CZO(111) orientation was confirmed by XRD data. Following the growth and characterization of thin films, ~1 monolayer of Pd was deposited on CZO(111) by thermal evaporation in a UHV chamber. After air calcination at 600 °C, the Pd/CZO films were reduced in 1% H2/N2 at 200°C, followed by annealing in N2 at 700 °C. Due to the additional reduced induced compressive stress, the films broke and formed a rough surface with 10-20 nm CZO mesas, as observed by HRTEM. However, films with a thickness above ~50 nm were stable, and Pd particles that formed on the surface became partially encapsulated by migration of CZO. In the films with a thickness above ~50 nm, the native compressive stress induced by the lattice mismatch between the film and substrate was relaxed by misfit dislocations at the interface, as revealed by high resolution scanning TEM images, to varying degree, depending on film thickness, but the extent of encapsulation appeared to be about the same for all film thicknesses. These results suggest that the driving force for the partial encapsulation resides in the nature of CZO, and is not simply a consequence of the lattice-mismatch-induced compressive stress in the films.

AS-TuP17 Cleaning and Characterization of InP Surface using Atomic Hydrogen and In/AlGaAs Heterostructure using Cl2/O2 Neutral Beam, J.S. Oh, K.S. Min, C.K. Kim, G.Y. Yeong, Sungkyunkwan University, Korea

Highly selective and low damage etching of the GaAs/AlGaAs heterostructure using Cl2/O2 Neutral Beam. Following atomic hydrogen cleaning, 340 °C and annealing at 400 °C, the surface was exposed to ~1 x 10-2 Torr of TMA at room temperature and annealed to 270°C. The TMA induces a surface reconstruction consistent with a bulk like bonding configuration between the Al atoms and the surface P atoms. The TMA passivation layer has horizontal rows of DMA. The cleaning and nucleation process does not disrupt the InP surface thereby providing a path for further scaling of the InP layer and the gate oxide.

AS-TuP18 Highly Selective and Low Damage Etching of GaAs/AlGaAs Heterostructure using Cl2/O2 Neutral Beam, J.S. Hammond, J.S. Oh, K.S. Min, C.K. Kim, G.Y. Yeong, Sungkyunkwan University, Korea

Growth of high-quality GaAs/AlGaAs multiquantum well (MQW) structures on GaAs substrates is essential in fabricating devices such as heterojunction superlattices, field effect transistors, injection lasers, and solar cells, etc. The GaAs on AlGaAs was etched using a low energy Cl2/O2 neutral beam and the schottky device channel which was fabricated on the exposed AlGaAs was compared with those fabricated after the etching using wet etching and a Cl2/O2 ion beam. The wet etching was conducted by a mixture of H2O and H3PO4. For the Cl2/O2 neutral beam etching, a neutral beam system composed of three-grid inductively coupled plasma (ICP)-type ion gun and a reflector installed just in front of the ion gun was used. Using a low energy Cl2/O2 ion beam or a Cl2/O2 neutral beam, highly selective etching of the AlGaAs layer to AlGaAs similar to wet etching could be achieved through the formation of Al2O3 on the exposed AlGaAs during the etching. When the electrical characteristics of the schottky devices were compared, the devices fabricated after the etching using the neutral beam showed the best electrical characteristics such as electrical stability, low leakage current, higher barrier height, etc. by showing low damage to the exposed AlGaAs surface. 


Anodic TiO2 nanotubes offer unique properties for a wide range of applications including energy conversion, photocatalysis and biomedical devices. It is widely accepted that the initial growth of the nanotubes is based on the formation of a compact anodic oxide followed by the formation of etching grooves and pores in the oxide. The mechanism of steady state growth of the nanotubes from the embryonic pores has, however, remained a topic of debate. To evaluate a flow model for the formation of the tubular structures, high spatial resolution Scanning Auger Spectroscopy data is used to elucidate the compositional variations across TiO2 nanotubes. The layers were fractured parallel to the axes of the nanotubes.
III-V compounds are candidates to replace Si as the semiconductor in

digital and optoelectronic devices. The high electron mobility and flexible bandgap of these materials make them attractive for high-speed electronics and optoelectronics. The poor ultimate resolution achieved with some of the characterization techniques traditionally employed for the assessment of the materials leads to limitations in the quality control of the production. The Sapphire/AlN/AlxGa1-xN optical constants. The PMSE delivers both unique and proven reliability for on-line quality control of production.

As the demand for high speed electronics remains ever increasing, the dimensions of MOSFET technology are continuously scaled down to the smallest possible levels. Traditional gate dielectrics such as SiO2 are unable to effectively stop electron tunneling from degrading device performance at thicknesses below 1-1.2 nm. We propose using high dielectric materials instead, which can effectively limit leakage current and have a band gap close to SiO2. This study will focus on the dielectrics Al2O3, HfO2, and Al2HfOx deposited using ALD. Several ratios of aluminum and hafnium in the Al2HfOx compound will be investigated to find a level that maximizes both electrical and mechanical properties of the film. The electrical properties of each dielectric will be characterized by XPS, AFM, spectroscopic ellipsometry, and by taking IV and CV measurements of fabricated transistors and capacitors. We will also investigate the effects of different annealing and deposition temperatures on the interface by TEM.

As-TuP23 Enhanced Green Emission from UV Down-Converting Ce3+-Tb3+ Co-Activated ZnAl2O4 Phosphor, K.G. Tshabalala, University of the Free State, South Africa

As-TuP24 Characterization of Al2Ga4-N Thin Film Light Emitting Diode (LED) Device by Spectroscopic Ellipsimetry, K. Uppiredi, L. Yao, HORIBA Scientific

The demand for efficient energy usage for lighting at industrial viable production costs catalyzing the rapid growth of global LED market. LED lighting is widely in use from LED-backlit TVs to solid-state lighting. The wall-plug efficiency, which characterizes its performance depends on the design and materials properties of thin film LED structure. The accurate determination of thickness and composition of the well and barrier layers is desired for quality control in the production. The Sapphire/Al2O3/AlxGa1-xN structure was characterized using a phase modulated spectroscopic ellipsometer (PMSE) at an angle of incidence of 70 degrees across the spectral range 0.6-6.5 eV. Using a three layer structure we accurately modeled the thickness of the layers in the spectral range from 0.6 to 4.5 eV and Al2Ga4-N optical constants. The PMSE delivers both unique performance and proven reliability for on-line quality control of production process.
Biofabrication and Novel Devices Focus Topic
Room: East Exhibit Hall - Session BN-TuP

Biofabrication and Novel Devices Poster Session

BN-TuP1 Photoluminescence Characterization of Polyphtiohyne Films Incorporated with Highly-Functional Molecules Such as Metallophthalocyanines, H. Koba, K. Onaka, H. Kato, S. Takemura, T. Hiramatsu, K. Shimada, K. Matsui, Kanto Gakuin University, Japan

Conducting polymer polythiophene (PT) films incorporated with highly-functional molecules such as phthalocyanines with different center metals were synthesized and characterized by X-ray photoelectron spectroscopy (XPS) measurements, photoluminescence measurements (PL) and time correlated single photon counting (TCSPC) measurements in order to obtain fundamental photoluminescence properties of various PT-phthalocyanine complexes prepared by different solvents. The electrochemical polymerization was performed in acetonitrile containing thiophene monomer and (ET)3NBF4 as a supporting electrolyte and the polymerization on an indium tin oxide (ITO) was conducted by applying positive voltage to the anode. The dopant molecules were iron phthalocyanine (FePc), copper phthalocyanine (CuPc) magnesium phthalocyanine (MgPc), lithium phthalocyanine (LiPc) and cobalt phthalocyanine (CoPc). Those molecules were doped in the polymer film by the diffusion method. The solvents used in the doping process were acetonitrile and toluene. At first, it was confirmed by XPS measurements that the metallophthalocyanines were introduced in the PT films. In the photoluminescence measurement, emission peaks were different in intensity and wavelength according to the dopant molecules. Those peaks were influenced by Soret and Q bands. The solvent used in the doping process also influenced the emission characteristics. In the case of CuPc using acetonitrile as a solvent, emission peaks which originated from Soret and Q bands were observed in the photoluminescence emission spectrum. On the other hand, emission peaks only due to Q band were observed in the cases of FePc, MgPc, LiPc and CoPc. Using toluene as a solvent drastically changed the emission characteristics. In the case of FePc, LiPc and CoPc, the emission peaks only due to Soret band were observed. As for CuPc, an emission peak due to Soret band was dominated. In the TCSPC measurements, it was confirmed that the number of life time components fitted to the decay curve ranged from 3 to 5 with several nanoseconds to several hundreds of nanoseconds. The life time and the number of components depended on the center metals of the doped phthalocyanine and the solvent used in the doping process. The present work clarified that the photoluminescence emission peak position, intensity and life time were varied by solvents and center metals of phthalocyanines.

BN-TuP3 Towards F3-ATPsynthase Based Hybrid Nanodevice Fabrication, J.K. Settle, M.L. Richter, C.L. Berrie, University of Kansas

Incorporation of biomolecules into nanoscale devices, termed nanoball surface. The resultant products showed relatively higher durability compared with that of catalysts prepared by conventionally-developed methods. In this work, we demonstrate surface decoration of chemically converted graphene with various metallic nanoparticles by the originally-developed solution plasma polymerization. Structural characterization and electrochemical activities of the all product was performed by a combination with TEM, XRD, AFM, Raman spectroscopy, and measurements of oxidation-reduction reactions [1]. The metallophthalocyanines were doped in the polymer film by the diffusion method. The solvents used in the doping process were acetonitrile containing thiophene monomer and (ET)3NBF4 as a supporting electrolyte and the polymerization on an indium tin oxide (ITO) was conducted by applying positive voltage to the anode. The dopant molecules were iron phthalocyanine (FePc), copper phthalocyanine (CuPc) magnesium phthalocyanine (MgPc), lithium phthalocyanine (LiPc) and cobalt phthalocyanine (CoPc). Those molecules were doped in the polymer film by the diffusion method. The solvents used in the doping process were acetonitrile and toluene. At first, it was confirmed by XPS measurements that the metallophthalocyanines were introduced in the PT films. In the photoluminescence measurement, emission peaks were different in intensity and wavelength according to the dopant molecules. Those peaks were influenced by Soret and Q bands. The solvent used in the doping process also influenced the emission characteristics. In the case of CuPc using acetonitrile as a solvent, emission peaks which originated from Soret and Q bands were observed in the photoluminescence emission spectrum. On the other hand, emission peaks only due to Q band were observed in the cases of FePc, MgPc, LiPc and CoPc. Using toluene as a solvent drastically changed the emission characteristics. In the case of FePc, LiPc and CoPc, the emission peaks only due to Soret band were observed. As for CuPc, an emission peak due to Soret band was dominated. In the TCSPC measurements, it was confirmed that the number of life time components fitted to the decay curve ranged from 3 to 5 with several nanoseconds to several hundreds of nanoseconds. The life time and the number of components depended on the center metals of the doped phthalocyanine and the solvent used in the doping process. The present work clarified that the photoluminescence emission peak position, intensity and life time were varied by solvents and center metals of phthalocyanines.

Graphene and Related Materials Focus Topic
Room: East Exhibit Hall - Session GR-TuP

Graphene and Related Materials Focus Topic Poster Session

GR-TuP2 Solution Plasma Assisted Surface Decoration of Chemically Converted Graphene Sheet with Various Metallic Nanoparticles, K. Sadase, N. Zettsu, T. Ueno, O. Takai, N. Saito, Nagoya University, Japan

Like the other newly discovered carbon nanomaterials, such as fullerenes and carbon nanotubes, graphene, a monolayer graphite, and its composites have been widely applied to the field of electrocatalysis due to that graphene have unique structures and inherent properties including high specific surface areas, chemical and electrochemical inertness, easy surface modification, higher electron conductivity and broad electrochemical windows. Thus, the use of graphene as substrates of catalysts provides a new opportunity for designing and constructing next-generation catalysts. Especially, graphene decorated with metallic nanoparticles can be employed as fuel cell electrocatalysts, in which they plays an important role in improving the charge-transfer efficiency and decreasing the overpotential of electrochemical reactions such as methanol oxidation and oxygen reduction.

However, there are still unresolved issues in practical use, such as durability. In situ growth method is the most widely used method for preparing graphene/metallic nanoparticle composites. Hydrophobic and/or electrostatic interactions are the main driving forces of adsorbing metallic nanoparticles on graphene surfaces. These relatively weak binding interaction often gave rise to desorption of the nanoparticles from the graphene surface during charging and discharging operation at high temperature.

Very recently, we reported a fabrication of Pt catalysts supported on carbon nanoballs, as well as characterization of their electrochemical activities (Saito et al., J. Vac. Sci. Technol. A, 27(4) pp.826-830). Plausible mechanism has remained a mystery, Pt catalysts were strongly attached to the carbon nanoball surface. The resultant products showed relatively higher durability compared with that of catalysts prepared by conventionally-developed methods. In this work, we demonstrate surface decoration of chemically converted graphene with various metallic nanoparticles by the originally-developed solution plasma polymerization. Structural characterization and electrochemical activities of the all product was performed by a combination with TEM, XRD, AFM, Raman spectroscopy, and measurements of oxidation-reduction reactions [1]. The metallophthalocyanines were doped in the polymer film by the diffusion method. The solvents used in the doping process were acetonitrile containing thiophene monomer and (ET)3NBF4 as a supporting electrolyte and the polymerization on an indium tin oxide (ITO) was conducted by applying positive voltage to the anode. The dopant molecules were iron phthalocyanine (FePc), copper phthalocyanine (CuPc) magnesium phthalocyanine (MgPc), lithium phthalocyanine (LiPc) and cobalt phthalocyanine (CoPc). Those molecules were doped in the polymer film by the diffusion method. The solvents used in the doping process were acetonitrile and toluene. At first, it was confirmed by XPS measurements that the metallophthalocyanines were introduced in the PT films. In the photoluminescence measurement, emission peaks were different in intensity and wavelength according to the dopant molecules. Those peaks were influenced by Soret and Q bands. The solvent used in the doping process also influenced the emission characteristics. In the case of CuPc using acetonitrile as a solvent, emission peaks which originated from Soret and Q bands were observed in the photoluminescence emission spectrum. On the other hand, emission peaks only due to Q band were observed in the cases of FePc, MgPc, LiPc and CoPc. Using toluene as a solvent drastically changed the emission characteristics. In the case of FePc, LiPc and CoPc, the emission peaks only due to Soret band were observed. As for CuPc, an emission peak due to Soret band was dominated. In the TCSPC measurements, it was confirmed that the number of life time components fitted to the decay curve ranged from 3 to 5 with several nanoseconds to several hundreds of nanoseconds. The life time and the number of components depended on the center metals of the doped phthalocyanine and the solvent used in the doping process. The present work clarified that the photoluminescence emission peak position, intensity and life time were varied by solvents and center metals of phthalocyanines.

GR-TuP3 Synthesis of Carbon Nanoballs, Covered by CNTs with Metallic Conductivity, K. Ohno, N. Zettsu, T. Ueno, O. Takai, N. Saito, Nagoya University, Japan

Carbon nanostructures, due to their unique electronic and extraordinary mechanical properties, have been receiving much attention for a wide variety of applications. Especially, highly-conductive carbon nanostructures is a growing candidate supports for heterogeneous catalysts, for use in fuel cells and in metal-air sources of current. Synergetic interaction of the catalyst-nanoparticles with carbon nanostructures exhibits better catalytic parameters than application of carbonaceous materials.

Very recently, we demonstrated a fabrication of platinum catalysts supported on carbon nanoballs (CNBs), as well as characterization of their electrochemical activities (Saito et al., J. Vac. Sci. Technol. A, 27(4) pp.826-830). Originally-developed CNBs is a highly crystallized nanoscale carbon powder with spherical shape. The CNBs can be synthesized by thermal decomposition of ethylene and hydrogens under argon stream. Diameter of CNB can be tuned from 200 to 700nm by changing the mixing ratio of H2: Platinum nanoparticles were densely loaded on the CNBs surface under modified solution plasma processing of CNBs and Pt ions with stabilizing agents. We obtained 1.6-fold increase in the electrochemical activity of Pt/CNBs system (0.26cm2/mg) compared with that of conventionally-used Pt/C system (0.16cm2/mg).

In this work, we propose newly developed CNBs, covered by carbon nanotubes (CNTs) with metallic conductivity in order to reduce contact resistance between neighboring CNBs in electrodes. We grew CNTs from various shaped Fe, Ni nanocrystals deposited onto an individual CNB, and subsequently decorated the surface of the CNBs with Pt nanoparticles as a catalyst by using solution plasma processing. The ratio of metallic to semiconducting CNTs of all our products was evaluated by Raman spectroscopy.
Thus we have been investigating the use of soluble precursor molecules which form the graphene templates at higher temperatures or in the presence of catalysis. In particular, one approach that is under investigation is the use of reversible Diels-Alder chemistry to produce soluble graphene precursor adducts. After solution coating of the materials onto appropriate substrates, the cyclodaddition can be reversed at modest temperatures to produce the desired polyaromatic precursor and subsequently processed into graphene through a further chemical or thermal consolidation. This presentation will review our progress to date in developing such techniques and will demonstrate the ability to form graphene from such approaches at relatively moderate temperatures. A discussion of the effect of choice of precursor on the ability to consolidate the material into graphene and its resulting quality will be presented.


Graphene is currently produced typically one of four routes: (1) some combination of mechanical and chemical exfoliation of graphite, (2) oxidation of graphite to form graphene oxide which is subsequently deposited onto surfaces and reduced to graphene, (3) high temperature processing of SiC, or (4) through high-temperature catalytic methods on metal surfaces using simple carbon precursors such as methane. Some applications for graphene would benefit from low temperature methods (<400 °C), and the first two methods mentioned above can inherently provide access to such low processing temperatures. However, simple exfoliation of graphite has only been demonstrated to produce relatively small, poorly controlled graphene flakes that are not easily processed due to their limited solubility in solvents and lack of methods to further assembly into larger structures. Therefore, we have been investigating the ability to utilize graphene oxide (GO) flakes as a soluble graphene precursor that can be assembled onto surfaces to form both continuous graphene films and directly form patterned graphene microstructures. We will present a process in which we deposit and pattern aminosilanes on a substrate, pattern them using lithographic techniques, and utilize such aminosilane patterns to control where GO assemblies on the surface and subsequently is reduced to form graphene. It will be demonstrated that by producing aminosilane monolayers that are dense and which possess a large fraction of non-hydrogen bonded amine terminal groups, continuous graphene oxide and graphene films and microstructures can be assembled on substrates. It will be shown by subjecting the resulting assembled graphene on such aminosilane layers to modest thermal treatments, that stable n-doped graphene can be produced. The materials produced via such methods will be discussed in terms of their spectroscopic (e.g. Raman) and electrical properties (e.g. I-V curves for FET devices, carrier concentrations, mobilities, etc.)

GR-TuP7 Graphene Layer-By-Layer Growth on Co3O4 (111) at 1000 K by Molecular Beam Epitaxy, M. Zhou, F. Pasquale, J. Kelleher, University of North Texas, A. Boosalis, M. Schubert, P.A. Dowben, University of Nebraska - Lincoln

We report layer-by-layer growth of macroscopically continuous and uniform graphene sheets on Co3O4(111) at 1000 K by carbon molecular beam epitaxy (MBE) from a graphite rod source. The direct growth of graphene on dielectric substrates is an essential step in the practical and scalable production of graphene-based devices. Co3O4(111) films 3 monolayers (ML) thick were formed from surface segregation of dissolved oxygen after deposition ~ 40 Å Co grown on Al2O3(0001) substrates at 750 K in UHV. Epitaxial Co3O4(111) films, as characterized by Auger spectroscopy and LEED, were formed by subsequent annealing to 1000 K in UHV, and exhibit a O-O nearest neighbor distance of 2.8 Å, in good agreement with literature. The evolution of the Auger electron C(1s) line shape from a graphite-like carbon MBE indicates sp2 hybridization, and layer-by-layer growth up to at least 3 ML average thickness. LEED spectra indicate that the sp2 (111) graphene overlayer is incommensurate with the Co3O4(111) substrate. The graphene-related diffraction spots remain sharp from a coverage of 0.4 ML up to 3 ML, indicating that the graphene sheets are azimuthally in registry with each other. Exposure of the 3 ML graphene/Co3O4(111)/Co(111) sample to ambient results in no observable change in Auger or LEED spectra, indicating macroscopically continuous graphene sheets. Subsequent acquisition of XPS spectra in a separate chamber yields a graphite-characteristic asymmetric C(1s) peak at 284.9 eV binding energy, indicating graphene → oxide charge transfer, as observed for graphene/SiC and graphene/MgO. A π → π* satellite feature is also observed. Spectromicroscopy measurements carried out in a separate system confirm the presence of a π → π* resonance, and similarities with the optical absorption of graphene/SiC are observed. Raman spectra acquired at different, macroscopically separated sample areas indicate a uniform 3ML graphene film thickness. These results also strongly suggest that other non-polar (111) transition metal oxide surfaces with similar O-O nearest neighbor distances may act as suitable substrates for graphene growth at moderate temperatures, opening the way to controlled direct growth of high quality graphene on a variety of dielectric substrates, with materials and processing technologies readily compatible with Si CMOS integration.

Helium Ion Microscopy Focus Topic Room: East Exhibit Hall - Session HI-TuP

Aspects of Helium Ion Microscopy Poster Session

HI-TuP1 From HIM to NIM: The Prospects of a Neon Ion Microscope, F.H.M. Rahman, L.A. Stern, J.A. Notte, Carl Zeiss NTS

From its time of its conception, the gas field ion source (GFIS) was operated with a variety of gas species - each considered for some particular virtue that depended on the particular application. However, practical issues such as vacuum, cost, and safety have prevented its commercial introduction. This presentation will review our progress over the past 50 years to develop a commercial ion source in the form of the ORION helium ion microscope. The neon ion beam normally incident upon aluminum at 30 keV, though sometimes at high temperature (100 °C), will sputter atoms from the surface at a rate that is comparable to the rate of collisions due to the incident beam. These sputtered atoms are emitted from the surface and enter the vacuum chamber, where they are detected by a variety of methods. The neon ion beam is then used to image the surface at high resolution and with high fidelity. Experimental results will be presented to characterize the basic properties of the focused ion beam from our prototype neon GFIS system. Imaging results will be presented to demonstrate our first cross-section milling and imaging characteristics.


We report the formation of solid-state nanopores using the highly focused ion beam and lithographic capabilities of a scanning Helium Ion Microscope (HIM). We will discuss several aspects of the fabrication process including the advantage of high sample throughput along with fine control over nanopore dimensions. We will compare characteristics of the resultant devices with those made by the established technique of transmission electron microscope milling and demonstrate the utility of our nanopores for biomolecular analysis.

HI-TuP3 Imaging and Identification of Self Assembled Monolayers using HIM, G. Hlawacek, A. George, J.E. ten Elshof, R. van Gastel, H. Zandvliet, B. Poelsema, University of Twente, The Netherlands

Helium Ion Microscopy (HIM) is a new and versatile tool for imaging and characterizing surfaces, buried interfaces, thin films and tackling many other problems in modern material science. HIM utilizes ionized Helium to scan the specimen surface. Secondary electrons created by the impinging ions allow to record morphology images with an unmatched lateral resolution of less than 0.35 nm. In addition, back-scattered ions carry the elemental information of the scattering partner – allowing for a elemental identification of the surface composition.

Here, we report on the visualization of thin self assembled monolayers (SAM) deposited on (001) silicon wafers, covered by a thin native oxide. In particular, SAMs formed by (3-Mercaptopropyl)trimethoxysilane (MPS) and Triethoxy-1H,1H,2H,2H-tridecafluoro-n-octylsilane (TDFOS) have been patterned into a rectangular stripe pattern using a two step gas-phase silanization process. The clever use of channeling into the underlying bulk (001) silicon, together with a work-function based evaluation of the
secondary electron data allows a clear assignment of different sample areas to the different chemical species. This is possible for both the electron and the ion generated image. The importance of channeling to distinctly and visibly tag the different SAMs will be demonstrated.

**HI-TuP4** Analysis of Metal Nanoparticles in Biological Tissues Specimens Using the Helium Ion Microscope, V.S. Smentkowski, L. Denuault, D. Wark, GE-GRC, L. Scipioni, D. Ferranti, Carl Zeiss SMT

The Helium Ion Microscope (HIM) is a newly introduced instrument that has a number of beneficial characteristics that are of importance for the analysis of biological/tissue samples, including: (1) the ability to perform high lateral resolution imaging, (2) high depth of field, (3) and the ability to analyze charging samples. In this poster, we summarize the first HIM analysis of spleen tissue samples that have been treated with a metal contrast agent. We show the advantages of HIM over techniques such as Scanning Electron Microscopy (SEM). The HIM analysis are complimented by surface analysis using Time of Flight Secondary Ion Mass Spectrometry (ToF-SIMS) in order to demonstrate that the contrast observed by HIM is indeed associated with the contrast agent.

**HI-TuP5** Fabrication of Carbon Nanomembranes by Helium Ion Beam Lithography, X. Zhang, H. Viekler, A. Beyer, A. Göltzhäuser, Bielefeld University, Germany

A helium-ion microscope can be used as beam writing tool on electron beam photoresists, such as hydrogen silsesquioxane (HSQ). It has been demonstrated to have a high resolution, a high sensitivity and a low proximity effect.

Here we report the fabrication of carbon nanomembranes from aromatic self-assembled monolayers (SAMs) with a helium ion beam as direct writing tool. Cross-linking of SAMs is achieved by exposure with helium ions which results in the formation of mechanically stable carbon nanomembranes. The required doses for cross-linking with helium ions are approximately one order of magnitude lower than with electrons. The cross-linked SAMs were transferred to either silicon substrates with an oxide layer for optical characterization or transmission electron microscopy (TEM) grids for preparing free-standing carbon nanomembranes. With helium ion based cross-linking we fabricated patterned nanomembranes as well. Furthermore, the proximity effect and the sample damage on the nano-scale pattern is investigated and discussed.

**HI-TuP6** Layer Thickness Homogeneity Determination via Rutherford Backscattering in Helium-Ion Microscopy, H. Viekler, K. Rott, A. Beyer, G. Reiss, A. Göltzhäuser, University of Bielefeld, Germany

The recently developed helium-ion microscopy allows remarkable surface resolution with the secondary-electron (SE) detector. Simultaneously, backscattered ions can be detected that allow imaging with a substantially higher elemental contrast. This Rutherford backscattered (RBS) ion contrast depends mainly on the elemental composition of the investigated sample surface. The escape depth of RBS ions is much larger than for secondary electrons. Thus whole layers with a wide range of thicknesses will contribute to a RBS ion image, whereas the SE image is far more surface sensitive, i.e. insensitive to buried parts under the sample surface.

In this contribution we examine RBS ion imaging as tool to characterize thickness variations of layered samples with well defined compositions. In a model example the homogeneity of a gold layer on a silicon substrate is investigated. The achievable spatial resolution for detecting buried inhomogeneities is analyzed. Furthermore we present examples with multiple layers.


Dedegradation of the photocathode materials in accelerator-based photoinjectors represents a challenge for sustained beam delivery in proposed fourth generation light sources. The quantum yield in most existing photocathodes degrades over time leading to machine downtime for quantum yield replenishing and in some instances to photocathode replacement. Several photocathode degradation processes have been proposed including ion back bombardment, photochemistry of surface adsorbed species and irradiation-induced surface and bulk defect formation. At present, no consensus exists within the user community as to the mechanisms of photocathode degradation. Better understanding of these mechanisms of existing photocathode materials could lead to improved emission properties and longer operating lifetime. Existing photocathode materials range from metallic (e.g. copper) to semiconducting (e.g. GaAs) with various structures, dopants, and surface preparations. Photocathode emission requirements include high electron yield and low thermal emittance at high repetition rate. The goal of this work is to thoroughly characterize the used photocathode samples obtained from Jefferson lab using helium ion microscope (HIM), Rutherford backscattering spectrometry (RBS) in channeling and random directions, secondary ion mass spectrometry (SIMS), atom probe tomography (APT) and atomic force microscopy (AFM) to understand the degradation mechanism. Four different GaAs samples (two control including one as prepared and the other as annealed but not used, and two used to delivered 1000 and 7000 Coulombs) were analyzed using these techniques. HIM images obtained at the damaged spot from the 7000 C sample clearly show that the surface at this spot is severely damaged. In addition, some cracks are clearly visible on the surface. HIM images collected at the tilt angle of 20° clearly show that these damage features are protruding above the surface of the photocathode samples at the center region of the spot. Stylus profilometer measurement on this spot reveals that the spot has peaks and valleys; the height of the main peak is around 7000 nm while the depth of the valleys ranges from 1000 to 3000 nm. It appears that the material in this area is melted. HIM images collected from all four samples clearly show that there is a systematic variation in the topography of the samples as a function of the number of use of the photocathodes. The larger the usage time the smaller the structures are. Detailed analysis of these samples using RBS, SIMS together with HIM will be discussed.

**In Situ Spectroscopy and Microscopy Focus Topic Session Room: East Exhibit Hall - Session IS-TuP**

**IS-TuP1** In Situ Infrared Spectroscopy of Oxidation Process of Amorphous Carbon Film, Depending on Substrate Temperatures, H. Shinohara, Y. Takaki, K. Haru, Y. Takami, Y. Matsuda, H. Fujiyama, Nagasaki University, Japan

There has been much interest in amorphous carbon films because they have a lot of useful properties: mechanical hardness, chemical inertness, and changeable electrical properties. The films can be deposited at low temperatures by using plasma process. The films have been used as coating materials for mechanical apparatus. The property of the film surface can be changed with the addition of the other atoms on the surface. The addition of oxygen atoms to the surface leads the surface hydrophilic. The hydrophilic property on the surface has advantages to the further surface treatment. Therefore, it is important to understand the oxidation process of amorphous films. One of the effective oxidation methods is oxygen plasma exposure. We fabricated the plasma oxidation process using in-situ infrared spectroscopy in multiple internal reflection geometry (MIR-IRAS). In this presentation, we focus on the dependence of oxidation process on the substrate temperatures. Infrared spectroscopic studies indicated that the oxygen plasma exposure induced the generation of OH components in the film. It means that carboxyl group would be formed by the exposure. With the increase of the substrate temperatures, the formation of OH components in the film was increased. Moreover, the hydrophilic property was decreased with the substrate temperatures. On the other hand, the etching rate due to the oxygen plasma exposure was increased with the substrate temperatures. It is suggested that the etching rate is increased with substrate temperature, compared with the preservation of the OH components in the film.

**IS-TuP3** In Situ TEM Studies of Nanoparticle Growth in a Fluorozirconate (ZBLAN) Glass Matrix, J. Johnson, University of Tennessee Space Institute

ZBLAN glass-ceramic materials are being developed as x-ray imaging plates. The materials are doped with europium and chlorine and can be heat treated in such a way that they form a novel nanocomposite material containing barium chloride nanocrystals, with the ability to convert x-rays into visible light. This property can be changed after illumination with a scanning laser beam in a photostimulated luminescence process. The ZBLAN glass only acts as an imaging plate upon annealing. As the annealing temperature and annealing time are increased, so a higher degree of nucleation of BaCl2 crystallites inside the glass matrix is observed. As a result, more crystallites are available to incorporate Eu2+ and hence increase the fluorescence intensity. However, a higher annealing temperature and a longer annealing time also lead to a larger degree of crystal growth, resulting in a loss of the x-ray sensitive property. This leads to in-situ electron microscopy of a ceramic-glass storage phosphor. The optimal annealing condition thus needs to compromise between the fluorescence intensity and the spatial resolution.
Here we present in situ TEM studies of ZBLAN glasses, being carried out to further understand the growth of nanoparticles inside a glass matrix under various heating conditions.

**Nanomanufacturing Science and Technology Focus Topic**

**Room: East Exhibit Hall - Session NM-TuP**

**Nanomanufacturing Science and Technology Poster Session**

**NM-TuP1** Nanoscopic Polymization of Polyamine on the Nanostructured Alumina Surface

**NM-TuP2** Dielectric Performance of Post Deposition Treated AlOx Films Prepared by Using Parallel-Plate Electrode PEALD

**NM-TuP3** Effect of Growth Temperature on Optical Properties of TiOx Films by Atomic Layer Deposition

**NM-TuP4** Fabrication of Double Nanohoneycombs (Pt/ZnO) with Controllable Size using Nanosphere Lithography and Plasma Enhanced Atomic Layer Deposition

**NM-TuP7** Fabrication of Nanopattern Sapphire Substrate by Nanosphere and Nanoimprint Lithography Technology

**Key:** PEALD, Direct plasma, High-k material, C-V, I-V, Dn, Parallel electrode
of sapphire substrate was nano-cone array structure with 400 nm diameter and 100 nm in thickness. The contact angles of two patterned sapphire substrates were measured to be 101.02° and 98.14° for nano-cone array structure, respectively. From the contact angle measurement results, it can be found that the surface property of sapphire substrate changed from hydrophilic, which contact angle was 24.46°, to be hydrophobic.

NM-TuP8 Fabrication of Single-Electron Transistor Utilizing Multi-Coated Self-Assembled Monolayer, N. Kwon, K. Kim, J. Chung, Sungkyunkwan Univ., Republic of Korea

We have fabricated quantum dots with the precise sizes from 30 nm to sub-10 nm at the controllable position. First, Au electrodes with the unique shape were obtained using a conventional lithography. Then, self-assembled multilayers, composed of alternating layers of α,ω-mercaptoalkanoic acids (~2 nm) and copper (II) ions, were deposited on Au electrode patterns to form the controllable gap between adjacent Au electrodes. After reaching the nanoanodization-scale gap, the second Au was deposited again. Finally, lift-off both e-beam resist and molecular resist were removed by lift-off, thereby resulting in quantum dot with nano-gap between gold electrodes. The physical properties were analyzed using scanning probe microscopy (SPM) and field emission scanning electron microscopy (FE-SEM). The electrical properties were evaluated using Keithley-4200.

NM-TuP9 Photoluminescence Studies of Nanostructured Alumina Surfaces Coated by Polythiophene Film and Copper Phthalocyanine A. Ishii, R. Nakashima, H. Kato, S. Takemura, H. Kobe, Y. Watanabe, T. Hiramatsu, Kanto Gakuin University, Japan

Photoluminescent properties of the nanostructured alumina surfaces and the surfaces coated by polythiophene (PT) nanofilm and Copper phthalocyanine (CuPc) were investigated. Nanostructures such as linked-crater structure and highly-oriented line structure were fabricated on an Al surface by a combined process of chemical and electrochemical treatments. The nanoanodized linked crater structure was fabricated on an Al surface by treatment with Semi Clean and successive electrochemical anodization in H$_2$SO$_4$ solution created a nanoanodized linked-crater structure on the surface. The crater size was estimated at 80-150 nm in diameter by dynamic force microscopy (DFM) measurements. Regarding the highly-oriented line structure, the anodization process applied to the original fiber-like surface was modified and new nanoanodized line structure on the Al plate. The anodization fabricated the fine line structure on the Al surface. The line distance was estimated at 40 nm. The fabricated nanostructured surfaces were identified as alumina by Fourier transform infrared spectroscopy (FT-IR) and x-ray photoemission spectroscopy (XPS) measurements. Conducting polymer polythiophene nanofilm growth on the nanostructured Al surface was conducted by an electrochemical synthesis method in an electrochemical cell. Polythiophene nanofilm was polymerized on the nanostructured Al used as an anode in acetonitrile containing thiophene monomer and (Et$_4$N)BF$_4$ by applying positive voltage to the anode. It was observed by DFM that nanofilm was grown along the crater structures or the line structures. CuPc deposition on the nanostructured surfaces was carried out by casting method. Photoluminescent properties of the nanostructured alumina surfaces and the surfaces topped by the nanofilm and CuPc were investigated. It was shown that characteristic ripples of several emission peaks appeared in the wavelength range of 350-550 nm in both cases of linked-crater and highly-oriented line structures while no ripple-shaped emission peaks were observed in the case of the original native Al oxide surface. The emission spectral profile was different in peak positions, number of ripples and intensities between two types of nanostructures. Photoluminescence measurements on the polythiophene nanofilm-grown linked-crater structure also showed that the rippled emission peaks clearly appeared. It was also clearly confirmed that the observed rippled emission patterns were significantly enhanced in both cases of CuPc deposited nanofilm-coated nanostructures. The mechanism of generation of the rippled peaks and the enhancement were discussed by considering nanoize effects and Al-O vibration modes.


The patterning of small contact holes is an ever present challenge in the field of photolithography. Recently, the importance of this challenge has expanded both because of the development of 3D NAND architectures such as BICS and because of the need to pattern even smaller contact holes for DRAM applications. As the required critical dimensions and pitches of contact holes become smaller, the methods typically used for lithography become insufficient. One approach to solving this problem is the use of criss-cross lithography. In criss-cross lithography, two sets of lines are patterned perpendicular to each other, and a freeze step is used to bind them in place. In the first portion of this work, criss-cross lithography is used to pattern 60 nm holes, with an argon freeze applied to harden the first layer of polymer before patterning the second layer. In this application, two types of problems were discovered: underexposure which causes patterns to become unstable due to gaps at the bottom of the structure and non-optimized dosing which results in patterns which, while circular before etch, exhibit X/Y directionality when any of a wide range of etch processes is applied to the pattern. We explain the causes of these two problems and demonstrate a consistent relationship between etch depth and X/Y directionality for these types of patterns.

In the later portion of this work we demonstrate the applicability of an alternate technique, thermal freeze, to the patterning of 60 nm contact holes. This technique has been applied and optimized such that it enables the criss-cross patterning approach to produce patterns that are consistent and circular even after the etch process. Along with demonstrating that thermal freeze can be used to produce criss-cross wafer, we also identify the exposure relationships required to achieve circular patterns.

**Nanometer-scale Science and Technology Division Room: East Exhibit Hall - Session NS-TuP**

**NS-TuP1 Electromigration Assisted Single Silver Nanowire Ammonia Sensing, W. Xing, S. Kyung, R. Penner, University of California, Irvine**

Due to small size, high sensitivity, and low power consumption, 1 dimensional nanowire sensors have attracted a lot of attention and may substitute for traditional thin film sensors in the future. To further improve the sensitivity, there is an obvious need to make sensors with a single nanowire and some seminal work has already been done using semiconductor nanowires in this direction. In this work, a single silver (Ag) nanowire was prepared on glass surface using Lithographically Patterned Nanowire Electrodeposition (LPNE) method. The produced Ag nanowire had a mean size of around 100nm and a height and range from 20-80nm and 100-400nm, respectively. Chemiresistor type sensors were fabricated from these single electromigrated Ag nanowires with 4 probe contacts, and their resistances upon exposure to different concentrations of NH$_3$ gas were investigated at room temperature. Furthermore, electromigration was used to create a narrow junction inside the nanowire, which enhanced the sensing performance dramatically. The electromigrated sensors showed a resistance change, $\Delta R/R_0$, that was large (90% for 7% NH$_3$), fast (<30s) and reversible with a detection limit of approximately 200ppm. In addition, the characterization of these nanowires by AFM, TEM, and SEM will be presented.

**NS-TuP2 Inhomogeneous Density Distribution of Silicon Dioxide Thin Films Thermally Grown on 1000 °C on Si (100) Surface, K. Odaka, A. Kurokawa, Y. Azuma, L. Zénà and T. Fujita, National Institute of Advanced Industrial Science and Technology (AIST), Japan**

We would report on the XRR (X-ray reflectivity) analysis to observe the density distribution of the SiO$_2$ thin film which is thermally grown on Si (100) at 1000 °C. The results show the non-uniform distribution in depth, and once the high density SiO$_2$ was formed during ramp-up it would remain under the following 1000 °C oxidation.

**Experimental** We measured XRR using an apparatus equipped with a shower of high purity N$_2$ to protect a specimen from contamination for a few hours of measurement. We analyzed the data with a fitting method [1], [2] which is well established to evaluate thicknesses, densities, and surface roughness of laminated layers of a specimen. We assumed a two-layer-fitting model consisting of a transition layer and an upper layer where each had uniform density in it. We oxidized a H-terminated Si (100) specimen of 15x15 mm$^2$ in dry O$_2$ flow at 1 atm. with a quartz tube furnace. The ramp rate was 33 K/min. We kept specimens at 1000 °C for 0 through 120 min to form SiO$_2$ films of 7.4 through 95.9 nm.

**Results** We measured XRR for the as grown samples. Then we etched the 95.9 nm sample with dilute HF solution and measured XRR at some thicknesses. The densities of the etched samples as a function of the thickness coincided with those of the as-grown samples. The transition layer was 1 nm thick and about 2.46 gcm$^{-3}$ in density for all the samples. The upper layer density changed from 2.35 to 2.25 gcm$^{-3}$ as the thickness increased. The upper layer thickness was obtained from the average value in its thickness according to our assumption. The density between the two-neighboring-measured points of the etched data might correspond to the real density distribution in depth. The results were as follows. The pre-formed layer of 7.4 nm including the transition layer which grew during the ramp-
up process had density of 2.35 g/cm³ or more. The density of the main layer formed at 1000 °C decreased rapidly to 2.25 g/cm³ at 20 nm, however the non-uniformity occurs mainly within 10 nm. The pre-formed layer remained stable at 1000 °C in 1 atm. for 120 min. The thermal SiO₂ films had stoichiometric composition. [3] The non-uniformity might be caused by change in structure of SiO₂-network.


**NS-TuP6 Nanoscale Surface Patterning for Controllable Metal Deposition. G. Smith, C.L. Berrie, University of Kansas**

Nanoscale patterning techniques using an atomic force microscope on self-assembled monolayers (SAMs) are used to controllably form metal nanofilms on gold and silicon surfaces. Nanoetching and nanografting are done on methyl-terminated SAMs to pattern in molecules with carboxyl termination. Organic self-assembled monolayers (SAMs) may be used to control the topography of nanofilms into methyl-terminated SAMs to achieve similar results. These nanofilms are then treated with an electroless metal plating solution to form metal nanofilms on selectively on the patterns. The patterns and features are characterized with atomic force microscopy. Controllable metal nanofilm formation has implications in nanoelectronics and nanodevice construction.


The CdSe quantum dots have been studied for various applications ranging from biomedicine imaging and sensing to hybrid solar cells. When the size of the nanocrystals reduces below the exciton Bohr’s radius, the energy levels become discrete (quantized). Consequently, the band gap of the quantum dots increases with decreasing the size of the dots. This is inherently reflected in the electronic structures of these quantum dots. In addition, these CdSe quantum dots are known to exhibit a phase transition from a stable hexagonal phase in larger dots to a metastable cubic phase at smaller dot sizes during their synthesis. As the size of quantum dot decreases, the number of energy associated with the surface increase leading to creation of vacancies, which results in the non-stoichiometric CdSe. However, the mechanism is not clear at atomic level. Therefore, we took a systematic approach to study the size-induced structural and electronic properties of CdSe quantum dots in toluene and drop-casted on Si by various in-situ and ex-situ imaging, spectroscopy and diffraction techniques to obtain the correlation between the quantum confinement and the corresponding stoichiometry, crystalline phases and the effect of surface ligands.


ZnO has increasingly been the focus for optical emission studies due to both its wide band gap of 3.37 eV and high thermal stability. In this research, vertically oriented ZnO nanowires (NWs) are grown upon silicon substrates using a novel, modified vapor-solid method within a vertical furnace. Electron-beam evaporation is then used with glancing angle deposition (GLAD) to functionalize the sides of the ZnO NWs with both Ag and MgO:Ag. Comparative characterization is then performed using photoluminescence measurements at varying angles to examine both surface-plasmon-enhanced emissions and to study the underlying coupling mechanisms responsible for the enhancements. This research may serve as the foundation for applications which range from high-efficiency LEDs to optoelectronic plasmon waveguides.
of integrated gas sensor device. To achieve a high-performance gas-sensor such experiments, we selectively grow SnO₂ nano-wires and verified measuring chamber to change the ionosorption rate. From the results of characterization is very actively performed because it is able to realize a


[1] Hyung Jin Kim, Yonghan Roh and Byungyou Hong, “Fabrication and characterization of DNA-templated conductive gold nanoparticle Chains, J.H. Heo, Korea Institute of Materials Science, South Korea, H.J. Kim, Sungkyunkwan University, South Korea

Recently, the Tin-oxide(SnO₂) has been widely used in the fabrication of various gas(O₂, CO, NO₂) detecting sensor. Basically, the SnO₂ is n-type semiconductor which has a wide band gap, about 3.99eV. Also, thin film of SnO₂ has been known as a transparent electrode and it is applicable for various electronic parts. Nowadays, research on SnO₂ nano-wire characterization is very actively performed because it is able to realize a micro/nano-electromechanical system with the fine alignment of SnO₂ nano-wire.

NS-TuP1 Preparation of Triethylamine Stabilized Silver Nanoparticles for Low-Temperature Sintering. S.L.C. Hsu, J.T. Wu, National Chia Kung University, Taiwan, R.O.C.

In this research, silver nanoparticles were synthesized by chemical reduction from silver nitrate using triethylamine as the protecting and reducing agents simultaneously. The average size of the silver nanoparticles was less than 5 nm, which allowed low-temperature sintering of the metal. X-ray diffraction (XRD), thermogravimetric analysis (TGA), and energy dispersive spectrometric (EDS) analysis results indicate that silver nitrate has been converted to silver nanoparticles completely. Using a 20 wt% silver nanoparticles suspension with thermal treatment at 150 °C, silver films with a resistivity of 8.09 x 10⁻⁵ W cm have been produced, which is close to the resistivity of bulk silver.

NS-TuP2 Selective Growth and Characterization of a SnO₂ Nano-Wire by Employing DNA-templated Gold Nano-particle Chains, J.H. Heo, Korea Institute of Materials Science, South Korea, H.J. Kim, Sungkyunkwan University, South Korea

Recently, the tin-oxide(SnO₂) has been widely used in the fabrication of various gas(O₂, CO, NO₂) detecting sensor. Basically, the SnO₂ is n-type semiconductor which has a wide band gap, about 3.99eV. Also, thin film of SnO₂ has been known as a transparent electrode and it is applicable for various electronic parts. Nowadays, research on SnO₂ nano-wire characterization is very actively performed because it is able to realize a micro/nano-electromechanical system with the fine alignment of SnO₂ nano-wire.

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Neutron Scattering Focus Topic
Room: East Exhibit Hall - Session NT-TuP

Neutron Scattering Poster Session


To understand the interfacial behaviors of materials under high pressure (P) and temperature (T) are of great importance and interest since lots of natural phenomena and practical applications involve those conditions. For instance, a metal surface/fluid interactions control success or failure of many attempts to engineer Earth’s subsurface for energy and/or environmental applications. The corrosion of metals and alloys in high subcritical aqueous systems, and especially in supercritical environments is an important safety issue in nuclear power plants.

However, due to the fact that most interface characterization techniques are difficult to implement at elevated P-T, little experimental attention has focused on solid/liquid interfaces at high P-T.

Neutron reflectometry (NR) is increasingly being used as a characterization tool to study the surface/interface of planar substrates. SPEAR at Lujan center is a Time-of-Flight (ToF) NR facility which is specifically designed to study solid-liquid interface and is able to in-situ monitor the surface/interface behavior with a space resolution of a few angstroms. The obtained real space model from reflectivity curve fitting can provide a lot of physical and chemical information about the interface.

One key gap to study the interfacial behaviors of materials under high P-T conditions is the lack of well designed pressure cell capable of handling P-T conditions close to or above supercritical conditions. To build up the capability of studying high P-T surface/interface, Lujan Center developed a special designed pressure cell which allows us to reach 200 MPa and 250 °C (in the future such cell will be equipped with with in-situ spectroscopic Raman and IR capabilities). Neutron is highly penetrating, which is able to “see through” high P-T aluminum cell walls and examining the surface/interface properties. Besides the pressure cell itself, the high P-T cell system includes three other subsystems: temperature, pressure and sample chamber environment control systems.


Bulk FeRh undergoes an antiferromagnetic to ferromagnetic phase transition as it is heated above room temperature. The addition of P lowers the phase transition temperature to that, in thin film form, the details of the phase transition can be studied while maintaining the same structural and morphological properties of the as-deposited film. The FeRh/Pd thin film was prepared by DC magnetron sputtering in an ultraclean sputtering system. A FeRh/Pd/Pt/FeRh/Pd trilayer was grown at 600°C on an a-axis sapphire substrate with a Rh seed layer and a Pt buffer layer. The epitaxial orientation of this 111-oriented thin film was confirmed by X-ray diffraction methods including standard high-angle diffraction, rocking curve analysis and pole figure analysis. The first-order metamagnetic phase transition and thermal hysteresis of the magnetic moment were examined by vibrating sample magnetometry. To study the detailed magnetic structure of a trilayer with a Pt spacer between two epitaxial films we applied polarized neutron reflectivity (PNR). PNR is used to detect the magnetic moment distribution in layered structures. Temperature-dependent PNR showed the dependence of ferromagnetic spin-splitting for the neutron reflectivity of the two spin polarization channels. Fitting of the PNR data shows a change of the spin splitting that is consistent with vibrating sample magnetometry data. PNR measurements at two different applied magnetic fields of 1 T and 0.5 mT revealed the dependence of magnetic splitting on applied magnetic field and a modification of the thermal hysteresis. This data confirms the strong field dependence of the magnetically stable state. Analysis of the off-specular neutron reflectivity data will show how the magnetic domains change with experimental conditions. The authors gratefully acknowledge financial support from DOE award DE-FG02-08ER46499. Research at Oak Ridge National Laboratory’s Spallation Neutron Source was sponsored by the Scientific User Facilities Division, Office of Basic Energy Sciences, U.S. Department of Energy.
SE-TuP2  Fabrication of Multilayer X-ray Gratings on Staircase Substrates. C. Liu, Argonne National Laboratory, S. Lynch, National Heart, Lung, and Blood Institute. L. Assoufid, Argonne National Laboratory, H. Wen, National Heart, Lung, and Blood Institute

Classical hard x-ray transmission gratings fabricated using lithography are limited by a small aspect ratio of less than 20 when the grating periods are smaller than 2 microns. State of the art x-ray phase contrast imaging techniques require high density transmission gratings with smaller periods and higher aspect ratio to cover larger energy range for thicker samples. This problem may be addressed with a sliced multilayer technique, similar to that used in multilayer Laue lenses and multilayer transmission mask gratings. To increase the acceptance area of the grating, it was proposed [1] to use a thin Si substrate that is asymmetrically etched to a staircase with each stair supporting a multilayer parallel to its surface. Using an x-ray beam shining through the layers at an oblique angle to the substrate and parallel to the layer surfaces, one hopes to have a large-area transmission grating with small multilayer periods. The first fabrication test is completed using W/Si multilayers with dc magnetron sputtering deposition. The layer thickness is designed to be the period/2N, where 2N is the total number of layers. Preliminary results and challenges are presented.


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SE-TuP3  Investigations on Physical Processes for Low Temperature Plasma Activated Wafer Bonding. T. Thomas Pluch, K. Hingerl, Johannes Kepler University, Austria. T. Dragoi, M. Wimpflinger, EV Group, Austria

Direct wafer bonding is a straightforward method of directly connecting wafers, with suitable (in terms of micro-roughness, flatness and cleanliness) surfaces, permanently to each other, by bringing them into contact and subsequently annealing them or simply storing them. The conventional process for hydrophilic oxidized Silicon surfaces (native as well as thermal oxide) is well understood, and explained the following way [1]:

Up to 100°C the substrate surfaces are held together via van der Waals interaction which is mediated by a few monolayers of water. In the range of 100-200°C the water diffuses away from the interface both along the interface and through the oxide into the crystalline bulk, where it reacts with the silicon and forms oxide. The increase of the bond strength from 50% to 100% of Si bulk strength is usually attributed to a closing of gaps at the interface, which starts at the softening temperature of the thermal oxide at around 900-1000°C, depending on whether dry or wet oxide was used.

Low temperature plasma activated direct wafer bonding is a process that lowers the required annealing temperatures necessary for reaching high bond strength. One example for such an improvement is a pair of native oxide – thermal oxide wafers, where bulk strength can be realized by plasma activation with subsequent annealing at temperatures below 200°C. At this temperature conventional wafer bonding reaches half of Si bulk strength, and is limited by gaps at the bonding interface. The mechanism behind this improvement compared to the non activated process is still under discussion.

To clarify the mechanism for this commercially available process, different bonding experiments were performed to evaluate the lifetime of the surface activation and the achievable bond strength when varying some of the boundary conditions of the process (substrates with different orientations, various plasmas, and lowering the annealing temperature).

By partly covering wafers during plasma activation, comparisons between the activated and non-activated regions could be made on single wafers. Therefore the influence of the slightly different substrates could be eliminated. Such wafers were then analyzed using atomic force microscopy, by spectroscopic ellipsometry, by Auger analysis and by X-ray photoelectron spectroscopy.

Finally a model for the mechanism, which was derived from the model for the conventional bonding process, and which explains the experimental results will be presented.

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surface alkoxy or amide is formed via the deprotonation of the corresponding alcohol or amine by surface atomic oxygen, which can subsequently nucleophilically attack the aldehydes (formed in situ from alkoxy or introduced directly) and produce the corresponding ester or amide. The product distribution of our low-pressure experiments matches remarkably well with gas phase reactions carried out in the ambient condition as well as liquid phase reactions, proves the generality of the mechanism.


Until recently it had been extremely difficult to experimentally address one of the most fundamental questions about graphene: How do the quantum physics of graphene near the Dirac point coincide with the Dirac point energy? Here we address this question by investigating graphene on a particularly interesting substrate, the carbon face of SiC, with high-resolution angle-resolved photoemission spectroscopy (ARPES). We present the first direct measurements of the self-energy in graphene near the neutrality point, and show that the many-body physics in graphene differ from those of an ordinary metal. These exciting findings set a new benchmark in our understanding of many-body physics in graphene and a variety of novel materials with Dirac fermions.

SS-TuP3 Edge Termination of Modified Graphene Oxide during Thermal Exfoliation. M. Aick, Y.J. Chabal, The University of Texas at Dallas

Nanopore formation in carbon materials (e.g. exfoliated nanostacks of graphite) has been widely studied through mechanical exfoliation, intercalation, electrochemical separation, chemical or thermal exfoliation of graphite oxide (GO) via expansion with partial oxygen removal. Amongst all these methods, exfoliation of modified graphene (GO), a solution-processable precursor compound where aromatic and heterocyclic rings with embedded oxygen functionalities exist, by thermal processing still remains elusive for the following reasons: (1) poor control of GO composition (initial oxygen content), (2) poor understanding of the chemical composition, (3) unknown role of oxygen, (4) strong interactions, and edge termination with oxygen. Infrared absorption spectroscopy coupled with in-situ thermal annealing process [1] makes it possible to examine the chemical changes taking place during thermal reduction to identify and understand interacting molecular environment and the edge functionalization. To unravel the complex mechanisms leading the removal of oxygen in GO, we have performed in-situ transmission infrared absorption spectroscopy (IRAS) measurements of graphene/graphite oxide (GO) thin and bulk films upon thermal annealing (60-850°C) in vacuum (10⁻¹⁰⁻ Torr). Control of the edge geometry of finite-sized modified graphene flakes depends very much on the control of the processing methods. This edge reconstruction further determines electronic, electric and mechanical properties of the exfoliated modified graphene flakes. Therefore, we not only perform studies deriving a thermal reduction mechanism, but also examine the edge reconfiguration with oxygen. We report here the observation of a surprisingly strong IR absorption band that occurs only upon thermal reduction of GO. After annealing at 850°C in vacuum, the strong enhancement of the new IR active absorption band is observed at ~800 cm⁻¹ [2]. The intensity of this band is 10-100 times larger than what is expected for the oxygen content of the reduced GO, namely between 5 and 8 at.%. This band is assigned to a specific oxidation state, involving oxygen located in the basal plane (forming C-O-C bonds) and at atomically straight edges of reduced graphene. The large enhancement in IR absorption is attributed to the direct participation of electrons, induced by the asymmetric C-O-C stretch mode displacement. These findings open new possibilities in the field of nanoelectronics for all sensor and energy storage applications. [1] M. Aick, et al. J. Am. Chem. Soc. (2011), in preparation. [2] M. Aick, et al. Nat. Mater. 9, 840-845 (2010).

SS-TuP4 Electrocatalytic Surfaces: Structure, Reactivity and Nanotemplating. X.F. Yang*, Lehigh University, B.E. Koel, Princeton University

Electrocatalysis in energy related applications such as fuel cells and hydrogen production is perhaps the most important enabling technology of the future. Pt-based electrocatalysts are widely used because of their exemplary performance, but these catalysts have serious drawbacks, e.g., cost, modest efficiency, and low durability, which limit fuel cell development. Our research explores non-Pt electrocatalysts or ultrathin-Pt film electrocatalysts to gain insight and discover materials that can replace Pt or greatly reduce Pt loadings while retaining or even increasing activity and/or stability. Our approach is to using model electrocatalyst surfaces with well-defined composition and structure to simplify and exert control on the system to improve our understanding of the surface phenomena that control electrocatalytic reactions. In this work, four types of model electrocatalysts were prepared in UHV: (i) Pd/Fe(111), (ii) Au/Pd/Fe(111), (iii) Pt on a faceted C/Re(11-21) nanotemplate, and (iv) Pd/Re(111) (poly). These surfaces were characterized using LEED, XPS, LEIS, and AES, and then their electrocatalytic activity for the oxygen reduction reaction (ORR), the hydrogen evolution reaction (HER), and ethanol oxidation (EO) reaction was measured. Significant surface segregation of Pd was discovered after clean Pd/Fe(111) was annealed at high temperatures in UHV. The surface structure strongly depends on the annealing temperature, with the formation of an atomically smooth, random substitutional alloy by heating to 1000 K, and the formation of Pd monomer and dimer adatoms by heating to 1250 K. The annealed Pd/Fe(111) surfaces exhibit the highest ORR activity. When a submonolayer amount of Au was deposited on Pd/Fe(111), the Au/Pd/Fe(111) surface was found to be highly active for the HER. In addition, a Pt monolayer on a Pt surface on a polytetrafluoroethylene substrate displayed great improvement in reactivity for electrochemical ORR and EO. In summary, we have investigated a range of non-Pt and ultrathin-Pt film model electrocatalysts that are more active than pure Pt and that point to new materials that could be used to reduce cost and improve activity by nanoeengineering novel electrocatalysts.


Oxide-supported bimetallic Pt-Au and Ni-Au clusters were studied as model catalysts for low temperature oxidation reactions. The growth, composition, and chemical activity of Ni-Au and Pt-Au clusters deposited at 300 K on TiO2(110) were investigated using scanning tunneling microscopy (STM), low energy ion scattering (LEIS), X-ray photoelectron spectroscopy (XPS), and temperature programmed desorption (TPD). The formation of the Au-TiO2 interface for low temperature oxidation was illustrated by a series of TPD and STM experiments that show a direct correlation between the number of Au-TiO2 interfacial sites and the activity on the surface. Bimetallic clusters were grown by first depositing Ni or Pt onto the surface in order to seed the more mobile Au at existing Ni or Pt clusters. The surfaces of the bimetallic clusters are significantly enriched in Au and are almost entirely pure Au for clusters with > 50% bulk Au composition. TPD of methanol and CO on bimetallic clusters that are highly enriched in Au at the surface still show significant activity characteristic of Ni or Pt on the surface, suggesting that methanol and CO are able to induce the diffusion of Ni and Pt to the surface of the clusters. Heating the bimetallic clusters above 600 K results in the selective encapsulation of Pt or Ni by a thin film of TiO2. Post-annealed Pt-Au and Ni-Au bimetallic clusters show promise for enhanced activity towards the conversion of methanol to formic acid.

SS-TuP6 Pattern Formation through Leveled Copper Etching after Dysfunctional Electropolishing. A.D. Pauric, P. Kruse, McMaster University, Canada

Electropolishing is a common industrial practice whereby a metallic surface is subjected to an anodic potential in an appropriate electrolyte to produce a microscopically smooth surface. It is typically performed in concentrated phosphoric acid with cell potentials ranging from between 1.3 and 2.3V. However, little research has been conducted in the parameter space outside the electropolishing regime. Previous research in our group using deviations from electropolishing conditions has characterized a wide range of fascinating surface structures including terraces, oxide nanotubes, stripes, and dimples. Our current work involves using the copper/phosphoric acid system as a model system to study deviations from electropolishing conditions and the resultant surface features. Upon exposing copper substrates to a high applied cell potential in concentrated phosphoric acid, etched surface patterns up to over a micron in depth are observed. Characterization of the patterns includes the use of atomic force microscopy (AFM), scanning electron microscopy (SEM), and surface enhanced Raman spectroscopy (SERS). The surface patterns evolve with variation in temperature and phosphoric acid concentration. A distinguishing feature is that the patterns are etched into rather than grown upon the copper substrate, promoting mechanical stability. Additionally, the tops of the surface features are level with respect to their surroundings. Potential applications include electrodes, current collectors for lithium ion batteries, catalysts, micro-cooling, and substrates for the fabrication of other surface morphologies.
SS-TuP7 Surface Chemistry of Atomic Layer Deposition of Manganese Thin Films. H. San, X. Qin, F. Zaera, University of California, Riverside. Manganese thin films potentially be used as Cu diffusion barriers in microelectronic devices, and may possibly be grown by atomic layer deposition (ALD), to produce highly uniform thin films with good conformality in high aspect-ratio structures. Here, the early stages of film growth of two precursors, methylcyclopentadienyl manganese tricarbonyl (CH₃C₅H₄Mn(CO)₃) and dimanganese decacarbonyl (Mn₂(CO)₁₀), were investigated by X-ray photoelectron spectroscopy (XPS) to assess their viability for ALD of manganese thin films. In most cases, only oxidized manganese could be deposited on silicon substrates covered with their native oxide layer. Only in depositions using (CH₃C₅H₄Mn(CO)₃) at relatively high temperatures (>300°C) it was possible to detect a low binding energy feature in the Mn 2p XPS that could be assigned to metallic Mn (although it is also possible to come from manganese silicate). It was also observed that the lowest binding-energy manganese species appeared only after the growth of a layer of oxidized manganese, and seems to form in the sub-surface. Electron-induced deposition was also studied and compared with the thermal process. The Mn₂(CO)₁₀ precursor was much more reactive, and could lead to multilayer deposition by itself at temperatures as low as 200°C.

SS-TuP8 Spin Effects on Metal Surface Reactions: O₂ on Ferromagnetic Pt. M.C. Escano, N.T. Quang, H. Nakanishi, Osaka University, Japan; E. Goyene, The University of British Columbia, Canada; H. Kasai, Osaka University, Japan. We studied O₂ chemisorption on Pt surface in the need to promote O₂ activation. Activation, in this case, is defined as lowered activation barrier for O₂ dissociation but minimized O adatom (O₂ ad) binding energies; or a significantly stretched O-O bond in a loosely bound molecular O₂. This kind of reaction is not to be attain since the O₂ molecules are able to interact with the Pt surface in its reaction path. However, such unique reaction is often sought in many electrochemical/chemical systems (i.e. fuel cell cathode catalyst, three-way automotive catalyst). Here, we show how the magnetic state of Pt can achieve this desired reaction. The model system involves a non-magnetic Pt layer pseudomorphically laid on top of magnetic surface (M=Fe(001), Co(001)). The inter-layer coupling of the Pt-M spins gives a transition temperature of Pt (Tₘ) at which the induced spin density functional theory, suggests that the magnetic ground state of the system is a ferromagnetically coupled Pt-M layer[1]. The induced spin moment of 0.5μB in the Pt layer is in agreement with x-ray Magnetic Circular Dichroism (XMCD) [2]. The hybridization of Pt-5d with the M-3d states give characteristic exchange splitting similar to the spin-reversal induced x-ray magnetic circular dichroism spectroscopy [3]. Using the Heisenberg spin Hamiltonian to determine the exchange interaction, we note that strong inter-layer coupling of the Pt-M spins gives a transition temperature of Pt that is lower than room temperature, in agreement with [2,3].

Potential energy surfaces obtained for O₂ dissociative adsorption on ferromagnetic Pt layer show much lower activation barrier for dissociation and also lowered O₂ ad binding energy. For molecular adsorption, the O₂ vibrational frequency is lower on ferromagnetic Pt as compared to paramagnetic Pt, despite the much lower binding energies on the former. We note that spin effects played significant role rather than charge transfer effects on the over-all structure and binding of the O₂ at transition state and on the surface [4]. This will be discussed in the meeting in terms of local density of states, charge transfer and spin density, and other parameters involved in the bonding and magnetic interactions.

References

SS-TuP9 X-ray Diffraction Study on Hydrogen-Induced Pd(110) Surface Reconstruction. M. Takahashi, S. Fujikawa, W. Hu, Japan Atomic Energy Agency, H. Tajima, Japan Synchrotron Radiation Institute. Absorption of hydrogen into a substrate begins with dissociated adsorption of hydrogen molecules. The mechanism of the transition from adsorption to absorption is an important knowledge for improving the efficiency of hydrogen storage materials. The aim of the present study is to verify the hydrogen absorption model in atomic scale through quantitative determination of structures of hydrogen-adsorbed Pd(110) by synchrotron X-ray diffraction. Experiments were performed at a synchrotron beamline 13XU at SPring-8 using a surface X-ray diffractometer integrated with a UHV chamber equipped with a cryostat. The sample was Pd(110) single crystal which was 10 mm in diameter and 3 mm in thickness. The clean Pd(110)-(1×1) surface was prepared by chemical polishing etching in HCl solution and cycles of Ar-sputtering and annealing at 900 K in UHV. For the hydrogen adsorption experiments, the hydrogen pressure was carefully controlled with a needle valve and a nude ion gauge. We measured five crystal truncation rod (CTR) profiles with increasing substrate temperature to room temperature from 57 K after the substrate was exposed to 10⁻⁷ Torr’s hydrogen. Before and after hydrogen adsorption at 57 K, the CTR profiles changed only slightly. More distinctive changes were observed when the sample temperature reached 180 K. At room temperature, the CTR profiles were accounted for by a structure model with disordering of surface atoms. According to past Low-energy electron diffraction, He diffraction and thermal desorption spectroscopy studies, hydrogen adsorption below a substrate temperature of 120 K induced the (1×2) surface reconstruction with 1.5 monolayer (ML) hydrogen through the (2×1) structure at a hydrogen coverage of 1 ML. When the temperature is raised to 500 K, the (1×2) reconstruction is broken down at low coverages and the 2×1 phase without hydrogen desorption. A comparison with our X-ray diffraction results and these past studies shows that the change of the CTR profiles at 180 K corresponds to the transition from the (2×1) to (1×2) reconstructions and concomitant incorporation of hydrogen into subsurface. These results provide us with an atomic-scale picture that explains how adsorbed hydrogen is absorbed into the Pd bulk.

SS-TuP10 Single Molecule Force Spectroscopy Studies on Nanoclay Surfaces. B. Ozkaya, G. Grandmeier, University of Paderborn, Germany. Sequential adsorption of oppositely charged polyelectrolytes leads to multilayered thin films via electrostatic self-assembly. Incorporation of sheet-like inorganic nanoparticles is a promising method to improve barrier and ion transport properties, as well as mechanical properties in such films. The interface chemistry between the nanosheets and the polyelectrolyte segments plays a crucial role in design of these hybrid materials. In the present study, we have investigated the effects of pH and ionic strength on the adsorption of single polyelectrolyte molecules on natural clay (Na-Montmorillonite) platelets by means of AFM-based single molecule force spectroscopy (SMFS). SMFS is one of the few methods where in-situ experiments can be performed to obtain precise and quantitative information on interaction forces at a molecular level. In order to perform SMFS on clay platelets (lateral size: 50-300 nm), a heterogeneous model surface has been obtained via electrostatic immobilization of exfoliated clay platelets on template stripped ultra-flat Au(111) surfaces. Probe molecules (polyallylamine) were covalently attached to the gold coated AFM cantilever. Equilibrium desorption plateaus of constant force were obtained from the successive force-distance measurements. By adjusting the ion valency in the electrolyte, selective information from clay platelets could be obtained. In the presence of monovalent ions, desorption plateaus of constant force in the range of ~50 pN were obtained from pH 3 to pH 7. At pH values above the pKₐ value of polyallylamine, confirmed with polarization modulation infrared reflection absorption spectroscopy (PM-IRRAS) measurements on cast polyallylamine-needle detectable desorption event even at low force-distance measurements. The constant negative surface charge of clay platelets simplifies the evaluation of effects of electrical double layer and polymer line-charge density on adsorption of the polyelectrolyte molecules. Overall, the results of pH and ionic strength dependent de-adhesion measurements provides valuable information on the interplay between surface properties and polyelectrolyte adsorption giving the basis for an improved understanding of the behavior of silicate nanoparticles in polyelectrolyte films.

SS-TuP11 Atomic Structure of Aluminium on Si(110): STM and First-principles Study of “4 × 6” Reconstruction. M. Yoshimura, D. Matsuoka, Toyota Technological Institute, Japan. With the miniaturization of semiconductor devices, low-dimensional structures such as quantum wires and quantum dots have recently attracted much attention. The “16 × 2” on a clean Si(110) surface has been noticed as an example, and unique surface spectroscopy for the fabrication of low-dimensional nanostructures, because the one-dimensional undulated terrace structure of monatomic height and of about 2.5 nm width are formed. Because the hole mobility of Si(110) surface is about 1.5 times as large as that of Si(100) typically used as substrates of present semiconductor devices, higher speed operation is expected [1]. Thus, the interaction between Si(110) and metal is very important [2,3]. Aluminium is the typical metal forming Schottky barrier with silicon and the Al-adsorbed structures have been well examined on Si(111) or Si(001) plane [4,5]. In contrast, little is known about the structures of the Al/Si(110), particularly in real-space. In this study Al/Si(110) surfaces are investigated in real space using STM and first-principle calculation.

The experiments were carried out in an ultrahigh vacuum (base pressure: 2.0 × 10⁻⁷ Pa). First-principles calculation was performed using VASP program [6]. Aluminium was deposited on a clean Si(110)-“16 × 2” at 600
In the empty state, alternative arrangement of zigzag rows and straight rows is visible in empty states. When filled states are prominent, the transition of defects in the zigzag row. They are assumed to be silicon substitutional defects, as is the case for Al/Si(111) [8]. Thus we conclude Al and Si atoms formed zigzag and straight rows, respectively. In addition, the straight row of the silicon was composed of the silicon pentagon [9] as revealed by high-resolution STM. Based on the above observation, an atomic structural model is proposed and discussion will be made based on the theoretical calculation.


SS-TuP12 Adsorption Dynamics of Ethylene on Si(001), M.A. Lipponer, N. Armbust, University of Marburg, Germany, M. Durr, HS Esslingen, Germany, U. Hofer, University of Marburg, Germany

The functionalization of semiconductor surfaces by means of organic molecules is of great interest due to possible applications in the field of nanoelectronics. However, only little information on the reaction dynamics of these systems is available. In this work, the adsorption dynamics of ethylene on Si(001) has been investigated. With ethylene being the most simple unsaturated organic molecule, the system serves as a model system for non-activated adsorption on semiconductor surfaces. In order to investigate its reaction dynamics, we employed a supersonic molecular beam which allows for the control of the kinetic energy of the impinging molecules. Additionally, surface temperature and the excitation of internal degrees of freedom varied when measuring the sticking coefficients as a function of relative surface coverage by means of King and Wells techniques.

With increasing kinetic energy of the impinging molecules, we find a decrease of the initial sticking coefficient as it is typical for non-activated reaction channels; the excitation of the internal degrees of freedom is shown to have a major impact on the reactivity. With increasing surface temperature, a decrease of the initial sticking coefficient is observed. Suprisingly, the monolayer surface coverage also decreases with increasing surface temperature. The results are discussed in the context of a reaction channel via a mobile precursor with long lifetime at low temperatures.

SS-TuP13 Surface Characterization of Polymeric Materials using TOF-SIMS and XPS, J. Lee, K.-J. Kim, Korea University, Republic of Korea, Y. Lee, Korea Institute of Science and Technology, Republic of Korea

TOF-SIMS and XPS are very useful techniques for the analysis of solid surfaces composed of organic and polymeric species. In order to widen the application of TOF-SIMS and XPS to archeology and plasma process, we analyze the dyed fabrics and the fluorocarbon thin films. First, the fabrics dyed with natural dyes and synthetic dyes were investigated with TOF-SIMS and XPS. Dyes investigated belong to various chemical groups, which include indigo, carthamin, crocin, shikonin, curcumin, purpurin and alizarin. TOF-SIMS and XPS spectra for the dyed textiles showed specific molecular ions and fragment ions from organic dyes as well as elemental ions from metallic mordants. Several ancient fabrics were also analyzed to identify the natural dyes. Secondly, Fluorine-containing hydrophobic thin films were obtained by two different plasma methods: inductively coupled plasma (ICP) and pulsed plasma (PP). Three kinds of fluorine-containing gases, as C-F, C-F-C, and C-F-C-F, are generated using plasma polymer films. Process parameters for plasma polymerization such as gas ratio, gas pressure, pulse frequency, and processing time were investigated. Surface analytical instruments such as TOF-SIMS, XPS, and AFM were used to characterize the fluorocarbon thin films generated by ICP and PP. In this work, TOF-SIMS was used to provide useful information about the chemical properties including surface composition and XPS was used to examine the chemical structure of dyed fabrics and fluorocarbon films.

SS-TuP14 Links Between the Surface Atomic Arrangement and Catalytic Properties of Bimetallic Alloys: A First Principles-based Investigation, J.A. Stephens, H.C. Ham, G.S. Hwang, University of Texas at Austin

Catalysts composed of more than one metallic element often exhibit remarkable activity and selectivity compared to their monometallic counterparts. These synergistic properties often can be explained in terms of two kinds of effects: modification of catalyst electronic structure due to interactions between dissimilar metal atoms (ligand effects) and the presence of mixed-metal surface sites that, because of their size and shape, promote some chemical reactions more than others (ensemble effects). Understanding in detail how ligand and ensemble effects operate in particular cases is an important step toward realizing the longer term goal of rational catalyst design. Experimental study has provided valuable insight into this problem, but progress has been hampered by the difficulty of studying reacting systems with atomic resolution. We employ the tools of molecular simulation in a two-pronged approach to complement these efforts. First, we use density functional theory to explore how particular atomic arrangements in the surfaces of bimetallic alloys influence their catalytic function. We have found, for example, that the experimentally-observed stability of Au-Pd catalysts to promote the direct synthesis of hydrogen peroxide may depend on the presence of surface Pd monomers surrounded by Au. Our calculations indicate that on larger Pd ensembles, O-O bond scission can more readily occur, leading to peroxide. We have similarly studied the oxygen reduction reaction and carbon monoxide oxidation on Au-Pd alloys. In the second part of our work, we attempt to predict how the metal atoms in catalyst surfaces are actually arranged using a combination of density functional theory, the cluster expansion method, and Monte Carlo simulation. From these simulations, we have obtained the temperature and pressure-dependent composition, size and shape distributions for the (111) and (100) surface facets of Au-Pd and Au-Pt alloys. Our results are in good agreement with available experimental observations.


Supramolecular motifs that provide controllable functionalization of surfaces on a nanometer scale offer significant advances in organic solar cells and other surface supported systems. Based on recent advances in two-dimensional organic self-assembly at the liquid/solid interface, we are developing complex surface architectures based on versatile nitrogen-rich heterocycles. This work focuses on the molecular design and subsequent adsorption of this class of compounds onto graphite (HOPG) utilizing liquid scanning tunneling microscopy to resolve the packing of the individual molecules. Current molecular design includes extended alkyl side chains, other substituents including amide linkages to facilitate intermolecular ordering. Exploration of photo-activated, tip-induced, and chemically stimulated switching of these molecules is underway to enable manipulation of patterning and packing.

SS-TuP17 Electrical, Physical, and Chemical Properties of the Metal to Amorphous Hydrogenated Boron Carbide Interface, M.S. Driver, S. Karki, A.N. Caruso, University of Missouri - Kansas City

Boron carbide (BC), as a semiconducting material, has been under scrutiny for several decades for its use in heterostructure devices toward applications in solid-state neutron detection and thermoelectric energy conversion. The heterostructure devices are typically heterojunctions in which p-type amorphous hydrogenated boron carbide (a-B:CH) is deposited onto n-type Si, with the Au contact instead of the a-B:CH face. Traditionally it has been believed that Cr forms an ohmic contact at the a-B:CH interface and that the heterostructure current–voltage rectification is a classical function of the p-n junction. However, we have found through photoemission studies of Cr overlayers on a-B:CH, that a complex series of interfaces is formed, involving various oxides and borides, which are likely contributing and could be a factor in the observed rectification. To follow up the Cr findings, we have explored Al, Cu, Au, Ag, and Ti using the same set of studies. This talk will provide an overview of the electronic, physical, and chemical interfaces between the above metals and a-B:CH in the context of understanding the (I(V) characteristics of the presumed p-n junction.

SS-TuP22 Electron Stimulated Reactions on Graphene-Coated Ru: Relevance to Extreme Ultraviolet Lithography (EUVL), B.V. Yakshinsky, R.A. Bartynski, Rutgers University

The contamination of optical surfaces in EUVL exposure tools, operating at 92 eV photon energy, results in degradation of the mirror reflectivity. We report studies of the thermal and electron-induced interaction of benzene and toluene vapors, typical background gases, with the Ru surface, as model cap layer for multilayer mirrors (MLM), using temperature programmed desorption (TPD), X-ray photoelectron spectroscopy (XPS), low energy ion scattering spectroscopy (LEIS), electron suggested desorption (ESD), low electron energy diffraction (LEED), and scanning tunneling microscopy (STM). A low energy electron source (100 eV) is used to simulate radiation excitations on the surface produced by EUV photons. Heating of adsorbed hydrocarbons leads to a stepwise dehydrogenation and buildup a self-limiting carbon monolayer. Electron bombardment of the bare Ru surface in the presence of gas phase hydrocarbons inevitably results in rapid accumulation of 1 ML of carbon or carbonaceous species. Subsequent
The effect of carbon in the structure of Mo$_2$B is increasing the superconducting transition temperature from 5.8 K to 7.5 K. Polycrystalline samples of the Mo$_2$B and Mo$_2$BC were synthesized by the arc melting technique. The samples were characterized by x-ray diffraction (XRD), x-ray photoelectron spectroscopy (XPS) and ultraviolet photoelectron spectroscopy (UPS). XRD results show that when carbon atoms are added to the structure of the Mo$_2$B compound a transition structural from body centered tetragonal to face centered orthorhombic is induced forming the Mo$_2$BC compound. The presence of the Mo$_2$B and Mo$_2$BC structures is confirmed with the band gap of 1.2-1.5 eV, whereas potential application in many technological fields. Copper oxide-based materials have been widely investigated due to their property of the F-DLC thin film under a vacuum showed 0.12 at a sliding voltage of 5 kV. PTFE doping was performed by using the electron-beam deposition method with PTFE evaporation rate from 0.05 to 0.2 nm/sec. The film thickness measured using QCM was kept at 200 nm. Film composition and microstructure were investigated by X-Ray photoelectron spectroscopy and Raman spectroscopy, respectively. The hardness was measured by an indentation method with a Knoop indenter. The friction coefficient was measured friction of a SUJ2 ball with a constant normal load of 98 N until the sliding distance reached to a length of 100 m. The friction coefficient property under the vacuum was measured in 5 x10$^{-4}$ Pa. Frictional properties changed under the vacuum and the atmosphere. Friction coefficient of the F-DLC thin film with PTFE deposition rate of 0.2 mm/sec showed 0.25 at a sliding distance of 100 m. Friction coefficient property of the F-DLC thin film under a vacuum showed 0.12 at a sliding distance of 100 m. It was found that F-DLC indicated a low friction coefficient in a vacuum, F-DLC can be expected to be used as one of space materials in the future.
respectively. Chromatic change of a methylene blue solution was applied to a photocatalytic property. Light irradiation to TiO2 films in a methylene blue solution was carried out by using a commercial sterilizing lamp as ultraviolet light and an artificial sun light as visible light. Transmittance of a methylene blue solution was measured by a spectrophotometer. Furthermore, photocurrent between the TiO2 film and a platinum electrode was measured by a volt-amper characteristic using an unresisted ammeter in a KCl solution of 0.5 mol/l.

The crystal structure of TiO2 turned from a rutile type into an anatase type with increase of O2 gas flow rate. Photocatalytic property and photocurrent property of an anatase type of TiO2 showed the high value. The effect of the Cu addition exhibited different behavior according to each crystal structure. As for the photocatalytic property, the effect of the Cu addition was observed in rutile type, while on the photocurrent property the effect of the Cu addition was observed in anatase type. It was considered that the charge separation between an electron and a hole was enhanced by adding Cu to the TiO2 surface.

SS-TuP28 Deviation from Wulff Structures for Pt Nanoparticles Supported on TiO2(110): A STM Study. F. Behafarid, B. Roldan Cuenya, University of Central Florida

This study reports the shape of micellar platinum nanoparticles (NPs) supported on TiO2(110) resolved by scanning tunneling microscopy (STM). Since micellar NPs are initially spherical in shape at room temperature, the 3D distribution of shapes obtained under high temperature annealing display structures typical of thermodynamic equilibrium, in contrast with kinetic shapes of physical vapor deposited NPs, which are normally 2D and dominated by the diffusion of metal atoms on the support. All of the NPs resolved show an epitaxial relationship with the support, evident by TiO2(110)[001] direction being one of the symmetry axis of the NPs. Analyzing these shapes reveals that in addition to hexagonal top (type A) and rhombic top (type B) NPs reported for evaporated NPs, there are two additional categories of shapes observed for micellar NPs (Type C and D). Type A and B consist only of 111 and 100 facets at the interface with the support and the free top surface. However, type C and D feature also 110 facets either at the interface with the support or at the free surface. This facet orientation was not observed in other studies due to its instability and high surface energy. The ratio of 100/111 facets obtained for these shape deviates from the Wulff structure. This deviation could be attributed to size effects, interface-induced strain, and/or possible TiOx adsorbates on exposed facets.

SS-TuP29 Slope Selection in Au Growth Fronts by Protrusion Interactions. J.L. Sacedón, A. González González, J.A. Aznárez, E. Rodríguez Cañas, E. Vasco, Consejo Superior de Investigaciones Científicas, Spain

Growth fronts of polycrystalline films composed by paraboloid-like surface protrusions (SPs) have been recently decomposed into elemental paraboloidal meridian zones, which allow a comprehensive statistical analysis of the film surface [1]. Using this analysis, local height distribution curves have been synthesized [2] and the interface width expressed as a function of the statistical parameters [3]. The method evidences that the average value of the slopes obtained at the SP border remained constant during the growth [2]. The analysis of distribution curves of the terrace width (~ the inverse of the slope) at the SP borders provides three terrace classifications typical of thermodynamic equilibrium, in contrast with kinetic shapes of physical vapor deposited NPs, which are normally 2D and dominated by the diffusion of metal atoms on the support. All of the NPs resolved show an epitaxial relationship with the support, evident by TiO2(110)[001] direction being one of the symmetry axis of the NPs. Analyzing these shapes reveals that in addition to hexagonal top (type A) and rhombic top (type B) NPs reported for evaporated NPs, there are two additional categories of shapes observed for micellar NPs (Type C and D). Type A and B consist only of 111 and 100 facets at the interface with the support and the free top surface. However, type C and D feature also 110 facets either at the interface with the support or at the free surface. This facet orientation was not observed before in other studies due to its instability and high surface energy. The ratio of 100/111 facets obtained for these shape deviates from the Wulff structure. This deviation could be attributed to size effects, interface-induced strain, and/or possible TiOx adsorbates on exposed facets.

SS-TuP38 UV Induced Photodesorption of O2 on Rutile TiO2(110): An Angular Imaging Study. D.P. Wilson, M.D. Kershis, Brookhaven National Laboratory, M.G. White, Stony Brook University and Brookhaven National Laboratory

The binding states of oxygen on TiO2(110) are important for many reactions, including the degradation of organic compounds and photodesorption. Experimental data and theoretical calculations have shown many different configurations for O2 on TiO2(110), including O2, O22-, and O2-. Defect sites in titania (bridging oxygen vacancy, Ti3+ interstitial), while necessary for the adsorption of oxygen, also play a role in the initial binding states. In this experiment, the angular distributions and images of oxygen photodesorption from a rutile TiO2(110) surface are studied under UHV conditions using a pump-probe Time-of-Flight (TOF) detection scheme to help determine the initial binding states of oxygen on this surface. Excitation occurs via photos of 3.7 eV photons followed by one-photon ionization using 13.05 eV photons. The delay time between the lasers can be varied according to the maximum desorption velocity of the oxygen molecules. Ions were detected using a dual microchannel plate and a phosphor screen. A CCD camera positioned behind a phosphor screen captured the light emission of the phosphor, allowing for the imaging of the desorption events. The spatial resolution of the images was limited by the motion of the ions through a Time-of-Flight (TOF) mass spectrometer to generate a probability distribution for detection that was used to compare images at different delays. Previous experiments on the oxygen velocity distribution on TiO2(110) showed 3 different “channels” for desorption, with two being “fast” and one being “slow.” The velocity distribution for the slow channel tracked with surface temperature, indicating that a trapping desorption mechanism dominated this channel. The “fast” channel, however, did not depend on temperature and were attributed to two different oxygen binding states on TiO2(110). Many different binding states of oxygen have been predicted and it is unclear which state is responsible for which channel. Images of the fastest channel and comparisons at different coverage and photon flux are shown.

The methyl radical velocity distributions from 4 different ketones (acetaldheyde, acetone, butanone, and acetoephone) were also investigated. For ketones where methyl radical desorption is not the preferred pathway (butanone), a 2+1 REMPI scheme was used for signal enhancement. Using these model systems allows for direct comparison of different properties of each molecule.

Thin Film Division

Room: East Exhibit Hall - Session TF-TuP

Thin Films Poster Session

TF-TuP1 Spatially Varied Orientation Selective Epitaxial Growth of CeO2(100) and (110) Areas on Si(100) Substrates by Reactive Magnetron Sputtering Utilizing Electron Beam Irradiation. T. Inoue, S. Shida, Iwaki Meisei University, Japan

Ceria dioxide is of great interest due to favorable properties as an electronic material, such as high dielectric constant of 26, high chemical stability, transmission in visible and infrared regions, and high efficient ultra-violet absorption. Epitaxial growth of CeO2 layers on Si substrates has been studied making the best use of a close lattice parameter matching relation. Recently, we have found that orientation selective epitaxial (OSE) growth (of 100) and (110) oriented CeO2 layers on Si(100) is capable by controlling surface potential distribution in reactive magnetron sputtering. Non-polar CeO2(110) grows on Si(100) with usual non-modified surface potential, whereas polar CeO2(100) grows on Si(100) with adequately bent surface potential. We are studying two OSE control methods, one is substrate bias application[1] and the other is low energy electron beam irradiation[2,3]. The latter has the attractive possibility of spatially varied two dimensional control of OSE grown regions.4] This paper describes experimental results of electron beam induced OSE growth with patterned scanning of 90 eV electron beams. The size and position of the electron beam irradiation area were controlled using absorbed electron current image observation. RHEED and XRD analyses proved the realization of two dimensionally controlled OSE growth of CeO2(100) and (110) areas in electron beam irradiated and non-irradiated areas, respectively. Precise XRD peak profile measurements revealed that there are considerably wide transition regions in between the above two areas, which contain both orientation components. For the application to two dimensionally patterned
hybrid orientation technology, the transition region width should be reduced significantly. Our experiments clarifed that the width of the transition region reduces proportionally with the logarithm of underlying Si substrate resistivity, reflecting the surface spread of potential distribution. These results will lead to sophisticated microelectronics devices using hybrid orientation technology. This work was supported by KAKENHI (20560024). A part of this work were conducted at the AIST Nanotechnology Network Japan of the MEXT, Japan.


**TF-TuP2 Influence of Growth-Rate on the Epitaxial Orientation and the Crystalline Quality of CeO2 Thin Films Grown on Al2O3(0001) by Molecular Beam Epitaxy**

M. Nandasiri, Western Michigan University, P. Nachimuthu, T. Varga, V. Shothathanandan, W. Jiang, S. V.N.T. Kuchibhatla, S. Thevuthasan

Cerium oxide (CeO2) is one of the extensively studied rare earth oxides; however, it continues to attract attention because of its potential use in medical biology, catalysis, intermediate temperature solid oxide fuel cells (IT-SOFC), and resistive oxygen gas sensors. Driven by the need for fundamental understanding of its unique properties, CeO2 thin films grown on various substrates by different methods at different experimental conditions have been extensively studied. However, the influence of growth-rate on the orientation and the crystalline quality of the CeO2 thin films is relatively unexplored. While understanding the influence of growth-rate, we evaluated the ability to tailor the orientation and the epitaxial quality of CeO2 films on Al2O3(0001).

CeO2 thin films were grown on Al2O3(0001) substrates at 650 °C with different growth-rates (1-10 Å/min) by oxygen plasma assisted molecular beam epitaxy (OPA-MBE). The growth rate induced epitaxial orientations and crystalline quality of CeO2 thin films were studied by in-situ reflection high energy electron diffraction (RHEED), atomic force microscopy (AFM), and x-ray diffraction (XRD) techniques. CeO2 grows as three-dimensional (3-D) islands and two-dimensional (2-D) layers at growth-rates of 1-7 Å/min and ≥ 9 Å/min, respectively. AFM images show average surface roughness of 5-10 Å, indicating the high-quality surfaces of CeO2 thin films. The formation of epitaxial CeO2(100) and CeO2(111) thin films occurs at growth rates of 1 Å/min and ≥ 9 Å/min, respectively. Glancing incidence XRD measurements have indicated that the films grown at intermediate growth rates (2-7 Å/min) consist of some polycrystalline CeO2 along with CeO2(100). As indicated by x-ray pole figure measurements, the CeO2 thin film grown at 1 Å/min shows six in-plane domains, characteristic of well-aligned CeO2(100) crystallites. As the growth rate increases, two in-plane domains and eventually disappear. At growth rates ≥ 9 Å/min, CeO2(111) film with single in-plane domain was identified. The formation of CeO2(100) 3D-islands at growth rates of 1-7 Å/min is a kinetically-driven process unlike at growth rates ≥ 9 Å/min, which result in an energetically and thermodynamically more stable CeO2(111) surface.

**TF-TuP3 Influence of Target Type on Surface Texture-etched AZO Films Prepared by Magnetron Sputtering for Solar Cell Transparent Electrode Applications**

T. Minami, T. Miyata, T. Hirano, Y. Noguchi, J. Nomoto, Kanazawa Institute of Technology, Japan

This paper describes the influence of sintered oxide targets on the surface texture formation as well as light management obtainable by wet-chemically etching transparent conducting Al-doped ZnO (AZO) thin films prepared by different types of magnetron sputtering deposits (MSD). For transparent electrode applications in thin-film solar cells based on CdTe, GeSe and Si, transparent conducting AZO thin films require an appropriate surface texture to improve photovoltaic properties through increased light scattering and subsequent light trapping. It was found that the suitability of the light scattering characteristics for thin-film solar cell applications was considerably dependent on both the type of AZO target and the type of MS deposition used.

The surface texture-etched AZO thin films were prepared by a d.c.- or r.f. (13.56 MHz) power-supersimpered d.c.- magnetron sputter deposition (dc-MSD or rf+dc-MSD) with sintered AZO targets that was followed by a heat treatment with rapid thermal annealing (RTA) and a subsequent wet-chemical etch. Five types of commercially available high-density-sintered rectangular AZO targets were used. The suitability of the light scattering characteristics for thin-film solar cell applications was evaluated by carrying out surface textureing of the samples with wet-chemical etching in a 0.1% HCl solution conducted after the heat treatment with RTA (at a temperature of 300-500°C for 1-5 min in air). It was found that the obtained surface texture was sensitive to the underlying Si substrate, and the haze value near to near-sintered were considerably affected by not only the AZO target used but also the type of MSD; the dc- and rf+dc-MSD were conducted using the five types of targets with properties that depended on the supplier. The surface-etched AZO thin films prepared by rf+dc-MSD and etched to an appropriate depth after RTA were found to exhibit a higher haze value and a lower resistivity than the targets exhibited by dc-MSD. It was also found from X-ray diffraction analyses that the as-deposited AZO thin films prepared by rf+dc-MSD exhibited better crystallinity than those prepared by dc-MSD; also, the crystallinity of AZO thin films, such as the c-axis orientation, was slightly improved by the RTA treatment. A high haze value generally above 70% in the range from visible to near infrared (at wavelengths up to 1200 nm), which is suitable for thin-film solar cell applications, was obtained in the best surface-etched AZO thin films: film thickness of 2 μm prepared by rf+dc-MSD with an appropriate target and etched to a depth of approximately 300 nm after RTA at approximately 500°C for 5 min.

**TF-TuP4 XPS and ToFSIMS Characterization of Laser Modified Films for Li-Ion Battery Cathodes**

M. Bruns, R. Kohler, J. Proell, C. Zieber, W. Pfleig, Karlsruhe Institute of Technology, Germany

The development of novel lithium-ion-ion batteries has become subject of great interest in recent years due to the necessity of improved performance for future mobile applications and energy storage systems. Therefore, a great deal of effort has been spent to develop strategies for the enhancement of battery lifetime, capacity, and cyclability. A self-evident goal to improve the electrochemical properties of cathode materials is to enhance the lithium intercalation rate by increasing the surface to bulk ratio. For this purpose, a very promising approach is the laser-assisted surface microstructure and structuring of commonly as a cathode material used lithium cobalt oxide and lithium manganese oxide.

The present study focuses on the characterization of non-structured and laser modified films before and after electrochemical cycling using a combination of complementary surface analytical methods. For this purpose rf magnetron sputtered lithium cobalt oxide and lithium manganese oxide were structured with laser radiation leading to a surface microstructure with increased active surface area. In particular, high repetition excimer laser radiation with a pulse width of 4-6 ns and 248 nm wavelength was used. Self-organized conical microstructures were formed using laser fluences between 0.5 J/cm² and 2.0 J/cm². A subsequent annealing process using high power diode laser radiation at 940 nm wavelength enables the adjustment of the required crystallinity.

X-ray photoelectron spectroscopy (XPS) provides the chemical composition in an atomic depth dependent manner. Therefore, XPS spectra of the surface and XPS sputter depth profiles. In addition, the conical topography was shown by scanning electron microscopy (SEM) and crystallinity was proven by X-ray diffraction.

**TF-TuP5 Effect of Temperature on the Native Oxide Consumption during the ALD of Ta2O5 and TiO2 on GaAs (100) Surfaces**

Y. Te. T. Gopum, UMBC

Thermal ALD processes for the deposition of TiO2 and Ta2O5 have been developed using amide precursors and H2O as reagents. The TiO2 films were deposited from tetrakis dimethyl amido titanium and H2O and the TiO2 films from pentakis dimethyl amido tantalum and H2O. The growth rate for both processes was ~0.6 Å/cycle at 200°C and ~2.5 Å/cycle at 250°C respectively. These temperatures represent the so-called ALD minimum. At these temperatures the existence of a reaction that leads to the consumption of the native oxide surfaces during depositions on GaAs (100) surfaces has been confirmed using x-ray photoelectron spectroscopy and high resolution transmission electron microscopy. To investigate the effect of temperature on the native oxide consumption rate two sets of samples with thickness 2 and 3 nm have been prepared at temperatures ranging from 100°C to 400°C. For both ALD chemistries a significant enhancement in the gallium and arsenic oxide consumption rate was observed for process temperatures above 300°C. For depositions performed at temperatures more than 50°C below the ALD minimum the consumption reaction is significantly slower.
Al-doped ZnO (AZO) thin films that were prepared for transparent electrode applications in thin-film Si-based solar cells must necessarily attain not only a decrease of plasma resonance frequency by lowering the carrier concentration while retaining a resistivity on the order of 105 Ωcm, but also a significant scattering of light incident on the film by surface texturing. In this paper, we describe the influence of doped Al impurity content on the electrical properties as well as the light management observable by surface texture-etched AZO thin films that were prepared with various Al contents by a dc magnetron sputtering (dc-MS) deposition and wet-chemically etched after a rapid thermal annealing (RTA) treatment. In addition, the stability of the resulting electrical properties for practical use in various environments was investigated for these AZO films prepared with various Al contents.
Transparent conducting AZO thin films were prepared with a thickness up to 2μm by a dc-MS deposition using sintered AZO targets that had various Al contents. The basic sputter depositions were carried out on OA-10 glass substrates at a temperature of 200°C in a pure Ar gas atmosphere at a pressure of 0.4Pa and a power of 200W. Surface texturing of the AZO thin films was carried out by wet-chemical etching in a 0.1mol/L HCl solution after the thin films were heat treated with RTA at a temperature of 300-500°C for 1-5min in air. It was found that the obtainable surface texture and electrical properties in texture-etched AZO thin films were considerably affected by the RTA treatment conditions as well as the Al content doped into the films. The transmittance in the near-infrared region of the resulting thin films was increased. A high transmittance above 80% at a wavelength of 1.2μm was obtained in 1-μm-thick-AZO thin films prepared under the following conditions: an Al content (Al/(Al+Zn) atomic ratio) of approximately 0.5at.% when the thin films were heat treated without RTA and an Al content below 1.5at.% when the thin films were heat treated with RTA. In addition, when the thin films were heat treated with RTA at 500°C for 1-5min, increasing the Al content doped into the AZO films enhanced the transmittance in the near-infrared region, irrespective of the film thickness. However, it should be noted that the optimal Al content doped into texture-etched AZO thin films that would be suitable for transparent electrode applications in thin-film Si-based solar cells was considerably dependent on the obtainable crystallinity in the films as well as whether the thin films were heat treated with RTA or not.

TF-TuP9  Structure and Photo-Functional Properties of N⁺ Irradiated TiO₂ Thin Film under Various Substrate Temperatures. H. Shukur, M. Sato, Kogakuin University, Japan, I. Nakamura, Tokyo Metropolitan Industrial Technology Research Institute, Japan, T. Takano, Kogakuin University, Japan
Titanium dioxide (TiO₂) has been fundamentally used as a photocatalytic application to decompose environmental pollution materials. TiO₂ can generate active oxygen by exposing to sunlight and also in various environmental and the low cost material, so that it is expected to use as an element of a clean energy system in the future. TiO₂ shows relatively high reactivity and chemical stability under UV light whose energy exceeds the band gap of 3.2 eV for the anatase crystalline phase and 3.0 for the rutile crystalline phase. Many techniques have been examined to increase its activity under visible light region.
N⁺ ion irradiation method has been used by many researchers because it can provide an activated TiO₂ in both of visible and UV light. However the collision between ion and film causes a defect in film structure as causing a decline in photo-functional property.
In this study structural, chemical and photo-functional properties of TiO₂ thin film with a rutile structure were studied after irradiation by various doses of N⁺ ion beam under various temperatures. Reactive magnetron sputtering method was employed to prepare TiO₂ thin film on glass substrate (coming #1737). Ti-O was sputtered from Ti target in an Ar/O₂ pressure of 0.4Pa and a power of 200W. Surface texturing of the AZO thin films was carried out by wet-chemical etching in a 0.1mol/L HCl solution after the thin films were heat treated with RTA at a temperature of 300-500°C for 1-5min in air. It was found that the obtainable surface texture and electrical properties in texture-etched AZO thin films were considerably affected by the RTA treatment conditions as well as the Al content doped into the films. The transmittance in the near-infrared region of the resulting thin films was increased. A high transmittance above 80% at a wavelength of 1.2μm was obtained in 1-μm-thick-AZO thin films prepared under the following conditions: an Al content (Al/(Al+Zn) atomic ratio) of approximately 0.5at.% when the thin films were heat treated without RTA and an Al content below 1.5at.% when the thin films were heat treated with RTA. In addition, when the thin films were heat treated with RTA at 500°C for 1-5min, increasing the Al content doped into the AZO films enhanced the transmittance in the near-infrared region, irrespective of the film thickness. However, it should be noted that the optimal Al content doped into texture-etched AZO thin films that would be suitable for transparent electrode applications in thin-film Si-based solar cells was considerably dependent on the obtainable crystallinity in the films as well as whether the thin films were heat treated with RTA or not.

The increasing in the Full Wave Half Maximum (FWHM) of XRD measurements with N⁺ ion irradiation under 25°C refers to declination of the crystal structure of the TiO₂ thin film. On the other hand HRFW decreased by increasing the substrate temperature and closed to the value of the as-deposited film prepared under 300 °C. Thus the improvement of photocatalytic property under visible light was increased from 9.1% at 25°C to 25.7% at 300°C under the same N⁺ irradiation (2.5x10¹⁷ions/cm²).

TF-TuP11  Study of Electronic Structure in In-Ga-ZnO Amorphous Semiconductor Films. Y. Li, Z. Liu, X. Hu, J. Ren, Xian Jiaotong University, China
Amorphous oxide semiconductor In-Ga-ZnO (IGZO) has unique electron transport properties such as large electron mobility (10-30 cm²/Vs) and good uniformity for active-matrix flat panel display applications [1-2]. Theoretical studies indicate the high electron mobility in amorphous IGZO is due to the spatially dispersed orbits of metal cations forming unaffected electron transport path. It has been reported that high electron mobility is originated from the strong ionic electronic structure compared to the covalent electronic structure in amorphous Si [3]. However, the transport mechanism to result in high electron mobility in amorphous structure has not been experimentally explained yet. The focus of this study is to use surface probe, optical as well as Raman spectroscopic techniques to investigate the electronic structure in IGZO. Amorphous IGZO thin films are being grown at room temperature by pulsed laser deposition. Initial Hall effect measurements indicate that amorphous IGZO films have electron concentration of 9×10¹⁷ cm⁻³ with Hall mobility of 16.8 cm²/Vs. Raman spectroscopy is being used to analyze the lattice arrangement, i.e. the amorphous state and the bonding properties in IGZO films. During film processing, hydrogen is being introduced into the material system to modulate the defect state levels and band gap structure in IGZO. X-ray photoelectron spectroscopy is being used to obtain the valence band spectrum of IGZO films. The energy state within the band gap is being analyzed using deep level transient spectroscopy. The correlation of growth parameters on optical properties as well as electronic structure will be described in this work.

Vacuum Technology Division
Room: East Exhibit Hall - Session VT-TuP

Vacuum Technology Poster Session & Student Poster Competition

We would show that a quartz tuning-fork type resonator can measure the viscosity and the molar mass of the gas in which the resonator is vibrating. The vibrating resonator has two kind of outputs which are frequency change (Δf) and impedance change (ΔZ). The Δf and ΔZ are defined as the shift from their origin measured at high vacuum. We reported that the Δf and ΔZ given as a function of pressure are independent, and then the measurement of Δf and ΔZ will give the gaseous viscosity with no need to measure the gaseous pressure [1, 2]. In this report we will show that the molar mass of the measurement gas can be given by measurement of Δf, ΔZ and pressure. In this experiment the temperature controlled measurements were essential because Δf and ΔZ are sensitive to the temperature variation. The measurement apparatus, such as the vibrating sensor, driving circuit for oscillation, mass flow controllers, gas accumulator and the pressure gauges, are in a temperature controlled chamber by 29±0.02°C. The impedance of the resonator was evaluated by the current passing through the sensor over a temperature range of 120 kPa and vacuum. The absolute pressure was measured with a capacitance manometer.

The results showed that the ΔZ(P) and Δf(P), give as a function of pressure, are larger for higher pressure. The ΔZ(P) and Δf(P) for Ar, N₂, O₂ gases do not have the intersection. However Ne gas, having smaller molar mass but larger viscosity, crossed the other curves. So we cannot distinguish the gas species simply by ΔZ(P) or Δf(P) measurements. To discriminate the gas...
species with their viscosity the ΔZ-M plot is useful. We found that the characteristic curves of ΔZ-M lied in the descending order of the viscosity, i.e., Ne, Ar, O2, and N2. These curves do not cross each above 1 kPa.

We found the molar mass can be derived with the vibrating sensor. The product of molar mass and pressure can be evaluated without pressure measurement. The molar mass can be given with additional pressure measurement. The results showed that above 10 kPa of the gas pressure deviation of measured molar mass is less than a few percent.


Scroll pumps are widely used in solar-optic and semiconductor industry for backing purpose. The performance of scroll pump could affect significantly the performance of pumping station. The performance of a scroll pump was predicted by using VacTran commercial software and the experiment were conducted to verify the prediction in this study. Specifically, the delivered pumping speed, conductance and delivered throughput of the pump were investigated. The experimental delivered pumping speed increased as the inlet pressure increased and reached to 29.6 m³/min at 11.62 mbar. The characteristic curves of pumping speed at a reasonable cost for most of the residual gases except CH4 and rare gases, which amount to less than an hundredth of the total outgassing rate. Sputter-ion pumps remove all gases, though with a lower pumping speed. As a consequence, an optimized design should be based on NEG assisted by sputter-ion pumps for the gases that are not adsorbed chemically. Two examples of such configuration are here described. In the first, a commercial NEG lump pump is installed on a dedicated set-up together with sputter-ion pumps of different nominal pumping speeds. We show that the ultimate pressure achieved in the system does not depend on the applied sputter-ion pump nominal pumping speed in the range 30 to 400 l/s-1, and that values in the XHV range can be reached. In the second, we consider the vacuum system of the long straight section of the Large Hadron Collider (LHC) where most of the vacuum pipes were coated by magnetron sputtering with thin Ti-Zr-V films using Kr as discharge gas; the guidelines for the choice of the location and quantity of the sputter-ion pumps are reviewed in term of sectorization criteria and CH4 and Kr outgassing rates.


A specialized cryopump known as a cryogenic viscous compressor (CVC) is being developed for the ITER vacuum system to pump the regenerated, hydrogenic, fusion reaction gases from the torus cryopumps and neutral beam cryopumps, to the tritium exhaust processing facility. Several of these pumps will operate in parallel and are staged to maintain continuous pumping during plasma operation. The CVC’s regenerator at a higher pressure (500 mbar) than the torus and neutral beam cryopumps, which allows the regenerates gas to be pumped by a tritium compatible scroll pump train, with sufficient speed to maintain the regeneration duty cycle. The CVC’s are cooled to operating temperatures by precooling the inlet gas with a 80kH cooled chevron heat exchanger, followed by a tube bank heat exchanger cooled with supercritical helium at 4.5K. Hydrogenic gas is frozen on the inner tube bank walls while helium impurity gas, a byproduct of the fusion reactions, passes through the CVC and is pumped by conventional vacuum pumps.

A conceptual design of the CVC has been developed and a representative prototype has been designed, fabricated, and is undergoing testing to verify the concept of a full scale CVC before detailed design is completed. While cooling is provided by either cold helium gas or supercritical helium, hydrogen with trace amounts of helium gas is introduced into the central column of the cryopump at 100 Pa and 80 K at flow rates of 8 mg/s. Heat transfer between the laminar flowing gas and the cold pump tube is being enhanced with the use of internal finned fins. Temperature and pressure measurements are made along the pump gas stream in order to benchmark with design heat transfers. Comparison with fluid dynamics code is under way. Modeling of the gas flowing into the pump and through the precooler heat exchanger and freezing sections is accomplished with the CFX computational fluid dynamics code[1]. The flows into the pump are at low temperature (~1 mbar) and are in a laminar, low Reynolds number regime, (Re < 300) that is handled well with the CFX code. As the gas begins to desublimate in the cold zone of the pump, it reaches a rarified gas regime where the CFX model for flow and heat transfer breaks down. The modeling results are being compared with the prototype testing and will be used to further optimize and ensure reliable operation of the full CVC in the ITER application.


Autoresonant Ion Trap Mass Spectrometers (ART MS) have demonstrated significant benefits when applied to vacuum quality measurement at ultra-high vacuum (UHV) levels. Vacuum quality monitors based on ART MS technology are known to deliver more accurate gas analysis at UHV levels than the other competitive mass spectrometry technology presently used for residual gas analysis. The speed, accuracy and remote-sensing capabilities of ART MS technology for vacuum quality measurement at UHV levels will be explained and several application examples will be presented. The low outgassing rates associated to ART MS sensors will be justified and explained in terms of surface area and power dissipation considerations.

Gas analysis results, data-acquisition rates and detection limit values will be listed, and compared against similar results obtained with legacy instrumentation including quadrupole-based residual gas analyzers. Instrument optimization strategies for UHV applications will be disclosed.

VT-TuP6 Combination of NEG and Sputter-Ion Pumps for Particle Accelerator Vacuum Systems. P. Chiggiaio, J.M. Jimenez, S. Meinier, I. Wevers, CERN, Switzerland, A. Bonucci, A. Conte, P. Manini, SAES Getters

NEG and sputter-ion pumps are usually combined in particle accelerators to attain UHV pressure specifications. NEG pumps provide very high pressure speed at a reasonable cost for most of the residual gases except CH4 and rare gases, which amount to less than an hundredth of the total outgassing rate. Sputter-ion pumps remove all gases, though with a lower pumping speed. As a consequence, an optimized design should be based on NEG assisted by sputter-ion pumps for the gases that are not adsorbed chemically. Two examples of such configuration are here described. In the first, a commercial NEG lump pump is installed on a dedicated set-up together with sputter-ion pumps of different nominal pumping speeds. We show that the ultimate pressure achieved in the system does not depend on the applied sputter-ion pump nominal pumping speed in the range 30 to 400 l/s-1, and that values in the XHV range can be reached. In the second, we consider the vacuum system of the long straight section of the Large Hadron Collider (LHC) where most of the vacuum pipes were coated by magnetron sputtering with thin Ti-Zr-V films using Kr as discharge gas; the guidelines for the choice of the location and quantity of the sputter-ion pumps are reviewed in term of sectorization criteria and CH4 and Kr outgassing rates.


Over the years, Nb/Cu technology, despite its shortcomings due to the commonly used magnetron sputtering, has positioned itself as an alternative route for the future of superconducting structures used in accelerators. Recently, significant progress has been made in the development of energetic vacuum deposition techniques, showing promise for the production of thin films tailored for SRF applications. JLab is pursuing energetic condensation deposition via techniques such as Electron Cyclotron Resonance and High Power Impulse Magnetron Sputtering (HiPIMS). As part of this project, the influence of the deposition energy on the material and RF properties of the Nb thin film is investigated with the characterization of their surface, structure, superconducting properties and RF response. It has been shown that the film RRR can be tuned from single digits to values greater than 400. This paper presents results on surface impedance measurements correlated with surface and material
characterization for Nb films produced on various substrates, monocryristalline and polycrstalline as well as amorphous.

VT-TuP10 Bulk-like Nb Films might be Possible with Coaxial Energetic Deposition for Superconducting RF Cavities, T. Tajima, High Energy Accelerator Research Organization (KEK), Japan and LANL, N.F. Haberkorn, L. Civale, Los Alamos National Laboratory, E. Valderrama, M. Krishnan, Alameda Applied Sciences Corporation

B_{pen}, the magnetic field at which magnetic vortices start to penetrate into Nb films prepared by coaxial energetic deposition (CED) technique was measured with a SQUID magnetometer. Unlike the films prepared by conventional sputtering technique that showed B_{pen} ~ 94 mT at 2.5 K, the CED films showed B_{pen} of 180-190 mT at 2.5 K, a value that is very close to the number for bulk Nb used for SRF cavities. This corresponds to an accelerating gradient (E_{acc}) of approximately 45-48 MV/m for the SRF cavities with B_{pen}/E_{acc} ~ 4 mT/(MV/m) such as those for the European XFEL or the ILC projects. These samples were coated on MgO, Sapphire and Borosilicate with RRR ranging between 21(Borosilicate) and 540 (MgO). The next step will be to coat on copper. If it is possible to fabricate Nb coated copper cavities that have similar performance to bulk Nb high-gradient cavities, this will lead to a significant cost saving since the cost of copper is about 2 orders of magnitude less than Nb. It will also have other benefits such as better thermal stability due to high thermal conductivity of copper and less susceptibility to ambient magnetic field than bulk Nb cavities as has already been shown by LEP Nb/Cu cavities at CERN.
Vital phenomena are engendered through the dynamic activity of biological molecules. Therefore, observing the dynamic behavior of biological molecules in action at high spatiotemporal resolution is essential for elucidating the mechanism underlying the biological phenomena. The dynamic biomolecular processes are now widely studied using single-molecule fluorescence microscopy. However, the fluorophores labeled biological molecules themselves are invisible in the observations even using super-resolution fluorescence microscopy. The structure of biological molecules has been studied using x-ray crystallography, NMR, electron microscopy, and atomic force microscopy (AFM) but the obtained structures are essentially static. Thus, the simultaneous assessment of structure and dynamics is infeasible. To overcome this long-standing problem, we make it possible to simultaneously record the structure and dynamics of biological molecules, we have been developing high-speed AFM for more than 15 years and at last it is now coming of age. Various AFM devices and control techniques were optimized or invented for high-speed scanning and low-invasive imaging. As a result, the imaging rate now reaches 10-30 frames/s for the scan range 250 × 250 nm, 100 scan lines, and the spatial frequency of a sample surface corrugation 0.1 nm [Prog. Surf. Sci. 83, 337-437 (2008)]. Remarkably, even delicate protein-protein interactions are not disturbed by the tip-sample contact. With this capacity of high-speed AFM, some biological processes are successfully captured on video, such as walking myosin V molecules along actin filaments [Nature 468, 72-76 (2010)], photo-activated structural changes in bacteriorhodopsin [Nat. Nanotechnol. 5, 208-212 (2010)], and cooperative GroEL-GroES interactions. The high-resolution movies not only provide corroborative ‘visual evidence’ for previously speculated or demonstrated molecular behaviors but also reveal more detailed behaviors of the molecules, leading to a comprehensive understanding of how they operate. Thus, the high-speed AFM imaging of functioning biological molecules has the potential to transform the fields of structural biology and single-molecule biology.

The development of powerful single molecule functional imaging tools has been critical to our understanding of molecular dynamics and structure-function relationships in (bio)molecular systems. Our lab’s focus is the design, implementation, and application of coupled imaging and spectroscopy is providing intriguing insights into the mechanisms of membrane disruption, receptor oligomerization, and protein-membrane interactions. We have devised several correlative approaches based on the integration of in situ atomic force microscopy with fluorescence and vibrational spectroscopies for extracting the orientation, conformation, and association dynamics of membrane-associated proteins in model membranes and in live cells. Some of the key challenges and opportunities afforded these new tools will be discussed.

References:

We detected the characteristic visible light emission from fluorescent protein molecules deposited on metallic silver (Ag) upon injection of tunneling electrons generated by a standard scanning tunneling microscope (STM) in ambient condition. A series of fluorescent proteins originating from jellyfish or coral, nowadays engineered to generate various colors of fluorescence by gene technology, is characterized with a β-barrel structure insulating the chromophore electronically from the surrounding. We purchased green, yellow, red and infrared fluorescent proteins (GFP, YFP, RFP, HcRed, molecular diameter ~ 5 nm), deposited on a bare Ag surface, and used a Ag tip set on a STM setup to generate fluorescence. Light from the gap was collected by an optical fiber and introduced to a grating spectrometer with a liquid N2-cooled CCD detector. On bare Ag surfaces, visible light was detected with the STM bias voltage within ±1.8 V in a modestly moisturized N2 atmosphere. The spectra were unstable in general, indicating light emission upon excitation of local plasmon [1], which depends on the changeable geometry of Ag tip. The wavelength onset of emitted light was equivalent to the STM bias voltage within ±3.0 V, obeying the law of quantum energy conservation. The fluorescent proteins were dissolved in pure water, drop-cast on the Ag substrate and air-dried to form multilayers. STM images mostly showed flat terraces with steps composed of the protein molecules. Within a 200 nm x 200 nm scanning area, the light emission spectra apparently involved the characteristic fluorescence peaks of proteins (GFP = 540 nm (2.30 eV), YFP = 550 nm (2.25 eV), RFP = 650 nm (1.81 eV)) over a background of weakened Ag plasmon spectrum. The same experiment with Au tips and Au(111) substrates was with almost no detection for the characteristic fluorescence of all the proteins. For clean Au(111), although visible light was detected, the above-mentioned plasmon energy conservation stood for the bias voltage only within ±1.9 V. The maximum energy of local plasmon on Au(111) is too small to excite the fluorescent proteins electronically. The characteristic fluorescence from proteins is considered aided by the plasmon excitation of the Ag substrate. The protein β-barrel structure reserves the lifetime of excited state towards light emission, insulating electronically from the metallic substrate against the radiationless de-excitation process of the present surface-adsorbate system.

References:
10:40am AS+BI+NS-WeM9 Determination of Molecular Polarization at Protein-Electrode Interfaces with Combined Optical, Transport, and Dielectric Scanning Probe Microscopy. A. Chen, K. Kathar-Guillpeau, B.M. Discher, D.A. Bonnell, University of Pennsylvania

Bio-molecule integrated electronic devices are of great interest recently. For such systems to be designed and fabricated, the optoelectronic properties of protein molecules in ambient environment must be understood at a fundamental level. Here we demonstrate a new scanning probe based technique: torsional resonance nanoimpedence microscopy (TR-NIM), which simultaneously probes transport and dielectric properties in conjunction with optical excitation. To make a controlled interface, we start by designing a peptide molecule with ability to control protein/electrode interface interactions, as well as incorporation of several different optically active cofactors, and we successfully patterned peptides on HOPG substrates. Using TR-NIM electronic transport and the effect of optical absorption on dielectric polarizability in oriented peptide single or multiple molecular layers is determined. This approach enables quantitative comparisons of the change in polarization volume between the ground state and excited state in both single and multiple molecular layers.

11:00am AS+BI+NS-WeM10 Scanning Local Capacitance Measurements with High Spatial and Dielectric Resolution. M.J. Brukman, S. Nanayakkara, D.A. Bonnell, University of Pennsylvania

Spatial variation of dielectric properties often dictates the behavior of devices ranging from field effect transistors to memory devices to organic electronics, yet dielectric properties are rarely characterized locally. We present methods of analyzing second harmonic-based local capacitance measurements achieved through non-contact atomic force microscopy. Unlike contact-based methods, this technique preserves tip shape and allows the same probe to realize high-resolution topographic imaging and scanning surface potential imaging. We present an improved analysis of the electrical fields between tip and sample, yielding high sensitivity to the capacitance-induced frequency shift.

The techniques are applied to thin-film strontium titanate and mixed-phase self-assembled monolayers to illustrate application to high dielectric constant hard materials and lower dielectric constant organic films. Conversion from frequency shift signal to dielectric constant $\kappa$ is demonstrated on both samples, with sub-5 nm spatial resolution and dielectric constant resolution between 0.25 and 1.

11:20am AS+BI+NS-WeM11 Parallel Momentum Conservation of Hot Electrons across a Metal Semiconductor Interface. J.J. Garramone, J. Abel, R. Balsamo, V.P. LaBella, College of Nanoscale Science and Engineering, the University at Albany-SUNY

Parallel momentum of electrons is a conserved quantity as the electron traverses a barrier between two materials which lead to refraction like effects in the electrons trajectories. Ballistic electron emission microscopy (BEEM) is a scanning tunneling microscopy (STM) based technique that injects hot electrons (E>EF) into a metal-semiconductor Schottky diode. A small fraction of these electrons will traverse the metal with little to no scattering and make it into the semiconductor and counted as BEEM current. This makes it an ideal technique to study parallel momentum conservation. However, direct observation of this effect has been rather elusive. To observe this effect the dependence of the attenuation length with hot electron energy of Ag on both the Si(001) and Si(111) substrates has been measured.

Samples consisted of nanometer thick Ag films that were deposited on HF cleaned Si(001) and Si(111) wafers and capped with 10 nm Au to prevent oxidation of the films. Attenuation lengths were extracted by measuring the BEEM current as a function of the metal overlayer thickness. The dependence of the attenuation length with tip bias (electron energy) displayed a sharp increase as the energy approached the Schottky barrier height for the Si(001) substrates and a slight decrease for the Si(111) substrates. This contrast is a direct result of parallel momentum conservation and the lack of zero parallel momentum states at the Si(111) interface when compared to the Si(001) interface. Additional insight into the relative contribution of both elastic and inelastic scattering can be obtained by fitting the data to a Fermi liquid based model.

Biomaterials in Medicine and Biology

Biomaterials in Medicine and Biology

Room: 108 - Session BI-WeM

Cells at Interfaces

Moderator: M.R. Alexander, University of Nottingham, UK

8:00am BI-WeM1 Real Time Analysis of Polymer Film Integrity Upon Exposure to Bacteria and Aqueous Medium. D.E. Barlow, J.C. Biffinger, Naval Research Laboratory, E.R. Petersen, Nova Research, Inc., J.N. Russell, P.E. Pehrsson, Naval Research Laboratory, W.J. Goodson, Air Force Research Laboratory

Polymer coatings are of great importance for protecting and imparting specific properties at the surfaces of man-made structures, but can be affected in many ways by the natural environments they must withstand. We have studied the effects of aqueous medium exposure and biofilm formation on antistatic polyurethane coatings in real time using in situ ATR-FTIR. The results show that the coatings are susceptible to water permeation and swelling, and deuterium exchange was also shown to occur within the films upon D2O exposure. When exposed to Pseudomonas fluorescens in M9 minimal medium, the coating interface became compromised as the pyruvate carbon source was depleted. Reasons for these changes will be discussed, including the role of water permeation and the potential for the bacteria to use the coating as a carbon source. While ATR-FTIR has been used in the past to study biofilm growth, these results also demonstrate the effectiveness of the method for assessing substrate impact, an often overlooked factor.

8:20am BI-WeM2 Early Stages of Bacterial Biofilm Formation – A Numerical Study of Bioadhesion on Biomaterials. D. Siegismund, A. Schroeter, S. Schuster, M. Rettenmayr, Friedrich Schiller University Jena, Germany

Biomaterials for implant purposes are increasingly applied in modern medicine e.g. to recover human body functions or for tissue substitution in general. Infections of these implants, called Biomaterial-centered infections
The adhesion of bacteria is thus the first crucial step for biofilm formation that is only incompletely understood. Interactions of bacteria with the surface are controlled by surface properties such as surface energy, surface chemistry and topography.

In the present work, a model for bacterial adhesion is introduced that describes the early stages of biofilm formation as a function of the surface properties. A two-dimensional Cellular Automaton (CA) / Finite Difference (FD) simulation model is combined with the predictions of the extended DLVO (Derjaguin, Landau, Verwey, Overbeek) theory that accounts for the interaction energies between the bacteria and the material’s surface. The model describes the mass transport of bacteria in an aqueous solution towards the material’s surface and the adsorption and desorption process, depending on the surface properties.

The adhesion process of different human pathogenic bacteria (Enterococcus faecalis, Staphylococcus aureus, Escherichia coli) on different biomaterial surfaces (titanium, stainless steel, polyethylene, poly(methylmethacrylate), polytetrafluoroethylene) has been investigated. Results are the surface coverage with bacteria and, where applicable, clustering of the bacteria due to their migration on the surface.

Excellent agreement with experimental findings from the literature and own adhesion experiments concerning the kinetics of the adsorption process is found. In addition, a realistic bond strengthening mechanism of bacteria on surfaces, as described in the literature, is reproduced by the model. By using a spatial pattern analysis of our own experimental data we show that physical processes occurring during initial stages of the adhesion process are essentially correctly incorporated in the model.

This work is the foundation for future studies using human tumor biopsy samples that will help elucidate the link between fatty acid composition within a tumor and the potential drug resistance of that tumor.


Biofilm formation leads to a 1000 times increase in antibiotic tolerance compared with planktonic bacteria and is associated with 80% of hospital acquired infections, resulting in $3.0\text{ billion in excess health-care costs}$ each year in the U.S alone. Thus, new materials for biomedical devices that prevent biofilm formation would offer enormous benefits to the health industry and improve patient welfare. However, our current understanding of bacteria-material interactions limits scope for rational design of such materials. Polymer microarrays are emerging as a key enabling technology for the discovery of new biomaterials. A method for forming polymer microarrays has been developed using contact printing to deposit nanolitre volumes of premixed acrylate monomer and initiator to defined locations on a poly(HEMA) coated glass slide with UV photo-initiation. This platform enables a large combinatorial space to be rapidly screened by a biological system: MDA-231 and MCF-7 cells. There is similar work currently being done that uses Ultra Performance Liquid Chromatography-MS and gene sequencing to start characterizing lipid membrane metabolism in breast cancer tumor tissue. The separation of the two cell lines across PC1 can be clearly seen in Figure 1. The entirety of the loads for PC1 can be seen in Figure 2, with cholesterol strongly loading towards the MDA-231 cells and many diacylglycerol (DAG) species loading towards the MCF-7 cells. Key differences were found in the levels of certain lipid constituents of the cell membrane, which may play a role in the ability of one cell type to be more drug resistant than the other. There are a variety of lipid components that have similar trends which are not discussed in this abstract but may play an important role in understanding this system.

This work is the foundation for future studies using human tumor biopsy samples that will help elucidate the link between fatty acid composition within a tumor and the potential drug resistance of that tumor.

10:40am BI-WeM9 Engineering Stem Cell Differentiation via Material Properties, T. McDevitt, Georgia Institute of Technology

Stem cell differentiation is sensitive to a variety of global and local environmental cues that impact cell fate decisions. Pluripotent stem cells (i.e. ESCs & iPSCs) are capable of recapitulating many aspects of early development and can serve as a robust cell source for the development of cell-based diagnostics and regenerative medicine therapies. ESCs are commonly differentiated as three-dimensional multi-cellular aggregates referred to as “embryoid bodies” (EBs), because of their ability to mimic the early morphogenic transformation of pluripotent cells into derivatives of the three germ lineages (ecto-, endo-, and mesoderm). In order to better understand and ultimately control ESC morphogenesis, we have focused on systematically engineering biochemical and biophysical parameters of the 3D EB microenvironment via the integration of different biomaterials and examining the emergent results on stem cell differentiation. Microparticles (MPs) of varying size (1-20 \mu m) and chemistries (i.e. PLGA, agarose, gelatin) were incorporated within 3D stem cell aggregates in a dose-dependent manner (~1 particle / 10 cells) without adversely affecting cell viability. Interestingly, the mere presence of relatively small numbers of different types of materials alone could modulate stem cell phenotype as evidenced by gene expression profiling and immunophenotype analyses. Delivery of morphogenic factors, such as retinoic acid (RA), bone morphogenic protein 4 (BMP4) or vascular endothelial growth factor (VEGF), to ESCs from incorporated MPs significantly impacted the differentiation of the cells to different lineages more than comparable soluble treatment methods. Altogether these results suggest that engineered biomaterials can direct the differentiation of stem cells through modulation of biochemical and/or biophysical properties of the 3D microenvironment. It is expected that the development of inherently scalable techniques to directly pluripotent stem cell differentiation will benefit...
the biomanufacturing of stem cell derivatives for regenerative cellular therapies and in vitro cell based diagnostic technologies, as well as enable engineering of tissues directly from stem cells.

11:20am BI-WeM11 Adhesion and Rolling of Leukemic Cells on Immobilized Hyaluronans, A. Rosenhahn, Karlsruhe Institute of Technology, Germany, C. Christophis, I. Taubert, G.R. Meseck, A.D. Ho, M. Grunze, University of Heidelberg, Germany

Adhesion and rolling on vessel walls are two processes which are relevant for the homing of hematopoietic cells. Especially in the case of acute leukemia, one key in successful therapy is the homing of the hematopoietic stem cells (HSC) to the bone marrow after transplantation. We investigated the interaction of HSC with the hyaluron binding motive and quantitatively studied the interaction of different leukemic cells with synthetic polysaccharide surfaces. For the experiments we applied a microfluidic shear force assay recently developed in our group [1]. Leukemic Jurkat and Kasumi-1 cells lacking CD44-expression showed no adhesion or rolling on the polysaccharides whereas CD44 expressing leukemic cells KG-1a, HL-60, and K-562 attached and rolled on hyaluronan. We find that at weak flow cells have a poor tendency to adhere and only if shear forces above a threshold are present, adhesion is mediated. While this effect is well known for leukocytes on hyaluronan expressing feeder layers, it is the first demonstration that the mechanism also occurs in leukemic progenitor cells toward synthetic hyaluronan coated surfaces. We also extended the study to hematopoietic progenitor cells and saw for the first time that also HSC with high degree of stemness show a flow induced interaction with hyaluronan.


Electronic Materials and Processing Division Room: 210 - Session EM-WeM

Low-k Materials and Devices
Moderator: B. French, Intel Corporation

8:20am EM-WeM2 Feature-Scale Modeling of Diffusion Barrier and Metal Seed Physical Vapor Deposition Processes, R.A. Arakont, J.-P. Trelles, D. Kim, M. Khabibullin, S. Nikonor, D. Zierzath, Intel Corporation

The continuous reduction of critical dimensions and the increasing complexity of interconnect structures has stressed the process requirements of metallization steps (i.e., deposition of diffusion barrier and metal seed, electrical contact, etch, and chemical mechanical polishing). Particularly, physio the vapour deposition (PVD) of the barrier and seed relies on a limited number of process parameters (e.g., target and bias power, reactor pressure) to satisfy increasingly tighter film requirements (e.g., coverage over high aspect ratio features, control of barrier thickness, proper aperture of the seed to prevent void formation during electroplating). Computational modeling has proven an efficient means to aid the design and development of PVD metallization steps by allowing pre-simulation analysis of the effects of feature geometry and process parameters (e.g., see [1]). A continuum-based process simulator, based on a level-set solver for multi-material topography evolution, is applied to the analysis of 2 successive metallization steps, namely the PVD of barrier-over-ILD and of seed-over-barrier. Primary (e.g., distributed along the domain boundary) as well as secondary (e.g., emitted and reflected from surfaces) fluxes are accounted for through a ray-tracing technique that ensures mass conservation. In contrast to Monte-Carlo methods, which allow the description of gas phase kinetics, our solver is based on the specification of finite-rate surface reactions, which provides smooth topography evolution and is suitable for the analysis of extended domains encompassing multiple features, as needed for the analysis of within-die pattern effects. Sputtering, attachment, and neutralization reactions are accounted for between all the neutral and ionic species and the entire set of surface and bulk species in the film to allow the description of inter-material interactions (e.g., re-deposition of sputtered barrier species over the seed). The simulator is used to analyze the effects of neutral and ionic flux distributions (e.g., thermal neutrals, directional ions dependent on bias power) and the sputtering yield characteristics (i.e., energy and angular dependence) on the obtained film. Figure 1 presents snapshots of simulation results of the two-step process over a dual damascene structure. The results show that the simulation of consecutive steps is essential for the realistic description of inter-process effects.


8:40am EM-WeM3 Ultralow-k PECVD pSiCOH Dielectrics and their Implementation in VLSI Interconnects, A. Grill, S.M. Gates, IBM Research, E.T. Ryan, GlobalFoundries, S. Nguyen, IBM Research

The performance of integrated circuits and their density has been improved continuously through the shrinking of the active devices according to Moore’s law. At the 0.25 mm technology node it became clear that new materials had to be introduced to reduce the RC of the interconnect which became a barrier to further improvement of the VLSI performance.

After IBM introduced Cu in 1997, low-k PECVD SiCOH dielectrics with a dielectric constant k=3.0 were introduced in 2003-2004 at the 90 nm node, after many other low-k dielectrics failed integration. Reduction or even maintaining of the interconnect capacitance at the decreasing dimensions of later technology nodes required the development of porous ultralow-k pSiCOH. The first generation of pSiCOH dielectrics with k=2.4 has been successfully integrated by IBM in 45 nm products, such as the Power 7 chip released in 2010. Material extendibility of porogen based pSiCOH has been demonstrated to k values as low as 2.0. However, the reduction of the dielectric constant is achieved for a given chemistry by increasing porosity in the films, resulting in a decrease of mechanical properties, degraded integrability of the dielectric, and potentially reduced reliability of the interconnect.

The original chemistry produced the pSiCOH films having a skeleton of mainly O-Si-O bonds, with nanometer sized pores stabilized by Si-CH3 groups (V1 type). These films are damaged by the integration processes and the degree of damage increases with decreasing k to k values as low as 2.0. However, the reduction of the dielectric constant is achieved for a given chemistry by increasing porosity in the films, resulting in a decrease of mechanical properties, degraded integrability of the dielectric, and potentially reduced reliability of the interconnect.

The talk will discuss the evolution of the low-k and ultralow-k dielectrics, the effects of the chemistry on the properties of the different types of the porous pSiCOH, and the behavior of such dielectrics during integration processing.

This work was performed by the Research Alliance Teams at various IBM Research and Development Facilities.

9:20am EM-WeM5 Limitations in Dielectric Constant Scaling for low-k a-SiC(N):H Diffusion Barriers in Nanoelectronic Applications, S. King, D. Jacob, Intel Corporation, M. Liu, D.H. Gidley, University of Michigan

As the semiconductor industry strives to keep pace with Moore’s Law, new materials with extreme properties are increasingly being introduced and tighter control of these material properties is being demanded. Low dielectric constant (i.e. low-k) materials are one specific example. Lower k (< 6) a-SiC(N):H materials are desired to replace a-SiNx:H (< 6.5) as the Cu diffusion barrier layer in order to reduce resistance-capacitance (RC) delays in nano-electronic Cu interconnect structures. Typical methods for producing low-k a-SiC:N:H materials consist of introducing controlled levels of nano-porosity via carbon doping during plasma enhanced chemical vapor deposition (PECVD) of a-Si-nx:H matrix materials. While lowering k, the introduction of nano-porosity can seriously compromise the moisture and Cu diffusion barrier performance for these materials. In this presentation, we demonstrate that critical thresholds in nano-porosity exist for the diffusion of water through low-k materials. Specifically, we utilize Fourier Transform Infra-Red (FTIR) spectroscopy, to show that the concentration and size of nano-pores formed in low-k a-SiC(N):H dielectric materials is controlled by the concentration of terminal Si-CH3 bonding versus Si-C/N network bonding. We further combine moisture diffusivity measurements with x-ray reflectivity (XRR) and positron annihilation lifetime spectroscopy (PALS) to demonstrate that low-k a-SiC(N):H dielectrics become poor moisture diffusion barriers at mass densities < 2.0 g/cm3 and when the pore size approaches that for the molecular diameter of water. The implications of these critical nano-porosity thresholds on continued dielectric constant scaling of low-k a-SiC(N):H diffusion barrier materials will be discussed as well as methods for overcoming these limitations.

9:40am EM-WeM6 Fundamental Characterization of Amorphous Hydrogenated Boron Carbide Toward its use as a Low-k Dielectric Material, B.J. Nordell, S. Karki, C. Clayton, M.S. Driver, M.M. Paquette, A.N. Caruso, University of Missouri-Kansas City

The development of stable and low-dielectric-constant (i.e., low-k) materials for interlayer dielectrics (ILDs) in ultra-large-scale integrated circuits has become an essential target for the semiconductor industry. Toward this end, several low-Z boron-based materials (e.g., boron nitrides
and boron carbonitrides) have been studied due to their exceptional thermal, mechanical, and chemical stability, which have exhibited $k$ values as low as 1.9–2.4. Amorphous hydrogenated boron carbide, in turn, may be an even more promising low-$k$ boron-based material on the basis of the lower polarity of B–C bonds relative to B–N bonds and the mesosporous porous icosahedral cage structure of boron-rich carbides which can exhibit significantly lower free volume, particularly for the low-density amorphous hydrogenated variants. Moreover, amorphous hydrogenated boron carbide films grown by radio frequency plasma-enhanced chemical vapor deposition methods from carbonaceous precursors exhibit resistivity’s $>1 \times 10^6$ Ω-cm. This talk will describe the causal relationship between the measured dielectric constant, resistivity, breakdown voltage, hardness, Young’s Modulus, and mass-density as a function of the PECVD growth parameters (power, pressure, substrate temperature, and gas flow) and film composition (notably hydrogen and oxygen content) in the context of establishing and optimizing amorphous hydrogenated boron carbide as a next-generation durable and resilient low-$k$ ILD.


In the demanding microelectronics industry there is a constant need to increase circuit density in multilevel Copper (Cu) back-end-of-line (BEOL) interconnects to improve the operating speed and reduce power consumption. With successful node-to-node scaling there arises the capacitance-resistance (RC) requirements for the BEOL is through the introduction of organo silicate glass (SICOH) materials with low dielectric constants ($k$-value) as interlevel dielectrics (ILD). At the 45 nm node, porosity was first introduced into the BEOL interconnect structures in the form of porous organo silicate films (p-SiCOH) with ultra low-$k$ ($\leq 2.55$) to further minimize the RC delay. These ULK materials incorporate a large number of methyl groups and pores into Si-O based network structures, although the initial precursors and final properties may vary. Both the SICOH and pSICOH ILD films are commonly deposited by plasma-enhanced chemical vapor deposition (PECVD). Since their introduction nanoporous ULK films have significantly increased the Cu BEOL interconnects RC-dominance. For ULK films, pore size cannot be kept constant and tend to crack as a result of elastic mismatch with the substrate. Meanwhile, pore collapse and carbon depletion occur when ULK is exposed to RF-plasma during etching or ashing. Subsequent moisture adsorption leads to the increase of effective $k$-value in Cu interconnects degrading RC performance. Therefore, the integration challenges of ULK are significant, such as plasma damage, chip packaging interaction, and dielectric/metal barrier compatibility. To achieve the careful optimization of ULK properties is crucial for successful process integration in the 28 nm BEOL node and beyond. This presentation will report on the ability to tune the material properties of ULK films with $k \leq 2.55$ through the use of new chemical precursors and simple processing optimization steps to meet the specific integration requirements. Three ULK material classes were evaluated, optimized, and characterized to compare electrical and mechanical properties, pore characteristics, FTIR, XPS, and thermal stability. The effect of UV Cure dose on the ULK film properties and correlation of the degree of plasma damage to the ULK chemical, physical, and structural properties will be discussed. We will show that balancing composition of the film to minimize damage for successful integration needs to be coupled with improving electrical and mechanical integrity for packaging compatibility.

11:20am EM-WeM11 Molecular Strengthening Mechanisms for Low-$k$ Dielectrics, R.H. Dauksardt, Stanford University

INVITED

Hybrid organic-inorganic glass films processed from small organosilane precursors exhibit unique electro-optical properties while maintaining excellent thermal stability. Processed using either sol-gel or plasma-enhanced chemical vapor deposition they have application in emerging CMOS, nanoscience and energy technologies. A fundamental challenge for their integration, however, remains their inherently mechanically fragile nature. This derives from the oxide component of the hybrid network and the presence of terminal hydroxyl and organic groups that reduce network connectivity. Also, to achieve ultra-low dielectric properties (i.e $k<2.4$) nanoporous forms of the hybrid films are required which further reduce mechanical properties. We describe the development of computational methods to address the fundamental relationship between molecular structure and resulting mechanical and fracture properties of organosilicate glasses. Using molecular dynamics and a simulated annealing approach, large distortion-free hybrid glass networks with well-controlled network connectivity can be generated. With this capability along with a novel fracture model and molecular dynamics simulations of elastic deformation, we are demonstrating the critical effects of network connectivity and nanoporosity on mechanical properties. The accuracy of our computational tools is confirmed through comparison to synthesized hybrid films where the molecular structure, connectivity and nanoporosity is carefully controlled. Having predictive models for how molecular structure affects mechanical properties offers the opportunity for computational design of new glasses and processes to achieve desired property profiles for guiding precursor selection. Thus in addition to the fundamental insights gained regarding structure-mechanical property relationships, we will present our efforts to apply these tools to design new neat and nanoporous glasses with exceptional mechanical properties and low density.

Energy Frontiers Focus Topic

Room: 103 - Session EN+EM+NS-WeM

Quantum Dot and Nanowire Solar Cells

Moderator: K. Leschkiti, Applied Materials Inc.

8:00am EN+EM+NS-WeM1 Hybrid Quantum-Dot/Organic Solar Cells Based on Silicon Nanocrystals, U. Kortshagen, C.Y. Liu, Z. Holman, J. Yang, University of Minnesota

INVITED

Organic solar cells based on bulk heterojunctions between acceptor and donor semiconductors have attracted significant attention due to their low cost, compatibility with roll-to-roll processing, and relative lack of health and environmental concerns. However, the stability of organic semiconductors under solar irradiation remains to be a challenge. Producing bulk heterojunction solar cells based entirely on inorganic materials thus has become an attractive proposition. In a first step into this direction, we have produced solar cells from silicon nanocrystals (Si NCs) and poly-$3$-hexylthiophene (P3HT). Silicon NCs $3–5$ nm in diameter were synthesized by dissociating silane gas in a nonthermal radio frequency plasma, and collected on a mesh downstream of the plasma. The silicon-hydride terminated NCs were dispersed with P3HT in 1,2-dichlorobenzene. Films of this blend with thicknesses of $100–200$ nm were spin-cast onto an indium tin oxide (ITO) substrate pre-coated with $50$ nm of poly$(3,4$-ethylenedioxythiophene)/poly(styrene sulfonate) (PEDOT:PSS). Metal electrodes (2 mm wide, $100$ nm thick) were then evaporated on top of the Si NC/P3HT film. The Si NCs were found to be efficient electron acceptors. The optimal weight ratio of Si NC to P3HT was $50\%$. The effects of annealing and different metal electrodes on Si NC/P3HT hybrid solar cells were studied. After annealing at $150^\circ$C, Si NC/P3HT solar cells exhibited power conversion efficiencies as high as $1.47\%$. The hole mobility in the P3HT phase extracted from space-charge-limited current measurements of hole-only devices increased from $2.5 \times 10^{-10}$ cm$^2$/V-s to $1.1 \times 10^{-9}$ cm$^2$/V-s after annealing, resulting in better transport in the solar cells. A quenching of the open circuit voltage and short-circuit current was observed when high work function metals are deposited as the cathode on Si NC/P3HT hybrid devices.

Devices with silicon-hydride terminated Si NCs were plagued by poor film morphology. Hence we studied functionalizing Si NCs with organic ligands. Functionalization with 1-occtene and 1-dodecene led to improved film morphology but the transport in the Si NC network became worse. However, excellent stability with respect to air exposure of these cells was observed. Efforts to improve the dispersability of Si NCs without inhibiting transport in the Si NC network are in progress. This work was supported primarily by the MRSEC Program of the National Science Foundation under Award Number DMR-0819885 and by the DOE Energy Frontier Research Center for Advanced Solar Photophysics.

8:40am EN+EM+NS-WeM3 Precision Engineering of Semiconductor Nanowires for Advanced Photovoltaic Devices, N. Shin, I.R. Musin, S. Sivaram, M.A. Filler, Georgia Institute of Technology

INVITED

Semiconductor nanowires offer exciting opportunities to engineer light absorption and carrier transport for ultrahigh efficiency photovoltaic devices. The precise control of crystal structure and geometry is required to achieve a desired behavior, especially in highly confined nanoscale systems.

In the ideal situation, the combination of nanowire diameter, lattice structure (e.g. diamond cubic, wurtzite), crystal orientation (e.g. $<111>$ vs. $<100>$), and sidewall faceting that yields the most robust device performance would be known and could be rationally synthesized. Unfortunately, an inadequate understanding of nanowire chemistry-structure and structure-property relationships prevents the accomplishment of this task with bottom-up syntheses at the present time. This presentation
will provide an overview of our recent efforts to bridge this knowledge gap. In our research, in-situ infrared spectroscopy is combined with an ultrahigh vacuum growth environment to fundamentally correlate nanowire chemistry with photophysics, while circumventing the sample degradation that can obscure the intrinsic properties of nanoscale structures. Group IV nanowires and their heterostructures are a model system and are fabricated with the vapor-liquid-solid (VLS) growth technique. The critical influence of surface-bound species near the three-phase boundary and their impact on nanowire crystal structure will be discussed in detail. This fundamental knowledge opens a generic and highly tunable route to engineer multiple classes (e.g. group IV and III-V, etc.) of semiconductor nanowires, heterostructures, and superstructures for advanced photovoltaic device applications.

9:20am EN+EM+NS-WeM5 Electrophoretic Deposition of CdSe Nanocrystals in Quantum Dot Sensitized Solar Cells, N.J. Smith, Middle Tennessee State University
Electrophoretic deposition (EPD) of nanocrystal thin films from the solution phase has received increasing attention due to its simplicity and ability to rapidly create controlled thickness films. In this presentation, we will discuss the impact of solvent choice, deposition time, and electrode voltage on the properties of CdSe nanocrystal films deposited by EPD for solar cell applications.

While hexanes are a commonly used solvent for depositing CdSe nanocrystals, resulting film formation by EPD is often not reproducible for different batches of nanocrystals. In this work, we report that adding acetone to solutions of nanocrystals in hexanes enables the consistent deposition of CdSe nanocrystal thin films of controllable thickness. These films form within a few seconds and require comparatively low electrode voltages for the deposition process. The addition of acetone appears to mitigate the effects of impurities in the precursor chemicals, the presence of solvents left over from nanocrystal synthesis, and variable environmental conditions that may otherwise compromise the quality of the resulting EPD nanocrystal films.

Quantum dot sensitized solar cells have been fabricated using the EPD process with CdSe nanocrystals solvated in hexanes with acetone. By modifying the EPD parameters, both the density and thickness of the nanocrystals films deposited on titanium dioxide coated ITO slides were affected. We will discuss the relationship between the EPD parameters and the resulting I-V characteristics and efficiency of the CdSe nanocrystal sensitized solar cells. Preliminary studies suggest that proper choice of EPD parameters can lead to more than one order of magnitude improvement in the solar cell performance metrics.

9:40am EN+EM+NS-WeM6 Selective and Highly Efficient Photo-Induced Activity Over Nano-Scale Sites in Porous Silicon: Potential Application for Hybrid Organic-Silicon PV, M. Asscher, The Hebrew University of Jerusalem, Israel, G. Toker, A. Nahor, O. Berger, S. Yitzchaik, A. Sáur, Hebrew University, Israel
Photo-induced processes over solid surfaces are of great basic and technological interest with applications in e.g. photolithography, sensing, catalysis and photovoltaics. Photo activity within porous silicon (PSi) following UV (193-266nm) irradiation has been studied. Unusually efficient, non-thermal, morphology and wavelength dependent photo-induced desorption (PID) of Xe was recorded. It is a selective process, revealing more than 3 orders of magnitude enhancement within pores, at the vicinal region of nano-slip tips, over atoms bound to flat chemistry surfaces. Remarkably large cross sections up to pPSi= 2·10^{-15}cm² were recorded, significantly larger than any previously published photo-induced events on solid surfaces at this wavelength range. Long lived, photo-induced positive charges (holes) located at inner surface nano-tips is proposed to stabilize transient negative Xe adsorbate ions as the precursor for this new photo-desorption process. These results were utilized for the construction of hybrid conducting polymer-porous silicon photovoltaic cell. Proof of concept and preliminary results are discussed.

10:00am EN+EM+NS-WeM9 High-Performance, Low-Cost Nanopillar Array Photovoltaics, A. Javey, University of California Berkeley
Semiconductor nanowires (NWs) hold great promise for fabrication of high-performance, low-cost solar cells. These materials have been extensively studied, both computationally and experimentally. The key advantage of NW solar cells resides in the ability to grow single crystalline NWs non-epitaxially on support substrates. Additionally, by varying the NW geometry, the electrical, optical and mechanical properties of the solar cells can be controlled, providing opportunities unavailable to planar thin-film solar cells. For example, ultrafast laser ablation in shared work on bottom-up CdS/CdTe nanopillar (NPL) solar cells, optical engineering of NPL arrays, and top-down fabrication of InP solar cells. Bottom-up NPL solar cells were fabricated on anodized aluminum oxide (AAO) templates. Due to the 3-D geometry, the templated NPL architecture orthogonalizes the light absorption and carrier collection directions, enabling cells to be optimized even for poor quality materials. First-generation CdS/CdTe solar cells on low-cost Al foil exhibited 6% efficiency at 0.6V open circuit voltage, VOC=0.54V, and 21 mA/cm² short circuit current, Jsc, despite significant (>30%) optical reflection from the top contact. Additionally, detailed electronic simulation was used to examine the design trade-offs involved with the NPL architecture. We found that even for a poor quality CdTe absorber layer, through proper design, moderate efficiencies of ~15% could be achieved. Additionally, all the processes used to fabricate top-down solar cells were roll-to-roll compatible. The combination of experiment and modeling show the promise of this method for fabricating low-cost, high-performance solar cells.

The AAO templated growth enables a high degree of control over the geometry and dimensions of the NPL arrays. For single-diameter Ge NPL arrays embedded in AAO, increasing the Ge material filling ratio both increases the reflectance and decreases the transmittance. The absorbance of an array is shown to strongly increase for increased diameter NPL arrays. Thus, by fabricating arrays of dual-diameter Ge NPLs (~100 nm and ~400 nm, incident light was absorbed with a film only 2 µm thick. This behavior is caused by the smaller NPL diameter at the interface between the material and incident light, and the higher diameter base then absorbs the light. Finally, radial p-n junction solar cells are fabricated from InP NPLs generated from a top-down etching procedure. The main challenge with radial p-n junctions is the ability to form ultra-shallow and conformal junctions along the radial axis. By utilizing a sulfur monolayer doping scheme, shallow junctions of ~10 nm were achieved. Finally, the highly electrically active dopant concentrations (~10^{19} cm^{-3}) are achieved. The fabricated solar cell exhibited a power conversion efficiency of 8.1%, a Voc=0.54 V, and a Jsc=25 mA/cm².

In summary, we have presented both top-down and bottom-up approaches for fabricating NPL solar cells, from both crystalline starting materials as well as low-cost Al foil. The templated AAO growth method also enabled optical engineering of NPL arrays, allowing a single material to maximize absorption and minimize reflection. Finally, the use of highly electrically active dopant concentrations was shown to enable radial InP solar cells.

11:20am EN+EM+NS-WeM11 Extremely Thin Absorber Solar Cells Based on CdSe-Coated ZnO Nanowires, H. Majidi, T.P. Le, G.W. Guglietta, J.B. Baxter, Drexel University
Solar cells can provide an abundant, clean, and sustainable source of electricity, but high costs have limited their implementation. The use of sensitized nanowire architectures may enable both low-cost manufacturing and high efficiency by decoupling the functions of light harvesting and charge transport into different materials. Sensitized solar cells consist of a bicontinuous interpenetrating network of electron- and hole-transporting materials with an interfacial absorber layer. In this architecture, interfacial recombination is the dominant loss process, so controlling the interfacial chemistry during deposition is critical.

We report on the use of n-type ZnO nanowire arrays sensitized with thin CdSe coatings and covered with p-type CuSCN in extremely thin absorber (ETA) solar cells. Low temperature, solution deposition methods were used for each material, offering the potential for inexpensive and scalable nanomanufacturing. Nanowire arrays provide direct pathways for electron transport as well as sufficient surface area for sensitization. The electrodeposited CdSe coatings are nanocrystalline and conformal with well-defined thickness. CuSCN is electrodeposited into the pore volume between nanowires. Morphology and microstructure of CdSe and CuSCN depend sensitively on bath chemistry and deposition potential. By controlling nucleation and growth rates, conformal and void-free materials can be deposited. A combination of solar cell measurements and ex situ materials characterization for both planar thin film stacks and nanowire arrays have been used to direct the selection of optimal ETA cell architectures. For example, ultrafast transient absorption spectroscopy demonstrates that interfacial electron transfer from photoexcited CdSe to the ZnO is much faster than recombination (~3 ps vs ~50 ps) for thin coatings. Planar solar cells were used to identify the optimal coating thickness of ~70 nm for these materials. External quantum efficiency measurements show efficient sensitization throughout the visible region of the solar spectrum. However, interfacial recombination limits overall energy conversion efficiencies.

137 Wednesday, November 2, 2011
Carderock Division

Calcium doped zinc oxide (Zn_{1-x}Ca_{x}O) is used as electron acceptor in the preparation of solution-processed bulk heterojunction solar cells. In this work, the specific photovoltaic device structure consisted of an InP/ZnS core-shell QD functionalized ZnO nanowire array that is embedded into a poly(3-hexylthiophene) (P3HT) hole transport matrix. The QD sensitized ZnO nanowires were characterized by optical absorption, confocal Raman and photoluminescence spectroscopy, as well as high resolution and scanning transmission electron microscopy. In addition, interface between QDs and ZnO NWs were studied by Atom Probe Tomography. QDs with different absorption bands were concurrently functionalized onto the same nanowire arrays in order to broaden the final device absorption bandwidth. The ZnO nanowire matrix was then patterned using inductively coupled plasma etching. The performance of planarized, quantum dot functionalized devices was subsequently compared to as-deposited and non-functionalized devices and the effect of QDs on device efficiency is presented. These include the study of the charge transfer mechanism using terahertz time domain spectroscopy.

Energy Frontiers Focus Topic

Room: 209 - Session EN+NS-WeM

Organic Photovoltaics

Moderator: R.A. Quinlan, Naval Surface Warfare Center, Carderock Division

Conjugated polymer/metal oxide hybrid photovoltaic devices have received considerable attention in recent years due to their potential for scalable, low-cost manufacturing. To improve device efficiency, most research has focused on increasing the fill factor and short circuit current. In this article, cadmium doped zinc oxide (Zn_{x}Ca_{1-x}O) is used as electron acceptor in the hybrid poly(3-hexylthiophene) /Zn_{x}Ca_{1-x}O bilayer solar cells to modify the band offsets and increase the open circuit voltage. XRD results show the sol-gel prepared Ca doped ZnO films have a hexagonal wurtzite phase with the QD lattice parameter being slightly larger than that of the ZnO substrate. The XRD pattern was used to model the electronic type of the ZnO material, and the Fermi level was calculated using independently determined band gaps. The Cd doped ZnO QD doped ZnO nanoparticles into bulk heterojunction solar cells to optimize device efficiency.

Spatially Localized Photocurrent Variation in Organic Photovoltaic Devices

8:40am EN+NS-WeM1 Electronically Monodisperse Single-Walled Carbon Nanotube Thin Films as Transparent Conducting Anodes in Organic Photovoltaics.

T.P. Tyler, R.E. Brock, H.J. Karmel, T.J. Marks, M.C. Hersam, Northwestern University

Carbon nanomaterial thin films are being increasingly investigated for use as transparent electrodes in a variety of optoelectronic devices [1-3]. These flexible and solution-processable films are ideal candidates for organic electronics, including organic photovoltaics (OPVs), where low production costs and mechanical robustness are essential. While carbon nanotube networks have begun to be explored in such devices, the inherent polydispersity of the samples—namely the mix of both semiconducting and metallic species—has prevented the realization of their full potential in these devices and convoluted a complete understanding of their implications on device operation. Herein we report the use of single-walled carbon nanotubes (SWNTs) sorted by electronic type via ultracentrifugation as the transparent anode in poly(3-hexylthiophene) (P3HT) [6,6]-phenyl-C_{61} butyric acid methyl ester (PCBM) organic photovoltaic devices. Through a combination of dip coating and nitric acid treatment we achieve film roughnesses comparable to typical transparent oxides such as indium tin oxide. While carbon nanotube films are often electronically doped during processing, either intentionally or as a byproduct of roughness-reducing acid treatments, we find that the application of the quintessential electron-blocking interlayer poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT:PSS) removes sufficient adsorbed dopant groups to return the semiconducting nanotubes to their original state, vastly reducing their contribution toward current collection and transport. This is observed by both sheet resistance increases and UV-NIR spectroelectrochemistry. Furthermore, using X-ray photoelectron spectroscopy reveals that the weakly-bound nitric oxide groups on the surface are almost entirely removed by PEDOT:PSS. By varying the semiconducting and metallic content in the electrodes, we find that metallic content greater than 70 percent yields devices with efficiencies 50 times greater than those comprised of almost entirely semiconducting SWNTs. This observation is counterintuitive considering that freshly acid-treated semiconducting SWNT films often possess a lower sheet resistance than their metallic counterparts [4]. This underscores the advantage of using metallic-enriched populations for transparent conductors, where unstable doping of semiconductor carbon nanotubes complicates processability and ultimately reduces device performance.

Numerous reports have established that improving the performance of bulk heterojunction (BHJ) organic photovoltaic (OPV) devices requires not only the development of materials systems with improved spectral response and high charge carrier mobility but also, (ii) understanding the morphology of these systems. Approaches for morphological characterization have included microscopic techniques such as scanning electron microscopy, transmission electron microscopy, and various atomic force microscopy techniques. Other methods have provided indirect information about active layer morphology by enabling the determination of charge carrier lifetimes. These approaches have included transient absorption, reflectance, transient photocurrent, and nonlinear optical techniques among others. While these methods have proven quite informative, the relationships between nanoscale morphology, device performance, and the underlying electrical characteristics of functioning devices are not yet fully understood.

Recently, impedance analysis has begun to be applied to BHJ OPV devices. These reports demonstrate that the impedance analysis framework established for dye-sensitized solar cells can, to some extent, be extended to bulk heterojunction devices in order to calculate average charge carrier lifetime, electron densities-of-states, and charge carrier concentrations. However, a detailed analysis of the impedance response of active devices, and its dependence on device processing history, morphology, and operating conditions is still needed. In this work we acquire and analyze the impedance behavior of operating P3HT:PCBM bulk heterojunction devices and we examine the dependence of impedance on illumination conditions, active layer composition, and annealing history. We also report a simplified equivalent circuit model that successfully describes bulk heterojunction devices over a range of illumination conditions and applied biases. We use this model to extract relevant device performance characteristics such as average electron lifetime and find, in agreement with other reports, that bimolecular recombination losses play a significant role in these devices. To this end, we demonstrate a correlation between device efficiency and lifetime, and describe how values extracted from the present equivalent circuit model can be used to optimize device performance with new materials systems.

Singlet exciton fission is a process by which a high energy singlet spontaneously decays into two low energy triplets. It is an example of a multi-exciton generation process that could allow the efficiency of solar cells to reach beyond the Shockley-Queslier limit. In this talk we examine singlet exciton fission in solar cells based on tetracene and diphenyltetracene (DPT). Notably we find that the photoreactions in each cell have opposite dependencies on the application of a magnetic field, suggesting that fission occurs in tetracene-based solar cells, but decreases the photocurrent in DPT-based solar cells. We report an organic semiconductor solar cell using tetracene and CuPC (Copper Phthalocyanine) as donors, and C60 as the acceptor. Tetracene absorbs photons in the 450-550 nm region, generating high energy singlet excitons which split into two lower energy triplets, potentially doubling the photocurrent in this part of the spectrum. CuPC extends the absorption of the solar cell to the red part of the spectrum. We also demonstrate bulk heterojunction tetracene-C60 devices and tested them for singlet exciton fission.

We observe that: (i) The addition of the CuPC layer does not affect the flow of excitons from tetracene to the CuPC – C60 junction significantly because of similarities in triplet energies of tetracene and CuPC, (ii) The application of a 0.5T magnetic field shows ~1.5% change in photocurrent, confirming singlet fission, (iii) Low temperature quantum efficiency measurements show a drop in the tetracene IQE (internal quantum efficiency), and a singlet exciton yield of 72% at room temperature. The drop in photocurrent at low temperature is expected since singlet exciton fission in tetracene is a thermally activated process.(iv) Singlet exciton fission has a very high yield even in bulk heterostructure devices and can be potentially used to improve the performance of polymer solar cells.

In comparison, we observe that solar cells consisting of DPT-C60 exhibit a large positive (+5%) effect of the magnetic field on the photocurrent, +5% at ~4T. We explain the anomalous magnetic field effect and demonstrate that the display be increased beyond 100% by biasing the device close to open circuit, potentially leading to applications as an anisotropic magnetic field detector.
While we find that incomplete partial dipolar monomolecular films can control solar cell behavior for single, poly- and nano-crystalline cells, for current to pass through the molecules, we need dense monolayers. Alkyl chain monolayers help form near-ideal Metal-Insulator-Semiconductor (MIS) diodes, with significant PV activity. Surprisingly, though we can actually make *MIS* cells without a separate *I*(nsulator) layer, (MIS) diodes, with significant PV activity. Surprisingly, though we can actually make *MIS* cells without a separate *I*(nsulator) layer, suggesting that 'MIS' effects are at least partly more "chemical than is often thought.

In this way we demonstrate a near-ambient, simple, potentially low-cost approach to make and modify semiconductor solar cells, using a monolayer of molecules, as short as two carbons, that self-assembles onto the semiconductor (absorber) surface, passivating and buffering it. Good passivation is necessary to express the molecule-induced interface dipole, which can change the semiconductor electron affinity by up to 1 eV. Good, stable interface passivation along with strong inversion allows minority carriers, generated by absorbed light, to move laterally within the semiconductor top layer, for collection by a minimal-area grid, deposited on the conducting polymer and also minimizes photocurrent losses, due to sheet resistance. Thus, ≤ 1 nm thick organic molecules appear to convey a unique advantage over inorganic passivation or buffer layers.

11:40am EN+NS-WeM12 Correlation of Interfacial Electronic Structures and Open Circuit Voltages in Organic Solar Cells. M.H. Chen, National Dong Hwa University, Taiwan, Republic of China. W.H. Tseng, Y.Y. Wang, C.T. Tseng, C.I. Wu, National Taiwan University

In this study, the interfacial electronic structures and energy band diagrams of polymer/fullerene bulk heterojunction (BHJ) solar cells are investigated. To study the device performance influenced by the interlayer, poly(3-hexylthiophene) (P3HT) mixed with 6,6-phenyl C61-butyric acid methylester (PCBM) are used as an active layer and bathocuprione (BCP) and calcium (Ca) are incorporated respectively with the cathodes. Since the mechanisms leading to the improvement of power conversion efficiency are more chemically and electronically complicated and have not been understood, the ultraviolet and x-ray photoemission spectroscopy (UPS and XPS) are used to investigate the properties of energy band, electronic structures and the interface chemistry at the interface. For the devices incorporated with BCP, highest occupied molecular orbital (HOMO) level of PCBM is pulled down about 0.3 eV with respect to the Fermi level after the deposition of BCP. It indicates that the BCP layer can modify the energy band by shifting the energy level of acceptors, causing the increase of built-in potential (Vbi). Thus, the open circuit voltage (Voc) attributed to the increase of Vbi resulted from BCP could be enhanced. Moreover, AFM image indicates that BCP molecules cannot form a uniform layer on the active layer surface, pointing out the band modulation is the key reason to the Voc improvement. For the devices with Ca as cathodes, the UPS results with deposition of Ca illustrate a 0.8 eV-downward shift in energy levels of P3HT, but not in those of PCBM. Therefore, the energy difference between the HOMO of P3HT and the LUMO of PCBM is widened, resulting in the increase of Vbi and the enhancement of device efficiency. Moreover, from the XPS spectra, there is an extra new peak appear at the lower binding energy about 162 eV, which suggests that the addition of electrons around the Ca atom. Deposition of Ca suspend the P3HT molecules after Ca deposition. The combinations of S and Ca spectra imply that the formation of Ca-S bonding at the Ca/P3HT interface, which is in good agreement with the findings of UPS spectra.

Exhibitor Technology Spotlight Room: West Exhibit Hall - Session EW-WeM

Exhibitor Technology Spotlight
Moderator: D. Surman, Kratos Analytical Inc.

10:20am EW-WeM8 Using the Apple IPAD with your PVD System Maintenance. C. Malocsay, Semicore
Rarely does one item change several industries at once like the Apple IPAD or other tablet computers. Semicore has followed other major companies that have taken this device and applied it to increase productivity and reduce costs. Not a replacement for your PC but considered as an extension of your computer just like a wireless mouse. Applications available at little or no cost can increase the users effectiveness in Thin film vacuum deposition systems and their day to day operation.

Graphene and Related Materials Focus Topic
Room: 208 - Session GR+MN-WeM

Graphene: Mechanical and Thermal Properties, Graphene MEMS and NEMS
Moderator: J. Rabe, Humboldt University Berlin, Germany

8:00am GR+MN-WeM1 Graphene Atomic Membranes: From Patchwork Quilts to Atomic Drums, P.L. McEuen, Cornell University

Graphene is the world’s first atomic membrane, a robust, one-atom thick freestanding layer of sp2-bonded carbon. The physical properties of these membranes straddle the border between soft and hard condensed matter. They are strong but highly flexible, with bending stiffness comparable to a lipid bilayer but stretching stiffness similar to diamond. Milk-scale polycrystalline graphene films can now be produced cheaply and easily, opening the door to applications in both science and technology. In this talk we will present new results on the structural and physical properties of this remarkable 2D material, including the first STEM images of graphene grain boundaries and the patchwork-quilt-like structure of graphene grains. We also discuss experiments on atomic drums made with graphene membranes that can be actuated and probed either electrically or optically. Unlike for traditional MEMs and NEMS, we find that stiction and entropy are key parameters in determining the drum’s vibration frequency.

8:40am GR+MN-WeM3 Nanomechanics of Graphene: Non-Linear Response, Fracture, and Crack Propagation, R. Perriot, Y. Lin, University of South Florida, X. Gu, Aalto University School of Science and Technology, Finland, V.V. Zhakhovsky, I.I. Oleynik, University of South Florida

Recent nanoindentation experiments on graphene have revealed its exceptional strength, making it an excellent candidate for the design of nano devices such as MEMS and pressure sensors. Therefore, it is critical to understand the mechanical properties of graphene, and its response to a wide range of loading pressures beyond the elastic regime. In this work we performed molecular dynamics (MD) simulations of the nanoinindentation of graphene membranes by a spherical indenter. The indentation curves (load vs. indentation depth) obtained from simulations revealed two regimes of response: linear for smaller, and non-linear for larger indentation depths respectively. The MD results are in good agreement with the theory of elastic plates and recent experiments. Using the kinetic theory of fracture, we were able to determine the breaking strength of graphene and its dependence on the indenter radius. MD simulations also provided an atomic-scale description of the breaking process, which occurs through crack formation and propagation in graphene.

9:00am GR+MN-WeM4 The Effect of the Environment on Electrical and Mechanical Properties of Graphene, K. Balotin, Vanderbilt University

Every atom of graphene, a monolayer of graphite, belongs to the surface. Therefore, the environment of graphene -- the substrate onto which graphene is deposited and the coating on top of graphene -- intimately affects the properties of graphene. In this talk, we demonstrate that both mechanical and electrical properties of graphene can be tuned by varying the environment of graphene.

To study the mechanical properties of graphene, we developed a novel technique that is based on measuring the temperature-dependent deflection of a "bimetallic" cantilever composed of graphene and silicon nitride or gold layers. We demonstrate that the built-in strain, the substrate adhesion force and even the thermal expansion coefficient of graphene depend on the substrate under it.

To study the electrical properties of graphene in various environments, we developed a technique to fabricate electrically contacted multiterminal suspended graphene devices that are submerged into liquids. We explore the dependence of electron mobility of graphene on dielectric constant and ionic concentration of liquids surrounding graphene. We find that ions in liquids can cause strong scattering in graphene and that very large values of mobility (~40,000 cm²/Vs) can be achieved in ion-free liquids.

9:20am GR+MN-WeM5 Nanoscale Friction and Adhesion Behavior of Graphene: The Effect of Sliding History, X.-Z. Liu, Q. Li, B. Zhang, R.W. Carpick, University of Pennsylvania

As a prominent example of a two-dimensional (2-D) material, graphene has drawn much attention because of its extraordinary physical properties. However, in contrast to its electronic and thermal properties, the mechanical and tribological properties of graphene remained poorly understood. These properties are interesting scientifically because of the extremely high...
That both the growth mechanism and the relaxation along the mismatched magnitudes of the structural strain and its local variation are significant and graphene is tightly grafted to the underlying interface layer and SiC amount of local strain. The large compressive strain implies that monolayer graphene and SiC substrate, is probed using Raman scattering. We show graphene films on silicon-carbide (SiC) (0001). The strain, due to the same area for a sufficient distance. This sliding-history dependence was not observed on bulk graphite or SiO2 substrates, and thus appears to be yet another unique feature of the tribological behavior of atomic sheets. These two observations strongly suggest that the geometric structure of the sheet and the contact area it makes with the tip is significantly affected by the sliding history.


10:40am GR+MN-WeM9 Molecular Dynamics Simulations of Melting of Graphene. B. Steele, V.V. Zhukhovsky, R. Perriot, I.I. Oleynik, University of South Florida
It has long been predicted by Peierls, Landau, and Mermin that infinite two-dimensional solids are unstable and should melt at any finite temperature. The stability of graphene, a two-dimensional layer of carbon atoms, is thus explained by the presence of an additional degree of freedom that allows it to buckle in the third direction normal to the 2-D perfect plane. We performed large-scale molecular dynamics (MD) simulations of graphene melting in order to provide a fundamental insight into the stability of graphene, as well as to investigate the nature of the defects naturally created by thermal excitations at high temperatures. We will discuss the types of defects appearing before melting, the atomic-scale mechanisms of melting, and the nature of the resulting carbon melt. In addition, the graphene melting in 3D space and 2D constrained melting is also discussed. The importance of a proper description of chemical bonding in graphene at high temperatures is illustrated using different interatomic potentials: the reactive bond order (REBO) potential and the newly developed screened environment dependent SED-REBO. Comparisons with other recent simulations of graphene melting (including those using the LCBOP2 potential for carbon) are also presented.

11:00am GR+MN-WeM10 Inhomogeneous Strain in Monolayer Epitaxial Graphene. D.A. Schmidt, Ruhr-University Bochum, Germany, T. Ohba, L.B. Biedermann, T.E. Beechem, S.W. Howell, G.L. Kellogg, Sandia National Laboratories
We report a large in-plane compressive strain (up to 0.5%) and its inhomogeneous variation at micrometer scale in single layer graphene films on silicon-carbide (SiC) (0001). The strain, due to the difference in lattice constants and thermal expansion coefficients of graphene and SiC substrate, is probed using Raman scattering. We show that both the growth mechanism and the relaxation along the mismatched symmetry of the graphene and underlying substrate can affect the exact amount of local strain. The large compressive strain implies that monolayer graphene is tightly grafted to the underlying interface layer and SiC substrate; otherwise it would delaminate to relieve the strain. The magnitudes of the structural strain and its local variation are significant and need to be taken into account for electronics applications based on the graphene-SiC(0001) system.

The Raman microscope was supported under BMBF grants 05K57PC2. D. A. S acknowledges support within the BMBF funded projects 05K57PC2 and 05K10PC. The work was also supported in part by the LDRD program at SNL and the US DOE Office of Basic Energy Sciences’ Division of Materials Science and Engineering (Contract No. DE-AC04-94AL85000). Part of this work was performed at CINT (Contract No. DE-AC04-94AL85000). SNL is a multi-program laboratory managed and operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Corporation, for the U.S. Department of Energy’s National Nuclear Security Administration under contract DE-AC04-94AL85000.

Helm Ion Microscopy Focus Topic
Room: 106 - Session HI+AS+B1+NS-WeM
Nanono and Bio Imaging with Helion Ion Microscopy
Moderator: A. Gölzhäuser, University of Bielefeld, Germany, V.S. Smentkowski, GE-GRC
8:40am HI+AS+B1+NS-WeM3 Helion Ion Microscopy Techniques for Imaging and Characterization of nano-Device Materials and Structures. S. Ogawa, T. Iijima, National Institute of Advanced Industrial Science and Technology (AIST), Japan
INVITED
This paper presents several imaging modes for nano-devices fabrication that may make HIM a tool of particular value to soft materials such as low-k dielectrics (low-k) with less transformation and more materials contrast which reflects damaged areas, and copper interconnect buried in dielectrics, and shows luminescence induced by the focused helium ion beam using the HIM for the first time.
Imaging of -100 nm pitch patterned low-k is important for LSI Cu/low-k interconnect processes, while SEM imaging results in changes to the low-k line edge roughness and shape by damage during an electron beam irradiation. The HIM could provide low-k dielectric secondary electron (SE) image with nm order resolution, deeper focus depth, less transformation because of three order magnitude lower thermal energy transfer into a unit volume of the low-k than the SEM under an appropriate operation condition1.
During the imaging, even at very low helium ion current, surfaces of samples were atomically etched off, as in a graphene patterning, and then blistering or physicaletch occurred with the increase of the helium ion current. This makes the interpretation of the HIM SE imaging difficult but helpful. Damaged areas at side walls of the low-k regions in a 140 nm pitch interconnect were successfully seen with a different contrast from non-damaged low-k regions at an “optimized” helium ion beam condition2, which was similar to a TEM/Valence EELS result. On the other hand, using the SEM, the damaged areas contrast in the low-k regions could not be imaged.
A new imaging mode, through the inter-level dielectric, of the underlying copper, was explored. Cu interconnect was seen through a 130 nm thick low-k dielectrics. The incident helium ions might generate secondary electrons(SEEs) at the buried Cu surface and the SEEs of 1-2 eV energy passed through the dielectric of a few eV band gap without any energy transfer, and then the image was obtained. Helium ion channeling at the Cu surface area varied the secondary electron quantity, and it might generate a crystal orientation contrast of the buried Cu metal.
Luminescence induced by the focused helium ion beam was studied using the HIM3. Helium ion beam of a few PA current was irradiated to a SiO2 film, and peaks in a spectrum were observed at around 281, 447, and 672 nm; these positions were different from those by a conventional SEM cathode luminescence. The further study will be presented.
L. Stern, W. Thompson and J. Notte of Carl Zeiss are acknowledged for their discussions in the Cu/low-k works.
9:20am HI+AS+B1+NS-WeM5 He Ion Image the Au (111) Herringbone Reconstruction. V. Veligura, G. Hlawacek, R. van Gastel, H. Zandvliet, B. Poelsem, MESA+ Institute for Nanotechnology, University of Twente, Enschede, The Netherlands
The herringbone reconstruction of the Au(111) surface was directly visualized using an Ultra High Vacuum Helium Ion Microscope. Ion channeling phenomena arise from the different atomic ordering in the outermost layer of the crystal. First, we investigated the channeling contrast from the bulk Au fcc structure by imaging polycrystalline Au on glass films. The contrast that was observed as a function of crystal orientation was found to conform to what is calculated from a simple hard sphere model. Consequently, the herringbone reconstruction was investigated. It is a periodic zigzag structure of the three different types of crystal stacking (fcc, hcp and bridge sites connecting these regions) and, ideally, has a
Imaging with the Helium Ion Microscope employed exhibits a low energy spread (<1 eV), small virtual source size (<1 nm) [2], enabling tall structures to be viewed in focus within a single characteristic of HIM is its remarkable capability to neutralize charge by the incident 30 keV helium ions, contributing to the HIM’s primary advantage in the imaging of solid samples: its high spatial resolution (0.25 nm) [2]. We have applied this novel technology across a broad spectrum of multidisciplinary applications (from basic materials science and semiconductor applications to the biological sciences) to assess its utility and possible advantages over alternative techniques.

One area where our investigations have gained significant traction is in the imaging of biological specimens. The utility of this instrument in addressing topics of the biological sciences lies in part to the HIM’s high spatial resolution. However, in the context of biological specimens, it is the ability to image non-conductive samples without the application of a metal (or other conductive) overcoat and without the need of a background gas (both of which degrade resolution and surface detail), which has proven to be a distinguishing attribute. This opens up a whole new range of biological problems that can be solved rapidly and with less risk of artifacts.

An equally compelling application is in the field of nano-structuring. The focused helium ions have the ability to directly modify the sample surface under a high ion flux (via surface sputtering). This enables the direct patterning of structures and promises great utility in the fabrication of sub-10 nm devices. It also provides a mechanism for high resolution patterning on nonconventional substrates (such as suspended graphene membranes), where resist-based lithographic techniques are not feasible. Our experiences in sub-10 nm pattern transfer for both graphene and plasmonics applications will be presented.


11:40 am HI+AS+BI+NS-WeM12 Application of Helium Ion Microscope on Semiconductor Surface Imaging and Metrology. H.K. Guo, National Institute for Materials Science, Japan, H. Itoh, National Institute of Advanced Industrial Science and Technology (AIST), Japan, K. Onishi, T. Iwasaki, D. Fujita, National Institute for Materials Science, Japan Scanning electron microscope (SEM) has been used in the semiconductor surface imaging and metrology for more than 50 years. Now, a new tool, Helium ion microscope (HeIM), is developed and applied to this work. SEM and HeIM are the same in some fundamental characteristics. But, the latter has advantages in smaller probe size, higher resolution, and greater depth of field. These abilities enhance the performance of the HeIM in the semiconductor surface imaging and metrology, such as imaging of low-k materials [1] and measurement of critical dimension of the semiconductor devices [2].

A standard sample for scanning probe microscope tip characterization [3, 4] was measured by using HeIM and atomic force microscope (AFM) as shown in Fig. 1 and Fig. 2. Line profile of the HeIM image in Fig. 1 shows high accuracy in edge definition of the sample. The contrast of the image is related to morphology and materials of the sample [5], the probe size of the Helium ion beam, direction of the sample and beam, charge distribution, and so on. All the aspects will be analyzed in our presentation. The AFM image of the sample shown in Fig. 2 is a dilation of the real surface topography of the sample due to the finite-size AFM tip [6]. With an erosion algorithm, the surface of the sample was reconstructed to be compared with HeIM measurement.


Magnetic Interfaces and Nanostructures Division Room: 105 - Session MI-WeM

Fundamental Problems in Magnetism
Moderator: C. Clavero, College of William and Mary

8:00am MI-WeM1 Fundamental Problems in Magnetism. W.H. Butler, The University of Alabama

In this presentation, we shall attempt to describe the fundamental magnetic properties, the physics that controls and limits them and the practical implications of possible improvement. The most important fundamental properties of magnetic materials are Curie temperature, saturation magnetization, and magnetic anisotropy. The maximum Curie temperature (Co) and maximum saturation magnetization (Fe2Co13) at room temperature have not increased in a century. We shall discuss the reasons for this and speculate on the prospects that either may be increased. The magnetic anisotropy is somewhat less refractory. We shall discuss the prospects and implications of significant increases. Although higher room temperature saturation magnetization would be very useful, there would also be interesting applications for a magnetic material with very low
magnetization, but very strong spin dependence of its transport properties. There are also interesting potential applications for insulating ferromagnets.

8:40am  MI-WeM3  Progress toward Understanding the Sign of Spin-Polarization at Interfaces in Organic Spin-Valves, G.J. Szczechowski, University of Alabama

In this talk I will present results from a systematic study to understand the role of LiF, Al2O3, and MgO barrier layers in organic spin-valves. The overall aim of this work is to better quantify the degree of spin-polarized electron injection and extraction at ferromagnetic metal/organic semiconductor/metal interfaces. In general work with ferromagnetic transition metals has a positive magnetoresistance. However, when one of the ferromagnetic layers is exchanged with La0.7Sr0.3MnO3, the sign of the magnetoresistance is inverted. In addition the spin-polarized tunneling measurements of the structural, electronic, and magnetic properties of these interfaces have been thoroughly investigated by cross-sectional transmission electron microscopy, photoelectron spectroscopy, and polarized neutron reflectometry. These results will be compared to other findings in the literature in order to summarize the current status of spin-polarized electron transport across organic semiconductor/insulator/ferromagnetic metal interfaces.

9:00am  MI-WeM4  Rational Design of New Spintronics Materials: From Topological Insulators and Spin Torque Applications, C. Feher, Johannes Gutenberg University Mainz, Germany

INVITED

Heusler compounds are a remarkable class of intermetallic materials with a 1:1:1 (often called Half-Heusler) or 2:1:1 composition comprising more than 1500 members [1]. Today, more than a century after their discovery by Fritz Heusler, they are still a field of active research. New properties and potential fields of applications emerge constantly; the prediction of topological insulators is the most recent example [2]. Surprisingly, the properties of many Heusler compounds can easily be predicted by the valence electron count or within a different phenomena is fundamental in order to tune the magnetic properties of nanoparticles for precise applications, from data storage to biomedical research.


10:40am  MI-WeM9  Spin-Split Bands in Non-Magnetic Systems, E. Fescov, Brookhaven National Laboratory

INVITED

For fundamental and technological reasons materials with a spin-split electronic band structure in proximity of the Fermi level are highly attractive. The possibility of separately manipulating the two spin channels introduces novel functional behaviors without counterpart in the corresponding spin-degenerate systems. A promising approach in this field consists in the exploitation of the spin-orbit interaction, which couples spin and orbital degrees of freedom. The Rashba interaction offers particularly interesting perspectives: at the surface of a crystalline material the breaking of the full translational symmetry gives rise to an effective (Rashba) electric field which splits in k-space the valence electrons with opposite spin orientations. For heavy elements, such as Bi, the Rashba interaction results in spin-polarized surface bands, detectable in angle-resolved photoemission experiments [1]. In this talk various cases of such systems will be presented. Bi-Ag(110) surface alloy provides direct evidence of the interplay between the anisotropic electronic properties of the substrate and the Rashba field splitting of the Bi-Ag alloy and the ferromagnetic splitting of the Fe bands.


11:20am  MI-WeM11  Unoccupied Electron States in Rashba Systems Studied by Spin-Resolved Inverse Photoemission, M. Donath, S.N.P. Wissing, A. Zumbüble, C. Eibl, A.B. Schmidt, Muenster University, Germany

We present the first spin-resolved inverse-photoemission measurements of the unoccupied part of the Rashba-split surface state on Au(111). This Shockley-type state is considered as the prototype of a Rashba-split electron state on a metallic surface. The spin splitting of the occupied part of this state was first indicated by spin-integrated photoemission data [1]. This pioneering work was followed by a spin-resolved study, which directly proved the spin structure of the state [2].

Our study complements the information on the spin character by following the surface state into the unoccupied energy region. The state crosses the Fermi energy as a function of the wave vector parallel to the surface and finally leaves the bulk-band energy gap. Our spin-resolved inverse-photoemission experiment stands out from conventional systems thanks to an improved energy and k resolution [3]. Our data confirm the spin character of the surface state, as far as it does not overlap with bulk states.
In addition, we show how the spin character is altered when the surface state becomes degenerate with bulk states.

Further Rashba systems with even larger spin splittings as well as topological insulators are currently investigated with our spin-resolved inverse-photoemission apparatus. We will provide a status report on our latest results.


11:40am MI-WeM12 MBE Growth of Topological Insulator Bi2Se3 on Epitaxial Graphene on 6H-SiC(0001), Y. Liu*, M. Weisert, L. Li, University of Wisconsin-Milwaukee

In this work, we report results on the MBE growth of Bi2Se3, a prototypical topological insulator, on epitaxial graphene on 6H-SiC(0001). In situ scanning tunneling microscopy indicates spiral growth, characterized by atomically smooth terraces 10 to 50 nm in width, separated by steps of 1-2 quintuple-layers in height. X-ray diffraction shows only the (003) family of diffraction peaks with a full width at half maximum of 0.1 degree. Raman spectroscopy reveals two characteristic peaks at 130.21 and 171.48 cm⁻¹, corresponding to the in-plane Eg2 and out-of-plane A1g2 vibrational modes, respectively. The close resemblance of the positions and line shapes of both these peaks to those of bulk Bi2Se3 attest to the very high quality of the film. These results and their impact on the properties of the topologically protected surface states of the Bi2Se3/graphene heterostructure will be presented at the meeting.

*Ying Liu applied for postdoctoral fellow award

Nanomaterials and Nanomanufacturing Science and Technology Focus Topic Room: 111 - Session NM+AS+MS-WeM Nanomanufacturing Issues: Metrology and Environmental Concerns Moderator: J. Johnson, University of Tennessee Space Institute, W. Collins, Fisk University

9:00am NM+AS+MS-WeM4 Particle Characterization Issues in Evaluating the Toxicity and Environmental Impact of Manufactured Nanomaterials, K.W. Powers, University of Florida

Nanomaterials and nanomaterials promise to revolutionize many key areas of science and technology, however, the environmental effects of nanomaterial enabled products need to be considered throughout their lifecycle, from manufacture to environmental disposal. As nanomaterials become more commonplace in commercial applications, there is a need to assess the potential health and safety effects on human and other biological organisms. Materials at the nanoscale often possess properties that are different from the equivalent bulk or molecular scale. It is clearly shortsighted to assume that toxicological profiles of nanomaterials are the same as in the bulk or molecular forms. As they address these issues, toxicologists often need assistance in understanding and accommodating many of the unique attributes of nanoscale materials as they begin to assess potential health and environmental effects. Though the interpretation of the biological markers of toxicity are well developed, there are a number of issues relating to dosage, size, shape, detection and characterization that are problematic. There is a growing consensus that the complexity of these issues requires a multidisciplinary approach to nanoparticle toxicology that includes medical personnel, environmental and physical scientists as well as engineers trained in particle technology.

Keywords: nanoparticles, nanocharacterization, nanotoxicology, toxicity,

9:40am NM+AS+MS-WeM6 Sampling for Airborne Nanoparticles and Selecting Respiratory Protection, S.M. Hays, Gobbell Hays Partners, Inc., J.R. Millette, MVA Scientific Consultants

As the manufacture and use of nanomaterials continue to increase, appropriate questions are raised about the release of airborne nanoparticles into the general environment and specifically into the breathing zone of people. The development of monitoring procedures specific to carbon nanotubes and other nanomaterials is crucial in determining the effectiveness of engineering controls and personal protection. This presentation will review experiments conducted to determine the efficacy of using standard air sampling methodology for sample collection with standard membrane filter cassettes and analysis using transmission electron microscopy (TEM).

Tests done to evaluate the use of cartridge style respirator filters in carbon nanotube aerosols will also be presented. These laboratory tests will be discussed in relation to air samples collected in a variety of actual field situations. Proposed methodologies for the analysis for nanotubes in settled dust will be presented. Finally, one author is chairing an ASTM committee that is developing a consensus method to collect and analyze airborne nanotubes. The current state of that committee’s work will be summarized.

10:40am NM+AS+MS-WeM9 Local Probes Enabling Science and Manufacturing, D.A. Bonnell, University of Pennsylvania INVITED

The last decade has witnessed significant advances in measuring nanoscale phenomena. These advances have enabled scientific discoveries and provided a framework to support some nanomanufacturing processes. Nevertheless, both scientific advance and to a greater extent manufacturing are limited by our current capabilities in nanoscale metrology. This talk will highlight some of the exciting advances in probe based metrology, project future developments and outline the challenges that are critical to realizing a robust nanomanufacturing sector. The outcome of a recent global assessment of Nano Metrology will also be summarized.


Optical measurement of film thickness requires knowledge of the complex refractive index (dielectric function) of each material in the film stack. Practical experience has shown that the dielectric function changes with film thickness for many poly crystalline metal films and single crystal semiconductor layers. (1, 2) Previous studies pointed to quantum confinement induced changes in the dielectric function of thin silicon nanofilms between 10 nm and 2 nm. Extra Thin silicon on silicon (ETSO) films were used for this study. These films are often referred to as crystalline silicon quantum wells (c-Si QW). Our most recent study shows that the dielectric function of c-Si QWs can be further altered by the presence of a dielectric layer above the nano silicon top layer. (3) Based on an elastic theory description of the acoustic phonon modes, the dielectric function of the c-Si QWs is found to be strongly influenced by electron–phonon scattering. We illustrate this point using low temperature measurements of the dielectric function of a series of c-Si QWs and by comparing room temperature measurements of the dielectric function of 5 nm c-Si QWs with native oxide, 10 nm SiO2, and 10 nm HfO2.

3. Evidence of phonon confinement effects on the direct gap transitions of nanoscale Si films, V.K. Kamineni and A.C. Diebold, submitted

Nanometer-scale Science and Technology Division Room: 203 - Session NS-WeM Carbon-Based Nanomaterials Moderator: M.C. Hersam, Northwestern University

8:00am NS-WeM1 Characterization of Large Area Graphene Crystallites Grown on Cu Foil Substrates, P. Tyagi, Z.R. Robinson, H. Geisler, C.A. Ventrice, Jr., University at Albany, H. Yang, T. Valla, Brookhaven National Laboratory, Y. Hua, R.S. Ruoff, University of Texas at Austin

Graphene growth on Cu foils by catalytic decomposition of methane forms predominantly single layer graphene films due to the low solubility of carbon in Cu. One of the key issues for the use of CVD graphene in device applications is the influence of defects on the transport properties of the graphene. In particular, the presence of grain boundaries within the graphene film will increase the probability for scattering of carriers, resulting in reduced mobilities. Therefore, an important goal is to develop techniques for growing graphene films with crystallites that have lateral dimensions of a few millimeters or larger. There are several factors that influence the size and orientation of the graphene crystallites such as the size and orientation of the grains within the metal foil, temperature

* Postdoc Award Finalist
gradients during growth, the hydrocarbon source pressure, and the growth temperature. By growing the graphene films using methane source pressures less than 50 mtorr, preanneal times of approximately an hour, growth temperatures of 1035 °C, and a tented Cu substrate geometry within a conventional tube furnace, graphene crystallites larger than a millimeter in size have been achieved.

Measurements of the graphene growth morphology and surface topography of the Cu substrate have been performed using scanning electron microscopy (SEM). The graphene crystallites show a dendrite pattern, and the Cu substrate typically shows a somewhat faceted structure at this growth temperature. Low energy electron diffraction (LEED) measurements show sharp diffraction spots but with multiple zero-order reflections, which results from the faceted structure of the Cu substrate after growth. Electron backscatter diffraction (EBSD) measurements have been performed on the Cu substrates to determine the crystallographic orientation and size of the substrate grains. Before growth, the average grain size is ~10 μm with a random orientation. After growth, the Cu substrate grain size is on the order of centimeters with a typical orientation towards the {100} surface termination.

Synchrontron-based angle-resolved ultraviolet photoelectron spectroscopy (ARUPS) measurements have also been performed to probe the electronic band structure of the graphene. A linear dispersion has been measured in the K direction with the Dirac point located near the Fermi level. Synchrotron-based angle-resolved ultraviolet photoelectron spectroscopy (ARUPS) measurements have also been performed to probe the electronic band structure of the graphene. A linear dispersion has been measured in the K direction with the Dirac point located near the Fermi level.

8:20am NS-WeM2 Gas Adsorption on Pt-Clusters Supported by Graphene, J. Knudsen, Lund University, Sweden, T. Gerber, University of Cologne, Germany, E. Graanäs, Lund University, Sweden, P.J. Feibelman, Sandia National Laboratories, K. Schulte, Lund University, Sweden, P. Stratman, C. Busse, T. Michely, University of Cologne, Germany, J.N. Andersen, Lund University, Sweden

Model systems of real catalysts consisting of nanoparticles deposited on substrates with a predetermined size distribution, making it difficult to link the adsorption properties to the atomic scale structure of the nanoparticles using averaging techniques. Metal nanoparticles grown on a graphene/ir(111) moiré film, however, show exceptionally well ordered arrays of nanoparticles with an extremely narrow size distribution [1, 2]. Further, it is possible to control the cluster size precisely by adjusting the amount of deposited material, since each moiré unit cell contains one cluster. The narrow size distribution and the easy control of cluster size make metal particles supported by graphene an ideal model system for adsorption studies with averaging techniques.

In this contribution we report on our studies on CO adsorption on such an ideal model system consisting of Pt-clusters grown on a graphene/ir(111) moiré film using photoemission X-ray spectroscopy (XPS), scanning tunnelling microscopy (STM), and density functional theory (DFT) [3]. For Pt/graphene without CO we observe pinning of the graphene film, as a shoulder at the high binding energy side of the C 1s peak observed for pristine graphene. DFT calculations reveal that this shoulder should be assigned to carbon atoms positioned below and in the vicinity of the Pt clusters, which all are displaced towards the ir(111) surface.

Upon CO adsorption we observe C 1s and the O 1s peak positions to be consistent with preferential adsorption in atop sites at the cluster step edges. When CO adsorbs on the clusters step edges. From real time STM movies taken during CO dosing we show that the unpinning of the graphene film leads to coalescences of the Pt clusters, when the clusters are smaller than approximately 10 atoms.

References:


The development of high-performance graphene-based nanoelectronics requires the integration of ultrafast and pinhole-free high-k dielectric films with graphene at the wafer scale. Here, we demonstrate that self-assembled monolayers of perylene-3,4,9,10-tetracarboxylic dihydride (PTCDCA) act as effective organic seeding layers for atomic layer deposition (ALD) of HfO2 and Al2O3 on epitaxial graphene on SiC(0001). The PTCDCA is deposited via sublimation in ultra-high vacuum and shown to be highly ordered with low defect density within the organic monolayer. The electron scanning tunneling microscopy. Whereas identical ALD conditions lead to incomplete and rough dielectric deposition on bare graphene, the chemical functionality provided by the PTCDCA seeding layer yields highly uniform and conformal films. The morphology and chemistry of the dielectric films are characterized by atomic force microscopy, ellipsometry, cross-sectional scanning electron microscopy, and X-ray photoelectron spectroscopy, while high-resolution X-ray reflectivity measurements indicate that the underlying graphene remains intact following ALD. Using the PTCDCA seeding layer, metal-oxide-graphene capacitors fabricated with a 3 nm Al2O3 and 10 nm HfO2 dielectric stack show high capacitance values of ~700 nF/cm2 and low leakage currents of ~5 x 10−9 A/cm2 at 1 V applied bias. These results demonstrate the viability of organically self-assembled monolayers as seeding layers for high-k dielectric films in graphene-based nanoelectronics.

9:00am NS-WeM4 Graphene-based Electronics and Optoelectronics, Ph. Avouris†, IBM T.J. Watson Research Center

Graphene a two-dimensional, single atomic layer material with linear electron dispersion has rather unique electrical and properties. There is currently strong interest in taking advantage of these properties for technological applications. In my talk I will review some of the key properties of graphene, how these are affected by environmental interactions and how they can be utilized in electronics and optoelectronics.

Specifically, I will discuss high frequency (>100 GHz) graphene transistors, their fabrication and operation, as well as related device physics aspects, such as carrier transport mechanisms, electrical contacts, temperature effects, energy dissipation, etc. Simple integrated graphene circuits will also be presented. I will then discuss key optical properties of graphene and how they can be combined with its excellent electrical properties and used in optoelectronics applications. Specific examples involving ultrafast graphene photodetectors and their applications in the detection of optical data streams will be presented.


Graphene has attracted a widespread of interest because of its unique structural and electronic properties however, manipulation of these properties is necessary before realizing its full potential as the next generation material in a broad range of electronic and sensing applications. Specifically, tailoring the surface chemistry of graphene by the addition of functional groups is an attractive way to simultaneously manage the conductivity and reactivity of this material. This work discusses the use of electron-beam generated plasmas to controllably functionalize graphene synthesized from different methods. Electron-beam generated plasmas, produced in a variety of background gases (e.g. N2, O2, SF6, NH3) were used to introduce functional groups in a range of atomic densities at the graphene surface, without damage to the underlying graphene structure. Plasma processing conditions and characteristics, as well as the resulting chemical, structural, and electrical properties of the functionalized graphene were examined. This work is supported by the Office of Naval Research.

10:40am NS-WeM9 Solution Plasma-Assisted Surface Functionalization of Chemically Converted Graphene Sheet toward an Enhancement of Solubility in Solution, N. Tsuda, T. Ueno, N. Zettsu, S. Cho, O. Takai, N. Saito, Nagoya University, Japan

Graphene, a one-atom layer of graphite, possesses a unique two-dimensional structure and excellent mechanical, thermal, and electrical properties. Thus, it has been regarded as an important component for making various functional composite materials. Graphene can be prepared through micromechnical exfoliation, epitaxial growth, and chemical vapor
deposition and electrochemical approaches. Different from these approaches, chemical synthesis of graphene using graphite, graphite oxide (GO), and other graphite derivatives as starting materials were tested to be effective for producing chemically converted graphene (CCG) from various precursors, such as graphite, carbon nanotubes, and polymers, in large scale and at low costs. Therefore, CCG is more suitable for synthesizing high-performance graphene based composites.

Graphene oxide can be chemically reduced to CCG. Hydrazine monohydrate was most widely used, mainly due to its strong reduction activity and the stability in aqueous media. Upon reduction with hydrazine, most oxygen-containing functional groups of graphene oxide are eliminated and the π-electron conjugation within the aromatic system of graphite is partially restored. As a result, the reduced graphene oxide (or CCG) is usually precipitated from the reaction medium because of the recovered graphite domains of CCG sheets increased their hydrophobic property and π-stacking interaction. The use of hydrazine as reducing agent also has several disadvantages. The trace residual may strongly decrease the performance of CCG in devices.

In this work, we demonstrate solution plasma-assisted surface functionalization of chemically converted graphene sheet in order to enhancement of solubility in both aqueous and organic solution. Solution plasma (SPP) is a plasma discharge in solution, which is expected a higher reaction rate under low-temperature conditions, and the greater chemical reaction versatility since the molecular density of liquid is much higher than that of gas phase.

A colloidal graphene oxide sheets was treated with SP in ammonium containing aqueous solution in order to make reduced CCG and functionalize CCG surface with primary amine group. A glow discharge was produced at bipolar-pulsed voltages with pulse width and frequency of 2 ms and 15 kHz, respectively. The all products were characterized by IR, Raman, spectroscopy, AFM, XRD, and TEM. Furthermore, we also demonstrate an introduction of organic and polymeric molecule as a second component onto the aminated CCG surface to insulate hydrophobic property and π-stacking interaction of neighboring CCG sheets in aqueous solution, and to be organic solvent solubilization.

11:00am NS-WeM10 Horizontally Aligned Carbon Nanotubes on Quartz Substrate for Electrolyte-Gated Chemical and Biological Sensing, S. Okuda, Y. Ohno, K. Maehashi, K. Inoue, K. Matsumoto, Osaka University, Japan

Electrolyte-gated carbon nanotube field-effect transistors (CNTFEt) based highly sensitive chemical and biological sensors were demonstrated. Dense, well-aligned CNTs grown on quartz substrates were utilized as channels of CNTFEt. Using the large number of CNTs is a simple strategy to realize excellent performance of CNTFEt. First, the pH dependence of CNTFEt was measured in buffer solution ranging from pH 4.0 to 8.3 by monitoring the drain current (I_D) in the CNTFET. Clearly and stepwise increases in I_D were observed against the change in pH in the solution. The detection limit for changes in pH was estimated to be 0.015, which is a superior characteristic to that of conventional CNTFEt. The result indicates that CNTFEt on quartz substrates can be used as highly sensitive pH sensors. Moreover, label-free biomolecule sensing was demonstrated. The target protein was a class of antibody, immunoglobulin E (IgE). To achieve the electrolyte-gated CNTFEt, we used graphene-modified CNTFEt with multi-channel. Then, binding event of target IgE onto the aptamers was detected. IgE-concentration dependence measurements revealed that we succeeded in detection of nM quantities of IgE. In conclusion, electrolyte-gated multi-channel CNTFEt will be useful for highly sensitive chemical and biological sensors.

11:20am NS-WeM11 Gas-phase Studies and Growth of Well-Defined Carbon Nanotubes, A. Kumar, P. Lin, R.M. Sangrana, Case Western Reserve University

Carbon nanotubes (CNTs) have attracted interest for a wide-range of technological applications including nanoelectronic, energy storage, and energy conversion devices. In many of these applications, the nanotube structure must be sufficiently controlled at the growth stage; however, current growth methods typically produce mixtures of tubes, including multi-walled, single-walled, and a range of different chiralities, that must be purified and separated to facilitate applications.

We are interested in controlling the properties of CNTs at the growth stage. We have recently developed a two-step process [1-3] to study and grow CNTs consisting of a microplasma reactor that controls the size- and composition of the nanoparticle catalysts and a flow furnace that nucleates and grows the nanotubes. In addition, the nanotubes are monitored in situ by aerosol measurements to provide real-time feedback and allow the reactor conditions to be rapidly optimized. By tuning the size of our catalyst, we have found that the fraction of single-walled CNTs in the as-grown product can be varied. Similarly, by tuning the composition of the catalyst, we have found that the chirality distribution of the as-grown nanotubes is changed. Here, we will present results for Ni-based bimetallic catalysts and the influence of the catalyst size and composition, as well as other growth parameters such as carbon feedstock and growth temperature, on nanotube growth. In addition to aerosol measurements, the nanotubes are collected and characterized by micro Raman spectroscopy and UV-Vis-NIR absorbance spectroscopy. The structure and properties of the nanotubes and their relationship to the catalyst and other growth parameters will be discussed in detail.


11:40am NS-WeM12 Visualizing Defect Distributions in Carbon Nanotubes using Linear Dichroism Signatures and Scanning Transmission X-ray Microscopy (STXM) and TEM-EELS, E. Najafi, A.P. Hitchcock, D. Rossouw, G. Botton, McMaster University, Canada

The X-ray Linear Dichroism (XLD) signal in spatially resolved X-ray absorption spectromicroscopy of individual carbon nanotubes (CNT) [1] has been shown to be sensitive to the local density of sp^2 defects along the nanotube. This dichroic signal is as strong for single-walled [2] as for multi-walled CNT, which is rather surprising given the much higher curvature in SWCNT than MWCNT. The link between the strength of the XLD of the C 1s → π* peak at 285.2 eV and defect density within the sampled area has been verified by intentionally inducing sp^2 defects by ion bombardment [3]. This XLD signal is potentially useful for guiding optimization of CNT synthesis and preservation of the quality of nanotubes through various processing steps used to make functional devices where defect distribution and character play an important role. However STXM has limited spatial resolution; 10 nm, state-of-art while this work was performed at ~25 nm. Recently we have demonstrated that Electron Linear Dichroic (ELD) signals similar to XLD can be measured in q-dependent C 1s electron energy loss spectroscopy carried out in an aberration compensated, monochromated transmission electron microscope. The signals are detected by operating in STEM mode, carefully arranging the conditions such that the spectrometer accepts a narrow range of off-axis scattered electrons (with a specific location, identified in diffraction mode), and using a tilt stage to change the orientation of the CNT relative to the incident and outgoing electron directions. STEM-EELS maps measured with 2 nm sampling over a portion of a MWCNT, and over a range of tilt angles provide quantitative maps of the ELD signal. The experimental conditions will be described and the defect mapping capability of this method will be demonstrated.

STXM measurements were carried out at the SM beamline at the Canadian Light Source, which is supported by the Canada Foundation for Innovation (CFI), NSERC, Canadian Institutes of Health Research (CIHR), National Research Council (NRC) and the University of Saskatchewan. We thank Chitra Karanakaran, Jian Wang and Martin Obst for their expert support of the CLS STXM. TEM-EELS was performed with the Titan-I system of the Canadian Centre for Electron Microscopy which is supported by CFI and NSERC.


Neutron Scattering Focus Topic Room: 207 - Session NT+AS+MI-WeM

Applications of Neutron Scattering I
Moderator: V. Lauter, Oak Ridge National Laboratory

8:00am NT+AS+MI-WeM A Deeper Look into Spintronic Material Systems with Neutrons and Synchrotron Radiation, T. Brueckel, Forschungszentrum Jülich, Germany

INVITED

The discovery of the Giant Magnetoresistance GMR effect triggered the evolution of Spintronics, i.e. information storage, information processing and information transport using the spin of the electron. While the first Spintronic devices were merely transition metal multilayers, the interest has shifted to include transition metal oxide systems, laterally structured films and magnetic nanoparticles. Scattering techniques applied to model systems...
are ideal to provide fundamental microscopic information on the spin and domain structure. In this contribution, we will give an overview highlighting the capabilities of modern neutron and synchrotron x-ray techniques. We will show that neutron scattering under grazing incidence is able to provide unique depth resolved information on magnetization, magnetic correlations and magnetization dynamics relevant for basic and applied research on nanostructured magnetic materials and how synchrotron x-ray scattering can provide complementary element specific information. Examples for current research on patterned metallic multilayers, thin transition metal oxide films and magnetic nanoparticles will be given.

8:40am NT+AS+ML-WeM3 Magnetic Properties of FePtRh Films and Multilayers Studied by Neutron Scattering D. Lott, J. Fenske, Helmholtz-Zentrum Geesthacht, Germany, G.J. Mankey, Univ. of Alabama, W. Schmidt, K. Schmatzel, Forschungszentrum Juelich, Germany, E. Tartakowskaya, National Academy of Science, Ukraine, H. Amalye, ORNL, F. Klose, A. Mulders, ANSTO, Menai, Australia, A. Schwery, Helmholtz-Zentrum Geesthacht, Germany, V. Lauter, ORNL Ordered FePt alloys with L1₂ structure are known as materials with FM order and a high magnetic moment of Fe providing a large magnetization. The large atomic number of Pt on the other hand results in a high magnetic anisotropy. If grown in thin films, the high anisotropy often results in perpendicular magnetization which is the preferred orientation for current magnetic recording systems. One way to control the magnetic properties in these materials is through the introduction of a third element into the crystal matrix e.g. Rh. When Rh is added to replace Pt in the equiatomic alloy, new magnetic phases emerge. Here neutron diffraction studies on the magnetic properties of different thick Fe₉₀Ptₓₐ,Rhₓ films in dependence on temperature and external magnetic fields allowed us to investigate the rich phase diagram of the system for thin films, e.g. the transition from the FM to AF state in the system with increasing Rh concentration. In particular films with a Rh concentration of about 10% show a temperature dependent AF-FM transition. From the neutron data it was moreover possible to determine the magnetic configurations in dependence on concentration, temperature and magnetic field on a microscopic scale. Based on the observed results a theoretical model considering the changes in the anisotropies could be developed. In a next step magnetic multilayer consisting of Fe₉₀Ptₓₐ,Rhₓ, bilayers with different Rh concentrations were grown and studied by polarized neutron reflectivity to investigate the magnetic interactions along the lattice matched interfaces. First results will be presented here.

9:00am NT+AS+MI-WeM4 Spectroscopic and Magnetic Characterization of the Spin-Crossover Transition in Thin Films of Fe(C₁₂H₈N₂)₂(NCS)₂, E.C. Ellingsworth, G.J. Szulczewski, The University of Alabama, Tuscaloosa, V. Lauter, Oak Ridge National Laboratory The octahedral complex bis(1,10-phenanthroline)dithiocyanate iron(II), Fe(phen)₂(NCS)₂, is known to exhibit an abrupt transition between a high and low magnetic spin state from 170 – 180 K in the bulk phase. As a result, Fe(phen)₂(NCS)₂ is an interesting organic semiconductor to study charge and spin transport in thin films. We synthesized and characterized Fe(phen)₂(NCS)₂ according to literature procedures. Thin films of Fe(phen)₂(NCS)₂ were made by vapor deposition onto a variety of substrates including Si, KBr, Au and Al and characterized by infrared and photoelectron spectroscopy, SQUID magnetometry, optical microscopy, and polarized neutron reflectometry. The films were found to be very sensitive to water vapor under ambient conditions, which complicates the structural, chemical and magnetic analysis of the films. However, appropriate capping layers can be deposited onto the Fe(phen)₂(NCS)₂ films to protect them from water vapor before removal from the vacuum system. The differences in the magnetic behavior of the thin films will be compared to the bulk phase.

9:20am NT+AS+MI-WeM5 Nanoscopic Magnetic Phase Separation at the SrTiO₃/SrₓCoₙO₃ Interface M. Sharma, M.A. Torija, Univ. of Minnesota, J. Gázquez, M. Varela, ORNL, J. Schmitz, C. He, Univ. of Minnesota, J.A. Borchers, M. Laver, NIST, S. El-Khatib, University of Sharjah, V. Lauter, H. Amalye, R. Gayot, ORNL, C. Leighton, Univ. of Minnesota INVITED

The remarkable functionality of complex oxides, when combined with the favorable lattice matching that is possible at their interfaces, provides many opportunities for new physics and applications. The perovskite manganites and cobaltites are excellent examples, being of interest in gas sensing, catalysis, and as electrodes in ferroelectric memory and solid oxide fuel cells. From the magnetism perspective they have potential for high conduction electron spin polarization, and a variety of functional ground states. However, the same delicate balance between phases that provides such impressive functionality also leads to a serious problem; it can be difficult to maintain desired properties (e.g. high spin polarization and conductivity) close to the interface with a dissimilar oxide. This is exemplified by magnetic tunnel junctions for example, where the interface spin polarization is suppressed and drops rapidly with temperature. In this work, using SrTiO₃(001)/La₁₋ₓSrₓCoO₃ [1] as a model system, we have combined epitaxial growth by high pressure oxygen sputtering with atomic-level structural characterization (including STEM/EELS imaging) [2], conventional magnetometry, electronic transport, small-angle neutron scattering, and polarized neutron reflectometry. We observe the usual degradation in magnetization and conductivity in the very thin film limit. We demonstrate that this is due to nanoscopic magneto-electronic phase separation in the interface region [3]. Essentially, nanoscopic ferromagnetic (FM) clusters form in an insulating non-FM matrix near the interface, resulting in reduced magnetization and conductivity, even at compositions that display no such phase separation in bulk. STEM/EELS depth profiling of the chemical composition reveals that this effect has a chemical origin, being due to subtle depth-wise variations in Sr and O content, resulting in reduced hole doping near the interface. Simple thermodynamic and structural arguments for the origin of these variations are provided, based on Sr dissolution energies and the critical link between strain state and O vacancy concentration provided by O vacancy ordering [2,3].

Work at UMN supported by NSF and DOE (neutron scattering). Work at ORNL supported by DOE. Work at UCM supported by the European Research Council.


10:40am NT+AS+MI-WeM9 Study of L1₀ Ordering in ⁰Fe/Pt Multilayers, K. Srikanti, Uge-Dae, Csr, India

Ordered L₁₀ FePt, FePd and CoPt alloy thin films have large magnetic anisotropy constants suitable for high-density recording media. The L₁₀ ordered phase is obtained with post growth annealing at high temperatures. However, the high temperature annealing leads to grain growth. The present work is an attempt to lower the transition temperature starting with multilayer precursors. The evolution of the structural and magnetic properties of Si(111)/Fe₀Pt(19Å)/Pt(25Å)/Pt(99Å) multilayers as a function of vacuum annealing at different temperatures is studied. The film thickness is selected to have equi-atomic stoichiometry. The multilayers are prepared by ion beam sputtering. X-ray reflectivity (XRR), X-ray diffraction (XRD), and magneto optical Kerr effect (MOKE) and conversion electron Mössbauer spectroscopy (CEMS) are used to characterise the as-deposited and annealed multilayers. Using XRR it is observed that due to intermixing FePt alloy formation takes place with annealing. The XRD indicated the presence of superstructure peaks at 350°C and above. Mössbauer measurements indicated a clear evidence for the strong exchange coupling between the soft fcc FePt and hard fct FePt phase. The results indicate that the multilayer structure does not transform directly to the ordered fct FePt, rather first an fcc FePt phase is formed and subsequently it gets converted in to the fct FePt phase as a function of annealing. A detailed study on stoichiometric FePt system embedded in Carbon matrix prepared by ion beam sputtering technique will also be discussed. Few preliminary results of polarised neutron reflectivity measurements on FePt/FeNi exchange spring magnets will be discussed.

Temperature sensitive hydrogel polymers are utilized as responsive layers in various applications. While the polymer’s native characteristics have been studied extensively, details concerning its properties during interaction with bio-related structures are lacking. This work investigates the interaction between a thermoresponsive polymer cushion and different lipid membrane capping layers probed by neutron reflectometry. N-isopropylacrylamide copolymerized with methacryloxybenzophenone first supported a lipid bilayer composed of 1,2-Dipalmitoyl-sn-Glycero-3-Phosphoethanolamine (DPPE) and subsequently 1,2-Dipalmitoyl-sn-Glycero-3-Phosphocholine (DPPC). The polymer-membrane systems were investigated above and below the polymer lower critical solution temperature (37 and 25 °C). While the same cushion supported each lipid membrane, the polymer hydration profile and thickness were markedly different for DPPE and DPPC systems. Since DPPE and DPPC have different bending rigidities, these results establish that the polymer-membrane interaction is critically mediated by the mechanics of the membrane, providing better insight into cell-hydrogel interactions. There has been increased interest in the effect of matrix elasticity on cell lineage specification. Polymeric matrices with known stiffness are utilized as supports to understand the physical effects of *in vivo* tissue microenvironment for therapeutic uses of stem cells. This work focuses on the influence of a capping layer on the mechanical properties of the underlying support.

**Plasma Science and Technology Division**

Room: 201 - Session PS+SE-WeM

**Atmospheric Plasma Processing and Micro Plasmas**

**Moderator:** P.L.S. Thambur, University of Texas at Dallas

8:00am PS+SE-WeM1 2011 AVS Peter Mark Award Lecture - Microscale, Atmospheric-Pressure Plasmas: A Platform for Nanomaterials Synthesis at Different Length Scales, M. Sankaran*, Case Western Reserve University

**INVITED**

Large-scale, low-pressure plasmas play an essential role in the processing of materials for a wide-range of applications including integrated-circuit (IC) manufacturing. In recent years, new challenges have arisen for these top-down approaches to materials processing. Advanced electronic devices will be comprised of nanomaterials such as nanoparticles and carbon nanotubes that cannot be fabricated by current plasma technology because of limitations associated with photolithography. In addition, emerging applications in sensors, energy, and medicine require nanomaterials that must be prepared from the “bottom-up” and assembled into macroscale structures. The aim of our research is to develop a new class of plasmas, termed microplasmas, for nanomaterials synthesis and assembly.

Microplasmas are electrical discharges formed in geometries where at least one dimension is less than 1 mm. As a result of their *pD* scaling (*p* is the gas pressure and *D* is the smallest dimension), microplasmas operate stably at atmospheric pressure. These properties open up unique opportunities for nanomaterials synthesis and assembly. For example, vapor-phase metal-organic precursors can be dissociated near ambient conditions to homogeneously nucleate metal [1] and alloyed [2] nanoparticles. The formation of well-defined metal nanoparticles in the gas phase allows direct introduction of these materials as catalysts for chiral-enriched carbon nanotube growth [3]. Recently, we have also coupled microplasmas with liquids to electrochemically synthesize nanoparticles from aqueous metal salts [4]. By extending this strategy to thin films, microscale patterns of nanoparticles are fabricated in a single step [5]. In this talk, I will discuss these topics in detail, highlighting the advantages of microplasma-based systems for the synthesis of well-defined nanomaterials at various length scales.


*Peter Mark Memorial Award Winner*

8:40am PS+SE-WeM3 Surface and In-Depth Modification of LDPE using an Atmospheric Plasma Torch. S. Abou Rich, P. Leroy, Université Libre de Bruxelles, Belgium, N. Wehbe, University of Namur, Belgium, N. Avril, L. Houssiau, University of Namur, Belgium, F. Reniers, Université Libre de Bruxelles, Belgium

In order to improve the adherence of a coating onto a polymer, the substrate surface must be properly functionalized. Plasma technologies are more and more used for that purpose. However, plasma treatments do not only modify the polymer surface, as active species can penetrate into its bulk.

In the present study, polyethylene samples are surface-functionalized by an atmospheric plasma torch, using argon and a mixture of argon-oxygen. The surface is characterized by dynamic water contact angle, atomic force microscopy and X-ray photoelectron spectroscopy. The changes in the bulk of the polymer have been characterized using infrared spectroscopy, angle-resolved X-ray photoelectron spectroscopy (ARXPS), and secondary ion mass spectrometry (SIMS).

It is shown that both plasma treatments (argon or argon-oxygen) lead to an increase in the hydrophilicity of the surface by the grafting of oxygen-containing polar functional groups. AFM reveals also a change in surface roughness induced by the plasma treatment, depending on the operational conditions. FTIR, ARXPS and SIMS evidenced that the plasma treatment also modifies a significant zone under the surface. The diffusion depth of oxygen as a function of the plasma parameters (power, treatment time, oxygen ratio in the gas phase) is determined.

The ageing of the plasma-modified polymers during storage in air is also studied.

9:00am PS+SE-WeM4 Surface Analysis of Polymers Treated by Remote Atmospheric Pressure Plasma. R.F. Hicks, University of California Los Angeles, E. Gonzalez, Intel Corporation, T.S. Williams, University of California Los Angeles

Atmospheric plasma treatment is a key process for improving the adhesive bond strength of polymers in many products, such as thermoplastic composites on jet aircraft and helicopters, molded plastic parts in automobiles, and plastic tubing and stents in implantable medical devices. In this study, the surface of high-density polyethylene (HDPE), poly(methyl methacrylate) (PMMA), and polyethersulfone (PES) were treated with a low-temperature, atmospheric pressure oxygen and helium plasma. The polymers were exposed to the downstream afterglow of the plasma, which contained primarily oxygen atoms and metastable oxygen molecules (*Δ* O2), and no ions or electrons. X-ray photoelectron spectroscopy of the HDPE revealed that initially 20% of the carbon atoms were converted into oxidized functional groups, with about half of these being carboxylic acids. Attenuated total reflection infrared spectroscopy of all three polymers was obtained in order to determine the types of functional groups formed by atmospheric plasma exposure. It was found that the polymers were rapidly oxidized with addition of alcohols, ketones, and carboxylic acids to the carbon backbone. Chain scission occurred on HDPE and PMMA, while on PES the aromatic groups underwent ring-opening and insertion of carboxylic acid. The implications of this work for forming strong adhesive bonds to these polymers will be discussed at the meeting.

9:20am PS+SE-WeM5 Deposition of SiO2 Films by Means of Atmospheric Pressure Microplasma Jets: Study of Deposition Mechanism, J. Benedikt, R. Reuter, D. Ellerweg, K. Ruegner, T. de los Arcos, A. von Keudell, Ruhr-University Bochum, Germany (INVITED)

Deposition of thin films with plasmas at atmospheric pressure is always a challenging task because of high collision rates, absence of ion bombardment, filamentary behavior of the plasma and limited knowledge of plasma chemistry. The preparation of high quality thin films is therefore still the main domain of low pressure plasmas. One of films, which can be prepared at atmospheric pressure, is SiO2. Hexamethyldisiloxane (HMSDO) and O2 (or N2O) are usually used as precursors fed into the plasma.

We have shown in the past that a good quality SiO2 films can be prepared by using microplasma jets driven by RF voltage and operated in Ar or He as plasma forming gas. Here we concentrate on the study of plasma chemistry and surface reactions leading to the film growth. The geometry of the microplasma jet and the localization of the plasma treatment allow studying of gas phase reactions and plasma-surface interaction separately. Molecular beam mass-spectrometry is used to measure HMSDO depletion and stable products in the gas phase. Depletion below 15% and limited fragmentation is observed even under conditions with high O2 density, which leads to formation of carbon free SiO2 films. The plasma-surface interaction is studied by application of several jets with different gas mixtures (He:HMDSO, He2O2, He:He/H2) to the same trace on the rotating substrate in controlled helium atmosphere. It is shown that surface reactions are responsible for the carbon removal from the grown film. Infrared spectroscopy, spectroscopic ellipsometry, and X-ray photoelectron spectroscopy measurements are performed to analyze film properties and surface evolution.
compare them with plasma measurements. A fluid model of gas flow and reaction kinetics in the effluent of the plasma is used to reproduce observed trends and measured deposition rates. Good agreement is achieved with relatively simple model of plasma chemistry and surface reactions.

11:00am PS+SE-WeM10 Synthesis of Li$_2$TiO$_3$ Nanoparticles Using an Atmospheric Pressure Plasma Jet, S.M. Chang, E.F. Rodriguez, H.C. Li, Y.J. Yang, N.L. Wu, C.C. Hsu, National Taiwan University, Taiwan, Republic of China

Nanocrystalline spinel Li$_2$TiO$_3$ (LTO) is a promising anode material for Li-ion batteries due to its zero-strain during intercalation/deintercalation and the 1.5 V (vs. Li+/Li) potential plateau. Development of a process that allows for rapid synthesis with controllable microstructure has been challenging. In this work, LTO nanocrystalline particles synthesized by an atmospheric pressure plasma jet (APPJ) will be presented. The APPJ is sustained using a repetitive pulsed power source with N$_2$ gas. Ti and Li ions-containing solution of specific formula is used as the precursor solution. The precursor is ultrasonically nebulized and is then carried into the downstream of the APPJ using a carrier gas. With this process, nanocrystalline LTO can be fabricated in one step with a short contact time (a few ms) between the precursor and the plasma jet without an additional annealing process. The LTO particle size can be effectively controlled using the precursor solution concentration and the carrier gas flow rate. 100-300 nm dense spherical particles can be fabricated under an optimized condition. The microstructure and porosity of the particles are strongly influenced by the pre-heat process for the nebulized precursor droplets prior to entering the jet downstream. Dense spherical and porous particles are fabricated under conditions with and without preheating, respectively. Finally, full battery performance test will be presented and key factors that dominate the fabricated nanoparticle characteristics will be discussed.

11:20am PS+SE-WeM11 Laser-Assisted Plasma Coating at Atmospheric Pressure: Production of Yttria-Stabilized Zirconia Thermal Barriers, Z. Ouyang, P. Raman, Y.L. Wu, L. Meng, T.S. Cho, D.N. Ruzic, University of Illinois at Urbana-Champaign

A laser-assisted plasma-coating technique at atmospheric pressure (LAPCAP) for use in thermal-barrier coatings (TBC) deposition has been developed. This technique allows PVD-quality depositions to be done at atmospheric pressure. The microwave plasma source employed has a working frequency at 2.45 GHz, and a maximum input power of 6 kW, and the attached plasma torch head has the ability to generate various types of atmospheric-pressure plasmas at the temperatures of room temperature (20°C) to more than 2,000°C. Optical emission spectroscopy (OES) technique has been used to spatially analyze some critical characteristics of plasma, such as electron density ($n_e > 10^{17}$ cm$^{-3}$), electron temperature ($T_e \approx 1$ eV), and plasma gas temperature ($T_p \approx 400-3,000$ K), under different operating conditions (gas type, input power and gas flow rate). A helium atmospheric plasma has been used to assist in Nd:YAG laser ablation (f = 100 Hz, Energy/pulse = 20 mJ at 266 nm; 120 mJ at 532 nm; 325 mJ at 1064 nm) of a 3% yttria-stabilized zirconia (3YSZ) target, to provide a more conformal deposit thickness, adhesion and control of the deposition on Rene N5 substrates with better lamination at a relatively higher deposition rate ($\approx 1,000$ nm/min), in comparison to traditional PVD methods. The morphology and characteristics of the films have been compared at three laser wavelengths (266 nm, 532 nm and 1064 nm), different laser energy densities (1-10 J/cm$^2$) and substrate temperatures (20-1100°C), using microscopy techniques such as scanning electron microscope (SEM), focused ion beam (FIB), X-ray photoelectron spectroscopy (XPS), and X-ray diffraction (XRD).

11:40am PS+SE-WeM12 Investigation on the Discharge Formation Mechanisms and Surface Analysis of SiO$_2$-like Layers on Polymers Synthesized Using High Current Dielectric Barrier Discharge at Atmospheric Pressure, M.C.M. van de Sanden, FOM-Inst. for Plasma Phys., Rijnhuizen & Eindhoven Univ. of Tech., Netherlands, A. Premkumar, Eindhoven Univ. of Tech., & M2I, Netherlands, S. Starostin, N. Wu, C.C. Hsu, N. Li, Y.J. Yang, N.L. Wu, C.C. Hsu, National Taiwan University, Taiwan, Republic of China

The dielectric barrier discharge is recognized as a promising tool for PECVD of thin films at atmospheric pressure. Emerging applications including encapsulation of flexible solar cells and flexible displays requires low costs production of transparent uniform and dense layers with low level of charged defects. Among the two discharging Townsend like discharge (TD) and glow like discharge (GD) the latter offers more flexibility for the high growth rates in plasma enhanced deposition. In this investigation we demonstrate the utilization of glow like discharge in, He free, industrially relevant gas mixture comprising Ar/N$_2$/H$_2$ for the deposition of high quality silica like films on large area polymeric substrates (PET or PEN) in a roll-to-roll configuration. While the discharge physics exhibiting the glow like behaviour is investigated via fast ICCD camera, voltage-current waveforms and optical emission spectroscopy, the deposited silica like films is comprehensively analyzed using AFM, SEM, XPS, SE and FTIR. The same evolution of the diffuse discharge, showed several phases starting from the initial ignition of the low current Townsend-like mode followed by the transition to glow like discharge which then undergoes lateral expansion providing uniform treatment of the whole substrate width. As a generic characteristic of the developed technology, it is observed that, irrespective of precursors (TEOS or HMDSO) or process gases (Ar/N$_2$ or air) employed, the films are smooth, both locally and globally, and of near stoichiometric silica with very low carbon content (< 2%). Detailed AFM morphology description and surface statistical analysis on SiO$_2$ dynamics showed that no film roughening in growth front and lateral directions observed and the synthesized layers (~350 nm) grow in a self-similar fashion following the topology of the substrate with no discontinuity. The films are grown with no defects, film are smooth incorporated during the deposition process and exhibit excellent barrier performances towards O$_2$ and H$_2$O permeation.

**Plasma Science and Technology Division**

**Room: 202 - Session PS+SS-WeM**

**Plasma Surface Interactions (Fundamentals & Applications)**

**Moderator:** C. Labelle, GLOBALFOUNDRIES

8:00am PS+SS-WeM10 Investigation of Sidewall Passivation Mechanisms in a CMOS-Compatible Plasma Etching Process for InP-based Photonic Devices, S. Bouchoule, CNRS-LPN, France, L. Vallier, CNRS-LTM, France, L. Gatiolova, G. Patriarche, S. Guilet, L. Le Gratiet, CNRS-LPN, France

Inductively coupled plasma (ICP) etching of IV-V semiconductors is now widely used for the development of high-performance emitters, and various chlorine- or HBr-containing chemistries have been proposed for the patterning of InP-based heterostructures required to reach the NIR region. Smooth and anisotropic etching is generally a key requirement, but only few studies exist on the understanding of the sidewall passivation mechanisms occurring during the etching of InP and related materials. We have shown for the Cl$_2$-H$_2$ and HBr chemistries that a silicon oxide layer acting as a lateral etch-inhibitor can build-up on the etched sidewalls of InP-based heterostructures, when a Si wafer is used as the sample tray. This configuration corresponds to most commercial ICP etch systems having an electrode diameter of 4-in or more, used to etch III-V samples of 2-in or less size. However, this may not be the case for future large surface processing of III-V when the III-V wafer will have the same size as the electrode or when III-V dies bonded onto a 200/300 mm wafer may to be etched, where most of the wafer surface is covered by a protecting layer that is not silicon. This may occur in III-V/Si photonic technologies. We have shown that high-aspect ratio etching of the photonic patterns via a SiO$_x$ sidewall passivation mechanism independent of the electrode surface can be obtained when a Si-containing gas such as SiH$_4$, or SiCl$_4$ added [JVSTB 29, 026001 (2011)]. A more detailed analysis of the plasma has shown that hydrogen may promote the deposition of a Si-rich passivation layer on the sidewalls of the etched patterns. SiOCl$_2$ sidewall passivation takes place during Si ICP etching using Cl$_2$-HBrO$_x$ chemistry in CMOS technology. We have therefore investigated SiCl$_4$/Cl$_2$/HBr/O$_2$/Ar plasma for the etching of InP dies in a 300-mm CMOS etching tool. This gas mixture provides SiO, and H species required for the build-up of a SiO$_x$ passivation layer on the InP sidewalls. We show that the passivation mechanism is enhanced when the HBr concentration is increased in the feed gas. We have performed a local analysis of the passivation layer deposited on the InP sidewalls using EDX spectroscopy coupled to TEM. We show that the nature of the passivation layer can be changed from a Si or nc-Si to SiO$_2$ depending on the hydrogen and oxygen concentrations in the gas mixture. Finally we demonstrate smooth and anisotropic etching of ridge waveguide and vertical Bragg reflector patterns in the CMOS etching tool.

8:20am PS+SS-WeM2 Coupling of Surface Mixed-Layer Kinetics and Monte Carlo Modeling for Profile Evolution in Patterning Complex Oxides, N. Marchack*, C.D. Pham, J.P. Chang, University of California Los Angeles

As the downsizing of integrated circuit devices continues, minute variations in the feature profiles from processing techniques such as plasma etching significantly affect device performance. With the increasing introduction of novel materials into integrated circuits, the need to predict post-etching profiles has become critical. Monte Carlo Modeling for Profile Evolution (MCM-PE) provides a means of predicting the profile evolution of diluted and complex multiple-layer systems. As shown in the work presented by Dr. Prasad in this session, MCM-PE has successfully predicted the etch profiles of InP for an ICP etch system. The approach, however, does not account for the effect of mixed-layer chemistry on the etch profiles. This work presents a modified MCM-PE approach to profile evolution in pattering complex oxides, where the mixed-layer chemistry is factored into the profile evolution. A method is also presented to evaluate the impact of different mixed-layer systems on the etch profiles, which can be used to derive useful design rules for optimal circuit performance.

* Coburn & Winters Student Award Finalist
surface, and the signal using photolysis of CCl₄ gave unphysically high values, probably due to poor knowledge of the laser spatial profile at the focal point. Therefore, we developed a new technique, based on 355nm (tripled YAG) photolysis of Cl₂ to generate a known density of Cl atoms. The variation of the absolute Cl density at the reactor centre was measured as a function of pressure and RF power in the range 3-90 mTorr and 20-500W. We also used the TALIF technique to determine the recombination coefficient, γCl₂, of atomic chlorine at the reactor walls from the rate of decay of the Cl density in the afterglow of a pulsed discharge. The signal to noise ratio is good enough to make measurements far into the afterglow (50 ms), when the gas has cooled to the wall temperature, making a precise measurement possible. We found that γCl₂ varies in the range 0.05-0.15, decreasing with increased pressure and RF power, and increasing as the gas residence time. We show that the latter effect is due to the increased proportion of O₂ due to inevitable small air leaks: the presence of 0.5% O₂
was shown to double the value of $\tau_c$. The origin of the pressure and power dependencies will be discussed.

Work partly supported by Agence Nationale de la Recherche project INCLINE (ANR-09 BLAN 0019)


11:00am PS+SS-WeM10 Silicon Etching Characteristics by Hydrogen Halide Ions (HCl+ and HBr+) and Ions of Desorbed Species (SiClx+).

T. Ito, K. Karahashi, Osaka University, Japan, S.-Y. Kang, Tokyo Electron Ltd., Japan, S. Hamaguchi, Osaka University, Japan

In recent reactive ion etching (RIE) processes for Si, halogen and hydrogen halide gases, such as Cl2 and HBr, have been widely used to achieve high selectivity, etching anisotropy, and high etching rates. Furthermore, in some highly selective silicon etching processes, higher gas-pressure processes have been found to be more effective. In higher-pressure systems, chemical compounds formed from the input gas and some of desorbed species containing Si may serve as additional etchants. To develop etching equipments based on such plasma chemistry, it is important to understand the evolution of etch products on silicon surfaces by energetic ion species associated with silicon and/or hydrogen containing species. To clarify the roles of SiClx, SiBrx, HCl, and HBr in silicon etching processes, we have employed a mass-analyzed ion beam system that can irradiate a sample surface with a specific ionic species under an ultra-high vacuum condition and evaluated the etching yields. The change in chemical nature of the substrate surface during the process can be observed in situ by X-ray photoelectron spectroscopy (XPS) installed in the reaction chamber. Time of Flight (TOF) measurement of species desorbed from the sample surface in a pulsed ion beam operation is also possible with the use of a differentially pumped quadrupole mass spectrometer (QMS). In this study, etching yields of silicon by Cl+, SiCl+, SiCl2+, Br+, H+, HCl+, and HBr+ ions were measured with incident energies of 100–1000 eV. A typical ion dose for each ion irradiation was 2–4E17/cm2. Yields by some of these ionic species have been known and our etching yield data are confirmed to be in good agreement with the earlier data. It is found that, for a given incident energy, the etching yield by SiCl1+ ions is higher than that by Cl+ ions whereas the etching yield by SiCl+ ions is lower than that of Cl+ ions, which may be accounted for by the number of Cl atoms and a possible deposition effect of Si. It has been also observed that deposition occurs under SiCl+ ion irradiation when the injection energy is lower than 300eV. Energy dependence of etching yields and effects of hydrogen will be discussed in detail.

11:20am PS+SS-WeM11 Interaction of Chlorine Plasma with SiCl, Coated Plasma Reactor Chamber Walls. R. Khare*, A. Srivastava, V.M. Donnelly, University of Houston

The interplay between chlorine plasmas and silicon chloride (SiClx) coated reactor walls has been studied by line-of-sight mass spectrometry in a radio frequency (rf) inductively coupled Cl2 plasma (ICP), using the spinning wall method. A bare silicon wafer was etched in a 400 Watt Cl2 ICP, with rf power applied to the stage resulting in a 1-10 VDC self-bias. Ethyl products were deposited on the plasma reactor walls and the rotating substrate surface, resulting in a thick layer of SiCl, that was characterized in situ by Auger electron spectroscopy. Some oxygen also incorporated into the film due to erosion of the fused silica discharge tube. The reactions of chlorine plasmas with this prepared surface were then studied by line-of-sight mass spectrometry. Without substrate bias, the chlorine plasma etches the SiCl layer to form products that result in detection of SiClx (x = 1-4) m/e components, as well as oxo-silicon-chloride products (m/e = 177, 247, 307, 361). In one experiment, after the deposition of dense SiCl layer on reactor and substrate surfaces, substrate rotation was stopped and the film was etched from the reactor walls with the chlorine plasma, leaving only the SiCl layer on 2/3 of the substrate surface that was out of the plasma. Upon resuming rotation, and exposing the SiCl2, Cl2 loaded surface to the Cl2 plasma, SiCl2 products were detected, but at suppressed levels, indicating that the evolution of eth products is a complex “recycling” process in which these species deposit and desorb from the walls many times, and repeatedly fragment in the plasma. These and other experiments will be discussed. This work is supported by the National Science Foundation and Lam Research Corporation.

* Coburn & Winters Student Award Finalist

11:40am PS+SS-WeM12 Numerical Simulation of Enhanced Oxygen Diffusion in Silicon as a Cause of Si Recess. K. Mizutani, M. Isobe, Osaka University, Japan, M. Fukasawa, T. Tatsumi, T. Hamaguchi, Osaka University, Japan

In a gate etching process, the formation of hollowed Si profiles around the polysilicon (poly-Si) gate can be severely degraded the device performance and reliability. In a typical process that causes Si recess, a plasma based on HBr and oxygen gases are used to etch polysilicon gates anisotropically. A recent study [1] based on multiple-beam and plasma experiments has shown that Si recess is caused by ion assisted oxygen diffusion, i.e., oxygen diffusion enhanced by hydrogen ion injections. In this study, we have used molecular dynamics (MD) simulations to understand the mechanism of enhanced oxygen diffusion in Si under such conditions. In the simulations, energetic hydrogen ions and oxygen ions (with kinetic energies close to room temperature) are simultaneously injected into a crystalline Si substrate initially covered with a native oxide layer. Simulation results are in good agreement with ion beam experiments performed under similar conditions given in Ref. [1]. In our simulations, O atoms are transported into the bulk Si due to momentum transfer from energetic hydrogen ions. In other words, the enhanced ion transport is not typical “diffusion” associated with thermal motion in solid. However, random walk characteristics of O atoms in Si under such conditions are interestingly similar to those of diffusion. In this study, we relate this oxygen transport to diffusion transport and present its effective diffusion coefficient as a function of hydrogen ion injection energy.

temperatures. The second derivative of the tunneling current was used to obtain information on the interfacial thermal coupling and energy transfer. This made it possible to perform experimental studies on the thermal conductance of metal-carbon interfaces, corresponding first principle calculation approaches for small gap interfacial thermal coupling, and discusses experimental and modeling results for different tip-surface combinations toward understanding near-field effects for thermal energy transfer.


Conventional single-color laser pump-probe methods for measuring thermal properties are limited by sample requirements that arise from considerations of surface roughness and compatible thermoreflectance transducers. Here we describe a new experimental arrangement for performing two-color time-domain thermoreflectance (TDTR). The technique is a variation of traditional pump-probe spectroscopy that is based on a femtosecond Ti:sapphire oscillator of fixed wavelength and an optical parametric oscillator (OPO), with the goal being to create an independently tunable probe wavelength. This method offers two advantages: 1) spectral filtering of diffusely scattered pump light (to prevent it from reaching the detector), and 2) generation of the thermoreflectance signal from different sample films.

The wavelength tunability of the system allows enhancement of TDTR signal generation for multiple thermoreflectance transducer materials. This wavelength-adjustable feature, in turn, facilitates direct measurement of the thermal transport properties of various thin films and substrates, which would be difficult with single-color femtosecond pump-probe systems. Demonstrated results include optimization of the probe wavelength for different metals, measurement of metal-graphite interfacial conductances on relatively rough samples, and two orders-of-magnitude calibration of thermal conductivity measurements using copper as a thermoreflectance transducer.

9:40am SE+SS-WeM6 Low-Friction V-alloyed ZrO2 Thin Films with Temperature Homogenization Functions for High Temperature Sliding Interfaces. O. Jantschner, C. Walter, C. Mitterer, University of Leoben, Austria

The effect of vanadium on reactively magnetron-sputtered zirconia coatings was investigated with respect to its structural and mechanical properties as well as its thermal management abilities for high temperature sliding interfaces. ZrO2 coatings with different V-content (0, 2.2, 5.8 and 17.4 at%) were co-sputtered from Zr and V targets using an Ar/O2 discharge. The X-ray diffraction pattern of the as-deposited coatings show a change in crystal structure from monoclinic (0-2.2 at% V) to cubic/trigonal (5.8 at% V) and finally X-ray amorphous structure at even higher V content (17.4 at% V). Hardness and Young’s modulus were evaluated by nanoindentation showing a decrease beyond 2.2 at% V from 17.4 to 7.5 GPa and from 230 to 150 GPa, respectively. The tribological investigations by ball-on-disc tests against alumina balls were carried out at three different temperature levels (25°C, 600 am) and (25°C, 600 am). Additional thermal analysis was performed to study the formation of tribofilms in the sliding contact. At 25°C, the coefficient of friction (COF) is about 0.2 for low V contents (≤2.2 at%). For higher V contents, the COF increases up to 0.5 and higher. At 600°C, the COF measured was between 0.4 and 0.8. At the even higher temperature of 800°C, the COF decreased to below 0.2 for V contents of 17.4 at%, where a self-lubricating film was formed in the sliding contact. Differential scanning calorimetry (DSC) measurements of virgin powder samples showed a characteristic exothermic peak at ~600°C which is due to the formation of a stoichiometric ZrV2O7 phase. This phase was found to decompose at ~800°C by an endothermic reaction in ZrO2 and V2O5. The re-runs of the DSC measurements indicated melting of V2O4 at 670°C.

In summary, alloying of V to ZrO2 coatings has on the one hand been proven to result in self-lubricous properties at temperatures above 700°C. On the other hand, the endothermic reactions needed for formation of the self-lubricous phase have the potential to reduce high local temperatures in the sliding contact, enabling thermal management abilities of these coatings.

10:00am SE+SS-WeM9 Thermal Characterization of Metal/Carbon Interfaces: Comparison of Metallized Nanotubes and Graphite. C. Muratore, S. Shenogin, A. Waite, A. Reed, J. Gengler, T. Smith, J. Hu, J. Bulman, A.A. Voevodin, Air Force Research Laboratory

Most applications of carbon nanotubes require contact with more ordinary materials, such as metals or polymers. Unfortunately, the extraordinary thermo-electro-mechanical properties of nanotubes are often neglected at the interface between the nanotubes and whatever they touch, resulting in a major shortfall between the measured and predicted performance of nanotube-based materials. One of the most troubling discrepancies in projected versus measured properties is found in thermal conductivity measurements on experimental nano-composite materials. For example, a continuous network of thermally conductive nanotubes (or about 1 percent, by volume) within an organic matrix (k = 0.3 W m-1 K-1) should yield a 30-fold increase in thermal conductivity over the pure matrix phase alone, based on simple effective medium theory. Despite this potential increase, experimental results typically show an increase of only a factor of 2 at best in composite films with nanotube fill levels.

To better understand the nature of interfacial resistance in carbon nanotubes, modeling and experimental studies investigating engineered interfaces on highly oriented pyrolytic graphite (HOPG) samples were conducted. This substrate was selected as a practical 2-dimensional analog for nanotube sidewalls to facilitate modeling and experimentation. Molecular dynamics simulations of heat transfer through metal-carbon interfaces were conducted, and measurements of thermal conductance at these interfaces were made by analysis of the two-color time domain thermoreflectance (TDTR) data from the samples. The TDTR analysis of the different metals on HOPG was made possible by having an optical parametric oscillator on the probe beam which allows for tuning the wavelength to match absorption bands for each metal studied. Comparison of simulation and experimental results between graphite and nanotubes is highlighted. Metal films were selected to identify effects of atomic mass, chemical interactions and mechanical properties. For example, metals known to exhibit in situ formation of an interfacial carbide layer when in contact with a carbon source and heated, such as titanium and boron, were investigated, and the effect of this carbide layer formation on interfacial conductance was examined. Graded and sharp interfaces were also considered with computational and experimental efforts.

11:00am SE+SS-WeM10 The Experiment of Surface Tension Driven Flow with Various Parameters on JEM/ISS. S. Yoda, Japanese Aerospace Exploration Agency, S. Matsumoto, JAXA, Japan, A. Komiyama, Tohoku University, Japan

The surface tension driven flow (Marangoni) experiments were carried out by using 50mm diam. with liquid bridge of Silicones oil under microgravity condition on Japanese Experiment Module on International Space Station. The experiments in these experiments we used liquid bridge length being correspondent to aspect ratio which is defined as liquid length/liquid diam., and temperature difference between hot and cold disks which sustains the liquid bridge. The particles coated by gold to fit with the density of the liquid silicones were inserted into the liquid bridge. Observing the movement of the particles by three CCD camera, we can determine the flow behavior of Marangoni with different temperature as 3 dimension observation of the flow. Moreover, two ultrasonic transducers were attached in the cold disk to measure the particles velocity. The number of experiments done on JEM were more than 40 times with around 6 hr for each experiment. The Marangoni number was changed with aspect ratio. The smaller aspect ratio around showed smaller critical Marangoni numbers, whereas the larger those were the larger numbers.


Thermal transport is restricted in systems including carbon nanotubes (CNT) due to high thermal interface resistance. We have substantially improved thermal transport at CNT-metal interfaces by functionalizing the contacting surfaces with aramid-propyl silane to form covalent chemical bonds bridging the CNT-metal gap. This strategy was suggested by molecular dynamics calculations of Hu et al. (1) We have characterized the resulting interface using multi-frequency and multi-wavelength time-domain thermoreflectance (TDTR) measurements, which are able to independently determine interface and bulk CNT contributions to thermal transport. TDTR analysis showed that thermal contacts were made between a functionalized Al surface and ~5% of the CNTs in an array grown on silicon. The intrinsic CNT interface conductivity exceeded 300 MW/m²-K, resulting in an effective thermal interface resistance of less than 0.8 mm²-K/W. Successful interface functionalization was independently verified by mechanical adhesion testing, which showed a correlation between interface strength and thermal transport, as discussed by Prashar. (2)

(2) Ravi Prashar, Appl. Phys. Lett. 94 041905 (2009)
Surface Science Division
Room: 107 - Session SS1-WeM

Atomic Control of Structure & Evolution
Moderator: T.S. Rahman, University of Central Florida

8:40am SS1-WeM3 Destabilization of Ag Nanoislands on Ag(100) by Adsorbed Sulfur. M. Shen, S.M. Russell, Iowa State University, D.-J. Liu, Ames Laboratory - US DOE.

Studies of chalcogen (O, S) interactions with coinage metal surfaces elucidate how those interactions affect mass transport on the surface and whether a general mechanism exists for these systems. Sulfur accelerates coarsening of Ag islands on Ag(100) at 300 K, and this effect increases with sulfur coverage over a range spanning a few hundredths of a monolayer, to nearly 0.25 monolayers. We propose that acceleration in this system is strongly tied to the formation of AgS2 complexes at step edges. These complexes can transport Ag more efficiently than Ag adatoms (due to a lower diffusion barrier and comparable formation energy), hence leading to enhanced coarsening. The mobility of isolated sulfur on Ag(100) is very low, so that complex formation is kinetically-limited at low sulfur coverages, and thus enhancement is minimal. However, higher sulfur coverages force the population of sites adjacent to step edges, so that formation of the complex is no longer limited by diffusion of sulfur.

9:00am SS1-WeM4 Pt Terminated Mono- and Multilayer CuPt Alloys Supported on Ru(0001) Single Crystals as Model System for Core Shell Particles. A.K. Engelfeld, R.J. Behm, Ulm University, Germany.

A popular concept to improve the catalytic activity of metal particles is the utilization of core shell particles. This means that a metal/ally core is encapsulated by an additional metal in the shell of the particle. An interesting example has recently been published by R. Srivastava et al. [1], who found that Pt enclosed alloy particles containing Cu, Co and Pt, have a much better activity towards oxygen reduction than pure Pt. Whereas the prerequisite concept is that it is proven to work, the "optimum catalyst" for oxygen reduction has not yet been found.

For a better understanding of the structure and the formation of such core shell particles we prepare nanostructured planar model surfaces under well defined conditions, such as in ultra high vacuum (UHV). The surfaces consist of mono- and multilayer CuPt alloys on a Ru(0001) single crystal. They can be prepared by subsequent evaporation of the single metals and annealing at elevated temperatures. By this means the composition of the core can be well defined. The shell can be achieved by terminating the alloy by an additional Pt layer.

In this work we focus on the preparation of mono and bimetallic CuPt layers. They are characterized via STM to elucidate the morphology as well as the atom distribution. From the atom distribution within the alloy we will discuss the dominant factor for alloy formation, in view of the different size of the atoms and difference in their intermetallic bonding. Furthermore we will elucidate the surface segregation behavior of Pt in the bilayer alloy during the alloying process.


Faceted surfaces can provide unique opportunities to explore how catalytic reactions respond to changes in the catalyst surface structure. In this study, using LEED, XPS, AES, and STM, we report on how the presence of surface carbon significantly modifies the surface structure of a Re(11-21) single crystal and causes faceting, i.e., an initially planar Re(11-21) surface becomes "nano-textured" to expose new crystal faces and form "pyramids" on the nanometer scale. In addition to describing these nanostructures, we identify different states of surface carbon and describe their dependence on the coverage of carbon. We also utilized the faceted Re(11-21) surface containing these nanoscale pyramids to explore for unusual catalytic properties. Here, we describe the synthesis of a model electrocatalyst by deposition of one monolayer of Pt on the faceted C/Re(11-21) surface and investigation of its performance for the hydrogen evolution reaction (HER). This Pt ML-C/Re(11-21) surface displayed higher activity for the HER than pure Pt. In particular, this is particularly promising since Re is only one-fifth the price of Pt. This study is the first application of using a nanoscale faceted surface as a template for electrocatalyst synthesis, and illustrates the potential for other such investigations. It is also of interest to further explore the catalytic activity of such faceted surfaces for heterogeneous catalytic reactions, e.g., selective reduction of NOx with NH3 and selective oxidation of methanol, in order to understand the various effects of facet size, orientation, and low-coordination sites that are available for reaction.

W.C. and R.A.B. acknowledge support under DOE Contract No. DE-FG02-93ER14331. B.E.K. acknowledges support by NSF Grant No. CHE-1129417.

9:40am SS1-WeM6 Nucleation and Growth of Ag Islands on the (1x3)x3R30° Phase of Ag on Si(111). A. Belianinov, Iowa State University & Ames Laboratory - US DOE.

Using STM, we measure densities and characteristics of Ag islands that form on the (1x3)x3R30° Ag phase on Si(111), as a function of deposition temperature between 50 and 300 K. Assuming that Ag diffusion occurs via thermally-activated motion of single atoms between adjacent sites, the data can be explained as follows. At 50-125 K, islands are relatively small, and island density decreases only slightly with increasing temperature; the island density does not follow conventional Arrhenius scaling, probably due to sol-gel mixture and/or to increase in the absorption and consumption of Ag atoms. At higher temperatures there is a transition to conventional Arrhenius scaling, from which a diffusion barrier of 0.20-0.23 eV can be derived. At 300 K Ag atoms can travel for distances on the order of 1 μm, and they nucleate preferentially at step bunches. We have used this information to nanopattern the surface with Ag. We have done this by creating artificial defects in the ν3-Ag structure with the STM tip, and then subsequently depositing Ag at 300 K. This leads to strong preferential aggregation at these sites.

10:00am SS1-WeM9 Surface Diffusion of In and Sn on Si(001) at Room Temperature. N. To, S. Dobrin, J. Nogami, University of Toronto, Canada.

A self-aligning nanostencil mask was used to pattern circular features of tin and indium on an atomically clean Si(001) substrate. The shadow mask limited material deposition to below where the membrane was open, leaving adjacent areas of clean surface for material to diffuse. STM was used to study the room temperature surface diffusion of these metals in UHV and DFT was used to calculate relevant activation barriers. The comparison of these two metals is significant since they have the same atomic structure in the first atomic layer when grown on Si(001). The diffusion of tin is limited in uniform motion with indium and remains so independent of indium coverage. Indium forms unstable 3D islands that dissolve over time and contribute to the spreading of a single atomic layer thick film on the surrounding clean surface. The difference in behavior between the two metals can be attributed to the energy balance between 3D islands and the 2D wetting layers, as well as differences in activation energy for diffusion of atoms on top of the first atomic layer of metal. These results also show the potential for stencil patterning to provide insight into aspects of thin film growth.

11:20am SS1-WeM11 The Effect of Surface Fluoride on the Crystalization and Photocatalytic Activity of Titania. J.J. Brauer, G.J. Szulczewski, University of Alabama.

A two-step strategy to improve the photocatalytic activity of titania is reported. First, nitrogen doped titanium dioxide, denoted N-TiO2, has been synthesized by sol-gel methods and used to increase the absorption of visible radiation. Second, surface hydroxyl groups of the as-synthesized powders are replaced with fluoride ions. The two-step strategy gives independent control of "bulk" doping and surface modification. The as-synthesized and annealed powders were characterized by x-ray photoelectron spectroscopy, x-ray diffraction, diffuse reflection UV/Vis spectroscopy, IR spectroscopy, and scanning electron microscopy. Surface fluorination has two important consequences: it lowers the temperature to crystalize the as-synthesized powders into the photoactive anatase phase and improves the retention of nitrogen-dopants upon annealing. The photocatalytic of the titania powders were characterized by assessing the ability to degrade aqueous solutions of methylene blue, a common dye molecule, with visible radiation greater than 420 nm. The photodegradation experiments show that the rate of methylene blue decomposition follows the trend: F- N-doped TiO2 > N-TiO2 > undoped TiO2. A mechanism to explain the observed effects will be presented.


An aluminum alloy, AA2024-T351 (Al 2024), with a composition having a higher proportion of Cu to Mg (Cu/Mg = 3.7), which is used in the
aerospace industry, has been studied for corrosion simulation development. Results show that the Al 2024 alloy’s microstructure and corrosion behavior is quite different from that reported for Al 2024 alloy with higher Mg, a more common composition (normally Cu/Mg = 2.9) [1]. Characterization of the Al 2024 indicates a dominant presence of 2nd phase (AlCuFeMnSi) intermetallic particles (>70%), but that the S phase (Al1CuMg) and θ phase (Al1Cu) precipitate particles are much smaller in population (~30%). Microscale corrosion studies show that open circuit (OC) and potentiodynamic polarizations (PP) values extracted from micropolarization curves [2] measured for Al 2024 (Cu/Mg = 3.7) 2nd, S, or θ matrix phase have a large variation within a similar range of values. These results are in contrast to those already reported for Al 2024 with a more common composition (Cu/Mg = 2.9) where the electrochemical potential values separate with respect to the alloy phase and the microstructure shows S phase particle to be in the majority (60%) [2]. Nanoscale dispersoid particles and, to a much lesser degree, nanoscale 2nd phase particles were also found present throughout the matrix of the Al 2024 studied. The impact of chromite and cerium based conversion coatings, which inhibit corrosion, on the microscale corrosion properties of the Al 2024 alloy has been studied. Electrochemical current density data extracted from micropolarization curves measured for Al 2024 2nd, S, and θ matrix phase expressed to electrolytically, containing chromate (Na2Cr2O7) or cerium based (Ce(NO3)3) molecules show the chromate coating to be more efficient as an corrosion inhibitor. However, other cerium based molecules, such as cerium dibutylphosphate (Ce(dpdb)) and cerium chloride (CeCl3), which have shown more promising results [3], deserve further investigation. [1] R.G. Buchheit, R.P. Grant, P.F. Hlava, B. McKenzie, G.L. Zender, J. Electrochem. Soc. 144 (1997) 2621, [2] T. Suter, R.C. Allikire, J. Electrochem. Soc. 148 (2001) B36, [3] S.J. Garcia, T.H. Muster, Ö. Özkazana, N. Sherman, A.E. Hughes, H. Terryn, J.H.W. de Wita, J.M.C. Mol, Electrochim. Acta 55 (2010) 2457.

Surface Science Division
Room: 109 - Session SS2-WeM

Chemisorption on Metal & Oxide Nanoparticles
Moderator: B. Roldan Cuenya, University of Central Florida

8:00am SS2-WeM1 Temperature Program Deposition and X-ray Photoelectron Spectroscopy Study of Cu on CeO2/YSZ (111). J.C. Lofaro, Jr., Stony Brook University, M.G. White, Stony Brook University and Brookhaven National Laboratory

Energy research has increased in importance in the past decade due to our growing understanding of climate science and rising oil prices. Many catalysts center around expensive and rare transition metals, such as Pt and Pd, supported on oxide substrates. However, copper, a relatively cheap and abundant metal, has been used as a heterogeneous catalyst in industrial settings for various chemical processes.[1] Recent works have shown that copper nanoparticles supported on metal oxides (ZnO, CeO2, TiO2) have higher activity for the water gas shift reaction (WGSR) as well as other important chemical reactions when compared to their individual components. Understanding how these complex catalysts work on a fundamental level will allow for the design and implementation of more efficient and selective systems in the future. Here, using a homemade thermal evaporator, a model system of copper nanoparticles deposited on CeO2 films (200 nm thick) grown on YSZ (111) single crystals is used. X-ray photoelectron spectroscopy (XPS) is used to characterize the oxidation state of supported copper nanoparticles and temperature programmed desorption (TPD) is used to probe their reactivity and thermal stability. Copper coverages ranging from 0.25ML to 1ML are investigated. Carbon monoxide and water are used as probe molecules since they are the reactants involved in the WGSR. We have found that copper's stability is highly temperature dependent and have found evidence of its encapsulation by the support.

4. X. Zhao, J. A. Rodriguez, J. Hrbeck, M. Perez, Surface Science, 2005, 600, 229.

8:20am SS2-WeM2 X-ray Photoelectron Spectroscopy and Scanning Tunneling Microscopy Characterization of the Active Edge Sites of MoS2 Nanoclusters. A. Tuxen, S. Parsons, J. H. Wirthe, J. V. Lauritsen, Aarhus University, Denmark

The atomic and electronic structure of MoS2 nanoclusters is of considerable interest due to the catalytic application of MoS2 in e.g. hydrotreating catalysis of crude oil and in photocatalysts and hydrogen evolution reactions. Previous atom-resolved STM results have shown in great detail that both the overall morphology and in particular the edge structure of MoS2 nanoclusters, which are known to contain the most catalytically active sites for hydrotreating and H2 dissociation, adopt a structure which is very dependent on the conditions under which the cluster are kept. Under sulfiding conditions, atom-resolved STM images show that the MoS2 nanoclusters expose fully sulfide edges, whereas activation by H2 or mixed H2/H2S exposures show that sulfur vacancies and S-H form on the cluster edges. In this work, we present in-situ STM results of single-layer MoS2 nanoclusters with a well known structure. The XPS studies done on well-characterized samples reveal a set of edge specific core level shifts in the Mo3d photoemission peak that can be uniquely associated with the fully sulfide edges, edge with S vacancies or fully reduced edges. The XPS fingerprint thus allows us to dynamically follow changes between the catalytically active states of MoS2 when exposed to sulfiding of sulfuroductive conditions. Preliminary in-situ XPS results on the same MoS2 samples obtained under a 10^-7 torr H2 atmosphere on pressure transient pressure drop experiments show that the MoS2 nanoclusters in its active state. To dynamically follow such structural changes at the MoS2 edges induced e.g. by hydrotreating reaction conditions we have here combined high-resolution x-ray photoelectron spectroscopy (XPS) and scanning tunneling microscopy (STM) studies of single-layer MoS2 nanoclusters with a well known structure. The XPS studies thus successfully show that XPS in combination with STM can be successfully used as a tool to characterize the chemistry of highly dispersed active sites of well-defined nanoclusters, such as the active edges on MoS2.

9:20am SS2-WeM5 The Structure and Chemical Activity of Two-dimensional Gold Islands on Single-layer Graphene on Single-crystalline MoS2. L. Liu, Texas A&M University, Y. Xu, Oak Ridge National Laboratory, Z. Zhou, Texas A&M University, Q. Guo, Chinese Academy of Sciences, China, Z. Yan, Y. Yao, Texas A&M University, L. Semidey-Flecha, Oak Ridge National Laboratory, D.W. Goodman, Texas A&M University


INVITED Among the many factors that influence catalytic activity of supported metal clusters, the effects of size and charge state of the clusters are most frequently discussed, e.g. in CO oxidation over supported gold. In this contribution, I will present results of our recent experimental efforts to characterize metal clusters on single-crystalline MoS2 surfaces, using Au on MgO thin films as examples. As a combination of various surface science techniques including low-temperature scanning tunneling microscopy, infrared spectroscopy, X-ray photoelectron spectroscopy, and electron paramagnetic resonance, the properties of supported Au atoms and clusters were addressed in detail. Starting with the unusual chemisorption behavior of single Au atoms and the perfect MoS2 surface, I will move on and show how the charge of the Au particles on the MgO surface, in most cases is inferred from distinct CO chemisorption features, may be influenced by modifications of the oxide surface, e.g. by their interaction with defects or hydroxyl groups, or by the MgO film thickness.

11.0.2 of the Advanced Light Source Berkeley show a characteristic sequence of sulfur reduction steps on the catalytically interesting edges followed by decomposition of MoS2 at higher temperatures. The present studies thus successfully show that XPS in combination with STM can be successfully used as a tool to characterize the chemistry of highly dispersed active sites of well-defined nanoclusters, such as the active edges on MoS2.
A new experimental approach to mapping the relationship between composition and catalytic activity in metal alloy catalysts is presented. We apply this methodology to study $\text{H}_2$-$\text{D}_2$ exchange reaction on Pt-Pd and Pd-Cu alloys. The activity measurements were performed on thin alloy films deposited in such a way that the elemental composition varied continuously across the film. We refer to these films as composition spread alloy films (CSAFs). CSAFs were prepared by co-deposition of Pd, Cu and Au onto a molybdenum substrate under UHV conditions. The top-layer and near-surface composition of the CSAFs were determined by low-energy ion scattering (LEIS) and X-ray photoemission spectroscopy (XPS) respectively. The activity of the alloy films in the $\text{H}_2$-$\text{D}_2$ exchange reaction was analyzed at atmospheric pressure using a 100-channel glass microfluidic device. During the spatially-resolved activity measurements, the microfluidic device delivers reactant gases to a 10x10 array of measurement points on the CSAF surface covering an area of 1 cm², each measurement point corresponding to a different catalyst composition. After coming in contact with the catalyst, the gases are withdrawn for mass-spectrometric analysis through a separate set of channels. The activity comparison among different catalyst compositions can then be established by correlating the XPS/LEIS data with the product concentration in each channel of the microfluidic device.

The doping of metal oxides has been explored in several investigations with important implications for tailoring and understanding CeO$_2$ catalysts or supports for production of hydrogen. But, it is not known how the structure or electronic structure of the Cr dopants on the adsorption behaviour of the doped oxide support. We assign this change in the electronic structure to charge transfer processes from the Cr centres into the Au growth mode to charge transfer processes from the Cr centres into the Au growth mode. In this work we investigate the use of a low-temperature scanning tunneling microscope (LT-STM) tip to manipulate individual hydrogen adatoms on the rutile TiO$_2$(110) surface at 77 K. We show that applied voltage pulses are effective for transferring hydrogen atoms from the surface to the STM tip with single-atom control. This tip-induced ‘desorption’ is useful for unambiguously distinguishing between surface hydroxyls (OH$_t$) and bridging oxygen (O$_{\text{br}}$) vacancies—two common surface defects whose appearance in STM images is quite similar. In addition we show that individual atoms can be redeposited on the surface precisely at selected $\text{O}_2$ sites, allowing for the controlled preparation of arbitrary hydrogen adatom assemblies. Such control is a prerequisite for the investigation of the structural dependence of surface photo- and thermal reactivity at the single-molecule level.
Thin Film Division
Room: 110 - Session TF1+EM-WeM

ALD/MLD: Hybrid Organic Films
Moderator: Q. Peng, Duke University

8:00am TF1+EM-WeM1 Vapor-Phase Fabrication of Organic-Inorganic Hybrid Thin Films Using Molecular Layer Deposition with Atomic Layer Deposition, M.M. Sung, Hanyang University, Korea

We report a vapor phase deposition method of high quality organic thin films, called molecular layer deposition (MLD). MLD is a gas phase process analogous to ALD and also relies on sequential saturated surface reaction which results in the formation of a self-assembled monolayer in each sequence. In the MLD method, the high quality organic thin films can be quickly formed with monolayer precision under ALD conditions (temperature, pressure, etc.). The MLD method can be combined with ALD to take advantages of the possibility of obtaining organic–inorganic hybrid thin films. The advantages of the MLD technique combined with ALD include accurate control of film thickness, large-scale uniformity, excellent conformality, good reproducibility, multilayer processing capability, sharp interfaces, and excellent film qualities at relatively low temperatures. Additionally, a vast library of materials is accessible by ALD methods, ranging from single elements to compound semiconductors to oxides, nitriles, and sulfides. Therefore, the MLD method with ALD is an ideal fabrication technique for various organic-inorganic nanohybrid superlattices.

8:40am TF1+EM-WeM3 Metalcone and Metalcone/Metal Oxide Alloys Grown Using Atomic & Molecular Layer Deposition Techniques, B.H. Lee, V.R. Anderson, S.M. George, University of Colorado, Boulder

A new class of films known as the “metalcones” can be grown using atomic layer deposition (ALD) and molecular layer deposition (MLD) techniques. Metalcones are unique inorganic–organic materials derived from the sequential, self-limiting reactions of metal and organic alcohol precursors. The first metalcones were the “alucones” based on trimethylaluminum and ethylene glycol (EG) and the “zirconones” based on diethylihexic and EG. Other metalcones can be fabricated with various properties using different metal precursors together with organic alcohols. This talk reports new metalcones known as the “zirconones” using zirconium tert-butoxide (ZTB) and EG. In addition, this talk will discuss two new tunable ALD/MLD films based on alloys of alucone MLD/Al2O3 ALD and zirconone ZrO2 ALD. Zirconene MLD films were grown by using zirconium tert-butoxide (ZTB) and EG. MLD growth was observed at temperatures ranging from 105 to 195°C. In situ quartz crystal microbalance and ex situ X-ray reflectance (XRR) measurements confirmed that the growth of zirconene MLD versus number of ZTB/EG reaction cycles. Zirconene MLD growth rates decreased versus temperature and varied from 1.6 Å per cycle at 105°C to 0.3 Å per cycle at 195°C. A constant density of ~2.3 g/cm3 was measured for all growth temperatures. XRR measurement also showed that zirconene MLD films were very stable under ambient conditions. The metalcones will have useful mechanical, optical and electrical properties that can be tuned by growing alloys of the metalcones and their parent metal oxides. For example, this tuning allows the density of alucone MLD/Al2O3 ALD alloys to be varied from 1.6 g/cm3 to 3.0 g/cm3. The density of zirconene MLD/ZrO2 ALD alloys could also be varied from 2.3 g/cm3 to 4.0 g/cm3. The reflective index of zirconene MLD/ZrO2 ALD alloys was also tuned continuously between 1.63 and 1.86. These new metalcone materials provide a tool set for engineering the functional properties of thin films. These materials can be grown with atomic control of thickness and excellent conformality. The metalcones and metalcone alloys can also be thermally annealed to remove the organic constituent and create porous metal oxide films.

9:00am TF1+EM-WeM4 Sequential Vapor Infiltration and Atomic Layer Deposition on Surfactant Films for Mesoporous Metal Oxide Films, B. Gong, D. Kim, G.N. Parsons, North Carolina State University

Mesoporous materials are critical for applications such as catalyst support, energy storage and conversion, and chemical separations. Conventionally, solution based approaches are employed for the preparation of these materials, and amphiphilic molecules are widely used as templates to form well defined pore size and surface area. In these methods, the interaction between the hydrophilic block of the surfactant molecules and the metal oxide precursor drives the self-assembly of ordered microcrystalline metal oxide hybrid materials, and porous inorganic structures were recovered after removal of the organic template. Recently, this selective interaction was also discovered during vapor phase sequential vapor infiltration and atomic layer deposition (ALD) of metal oxide onto polymers, where precursor infusion and reaction depends strongly on the interaction between the precursor and polymer starting substrate. We therefore believe that sequential vapor infiltration or ALD on amphiphilic surfactant molecule films could also yield ordered inorganic/organic hybrid materials and porous metal oxides.

In this work, mesoporous aluminum oxide and titanium oxide materials were prepared by the vapor phase infiltration and ALD on two kinds of surfactants: the ionic cetyltrimethylammonium bromide (CTAB) and poly ethylene oxide based nonionic surfactant F127 (PEO-PPO-PEO). The nitrogen adsorption and desorption measurement was employed to measure the specific surface area and the pore size distribution of the resulted material. In-situ FTIR was used to monitor the chemistry change during the vapor infiltration, SEM TEM and XRD were used to characterize the structure and the morphology of the porous material.

We find that both sequential vapor infiltration and ALD produced mesoporous materials. However, the vapor phase infiltrated films show a higher density for the same size and surface area as compared to ALD produced material. These results demonstrate a new vapor phase approach for well defined mesoporous materials, which would potentially important for many advanced applications.


Organic electronics constructed on flexible substrates stand to revolutionize the display and lighting markets due to the low potential cost of manufacturing inherent to roll-to-roll manufacturing. A significant downside to organic electronics is their sensitivity to atmospheric oxygen and moisture. Various encapsulation techniques have been demonstrated, some of which have achieved Water Vapor Transmission Rates (WVTR) better than 1e-6 g/m2/d, which has been suggested to be the maximum allowable rate to give a 10-year usable lifetime of 10,000 hours for Organic Light Emitting Diodes (OLEDs). However, most of the encapsulation technologies that provide these low WVTRs lack the flexibility of the underlying plastic substrate or the organic electronics, limiting the actual flexibility of the final product.

Here we will report on the flexibility of aluminum-, hafnium-, and zirconium-based ALD films deposited on representative substrate material samples of polyethylene napthalate (PEN) at 115°C. Metal precursors used were trimethylaluminum (TMA), tetakis(dimethylamino)hafnium (TDMAH), and tetakis(dimethylamino)zirconium (TDMAZ). Water was used as the ALD co-reagent for depositing hybrid organic films. Various nanolaminate combinations of the inorganic/organic materials were also investigated for their response to various levels of strain. Inorganic Al2O3 films subjected to 2% strain were observed to begin cracking at thicknesses below 27µm. A 2% strain did not cause any cracking on organic films deposited with TMA and glycerol to the thickest film studied at 245µm. Organic films were always observed to be more flexible than inorganic films of the same thickness. Nanolaminates were observed to have flexibility intermediate to their pure inorganic and organic constituents.

WVTR measurements were performed on inorganic, organic, and nanolaminate aluminum films. A WVTR of 6.1e-6 g/m2/day was obtained for a 100nm inorganic Al2O3 film. A substantially more flexible film, TMA + glycerol film gave a WVTR of 9.2e-6 g/m2/day.

The technical issues associated with uniformly delivering glycerol to a 115°C ALD reactor will also be discussed.

9:40am TF1+EM-WeM6 Polymer Wires Containing Quantum Dots with Different Lengths Grown by Molecular Layer Deposition: Potential Applications to Sensitization in Photovoltaics, T. Yoshimura, R. Ebihara, A. Oshima, Tokyo University of Technology, Japan

Introduction: Molecular layer deposition (MLD) grows tailored polymer wires with designated molecular arrangements by connecting different kinds of molecules with monomolecular steps. We grew polymer wires with quantum dots (QDs), called “polymer multiple quantum dot (polymer MQD),” by MLD using three kinds of molecules, tetraphthalaldehyde (TPA), p-phenyleneediamine (PPDA) and oxalic dihydropyridine (ODH), and proposed their potential applications to sensitized photovoltaic devices.

Polymer MQDs: As a preliminary work, polymer MQDs of OTPTPT, OTPT and OT were grown. In OTPTPT, molecules are connected in a sequence of -ODH-TPA-PPDA-TPA-ODH--. The region

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between two ODHs is a QD of ~3-nm long. In OTPT, ~2-nm-long QDs are constructed with a molecular sequence of -ODH-TPA-PPDA-TPA-ODH-... In OT, ~0.8-nm-long QDs are constructed by connecting ODH and TPA alternately. A polymer MQD with three kinds of QDs, “3QD-MQD,” in which OT-like QD [OT], OTPT-like QD [OTPTP] and ODPDPA-like QD [OTPTP] are formed in one polymer wire, was grown with a sequence of -ODH-TPA-ODH-PPDA-TPA-ODH-PPDA-PPDA-TPA-PPDA-TPA-ODH-... For polymer MQD growth, we used the carrier-gas type MLD, where carrier gas of nitrogen was employed to give molecular gas blows onto substrates through valves for molecular gas switching.

[Light Absorption Spectra] Light absorption measurements revealed that the peak energy of the absorption spectra shifts toward the high energy side with decreasing the QD length, namely, in the order of OTPTPT, OTPT, and OT. This effect is attributed to the quantum confinement. The absorption spectrum of 3QD-MQD is broad extending from ~480 nm to ~300 nm, which is a superposition of the spectra of [OTPTPT], [OTPT], and [OT]. This result indicates that the polymer MQD with different QD lengths could be an efficient sensitizer. In addition, the molecular orbital calculation confirmed that the electron density is high in the QD regions of 3QD-MQD.

[Proposal of Polymer MQD Sensitization] In the polymer MQD sensitization, polymer MQD is on a ZnO surface as a sensitizer. QDs exhibit narrow absorption bands comparing with bulk semiconductors like Si due to their zero-dimensional characteristics. The absorption peak energy can be adjusted by QD lengths. Therefore, polymer MQD containing different-length QDs enables us to divide the wavelength region for light absorption into narrow regions. Each QD absorbs lights of wavelengths matched to its own energy gap, and injects the excited electrons into ZnO. This might suppress the energy loss arising from the heat generation in the light absorption process. The structure is regarded as a molecular tandem structure.

10:40am TF1+EM-WeM9 Hybrid Materials by Vapor Phase Infiltration, M. Knez, Max-Planck-Institut für Mikrostrukturphysik, Germany

Atomic layer deposition (ALD) is a thin film deposition technique which was developed in the 1970s to meet the needs for processing thin film electroluminescent displays (TFEL). Technically and chemically it is similar to chemical vapor deposition (CVD). However, in contrast to CVD, ALD incorporates as a specific feature the separation of the chemical reaction into two half-reactions. The ALD is not only able to perform thin film coatings of inorganic materials, but also allows coatings with organic-inorganic hybrid materials and, due to the separated exposure of the substrate to the precursors, infiltration of soft matter.

This talk will show top-down approaches to hybrid organic-inorganic and bio-inorganic materials obtained by infiltration with metals from the vapor phase. With tiny amounts of metals infiltrated, biological materials, such as spider silk or collagen, can positively change their mechanical properties after being treated with pulsed vapors of metal-organic precursors. The improvement of mechanical properties is related to changes in the molecular structure of the protein-based materials. However, not only biopolymers undergo changes after infiltration. With the example of some synthetic polymers, analogous routes to modify their mechanical properties will be shown.

11:20am TF1+EM-WeM11 Titanicone Molecular Layer Deposition Using TiCl4 and Sugar Alcohols and Porous TiO2 Films Produced by Annealing, R.A. Hall, A.I. Abdulagatov, S.M. George, University of Colorado, Boulder

Metalcone molecular layer deposition (MLD) can be performed using metal precursors and organic diols or triols. The first metalcone MLD films were the alucones and zircones grown using trimethylaluminium and diethylylzinc, respectively, with ethylene glycol (EG). In this work, we report the growth of titanicone MLD films using TiCl4 and two sugar alcohols: EG and glycerol (GL). The titanicones may have useful photocatalytic properties and may form valuable porous TiO2 frameworks upon annealing to remove the organic constituent. Titanicone films were grown using TiCl4 and EG at temperatures between 90-135°C. Quartz crystal microbalance (QCM) measurements observed a growth rate of ~83 ng/cm²-cycle from 90 to 115°C before decreasing significantly at 135°C. X-ray reflectivity (XRR) studies obtained a growth rate of 4.5 Å/cycle with a density of 1.84 g/cm³ at 115°C. Titanicone films were grown using TiCl4 and GL at higher temperatures between 130-210°C. The GL is believed to lead to more cross-linking that stabilizes the MLD film. QCM measurements observed growth rates that varied slightly from 49 ng/cm²-cycle at 130°C to 34 ng/cm²-cycle at 210°C. XRR studies yielded a growth rate of 2.2 Å/cycle at 150°C. QCM measurements revealed that the surface chemistry for titanicone MLD was self-limiting. XRR studies indicated that the titanicone films were stable in air. The titanicone films were absorptive in the ultraviolet and consistent with an optical bandgap of ~3.5 eV. Annealing the titanicone films removed the carbon component and yielded porous TiO2 films. Ultraviolet exposure also appeared to be able to produce porous TiO2 films. The ability to deposit conformal porous TiO2 films on high surface area substrates could produce “super” high surface area substrates. These substrates may serve as TiO2 scaffolds for dye-sensitized solar cells or photocatalytic membranes.
Exhibitor Technology Spotlight

Moderator: Langley


Granville-Phillips®, the Instrumentation Center for Brooks Automation Inc., recently introduced the VQM830 Vacuum Quality Measurement (VQM) System for gas analysis at high and ultra-high vacuum levels that is comprised of a high-speed Autoresonant Ion Trap Mass Spectrometry (ART MS) Sensor and High Performance VQM Controller. With advantages in speed, power consumption, simple calibration, accurate low mass reporting, and UHV performance, the VQM surpasses the traditional RGA products. In addition to explaining the advantages of this next generation mass spectrometer, new additions to the VQM family since initial product shipment will be covered.

12:40pm EW-WeL3 Combining NEG and Sputter Ion Pump Technologies to Meet the Challenges of UHV-XHV Systems, B. Garcia, F. Siviero, A. Conte, L. Viale, A. Bonucci, P. Manini, L. Caruso, A. Cadoppi, SAES Getters

Current UHV and XHV vacuum technology requires better vacuum, lower power consumption and smaller components. The NEXTorr pumping system meets these requirements by providing large pumping, good gas capacity, low power consumption and vibration free pumping in an unprecedented small size. Specifications and data will be presented to discuss advantages of the NEXTorr pumping system.

1:00pm EW-WeL4 EW - No Title - EW - No Title, S. Palmer, Agilent - Varian Vacuum Division

1:20pm EW-WeL5 Faster, Higher Resolution and More Accurate Imaging with the Cypher™ Atomic Force Microscope, K. Jones, Asylum Research

Within four years of the invention of the AFM, micro-fabricated cantilevers with integrated tips appeared, saving early practitioners from the joys of hand-assembling their cantilevers. However, even though many researchers soon understood the benefits of further miniaturization of the lever, standard commercial levers remained at the same 100 to 300 µm size for nearly the next two decades, in part because making a commercial instrument capable of using much smaller levers presented significant technical challenges. Within the past few years, commercial instruments like the Cypher AFM, from Asylum Research, have appeared which are capable of using cantilevers as small as 10 µm in length and with resonance frequencies 5 MHz and higher and those levers are now readily available. Small levers bring two major benefits to AFM. The first is much smaller thermal noise, enabling quieter force measurements and higher resolution imaging with Angstrom-scale cantilever amplitudes. The second is a major speed boost for AC modes (tapping, non-contact) in both air and liquid. When coupled with other instrumental improvements such as a high-speed scanner, the shorter levers allow scanning with good tracking at rates 20X to 40X what was possible with conventional levers. I will talk about the technical details behind both these improvements. I will also present images and movies highlighting the improvements, including images showing individual vacancy defects on crystals and movies showing fast scanning on polymers, crystals, and biological samples.


SpringerMaterials is an invaluable database for research both exploring and requiring vacuum science, as well as for the development of equipment that harnesses vacuum technology. Use this online resource to search for data on materials’ interactions with photons and electrons, molecular constants, coupling constants via nuclear magnetic resonance data, band structures via Photoelectron Spectroscopy, electronic transport, thermal and optical properties for a huge range of semiconductors, surface sciences and properties, characterization methods, metallic and organic thin films, particle detector systems, and so much more! With over 100,000 critically evaluated documents on properties of about 250,000 different substances, a robust metaccontent system and advanced search engine, you are sure to find what you are looking for vacuum research and development.
Magnetic and Electron Correlation Effects in Actinides and Rare Earths
Moderator: J.G. Tobin, Lawrence Livermore National Laboratory

2:00pm AC+MI-WeA1 Electronic Structure Theory of Complex Ordered Actinide Materials. P.M. Oppeneer, Uppsala University, Sweden

Actinide materials display many complex and correlated behaviors that originate from the special properties of the open f-shell atom embedded in a specific material’s environment. First-principles investigations provide a route to assess these anomalous phenomena in a materials specific way, providing direct, fundamental insight.

Here we consider recently obtained ab initio modeling results for actinide materials that are in the focus of current interest: actinide oxides, such as NpO2, PuO2, and higher-oxides, U3O8, and Np2O5, the hidden order (HO) material URu2Si2, and correlated plutonium compounds.

NpO2 is one of the very few materials in which complex multipolar order has been identified. Using the density-functional theory (DFT)-based LDA+U method we provide a first-principles theory of multipolar order and superexchange in NpO2. DFT+U calculations offer a precise microscopic description of the 3q-antiferro ordered phase. We find that the usually neglected higher-order multipoles (electric hexadecapoles and magnetic triakontadipoles) are at least equally significant as the electric quadrupoles and magnetic octupoles [1].

We further investigate light actinide oxides in higher oxidation states, such as U3O7, Pu2O7, and NpO7, for which non-collinear magnetic ordering is predicted. The possible further oxidation of PuO2 to PuO2+ is studied using DFT+U calculations in combination with x-ray absorption measurements [2].

The Pu monochalcogenides are intriguing materials, in which a correlated temperature gap develops, reminiscent of the behavior seen in Kondo insulators. Using dynamical mean field theory (DMFT) in comparison to LDA+U calculations, we show that dynamical self-energy fluctuations are important for the formation of an unusual gap. Static approximations to the self-energy as the LDA+U fail to provide a gap.

For URu2Si2 we report extensive electronic structure investigations [3], using full-potential LSDA, LSDA+U, and DMFT approaches to assess the origin of the hidden order. Our investigation show that the itinerant f-electron picture provides an excellent description of the materials properties of this fascinating compound. The Fermi surface which is crucial for the HO transition and the occurrence of unconventional superconductivity is accurately given. Our study points to the formation of long-lived spin fluctuations that are the driving quasiparticles for the HO.


4:00pm AC+MI-WeA7 Advanced x-ray Spectroscopies on 4f and 5f Systems. J. Bradley, M. Lipp, Lawrence Livermore National Laboratory, A. Sorini, SLAC National Accelerator Laboratory

Photon-in photon-out x-ray spectroscopies allow for a bulk-sensitive, high-pressure compatible look at rare earth and actinide electronic structure. The techniques couple to well-defined and meaningful quantum mechanical observables, including orbital orientation number and magnetic moment. These observables are key differentiators between theoretical treatments of strongly correlated systems, and they also provide meaningful and correct intuitive understanding. Here we will present an selection of measurements, both recent and high pressure, that exemplify the kind of insight these techniques can provide. In particular, we will address the question of rare earth volume collapse, where considerable controversy has existed between competing (Mott vs. Kondo) theoretical treatments.


Hard X-ray Photoelectron Spectroscopy (HAXPES) with 7.6 keV photons has been performed on single crystals of UPd3, UGe2, and USb2 at the European Synchrotron Radiation Facility, France. In the case of UPd3 we studied the localization transitions which occur under pressure. An 800 meV splitting within the Sb 3d core level was observed. The splitting of the Sb core levels is attributed to manifestations of two distinct Sb binding sites within the USb2 single crystal as supported by consideration of interatomic distances and enthalpy-of-formation. Photoelectron mean-free-path vs oxide layer thickness considerations were used to model the effectiveness of HAXPES for probing bulk features of in-air cleaved samples.

4:40pm AC+MI-WeA9 Actinide Dioxides under Pressure. L. Petit, Daresbury Laboratory, UK

The self-interaction corrected local spin density approximation is used to investigate the oxidation of actinide dioxides under pressure. The methodology enables us to determine the ground state valency configuration of 5f electrons and to study the localization delocalization transition that occurs under pressure. We argue that this delocalization facilitates the oxidation of the actinide dioxides and present results for the estimated transition pressures.


The electronic structure of Pu materials is directly tied to the details of the 5f electron bonding and hybridization. In compounds where direct 5f-5f bonding is negligible due to crystal structure and wavefunction overlap, hybridization is the key component for 5f electron influence on electronic properties. We examine two strongly correlated materials, PuCoGa5 and PuO2 that span the range of interesting materials from Mott insulator to heavy fermion superconductor. The synergy between synthesis, spectroscopy and modeling has provided a unique opportunity to explore details of the energy and crystal momentum dependence of Pu compound electronic structure through angle-resolved photoemission on single crystal samples and advanced modeling based on theories beyond density functional theory.

The strength of the 5f electron hybridization may be quantified through dispersion in 5f electron peaks from the angle-resolved photoemission. In the case of PuO2, we see over two eV of dispersion in the hybridized (O 2p - Pu 5f) valence band. For PuCoGa5, the quasiparticle peak at the Fermi energy shows 50 meV or more of dispersion in reciprocal space over a range covering slightly less than half the zone center to zone boundary. We are unable to follow the peak dispersion beyond this point as it crosses above the Fermi energy. These energy dispersions place significant constraints on models, which might be used to describe the electronic structure of these strongly correlated materials. For PuCoGa5, models, which place the 5f electrons in a localized configuration without significant hybridization, would not agree with the experimental results. In the case of PuO2, the dispersion measured in photoemission agrees well with the
hybrid functional calculations for PuO₂ and further support the increase in hybridization moving from ionic UO₂ to covalent PuO₂.

5:20pm AC+ML-WeA11 Structure and Magnetic Properties of Actinide-Based Thin Films. L. Havela, Charles University, Czech Republic, N.-T. Kim-Ngan, Pedagogical University Cracow, Poland, A. Adamska, Charles University, Czech Republic, A.G. Balogh, TU Darmstadt, Germany, T. Gouder, European Commission, JRC Institute for Transuranium Elements, Germany INVITED

Actinide-based sputter deposited films were so far used in the context of surface-science studies (such as [1]) and for exploration of electronic structure by photoelectron spectroscopy (e.g. [2,3]). In addition, sputter deposition was used in attempts to synthesize amorphous uranium alloys for ex-situ studies of magnetic properties. Such early (late 1980's) attempts in U, oxidation has to be suspected for films prepared in HV conditions.

diagnostics of the deposited material. Considering strong electropositivity ex-situ studies of magnetic properties. Such early (late 1980's) attempts in conditions (deposition rate, substrate type and temperature). Employing situ studies over months. The reason can be seen in pronounced UN films have a long-term stability, which allows comfortably to make exdiagnostics by XPS, Glancing Angle XRD, and RBS, it was established that [6]. We have used sputter deposition to investigate structure and magnetic properties of various U-based compounds as a function of deposition conditions (deposition rate, substrate type and temperature). Employing diagnostics by XPS, Glancing Angle XRD, and RBS, it was established that UN films have a long-term stability, which allows comfortably to make ex-situ studies over months. The reason can be seen in pronounced compressive residual strains, imposed during the deposition, which prevent progressing the surface oxidation into the bulk of several hundred nm thick films. Departing more from a fully crystalline state, the antiferromagnetism of UN is masked by a weak ferromagnetism, as usual for nanograined AF structures, and finally both moments and their order disappear [7]. Similar suppression of magnetism was found for ferromagnetic US [8].

Recently we undertook sputter-deposition experiments on Fe-rich U-Fe alloys derived from the Laves phase UFe₂, which combines the 3d and 5f magnetism in a compound with a relatively high Curie temperature (Tc = 162 K). An Fe-excess is expected to increase the Tc value markedly. Nanocrystalline material obtained up to the stoichiometry UFe₂, by sputtering, with the excessive Fe atoms entering the U sublattice has Tc enhanced up to 230-240 K. More Fe leads to the segregation of α-Fe. We succeeded to synthesize amorphous films by U and Fe co-sputtering, with stoichiometry up to UFe₂, Tc is enhanced up to 450 K in this case.


Applied Surface Science Division
Room: 102 - Session AS-WeA

Correlative Analysis - A Multi-technique Approach for Identification and Structure-Property Relationships Moderator: K. Artyushkova, The University of New Mexico

2:00pm AS-WeA1 Complementary Ultra Thin Film Analysis using Low Energy Ion Scattering (LEIS) and TOF-SIMS. T. Grebel, P. Bruneer, ION-TOF GmbH, Germany, N. Havецroft, ION-TOF USA, Inc., H. Brongersma, E. Niehuis, ION-TOF GmbH, Germany

Ultra-thin film structures have become increasingly important and simultaneously gained complexity with regard to the number of layers and the elemental composition. Understanding the processes occurring during deposition is crucial for improving the film quality. Especially during the first stages of film growth, analytical techniques with high surface sensitivity and good detection limits are required to study the growth process.

Low Energy Ion Scattering (LEIS) lends itself well to these tasks with its ultimate surface sensitivity of a single monolayer and detection limits of down to 10 ppm. This is accomplished by bombarding the surface with noble gas ions of a few keV and measuring the energy loss of the backscattered ions at a fixed scattering angle. The energy spectrum is converted into a mass spectrum of the elements present at the sample surface. The absence of matrix effects allows a straightforward quantification.

Besides the composition analysis of the outermost atomic layer, depth profiling is available via two distinct methods. Static depth profiling exploits the fact that ions scattered in deeper layers lose additional energy proportional to the penetration depth. As this process involves neutralization and re-ionization, intensities are lower than for the ions scattered at the surface. Thus, these ions can be distinguished, giving information about the elemental distribution in the first few nm in a nondestructive way. Alternatively, dynamic depth profiling is available by using a second, low energy sputter ion beam to erode the surface while recording surface spectra at different depths. This yields a quantitative, high depth resolution depth profile. By observing the change in the static in-depth signal during sputtering, the sputter rate can be intrinsically and continuously determined.

The unique advantages of LEIS complement established techniques like TOF-SIMS. The latter is often hampered by sputter transients at interfaces and difficult quantification especially of matrix species, but excels as far as the detection of trace elements or the gaining of chemical composition information is concerned. We applied both LEIS depth profiling modes to a number of thin film sample systems. Hereby we show the possibilities arising from each of the two modes, as well as from the combination with TOF-SIMS. Specifically, we worked on model samples relevant to the semiconductor industry (high-k, SiGe). Some of these samples were designed for studying the response function of the in-depth signal in order to improve the understanding and to allow the application to real-world samples, e.g. to correct for varying erosion rates.


Surface treatment of polymers produces materials that exhibit a wide range of surface compositions, properties and structures. The chemical and structural characteristics of these novel materials can be exploited for the fabrication of devices for bio-medical and electronics applications. Additionally, the wear-resistant properties of steel can be modified by coating the surface with a diamond-like carbon (DLC) film.

The combination of a variety of complementary surface-sensitive electron spectroscopies maximises the information available to the analyst for full quantification of surface characterisation of polymer surfaces and DLC films. The silicon content of a DLC film can affect its hardness, for example, and this is the ideal technique for chemical quantification of the silicon. The concentration of hydrogen in a DLC film also modifies its wear properties, but XPS cannot quantify this element. It is possible, however, to detect and quantify hydrogen using Reflective Electron Energy Loss Spectroscopy (REELS). When used together, XPS and REELS can provide a total quantification for polymer surfaces and DLC films. This presentation will show how Thermo Scientific tools can be used to investigate the chemistry and structure of various polymer and DLC samples. Chemical changes produced by surface treatments were examined by high energy resolution XPS and argon profiling (both monomer and gas cluster). Complementary REELS measurements were used to examine the level of carbon unsaturation at the uppermost surface of each sample and to detect and quantify hydrogen.

2:40pm AS-WeA3 Challenges Associated with Mathematically Correlating Data from Multiple Surface Characterization Techniques. K.G. Lloyd, D.J. Walls, L. Zhang, J.P. Wyre, DuPont Corporate Center for Analytical Sciences INVITED

There are now many examples of multivariate analysis of surface-specific technique data [1,2]. These include multivariate statistical methods such as Principal Components Analysis (PCA), Partial Least Squares (PLS), or Multivariate Curve Resolution (MCR) applied to so-called “hyperspectral” mapping data, in which hundreds of channels of spectral data are collected at each pixel of a two-dimensional pixel array spanning an area of interest. The idea of trying to mathematically correlate different sets of mapping data from the same area is not new [3], and falls under the broader category of ‘image fusion’ used in conjunction with remote sensing applications [4]. However, this approach is not prevalent in the surface science literature, with the notable exception of Falgun and Artushkov [5,6]. There are good reasons for this, from both the experimental and modeling perspectives. This talk will discuss the challenges associated with mathematically correlating spectroscopic and mapping data from multiple surface-specific techniques. Examples from the literature and the analytical lab will be discussed.

161 Wednesday Afternoon, November 2, 2011

4:00pm AS-WeA7 Multi-technique Characterization for Interfacial Analysis, Depth Profile and Chemical Imaging S.Y.N.T. Kuchibhatla, V. Shashikanth, B.W. Arey, C.M. Wang, Carnegie Mellon University, T. Prosa, Cameca Instruments Inc.

Nano尺度的界面分析在多种应用中包括燃料细胞、LEDs等。此外，我们将围绕EMSL、西北太平洋国家实验室来阐述感兴趣的影响因素的使用。例如，辐射等环境因素，特别是在核材料中具有重要意义。我们将在未来开发更多使用控制界面的材料。然而，基本的理解要求开发下一代具有检测到的界面的材料，这在很大程度上依赖于文献。在这个过程中，我们将其用于结合分析更多的技术，并且根据需要，选择新发展的技术和化学分析（更好的化学敏感性及高分辨率技术）来实现这一目标。原子探针成像（APT）是一种新发展技术，它以高分辨率扫描电子显微镜为基础，可以提供三维化学图像。

4:20pm AS-WeA8 Characterization of Lubricant Coated Cartridges Using Multiple Surface Analytical Techniques, X. Dong, Z. Xiao, C. Kemp, Eli Lilly and Company

润滑剂的性能评估是一个挑战，因为润滑剂的分布，包括那些使用塑料或玻璃材料的药物容器。克服这一挑战的方法之一是使用常规透射椭偏仪来考察润滑剂的分布。开发新型的微堆叠薄层 TLC 板，并且常常更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也更高。微堆叠薄层 TLC 板的性能通常在商业上可获得高性能 TLC 板，有时更高，也
In this work we report unusual pressure-composition isotherms for H2/Pd nanorings where a double plateau isotherms are observed during the charging process and a single plateau isotherm during the hydrogen desorbing on a solid support. Typically tens of secondary ions are ejected from each impact. They are identified with time-of-flight mass spectrometry and recorded individually. This approach reveals molecules co-located within the 10-20 nm diameter area of emission from one Au nano-impact. We demonstrate that the event-by-event bombardment-detection mode is sensitive to the chemical and/or physical nanoscale separation of molecular species. The performance is illustrated with the determination of the relative abundance of the oxide layer in the near surface of 50-100 nm nanoparticles; the nature and abundance of different nano-objects (5-20 nm in diameter) in mixtures of nano-sized solids; the composition of bio-objects such as a bacteriophage including the amino acids of the proteins surrounding the phage and the bases from the encapsulated DNA. The distinct feature of the nanoprobe technique presented here is in the detection of co-emitted ejecta from individual projectile impacts which allows to test chemical composition, in a nonimaging mode, yet at an exquisite level of spatial resolution. Moreover the co-emission of fragment and parent ions enhances the accuracy of molecular identification.

Support worked by NSF grant CHE-0750377

2:40pm BI+AS+NS+SS-WeA3 Strategies for Studying the Surface Chemistry of Engineered Nanoparticles with SIMS. C. Szakal; J. MCCAINTY, National Institute of Standards and Technology, K. Louis, R.J. Hamers, University of Wisconsin-Madison, R.D. Holbrook, National Institute of Standards and Technology

The environmental toxicity of engineered nanoparticles (ENPs) is of increasing importance as these materials become more widely used in manufacturing processes and consumer products. Nanoparticles have extremely high surface-to-volume ratios, which makes the surfaces more critical than their corresponding bulk materials in terms of reactivity, aggregation, and toxicity to various life forms. Therefore, it is critical that we develop methods to distinguish small chemical changes on nanoparticle surfaces in order to understand how these materials will interact outside of controlled laboratories. Conventional approaches of nanoparticle characterization have focused on high resolution morphological imaging (TEM, SEM) and elemental property measurements such as surface charge. However, chemical information is generally only inferred from these materials with most current methods. If it is possible to obtain both elemental and molecular information from ENP surfaces, we may be able to determine the eventual fate of ENPs in the environment.

We have developed a comprehensive approach for studying the surface chemistry of ENPs, including: (1) preparation of ENPs to controllably study desired variables, (2) development of methods such as time-of-flight secondary ion mass spectrometry (ToF-SIMS) and environmental scanning electron microscopy (ESEM) to probe small changes in ENP surface chemistry and/or aggregation, and (3) development of methods to improve the speed and reproducibility of ENP aggregation for batch studies. These approaches will be utilized as the basis of future toxicity studies of selected ecosystems.


Nanostuctured materials have been proposed as a solution for the control of biological interactions at interfaces with nanometer spatial resolution, but important challenges still remain to be addressed. Of the established fabrication techniques, electron beam lithography is expensive, and requires exposure under vacuum, while scanning probe methods are slow and (with few exceptions) do not permit fabrication over large areas. In contrast, interferometric lithography (IL) is a simple approach that uses inexpensive apparatus to fabricate sub-wavelength structures over macroscopic areas. When two coherent laser beams interfere, they yield a sinusoidal pattern of intensity that may be used to modify photoresists. Previously IL has been investigated extensively for semiconductor nanofabrication, but our recent data show that combined with self-assembled monolayer resists it provides a fast, simple method to create molecular nanostructures over macroscopic areas. Illustrations will be provided of bianofabrication using interferometric modification of protein-resistant (oligo ethylene glycol) functionalised surfaces, where feature sizes as small as 30 nm (3/8) have been achieved over square cm areas, and the controlled growth of protein-resistant brush structures from patterns of initiators for atom-transfer radical polymerisation. The

In this work we present a simple model, based on the observed heterogeneity of the nanorings (as seen in TEM) and by taking into account the defect induced lattice strain, that accounts for the unusual behavior of the observed isotherms.

4:00pm BI+AS+NS+SS-WeA7 Surface Functionalization and Analysis of Functional "Soft" Nanostuctures: From 2 to 3 Dimensions. H. Schönher, University of Siegen, Germany

INVITED

The local properties of soft matter, e.g. for the fabrication of functional biomimetic interfaces or nanoarchitectures, are of tremendous importance for ultimate functionality. In this presentation, the closely interrelated areas of surface chemical functionalization / engineering and analysis of properties will be discussed based on three key examples. These examples include: (i) synthesis and modification of polymer brushes with particular focus on the nanomechanical properties, (ii) ultra small diameter nanotubes obtained by the layer-by-layer assembly of polyelectrolytes inside a sacrificial porous template and (iii) block copolymer nanocapsules that are developed for advanced wound management. In all examples, confinement effects are expected to play a significant role in determining e.g. the mechanical properties of the microstructured components in terms of their macroscopic properties.

For thin polymer films (2D) the dependence of the mechanical properties on the film architecture was unraveled. Compared to spin-coated films, brushes synthesized on gold surface by surface initiated polymerization showed higher elastic moduli, which is attributed to entropy effects. Upon chemical crosslinking tunable elastic properties are obtained, which provides interesting pathways for the fabrication of defined cell - surface contacts.

Similarly important are defined nanoscale objects that can be obtained via the replication of small templates by the so-called layer by layer (LBL) deposition of polyelectrolytes (G. Decher Science 1997, 277, 227, LBL deposition in porous Anodic Aluminum Oxide (AAO) was very recently extended to the 100 nm length scale due to an alleged entropic barrier caused by adsorbed polyelectrolytes close to the pore orifice [Y. Cho et al. Small 2010, 6, 23, 2683.]. However, in contrast to this report, we show that the adsorption of polyelectrolytes on the top plane of the AAO and polymer sedimentation have been identified as main bottlenecks. Suppressing these processes enabled us to produce free standing polymer nanotubes with external diameters of < 55 nm.

Finally, first steps in the development of active nanocapsules filled with a reporter dye or an antimicrobial agent for applications in burn wound management will be presented. In particular the case of burn wounds and the devised biomimetic strategy of BacterioSafe will be introduced. Subsequently, the fabrication and characterization of a polystyrene-block-poly(acrylic acid) amphiphile-based model vesicle system, in particular the loading and release behavior and mechanical properties will be discussed.
fabrication of metallic nanostructures over macroscopic regions, including Ti structures as small as 35 nm, and gold nanostructures of controlled size and periodicity will also be demonstrated.

5:00pm BI+AS+NS+SS-WeA10 Functionalization of Mesoporous Silicon Biosensors to Achieve Tunable DNA Bioelectrode Density. J. Lawrie, R.R. Harl, B.R. Rogers, P. Laibinis, S.M. Weiss, Vanderbilt University

Porous silicon has become a widely studied material for sensing over the last decade based on its large surface to volume ratio and easily tunable morphology. With growing interest in the detection and analysis of genetic material, DNA oligos have become an increasingly important bio-recognition element in porous silicon and many other sensor platforms. As aptamers, nucleic acids serve as high affinity bioelectrodes to a wide range of small molecules and biological materials, opening up a number of potential applications in environmental science, chemical and biological defense, and medical diagnostics. In this work, tuning of the porous silicon surface chemistry is described. Controlling surface silanization, bioelectrode density, and bioelectrode charge and secondary structure enables the fabrication of reusable, label-free optical sensors toward specific nucleic acid targets. Detection limits in the nanomolar range have been demonstrated.

We have previously shown that in situ DNA synthesis via the phosphoramidite method in porous silicon produces high bioelectrode coverage for label-free optical biosensing applications. Low hybridization efficiency, despite high sensitivity, for such sensors indicated that tuning the receptor surface density could further improve detection limits. To modify surface receptor density, two-component trichlorosilane monolayers were deposited from solution onto porous silicon. One monolayer component remained active to phosphoramidite chemistry while the second component was inert. This method enabled a range of surface probe densities to be achieved and controlled via silanization conditions. Monolayer composition and DNA receptor density were verified using XPS, contact angle, and UV-Vis spectrophotometry. For a 16mer DNA oligo bound within a porous silicon waveguide, detection of the complementary target nucleic acid was maximized when 25% of the internal pore surface area was active toward DNA synthesis. Tuning surface DNA density increases sensitivity by a factor of 2-3. Label-free, target-specific detection of oligos was observed at concentrations of 25nM.

We will present results from hybridization efficiency studies in which DNA bioelectrode surface density, length, and secondary structure are varied. These parameters are vital to nucleic acid aptamer sensing strategies in label-free optical biosensors. Predicting appropriate receptor surface density for aptamer sensors based upon oligo sequences will provide advantages in achieving fast and sensitive waveguide sensors for detection in complex media.

Acknowledgements: This work is supported in part by the Army Research Office (W911NF-08-1-0200).


This presentation describes the fabrication and characterization of a novel composite membrane that consists of two types of nanoporous materials, namely, nanoporous gold leaf (NPGL) and nanoporous alumina, and a selective poly(perfluorohexyl norbornene) (pNBF6) polymer. Integration of the three materials is achieved by means of silane and thiol chemistry, and surface modification. The use of surface-initiated ring-opening metathesis polymerization (SI-ROMP). The former two provide functionalization of the nanoporous substrates, and the latter promotes the generation of the polymer film within and atop of the alumina-NPGL membrane. The synthetic process is versatile in that simultaneous or selective growth of the polymer film in each nanoporous layer is straightforward. The use of SI-ROMP allows tailoring of the extent of polymerization of pNBF6 throughout the structure by varying polymerization time. Advancing contact angle measurements show that the surface of these composite membranes exhibits both hydrophobic and oleophobic behavior. Scanning electron microscopy (SEM) images indicate that the thin polymer films cover the porous substrates entirely. Results to date on electrochemical impedance spectroscopy measurements revealed six deep levels in Mg$_{x}$Zn$_{1-x}$O samples while only three deep levels were observed in In$_{x}$Ga$_{1-x}$N samples. The frequency dependence of the capacitance caused by the high series resistance of the SDs required low frequency lock-in based capacitance measurements for accurate determination of the deep levels. (Donor-acceptor complexes contribute to near- and sub-surface carrier density. Defects also couple to nanostructures, which form spontaneously on ZnO polar surfaces and defects, such as ZnO clusters, can be excited by light.)

These results reveal the interplay between ZnO electronic defects, dopants, polarity, and surface nanostructure, and they highlight new ways to control ZnO Schottky barriers and doping.

2:40pm EM-WeA3 High Level of Mg Alloying Effects on the Deep Level Defects in Mg$_{x}$Zn$_{1-x}$O. E. Gur, The Ohio State University, G. Tabares, Ciudad Universitaria, Spain, A. Arehart, The Ohio State University, J.M. Chauveau, University of Nice Sophia Antipolis, France, A. Hierro, Ciudad Universitaria, Spain, S.A. Ringel, The Ohio State University

The Mg$_{x}$Zn$_{1-x}$O material system is very promising for ultraviolet emission and detection applications because of its direct band-gap, band-gap tunability, low lattice mismatch, and low polarization field. Using molecular beam epitaxy, a-plane 1.5 micron thick $n$-type Mg$_{x}$Zn$_{1-x}$O films ($x=0, 0.26, 0.36, 0.43, 0.52$) were grown on r-plane sapphire. As-epitaxially grown n-type ZnO Schottky contacts were used to perform deep level optical spectroscopy measurements (DLOS). The Schottky diodes (SD) exhibited rectification with low leakage currents and barrier heights between 1.00-1.19 eV from Mg$_{x}$Zn$_{1-x}$O to ZnO. The frequency dependence of the capacitance caused by the high series resistance of the SDs required low frequency lock-in based capacitance measurements for accurate determination of the deep levels. (Donor-acceptor complexes contribute to near- and sub-surface carrier density. Defects also couple to nanostructures, which form spontaneously on ZnO polar surfaces and defects, such as ZnO clusters, can be excited by light.)

These results reveal the interplay between ZnO electronic defects, dopants, polarity, and surface nanostructure, and they highlight new ways to control ZnO Schottky barriers and doping.

Electronic Materials and Processing Division Room: 210 - Session EM-WeA

Defects in Electronic Materials
Moderator: B.D. Schultz, University of California, Santa Barbara


INVITED Native point defects in semiconductors have until now not been considered a means to control formation or doping in devices and materials. In this work, tuning of the porous silicon surface chemistry is described. Controlling surface silanization, bioelectrode density, and bioelectrode charge and secondary structure enables the fabrication of reusable, label-free optical sensors toward specific nucleic acid targets. Detection limits in the nanomolar range have been demonstrated.

We have previously shown that in situ DNA synthesis via the phosphoramidite method in porous silicon produces high bioelectrode coverage for label-free optical biosensing applications. Low hybridization efficiency, despite high sensitivity, for such sensors indicated that tuning the receptor surface density could further improve detection limits. To modify surface receptor density, two-component trichlorosilane monolayers were deposited from solution onto porous silicon. One monolayer component remained active to phosphoramidite chemistry while the second component was inert. This method enabled a range of surface probe densities to be achieved and controlled via silanization conditions. Monolayer composition and DNA receptor density were verified using XPS, contact angle, and UV-Vis spectrophotometry. For a 16mer DNA oligo bound within a porous silicon waveguide, detection of the complementary target nucleic acid was maximized when 25% of the internal pore surface area was active toward DNA synthesis. Tuning surface DNA density increases sensitivity by a factor of 2-3. Label-free, target-specific detection of oligos was observed at concentrations of 25nM.

We will present results from hybridization efficiency studies in which DNA bioelectrode surface density, length, and secondary structure are varied. These parameters are vital to nucleic acid aptamer sensing strategies in label-free optical biosensors. Predicting appropriate receptor surface density for aptamer sensors based upon oligo sequences will provide advantages in achieving fast and sensitive waveguide sensors for detection in complex media.

Acknowledgements: This work is supported in part by the Army Research Office (W911NF-08-1-0200).

2:20pm EM-WeA2 Fabrication of Alumina Nanoporous Membranes. J. Kulesa, J. Pappas, S. L. Suib, The Ohio State University, R. Solomon, The Ohio State University, I. M. Zaworotko, J. Zhang, J. J. Song, Z. Qiang, Univ. of Dayton, D. C. Look, Air Force Research Lab

Fabrication of metal nanostructures over macroscopic regions, including Ti structures as small as 35 nm, and gold nanostructures of controlled size and periodicity will also be demonstrated.
SrTiO₃ and related perovskite materials are increasingly being utilized in a wide range of electrical applications. Furthermore, SrTiO₃ is often used as a substrate for growth of thin film structures. Thus, knowledge of its defect structure and, in particular, the dependence of its properties on understanding their impact on conductivity and optical phenomena. In the present work we utilize the limited penetration depth of x-rays to study the near-surface properties of SrTiO₃ substrates and epilayers and compare the results to analogous measurements using electrons. For an incidence angle of 10 degrees, the x-ray penetration depth varies between 22 and 207 nm as the energy changes from 600 to 1500 eV. For electrons the mean penetration depth varies from 3 to 90 nm as the energy changes from 0.5 to 5 keV. Thus, by obtaining x-ray excited optical luminescence (XEOL) and cathodoluminescence (CL) spectra as a function of x-ray or electron energy, it is possible to probe the near-surface region with nm-scale resolution. We will present energy-dependent XEOL and CL data from several SrTiO₃ samples. The luminescence spectra show features that can be assigned to Ti³⁺ (~1.6 eV) and oxygen-related (~1.9-2.9 eV) defects, but the relative intensities differ between XEOL and CL. We will discuss the results in terms of the respective mechanisms, previous CL studies [1,2] and theoretical models.

This work was performed at the Advanced Photon Source and was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Contract No. DE-AC02-06CH11357. We gratefully acknowledge Ohio State University support from Army Research Office Grant W911NF-10-1-0220


4:00pm EM-WeA7 Quantifying the Surface Generation Rate for Bulk Point Defects in TiO₂. K. Pungan-Okimoto, A. Hollister, P. Goral, E.G. Seebauer, University of Illinois at Urbana Champaign

The defect and surface properties of titania have been well-studied for its potential use in memory resistors and nanoelectronic gas sensors but the control of these properties in such fields has still not been fully realized. Such control of defect mobility and concentration in titania is essential for improving the operation of these technologies. Previous work in our research group has found that controlling the rutile (110) surface has opened up a new pathway for oxygen self-diffusion via an oxygen-interstitial mechanism in conditions where oxygen vacancies and titanium interstitials are thought to dominate. The present work employs the detailed diffusion-reaction network of point defects in rutile to simulate interstitial-mediated self-diffusion of oxygen and to calculate a generation rate of bulk point defects at the surface. The model explicitly incorporates gas adsorption onto the (110) surface, interstitial generation at the surface/near surface, and diffusion of the oxygen in an interstitial-mediated mechanism through the bulk. In addition, the model employs techniques drawn from systems engineering to estimate the key kinetic parameters. Simulated diffusion profiles were compared to experimental profiles obtained by exposing annealing single-crystal rutile to isotopically labeled oxygen gas and then measuring isotopic oxygen concentration with secondary ion mass spectrometry. The simulated diffusion profiles matched experimental results in verifying the oxygen interstitials do indeed mediate oxygen self-diffusion in our experiment. More importantly, our model allows us to calculate the oxygen interstitial generation rate at the (110) surface, a first for titania.

4:20pm EM-WeA8 Metastable Defects and Recoverable Degradation in InAs/AlSb HEMTs. X. Shen, S. DasGupta, R.A. Reed, R.D. Schrinpf, D.M. Fleetwood, S.T. Pantelides, Vanderbilt University

It is known that some defects undergo a structural transformation to a metastable configuration when capturing a charge carrier. However, their role in affecting device performance are not widely recognized. Here we show that metastable oxygen defects can cause recoverable degradation of InAs/AlSb HEMTs (high-electron mobility transistors).

Recently, we performed electrical stress tests on InAs/AlSb HEMTs and observed a recoverable degradation in some of the devices. The degradation is manifested as negative shifts of the transconductance peak and threshold voltage, which nearly completely recover after two days at room temperature. The recoverable nature of the degradation suggests that metastable defects are involved. The threshold shift indicates an increase of donor concentration or a decrease of acceptor concentration in the device.
boundaries at the distances where defect-defect and defect-GB interactions are found to be significant. GB-558 consists of pentagon and octagon pairs, whereas in GB-575 it is formed by repeated pentagon-heptagon pairs. We show that recombination of vacancies and interstitials can occur at grain boundaries resulting in efficient annealing of defects. The compression (stretch) for GB-558 is up to 3%, while in GB-575, the stretch reaches 9% at the C-C bonds between pentagon and heptagon or between hexagon and heptagon. We find that single vacancies can diffuse towards a grain boundary, attach to a pentagon, then merge into the grain boundary to release the compressive strain and cause local reconstruction. Molecular dynamics simulations show that if a single vacancy exits at 1nm from the grain boundary GB-558, it diffuses towards the grain boundary and attaches at the pentagon ring after 160 ps at 3000 K. It takes another 40 ps to merge into the grain boundary. Similar phenomena occur in GB-575. Stretched C-C bonds on the pentagon, on the other hand, accumulate interstitials. An interstitial that was initially positioned 7 Å above the graphene sheet, was adsorbed on the grain boundary. Once the vacancy coalesces into the grain boundary, the carbon adatom fills the reconstructed vacancy and recovers the original grain boundary structure. This recombination takes only ~ 0.5 ns at 2000 K. The results suggest a new mechanism of defect annealing with a two-step procedure; i.e. point defects (single vacancies and interstitials) diffuse and recombine locally at grain boundaries. This work was supported by DTRA Grant No. HDTRA1-10-1-0016, the US Department of Energy, Basic energy Sciences, and the William A. and Nancy F. McMinn Endowment at Vanderbilt University. The calculations were performed at ORNL’s Center for Computational Sciences and the Air Force Research Laboratory DoD Supercomputing Resource Center.

5:20pm EM-WeA11 X-ray Induced Defect Formation in Graphene. E.X. Zhang, A.K.M. Newaz, S. Bhaduri, M.L. Alles, D.M. Fleetwood, R.D. Schrimpf, K. Bolotin, R.A. Reed, R.A. Weller, S.M. Weiss, S.T. Pantelides, Vanderbilt University. Graphene is a truly two-dimensional material that consists of carbon atoms connected by sp2 bonding and arranged in a honeycomb lattice. Due to its unique properties and potential applications in future electronic devices, graphene has received much attention recently from the scientific community. Initial studies of the effects of electron-beam and proton irradiation have been performed on graphene materials, but there remain significant questions about the nature of the conductivity and of the defects. In this work, the responses of graphene materials to 10-keV x-ray radiation are evaluated with Raman spectroscopy. A defect related peak (D-peak) appears (~1345 cm⁻¹) after x-ray irradiation in air; the intensity of the D-peak increases with increasing total dose, and decreases with post-irradiation vacuum annealing at 350 °C. (See supplemental figures.) These results suggest that new defects can be created and/or that weak bonds at defect precursor sites can be scissored and decorated with impurities (e.g., H2, O2, during x-ray exposure). The charge neutral point of graphene transistors made from similar material shifts positively with increasing x-ray dose, suggesting that similar defects may contribute to shifts in the Dirac point and degradation in the conductivity.

It has been shown previously that ozone can react with C-C bonds to degrade graphene layers. We have measured significant ozone generation during x-ray irradiation, suggesting that the x-rays may not directly create the defects sensed by Raman, but that x-ray generated ozone may lead to at least some of the observed defects. However, it is likely that there is an additional source of degradation, since in previous work the ozone reaction typically has been shown to lead to irreversible degradation, but we find that vacuum annealing at elevated temperature leads to partial recovery. Hydrogen annealing treatments of graphene have shown reversible degradation. Thus, it seems likely that hydrogen released by x-ray exposure in the SiO2 substrate reacts with defect precursors (e.g., weak bond sites) in the graphene. Hydrogen reactions with C-C bonds can enhance the intervalley scattering, thereby increasing the D line intensity. Much of the hydrogen may be driven out by annealing in vacuum at 350 °C, leading to recovery of the degradation, consistent with our experimental results. This work was supported by the DTRA Basic Research Program through Grants HDTRA1-10-1-0016 and HDTRA1-10-1-0041.

5:40pm EM-WeA12 Influence of Point Defects on the Properties of Highly Mismatched Alloys. R.L. Field III, T. Dannecker, Y. Jin, C. Kuziuk, C. Udovic, University of Michigan. It has been suggested that alloy films composed of highly immiscible solute atoms in a solvent, termed “highly-mismatched alloys” (HMAs), are promising for energy conversion devices due to their ability to efficiently absorb light and heat, and to subsequently transport charge carriers. The properties of HMAs are often described with models focusing on the influence of individual solute atoms, assuming that all solute atoms “see” the same atomic environment. In the case of GaAsN alloys, the single local environment models predict a N composition-dependence of the energy band gap which agrees qualitatively with experiment. However, such models do not quantitatively explain several extraordinary electronic and optical properties. In this talk, I will discuss our investigations of the growth [1], structure [2-3], and properties [4-6] of GaAsN alloys. I will focus on correlations between the presence of N-N, As-N, and N-Si pairs [2-5-6] and extraordinary physical phenomena, including non-monotonic composition-dependent effective masses [7] and persistent photoconductivity [8]. I will also discuss our preliminary work on complementary alloys in the GaAsBi system.

layers deposited on top of the completed CIGS solar cells we have demonstrated that initially 11.1% efficient CIGS solar cells lose less than 7% of this peak efficiency and still exhibit efficiencies greater than 10% (factor of 10%) after 150 hours at 85 °C and 85% relative humidity. In comparison, under identical test conditions, the solar cells without the SnO2 layer lost nearly 80% of their initial efficiency within 24 hours after commencing the test. We studied the effects of deposition conditions and film thickness for different film structures on the solar cell stability in damp-heat tests. The deposited SnO2 films are amorphous when deposited at room temperature or when the films are thin, but show increased crystallinity for thicker films and films deposited at 150 °C. We found that solar cells coated with morphopolymorph SnO2 films exhibit better heat stability than those coated with polycrystalline films. By polycrystalline we mean films that consist of nanocrystalline SnO2 embedded in amorphous SnO2. We attribute this difference to the lack of grain boundary diffusion in polymorphous SnO2 films. Replacing the crystalline ZnO window layer with a SnO2 film can provide further protection of the CIGS solar cells. We demonstrate a 8.2±0.2% efficient CIGS solar cell with a SnO2 window layer. Same solar cell fabrication process and CIGS film with ZnO window layer resulted in 8.2±0.6% overall efficiency. The open circuit voltages of the two cells were nearly the same indicating that the band alignment with the SnO2 film is suitable for CIGS. These SnO2 films were deposited using magnetron sputtering at 5 mTorr and 150-250 W RF power using Ar as the sputtering gas without substrate heating.


A key criteria for achieving terawatt-compatible photovoltaic (PV) technology is the ability to fabricate high power conversion efficiency (>10%) solar cells using a low-cost process (< $1/watt) and readily available constituents. This talk will discuss recent developments that have enabled the demonstration of the first CZTSSe solar cells with certified power conversion efficiencies of over 10%, using a glass/Mo/CZTSSe/Cds/ZnO/ITO structure and a simple liquid-based deposition approach. We also present a device characterization study that compares the CZTSSe devices with higher-performing CIGS analogs, elucidating some of the key performance bottlenecks in CZTSSe cells, including dominant buffer-absorber recombination, high series resistance and a relatively short minority carrier lifetime. These studies help to elucidate key areas for improvement for the CZTSSe cells in the effort to develop a high performance pervasive technology.

4:00pm EN1+TF-WeA7 Solar Cells from Colloidal Dispersions of Cu2ZnSnS4 Nanocrystals. A. Khare, Y. Li, B. Chernomordik, B.S. Tousan, A.W. Willis, D.J. Norris, E.S. Aydil, University of Minnesota

Copper zinc tin sulfide (Cu2ZnSnS4, or CZT) is emerging as a promising photovoltaic material for thin film solar cells. CZT has a band gap of ~1.4 eV, the ideal value for converting the maximum amount of energy from the solar spectrum. In addition, CZT has a high absorption coefficient (~105 cm−1 in the visible region of the electromagnetic spectrum) and its constituent elements are all abundant in the earth’s crust and are non-toxic. We have synthesized CZTS nanocrystals from metal dithiocarbamate complexes. The diameter of the nanocrystals can be varied from 2-7 nm by changing the temperature and synthesis time. A suite of methods including Raman spectroscopy, optical absorption, electron energy loss spectroscopy and X-ray diffraction were used to characterize these nanocrystals and show that they are phase-pure CZTS. Nanocrystals with diameters less than 3 nm exhibit quantum confinement. These quantum dots were used to assemble quantum dot solar cells. In a second approach to making solar cells, thin films of CZTS nanocrystals are annealed to form thin films with large grains. Nanocrystals melt at temperatures much less than the bulk temperature and recrystallize to yield larger CZTS grains. These films were then used for making conventional thin film solar cells.


Iron pyrite (FeS2) has been considered one of promising materials for use in solar cells due to its large absorption coefficient, suitable band gap and elemental abundance. In-lab X-ray photoelectron spectroscopy and tunable-energy synchrotron X-ray photoelectron spectroscopy were used to explore the surface structure of pyrite thin films by metal organic chemical vapor deposition (MOCVD). The influence of sodium diffusion on the growth of pyrite thin films on glass substrates was examined. By using synchrotron X-ray photoelectron spectroscopy, the different types of sulfur chemical states on the surface of pyrite thin films were resolved. The mechanism of pyrite oxidation after exposure to different oxidizing environments indicated that the surface monosulfide species were oxidized first. In addition, the bands of pyrite thin films are modified by combining valence band spectroscopy with X-ray absorption spectroscopy compared to conventional ultraviolet-visible absorption spectroscopy. The discrepancy between the two measurement techniques will be discussed.

4:40pm EN1+TF-WeA9 Effect of the Use of a c-CdS Nanocrystalline Layer on the Photovoltaic Characteristics of the Screen Printed CdS/CdTe Heterostructure. L.G. Rangel-Chavez, UAM-Azcapotzalco, Mexico, M. Garcia-Aguirre, F.A. Cuevas-Ortiz, Cinvestav-IPN, Mexico, M.I. Neria-Gonzalez, Instituto Tecnologico de Estudios Superiores de Ecatepec, Mexico, M.A. Melendez-Lira, Cinvestav-IPN, Mexico

The fact that the efficiency for the CdS/CdTe system has remained without any change for several years until now requires employing new approaches in order to increase it. In this work it is reported the effect on the photovoltaic characteristics of the CdS/CdTe system by employing as substrate CdS the one obtained by pyrolysis heat treatment of heavy metals in residual waters. By using a novel sulfate-reducing bacterium (Desulfovibrio alakensis 6SR) particles of the CdS semiconductor are obtained. Structural and optical characterization by XRD, UV-vis, Raman and photoluminescence spectroscopies show that nanoparticles of cubic CdS are obtained. The nanoparticles of cubic CdS are deposited on CdTe films by screen printing and then subject to a mild thermal treatment. It is expected that the employment of CdS with cubic rather than hexagonal crystal structure will reduce the number of cadmium vacancies if combing the transport of minority carriers through it. We will report the effects on the spectral response and filling factor obtained for the CdS/CdTe heterostructure by employing nanostructured cubic CdS as compared to those obtained employing hexagonal CdS.

* Work partially supported by CONACyT-Mexico.

5:00pm EN1+TF-WeA10 Surface Structure and Chemistry of AgInSe2 Studied by Scanning Tunneling Microscopy. P. Peña Martín, J.W. Lyding, A. Rockett, University of Illinois at Urbana-Champaign

Chalcopryte semiconductors such as (Cu2Ag)(In,Ga)(S,Se)2 show great promise in thin film solar cells as they exhibit high optical absorption and excellent performance even as polycrystalline layers. The alloy AgInSe2 (AIS) is a promising candidate for solar applications, as it has a nearly ideal energy gap (1.2 eV), high absorption coefficient, and shows sharper photoluminescence emissions than do the Cu-containing alloys. The surface of the material forms the heterojunction and determines many aspects of device performance. These semiconductors also contain a large number of intrinsic point defects, which are probably responsible for minority carrier recombination in the depletion region. Therefore understanding the surface and near surface nanostructure and nanochemistry are critical to device optimization. To characterize the structure and chemistry near the surface, we used ultra high vacuum scanning tunneling microscopy (UHV-STM) to obtain atomic-scale topographic and electrical information. We report the first atomically-resolved STM images and current-voltage profile measurements of AIS. Epitaxial layers were grown on epi-ready substrates of p-GaAs(111)A by hybrid technique, in which Ag and In are sputtered concurrently with effusion cell evaporation of Se. The resulting film was transported to a N2 ambient to the STM laboratory, mounted on normal lab air as quickly as possible, and introduced to vacuum, with a total film pressure of ~10−9 Torr. STM images were acquired with a Pt/Ir tip operating in the constant current mode, with topograph and contour images acquired with the approach curve set to 15 mV and 10−9 Torr. Topographic images reveal atomically-resolved regions of the surface layer by layer exposure. We attribute these to the polar metal-terrace (0112) plane, 0.34 and 0.36 nm along rows oriented 60° from each other. This indicates that the surface does not reconstruct, unless by swapping one type of metal atom for another. Current-voltage spectra confirm that the material exhibits n-type behavior with an energy gap close to the bulk value of 1.2 eV. Some regions exhibit more fluctuations in the bandgap for a series of
Copper zinc tin sulfide (Cu2ZnSnS4 or CZTS) is a potential candidate for next generation thin film solar cells because it contains abundant and nontoxic elements and exhibits high light absorption. Thin films of CZTS are typically synthesized by sulfidizing a stack of zinc, copper and tin films. In addition to CZTS, a variety of binary and ternary metal sulfides can form and distinguishing among phases with similar crystal structure can be difficult. Herein, we show that confocal Raman spectroscopy and imaging can distinguish between CZTS and the other binary and ternary sulfides [1]. Specifically, Raman spectroscopy was used to detect and distinguish between CZTS (338 cm⁻¹), Cu2SnS3 (298 cm⁻¹) and Cu5SnS6 (318 cm⁻¹) phases through their characteristic scattering peaks. Confocal Raman spectroscopy was then used to image the distribution of coexisting phases and is demonstrated to be a useful tool for examining the heterogeneity of CZTS films. We show that, during sulfidation of a zinc/copper/tin film stack, ternary sulfides of copper and tin, such as Cu5SnS6, form first and are then converted to CZTS. The reason for formation of Cu5SnS6 as an intermidary to CZTS is the strong tendency of copper and tin to form intermetallic alloys upon evaporation. These alloys sulfidize and form copper tin sulfides first and then eventually convert to CZTS in the presence of zinc. As a consequence, films sulfidized for 8 hours at 400 °C contain both CZTS and Cu5SnS6 while films sulfidized at 500 °C contain nearly phase-pure CZTS. In addition, using Cu Ka radiation, we identify three CZTS X-ray diffraction peaks at 37.1° ([202]), 38.8° ([211]) and 44.9° ([020] and [213]), which are absent in ZnS and very weak in Cu5SnS6.


Thin film solar cells based on Cu(In,Ga)(S,Se)2 and CdTe have demonstrated significant improvement in last few years and they are also being transferred to production level. However, both CIGS and CdTe based thin film solar cells are hindered by potential environmental hazard issues and scarcity issues associated with the constituent elements, mainly Te, In, Ga and to some extent Se. Recent research trends are moving towards finding alternatives based on earth-abundant and non-toxic elements. An alternative material Cu(Zn,Sn)(S,Se)2 is being explored by the thin film photovoltaics community which contains earth abundant materials like Zn and Sn. CZTS structure can be derived from CuInS2 chalcopyrite structure by replacing one half of the constituent indium atoms by zinc and other half by tin. The resulting bandgap varies in the range of 0.8 eV for a selenium structure to 1.2 eV for a sulfide structure. Copper and zinc atoms in earth’s crust are 50 and 260 ppm respectively and while abundance of zinc and tin is 75 and 2.2 ppm in respectively. As compared to this, indium in earth’s crust is 0.049 ppm and selenium 0.05 ppm. CZTS also has a large absorption coefficient in the order of 10⁶ cm⁻¹. There are vacuum-based as well as non-vacuum-based approaches for the synthesis of CZTS solar cells. Among vacuum based approaches, generally sulfurization in H2S atmosphere is carried out on sputtered CuZnSn precursors. However, there are toxicity issues involved with the use of H2S gas. We subscribe to the philosophy of sticking to a non-toxic approach of synthesizing thin film solar cells and for the first ever time introduced a non-toxic sulfur precursor called di-tertiray-butyl-disulfide (TBDS) for the sulfurization of CZT layer. Initial results are quite encouraging and device quality CZTS cells are already being synthesized. The CZTS cells are characterized using materials characterization techniques such as SEM, XRD, XPS and photovoltaic parameters are being extracted under AM 1.5 conditions and also analyzed using quantum efficiency measurement. This is the very first attempt to successfully synthesizes CZTS solar cells using a non-toxic sulfur source.
Titanium dioxide (TiO$_2$) thanks to its interesting properties as nontoxicity, low cost and high chemical stability, has been extensively investigated for photocatalytic applications. The degradation properties of TiO$_2$ nanofibers (NFs) have been shown interesting degradation properties, which are much better than those obtained using other methods. In the recent research is to be able to prepare photocatalyst which can be active by absorbing the visible light, in order to increase the application of its nanoscale properties. The high surface area and the nanoscale dimension, capable of light absorption in a small fraction of the solar spectrum can be used to promote the light conversion of the solar spectrum into other forms such as heat or electricity. In this paper we will show the photocatalytic results obtained depositing TiO$_2$ N-doped nanofibers on conductive substrates, using different deposition and post-deposition parameters (solution composition, annealing temperature and atmosphere) and have been characterized by X-ray photoemission spectroscopy (XPS), X-ray diffraction (XRD) and Secondary Electron Microscopy (SEM). The photocatalytic properties have been studied recording the variation of the optical absorption of MB when the sample is illuminated by an halogen lamp (visible spectrum).

The annealing process determines a partial loss of nitrogen and the formation of the Anatase crystalline phase. The TiO$_2$ N-doped NFs have shown interesting degradation properties, which are much better than those observed when TiO$_2$ NFs are used.

Fabrication of Palladium Nanoscale Structures for Hydrogen Sensing Applications, D. Rodríguez-Vindas, University of Puerto Rico at Rio Piedras, C. Ortiz, V. Panotian, University of Puerto Rico at Cayey, W. Otaño, University of Puerto Rico at Cayey and Institute for Functional Nanomaterials. Palladium (Pd) metal is one of the most prominent materials studied for the detection of hydrogen gas. Hydrogen rapidly dissociates on its surface and diffuses into subsurface layers forming palladium hydride with consequent changes in optical, mechanical and electrical properties that are easily detected. Materials with nanoscale morphologies are promising to improve sensor performance. They provide large surface areas for adsorption, and a smaller crystallite size reducing the time needed for “bulk” diffusion. The amount of sites available for hydrogen adsorption per Pd atom is also higher in the surface and subsurface layers resulting in higher sensitivity. In this project, Pd nanoribbons and nanoshells are prepared by magnetron sputtering deposition on top of the mat of polymer fibers, which act as a template that shapes the morphology of the palladium being deposited while providing support to the metallic scaffold that is created. Sputtering is a line-of-sight deposition process and the fibers become a variable-angle-substrate for the incoming Pd flux. A larger amount of palladium is deposited on top of the fiber where the incoming flux is perpendicular to the surface compared to the sides where the flux is incident at a glancing angle. The top and sides of the fibers shadow their bottom parts closer to the substrate preventing any substantial deposition there. The end result of the deposition is the formation of Pd nanostuctures, thicker in the middle region than at the edges, with a large void network. Process parameters such as deposition time, sputtering pressure, and power can be used to produce nanostuctures with different thicknesses and morphologies. The high sensitivity and response time shown to 1% or less of hydrogen in nitrogen is understood to result from the reduced dimensions combined with this unique nanostucture. A description will be given of the conductance changes with hydrogen concentration as result of the competing mechanisms of percolation and scattering. This work shows the use of nanotechnology combined with microporous processing approaches to introduce new sensor geometries with different behavior and morphology than simple thin film or wire approaches.
However for graphitization in a Si-rich environment we observe a 3×3 interface in agreement with other groups. C-face depends on the means of graphene formation. For graphitization in a disilane environment. It is found that different interface structures occur which indicates the existence of localized electronic states originated from images of the domain boundary, the chain of protrusions was observed, the pentagon or heptagon structures at the boundary.


4:40pm GR-WeA9 Thermionic Emission of Graphene on Metal Surfaces. E. Starodub, N.C. Bartelt, K. McCarty, Sandia National Laboratories. We employ low-energy electron microscopy (LEEM) to study the thermionic emission of graphene on representative metals, Ru(0001) and Ir(111). In traditional LEEM, an image is produced using low-energy electrons reflected from a surface. The unique capabilities of LEEM allow us to image directly electrons thermionically emitted from graphene at high temperatures, above 1100K, using the same imaging optics. Due to the strongly anisotropic nature of graphene, we can observe rotational symmetry across the high symmetry directions. However, this broken mirror symmetry will be very difficult to detect using laterally averaging techniques such as SXRD because of the presence of two chiral enantiomers and two terminations of the hep substrate. Using LEEM and µLEED we have successfully obtained diffraction information from an area with a diameter of 250 nm within a single surface termination. While no evidence is found with such high spatial resolution diffraction measurements that confirm chiral features within the supercell unit cell, we do in fact observe rotation or twist of the mirror-like superposition over long length scales. Moiré twist causes the entire ensemble of satellite diffraction spots along each integer order spot to rotate as a group about their respective stationary foci. When the µLEED beam is scanned across the surface, the rotation angle undulates randomly. The data also suggest that the ground state configuration comprises a graphene layer that is slightly rotated with respect to the underlying substrate surface. Conceivably, the orientational variations that are observed here are due to sub-micron length scales might produce a signature in laterally averaging measurements that could be construed as evidence of chiral features on the short length scale within the unit cell. [1] Emtsev et al., Phys. Rev. B 77, 155303 (2008). [2] Varchon et al., Phys. Rev. Lett 99, 126805 (2007). [3] Ouda et al., Phys. Rev. B 82, 041411 (2010). [4] Hibino et al., Phys. Rev. B 77, 075413 (2008).
which enables the imaging of states within a few meV of the Fermi level that are not accessible with conventional W tips. Modeling of these tips using X/W(110) (X=Cr, Fe, W) by first-principles calculations indicates that states responsible for enhanced tunneling are located 0.4–0.6 eV above and below $E_F$ for Fe/W tips, and 0.3 eV above $E_F$ for Cr/W tips. Further calculations show that the formation of an apex atom is not stable for W/W(110) or Fe/W(110) tips, but is stable for Cr/W(110) tips, resulting in point-like iso-density of states contours that are ideal for the selective imaging of the complex electronic properties of the epitaxial graphene on SiC(0001).
Magnetoelectric antiferromagnet, which can be used to induce a switchable boundary magnetization carried by a magnetic material. This concept is particularly useful for applications in non-volatile magnetoelectric memory and spin field-effect transistors. The Neel temperature of the Cr2O3 material is 307 K, which is too high for these purposes. However, for these purposes, the Cr2O3 Neel temperature of 307 K is too low. Using first-principles calculations, we predict that it can be increased by introducing boron as a substitutional dopant in the anion sublattice, which sharply enhances the Curie temperature from 69 K up to as much as 170 K.

Quantum Dots

Using atomic force microscopy, in situ scanning tunneling microscopy, transmission electron microscopy, and in situ scanning Auger spectroscopy, we have grown heteroepitaxial Ge1-xMnx islands bound by {105} facets observed following the introduction of one or more substitutional Gd atom, as well as with an oxygen misfolding and aggregation is unknown, but evidence suggests that tau in AD brains may abnormally interact with the neuronal cell membrane. Using neutron scattering, we can experimentally study the mass and surface fractals discovered in the sedimentary rocks, attractive colloidal glasses and current presentation is focused at the mass and surface fractals discovered experimentally in the sedimentary rocks, attractive colloidal glasses and natural and man-made materials exhibiting multi-level morphology. There is much interest in Gd-doped EuO as a half-metal that could be used as a spin injector. Gd doping (and, more controversially, O deficiency) sharply enhances the Curie temperature from 69 K up to as much as 170 K. I will report the results of first-principles studies of exchange interaction in Gd-doped EuO. In the virtual crystal approximation, the indirect exchange through the conduction band qualitatively explains the observed doping dependence of the Curie temperature. We also considered EuO supercells with one or more substitutional Gd atom, as well as with an oxygen vacancy, and found deviations from the virtual-crystal behavior that can be associated with local lattice relaxations. Time permitting, I will also show how first-principles calculations for large supercells with noncollinear spins can be used to analyze the spin-disorder contribution to the electric resistivity of magnetic metals.

5:20pm MI-WeA11 Magnetic, Structural and Morphological Characterization of Self Assembled Dilute Magnetic MnGe1-x Quantum Dots, J.K. Kassin, J.A. Floro, P. Reinke, C.A. Nolph, University of Virginia

Group IV dilute magnetic semiconductors (DMS) are candidates for the development of spin based devices due to their compatibility with the traditional semiconductor technology. We have grown heteroepitaxial Ge1-xMnx quantum dots (QDs) on Si (001) by molecular beam epitaxial co-deposition, with x ranging from 0.02 – 0.22. There is evidence in the literature for room-temperature ferromagnetism in Mn-doped Ge quantum dots. Using atomic force microscopy, in situ scanning tunneling microscopy, transmission electron microscopy, and in situ scanning Auger mapping, our goal is to clearly ascertain how and where Mn incorporates in our films, especially where the magnetically-active Mn resides, and in so doing to contribute to our understanding of the basic origin of ferromagnetic (FM) ordering in this system. Morphology of the QD’s up to 5 at.% nominal Mn atomic fraction mirror those observed in pure Ge QDs grown at identical temperatures and deposition rates. The standard “hut cluster” islands bound by [105] facets are observed followed by the introduction of dome clusters at larger Ge thicknesses. Noticeable changes in morphology, QD’s density and mean volume become apparent for the highest Mn contents. Further increase in Mn content promotes introduction of rods believed to be a germaine phase. Field cooled hysteresis loops obtained by vibrating sample magnetometry with an in-plane external magnetic field demonstrate ferromagnetic behavior at 5K, with a maximum magnetization saturation of 2.1μB per Mn ion and a coercivity of 250 Oe recorded for x=0.02. Ferromagnetism disappears above 70K, and is not improved by increasing average Mn content of the films. The saturation moment on a per atom basis is seen to decrease with an increase in Mn content. While we cannot yet isolate any specific island type, or the wetting layer, as being primarily responsible for the ferromagnetism, we have shown that dome clusters are not a prerequisite to ferromagnetism. This work is supported by the National Science Foundation under grant number DMR-0907234.

4:40pm MI-WeA9 Concepts based on Magnetoelectrics and Half-Metals for Spintronic Applications, K.D. Belashchenko, University of Nebraska-Lincoln

I will discuss two new concepts of interest for voltage-controlled magnetism and for efficient high-current spin injection in semiconductors, as well as our computational studies of magnetic materials that may be employed in such devices.

First, I will explain the concept of a boundary magnetization carried by a magnetoelectric antiferromagnet, which can be used to induce a switchable magnetization. This effect, demonstrated experimentally using magnetoelectric Cr2O3, can be utilized in non-volatile magnetoelectric memory and spin field-effect transistors. We have grown heteroepitaxial Ge1-xMnx islands bound by {105} facets are observed followed by the introduction of dome clusters at larger Ge thicknesses. Noticeable changes in morphology, QD’s density and mean volume become apparent for the highest Mn contents. Further increase in Mn content promotes introduction of rods believed to be a germaine phase. Field cooled hysteresis loops obtained by vibrating sample magnetometry with an in-plane external magnetic field demonstrate ferromagnetic behavior at 5K, with a maximum magnetization saturation of 2.1μB per Mn ion and a coercivity of 250 Oe recorded for x=0.02. Ferromagnetism disappears above 70K, and is not improved by increasing average Mn content of the films. The saturation moment on a per atom basis is seen to decrease with an increase in Mn content. While we cannot yet isolate any specific island type, or the wetting layer, as being primarily responsible for the ferromagnetism, we have shown that dome clusters are not a prerequisite to ferromagnetism. This work is supported by the National Science Foundation under grant number DMR-0907234.

Neutron Scattering Focus Topic Room: 207 - Session NT+AS-WeA

Moderator: J. Majewski, Los Alamos National Laboratory

3:00pm NT+AS-WeA4 Probing Fractals by the Combined Ultra-Small- and Small-Angle Neutron Scattering (USANS/SANS) Technique, M. Agamanil, Oak Ridge National Laboratory

Many natural and man-made materials exhibiting multi-level morphology (atoms – molecules – aggregates – agglomerates), in other words, existence of intermediate structural units between atomic/molecular and macroscopic levels, usually call hierarchical structures. The combined USANS/SANS is one of the best techniques using at present time for characterization of the hierarchical structures, which in many cases shows fractal behavior. The current presentation is focused at the mass and surface fractals discovered experimentally in the sedimentary rocks, attractive colloidal glasses and aggregates of soot particles in MCT-30 engine oil. Some of the fractal structures, particularly the surface fractals in rocks, are extended over three orders of magnitude in the length scale; therefore, application of the combined USANS/SANS technique, which covers the Q-range extended over five orders of magnitude in the reciprocal space (2πkq Å−1 < q < 1 Å−1), is required to obtain complete structural information for complicated hierarchical structures with fractals.

4:20pm NT+AS-WeA8 Interaction of Alzheimer’s Disease Tau Protein with Model Lipid Membranes, E.M. Jones, Univ. of New Mexico, M. Dubey, Los Alamos National Lab, P.J. Camp, B.C. Givler, Univ. of New Mexico, J. Biernat, E. Mandelkow, Max Planck Unit for Structural Biology, Germany, J. Majewski, Los Alamos National Lab, E.Y. Chi, Univ. of New Mexico

* Falicov Student Award Finalist
enzymes by total internal reflection fluorescence (TIRF) microscopy. Measurements of changes in mass and film stiffness using a quartz crystal microbalance (QCM), and visualization of the motion of individual lipid monolayers at the air/water interface and supported lipid bilayers as model membrane systems, we characterized the interaction between 4 τ assays with membranes of different lipid compositions and elucidated the structure of the protein-membrane films using a combination of biophysical techniques, including pressure-are isothersms, fluorescence microscopy, and x-ray and neutron scattering. Our data show that the full length human tau (hTau40) and its constructs are highly surface active and exhibited strong association with negative DMPG lipids and induced morphological changes observed with fluorescence microscopy, while exhibiting weaker and no interactions with positive DMTAP and neutral DMPC lipids. To elucidate molecular-scale structural details, we used X-ray scattering techniques to study tau and lipid monolayer association. X-ray reflectivity modeling indicated hTau40’s presence under a DMPG monolayer and partial insertion into the lipid headgroup region, while grazing incidence X-ray diffraction data showed hTau40 insertion disrupting lipid packing. We also used neutron reflectometry assays to investigate hTau40’s ability to disrupt lipid bilayers. The protein completely disrupted a DMPG bilayer while not affecting a neutral DPPC bilayer. These results indicate hTau40 has a propensity to interact with a negatively charged membrane surface and disrupt lipid packing, suggesting a possible protein-aggregate induced mechanism for aggregation and toxicity.

5:00pm NT+AS-WeA10 Stabilization of a Lipid Multilayer System by Polysaccharides. M. Kreuzer, M. Strobl, University of Heidelberg, Germany, M. Reinhardt, R. Steitz, Helmholtz-Zentrum Berlin für Materialien und Energie, Germany, R. Dahint, University of Heidelberg, Germany

Hyaluronic acid (HA) is a high molecular weight polysaccharide. It is involved in a wide range of processes in the human body, such as wound healing, tumor progression and joint lubrication. Here we show that HA also stabilizes a lipid multilayer system at physiological conditions. The observed effect may be an important contribution to joint lubrication as lipid films covering the cartilage of natural joints are assumed to reduce internal friction. Neutron reflectometry investigations were carried out at V6 and the new BioRef neutron reflectometer at Helmholtz-Zentrum Berlin. Measurements against excess D₂O verified, that an oligomannuram DMPC lipid bilayer coating remains stable on a silicon substrate at 21 °C in its ordered state (Lα, Pα) with a d-spacing of 66 Å, but detaches almost completely from the solid support at 38 °C in its chain-disordered state (Lβ'). By contrast, oligomannuram lipid bilayers remain stable on a substrate at 38 °C when incubated with a solution of HA in D₂O. Lamella transformations occur over time, resulting in a new lamella phase with a d-spacing of 233 Å. This effect has to our knowledge not been reported before on solid-supported oligomannuram systems. We will discuss potential consequences of the “new” lamella phase with respect to further insight into joint lubrication.

5:20pm NT+AS-WeA11 Neutron Reflectometry, QCM-D, and TIRF Study of the Interaction of Endoglucanases with Films of Amorphous Cellulose. M. Kent, Sandia National Laboratories

We have made electron spin resonance (ESR) and current density versus voltage measurements on a moderately extensive set (over 50 films) of dielectric/silicon structures involving materials of importance to low-κ interconnect systems. Most of the dielectrics investigated involve various compositions of SiOC:H. In addition we have also made measurements on other dielectrics including SiO₂, SiCN:H and SiN:H, some of which are utilized as ESLs. In our study we have made ESR and current density versus voltage measurements both before and after exposing the dielectrics to UV light (hc/λ ≤ 5 eV), and films that have experienced an industrial UV curing process. We observe extremely gross differences in the ESR spectra and leakage current versus voltage response of these low-κ films. We find that UV exposure consistently increases both the density of paramagnetic defects and the leakage current density at a given field. Paramagnetic point defects observed in these films include, E’ centers, silicon dangling bond defects in which the silicon is back bonded to oxygen, the 74 gauss doublet which is E’ center complexed to a hydrogen atom, the 10.4 gauss doublet which is a hydrogen coupled E’ center, the K-center which are silicon vacancies back bonded to three nitrogens, and possibly silicon and carbon dangling bond centers and likely organic radicals. We have also made electrically detected magnetic resonance (EDMR) spin dependent trap assisted tunneling measurements on some ILD films. The close correspondence between the ESR and SDT result establishes a direct link between the defects observed in ESR and the defects responsible for the increased tunneling currents. We have also observed a correspondence between ESR amplitudes and leakage currents. Our preliminary results suggest that the UV curing process creates paramagnetic centers which take part in trap assisted tunneling. Our results indicate quite clearly that the processing parameters have extremely gross effects upon defect densities within these films.

Plasma Science and Technology Division
Room: 202 - Session PS+EM-WeA
Low-K Materials & Integration
Moderator: S. King, Intel Corporation

2:00pm PS+EM-WeA1 Electric and Optical Characterization of Leakage and Breakdown in Low-k Dielectric Materials. J.M. Atkin, R. Laibowitz, Columbia University, T.M. Shaw, IBM T.J. Watson Research Center, T.F. Heinz, Columbia University

Low-k dielectric thin films are finding increased use in integrated circuits for the faster signal speed that they permit. These materials, however, have higher leakage currents and shorter lifetimes than SiO₂-based dielectrics. With the continued push to lower values of k, these problems are becoming more acute.

In this paper, we present results of several complementary characterization techniques for determining key physical properties, such as trap densities and barrier heights, that influence leakage and time-dependent dielectric breakdown (TBBD) phenomena. Electrical characterization techniques include impedance spectroscopy and the measurements of transient currents. In addition, we make use of distinctive optical characterization techniques to obtain specific information about the underlying material properties. Internal photoemission (or photocurrent) spectroscopy yields information on interfacial barrier heights from the photon energies required to induce a current. Optical second-harmonic generation (SHG) provides a sensitive, non-contact method for measuring both photo-driven and spontaneous charge transport. From these methods, we show that the increased leakage current with low-κ materials is due to electrical breakdown, can be directly correlated with increased trap densities. Conduction models accounting for the early failure mechanism will be discussed. Partial support for this work from the Semiconductor Research Corporation is gratefully acknowledged.

2:40pm PS+EM-WeA3 Electron Spin Resonance Study of Low-K Dielectrics and Etch Stop Layers. B.C. Bittel, P.M. Lenahan, T.A. Pomorski, Penn State University, S. King

The electronic properties of thin film low-κ interlayer dielectric (ILD) and etch stop layers (ESL) are important issues in ULSI development. However as the semiconductor industry looks to transition to 16 nm and beyond technology nodes, numerous concerns with low-κ materials need to be addressed. Leakage currents, time dependent dielectric breakdown and stress induced leakage currents are critical problems that are not yet well understood in ILD. A topic of current interest is ultraviolet light (UV) curing of low-κ materials. We have made electron spin resonance (ESR) and current density versus voltage measurements on a moderately extensive set (over 50 films) of dielectric/silicon structures involving materials of importance to low-κ interconnect systems. Most of the dielectrics investigated involve various compositions of SiOC:H. In addition we have also made measurements on other dielectrics including SiO₂, SiCN:H and SiN:H, some of which are utilized as ESLs. In our study we have made ESR and current density versus voltage measurements both before and after exposing the dielectrics to UV light (hc/λ ≤ 5 eV), and films that have experienced an industrial UV curing process. We observe extremely gross differences in the ESR spectra and leakage current versus voltage response of these low-κ films. We find that UV exposure consistently increases both the density of paramagnetic defects and the leakage current density at a given field. Paramagnetic point defects observed in these films include, E’ centers, silicon dangling bond defects in which the silicon is back bonded to oxygen, the 74 gauss doublet which is E’ center complexed to a hydrogen atom, the 10.4 gauss doublet which is a hydrogen coupled E’ center, the K-center which are silicon vacancies back bonded to three nitrogens, and possibly silicon and carbon dangling bond centers and likely organic radicals. We have also made electrically detected magnetic resonance (EDMR) spin dependent trap assisted tunneling measurements on some ILD films. The close correspondence between the ESR and SDT result establishes a direct link between the defects observed in ESR and the defects responsible for the increased tunneling currents. We have also observed a correspondence between ESR amplitudes and leakage currents. Our preliminary results suggest that the UV curing process creates paramagnetic centers which take part in trap assisted tunneling. Our results indicate quite clearly that the processing parameters have extremely gross effects upon defect densities within these films.
Defect concentrations in low-k organosilicate glass [SiCOH] films deposited on high-resistivity silicon were measured with electron-spin resonance. Both plasma exposure and ultraviolet exposure were used. During argon electron-cyclotron resonance plasma exposure, ion and photon bombardment increased the measured defect concentrations. Ultraviolet lamp exposure was also shown to increase the defect concentrations. SiCOH samples with several dielectric constants were examined showing that as the value of the dielectric constant was lowered, the defect concentrations were shown to increase significantly.[i] In addition, the nature of the defects in SiCOH was investigated using air and nitrogen plasmas. The defects were found to be silicon dangling bonds. Air-plasma exposure increases the defect concentrations by breaking silicon-hydrogen bonds, measured by Fourier-transform infrared spectroscopy. Nitrogen-plasma exposure as well as free-radical exposure have only a small influence on the bond breaking. It was also shown that UV curing improves the chemical-damage resistance of the dielectric.

Work Supported by the Semiconductor Research Corporation under contract 2008-KJ-1781 and the National Science Foundation under Grant CBET-1066231.


Time dependent dielectric breakdown (TDDB) is a major concern for newly emerging low-k organosilicate (SiCOH) dielectrics. TDDB degradation can be caused by changes in electrical, chemical, and mechanical properties of the dielectric materials.[i] [ii] [iii] [iv] In order to examine the effect of plasma exposure on TDDB degradation, time-to-breakdown measurements were made on porous SiCOH before and after exposure to a variety of plasma exposure conditions. Plasma parameters were changed between exposures such that each sample was subjected to different charged particle and vacuum ultraviolet photon fluxes. In order to determine how TDDB degradation was affected by each of them during plasma exposure, ion and photon bombardment were affected by each of them during plasma exposure. By utilizing a capillary-array window to separate charged particle and photon bombardment, it is possible to show that each process is responsible for causing different types of TDDB degradation. A constant voltage TDDB measurement technique was implemented to analyze unexposed, VUV-irradiated and plasma (charged-particle and photon bombardment) exposed samples to examine the degradation in TDDB. It was observed that the time to breakdown reduces as the electric field in the material decreases, which is consistent with what has been previously predicted. It was also found that the unexposed samples exhibit longer time-to-breakdown, indicating highest reliability. Capillary-array-window covered samples exhibited marked degradation in leakage currents and time-to-breakdown relative to the unexposed samples. However, samples exposed to both charged particle and VUV photon bombardment exhibited the most significant degradation, resulting in substantially reduced breakdown times and increased leakage currents. Thus both charged particle and photon bombardment degrade TDDB.

This work has been supported by the Semiconductor Research Corporation under contract 2008-KJ-1871 and by the National Science Foundation under Grant CBET-1066231.

through the use of these high performance generators, a more holistic analysis of current power delivery systems seems to indicate that the characteristics and quality of the power delivery system could be partly limited by power delivery system architectures based on autonomous building blocks. This observation suggests that a higher level of cooperation in communication, measurement and control among the components of the power delivery system could further benefit the development of next generation processes and tools. In this presentation we will show results [1]. As such, a new generation of power delivery systems where the matching networks and generators are fully integrated from a communication, measurement and control point of view. Although certainly possible, this integration scheme does not necessarily imply full physical integration. The performance of the integrated hardware combined with high performance control algorithms on both fixed and dynamic loads will be discussed in terms of electrical measurements performed at the loads under various operating conditions, including match and frequency tuning while pulsing. The results will demonstrate the potential for these integrated systems to enable processes with stringent requirements such as highly accurate power regulation in a wide dynamic range, faster power stabilization during transients, and minimal reflected power, both in pulsing and CW modes.


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An increase in the chamber pressure results in a decrease in the maximum RF electric field that can be sustained without breakdown as the pressure approaches a few mTorr. The breakdown event leads to formation of a plasma in the structure, and the addition of an external magnetic field causes the formation of a plasma at lower pressures. Ultra-violet light, with an energy greater than that of the work function of the electrode material, has been shown to induce electron emission from the surface and initiate multipactor discharges. In addition, we are using optical emission spectroscopy to determine the magnitude of the DC and RF electric fields near the electrode structure by utilizing the dynamic Stark effect. Experimental details and future research directions will be presented.

4:00pm PS-WeA7 A Narrow Ion Energy Distribution Bias System. V. Brouk, Advanced Energy Industries, S.C. Shannon, North Carolina State University, D.J. Hoffman, D. Carter, W. Hattel, Advanced Energy Industries

Conventional bias systems use sine wave voltage systems to achieve ion energy distribution functions for the creation of thin films. By combining dual frequency sine waves, the mean energy and its spread can be independently controlled. Arbitrary wave-shaping has been suggested to create single energy near-delta function distributions. We investigate a system where the waveform is defined a priori where two elements are run in a feed forward system to control the instantaneous IEDF. We evaluate the effectiveness of this system in an argon/oxygen plasma at typical operating pressures inside the 10-150 mTorr range at plasma densities in the low 10^16 cm^-3 range.


Inductively coupled plasma (ICP) sources, commonly used for semiconductor and conductor etching, embody the concept of functional separation between plasma production and ion energy control, wherein the inductive coupling through the coils is only responsible for the plasma generation while the bias determines the ion energies. Plasma etching of microelectronics structures at advanced technological nodes (~ 3x nm), especially complicated structures such as multi-gate MOSFETs and 3D microelectronics structures at advanced technological nodes (~ 3x nm), are placing great emphasis on control of ion energy distributions (IEDs) to finely discriminate etching thresholds. Sinusoidal biases typically provide broad IEDs, making such control difficult to achieve. One promising alternative is the non-sinusoidal bias waveforms, which have been demonstrated to provide such control. However, there are issues associated with passing non-sinusoidal signals through the finite impedance of the match, transmission line, and the substrate. Multi-frequency bias is compatible with current manufacturing hardware, and has been successfully used for controlling IEDs in capacitively coupled plasmas. This approach may be enabled to use IED control in ICP etchers as well. We investigate the impact of multiple bias frequencies in this paper. A high frequency applied in addition to a relatively low frequency bias causes the sheath potential to vary in a complicated manner due to the non-linear nature of the sheath. As a result, IED exhibits a complex dependence on relative bias voltages and frequencies. In this work, the effect of applying bias at multiple frequencies will be discussed using results from a computational investigation. The 2-dimensional plasma equipment model, HPEM,[4] has been modified to enable power deposition at multiple frequencies on the same electrode. Results will be discussed for Ar/Cl2 plasma utilizing a 13.56 MHz bias in addition to a bias at a different frequency in an ICP chamber. The additional frequency is varied over a wide range and its consequences assessed on the ion and radical flux and IEDs incident on the wafer. We found that, in addition to modulating the IEDs, the flux composition is different depending on the frequency due to the secondary plasma generation by the alternate frequency.


Low pressure (sub-20 mTorr) capacitively coupled plasmas (CCP) are playing an increasingly important role in technological applications. As the mean free path becomes commensurate with the discharge dimensions, the fluid assumptions inherent in plasma and sheath models start to break down and ought to be reexamined. We focus on one aspect of the CCP operation in particular: the energy distribution of these electrons close to the surfaces, and use kinetic particle-in-cell (PIC) models to understand the temporal behavior of the EED. Kinetic results are compared to fluid representation of the EED at electrodes to identify deficiencies in the fluid model at low pressures and propose solutions.

The sheath at the plasma-surface interface ensures that the electrons remain confined in the bulk plasma. However, during certain phases of the radio-frequency (RF) cycle in a CCP, the sheath collapses and the electrons exit at the electrodes. Energy distribution of these electrons contains useful information about the bulk plasma and the sheath. One can probe into the energy characteristics of these electrons using dc probes embedded in the electrode. Analysis of the resulting probe data can be used to determine the electron temperature, the electron density, and the EED in the bulk plasma. If a fluid model is used for this analysis, the electrons are assumed to be governed by the Boltzmann relation where their density and flux depend exponentially on the sheath voltage. Electrons are however highly non-equilibrium near the sheaths in CCPs and the Maxwellian distribution assumption (implicit in the Boltzmann relation) is questionable. Furthermore, most probe analysis models are dc-based. Low pressure situations demand further scrutiny as even the bulk plasma EED tends to become non-Maxwellian.

1 and 2-dimensional PIC model of CCPs are used for this investigation. These models consider plasma chemistry using the Monte Carlo technique. Simulations are done for Ar and Ne plasmas under a variety of conditions (13.56 – 60 MHz RF frequency, RF voltage of 100 – 500 V, 5 – 100 mTorr gas pressure). The 1-dimensional PIC model is used to examine the EED at the electrodes where the sheath undergoes substantial variation during the RF cycle. The 2-dimensional model is used to investigate the EED at small metal surfaces (e.g., a probe) away from the primary electrodes. DC voltage is also applied to the probe electrode in the 2-dimensional simulations. It is found that, in addition to a non-Maxwellian contribution from electrons adjacent to the sheath, the EED also contains high energy electrons which are the remnant of electrons that were accelerated at the opposite sheath.

5:00pm PS-WeA10 The Control of Electron Shading and Plasma EED in a DC/RF Parallel-Plate Etcher. L. Chen, Tokyo Electron America

There are several principles in the consideration of plasma etcher design. This paper addresses two important areas: (1) the ability of tailoring the electron energy distribution function (EED), (2) the ability of adjusting the charging and neutralization of surface features (the electron shading effect). Stochastic heating by high frequency RF (VHF) energizes the Maxwellian bulk into the energetic tail population for efficient ionization. Such energetically bottom-up heating also indiscriminately populates the below-ionization energetic group, leading to a downstream dissociation and VUV production. A generic DC/RF system has a RF biased wafer-electrode with a high-negative DC superimposed opposing electrode. The DC/RF system dominates its electron heating with an energetically top-down process. The secondary electrons emitted from the high-negative DC surface disseminate the beam-energy into a distribution of energetic-electrons through collisions and more importantly, various beam-wave instabilities. These energetic electrons are trapped between the sheaths of the two parallel plates when the RF sheath field is sufficiently strong, dissipating their energies mainly into ionization. The energetic part of the EED reveals (in descending order) a group of ballistic-electron associated with the applied –DC voltage, an energy continuum, and finally stop at a lower medium-energy peak in the range of ~ 40eV to 300eV depending on the process. The data show extremely efficient ionization by these energetic electrons and as a result, the Maxwellian bulk remains relatively unchanged at T_e ~ 1.8eV regardless the bias RF power and process pressure. Such energetically decoupled EED enables increased ionization without increasing molecular dissociation. In one RF period, the trapping of these energetic electrons diminishes as the RF sheath collapses. By pulse-syncing the –DC voltage and the bias RF voltage, ion-bombardment excitation of the wafer surface can alternate with high-energy electron-bombardment neutralization of the wafer surface. Ion-bombardment of the wafer surface occurs when the bias RF pulse is high and the –DC pulse is at a medium level. As the RF and the DC pulses are synchronously altered to a low RF voltage and a high-negative DC voltage, ion-bombardment excitation of the wafer surface diminishes and the wafer surface sees an increased current of energetic (>25eV) electrons. High aspect ratio SEM data show that the signatures of electron shading (e.g., bending and twisting of the
features) are eliminated when synchronous DC/RF pulsing is implemented allowing energetic electrons to reach the bottom of high aspect ratio features.

5:40pm PS-WeA12 Negative Plasma Potentials Produced by Electropositive Plasmas in a Multi-Dipole Chamber, N. Hershkowitz, University of Wisconsin-Madison, L. Okusaz, Suleyman Demirel University, Turkey, J.P. Sheeahan, University of Wisconsin-Madison

Negative plasma potentials were obtained in DC hot filament unmagnetized electropositive argon plasma in two configurations. For sufficiently low plasma density (~10^10 cm^-3) bounded by conducting walls, double layers provide ion and electron confinement near the walls. Similar results were observed in higher density plasmas (~10^11 cm^-3) when a thin dielectric coating of oil covered the surface of the conducting walls. The potential profiles, measured using emissive probes in the limit of zero emission, from the center of the plasma to the potential minima are quite similar in shape to those observed when the plasma has positive plasma potentials. The primary electrons emitted from the filaments are important for charge conservation and for modification of the Bohm criteria but are not important for current balance.

This work was supported by US Department of Energy grants No. DE-AC02-93CH11466 and No. DE-FG02-97ER54437, and the Fusion Energy Sciences Fellowship Program administered by Oak Ridge Institute for Science and Education under a contract between the U.S. Department of Energy and the Oak Ridge Associated Universities.

Advanced Surface Engineering Division Room: 104 - Session SE+PS-WeA

Atmospheric Pressure Plasmas
Moderator: H. Barankova, Uppsala University, Sweden

2:00pm SE+PS-WeA1 An Investigation of the Influence of Hybrid Current Modes on the Plasma Behavior during Plasma Electrolytic Oxidation (PEO) Coating Process on Mg and Mg-Alloys, R.O. Hussein, D.O. Northwood, X. Nie, University of Windsor, Canada.

The increased use of magnesium alloys is considered one of the more promising methods for light-weighting in the automotive industry since, for a given strength level, Mg represents a 57% weight reduction over steel and 8% weight reduction over aluminum. However, due to its high chemical and electrochemical activity, magnesium has poor corrosion resistance in aqueous and other environments. In order for Mg and its alloys to find increased usage, there is a need to surface engineer these materials to improve corrosion and wear resistance. Plasma Electrolytic Oxidation (PEO) is an electrochemical process working at atmospheric pressure that uses an environmentally-friendly aqueous electrolyte to oxidize the metal surfaces to form ceramic oxide coatings which impart a high corrosion and wear resistance. The properties and structure of PEO coatings are dependent on parameters such as the electrolyte composition of the electrolyte and the process conditions including current density, current mode and processing time. In this study we investigated the effect of current mode on plasma temperature and coating properties of PEO coatings formed on pure magnesium and an AM60B magnesium alloy (mass fraction: Al 5.6-6.4%, Mn 0.26-0.4%, Zn ≤ 0.2%, balance Mg). Unipolar, bipolar and hybrid (combination of both) current modes were used in this work. Optical Emission Spectroscopy (OES) was employed to study the plasma species, and electron temperature of the plasma. The morphology and microstructure of the coatings were investigated using Scanning Electron Microscopy (SEM), Potentiodynamic polarization in a 3.5% NaCl solution was used for the corrosion investigations.

2:20pm SE+PS-WeA2 Aging Mechanism of the Hydrophilic Silicon (100) Native Oxide Surface, T.S. Williams, R.F. Hicks, University of California Los Angeles

The wetting behavior of surfaces is important in many applications, such as for example, microfluidic devices that are fabricated on silicon wafers. In this study, silicon native oxide surfaces were cleaned with a radio frequency, atmospheric pressure helium and oxygen plasma and with ammonium hydroxide, hydrogen peroxide, and deionized water in a 1:1:5 ratio (RCA SC-1). Both processes created a hydrophilic state with water contact angles <10° and 16.2±1.7° respectively. During subsequent storage in a chamber purged with boil off from a liquid nitrogen tank, the water contact angle increased over several days at a rate dependant on the cleaning method used. Internal reflection infrared spectroscopy revealed that the change in water contact angle was due to the adsorption of organic molecules with an average hydrocarbon chain length of 10±2. The rate of the adsorption process decreased with the fraction of hydrogen-bonded hydroxyl groups on the surface relative to those groups that were isolated. On Si (100) surfaces that were cleaned by RCA SC-1 and the plasma, 96% of the silanol groups were hydrogen bonded. Several methods have been explored for keeping the silicon dioxide surface in a hydrophilic state for extended periods of time, and these will be presented at the meeting.

2:40pm SE+PS-WeA3 Polyimide Surface Treatment to Hydrophobic Surface with Self-Assembled Mask Layer for Direct Inkjet Patterning Process, J.B. Park, G.Y. Yeon, Sungkyukwan University, Republic of Korea

The inkjet printing technology is emerging as one of the important process technologies for electronics, because it can significantly reduce the manufacturing process cost, materials waste, and number of process steps. Especially, many researchers investigating the inkjet-printed process have focused on attention to two considerations: (1) conductive film or polymer film, and (2) the technique is not suitable for applying to organic transistors yet, because its pattern width is limited to several dozens µm by statistical variations of the flight direction of droplets and their spreading on the substrate.

In this study, polyimide film was treated to hydrophobic surface using modified atmospheric pressure plasma system (double discharge system). Especially, for keeping the surface characteristics permanently, surface of the polyimide film was etched and textured with self-assembled mask layer to form the textured polymer surface. The texturing process is progressed for 4 process step. The mask layer(HMDS) is deposited and agglomeration/oxidation by using atmospheric discharge of HMDS/He gas mixture through 1st step and 2nd step and Etch and texturing process is progressed with atmospheric discharge of He/O2 gas mixture in 3rd step. And the mask layer is removed in 4th step using NF3/H2 gas mixture of removing atmospheric plasma. In the 2nd process, the HMDSO mask layer is formed specific tissue layer as the input power was increased. And the morphology is also varied with the whole size of the HMDSO mask layer. And this whole process is possible because of perfect etch selectivity between the polyimide film and HMDSO layer. The contact angle of textured polyimide film was measured over the 100 degree and varied with textured surface morphology. And keeping the pattern width of inkjet printing was clearly improved compare with normal polymer surface.

3:00pm SE+PS-WeA4 In Situ Fabricating Blue Ceramic Coatings on Al Alloy by Plasma Electrolytic Oxidation, Z.J. Wang, R.O. Hussein, X. Nie, H. Hu, University of Windsor, Canada

In-situ formation of novel blue ceramic coatings on Al alloy with a controllable blue color was successfully achieved using a plasma electrolytic oxidation (PEO) process working at atmospheric pressure. This novel blue ceramic coating overcomes the shortcomings of traditional surface treatments resulting from traditional dyeing process by depositing organic dyes into the porous structure of anodic film, such as poor resistance of abrasion and rapid fading when exposed to light. X-ray diffraction, scanning electron microscopy and energy dispersive spectroscopy were employed to characterize the microstructure of the blue ceramic coating. The main compositions of the coating are CoAl2O4 and Al2O3. This work shows that the working current density plays significant roles on CoAl2O4 phase in the ceramic coating which has a controllable influence on the coating color. Low current density reduces the percentage of CoAl2O4 component in the coating, and light blue ceramic is produced. On the other hand, higher current density increases CoAl2O4 component in the coating, hence deep blue ceramic is fabricated. Electrochemical test, ball-on-plate sliding wear tester and thermal shock method were utilized for the corrosion, wear and thermal shock resistance analysis of the fabricated ceramic coatings. The results indicate that the developed blue ceramic coating via PEO process superiorly improves the tribological property, anti-corrosion property and thermal shock resistance. The fabricated blue ceramic coating tends to be applied to colored light cast alloys for applications used in relatively harsh and severe working environments.

4:00pm SE+PS-WeA7 Cold Atmospheric Plasma Sources for Treatment of Cell-Containing Surfaces, M.G. Kong, Loughborough University, UK

INVITED

Low-temperature gas discharges generated at atmospheric pressure, commonly known as cold atmospheric plasmas, are a relatively new member of the processing plasma family. Without the need for a vacuum chamber, the plasma can be driven by more cost-effective route to material processing and open up opportunities for chamber-less processes and vacuum-incompatible materials such as moist objects. With a mean electron energy at a few eV and a gas temperature close to room temperature, cold atmospheric plasmas are ideally placed for effective chemical dissociation and hence for a wide range of materials processing applications. In this

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contribution, a review of the current cold atmospheric plasma sources will be presented in terms of their underlying science and their current technological capability. These include the traditional dielectric barrier discharges, radio-frequency glow discharges, and cold atmospheric plasma jets and jet arrays. Through discussion of their plasma characteristics, their applications are exemplified through treatment of cell-containing surfaces, including living tissues. The interaction of cold atmospheric plasmas with individual microorganisms and indeed microbial communities will be discussed, and its implications to treatment of skin diseases and wounds will be presented. This will also be supported with data of plasma interaction with mammalian cells.


In this paper, the spatial distribution of atomic oxygen density in a 60-Hz non-equilibrium atmospheric pressure plasma[1] is diagnosed by two-photon absorption laser induced fluorescence (TALIF)[2]. The plasma unit is made of ceramics comprised three regions: gas diffusion region, main discharge region, and plasma jet in the open air. The discharge gases were Ar and a small amount of O2. The plasma could offer electron density as high as 1015 cm-3 with a low gas temperature[1,3], and have been used for various applications such as industrial or biomedical plasma. We will present the fundamental research has been carried out on the model transition-metal carboxylate bonds effectively convert photons into electric energy. In last years, surface plasmon has been commercialized for surface treatment processes because of its high performance characteristics has produced correlations relating power and charge transfer to various electrical and geometrical parameters of the system which will be discussed in this paper.

5:00pm SE+PS-WeA10 Dense Atmospheric Pressure Discharges for Surface and Gas Treatment. M.J. Kelly, B.D. Schultz, W.M. Hooke, International Technology Center.

Dielectric barrier discharge (DBD) plasmas have been formed in atmospheric pressure gases exhibiting peak currents in excess of 100 amperes. Power densities during the pulse routinely exceed 100 kilowatts per cubic centimeter for moderately sized electrodes (~100 square centimeters) with ionization densities of 1010 per cubic centimeter. Charge transfer of 100 microcoulombs per pulse has been repeatedly generated at frequencies up to 100 hertz, and the charge delivery is found to scale in proportion to the electrode area for a given dielectric consistent with a homogeneous discharge. Diffuse discharges have been formed over larger areas (exceeding 1 meter in length and 500 square centimeters) as well. These results were obtained using a custom high voltage driving source and in the absence of helium, argon, or any other easily ionized gas. The source readily achieves an overvoltage in excess of the DC breakdown voltage prior to the onset of breakdown in which 20-30 kV is delivered with rise times shorter than the lag time between the pulse crossing the threshold voltage and the onset of a discharge. Electrical modeling of the discharge characteristics has produced correlations relating power and charge transfer to various electrical and geometrical parameters of the system which will be discussed in this paper.
decades numerous IR investigations of oxide powders, including the different modifications of TiO2, have been reported. An unambiguous assignment of the bands in the complex spectra in connection with the properties of the adsorbed species is not possible on the basis of data recorded for well-defined reference systems, e.g. surfaces of single crystals. Unfortunately, studies on oxide single crystals are extremely scarce due to the fact that the sensitivity of reflection IR-spectroscopy for molecular adsorbates is two orders of magnitude lower for oxides compared to metal single crystals. Only recently was it possible to overcome these technical problems by employing a novel, optimized spectrometer.

Here, we will demonstrate the performance of this highly sensitive IRRAS-setup by presenting high-quality IR-spectra obtained for two molecules, benzoic acid and terephthalic acid, adsorbed on rutile TiO2 (110). Owing to the fact that many Dye Sensitized Solar Cells (DSSCs) consist of dyes grafted to the oxide support via carboxylate groups determining and controlling the adsorption of carboxylic acids on oxide substrates is fundamental to understanding the energy transfer from the molecule to the substrate. For the present experiments, monolayers of terephthalic acid (TPA) and benzoic acid (BA) were first deposited under UHV-conditions on a rutile TiO2 (110) surface at room temperature. Subsequently the sample was transferred in the main chamber and subjected to an analysis in a highly sensitive UHV IRRAS system. While for BA the expected bidentate carboxylic acid binding is observed, for TPA the presence of two carboxylic acid groups leads to interesting complications. The IR-spectra allow, in particular, answering the question whether for the flat-lying TPA species observed in scanning probe techniques the carboxylic acid group is still protonated, a question which could not be answered by the results from x-ray absorption spectroscopy.


References:
The chemical reactivity was characterized by the adsorption and decomposition of various molecules such as methanol, water and acetaldehyde. Reaction products were monitored by TPD and surface intermediates were determined by soft x-ray photoelectron spectroscopy. In general the CeO2(100) surface was found to be more active, i.e. molecules adsorbed more readily and reacted to form new products, especially on a fully oxidized substrate. However the CeO2(100) surface was less selective with a greater propensity to produce CO, CO2 and water as products. The differences in chemical reactivity are discussed in light of possible structural terminations of the two surfaces.

Research sponsored by the Division of Chemical Sciences, Geosciences, and Biosciences, Office of Basic Energy Sciences, U.S. Department of Energy. Portions of this work were conducted at the National Synchrotron Light Source, Brookhaven National Laboratory, and Oak Ridge National Laboratory's Center for Nanophase Materials Sciences, which are sponsored by the Office of Basic Energy Sciences, U.S. Department of Energy.

4:40pm SS-WeA9 Adsorption and Photo-Reactivity of CO on TiO2(110), N.G. Petrik, G.A. Kimmel, Pacific Northwest National Laboratory

We have studied the low-temperature adsorption and reactions of CO on reduced, oxidized, hydroxylated, and electron-irradiated TiO2(110) using temperature programmed desorption, photon-stimulated desorption (PSD) and reflection-absorption infrared spectroscopy (RAIRS). Changing the condition of the crystal surface and the adsorbate coverage provides insight into the interactions of adsorbed CO with 5-fold coordinated Ti sites (Ti5c), bridging hydroxyls (BBO) sites, and defect sites (oxygens, F, vacancies, bridging hydroxyls and radiation-induced surface defects). Infrared spectra were obtained for light with the plane of incidence parallel and perpendicular to the [001] azimuths of TiO2(110). For adsorption on Ti5c sites, the RAIRS spectra are consistent with CO adsorbed nearly perpendicular to the surface. For adsorption on BBO sites, the molecules adsorbed parallel to the surface and perpendicular to the rows of BBO atoms. The reactivity of various molecular adsorption forms of CO is probed using PSD. In CO photo-oxidation, the PSD yields of CO and CO2 change dramatically with initial CO coverage, indicating the importance of the relative position and orientation of O2 and CO molecules for the photochemical reaction.


Understanding the fundamental aspects of CO2 adsorption and reaction on well-characterized oxide surfaces is critical in providing fundamental understanding on how to control catalytic carbon sequestration and CO2 conversion to fuels. A model oxide surface, rutile TiO2(110) is used to investigate the adsorption properties of CO2 using scanning tunneling microscopy (STM) and atomic force microscopy (AFM) images obtained before and after in-situ exposures of CO2 at 50 K reveal that the CO2 molecules preferentially bind in bridge-bonded oxygen vacancy (BVO) defect sites. We show that electron injection from the STM tip can induce CO2 reduction to CO and V0 annihilation. After the saturation of V0’s, CO2 molecules preferentially adsorb on five-coordinate Ti sites, where they remain mobile even at 50 K. The mobility of these molecules can be correlated with the incorporation of CO2 onto TiO2 surfaces. The adsorption behavior of CO2 on hydroxylated TiO2 surfaces will also be presented.

5:20pm SS-WeA11 Interaction of ZnO-supported Cu Oxides with CO and CO2, Y. Zhang, F. Wang, M. Le, M. Ren, J. Flake, P. Sprunger, R. Kurth, Louisiana State University

Cu and Cu-oxide nanoclusters supported on ZnO are prototypical catalysts for the electrochemical reduction of CO and CO2 to methanol. In this report we describe the interaction of CO and CO2 with Cu oxide nanoclusters on ZnO(1010) with a combination of surface sensitive tools including STM for structural information, EELS for electronic and vibrational studies as well as synchrotron-based photoemission for electronic properties. EELS deposited on ZnO(1010) is oxidized with a combination of O2-exposure and annealing procedures to result in two distinct Cu-oxide (CuI and CuII) clusters, which preferentially nucleate and grow at step edges. Photoemission shows a large charge transfer between the oxide cluster and the substrate surface as well as significant band bending. It is believed that the CO2 adsorption, forming a carbonate species, and consequent reduction, is coupled to the induced defects and electronic perturbation of the CuO/ZnO nanoclusters, absent in the case of Cu/ZnO nanoclusters. In addition to vibrational EELS and TDS to characterize the adsorption of the CO and CO2 adsorption species, similar results from Au on ZnO(1010), which shows a lack of cluster formation growth, will be compared and contrasted.

This material is based upon work supported as part of the Center for Atomic Level Catalyst Design, an Energy Frontier Research Center funded by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences under Award Number DE-SC0001058.


Chemically selective sensors are required for detection of chemical warfare agents with ever increasing demands on the selectivity, sensitivity, lifetime, speed, and reduced power consumption of these devices. Strategies for reducing the scale of these sensors have been explored to produce microfabricated Nitrogen-Phosphorus Detectors (NPDs) to accommodate these many requirements. The device integrates sol-gel derived alkali metal silicate thin films on low thermal mass silicon substrates for field portable gas chromatography applications. In spite of the long history of NPDs, the details of the chemically-mediated emission related to their selectivity are not well understood. The NPD signal current ultimately depends on the transfer of electrons across the surface potential barrier of the thermionic cathode emission. Two mechanisms have been described in the literature to account for the chemically-selective ionization observed in NPDs: (a) gas-phase ionization models and (b) surface mediated electron emission. The latter mechanism has been the focus of our measurements of the surface work function of candidate emitter materials as a function of composition, structure, temperature, and ambient atmosphere. Specifically, both the local work function variations by scanning probe measurements and effective average work function by measuring total emission will be discussed.

Thin Film Division

Room 109 - Session TF1+EM-WeA

Nonvolatile Memory

Moderator: T. Karabacak, University of Arkansas at Little Rock

2:00pm TF1+EM-WeA1 Application of Amorphous Zinc Tin Oxide for Memristor Devices, G.S. Herman, J.S. Rajachalambarum, S. Murali, J. Couley, Oregon State University, S.P. Sanghavi, P. Nachimuthu, V. Shathanandan, T. Farga, S. Thvuthasain, Pacific Northwest National Laboratory

Amorphous zinc tin oxide semiconductor materials have been studied primarily as the active semiconducting material for thin film transistors with applications including transparent and flexible electronics. Due to the amorphous nature of these materials excellent uniformity can be obtained over large area, while still having reasonably high electron mobilities (~10 cm²/Vs). Furthermore, considerable control over the electrical properties can be maintained, where insulating, semiconducting, and conductive properties can be obtained by varying the processing conditions. We have recently used sputter-deposited zince tin oxide as a bipolar switching element in memristor devices. Memristors are a two-terminal nonvolatile data memory device that is very promising for the replacement of silicon-based Flash. A significant benefit is that memristors can be manufactured with high areal densities with potentially low manufacturing costs. A variety of oxide-based materials are being evaluated for memristors, however many of these are nanocrystalline which can lead to non-uniformity of devices. We have found that the switching properties of zinc tin oxide are closely related to the process conditions, the electrical test conditions, and the electrode materials. In this presentation we will discuss the physical and electrical characterization of the oxide films and the switching mechanisms for these materials.

2:20pm TF1+EM-WeA2 PE-MOCVD of GeTe Materials for Phase Change Memory Applications, E. Despiau-Pujo, L. Dussault, C. Vallée, LTM/CRNS-UJF, France, E. Gourevest, ST Microelectronics, France, D. Jourde, S. Maitrejean, P. Michallon, CEA Leti Miniatures Campus, France

Phase Change Random Access Memory (PCRAM) are very promising candidates for next generation of non-volatile memories. Those devices store information using the high electrical contrast between the amorphous and crystalline phases of chalcogenide alloys such as Ge2Sb2Te5 (also called GST) which are mainly deposited by physical vapor deposition (PVD) [1].
However, two important drawbacks have to be overcome: a too short archival life for GST at high operating temperatures and a too high power consumption. To overcome the binary compound GeTe seems to be a promising candidate for high temperature applications since it has an estimated archival life up to ten years at 110°C. A pronounced increase in thermal stability of N- and C-doped GeTe was also demonstrated [2].

Concerning the power consumption, it was shown that a way to reduce the high operating currents is to confine the material. In this way, several groups try to develop a new deposition process to achieve the gap filling of the confined structure. Atomic Layer Deposition (ALD) of GST has shown very good step coverage but the deposition rate remains very low. Therefore, we proposed to study the deposition of GeTe in a shower-head type 200mm plasma-enhanced pulsed liquid injection CVD reactor (AltaCVDD200). Ge and Te liquid precursors are introduced into the deposition chamber as vapours through a pulsed injection system and an evaporating furnace. A capacitatively-coupled RF plasma is applied between the shower head and the substrate heater to promote homogeneous deposition. Chamber walls are maintained at 70°C to avoid condensation of precursors vapours. The deposition chamber is mounted on a cluster tool which allows quasi in situ analysis of the deposited films by angle-resolved XPS while the plasma is analyzed by optical emission spectroscopy (OES).

In this paper, we investigate the optical emission spectra of high pressure (2-20 Torr) Ge- and Te-containing plasmas. Reference UV-visible emission spectra (200-800 nm) are recorded for various conditions of RF power, pressure and flow rates. The role of H atoms and C contamination in the deposition (due to the decomposition of the precursors) and etching of Ge and Te atoms at the growing film surface. Role of C contamination in the phase change properties is also discussed and phase transitions of plasma deposited GeTe-C materials are compared with sputtered GeTe-C.


2:40pm TF1+EM-WeA8 Embedded HfO2 based 1T1R Cells for Future RRAM Applications. Ch. Wenger, T. Berndt, Ch. Walczyk, D. Walczyk, M. Malgorzata, HHP, Germany

The integration of various functionality to (Bi)CMOS circuits is in the focus of the “More than Moore” approach. Here, we demonstrate the incorporation of nonvolatile memories (NVM) into the Back end of line (BEOL) of Bi(CMOS) circuits. The added functionalities open new technological possibilities for high value microelectronics systems.

Embedded nonvolatile memories (NVM) with high-density, high-speed, and low-power are attractive for a growing number of applications. One promising candidate for next-generation nonvolatile memories is based on the electrically switchable resistance change between a high (OFF-state) and a low (ON-state) resistive state of a metal-insulator-metal (MIM) structure. This approach is often termed resistance random access memory (RRAM) technologies. Due to the cost effectiveness and BEOL compatibility with (Bi)CMOS technologies, this approach is highly attractive. By combining the MIM devices (R) with selection transistors (T), the 1T-1R cells offer good scalability, long retention time, and rapid read/write times. In this letter, the reliable bipolar resistive switching of TiN/HfO2/TiN devices embedded into 1T-1R cells is demonstrated.

The current-voltage (I-V) characteristics of the TiN/HfO2/TiN diodes and 1T-1R cells were studied by DC voltage sweep measurements. By applying a positive voltage at the top electrode, the resistance is reduced for V > Vsel. The OFF-state is switched into the ON-state when a negative voltage is applied beyond Vreset. In case of the 1T-1R cell, the forming and set processes can be controlled by changing the gate voltage (V_G) of the select transistor.

4:00pm TF1+EM-WeA7 Synthesis and Characterization of Multiferroic Oxides by Radical Enhanced Atomic Layer Deposition, C.D. Pham, J.H. Choi, J.P. Chang, University of California Los Angeles

Multiferroic materials exhibit two or more forms of ferroic order such as (anti-)ferroelectricity, (anti-)ferroelasticity, and ferromagnetism. Multiferroic materials containing both ferroelectricity and ferromagnetism will exhibit some amount of magnetoelectric coupling which is a desirable aspect for the future of non-volatile memory, as these materials could potentially be used for devices that will be written magnetically and read electronically or vice versa, as well as the prospect of four-valued memory. Materials which exhibit magnetoelectric coupling have been well studied, however, the synthesis methods may not easily translate into large scale integration. One possible route for synthesis on a commercial scale, atomic layer deposition (ALD) is a thin-film processing technique which involves alternatingly flowing non-self reacting precursor vapors or gases onto a substrate. As a result of the self limiting reaction, the precursors only form a single monolayer per cycle. The sequential and self-limiting nature of the deposition is used to deposit thin films with good compositional control, high conformity, high uniformity, and excellent thickness control.

To create multiferroic crystal structures, a 1:1 stoichiometric ratio between cations is desired with low contamination by organic ligands in order to form the crystal phases that permit multiferroicity. Therefore, in this work, multiferroic YMnO3 and BiFeO3 on various substrates are synthesized by radical enhanced atomic layer deposition (RE-ALD) using Y(mhdx) (mhdx = 2,2,6,6-tetramethylheptyl-pentane-3,5-dione), Mn(mhdx), Fe(mhdx), and Bi(mhdx) as metal precursors and oxygen radicals as the oxidizer. By varying the cycle sequences, controlled composition is demonstrated and verified through XPS. Growth rates are shown on a thickness per cycle basis as a function of deposition temperature, precursor pulse times, and substrate. The crystal structure as well as ambient environment are examined by XRD and extended x-ray absorption fine structure spectroscopy (EXAFS) respectively and are accompanied by TEM micrographs. Finally, magnetometric measurements made using a super conducting quantum interference device (SQUID) magnetometer, zero-field cooled and field cooled (ZFC-FC) M vs. T and M vs. H are shown on 1:1 YMnO3 stoichiometric films showing a Néel temperature TN = ~45 K and a coercive magnetic field Hc = 130 Oe for Sf(111) and Hc = 500 Oe for YSZ(111).

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References:

We have previously reported on fully perpendicular Co/Pd multilayers (ML)-based CoFeB/MgO/CoFeB magnetic tunnel junctions (MTJ’s). However, Co/Pd ML-based MTJ’s have rarely exhibited TMR ratios greater than about 10%. This has been attributed to the inability to pull a sufficiently thick CoFeB layer perpendicular on top of MgO as well as the incomplete capping of CoFeB from MgO owing to the adjacent fcc Co/Pd ML. Recent results have generated great interest in MTJ’s with pinned perpendicular synthetic antiferromagnets (SAF), of the form AP1/Ru/AP2 where AP1 and AP2 are Co-based multilayers, for instance, Co/Ni or Co/Pd. We report on fully perpendicular MTJ’s with a thin CoFeB free layer and a Co/Pd(Pt) ML-based SAF pinned layer. For Co/Pd ML SAF’s, strong antiferromagnetic coupling was seen at 1Ru of 1.1nm, with a coupling strength of 0.017 mJ/m2. For Co/Pt ML SAF’s the optimum antiferromagnetic coupling was found at slightly higher Ru thickness of 1.3 nm, with a coupling strength of 0.013 mJ/m2. Improved MTJ properties are expected from using a thin Ta-seeded CoFeB bottom free layer, along with a thin, amorphous Ta layer used to transition from bcc CoFeB to fcc Co/Pd(Pt) for the top pinned layer. The full stack is of the form: bottom Ta2(2)/CoFeB(1)/MgO(0.6)/CoFeB(0.8)/Ta(0.3)(Co(0.3)/Pd(1))/CoFeB(0.5)/Ru(1) or 1.3/(Co(0.3)/Pd(1)) or Co(0.5)(Pt(2))/CoFeB(0.3)/Co(0.3) or Pd(1) or Ru(1.1) or 1.3/(Co(0.3)/Pd(1)) or Co(0.5)(Pt(2))/9/top lead. CPT measurements indicated TMR values as high as 20% for as-deposited stacks. Magnetometry of blanket stacks showed a large separation in the switching fields of free and pinned layers, with free layer switching close to zero field and pinned layer switching at 0.8-1.8 kOe (Fig.1). This symmetric extended plateau of constant magnetization offers a large dynamic range over which the magnetic configuration remains stable. The stacks were patterned into MTJ’s, annealed at 240°C in an in-plane field of 0.5 T, and characterized magnetically and electrically.

Acknowledgements:
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References:

INVITED

Phase change memory (PCM) and spin-torque magnetic random access memory (ST-MRAM) have recently garnered significant interest for future non-volatile memory applications due to their promise for scalability beyond that of conventional DRAM and flash memory technologies. In addition, both PCM and ST-MRAM have potential to enable improvements in programming speed, low voltage operation and high endurance, as compared to current devices. However, the introduction of each technology not only brings with it new device challenges, but the fabrication process itself can also alter the properties of the materials, leading to performance degradation. In this work, we explore the effects of integration and processing on Ge2Sb2Te5 (GST) and CoFeB for PCM and ST-MRAM devices respectively, with relevance to scaling the 14nm node and beyond. Using multiple analysis techniques including time resolved laser reflectivity and X-ray diffraction, we determine that exposure to various etch and ash plasma chemistries can cause a significant reduction in the (re)crystallization speed of GST, while concurrently increasing the transition temperature from the rocksalt to the hexagonal phase. Furthermore, by utilizing optical emission spectroscopy, X-ray absorption spectroscopy (XAS) and X-ray photoelectron spectroscopy (XPS) we are able to link the changes in crystallization to etch-induced modification of the GST stoichiometry. For CoFeB materials, vibrating sample magnetometry is used to determine the effect that plasma exposure (as used for back patterning and encapsulation) has on the magnetic properties. We find that even when relatively thick capping layers are present, magnetic moment can still be significantly degraded. The severity of this effect is highly dependent on plasma chemistry. Further results of process-induced material modification and device degradation as a function of etch, encapsulation, and thermal processing will be presented, focusing primarily on the impact on scaling of these technologies. Finally, potential integration and processing solutions to circumvent these issues will also be discussed.

Thin Film Division
Room: 110 - Session TF2+EM-WeA

Nanostructuring Thin Films
Moderator: A.V. Melechko, North Carolina State University

2:00pm TF2+EM-WeA1 Templated Solid-State Dewetting for Patternning of Films, C.V. Thompson, J. Ye, A.L. Giemann, Massachusetts Institute of Technology

INVITED

Most crystalline thin films are metastable in the as-deposited state, and will dewet to form islands when they are heated to temperatures that lead to sufficiently high atomic mobilities. This can happen well below the melting temperature of the film, so that the material remains in the form of a crystalline solid throughout the dewetting process. When unpatterned films dewet on flat substrates, the resulting islands have widely varying sizes and spacings. However, we have shown that substrate patterning can be used to control solid-state dewetting processes and to produce ordered arrays of monodispersed and crystallographically aligned islands from polycrystalline films. Recent experiments on patterned single-crystal films have allowed independent study of the various mechanisms that control structure evolution during solid-state dewetting. These include fingering instabilities, edge faceting, corner instabilities, pinch-off processes, and Rayleigh-like instabilities. Surface energy anisotropy plays a very important role in these processes, and for single-crystal films, leads to the formation of crystallographically aligned complex patterns of lines and islands that can be reproducibly controlled through pre-patternning.

2:40pm TF2+EM-WeA3 Dynamics of Solid Thin-Film Dewetting in the Silicon-On-Insulator System, E. Bussmann, F. Cheynis, F. Leroy, P. Müller, CINaM-CNRS, France

Thin-film dewetting is a process wherein a film on a substrate spontaneously agglomerates into 3D islands, which in some instances are ordered. A detailed understanding of the mechanism and dynamics of dewetting is crucial, either to avoid the agglomeration, or to engineer organized arrays of nanostructures. Silicon-on-Insulator (SOI) films, which are promising substrates for microelectronics, undergo dewetting when annealed at >700°C under ultrahigh vacuum conditions. The Si film spontaneously transforms into an assembly of ordered nano-sized Si islands. Previous ex-situ studies of dewetted SOI films provided a qualitative description of the dewetting process [1-4]. However, the dewetting dynamics, as well as the thermodynamic driving forces and atomistic mechanisms at work, remained largely unclear. We simultaneously measure the real-time dewetting dynamics and the motion of surface atomic-steps (surface self-diffusion) using low-energy electron microscopy (LEEM) [5]. We observe the following scenario: (i) dewetting voids nucleate at defects in the Si(001) layer. In the early stages of dewetting, the area of the opening voids grows linearly with time, and the Si ejected from the voids accretes into a rim surrounding the dewetted area. (ii) As dewetting progresses, the rim undergoes an instaibility that leads to the formation of elongated Si fingers. Once the first fingers have formed, the void area grows as the square of time. (iii) Finally, the Si fingers undergo a Plateau-Rayleigh instability, breaking apart into 3D Si nano-islands. We compare our measurements of the morphological evolution of dewetting to a simple analytical model for dewetting void growth (based on surface diffusion, nucleation on the top of the 3D structures, and mass-conservation), and to Kinetic Monte Carlo simulations. The KMC simulations reproduce the qualitative features of the complex void shape evolution in detail, while the analytical model of void growth allows us to connect the void growth rate with the dewetting driving force. These approaches unambiguously show that the SOI dewetting process is surface-diffusion-limited and driven by surface and interface free-energy-minimization.


3:00pm TF2+EM-WeA4 Self and Directed Assembly of Thin Metallic Films by Pulsed Laser Induced Dewetting, Y. Wu, University of Tennessee, J.D. Fowlkes, Oak Ridge National Laboratory, L. Kondic, New Jersey Institute of Technology, J. Diez, Universidad Nacional del Centro de la Provincia de Buenos Aires (UNCIPBA), Argentina, N.A. Roberts, P.D. Rack, University of Tennessee

The synthesis and assembly of functional metallic nanomaterials is critical for realizing many important applications of nanoscience and nanotechnology, and metallic thin film dewetting has been an effective and low-cost approach to this end. In this study, we investigated dewetting of metal thin films via pulsed nanosecond laser melting. We also explore the dewetting and nanopattern formation of nanolithographically pre-patterned thin films of various shapes to understand initial and boundary conditions in guiding the assembly. More recently, nanolithography was used to impose the perturbation which ultimately led to an organized nanoparticle array. Specifically, liquid-phase pulsed laser induced dewetting (PLiD) was used to transform metallic thin film strips into nanoparticle arrays. We demonstrated that the assembly accuracy and precision could be drastically improved by merely imposing a synthetic sinusoidal perturbation onto the lateral surfaces of the thin film strip. The synthetic perturbations in the strip translated into an unstable varicose oscillation on the rivulet during retraction – a precise nanoparticle diameter and pitch emerged thereby superseding the otherwise naturally evolving modes predicted by the modified Rayleigh-Plateau instability. A nanoscale, synthetic perturbation was usefully imposed to “nudge” the natural, self-assembly dispersion toward significantly higher order.

4:00pm TF2+EM-WeA7 Directed Self-Assembly of Ge Heteroepitaxial Quantum Dots with sub-35nm Spacing, C. Petz, University of Virginia, D. Yang, J. Levy, University of Pittsburgh, J.A. Floro, University of Virginia

Artificially ordered Ge quantum dot (QD) arrays, where confined carriers can interact via spin coupling, may create unique functionalities such as spin-based logic gates. In the development of such arrays for quantum computing requires fine control over QD size and spatial arrangement on the sub-35 nm length scale. We employ fine-probe electron-beam irradiation to locally decompose ambient hydrocarbons onto a bare Si (001) surface. These carbonaceous patterns are annealed in UHV, forming ordered arrays of nanoscale SiC precipitates that serve as templates for subsequent Ge quantum dot nucleation via strain-induced self-assembly during heteroepitaxy. The nanoprecipitates effectively reduce the critical thickness for Ge QD formation to below the 3-4 monolayers typical of Stranski-Krastanov growth in the Ge/Si (001) system. Thus, Ge QDs in the Si-patterned regions nucleate prior to formation of randomly located QDs
in the unpatterned areas. It is critically important to ascertain the variability in Ge QD size and placement, and ultimately to determine the crystalline quality and interparticle small Ge dots on SiC nanoprecipitates. Using atomic force microscopy and cross-sectional transmission electron microscopy, we investigate the patterned surface morphology and internal structure of patterned QDs to develop a fundamental understanding of the Ge adatom behavior in the vicinity of local high lattice-mismatch nanoprecipitates. We find that Ge self-assembly at SiC sites depends on the QD spacing and diffusion. Local strain gradients from the SiC surface may repel Ge to higher lattice mismatched SiC sites. Support from the DOE Office of Basic Energy Sciences is gratefully acknowledged under grant number: DE-FG02-07ER46421.


Unusually shaped micron- and nanometer-sized particles are becoming key components in catalytic and biological applications. Zeolite particles are typically modified for ion-exchange and catalytic applications. Fe₂O₃ nanoparticles are widely used in biological applications such as MRI imaging and site-specific drug delivery, thereby creating a need for surface functionalization techniques to ensure biocompatibility. Plasma processing can effectively modify and implant functional groups onto flat substrates, and is non-directional. Thus, the primary focus here is on applying these plasma processes to unusually shaped materials and understanding the gas-phase chemistry and surface reactions that make these coatings viable. Previous work in our lab employed plasma-enhanced chemical vapor deposition to create composite SiO₂/Fe₂O₃ nanoparticles. Here, we use plasma processing methods for the modification of zeolite surfaces and functionalization of Fe₂O₃ nanoparticles. Compositional and morphological data demonstrate that the conformal treatment of particles was achieved and that the use of PECVD methods allowed for advanced control over surface modification and specific tailoring of the structure, composition, and growth characteristics of any deposited film. Insight into the modification and deposition process is provided by atomic optical emission spectroscopy (AOES) and laser induced fluorescence spectroscopy (LIF), which allow characterization of the gas-phase species and their energetics (i.e. internal energies) for each system. To further investigate the functionalization of nanoparticle surfaces, additional studies explore the contributions of gas-phase OH radicals to the creation of SiO₂/Fe₂O₃ composite nanoparticles. Scattering coefficients and gas-phase density measurements derived from our imaging of radicals interacting with surfaces technique (IRIS) provides additional insight on the molecular-level chemistry occurring at the interface between gaseous plasma species and nanoparticle substrates. The operation and design of an in-house rotating drum reactor will be discussed as a potential method for adapting the composite nanoparticle fabrication to an industrial scale.

4:40pm TF2+EM-WeA9 Role of Ion Flux on Alignment of Carbon Nanofibers Synthesized by DC Plasma on Transparent Insulating Substrates, R.C. Pearce, North Carolina State University, A.W. Vasenkov, CFD Research Corporation, D.K. Hensley, M.L. Simpson, T.E. McKnight, Oak Ridge National Laboratory, A.V. Mel'chikov, North Carolina State University

A key factor to the implementation of vertically aligned carbon nanofibers (VACNFs) in devices is a more fundamental understanding of how to control fluctuations in the growth direction of the fibers. It has previously been hypothesized that the electric field is the primary factor in determining the orientation of fibers grown catalytically using plasma enhanced chemical vapor deposition (PECVD). Here we present results of carbon nanofiber synthesis on insulating substrates by de plasma in the vicinity of grid electrodes. To perform these observations, VACNFs were grown on a fused silica substrate using direct current PECVD. To maintain continuous glow discharge above the substrate, a metal grid electrode layer (Cr) was deposited over silica with windows of exposed silica ranging in size from 200 µm to 1 mm. Observed trends in nanofiber alignment at the window-electrode interface suggests that the alignment is governed by the direction of the ion flux rather than the electric field at the substrate level. The proposed alignment mechanism is that ion sputtering of the carbon film on a catalyst particle the growth direction of the nanofibers. With this development, fiber growth direction can be better manipulated through changes in ionic flux direction, opening the possibility for growth of nanofibers on substrates with unique geometries.

5:00pm TF2+EM-WeA10 High-Performance Poly-3-alkyliophene-Carbon Nanotube Composites for Transparent Electrodes, S.E. Hellstrom, R.Z. Jin, R.M. Stoltenberg, Z. Bao, Stanford University

Flexible transparent electrodes are crucial for flat panel display and solar cell technologies. While carbon nanotube network electrodes show promise, their fabrication often involves insulating surfactants which worsen conductivity. As an alternative, we show that small amounts of conjugated semiconducting polymer added to nanotube dispersions enables straightforward deposition of uniform electrodes by spin-coating or drop casting. After doping, electrodes as good as 120 ohm/sq with 81% transmittance at 550 nm are obtained. Tuning system chemistry and deposition parameters allows control of tube bundle size, density, and alignment, and these may be correlated with electrode performance.

We also employ the selectivity of P3HT:CNT composites in wetting of and adhesion to different dielectric surfaces, to simultaneously fabricate and pattern organic electrodes with unprecedented nanotube density gradients and highly-tunable feature resolutions. We employ these to drive the vertical alignment and C60 transistors that are competitive with those made using Au electrodes. We thereby suggest that this material system and fabrication technique has promise in applications requiring flexible, semitransparent, low-cost complementary circuits.


Silicon-Germanium semiconductor alloys are important in improving Group IV opto- and microelectronics. Typically, SiGe is used as a stressor to introduce strain into Si, thereby altering the electronic band structure [1, 2] or changing optical properties [3]. The SiGe stressor can be incorporated locally to strain small areas of Si uniaxially [2], or relaxed SiGe can be used as a uniformly strained Si layer for growth of SiGe quantum dots over large areas [1]. We will focus on the latter: using relaxed SiGe to induce global biaxial strain in Si. Conventional methods for creating relaxed SiGe substrates involve epitaxial growth on Si substrates and relaxation of the alloy via dislocations. The density of defects that reach the top relaxed SiGe layer can be limited though various techniques [4], but strain inhomogeneities and surface roughness created by the dislocations remain. We demonstrate the fabrication of SiGe nanomembranes (NMs) exhibiting defect-free, smooth, single-crystalline sheets of SiGe alloy. A thin SiGe layer (less than the kinetic critical thickness for dislocation formation) is grown on a silicon-on-insulator (SOI) substrate with molecular beam epitaxy (MBE), followed by a Si capping layer of similar thickness to the Si template layer of the SOI. The SiO₂ layer of the SOI is selectively etched away, leaving the Si/SiGe/Si trilayer heterostructure free to strain share [5]. The Si layers of the trilayer are then selectively etched away, leaving a fully elastically relaxed SiGe NM. These SiGe NMs can be transferred to new handling substrates, bonded, and used as templates for growth of new defect-free materials.

A specific application involving strained Si/relaxed SiGe heterostructures is fabrication of 2-dimensional electron gas devices (2DEGs) that can be patterned and gated to confine individual electrons into quantum qubits with long coherence times. These devices are attractive for their demonstrated changes in the electrostatic potential and thus require superb material quality. We compare the material quality of strained Si/SiGe heterostructures grown on SiGe NMs with those created on SiGe substrates relaxed via dislocations. Research supported by DOE. Facilities support from NSF-MRSEC is acknowledged. DMP is supported by a NSF Graduate Research Fellowship.


5:40pm TF2+EM-WeA12 Multilayer Barrier Coatings for Organic Photovoltaics, A.M. Coclite, K.K. Gleason, Massachusetts Institute of Technology

Barrier coatings, which prevent the permeation of water into OPV devices fabricated on flexible plastic substrates, are essential to extend the device lifetime. Such protective coatings are made of multilayer stacks where multiple dense, inorganic layers are alternated with soft, organic ones. The inorganic layer contains inevitably some pinholes and defects. The roles of the organic layer are (i) creating a tortuous and longer path around the defects of two successive inorganic layers (ii) filling the pores of the inorganic underlayer limiting the propagation of defects from one inorganic layer to the other and (iii) smoothing the substrate surface roughness.

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In the past, we obtained good barrier properties (WVTR=10^{-2} g/cm²/day) with a hexamethyldisilazane (HMDS) polymer using a plasma-enhanced chemical vapor deposition (PECVD) process. For metal deposition, a non-stick surface is critical to reduce friction and wear. For this reason, we developed a new fluorinated alkynyl-derived SAM (self-assembled monolayer) on Si(111) surfaces. This SAM was prepared by covalently bonding a fluorinated alkynyl monolayer on the Si(111) surface. The resulting surface is highly hydrophobic and exhibits excellent friction properties.

A combination of spectroscopic (XPS, IR), microscopic (AFM), and contact angle measurements shows how the monolayer deposition on Si(111) surfaces results in the formation of a highly hydrophobic and low-friction surface. These results are also supported by in-situ friction measurements, which show a significant reduction in friction coefficient for the fluorinated monolayer compared to the bare Si(111) surface. The mechanism for this reduction in friction is currently under investigation.

The deposition of the fluorinated monolayer on Si(111) surfaces can be achieved using a combination of plasma-enhanced chemical vapor deposition (PECVD) and thermal evaporation techniques. The resulting monolayer is highly hydrophobic and exhibits excellent friction properties. These results are in agreement with previous studies, which have shown that fluorinated monolayers can significantly reduce friction and wear.

To summarize, the development of a new fluorinated alkynyl-derived SAM on Si(111) surfaces provides a promising approach for the design of low-friction and wear-resistant surfaces. Further work is needed to optimize the deposition conditions and to investigate the long-term stability of the fluorinated monolayer on Si(111) surfaces.
extend the operational life indefinitely and to potentially negate any mechanical contributions to wear, a stable lubricating tribofilm of sufficient chemistry and thickness must be maintained during operation. However, because sliding occurs in a buried interface, it has proven challenging to determine what materials processes are actively enabling stable performance and/or what to add to the system to improve lubrication. In this talk, results from a number of ultra-low wear systems (polymers, metals, and ceramics) that have been studied using a variety of active and in situ tribological instrumentation will be presented, along with a discussion of the various mechanisms that we believe to be responsible for this unique behavior.

5:00pm TR-WeA10 Study on the Fatigue Wear Behaviour of TiN and WC DLC-coated Stainless Steel under Inclined Impact-Sliding Load Tests. Y. Chen, X. Nie, University of Windsor, Canada

WC/C diamond-like carbon (DLC) coatings and Titanium Nitride (TiN) coatings are widely used in industrial machinery and tools. Both of the two coatings have extremely hard surfaces comparing to the stainless steel substrate SS316L. The W: DLC coating gives low coefficients of friction against a number of counterfaces, has relatively high lubricity and resistance to adhesive wear comparing to the TiN coating. In this study, a new method which is called cycled inclined impact-sliding test is introduced and utilized to study the coating durability under a combined force of an Impact force Fi and Pressing force Fp (Fi/Fp=200N/400N and 200N/200N) and the fatigue wear behaviors after up to 1000 cycles impact tests. A 10mm steel (AISI 52100) bearing ball is used as the impact indenter. Due to the low coefficient of friction of W-DLC coating against steel counterface, greater impact cycle was endurable before the failure of the coating when impacted in dry air condition. Under inclined impact and sliding forces, fatigue cracking was first initiated, followed by chipping and pealing of the coatings. The SEM showed that different types of fatigue wear cracks were found which distributed in the different areas (head and tail parts) of the impact scars. Most of those scars, observed by 45º tilted cross-sectional SEM, penetrated the coating and caused hardening of the under layer (interface layer). Material transfer from the indenter ball could also be detected by EDX in some areas of the impact scar on both of the coating materials.


In an effort to reduce the friction between sliding components scientists and engineers have developed a multitude of lubrication schemes. One of the most intriguing concepts is referred to as 'structural lubricity', where flat surfaces are thought to slide past each other virtually frictionless if their atomic structures are incommensurate. In this talk, we analyze the fundamental mechanisms that govern the area-dependence of friction in extended but atomically flat contacts of dissimilar materials with a particular emphasis on the relation between structure (crystalline vs. amorphous) and friction. The resulting sublinear power laws, which link mesoscopic friction to atomic principles, are then confirmed by measuring the sliding resistance of gold and antimony particles on graphite [1,2]. The findings suggest that engineering surfaces with unprecedented low friction can be realized.


5:40pm TR-WeA12 Auger Surface Analysis of Deposits Formed on Magnetic Tape Recording Head Surfaces. F.E. Spada, University of California, San Diego, D.F. Paul, J.S. Hammond, Physical Electronics

The 2008 International Magnetic Tape Storage Roadmap\1 projects that the total magnetic spacing between the recording head and the tape magnetic layer must decrease from the current 43 nm spacing to about 23 nm by the year 2018 in order for tape to maintain its cost advantage as an information storage medium. Because tape drives are contact recording systems, interactions between head materials and components in the tape magnetic layer can detrimentally affect the head-tape separation via deposition formation on head surfaces as well as preferential erosion of critical recording head elements. Understanding the nature of these interactions is therefore essential for mitigating undesirable increases in the magnetic spacing. This study shows that deposition and erosion phenomena in tape heads can be varied at the local level by changing the electrical configuration of adjacent pole tip structures in multichannel heads, and that the composition of the head deposits depends on the electrical configuration of the pole tips. Using atomic force and electric force microscopy, we show that conductive deposits form on the “trailing edge” of pole tips which are electrically connected to earth ground or to the head substrate. The conductive deposits become non-conductive further “downstream” from the pole tips. Deposits adjacent to electrically isolated poles are always non-conductive. Auger analysis shows that the surfaces of the conductive deposit regions contain high levels of Fe and Co, and small amounts of P and Y, whereas the surfaces of the non-conductive deposits contain predominantly P and Y, with very low levels of Fe. Because all of these elements are present in the magnetic coating of the tape, and because the compositions of the deposits on heads having NiFe pole tips is similar to those on heads having CoZrTa pole tips, these results suggest that the deposits originate from components in the tape and not from metallic structures in the tape head.


*Supported by the Information Storage Industry Consortium Tape Program
Actinides and Rare Earths Focus Topic
Room: 207 - Session AC+SS-ThM

The Surface Science of Actinides and Rare Earths
Moderator: R. Schulze, Los Alamos National Laboratory

8:00am AC+SS-ThM1 The XPS of Heavy Metal Oxides: New Insights Into Chemistry. P.S. Bagus, University of North Texas, E.S. Ilton, Pacific Northwest National Laboratory, C.J. Nelin, Consultant

The XPS of Rare Earth and Actinide oxides are commonly used to obtain information about the oxidation state of the metal by taking various features of the spectra as fingerprints of the metal oxidation state. However, it is possible to obtain detailed information about the nature of the chemical interactions from these features by using the predictions of rigorous theoretical analyses. One of our important concerns is to make direct assessments of the covalent character of the metal-ligand interaction; i.e., the mixing of O(2p) with partly occupied, or unoccupied, metal levels to form bonding and anti-bonding orbitals. We relate this covalent character to the XPS features. In particular, we investigate the connection between the covalent character of the interaction and the satellite intensity. We also investigate the characterization of the satellites and discuss their assignment as shake-off satellites, assignment that is naturally consistent with the natural mixing of the metal and oxygen levels. Furthermore, we examine how vibrational excitations can lead to broadening of the XPS features and suggest that the observed broadening of XPS peaks may contain, hitherto not utilized, information about the chemical interactions in an oxide. Our focus will be on the XPS of two Rare Earth oxides, CeO2 and LaAlO3, and two actinide oxides, UO2 and PuO2, these systems have different electronic character that permits the mechanisms discussed above to be explored and compared. Our theoretical analyses are based on relativistic molecular orbital wavefunctions, Wf’s, for both initial states, before ionization, and final states, after ionization. The Wf’s are for materials models that contain explicit cations and anions embedded in a point charge field. With the variationally optimized orbitals for these Wf’s, covalent mixing is naturally taken into account. The cluster Wf’s include one-body and many-body effects and do not use parameters that are adjusted to make calculated relative energies and intensities fit to experiment.

Thursday Morning, November 3, 2011


An understanding of the oxidation and corrosion processes of plutonium metal at room temperature is important to the safe, effective use of this material. We are currently studying the oxidation of plutonium metal at room temperature. The oxidation rate for the δ-phase stabilized, plutonium/gallium alloy (a commonly employed alloy) can be significantly affected by a number of parameters including the gallium content and the composition of the oxidizing atmosphere (O2, O2/H2O, H2O). The nature of plutonium oxidation typically has been thought of as the growth of a dioxide (PuO2) overlayer on the metal to a thickness which the film begins to be explored and compared. Our theoretical analyses are based on relativistic molecular orbital wavefunctions, Wfs, for both initial states, before ionization, and final states, after ionization. The Wfs are for materials models that contain explicit cations and anions embedded in a point charge field. With the variationally optimized orbitals for these Wfs, covalent mixing is naturally taken into account. The cluster Wfs include one-body and many-body effects and do not use parameters that are adjusted to make calculated relative energies and intensities fit to experiment.

Historically, the oxidation/corrosion of plutonium has been studied by oxygen uptake of samples at elevated temperatures inferred from mass gain measurements. Accuracy of these experimental setups likely limited the oxygen uptake of samples at elevated temperatures inferred from mass gain measurements. It is unclear how pressure affects the oxidation process – particularly in the context of the formation of U3O8, with and without the δ-phase stabilized, δ-plutonium alloy. The gallium content relative to plutonium is δ-plutonium alloy. The gallium content relative to plutonium is


The oxidation of uranium dioxide has received much experimental and theoretical attention over the last several decades in large part because of its relevance to the operation and storage of uranium-based nuclear fuel. The oxidation process is inherently complicated, involving the formation of multiple different phases via distinct mechanisms even at relatively low temperatures. In the range of a few hundred degrees centigrade oxidation is generally assumed to be a two step process:[1] UO2 → UO3/2(UO2) → UO3. At low pressures the intermediate phases adopt crystal structures that are modifications of the UO2 fluorite structure and are slightly denser. By contrast, UO3 forms a considerably less dense orthorhombic structure (by some 23%). The large volume expansion resulting from the oxidation of UO2 to UO3 is a potentially serious concern in the event of oxidation of a fuel element, with consequent splitting of protective sheaths and the spalling of powder.

While attention has been focused on the oxidation of UO2 at elevated temperatures, the associated experiments have all been performed at low partial pressures of oxygen. It is unclear how pressure affects the oxidation process – particularly in the context of the formation of U3O8, with and without the large volume change with respect to UO2. We have examined the oxidation of a nominal single crystal of UO2 in pure oxygen at elevated pressures up to approximately 0.9 GPa (9000 atm) and temperatures of up to 450 °C. In situ Raman scattering measurements were made as a function of temperature in order to monitor the oxidation. Recovered material was examined using electron based techniques including SEM, STEM, and electron diffraction and also using x-ray photoelectron spectroscopy. Material synthesized under high pressure has a Raman spectrum that is different from both the UO2 starting material and the common form of U3O8. Also, compared with common U3O8, we find that it has fewer crystalline defects and mostly adopts a hexagonal rather than orthorhombic form. Figure 1 of the supplemental document compares Raman spectra of UO2 in oxygen with synthesized material. Fig. 2 compares electron diffraction obtained from recovered material with U3O8 synthesized at ambient pressure.

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Preferential uranium hydriding occurs frequently in narrow gaps. There are different hypotheses about its causes, one of which could be radiation-induced chemistry in gaps. Both 235U and 238U generate ionizing radiation capable of vibrationally or translationally exciting, ionizing, or dissociating H2. Here we use the MCNPX radiation transport code to calculate the energy dependent electron flux generated from the 235U, 238U, and 238U decay and from photoelectrons generated by brems-strahlung. We apply the code to gaps occurring in two 235U cylindrical pieces welded together and filled on the inside with LiD with a 100µm gap between 235U and LiD, and having a 100µm gap in 238U itself, typical of a joint welded uranium shell. The MCNPX Monte Carlo Code – as configured now tracks the life cycle of electrons throughout the material and calculates the electron flux as a function of energy, putting results into “energy bins” 1 keV wide. We find that at 2 keV (±5 keV, the last energy bin) the calculated electron flux in the U-U gap is approximately 19 times larger than in the U-LiD gap, and fifty two times larger than in the U-air gap on the outside of the cylindrical

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shells. Cross-sections for electron-hydrogen collisions peak, however, below 1 keV energy. We establish the upper limit of the effect of electron-hydrogen collisions by extrapolating the MCNPX electron flux results from the last bin to energies as low as 1 eV by fitting a function to the flux between 2 and 20 keV. To calculate the fraction of H\textsubscript{2} vibrationally or translationally excited, ionized, or dissociated per cm\textsuperscript{3}s, we integrate the product of the energy dependent cross sections (listed in reference [2]) and the energy dependent electron flux over the relevant energy range. The fraction of H\textsubscript{2} molecules calculated to be dissociated is small, but significant during long-time exposure. Future work will extend the MCNPX code below 1keV (as is done for biological radiation damage), to avoid energy extrapolation.

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References


Recent studies showed a colossal enhancement in the ionic conductivity of multi-layer oxygen-ion conducting thin films compared to most commonly used solid oxide fuel cell (SOFC) electrolytes. It has been observed that, the oxygen ionic conductivity of nano-scale hetero-structures increases with the increasing number of layers. However, some of these results were questioned due to the inability to distinguish electronic and ionic conductivity. Thus, here we investigated the scandia stabilized zirconia and samaria doped ceria (SDC/ScSZ) multi-layer system to understand the oxygen-ion conductance in multi-layer hetero-structures.

In this study, the growth of SDC and ScSZ multi-layer thin films was carried out using the optimized growth conditions, dopant concentrations and film properties established for single-layer SDC and ScSZ epitaxial thin films. The epitaxial SDC and ScSZ multi-layer thin films were grown on Al\textsubscript{2}O\textsubscript{3}(0001) substrates by oxygen plasma-assisted molecular beam epitaxy (OPA-MBE) at 650°C. The number of layers in the multi-layer hetero-structures was varied from 2 to 20 by keeping the total film thickness constant at 140 nm. Following the growth, thin films were characterized by various in-situ and ex-situ characterization techniques including reflection high energy electron diffraction (RHEED), x-ray diffraction (XRD), Rutherford backscattering spectrometry (RBS), and x-ray photoelectron spectroscopy (XPS). The initial streaky RHEED pattern indicates the epitaxial growth of SDC/ScSZ multi-layer thin films. Furthermore, RHEED patterns indicate the transition of the initial smooth surface to a rough surface with the increase in number of layers. The individual layer thickness was found to be approximately 7 nm for the twenty-layer film as confirmed by x-ray reflectivity data. RBS was also used to find the composition and thickness of the films. HRXRD patterns of ScSZ/SDC thin films exhibit only CoO\textsubscript{3}(111) and ZrO\textsubscript{2}(111) reflections, indicating the growth of epitaxial SDC(111) and ScSZ(111) multi-layers. XPS depth profile confirmed the uniform dopant concentration in both SDC and ScSZ layers, which was found to be 7 and 6 atom % for Sc and Sm, respectively. In the near future, the ionic conductivity of SDC and ScSZ multi-layer thin films will be measured by four probe conductivity method.

Applied Surface Science Division Room: 102 - Session AS-ThM

Analysis of Insulators and Challenging Samples Moderator: D.J. Gaspar, Pacific Northwest National Laboratory

TOF-SIMS has emerged as an important tool for imaging mass spectrometry of biological samples due to its unique capability to detect molecular ion fragments and elemental ions at sub-micron spatial resolution and without the sample treatments required by e.g. the MALDI technique. For many biological specimens, the ability to image samples having a large degree of surface topography is also highly desired. The resulting elemental and molecular images provide important information regarding the composition of biointerfaces, for example between plant tissues and their natural environment. This TOF-SIMS study involves the differentiation and characterization of epicuticular waxes present at the surfaces of Arabidopsis thaliana organs including the flower, stem, adaxial (top) leaf surface and the abaxial (bottom) leaf surface.

High spatial resolution images obtained from a flower petal, an adaxial leaf surface, and an abaxial leaf surface revealing spores, trichomes and responsive pores, respectively were acquired using TOF-SIMS to image molecular ion fragments with a spatial resolution of < 0.3 microns. Total ion images, and molecular fragment ion images of epicuticular wax components, demonstrate the capability to image entire organ surfaces without topographical artifacts. High mass range spectra in both the positive and negative modes were generated using TOF-SIMS.
and negative secondary ion polarities reveal that the epicuticular surface of each Arabidopsis thaliana organ is comprised of distinct wax components. Mass spectra acquired from specialized cells forming the 1 μm × 7 μm respiratory pores of the abaxial leaf surface indicate that, even within a single organ surface, the epicuticular wax composition may vary. The differences in wax composition on each of the interrogated organs of Arabidopsis thaliana will be presented and discussed. Structural assignments for characteristic mass spectrometric features related to the wax composition will also be presented and discussed.


This research provides the first in situ detection of aqueous surfaces using time-of-flight secondary ion mass spectrometry (ToF-SIMS). ToF-SIMS provides molecular information with super-low detection limits, making it a great choice to detect short-lifetime chemical reaction intermediates in aqueous solutions. However, it is difficult to detect aqueous solutions by ToF-SIMS because ToF-SIMS is a high vacuum technique. The vapor pressure of water is about 2-3 kPa at room temperature. We designed and fabricated a self-contained microfluidic module, allowing in situ analysis of aqueous surfaces by ToF-SIMS. Microfluidic channels were prepared on top of a block of polydimethylsiloxane (PDMS) using a patterned silicon template with soft lithography techniques, and then a ~100nm thick silicon nitride film was placed on top of the microfluidic channels, on which a small hole with a diameter of ~2-3 microns could be drilled using a focused primary ion beam in a ToF-SIMS instrument. De-ionized water, heavy water, or a 0.5% (weight ratio) formic acid aqueous solution, a 0.5% glutamic acid aqueous solution were successfully analyzed through the small hole using our ToF-SIMS instrument (IONTOF-V). The pressure in the analysis chamber was usually at ~5E-7 mbar during measurements, and this value was sufficient for ToF-SIMS analysis. Our device could be safely staying in vacuum for more than 8 hours, and 1 hour continuous SIMS measurements were practical. Molecular signals (M−H+) from formic acid (HCO2−), glycerol (CH3(OH)2), and glutamic acid (CH2NO2) were clearly observed in the small hole, but not elsewhere. Based on current data, the detection limits of formic acid, glycerol, and glutamic acid were estimated to be <0.01% (weight ratio).

8:40am AS-ThM3 Challenges of the XPS Analysis of the Ionic Liquid [BMIM][PF6], R.R. Hartl, B.D. Booth, G.K. Jennings, B.R. Rogers, Vanderbilt University

Room temperature ionic liquids (ILs) are ionic compounds that are liquids at ambient conditions due to a bulky, asymmetric organic ion that inhibits crystalline packing. The inhibition of crystalline packing results in ILs having a vapor pressure that is negligibly low. The properties of ILs make them candidates for applications such as nonvolatile solvents, electrolytes, and lubricants. The ionic liquid pair 1-butyl-3-methylimidazolium hexafluorophosphate [BMIM][PF6] is being investigated for use as a lubricant in micro-electrical mechanical systems (MEMS). Surfactant properties are important in tribological applications like lubrication and the nature of the surfaces and intentional as well as inadvertent coatings (shells) on nanoparticles. Some of the methodology and special capabilities developed in EMSL for controlling sample environments, the value of more rapid and enhanced data analysis, and the importance of procedural guidelines and standards will be noted. Although many of the topics discussed relate to a variety of surface analysis methods, the talk will focus on XPS and use TOF-SIMS to provide an example of challenges associated with measurement of hydrogen. The importance of complementary measurements by a variety of methods (including XRD, TEM, APT, SEM and RBS) will be highlighted.


Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently. Electron emissions from poorly conductive specimen stimulated by photon and electron injection results in charging phenomena. These can cause distortion of data in electron spectroscopy and image artifacts in scanning electron microscopy because surface potential which influence the motion of emitted electron is modified by charging and change time dependently.

Nafion based membranes are being developed for applications in fuel cells. These fuel cell membranes are multi-layer structures with a permeation membrane and two thin layers of noble metal in a polymer matrix acting as the anode and the cathode of the fuel cell. The cross-sectional chemical analysis of these membranes, both new and used, may be extremely useful in improving the performance of the fuel cells. In this study we will highlight new and advanced scanning XPS microprobe based techniques to provide cross-sectional line scans and high resolution chemical mapping of these membranes. The line scan and chemical mapping results elucidate the modifications of the membrane chemistry as a function of use.


Organic light emitting devices (OLEDs) have the potential to dramatically change the way we light our living spaces. OLEDs offer the potential for high efficiency lighting from a large area source, with the possibility of both transparent and flexible lighting as well. Currently, OLED displays are found commercially in cell phones and in televisions in limited production. One of the key barriers holding back widespread adoption of OLEDs for both lighting and other products is the challenge of generating long-lived devices, particularly for high-efficiency phosphorescent blue OLEDs (a necessary component for a high-efficiency white OLEDs). Challenges in the design of stable materials are hampered by our relatively poor ability to measure degradation products in situ, or in ways that ensure we have not damaged the material in the measurement process. To this end, we have generated thin films of single blue phosphorescent OLED components, multilayer films, and working OLEDs before and after aging. These components including the well-studied bis[2,4,6-difluoro-phenyl]pyridyl-N,C2′′Iridium (III) (F1rpic), 2,8-bis[(diphenylphosphoryl)dibenzo thiophene (PO15), 4-(diphenylphosphoryl)-NN-diphenylaniline (HM-A1) and di-[4-(N,N-ditolyl-amino)-phenyl]cyclohexane (TAPC). Based on previous experiments in our lab and literature reports, F1rpic and PO15 are known to undergo degradation reactions under conditions similar to those used in this series of experiments. Analysis of these films and systems by gas cluster ion beam secondary ion mass spectrometry (GCIB-SIMS) and other methods show degradation products, which permit the development of useful models for their degradation pathways. These results clearly demonstrate the utility of GCIB-SIMS in the characterization and analysis of OLED devices and materials, and should enable better design and screening of stable materials for high-efficiency OLEDs.

Biomaterial Interfaces Division
Room: 108 - Session BI-ThM

Biomedical Materials
Moderator: S.L. McArthur, Swinburne University of Technology, Australia

8:20am BI-ThM2 Surface Characterization: A Critical Component in Understanding the Biocompatibility of Biomaterials. L. Salvati, S. Vass, DePuy Orthopaedics INVITED

A biomaterial is defined as “any synthetic material or device—e.g. implant or prosthesis-intended to treat, enhance or replace an aging or malfunctioning native tissue, organ or function in the body.” David Williams, states that “Biocompatibility is largely about the chemical interactions that take place between the materials and the body fluids, and the physiological responses to these reactions.” These reactions are dominated by the initial events at the molecular level, the interface, thus it is not hard to see the relationship between the implant surface properties and it’s in vivo. It is clear that the performance of a biomaterial is directly linked to the surface chemistry, composition and topography of the device. However, despite the preponderance of evidence, biomedical device companies as a whole do not utilize surface analytical methods nearly enough. Why is that?

In orthopaedics, most of the implants are fabricated from metals which directly contact biological fluids that are typically complex aqueous mixtures. Consequently corrosion and/or corrosion prevention are important considerations in device manufacturing. For the most part, orthopaedics manufacturers utilize the same passivation methodologies used to impart corrosion resistance to stainless materials. Numerous researchers have characterized the impact of acid passivation on stainless materials, but little has been done to study the effect of these treatments on the CoCr alloys. One of the topics in this presentation deals with the effects of passivation on CoCr alloys. Specifically, the presentation will correlate specific surface treatments with surface chemistry and ultimately to metal ion release. The study will also address questions related to the use of delicate materials such as oxygen replacement for nitric acid passivation. There is no argument that the process is much greener, but there are plenty of questions as to its effectiveness. Considering that the “changes” imparted by the passivation solutions effect the outer most 10-100A, the only means to characterize the affects of passivation requires surface analysis methodologies. The importance of surface analysis methods, especially XPS, will be highlighted by this example.

In addition to the discussion of metal alloy passivation, this presentation will also deal with other surface related issues that could impact the biocompatibility of biomaterials. The talk will show the potential impact of packaging materials and cleaning processes on the surface chemistry and composition of biomaterials.


Controlled release of active pharmaceutical molecules from biocompatible polymers over defined time periods is an area of intense study. Present applications include drug eluting stents and other drug delivery systems. One of the most important parameters which govern drug dosing is the drug concentration depth profile in the supporting polymer matrix. In a previous study we have shown that combining X-ray photoelectron spectroscopy (XPS) with a coronene ion source is a very powerful tool for investigating the drug distribution with depth of a model system [1]. The use of cluster ion sources for sputter depth profiling of thin film or multilayer organic materials during XPS analysis has become routine. A wide range of organic systems are amenable to profiling and there is a good understanding of the experimental parameters which contribute to successful analysis. Here we report on extending the aforementioned study to materials which closely resemble real world samples intended for use in vivo.


RF magnetron sputtering of nylon 6.6 was used for the deposition of nitrogen rich films. Deposition was followed by N2 H2 plasma post-treatment to enhance primary amine concentration on the surface. Maximal reached NH3 concentration was 11 % with aminoslectivity 13.5 %. The films exhibited small negative z-potential at basic pH with isoelectric point ~ pH 4.5. Bio properties of the films were tested by QCM in terms of ability to adsorb different proteins and their antigens. The interaction between the film and the buffer solution was also studied and compared to other films such as poly-acrylic acid and PEO plasma polymers. The results show that these films are stable and can be used as a platform where positively charged surfaces are needed.

10:40am BI-ThM9 Surface Analysis in Biotech & Pharma: A Surfet of Frontiers. E. Johnston, Genzyme INVITED

Surface and interfacial analytical tools continue to provide new value and find unexpected new uses in the biotech and medical device industries. Some uses are investigational in nature and help solve critical problems within manufacturing and quality control. Other applications fall squarely within the realm of R&D - tilting the balance between feasibility of a product or obsolescence of a project, or providing fresh insight into the nature of biomaterial/biological interactions. By way of example, a study will be presented illustrating how TOF-SIMS was used to image a phosphate-binding drug particle in the complex matrix of the rat gastrointestinal tract. Sample preparation was challenging due to the highly hydrated nature of the tissue material. The results yielded surprising information about the ions that bind to this cationically charged particle and opened new avenues for inquiry and study.
that heparin behaves identically to H2O on hydrophobic surfaces. Heparin, present during surgery, has to be investigated. Our results show coagulant agent in the eye. Thus fibrinogen can indeed prevent fogging, but removed after application on the IOL’s, thus remaining as a potent IOL’s in various dilutions does prevent fogging. However, it cannot be failure inferred to be due to blood proteins on IOL’s.

The work solves the problem by modifying water affinity of IOL’s using a condensation of bodily fluids. New, high performance accommodating secondary eye surgery. Secondary surgery performed after implantation of % of cataract patients suffer subsequen tly from diabetic and other retinal Over 15 million cataract surgeries are performed each year worldwide. 2-6

of stimuli-responsive polymer brushes were prepared on solid substrates by a “grafting-to” procedure. The brush properties (e.g. swelling/collapsing) as controlled adsorption and release of biomolecules. For that, different types of stimuli-responsive polymer brushes were coated on silicon wafers, carefully equilibrated and measured in dependence on temperature (up to 250°C) in an inert gas atmosphere. The Tg data were determined from the ellipsometric results and discussed.

In the second part, “smart” surfaces will be presented which can be used for controlled adsorption and release of biomolecules. For that, different types of stimuli-responsive polymer brushes were prepared on solid substrates by a “grafting-to” procedure. The brush properties (e.g. swelling/collapsing) as well as the resulting adsorption/desorption of model proteins (e.g. Human Serum Albumin, Chymotrypsin) can be switched in an appropriate aqueous medium with temperature (PNIPAAm) and/or pH (PAA-P2VP). The corresponding processes at the solid-liquid interface were studied in-situ by spectroscopic Vis-IR-ellipsometry 3,4/.

Thus, the role of hydro-affinity of blood proteins preventing coagulation, heparin, present during surgery, has to be investigated. Our results show that heparin behaves identically to H2O on hydrophobic surfaces. Heparin simply de-wets on silicone IOL’s and hydrophobic acrylic lenses. It does not prevent fogging on IOL’s nor interfere with our anti-fogging emulsion. Fibrinogen is the other protein investigated because it enhances blood coagulation and is often present in trauma situations. Fibrinogen applied to IOL’s in various dilutions does prevent fogging. However, it cannot be removed after application on the IOL’s, thus remaining as a potent coagulant agent in the eye. Thus fibrinogen can indeed prevent fogging, but is not viable since it cannot be removed after application like Vitreox™. Fibrinogen could explain why some IOL’s fog while others do not during emergency secondary eye surgery.

Vis-Ellipsometry (SE) is not only possible to investigate optical properties of these films but also the composition of it, e.g. volume fraction of Fe3O4-NP.

We studied two types of films and present results for a thin NP-composite film prepared by pre-mixing of the PNiPAAm with hydrophobic Fe3O4-NP, spin-coated and grafted to a silicon substrate (System 1) and a film prepared by adsorption of hydrophobic functionalized Fe3O4-NP onto PNiPAAm brushes (System 2). The former system was chosen to develop an optical model starting from a simple two component effective-medium-approach (Maxwell-Garnett-EMA) using the optical constants of the pure polymer measured by SE and Fe3O4 with averaged data taken from three recent publications. The SE best-fit-results were validated against scanning electron microscopy (SEM), atomic force microscopy (AFM) and X-ray photoelectron spectroscopy (XPS).

After transferring this optical model to System 2, the adsorption of hydrophobic functionalized Fe3O4-NP onto PNiPAAm brushes was investigated and will be discussed with additional results obtained from contact angle (CA) and phase transition (LCST) measurements.

9:00am EL+AS+EM+MS+PS+TF-ThM4 The White Scarab Beetle Cyphochilus insulanus –Scattering and Polarization Properties. C. Akerlind, Swedish Defence Research Agency / Linköping University, Sweden, H. Arwin, Linköping University, Sweden, T. Hallberg, H. Kariis, Swedish Defence Research Agency, Sweden, J. Landin, K. Järrendahl, Linköping University, Sweden

Three methods were used to characterize the optical properties of the light scattering white scarab beetle Cyphochilus insulanus. Spectral directional hemispherical (DH) reflectance measurements in the wavelength region 250 nm to 25 µm were performed using integrating spheres. The general spectral appearance shows a relatively strong reflectance band in the range 400 - 1600 nm. The Bidirectional Reflection Distribution Function (BRDF) was measured over a semi-circle in the plane of incidence at the wavelengths 633 nm and 3.39 µm, using p- and s-polarized light. In the visible the BRDF data shows a near Lambertian behaviour with a constant BRDF for most angles, i.e. the light is diffusely scattered. For large incident angles and in the infrared the BRDF is more specular. Full Mueller-matrix spectroscopic ellipsometry (MMSE) measurements were performed in the wavelength range 250 - 1000 nm at angles of incidence between 45° and 75° using a dual rotating compensator ellipsometer. The Mueller data show that the detected light is partially polarized in general but there is no linear or circular polarization P even for nonpolarized incident light. The angular dependence of P was pronounced for incident p-polarized light and was very low near the Brewster angle. In comparison, P was high for all measured angles for incident s-polarized light. Close to the Brewster angle, the reflected light showed a high ellipticity for incident p-polarized light. The optical data is used to model the structural and optical response of the beetle cuticle. With input from the DH and BRDF-data analysis of the Mueller Matrix data was made using Fresnel-based layer modelling. 2-, 3- and n-phase models are compared. The obtained optical properties are also used to calculate color data in terms of chromaticity and whiteness. The scattering properties of the Cyphochilus insulanus cuticle are also discussed by combining the BRDF and MMSE data as well as recently obtained Mueller matrix imaging polarimetry results.


Recent improvements in nanofabrication strategies have led to the development of patterned surfaces with nanoscale features. Incorporating biomolecules such as proteins, DNA, drugs, and even whole cells could allow for functionalization of nanostructured surfaces for biological applications including biosensing, tissue engineering scaffolds, and drug and gene delivery. Along with the nanofabrication of biological devices, there is a need to develop instrumentation capable of probing and characterizing the dynamic evolution of these bio-functionalized interfaces. Spectroscopic ellipsometry combined with quartz crystal microbalance with dissipation (SE-QCM-D) is a non-destructive optical/mechanical characterization technique that reveals dynamic properties, including average film thickness (with sub-angstrom resolution), adsorbed mass, and porosity.

Nanostructures in the form of sculptured thin films (STF) were fabricated by glancing angle deposition via electron beam evaporation of titanium onto gold-covered quartz sensors. The sensors were then mounted within an SE/QCM-D liquid cell, and proteins of varying sizes were deposited and characterized, in-situ. Protein adsorption was detected shortly after introducing the protein solutions by SE and QCM-D as a change in the optical response and decrease of vibration frequency, respectively. QCMD reported greater adsorbed mass for larger proteins (fibronectin) than smaller proteins (bovine serum albumin). The adsorbed mass of proteins within the nanosized scaffold was larger than that of flat protein deposits, confirming that the STFs are capable of trapping proteins. Analysis of the anisotropic optical response from the nanostructures, which is very sensitive to environmental changes, adds complementary information on protein adsorption; the optical quantification is in agreement with QCM-D results. In the adsorption of other biomolecules, such as cells and DNA complexes, has also been accomplished. The use of combinatorial SE/QCM-D to characterize and monitor the attachment of biomolecules on complex nanotopographies will improve the design and fabrication strategies for a wide array of biotechnological devices.

9:40am EL+AS+EM+MS+PS+TF-ThM6 Characterization of Multilayer Organic Thin Film for Use as An Aptamer Biosensor with Hybrid Spectroscopic Ellipsometry and Quartz Crystal Microbalance with Dissipation. J.Y. Gerassimov, K.B. Rodenhausen, H. Wang, R.Y. Lai, M. Schubert, University of Nebraska - Lincoln

DNA aptamer molecules passivated by alkanethiols can be used for biological detection and screening. Through the use of spectroscopic ellipsometry (SE, optical) and quartz crystal microbalance with dissipation (QCM-D, mechanical) techniques, selective binding of analytes to chemisorbed aptamer probes can be observed in-situ. The system analyzed consists of a gold-coated quartz substrate, a multilayer organic thin film (containing aptamer probe, alkanethiol, and single-stranded DNA analyte), and physiological buffer solution. The attachment and detachment of material, the hybridization efficiency of the aptamer probes, and changes in the porosity of the multilayer organic thin film were all determined by SE/QCM-D.

In this contribution, we present the real-time SE/QCM-D characterization of (a) the formation of the aptamer probe layer, (b) the subsequent chemisorption of all other layers, and (c) the interrogation of single-stranded DNA that is non-complementary or complementary to the sequence found on the aptamer probe. The aptamer DNA sequence encodes codon 12 of the K-ras gene; mutations of this gene are frequently found among pancreatic cancer patients. We found that introduction of either complementary or non-complementary DNA caused increases of the multilayer organic thin film thickness. However, our SE/QCM-D analysis showed that the porosity of the corresponding organic thin film depended differently depending on the compatibility of the DNA analyte. The SE/QCM-D technique provides evidence for different surface attachment mechanisms and can be useful in characterizing biological interfaces.

10:40am EL+AS+EM+MS+PS+TF-ThM9 Contamination Processes of EUV Optics Characterized by Spectroscopic Ellipsometry. L.J. Richter, C. Tarrso, S. Granham, S.B. Hill, T.B. Lucartorio, National Institute of Standards and Technology, N.S. Faradzev, University of Virginia

Extreme ultraviolet (EUV) lithography using 13.5 nm light is emerging as a viable tool for semiconductor fabrication at design rules below 32 nm. Tool performance critically depends on limiting and mitigating degradation of the EUV optical elements. A primary degradation mechanism is the EUV induced deposition of carbon from ambient species originating from outgassing of the unbaked vacuum system and/or outgassing from the EUV irradiation of the resist. Using the high brightness of the Synchrontron Ultraviolet Radiation Facility (SURF-III) at NIST both fundamental studies of EUV induced contamination and practical (“witness plate”) studies of resist out gasgging are performed. Typical deposits are hydrogenated amorphous carbon features with peak thicknesses of about 1 nm and nominal lateral extent of 1 mm. Both ex-situ, small spot mapping spectroscopic ellipsometry (SE) and in situ single-wavelength imaging nulling-ellipsometry are used for rapid, sensitive, contamination metrology. Fundamental studies of contamination by admitted gases indicate that the process is a complex function of both gas pressure and photon dose. Correlations between SE and XPS are suggestive that, at high EUV fluxes, densification can occur resulting in spatial variation in the deposit dielectric function. The use of principal component analysis of the SE images to highlight the spatial diversity will be discussed.

11:00am EL+AS+EM+MS+PS+TF-ThM10 Characterization of Organic Solar Cells Materials and Structures by Spectroscopic Ellipsometry. J.P. Piel, L. Kitzing, A. Bondaz, C. Defranoux, SEMILAB-SOPRALAB, France

Spectroscopic Ellipsometry (S.E) is a well known optical technique widely used for the characterization of all types of thin films for determination of film thickness and optical indices on glass or plastic substrates. S.E. is also being applied to the characterization of materials and multilayer structures of organic materials like organic light-emitting diodes (OLEDs) or Organic Solar Cells.
We present the determination of the refractive indices of organic Solar Cells materials like P3HT, PCBM, Pentacene, Perylene, and their blends. Complex organic materials can be analyzed accurately and fully characterized from their absorption bands in the visible and UV range (190nm to 900nm).

Transmission and absorption can be also measured at the same time and be used to determine the optical properties of these materials. Using these refractive indices, analysis of real multi-layer stacks can be done. The refractive indices can be used afterwards to automatically optimise and balance the energy flow dissipation inside an encapsulated media. We will demonstrate how we can measure single layer properties and multi-layer stacks, through encapsulated samples, from the back side of the substrate. This technique can be applied to test structure or real Organic Solar Cell monitoring. Backside measurement combined with a water vapoour cell used for ellipsometry porosimetry is used to test the efficiency of the thin film encapsulation.

We also present the characterization of ITO and ZnO transparent electrodes by S.E. and how near infra-red ellipsometry is used to infer the electric behaviour on encapsulated samples.

Electronic Materials and Processing Division Room: 210 - Session EM+TF-ThM

Hybrid Electronic Materials and Interfaces

Moderator: A.J. Muscat, University of Arizona

8:00am EM+TF-ThM1 Chemical Modification of Surfaces for Biological Applications. W.G. McGimpsey, Kent State University

INVITED

The chemical and physical characteristics of surfaces can have profound and useful effects on the behavior of biological systems. We describe here several methods for modifying the surfaces of a variety of biocompatible materials, including metals, glass and polymers, in order to produce desired chemical and physical properties. The effects of surface chemistry and surface morphology on the growth and proliferation of mammalian cells as well as bacteria are also reported. Application investigated include control of the growth of microbial biofilms, the growth of three-dimensional tissue assemblies, the patterned growth of neurons and other cell types and the functionality of primary neurons deposited on surfaces.

8:40am EM+TF-ThM3 Modification of Oxide-Free Silicon Surfaces with Phosphonic Acid Self-assembled Monolayers. P. Thissen, T. Pékőto, A. Vega, Y.L. Chabal, University of Texas at Dallas

Phosphonic acid self-assembled monolayers (SAMs) are being considered as versatile surface modification agents due to their ability to attach to surfaces in different configurations, including mono-, bi- or even tri-dentate arrangements. Different chemical strategies for grafting SAMs on oxide-free silicon have been developed. Recently, a novel method for preparing OH-terminated, on otherwise oxide-free silicon has been reported [1] and further expanded for this work. This atomically flat surface contains precisely 1/3 OH and 2/3 H termination groups.

Using this model surface, we demonstrate that the phosphonic group of organic molecules can be chemically grafted to the OH group on the surface using a single chemical step, leaving the Si-H termination unaffected, without oxidation of the Si surface. We also show that the nature of solvents is important as it can act as a catalyst. The perfection of the surface (that remains atomically flat throughout the modification) makes it possible to use first principles DFT-based calculations to model the IR and XPS data obtained for this surface. Thus, a detailed structure for the SAMs can be derived on an atomic level. It is found that phosphonic acids are chemically attached to the Si(111) surface as monodentate via Si-O-P bond upon reaction with the OH groups. The remaining groups of P=O and P-OH are further oriented by using the Drude behaviour on encapsulated samples.


9:00am EM+TF-ThM4 Competing Effects of Interfacial Organic Layers on the Nucleation of Inorganic Thin Films Deposited Via Atomic Layer Deposition. K.J. Hughes, J.R. Engstrom, Cornell University

Over the past several years a number of groups have been investigating the use of interfacial organic layers (IOLs) in the form of self-assembled monolayers (SAMs) and/or surface grown or bound oligomers to promote the subsequent growth of inorganic thin films. Work in our group has initially focused on determining the relative importance of the type, density and orientation of the organic functional groups present in IOLs on the subsequent growth of the thin film, particularly via ALD [1]. Here we shall report on two perhaps less well-studied aspects concerning the use of SAMs or IOLs to promote nucleation and growth via ALD: (i) the effects of the underlying substrate on the IOL, and its ability to promote growth; and (ii) given the same IOL/substrate combination, the effects on ALD for a series of inorganic thin film deposition processes.

To examine these systems we have examined a wide variety of experimental tools, including high resolution low energy viscous flow ALD tool, and also an ultrahigh vacuum (UHV) molecular beam based ALD process, where we employ in situ x-ray photoelectron spectroscopy. Concerning the first of these, we have examined the effect of a thin (< 10 Å) organic layer, poly(ethylene-imine) (PEI) on the ALD growth of TaN, where the underlying substrate is a chemically oxidized SiO2, or a porous SiO2, based low-dimensional thin film. Here we observe essentially the same result from conventional viscous flow, and molecular beam UHV ALD: PEI acts to attenuate TaN, ALD on SiO2, while it enhances growth on a low-k dielectric thin film. From a practical point of view, most importantly, we find that PEI stops infiltration/penetration of the TaN ALD thin film into the porous low-k dielectric. Concerning the second major issue, we have examined the effect of PEI on the subsequent growth via ALD of a series of inorganic thin films, namely: Al2O3, HfO2, Ta2O5, and TaNx. Here PEI has the effect of attenuating growth to different degrees depending on the subsequently grown ALD thin film, leading to no attenuation of growth for Al2O3, but significant incubation periods, in increasing order, for Ta2O5, HfO2 and TaN. The length of the incubation time is found to correlate with both the total enthalpy change of the overall ALD reaction, as well as the net internal energy change of a single ligand exchange reaction representative of the second half of the ALD reaction, suggesting that these factors associated with the ALD process play a key role in determining the length of the incubation period caused by PEI.

form the needed structures. Further the micron-scale UV photopatterning of SAMs does not require a clean room or expensive lithography equipment.

9:40am  EM+TF-ThM6  Sensing Mechanism for Peroxide and Hydroperoxide Vapors in Phthalocyanine Thin Film Transistors. J. Royer, E. Kappe, W. Trogler, A.C. Kummel, University of California San Diego

Organic thin-film transistors (OTFTs) are promising candidates for selective chemical sensors due to numerous chemical and electrical parameters which govern sensor response. Analyte selectivity can be obtained using multiparameter electrical monitoring of a single OTFT which is sensitive to changes in mobility, $L_i/L_o$ ratio, and/or threshold voltage. The present study demonstrates selective hydrogen peroxide and organic peroxide sensors based on irreversible metal-phthalocyanine (MPC) OTFT threshold voltage shifts. The irreversible threshold voltage shift is not evident with non-oxidizing analytes such as di-methyl methylphosphonate (DMMP) and common background analytes such as water vapor. Furthermore, the threshold voltage shift responds linearly to the dose time which permits dosimetric sensing analysis. A proposed mechanism for peroxide sensing is determined using simultaneous monitoring of mobility and threshold voltage. The data reveal reversible mobility and irreversible threshold voltage response. Mobility response time is fast and saturates quickly whereas threshold voltage response is dosimetric, and irreversible, suggesting an accumulation of uncompensated positive charge in the MPC film. The results are consistent with a dual response adsorption/decomposition mechanism in which the peroxide reversibly decreases mobility through a molecular chemisorption event and irreversibly shifts threshold voltage due to electron transfer from the MPC to peroxide. The electron transfer from the MPC to peroxide dissociates the peroxide ion for logic operation on a glass substrate is very impressive. As a more revolutionary approach to realize advanced thin-film devices using both oxide and organic layers, organic/inorganic hybrid layer techniques were also employed, so that hybrid p-n diodes, image sensors, nonvolatile memory, and complementary inverters have recently been demonstrated. These hybrid approaches are an interesting and attractive way to extract a unique device performance which may not be possible with organics or with inorganics alone, compensating some weakness of organics with inorganics or vice versa. The most representative example among the organic/inorganic hybrid devices may be complementary thin-film transistor (CTFT) inverters with an organic p-channel thin-film transistor (TFT) and an oxide n-channel TFT. In particular, a vertically stacked CTFT (VS-CTFT) inverter for logic operation on a glass substrate is very impressive, since the vertical stacking of organic p-TFT on oxide n-TFT improves device integration or device area reduction. For the VS-CTFT inverter for logic and photo-gating, we used a thermally-evaporated p-channel pentacene layer, sputter-deposited n-channel GaZnSn:based oxide (GZTO), and atomic layer deposited (ALD) Al$_2$O$_3$ dielectric. Our VS-CTFT inverter nicely displays effective photo- and electrical-gating with a high voltage gain, dynamically operating in the low-voltage regime of 3.5, and 8 V (the thin pentacene channel receives blue photons). As a more advanced hybrid approach, we have successfully fabricated transparent image pixels that operate at 3 V as composed of pentacene thin-film transistor (TFT) and semitransparent pentacene/ZnO photodiode with a transparent top electrode in the interest of a light detectable smart functional windows. Our transparent pixels was equipped with an additional 6,13-pentacenequinone phosphor layer as deposited on the pentacene/ZnO photodiode. The organic phosphor transmits most of visible photons but absorbs ultra-violet (UV) photons to convert them to yellow-green photons, so that the emitted yellow-green lights excite the pentacene/ZnO diode under a reverse bias state. Our approach to the transparent pixel adopting such an organic phosphor layer certainly makes the pixel operation efficient under not only visible photons but also UV, protecting the organic pentacene from direct UV.

10:40am  EM+TF-ThM9  Organic/Oxide Hybrid Thin-Film Applications for Photo-detector Cells and Complementary Inverters. S.I. Im, Yonsei University, Republic of Korea

Due to the distinct advantages and functionalities, oxide and organic devices on glass or flexible substrates have extensively been studied in such basic forms as thin-film transistors (TFTs) and light emitting diodes. As a more revolutionary approach to realize advanced thin-film devices using both oxide and organic layers, organic/inorganic hybrid layer techniques were also employed, so that hybrid p-n diodes, image sensors, nonvolatile memory, and complementary inverters have recently been demonstrated. These hybrid approaches are an interesting and attractive way to extract a unique device performance which may not be possible with organics or with inorganics alone, compensating some weakness of organics with inorganics or vice versa. The most representative example among the organic/inorganic hybrid devices may be complementary thin-film transistor (CTFT) inverters with an organic p-channel thin-film transistor (TFT) and an oxide n-channel TFT. In particular, a vertically stacked CTFT (VS-CTFT) inverter for logic operation on a glass substrate is very impressive, since the vertical stacking of organic p-TFT on oxide n-TFT improves device integration or device area reduction. For the VS-CTFT inverter for logic and photo-gating, we used a thermally-evaporated p-channel pentacene layer, sputter-deposited n-channel GaZnSn:based oxide (GZTO), and atomic layer deposited (ALD) Al$_2$O$_3$ dielectric. Our VS-CTFT inverter nicely displays effective photo- and electrical-gating with a high voltage gain, dynamically operating in the low-voltage regime of 3.5, and 8 V (the thin pentacene channel receives blue photons). As a more advanced hybrid approach, we have successfully fabricated transparent image pixels that operate at 3 V as composed of pentacene thin-film transistor (TFT) and semitransparent pentacene/ZnO photodiode with a transparent top electrode in the interest of a light detectable smart functional windows. Our transparent pixels was equipped with an additional 6,13-pentacenequinone phosphor layer as deposited on the pentacene/ZnO photodiode. The organic phosphor transmits most of visible photons but absorbs ultra-violet (UV) photons to convert them to yellow-green photons, so that the emitted yellow-green lights excite the pentacene/ZnO diode under a reverse bias state. Our approach to the transparent pixel adopting such an organic phosphor layer certainly makes the pixel operation efficient under not only visible photons but also UV, protecting the organic pentacene from direct UV.

11:40am  EM+TF-ThM12  Electroless Deposition of Metals on SiO$_2$ Surfaces Modified by a Self-Assembled Monolayer. R. Jain, A. Ng, A.J. Muscar, A. Villan, R. Har-Lavan, O. Taffe, P. Joshi, R. Kazaz, D. Cohen, Weizmann Institute of Science, Rehovot Israel

The Schottky-Mott model predicts that the eventual energy barrier formed when metal and semiconductor are brought into intimate contact, will be equal to the difference between the metal work function and the semiconductor's electron affinity (for n-type). 60 years of extensive experiments have clearly shown that this ideal picture, that seems to hold for wide bandgap semiconductor such as ZnO, GaS etc., is far from being adequate for the more covalent narrow bandgap Si, Ge, InP, and GaAs. Following Bardeen, who attributed this deviation from the model to surface states energetically located within the semiconductor's forbidden gap, different explanation were raised as for the source of those surface states.

One commonly used model for surface states formation at the interface of semiconductor and metal is the intrinsic Metal Induced Gap State (MIGS) model, stating that gap states are inevitably formed due to the decay of metal electronic states' wavefunction into the semiconductor's bandgap. We have examined the presence of such MIGS using Hg, which is one of the few metals that doesn't interact chemically with Si, as an electrode. Furthermore, in order to eliminate surface states which are just due to Si dangling bonds, we have used state of the art mixed molecular monolayers of hydroquinone and alcohols that were shown to be the best chemical passivation for Si surfaces.

Using alcohols of different alkyl chain lengths we succeeded to effectively change the electron affinity of the Si over 400 mV range while maintaining firm surface passivation. Current-voltage measurements of diodes formed that way, with Hg on organically modified Si surface, demonstrated near ideal Schottky-Mott characteristics with index of interface behavior $S=0.9$ (compared to a common value for Si $S=0.1$).
Of all metal fluorides, iron-based compounds are the most promising to conversion mechanism using surface science tools. We have produced model oxyfluoride systems in order to characterize the known about the electronic structure of metal oxyfluorides, particularly the energy gap and thus a higher electronic conductivity. Currently, little is use of metal oxyfluorides, which are characterized by a slightly smaller efficient charge and discharge. One path to overcome this limitation is the Transition metal fluorides have recently gained interest as possible electrode materials in lithium ion conversion batteries. Owing to their large band gaps, they operate at high voltages and enable high energy densities.

It has been found that oxygen insertion into the FeF2 matrix can be controlled until complete oxidation occurs. As expected for a Mott-Hubbard insulator, the valence band of FeF2 can be interpreted using a simple crystal field approach. In the case of FeO2, strong charge transfer effects need to be taken into account in order to interpret the band edges. To explore the conversion process, Li has been evaporated onto these iron oxfluorides in-situ. Preliminary results addressing the reactivity of lithium at the surface of these materials will also be presented.

Modified graphene is one of a promising candidate with its high theoretical surface area (2630 m$^2$/g). EDLC working principle also depends on how effectively it can polarize the electrolyte solution. In principle, ionic liquids are promising electrolyte systems with their high non-volatility, non-flammability, thermal stability and good solvating ability. This study focuses on the intercalation studies of ionic liquids such as N-methyl-N,N,N-tris(2-hydroxyethyl)ammonium iodide, N-methyl-N,N,N-tris(2-hydroxyethyl)ammonium methylsulfate, 1-butyl-3-methylimidazoliummethylsulfate and N-octyl-N-methylpiperidinium methylsulfate. To achieve a high surface area material with modified graphene, it is essential to understand the interfacial interactions upon intercalation of ionic liquids.

We study powder X-Ray Diffraction Analysis to characterize the interlayer distance of modified graphene. For instance, its d-spacing increases by about 4-12 Å after intercalating as-synthesized graphite oxide (GO, ~9Å) with N-methyl-N,N,N-tris(2-hydroxyethyl)ammonium methylsulfate (IL) at room temperature. Annealing at 300-500°C, (002) orientation of GO disappears as a result of thermal exfoliation. In addition, in-situ Infrared Absorption Spectroscopy (IRAS) measurements were performed to explore the intercalation chemistry of modified graphene in the presence of ionic liquids. Therefore, we perform thermal reduction studies in vacuum coupled with IRAS measurements to characterize the chemical interactions during thermal exfoliation of reduced GO. After annealing GO intercalated with this IL, the loss of C-N and C-O containing species is identified at ~1000-1500 cm$^{-1}$ and 800-1200 cm$^{-1}$ with contribution from C-OH groups at 3000-3700 cm$^{-1}$. Complete removal of these species is observed after a ~500°C anneal resulting in a weak infrared absorbance intensity of sp$^2$-hybridized C=C species at ~1580 cm$^{-1}$. The presence of new formation of sheet-to-sheet linking or bonding motifs was also studied with X-ray Photoelectron Spectroscopy (XPS).

*Research supported by the U.S. Department of Energy, Office of Basic Energy Sciences, Division of Materials Sciences and Engineering under Award DE-SC001951.*

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**EN+NS-ThM2** 8:20am  Electronic Structure and Chemical Composition of Candidate Conversion Material Iron Oxyfluoride, **R. Thorpe**


Transition metal fluorides have recently gained interest as possible electrode materials in lithium ion conversion batteries. Owing to their large band gaps, they operate at high voltages and enable high energy densities. However, this large band gap inhibits charge conduction and thus impedes efficient charge and discharge. One path to overcome this limitation is the use of metal oxyfluorides, which are characterized by a slightly smaller energy gap and thus a higher electronic conductivity. Currently, little is known about the electronic structure of metal oxyfluorides, particularly the relation between chemical structure, composition, and energy gap. Hence, we have produced model oxyfluoride systems in order to characterize the conversion mechanism using surface science tools.

Of all metal fluorides, iron-based compounds are the most promising to maximize energy density. Ultra-thin FeF$_2$ films have been synthesized via the fluorination of clean Fe foil exposed to XeF$_2$, following a self-limited Mott-Cabrera mechanism. The FeF$_2$ films have then been sequentially exposed to a partial pressure of O$_2$ of 2x10$^{-6}$ Torr at 285°C in order to produce oxyfluoride. Using x-ray and ultraviolet photoemission as well as inverse photoemission, we have probed the electronic structure of these FeO$_2$F$_x$ samples and characterized the occupied and unoccupied states near the band gap of the material.

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**EN+NS-ThM10** 11:00am  Exploration of the Effects of Si Nanowire Length and Doping on Li-Ion Battery Anode Performance, **F. Rudi**

F. Rudi, V. Chakrapani, M.A. Filler, P.A. Kohl, Georgia Institute of Technology

Silicon nanowires have recently garnered significant attention as a potential candidate to replace graphite as the negative electrode in a lithium-ion battery. Silicon’s earth abundance, extensive knowledge base, and its theoretical capacity of 4200 mAh/g make it an attractive material for this purpose. While the high incorporation of lithium presents a problem as silicon undergoes a significant volume change upon intercalation (up to 400%), nanowires permit facile radial strain relaxation and allow lithium without pulverization. In this work, we report on the first systematic study of nanowire length and doping on the cycling performance of NW electrodes. Silicon nanowires were grown on stainless steel substrates via the vapor-liquid-solid technique in a cold-wall low pressure chemical vapor deposition reactor.
pressure chemical vapor deposition reactor. Growth times ranging from 5-
60 minutes and doping concentrations between 10^5 – 10^7 atoms cm^-2 were
studied. The half cells were cycled against lithium metal between (1) 0.01
and 2.0 V and (2) 0.07-0.7 V at a C/20 rate for 20 cycles. Cycling
performance at different depths of discharge was also studied. Nanowires
grown at short and long times both exhibit lower capacities than those
grown at intermediate times. We attribute this effect to the loss of nanowire
contact at the interface between the stainless steel and nanowire array for
long nanowires, while the percolation network formed by short nanowires is
not sufficiently robust to prevent loss of electrical connectivity upon wire
breakage further from this interface. Nanowires at higher doping
concentrations were not found to improve cycling performance drastically,
which we attribute to the dopant effects on the growth mechanism of the
nanowires and placement of dopant atoms in the silicon matrix that may have
inhibited lithium atom insertion. We will present novel electrode fabrication
routes that overcome these two challenges.

1. J.Y. Huang et al., Science 330, 1515-1520 (2010); Nano Lett. (revised);
ACS Nano (in press).

Graphene and Related Materials Focus Topic
Room: 208 - Session GR+NS+PS+SS-ThM
Graphene: Surface Chemistry, Functionalization,
Plasma Processing and Sensor Applications
Moderator: G.G. Jernigan, U.S. Naval Research Laboratory
8:00am GR+NS+PS+SS-ThM1 Tailoring Graphene’s Properties
through Chemistry, J.T. Robinson, Naval Research Laboratory INVITED
Graphene’s unique electronic transport properties have motivated intensive
research and development to mold it into the electronic material of the
future. However, graphene can be much more than an electronic switch. Its
high structural integrity and chemical flexibility enable extensive control of
its optical, mechanical, and electronic properties. The most scalable and
inexpensive route to modify these properties is chemical functionalization.
Consequently, chemically modified graphenes (CMGs) have emerged as a
system of materials whose many attractive properties complement and
extend those of unmodified graphene.

In this talk I will describe efforts at NRL to synthesize and characterize new
CMGs as well as first steps towards applications such as sensors and
nanomechanical resonators. To begin, I will discuss the interaction of small
molecules (CCL4, CS2, H2O and acetone) with single-layer graphene under
steady-state conditions using infrared multiple-internal-reflection.
Adsorption-induced changes in the IR spectra suggest the formation of in-
plane strain, where we observe important differences arising between
species that form liquid-like layers under steady-state conditions and those
that do not. Second, I will not synthesize CMGs, e.g. a well-known derivative
of graphene that has a rich ensemble of oxygen-based functional groups and
related defects. These defects are readily tunable through chemical or
thermal treatments and facilitate the formation of vapor and bio-sensors
with parts-per-billion and nanomolar sensitivities, respectively. Third, I will
discuss the stoichiometric addition of fluorine atoms to graphene and
describe their resulting properties. Experimentally, chemically modified
graphene derivatives become highly resistive and optically transparent,
while DFT calculations show band gaps open depending on the fluorine
coverage and ordering. Finally, through combining these two material
systems, I will discuss the fabrication and performance of CMG-based
nanomechanical resonators. Through chemical modification, the frequency
of CMG-based resonators is tunable over 500% and their quality factors can
exceed 20,000 at room temperature.

9:00am GR+NS+PS+SS-ThM4 Novel Strategies for the Chemical Functionalization of Graphene: Towards Graphene/Molecular
Nanosheet Heterostructures, A. Turchanin, C.T. Nottbohm, Z. Zheng, M.
Schnietz, A. Beyer, University of Bielefeld, Germany, M. Heilmann, M.
Sauer, Julius-Maximilians-University Würzburg, Germany, A. Göltzlhüser,
University of Bielefeld, Germany
Chemical functionalization of graphene is essential for implementations of the
2D carbon sheets in various functional devices (e.g. chemical and
biochemical sensors, nanoelectromechanical components, etc.) and for
tuning their electrical properties. However, the functionalization is difficult
to achieve due to the challenge of introducing sharp, non-wetting,
high structural quality. On the other hand, ultrathin (~1 nm) molecular
nanosheets made from self-assembled monolayers (SAMs) possess well-
defined chemical groups intrinsically. Moreover, due to the directionality of
the constituting molecules both faces of the free-standing nanosheets -Janus
nanomembranes- can be independently and specifically functionalized.
Simple mechanical stacking of the nanosheets allows fabricating ultrathin
layered structures with tunable physical and chemical properties. Upon
annealing these stacks are converted into graphene sheets with adjusted
thickness. The engineering of graphene/nanosheet heterostructures opens up
novel routes towards chemically functionalized graphene sheets for
functional applications. A potential of this approach will be discussed.
Z. Zheng, C.T. Nottbohm, A. Turchanin, H. Muzik, A. Beyer, M.
Heilmann, M. Sauer, A. Göltzlhüser, "Janus nanomembranes: A generic
8493-8497
C.T. Nottbohm, A. Turchanin, A. Beyer, R. Stosch, A. Göltzlhüser,
"Mechanically stacked 1 nm thick carbon nanosheets: 2D layered materials
with tunable optical, chemical, structural and electrical properties", Small 7
(2011) 874-883
A. Turchanin, D. Weber, M. Bünfeld, K. Kisielski, M. Fustel, K. Efetov,
R. Stosch, T. Weimann, J. Mayer, A. Göltzlhüser, “Conversion of self-
assembled monolayers into nanocrystalline graphene: structure and electric
transport”, ACS Nano 5 (2011) DOI: 10.1021/nn200297n

Thursday Morning, November 3, 2011 194
9:20am  GR+NS+PS+SS-ThM5  Biosensors Based on Chemically Modified Graphene, R. Stine, J.T. Robinson, P.E. Sheehan, C.R. Tamanaha, U.S. Naval Research Laboratory
The sensitive and specific detection of biomolecules without using a label is a long-standing goal of the biosensors community. Several promising advances of the past several years formed biological field effect transistors (bioFETs) that have as the gate nanoscale materials such as nanowires and carbon nanotubes. The nanoscale dimensions of these materials allow the small changes associated with biomolecules to significantly change the conduction through the gate. These conduction changes can be correlated with solution concentration to give precise readouts. While bioFETs are a promising way forward, there are many processing difficulties associated with these 1-D materials that inhibit large scale, reproducible fabrication of devices. Here, we will discuss our efforts to develop biosensors based on 2-D chemically modified graphene. These devices impart the sensitivity gains seen with other nanoscale materials, but offer a configuration amenable to processing techniques that are common in the semiconductor industry. We will focus primarily on chemically modifying graphene for attachment of biomolecular probes. Devices utilizing both graphene and graphene oxide will be covered, and surface spectroscopic studies of the material modification will be discussed. Successful results for the detection of specific DNA hybridization will also be presented, with detection limits that compare favorably with the best results reported from nanowire bioFETs.

Acknowledgements: R.S. is an employee of Nova Research Inc., Alexandria, VA, USA. This project received support from the Defense Threat Reduction Agency-Joint Science and Technology Office for Chemical and Biological Defense.

9:40am  GR+NS+PS+SS-ThM6  Controllable Defect Healing and N-doping of Graphene by CO and NO Molecules, B. Wang, Vanderbilt University, S.T. Pantelides, Vanderbilt University and ORNL
Point defects alter strongly the physical and chemical properties of graphene, e.g. they degrade electrical transport and enhance chemical reactivity. Defects could also be used to achieve graphene functionalization, e.g. N atoms, as n-type dopant, can be introduced to obtain n-type graphene. Thus, controllable defect healing and N-doping in graphene would be very valuable for potential device applications. Here we report first-principles calculations of dynamic reactions that suggest a procedure for defect healing and N-doping with fast dynamics and low thermal budget. Vacancies in graphene can be healed by sequential exposure to CO and NO molecules. A CO molecule gets adsorbed at a vacancy site and a NO molecule subsequently removes the extra O by forming NO₂, which desorbs quickly. Another NO molecule gets adsorbed at the vacancy site removing the extra oxygen atoms. N-doping results in a defect-free graphene sheet. Controllable N-doping can be achieved by sequential vacancy creation (e.g. by electron or ion beam) and subsequent exposure to NO molecules at room temperature. NO molecules are trapped at vacancies and other NO molecules remove the extra O atoms simultaneously, leaving N atoms incorporated in graphene. Both reactions (healing and doping) are exothermic. We suggest that a combination of CO and NO molecules can potentially provide simultaneous healing and doping. Adjusting the ratio could fine-tune the N-doping level. The proposed strategy introduces no extra defects and is promising for graphene-based electronic materials in radiation environments. Finally, we propose that NH₂, which is normally used in experiments to introduce N atoms, may be not a good choice for N-doping since the dissociated H atoms can be trapped at vacancies and act as impurities that increase the resistivity of graphene.

This work was supported by DTRA Grant No. HDTRA1-10-1-0016 and the Williamson and Nancy F. McMinn Endowment at Vanderbilt University. The calculations were performed at ORNL’s Center for Computational Sciences.

10:04am  GR+NS+PS+SS-ThM9  Aptamer Modified Graphene Bio Sensor, K. Machushi, Y. Ohno, K. Matsumoto, Osaka University, Japan
Since graphene has high mobility and a large surface area, it is suitable for the application of the high sensitive sensor. In the present paper, we have first succeeded in the selective detection of the bio molecule such as IgE using the aptamer modified graphene FET.

The graphene was formed using the conventional mechanical exfoliation method on the SiO₂/Si substrate. The source and drain electrode were formed by the electron beam lithography and Ti/Au evaporation. The silicon rubber pool was formed on the fabricated graphene FET, and phosphoric buffer solution was poured into the silicon rubber pool. The Ag/AgCl reference electrode was introduced into the phosphoric buffer solution, which works as a top gate electrode for the graphene FET.

As a first step of the biosensor, three bio molecule such as Immunoglobulin E(IgE), Streptavidin(SA), and Bovine serum albumin(BSA) were introduced into the phosphoric buffer solution of pH of 6.8, and the change of the drain current of the graphene FET was detected. In this case, the IgE and SA shows the decrease of the drain current, while the BSA the increase of the drain current. Because, in the phosphoric buffer solution of pH of 6.8, IgE and SA are negatively charged, while BSA negatively charged. Therefore, the hole current of the graphene FET change the drain current following the charge of the bio molecule. As a result, bare graphene FET can detect the bio molecule following the charge of the molecule, but it does not have the selectivity

As a second step, in order to get the selective sensing of the bio molecule, the surface of the graphene was modified by the IgE aptamer, which was connected to graphene using the linker(-l-lysinebutanato acid succinimidyl ester) and the IgE was known to selectively couple to IgE. When the BSA and SA were introduced into the phosphoric buffer solution on the aptamer modified graphene FET, there occurred no change in the drain current, while the IgE was introduced in the solution, the drastic decrease of the drain current was observed. This means the BSA and SA do not couple to IgE aptamer, and only IgE couple to the IgE aptamer on the graphene FET. Therefore, the selective sensing of the IgE was successfully carried out.

We have first succeeded in the selective sensing of IgE using the modified graphene FET.

11:10am  GR+NS+PS+SS-ThM10  A Molecular Dynamics Study of Chemical Modification of Graphene Oxide Sheets, T. Liang, B. Devine, S.R. Phillpot, S.B. Sinnott, University of Florida
Graphene, the single-layered graphite, has attracted tremendous attention owing to its fascinating physical properties. One of the main obstacles in this field is to find an efficient and consistent approach to produce graphene sheets at large quantities. In addition to the mechanical exfoliation method, many chemical approaches have been developed to synthesize graphene on a large scale. The key intermediate product in these chemical approaches is the graphene oxide sheets, which are often heavily oxygenated with hydroxyl or epoxide functional groups on the surface and carboxyl or carbonyl groups at the edge. However the energetic and kinetics associated with graphene oxide sheets had not been elucidated in detail due to the inherent chemical complexity of the system. Here, a new dynamic charge empirical potential is presented that is used in classical molecular dynamics simulations to elucidate the dynamics of graphene oxidation and the resulting influence on their mechanical and structural properties. In addition, the oxygenating and hydrogenating processes of defective graphene sheets at room temperature in addition to elevated temperatures are presented. The findings are compared to the results of first principles density functional theory findings and to experimental data.

11:20am  GR+NS+PS+SS-ThM11  Enhancing and Controlling the Chemical Reactivity of Epitaxial Graphene via Growth Induced Strain, J.E. Johns, Northwestern University, Md.Z. Hossain, Gunma University, Japan, M.C. Hersam, Northwestern University
The high electrical and thermal conductivity of graphene, as well as its two dimensional nature, has led to its rapid integration into a wide range of practical applications including high frequency analog transistors and transparent conductors. However, many other potential applications, such as excitonic switches, pseudospin devices, or digital logic circuits, require covalent chemical modification of graphene. Due to the chemical inertness of its pi bonded network, previous methods for covalently modifying graphene have required extreme, irreversible conditions including acidic treatments, high energy radical polymerization, and ion beam implantation. Here we present an alternative method for increasing the chemical reactivity of graphene by systematically altering the compressive strain of epitaxial graphene (EG) on SiC(0001). Depending on its annealing history, EG has been shown to have a compressive strain of 0% to 1% due to a mismatch of thermal expansion coefficients with the underlying buffer layer and silicon carbide substrate. Using differing thermal treatments, we show that the amount of strain in epitaxial graphene can be tailored, as verified by characteristic peak shifts of the 2D Raman band. The resulting chemical reactivity of the strained EG is studied at the atomic-scale using ultra-high vacuum scanning tunneling microscopy following reversible gas phase reactions of EG with oxygen and fluorine. These results suggest a new method for controlling the electronic properties of graphene, and provide fundamental insight into the nature of chemical bonding on EG.

Graphene, a sp²-structured monolayer of carbon atoms, has attracted much interest for its fundamental science and its potential in many device applications. By tailoring its surface chemistry, material properties can be regulated and thus broaden the number of potential applications. In this work, we demonstrate that by chemically functionalizing graphene the
electrical properties and its interaction with adsorbates may be controlled. Electron beam generated plasmas produced in ammonia-containing gas, is used to controllably introduce nitrogen and primary amines. A study of the chemical, electrical and structural properties of the chemically-modified graphene at different functional group concentrations is discussed. In addition, the use of amine-functionalized graphene as a bio-sensing platform for DNA detection using a field-effect-transistor-based sensor is demonstrated. This work is supported by the Office of the Naval Research.

**Magnetic Interfaces and Nanostructures Division**

**Emerging Magnetic Characterization and Results**

**Moderator:** A.N. Caruso, University of Missouri-Kansas City

8:00am **MI-ThM1** The X-Ray View of Ultrafast Nano Magnetism. *H.A. Durr, SLAC National Accelerator Laboratory* 

**Invited**

Polarized soft x-rays have been used over the past 20 years to obtain fascinating new insights into nanoscale magnetism. The separation of spin and orbital magnetic moments, for instance, enabled detailed insights into the interplay of exchange and spin-orbit interactions at the atomic level. X-ray and photoelectron imaging techniques have revolutionized our understanding of magnetism of the ULTRA SMALL. In addition the now available polarized soft x-ray pulses with only few ps duration allow us to observe the magnetic interactions at work in real time, i.e. they open the door to study ULTRA FAST magnetism. The ultimate goal of such studies is to understand how spins may be manipulated by ultrashort magnetic field, spin polarized current or light pulses. In this talk I will give an overview of achievements and the current status of probing magnetism of the ultra small and ultra fast using x-rays from synchrotrons [1-3] and more recently from x-ray free electron lasers.


8:40am **MI-ThM3** Spectroscopy of Magnetic Thin Films. *S.N. Gilbert, N.H. Tolk, Vanderbilt University*

Recent studies of magnetic thin films and spintronic devices will be presented. Time-resolved Kerr Effect measurements of ferromagnetic/antiferromagnetic interfaces as a function of film layer thickness and antiferromagnetic spin orientation will be discussed. Magnetic and time-resolved spin characterization of novel spintronic devices and materials will also be shown.

9:00am **MI-ThM4** Detection and Control of Electronic Phase Competition in Complex Oxides. *T.Z. Ward, Oak Ridge National Laboratory* 

**Invited**

Electronic phase separation is present in many complex material systems and has been linked to colossal magnetoresistance, high Tc superconductivity, and multiferroicity. Here, nanometer to micron sized regions of vastly different electronic and magnetic properties can coexist and evolve with changing external conditions. In this talk I will discuss recent work on fabricating single crystal wires of electronically phase separated manganites to a size comparable to the domains of the electronic phases residing in the material; thereby allowing finite emergent regions to dominate device characteristics. This has given us a means to probe, observe and exploit properties which are *hidden* in unconfined systems. Transport measurements on simple confined structures reveal new properties such as ultrafast jumps in resistivity, a reemergent metal-insulator transition, and discrete resistive hopping that are unseen in larger samples. We have found that these properties are also tunable through doping, strain, electric field and magnitude of confinement. This ability to control key elements of the underlying complex electronic correlations and observe the resulting changes in a material’s behavior help answer questions about the fundamental physics that rule emergent phenomena in complex materials while opening the door to new device functionality.


Antiferromagnets play a critical role in spintronic applications such as pinning layers in magnetic memories. The development of spin-polarized scanning tunneling microscopy and spectroscopy (SP-STM/STS) has shown its unprecedented power in resolving the local spin and domain structures of antiferromagnetic surfaces down to atomic level. While most efforts have been made on imaging metal surfaces at cryogenic temperatures, only a few have been devoted to the study of room-temperature magnetic systems. Here we apply SP-STM/STS to study the local spin and magnetic properties of a technologically driven material system which exhibits layer-wise antiferromagnetism with a very high Néel temperature (T_N ~ 900 K). Mn$_2$N$_2$(001) thin films have been grown on MgO(001) substrates using ultra high vacuum plasma-assisted molecular beam epitaxy and transferred in situ to a home-built room-temperature SP-STM for magnetic imaging. Results have shown that the surface exhibits a *topological spin pyramid* structure with alternating single Mn- and double MnN- layers, while opening the door to new device functionality.


Manganese doped, magnetic germanium quantum dots are predicted to be important building blocks for the future of spintronic devices. The combination of quantum confinement and carrier mediated ferromagnetism makes these structures particularly interesting. The goal of this work is to understand and control the Mn environment within the Si(100), Ge wetting layer and Ge quantum dot (QD) systems and understand how it influences the magnetic properties. Samples were investigated primarily using scanning tunneling microscopy followed by magnetic analysis using a vibrating sample magnetometer and one sample with x-ray magnetic circular dichroism. An important materials question is the competition to form secondary phases in the Mn-doped Ge(105)films that will be discussed. (1) The investigation of the stability and evolution of Mn nanostructures on a Si(100)-(2x1) reconstructed surface as a function of annealing temperature up to temperatures typical for Ge QD growth. At an annealing temperature of approximately 316°C, Mn adatoms move into Si sub-surface sites and we observe an electronic effect consistent with acceptor dopants. (2) The use of a surface driven approach where Mn is deposited on the Ge QD surface and forms well-defined islands on the QD and wetting layer surface. We observed the behavior of the Mn islands during STM measurement with increasing annealing temperatures and how the islands evolved via ripening and migration across the surfaces. In addition the structure and bonding of the MnGe layers specifically in Ge(105) films will be discussed. (3) The co-deposition of Ge and Mn throughout the Ge QD growth process. For route (3) the highest Mn concentration is 23% which results in only minor perturbations in the Ge QD growth (fewer and smaller Ge QDs), albeit

*Falicov Student Award Finalist*
secondary phases form on the surface. Lower concentration samples (5% and 8% Mn) yielded high quality quantum dots and no observable secondary phases on the surface. We presume that when secondary phases form, the majority of the Mn deposited is consumed to form the secondary phases. The competition to form secondary phases is investigated further utilizing scanning auger microscopy to map Mn and low energy electron microscopy to study the growth sequence as a function of Mn concentration. Magnetism results from one particular sample (Mn0.05Ge0.95 QD) indicate a ferromagnetic material with a Curie temperature above room temperature. We’d like to acknowledge our funding support from NSF CHE-0828318 and DMR-0907234.

11:00am MI-ThM10 Novel Iron-Induced Structures on Gallium Nitride (0001) and (000-1) Studied Using Scanning Tunneling Microscopy and First Principles Theory, W. Lin, Ohio University Nanoscale and Quantum Phenomena Institute, H.A.H. Al-Britten, Ohio University Nanoscale and Quantum Phenomena Institute and KAIST, King Saud Univ., Saudi Arabia, K.K. Wang, A.V. Chinchore, M. Shi, Y. Liu, N. Takeuchi, A.R. Smith, Ohio University Nanoscale and Quantum Phenomena Institute

There is much interest in the field of spintronics in which magnetic phenomena are combined with electronic properties to form a new class of materials with added device functionality. An essential area is that of magnetic materials for data storage. First-principles theoretical calculations have been carried out using a custom-designed, home-built molecular beam epitaxy/scanning tunneling microscopy (MBE/STM) facility. Growth of iron on GaN is carried out using an Fe diffusion cell and at a substrate temperature which is carefully selected in order to produce the highest quality atomically-smooth Fe-induced structures. It is found that the Fe-induced structures on Ga-polar GaN(0001) strongly depends on the presence of the pseudo-1×1 surface structure as a starting surface, and that under the correct conditions a clear Fe-induced structure grows outward from the GaN step edges, as revealed in scanning tunneling microscopy images. First-principles theoretical calculations have been carried out which suggest a low-energy model for the 6/6 structure consisting of Fe atoms embedded within the pseudo-1×1 layer and with Ga adatoms at the top. The results for N-polar GaN(0001) are quite different. In this case, deposition of Fe onto a Ga-rich surface results in the formation of uniform Fe-induced islands having a 4/2 zigzag row structure. The zigzag rows orient along the high symmetry [1120] directions of the surface.

Efforts are also underway to investigate the chemical stoichiometry, and electronic and magnetic properties of these Fe-induced structures and to explore the evolution of these monolayer films as additional Fe and/or Ga is added to the surface. This work has been supported by the U.S. Department of Energy, Office of Basic Energy Sciences (Grant No. DE-FG02-06ER46317). Additional support from the National Science Foundation (Grant No. 0730257) is also acknowledged.

Y.L. is now at Los Alamos National Laboratory, Los Alamos, NM.

N.T. was a visiting Presidential Scholar from the Universidad Nacional Autonoma de Mexico during 2010-11.


11:20am MI-ThM11 Scanning Tunneling Microscopy and Spectroscopy Performed on Single Mn Monolayer on Wurtzite (000-1) GaN, A.V. Chinchore, K.K. Wang, A.R. Smith, Ohio University, P. Ferrari, A. Barraul, University of Buenos Aires, Argentina

The III-V diluted magnetic semiconductors (DMS) are a new class of materials with promising applications in spintronics.[1] The low solubility of transition metal atoms into III-V semiconductor host has been a key concern in the successful development of DMS. This low solubility however was used to advantage by Lu et al. to develop an ideal magnetic semiconductor bi-layer.[2] Wang et al. recently reported high density 2D Mn-Ga stripe phases on Ga-Polar GaN(0001) surface, with interesting atomic spin arrangement. [3] The N-Polar GaN(0001) 1×1 structure offers an added advantage over the Ga-Polar structure, as the Mn atoms deposited on this surface are in closer proximity to the N atoms favoring the GaMnN bonding.

We have conducted a series of experiments aimed at understanding the behavior, electronic and magnetic properties of Mn atoms on N-Polar GaN(0001) 1×1 surface. The experiments were performed in a custom built MBE-STM system with in-situ sample transfer ability. The growth is monitored with reflection high energy electron diffraction (RHEED). The standard GaN(0001) 1×1 surface was prepared and was exposed to sub monolayer doses of Mn at various temperatures. It was observed that the behavior of Mn atoms on GaN(0001) 1×1 surface is highly sensitive to the substrate temperature. The temperature Mn deposition, T, ~ 100 °C, led to the formation of a metastable 3×3 structure which transformed to a more stable 3×3 R30° structure, when the sample is heated to T, ~ 120 °C, as confirmed by RHEED. It was observed that the 3×3 3 R30° structure is stable up to 750 °C. The temperature dependent behavior of the structures suggests that the Mn atoms are physisorbed while forming the metastable structure and that they are chemisorbed in the case of the stable 3×3 3 R30° structure. The STM measurements performed on the Mn 3×3 structure showing the metastable nature of the structure are presented as well are the STM and STS results showing the characteristics of 3×3 3 R30° surface. RHEED simulations confirming the surface atomic arrangement for the structure are presented. The theoretical calculations are performed using the first principles and the Tersoff-Hamann simulation method. The results indicate that the Mn atoms push the Ga atoms laterally in the surface ad-layer forming bonds directly with the bilayer N atoms. The funding from NSF and DOE for the project is greatly acknowledged.

References.


11:40am MI-ThM12 Designing of Engineered Multiferroic Composites by Radical Enhanced Atomic Layer Deposition, J.H. Choi, T.E. Quckel, S. Tolbert, J.P. Chang, University of California Los Angeles

Multiferroic materials induced polarization under external magnetic field H, or induced magnetization under external electric field E. Magneto-electric (ME) phenomena in multiferroic materials holds considerable promises because of their potential applications in spintronics, such as magnetoelectric sensors, magneto-capacitive devices, and electrically driven magnetic data storage. The ultimate goal for practical device application of multiferroic materials is dependent on how to create strong ME coupling between different types of ferroic order. The strictive interaction between the piezoelectricity of the ferroelectric (FE) phase and the magnetostriiction of the ferromagnetic (FM) phase lead to produce larger ME coefficients than single phase multiferroic materials. Thus, the research has been directed towards designing engineered multiferroic composite materials in the form of horizontal multilayer (2-2), vertical superstructures (3-1) or other nanoparticle composite structures (3-0) in a precise controlled manner. In this work, the BiFeO3 (BFO) and Pb(ZrTi)O3 (PZT) thin film were synthesized by radical enhanced atomic layer deposition (RE-ALD). RE-ALD is a gas-phase technique in which precursor vapors are pulsed alternately into the reaction chamber and the thin film growth proceeds through surface reactions in a self limiting manner. The advantages of ALD include excellent conformality, simple and accurate thickness control and good uniformity on large areas. In order to demonstrate conformal deposition of engineered multiferroic materials in the form of 3-0 or 2-2 configuration, PZT and BFO was deposited onto a mesoporous CoFe2O4 (CFO) substrate by RE-ALD. The mesoporous CFO films were found to be fully filled by ALD PZT and BFO. The composition and crystal structure of the PZT-CFO and BFO-CFO systems were confirmed by X-ray Photon Spectroscopy and X-ray Diffraction (XRD), respectively. More detail crystal structure were investigated by synchrotron XRD and extended x-ray absorption fine structure spectroscopy (EXAFS). The magnetic and ferroelectric properties of the PZT-CFO or BFO-CFO systems were characterized by a superconducting quantum interference device (SQUID) magnetometer and piezoresponse force microscopy (PFM). Magnetic properties such as coercive magnetic field (Hc) and saturated magnetization (Ms) were systematically analyzed on composite systems and the pure CFO substrate. In addition, P-E loops for PZT-CFO and BFO-CFO thin films were measured at room temperature and the saturation polarization (P_s) and coercive field (E_c) were investigated with respect to thickness and crystal plan.
8:20am **NS-Thm2** Chemical Modification and Patterning of Self Assembled Monolayers using Scanning Electron and Ion-Beam Lithography, M.J. Perez Roldan, C. Pascual Garcia, G. Marchesini, D. Gilliland, G. Ciccione, P. Colpo, F.J. Rossi, European Commission, JRC Institute for Health And Consumer Protection, Italy

We present chemical modification of self assembled monolayers (SAMs) using electron and ion-beam lithographies. We used thiolated polyethylene oxide (PEO) SAMs which we used to fabricate chemically contrasted patterns at the nanoscale. Patterned surfaces were characterized by X-ray photoelectron spectroscopy (XPS), time of flight-secondary ion mass spectrometry (ToF-SIMS). Results showed a chemical modification of surfaces patterned by means of electron beam (e-beam) lithography and a removal of PEO SAMs on the areas treated with the ion beam. The chemical modification of PEO SAMs converted the non-fouling surfaces on fouling surfaces.

8:40am **NS-Thm3** Interlocking Pinwheel Chains Formed by Self Assembly of Aromatic Cyanides, M. Luo, W. Lu, E. Chu, D. Kim, Z. Cheng, D. Sun, K. Cohen, Y. Zhu, J. Wyrick, University of California, Riverside, T.L. Einstein, University of Maryland, College Park, L. Bartels, University of California, Riverside

As part of a bottom-up strategy, molecular self assembly can be a promising technique to create surface patterns with ultimately small feature sizes in an economic efficient fashion. Understanding of the factors which guide molecules into different patterns thus become an important goal for prediction and control of molecular patterns structures.

Here we present the formation of interlocked arrays (‘gear chains’) of pinwheels through self-assembly of 3-phenyl-propynenitrile (PPN) molecules on a Cu(111) surface. Variable temperature scanning tunneling microscopy (STM) reveals upon molecular deposition a pattern of small hexagonal features, which coalesce into sequences of larger, interlocking pinwheel-shaped structures. The pinwheels have an outer diameter as large as ~4nm. The driving force of this entropically disfavored pinwheel formation is discussed.

9:00am **NS-Thm4** Electronics and Mechanics of Single Molecule Circuits, L. Venkataraman, Columbia University

**INVITED** Understanding and controlling electron transfer across metal/organic interfaces is of critical importance to the field of organic electronics and photovoltaics. Single molecule devices offer an ideal test bed for probing charge transfer details at these interfaces. Results from these single-molecule measurements can be directly related directly theoretical models, unlike measurements at the ensemble level. The ability to fabricate single molecule devices and probe electron transfer reliably and reproducibly has enabled us to study and model transport through them.

In this talk, I will review the scanning tunneling microscope break-junction technique we use to measure electronic transport through single molecule junctions. I will discuss our measurements using novel metal-molecule link chemistries, including amines, phosphines[1] and results from recent work using tri-methyl tin linkers, which yield direct Au-C coupled single molecule junctions[2]. I will show how the intrinsic molecular properties influence measured single molecule conductance and bond rupture forces[3]. Finally, I will show how a mechanically controlled binary single molecule switch can be created using bipyridine molecules[4].


9:40am **NS-Thm6** Complex Rotation Mechanisms of a Molecular Machine Probed by STM, H. Kersell, U.G.E. Perera, Y. Zhang, University of California, Berkeley

**invited**

The complex rotation of a ruthenium based double-decker molecular rotor is resolved via ultrahigh vacuum low temperature scanning tunneling microscopy. The study was performed at temperatures of 4.2 K and 77 K on a Au(111) substrate. Inelastic electron tunneling (IET) was utilized to induce stepwise rotation of the molecule with respect to the surface. Subsequent rates of molecular switching, induced via tunneling currents, display sets of discrete energy minima with respect to the molecular stator and to the surface. The molecular rotator is composed of a set of semi-rigid arms connected to semi-rigid components, upon which IET induced rotation is observed to change conformation to the particular energy minima of the rotor. The resulting STM images were compared to calculated images of the same molecule. Additionally, the rotator was dissociated, revealing an intact stator adsorbed on the Au(111) surface. We acknowledge the financial support of US-DOE; DE-FG02-02ER46012, and NSF-PIRE; OISE 0750257 grants.

10:40am **NS-Thm9** Self-Assembled Double Strand DNA Monolayers as Spin Filters, Z. Xie, S.R. Cohen, T.Z. Markus, R. Naaman, Weizmann Institute of Science, Rehovot Israel

Spin control provides new and interesting opportunities for control and study of the factors governing electron transport. Recent work by Naaman and co-workers has shown that self-assembled monolayers of double stranded DNA (ds DNA) can act as a spin filter for electrons photoemitted from a gold substrate.[1,2] This phenomenon depends on the helicity of the ds-DNA, which leads to spin polarization and consequent capture of filtered electrons that tunnel back to the substrate. In this work, this effect is investigated for electron flow between two electrodes, a bottom Ni electrode to which one strand of the DNA is bound, and a top gold nanoparticle electrode which serves to identify the ds-DNA and provide good electrical contact through binding to the complementary strand. The current characteristics are measured by conductive scanning probe microscopy, as applied in a previous study of electron transport in DNA monolayers.[3] A magnetic field of approximately 0.3 T at the surface is provided by a permanent magnet placed below the sample. The results are consistent with the photomission work, namely marked differences in the current flow depending on magnetic field alignment. Furthermore, this effect depends on length of DNA chain, with longer chains providing a more significant effect relative to shorter ones. These experimental findings, together with a physical model will be presented.


11:00am **NS-Thm10** Controllable Phase Transition Using a Probing Tip, Q. Li, Oak Ridge National Laboratory

Molecular Self-assembled monolayers have applications in many different fields, such as sensing, lubrication and molecular electronics. In our study, a movie containing more that 50 STM images is acquired to investigate the dynamic behavior of the phenyl-acetylene molecular assembly process on Au(111) surface at LN2 temperature. Besides that, we found the phenyl-acetylene molecules are very sensitive to the polarity of the bias voltage. Therefore, phase transition between molecular ordered and disordered structure can be controllably achieved by applying positive or negative bias voltage. Systematically study of the phase transition process turned out it may be related to the charging effect of the molecules.


Novel nanocarbons such as fulleranes, nanotubes, graphene, and nanographite reside at the cutting edge of nanotechnology and technology. This paper presents a fundamental study of how size, shape, chemical functionalization, and bond strain affect electronic structure in several benchmark series of chemically pure, novel carbon-cage compounds ranging from diamondoids (a fully sp3 form of nanodiamond) to cubane. Size and shape are studied with the diamondoid series from adamantane to hexamantane, where the observed gap changes are primarily due to evolution in occupied states, as measured with photoelectron spectroscopy (XPS & UPS). Bond strain is studied with dodecahedrane, octahedrane, and cubane, where increasing bond strain leads to two major changes in the near-edge x-ray absorption fine structure (NEXAFS) spectra. First, a broad C-C σ* resonance in the absorption splits into two more narrow and intense resonances with increasing strain. Second, the first manifold of states previously associated with tertiary C-H σ* in the diamondoid series appears to broaden and shift to lower energy. This feature is more than twice as intense in cubane as octahedrane, even though these two molecules have similar stoichiometries (C20H2 vs. C20H3). We attribute the additional intensity to π* states, indicating a high degree of p interaction between parallel C-C bonds in the cubane.

**Thursday Morning, November 3, 2011**
Spin-capable multi-walled carbon nanotube (MWCNT) forests that can form webs, sheets, and yarns provide a promising means for advancing various technologies [1-4]. The important factors enabling the growth of the spin-capable forests are still not well understood. Growing spin-capable CNT forests depends on several growth factors such as the catalyst film thickness, the growth temperature, and the carrier and reactant gases [5-9]. Other factors still remain to be investigated more thoroughly. These include the flow rate (or ratio) of the reactant gas, the reactant gas species, and the pressure.

Herein we show how both the spinning capability and morphology of MWCNT forests are changed significantly by controlling the acetylene (C2H2) concentration and the thermal ramp rate. The acetylene gas flow is varied in the range of 0.25 ~ 7.5 % in volume. The MWCNTs grown at C2H2 concentrations between 1.5 ~ 3.5 % are well-aligned and are spine-capable. The well-aligned forests have higher areal density and shorter distances between the CNTs caused by strong Van der Waals interactions. CNTs grown at C2H2 concentrations under 1.5 % or over 3.5 % are curled and have random orientation. The resulting forests have reduced areal density and have poor spinnability. The thermal ramp rate is varied from 30 °C/min to 70 °C/min. Only the CNT forests grown with 50 °C/min condition are well-aligned and spinnable due to high areal density and closer spacing between adjacent CNTs. This condition alone results in Fe nanoparticles which have the proper size and density to produce spin-capable CNT forests.

Figure 1 shows SEM images and picture of spin-capable CNTs grown at 1.5 vol.% of acetylene and 50 °C/min on 70KΩ eq. film at 780°C for 5min with mixture of He, H2, and C2H2. The spinnable CNTs of 330 μm have good alignment which is dependent on the ability to form ribbons. From a 1 x 1 cm substrate, the CNTs can form a 4 m length sheet.


**Plasma Science and Technology Division Room: 202 - Session PS+TF-ThM**

**Plasma Deposition and Plasma Enhanced ALD Moderator:** S.-P. Tay, Mattson Technology Inc.

8:00am PS+TF-ThM1 High Quality SiNx by Microwave RLSA Plasma Enhanced Atomic Layer Deposition, **T. Karakawa**, M. Oka, N. Fukugi, H. Ueda, T. Nozawa, Tokyo Electron Technology Development Institute, INC, Japan

Shrinking critical dimensions of Ultra Large Scale Integration (ULSI) and optical device structures continue to drive advances in semiconductor fabrication processes. Three dimensional (3D) and metal gate structures for example, require low temperature dielectric layers (e.g., SiO2 and SiNx) with dimensions and film quality that may be met only by Plasma Enhanced Atomic Layer Deposition (PEALD) [1]. In addition to film quality and conformality, minimizing plasma damage in the PEALD process sequence is imperative. Prior to this work we characterized PEALD SiO2 deposition in a Radial Line Slot Antenna (RLSA) plasma source using bis-tetraetyl-buthyl-amino-silane (BTBAS) as a precursor [2]. In this study, we determined RLSA ALD process conditions favorable for SiNx film formation. The Si ALD precursor was dichlorosilane (DCS) and nitridation employed a NH3, N2 and Ar RLSA plasma. The wafer temperature was controlled below 400°C during the ALD process. Precursor adsorption time, temperature, plasma power were varied in order to determine the RLSA Plasma conditions resulting in the best SiNx ALD film quality. We obtained very high quality SiNx films having almost the same HF wet-etching rate as thermal LP-CVD SiNx (720°C) film. In this presentation, the results of Angle Resolved X-ray Photoelectron Spectroscopy (ARXPS) analyzed RLSA ALD deposited SiNx films will be presented. We found that the sub-nitride bonding ratio of the SiNx film was strongly correlated with the HF wet-etching rate, a measure of the film quality. The lower sub-nitride bonded SiNx film such as Si3N4 was created by the RLSA plasma at low temperature with low plasma density. The reason so little damage occurs is the low rate of ion bombardment on new Si3N4 surfaces during plasma nitridation.

Quantum dot sensitized solar cells using semiconductor nano-particles have attracted much interest because they are expected to have a high efficiency and a low manufacturing cost. Narrow band-gap semiconductors such as CdS, PbS, and CdSe are employed as sensitzers, and they transfer photogenerated electrons to them to large band-gap semiconductors such as TiO2 under light excitation. Our interest has been concerned with quantum dot solar cells using Si coumpound nano-particles because Si is abundant and has little toxicity. We have succeeded in producing Si nano-particles of a narrow size dispersion using a multi-hollow discharge plasma CVD method [1], and have applied them to Si quantum dot sensitized solar cells [2]. In our CVD system, discharges were sustained in 8 small holes of 5 mm in diameter at SiH4 and H2 flow rates of 2 and 48 sccm. Si nano-particles were nucleated, grew in SiH4/H2 plasma produced inside small holes, and were transported to the downstream region by neutral gas flow. We also performed surface nitridation of Si nano-particles to terminate dangling bond of the surface. Our experiments clearly demonstrated advantages of nitridation of Si nano-particles on the device performance; the short circuit current of Si QDs sensitized solar cells showed 1.3 times higher value by the nitridation and a photon to current conversion efficiency (PCE) achieved a high value of 40% at short wavelength of 350nm [3, 4]. Moreover, quantum dot sensitized solar cells using FeSi nanoparticles show better performance than those using Si nanoparticles. We will compare characteristics of three kinds of quantum dot sensitized solar cells using Si, SiSiN core shell, and FeSi nanoparticles and discuss relationship between optical and electrical properties of the nanoparticles and the device performance.


In this contribution, we report on the control of the composition and structure of films deposited in Ar/organosilicon precursor mixtures by using a remote plasma, i.e. the expanding thermal plasma, CVD process. The characterization of the deposited layers has been carried out by means of Fourier-transform infrared spectroscopy (FTIR), spectroscopic ellipsometry and X-ray photoelectron spectroscopy (XPS). The characterization of thin films showing a tunable chemical composition and optical properties has allowed identifying the main dissociation paths of the deposition precursor as controlled by the argon ions and electrons emanating from the plasma source, i.e. a cascaded arc, in the downstream region, where the monomer is injected. In particular, Ar ions are responsible for the charge exchange reaction with the monomer and electrons participate to the dissociative recombinations with the molecular ions generated in the first reaction, as already proven in the case of other molecular gases [1]. An optimum in the monomer structure retention of 35 % has been observed under conditions of low plasma reactivity, i.e. high monomer flow-to-(Ar+,e-) flow rate ratio.

[1] M. Creatore, Y. Barrell, J. Benedikt, M.C.M. van de Sanden, M. Creatore, Eindhoven University of Technology, the Netherlands

Organosilicon polymeric thin films (SiCxHyOz) have attracted considerable interest due to their wide range of applications such as interlayers in gas/moisture diffusion multi-layer systems, low dielectric constant interconnect materials in microelectronic circuits and biocompatible coatings for medical implants, to name a few. An accurate control of the optical and electrical properties of the nanoparticles and the device characteristics of three kinds of quantum dot sensitized solar cells using Si, SiSiN core shell, and FeSi nanoparticles and discuss relationship between optical and electrical properties of the nanoparticles and the device performance.

[1] M. Creatore, Y. Barrell, J. Benedikt, M.C.M. van de Sanden, M. Creatore, Eindhoven University of Technology, the Netherlands

In this work, plasma-enhanced atomic layer deposition (PE-ALD) of Lanthanide series gate dielectrics is being investigated. The study aims to demonstrate the uses of second generation high-k oxide materials in high-performance CMOS technology. We have characterized a number of ALD oxide and nitride dielectrics deposited on various substrates using continuous wave (cw) and pulsed excimer lamp-based ALD. In the O2 plasma-based ALD reactions, we have employed a single wafer R&D plasma-assisted ALD reactors. In the O2 plasma employed, vacuum ultraviolet (VUV) photons with energies up to 9.5 eV were detected and these photons were found to be able to generate electronic defects at thin film interfaces. This was demonstrated by experiments in which Al2O3 passivated Si(100) samples were exposed to O2 plasmas. By exposing the samples through quartz and MgF2 windows, the role of ions was excluded and the specific role of the high energy VUV photons was confirmed unambiguously. Furthermore, during regular ALD conditions, an ion energy of ~30 eV was measured. This energy is sufficient to contribute to the ALD process by, e.g., the displacement of lattice atoms and enhancement of the ALD surface reactions, however, it is low enough to prevent substantial damage to the deposited layers. The impact of the ions was further explored by enhancing the energy of the ions through the implementation of substrate biasing, either through substrate self-biasing or by RF biasing. By enhancing the ion energy up to 230 eV, these experiments demonstrated that at 300°C the crystallinity of TiO2 films can be changed from the anatase to the rutile crystalline phase. Moreover, at a substrate temperature of 200°C the rutile phase can be obtained when employing substrate biasing while normally amorphous TiO2 is obtained. These results are particularly significant as generally the deposition of rutile TiO2 is difficult to achieve by ALD due to substrate temperature limitations imposed by the precursors used. It is therefore evident that substrate biasing is a promising method to extend the possibilities of ALD.

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window of self-limiting ALD growth of 1.4 Å/cycle was observed at 250°C and below under certain precursor dose conditions. As the temperature increased, high-quality films are deposited, but the growth mechanism appears to become CVD-like. At 250°C, the refractive index of the film is stable at ~1.80 regardless of other deposition conditions, and the measured dispersion characteristics are comparable to those of bulk Gd2O3. The electrical characteristics of the films, such as fixed charge and dielectric constant, are extracted from C-V measurements using TiN metal gate capacitors, and will be reported.

The plasma etching rate of the ALD Gd2O3 film in a high-density helicon reactor is very low. Little difference is observed in etching rate between Cl2 and pure Ar plasmas, suggesting that physical sputtering dominates the etching process. A threshold bias power exists below which etching does not occur, thus it may be possible to etch a metal gate material and stop easily on the Gd2O3 gate dielectric.

One focus of the presentation will be the preparation and full characterisation of bimetallic Pd/Rh-CNT hybrids. The x-ray diffraction (XRD) and TEM (EDX) analyses were used to confirm that the deposited nano-particles are indeed truly Pd/Rh bimetallic, excluding the possibility of a simple physical aggregate/mixture of the two metals. Complementary analytical tools such as x-ray photoelectron spectroscopy (global information) and scanning transmission x-ray microscopy (truly local information) reveal that the particles contain a metal/oxygen ratio dependent of the processing gas; they testify also of the possibility of a nano-particle core-shell structure and of a reorganisation of its structure depending of the processing gas.

**Plasma Science and Technology Division**

**Room: 201 - Session PS-ThM**

**Neutral Beam and Low Damage Processing**

**Moderator:** S. Bouchoule, CNRS-LPN

**8:00am PS-ThM1 2010 Plasma Prize Lecture - Super-low Damage Top-down Processing for Future Nanoscale Devices. S. Samukawa**, Tohoku University, Japan

*For the past 30 years, plasma process technology has led in the efforts to shrink the pattern size of ultralarge-scale integrated (ULSI) devices. However, inherent problems in the plasma processes, such as charge build-up and UV photon radiation, limit the process performance for nanoscale devices. To overcome these problems and fabricate nanoscale devices in practice, we have proposed damage-free neutral-beam process. In this process, we introduce a developed damage-free etching technique: a designable deposition of super low-k SiOC film and low-temperature Si oxidation (thin SiO2) processes using neutral beams and discuss the actual applications of neutral beam processing for future nanoscale devices (such as, Fin-MOSFET, and Quantum Dot Solar Cell). Neutral beams can perform atomically damage-free etching, deposition and surface modification. Then, the neutral beam process can precisely control the atomic layer chemical reaction and defect generation. This technique is a promising candidate for the nano-fabrication technology in future nanoscale devices.*


To create high performance metal-oxide semiconductor devices, it is necessary to reduce variations in the critical dimension, the etching profile, and the amount of damage caused. Recent advancements in plasma processing for the gate electrode, sidewall, and high aspect contact hole have highlighted the importance of fully understanding how plasma induces damage and how to control this damage. We also need to find a way to quantitatively predict the damage depth profile using a numerical simulation that takes a realistic surface reaction into consideration, because it is quite difficult to observe the distribution of damage in the patterns with high aspect ratios. One common numerical simulation method, molecular dynamics (MD) calculation, unfortunately has a very limited range and cannot simultaneously consider a time-dependent etching profile in the 100 nm scale.

We developed a numerical simulation method for the distribution of plasma-induced physical damage to the SiO2 and Si layers during fluorocarbon plasma (C4F8/O2/Ar) etching. In our method, the surface layer is assumed to consist of two layers: a C-F polymer layer and a reactive layer. Physical and chemical reactions in the reactive layer divided into several physical and chemical components to the etching as well.

We used our simulation method to calculate the SiO2 etch rate, the thickness of the C-F polymer layer (Tc-F), the selectivity of SiO2 to Si layer, and the O2 etch rate of both the C-F polymer layer and the Si layer during C4F8/O2/Ar plasma etching in the steady state. Results demonstrated that calculation of the absolute values as well as their behaviors were consistent with those of our experimental data. We also successfully predicted depth profiles of physical damage to the Si and SiO2 layers in the steady state introducing our re-gridding method, which were affected by the Tc-F value. When we calculated the time-dependence of the amount of Si damage, we found that depth of the damage generated in the Si and the depth of the etching step during the SiO2/Si layer etching, in spite of the high selectivity. After that, the amount of damage was gradually decreased by etching and finally became constant.

These results demonstrate that the Tc-F value and the over etching time must be carefully controlled by process parameters to reduce the amount of damage during fluorocarbon plasma etching.

**9:00am PS-ThM4 Theoretical Analysis of Electron Transfer during the Process of Neutral Beam Generation**

**T. Inoue, H. M. Nakaya, T. Jiwasaki, K. Ono, Y. Irie**, Mirazizo Information & Research Institute, Inc., Japan, S. Ueki, BEANS Project 3D BEANS Center, Japan, O. Nukaga, Fujikura Ltd., Japan, T. Kubota, Tohoku University, Japan, M. Sugiyama, University of Tokyo, Japan, S. Samukawa, Tohoku University, Japan

We have developed numerical simulation software named QuickQD[1,2] that calculates the time-evolution of wave functions of electrons based on the First principles Quantum Mechanics. We have applied QuickQD for analyzing neutral-beam generated by the neutral-beam etching system developed by Samukawa, et al [3]. Positive or negative ions passing through a graphite aperture of this etching system are converted to neutral atoms by exchanging their valence electrons during a collision with the aperture sidewall.

Our numerical model consists of an ion (Cl- or Cl2+) and 24 carbon atoms forming a graphite sheet. The ion has been moved to collide with the graphite sheet and then bounce back to its original position. QuickQD has simulated the time-evolution of several ten wavefunctions of both the ion and carbon electrons during the whole process of the collision. We have evaluated the distribution of electron density around the ion, and we have determined the probability of neutralization. We have obtained some numerical results that agree well with experiment results, for example, a negative Cl- ion transfers its valence electrons to the graphite and is converted to a neutral Cl atom with high probability, meanwhile a positive Cl2+ ion receives valence electrons from the graphite and is converted to a neutral Cl2 molecule with low probability[4].

To understand the quantum process of electron transfer and the difference of neutralization efficiencies between a negative ion and a positive ion, we have investigated the behavior of each time-evolving electron wave function. We have found that some electrons that existed in the orbital of
BEANS Project, Japan

analyze the energy and angular distribution passing through the aperture.

To realize more practical neutral-beam etching system, we also have to neutralize by the collision with aperture sidewall. A negative ion transfers between incident particles and side wall of aperture. We determine initial aperture inlet, and have to describe the scattering process for collision.

In this study, we calculate the trajectory of particles generated by Monte-Carlo method, and analyze the energy and angular distribution of neutral beam at aperture outlet. To analyze energy and angular distribution at outlet, we have to determine the initial conditions of incident beam at aperture inlet, and have to describe the scattering process for collision between incident particles and side wall of aperture. We determine initial beam condition at inlet using the experimental data fitting. And we use the inelastic scattering model known as Hard-Cube model [3] for treating collision between particles and side wall of aperture. Particles injecting into aperture which have broad angular distribution are collimated by the inelastic scattering process.

Using above simulation scheme, we analyze the energy and angular distribution for specific aperture configuration (i.e. aperture aspect ratio) and compare with experimental data. We also analyze the particle distribution of neutral beam at silicon wafer injecting from aperture outlet and discuss the uniformity of neutral beam etching. Using above angular distribution of neutral beam and etching simulator, we predict etching shape by neutral beam generated by various aperture configurations (i.e. aperture aspect ratios).


9:40am PS-ThM5 Energy and Angular Distribution Analysis for Neutral Beam and Application for Etching Simulation. S. Ohsuka, N. Watanabe, T. Iwasaki, K. Ono, Mizuho Information & Research Institute, Inc., Japan; Y. Iriye, O. Nakagou, S. Ueki, BEANS Project 3D BEANS Center, Japan; T. Kubota, Tohoku University, Japan; M. Sugiyama, University of Tokyo, Japan; S. Samukawa, Tohoku University, Japan

The neutral-beam etching system developed by Samukawa et al. [1] has a carbon plate which has numerous apertures, where positive or negative ions pass through. In this system, most of those ions passing through the apertures are efficiently converted to neutral atoms. We consider ions are neutralized by the collision with aperture sidewall. A negative ion transfers some of its valence electrons to the aperture sidewall by the collision, and a positive ion receives some valence electrons from the aperture sidewall. The dynamical process of electron transfer by the collision can be described by the Quantum Mechanics as a time-evolution of wave-function during the collision, and we have developed computational simulation software named QuickQD [2].

To realize more practical neutral-beam etching system, we also have to analyze the energy and angular distribution passing through the aperture. Those distributions characterize the ability of neutral beam etching. In other words, to achieve efficient etching system, we have to develop optimized aperture configurations (i.e. arrangement and aspect ratio of apertures) and get conditions for suitable energy and angular distribution of neutral beam.

In this study, we calculate the trajectory of particles generated by Monte-Carlo method, and analyze the energy and angular distribution of neutral beam at aperture outlet. To analyze energy and angular distribution at outlet, we have to determine the initial conditions of incident beam at aperture inlet, and have to describe the scattering process for collision between incident particles and side wall of aperture. We determine initial beam condition at inlet using the experimental data fitting. And we use the inelastic scattering model known as Hard-Cube model [3] for treating collision between particles and side wall of aperture. Particles injecting into aperture which have broad angular distribution are collimated by the inelastic scattering process.

Using above simulation scheme, we analyze the energy and angular distribution for specific aperture configuration (i.e. aperture aspect ratio) and compare with experimental data. We also analyze the particle distribution of neutral beam at silicon wafer injecting from aperture outlet and discuss the uniformity of neutral beam etching. Using above angular distribution of neutral beam and etching simulator, we predict etching shape by neutral beam generated by various aperture configurations (i.e. aperture aspect ratios).


10:40am PS-ThM9 Improvement in the Evaluation Technique for Plasma-Etch Si Damage using Photoreflectance Spectroscopy with Temperature Control. A. Matsuda, Y. Nakahoko, T. Takao, K. Eriyuchi, K. Ono, Kyoto University, Japan

In etching processes during MOSFET fabrication, bombardment of high-energy ions from plasma creates damaged structures in layers near the substrate surface, characterized by defect sites in the crystalline structure such as vacancies, interstitials, displacements, and dangling bonds. The negative effects of the damage on device characteristics (such as drain current degradation) have become increasingly significant as devices scale down rapidly [1]. To realize low-damage process, an advanced damage evaluation technique is essential. Photoreflectance spectroscopy (PRS) is an optical technique capable of detecting damage with high sensitivity. The surface is perturbed with an electric field by a modulation laser beam, and the reflectivity of a probe beam is measured. PRS has been studied as an advanced technique for contactless damage evaluation.

In this study, we employed an improved technique, where the temperature of the sample is controlled with liquid N₂ during PRS measurement. N-type Si (100) wafers were exposed to an inductively coupled plasma using argon gas, under various rf-bias powers to control the incident ion energies (E). At room temperature, the amplitude of the spectral peak decreased with the increase in E. This feature is explained in earlier literature that carriers trapped at defect sites in the amorphous carbon layer in a decrease of the amplitude [2]. When the sample was cooled to 90 K, the spectra were enhanced and the peak of the high-E sample was revealed. By fitting the spectra to a functional form [3] and modeling the parameters’ temperature dependences, we found that the amplitude enhancement is related to the temperature dependence of the spectral broadening parameter (Γ). We also found that, at a fixed temperature, Γ for damaged surfaces were larger than that of the control sample. These results show that the damage causes changes in Vₐ and Γ, but by controlling the sample temperature, we were able to lower Γ, which increased the amplitude and revealed the peak.

The PRS-based damage evaluation technique discussed here expands the range of the applicable plasma damage conditions. Furthermore, temperature dependences of the spectra and its parameters (e.g. Γ and optical band gap) give us an insight into the band structures of damaged Si. This PRS-based technique is expected to be potentially useful as a future in-situ monitoring technique.

temperature oxidation process to form SiO\(_2\) films, residual compression stress between the Si substrate and SiO\(_2\) remains due to the difference in the thermal expansion coefficient between Si and SiO\(_2\). This residual stress degrades the sub-threshold characteristics of MOSFETs because it increases interfacial state density. Especially, in case of future 3-dimensional transistors such as FinFETs, stress concentration occurs at corner and edge of 3D structures, which may cause drastic increase of leakage current. There have been recent active investigations on the thermal oxidation of Ge to enable the fabrication of high-mobility Ge MOS transistors. After high-temperature thermal processes, however, Ge oxide has poor thermal stability and a large amount of suboxide.

We developed an alternative oxidation process to solve these problems using a low temperature neutral beam (NB) technique to form the gate dielectric film. In this study, we investigated the mechanism to form thin oxide films using a low-temperature neutral beam oxidation (NBO) process. Arhenius plot has shown that activation energy of NBO was extremely low, which enabled low-temperature oxidation at 300°C or even at room temperature. This should be because bombardment energy of oxygen beam assisted the oxidation reaction. Also, we investigated that the suboxide in thin oxide film using NBO process. As a result, there was little suboxide at the interface between oxide films and semiconductor using NBO process even at low-temperature. These results demonstrate the outstanding potential of the low-temperature NBO process for fabricating gate dielectric films.


This paper reports an attempt of low-temperature recovery for gallium nitride (GaN) damaged by the exposure of a chlorine-based etching plasma. The work is motivated by the challenge for manufacturing highly-reliable GaN-based devices which have a great deal of potential in optical, high-frequency, and high-power electronics field. So far, high temperature annealing or nitrogen plasma were used as the post-process. However, the high-temperature treatment could not restore the reduced atomic ratio of nitrogen on the damaged surface. The nitrogen plasma introduces damage by the ion bombardment. In this study, a high-density radical annealing in nitrogen on the damaged surface. The nitrogen plasma introduces damage to the GaN by the ion bombardment. So, the process was newly proposed and evaluated in all in-situ experimental system consisted of etching, radical annealing, and surface analysis.

Radicals and ions extracted from a chlorine plasma were irradiated to n-GaN with an ion dose of 5 \times 10^{16}cm^{-2} at 500 eV [1]. Then the radical annealing using nitrogen radical (N\(^+\)) or hydrogen radical (H\(^+\)) with a radical dose of 1 \times 10^{16}cm^{-2}was applied using a high-density radical source [2] at room temperature. The X-ray photoelectron spectroscopy (XPS) and atomic force microscopy (AFM) evaluation carried out after each process.

In N\(^+\) annealing case, the surface chloride was removed and N/Ga was improved, but there was no change in the composition ratio of oxygen (O). On the other hand, hydrogen radical (H\(^+\)) could efficiently remove O, and N/Ga was improved greatly. Other characteristic findings were that Ga-Ga (metallic Ga) was observed in the Ga 3d spectra and the surface roughness increased from 0.335 to 6.464 nm rms.

As a result, the radical annealing at room temperature was shown to be very effective for improving GaN stoichiometry to achieve a better interface, whereas the condition should be optimized very carefully.

Acknowledgments

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* Coburn & Winters Student Award Finalist

Advanced Surface Engineering Division
Room: 104 - Session SE-ThM

Nanostuctured Thin Films and Coatings
Moderator: C. Mitterer, University of Leoben, Austria


Memristors, nonvolatile bipolar resistive switching devices first intentionally fabricated in 2008 [1], have attracted attention for use in a wide range of applications. To date, most memristors have been fabricated from either TiO\(_x\) [1] or TaO\(_x\) [2]. However, it is necessary to explore other dielectric materials, because the memristive properties of these dielectrics have not yet been explored.

In this study, memristors were fabricated on both silicon and flexible polyethylene teraphthalate (PET) substrates using HfO\(_2\), HfSiO\(_x\), and ZrSiO\(_x\) thin films and thermally evaporated aluminum contacts. The dielectric thin films were formed from sol-gel precursors synthesized based on the recipe of Meena et al. [3]. Following synthesis, the precursors were spun on to substrates, exposed to an oxygen plasma, and annealed in air at 150°C for 24 hours. Memristors with cross sectional areas ranging from 2\times2 mm\(^2\) to 0.5\times0.5 mm\(^2\) were fabricated.

The dielectric films were extensively characterized by using optical profilometry and microscopy, x-ray photoelectron spectroscopy (XPS), and deep ultraviolet spectroscopic ellipsometry (SE). The optical profilometry measurements demonstrate that the films show no evidence of significant inhomogeneities due to the surface-based sol-gel synthesis methods. According to the XPS measurements, the films are \(\approx\)50% carbon. Such high levels of carbon are reasonable given the synthesis approach employed. and
similarly high levels of carbon have been observed previously in TiO₂ films synthesized from sol-gels [4]. From the SE measurements, it is evident that the bandgaps of the HSO₃, H₂SO₃, and ZrO₂ films are similar to their predicted values, but the densities of the films are ~0.5 due to the high amounts of excess carbon. Transmission electron microscopy measurements of the films will also be discussed.

Current-voltage (I-V), capacitance-voltage (C-V), capacitance-frequency (C-f), and conductance-frequency (G-f) measurements were performed at room temperature in a commercial probe station. The I-V measurements indicate that devices fabricated from these films exhibit bipolar resistive switching. Furthermore, optical microscopy images indicate that the switching events are often accompanied by significant changes to the morphology of the active region of the device, particularly when the switching occurs at high voltages. These changes suggest the formation of localized conduction pathways through the dielectric. The C-f and G-f measurements indicate that the capacitive properties of the devices also change following switching events. Additionally, after 24 days, retention tests show the devices are still in the ON state, thus, they may be considered nonvolatile.

References
mode both the Nb/C ratio and the Ge content was varied to study the effect of composition. XRD, TEM, and SEM showed that the NbC films are columnar and exhibit a preferred 111 crystallographic orientation. With increasing Ge content up to 10 at.%, the NbC grain size was reduced to ~5 nm with a random orientation. While the nc-NbC/Ge nanocomposites have no discernable amorphous tissue phase, XPS showed predominantly Ge-Ge with only little Ge-C bonds. In contact resistance measurements the films performed well (1.7-3.7 mΩ) in comparison to Ag reference (1.2 mΩ) at a contact force of 10 N.


11:20am SE-ThM11 Tuning the Properties of Chromium Oxynitride Coatings. L. Castaldl, Oerlikon Balzers AG, Liechtenstein, J. Patscheider, EMPA, Switzerland, V. Slikkover, ETH Zurich, Switzerland, D. Karapov, A. Reiter, Oerlikon Balzers AG, Liechtenstein

CrO,N thin films deposited by reactive cathodic arc evaporation were investigated for concentration levels oxygen between 0 and 0.1. The present study shows some of the possibilities for tuning the crystallographic and electronic properties of chromium oxynitride films by progressively increasing the oxygen concentration. Upon oxygen addition the CoO,N films remain in the face-centered cubic B1 structure of CrN for oxygen fractions of up to x = 0.70. At higher oxygen contents the properties of the films approach those of CrO and crystallize in the corundum structure. The gradual decrease of the stress-free lattice parameter of the films with the B1 structure at higher oxygen concentrations is consistent with oxygen substitution for progressively nitrogen atoms. The almost random orientation of oxygen-free CrN films transforms to a distinct (001) preferred orientation with the increase of oxygen concentration in the films. XPS investigations prove enhanced charge transfer from Cr to O 1s with increasing x with respect to N by changes in the core-level binding energies (BE) of the Cr 2p1/2 and N. A shift of the valence band edge toward higher BE indicates a decrease of electrical conductivity of the films with increasing oxygen content.

11:40am SE-ThM12 Synthesis of Al-Cr-O and Al-Cr-O-N Thin Films in Corundum-Type Structure by Reactive r.f. Magnetron Sputtering. M. Stueber, Karlsruhe Institute of Technology (KIT), Germany, D. Dieckle, Walter AG, Germany, H. Leiste, S. Ulrich, Karlsruhe Institute of Technology (KIT), Germany

The PVD synthesis of wear and oxidation resistant aluminum oxide and derivative coatings is currently attracting large scientific and technical interest. Ternary Al-Cr-O thin films with mechanical properties comparable or superior to binary Al-O thin films can be deposited at moderate deposition temperatures. New coatings from the quaternary Al-Cr-O-N system could even offer increased strength, hardness and toughness. A combinatorial approach to the growth of Al-Cr-O-N thin films by means of reactive deposition/magnetron sputtering will be presented. For specific deposition conditions well adherent, nanocrystalline Al-Cr-O-N thin films with high Vickers hardness and elastic modulus values were grown at non-equilibrium conditions on cemented carbide and silicon substrates. Detailed results on the coatings composition, constitution, microstructure and properties will be presented and discussed in comparison to ternary Al-Cr-O thin films deposited under identical conditions.

Surface Science Division Room: 107 - Session SS-ThM

Oxide Surface Structure & Reactivity

Moderator: G.A. Kimmel, Pacific Northwest National Laboratory

8:00am SS-ThM1 Bond Activation of Alkanes on CaO(100). A. Chakradhar, U. Burghaus, North Dakota State University

The molecular and dissociative adsorption kinetics, respectively, of ethane, butane, pentane, and hexane are studied on CaO(100) using thermal desorption spectroscopy (TDS) and Auger electron spectroscopy (AES). A three-step adsorption/desorption cycles, AES scans show no carbon residuals on the surface. Molecular adsorption as well as bond breaking is observed for all alkanes studied, except for ethane. Molecular desorption of these alkanes is characterized by two TDS peaks while recording the parent mass. These two features are assigned to different adsorption sites/configurations of the alkanes. In addition, the binding energies of the bound alkanes, based on their desorption temperatures, are calculated by using Redhead equation. The bond activation is evident from desorption of hydrogen and mostly methane and ethylene fragments in multi-mass TDS experiments. Bond activation of hydrocarbons is pertinent for the petroleum industry, providing transportation fuels and other petrochemical products.

11:40am SS-ThM3 Organometallics as Probes of Functional Group Distribution on Oxide Surfaces. S.L. Scott, University of California, Santa Barbara

The reactivity of the hydroxyl-terminated surface of silica is important for constructing tailored interfaces in many applications, including catalysis, separations, and microelectronics. Amorphous silicas are particularly interesting because of their high surface areas. The hydroxyl content of these surfaces can be reduced thermally, but it decreases slowly because the hydroxyls have very low mobility. Their mutual condensation requires proximity, and, frequently, the formation of strained siloxane rings. Since these silicas have no long-range order, it is often postulated that the hydroxyl groups are distributed randomly, and that when the nominal hydroxyl density is low (e.g., after thermal treatment at temperatures exceeding 500 °C), these hydroxyls can be considered essentially isolated. Curiously, the Ga K-edge EXAFS shows that the reaction of Ga(CH3)3 with such a silica pretreated at 800 °C generates only digallium sites, requiring that the hydroxyls be organized pair-wise, likely as vicinal silanols linked by a single siloxane bond. The origin of the preferred vicinal disposition may be hydrolysis of a strained siloxane bond. Where isolated hydroxyls do exist, they appear to be located on reactive 2-rings whose reactivity resembles that of a masked silanol. Consequently, it is unwise to assume that site isolation results from grafting onto these silicas, although the vicinal disposition of silanols provides an opportunity to create tailored bifunctional surfaces.

9:20am SS-ThM5 Are MgO Thin Films More Reactive Than Bulk MgO Surfaces?, G. Cabailh, R. Lazzari, H. Cruguel, J. Jupille, UPMC and CNRS, France, L. Savio, IMEM-CNR, France, M. Smerieri, A. Orzelli, L. Vattuone, M. Rocca, Università di Genova, Italy and IMEM-CNR, France

The ubiquity of the interface between water and oxide materials has prompted a tremendous activity to determine the adsorption mechanisms at the microscopic scale on crystalline surfaces of simple oxides. Among these, magnesium oxide MgO offers the advantage of having various morphologies of good crystalline quality, involving cleaved crystal surfaces, high surface area samples and supported films, all of these dominated by the low index (100) orientation. On bulk MgO, the fivefold coordinated atoms of the basal (100) surface do not dissociate isolated H2O molecules. Conversely, H2O is easily dissociated at low-coordinated sites such as steps and kinks. A puzzling case is the adsorption of H2O on metal-supported MgO films in the monolayer range of which coordinate OH groups has been estimated to 60 to 70% of a monolayer [1,2]. Similar OH coverages were obtained by aging freshly prepared films in the residual atmosphere of the vacuum chamber [1], although the observation was contradicted by the finding of a more modest effect that cast doubt on OH being the cause of aging [2]. The extraordinary uptake of OH groups was attributed to the peculiarities of the electronic properties of the thin supported MgO films [3]. However, density functional theory hardly supports this view. Little changes in both the electronic structure and the capacity to dissociate H2O are predicted for monolayer-thick MgO(100) islands that, at variance with experiment, are only expected to dissociate H2O molecules along their borders [4].

In an attempt to solve the discrepancies between experiments and between experiment and simulation, MgO films of different stoichiometry were grown on MgO(100) by reactive deposition of Mg in an O2 partial pressure mixture, prior to being exposed to H2O vapor and/or aged in vacuum [5]. Films were observed by scanning tunneling microscopy (STM) and water uptake was analyzed by x-ray photoemission spectroscopy (XPS). The stoichiometry
and, consequently, the chemical activity towards hydroxilation of the MgO(100) films, was shown to strongly depend on the O$_2$ pressure during the film growth. Oxygen-deficient films undergo dramatic oxygen uptake either by exposure to H$_2$O or by aging in vacuum. Conversely, on stoichiometric MgO islands, XPS analysis and STM images are consistent with the prediction that H$_2$O only dissociates at the island edges.


9:40am SS-ThM6 The Adsorption of Silver on Fe$_2$O$_3$(111) Studied by Adsorption Microcalorimetry, LEIS, and AES. J.C. Sharp, Y.K. Yoo, C.T. Campbell, University of Washington  
Noble metals supported on oxide surfaces are of interest due to their catalytic activity and their sintering resistance is a subject of concern in terms of their long-term stability under reaction conditions. The heat of adsorption of silver on Fe$_2$O$_3$(111) grown on Pt(111) was measured using adsorption microcalorimetry, and its growth morphology was measured with low-energy ion scattering spectroscopy (LEIS) and Auger electron spectroscopy (AES). The initial sticking was 0.96 rising to 0.99 after 1 monolayer of Ag deposited. The LEIS and AES data were fitted to a hemispherical cap model for the Ag nanoparticles, which were found to grow with a particle density of 4 x 10$^{15}$ particles per cm$^2$. The initial heat of adsorption was found to be 220 kJ/mol, rising to 285 kJ/mol after 1 monolayer of deposited Ag. From these data, we extracted the energy of metal atoms versus the Ag particle size to which they attach on Fe$_2$O$_3$(111), and compare it to results on other single crystalline oxide surfaces.

Titanium(V) oxide (TiO$_2$), acting as a stable support for photocatalysts and sensitizers, has applications in catalysis and energy science. Metal nanoparticles supported on TiO$_2$ have been shown to have unusual reactivity. The deposition of metal nanoparticles can alter the electronic properties of the TiO$_2$ nanoparticles. To understand the influence of metal deposition, we photodeposit platinum (Pt) onto ordered linear arrays of TiO$_2$ nanoparticles. The Pt (0.3 nm) deposition at the step edges of highly oriented pyrolitic graphite. X-ray photoelectron and absorption spectroscopies were used to explore the electronic structure of the TiO$_2$ nanoparticle arrays with and without photodeposited Pt. The titanium L-edge spectra, supported by multiplet calculations, provide crucial information about crystal field effects, atomic multiplet interactions, and the phase of the TiO$_2$ particles. Valence photoelectron results and band structure calculations using density-functional theory indicate a narrowing of the TiO$_2$ band gap when Pt is loaded onto the TiO$_2$ nanoparticle surface. This suggests that Pt photodeposition onto linear TiO$_2$ nanoparticle arrays may enhance the solar absorption of TiO$_2$ due to narrowing of the TiO$_2$ bandgap.

MgAl$_2$O$_4$(100) is a promising material for substitution of ZnO. As a representative TCO for liquid transportation fuels: Sunshine to Petrol. Sandia is a multi-program laboratory operated by Sandia Corporation, a wholly owned subsidiary of Lockheed Martin Company, for the U.S. Department of Energy’s NNSA under contract DE-AC0494AL85000.  
Nevertheless studies on GZO films grown by ALD at low temperature condition which can be applied to flexible devices were still not carried out as far as we know of its potential. In this study, we investigated the growth characteristics and film properties of low temperature ALD (LT-ALD) GZO films by varying deposition method. Field emission scanning electron microscopy (FE-SEM) observation of the GZO films deposited on 51 via patterns showed that the film has excellent conformality with over 95 % coverage even at room temperature growth. Additionally, the electrical and microstructural properties were assessed by various analytical techniques including X-ray photoelectron spectroscopy (XPS), X-ray diffraction (XRD) and atomic force microscope (AFM). Also spectrophotometer was used to measure a transmittance of the film and showed high transmittance that could be applicable to transparent devices.

8:20am TC+AS+EM-ThM2 Enhancement of C-Axis Orientation of Ga-doped ZnO Films Deposited on Unintentionally Heated Glass Substrates using Nanosheet Seed Layers. H. Makino, Kochi Univ. of Tech., Japan, T. Shibata, NIMS, Japan, N. Yamamoto, Kochi Univ. of Tech., Japan, T. Sasaki, NIMS, Japan, T. Yamamoto, Kochi Univ. of Tech., Japan

Ga-doped ZnO (GZO) film is one of promising candidates as substitute for ITO transparent electrodes in optoelectronic devices. Control of structural properties, especially c-axis orientation, is crucial issue to improve electrical properties of polycrystalline GZO films on glass or plastic substrates [1]. Recently, nanosheet seed layers were proposed to control crystal orientation of oxides films on amorphous substrates [2]. In this study, we employed a seed layer of tungsten oxide nanosheets with two-dimensional hexagonal lattice structure for deposition of GZO thin films on unintentionally heated glass substrates. The nanosheets were assembled on glass substrates by Langmuir-Blodgett method. The GZO films with thickness of 100 nm were deposited by an ion-particle with different ambient and deposited unintentionally heated glass substrates with and without the nanosheet seed layers. The crystal structural properties were characterized by x-ray diffraction measurements. The c-axis orientation of the GZO films was drastically enhanced by the nanosheet seed layers. The intensity of (002) diffraction peak of GZO films deposited with the seed layers was about 40 times as strong as that deposited without the seed layers. The degree of c-axis orientation was evaluated by the (002) x-ray rocking curve (XRC). The full-width half-maximum of XRC of the GZO films on the nanosheet seed layers was 2.6 °, which is even lower than that of GZO films deposited on bare glass substrate at 200 °C.

The electrical properties were characterized by Hall effect measurements at room temperature. The GZO films deposited on the nanosheet seed layers showed the resistivity of 2.9 x10³ Ωcm with the Hall mobility of 24 cm²/Vs and the carrier concentration of 9.0 x10²⁰ cm⁻³. On the other hand, the GZO film deposited without the nanosheet seed layers showed the resistivity of 5.0 x10⁴ Ωcm with the Hall mobility of 17 cm²/Vs and the carrier concentration of 7.2 x10²⁰ cm⁻³. Both the Hall mobility and the carrier concentration were improved by the nanosheet seed layers. [1] T. Yamada et al., J. Appl. Phys. 107, 123534 (2010). [2] T. Shibata et al., Adv. Mater. 20, 231 (2008).

8:40am TC+AS+EM-ThM3 Multi-component Transparent Conducting Oxides: Progress in Materials Modeling S.-H. Wei, National Renewable Energy Laboratory

INVITED

Transparent conducting oxides (TCOs) play an essential role in modern optoelectronic devices through their combination of electrical conductivity and optical transparency. We review recent progress in our understanding of multi-component TCOs formed from solid solutions of ZnO, In₂O₃, Ga₂O₃ and Al₂O₃, with a particular emphasis on the contributions of materials modeling, primarily based on Density Functional Theory. In particular, we highlight three major results from our work: (i) the fundamental principles governing the crystal structures of multi-component oxide structures including (In₂O₃)(ZnO)n, named IZO, and (In₂O₃)(Ga₂O₃)(ZnO)n, named IGZO; (ii) the relationship between elemental composition and optical and electrical behavior; (iii) the origin of high-performance of amorphous oxide semiconductors. From these advances, the challenge of the rational design of novel electronic materials is discussed.

9:20am TC+AS+EM-ThM5 Composition Control of Electron Beam Deposited Nb-TiO₂ Thin Films. N.A. Beckers, T.R. Tucker, University of Alberta, Canada M.D. Fleischauer, NRC-National Institute for Nanotechnology, Canada, M.J. Brett, University of Alberta, Canada

Nb-doped TiO₂ has been identified as a potential indium-free transparent conductor, and has been fabricated by pulsed laser deposition and sputtering with good success. Other deposition methods, such as electron beam evaporation, are of interest for this material but have had limited demonstration to date. It would be advantageous to be able to use electron beam evaporation because in addition to planar films, the collimated flux allows for structured thin films via glancing angle deposition (GLAD). Composition control is essential for doped functional materials, which is typically difficult to obtain through physical mixing of source materials. Here we show that using a sol gel approach to prepare the source materials provides a possible route to circumvent this issue. The facile, solution based sol gel process for the synthesis of ceramic and glassy materials allows for precise composition control by altering the amounts and ratios of the metal oxide precursors. A sol gel methodology was used to prepare a series of Nb-doped TiO₂ deposition films with the following compositions: Nb₀₂₋₀₆Ta₂O₇O₂, Nb₁₂Ta₂O₇O₂, and Nb₂Ta₄₋₅O₇O₂. We will show that XPS results confirm the composition of the electron beam deposited films and reflects the composition of the source materials. Precise suppliers of Nb₂O₅ and TiO₂ do not show the same transformation as the sol-gel derived source material. Details on the effects of the post-deposition annealing environment on the thin film optical and electrical properties will also be presented.

References:

9:40am TC+AS+EM-ThM6 Laboratory and Production-Scale Low-Temperature Transparent Conducting Oxide Deposition. E. Ritz, University of Illinois at Urbana Champaign, G.B. Rayner, Kurt J. Lesker Company, D. Andreyczuk, Urbana Champaign, E. Dockstader, Kurt J. Lesker Company, D.N. Ruiz, University of Illinois at Urbana Champaign

Transparent conducting oxides (TCOs) are a class of materials that are becoming increasingly inarguable in our daily lives due to their use in electronic displays and mobile devices. There is a strong need to develop an economical deposition technique that allows for high transparency films with high electrical conductivity while replacing costly materials such as Indium Tin Oxide (ITO) with alternatives such as Aluminum-doped Zinc Oxide (AZO). In addition, a low-temperature deposition method would allow creation of TCOs on flexible plastic substrates, such as polyethylene terephthalate (PET). By using a dual DC magnetron system with a secondary RF antenna running at 13.56 MHz, a process has been developed that can deposit TCO films without significantly heating the substrate while maintaining high transmission and electrical properties. This capability has been demonstrated on a small-scale experimental setup utilizing 3-inch diameter circular magnetrons as well as a prototype production-scale chamber operating with 18x3.5 inch rectangular magnetrons aimed at flexible photovoltaic manufacturing. Using an immersed inductive RF antenna, ionization fraction can be increased to over 80%, measured by a gridded energy analyzer, and plasma density increased by an order of magnitude from 10⁵ cm⁻³ to 10⁶ cm⁻³, as measured by Langmuir probe. The secondary plasma deposits energy in the film without heating the substrate above 100°C while still achieving film resistivity on the order of 10⁻¹⁰Ωm-cm (measured by four-point probe method) and transparency of greater than 90% in the visible wavelengths (measured by spectrophotometry.) Adjusting the RF power (0-1000W) and the oxygen content (0-5%) in the plasma enables the ability to tune the film transparency and conductivity to desired levels. Crystal formation of films analyzed by x-ray diffraction (XRD) and elemental composition determined by x-ray photoelectron spectroscopy (XPS).

10:40am TC+AS+EM-ThM9 Optical and Electronic Properties of Photonic Crystal Based Transparent Conductors. S. Narayanan, M. Bockstaller, L. Porter, Carnegie Mellon University

Transparent conductors are becoming ubiquitous in a host of civil and military applications, including transparent electrical contacts in solar cells and LEDs, heated glass for aircraft and automobile windows, and electrochromic devices and smart windows. However, finding abundant materials with optimal electrical and optical properties and that can be produced economically is a particular challenge. Moreover, the supply and large demand, of late, for indium has inspired focused research on finding alternatives to ITO as a transparent conductor. We report here a novel approach to control optical properties such as absorption, transmission and reflection in multilayered structures (based on[1-2]) with absorbing components. Appropriate combination of materials could, among others, allow for applications like transparent electrodes, transparent electromagnetic shielding, flexible transparent conductors, etc. Through this study, we have demonstrated the validity of this approach using a few different materials combinations including polymer/metal and metal/ceramic systems. In these realizations the approach was shown to
increase the transparency in the visible frequency range by ~ 3 orders of magnitude as compared to the reference materials. For example, transmittances of 30-50% of incident light in the visible region were measured for films containing a polymer (polystyrene – PS) and an amount of metal (gold – Au) that was 3-4 times as thick as its skin depth (~ 40-50 nm). We have also found compatibility between the observed experimental results and numerical simulations. Apart from enhanced optical transmittance, resistivity values of ~ 10^8 Ohm-cm (comparable to Au films having ~10 Ohm-cm) have also been discovered in structures having insulating components. Our ongoing and future work is focused on alternative structures to enhance conductivity in the transverse direction as well as incorporate flexibility in the same. References: [1] M. Scalora et al., J. Appl. Phys. 83 5 (1998) 2377-2383 [2] M. Scalora, M. J. Bloemer, C. M. Bowden, Optics and Photonics News 10 9 (1999) 24-27

11:00am TC+AS+EM-ThM10 Effect of Plasma Treatment and Annealing on the Electrical Properties of Spin-Coated Colloidal ITO Films, S.M. Joshi, G.W. Book, R.A. Gerhardt, Georgia Institute of Technology

Colloidal ITO based inks may be an attractive route to direct writing transparent circuits and also be useful for the fabrication of transparent conductive ITO films on complex shaped substrates. The presence of stabilizing ligands and minimal contact between the ITO nanoparticles in the deposited films can be a challenge in obtaining the optimum electrical properties. This study investigates the effect of plasma treatment and annealing on the electrical properties of colloidal ITO films. Crystalline colloidal ITO nanoparticles were synthesized in-house by a non-aqueous technique. The solutions were spin coated onto glass and quartz substrates and their electrical and optical properties were evaluated. All films were found to be completely transparent, while the as-deposited films had resistivities more than 10^8 ohm-cm. Plasma treatments were shown to be effective in removing residual organics in the films, and even without annealing, some recipes were able to reduce the film resistivity by more than four orders of magnitude. Plasma treatments, when done in combination with annealing, resulted in films with resistivities less than 1 ohm-cm.


The field of flexible electronics has expanded tremendously over the past few years. Similar to what happened in silicon integrated circuit technology 40 years ago, flexible electronics are now at a point where system design and product integration will drive the technology. Flexible electronics will likely push the limits of material performance, process integration, circuit design, and system integration to demonstrate the full potential of flexible electronics. In general, key components for any flexible electronic application include thin film transistors. In order to be competitive with state-of-the-art a-Si:H thin film transistors, any other thin film transistor technology must show reproducible transistor parameters such as mobility, threshold voltage, drive current and reliability.

A grand challenge in flexible, thin-film-transistor (TFT) circuitry is the development of complementary metal oxide semiconductor (CMOS) circuits. Although flexible digital circuits, flexible sensors and substrates and solar cells have already been demonstrated, the missing technology piece that must be developed is flexible analog circuitry. For example, an operational amplifier will enable the interface to most sensors and actuators, significantly expanding the functionality of flexible electronic systems. In this paper, we will present n- and p-type chalcogenide-based materials that can be used as the building blocks for analog CMOS-based circuits. In particular, we will introduce the use of chemical bath deposition as an alternative to deposit these materials and will discuss the correlation between device characteristics and materials properties. Photolithography-based chalcogenide-based TFTs processed by chemical bath deposition achieved mobilities in the order of 10-25 cm^2/V-s. In addition, we have demonstrated hybrid CMOS for a-Si-Pentacene, CdS-Pentacene and CdS-TIPS Pentacene. We also present the impact of semiconductor thickness, gate dielectrics and contact in device performance. In addition, NAND, NOR and Inverters are demonstrated using chalcogenide-based materials integrated with either a-Si or pentacene. Device processing is carried out at a maximum processing temperature of 110ºC, which is compatible with most plastic substrates.

Thin Film Division
Room: 109 - Session TF1-ThM

Post-Deposition Processing and Characterization of Thin Films
Moderator: C. Vallée, LTM/CNRS-UFJ, France

8:00am TF1-ThM1 Protrusions, Surface Grains and Extended Single-Crystalline Plates, A. González-González, Instituto de Ciencia de Materiales de Madrid, Consejo Superior de Investigaciones Científicas, Spain, G.M. Alonzo Medina, A.I. Oliva, Centro de Investigaciones y de Estudios Avanzados del IPN, CINVESTAV Unidad de Mérida, Mexico, C. Polop, Universidad Autónoma de Madrid, Facultad de Ciencias, Spain, E. Rodríguez Cañas, J.L. Sacedón, E. Vasco, Instituto de Ciencia de Materiales de Madrid, Consejo Superior de Investigaciones Científicas, Spain

The processes controlling shape transformations during post-deposition treatments attract great interest due to their technological implications. As an example, residual stress relaxation during thermal annealing could influence component lifetime and performance in applications ranging from microelectronics to mechanical coatings, affecting their functional properties [1]. In this work, we investigate the morphology evolution during thermal annealing at 0.2 T_melting in polycrystalline Au(111) films using atomic force [scanning electron] (AFM [SEM]) microscopies and x-ray diffraction in order to follow both shape and structural transformations of surface features for different annealing times. Before annealing, a high density of round-like surface grains—which are surface components of an underlying polycrystalline microstructure—is observed. During annealing, surface grains merge together into “multigrain” structures that expand laterally up to a saturation size, suggesting a size-dependent phenomenon controlling surface recrystallization. By comparison with Au polycrystalline growth fronts for the same temperature range [2] (T zone of the zone models [1]), we can relate the annealing-generated multigrain structures to the incipient formation of extended sub-micrometric-sized plates that are observed for thinner films. They contain many small surface grains with low-angle or no inner grain boundaries [2]. The small amount of material to be recrystallized for each component favors the multi-component extension of recrystallized zone. Interestingly, results here reported concerning the recrystallization phenomenon are discussed on the basis of local interactions between surface grains triggering plausible elastic/plastic mechanisms for stress accommodation (grain zipping and shear strain) and relaxation by surface diffusion processes [1,3], which in principle seems not to be very different from those expected in the coalescence stage at the T zone.


8:20am TF1-ThM2 Characterization of Mg Acceptors in GaN:Mg Grown by Metal Modulated Epitaxy and MOCVD, J.E. Lowder, M.W. Moseley, B. Gunning, W.A. Doolittle, Georgia Institute of Technology, M.E. Zwanut, J. Dashdorj, University of Alabama

Acceptor doping of III-Nitrides has been the subject of many studies due to the relative low hole concentration (~10^17-10^18 cm^-3) material commonly grown in contemporary devices. This limitation is thought to be one of the contributing factors to efficiency droop in LED’s. Recently, extremely high hole concentration (p>4x10^17 cm^-3) material with very high (~50%) ionization efficiency (see Fig. 1) has been demonstrated by metal modulated epitaxy (EME), an application of MBE where surface chemistry is controlled via shuttering of the source material [J. Appl. Phys. 106, 014905]. The samples in this study are characterized by temperature dependent Hall measurement to elucidate the reduction in the acceptor activation energy. Electron paramagnetic resonance spectroscopy (EPR) was used to show the microscopic view of the Mg acceptor without influence of the surrounding crystal as well as the effect of annealing temperature on the EPR signal. It is found that the primary transport mechanism is likely due to impurity band conduction, consistent with a distributed acceptor band as opposed to an isolated acceptor energy level as is generally observed in p-type GaN. Likewise, the interaction of the Mg with hydrogen shows different annealing behavior when compared to MOCVD p-type GaN.

The samples analyzed by EPR were annealed at various temperatures in forming gas (H2:N2). Fig. 2 shows the relative intensity from the EPR signal of the neutral Mg acceptor as a function of forming gas anneal temperature.
Thermal annealing is a critical but often neglected step in semiconductor fabrication. It has been routinely used to activate the ion implanted dopant and at the same time, to restore the single crystal structure. However, it is also a powerful tool in repairing damages in bulk films as well as at film interfaces, which are caused by the plasma-related deposition or etching process. In addition, the thermal annealing step can change film characteristics, which affects the subsequent process result. In this talk, we will review some of his recent work related to thermal annealing, which involves three types of solid-state devices and thin films. First, an example of repairing the RIE damaged a-Si:H TFTs with a low temperature thermal annealing process will be given. This step reduced defects in the bulk a-Si:H and SiNx layers as well as at the gate-dielectric interface. This result has been interpreted in all TFT LCD productions around the world. In the second example, it will be shown that the thermal annealing step changed the grain size of the sputter deposited copper thin film. It resulted in the change of the copper-C1 reaction mechanism and therefore, the copper consumption rate. This is critical to a new plasma-based copper etch process that has been interpreted in large-area TFT LCDs and BiCMOS chips. In the third example, it will be shown that the post deposition anneal (PDA) condition is the key to the realization of a nonvolatile memory device, i.e. the nanocrystals embedded high-k capacitors or MOSFETs. Furthermore, an additional low temperature annealing step removed the defects generated in the gate and back-contact sputtering processes. In summary, the thermal annealing effects are non-negligible because they are directly involved in product reliability and yield.


8:40am TF1-ThM3 Post Deposition Annealing Effects on Thin Film Material, Process, and Device Properties. Y. Kuo, Texas A&M University

INVITED

Thermal annealing is a critical but often neglected step in semiconductor production. It has been routinely used to activate the ion implanted dopant and at the same time, to restore the single crystal structure. However, it is also a powerful tool in repairing damages in bulk films as well as at film interfaces, which are caused by the plasma-related deposition or etching process. In addition, the thermal annealing step can change film characteristics, which affects the subsequent process result. In this talk, we will review some of his recent work related to thermal annealing, which involves three types of solid-state devices and thin films. First, an example of repairing the RIE damaged a-Si:H TFTs with a low temperature thermal annealing process will be given. This step reduced defects in the bulk a-Si:H and SiNx layers as well as at the gate-dielectric interface. This result has been interpreted in all TFT LCD productions around the world. In the second example, it will be shown that the thermal annealing step changed the grain size of the sputter deposited copper thin film. It resulted in the change of the copper-C1 reaction mechanism and therefore, the copper consumption rate. This is critical to a new plasma-based copper etch process that has been interpreted in large-area TFT LCDs and BiCMOS chips. In the third example, it will be shown that the post deposition anneal (PDA) condition is the key to the realization of a nonvolatile memory device, i.e. the nanocrystals embedded high-k capacitors or MOSFETs. Furthermore, an additional low temperature annealing step removed the defects generated in the gate and back-contact sputtering processes. In summary, the thermal annealing effects are non-negligible because they are directly involved in product reliability and yield.


9:20am TF1-ThM5 Semiconductor Thin Film Metrology using Coherent Acoustic Phonon Spectroscopy. A.D. Steigerwald, K. Varga, A.B. Hmelo, Vanderbilt University; L. Feldman, Rutgers University; N.H. Tolk, Vanderbilt University

Here we discuss the use of coherent acoustic phonon spectroscopy as a noninvasive and nondestructive measurement tool for semiconductor thin films. Specifically we emphasize its usefulness in studying the optoelectronic structure of materials with various types of structural defects. We discuss in-depth studies of ion-implanted GaAs, demonstrating that the technique may be used to quantitatively determine depth profiles of lattice disorder. Our optoacoustic measurements are shown to be 2-3 orders of magnitude more sensitive in defect concentration than channeling techniques, and establish a quantitative dependence between the change in optical response and defect concentration between 10^14-10^15 defects/cm^2. Further, we demonstrate the entire range over which the coherent acoustic phonon technique is applicable in defect studies, and show results ranging from no noticeable change in optical response to complete damping of the phonon wave. We also discuss the electronic nature of the CAP response, which can provide insight into the interplay between lattice disorder and electronic structure.

9:40am TF1-ThM6 On the Phase Formation of Reactively Sputtered ZrO2 Thin Films. R. Snyders, R. B. Rogers, R. B. Wood, Vanderbilt University

Boron doped silicon samples (0.01 Ω-cm - 0.02 Ω-cm) were irradiated at dose rates ranging from 5.8 krads(SiO2)/min to 31.5 krads(SiO2)/min using a 10 keV x-ray source (ARACOR Model 4100). The samples were cleaned in dilute HF acid to remove the initial native oxide and obtain a clean surface prior to irradiation. The resulting silicon oxide thickness was estimated using ellipsometric ellipsometry (J.A. Woodam-M-2000). Different models for the optical properties of the surface layer were evaluated. Oxide growth on the irradiated samples was compared to oxide growth on control samples, which were placed in ambient air at room temperature. The ellipsometry analyses suggest that the oxide growth on the irradiated samples was greater than that on the control samples. Initial XPS analysis showed that the oxide layers on the irradiated and control samples were chemically different, suggesting that differences observed in the ellipsometry analyses may be due, in part, to differences in the optical properties of the oxide layers and not purely due to a change in thickness.

We will present the results of the oxidation study in addition to XPS and AES characterization of the resulting oxide films. The impact of heating the silicon substrate up to 150°C during x-ray irradiation will also be discussed. A mechanism to explain the experimental observations is proposed based on ozone concentration measurements performed during irradiation. The possible generation of atomic oxygen, due to dissociation of molecular oxygen, by the high energy x-rays is suggested as a key factor in the observed x-ray irradiation induced silicon oxidation.
increase in contact resistance (from 30 mΩ to 200 mΩ) to the microstructure is reflected in the properties of the films with an abrupt transition embedded in an amorphous matrix phase. The size of the carbide grains measured using a four wire set-up with an Au-coated probe pressed against the films using solid-state magic angle spinning nuclear magnetic resonance (MAS-NMR) and Fourier transform infrared (FTIR) spectroscopies, backed by density functional theory (DFT) molecular structure calculations. The comparison of experimentally observed spectral features with theoretical predictions for model molecular compounds provides valuable insight into the different local chemical environments and intermediate-range networks that make up the a-B,C:H, films. We demonstrate how applying these combined analyses provides an important stepping stone to understanding and optimizing the chemical, electrical, and mechanical properties of a-B,C:H films for next-generation device fabrication.

Our results show that films with a silicon content less than 20 at.% exhibit a nanocomposite (nc-NbC/a-SiC) structure with nanocrystalline NbC grains embedded in an amorphous matrix phase. The size of the carbide grains decrease with the carbon content increase. A transition to a completely amorphous structure occurs with a Si content above 20 at.%. This transition in microstructure is reflected in the properties of the films with an abrupt increase in contact resistance (from 30 mΩ to 200 mΩ at 1 N) and a change in the mechanical behavior of the thin films. The XPS spectra show a change of chemical bonding from mainly C-Nb to a significant amount C-Si bonds as the Si and C content increase in the films. This results in harder films and an increase in the electrical resistivity. The amorphous films can be described as a metal carbide-based glass and the formation of this type of structures show that, with optimised processing, CVD Mn-based barriers are promising candidates as Cu barriers in advanced interconnects.

In conclusion, voltage ramp and TDDB measurements on planar capacitors structures show that, with optimised processing, CVD Mn-based barriers are promising candidates as Cu barriers in advanced interconnects.


Thin Film Division

Modeling and Analysis of Thin Films
Moderator: P.D. Rack, University of Tennessee Knoxville

8:00am TF2-ThM1 Aluminum Molecular Model for DSMC Simulations of Thin Film Deposition, A. Venkattraman, A. Alexzenko, Purdue University

The direct simulation Monte Carlo (DSMC) technique has been shown to be able to predict various properties of thin films grown using vacuum deposition methods such as CVD, PVD, EBPVD. Such simulations can also provide information about the energy distribution and orientation of vapor molecules striking the substrate which are critical inputs to the prediction of the grain size, residual stress and other properties of the deposited films. One of the most important inputs to a DSMC simulation is the molecular model that determines the interaction between the simulated particles. The variable hard sphere (VHS) model that is widely used due to its combination of simplicity and accuracy is typically determined by fitting to viscosity data obtained from experiments. In the absence of direct measurements of transport coefficients for metal vapors such as Aluminum, one needs to resort to other techniques to determine a set of accurate molecular model parameters. In this work, we compare DSMC simulations with Aluminum thin film deposition experiments to determine the VHS model parameters. The growth rate of the thin films at the substrate location depends strongly on the transport properties of the metal vapor – viscosity being one of the most important – and hence can be compared with the DSMC simulations to determine a suitable molecular model. In a similar analysis for copper published earlier, we used experimental data available in literature while the experiments for this study are performed in the electron-beam evaporator in the Birck Nanotechnology Center at Purdue University.


This paper describes three-dimensional flow/thermal/chemistry modeling efforts to study the silicon dioxide deposition using in situ gas phase reaction mechanism (ISSG). This 3D model incorporates 27 step gas phase reaction mechanism which is responsible for ISSG chemistry. The complex 3D geometry, supersonic flow, detailed chemistry pose major challenges to the model convergence and results in unrealistic results due to H2 and O2 exothermic

Acknowledgement: This work was supported in part by the DTRA Basic Research Program (Grant No. HDTRA1-10-0041).

10:00am TF1-ThM11 Characterization of Amorphous and Nanocomposite Nb-Si-C Thin Films Deposited by dc-Magnetron Sputtering, N. Nedfors, Uppsala University, Sweden, O. Tengstrand, Linköping University, Sweden, A. Flink, Impact Coatings AB, Sweden, A.M. Andersson, ABB AB, Corporate Research, Sweden, P. Ekland, L. Hultman, Linköping University, Sweden, U. Jansson, Uppsala University, Sweden

Thin films of Me-Si-C (Me = early transition metal) have interesting multifunctional properties, see, e.g., [1]. We have in the present study investigated thin films in the Nb-Si-C system deposited by dc-magnetron sputtering using elemental targets. The microstructure and composition of the films have been characterized by x-ray diffraction, x-ray photoelectron spectroscopy (XPS) and transmission electron microscopy. Nanoindentation measurements provide valuable insight into the different local chemical environments and intermediate-range networks that make up the a-B,C:H, films. We demonstrate how applying these combined analyses provides an important stepping stone to understanding and optimizing the chemical, electrical, and mechanical properties of a-B,C:H films for next-generation device fabrication.


Diffusion barrier layers with a uniform thickness and good step coverage in narrow lines are needed to enable the continuing scaling of Cu interconnects. Since physical vapor deposition (PVD) processes have limitations in respect of conformality, atomic layer deposition (ALD) and chemical vapor deposition (CVD) might become the preferred alternatives. Among others, Mn-based CVD barriers have been proposed [1], but up to now, Cu barrier properties for such films have not been proven electrically.

In this study, MnOx layers were deposited by CVD, on top of an O3/TEOS SiO2 layer. Deposition was done at two different temperatures: 200 °C and 350 °C. The effect of a post-plating anneal of one hour at 300 °C or 430 °C was studied. A test structure based on planar capacitors was used, in which first wide areas were patterned, followed by the deposition of the oxide and CVD MnOx layers, metallization, and CMP, respectively [2]. After passivation, voltage ramp (at 25 °C and 100 °C) and TDDB measurements (at 100 °C) were conducted. The electrical data were compared with those from a known good TaN/Ta-barrier reference system. Voltage ramp measurements at 25 °C show a similar behaviour for all Mn-based films under study, except for the one deposited at 350 °C, with a post-plating anneal at 430 °C. For the latter a leakage current of about 1 order of magnitude lower is found, comparable to the leakage current of our TaN/Ta reference. For the film deposited at 200 °C, and annealed at 430 °C, voltage ramp measurements only showed shorts.

TDDB lifetimes were in all cases higher than for a reference without barrier, but lower than for a TaN/Ta reference. For the Mn-based films deposited at 350 °C, a post-plating anneal at 430 °C clearly improved the reliability properties. In this case, the extrapolated lifetime at user conditions (using the E-model), is above 10 years. A γ value of -3.4 cm/MV was found. However, for the films deposited at 200 °C, an anneal at 300 °C already degraded the reliability properties.

In conclusion, voltage ramp and TDDB measurements on planar capacitors structures show that, with optimised processing, CVD Mn-based barriers are promising candidates as Cu barriers in advanced interconnects.

reaction. Complex 3D geometry causes CFD model mesh size to become more than a million cells. The computational times to include all chemical reactions on a complex flow problem are exorbitant. Thus a simple 2D RTP chamber model was built to examine the validity of gas phase reaction mechanism by comparing in-house simulation results with Professor Robert J. Kee, Colorado School of Mines [1] and to set the solver parameters for stable solution in 3D. After a working 2D model, simple representative 3D model was built with very good quality structured mesh and further geometric complexity was added in steps. The oxygen radical distribution predicted from the model matched very well with the oxide growth uniformity over a wide range of chamber pressure, gas flow rate, hydrogen fraction, and gas distributor geometry. At 5 torr, temperature distribution is dominated by wafer temperature as gas phase reaction is weak. At higher pressure exothermic gas phase reaction causes higher temperature above the wafer. There is weak gas phase reaction at 5 torr, resulting in low O atom number density. The O radicals diffuse in the chamber due to higher velocity at 5 torr where they recombine. At higher pressure flame ignites right at the edge of the wafer due to higher residence time resulting in very high O atom number density at the edge. This well calibrated simulation model was used in understanding and expanding process space by optimizing on several hardware and process variables using virtual prototyping before building the hardware.

8:40am TF2-ThM3 Composition and Finite Size Effects in Thin Magnetic Films for Data Storage Applications: Magnetic and Transport Properties, O.N. Mryasov, University of Alabama INVITED

Scaling of magneto-resistance (MR) and resistance area product (RA) with thickness is one of a critical materials specific properties of hard disk drive sensors. We consider fundamental aspects of MR-RA scaling with two dependence of RA are consistent with experimental [7]. We also present results of direct spin dependent electronic transport simulations for two metal systems such as TiN. The currently limited number of TiN MEAM simulations of TiN thin films growth, difficult case studies presented, which successfully reproduce experimentally documented crucial processes in the initial stages of TiN nucleation, known to dramatically affect growth modes, and ultimately, properties of thin films. The implications of these results, and perspectives for large scale simulations of this extremely important magnetic material system, are discussed.

9:40am TF2-ThM6 Hard, yet Tough, Transition Metal Nitride Thin Films by Alloying and Valence Electron Concentration Tuning, D.G. Sangiovanni, V. Chirita, L. Hultman, Linkoping University, Sweden

Improved toughness in hard and superhard thin films is a primary requirement for present day ceramic hard coatings, known to be prone to brittle failure during in-use conditions, in modern applications. Based on the successful approach and results obtained for TiN- and VN-based ternary thin films [1,2], we expand our Density Functional Theory (DFT) investigations to TiAlN-based quaternary thin films. (TiAl)1-xWxN thin films in the B1 structure, with 0.06 ≤ x ≤ 0.75, are obtained by alloying with W = V, Nb, Ta, Mo and W, and results show significant ductility enhancement. Hence the toughness of TiAlN-based thin films. Importantly, these thin films are also predicted to be hard/superhard, with similar and/or increased hardness values, compared to TiAIN. For (TiAl)1-xWxN these results have experimentally been confirmed recently [3]. As previously demonstrated [1], the ductility increase originates in the enhanced occupancy of d-t2g metallic states, induced by the valence electrons of substitutional elements (V, Nb, Ta, Mo, W). This effect is more pronounced with increasing valence electron concentration (VEC), and, upon shearing, leads to the formation of a layered electronic structure, consisting of alternating layers of high and low charge density in the metallic sublattice. This, in turn, allows a selective response to tetragonal and trigonal deformation: if compressive/tensile stresses are applied, the sublattice responds in a "hard" manner by resisting deformation, while upon the application of shear stresses, the layered electronic arrangement is formed, bonding is changed accordingly, and the structure responds in a "ductile/tough" manner as dislocation glide along the [110]<110> slip system becomes energetically favored [2].


10:40am TF2-ThM9 Using Crystallographic Space Group-Subgroup Relations to Analyze Phase Selection and Transition in HfO2 and Hf-based Ternary Oxide Films, C.R. Aita, University of Wisconsin-Milwaukee

HfO2 and Hf-based ternary oxides are important candidates for ultrathin high permittivity dielectric applications. However, technological insights of their use in actual devices far outstrip our knowledge of the fundamental science that governs phase selection and transition in these materials. The latter is important for predicting both initial device performance and long term stability. One big issue is that pure HfO2 readily forms nanocrystallites in thick films. These crystallites exhibit finite size effects on two different length scales: (1) Two metastable phases initially form in crystallites <7nm in size [1] and transform to monoclinic (m) HfO2, the standard state, as crystallites grow. (2) Upon transformation from the metastable, m-HfO2 nanocrystallites whose size is <= 11 nm exhibit a lattice expansion concurrent with surface dipole repulsion [2]. A second issue involves the stability of Hf-based ternaries that are either intentionally grown or inadvertently form as a result of cation mixing during thermal processing or heattreatment device use. These questions are being addressed from an experimental viewpoint through controlled isochronal and isothermal annealing studies. In this paper, we use crystallographic space group-subgroup analysis to examine phase selection and transition in three sputter deposited nanolaminates, HfO2-Al2O3, HfO2-TiO2, and HfO2-ZrO2. The goal is demonstrate how this tool connects phase transitions between seemingly unrelated structures by symmetry considerations. We show that several important transitions observed in these materials are 2nd order and can be described by a simple relation between a parent group of higher symmetry and a daughter group of lower symmetry. Using the suite of programs in the Bilbao Crystallographic Server [3], first, conjugacy classes associated with
the parent and daughter transition are identified, and then using the operations within each class, the general atom positions of the parent are decomposed into cosets of symmetry elements expressed the daughter's basis. Symmetry elements that are "lost" in the decomposition are used to identify a twin domain structure in the daughter resulting from the transition. Using these formalisms, we discuss metastable phase—mHO₂ transition in pure HHO₂, the robustness of an entropy-stabilized HFAl-oxide phase, and the initiation HFO₂ demixing.

Support from UWM Foundation Catalyst Grant / Rockwell Automation Charitable Trust.


11:00am TF2-ThM10 Experimental and Theoretical Investigations Using SiO₂ Nanotemplates to Relieve Stress Caused by Thermal Expansion Coefficient Mismatch in Epitaxial Germanium Grown on Silicon, S. Ghosh, D. Leonhardt, S.M. Han, University of New Mexico

High-quality Ge-on-Si (GoS) heterostructures are pursued for many applications, including near-infrared photodetectors, high-mobility devices with Si-based integrated circuits, and virtual substrates for III-V multijunction solar cells. Growing low-dislocation-density GoS and subsequently integrating III-V layers present two significant engineering challenges: lattice mismatch and thermal expansion coefficient mismatch. The materials engineering solutions to circumvent the lattice mismatch include metamorphic growth, graded buffer layers, selective epitaxial overgrowth, aspect ratio trapping (ART), and a variety of defect filtering strategies. The ART technique, in particular, utilizes high-aspect-ratio holes or trenches etched through dielectric films to trap dislocations, greatly reducing the dislocation density. However, one shortcoming of ART is that it has been demonstrated to be effective only for small holes or narrow strips with dimensions less than 1 μm. In this study, we demonstrate that a combination of ART with selective epitaxial growth can produce large areas of high-quality GoS. We focus on the use of SiO₂-based templates with nanoscale windows placed on GoS to relieve the thermal stress. We observe that voids formed around the top and sidewalls of SiO₂ template deposited by chemical vapor deposition, further relieving the thermal stress. The same templates also filter threading dislocations propagating from the underlying Ge-Si interface. The Ge layer grown and coalesced over the template is analyzed by transmission electron microscopy and etch pit density measurements. When the template is used, the threading dislocation density near the Ge film surface is approximately <10³ cm⁻², while the twin defect density is approximately <5×10¹⁰ cm⁻². Finite element modeling based on a commercial software package COMSOL is used to calculate the thermal stress occurring in the epitaxial Ge due to differences in thermal expansion coefficients among Ge, Si, and SiO₂. The simulation results, comparing Ge grown on Si with and without SiO₂ templates, show that the nanoscale templates can effectively reduce the thermal stress. The resulting stress results obtained using the simulation model corroborate the experimental observations. In summary, the simulation results suggest that the SiO₂ nanotemplates can reduce the stress caused by the thermal expansion coefficient mismatch, while simultaneously reducing the lattice-mismatch-induced dislocations in Ge grown on Si.


We demonstrate that surfactant-assisted epitaxy is a useful method for stabilizing the growth of [111] CaO films on (0001) GaN. Surface free energies, calculated by ab initio thermodynamics, for configurations of CaO surfaces with varying surfactant coverage will be presented. These results explained the recently reported experimental observation that incorporating water vapor during CaO deposition produces a hydroxylated surface. Hydroxylisation changes the preferred habit of CaO from (001) to (111), which enables layer-by-layer growth of (111) CaO on (0001) GaN. Together with experiment, these results demonstrate a new approach, applicable to numerous materials systems, where chemical boundary conditions are engineered to regulate the growth mode. Unique opportunities to integrate highly heterogeneous materials of dissimilar structure and symmetry are consequently available.

Thursday Morning, November 3, 2011  212
8:20am TR+AS+SS-ThM2 Surface Alterations Effects on Ice Adhesion Strength. C. Ellis-Terrell, M. Miller, Southwest Research Institute, M. Zou, University of Arkansas at Fayetteville, P. Wei, Southwest Research Institute, S. Beckford, University of Arkansas at Fayetteville, G. Hatton, Shell Global Solutions, Inc.

Ice adhesion is a serious problem in areas such as the oil, gas, and automotive industry, telecommunications and power line transmission. There is a significant amount of research directed towards designing a coating to reduce ice accumulation. This study focuses on measuring the effects of surface roughness and surface energy on ice adhesion strength. Surface texturing ranged from high to low surface roughness. A sandblasting technique was applied to the aluminum surface creating a high surface roughness. Surface energy changes were created by depositing a silicon doped hydrocarbon film, using plasma enhanced vapor deposition. A custom built apparatus was employed to specifically measure the adhesion force of an ice droplet. The results illustrate that the smoother as-received surfaces have lower ice adhesion strength than the rougher sandblasted surfaces.


Oxides such as SiO2, Al2O3 and HfO2 are typically used together with Si in many high-performance electronic devices, including metal-oxide-semiconductor (MOS) devices/junctions and micro- and nano-electromechanical systems (MEMS/NEMS). The lack of precise control over mechanical properties can lead to the degradation of these materials. It is therefore critical to understand the nanometer-scale mechanical properties of materials or complex systems being considered for use in electronic devices. Nanoindentation and nanoscratching are important methods for investigating the mechanical behavior of small volumes of materials, such as thin film systems. Here, classical molecular dynamics simulations are used to examine the responses to nanoindentation and nanoscratching of thin films of SiO2 and HfO2 on silicon substrates. The goal is to determine the influence of thin film types and the structure of thin film and substrate interface on the responses. Because these systems consist of heterogeneous interface with significant changes in bonding as one crosses from one side of the interface to the other, the empirical charge optimized many-body (COMB) potential as implemented in large-scale atomic/molecular massively parallel simulator (LAMMPS) program is used to model the structural evolution, mechanical response and charge transfer in these systems in response to a nanometer-scale spherical indenter. Aspects of the SiO2/Si and HfO2/Si interfaces during nanoindentation and nanoscratching, including the mechanisms by which fracture and plasticity occurs, will also be addressed. We gratefully acknowledge the support of the National Science Foundation through grant numbers DMR-0426870 and DMR-1005779.

9:00am TR+AS+SS-ThM4 Accelerated Molecular Dynamics Simulations of Nanoscale Friction. W.K. Kim, University of Minnesota, M.L. Falk, Johns Hopkins University

Accelerated molecular dynamics simulations are implemented to model the sliding process of atomic force microscope experiments and to lower the sliding speeds below those in a conventional MD simulation. In this study the hyperdynamics method, originally devised to extend MD time scales for non-driven systems, is applied to the frictional sliding system. This technique is combined with a parallel algorithm that simultaneously simulates the system over a range of slider positions so that the overall acceleration rate is approximately the number of processors multiplied by the boost factor from the hyperdynamics method. The new methodologies are tested using two-dimensional and three-dimensional Lennard-Jones AFM models. The methodology is then applied to simulated sliding between an oxidized silicon tip and surface achieving a range of six decades of velocity and reproducing the experimentally observed velocity dependence of the friction force. In doing so we learn something new about this system and about friction between amorphous surfaces in general. Unlike in the crystalline case, as increasing force is applied to the amorphous tip intermediate states arise. These intermediate states serve as critical transition pathways. The emergence of such states leads to the amorphous tip intermediate states. These intermediate states serve as successful descriptors of both the experimental and the simulated data.


The behavior of nanoscale contacts is complex and often cannot be understood through continuum mechanics alone. Here, parallel molecular dynamics (MD) simulations using the AIREBO potential for hydrocarbons to model indention and friction, are used to investigate nanoscale contacts of carbon-based materials, such as diamond, DLC, and ultrananocrystalline diamond (UNCD). Specifically, the contact of carbon-based AFM probes is simulated to understand the effects of experimental parameters, including tip geometry and material selection, on the adhesion between the tip and sample. Results from the MD simulations will be compared to and discussed within the context of the complementary atomic force microscope experiments and finite element simulations. The tribological response of carbon-based materials is very sensitive to environmental conditions. For example, the presence of water has been shown to negatively impact the friction performance of hydrogenated DLCs but to improve the performance of nanocrystalline and ultrananocrystalline diamond. We have been working to develop a potential energy function that is capable of modeling carbon-based materials in the presence of water. This talk will also outline our current efforts at potential development.

10:40am TR+AS+SS-ThM9 Modeling the Pressure Dependence of Shear Strength in Sliding, Boundary-Layer Friction. M. Garvey, M. Weinert, W.T. Tysoe, University of Wisconsin-Milwaukee

The pressure dependence of the shear strength of model alkali halide lubricant systems has been investigated at the density functional theory level. This is compared to the experimental dependence given by $S = S_0 + P^m$ where $S$ is the shear strength and $m$ is the mness of pressure dependence. Theoretical shear strengths were calculated and shear is found to occur between the film and the sliding interface. The heights of the potentials were calculated as a function of compression, allowing the lateral force to be calculated as a function of pressure. The calculated values of $S_0$ and $m$ were in good agreement with experimental data.


Inorganic fullerene (IF)-like MoS2 nanoparticles have been shown to be good lubricating and anti-wear additives when dispersed in a base oil. This improved tribological performance appears to be a result of the size and structure of the nanoparticles along with the test conditions. Possible lubrication mechanisms include pure rolling to sliding to the exfoliation of lamellar MoS2 sheets inside the contact. In situ transmission electron microscopy (TEM) experiments have been used to manipulate individual MoS2 nanoparticles and investigate their responses to compression and friction in both different contact configurations. However, the tribological response of the MoS2 nanoparticles make distinguishing the properties which affect the lubrication mechanism exceedingly difficult; thus, a computational approach is used to more fully understand the most important mechanisms. Therefore, classical molecular dynamics (MD) simulations of individual nested MoS2 nanoparticles are performed where they are subjected to compression and shear forces between sulfur-terminated molybdenum surfaces. Two specific nanoparticle configurations are considered, with both structures containing three layers. The first configuration is a curved, ellipsoidal MoS2 nanoparticle structure with a major and minor diameter of approximately 8.9 and 6.6nm, respectively. The second nanoparticle configuration is an octahedron with grain boundaries that are approximately 6.2 nm in length. MD simulations of these structures indicate the role of curved and faceted morphologies as well as grain boundaries on the rolling/sliding behavior and nanoscale exfoliation of the particles. The results are used to interpret the experimental TEM findings and predict the dominant mechanisms associated with enhanced lubrication through the addition of these particles to base oils. This work is supported by the Office of Naval Research.


The primary problem with lateral force microscopy (LFM) has been the difficulty in calibrating the cantilever and tip in order to obtain quantitative friction data. Two recent review articles and several research articles have expressed this difficulty and the need for a simple, universally-accepted method [1,2]. The available procedures have numerous limitations. Some require specialized samples or setups. Others are difficult to perform. A
number are indirect, or only suitable for certain cantilevers. Several risk
damage to the tip or sample, or both, and might require the geometry of the
cantilever, which can be hard to measure. We present a procedure that
alleviates these problems [3]. The linear relationship between the detected
voltage and lateral force is exploited to obtain the slope (calibration factor)
and intercept that convert voltage to lateral force. The method is
independent of sample shape, probe shape, and scan parameters (load force,
gain, and scan rate). The accuracy was investigated on an order-of-
magnitude level and was within 50% of torsional spring constants obtained
from geometry, and the precision was under 10%. Small scan areas were
also found to produce accurate calibration factors and could help to limit
tip-sample wear. Quantification of nano-Newton friction forces might now
become routine.

73-104.
M. Stevens, D. Liu, and N.A. Burnham, “Shape-Independent Lateral Force
Calibration,” submitted April 2011.

11:40am TR+AS+SS-ThM12 Atomic Stick-Slip Friction Studied by
Optimally-Matched Accelerated MD Simulations and AFM
Experiments, Y. Dong, Purdue University, Q. Li, R.W. Carpick, University
of Pennsylvania, A. Martini, Purdue University
Atomic-scale stick-slip friction of platinum on gold (111) surface is
quantitatively studied both experimentally and through optimally-matched
accelerated molecular dynamics (MD). In order to make a direct
comparison between simulation and experiment, many other factors are
matched as closely as possible, such as misalignment, size effect of the tip,
cantilever compliance, normal load and so on. The Parallel Replica
Dynamic Method (ParRep) is used to accelerate the simulation so scan
velocities can be decreased to scales approaching those used in atomic force
microscope experiments. A logarithm dependence of friction on scanning
velocity is observed both in Atomic Force Microscope (AFM) and MD
reveals that at low speed the atomic friction lies in thermal activation
regime. A further comparison shows that AFM and MD provide consistent
energetics, which supports that MD can be used to interpret AFM results;
but attempt frequencies differ by orders of magnitude, which is attributed to
the inertia discrepancy.
2:00pm AC+TF-THA1 Plutonium Sorption and Reactivity at the Solid/Water Interface. M. Schmidt, P.A. Fenter, S.S. Lee, R.E. Wilson, L. Soderholm, Argonne National Laboratory

Reliable long-term predictions about the safety of a potential nuclear waste repository must be based on a sound, molecular-level comprehension of the geochemical behavior of the radionuclides. We apply in situ crystal truncation rod (CTR) measurements and resonant-anomalous X-ray reflectivity (RAXR) in combination with alpha-spectrometry to elucidate the sorption behavior of tetravalent actinides on muscovite under varying solution conditions.

Key retention mechanisms, particularly in clay formations, are surface mediated processes. In order to be able to understand these processes analytical techniques that allow selectively probing the mineral/water interface and elucidating processes at the interface under in situ conditions are required. X-ray reflectivity techniques (CTR, RAXR) have proven to be valuable tools for geochemical studies concerning the sorption behavior of metal ions [1]. More recently they have also been applied to study the sorption behavior of actinides [2].

From CTR measurements the complete adsorption structure, consisting of adsorbed water and the ions adsorbed as inner sphere, outer sphere or extended outer sphere complex can be derived. RAXR extends this approach by providing elemental specificity to the CTR measurements, thus characterizing the contribution of a particular element to the structure.

Using a purpose-built sample cell for radiological experiments, X-ray reflectivity data was collected in situ from muscovite (KA(1/2)(OH,F)(1/2)(SiAl)O(3)) in contact with the actinide-bearing solutions varying in composition (ionic strength, actinide concentration and speciation, background electrolyte). The data is complemented by precise quantitative analysis by means of alpha-counting experiments. The results clearly show the strong influence of the actinides’ aqueous chemistry on their sorption behavior. It will be shown that a full description of the sorption behavior requires parameters such as the hydration enthalpy, complexion constants, hydrolysis constants, and polymer formation constants. At low actinide concentrations the formation of hydrated surface complexes is observed in good agreement with the large hydration enthalpies of the highly charged ions. At large excess of background electrolyte sorption occurs under preservation of the complexion by the anion. When a threshold metal ion concentration is exceeded sorption of polymers is observed which grow up to 150Å in size, while covering only small fractions of the surface.

1. Fenter, P., Reviews in Mineralogy and Geochemistry 2002, 49, 149-220.

2:40pm AC+TF-THA3 Crystal Chemistry of Thorium Oxy Compounds Containing Tetrahedral Oxyanions. A.J. Albrecht, P.C. Burns, University of Notre Dame

Here we examine the crystal chemistry and structural topologies of Th compounds containing various cations that are coordinated by oxygen atoms in tetrahedral arrangements. We define the structural unit to be the part of the structure that consists of those polyhedra that contain higher valence cations, with an emphasis on their connectivity. In most cases, structures also contain interstitial units that balance the charge of the structural unit. In the compounds under study, which include both new structures and those from the literature, we find structural units ranging from clusters and chains, through sheets, to extended frameworks. The structural units in thorium compounds are of particular interest because they provide insights into the possible structures of compounds consisting of tetravalent transuranium cations, especially Np and Pu.


We report on the solution based synthesis of epitaxial thin films of neptunium oxide and plutonium oxide. Actinides represent a tremendous challenge to first principle calculations of orbital energies due to the complicating features that arise from f orbital interactions. Theoretical development from first principle calculations relies on predictions of continuous materials with no boundaries. To test these theories requires experimental results using high quality single crystals. For many materials this can be a straight forward process. For actinide oxides the experimental work is complicated by the radioactivity that makes CVD processes challenging from a safety perspective and by the recalcitrant nature of the oxides. The challenge of modeling actinide oxides is best represented in Mott insulators such as UO2. Theoretical calculations that worked with transition metal oxides predict UO2 to be a metal as opposed to a metal with a 2.3 eV band gap. Recent reports predict the same metallic behavior for NpO2. Many theoretical reports have highlighted the need for experimental work on single crystals. In the absence of single crystals they rely on work done on powders as in the case of PuO2. We report here on epitaxial films of PuO2 that are thin enough to measure the optical band gap directly. XRD has been used to determine the alignment of the PuO2 relative to the substrate and EXAFS confirms the stoichiometry.

3:40pm AC+TF-THA6 U(VI) Uranyl Cation-Cation Interactions in Framework Germanates. J.M. Morrison, P.C. Burns, University of Notre Dame

The crystal structure determinations of four highly complex U(VI) germanate framework compounds reveal three distinct U(VI) sites—two that are coordinated by five equatorial oxygen atoms each to form pentagonal bipyramids and one that is coordinated by six oxygen atoms to form a distorted octahedron without the uranyl ion [1]. This is uncommon as most inorganic U(VI) compounds contain the uranyl ion, (UO2)\(^{2+}\). Also present are cation-cation interactions which occur when an oxygen atom of the uranyl ion also acts as an equatorial oxygen atom in a neighboring U(VI) polyhedron. CCI are present in fewer than 2% of U(VI) compounds. Finally, a disordered system with the possibility of one-dimensional Ge\(_2\)O\(_5\)\(^{2-}\) chain has led us to suggest three structural models with regard to Ge coordination. The interesting structural and chemical complexities of these CCI-bearing compounds will be presented along with a discussion of the densities of CCI-bearing U(VI) compounds.


4:20pm AC+TF-THA8 The Crystal Chemistry of Uranyl Selenates and their Relations with Uranyl Sulfates. E.M. Wylie, P.C. Burns, University of Notre Dame

Uranyl sulfate minerals have been studied for several decades owing to their importance in understanding ore genesis, as well as the interaction of uranium mine and mill workings with the environment. In contrast, no uranyl selenate minerals have been described, although several uranyl selenates are known. We are exploring the crystal chemistry of synthetic uranyl selenates produced under mild hydrothermal techniques. We have obtained crystals of several compounds and characterized them with single-crystal X-ray diffraction. They contain a sheet of edge-sharing uranyl pentagonal bipyramids and selenate tetrahedra that is topologically identical to those found in the zippelite group of uranyl sulfate minerals. These synthetic materials provide further insights into layered uranyl phases, including the relationships between the configurations of the structural sheets and their corresponding interlayer complexes. These structures will be placed in the context of known synthetic and natural uranyl sulfate and selenate compounds.

4:40pm AC+TF-THA9 Radiation-Induced Degradation of Photoluminescence in YAG:Ce. S.G. Gollub, D.G. Walker, S.L. Weeden-Wright, Vanderbilt University

Thermographic phosphors are ceramic based materials whose photoluminescence is temperature dependent. We fabricated the phosphor YAG:Ce\(_{0.1}\) to determine the influence of the cerium ions and the cerium-carbonate environment on the photoluminescence spectra. In particular we investigated the effects of non-ionizing radiation on the photoluminescent spectra. No change to the spectrum was observed after exposure to 1 Mrad of x-ray radiation. Because x-rays are typically ionizing, we did not expect to see any significant degradation. When the material was bombarded with protons, which are known to cause displacement damage in many materials, a degradation was observed. Results for a damage cross section and stopping power were commensurate with predicted values using SRIM. Results are shown as a function of proton energy and dose.
2:00pm EL+AS+EM+MS+PS+TF+ThA1 Applications of Ellipsometry in Photovoltaics, D. Levi, National Renewable Energy Laboratory

With the growing possibility of anthropogenic-induced climate change there has come increasing concern over energy-related emissions of carbon dioxide into the atmosphere. The search for low or no-carbon energy sources has intensified. This has lead to a twenty first century gold rush into photovoltaics which will yield electricity from photovoltaics is still a long ways from economic competitiveness with fossil fuel-based electricity sources. The U.S. Department of Energy recently announced the Sunshot program, with the expressed goal of $1/Watt installed cost for utility scale PV plants by 2017. This aggressive goal will require radical advances in new and existing PV technologies.

Spectroscopic Ellipsometry for Photovoltaics, Metals and Oxide Thin Films

Moderator: M. Creatore, Eindhoven University of Technology, the Netherlands

Spectroscopic Ellipsometry Focus Topic Room: 209 - Session EL+AS+EM+MS+PS+TF+ThA

EL+AS+EM+MS+PS+TF+ThA3 Comparison between Ex Situ and Real Time Spectroscopic Ellipsometry Measurements of Structurally Graded Si:H Thin Films, N.J. Podraca, University of Toledo

Analysis of spectroscopic ellipsometry measurements of graded thin films remains challenging, although analysis procedures and software have improved over the past several decades. Practical use of these processes remains somewhat time consuming and is often not fully utilized by the casual user. In this work, ex situ ellipsometric spectra collected for static ex situ measurement at the end of the deposition, obtaining ε and structural parameters of the film becomes less precise. Specifically, sensitivity to the variation in the nanocrystallite fraction with thickness may be lost and inaccurate ε for the component materials may be obtained. This works seeks to compare the structural and optical properties of (a+nc)-Si:H obtained by RTSE and VIA with those from analysis of static ex situ spectra with models using different structures, parameterizations in ε, and spectral range restrictions. These comparisons will be used to verify appropriate structural and dielectric function models to more accurately analyze structurally graded thin films under different material and measurement circumstances.

Cell Characterization by Phase Modulated Spectroscopic Ellipsometry

3:40pm EL+AS+EM+MS+PS+TF+ThA4 Real-Time Spectroscopic Ellipsometry of Cu(In,Ga)Se2 Thin Film Deposition: Copper Transition in 3-Stage Co-Evaporation Process, D. Attygalle, University of Toledo, F. Romer, University of Toledo, J. Marsillac, Old Dominion University, R.W. Collins, University of Toledo

With record efficiencies above 20%, Cu(In,Ga)Se2 (CIGS) based solar cells have shown the greatest potential for success among the thin film photovoltaics technologies. Thermal co-evaporation of individual elements has proven to produce extremely high quality CIGS materials, provides a high level of flexibility, but also generates greater challenges in process optimization. The limitations of existing process monitoring capabilities, hence the challenge of correcting process fluctuations in real time, has led the industrial community toward more controllable CIGS deposition processes. Real time spectroscopic ellipsometry (RTSE) can be used successfully in the monitoring of complicated processes -- including CIGS film preparation by co-evaporation using precursor films of (Inx,Ga1-x)2Se3. Information extracted from RTSE includes the evolution of bulk layer and surface roughness layer thicknesses, the composition and phase, as well as the layer dielectric functions, all of which can assist in understanding the fabrication process and in optimizing solar cells. In this study, the focus is on the transitions of Cu-poor to Cu-rich CIGS and vice versa by observing the changes in the (ψ, Δ) spectra obtained by RTSE. The commonly used monitoring method, which involves observing the changes in emissivity of the film, strongly depends on the apparatus design, the substrates, and the bulk layer thickness. Since a CIGS film is prepared by exposing a precursor film of (Inx,Ga1-x)2Se3 to Cu and Se fluxes, thereby becoming Cu-rich, a semi-liquid Cu2-,Se phase is believed to form on top of a bulk layer consisting of mixed phases of CuIn(x),Ga1-x)2Se3 and Cu2-,Se [1]. A multilayer optical model, with appropriate effective medium approximation layers to represent this structure, has shown good agreement with the observed (ψ, Δ) spectra. Since RTSE is highly sensitive to monolayer-level changes in the top-most layer, RTSE gives superior sensitivity in Cu-rich to Cu-poor end point detection, which occurs when the top Cu2-,Se phase drops below detectable limits. Furthermore this method is less affected by the substrate and bulk layer thickness. Although careful analysis of RTSE can give a wealth of information about CIGS material properties and their evolution, this type of end point detection can be successful simply by monitoring the real-time changes in the (ψ, Δ) spectra.

range 190 – 2100 nm (0.6-6.5 eV). Two different analysis protocols were used to model the P3HT:PCBM blend structure. In the first protocol, effective medium approximation was used to represent the optical constants of the film, whereas in the second one the blend was treated as one single homogenous material. The approach renders investigation of final morphology and composition.

4:00pm EL+AS+EM+MS+PS+TF-ThA7 In Situ Spectroscopic Ellipsometry during Atomic Layer Deposition of Pt, Pd and Ru. N. Leik, J.W. Weber, M.J. Weber, A.J.M. Mackus, H.C.M. Knoops, W.W.M. Kessels, Eindhoven University of Technology, Netherlands

The precise thickness control of atomic layer deposition (ALD) and its conformal growth make ALD the method of choice for nanometer thin film deposition. Platinum-group metals such as Pt, Pd and Ru have many applications in the areas of nanoelectronics and catalysis and recently there has been considerable interest to deposit films of these materials by ALD. Spectroscopic ellipsometry (SE) is a non-destructive technique that can be used in situ during ALD to precisely monitor the thickness of the films. SE also provides information on the optical and electrical properties of the films which is very relevant for their applications. Choi et al. [1] previously investigated the dielectric functions of Pt-group metal films with a thickness of ~400 nm as prepared by physical vapor deposition. For the aforementioned applications, however, the films are required to be much thinner, which leads to differences in film morphology as well as to dielectric functions that can be different from those of bulk films. In the spectroscopic ellipsometry work to be presented in this contribution we have therefore focused on films with thicknesses from 5 nm to 35 nm. In situ data was obtained during ALD in the photon energy range of 0.7 – 6.5 eV. Using a Kramers-Kronig consistent B-spline model to account for the thickness-dependent dielectric functions, we were able to obtain accurate ALD growth-per-cycle values for Ru, Pt and Pd (1.00 ± 0.06 Å, 0.47 ± 0.04 Å, 0.14 ± 0.02 Å). Furthermore, the contributions from free-carriers (Drude term) and interband absorptions (Lorentz-oscillator contributions) were investigated by combining the SE data with FT-IR reflectance data such that the photon energy range of 0.04 eV – 6.5 eV was covered. In this range it was possible to represent each film with a unique Drude-Lorentz model although some ambiguities about the Lorentz oscillator contributions remained in the case of Ru. It will be shown that the extracted thicknesses and electrical resistivities from this model are in line with data obtained from X-ray reflectometry and four-point probe measurements (for example Ru: $\rho_{SE}$ ~23 $\mu\Omega$cm and $\rho_{PP}$ ~16 $\mu\Omega$cm). Furthermore, in the case of Ru also the influence of the film roughness will be addressed.


The fabrication of three-dimensional metal nanostructures with tailored geometry is one of the central challenges of nanotechnology because geometrical and material parameters are responsible for the optical, electronic, and mechanical properties of nanostructured thin films. Engineered artificial sculptured thin films (STFs) with designed anisotropies are potential candidates for applications in various fields such as optics, magneto-optics, as well as chemical and biological sensing and detection. However, in order to utilize metallic nanostructures for novel applications their size-, structure-, and material-driven physical properties have to be understood and quantified.

We utilize glancing angle electron-beam deposition, which exploits physical vapor deposition for the simultaneous deposition of metallic thin films at a tilted angle (~85°) to the substrate. In this way, films with thicknesses of micrometers can be deposited with a uniform thickness of ~400 nm. The thin films are then coated with a metal thin film by means of atomic layer deposition (ALD) to form metal STFs. By controlling the ALD process, the thickness of the deposited metal layer can be controlled. This allows the fabrication of nanostructured thin films with a thickness of ~400 nm. The thin films are then coated with a metal thin film by means of atomic layer deposition (ALD) to form metal STFs. By controlling the ALD process, the thickness of the deposited metal layer can be controlled. This allows the fabrication of nanostructured thin films with a thickness of ~400 nm. The thin films are then coated with a metal thin film by means of atomic layer deposition (ALD) to form metal STFs. By controlling the ALD process, the thickness of the deposited metal layer can be controlled. This allows the fabrication of nanostructured thin films with a thickness of ~400 nm. The thin films are then coated with a metal thin film by means of atomic layer deposition (ALD) to form metal STFs. By controlling the ALD process, the thickness of the deposited metal layer can be controlled. This allows the fabrication of nanostructured thin films with a thickness of ~400 nm.

4:40pm EL+AS+EM+MS+PS+TF-ThA9 Ellipsometric Characterisation of Porous Aluminium Oxide Supports. F. Ogiergo, N.E. Benes, H. Wormeester, IMEC, Leuven, Belgium

Porous aluminium oxide is widely used as a support material for thin film inorganic micro- and mesoporous membranes. Such membranes are used in energy-efficient gas separation, pervaporation and nanofiltration processes. Ellipsometry can be used to determine material properties of the thin membranes, as well as to determine loading and provide a detailed interpretation of the ellipsometry data requires a detailed knowledge of the porous aluminium oxide support. This support is made of aluminium oxide particles that are sintered together. In between the particles voids are present that amount to ~38% porosity. We have studied the influence of the size of the voids on the optical response of the support material. For this study, voids with a diameter of around 60, 80 and 160 nm were used. We noted a strong dependence of the normal incidence specular reflection on the void diameters and a subsequent increase in off specular reflection. In ellipsometry, only a limited depolarization of the specular reflected light was noted in the wavelength range between 300 and 1750 nm. The angle dependent ellipsometry measurements showed that the optical properties of these supports can not be obtained from a direct inversion. The reason for this is that the interface the more the voids are cut, which leads to a distribution of openings at the surface, i.e., a substrate with a very rough surface. This roughness was modelled with a graded porosity changing from ~38% in the bulk to ~75% at the outer surface. This measured variation in porosity is very similar to the cumulative height distribution of the surface layer obtained from AFM. The validity of this graded porosity model was verified by the analysis of a sample with a thin polysulfone (PSU) layer deposited on the support. The PSU layer partly fills the open pores at the surface. This results in an interface with a graded variation in aluminium oxide, void and PSU.

The proper treatment of the surface layer also provides the optical properties of the porous aluminium oxide bulk material itself. These optical properties can in a limited wavelength range be modelled with Bruggeman’s effective medium approximation. As a consequence of the size of the inclusions, their diameter is no longer negligible with respect to the wavelength of light in the UV part of the spectrum. For the material with the largest pore size, also a large part of the visible range has to be excluded. A more elaborate approach than the standard effective medium approach has to be used in this case.

common practice to encapsulate the thin film with a more environmentally stable material. In this study, the material chosen was SiO₂ grown in the same deposition chamber, pre-atmospheric exposure, via rf sputtering.

Vanadium dioxide (VO₂) is a strongly-correlated electron material with a well-known semiconductor-to-metal transition (SMT) that can be induced thermally (Tc ≈ 68°C), optically, or electrically. Recently, VO₂ films have attracted attention as a component in active metamaterials, especially in conjunction with metal nanostructures. Since these structures are highly sensitive to the dielectric properties of the embedding material, the SMT of VO₂ can be used to tune the optical response of the structure. Accurately modeling the behavior of these structures requires detailed knowledge of the dielectric function of VO₂ as it undergoes the SMT; however, previous measurements of the optical constants of VO₂ reveal significant variations between experiments.

To understand systematic variations due to growth conditions, films of VO₂ were deposited on either silicon, glass, or sapphire substrates by pulsed laser ablation of vanadium metal targets in 10 mTorr oxygen (O₂) background gas, followed by annealing at 450°C in 250 mTorr of O₂. Anneal times were varied from 30 to 90 depending on film thickness; deposition thickness was varied from 20 nm to 200 nm. For each sample, temperature-dependent spectroscopic ellipsometry measurements at optical and near-infrared wavelengths were conducted to determine the dependence of the optical constants on film thickness, substrate and crystallinity, and temperature.

Bruggeman and Maxwell-Garnett effective-medium formulations were used to account for three constituent materials: semiconducting VO₂, metallic VO₂, and vanadium pentoxide (V₂O₅). The effective dielectric functions were modeled using Lorentz and Tauc-Lorentz oscillators. Our results show that the contribution of V₂O₅ to the effective dielectric function increases with annealing time, consistent with previous studies. The results are also substantiated using Rutherford backscattering, X-ray photoelectron spectroscopy and X-ray diffraction.

Energy Frontiers Focus Topic Room: 102 - Session EN+MS+VT-ThA

Photovoltaics Manufacturing
Moderator: V. Ku, Satcon Technology Corporation

2:00pm  EN+MS+VT-ThA1  Waste Not, Want Not. L.V. Maness, Jr., South Park Platinum, Inc.  INVITED

Rapidly increasing demand for certain geological commodities, in particular semi-conductors, rare earth elements (REE) and those with other desirable characteristics, are making the extraction of formerly ignored elements of significant financial interest for miners, recyclers and other commodity producers. This need is tempered by legal and regulatory requirements that minimize pollution. Historically, miners, smelters and others have used almost all the physical, chemical and electromagnetic characteristics of ores as a means of extraction and concentration – with the exception of the unique benefits afforded by processing in vacuums. Vacuum use enables the clean separation and capture of many commodities that heretofore were wasted, either up smokestacks, in water, or in waste dumps, etc. As a side-benefit, the coating with certain REE, etc., of heated surfaces will enable the selective emission of optimized thermal frequencies for use in generating electricity using Thermal Photovoltaics (TPV) technologies: this major improvement in TPV efficiency will result from the suppression in a vacuum of transfer of energy via conduction and convection. In addition, for the many commodities that vaporize selectively in vacuums, a vacuum-smelter would enable the non-polluting capture of all processed materials. Such capabilities will enable revolutionizing the economics of many operations, since the removal of one component from a mix will increase proportionally the relative percentages of the other components. This development will bring closer the goal of “No Waste Mining” and of the separation and use of certain otherwise useful constituents, such as arsenic, which are considered harmful, are tightly regulated and whose disposal is presently a very high cost-item.


INVITED
The U.S. Photovoltaic Manufacturing Consortium (PVMC) – a $300 million partnership between SEMATECH and the College of Nanoscale Science and Engineering (CNSE) of the University at Albany – will bring together the entire supply chain of companies to enable the development of advanced PV processes. Created as part of the U.S. Department of Energy’s (DOE) SunShot initiative, which is designed to reduce the cost of photovoltaic solar energy systems by about 75 percent over the next decade, the PVMC will engage over 40 companies and organizations from throughout the solar community. Through PVMC, SEMATECH and CNSE will spearhead a unique research and development collaboration through which industry, academia and government will accelerate the development, commercialization and manufacturing of next-generation copper indium gallium selenide (CIGS) thin film PV manufacturing technologies, increasing performance while driving down the cost and risk of bringing them to the marketplace. By integrating the industrial research consortium and manufacturing development facilities models, PVMC offers lab-to-fab capabilities that will support rapid commercialization of new technologies and incubation of new start-up firms. PVMC offers the Federal government an unparalleled opportunity to promote the competitiveness of the U.S. PV industry.

4:20pm  EN+MS+VT-ThA8  Potentials and Challenges for High Efficiency Multi Junction Solar Cells. Bedair, North Carolina State University

INVITED
The developments of multi junction solar cell will be outlined. The major challenges facing this structure will be discussed. There are several current approaches to improve the efficiency of MJ solar cells. They include: lattice matched structure, metamorphic s structure, inverted structure and GaAsN based approach. We will discuss the concept of strain balanced strained layer super lattices in improving the conversing efficiencies of lattice matched MJ solar cells.

The advantages and the limitations of each of these approaches will be presented and discussed. We also discuss the limitations facing the applications of MJ solar cell at high solar concentrations. We will address the tunnel junction issues for concentration exceeding 1000 suns. Series resistance and heat dissipation issues will also be outlined.

Finally the cost issues will be addressed with their limitations on the potential applications of this approach.

Energy Frontiers Focus Topic Room: 103 - Session EN+NS-ThA

Nanostructures for Energy Storage and Fuel Cells II
Moderator: J. Lewis, RTI International

2:00pm  EN+NS-ThA1  Charge-Storage Processes in Model MnO₂-Li-HOPG Systems: UHV-STM Investigations. S.C. Bharath, W. Song, J.E. Reutt-Robey, University of Maryland, College Park, K.R. Zavadil, Sandia National Laboratories

Nanostructured materials have the potential to substantially improve the speed, efficiency, and cyclic lifetime of energy storage systems such as the lithium-ion battery (LIB). Nanometer-scale oxide cathodes elements allow for greater extent of lithium incorporation due to improved strain accommodation relative to conventional cathode structures. Exposed surface facets in structures with high surface/volume ratio will act as gateways to lithium insertion, playing an important kinetic role in charge storage. However, detailed mechanisms of lithium insertion and their dependence on oxide facet orientation and grain size and shape are largely unknown. As a platform for fundamental investigations of charge-transfer processes in nanocristalline materials, we have synthesized a MnO₂-Li-HOPG model system. This system consists of low-dimensional β-MnO₂ and cubic spinel Li₁ₓMn₂₋ₓO₄ nanocrystallites, synthesized by the reactive co-deposition of elemental Mn and molecular oxygen on native and modified HOPG (C(0001)) surfaces with ultrahigh vacuum conditions. Nanostructural phase and composition has been confirmed using single particle electron diffraction, as well as TOF-SIMS and scanning AES. Morphology of individual nanocrystallites, and their registration to the C(0001) support, have been determined with ambient AFM and UHV STM methods. The morphological response of β-MnO₂ nanocrystallites to Li⁺ insertion has been preliminarily explored under solid-state conditions with UHV-STM. The sensitivity of these methods are currently limited by low β-MnO₂ – C(0001) electrical conductivity, and efforts to overcome this limitation will be described. Additionally, alkali seeded growth of MnO₂ –
can distinguish possible H₂ binding sites based on the perturbation of the (IR) absorption spectroscopy to characterize its interaction. IR spectroscopy is a powerful tool for studying the incorporation of hydrogen into MOFs using infrared spectroscopy. This work explores the incorporation of hydrogen into MOFs using infrared spectroscopy. Feasibility studies and characterization of the incorporation of hydrogen into MOFs using infrared spectroscopy are possible to enhance the interaction of molecular hydrogen with the host. Organic Frameworks (MOFs) are promising candidates for hydrogen storage materials because their high surface area and porosity facilitate high hydrogen storage.

Hydrogen storage is one of the most challenging problems in hydrogen-related research. University of Texas at Dallas funded by the U.S. Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences (BES) under award DESC0001160. Supported by the Science of Precision Multifunctional Nanоструктуры для хранения водорода (NENES), an Energy Frontier Research Center funded by the U.S. Department of Energy (DOE), Office of Science, Office of Basic Energy Sciences (BES) under award DESC0001160.

2:20pm EN+NS-ThA2 The Influence of Surface Chemistry as a Function of Salt Composition on the Rate Capability of LiNi₅MnₓO₂. Composite Electrodes for Li-ion Rechargeable Batteries as Investigated using XPS, R.A. Quinlan, Naval Surface Warfare Center, Carderock Division, Y.C. Lu, Massachusetts Institute of Technology, A.N. Mansour, Naval Surface Warfare Center, Carderock Division, Y. Shaoo-Horn, Massachusetts Institute of Technology.

LiCO₂ is currently the most widely used cathode material in commercial Li-ion battery technology because of its high working voltage, structural stability and long cycle life. However, cobalt is expensive and there are safety and toxicity concerns. Therefore, there has been a consensus to decrease or remove the cobalt in the starting material. Both the quenched and annealed electrodes were cycled in the voltage range of 2.0 V – 4.6 V in 1M LiPF₆/EC:DMC (1:1) and LiClO₄/EC:DMC (1:1:1). The composite electrodes (in the discharged state) were disassembled in an Ar environment glove box and transported to the XPS analysis chamber without exposure to ambient. Cycling performance shows enhanced capacity retention and device performance for annealed electrodes cycled in LiPF₆ with little to no enhancement observed for annealed electrodes when cycled in LiClO₄. Our initial analysis indicates the formation of both Ni and Mn fluorides/oxofluoride species on the surface of electrodes cycled in LiPF₆. Additional changes in the surface chemistry and the influence on the rate capability will be discussed.

2:40pm EN+NS-ThA3 Hydrogen Storage in Metal Organic Frameworks (MOFs). N. Nijem, University of Texas at Dallas, L. Kong, H. Wu, Y. Zhao, J. Li, D.C. Langreth, Rutgers University, Y.J. Chabal, National University of Texas at Dallas.

Hydrogen storage is one of the most challenging problems in hydrogen-based energy technologies. One of the goals of hydrogen storage is the ability to store a high volumetric density of hydrogen at room temperature. As a result, studies exploring molecular hydrogen interaction in storage materials are important to facilitate further development of materials. Metal-organic Frameworks (MOFs) are promising candidates for hydrogen storage because of their high surface area and porosity. High capacity physisorption on specific sites of the structures and because many options are possible to enhance the interaction of molecular hydrogen with the host. This work explores the incorporation of hydrogen into MOFs using infrared (IR) absorption spectroscopy to characterize its interaction. IR spectroscopy can distinguish possible H₂ binding sites based on the perturbation of the internal H₂ stretch mode. IR measurements are performed on saturated metal center MOFs varying the ligand and/or the metal center and on unsaturated metal center MOFs-74-M (M=Zn, Mg and Ni). We combine metal center MOFs varying the ligand and/or the metal center and on unsaturated metal center MOFs-74-M (M=Zn, Mg and Ni). We combine metal center MOFs varying the ligand and/or the metal center and on unsaturated metal center MOFs-74-M (M=Zn, Mg and Ni) and we show that dipole moments of adsorbed H₂ depends greatly on parameters such as geometry of adsorption site and H₂-H₂ interactions. Measurements performed at low temperatures on MOF-74 show that IR shifts of H₂ is greatly red shifted (an additional ∼30 cm⁻¹) due to H₂-H₂ interactions on close proximity adsorption sites, and that dipole moments of adsorbed H₂ can appreciably vary with loading. Our analysis indicate that the intensity of H₂ IR band cannot always be a measure of the amount of adsorbed H₂, therefore methods such as variable temperature IR (VTIR) used to deduce binding energies cannot always be implemented.
quantum dots (QDs) during the initial nucleation cycles of ALD PbS[5]. These QDs were deposited directly on nanowire surfaces, suggesting the ability to combine light trapping in nanostructured materials with quantum confinement effects.


4:20pm  
EN+NS-ThA8  

Successful water management in proton exchange membrane (PEM) fuel cells requires a delicate balance of fuel, oxidant and water transport through a variety of length scales from ~1 nm in fuel channels, through ~100 nm in the gas diffusion media, to ~1-10 nm in the catalyst layer. The combination of a high degree of porosity in the reaction zone, complex composition, heterogeneous wetting properties, and the presence of water in two phases (gas, liquid) makes optimization of the performance of PEM fuel cell challenging. We are studying water distributions in situ in thin sections of membrane electrode assemblies (MEA) equilibrated with water vapor at controlled relative humidity (SRH) by in situ scanning transmission X-ray microscopy (STXM). The intrinsic soft X-ray absorbance properties of the constituent materials allow direct in situ visualization of water uptake into MEAs and the differentiation and mapping of the gaseous and liquid/sorbed water. The method provides maps of liquid and gaseous water distributed over the catalyst layer, with coating down to sub-micron and can be applied to MEAs as well as the polymer electrolyte membrane. Condensation at specific sites is observed when the relative humidity at the MEA is higher than ~80%.

Research funded by AFCC and NSERC. Measurements were also made at the Canadian Light Source (supported by NSERC, NRC, CIHR, and the University of Saskatchewan) and at the Advanced Light Source (supported by the Division of Basic Energy Sciences of U.S. DoE.)

4:40pm  
EN+NS-ThA9  
Optimization of the Delta Phase in Bismuth Oxide Thin Films. P. Silva-Bermudez, O. Garcia-Zarco, Universidad Nacional Autónoma de México. E. Camps, L. Escobar-Arceon, Instituto Nacional de Investigaciones Nucleares, Mexico, S.E. Rodil, Universidad Nacional Autónoma de México

Bismuth oxide Bi$_2$O$_3$ has interesting technological applications, which have not been largely used due to the particular polymorphism of the material. Bismuth Oxide has five polymorphic forms: α, β, γ, δ and ε -Bi$_2$O$_3$. Among them, the low-temperature α and the high-temperature δ phases are stable and the others are metastable phases, as has been established by bulk solid-state studies. Each polymorph possesses different crystalline structures and various electrical, optical and mechanical properties. The face-centered cubic δ -Bi$_2$O$_3$ is stable over a narrow temperature range 729–825 °C (melting point) and it has the peculiarity of being among the few materials presenting high ionic conductivity at moderate temperatures (600–700°C). In this research, we explore how to obtain δ -Bi$_2$O$_3$ thin films as high-ionic conductivity for the development of micro solid state fuel cells. However, the first challenge is to find the deposition conditions of the magnetron sputtering system to ensure the formation of the desired δ -Bi$_2$O$_3$, which is only thermodynamically stable at high temperatures. Based on previous results of Fan et al. (Fan 2006), we choose as the deposition variables the substrate temperature (room temperature to 300 °C) and the power (100-200 W). Our target was pure Bi$_2$O$_3$, but the first results indicated that it was necessary to compensate oxygen losses; therefore the atmosphere was a mixture of Argon and Oxygen, where the Oxygen flow was 20% of the total. The results from the different characterization techniques suggested that substrate temperatures between 150 and 200°C are appropriate to obtain the δ -Bi$_2$O$_3$ phase at a high deposition rate, between 1.5 to 2 nm/s. X-ray diffraction (XRD) and Raman spectroscopy, it was possible to clearly prove the presence of the δ-phase. The explanation for the stabilization of the high temperature phase might be related to the 2-dimensional confinement and/or effect of the small crystalline size. The physical properties of the δ -Bi$_2$O$_3$ thin films were further investigated, optical properties by transmission spectroscopy and ellipsometric spectroscopy, the ultraviolet-visible range, surface resistivity by the four-points method, composition by X-ray photoelectron spectroscopy and X-ray energy dispersion.


5:00pm  
EN+NS-ThA10  

Piezoelectric materials may arguably be the most functional, versatile and widely used materials with a wide range of applications including mechanical sensors, actuators, energy storage and energy harvesting devices. The well established applications are largely based on inorganic piezoelectric materials. The successful employment of polymeric piezoelectric materials, such as polyvinylidene fluoride (PVDF) and its related co-polymers, although light weight, flexible, optically transparent and cost effective, are limited by relatively low piezoelectric coefficients, thermal stability, and durability. The focus of the group is to develop a multilayered piezoelectric PVDF system for improved energy harvesting and energy storage efficiency. These systems are fabricated using enabling technology in co-extrusion which allows more cost effective and large area device production. As opposed to traditional film-based techniques. Many efforts have been made by the team to fabricate these micro- and nano-layered systems resulting in much improved device performance. A three-time improvement of capacitive electrical energy density has been demonstrated. The focus of this research is to understand the physics of why these multilayered systems perform better than a single layer by developing a characterization technique using both confocal second harmonic generation (EFISH) laser spectroscopy. Our results have shown that SHG is a very sensitive, non-destructive and versatile technique that can be used to study the piezoelectric and structural properties of layered systems. When combined with EFISH this technique allows the interrogation of electrical properties within the individual layers and at the interfaces between the layers. This information can be used to design an interrogant based on these techniques. Many of these materials (e.g. current collectors found in solar cells, batteries, and charge storage) produce electronic and dielectric effects that can be studied using in situ techniques. These effects can be identified and measured by EFISH laser spectroscopy. Our results have shown that SHG is a very sensitive, non-destructive and versatile technique that can be used to study the piezoelectric and structural properties of layered systems.
technologically relevant band gap in graphene. We are particularly interested in a novel form of semiconducting graphene that we call nanoribbons, which consists of graphene perforated by regular hexagonal arrays of nanoscale holes. Unlike nanoribbons, nanoperforated graphene advantageously retains a large-area two-dimensional form factor. In this talk, we will discuss the inter-relationship between the physical structure of nanoperforated graphene and its electronic properties, with specific emphasis on how its band gap experimentally varies with feature size and how charge transport is affected by structure (including the role of edge defects and the observation of single-electron charging effects). We will also report on efforts in our group to realize nanostructured graphene materials with well controlled edge structure and superior properties via scalable and rationally controlled bottom-up growth that avoids top-down etching without sacrificing arbitrary pattern forming ability.

2:40pm GR+TF+NS-ThA3 Quantum Pumping in Graphene Nanoribbons, T. Kauer, Ohio University, L. Arrachea, Universidad de Buenos Aires, Argentina, N. Sandler, Ohio University

The interest in the development of devices at the nanoscale has intensified the search for mechanisms that provide tailored control of transport properties while reducing effects of heat dissipation and contact resistance. For instance, charge pumping is one of the current generating methods that allows for minimizing the effects of contact resistance. Charge pumping is the mechanism used to generate DC currents in open-quantum systems by applying local de-phased time-dependent potentials. We analyze the properties of non-equilibrium zero-bias current through nano-ribbons using tight-binding Hamiltonians and the Keldysh formalism. This theoretical treatment, based on non-equilibrium Green's function techniques, is the most appropriate one to address questions for systems in non-linear, out of equilibrium conditions. We develop a numerical implementation for the models described below in a wide range of non-equilibrium regimes.

After reviewing results for quantum pumping in a one-dimensional chain attached to two reservoirs, with two local single-harmonic potentials oscillating in time, we study finite-width ribbons of square and graphene lattices. The transmission function reveals the value of the resonant frequency and explains how the quantum charge pumping works. We analyze the dependence of the DC current as a function of different parameters such as chemical potential, pumping amplitude, frequency, etc. In addition, the behavior of the current is explained. Pumped currents can also be generated by application of laser fields. We present the comparison between these two pumping methods. Possible extensions for disordered systems will be discussed.

3:00pm GR+TF+NS-ThA4 First-principles Study of Field Emission from Graphene Nanoribbons, J. Driscoll, K. Varga, Vanderbilt University

A real-space, real-time implementation of time-dependent density functional theory [1,2,3] is used to study electron field emission from graphene nanoribbons. The structures are shown to be good field emitters with spatial variation of the emission current influenced by the presence of passivating hydrogen. The nanoribbons are seen to produce slightly lower currents than nanotubes formed from the ribbons. Spin-polarized field emission from carbon nanotubes has been calculated with and without Fe adsorbates (atoms and clusters). It was observed that various adsorbates cause the separation of density into spin-polarized regions. The calculations predict that carbon nanotubes with various adsorbates can be used as spin-polarized current sources. The spin-polarized results for nanotubes will be compared to similar calculations for graphene nanoribbons.

References

Acknowledgments
This work is supported by NSF grant CMMI0927345.

3:40pm GR+TF+NS-ThA6 Quantum Transport Properties of Modified Graphene Nanoribbons with Boron Nitride Domains at the Nanoscale, A. Lopez-Bezanilla, Oak Ridge National Laboratory

Carbon-based systems are being widely investigated as potential candidates for nanoelectronic interconnects and transistors. The control of electric current is, therefore, an important challenge in nanostructures engineering. The possibility of creating hybrid one-atom thick layers containing C, B and N atoms has attracted much attention as they can provide an efficient way to create new materials with properties complementary to those of graphene and h-BN. Here we present a theoretical methodology and study of charge transport through GNRs with BN domains randomly distributed along the ribbon surface. We resort to both first principles calculations to obtain suitable parameterization of the electronic structure, and a transport approach based on the ab initio results to explore conduction regimes through large and disordered systems. The quantum transport modeling is based on the Green’s function formalism, combining an iterative scheme for the calculation of transmission coefficients with the Landauer’s formula for the coherent conductance.

Our results describe how the conductance of the hybrid systems is altered as a function of incident electron energy and BN domain density. We explore the transport regimes comparing different degrees of BN doping and BN domain size for ribbons of various widths and lengths on the order of the micrometer. A comparison with other types of defects such as atoms in epoxy configuration and functional groups covalently attached to the ribbon surface will also be discussed.

4:00pm GR+TF+NS-ThA7 Simple and Scalable Route for the ‘Bottom-Up’ Synthesis of Few-Layer Graphene Platelets and Thin Films, K. Coleman, University of Durham, UK

Graphene has generated much interest owing to its exceptional electronic properties and high mechanical strength. This has enabled new types of electronic devices and composite materials to be envisaged. The main problem is the availability of the material and the difficulties associated with large-area synthesis. Here we present a simple, consistent and scalable chemical vapour deposition method involving metal alkoxides in ethanol to produce few-layer graphene platelets. The graphene platelets have been fully characterised using TEM, SEM, AFM, XPS and XRD. The methodology used has the added flexibility in that it can be used to grow conducting transparent thin films on inert substrates such as silicon wafer and quartz glass. Importantly, no heavy metal catalysts were required to produce the few-layer graphene platelets or graphene films and all non-carbon by-products are soluable in water.

4:20pm GR+TF+NS-ThA8 Approaching the Intrinsic Bandgap in Suspended High-Mobility Graphene Nanoribbons, M.-W. Lin, C. King, Wayne State University, L.A. Agapito, N. Kioussis, California State University Northridge, Y. Zhang, M.-C. Cheng, Wayne State University, W.L. Wang, E. Kaxiras, Harvard University, Z.X. Zhou, Wayne State University

We report the first variable-temperature electrical-transport study of suspended ultra-low-disorder GNRs with nearly atomically smooth edges. Suspension of the GNRs not only removes the substrate influence but also allows a thermal removal of impurities, including those trapped at the interface between the GNR and the substrate, leading to a substantial increase of the carrier mobility. We observe high mobility values over 3000 cm2 V-1 s-1 in GNRs that are ~3-4 nm wide, the highest reported to date on GNRs of similar dimensions. Furthermore, we demonstrate that the activation gap extracted from the simple activation behavior of the minimum conductance and residual carrier density at the charge neutrality point approaches the intrinsic bandgap in ultra-low-disorder GNRs. Comparison of the bandgap values of multiple samples shows that the bandgaps in our ultra-low-disorder samples are closely proportional to the ribbon width, consistent with theoretical predictions. On the other hand, non-negligible disorder in GNRS obscures the observation of the intrinsic bandgap in transport measurements. In addition, the size of the bandgap derived from the transport measurements is in quantitative agreement with the results of our complementary tight-binding calculations for a wide range of chiral angles characterizing the GNR structure, suggesting that the underlying electronic origin of bandgap enhancement is the magnetism of the zigzag edges.

4:40pm GR+TF+NS-ThA9 Fabrication of Polymer-Protected Graphene Nanoribbons by Thermal Dip-Pen Nanolithography (tDPN), W.K. Lee, J.T. Robinson, R. Stine, A.R. Laracuente, Naval Research Laboratory, W.P. King, University of Illinois at Urbana Champaign, P.E. Sheehan, Naval Research Laboratory

The lithographic patterning of graphene nanoribbons (GNRs) to engineer band gaps has gained much attention as one path to realizing graphene-based devices. We employed thermal dip-pen nanolithography (tDPN) to pattern GNRs on CVD single-layer graphene (SLG) that had been transferred onto a SiO2 substrate. In tDPN, a heatable AFM cantilever regulates the deposition of an ink through controlled melting, much like a nanoscale soldering iron. tDPN has been successful at depositing polymers ranging from semiconductors to insulators on a variety of surfaces. To create the nanoribbons, we deposited poly styrene (PS) ribbons via tDPN on a SLG film between the source and drain electrodes. The areas of the graphene not protected by the polymer were then modified to isolate thin graphene nanoribbons. We show that the PS protected ribbon was the only
Thin and bulk films upon thermal annealing (60-850°C) in vacuum absorption spectroscopy (IRAS) measurements of graphene/graphite oxide removal of oxygen in GO, we have performed the edge functionalization. To unravel the complex mechanisms leading to spectroscopy coupled with interactions, and edge termination with oxygen. Infrared absorption spectroscopy coupled with in-situ thermal annealing process [1] makes it possible to examine the chemical changes taking place during thermal reduction to identify and understand interacting molecular environment and the edge functionalization. To unravel the complex mechanisms leading to the removal of oxygen in GO, we have performed in-situ transmission infrared absorption spectroscopy (IRAS) measurements of graphene/graphite oxide (GO) thin and bulk films upon thermal annealing (60-850°C) in vacuum (10⁻¹⁰⁻¹⁴ Torr). Control of the edge geometry of finite-sized modified graphene flakes depends very much on the control of the processing methods. This edge reconstruction further determines electronic, electric, optical and mechanical properties of the exfoliated modified graphene flakes. Therefore, we not only perform studies deriving a thermal reduction mechanism, but also examine the edge reconfiguration with oxygen. We report here the observation of a surprisingly strong IR absorption band that occurs upon thermal reduction of GO. After annealing at 850°C in vacuum, the strong enhancement of the new IR active absorbance band is observed at ~800 cm⁻²[2]. The intensity of this band is 10-100 times larger than what is expected for the oxygen content of the reduced GO, namely between 5 and 8 at%. This band is assigned to a specific oxidation state, involving oxygen located in the basal plane (forming C-O-C bonds) and at atomically straight edges of reduced graphene. The large enhancement in IR absorption is attributed to the direct participation of electrons, induced by the asymmetric C-O-C stretch mode displacement. These findings open new possibilities in the field of nanoelectronics for all sensor and energy storage applications. [1] M. Acik, et al. J. Am. Chem. Soc. (2011), in preparation. [2] M. Acik, et al. Nat. Mater. 9, 840-845 (2010).

5:20p GR+TF+NS-ThA11 Study of Ridges on Epitaxial Graphene on 6H-SiC(0001). Y.Y. Li, Y. Liu, L. Li, University of Wisconsin-Milwaukee
The graphitization of hexagonal SiC surfaces provides a viable alternative for the synthesis of wafer-sized graphene for mass device production. During the later stages of growth, ridges are often observed on the graphene layers as a result of bending and buckling to relieve the compressive strain between the graphene and SiC substrate, which also introduce ripples in the otherwise atomically flat graphene sheet. In this work, we show, by atomic resolution STEM imaging, that ridges are in fact bulged regions of the graphene layer, forming one-dimensional (nanowire) and zero-dimensional (quantum dot) nanostructures. We further demonstrate that their structures can be manipulated and even new ones created by the pressure exerted by the STEM tip during imaging. These results and their impact on the electronic properties of epitaxial graphene on SiC(0001) will be presented at the meeting.

MEMS and NEMS Group Room: 105 - Session MN-ThA

Multi-scale Interactions of Materials and Fabrication on the Micro- and Nano-scale
Moderator: A.V. Sumant, Center for Nanoscale Materials, Argonne National Laboratory

2:00p MN-ThA1 Heterogeneous Microsystem Integration with Self-Assembly, K. Bohringer, University of Washington INVITED
Self-assembly is the spontaneous and reversible organization of components into ordered structures, representing an alternative to the conventional manufacture of systems made of components from milli to nano scales. First commercial applications of self-assembly have appeared in recent years, for example in the fabrication of radio frequency identification (RFID) tags. However, the full impact of this new approach towards hetero system integration will only be realized once self-assembly can be programmed on demand. This presentation gives an overview of several projects that aim at programmable self-assembly. A key concept is the "programmable surface" an interface whose properties can be controlled with high spatial and temporal resolution. Several crucial topics are discussed: real time control of interfacial properties; optimization of binding site designs; and algorithms for the modeling and control of self-assembly. Promising novel manufacturing methods are emerging that combine the precision and reproducibility of semiconductor fabrication with the scalability and parallelism of stochastic self-assembly and with the specificity and programmability of biochemical processes.

Recently, dip soldering has been used as a mechanism for driving the assembly of three-dimensional (3D), micromachined structures. Solder is deposited on adjacent metallic faces of planar polyhedral patterns, bridging the small gaps between individual faces. When all but one face of a polyhedral pattern are freed from the substrate and solder is reheated to a liquid state (reflow), the free faces of the pattern fold upwards, out of the plane, to form the desired polyhedron. The wetting of solder with regards to coverage of metallic faces has been described previously, but the lateral bridging between the metal faces remains relatively unexplored. The goal of this work is to characterize the parameters influencing the bridging and folding process for two different ways of dip-soldering: face and edge soldering. Face soldering refers to the complete wetting of metal faces while edge soldering refers to selectively applying solder on the edges of a face that come in contact with other faces when folded. Our work explores bridging yield for various gap sizes and face thicknesses for eight different polyhedral patterns. Experiments show that the thickness and gap size strongly influence successful bridging. Experiments also show that improved control over the bridging process increases the yield of folded structures. In particular, gap size is positively correlated to face thickness for successful folding. Moreover, face soldering results in higher yields than edge soldering for all patterns.

Graphene’s unparalleled strength, stiffness, and low mass per unit area make it an ideal material for nanoelectromechanical systems (NEMS), but graphene resonators have been challenging to fabricate in large numbers and have exhibited poor quality factor. Here, we present simple methods of fabricating large arrays of graphene resonators from CVD-grown graphene and discuss their properties. We focus on circular graphene resonators with diameter of up to 30 microns, for which we observe highly reproducible resonance frequencies and mode shapes, as well as a striking improvement in the membrane quality factor with increasing size. The largest graphene resonators display quality factors as high as 2400 ± 300, about an order of magnitude greater than previously observed quality factors for monolayer graphene. These measurements shed light on the mechanisms behind dissipation in monolayer graphene resonators and demonstrate that the quality factor of graphene resonators relative to their thickness is high compared to nanomechanical resonators demonstrated to date. We conclude by providing an outlook for graphene NEMS and their applications.
We will detail work to develop a protocol for improved encasement of nanorods overhanging the cantilever leading edge to protect against damage, as well as our progress in implementing the nanomagnet-tipped cantilevers in MRFM experiments to rapidly detect single electron spins.


Electrochemical deposition is an attractive method for batch processing of magnetic films in patterned structures. Electroplating is a relatively simple process with a wide variability and control of film thickness and good scalability and compatibility with most of the MEMS microfabrication processes. Additionally, electrochemical deposition allows for the controlling of magnetic film anisotropy a key factor for the design of devices that operate in-plane or out-of-plane. Interest has been shown in CoNiP thin films for use in a number of MEMS applications however, the issues of integration were not addressed.

The integration of CoNiP magnetic films into MEMS sensors was studied. Through-mask electrodeposition of 1-2 μm thick magnetic films from concentrated ammonium chloride electrolyte was carried out at current densities of 30-150 mA/cm2 using both direct current and pulse plating modes. The effects of current density, seed layer, passivation layer, pattern size and geometry on magnetic properties and feature-scale thickness distribution were investigated. Geometries included various arrays of micron scale stripes and dots, and large 1-4 mm2 square areas. Feature scale profiles and magnetic properties of the films are influenced by current density as well as by feature size and geometry. Magnetic properties of CoNiP films after post-electrodeposition processing remain in the range suitable for sensor operation and are therefore shown to be suitable for integration in MEMS sensor.

Micropatterned CoNiP magnetic thin films have been integrated into silicon-on-insulator (SOI) MEMS devices. The patterned micromagnets – large scale arrays, stripes and dots – were chosen for feature-scale thickness distribution in relation to pattern geometry and current density, the effects on magnetic properties due to post-electrodeposition processing and their compatibility with standard MEMS process chemicals. Thickness distribution is strongly correlated with pattern geometry and current density. Magnetic properties remain in a range suitable for integration into MEMS devices following post-electrodeposition fabrication processes such as lithography, sputtering and etching.

We will present our preliminary efforts in building an integrated NOMS large-scale-integrated lab-on-a-chip NEMS sensing arrays.
nanophotonic NOMS chips in vacuum via free-space focusing onto grating couplers. The external-to-vacuum optics arrangement gives independent control over the position and input/output angles of both the input and output laser beams. This geometry allows us to directly compare photonic readout of NEMS motions with conventional free-space Fabry-Perot interferometry. Finally, we will update our progress in 3D integration of NEMS and photonics.

**Nanometer-scale Science and Technology Division**

**Room: 203 - Session NS-ThA**

**Biological Nanomaterials**

**Moderator:** N.A. Burnham, Worcester Polytechnic Institute

2:20pm **NS-ThA2 Biologically-Inspired Reversible Adhesives: Where Are We Now?**

S. Gorb, Zoological Institute at the University of Kiel, Germany

**INVITED**

Biological hairy attachment systems demonstrate their excellent adhesion and high reliability of contact. The structural background of various functional effects of such systems is discussed in the present paper. Additionally, it is demonstrated here, how comparative experimental biological approach can aid in development of novel adhesives. Experimental studies show that the effective elastic moduli of fiber arrays and spatiulike terminal elements are low, and this is of fundamental importance for adhesion enhancement on rough substrata and for an increased tolerance to defects at the level of individual contacts. Based on the broad structural and experimental studies of biological attachment devices, the first industrial bioinspired reversible adhesive foil was developed, which adhesive properties were characterised using variety of measurement techniques and compared with the flat surface made of the same polymer. The microstructured foil demonstrates considerably higher pull-off force per unit contact area. The foil is less sensitive to contamination by dust particles, and after washing with water, its adhesive properties can be completely recovered. This glue-free, reversible adhesive is applicable in dynamic pick-and-drop processes, climbing robots, and other systems even under vacuum conditions. The foil represents therefore a considerable step towards development of industrial dry adhesives based on the combination of several principles previously found in biological attachment devices.

3:00pm **NS-ThA4 Ultrastable Superparamagnetic Nanoparticle Design for Membrane Assembly and Triggered Release**

E. Amstad, M. Teztor, ETH Zurich, Switzerland, E. Reinhult, University of Natural Resources and Life Sciences Vienna, Austria

Application of superparamagnetic iron oxide nanoparticles as biomedical imaging contrast agents and as actuators in smart materials, e.g. for drug delivery and release, require them to retain high stability even in extremely dilute suspensions, high salt and at elevated temperatures. These requirements can only be met by sterically repulsive stabilization through irreversibly binding, low molecular weight dispersants of e.g. poly(ethylene glycol) or a similarly irreversibly bound organic shell which stabilizes the nanoparticle into another matrix material.

We have recently demonstrated that we can stabilize magnetite nanoparticles which fulfill these stability criteria using self-assembling dispersants with nitrocatechol anchors (1-2). This allows us free control over the dispersant type by simple co-adsorption of dispersants to as-synthesized core Fe₃O₄ particles. Combined with independent control over the Fe₃O₄ core size in the range 3-15 nm a versatile toolbox for assembly of various smart materials and for biomedical applications has been created. This presentation is focussed on recent results demonstrating and characterizing assembly of such nanoparticles into membranes of stealth liposomes (3). We show that there are strict requirements for the size of particles that can be assembled into lipid bilayer membranes and that a requirement for efficient assembly and actuation as well as liposome stability is to ensure stability of the hydrophobic shell surrounding the nanoparticle within the membrane. Encapsulated molecules were released multiple times by application of short bursts of alternating magnetic fields through a localized phase change in the membrane without heating of the surrounding aqueous environment. This allowed control of both timing and dose release. The highest efficiency of phase change and encapsulation was obtained for irreversibly stabilized superparamagnetic iron oxide nanoparticles with diameters <6 nm inserted into the lipid membrane.

1. E. Amstad et al., Nano Lett, 9;4042 (2009)

3:40pm **NS-ThA6 Nanoscale Electrical Interaction between Carbon Nanotubes and DNA, Y. Cao, Y. Xu, Vanderbilt University**

Carbon nanotubes-bioconjugated systems have emerged as one of the most promising materials for biological and biomedical applications, such as biosensors, drug delivery, and imaging. Recently, Carbon nanotubes (CNTs) have shown the ability to protect bound DNA cargos from enzymatic cleavage both during and after delivery into cells. This ability may result from the interaction between CNTs and DNA, which makes DNA resistant to nuclease degradation. Therefore, it is important to study the interaction between CNTs and DNA. In this work, we have developed a nanoscale optoelectronic probing system by combining highly-sensitive CNT transistors with advanced dual-trap optical tweezers to investigate the interaction between CNTs and DNA at the single-molecule level. We tightly bonded both ends of a DNA molecule with microbeads, which could be held and manipulated by optical tweezers. When the DNA molecule moved close to a suspended CNT transistor, the negative charge from the DNA molecule would change the local electrostatic environment around the CNT. Through scanning photocurrent measurements, the electrical coupling between individual DNA molecules and CNTs could be investigated.

4:00pm **NS-ThA7 Surface Functionalization of Nanomaterials: From Heterogeneous Catalysis to Nanoparticle Drug Delivery**

W. Gao, Brigham and Women’s Hospital and Harvard Medical School

**INVITED**

The advent of nanotechnology has vastly advanced our fundamental understandings on nanomaterials, in particular their surface properties. It has also revolutionized the way we functionalize these materials to exploit novel properties and applications. For example, metallic nanoclusters can be processed into different morphologies on support surfaces and subsequently allow desired reaction pathways to occur. In addition, a great number of metal-oxides have been grafted onto single-crystal surfaces. It is thus important to understand the functional effects of such systems is discussed in the present paper.

The highest efficiency of release and encapsulation was obtained for irreversibly stabilized superparamagnetic iron oxide nanoparticles with diameters <6 nm inserted into the lipid membrane. 224
Plasma Science and Technology Division
Room: 202 - Session PS+SS-ThA

Plasma Surface Interactions (Fundamentals & Applications) II

Moderator: A. Kumar, Case Western Reserve University

2:00pm PS+SS-ThA1 Plasma Prize Lecture - The Role of Atomic Hydrogen on Plasma Synthesis of Carbon Nanotubes, E.S. Aydil*
University of Minnesota

INVITED

Hydrogen containing discharges are used widely in plasma synthesis of a variety of nanstructures including nanoparticles and carbon nanotubes. We developed a method for measuring the H-atom flux at the plane of the substrate surface during H₂ plasma exposure. Our method is based on infrared absorption of the change in free-electron absorption in a polycrystalline ZnO film when this film exposed to H atoms. Hydrogen acts as an electron donor in ZnO, and thus the concentration of hydrogen-generated free carriers can be extracted from their absorption in the infrared. The change in the concentration of free carriers can in turn be related to the flux of H atoms impinging on a ZnO film placed on the substrate plate. Using this and a suite of additional plasma and material characterization techniques, including optical emission, infrared, and Raman spectroscopy and electron microscopy we systematically investigated the interrelation among plasma gas phase composition, catalysts morphology, catalyst structure, and carbon nanotube structure in plasma enhanced chemical vapor deposition of carbon nanotubes. The structures of carbon nanotubes grown from catalytic nanoparticles via PECVD in CH₄/H₂ mixtures show a strong dependence on the H₂/CH₄ ratio in the feed gas. Hydrogen plays a critical role in determining the final carbon nanotube structure through its effect on the catalyst crystal structure and morphology. At low H₂/CH₄ ratios, iron catalyst nanoparticles are converted to Fe₃C and well-graphitized nanotubes grow from ductile bcc iron nanocrystals that are easily deformed into tapered nanocrystals that yield nanotubes with thick walls. In the limit of pure hydrogen the cylindrical graphene walls of a nanotube are etched and amorphized by the H atoms. Etching is not uniform across the length of the CNT but rather, small etch pits form at defective sites on the CNT walls along the entire nanotube length. Once an etch pit is formed, etching proceeds rapidly, and the remainder of the CNT is quickly etched away.

2:40pm PS+SS-ThA3 CF and CF₂ Contributions to Plasma-Enhanced Chemical Vapor Deposition of Fluorocarbon Films in CF₆ Systems, M.F. Cuddy, E.R. Fisher, Colorado State University

Inductively coupled fluorocarbon (FC) plasmas produced from CFₓ, (x ≥ 1) precursors are widely employed in industrial processes ranging from circuit fabrication to deposition of low-K optical coatings. The utility of the plasma system is largely dictated by the x/y ratio, specifically in that precursors with lower ratios tend to more efficiently deposit FC films. Film growth is thought to be related to the behavior of radical species within the plasma, such as CFₓ and CFₓ. We report here on gas-phase behavior of these radicals in FC plasma systems with precursor x/y ratios ≤ 4, including relative gas-phase concentrations and kinetics and subsequent contributions to Si wafer processing. Our imaging of radicals interacting with surfaces (IRIS) experiment reveals that the propensity for scatter of CFₓ and CFₓ declines dramatically with decreases in precursor x/y ratio. Similarly, with decreasing x/y ratios, we observe increases in FC film surface energies, suggesting that avenues to tailor specific film properties are feasible. High-resolution x-ray photoelectron spectra and surface sum frequency vibrational spectra which corroborate the relationship between the choice of precursor and characteristics of deposited FC films will also be discussed. Ultimately, this work aims to establish a connection between species behavior near surfaces and resulting film properties during FC plasma processing.

3:00pm PS+SS-ThA4 Polymer Surface Modification: Real-time In Situ Electron Spin Resonance Study for Plasma Processes, K. Takeda, H. Kondo, M. Sekine, M. Hori, Nagoya University, Japan

A comprehensive understanding of interaction between plasmas and soft-materials is essential for advanced plasma processing technology. Simultaneous measurements of the gas-phase ESR signals and the surface dangling bond signal [1,2], and kinetics analysis of radical formation were performed using in situ real-time electron spin resonance (ESR). Chemical reactions under plasma are very complicated, due to the simultaneous irradiation of electrons, ions, radicals, and photons. Therefore, the individual contributions of each of these reactive species in the plasma must be elucidated.

An ESR system was connected to a plasma discharge system (2.45 GHz, 50 W) using a quartz tube with an inner diameter of approximately 9 mm. Gas (H₂/O₂, etc.) was flowed into the quartz tube and the pressure was maintained at approximately 10 Pa in the down-flow region. ESR measurements were conducted using a standard X-band (9 GHz) spectrometer (Bruker Biospin, EMX plus) with a microwave resonator. The quartz tube and polymer (PTFE, PMMA, etc.) film sample were inserted inside the ESR cavity in the down-flow region, typically 20 cm from the plasma discharge.

Individual contributions from gaseous radicals and plasma emission light have been studied in a similar manner using pallets for plasma process evaluation (PAPE) [3].

For irradiation with both atomic H and VUV on the PTFE film, significant acceleration in the rate of C-DB formation was observed at the beginning of irradiation. Moreover, the surface radicals produced immediately changed to peroxy-radicals when the treated PTFE films were exposed to air. This suggests high reactivity of C-DB with oxygen and the peroxy-radical species can contribute to enhancement of the surface biocompatibility.

In summary, during exposure of gaseous radicals on the polymer surface, the in situ real-time ESR technique was demonstrated as a new experimental approach to the microscopic understanding of chemical reactions on surfaces with gaseous radicals during plasma processes. We have successfully obtained information regarding the reaction mechanism with radicals generated by plasma induced surface interactions.

ACKNOWLEDGMENT

This work was supported in part by the Knowledge Cluster Initiative (Second Stage) of the Tokai Region Nanotechnology Manufacturing Cluster.

REFERENCES


3:40pm PS+SS-ThA6 Control of Hydrocarbon Surface Density during H₂/Ar Low Temperature Plasma Interaction, N. Fox-Lyon, G.S. Oehrlein, University of Maryland, College Park, N. Ning, D.B. Graves, University of California, Berkeley

Control of surface properties of hydrocarbon materials during interaction with H₂/Ar low temperature plasma has applications in diverse areas including: thin film synthesis, electronic device manufacturing, nuclear fusion reactor design, and plasma sterilization. Plasma processing/exposure of hydrocarbon materials can induce large changes in surface chemistry and morphology due to interaction with ions, reactive neutrals, and UV/VUV photons. Hard hydrocarbon materials such as amorphous hydrocarbon (a-C:H) and diamond-like carbon (DLC), suffer a loss of density and a thick modified layer with reactive plasma etching through reactant saturation of the surface and an increase in density and formation of a thin modified layer by inert plasma etching through selective sputtering of H. Whereas soft hydrocarbon materials (polymers, biomaterials) exposed to reactive and inert plasmas have been shown to increase in density and become chemically modified. In this work we have explored the plasma/surface interaction with hard/soft hydrocarbon films using different H₂/Ar gas feedstock compositions along with different ion energy/fluence to the surface. The time dependent changes in optical properties and the etch yields were found using real time in-situ ellipsometry. Using multilayer films (e.g., soft a-C:H over hard a-C:H) we monitored dynamic changes in the penetration/modification depths for different plasma chemistries.

Shallow modifications by inert plasmas (such as Ar) are well understood. Ar plasma depletes H from the surface of a-C:H films through selective sputtering, scaling with ion energy. H₂ plasmas have been shown to cause deep hydrogenation of the surface not predicted with TRIM models. Mixing the hydrocarbon gas (Ar/H₂) can be used to control the surface density from H depleted to H saturated. To better understand the roles of ion mass, etch rate and diffusion on the depth/degree of modification seen for H₂ plasmas, we performed comparable studies using D₂. We find that increasing the mass of the ions (by using H isotopes such as D) causes a large change in the etching and surface modification behavior. The etch rate of hard a-C:H in D₂ plasma is ~2 times the rate in H₂ plasma, and surfaces show a lower degree of modification (hydrogenation) than for H₂ plasma. In soft

* 2009 Plasma Prize Winner
hydrocarbon materials exposed to D$_2$, H$_2$, and Ar plasmas (listed in order of modification, least to greatest), we find that the density increase is dependent on the ion chemistry, energy, penetration depth, and mass. The data will be compared with molecular dynamics simulation results.

4:00pm PS+SS-ThA7 Atomic Force Microscopy Determination of the Elastic Modulus of Nanometer Thick, Ultra-Stiff Modified Layers after Plasma Etching of a Polymer Film. T. Lin, University of Maryland, College Park, H.C. Kan, National Chung Cheng University, Taiwan, Republic of China, R.L. Bruce, G.S. Oehrlein, R.J. Phaneuf, University of Maryland, College Park.

We report on a determination of the elastic modulus for ultrathin (~2 nm) stiff modified layers produced by argon plasma etching of the model photoresist polymer, poly(ether) (PS). Measured force curves allow a direct determination of the effective modulus of the damaged layer plus poly(ether) underlayer within a model which accounts for adhesive forces. The elastic modulus of the damaged layer is obtained via comparison with numerical simulations for contact between a spherical AFM probe and a bilayer-structured film system in a Hertzian mechanics model. Our results show directly that an extremely stiff modified layer is formed, with the modulus increasing with Ar-ion energy during etching, in good quantitative agreement with estimations based upon measurements of the dominant corrugation wavelength and buckling theory.


Higher aspect ratio of DRAM capacitor is required in order to maintain enough capacitance as the device structure is scaled down. The electrode of DRAM capacitor is fabricated in mold by RIE – typically dielectric. Thus, it is necessary to develop new techniques to fabricate higher aspect ratio dielectric mold for further DRAM scaling. RIE challenges for mold fabrication are high dielectric etch selectivity to mask, minimum bowing, and sustainment of enough bottom CD.

Generally speaking, mold etch process is optimized by accurate polymer control. However, conventional polymer control technique became marginal in high aspect ratio capacitive DRAM structure. Even slight excessive polymer makes bottom CD smaller, and sometimes clogs the top portion of mold. On the other hand, slightly lacking polymer bridges neighboring molds at the top and/or sidewall portion due to lack of etch selectivity to mask and/or bowed etch profile.

In this paper, we suggest DC superimposed capacitively-coupled plasma (CCP) etch technique as a breakthrough of high aspect ratio dielectric etch. High energetic electrons emitted and accelerated by superimposed DC hardens the sidewall polymer, and prevents etch profile from bowing. Bottom CD can be enlarged in the over etch step after etch front reaches etch stop layer. In general, bowing etch profile becomes significant during the over etch step. However, this effect can be avoided with stiff polymer enhanced by superimposed DC.

In conclusion, etch selectivity/durability of the undesired etch material/portion can be enhanced by superimposed DC as long as polymer is generated in the desired area by etch process optimization. Superimposed DC allows us to enhance etch selectivity/durability not with thickness of polymer but stiffness of polymer.


Plasma surface modifications have the ability to improve performance and realize new applications for polymer membranes in a variety of areas including: microfiltration, gas separation, and water treatment. Past studies in our labs have shown that although some polymer membranes can be successfully modified and are stable after treatment, many partially revert to their untreated state when aged. We are developing a technique that utilizes inductively coupled H$_2$O plasmas to implant polar functional groups onto the surface of track-etched polycarbonate membranes free of wetting agents. Water contact angle results on freshly treated and aged membrane samples show that the treated membranes have improved wettability compared to untreated samples and that the effects persist after treatment. Analysis of freshly treated samples using x-ray photoelectron spectroscopy (XPS) shows changes in surface chemistry, is consistent with analysis of XPS spectra of the C1s region shows that the fundamental polycarbonate structure is maintained near the surface. Treatment effectiveness under different plasma conditions, aqueous flux used to evaluate the performance of modified membranes, analysis of gas phase plasma species using optical emission spectroscopy (OES) to probe the processes that lead to surface modification and a comparison of a system using a mixture of O$_2$ and NH$_3$ as feedgasses will be discussed along with comparisons to other polymer membrane materials treated under similar conditions.


Fast and easy tunable patterning of surfaces has become of growing interest in the last couple of years in different fields like surface functionalization, thin film coatings and biomedical applications.

The concept of Plasma Printing combines the advantages given by the non-equilibrium character of pulsed corona discharges (tunability of surface chemistry), their creation and propagation along small volumes (local plasma treatment) and the mobility of the plasma source (i.e. plasma gun or plasma printer (inline processing)). This unique combination allows to treat (in real time) surfaces with arbitrary patterns design with limited restrictions unlike in the use of atmospheric pressure (dielectric barrier discharge) DBD for patterning of surface (also called DBD stamping).

InnoPhysics developed a proprietary Digital-on-Demand PlasmaPrint hardware solution that enables software patterned surface functionalization, etching and deposition of functional coatings on thin (plastic) substrates. A few kW sinusoidal pulse is applied on the electrode gap (pin to plate geometry configuration) in the 50-100 kHz range which generates a 1-10 μs pulsed plasma of about ~ 200 μm diameter in contact with the surface.

In order to assess the performances of the setup, a parametric study of polymers with respect to gas mixtures was performed to detect optimum of surface hydrophilization as well as selective chemistry groups grafting like OH, NH and CO.

PE, PET, FEP as well as PTFE (and PCTFE) were treated by nitrogen and different admixtures of gas/solution namely oxygen, ethanol, water and ammonia (NHLOH solution). The surfaces were analyzed by water contact angle (WCA), x-ray photoelectron spectroscopy (XPS) and IR spectroscopy in attenuated total reflectance mode (FTIR ATR).

WCA was used to measure the surface energy of the surface and significant improvements of wetting properties were found for a few seconds of treatment time (i.e. ~30” in the case of PET treated by N$_2$/NH). Grafting of N species up to a few percents was also found for various plasma compositions. The deconvolution of high resolution C1s, N1s and O1s spectra combined with the analysis of the IR spectra for the same conditions allowed eventually to get more insight in the chemical groups grafting at the surface following the plasma treatment.


In this presentation we will report our work on the design and experimental results of a plasma unit to clean the backside of substrates. Backside contamination of substrates can limit the quality of the front side process. Outgassing of backside contaminants can influence the front side process, or particles on the backside can influence the image quality in a wafer stepper. The complete backside cleaning unit uses three modules to remove both organic and an-organic particles as well as organic contamination layers. This is all done in a high vacuum environment. To remove particles we will use a commercial available tacky roller. The second method to remove particles uses an UV lamp that will charge the particles on the substrate and a strong electric field to remove these charged particles from the substrate. To remove organic contamination, for example resist residues after processing, we will use a RF plasma. The design of the RF plasma unit makes continuous cleaning over a large surface area possible. The gas in the plasma unit itself will have a pressure of 0.5 to 1 mbar. The vacuum chamber, in which the complete backside cleaning unit is placed has a pressure of around 10$^{-5}$ mbar when the plasma unit is in use. This difference in pressure is achieved by placing the inlet of the gas in the RF plasma unit and making the outlet to the vacuum chamber very small. This design leads to a large expansion of the gas at the outlet of the RF plasma unit. The
benefits of this design are that the pressure within the RF plasma unit is high enough for creating plasma and the pressure within the vacuum chamber is low enough to prevent that the whole vacuum chamber is filled with plasma.

This project focuses on backside cleaning of wafers within the "European Equipment & Materials Initiative for 450 mm" (EEMI450) under the ENIAC research program.

Within this European project, TNO is responsible for addressing contamination control issues. The concept of this backside cleaning unit can also be used to clean substrates used in roll to roll lines for solar cells or other processes that need continuous cleaning at high speed.

Plasma Science and Technology Division

Room: 201 - Session PS-ThA

Plasma Diagnostics, Sensors and Control II

Moderator: J.-P. Booth, CNRS/Ecole Polytechnique, France

2:00pm PS-ThA1 Characterization of Atomic Oxygen Emission by PROES and Ion-Flux Measurement in an ECR Plasma Etccher. V. Milosavlievic, Dublin City University, Ireland and University of Belgrade, Serbia, B. Dolajil, D. Gahan, Impedans Ltd., Ireland, N. Mac гарa, Dublin City University, Ireland, MB. Hopkins, Impedans Ltd., Ireland, S. Daniels, Dublin City University, Ireland

For many years, optical emission spectroscopy (OES) has been successfully used for the measurement and control of plasma products in industrial plasma reactors. We have extended this technique using phase resolved optical emission spectroscopy (PROES), in a industrial electron cyclotron resonance (ECR) plasma etccher.

Experiments were conducted in a pure argon discharge with a SiO2 wafer on the biased electrode. Argon ion bombardment of the wafer liberates oxygen atoms to the discharge. Therefore, oxygen is only present at the beginning of a discharge in solid state, i.e. in the SiO2 lattice. The ECR etccher used in this experimental study has a 2.45 GHz microwave generator with a maximum power of 2kW, W variable magnetic field of up 90 mT and 2MHz RF bias with maximum power of 250 W. The SiO2 wafer is mechanically clamped to the chuck to which the RF bias is applied.

In order to study the behavior of oxygen in the sheath region above the wafer an iCCD camera and high resolution spectrograph are employed and the iCCD camera is gated with respect to the 2MHz RF bias frequency. The production of oxygen is mostly due to RF voltage oscillation across the wafer induced by argon ion bombardment of its surface. The atomic oxygen spectral line intensity, from the 777 triplet, is monitored with respect to phase of the RF bias. Ion energy distribution functions, at the wafer surface, are measured using a floating retarding field energy analyzer (RFEA). The floating RFEA is placed on the rf biased wafer surface and signal cabling is taken out through the reactor vacuum pump iunnel. This prevents the need for any modification to the reactor configuration. The RFEA sensor is 7 cm in diameter and the wafer on which it sits is 200 mm in diameter resulting in significant exposure of the wafer to ion bombardment. Phase resolved measurements are made using the iCCD camera which is operated with a repetition rate of 2 MHz synchronously (triggered) with the RF bias. The integration gate of 3.90625 ns is locked to a fixed phase position within the RF cycle (500 ns). This gives exactly 128 intervals over the 2pi RF cycle. Spectra are resolved between the fixed phase and the gate allows one to cover the complete RF cycle. We record strong correlation between the ion-flux and the PROES data.

This work was a partly funded by SFI under the Precision project.


Passive, non-invasive optical emission measurements provide a means of probing important plasma parameters without introducing contaminants into plasma systems.* Due to the dominant role of electron-impact collisions in gas-phase reactions, our investigation focuses on characterization of the electron energy distribution function (EEDF). In particular, we highlight the ability to observe EEDFs under non-equilibrium conditions in which the EEDF deviates from the Maxwell-Boltzmann form. The energy dependence of the EEDF, which varies with plasma generation method and operating conditions, has significant implications for gas phase reaction rates and is thus critical to the predictive control of plasma process outcomes. EEDFs are determined using measurements of argon emission intensities in the 650-1150 nm wavelength range and measured metastable and resonance levels. Results using argon emission spectra will be presented for an inductively-coupled plasma (ICP) over a wide range of operating conditions (pressure, RF power, Ar/Ne/N2 gas mixtures), which show a depletion of the EEDF relative to the Maxwell-Boltzmann form at higher electron energies, in good agreement with measurements made with Langmuir probes and predictions of a global discharge model. These results are consistent with predictions of electron kinetics and can be explained in terms of reduced lifetime for energetic electrons due to wall losses and inelastic collisions. For Ne/Ar plasmas, analysis of neon emission spectra in addition to the argon analysis provides enhanced sensitivity to the presence of high-energy electrons. This example highlights the potential utility of this method as a tool for probing kinetics of many types of low-temperature plasma systems, which are typically characterized by non-Maxwellian EEDFs.


This work was supported by the Wisconsin Alumni Research Foundation (WARF) and by NSF Grant CBET 0714600.

2:40pm PS-ThA3 Two Dimensional Laser-Collision Induced Fluorescence Measurements in Low Pressure Plasmas. E.V. Barnat, Sandia National Laboratories

We describe a method to conduct optical emission spectroscopy (OES) measurements, electron beam excitation, that can be adopted to study and quantify process chemistry species. Our method and experiment, designed to be incorporated as a diagnostic system in process tools, relies on extracting electron beam from an inductively coupled plasma. First we will present and discuss results that show electron energy dependent cross section measurements in gas mixtures specifically Fluorocarbon process chemistries. Energy dependent optical excitation cross sections of Fluorine, Oxygen and ionic species as measured with this method will be presented.

Energy dependent optical excitation cross sections of Fluorine, Oxygen and ionic species as measured with this method will be presented. This work was supported by the Department of Energy Office of Fusion Energy Science Contract DE-SC0001939*.

3:00pm PS-ThA5 Two dimensional Laser-Collision Induced Fluorescence Measurements in Low Pressure Plasmas. S. Daniels, NCPST Dublin City University, Ireland

We describe a method to conduct optical emission spectroscopy (OES) measurements, electron beam excitation, that can be adopted to study and quantify process chemistry species. Our method and experiment, designed to be incorporated as a diagnostic system in process tools, relies on extracting electron beam from an inductively coupled plasma. First we will present and discuss results that show electron energy dependent cross section measurements in gas mixtures specifically Fluorocarbon process chemistries. Energy dependent optical excitation cross sections of Fluorine, Oxygen and ionic species as measured with this method will be presented.

This work was supported by the Department of Energy Office of Fusion Energy Science Contract DE-SC0001939*

3:40pm PS-ThA6 Controlled Electron Beam Excitation Method to Study Process Chemistries. P.L.S. Thamban, G. Padron-Wells, University of Texas at Dallas, J. Hsieh, Verity Instruments Incorporated, M. Kroeker, University of Texas at Dallas, J. Hosch, Verity Instruments Incorporated

We describe a method to conduct optical emission spectroscopy (OES) measurements, electron beam excitation, that can be adopted to study and quantify process chemistry species. Our method and experiment, designed to be incorporated as a diagnostic system in process tools, relies on extracting electron beam from an inductively coupled plasma. First we will present and discuss results that show electron energy dependent cross section measurements in gas mixtures specifically Fluorocarbon process chemistries. Energy dependent optical excitation cross sections of Fluorine, Oxygen and ionic species as measured with this method will be presented. The controllable excitation method and its applications to quantitative measurements of species in process chamber/exhaust will then be presented. Comparative measurements of species densities as measured with Fourier Transform Infrared Spectroscopy (FTIR) and e-beam excitation will be presented.

This project is funded by NSF-Grant (CBET-0922962) and Verity Instruments.

4:00pm PS-ThA7 Experimental Implementation of Robust Multivariable Real-time Feedback Control Design for RIE Plasma Processing System. Y. Zhang, B.J. Keville, A. Holohan, S. Daniels, NCPS T Dublin City University, Ireland

A robust multivariable real-time feedback control strategy for improving output characteristics of a reactive ion etching (RIE) plasma system is presented. Semiconductor fabrication is one of the major applications of low-pressure plasmas. During the course of manufacturing of semiconductor devices, it is often necessary to etch dielectric and/or metal layers to provide features in the layers for subsequent semiconductor processing steps. Reducing process variation is becoming ever more critical and challenging due to shrinking device geometries and an increase in wafer size. Developments in process control are struggling to keep pace with these more stringent demands due to the fact that most semiconductor manufacturing tools are run in open loop mode. In this case,
4:20pm PS-ThA8 Real Time, Multivariable Control of an SF6/O2/Ar Plasma, B.J. Keville, M.M. Turner, Dublin City University, Ireland Plasma deposited thin film electronics, SF6, mixed with oxygen and argon has been used for silicon etching in microelectronics manufacturing. Fluorine atoms produced by dissociation of SF6 etch Si with very high rates. Lateral etching, which reduces feature anisotropy, may be inhibited by the formation of a silicon oxide passivating layer on feature sidewalls. It has been demonstrated experimentally that feature profile shape is determined to a large extent by the balance between O and F radical densities at the surface of the substrate. In general, etch recipes are specified in terms of inputs such as gas flow rates, RF power and pressure and processes are run ‘open loop’. ‘Chamber matching’, which entails ex situ statistical analysis of metrics such as etch depth, uniformity, anisotropy and selectivity, is required to ensure that each chamber produces acceptable results. However, process reproducibility may be degraded due to real-time disturbances such as MFC and match network drift, wall seasoning and substrate loading. An alternative approach which would reduce the need for chamber matching and reduce process sensitivity to disturbances would be to specify a recipe in terms of plasma parameters such as O and F radical densities, and the fluxes and energies of ions at the wafer surface and to regulate these in real time by adjusting the inputs with a suitable real time control algorithm. This presentation describes how a real time, multivariable control algorithm for an SF6/O2/Ar plasma plasma may be designed with the aid of a control-oriented process model. The stability and efficacy of the control algorithm is demonstrated using a model of the process and a variety of simulated disturbances. Experimental implementation of the control algorithm on a laboratory capacitively coupled plasma is described.

5:00pm PS-ThA9 Maxwell Demon and its Instabilities, C.S. Yip, N. Hershelwitz, University of Wisconsin-Madison Previous experiments[1] have shown that in a low pressure, low temperature plasma, positively biasing an array of thin wires can increase electron temperature by creating an angular momentum trap to absorb cold electrons. In this experiment, such a Maxwell demon device was reproduced by welding 0.025mm tungsten wires onto stainless steel shafts, which were then covered with ceramic. This device was used to more than double plasma electron temperatures in a multi-dipole chamber operating in the mTorr regime. Moreover, the demon is observed to reduce the cold electron population in a plasma with a bi-Maxwellian electron distribution, leaving a single Maxwellian electron distribution. However, at high positive voltage, instabilities in the kHz range prevent acquisition of meaningful temperature data. The conditions of this instability are investigated by varying neutral pressure, plasma density and applied voltage up to 1500V in an argon plasma.

References

5:00pm PS-ThA10 Reliable Arc Detection and Arc Mitigation in RF Plasma Systems, D. Coumou, R. Choumey, MKS, ENI Products

1. Introduction
Arc disturbances in an RF plasma source are typically short duration transients arising from discharges between the plasma and the electrode, the plasma and the chamber sidewall, or discharges within the plasma that are induced by the build-up of polymer structures. When these transients occur, a reliable means is necessary to detect the presence of the arc and to interrupt the RF power delivery system to mitigate the arc event. We present a novel solution of arc detection using suitable tools from a communications equivalent paradigm that supersedes conventional heuristic methods. The proposed arc detection scheme is a quantitative approach measuring the relative arc energy of the plasma arc transient. A receiver operating characteristics (ROC) curve demonstrates the robust detection of arc transients relative to a ground truth source and yields insightful information contrasting the detection of arc disturbances in different RF sensor configurations in the RF power delivery system. An arc event is detected, arc mitigation is deployed based on suppressing the RF power with duration proportional to the detected arc energy. The rapid, and if necessary, repeated control of the RF source results in a reduction in the plasma potential to extinguish the arc source and alleviate subsequent damage. Results from PECVD and PVD tools corroborate the impact of this new scheme to significantly ameliorate thin-film manufacturing.

Advanced Surface Engineering Division
Room: 104 - Session SE+PS-ThA

Pulsed Plasmas in Surface Engineering
Moderator: J. Patscheider, EMPA, Switzerland

In this study, WO3 films have been synthesized using reactive HiPIMS of a metallic tungsten target in Ar/O2 mixtures. A comparison is made between results obtained in HiPIMS with those obtained with a conventional DC reactive magnetron sputtering (RDCMS) at identical mean power (PD). We discuss the influence of the pulse duration (t) and the target voltage (VD) on both the film deposition rate (RD) and the hysteresis behaviour. During the HiPIMS experiments, t is varied between 10 and 50 µs and VD between 800 and 1500 V. PD is kept constant by adjusting the frequency. In reactive mode, for a given value of PD, RD increases as t and VD are increased. Comparing the HiPIMS data with those recorded during the RDCMS process, it is found that for t = 50 µs and VD = 1500 V, RD in HiPIMS is larger than for the RDCMS discharge. In order to understand this behaviour, the target current waveforms associated with these working conditions have been studied. For this particular condition (t = 50 µs, VD = 1500 V), the discharge current waveforms in metallic and reactive mode are similar. This observation would reveal that the target surface chemistry is identical, although the discharge is ignited either in a pure Ar or in an Ar/O2 mixture.

The ion flux composition was also studied with a mass spectrometer located in front of the magnetron target. Time-resolved and time-averaged measurements were carried out.
control, developed by us, was used for high-rate reactive deposition of an argon-oxygen gas mixture was around 2 Pa. An effective reactive gas flow rate of 500 Hz at duty cycles ranging from 2.5 to 10%. The total pressure of the process was 15 Pa. The substrate was biased to a high negative rf potential, during their etching using a Modulated Pulse Power (MPP) Hollow Cathode Magnetron. The plasma conditions during the continuous and pulsed phases can be varied independently. The plasma source has been modified to allow for combined pulsed/continuous operations [1]. Parallel to the DC power supply, the plasma source is connected to a capacitor bank (5 kV, 8.4 mF, 100 kJ), which is discharged in the source to transiently increase the input power. Peak surface heat fluxes in excess of 1 MW/m² have been generated. Distinct discharge stages were normally observed in one MPP pulse packet. Time-dependent plasma parameters were investigated using a triple Langmuir probe to help understand the MPP discharge characteristics and its performances. Plasma behaviors were shown to closely depend on the pulse waveforms and various other parameters including pulse current, repetition frequency, pressure, and distance from the target. A high electron density \( n_e \) of \( 3 \times 10^{16} \) m\(^{-3} \) and an electron temperature \( T_e \) of 5 eV during the pulse were obtained at the substrate level, with an average power less than 8 kW. Compared with the DC magnetron sputtering at the same average power, the pulsed plasma density was an order of magnitude higher, which resulted in an enhanced ionization of the sputtered flux. As measured by an electrostatic gridded energy analyzer combined with a quartz crystal microbalance, the Cu ionization fractions above 30% were easily achieved by the MPP sputtering on the substrate level, twice higher than those by the DC sputtering. Increasing the pulse duty ratio or reducing the pressure resulted in a stronger ionization. The performance of Cu deposition in narrow trenches (70-100 nm) using the MPP sputtering was further studied, which exhibited an improvement over the DC sputtering. Stronger pulses with higher duty ratios and a lower gas pressure were preferred to reduce the overhang and achieve better step coverage and bottom coverage.

1.2×10⁻¹ mbar. The current work has shown that the concept of combining a 2.5:1 increased from 15% to 20% as pressure increased from 2.95×10⁻² to 1.55×10⁻¹ mbar. The coalescence threshold of unbiased vias with width 0.36 µm and aspect ratio of 1.2:1 was studied with cross sectional SEM. The effect of peak duty cycle of < 25% is used to achieve high peak powers and plasma deposition based on sputtering without magnetic fields. A plasma is generated in front of the target via an inductively coupled coil driven with a 13.56 MHz radio frequency (RF) power supply. The target is then biased to a high voltage to initiate sputtering. In order to ionise significant fractions of the sputtered flux, the RF power density is pulsed with peak values in excess of 30 Wcm⁻² to produce plasma density of the order of 10¹⁴ cm⁻³. A low duty cycle of < 25% is used to achieve high peak powers and plasma densities at low average power. The degree of ionisation of ICIS of Cu and Ti in Ar atmosphere were evaluated using optical emission spectroscopy and atomic absorption spectroscopy and the film microstructure and compositional data gather from XPS and X-ray diffraction (XRD) to the optical transmission and optical dispersion obtained by UV-Vis spectroscopy and spectroscopic ellipsometry, respectively.

4:40pm SE+PS-ThA9 Inductively Coupled Impulse Sputtering (ICIS): A Novel Technique for Ionised PVD. A.P. Ehiasarian, D. Loch, Sheffield Hallam University, UK.

One limitation of magnetrons is their use of inhomogeneous magnetic fields which constrains deposition of magnetic materials to thin targets, complex (magnetic) alloys by erosion-dependent stoichiometry, and oxides by build-up of arc-prone insulating layers on the target edge. Inductively Coupled Impulse Sputtering (ICIS) is a new technology for physical vapour deposition based on sputtering without magnetic fields. A plasma is generated in front of the target via an inductively coupled coil driven with a 13.56 MHz radio frequency (RF) power supply. The target is then biased to a high voltage to initiate sputtering. In order to ionise significant fractions of the sputtered flux, the RF power density is pulsed with peak values in excess of 30 Wcm⁻² to produce plasma density of the order of 10¹⁴ cm⁻³. A low duty cycle of < 25% is used to achieve high peak powers and plasma densities at low average power. The degree of ionisation of ICIS of Cu and Ti in Ar atmosphere were evaluated using optical emission spectroscopy and atomic absorption spectroscopy and the film microstructure and compositional data gather from XPS and X-ray diffraction (XRD) to the optical transmission and optical dispersion obtained by UV-Vis spectroscopy and spectroscopic ellipsometry, respectively.

Surface Science Division
Room: 107 - Session SS-ThA

Semiconducting & Ferroelectric Surface
Moderator: J. Millichick, University of Michigan

2:00pm SS-ThA1 Polarization Dependence of the Surface and Interfacial Chemistry of Ferroelectric Oxides. E.J. Altman, Yale University INVITED

In analogy to ferromagnets, ferroelectric materials develop remnant macroscopic electric fields that can be switched by applying an external field. The bulk electronic polarization is stabilized by compensating charges at the surface that can be supplied by adsorption or electronic and structural reconstructions. Because opposite compensating charges are required on oppositely poled surfaces, ferroelectric materials offer unique opportunities to create surfaces with switchable chemical properties. Further, thin ferroelectric films would be expected to switch their polarization in the presence of molecules that preferentially adsorb on one of the polar surfaces, suggesting a new avenue for chemical sensing. It will be shown that the adsorption of polar molecules such as alcohols and carboxylic acids depends on the ferroelectric polarization direction, with stronger adsorption on positively polarized. The differences in adsorption strengths are comparable to the energy required to switch 20 nm thick ferroelectric films, thus ferroelectric chemical sensing is feasible. This possibility is further explored through in situ measurements of changes in polarization of thin epitaxial titanate stack ferroelectric films in response to oxidizing and reducing environments. A limitation to switchable chemistry, however, is the low reactivity of typical ferroelectric oxides. Efforts to increase the reactivity through deposition of catalytic metals fail because the metals form three-dimensional clusters whose surfaces are too far from the ferroelectric substrate to be affected. Results of a combined experimental/theoretical approach to identifying catalytic oxides that form stable, atomically thin layers whose reactivity is influenced by the polarization direction of the ferroelectric will be presented. Specific examples include Cr₂O₃/LinBo₃ and perovskite layers on ferroelectric titanates.

2:40pm SS-ThA3 Surface Reactions of BaTiO₃ (001): STM Study. E.H. Morales, D.A. Bonnell, University of Pennsylvania

BaTiO₃ is currently used in Random Access Memories and is a versatile material with many potential applications. Ferroelectricity at a surface provides unique possibilities to examine mechanisms of molecular adsorption. Experimentally, the polarization can be manipulated in situ by negatively and positively poling the surface. In this manner, the interaction with foreign gaseous species can be controlled. It is necessary to develop fundamental understanding of BaTiO₃ surfaces in order to control the ferroelectric polarization and consequent interactions with adsorbates and in devices. Indeed advances in surface techniques allow for molecular adsorption of a variety of molecules; however, the mechanistic processes are not yet understood. Here we present STM of atomically resolved reconstructions, (5√x5√y) R26.6° and (3×1) and relate water adsorption to local polarization. Structural variations and adsorption sites will be discussed in terms of Density Functional Theory predictions. Preliminary results of the interaction of BaTiO₃ (001)'s surface and CO₂ will be presented as well.

3:00pm SS-ThA4 Polarization Dependent Interactions at Ferroelectric Surfaces. J. Giarra, M. Zhao, J.M. Vohs, E.H. Morales, D.A. Bonnell, University of Pennsylvania

Ferroelectric polarization at surfaces has been exploited over the last several years to mediate reactions in aqueous solutions. More recently studies have shown that ferroelectric polarization at surfaces affects molecular adsorption. The study presented here uses a variety of surface techniques (scanning tunneling microscopy and spectroscopy, low energy electron diffraction, and programmed desorption) to examine molecular adsorption on ferroelectric surfaces. Various molecules and various substrates are chosen to determine the effect of polarization orientation and magnitude on physisorption, disassociative chemisorption, defect mediated adsorption and chemical reactions. Specifically, results of interactions of small molecules on LinBo₃ and BaTiO₃ will be presented. We show that ferroelectric polarization can affect both physisorption and chemisorption bond energies as well as the sticking coefficients.

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INVITED

Scaling of gate oxides on MOSFETs requires nucleating oxide ALD in every unit cell of the semiconductor channel surface without disrupting the semiconductor surface (high mobility) while eliminating dangling bond states (low defect density). As an example, the TMA reaction products (dimethyl aluminum, DMA) spontaneously form a layer on the Ga-polar surface of charged aluminum nucleation in each unit cell. For integration of III-V MOSFETs on 300 mm silicon wafers, it is possible that a gate last process may be required for which As-decapping is not suitable. A combination of atomic hydrogen dosing, annealing, and TMA dosing is observed to produce an ordered unpinned passivation layer on air exposed InGaAs(001)(+4+2) surface with only monatomic steps. This TMA passivation layer on a cleared air exposed III-V has the same horizontal rows of dimethyl aluminum (STM) and the same unpinned interface (STS) observed with the TMA/decapped InGaAs(100). While TMA directly reacts to create a high density of sites to nucleate gate oxide ALD on III-V surface, for Ge(100) surface functionalization is required. The direct reaction of TMA on a Ge(100) surface and the effect of monolayer H2O pre-dosing were investigated. At 300K, a saturation TMA dose produced 0.8 ML semi-ordered ALD (100)Ge surfaces due to the dissociative chemisorption of TMA. XPS analysis of the 200°C annealed Ge(100) surface showed that only carbon content was reduced during annealing, while the Al coverage was maintained at 0.15 ML, consistent with the slow density of nucleation sites due to site blockage by reaction products. Conversely, saturation TMA dosing at RT on the monolayer H2O pre-dosed Ge(100) surface followed by annealing at 200°C formed a saturation layer of ALD bonds with an Al coverage a factor of two greater than the TMA only dosed Ge(100). The Ge-OH sites catalyze the dissociative chemisorption of TMA that exceed 300K to create a semiordered surface of Ge-O-Al bonds with an Al coverage double the coverage from a TMA surface without the H2O preadsorbed.

4:20pm SS-ThA8 Electrostatic Coupling of Surface Charge to Bulk Defect Behavior in Metal Oxides, P. Goett, K. Pangerl-Okamoto, A. Hollister, E.G. Seebauer, University of Illinois at Urbana-Champaign.

The technologically useful properties of semiconductor oxides such as titania and zinc oxide often depend on the concentration and diffusion of point defects. Near-surface effects are particularly important in nanoscale devices because most of the bulk is located in the vicinity of the surface. Past work in our laboratory with silicon and titania has shown that semiconductor surfaces serve as efficient pathways for generation and annihilation of point defects in the underlying bulk. Surfaces can, in addition, support electrically charged defects which create near-surface strong electric fields that can influence the local motion of charged defects resulting in the formation of space-charge layers. The electric field-driven accumulation or depletion of charged oxygen defects in such space-charge regions in metal oxides have direct implications on the performance of nanoscale devices such as gas sensors and memory resistors. Oxygen diffusion behavior was studied by exposing natural-abundance single-crystal rutile to isotopically labeled oxygen gas. The resulting profiles were up at the surface can be quantified by an electric potential (V_i). The profiles accumulation or depletion of charged oxygen defects in such space-charge material (V_i). The profiles accumulation or depletion of charged oxygen defects in such space-charge regions by rectifying contacts is a prerequisite for active devices. In the emerging field of transparent electronics, only metal-insulator-semiconductor field-effect transistors (MISFETs) were considered so far. In
this contribution transparent, high-performance MESFETs, inverters etc. based on ZnO and related ternaries are presented. We discuss design pros and cons as well as limitations regarding device performance, reliability and stability. 

The influence of the contact metal and dielectric passivation layers on the properties of ZnO Schottky diode, used as gate electrode within the MESFETs, as well as sources of non-idealities will be highlighted.


Amorphous oxide semiconductors (AOSs) are expected as an alternative to amorphous-poly-Si for thin-film transistors (TFTs) in next-generation flat-panel displays (FPDs) because AOS TFTs have many advantages such as large field-effect mobilities (>10 cm²/Vs) and low-temperature process [1]. For more advancing AOS optoelectronic technology, it is important to develop more various devices other than TFTs, and to study some remaining issues such as operation characteristics of AOS devices under light illumination.

In this study, we fabricated good and stable metal-AOS Schottky contacts made of amorphous In-Ga-Zn-O (a-IGZO) and bottom Pt electrodes at temperatures below 200°C even though it is generally difficult to make high performance oxide Schottky junctions [2]. It was found that the a-IGZO/Pt Schottky contacts have an ideality factor n ~ 1.1 and a Schottky barrier height φb ~ 0.9 eV, which were evaluated from their J-V curves using the thermionic emission model. From C-V results, the Schottky junctions operated in the full-depletion condition, whose C corresponds to the geometrical capacitance of the a-IGZO layer, and relative permittivity εr of a-IGZO was obtained approximately 13. However, the results of temperature dependences of J-V characteristics were unexplained if we take a simple uniform Schottky barrier model; we found that the barrier potential fluctuations model [3] explained them well, and the mean barrier height φb ~ 0.9 eV and the net electron affinity γ of a-IGZO at 4.2 eV were obtained. On the other hand, the Schottky contacts showed very small open circuit voltages (Voc) ~ 0.1 V and AM1.5 light illumination, which are far smaller than the built-in potential (Vbi ~ 0.4 eV) estimated from the C-V measurements in dark. We also observed that Voc decays with time after starting the light illumination. We will discuss the mechanism of the small Voc based on these results.


Printed electronics is a rapidly growing industry and within this emerging field there are three required material categories critical to fabricating active and passive circuitry - insulators (dielectrics), semiconductors (polymers), and conductors (metals). The increased interest in printable electronics as alternatives to silicon-based technologies is fueled by the promise of large-area, flexible, and ultra-low-cost devices. To enable the growing demands of printing processes this work develops metal-containing inks for the deposition of the coating metals - copper, silver, and gold. These metals are chip components ranging from interconnects to source and drain contacts in organic field effect transistors. The liquid ink approach is based on fundamental advances in coordination chemistry to fabricate discrete metal complexes that can be heated or irradiated to yield metallic films. Ultimately, inkjet printing technologies were employed to deposit these metal inks in specific, predetermined patterns that were directly transformed into active and passive devices. The versatility of this approach holds the possibility of printing any metallic design and pattern on virtually any type of substrate.

3:40pm TC+EM+NS-ThA6 A New Application for a-IGZO TFTs: An Addressable Microfluidic Electrowetting Channel Device, J. Noh, J.H. Noh, University of Tennessee, E. Krest, Heisekenf, University of Cincinnati, P.D. Rack, University of Tennessee

An electrowetting (EW) microfluidic platform designed for control and transport of aqueous and polar species has been fabricated on passive electrodes as well as an active matrix thin film transistor (TFT) array. To drive the EW devices we integrated the micro fluidic platform on a base-plane of transparent TFTs. Specifically, we have used an InGaZnO (IGZO) active layer for the TFT device which has superior performance and offers the benefit of transparent devices for biological and display applications. The TFTs are fabricated with a bottom-gate staggered structure with Cr gate and SiO2 gate dielectrics deposited via plasma enhanced chemical vapor deposition (PECVD). The a-IGZO semiconducting active layers are deposited using rf magnetron sputtering in a reactive Ar-O2 atmosphere. Finally, source and drain contacts were formed with evaporating Ti/Au. Finally the device is annealed in a N2ambient for electrical activation. For the EW device integration, Al electrodes are have been deposited various passivation layers. Subsequently a top dielectric and a hydrophobic Fluoropel layer are applied. In this presentation we will review the process flow and will discuss the materials integration issues of EW device and its effect on the TFT performance. We will illustrate the EW characteristics based on standard planar electrowetting on dielectric (EWOD) platforms and compare them to a new concept we have termed the “Laplace Barrier” which includes post arrays and enhances electrowetting characteristics.

4:00pm TC+EM+NS-ThA7 Amorphous Oxide Semiconductor Thin-Film Transistors, J.F. Wager, K. Hoshino, Oregon State University, B. Yeh, R.R. Hoffman, Hewlett Packard, Palo Alto, CA

Amorphous oxide semiconductor (AOS) thin-film transistors (TFTs) are transitioning towards commercialization for active-matrix liquid crystal display flat-panel display backplane applications. Additionally, AOS TFTs offer an attractive approach to printed electronics. The primary focus of this talk will be to discuss our novel approach to top-side passivation of bottom-gate indium gallium zinc oxide (IGZO) and zinc tin oxide (ZTO) AOS TFTs. Device performance between passivated and unpassivated AOS TFTs will be compared. Passivation mechanisms will be considered in the context of induced-gap state and device physics electrostatic modeling.

4:40pm TC+EM+NS-ThA9 Why Optimum Oxygen Pressure Range Exists for Fabricating Amorphous In-Ga-Zn-O Thin-Film Transistor and How it Should Be Optimized, K. Ide, J. Nomura, T. Kamiya, H. Hosono, Tokyo Institute of Technology, Japan

Amorphous oxide semiconductors (AOSs) represented by amorphous In-Ga-Zn-O (a-IGZO) are expected for large-area high-performance flexible electronic devices, because AOSs have large electron mobilities greater than 10 cm²/Vs and may even be fabricated at room temperature (RT). In particular, a-IGZO has good controllabilities of carrier concentration, and thin-film transistors (TFTs) exhibit superior properties in large-area and high-temperature applications. In this study, we investigated effects of oxidation on operation characteristics of a-IGZO TFTs. Bottom gate, top-contact a-IGZO TFTs were fabricated on SiO2/c-Si substrates by RF magnetron sputtering. Sputtering conditions were the RF power of 70 W and the total pressure of 0.55Pa. Two oxidation treatments were examined; (i) ozone annealing and (ii) varying a mixing gas ratio of Ar : O3 from 18 : 2 to 19.8 : 0.2 in standard cc per minute (scm) during the channel deposition. For the ozone annealed TFTs, annealing at ≤ 250°C produced good TFTs, while annealing above 300°C caused large hysteresis and S slope. After applying a high VGS larger than 40 V, the transfer characteristics showed the large VDS of 40V and the small hysteresis. Trap state around Fermi level of the large S state and the large VDS state were -4x1017 and -1x1017 cm⁻²eV⁻¹, which were estimated by C-V analysis. The large VDS state is very stable in the dark, but the TFT recovers to the initial large S state by light illumination. The photoresponse measurements revealed that deep trap states were formed at 2.3 eV below the conduction band minimum by applying a high VDS. Thermal desorption spectra showed that weakly-bonded excess oxygenes were incorporated in the a-IGZO layer. From these results, we built a subgap DOS model of the trap states of the excess oxygenes. We also confirmed similar behaviors in a-IGZO TFTs fabricated in high oxygen pressure conditions. This study reveals that the control of oxygen stoichiometry is important for obtaining good performance and stability of AOS TFTs.

5:00pm TC+EM+NS-ThA10 Effects of Low-Temperature Annealing and Deep Traps in Operation Characteristics of Amorphous In-Ga-Zn-O Thin-Film Transistors, T. Kamiya, Y. Kikuchi, K. Ide, J. Nomura, H. Hosono, Tokyo Institute of Technology, Japan

Amorphous oxide semiconductors (AOSs) represented by a-In-Ga-Zn-O (a-IGZO) are expected for channel materials in thin-film transistors (TFTs) for next-generation flat-panel displays such as jumbo-size / fast / ultrahigh-resolution liquid-crystal displays and organic light-emitting diode displays. These are also expected for flexible electronics devices because they may be fabricated on unheated substrates, and thus produce flexible displays and circuits on inexpensive plastic substrates such as PET. On the other hand, it is known that, although room-temperature fabrication is possible for AOS TFTs, post-deposition thermal annealing at ≥300°C is better employed to obtain good stability. To employ this technology to the flexible electronics, the annealing temperature should be lowered to 200°C or far below. We reported that wet O2 annealing produces the best performance TFTs when
annealed at ≥300°C, while it caused serious negative threshold voltage (V_{th}) shift at ≤200°C. In this paper, we report the origin of the negative V_{th} shift by employing photoresponse spectroscopy of TFT characteristics. It revealed that the near-valence band maximum (VBM) states are reduced significantly even by the low-temperature 200°C annealing, and implied that the negative V_{th} shift originates from free electrons released by annihilation of the near-VBM states.

5:20pm TC+EM+NS-ThA11 β-alumina (SBA): A Promising High Dielectric Constant Gate Material for Solution Processed, Transparent and Low Voltage Transistor Devices, B. Zhang, Y. Liu, H. Katz, Johns Hopkins University

β-alumina (SBA) has been discovered as a promising high dielectric constant gate material for solution processed, transparent and low voltage transistor devices. Some experimental evidence indicates that the mobile Na ion within two spinel blocks made by Al and O is responsible for the high dielectric constant. Transistors (W/L ratio 10) using SBA as gate layer and zinc tin oxide (ZTO) as active layer only need 2V to obtain 0.7mA output current. SBA material is compatible with organic semiconductors such as PQT12 and pentacene as well. Some key issues regarding using SBA for real applications, such as device stability in the ambient atmosphere, response under high frequency, and threshold voltage shift under gate bias have also been studied. It is found that encapsulating the device with CYTOP fluorinated polymer is an effective way to increase the operational stability of the devices in the ambient environment.

Thin Film Division
Room: 110 - Session TF+EM+SS-ThA

Applications of Self Assembled Monolayers
Moderator: M.R. Linford, Brigham Young University

2:00pm TF+EM+SS-ThA1 Organic Monolayers on Silicon-rich Substrates: Methods and Mechanisms. H. Zuilhof, Wageningen University, Netherlands

INVITED
SAMS on silicon-containing substrates including silicon, silicon nitride and glass widely expand the applicability of such materials. The presentation will focus on 3 recent developments:

1) The surface modification of H-terminated Si has functioned as a stepping-stone for the construction of a wide-range of hybrid materials. Recent synthetic improvements and detailed mechanistic studies have shown how to optimally construct such modified substrates. These results will be presented in the light of further developments.

2) Silicon nitride can be modified by covalent SAM attachment to achieve chemically highly robust systems. Biofunctionalization thereof with oligosaccharides and antibodies provides novel diagnostic applications in terms of specific bacterial capture in combination with generally antifouling substrates when combined with lithographic structuring of the material. Proof-of-principle and remaining challenges will be demonstrated based on recent experimental progress.

3) A photochemical method we recently developed to modify glass (SiO2), and the application thereof within the field of modified glass microchannels will be shown, e.g. via the on-chip embedding of enzymatic cascade reactions using a combination of local SAM attachment and DNA-enzyme hybrids. Finally, generalizations to the modification of other oxide surfaces will be provided.

Key references:

2:40pm TF+EM+SS-ThA3 Molecularly Modulated Electrical Transport at Chemically Passivated Silicon Surfaces, G. Dubey, NRC-SIMS, Canada, F. Rosei, INRS-EMT, Canada, G.P. Lopinski, NRC-SIMS, Canada

Adsorption of charged or polar species on semiconductor surfaces can modulate the electrical properties through long-range field effects. Hydrogen-terminated silicon-on-insulator (SOI-H) is an interesting model system for investigating this sensitivity to surface processes [1,2]. Accumulation mode SOI-H pseudo-MOSFETs [3] have been used to probe molecular adsorption and reaction events. Current-voltage characteristics of such n-channel devices are found to be sensitive to the environment, with the accumulation threshold voltage, or flat-band voltage (V_{fb}), exhibiting large reversible changes upon cycling between ambient atmosphere, high vacuum (<10-7 Torr) and exposure to water and pyridine vapour at pressures in the torr range. Both these adsorbates act as effective electric donors, shifting the flat band potential to more negative values. The field-effect mobility is found to be comparatively less affected through these transitions. Adsorption of the well known electron acceptor tetracyanoethylene (TCNE), is shown to cause depletion, with ppm levels of TCNE vapour in ambient atmosphere found to rapidly decrease the saturation current by over two orders of magnitude. The effect is only partially reversible on the hydrogen terminated surface, due to the accumulation of strongly bound TCNE molecules on the surface. In addition, oxidation of the H-terminated surface is seen to result in irreversible shifts in both the flat-band voltage and field-effect mobility. In order to passivate the surface from these irreversible processes, a photochemical gas phase reaction [4] with dichloromethane under argon was used to form a decyloxy monolayer on the SOI(100)-H surface. Formation of this monolayer is found to result in a relatively small shift of threshold voltage and only a slight degradation of the field effect mobility. Decyl passivation only slightly decreases the response of the FET to TCNE adsorption while significantly improving the reversibility of the response. These results suggest that alkyl monolayer dielectrics formed by the gas phase photochemical method can function as good passivating dielectrics in field effect sensing applications.


3:00pm TF+EM+SS-ThA4 Micrometer- and Nanometer-Scale Patterning of Azide-Functionalized Self-assembled Monolayers on Gold and Aluminum Oxide Surfaces, O. El Zubir, I. Barlow, G. Leggett, N. Williams, University of Sheffield, UK

Self-assembled monolayers (SAMs) have been prepared by the adsorption of [11-(4-azido-benzoyl amino)-undecyl] phosphonic acid on aluminum oxides and by the adsorption of [11-(4-azido-acrylamino)-undecyl] phosphonic acid on gold. Exposure to near UV radiation causes elimination of nitrogen and the creation of a reactive nitrone species. Quantitative studies by X-ray photoelectron spectroscopy (XPS) and contact angle measurements show that the reaction is complete after an exposure of 20 J cm–2. The reaction enables selective introduction of primary amines to the surface. By exposure of the azide under a solution of the amine, derivatization may be carried out with high efficiency, providing a simple pathway for the creation of a variety of multiple-component surface chemical patterns. Micropatterned surfaces were fabricated by exposure the SAMs to UV-laser (λ=325 nm) through a mask in presence of amines. Photopatterning was carried out at the nanomater scale by using scanning near-field photolithography (SNP) in which a scanning near-field optical microscope (SNOM) coupled to a UV laser (λ=325 nm) is utilized as the light source. Selective modification of azide terminated monolayers on aluminum oxide by protein-resistant molecules enabled the fabrication of protein nanopatterns that could be imaged by fluorescence microscopy.

3:40pm TF+EM+SS-ThA6 Free-standing, SAM-based, Hybrid Biocompatible Nanomembranes for Biological Applications, N. Meyerbroeker, W. Eck, M. Zharnikov, Universitat Heidelberg, Germany

Functional monomolecular films - so-called self-assembled monolayers (SAMs) - represent a broad platform for nanofabrication, sensor design, and molecular electronics, as well as a framework for Chemical Lithography, and a model system for organic interfaces and molecular, macromolecular and biological assemblies. These films can be prepared on different substrates which provide necessary support and guarantee the persistence of the 2D molecular interaction. Since the intermolecular interaction within the SAMs is relatively weak, such films cannot exist without support, dissipating into the individual constituents upon the separation from the substrate. However, as far as the individual molecules within a SAM can be cross-linked extensively by physical means such as electron irradiation, the resulting quasi-polymer film can be separated from the substrate and exist as a free-standing monomolecular membrane on its own. In this presentation we describe the formation of such free-standing membranes on the basis of aromatic SAMs. In contrast to the previous work, we used not only non-substituted but nitro- and nitrile-substituted SAM constituents which, due to either nitro-to-amine or nitrile-to-amine transformation, become chemically reactive after the irradiation treatment and can be subsequently coupled to further species. As such species we used oligo(ethylene glycol)-based molecules with a specific anchor group providing the coupling to the amine groups of the monomolecular template. After the attachment, these templates and respective hybrid membranes

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become protein-repelling and as such can be used as a non-disruptive and highly transparent support for proteins and cells in transmission electron microscopy (TEM) experiments. Whereas the ultimate thinness of this support guarantees a high imaging quality, protein-repelling ensures the lack of protein denaturing, which extends essentially the possibilities of TEM experiments in their specific application to sensitive biological targets.

4:00pm TF+EM+SS-ThA7 Development of Nanoscale Heterostructures: From Single Component Nanostructures to Multicomponent Nanosystems, N. Chope, The University of Alabama, Tuscaloosa

Chemical vapor deposition (CVD) and solution synthesis was coupled to realize a unique surfactant-free approach for the direct nucleation of nanoparticles on 1-D nanostructures. Towards this end, skeleton nanostructures such as carbon nanotubes and oxide nanowires were successfully utilized as a selective nucleation sites for nanoparticles. Carbon functional groups to chelate with metal ions [1], and thus can incorporate d-orbital transition metals with the SAMs. One challenge in molecular electronics is the reliable formation of a top contact to the organic layer. Fabrication techniques involving metal evaporation of the top contact often result in penetration to the substrate [2] or into the SAM [3], either of which influences the measured electrical properties across the junction. Here, we utilize flip chip lamination (FCL), a soft metallization technique by nanotransfer printing, to form a top contact onto SAMs on Au [4]. Based on previous work by Ulman et al. [5] and Allara et al. [6], we have incorporated metal ions (Cu and Ni) with mercaptohexadecanoic acid (MHA) and formed a molecular junction by FCL. SAMs of MHA were deposited on Au on Si and Au on polyethylene terephthalate (PET) substrates. The MHA/Au/Si was exposed to metal ion (M) containing solution. The M/Au/Si samples were then laminated to MHA/Au/PET to create a ‘molecular sandwich’ which resulted in the following structure: PET/Au/MHA-M-MHA/Au/Si where the PET substrate is removable.

The SAMs on Au were investigated by using X-ray photoelectron spectroscopy (XPS) and p-polarized reflection absorption infrared spectroscopy (p-RAIRS), and both methods confirm the initial presence of carboxylic acid on the (pre-FCL) surface. Successful incorporation of metal ions within the SAMs was directly confirmed by XPS (occupying both the −COOH sites, and indirectly by p-RAIRS with the appearance of C=O bands in an acid salt environment. Electrical and physical characterization (using backside p-RAIRS and near edge X-ray absorption fine structure (NEXAFS)) measurements to investigate the monolayer after FCL are currently ongoing. With these results, we are able to obtain a thorough picture linking electrical properties with physical characterization of the buried molecular junctions. [1,2]

References

5:00pm TF+EM+SS-ThA10 Examining the Role of Laminin-Derived Peptides in Neural Cell Attachment, N.A. LaFrang, J.A. Maurer, Washington University in St. Louis

Laminin-1 is a 900KDa glycoprotein that is a major component of the extracellular matrix (ECM). Laminin is known to be involved in many cellular processes including cell adhesion, migration and differentiation in vitro and is often used in in vitro experiments to encourage neuronal attachment. While laminin contains some integrin-receptor binding sites, multiple domains on the protein including the peptide sequences iKVAV, YIGSR and RGD have also been shown to bind to non-integrin neuronal receptors, encourage cell adhesion, and encourage neurite outgrowth in the absence of the full protein structure. However, previous experiments performed using these peptides as substrates for neuronal culture fail to consider the effects of excretion of the cells own ECM proteins onto the surface, which we have shown can occur on surfaces with high peptide concentrations. By utilizing self-assembled monolayers (SAMs), we have created a substrate that prevents the non-specific adsorption of proteins excreted by neurons, while introducing a low-concentration of the peptide sequences to encourage neuronal attachment.

A pair of alkanethiol monomers for assembly on a gold substrate have been designed which contain a tetra-ethylene glycol moiety at the tail to prevent the non-specific adsorption of proteins. One of the monomers is terminated with an azide moiety that can be used to specifically attach molecules with an alkyne group by a copper-mediated azide-alkyne cycloaddition (CuAAC) reaction, also known as the “click” reaction. By introducing 5-pentynoic acid at the N-terminus of each of the peptides during synthesis, we have introduced a bioorthogonal attachment point for reaction with the azide-terminated monomers on the surface. Once prepared, these surfaces have been confirmed to be resistant to adsorption of proteins by quartz-crystal microbalance (QCM) experiments. Primary culture neurons dissected from embryonic mice have been cultured on the surface and the effects of the peptides on neurite outgrowth have been quantified. To better understand the role of these peptides in cell adhesion, neurons transfected with a plasmid encoding EGFp-vinculin or EGFp-paxillin have been cultured on the surface and the focal adhesion morphologies have been observed in live-cell imaging experiments. These results are compared to those prepared for a surface where the full laminin protein has been adsorbed. Together, these experiments help to design a surface resistant to non-specific protein adsorption that allows for neuronal adhesion, sheds light on laminin-mediated attachment, and aids in developing better small-molecule mimics of receptor-mediated cellular adhesion.

5:20pm TF+EM+SS-ThA11 Contact-Free Electrical Characterization of Molecular Layers using CREM, H. Cohen, Weizmann Inst. of Science, Israel

While generally recognized as a powerful analytical tool, XPS is insensitive to hydrogen atoms and, in many cases, to fine variations in the environment of carbon atoms. As such, XPS characterization of organic molecular layers is rather limited, in particular under demands for high overlayer quality. On the other hand, a recent XPS-based technique for chemically resolved electrical measurements (CREM) proposes high sensitivity to even small amounts of defects or organization imperfections. [1,2]

Here, CREM application to monolayers self-assembled on metallic or semiconducting substrates is overviewed, demonstrating some of the unique capabilities offered by this non-contact probe. Damage evolution under irradiation is specifically discussed; two of its extreme limits being modeled, yielding the effect of corresponding defect sites on the CREM-derived I-V curves. Potential applications to molecular electronics, approaching atomic resolution in the electrical data, will be discussed.

References

Triboology Focus Topic
Room: 111 - Session TR-ThA

Advanced Tribological Materials
Moderator: S. Perry, University of Florida

2:00pm TR-ThA1 Nanomechanics and Nanotribology of ZrB2 Thin Films Deposited by DC Magnetron Sputtering, E. Broitman, H. Högborg, L. Hultman, Linköping University, Sweden

The microstructure, nanomechanical and nanotribological properties of ZrB2 thin films grown by DC magnetron sputtering have been studied as a
function of Ar pressure, substrate bias, and substrate temperature. Films, ~500 nm thick, were deposited onto Si (001) and Al2O3 (0001) substrates from a compound target using an industrial chamber CC-800/9 from CemeCon operated at a fixed target-to-substrate distance of 7 cm. X-ray diffraction patterns show that 0001-oriented films can be obtained on both substrates at a substrate bias of ~80 V without any external heating. Transmission electron microscopy of samples grown at different conditions reveal the presence of an amorphous 100-300 nm thick layer close to the substrate, followed by the nucleation of ZrB2(0001). The same oriented structure appears for samples grown up to 150 °C, but at higher temperatures this phase is gradually recovered. At 500 °C, cross-sectional scanning electron microscopy shows a columnar microstructure with re-nucleation during the growth. For films grown at 100 °C, little impact on the texture is observed when the substrate bias is changed from floating to ~200 V.

Nanomechanical and nanotribological properties measured with a Hysitron Triboindenter 11 TI 950 reveal that the films have high hardness and elastic recovery, and low friction. For films grown at low temperature, the hardness, reduced elastic modulus, and elastic recovery decrease from 25 to 19 GPa, 290 to 200 GPa, and 96 to 92%, respectively, when the amorphous interface increases from 100 to 300 nm. Nano-frictional tests were done in a load-controlled feedback mode using a force of 1 mN; a total of 40 reciprocating passes were performed for each test using a diamond 90° probe with a 1 μm tip radius. The friction tests reveal a friction coefficient µ in the range 0.10-0.13 for ZrB2 samples grown at different conditions, in contrast of µ = 0.6 for a pure Zr film.

2:20pm TR-ThA2 Structural and Tribological Properties of CrNMPP/TiNDCMS Multilayer Coatings, J. Paulitsch, C. Maringer, D. Holzer, P.H. Mayrhofer, Montanuniversität Leoben, Austria
Deposition processes like the high power impulse magnetron sputtering (HIPIMS) indicate high metal ion ratios in the plasma, which result in increased structural and mechanical properties. The galvanic CrNMPP/TiNDCMS multilayer deposition technique, compared to direct current magnetron sputtering (DCMS), narrows the industrial application range of this technology. The modulated pulse power (MPP) deposition technique on the other hand uses multiple complex pulsing steps to increase the metal ion ratio in the plasma without dramatically reducing the deposition rates as compared to DCMS.

Recently we showed that a multilayer architecture of CrN and TiN, deposited using the hybrid HIPIMS/DCMS deposition technique, results in coatings exhibiting high friction coefficients in the range 0.14 for diamond-like carbon (DLC) coatings when tested at RT and ambient air conditions. Here we show results of MPP/DCMS deposited CrN/TiN multilayer coatings indicating comparable mechanical and tribological properties, hardness values around 25 GPa and coefficient of friction below 0.05. Furthermore, investigations on their dependence to the atmospheric conditions used during dry sliding as well as theoretical investigations of the layered structure using density function theory simulations were carried out.

2:40pm TR-ThA3 Tribological Properties of Plasma Electrolytic Oxidation (PEO) Coatings on an Aluminum A356 Alloy, J.F. Su, X. Nie, University of Windsor, Canada
To reduce the fuel consumption and pollution of passenger vehicles, the aluminium engines have been increasingly used throughout the last 30 years. Since most technical aluminium alloys provide only poor wear resistance, various technical solutions exist to generate a wear-resistant cylinder bore surface against the sliding piston ring. A Plasma Electrolytic Oxidation (PEO) process has been developed in our group to produce oxide coatings on an Al alloy A356 for Al engine block, to battle against the wear attack. Primary results showed that the PEO coatings, thinner than 8 μm, are promising candidates to resist wear at elevated temperature. In this work, further study was carried on a serial of PEO coatings on the A356 alloy. The microstructure, coating thickness and tribological properties were tailored by adjusting the PEO process parameters. The tribological performance of the PEO coatings was better than a Plasma Transferred Wire Arc (PTWA) coating which is currently used for engine applications. Based on this study, selections on optimal thickness and morphology of PEO coatings for better wear resistance were proposed.

3:40pm TR-ThA6 Tribochemical and Microstructural Evolution during Friction and Wear of Nanocoating Coatings, J. Tschirch, The University of North Texas
Friction and wear mitigation is typically accomplished by introducing a shear accommodating layer (e.g., a thin film of liquid) between surfaces in sliding and/or rolling contacts. When the operating conditions are beyond the liquid realm, such as in extreme environments, attention turns to solid coatings. The focus of this talk is how contacting surfaces and subsurfaces change both structurally and chemically in order to accommodate interfacial shear for two multifunctional coating systems: nanocomposite MoS2/Sb2O3/Au and Ni/TiC/graphite. It was determined that the coatings exhibit velocity accommodation modes (VAM) of interfacial sliding and intergranular shear, as determined by advanced electron microscopy (3-D focused ion beam serial cross-sectioning, HAADF-STEM, and HRTEM) and spectroscopy (Raman, Auger and EDS wear maps) techniques.

In the case of amorphous-based MoS2/Sb2O3/Au nanocomposite sputtered coatings, the main mechanism responsible for low friction and wear in both dry and humid environments is governed by the interfacial sliding between the wear track and the friction-induced transfer film on the counterface ball. In dry environments, the nanocoatings have the same low friction coefficient as that of pure MoS2 (~0.007). But unlike pure MoS2, coatings which wear through in air (50% RH), the composite coatings showed minimal amount of wear with wear factors of ~1.2-1.4 × 10^-5 mm^3/Nm in both dry nitrogen and air. Cross-sectional TEM of wear surfaces revealed that frictional contact resulted in amorphous to crystalline transformation in MoS2 with 2H-basal (0002) planes aligned parallel to the sliding direction. In a humid wear environment and subsurface regions exhibited islands of Au. The mating transfer films were also comprised of (0002)-oriented basal planes of MoS2 resulting in predominantly self-mated ‘basal-on-basal’ interfacial sliding, and thus low friction and wear.

In the case of laser deposited Ni/TiC/graphite composite coatings, it was determined during sliding that a wear-induced tribochemical and structural change from microcrystalline graphite to amorphous carbon/nanocrystalline graphite hybrid layer resulted in decreased friction and wear. Other novel insights of determined friction mechanisms, such as a mechanically mixed layer developed consisting of predominately refined nanocrystalline Ni grains (~10 nm grain size) and disordered carbon below this hybrid layer. The formation of these low interfacial shear strength films and recrystallized zones were responsible for intrafilm shear VAM to achieve low friction coefficients (~0.09) and wear factors (~6.8 x 10^-6 mm^3/Nm).

4:20pm TR-ThA8 Study of Failure Mechanisms of a PVD TiAlN Coating by an Impact-Sliding Tester, J.F. Su, X. Nie, University of Windsor, Canada
Physical vapor deposition (PVD) coatings usually have high hardness and wear resistance and have been considered as necessary top layers of a wide variety of mechanical components to battle the wear problems. One of applications of hard PVD coatings is used as much-needed protective top layer on the surfaces of stamping dies thereby to extend the tool life and improve the quality of the stamped products. Impact fatigue tests have been carried out to investigate the failure behavior of coating-substrate systems under simulated stamping force conditions in our group. However, machining forces on the work pieces are not always only perpendicular to the surface. Tangential or parallel forces are usually involved such as in stamping, milling and turning. In this work, an impact-sliding fatigue tester was proposed as an experimental technique to investigate the failure behavior of coating-substrate systems under shear combined conditions. Each impact-sliding cycle consisted of a 200 N impact force and a 200 N pressing force, respectively. One PVD TiAlN coating on D2 substrates was tested at this combination of impact/pressing loads for 1,500 cycles. Impact-sliding wear track was then observed at cross section obtained after conventional and microdischarge machining (EDM). Coating failure mechanisms were found to be crumbling, peeling, fatigue cracks and material transfer. Fatigue cracks were particularly discussed from the point of view of fracture toughness.

As mechanical devices have shrunk to microscopic sizes, the need for a more fundamental understanding of friction and other surface phenomena has become urgent. While the emerging technology of microelectromechanical systems (MEMS) shows promise as the mechanical counterpart to integrated circuits, progress remains slow as structural materials and lubricant strategies continue to be developed. We report on the results of a collaborative effort to study the frictional properties of organic monolayers deposited on metal oxide surfaces. These interfacial systems have the potential to offer an alternative to silicon-based device fabrication. Both a nanoindentor-quartz crystal microbalance (NI-QCM) as well as an atomic force microscope (AFM) in lateral force mode have been used to perform tribological experiments at sliding velocities spanning the range from microns per second to meters per second. Our studies have investigated two different self-assembled monolayers chemisorbed onto platinum oxide surfaces with realistic contact roughnesses and sizes: octadecylphosphonic acid (ODP) and octadecyloctachlorosilane (OTS). Both monolayers are observed to exhibit substantially reduced friction as
compared to the bare interface, at both low as well as high sliding speeds. However, the films appear to fail upon exceeding a threshold contact pressure. We compare the tribological responses of the bare and monolayer coated interfaces of different systems and discuss insights into the molecular-level mechanisms responsible for the observed behaviors.

5:00pm  TR-ThA10  Nanotribological Characterization of Percolating Lead Films Above and Below $T_c$, K. Stevens, J. Krim, North Carolina State University

Friction at the nanoscale shows a strong and complex relationship to surface roughness and atomic disorder [1]. Recent research in superconductivity dependent friction [2-5], along with reports that quantum size effects [6] can influence diffusion (and thus friction) of adsorbed layers, has motivated our investigation. In particular, we have performed friction measurements of adsorbed nitrogen and helium films sliding on nanostructured lead films substrates that have been deposited on titanium, a substrate that lead does not wet. Varying the lead coverage results in a spectrum of percolated morphologies. We prepare these films on a quartz crystal microbalance (QCM) and probe their topologies by means of adsorption onto the surface [7].

Measurements have been recorded on nanoclustered lead films with coverages crossing the critical concentration for percolation. We study the substrate in the superconducting and normal states, which allows us to isolate and quantify the contribution of electronic and phononic dissipation to the total friction present [2]. Submonolayer adsorbate coverages have allowed us to probe the edge effects of surface nanoclusters, while multilayer coverages have let us explore the strength and proximity effects of surface roughness. We compare our measurements to those reported by Pierno et al. on films of ordered Pb(111) terraces, where atomic step edges are present [3], and conclude that the variation in reported values of friction on nanostructured lead is due to phononic effects at the step edges.

Funding provided by NSF DMR.


5:20pm  TR-ThA11  Advanced SP3EC Carbon Nanocomposite Coatings, J. Larson, United Protective Technologies

United Protective Technologies (UPT) has developed a room temperature plasma assisted chemical vapor deposition (PACVD) coating process to build carbon based coatings. Versions of this coating are in use for infrared optical applications, galvanic corrosion barriers and tribological modifications to critical components. The process used to produce SP$^3$EC™ coatings is compatible with a wide range of materials including semiconductors, metals, polymers and composites. Deposition parameters for the SP$^3$EC™ process can be controlled to produce a wide range of carbon based thin films, ranging from 120 nm diamond crystalline grains to low friction glassy amorphous carbon films. Layered “nano-composite” structures constructed of these films have been proven to improve the life, performance and reliability of components under high wear conditions and corrosion conditions. Additionally, the SP$^3$EC™ process is non-toxic and environmentally friendly.

Current applications for SP$^3$EC™ coatings include targeting optics with improved durability for the AH-64 along with corrosion and wear coatings for UH-60 rotor components. Tribological coatings for improved efficiency of helical gears have been developed in conjunction with advanced wear coatings for aluminum and steel components. This presentation will include details on the mechanical and barrier characteristics of SP$^3$EC™ coatings for various applications, current process capabilities and developmental applications of this coating process.
A commercially available amino-terminated PDMS was fractionated by size exclusion chromatography (SEC) in the molecular weight from \( M_n = 7000 \) to 100,000 with narrow molecular weight distributions \((M_w/M_n=1.2)\), and transformed to a triethoxysilane (TES) end-functionalized PDMS on both chain ends. The silicon wafer was immersed in dichloromethane solution of the telechelic TES-terminated PDMS at 293 K for 24 h to immobilize both chain ends of the PDMS. The surface morphology of the telechelic PDMS-immobilized silicon wafer was observed by atomic force microscopy (AFM) under air at room temperature. The root mean square (rms) of the surface roughness was 0.6 nm in a 2 x 2 \( \mu \text{m}^2 \) scanning area. The smooth morphology indicated that loop structure was formed without cross-linking and gelation of TES. Mono-functionalized PDMS was also synthesized to fabricate a linear brush on the silicon wafer. The thickness of these brushes in a dried state determined by ellipsometry increased from 2 to 5 nm with an increase in the molecular weight of \( M_n = 5000 \) to 100,000. The loop and linear polymer brushes with same molecular weight showed the almost same thickness in a dried state. Apparent difference derived from a topology effect was not observed in a dried state. AFM measurement of the swollen thickness of the loop and linear brushes in a solution is in progress.

Polylactic acid (PLA) was chosen as the scaffold material due to the fact that it has most of the desired TE scaffold properties. A PLA nonwoven web from Ahlstrom Nonwovens was used as the material. Surface modification was performed by thermally initiated free radical polymerization using vinyl monomers such as maleic acid and maleic anhydride. Potassium persulfate was used as the initiator and Mohr’s salt was added to reduce the extent of homopolymerization. The grafted surfaces were coated with a bioactive coating of collagen using a spacer molecule called genipin. The grafted surfaces were then evaluated via Fourier transform infrared spectroscopy (FTIR), contact angle measurements, X-ray photoelectron spectroscopy (XPS), and dyeing with a basic dyestuff and using visible spectrophotometry. The FTIR spectra show the grafting of carboxylic acid groups of maleic acid onto the surface of the PLA. The effect of monomer concentration is seen in the contact angle measurements. With increase in the monomer concentration, the contact angle fell, indicating an increase in the hydrophilicity of the material. The collagen was also successfully coated onto the grafted surface which is confirmed by the FTIR spectra showing the characteristic amide bands. Further evaluation of the surfaces is continuing with dyeing, XPS, and measuring the biological performance. In addition, other surface modification methods like plasma, atom-transfer radical polymerization (ATRP), are being undertaken.

Development of flexible coating strategies for angiogenesis promotion is critical for the health care industry to effectively treat chronic non-healing wounds. This need will continue to intensify in light of the increasing number of patients diagnosed with diabetes and an aging population. In addition, such strategies are required within the tissue engineering community to overcome issues associated with engineered materials failing to engraft as a result of delays in neovascularisation. An important requirement for a pro-angiogenic biomaterial is the ability to maintain a regulated release of bioactive growth factors to the wound site through the use of heparin. A number of strategies for the use of natural and synthetic heparin-mimetics have been developed, however in general they fail to provide a consistent method to immobilize heparin to the wound site. Therefore, new strategies to effectively immobilise heparin for the release of bioactive growth factors are being developed. This work describes a layer-by-layer (LBL) approach that provides a platform for immobilising a significant amount of heparin to the substrate of choice. The LBL coating prevents desorption of immobilised heparin as a result of ‘covalent bond-like’ interactions from the electrostatic attraction between the multiple layers of polyelectrolytes. Employing plasma polymerisation, a thin functional coating was applied to the substrate to provide the necessary surface charge in order to build the LBL architecture. In this particular case, plasma polymerised acryl acid (ppAAc) was deposited onto Si wafer and two different tissue scaffolds, a commercially available polypropylene type, and an electrosprun PLGA (75:25) type previously reported in the literature. The LBL structure was then created by physically absorbing alternative layers of polyethylenimine (PEI) and poly(acyrylic acid) (pAAc), and the desired number of layers were achieved, heparin was immobilised to the structures. The kinetics of the fabrication procedure was examined using a quartz crystal microbalance with dissipation (QCM-D) system, while the resulting structures were probed using X-ray Photoelectron Spectroscopy (XPS) and Atomic Force Microscopy (AFM). The effect of pH of the LBL solutions (PEI and pAAc) on immobilising structure and potency to immobilise certain biologically active molecules were examined. This technique allowed for the immobilisation of a significant amount of heparin to the test substrates explored, more so than that previously reported using similar approaches. This approach therefore provides an inexpensive and easily expandable coating methodology for applying a pro-angiogenic interface to tissue engineered materials.
time. The mass increase versus concentrations of dNTP allows us to determine the reaction constant, which reflects the growth kinetics of DNA polymerization on the surface. In addition, we examined the effect of grafting density on the polymerization reaction and the conformation of DNA brushes.

Bi-Thp5 Plasma-based Approaches for Biointerface Preparation, E.H. Lock, S.H. North, S.G. Walton, C.R. Taitt, Naval Research Laboratory (NRL)
The ability to predict and manipulate biomolecule behavior at the biointerface determines the success of biomaterials in applications ranging from biosensing to medical devices and therapeutic products. However, precise biointerface engineering will remain elusive until the roles of physical and chemical properties of surfaces on abiotic and biotic interfacial interactions are well understood. We have shown that plasma treatment of polymers generates chemically reactive surfaces for successful silanization and biomolecule immobilization [1, 2]. The focus of this work is to investigate the influences of surface chemistry and surface morphology on biomolecule attachment. We fine-tune our plasma system to favor the production of specific functional groups that promote subsequent biomolecule attachment. The effects of surface morphology on biomolecule immobilization are also assessed. Plasma diagnostics and modeling allows us to elucidate the effects of plasma parameters (plasma density, electron temperature and the resulting kinetic energy) on the polymer modifications. The work was supported by the Office of Naval Research.

References:

Bi-Thp6 A Novel ALD Al2O3-Parylene Bi-Layer Encapsulation for Biomedical Implantable Devices, X. Xie, L.W. Rieth, F. Soltzbarber, University of Utah
Atomic layer deposited (ALD) Al2O3 has been widely used as encapsulation material for organic LEDs and solar cells due to its low water vapor transmission rate (WVTR) (~5×10^-5 g·H2O/m²·day). However, its coating performance for implantable devices still needs investigation. Parylene has been commonly applied as encapsulation for implantable devices, such as Utah Electrode Arrays (UEAs). The idea of combing Al2O3 and parylene is based on the concept that Al2O3 works as moisture barrier and parylene as ion barrier. In this paper, Al2O3 was deposited by both thermal and plasma-enhanced ALD on interdigitated electrodes (IDEs) for comparison. AFM micrographs (Fig. 1) show that Al2O3 films deposited on silica substrate (RMS surface roughness of 0.17 nm) by thermal and plasma-enhanced ALD have RMS surface roughness of 0.51 nm and 0.48 nm, respectively. XPS shows that Al2O3 films had an oxygen to aluminum ratio of 1.2 while thermal ALD Al2O3 is 1.09, indicating that the former is closer to Al2O3. A 6-μm thick parylene-C layer was deposited by CVD using Gorman process on top of Al2O3 and saline A-174 (Mentivaent Environmental Materials) was used as adhesion promoter. The samples were soaked in 1× PBS at 37 °C and 57 °C for accelerated lifetime test. Electrochemical impedance spectroscopy (EIS) and chromoanometry were used to evaluate the performance of the encapsulation. Preliminary data shows that the leakage current (Fig. 2) remained very small (~7 pA) and the electrochemical impedance (Fig. 3) was consistently high (~3 MΩ at 1 kHz) after 5 days of soaking test at 57 °C (equivalent to at least 20 days of soaking test at 37 °C). Comparing with parylene and Al2O3 control samples, the Al2O3-parylene coated sample showed lower leakage current (Fig. 2). The impedance for three different types of samples was almost the same. However, the phase of parylene-C coated sample slightly declined after 5 days of soaking test (Fig. 3), suggesting that sample coated with Al2O3 had lower WVTR. No obvious difference has been observed yet for samples soaked at different temperatures since the soaking period is relatively short. Parylene loss one tenth the day of soaking test for one sample, which might be caused by the poor adhesion between Al2O3 and parylene. In conclusion, preliminary results shows that the Al2O3-parylene bi-layer encapsulation scheme is promising encapsulation in terms of leakage current and electrochemical impedance. Long-term soaking tests are being performed to further investigate the functionality of this novel encapsulation scheme.

Bi-Thp7 Complementary Electronic and Vibrational Circular Dichroism Analysis of Bovine Serum Albumin Adsorbed on Hydroxyapatite Microspheres, K.P. Fears, Naval Research Laboratory, D.E. Day, Missouri University of Science and Technology, D.Y. Petrovsky, International Iberian Nanotechnology Laboratory, T.D. Clark, Naval Research Laboratory
Bovine serum albumin (BSA) is a widely studied globular protein that contains ca. 68% of alpha-helices and <2% of beta-sheets in its native conformation. The well characterized secondary structure of BSA is commonly used as a benchmark for electronic (ECD) and vibrational (VCD) studies of proteins in solution. Both ECD and VCD indicated a substantial loss of helical structure accompanied by an increase of beta-sheet character in BSA thermally denatured in solution. In surface adsorption experiments, hydroxyapatite microspheres were incubated in solutions with low (1.0 mg/ml) or high (50.0 mg/ml) concentrations of BSA for 10 hours, then tripod rinsed and re-suspended in buffer for analysis. The ECD spectra were similar for BSA adsorbed from low and high concentration solutions, both showing a sizeable increase in beta-sheet character upon adsorption, while being dominated by alpha-helical features. The VCD spectra also exhibited stronger peaks in the beta-sheet region upon adsorption of BSA on hydroxyapatite. VCD signal enhancement, however, was observed upon adsorption from the high concentration BSA solution, indicating the formation of macroscopic chiral structures. The analysis of proteins adsorbed on surfaces thus can be enhanced by taking advantage of the complementary sensitivities of ECD and VCD spectroscopies to the secondary structures of biomolecules.

Bi-Thp9 Binding Affinities of wt and H93R PTEN to Lipid Membranes Containing PS and PI(4,5)P2, S. Shenoy, Carnegie Mellon University, Gericke, Kent State University, A.H. Ross, School of Medicine, University of Massachusetts Medical School, M. Lösche, Carnegie Mellon University and National Institute of Standards and Technology
PTEN is a phosphatidylinositolphosphate (PIP) phosphatase frequently mutated in human cancer [1]. By lowering PI(3,4,5)P3 levels in the plasma membrane, it functions as an antagonist to PI3-kinase in the regulatory circuit that controls cell proliferation and survival. wt PTEN has only weak affinity to neutral zwitterionic phosphatidylcholine (PC) membranes but a strong interaction with anionic lipids. Its C2 domain was shown to bind in a Ca2+ independent manner to phosphatidylinerine (PS) and phosphatidylglycerol (PG), whereas a short N-terminal domain binds specifically to PI(4,5)P2 [2,3]. H93R PTEN is an autism related mutant which has decreased phosphatase activity [4].

Using Surface Plasmon Resonance (SPR), we characterized the affinity of wt and H93R PTEN to tethered bilayer lipid membranes (tBLMs) that contain PC and PS, PC and PI(4,5)P2, and PC, PS and PI(4,5)P2. As compared with wt PTEN, we find that the H93R mutation is sufficient to cause significant increases in the protein’s association with lipid membranes containing PS. PI(4,5)P2 enhances the apparent binding constant for both proteins and leads to intriguing binding kinetics of the protein to the membrane. The binding of either protein to membranes containing both PS and PI(4,5)P2 shows a biphasic behavior, suggesting two independent binding sites. This supports the hypothesis of non-competitive binding of the protein to PS and PI(4,5)P2 [5]. We also performed neutron reflectivity experiments to determine the structure and orientation of PTEN bound to the membrane.


Bi-Thp10 Membrane Binding and Structure for Cytoplasmic Domain of Zeta Subunit of T Cell Receptor, P. Shekhar, Carnegie Mellon University, F. Heinrich, Carnegie Mellon University and National Institute of Standards and Technology, K. Zimmerman, University of Massachusetts Medical School, M. Lösche, Carnegie Mellon University and National Institute of Standards and Technology, L.I. Stern, University of Massachusetts Medical School
The cytoplasmic domain of the T-cell receptor zeta subunit, ζζ, a cell surface protein complex responsible for binding peptide fragments of foreign antigens bound to major histocompatibility complex (MHC) proteins, is sufficient to couple receptor ligation to intracellular signaling cascades [1]. These domains carry immunoreceptor tyrosine-based activation motifs (ITAMs), i.e. signaling motifs that are phosphorylated by tyrosine kinases following receptor crosslinking. The phosphorylation of ITAMs is a first and obligatory step in signal transduction. ζζ has been shown to be unstructured in aqueous solution and to assume a helical
conformation in the presence of anionic lipid vesicles [2,3]. Membrane binding and membrane-induced conformational changes likely plays an important role in signal transduction, but no direct structural information on these functionally important lipid-bound states was available so far. Using a synthetic membrane model, i.e., fluid lipid bilayers tethered to planar solid supports [4,5], we report surface plasmon resonance (SPR) results on the binding kinetics and neutron reflectivity investigations of the association of the disordered ζ_s with membranes. We determine the extent to which the protein penetrates into the bilayer and discuss structural details of the ζ_s-lipid interaction.


Spectroscopic Ellipsometry Focus Topic
Room: East Exhibit Hall - Session EL-ThP

Spectroscopic Ellipsometry Poster Session


Tungsten (W) and titanium (Ti) films are widely used in electrochemistry, microelectronics, energy conversion and nanotechnology. In integrated optics, the nanometric Ti films are used as a source for doping LiNbO3 and LiTaO3 substrate and optical waveguide fabrication by thermal diffusion. Because effective refractive indices of the waveguide modes are strongly dependent on the optical profiles in doped layer, precise control of Ti film thickness (h) is needed in the range ~10-50 nm. Ellipsometry can be successfully applied for nondestructive determination of the thickness of a dielectric and semi-transparent metal film when optical constants of the material are known. Regrettably, noticeable scattering was found for optical constants reported earlier in literature for W and Ti films and crystals. As it seems, this scattering appeared due to different film quality and surface state. The focus of the present work is centered on W, Ti and W-Ti film fabrication and evaluation of their optical parameters with spectroscopic ellipsometry. Tungsten and tungsten-titanium films were prepared by magnetron sputtering deposition in vacuum below 10.5 Torr. Titanium films were fabricated by thermal evaporation method in vacuum below 10.5 Torr. The substrate temperature was T=100 °C. For precise determination of optical parameters, thick metal films (h~100 nm by as determined from optical interferometry) were prepared on silica substrate. To increase the metal adhesion, the substrate was subjected to RCA chemical cleaning just before insertion into vacuum chamber. Structural parameters of metal films were calculated with reflections high-energy electron diffraction (HEED).

Surface micromorphology was controlled with atomic force microscopy (AFM). Spectral dependencies of refractive index n(λ) and extinction coefficient κ(λ) were determined with the help of spectroscopic ellipsometry in the spectral range, λ~250-1030 nm. A relation between optical constants of pure metal W and Ti and mixed metal W-Ti films is discussed.

EL-ThP2 Temperature Dependences of the Dielectric Response of InSb Measured by Spectroscopic Ellipsometry. J.J. Yoon, T.J. Kim, S.Y. Hwang, M.S. Divhare, Y.D. Kim, Kyung Hee University, Republic of Korea, Y.C. Chang, Academia Sinica, Taiwan, Republic of China

InSb is a promising material for optical devices, particularly for high-frequency and nonlinear-optical applications. InSb has a high electron mobility and offers excellent design flexibility as a result of its large conduction band offset in heterostructures. Consequently, InSb offers significant potential for devices such as quantum-well lasers, laser diodes, and heterojunction bipolar transistors. A knowledge of the dielectric function at various temperatures is required for optimizing the properties for specific device applications. In-situ control of growth is also becoming an important technique. Therefore, the dielectric function at growth temperatures is also needed. On the other hand, critical point (CP) energies can be better identified from low-temperature data, where the decreased electron-phonon interaction allows separation of CP structures that are nearly degenerate at room temperature.

Although the optical properties of InSb have been well studied, there are only a few reports of their temperature dependence in the 1.2 to 5.6 eV spectral ranges [1]. Here, we report results of an investigation of the temperature dependence of the dielectric response of InSb from 22 K to 700 K and from 0.74 to 6.57 eV.

Spectroscopic ellipsometric (SE) data were obtained on a bulk semiconducting InSb (100) substrate. The cryostat consisted of a stainless-steel chamber with high-quality stress-free fused-quartz windows. To avoid condensation at low temperatures, the sample was maintained in ultrahigh vacuum during measurement. SE data were obtained at an angle of incidence of 70.41° using a conventional rotating-compensator system with a diode-array detector. The influence of the oxide layer was removed mathematically by a multilayer calculation. In the E1 energy region only the four structures are clearly resolved at 300 K. However, at 25 K the E1 and E2 structures are seen to consist of five CPs. We identified the origin of these structures with band-structure calculations using the LASTO method. Separation of the E1', E2'+Δ1, E3, E5+Δ2, E6', E1', E1' +Δ1 CPs was clearly found in the region of the E2 peak. Two saddle-point transitions, Δ1-Δ1* and Δ1'-Δ1*, are clearly seen. We also determined the temperature dependences of the newly observed transitions near 5.9 eV. These results will be useful in a number of contexts, including the design of optoelectronic devices based on InSb, as data for improved band structure calculations, and for in-situ monitoring.


EL-ThP3 Tailored Helical Nanostructures Investigated with Mueller Matrix Ellipsometry. R. Magnusson, J. Birch, C.L. Hsiao, P. Sandström, H. Arwin, K. Järenäk, Linköping University, Sweden

Metamaterials showing chiral features in the optical spectral range have been fabricated with the aim to obtain polarized reflection with high ellipticity.

A series of tailored anisotropic and transparent structures of helical Al-xIn_xSb nanorods were grown using UHV magnetron sputtering on sapphire substrates. Due to an internal in-plane composition gradient across the crystalline structure, the nanorods will tilt relative to the substrate normal. By rotating the substrate step-by-step around its normal during deposition ‘staircase’ helical structures are obtained. The layer thickness for each step is controlled to tailor the nanorods. Samples with different pitch and layer thickness and with right-handed as well as left-handed chirality were grown.

Ellipsometric measurements were performed using a dual rotating compensator ellipsometer providing the full Mueller matrix in the spectral range 245-1700 nm at multiple angles of incidence and 0-360° sample orientation. The relation between the optical characteristics of the samples, specifically the ellipticity, and structural parameters such as number of layers, layer thickness and nanorod pitch of the samples, was studied. For certain wavelengths, near circular polarization is observed both for right- and left-handed chiralities.

Based on the Mueller matrix data, descriptions of the polarization states and degree of polarization in reflection for different incoming states of polarization will be presented.
Electronic Materials and Processing Division
Room: East Exhibit Hall - Session EM-ThP

Electronic Materials and Processing Poster Session
EM-ThP1 Novel Quantized Edge States on Ag(111). A.R. DiLullo, Ohio University, D. Acharya, Pacific Northwest National Laboratory, N. Takeuchi, Universidad Nacional Autonoma de Mexico, S. Ulloa, S.-W. Hla, Ohio University
A one-dimensional electronic edge state has previously been observed using ultra-high-vacuum low-temperature scanning tunneling microscopy (UHV-LT-STM) methods at a sample tunneling bias of approximately 4V on Cu(111) and explained in terms of an image state at the step with energy modified by a dipolar potential[1]. Presented here are new measurements using sample bias dependent UHV-LT-STM imaging techniques of step edges on a Ag(111) surface which show numerous higher order states with correlated energy spacing. Measurements are additionally made at tip induced vacancy locations (holes in the surface) to examine effects of confinement on the newly observed one dimensional states. Experimental results are supported by density functional theory (DFT) calculations, and a framework for explanation of the newly observed energy correlated one dimensional and confined states is explored.

Supporting funding is supplied in part by the grants DOE DE-FG02-02ER46012 and NSF OSIE 0730257.


EM-ThP2 Fabrication of Multilayered Bi2Te3/Bi2Te3-xSe Thin Film Cooling Devices. M. Hines, J. Lenhardt, Alabama A&M University, M. Lu, Brookhaven National Laboratory, Z. Xiao, Alabama A&M University
In this paper, we report to fabricate multilayered Bi2Te3/Bi2Te3-xSe thin film cooling devices using the microfabrication techniques. The multilayered Bi2Te3/Bi2Te3-xSe thin films will be grown using the e-beam evaporation. The in-plane and cross-plane micro cooling devices will be fabricated using the standard circuit (IC) fabrication process; pn junction diodes will be fabricated as thermometers for the measurement of temperature in the devices. The electrical and thermal properties of the e-beam-grown Bi2Te3/Bi2Te3-xSe thin films and the cooling efficiency of the fabricated cooling devices will be measured, and the measurement results will be reported in the conference. The developed devices could be a good candidate for the application of high-efficiency solid-state micro-cooling.

EM-ThP3 Solution-Based High Performance and Fully Patterned Chalcogenide Thin Film Transistors. J.J. Mejia, A. Salas-Villaseñor, A. Carrillo-Castillo, B.E. Gnade, M.A. Quevedo-Lopez, University of Texas at Dallas
In recent years, the development of novel processes for inexpensive and flexible electronics has become an increasing research area where low-cost and low temperature deposition techniques are key point to fabricate large area and flexible circuits. Here, we demonstrate fully photolithography defined thin film transistors using cadmium sulfide (CdS) and lead sulfide (PbS) as n-type and p-type semiconductors, respectively. These chalcogenides materials are deposited using chemical bath deposition (CBD) which is a low cost solution-based process that requires temperatures below 70° C. Extracted mobility for CdS was 25 cm2/V-s and 0.14 cm2/V-s for PbS. These mobilities are among the highest reported for a fully patterned TFT made with either CdS or PbS as semiconductor. The maximum temperature used in the complete fabrication process was kept below 100° C. In addition, we studied how the device performance (mobility, threshold voltage and contact resistance) is affected depending on the semiconductor thickness, thermal annealing and thermal oxidation in air.

Our fabrication approach can be integrated in complex designs such as CMOS logic gates, pixel arrays, etc., complying with all the requirements for a flexible electronics technology.

EM-ThP4 Novel Materials and Device Structures for Solid-State Charged Particle Detectors. J.W. Murphy, J.J. Mejia, B.E. Gnade, M.A. Quevedo-Lopez, University of Texas at Dallas
Detectors for charged particle spectroscopy are typically fabricated from crystalline silicon, germanium, or gallium arsenide, which are high-cost materials. However, if the purpose of the device is particle detection rather than spectroscopy, then it is possible to use less expensive materials and deposition techniques to fabricate electrically-sensing, solid-state particle detectors. Particularly, we are interested in detecting alpha particles emitted from a 210-Polonium source. In this work we investigate ZnO nanostructures in combination with poly3-hexylthiophene (P3HT) to form hybrid inorganic/organic p-n diodes which are as charged particle sensors. The ZnO nanostructures are grown from solution on a Cr electrode, and the P3HT is subsequently deposited via the drop-casting technique to achieve bi-layer film thicknesses on the order of microns. We evaluate the devices' performance in terms of leakage current, capacitance, and alpha particle detection efficiency as a function of thickness, DC bias, and annealing treatment. The structures are investigated using x-ray diffraction and cross-sectional electron microscopy. We also study the materials degradation upon exposure to radiation by monitoring the leakage current.

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EM-Thp5 Structural and Electrical Characteristics of TaN Film Deposited by DC Sputtering for MOS Capacitor and Schottky Diode Upper Electrodes, L.P.B. Lima, J.A. Diniz, State University of Campinas, Brazil. It was reported by Su et al. in (ACS Nano, 4, 5285-5292, 2010) that the use of high-temperature alcohol vapor for reducing GO increases the conductivity dramatically which improves the graphitic domains. However, details of understanding reduction mechanisms behind the interactions of alcohol molecules within the reduced defective sites of reduced GO still remains elusive.

EM-Thp7 Exploring Thermal Reduction Efficiency of Reduced Graphene Oxide with Alcohols Studied by First - Principles Calculations and Infrared Spectroscopy, R.M. Abolfath, C.G. Gong, M. Aick, Y.J. Chabal, K. Cho, The University of Texas at Dallas Graphite oxide (GO) is convenient to be used as a precursor for functionalization studies and explore the chemistry in solution. Since GO is solution-processable and hygroscopic, tuning the chemical properties by reduction thereof tailors the electronic and electric properties of thermally/chemically reduced GO. The reduction processes of GO sheets have gained much interest since improvement and systematic investigation of the graphitic structure-electrical property relationship is particularly required for graphene nanoelectronics applications. The excellent electrical conductivity of the reduced GO sheets therefore promises potential electronic applications.

EM-Thp6 TiAlO and TiAlON Obtained by e-Beam Evaporation with Additional Electron Cyclotron Resonance (ECR) Plasma Oxidation and Oxynitridation on Si for MOS Gate Dielectric, J. Miyoshi, A.R. Silva, F.A. Cavarsan, J.A. Diniz, L.P.B. Lima, State University of Campinas, Brazil. We have explored a variety of candidate interface structures and confirmed that the carbonyl formation which tends to enlarge the etch hole upon annealing can be blocked.

EM-Thp8 Characterization of Ion Implantation-Induced Vacancy Defects and Graphitization in Diamond Lattices by Coherent Acoustic Phonon Spectroscopy, J.M. Gregory, A.D. Steigerwald, Vanderbilt University, H. Takahashi, Japan Advanced Institute of Science and Technology, Japan, H.N. Toi, Vanderbilt University. We describe implantation damage experiments on single-crystal diamond samples and subsequent characterization using optical pump-probe techniques. Samples are irradiated using He+ ions with energies on the order of 1 MeV, generating lattice defects and buried layers of graphitic carbon under high pressure. Using the time-resolved optical technique known as coherent acoustic phonon spectroscopy, these samples are characterized in a depth-dependent manner. The resulting oscillation patterns reveal clues about the optical and electronic structure of the defects and the graphitic layers.

EM-Thp9 New Reconstruction Mechanism of Dense SiC(111) on Sparse Si(110) Interface, E. Abahure, University of Tokyo, Japan, J.-I. Iwata, University of Tsukuba, Japan, A. Oshiyama, University of Tokyo, Japan. SiC grown on Si substrates is a promising candidate for mass production of Graphene, an emerging material in technology. SiC(111) on Si(110) with particular alignments of the two planes offers almost perfect matching of the lattice periodicity, albeit dense SiC and sparse Si. We here report the total-energy electronic-structure calculations based on the Real-Space Density Functional Theory (RSDFT), that elucidate new reconstruction mechanism of the dense 3C-SiC(111) on the sparse Si(110) interface with the relaxed superstructures, displaying undulating structures near the interface. We have explored a variety of candidate interface structures and 241 Thursday Afternoon Poster Sessions

Reference:
reached two distinctive types called type I and II which shows bistability at the interface. In type I, the silicon-silicon interface energy is calculated as 9.89 eV and that of silicon-carbon is 10.84 eV. We argue that in both types, the silicon-silicon interface is energetically favorable compared with the silicon-carbon interface.

EM-ThP10 The Influence of the Layer Thickness on the Optoelectronic Properties of InN, M.K.I. Senevirathna, S. Gamage, R. Atalay, Georgia State University, J. Hong, Georgia Institute of Technology, N. Dietz, A.G.U. Perera, Georgia State University

Indium nitride (InN) and indium-rich group III-nitrides may have great potential for high efficient energy conversion devices such as solar cells, high speed optoelectronic devices, and various types of light emitting device structures. Scientists are exploring several different growth methods and various characterization methods to improve the material quality and to understand the structural, optical and electrical properties of InN epilayers. However, till today, the growth of high quality InN alloys and epilayers is still a challenge, mainly due to low InN dissociation temperature and due to stoichiometry instabilities at optimum growth conditions. InN epilayers exhibit significant different physical properties depending on the growth techniques (PAMBE, MBE, MOCVD, etc.) and the substrate material used. At present, low-pressure CVD based growth methods are limited to InN growth temperatures at or below 600°C, which creates problems related to a suited nitrogen precursor, since the ammonia decompositon at these growth temperatures is insufficient. To stabilize InN at higher growth temperature, we explored the growth of InN by high-pressure chemical vapor deposition (HPCVD) at 10 bar and 15 bar reactor pressures. Under these growth conditions the growth temperature can be increased to around 800 °C, resulting in improved ammonia decomposition and smaller group III/N precursor ratio. This contribution presents results on the effect of the layer thickness on the physical properties of epitaxial InN layers. All InN layers where grown on GaN/sapphire (0001) templates under identical growth conditions, only the growth time was varied. Fourier transform IR reflectance (FTIR) spectroscopy was used to analyze the film thickness and the optoelectronic layer properties. We will present results on the free carrier concentration and mobility as a function of layer thickness. The reflectance spectra were simulated using a Lorentz-Drude model and a multilayer stack model, which allows determining the phonon frequencies, dielectric function, plasma frequency, and damping parameters. From these, the free carrier concentration and mobility for each layer can be calculated. The crystalline quality of the epilayers has been characterized by XRD 2theta-omega scans and by Raman spectroscopy analysis.

EM-ThP11 The Effect of Reactor Pressure on the Optoelectronic Properties of InN Epilayers Grown by HPCVD, M.K.I. Senevirathna, S. Gamage, M. Buegler, R. Atalay, Georgia State University, J. Hong, Georgia Institute of Technology, N. Dietz, A.G.U. Perera, Georgia State University

The dependency of the optoelectronic and structural properties of InN epilayers on the reactor pressure is presented. The InN epilayers were grown by high-pressure chemical vapor deposition (HPCVD) varying the reactor pressure from atmospheric pressure from atmospheric pressure to 18.5 bar. The optoelectronic and structural properties such as free carrier concentration and mobility have been studied using Fourier transform IR reflection spectroscopy. The film thickness, growth rate, free carrier concentration and carrier mobility of the InN layer are obtained by simulating the IR reflectance spectra, using a multilayer stack structure. The bulk and surface structural properties of InN layers were held constant for ten minutes before cooling back to room temperature. Experiments were performed with increasing reactor pressure from atmospheric pressure to 18.5 bar. No significant variation of the properties of the films was observed until the samples were annealed to 600 °C. However, during isothermal heating to 650 °C or above, the evaporation of molecular nitrogen commenced around 450 °C and an abrupt increase occurred above 600 °C indicating serious degradation of the film. The stability of InN layers was found to be dependent on V/III ratio, being more stable for the film grown at higher V/III ratio. Growth rate of up to 200 ms for InN epilayers with V/III ratio of x=2. Furthermore, photoluminescence measurements and x-ray diffraction (XRD) are used to study the structural properties and the surface chemical composition using Fourier Transform Infrared spectroscopy (FTIR) and transmission spectroscopy. In order to manage the growth processes in chemical vapor deposition, determination of the crystalline quality and orientation which also affect the surface configuration. The bulk and surface structural properties of InN layers grown by high-pressure chemical vapor deposition (HPCVD) were investigated by Raman spectroscopy, x-ray diffraction (XRD), and high resolution electron energy loss spectroscopy (HREELS). HREELS spectra of InN epilayers on GaN/sapphire substrates showed surface NH species and nitrogen-termination. For this layer Raman spectroscopy and x-ray diffraction (XRD) showed c-axis film orientation. In contrast, HREELS spectra of InN grown directly on sapphire with a higher group V/III ratio of 3000 revealed that the dominant surface species is NH. For this sample, the Raman and XRD analysis indicate the existence of nitrogen-rich epilayer. Two factors suggested to account for the appearance of these tilted planes are slower surface diffusion under more nitrogen-rich conditions and strain due to the larger lattice mismatch when growing on sapphire substrates.
EM-ThP16 P-Si / N-type GaN Heterojunction. J. Kim, J. Ahn, Korea University, Republic of Korea

P-N heterojunctions from P-Si thin film / N-GaN microstructures have been demonstrated. To control the position of GaN microstructures, a mechanical method or electric-field assisted method was employed. Firstly, thermally-grown SiO2(300nm) on top of p-Si substrate was wetetched to define the mesa structures. Then, the metal was deposited on both front (on top of the SiO2) and backside. Finally, GaN microstructures were moved to form p-n heterojunction between thin film and bulk microstructure. The crystals and morphology were obtained by the co-biased, the electron luminescence from p-n heterojunction was observed. The details of the fabrication and the characterizations of P-Si thin film / N-GaN microstructures will be presented.

EM-ThP17 CIGS Thin Film Prepared by One-Step Sputtering Process by Using a Powder Target. S. Oh, G. Cho, N. Kim, Chosun University, Republic of Korea

CIGS thin film with cells reported the highest efficiency up to 20% in the thin film solar cells which is generally considered as the most promising thin film solar cell. It has been fabricated by using the three stage process in evaporation process with four sources of Cu, In, Ga, and Se pure elements. This preparation method requires very strict controls during the deposition process of CIGS thin film. Recently, there are two widely used methods for CIGS thin film fabrication: selenization with H2Se or Se after sputtering and thermal co-evaporation. Although the sputtering process was the most drawbacks including the additional manufacturing steps, higher materials costs, and the less efficiency. The co-evaporation for CIGS thin film is hard to manage the sources in sequence and to deposit over the large area. For this reason, the novel method for CIGS preparation was demonstrated by using the one-step sputtering process with the powder target. This method has advantages of the production of target, change of stoichiometry, and easy doping of impurities. The composition of the powder-sputtered CIGS thin film with 1 μm-thickness were analyzed and compared to that of the starting powder target with a composition of Cu, In, Ga, and Se at 25, 15, 10, and 50 at% by electron probe microanalysis (EPMA). The crystal structure, surface morphology, optical properties, and electrical characteristics of CIGS thin film were analyzed by using X-ray diffraction (XRD), atomic force microscope (AFM), UV-Visible spectrophotometer, and a Hall effect measurement system.


Thin-film transistors (TFTs) using amorphous oxide semiconductors (AOSs) as an active layer have shown remarkable performances including low-temperature processing, high transparency, environmental stability, and high mobility for alternative display backplanes of active-matrix organic light emitting displays (AM-OLEDs), thin-film-transistor liquid crystal displays (TFT-LCDs) and opto-electronic components. Oxide material combinations such as InZnO (IZO), InGaZnO (IGZO), ZnSnO (ZTO), and GaZnSnO (GZTO) have been vastly studied for enhancing the mobility and stability. Though the mobility of the amorphous oxide TFTs has been much improved for several years, the instability under prolonged on-current status is still critical issue to substitute for Si-based technology. Recently, other elements of high oxygen affinity such as Al, Zr, Si, and Hf were added to oxide based TFTs for improving the stability characteristics, which, however, results in a mobility reduction problem. In order to overcome the problem, we used Ta dopant. This work reports on the characteristics of bottom-gate Ta-doped InSnO (ITO) TFTs. An amorphous ITO:Ta channel layer was deposited at room temperature by sputtering. The devices were annealed at 50 ~ 200°C in oxygen condition for 1 hour. Optical transmittance data in the visible range reveal average transmittance higher than 80% including the glass substrate. The field-effect mobility of 3.5 cm2/V-s, the ON/OFF ratio of >104, and the sub-threshold slope of 81 V/decade are observed in the n-channel ITO:Ta TFTs. Effects of ITO:Ta preparation conditions on the resulting TFT performances were discussed. Stability experiments and further device optimizations through processes are under investigation. The detailed results will be revealed in this presentation.

EM-ThP19 Infrared Absorption Enhancement in Silicon via Silicidation and Nanoparticle Incorporation. C.M. Gonzalez, R. Suchan, G. Duscher, R. Kalyanaraman, P.D. Rack, University of Tennessee, Knoxville

Conventional silicon photovoltaics lack the ability to absorb the full electromagnetic spectrum arriving from the sun. Nickel silicides are small band-gap semiconductors that effectively engage the near-infrared region, whereas bare silicon does not. Ultra-thin (~100nm) silicon structures incorporated with nickel silicides have been synthesized and characterized and have shown enhanced absorption from 750-3000nm wavelengths. Four different structures were constructed via a combination of sputtering, co-sputtering and their optical and electrical characteristics were compared. In one case, the nickel is deposited and reacted between the p and the n+ silicon regions; in another, nickel is co-sputtered along with p-Si and co-sputtered with n+ silicon; and finally nickel is co-sputtered with silicon followed by laser irradiation to form nickel silicide nanoparticles via pulsed laser induced dewetting. This last structure contains non-patterned nanoparticles (<50nm) in close proximity to a p-n junction after capping the former with the p+n silicon. In this presentation we will correlate the material composition and micro and nanostructure by STEM and EELS to the observed optical and photovoltaic responses and demonstrate effective media approximations for the observed optical properties.

EM-ThP20 Fabrication of Shape-Controlled Metal Nanodot Array by Electrostatically-Driven Self-Assembly as well as their Charge Injection Properties. R. Sumi, H. Watanabe, Osaka University, Japan, T. Hosoi, H. Nagoya University, Japan

Nanometer-sized inorganic particles play an important role in modern nanotechnology owing to their superior characteristics that can lead to fascinating function that are difficult to be realized using conventional bulk materials. Recently the development of shape-controlled synthesis techniques make possible to obtain various well-defined shapes of nanoparticles with high yield. To determine their surface-to-volume ratio and crystal structure are expected to lead to improvement in performance in given application.

In this paper, we demonstrated fabrication of various metal nanodots MOS capacitors for floating nanodot gate memory using a evaporation-based colloidal self-assembly in order to attain a tight control over the size, shape, and density of metal nanodots, as well as the study of their effects on the charge injection characteristics of the nanodot array. We synthesized Au nanoparticles with 2nm diameter by solution plasma processing in aqueous solution. The surface was modified with organic surfactants which tuned their zeta-potential to be approximately -40mV. We have recently proposed a versatile method for the fabrication of self-assembled metallic nanodot arrays onto a SiO2/Si substrate with dimension of 50 x 100μm2 by using a newly developed electrostatically-driven self-assembly. The substrate surface was modified with amino-silane agents prior to assembly. In order to make MOS capacitors, Au nanodot array as a charge trapping layer, the Au nanodot array was embedded in a gate oxide.

By precise control of the velocity of the leading edge of a liquid slug, the volume ratio of the particles, and the deposition rate, we were able to reproducibly form an array consisting of a single layer of Au nanodot array with density of 1012particles/cm2. We further demonstrated the charge injection characteristics of the Au nanodot array, embedded in the ultra-thin SiO2 layer consisting of both thermally grown tunnel oxide and RF-driven sputtered control oxide layer. Counterclockwise hysteresis was observed reproducibly, whereas there was no hysteresis in the C-V curve of the MOS capacitor without Au nanodots. This hysteresis indicates the charging and discharging of the embedded Au nanodots. This electron confinement caused the flat-band voltage shift observed as the capacitance hysteresis.

EM-ThP21 Small-Molecule Scaffolds for Directed Self-Assembly. P. Manchero-Passo, A.J. Muscat, University of Arizona

Self-assembled monolayers (SAMs) are often used as scaffolds for the deposition of a subsequent layer. This process allows the surface properties to be tailored to a specific application and facilitates the construction of functional nanostructures. For instance, copper electrodes and graphene have been deposited on carbonyl-terminated and on 3-aminopropyltriethoxysilane (APTES) monolayers, to create metallic electrodes for electronic devices and a lubricating layer for nanoelectromechanical systems, respectively. Small silane molecules in particular could be used to direct the self-assembly of another layer and keep it in close proximity to a silicon surface. Vinylichlorosilane (VTCs, CH2=CH-SiCl3) is a candidate and contains a carbon-carbon double bond. A vinyl C=C bond is a versatile functional group because it can be reacted in situ to generate a variety of terminations, including -COOH, -SH, and -NH2, and could also direct the assembly of new molecules on the surface. While VTCs SAMs have been studied previously, a reproducible and self-limiting process to form a single monolayer from solution has not yet been achieved. In this work we adsorbed VTCs from solution on a Si(100) substrate and characterized the layer with ellipsometry, X-ray photoelectron spectroscopy (XPS), and atomic force microscopy (AFM). The metal nanodot arrays were cleaned and hydroxylated in a piranha solution. After drying, the samples were dipped for different times into a 1:1000 solution of VTCs in toluene, hexane, chloroform and acetone. The thickness of the monolayer was monitored ex situ using ellipsometry. Toluene and hexane yielded
multilayers with a thickness greater than 10 Å, whereas chloroform and acetone produced more consistent monolayers with a thickness of about 6 Å. A bromination was performed to demonstrate the presence of the unsaturated bond. We are in the process of making XPS and AFM measurements to determine whether VTCS is covalently attached to the Si substrate and to assess whether the vinyl group is suitable to graft molecules to the substrate.

EM-ThP22 Deposition of Organic and Inorganic Hybrid Laminates using Ozone-Based ALD (Atomic Layer Deposition), J. Huang, M. Lee, NCD Technology, J. Huang, M. Lee, University of Texas at Dallas, P.-R. Cha, University of Texas at Dallas and Kookmin Univ., Korea, J. Kim, University of Texas at Dallas

Recently, organic materials have been attracting attention due to the large area, low cost, and flexibility. On the other hand, inorganic conducting materials have several benefits such as high conductivity, functional properties, and rigid characteristics, etc. A novel technique for building an organic and inorganic hybrid stack has been introduced by M. M. Sung et al. using a modified ALD method. It has advantages to minimize the defects during growth of organic and inorganic films because hybrid films are deposited by an in-situ process through the sequential self-limiting surface reaction like atomic layer deposition (ALD), while maintaining their flexibility and functionality.

In this study, we investigate the growth characteristics of organic/inorganic hybrid film. Organic films have several properties such as high flexibility, functional properties, and rigid characteristics, etc. Novel technique for building an organic and inorganic hybrid stack has been introduced by M. M. Sung et al. using a modified ALD method. It has advantages to minimize the defects during growth of organic and inorganic films because hybrid films are deposited by an in-situ process through the sequential self-limiting surface reaction like atomic layer deposition (ALD), while maintaining their flexibility and functionality.

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References

EM-ThP23 Simple Methods to High Quality Self-Assembled Monolayers of Alkylsilanes on Oxides, B.D. Gates, Y. Gong, A. Liu, H.W. Ng, M. Wang, Simon Fraser University, Canada

The interfacial properties of a material can be easily modified by modifying its surfaces with self-assembled monolayers. These single-molecule thick coatings can be used to prevent non-specific adsorption of biomolecules, protect the underlying material from chemical or physical attack, serve as a thin layer to prevent or direct the deposition of a coating material, and reduce friction between two materials. Interest in the use of self-assembled monolayers continues to grow. The applications being pursued for these molecular coatings are increasingly interested in the robustness and uniformity of these films. An important aspect of their resilience is the density and size of defects within the monolayer, which can have a large role in their overall performance. One type of monolayer for which it is particularly challenging to achieve high quality coatings are those based on the assembly of alkylsilane molecules. Often these coatings are pursued for the modification of oxidized surfaces, but form either a submonolayer or a multilayered film. A primary goal of the work that will be presented is to further optimize the conditions in which monochlorotri[propyl]silane-based molecules are assembled into monolayers. Considerations for the development of these refined methods include the ease of scaling the technique to cover larger areas and the ability to avoid stringent requirements for environmental control during growth of the monolayer. Additional work is being pursued to identify and repair defects within the silane-based films. One approach that will be discussed uses solvent extraction techniques to remove adsorbed molecules, followed by further assembly of alkylsilane molecules onto the unprotected regions of these surfaces. The 'repaired' surfaces exhibit improved resistance to chemical attack. A key aspect of this work has been the development of new analytical methods for determining the density and size of the defects. These methods are used to also monitor the success of repairing the defects. The talk will cover these and other aspects of our progress towards developing simple methods that will produce high quality monolayers of alkylsilane-based molecules.


More cell phones are damaged by water than by any other means, and this damage often requires the devices to be discarded. The number of damaged phones is also increasing because these phones are now taken almost everywhere. Chemical vapor deposition may provide a solution to this problem, which is large (100.9 million Smartphones were shipped in Q4 2010). For example, the process may be connected to a hydrophobic monolayer or multilayer of fluorosilanes. Bonding of the fluoroilane may be improved using of a primary adhesion layer, which may be a different silicone monolayer, e.g., an isocyanatotiolane, and/or by introduction of hydroxyl groups via plasma treatment. The latter process is typically rapid and economical and can take place both on oxide and polymeric materials. The presence of OH groups can be assessed by XPS, ToF-SIMS and ATR-FTIR. The density of surface hydroxyl groups can be varied by changing the proportions of etch gases, the time and intensity of the plasma treatment, and the system base pressure. The hydrophobicity of the surface can be characterized by contact angle goniometry and XPS and ToF-SIMS analysis of fluorine. Resistance to abrasion can be tested with a Martindale abrasion tester. This work can be further extended to touch screen panels in equipment used under water.

EM-ThP25 Impact of Substrate Type on the Formation of Si Nanocrystals in the Annealed SiOx, and SiOx:Er,F Films, M.V. Sopinsky, M.V. Sopinsky, V.V. Strelochuk, I.L. Veligura, N.A. Vlasenko, V. Lashkaryov Institute of Semiconductor Physics, Ukraine

Erbium-doped nc-Si/SiOx nanosystems are considered as possible candidates for the manufacturing of light-emitting sources compatible with silicon technology [1]. Fluorine, like other halogens, has a tendency to enhance the crystallization of SiOx films on c-Si (Ref. 2). Hence, the introduction of F will intensify the transformation of amorphous silicon nanoinclusions formed on initial stage of SiOx thermal decomposition into Si nanocrystallines, and reduce the low-temperature threshold of this process. Lowering of the process temperature is the essential advantage of every technology. In case of SiOx:Er,F films this is principally important because at T > 900°C the segregation of erbium ions into metallic clusters takes place. Consequently, the final stage of forming the erbium-doped light-emitting system should not exceed 1000°C.

In this work the silicon nanoparticle formation due to the structural-phase transformations in the SiOx and SiOx:Er,F films evaporated onto c-Si and sapphire substrates is studied. These transformations are induced by moderate temperatures annealing (650–1000°C). For this study the methods of Micro-Raman scattering and multi-angle ellipsometry are used. It was established that more intensive formation of a-Si nanoparticles takes place in the doped films as compared to undoped SiOx films on both c-Si and sapphire substrates. Besides, it was found that the crystallization of a-Si nanoparticles is much more intensive and occurs at lower temperatures (750°C) in the annealed SiOx:Er,F films on sapphire substrate as compared to the annealed SiOx:Er,F films on c-Si substrate.

Analysis of available literature data which deal with behaviour of Er and F in silicon, SiOx, and SiOx:Er,F systems allows explaining the obtained results by the influence of F. In our opinion, the mechanism of F influence on the crystallization of a-Si nanoparticles is similar to the mechanism of h-hydrogen-induced crystallization of amorphous Si thin films [1]. Fluorine diffuses into Si–O and c-Si extremely fast. Therefore, during the annealing of SiOx:Er,F films deposited on c-Si the significant part of mobile fluorine easily diffuses into c-Si substrate and its concentration in the film drops considerably. On the contrary, F very slowly diffuses into Al2O3 – even the thin native aluminum oxide layer provides very good protection against fluorine diffusion. Accordingly, fluorine concentration in the film on substrate stays significant, and its crystallization action manifests itself stronger.

Conclusions
1) It is shown that the formation of erbium-doped nc-Si/SiOx nanosystems at the temperatures T ≥ 750°C is possible by annealing of SiOx:Er,F films on sapphire substrate.

2) This work, together with the work [2], demonstrates the wide variety of mechanisms by which a substrate can impact upon the formation of nc-Si/SiOx nanosystems.


Energy Frontiers Focus Topic
Room: East Exhibit Hall - Session EN-ThP

Energy Frontiers Poster Session

Recently, photovoltaic cell is accepted as new green technology replacing fossil fuel. Because of a next-generation photovoltaic cell with flexibility, portability and low cost, organic thin film solar cell (OTFSC) is currently the subject of great interest and intensive study. OTFSC is composed of organic electronic materials, which have differences in electron affinity and ionization energy, between two metallic conductive electrodes. The layer of the material with higher electron affinity and ionization potential is electron acceptor, and the other layer is electron donor. The hole-electron pairs (exciton) generated in either material, when OTFSC was photoinduced. When most of excitons reach the interface of the electron acceptor and electron donor, excitons break into carriers of electrons and holes efficiently. Electrons and holes move in the acceptor and donor, respectively. The carriers are collected by the electrode in contact with each layer. In the latest report, OTFSC has exceeded 8 percent of conversion efficiency. However, it was difficult to increase the area of OTFSC with high conversion efficiency. For instance, the conversion efficiency depended on the layer defects, which was caused by contamination of dust and H2O2 oxidation and inhomogeneous coating. The defects influenced the carrier lifetime in OTFSC. Hence, the inspection technique, which was evaluated in two dimensions, was required to measure the carrier lifetime. The present inspection technique (Laser beam induced current method) had to contact electrodes with electric wires and pass an electric current through OTFSC. Because the carrier lifetime was measured by the decay of electric current, the value was influenced by every interface of circuitry in OTFSC. Therefore, it was necessary to detect the pure carrier lifetime in the organic semi conductivity layer. In this study, to investigate the pure carrier lifetime, we developed the analysis system employing micro wave photoconductivity decay (μPCD). The carrier lifetime was discussed from the viewpoints of electrochemistry and quantum mechanics.

Thermionic electron emission from nitrogen doped hydrogen terminated diamond film had been evaluated 18 months after it had been deposited on a 25 mm diameter molybdenum substrate by microwave plasma assisted chemical vapour deposition. A hydrogen enriched surface layer, enabling an easy electron field was stable but relatively low concentrated. The AZO thin film and the thin-film buffer layer were prepared by pulsed laser deposition (PLD) using an ArF excimer laser. As an example of AZO n-doped ZnO heterojunction solar cells, AZO/ non-doped ZnO (ZO)/Cu2O solar cells were fabricated by inserting a ZO thin film as the buffer layer. It was found that the obtainable conversion efficiency in AZO/ZO/Cu2O heterojunction solar cells increased markedly as the deposition temperature of the AZO and ZO thin films was decreased from approximately 300 °C to room temperature (RT), i.e., non-intentionally heated Cu2O sheets. In addition, when the ZO thin films were prepared on non-intentionally heated Cu2O sheets by PLD, the efficiency (η) increased considerably as the introduced O2 gas pressure was increased; η approximately 3% with a ZO thin film layer deposition at an O gas pressure of 0.1 Pa, gradually increased as the O2 gas pressure was increased up to approximately 1.2 Pa, and then decreased markedly at a pressure of approximately 1.5 Pa. The optimal thickness of the ZO thin film layer was in the range from 30 to 50 nm. The obtained increase of η is attributed to an improvement of film quality; in contrast, the decrease of η as the AZO film thickness increased up to approximately 50 nm may be attributable to the short lifetime of minority carriers in the n-ZO thin film. An AZO/ZO/Cu2O heterojunction solar cell fabricated under optimal preparation conditions exhibited an efficiency of 3.83% under simulated AM1.5G solar.

Energy Frontiers Poster Session

EN-ThP4 Effect of Inserting a Thin Buffer Layer on Obtainable Efficiency in n-ZnO/p-Cu2O Heterojunction Solar Cells. T. Minami, T. Miyata, Y. Nishi, J. Nomoto, Kanazawa Institute of Technology, Japan
The effect of inserting a thin-film buffer layer on the obtainable efficiency in n-ZnO/p-Cu2O heterojunction solar cells was investigated with a transparent conducting Al-doped ZnO (AZO) thin film/Cu2O sheet structure. To improve conversion efficiency, various heterojunction solar cells were fabricated by forming an AZO/thin-film buffer layer/Cu2O structure on the front surface of thermally oxidized Cu2O sheets that function as the active layer as well as the substrate. It was found that achieving higher efficiency Cu2O-based heterojunction solar cells fabricated by using various thin Cu2O sheet substrates. It was found that the surface of the Cu2O sheets always be treated using a low-damage deposition technology at a low deposition temperature. The Cu2O sheets, with electrical properties such as resistivity on the order of 10² Ωcm, hole concentration on the order of 10¹³ cm⁻² and Hall mobility above 100 cm²/Vs, were prepared by a thermal oxidation of copper sheets under appropriate conditions. The AZO thin film and the thin-film buffer layer were prepared by pulsed laser deposition (PLD) using an ArF excimer laser. As an example of AZO/thin-film buffer layer/Cu2O heterojunction solar cells, AZO/ non-doped ZnO (ZO)/Cu2O solar cells were fabricated by inserting a ZO thin film as the buffer layer. It was found that the obtainable conversion efficiency in AZO/ZO/Cu2O heterojunction solar cells increased markedly as the deposition temperature of the AZO and ZO thin films was decreased from approximately 300 °C to room temperature (RT), i.e., non-intentionally heated Cu2O sheets. When the AZO thin films were prepared on non-intentionally heated Cu2O sheets by PLD, the efficiency (η) increased considerably as the introduced O2 gas pressure was increased; η approximately 3% with a ZO thin film layer deposition at an O gas pressure of 0.1 Pa, gradually increased as the O2 gas pressure was increased up to approximately 1.2 Pa, and then decreased markedly at a pressure of approximately 1.5 Pa. The optimal thickness of the ZO thin film layer was in the range from 30 to 50 nm. The obtained increase of η is attributed to an improvement of film quality; in contrast, the decrease of η as the ZO film thickness increased up to approximately 50 nm may be attributable to the short lifetime of minority carriers in the n-ZO thin film. An AZO/ZO/Cu2O heterojunction solar cell fabricated under optimal preparation conditions exhibited an efficiency of 3.83% under simulated AM1.5G solar.

The solar cell efficiency is hardly connected with device fabrication processes, such as texturing silicon surface, which increases the surface absorption area of incident radiation on the cell and, consequently, reduces the overall reflectance. Notably, the front surface of the silicon substrate are fabricated with texturing surfaces to increase the device efficiency. These surfaces are obtained by anisotropic wet etching in KOH and NaOH alkaline solutions [1]. The disadvantage of these solutions are K⁺ and Na⁺ ions from KOH and NaOH, which are mobile charges in Metal-Oxide-Silicon structures. Thus, these solutions are not compatible for CMOS technologies which is used for integration circuits and devices on silicon surface. As an advantage, using these alkaline solutions, the <111> planes of silicon substrate with (100) crystallographic orientation are exposed, resulting in groups of pyramid on silicon surface. The pyramids induce many reflections and scatterings of the incident radiation, which increase the light absorption on the surface. This effect is defined as light trapping [1]. In this work, the silicon substrate texturing is carried out by an anisotropic wet etching, with an alkaline solution based on NH4OH, because this solution can form pyramids on the surface, does not release alkaline.

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products/reactants on the silicon surface and is fully compatible with the CMOS technology. With this solution (concentration of 9 wt% of NH₄OH and four magnetic stirring velocities) the nuclei of pyramids of silicon substrates with (100) crystallographic orientation are exposed and these planes can form some pyramids on surface. Scanning Electron Microscopy analysis presents that these pyramids are observed all long on the surface, indicating that the wet etching was uniform on the substrate. Furthermore, the pyramid height values are between 3.5 and 7.2 µm, and the silicon etching wet rate was about 0.7 µm/minute. The reflectance measurement shows that the surfaces with and without texturing present reflectance values of about 16% and 38%, respectively. The reflectance values of about 16% with texturing are similar to the results from references [1], which have used KOH solution. This result indicated that our texturing process based on NH₄OH, which is fully compatible with the CMOS technology, can be used to fabricate Si-based solar cells [2].

REFERENCES


Recepients of the importance of developing the future energy is recognized in various industries. In one of the green innovations, there are great expectations to develop organic photovoltaic cells; Dye sensitized solar cell (DSSC) and Organic thin film solar cell (OTFSC). Organic photovoltaic cells with inexpensive and low resource constraint are currently the subject of great interest and intensive study. DSSC is composed of photo-sensitized anode using molecular dye placed on a porous layer of titanium dioxide nanoparticles, and an electrolyte solution. The photo electrochemical system was known as the Grätzell cell. Electrons at the dye excited because of sunlight move into the titanium dioxide, and are collected by the electrode. On the other hands, OTFSC is composed of organic electronic materials, which have differences in electron affinity and ionization energy, between two metallic conductive electrodes. The layer of material with higher electron affinity and ionization potential is electron acceptor, and the other layer is electron donor. The hole-electron pairs (excitons) generated in either material, when OTFSC was photoinduced. When most of excitons reach the interface of the electron acceptor and electron donar, excitons break into carriers of holes and electrons efficiently. Electrons and holes move in the acceptor and donor, respectively. The carriers are collected by the electrode in contact with each layer of the anode solar cell, the time would influence the performance of the solar cell, because the carriers (electrons) move through the semiconductor and reach the electrode. Therefore, we investigated the carrier life time of several organic photovoltaic cells by using micro wave photoconductive decay (μPCD). The difference of the carrier life time was discussed from the viewpoints of electrochemistry and quantum mechanics.

EN-ThP8 Making Molecular Multilayers using “Click” Chemistry: Growth, Characterization, and Application in p-Type Dye Sensitized Solar Cells, P.K.B. Palomaki, P.H. Dinolfo, Rensselaer Polytechnic Institute

Bottom-up approaches to creating molecular multilayer assemblies using layer-by-layer (LbL) techniques give one the ability to tailor the surface properties of an interface through molecular control. LbL assembly methods can provide molecular level control of structure in one dimension from simple solution deposition processes. We have developed a versatile LbL fabrication method using copper(I)-catalyzed azide-alkyne cycloaddition (CuAAC) or “click” chemistry in the construction of multilayer assemblies on oxide surfaces.1,2 These reactions occur in minutes at room temperature allowing for the facile creation of complex, covalently attached multilayer assemblies. Multilayers containing synthetic porphyrins, perylenes, and mixtures of the two have been constructed in order to highlight the versatility of this method. The fabrication of this new type of molecular multilayer thin film will be discussed along with characterization of the photophysical, electrochemical, and structural properties. Initial results pertaining to the use of these films as light harvesting systems in p-type dye sensitized solar cells will be presented.

EN-ThP9 Powder-Sputtered Cu(In,Ga)Se₂ Thin Films: Nonstoichiometric Influences on Optical and Electrical Characteristics, N. Kim, J. Jeon, W. Lee, Chosun University, Republic of Korea

The thin film solar cell efficiency of 19.2% (0.41 cm²) was accomplished by using Cu(In,Ga)Se₂ (NREL, 2003), which has been fabricated by using the three stage co-evaporation process. There are two general methods for Cu(In,Ga)Se₂ thin film fabrication: selenization after sputtering and thermal co-evaporation. The novel method of powder-sputtering was employed for the production of Cu(In,Ga)Se₂ to deposit the thin film over the large area with a good uniformity without selenization of toxic H₂Se because the co-evaporation method requires very strict controls during the deposition process of Cu(In,Ga)Se₂ thin film. This method has the advantages of the production of target, change of stoichiometry, and easy doping of impurities. The composition of the starting powder target was changed by x = 0.1, 0.3, 0.5, 0.7, and 0.9 in the deposition process of Cu(In,Ga)Se₂ thin film while Cu/(In+Ga) ratio was ‘1’. The compositions of the starting powder target and the deposited samples with 1 µm-thickness were comparatively analyzed by electron probe microanalysis (EPMA). The crystal structure, surface morphology, optical properties, and electrical characteristics of Cu(In,Ga)₁₋ₓSe₂ thin films were analyzed by using X-ray diffractometer (XRD), atomic force microscope (AFM), UV-Visible spectrophotometer, and a Hall effect measurement system to investigate the influence of nonstoichiometry on the characteristics. The optimized composition was selected through the comprehensive analysis of the characteristics.

EN-ThP10 Indium-Doped CdS Thin Film by He-Ne Laser Exposure for CIGS Solar Cells, K. Myung, N. Kim, W. Lee, Chosun University, Republic of Korea

Cadmium sulfide (CdS) is a direct and wide band gap (2.42 eV at room temperature) II- VI semiconductor, which is a very desirable window layer for many photovoltaic solar cells including CIGS solar cell. CdS thin film has been paid to the considerable attention due to its band gap, high absorption coefficient, reasonable conversion efficiency, stability and low cost. It is generally known that the undoped CdS thin film has the high electrical resistivity. Some impurities were employed for doping CdS to produce less resistive films. Especially, indium (In) showed the improvement of resistivity and optical transmittance when it was doped into the sputtering-deposited CdS thin film. In this study, the structural, optical, electrical, and morphological properties of indium-doped CdS thin films prepared by the effective method with He-Ne laser at room temperature. Indium was deposited on the 200 nm-thickness of CdS thin film by the sputtering method with 10 nm-thickness. He-Ne laser (632.8 nm) was exposed with a small energy by a change of exposure time. The remained indium layer was removed with the chemical solutions. The doping effects of indium on the CdS thin films were investigated by analyzing the structural, optical, electrical, and morphological properties of CdS thin films by using X-ray diffraction (XRD), Auger electron spectroscopy (AES), UV-Visible spectrophotometer, Hall effect measurement system, and atomic force microscope (AFM). AES depth profile was employed to examine the indium-doped CdS thin film with a change of He-Ne laser exposure time. The doping uniformity and amount of indium into the depth of CdS thin film was compared to the characteristics of CdS thin film. The optimization of indium-doping process was achieved to enhance the electrical and optical characteristics including resistivity and optical transmittance.

EN-ThP11 Three-Terminal Tandem Devices for PV, TPV and CPV Applications, M. Emziane, Masdar Institute, United Arab Emirates

One of the key approaches for achieving very-high efficiency for photovoltaic (PV) devices is to use, instead of single junctions, double or multi-junction devices due to an extended solar spectral coverage leading to an enhanced overall power output from the multi-junction PV devices. In this study, we investigated two-junction solar cells based on group IV materials, i.e. Si and Ge, for top and bottom sub-cell, respectively. Unlike the conventional tandem configuration that was shown to deliver a slightly lower output, these double-junction cells were assembled with three terminals and operated independently [1, 2]. The main advantages of choosing these materials and device configuration are: (i) cheapest, largest and easiest group IV substrates; (ii) extended spectral coverage leading to more photons being converted; and (iii) no
current matching or the associated tunnel junctions between the two sub-cells.

We have undertaken a comprehensive modeling analysis for the device optimization and performance prediction. n/p-n/p as well as p/p-n device structures were investigated and optimized with regard to the thicknesses and doping levels of both top and bottom active junctions that lead to the highest device performance. Due to the split of the incident solar spectrum between the top and bottom sub-cells, the latter only receives the light to which the former is transparent (mainly in the near infrared) and therefore behaves differently from the single-junction cell counterpart.

Optimal current-voltage and power-voltage characteristics were generated for individual cells together with the corresponding quantum efficiency spectra. The applications of these devices in conventional PV, concentrator PV and thermal PV were assessed and the output PV parameters were predicted as a function of the simulated operating conditions.

References:


Non-pulsing DC power supplies for highly arcing processes are gaining more and more interest slowly pushing out older pulsing DC power supplies as more efficient and more cost optimal. The importance of arc management in DC power supplies is taking on a new meaning as most advanced DC generators are more and more successful in sputtering highly arcing and difficult materials, providing high deposition rates at competitive investment.

It has been developed as the answer to the market’s demand to limit arc energy during an arc occurrence in sputtering. Industrial implementation has already been successfully performed with air-cooled DC power supplies that have been on the market for years. The possibility of using DC power supplies for sputtering highly-arcing materials is the result. It is the beginning of a new age in magnetron sputtering technology. Further improvement and optimization by using more efficient water cooling and faster transistors has resulted in a new generation of power supplies with state-of-the-art arc management parameters. A complementary feature of the newly-developed arc management circuitry is software that supports efficient arc handling: self-adjusting arc management settings and an internal oscilloscope.

EN-ThP13 Photogenerated Current Enhanced by Surface Plasmon Resonance in Metal Grating. D.J. Lee, Inha University, Republic of Korea

Photogenerated current enhanced by surface plasmon resonance excited in metal grating was confirmed in Au/p-type a-Si Schottky structures. Conduction electrons in metal absorb the incident photon of energy hv (h is the Plank’s constant and v is the optical frequency) through the a-Si film, gaining enough energy to cross over the Schottky barrier into the semiconductor, where they are swept across the depletion region and collected as photocurrent under reverse bias. In addition, the surface plasmon polariton is excited in metal grating fabricated on the photore sist layer, and excited electrons will flow over the Schottky barrier. This additional photocurrent added to that directly induced by incident light and enhanced the responsivity of the photodetector.

EN-ThP14 The Science of Precision Multifunctional Nanostructures for Electrical Energy Storage. A. Predith, University of Maryland, College Park

Nanostructures for Electrical Energy Storage (NEES) is a five-year Energy Frontier Research Center investigating nanoscale behavior in lithium-ion battery materials. Funded by the US Department of Energy, the Center studies the properties of composite nanostructures containing an oxide or silicate as a charge storage material and low dimensional forms of carbon as an electronic conductor or mechanical support. NEES is particularly interested in nanostructures that are precise and regularly ordered. These structures provide a unique test bed for uncovering to the science of electrochemistry at the nanoscale.

The two Science Thrusts of the Center investigate nanotubes, nanowires, and layers of MoO2, silicon, and carbon electrode materials as well as nanomembranes in an electrolyte. They examine the mechanical, electrochemical properties, and interfacial phenomena that arise in heterogeneous material combinations. The two Enabling Thrusts of the Center develop model systems and characterization instruments for the nanoscale. Ultra high vacuum scanning probe microscopy and first principles modeling provide for the study of pristine systems, and in situ transmission electron microscopy and microelectromechanical systems are allowing members of NEES to observe electrochemical processes as they unfold.

NEES is a collaboration of 19 senior investigators, ~35 postdoctoral associates and graduate research assistants, and many staff members at six institutions. The Center integrates their perspectives to pursue a comprehensive examination of electrical energy storage using precision, multifunctional nanostructures. Members include Philip Collins (University of California-Irvine), Michael Fuhrer (University of Maryland), Sang Bok Lee (UMD), Charles Martin (University of Florida), Reginald Penner (UCI), Zuzanna Siwy (UCI), John Cummings (UMD), S. Thomas Picraux (Los Alamos National Laboratory & Center for Integrated Nanotechnologies), Mark Reed (Yale University), Chunsheng Wang (UMD), YuHuang Wang (UMD), Kevin Leung (Sandia National Laboratories), Janice Reutt-Robey (UMD), Kevin Zavadil (SNL), Reza Ghodssi (UMD), Jianyu Huang (SNL, CINT), Robert Hwang (SNL), Gary W. Rubloff (UMD), John Sullivan (SNL, CINT), and Ashley Predith (UMD).


The long-term success of photosynthetic organisms has resulted in their global superabundance, which is sustained by their widespread, continual mass-production of the integral proteins that photocatalyze the chemical processes of natural photosynthesis. Here, a fast, general method to assemble wet and dry solar cells composed of one such photosynthetic protein complex, Photosystem I (PSI), onto a variety of electrode substrates, including gold, nanoporous gold, and graphene is reported. The resulting PSI films consist of a protein matrix that is permeable to electrochemical mediators in a wet cell and conducting for electron transfer in a dry cell. These results demonstrate how the assembly of micron-thick coatings of PSI on non-biological substrates yields a biohybrid ensemble that manifests the photocatalytic activity of the film’s individual protein constituents, and represent significant progress toward affordable, bioinspired renewable energy conversion platforms.

EN-ThP16 Thickness Dependent Photo-Voltaic Effect In Ferroelectric PLZT Thin Films. S. Kotru, H.V. Nampoori, The University of Alabama, Tuscaloosa

Ferroelectrics are emerging as potential candidate materials for energy harvesting. Among the class of ferroelectrics, Lanthanum doped Lead Zirconate Titanate (PLZT) is one which possesses high electro-optic coefficient and exhibits high optical transparency thereby makes it an attractive material for PV applications. In this work ferroelectric thin films of Pb0.95La0.05 (Zr0.54Ti0.46)O3 (PLZT) of various thicknesses were fabricated using chemical solution deposition method. Structural and ferroelectric properties of these films were studied using standard characterizations methods. Capacitor type solar cells were fabricated from these films using Pt as the top electrode. The IV curves were measured for each device where the thickness of PLZT film ranged from 70 nm to 350 nm. For devices, made with films less than 150 nm thickness, no PV response could be measured as an electrical contact could not be made on such films. From the PV curves measured on films with 140 to 350 nm thickness, the photocurrent density was seen to decrease from 597 to 64 nA/cm2, where as no major change was observed for the open circuit voltage. Thus thinner films are seen to enhance the PV response in ferroelectric based capacitor type solar cells.


Chemical-vapor-deposited diamond is an attractive material for electron field emitters because of its low or negative electron affinity, mechanical strength, and chemical inertness. Arrays of ultra-sharp diamond tips with a radius of curvature less than 5nm have been fabricated [1] and show significant improvement in emission brightness and turn-on field compared
The influence of different Ni\textsuperscript{2+} and Cu\textsuperscript{2+} concentrations on size serves as self-defensive mechanism of these bacteria against toxic influence. The method of measurement has been investigated. The ability of this bacteria to reduce – H\textsubscript{2}S, and formation of the unsoluble metal sulfides. It also precipitate toxic heavy metals compounds. It’s a result of their biogeochemistry of aquatic environments because of their ability to reduce ions that have an overwhelming harmful effect towards all living organisms.

Nowadays MFCs is considered to be applied as highly effective and self-sustaining model of wastewater treatm ent which contains energy in the matter oxidation are self-sustaining. Produce electricity represents a highly effective method for bioenergy. The use of renewable fuels requires the development of alternative sources of energy such as waste biomass for environmental benefits. MFCs investigations have detected substantial interest in development of different devices for generating electricity and removing wastes. The development of processes that can apply bacteria of normal environmental microflora to produce electricity represents a highly effective method for bioenergy production as bacteria are self-replicating, and thus the catalysts for organic matter oxidation are self-sustaining.

EN-ThP18 Microbial Fuel Cells Development on the Base of Desulfurobium Acetoxidans Bacteria. O.M. Vasyliv, O.I. Bilyi, S.O. Hnatish, Ivan Franko National University of Lviv, Ukraine

Microbial fuel cells (MFCs) are able to capture the electrons produced during different biochemical microbial processes to generate electricity. They offer the promise of a new sustainable source of energy, produced from biodegradable and reduced compounds. Increased shortcoming of unrenewable fossil fuels causes the need for it’s less dependence. The enhance of use of renewable fuels requires the development of alternative sources of energy as waste biomass for environmental benefits. MFCs investigations have detected substantial interest in development of different devices for generating electricity and removing wastes. The development of processes that can apply bacteria of normal environmental microflora to produce electricity represents a highly effective method for bioenergy production as bacteria are self-replicating, and thus the catalysts for organic matter oxidation are self-sustaining.

Magnetic Interfaces and Nanostructures Division Room: East Exhibit Hall - Session MI-ThP

MI-ThP1 Investigations of Ni and Co Magnetic Overlayers at the Advanced Photon Source. G.D. Waddill, T. Konmes, Missouri University of Science and Technology, S.W. Yu, J.G. Tobin, Lawrence Livermore National Laboratory

Magnetic overlayers and bilayers of Ni and Co on Cu(001) have been investigated as a function of coverage, using X-ray Magnetic Circular Dichroism in X-ray Absorption Spectroscopy (XMCD-XAS) and Photoelectron Spectroscopy (PES). These studies were pursued at Beamline 4 at the Advanced Photon Source (APS).

Lawrence Livermore National Laboratory is operated by Lawrence Livermore National Security, LLC, for the U.S. Department of Energy, National Nuclear Security Administration under Contract DE-AC52-07NA27344. This work is funded by the DOE Office of Science, Office of Basic Energy Science, Division of Materials Sciences and Engineering. The Advanced Photon Source (APS) is supported by the Director, Office of Science, Office of Basic Energy Sciences.


Thin Co films possess high value of saturation magnetization, low coercivity and uniaxial magnetic anisotropy. These properties are necessary for high density storage devices, magnetic field sensors and applications for UHF electromagnetic radiation. Consequently Co is perspective material for such employments. There are many influences to affect magnetic film growth. The latter allows one to govern their properties in wide range. Comparative studies of microstructure and magnetic properties (static and dynamic) are essential to develop new approaches for directional obtaining of materials with given frequency dispersion of magnetic permeability. The results of a comparative study of microstructure (TEM, GIXD), static (VSM) and dynamic (in the range of 0.01 to 10 GHz) magnetic properties of thin Co films are presented. The objects studied were rectangular films (60 x 600 mm) of 20 mm thick on poly(ethylene terephthalate) substrate. High purity (99,95%) cobalt disk target and a rotating drum as a substrate holder was used to obtain Co films.

Under a certain sputtering conditions two different types (with ferromagnetic resonance at 2 and 6 GHz) of Co films were obtained. Both types of films are of nanocrystalline structure. The grain size of crystalline size (from 7 to 90 nm) along with the increase of coercivity (from 17 to 110 Oe) and the increase of resonant frequency (from 2 to 6 GHz) was discovered.
Uniaxial crystalline texture discovered was found to effect resonant frequency. When the texture axis is in the film plane, mechanical bending of Co film may cause resonance frequency varying up to 1 GHz. Perhaps the effect is due to magnetostrictive mechanism [1]. The relationship obtained allows one to develop new pathways for magnetic materials production.


MI-Thp3 A Facile and Controllable Two-Step Electrodeposition Technique in Synthesis of Nanostructures of Metal Oxides on Carbon Nanotubes S. J. Yang, S. Ganasekaran, University of Wisconsin-Madison The nano dimensions of materials are comparable to the size of the target analyte biomolecule, higher catalytic reaction, better affinity binding or more efficient molecule-capturing may occur, leading to high sensitivity. And it is possible to use nanoparticle tags for designing electrical bioaffinity assays with remarkable sensitivity and multiplexing ability. So far, efforts have always been made to design novel nanomaterials useful in solving emerging bioanalytical problems such as rapidness, anti-interfering ability, specificity, stability and sensitivity. Synergies of nanocomposite materials, generally retaining the functional properties of each component and possibly yield synergistic effects via cooperative interactions, have exploited a new area to miniaturize and optimize nano-scale sensors and electronics. The synergistic interactions of these new features include but not limited to increased surface area, enhanced electrocatalytic activities, improved biocompatibility, promoted electron transfer and better invarianlarity against intermediate species. A lot of efforts have been made to fabricate nanocomposite materials of metals/metal oxides nanostructures and carbon materials, using a number of techniques, including sputtering, sol-gel, hydrothermal, microwave, and electrodeposition from different precursor solutions containing complex agents. Among these, electrodeposition is the easiest, most controllable, environment-friendly and robust technique for synthesis of metal/ metal oxides NPs, in which, the size, density, composition and even the shape of NPs could be well-controlled by electrodeposition potential, time, concentration and composition of metal precursor solutions.

Herein, we report a general two-step approach of electrodeposition useful in facile, controllable and green electrochemical synthesis of metal oxide NPs onto carbon supports, using carbon nanotubes (CNTs) as an example. First, metal nanostructures were electrochemically deposited onto carbon supports at a constant potential with the density, shape and electrocatalytic activities of the produced nanostructures well-controlled by the time and deposition potential applied as well as the concentration of the precursor solution. Then the as-deposited metallic nanostructures were oxidized into metal oxide nanostructures by repetitive potential cycling with extent of oxidation and generation of metal oxides controlled by the number of potential circles.

The as-synthesized metal oxides-CNTs composites were characterized and applied as a glucose sensor for illustration of their electrocatalytic properties.

MI-Thp4 Magnetic Properties and Size Control of Zn0.95Mn0.05O Nanorods Deposited by Pulsed Laser Deposition. T.C. Wu, Y.C. Teh, D.R. Liu, D.Y. Chuang, National Applied Research Laboratories, Taiwan, Republic of China

The well-aligned ZnO nanorods with 5 at.% of Mn doping (Zn0.95Mn0.05O) were deposited on silicon (100) substrates by pulsed laser deposition at three different substrate temperatures ranging from 600 °C to 700 °C, while the structure with and without a ZnO seed layer were both considered. The magnetic and structural properties of Zn0.95Mn0.05O nanorods has been characterized by x-ray diffraction (XRD), energy-dispersive X-ray spectroscopy (EDX), scanning electron microscopy (SEM), high-resolution transmission electron microscopy (HRTEM), and superconducting quantum interference device (SQUID). We demonstrate that the distribution and diameter of well-aligned Zn0.95Mn0.05O NPs are controllable, which strongly depend on the substrate temperature. Also, the magnetic properties of Zn0.95Mn0.05O NPs are controllable and strongly depend on the morphology of Zn0.95Mn0.05O NPs, and are thus appropriate for further applications.

MI-Thp5 Characterization of Metal Oxides Tunnel Barriers for use in a Non-Local Spin Detection Device. A. Matsubayashi, College of Nanoscale Science and Engineering, the University at Albany-SUNY

Metal oxides can be utilized as interfaces layers between ferromagnetic metals and graphene to achieve spin injection into graphene. Utilizing the spin of the electron as well as its charge has the potential to be utilized for logic devices in the post CMOS era. The goal of our research is to inject and readout spins using a non-local measurement device. However the efficient spin injection has been realized its difficulty due to the conductivity mismatch problem1,2. In order to achieve the efficient spin injection, it has been discovered that the insertion of a few nanometers of a tunnel barrier between the ferromagnetic metal and the graphene increases the contact resistance and measured spin lifetime3. However, non-uniformity of the tunnel barriers (pinholes)4 lowers the quality of the interface barrier. In this study, we investigate the fabrication of tunnel barrier on graphene using various metal oxides such as aluminum oxide grown under UHV conditions on the graphene. Graphene samples were loaded into an ultrahigh vacuum MBE (Beam Epitaxy) machine. D sized thickness of metals were deposited from a Knudsen cell. Samples were then transferred back into the load lock and exposed to approximately 130 mTorr of pure O2 for 20 min. Several measurements were performed including scanning electron microscopy, X-ray photoelectron spectroscopy, and angle resolved XPS characterize the electrical and structural quality of the films and their suitability for to be utilized as a tunnel barrier in graphene spin measurements.

References:

MI-Thp6 In Situ Scanning Tunneling Spectroscopy on Ordered, Epitaxial La0.83Pr0.17Ca0.83MnO3 Films. Z. Gai, Oak Ridge National Laboratory, M. Gao, Chinese Academy of Sciences, P.C. Snijders, H.W. Guo, T.Z. Ward, Oak Ridge National Laboratory, H.J. Gao, Chinese Academy of Sciences, J. Shen, Fudan University, China

Scanning probe microscopy has been shown to be a potent tool to investigate the structural, electronic, magnetic properties as well as their spatial distributions of strongly correlated perovskite manganites. Differences in tunneling spectra were interpreted to originate from the metallic and insulating phases in the material. However, questions were raised and still remain whether SPM on manganites probes bulk characteristics or is limited to surface properties. Complicating factors in this discussion are that most scanning probe studies either were performed on rather rough or grainy (and thus not very well ordered) surfaces, or on surfaces that were contaminated due to exposure to ambient air. Such ill-defined surfaces exhibit many localized surface states that often give rise to non-equilibrium occupation of states, and associated artifacts in scanning tunneling spectroscopy (STS) data, often masking or overwhelming possible bulk contributions. However, an STM does probe the electronic structure at the surface and this can include both surface and bulk contributions. Therefore, even on ideally ordered surfaces such as in situ grown and studied epitaxial manganite films, it is still unclear whether surface states (or even surface ground states that are different from their bulk counterparts) that are induced by the broken symmetry of the surface will overwhelm the bulk contributions in STS. We report temperature dependent scanning tunneling spectra recorded on in situ grown, single crystalline epitaxial La0.83Pr0.17Ca0.83MnO3 (x=0.3) (LPCMO) films with different thicknesses: 100 nm and 25 nm. On the 100 nm LPCMO film, the tunneling spectra show a higher metal-insulator transition temperature as compared to the 25 nm LPCMO film. Consistently, bulk transport data for the two thicknesses of LPCMO films also show different metal-insulator transition temperatures, thus revealing that tunneling spectroscopy on manganites is sensitive to the bulk electronic structure and not limited to surface effects.

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In a search for new multiferroic materials where the direction of magnetization can be switched by an applied electric field, we have looked for materials in which polarization and magnetization are strongly coupled. Recent theory calculations predicted that the family of compounds MTiO3 (M = Mn, Fe, Ni), in a certain polymorphic structure (aenctic R3c), are promising candidates where a polar lattice distortion can induce weak
ferromagnetism. Guided by these insights, a rhombohedral phase of NiTiO₃ has been prepared in epitaxial thin film form, whose structure is very close to that predicted to be a multiferroic. The synthesis of such new epitaxial films, their full structural characterization along with our first-principles DFT calculations to predict the desired NiTiO₃ structure and its stability are reported.

**Plasma Science and Technology Division**

**Room: East Exhibit Hall - Session PS-ThP**

**Plasma Science and Technology Poster Session**


We report on an investigatory study into the anomalous copper contamination observed on silicon test wafers run in hydrogen plasma processing systems using substantially oxygen-free plasmas. The copper is believed to originate from trace copper constituents of aluminum and other alloys used in construction of the vacuum system components. The transport of the copper to the wafer surface is consistent with the formation of meta-stable CuH.

Several plasma-ash systems from multiple vendors were tested using VPD/ICP-MS on silicon test wafers processed using hydrogen plus nitrogen gas mixtures. Anomalously high copper contamination levels in excess of 5x10⁵ atoms/cm² were measured. The copper contamination was found to increase linearly with plasma process time and the rate was highly system dependent. Clean systems tended to have higher contamination rates as compared to systems that have run large amounts of mixed processes, suggesting carbon or hydrocarbon deposits on chamber walls can somewhat suppress the CuH formation. Cycling these systems with oxygen plasmas increased the copper contamination rate. Additionally the contamination rate increased with increasing chamber wall temperature suggesting strongly that the chamber construction materials are the source of the contamination. Additionally several systems exhibited increases of cadmium, germanium and nickel. Each of these metals also has stable or meta-stable metal hydrides.

It was found that adding oxygen to the plasma in excess of 1% by volume, suppressed completely the anomalous copper contamination. VPD-ICP-MS measurement showed that 1% or higher oxygen concentration reduced copper contamination to below 1x10⁴ atoms/cm².

**PS-ThP2 Real-Time Monitoring of Reactors Wall Status during Silicon and Metal Gate Etching. A.P. Marchelli, G.M. Pietrogrande**, MICRON Semiconductor, Italy, Italy

One major challenge in plasma etching processes for sub-65nm integrated circuits is to assure the wafer to wafer repeatability. Particularly in plasma processes etching for IC fabrication, with metal gate, organic or metallic residues are deposited on the chamber walls. Because of the deposition of these layers a large and uncontrolled shifts in the etch process could be observed. To avoid this problem dedicated dry cleaning of the etching chamber must be developed but this methodology requires the precise knowledge of the layer coated on the reactor walls (1). This is not easy because of complexity of the stacks that need several steps of etching with different chemistries. Particularly the sharing of the equipments among many different processes increases the complexity of layer coated. In order to assure that every wafer has been etched correctly a real-time monitoring of plasma impedance has been developed. The experiments were performed in a low pressure inductively coupled plasma reactor supplied with 13.56 MHz RF power. Through the monitoring of dedicated equipment parameter tightly related to the plasma impedance is possible to active a dedicated fault detection control. During the wafer process, the check of this parameter along dedicated step that use only oxygen, it is possible to get the low pressure inductively coupled plasma reactor supplied with 13.56 MHz power to the coil wrapped the glass tube. Hydrogen plasma was exposed to chemical oxide removed Si(100) surfaces. After the oxide layer was etched, the surface was terminated with hydrogen. Moreover, the surface was distorted, so that the atomic arrangements on the surface were changed into amorphous layer. With the increase of substrate bias which was fed with RF (800 kHz) power, the formation of the amorphous layer was increased.

Furthermore, we noticed that the main peak position was changed with the exposure in this process; in the initial stage, the main peak was observed between 2000 and 2030 cm⁻¹. In the middle stage the main peak was located at 2060 cm⁻¹. In the final stage the main peak was located at 2100 cm⁻¹. From the assignment of the peak positions, amorphous SiH components were formed in the initial stage; amorphous SiH₂ components were formed in the final stage. The peak at 2060 cm⁻¹ observed in the middle stage can be assigned to the distorted SiH or hydrogen terminated vacancy. It is suggested that the kinds of hydride components are changed with the exposure.


Hydrogen plasma generates hydrogen radicals and ions in it. Both of them are important to Si technologies, such as the deposition of Si-related films, the etching of the films and so on; hydrogen radicals generate the adsorption sites by abstracting hydrogen from hydrogen terminated Si surface, while hydrogen radicals terminated dangling bonds of Si films. On the other hand, hydrogen ions also play an important role in etching process. Then, it is important to investigate the reaction process of hydrogen plasma with Si surfaces. We investigated the reaction processes at the different substrate bias conditions. The processes were investigated with infrared spectroscopy in multiple internal reflection geometry (MIR-IRAS). Hydrogen plasma was generated by feeding RF (13.56 MHz) power to the coil wrapped the glass tube. Hydrogen plasma was exposed to chemical oxide removed Si(100) surfaces. After the oxide layer was etched, the surface was terminated with hydrogen. Moreover, the surface was distorted, so that the atomic arrangements on the surface were changed into amorphous layer. With the increase of substrate bias which was fed with RF (800 kHz) power, the formation of the amorphous layer was increased.

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**PS-ThP4 Comparative Study on Atmospheric Pressure Plasma Modification on Packaging Material using Microwave and DBD Sources. J.M. Hong, Y.L. Wu, T.S. Cho, D.N. Ruzic**, University of Illinois at Urbana Champaign

Atmospheric plasma treatment is a very promising way to get hydrophilic or hydrophobic surface characteristics for many applications. In the food manufacturing industry there have been various kinds of technical approaches for prohibiting oxygen or water vapor penetration into products in order to preserve freshness and prolong shelf life. In this study two different types of atmospheric pressure plasmas, microwave and dielectric barrier discharge, are used on paraffin-wax coated paper. Different process recipes were applied on the packaging material and the surface modification results were evaluated. In the case of the microwave torch, a 10% improvement in hydrophobicity in terms of water contact angle measurement result and relative permeability change for water vapor was measured. The water contact angle was increased to 116±6 from 105±7 through applying a plasma treatment using a He/N₂ mixture. For the perpendicular three replicates for each conditions were prepared for each sample and weighed periodically. Each sample was prepared with a cup containing 2.0g of de-ionized water covered with treated or untreated packaging paper. According to its different treatment condition, the water loss of the plasma treated sample was 7-9% less than untreated one. This means the plasma-treated material will do better as a protective barrier against water vapor transmission. With the DBD remote plasma, a highly efficient hydrophilic process was used using nitrogen gas with small amount of air or hydrogen insertion. In order to understand the different surface reaction for the two plasma sources, the surface morphologies of treated samples were characterized by scanning electron microscopy and chemical properties were evaluated.


Platinum nano-particles (NPs) and platinum thin films were deposited by using inductively coupled plasma atomic layer deposition (PEALD). In this PEALD process, the Ar/O₂ reaction plasma was applied to decompose the ligands of Pt precursor (Me₅CpPtMe₃) at varied substrate temperatures between 150 to 300 °C. The partial pressures of Ar and O₂ were held at 1.5 and 0.5 Torr, respectively during the PEALD Pt NPs process. Next the PEALD Pt thin films deposited by using the lower pressure of Ar and O₂ at 0.3 and 0.1 Torr. X-ray diffraction (XRD) patterns show that Pt NPs and thin films are polycrystalline and a preferred orientation along the (111) plane of the fcc structure. Based on the Transmission Electron Microscopy (TEM) measurement, the growth rate of Pt NPs size and Pt thin films thickness is about 0.4 Å/min/cycle. The Pt thin film film 4f XPS spectra consists with the Pt 4f data that observed metallic Pt peaks deposited at 200 °C on Si wafer using Ar/O₂ plasma of 400 W. The step coverage was approximately 100% coated in 170 nm (open diameter) trench structures with 3.5 : 1 aspect ratio. The
low resistivity of Pt NPs and Pt thin films prepared by using 600 and 200 cycles was 1.83×10⁻⁸ and 1.62×10⁻⁷ Ω·cm, respectively.

Keywords: indutively coupled plasma, PEALD, platinum NPs

PS-ThP7 Properties and Performance of TiN, TiCN Coatings on Stainless Steel for Commercial Production Equipment. Y.L. Wu, J.M. Hong, T.S. Cho, D.N. Ruzic, University of Illinois at Urbana Champaign Commercial production equipment coated with TiN and TiCN films were studied experimentally and their material properties and performance were characterized in detail. The goal is to produce a titanium nitride and titanium carbide nitride coating on bare stainless steel surfaces that can resist material adhesion on the equipment surfaces. TiN and TiCN were chosen as the coating material due to their superior performance of low friction coefficient and high wear resistivity. Previous studies have shown that the friction coefficient of the coatings could be as low as 0.15 and wear rate of 6×10⁻⁸ mm²/N·m whereas the friction coefficient was 0.6 and the wear rate was about 3 orders of magnitude higher for the bare stainless steel surfaces. In this study the TiN and TiCN coatings were made using magnetron reactive sputtering using a titanium target of 300mm in diameter and 1cm thickness with methane and nitrogen as reactive gases. The coating properties were characterized tribology testing. Adhesions of the coatings on equipment surfaces were determined with single-pass scratch tests and the friction coefficients and wear rates were found using pin-on-disc tests. Resistivity on gum base materials were also characterized with the amount of adhered gum based materials and contact angle measurements.

PS-ThP8 The Dry Etching Properties of Nα0.5K0.5NbO3 Thin Films in Inductively Coupled Plasma. Y.H. Joo, J.C. Woo, C.I. Kim, Chung-Ang University, Republic of Korea

Recently, the ferroelectric materials are possible to study many fields for the application of the ferroelectric materials. The application of ferroelectric materials in random access memory devices tends to replace both magnetic core and magnetic bubble memories. And observed high electric field tunability with low losses makes this material suitable candidate for various voltage tunable microwave devices. Ferroelectric materials usually have high dielectric constants. From several published works, it was found that the sodium-potassium niobate, Na0.5K0.5NbO3(NKN) thin films grown on oxide substrate have a great dielectric and piezoelectric characteristics. Ferroelectric NKN thin films attract attention for their superior dielectric, piezoelectric, and crystalline properties. Therefore, we can use NKN thin films for many applications including non-volatile memory, sensors, optical switches, and ultrasonic transducers. The preferential development of anisotropic etching process for NKN thin films is desirable for highly intergrated ferroelectric application. Unfortunately, a small number of papers on NKN etch behavior published. The etch characteristics are not known appropriately. This fact makes some difficulties for integration of NKN in the microelectronic technology. In this work, we investigated etch characteristics of the NKN thin films in inductively coupled plasma. Both NKN etch rate and NKN/SiO2 etch selectivity were measured as functions of gas mixing ratio, input RF power, and dc bias voltage. The chemical reaction between plasma and NKN thin film surface were investigated by x-ray photoelectron spectroscopy (XPS). Field emission Auger electron spectroscopy (FE-AES) was used for the elemental analysis of the etched surfaces. Field emission scanning electron microscopy (FE-SEM) was used to investigate the etching profile.

PS-ThP9 The Dry Etching Properties of IZO Thin Films in High Density CF₄/Ar Plasma. J.C. Woo, C.I. Kim, Chung-Ang University, Republic of Korea

The Indium zinc oxide (IZO) is a wide band gap transparent conducting oxide that has provoked a great deal of interest recently, due to its many potential applications. It allows the fabrication of devices with a low operating voltage such as the design of such devices will lead to more efficient next generation of display technologies based on transparent and flexible devices. IZO thin films have attracted a great deal of attention because of their excellent optical transmission, high conductivity, chemical stability, thermal stability and low compressive stress. Therefore, amongst the various patterning techniques used for IZO thin films, plasma etching is preferred, because it allows for high resolution pattern transfer for optoelectronic device structures. Although the plasma characteristics of IZO thin films have been well optimized, the development of an efficient pattern transfer process remains to be achieved. Among the various patterning techniques, the dry etching process has several advantages over the conventional wet etching process, including high resolution and easy process automation. Moreover, there have only been a few reports on the IZO dry etching using the high density plasma sources favored by modern microelectronic technology. As a result, the influence of the process parameters on the IZO etch rate has not been explored in detail and the IZO etch mechanism is not quite clear.

In this work, we investigated the etching characteristics of IZO thin film using a high density plasma (HDP) system. The etching characteristics of the IZO thin film were investigated in the terms of the selectivity of IZO over SiO2 as a function of the etch chemistry. The chemical states on the etched surface were investigated by X-ray photoelectron spectroscopy (XPS). Field emission Auger electron spectroscopy (FE-AES) was used for the elemental analysis of the etched surfaces.

PS-ThP10 The Dry Etching Properties of TaN Thin Film Using Inductively Coupled Plasma. J.C. Woo, Y.H. Joo, C. Li, C.I. Kim, Chung-Ang University, Republic of Korea

***PLEASE NOTE YOU MUST IDENTIFY A DIFFERENT PRESENTER FOR THIS ABSTRACT. YOU MAY PRESENT ONE PAPER ONLY (ORAL OR POSTER) AT THE CONFERENCE. J. WOO IS CURRENTLY LISTED AS PRESENTER FOR ABSTRACT #12955***Recently, to overcome these problems, conventional poly-Si gates on high-k dielectric layers could be replaced by metal gates on high-k dielectric materials. Metal electrode/high-k gate stack structures, etching of the metal electrode against a high-k dielectric layer using etching processes can be employed. In this approach, a very high etch selectivity of the metal gate electrode to the high-k dielectric is required because of the very thin nature of high-k dielectric materials. Therefore, a very high selectivity of a metal gate electrode to the high-k dielectric is also needed to minimize Si damage loss at ultrashallow source/drain regions.

In this work, the TaN thin films were etched in C2/BC3/Ar and O2/C2/BC3/Ar plasma. The effects of adding O2 to the C2/BC3/Ar plasma were investigated for the purpose of improving the etch selectivity of the TaN to SiO2 layer. The etching characteristics of TaN thin films were investigated in terms of etch rates and selectivity as a function of the gas mixing ratio. X-ray photoelectron spectroscopy (XPS) and Auger electron spectroscopy (AES) were used for elemental analysis of the etched surfaces.


The availability of particle counters which may be operated at ambient pressures of up to 2×10⁴ Torr is highly desirable in many current research applications. In one field in particular, it allows mass spectroscopy of gas analysis and processing plasmas to be performed using pressures of this order in the quadrupole mass spectrometer (QMS). These pressures are much closer to those of many processing plasmas so that the sampling of neutral species, in particular, from the plasma is improved. The sampling of ion and neutral species from magnetron plasmas is a good example. The particle counter used in the present investigation, could be mounted axially so as to have a direct line-of-sight view of the sampling orifice of the QMS. Consequently, energetic neutral species such as metastable atoms of helium, which are produced in helium plasmas and have long lifetimes against spontaneous de-excitation, may travel to the detector and have sufficient energy to be counted there. The detection of metastable helium may be of importance in the study of reaction processes during hydrogen (H/HDeuterium) (D₂) plasma, of which the residual fusion byproduct is helium ash. Furthermore, collision processes in the ionisation source of the QMS, (including Penning ionisation), which are insignificant at the more usual source pressures of below 1×10⁴ Torr, generate product ion species whose study helps the interpretation of the processes occurring in the plasma reactor. Typical data from neutral gases and plasmas in a range of gas mixtures which include helium or argon with molecular gases such as hydrogen and nitrous oxide are shown. The majority of the data presented consist of electron impact threshold ionisation efficiency curves obtained by scanning the energy of the electrons in the QMS source. The results presented are discussed in terms of processes which include collisions between metastable species such as He(2S) and He(2P) at 20.61 eV and 19.82 eV respectively [1] having radiative lifetime states of 20+2 μs and 2 μs respectively and other plasma constituents. The measurements suggest new avenues of research into gas analysis and processing plasmas in which energetic, long-lived, metastable species may have important roles.


Wave-cutoff method is an electron density measurement tool. Most of the diagnostic tools, including cutoff probe, need a few seconds to measure the plasma parameters. In this presentation, a fast measurement method named Fourier cutoff probe method is newly introduced. A wave-cutoff probe system consists of two antennas and a network analyzer. A network analyzer provides the transmission spectrum and the reflection spectrum by frequency sweeping. The plasma parameters such as electron density can be obtained through these spectra. The frequency sweeping time, the time resolution of the wave-cutoff method, is about 1~10 seconds.

We proposed a new method to use an impulse generator and an oscilloscope instead of a network analyzer. This method was named Fourier cutoff probe method. A short impulse signal with a few GHz band and its transmitted signal were detected by an oscilloscope, and the Fourier analysis of the signals made a transmission spectra of plasma. The data acquisition time can be reduced with this method.

In this work, 3 diagnostic methods, Langmuir probe, network analyzer cutoff probe, and Fourier cutoff probe, were also compared. The measurement results were well matched. This method provides electron density in time resolution of 10 nanoseconds, and it took several seconds to make a measurement. This technique is found to be very useful especially in pulsed plasma. Fourier cutoff probe was 40 times faster than Langmuir probe in a pulsed plasma measurement.


Measurement of plasma density, a key parameter that control the property of processing plasmas and hence the processing results, has been the focus of extensive studies in recent years, not only for characteristic of the plasmas but also for development of tools for monitoring of the plasma based processes. In this study, a multi-mode plasma absorption probe was investigated by employing a full wave electromagnetic simulation using a finite element code, High Frequency Structure Simulation (HFSS, ANSOFT CORP). The spherically-shaped probe was formed by two isolated hemispherical conductors, which was enclosed by a layer of dielectrics (alumina). In the simulation analysis, a coaxial line was attached to the spherical probe head and the "driven mode" was adopted, i.e., the wave was fed into the coaxial line and the absorption spectrum, reflection coefficient vs frequency, was then analyzed. The probe was in a plasma, characterized by the collisional electromagnetic plasma permittivity, determined by plasma density, temperature and plasma frequency and momentum transfer collisional frequency. Simulation results show that the absorption spectrum exhibits a number of resonances, as a result of cavity-like structure resonant modes, as well as the plasma density, similar to the results reported in a previous work based on an electrostatic analysis. The electromagnetic analysis adopted in this work, however, reveals that the wave also propagates along the interface between the plasma and the coaxial cable, similar to the characteristics of a conventional plasma absorption probe. This effect thus contributes to resonant peaks in the absorption spectrum that did not appear in the electrostatic analysis. A parametric study by varying the plasma density, probe size and dielectric thickness have been conducted and the results will be presented.

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As the size of the semiconductor device is drastically scaled down to nano-scale, the etching technology of high aspect ratio contact (HARC) is increasingly important. To etch the nano patterned HARC, the multi-layer resist (MLR) structure conventionally composed of a PR, SiON layer, and amorphous carbon layer has been developed because it is difficult to etch HARC directly using a photoresist mask only. However, even though the etching of HARC tends to show higher etch selectivity and proper etch profile with MLR structures, for the next-generation nano-scale dynamic random access memory devices, the etching of nano-scale SiO2 HARC using fluorocarbon plasmas is becoming more and more important.

Nanoscale SiO2 contact holes were etched by using C4F8/CHF3/O2/Ar gas mixtures in dual-frequency capacitive coupled plasmas (DF-CCPs) where 60 MHz power is coupled to the top electrode while 2 MHz bias power is applied to the bottom electrode. We added CHF3 and O2 gases to control the etch rate and improve the etch selectivity. Proper additive gas ratios are needed to get the high etch selectivity and maintain CD size. We measured radial intensity, etch rate and SiO2 surface by using optical emission spectroscopy (OES), X-ray photoelectron spectroscopy (XPS), field emission scanning electron microscopy (FE-SEM) respectively.

In this study, we confirm that the etch rate and etch selectivity are varied as a function of CHF3/O2 gas ratio. The SiO2 and amorphous carbon layer(ACL) etch rate are increased with increasing the HF power due to the introduction of gas dissociation. But as increasing the HF power, etch selectivity is saturated. In this study, Highest etch selectivity of about 6 is obtained at HF power 300 W, LF power 500 W, Ar/CH4/CHF3/O2 = 150/20/25/5 sccm, process pressure of 20 mTorr.

PS-ThP15 An Experimental Study on Large-Area Multi-Electrode Discharge for the Deposition of Microcrystalline Silicon Thin Film for Solar Cell. Y.S. Lee, S.H. Seo, H.Y. Chang, Korea Advanced Institute of Science and Technology, Republic of Korea

Recently, there are many researches in order to increase the deposition rate (D/R) and improve film uniformity and quality in the deposition of microcrystalline silicon thin film. These two factors are the most important issues in the thin film solar cell, and for the purpose of that, several process conditions, including the large area electrode (more than 1.1 X 1.3 (m2)), higher pressure (1 ~ 10 (Torr)), and very high frequency regime (VHF, 40 ~ 100 (MHz)), have been needed.

But, in the case of large-area capacitively coupled discharges (CCP) driven at frequencies higher than the usual RF (13.56 (MHz)) frequency, the standing wave and skin effects should be the critical problems for obtaining the good plasma uniformity, and the ion damage on the thin film layer due to the high voltage between the substrate and the bulk plasma might cause the defects which degrade the film quality.

In this study, we will propose the new concept of the large-area multi-electrode (a new multi-electrode concept for the large-area plasma source), which consists of a series of electrodes and grounds arranged by turns. The experimental results with this new electrode showed the processing performances of high D/R (1 ~ 2 (nm/sec)), controllable crystallinity (~70% and controllable), and good uniformity (less than 10%) at the conditions of the relatively high frequency of 40 MHz in the large-area electrode of 280 X 540 mm2. And, we also observed the SEM images of the deposited thin film at the conditions of peeling, normal microcrystalline, and powder formation, and discussed the mechanisms of the crystal formation and voids generation in the film in order to try the enhancement of the film quality compared to the cases of normal VHF capacitive discharges. Also, we will discuss the relation between the processing parameters (including gap length between electrode and substrate, operating pressure) and the processing results (D/R and crystallinity) with the process condition map for µc-Si:H formation at a fixed input power and gas flow rate.

Finally, we will discuss the potential of the multi-electrode of the 3.5G-class large-area plasma processing (650 X 550 (mm2)) to the possibility of the expansion of the new electrode concept to 8G class large-area plasma processing and the additional issues in order to improve the process efficiency.

PS-ThP16 Qualitative Research on Low-Damage Neutral Beam Etching Effect of Mechanical Properties. Y. Nishimori, U. Shinji, K. Tomohiro, BEANS Project, Japan, S. Masakazu, University of Tokyo and BEANS Project, Japan, G. Hashiguchi, Shizuoka University and BEANS Project, Japan

The Neutral Beam Etching (NBE) system has been expected to provide a damage-free top-down process for organic and inorganic materials and facilitate the practical development of future Nano scale devices.

In order to examine the mechanical damage-free effect by NBE, we have conducted NBE surface treatment to microcantilever formed over an 8 inch wafer from the upper surface several times, and measured the resonance frequency and the quality factor (Q-factor) of 52 cantilevers after each treatment.

The NBE treatment removes about 50nm, 100nm, and 500nm thick silicon surface layer, respectively.

The Q-factor of cantilevers is attributed by summation of various kinds of energy dissipation mechanisms such as air damping, thermo elastic damping, support, and surface dissipation.

Based on the theoretical consideration of mechanical vibration, we have designed the microcantilever having 30 um in width, 150 um in length and 2 um in thickness, by which the surface energy dissipation become dominant compared to other ones in high-vacuum condition.
In general, Q-factor is a function of the resonant frequency (fr) so that we can’t argue the effect of NBE treatment from the Q-factor change as it is. Therefore, the use of values Q-factor divided by resonant frequency (Q/fr) value) which becomes independent of the thickness in case that the Q-factor is dominated by the surface dissipation and is a good parameter to represent surface dissipation.

From the viewpoint of transition of Q/fr value taken after each NBE treatment, surface dissipation of the cantilever was increased with each treatment process and saturated after adequate NBE treatment. This indicates that surface damage introduced during fabrication process can be removed completely by slight surface etching using NBE.

In the conference, we will report the precise results of the NBE treatment experiment.

PS-ThP17 Free-standing Anodic Aluminum Oxide on Silicon Substrate by using Neutral Beam Etching C.K. Kim, K.S. Min, J.S. Oh, G.Y. Yeom, Sungkyunkwan University, Republic of Korea

When the Anodic Aluminum Oxide (AAO) is used as the solid mask, compared to the patterning using electron-beam lithography and block copolymer, the patterning using AAO is cheaper, more reliable, and easier in controlling the size and shape. However, due to the stress at the interface between the aluminum and the substrate, the alumina barrier layer is formed at the bottom of the AAO holes during the formation of AAO and it prevents direct physical and electrical contact to the substrate.

In this work, to study the effect of surface charging on the removal of the barrier layer, the Cl2/BCl3 neutral beam etching has been used and the results were compared with the AAO etched by an ion beam etching. By the neutral beam etching with BCl3 containing gas mixtures, the barrier layer was successfully removed due to the formation of volatile BOxCl3 during the etching. However, when the barrier layer was etched using the ion beam with BCl3 containing gas mixtures, even though AAO itself is etched, the barrier layer located at the bottom of the AAO pore was not easily etched due to the charging of the AAO pore similar to the case of conventional reactive ion etching.

PS-ThP18 Synchronous Pulsed Inductively Coupled Plasma Source for Controllable Charged Process, K.S. Min, J.S. Oh, C.K. Kim, G.Y. Yeom, Sungkyunkwan University, Republic of Korea

A high-density ion source derived from an inductively coupled plasma is a useful tool in the fabrication of devices. A disadvantage of the technique is that charge-induced damage can cause irreversible changes in the device properties. In order to avoid or to compensate for the non-uniform accumulation of positive and negative charges near the treated surface, a number of low-damage processes have been proposed. The most promising of these processes uses high-energy 10–1000 eV beams of fast atoms and radicals for surface treatment.

This letter reports on a different method of negative beam formation that has potential application to neutral beam etching. In particular, we obtain a flux of fast neutral atoms with narrow energy and angular distribution from an initially negative ion beam which is extracted from the discharge volume. By increasing the pulse frequency of the antenna voltage to 20 kHz and applying a synchronized bipolar potential to the accelerating electrodes, it is possible to produce a high-density negative ion beam whose energy level is determined by the amplitude of the extraction potential. During the discharge, a positive potential is applied to the extraction electrode and a negative potential to the focusing electrode. In the afterglow period, these potentials are switched, allowing high-energy negative ions to escape the source. The lower and upper limits of the bipolar extraction potential on the two electrodes can be varied independently that the ratio of positive to negative ions in the resulting beam can be adjusted from 0% to 100%.

PS-ThP19 Initial Studies of a-Si Deposition and Nano-Crystallization using a Novel Reactor with Rotating Substrates, Z. Chen, J.A. Mucha, V.M. Donnelly, D.J. Economou, University of Houston, Y. Lee, A.J. Akey, I.P. Herman, Columbia University

Amorphous silicon deposition and nano-crystallization experiments were performed in a novel reactor with rotating substrates. Separate plasma sources were used for deposition (a capacitively coupled silane/helium plasma) and crystallization (an inductively coupled hydrogen plasma). Substrates were exposed sequentially and repeatedly to the deposition and crystallization plasmas for different times by varying the rotation speed. Radical concentrations in the plasma were monitored by optical emission spectroscopy while stable products were measured using a mass spectrometer. Films were characterized by spectroscopic ellipsometry (thickness, optical constants), Fourier transform infrared absorption (silicon-hydrogen bonding, hydrogen concentration), Raman spectroscopy and X-Ray diffraction (crystalline structure, degree of crystallization).

Results will be presented for different plasma operating conditions, substrate temperatures, and exposure times in the deposition and crystallization plasmas. The effect of “cross-talk” between the two plasmas and methods to prevent it will also be discussed.

Supported by the University of Houston GEAR Program and DoE


An Electron Beam Exciter device has been developed to perform a controllable optical emission spectroscopy diagnostic system. In this design, a small to moderate amount of current extraction (0–5 mA) is necessary to achieve the desired emission signal to produce reliable optical measurements utilized in etch process monitoring and control. Current extraction from this downstream plasma system is achieved in a Non-ambipolar Flow regime.2 Non-ambipolar flow occurs when all electron transport is from the excited plasma system, exits through a uniform electron sheath formed about the perimeter of a Nozzle Aperture (area Ae) while the ions are lost to the Exciter ICP grounded wall (area Ai). Electron extraction from the exciter ICP is maximized when the ratio of the ion loss collection area (Ai) to the electron loss collection area (Ae) is proportionally equal to the square root of the ratio of the ion mass to the electron mass, and the ion sheath potential drop to the Exciter ICP grounded wall is much larger than Tc/e.2,3 To investigate the degree of Non-ambipolar Electron flow in our device, we have tested a set of manufactured Nozzle extractors with different geometrical dimensions so as to discern the influence of Ai/Ae on the electron extraction process in low pressure (5-35 mTorr) Ar discharges. Strong agreement is observed between the measured plasma parameters at the ICP extraction region and the total amount of extracted current. In addition there is a measurable energy dependent response in the ratio of h(E)/h(E) indicative of electron-atom/molecule collision cross section for ionization. We have measured this ratio response in Ar and O2 atmospheres. This project is funded by NSF-Grant (CBET -0922962) and Verity Instruments.

3S. D. Baalrud; N. Hershkowitz; B. Longmier. Physics of Plasmas, 2007, 14, 042109 (1-6)

PS-ThP21 Fluid Model Simulation of Controllable E-beam, D. Urrabazo, G. Padrón-Wells, P.L.S. Thamban, M.J. Goeckner, University of Texas at Dallas

We have developed a new chemical diagnostic based on a controlled electron beam. Specifically we make use of an inductively coupled plasma, electron extraction optics and traditional optical emission spectroscopy. In the work reported here, a computer simulation was developed which allows the use of three different models: ambipolar, classical, and revised classical. Here, we investigate both the strengths and weakness of the various models as well as determine which system characteristics can be used to enhance the system performance. These simulation results will be compared to electrical probe measurements gathered from the experimental apparatus. This work is supported in part by NSF (Grant CBET- 0922962) and Verity Instruments.


Microwave plasma sources are widely used in industrial plasma processing devices for their favorable characteristics. In particular, electron cyclotron resonance (ECR) reactors offer a number of desirable characteristics for plasma etching and deposition, including higher plasma density, lower ion energy, lower pressure operation, and higher uniformity compared with the conventional type plasmas. ECR devices designed for commercial use usually provide the user with very limited access for controlling and characterizing the plasma conditions.

Retarding field energy analyzers (RFEAs) are generally used to measure the ion energy distribution function (IEDF) at electrically grounded surfaces. An RFEA can also be used to measure IEDFs at the surface of radio-frequency (RF) driven electrodes by electrically isolating it from the ground reference. The RFEA then “follows” the RF bias without disturbing it. Using the same principle, but applying grid bias signals of opposite polarity, one can also measure the electron energy distribution function (EEDF) at
the surface of RFEA. From the measured IEDFs and EEDFs it is possible to calculate the respective ion and electron flux and energy. These quantities are the governing parameter in plasma surface processing. Therefore, the metrological capabilities of the RFEA together with its non-perturbating in situ operation makes it a convenient diagnostic tool for characterizing plasmas in devices like ECR etching reactors.

In this work we present IEDF measurements at the surface of an RF driven electrode using a RFEA installed in an industrial ECR etching reactor. The RFEA is mounted at the RF biased electrode (on which the SiO$_2$ wafer is mounted) and is connected through high input power low pass filters to the ground referenced grid bias signals. The RFEA installation does not require any modifications to the reactor since the signal cabling is taken out to atmosphere through an existing unused port located in the vacuum pump tunnel. The discharge control parameters were varied as follows; pressure 5 to 20 mTorr, flow rate 10 to 50 sccm of pure argon and oxygen, microwave input power 750 to 1500 W, RF input power to 250W, and a variety of magnetic fields and target voltages. The radial location of RFEA has also been varied. The measured fluxes and energy distribution of the ions and electrons impacting the wafer surface are presented and discussed.

PS-ThP23 Simulations of the Radial Line Slot Antenna Plasma Source. P. Ventszek, Tokyo Electron America, S. Mahadevan, L. Raja, Eeseg Technologies, T. Iwao, Tokyo Electron Technology Development Institute, INC., L. Chen, M. Funk, B. Lane, R. Sundararajan, Tokyo Electron America, Y. Yoshihara, Tokyo Electron Technology Development Institute, INC., J. Zhao, Tokyo Electron America, T. Nozawa, C. Tian, K. Ishibashi, Tokyo Electron Technology Development Institute, INC.

The Radial Line Slot Antenna (RLSA) plasma source couples microwave power through a slot antenna structure and window to a plasma characterized by a generation zone adjacent to the window and a diffusion zone that contacts a substrate. The diffusion zone is characterized by a very low ion temperature. This property of the source underlines its use for etch applications and thin film processing for which low ion energy is desirable. Models of the RLSA source typically include ambipolar diffusion as an approximation enabling 2D and 3D simulations to be feasible. While such models have predictive value, they also have shortcomings when surface charging, capacitive coupling through a substrate or electroreflectivity are important. In this presentation we describe a self-consistent fluid model of the plasma kinetics of the RLSA source and demonstrate 3D source characteristics for benchmark argon and oxygen plasmas with and without capacitive coupling. Comparisons between the ambipolar model and self-consistent model are made.

PS-ThP24 Dual-Coil, Dual-Frequency ICP Source for Plasma Processing. V. Nagorny, D. Lee, Mattson Technology, Inc.

RF plasma sources used in modern plasma etch applications are required to provide high plasma uniformity and variable plasma controls, without independent plasma profile, plasma density and ion energy controls. They must be able to sustain a stable plasma in a very different gases and under very different conditions (gas flow, gas pressure, etc.). Finally, going forward the system has to produce a minimum impact on environment (energy consumption, EM emission). Mattson Technology has developed an etch tool with a new and efficient ICP source that uses two coils for operation at significantly different frequencies (13.56 and 2 MHz) by utilizing a ferromagnetic core in a low frequency coil. Tests on the tool revealed that:

- It provides high-density plasma generation, with no capacitive coupling between plasma and the source;
- There is no interference between higher and lower frequency RF subsystems;
- It provides good plasma and process profile control;
- It has wide processing window (gas pressure, gas flow, gas composition, source and bias power);
- High efficiency and stability of operation with both electropositive and electronegative gases.


Organic photoresist in semiconductor photolithographic process has been removed by downstream oxygen plasma which is generated in a quartz chamber or tube. In case of stripping heavily ion-implanted photoresist whose surface is carbonized by energetic ions, it is sometimes necessary to add CF$_4$ gas to oxygen. Moreover, hydrogen based plasma begins to be used to strip photoresist coated on low-k dielectric layers, where oxygen plasma is known to degrade low-k dielectric films. However, fluorine or hydrogen ions can attack the quartz walls and generate particles. To solve this problem, plasma generation chamber should be made of materials inert to fluorine or hydrogen ions. In this work we have designed and constructed a plasma source chamber made of ceramic tube. A 12 cm diameter ceramic tube is surrounded by a Faraday shield and wound by a 4 turn helical copper coil. The ICP source operates at 13.56 MHz RF power. There are also two turn dummy coils, which are electrically grounded, at the top and bottom sides of the 4 turn power coil. All coils have hollow square cross-section and cooling water flows through the coils. At the bottom of the 25 cm long tube an electrically grounded baffle with many small holes are located and only radicals are fed to the stripping process chamber for 300 mm wafers. Because ceramic materials generally have larger thermal expansion coefficients than quartz, larger thermal stress exists in the ceramic tube across tube thickness and at gaps between cooling coils, which results in breakage of ceramic tube. It is very important to achieve good thermal contacts between cooling water tubes and tube wall and thus small temperature gradient in the tube. We have investigated the temperature profiles and thermal stress to the ceramic tube of ceramic tube materials (Al$_2$O$_3$ and AlN). Types of ceramic paste to glue cooling coils to the ceramic tube, size of cooling coils, thickness of ceramic tube, and RF impedance matching methods. It is found that AlN which has larger thermal conductivity than Al$_2$O$_3$ shows no breakage of tube. Size of cooling coils and types of ceramic paste have no effect on tube breakage. In case of Al$_2$O$_3$, thinner tube survives from breakage. It is confirmed that little side wall attack is observed after 1000 runs of photoresist strip by H$_2$/N$_2$ plasma.


To produce semiconductors at low cost, the transition to 450 mm-sized wafers is being considered. For semiconductor fabrication on 450 mm substrates, the etching performance is expected to equal or exceed 300 mm substrates with improved productivity. An ICP plasma source capable substrate apparatus was developed on a proven plasma source using microwave electron cyclotron resonance (ECR) currently utilized for 300 mm equipment. We have evaluated plasma uniformity by using: 1) Single Langmuir probe with -50 V bias to measure ion current flux (ICF) distribution in the plasma reactor. The plasma generated area and the spatial distribution of ICF were scanned in the radial and z-axis directions in the reactor. 2) A 41-point wafer stage to measure ICF distribution, 3) Etch rate uniformity of poly-Si and SiO2 using coupon samples attached on 450 mm bare Si wafers. We used HBr/Cl2/O2/Ar gas and 0.1-10 Pa pressure range as test conditions.

We found that the high density plasma generated at the ECR zone and the shape of the plasma was ring-shaped. This ring-shaped plasma diffused and became increasingly uniform, finally becoming almost flat at a height of 25 mm above the wafer stage. To evaluate ICF distribution of the wafer stage, 41 pads were used to measure plasma density distribution using about 500 test conditions. By controlling the magnetic fields, non-uniformity of ICF below ±5% was achieved for several test conditions of gas and pressure. Poly-Si and SiO2 etching rate were observed to be very similar to rates seen using 300 mm substrate equipment with non-uniformity below ±1% within a wafer.

In this preliminary study, though we need more process evaluation by using whole blanket wafers and patterned wafers, the microwave ECR plasma apparatus for 450 mm wafers achieved the same level of process performance as the 300 mm equipment.


Increasingly stringent demands for maximum device speed and control of random dopant fluctuation effects dictate changes in device architecture, with FinFET adoption as an enabling technology. Drifting from 22nm and beyond the 15 nm technology node. Three-dimensional architectures pose a fundamental challenge for traditional ion beam implant processes, since, depending on the aspect ratio of the structure, certain sections can be inaccessible to the dopant beam. Plasma immersion implantation (Plasma Doping), which has been adopted as an enabling, production-worthy technology in the ultra high dose and low voltage regime for advanced DRAM device nodes, can be an alternative capable of overcoming these limitations. In this work, experiments are carried out using a VISta PLAD system, which uses an ICP source to generate a discharge and a series of negative dc voltage pulses applied to the substrate to accelerate ions towards it. Silicon trenches are implanted using an AsH$_3$-containing gas mixture in the PLAD reactor. The presence and activation of the dopant species is monitored by physical (Secondary Ion Mass Spectrometry, Transmission Electron Microscopy, Energy Dispersive X-ray spectroscopy) and electrical methods, and discussed in conjunction with analysis of the gas phase in the plasma chamber. Our experimental results indicate the occurrence of a combination of implant and ion assisted deposition processes. It is shown that balancing the constituents of a plasma doping process and plasma characteristics such as

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as electron temperature and gas phase composition is required to allow dopant incorporation into sidewalls of 3D structures.

PS-ThP28 Study of Low-k Dielectric Damage in a Radial Line Slot Antenna (RLSA) Reactor. Y. Susa, Tokyo Electron Technology Development Institute, INC., R. Sundararajan, J. Zhao, L. Chen, Tokyo Electron US Holdings Ltd., T. Nozawa, Tokyo Electron Technology Development Institute, INC., J.A. Macha, D.J. Economou, V.M. Donnelly, University of Houston, R. Wise, IBM Research

Low-k dielectrics are widely used as insulating materials in ULSI circuits. In BEOL processes, etching of low-k layers can cause serious damage, especially during photoresist ashing. In this study, a Radial Line Slot Antenna (RLSA) microwave plasma source was used in an effort to minimize damage to low-k dielectrics as a result of photoresist ashing. The focus of the work was ashing in CO₂-containing gas mixtures of test wafers partially covered with blanket low-k and photoresist layers. Damage to the SiCOH (k=2.5) low-k dielectric was characterized using Spectroscopic Ellipsometry, FTIR and XPS. It was found that low pressure (~ 5 mTorr) and high bias voltage (~2000 V) were beneficial in terms of minimizing low-k damage. XPS revealed that, under these conditions, an oxide “crust” formed on the surface of the low-k, that apparently protected the underlying material. Low pressure favors an enhancement of the ratio of the ion-to-O- atom flux. Energetic ions can be instrumental in forming this oxide crust by near surface ion implantation. The oxide hinders diffusion of the (already low density) O atoms into the low-k film resulting in minimal damage.


New pulsed plasma generator for reactive magnetron sputter processes was developed. (old sentence)

A new pulsed plasma generator for reactive magnetron sputter processes has been developed. (new sentence)

Compare with existing pulsed DC plasma generators new generator has high current capabilities in the range of 100 – 300 A. The new pulsed plasma generator has high current capabilities in the range of 100 – 300 A. (new sentence)

In proposed method near arc free magnetron discharge in reactive atmosphere can be generated by adjusting the frequency and amplitude and shape of the voltage pulses. (old sentence)

In the proposed method, near arc free magnetron discharge in reactive atmosphere can be generated by adjusting the frequency and amplitude and shape of the voltage pulses. (new sentence)

The application of the new pulsed plasma generator for reactive sputtering of Si, SiO₂, AlN and TiO₂ films at ENDURA 200 mm tool (AMAT) will be discussed. (old sentence)

The application of new pulsed plasma generator for reactive sputtering of Si,Ni, SiO₂, AlN and TiO₂ films at ENDURA 200 mm tool (AMAT) will be discussed. (new sentence)

Transparent Conductors and Printable Electronics
Focus Topic
Room: East Exhibit Hall - Session TC-ThP

Transparent Conductors and Printable Electronics
Poster Session

TC-ThP1 A Combinatorial Thin Film Sputtering Approach of the Synthesis and Characterization of Al₂O₃·TiO₂ High-k Dielectrics for Oxide TFT Application. J.H. Noh, J. Noh, P.D. Rack, The University of Tennessee

For the last decade, oxide based thin-film transistors (TFTs) have been extensively investigated because of their transparency, high mobility, low process temperature, which are expected to serve as the basis for optoelectronic and flexible devices. However, most of the work on oxide TFTs still rely on conventional dielectrics from Si technology, such as plasma-enhanced chemical vapor deposited (PECVD) SiO₂ or SiN, with process temperature of 250-300°C. For high performance, low-cost and flexible electronics, high-k dielectrics at low process temperatures are needed. A sputtering approach is used for the synthesis and characterization of Al₂O₃·TiO₂ high-k dielectrics because a combinatorial thin film sputtering approach can yield a wide range of compositions via a single co-sputter deposition process. The composition range of the films were simulated using a co-sputtering simulation and compared favorably to compositions measured by the wavelength dispersive spectrometer (WDS). The TFTs were fabricated with a bottom-gated staggered structure using amorphous indium gallium zinc oxide (a-IGZO) and InO₃ as the semiconducting active layer. Standard I-V and C-V data on the dielectric multilayers will be compared as a function of composition and, finally, the TFTs’ performance will be presented according to the relative contents of TiO₂ and Al₂O₃.


We fabricated sub-micron organic thin film transistors on polyethersulphone (PES) substrate using ink jet printing combined with an imprint method. The channel lengths of OTFTs were in the range between 5000 and 50 nm (polysilPentacene (TIPS) OTFTs) using this fabrication technique. TIPS pentacene was used as an active material and Polyvinyl alcohol (PVA) was chosen as a gate insulator. TIPS pentacene was printed by jetting onto the confined channels which were prepared using imprinting. The surface of confined channel was modified by UV irradiation in order to enhance the crystallinity of tips pentacene. The physical properties were analyzed using SPM, SEM, and XRD. The electrical properties were extracted from the transfer characteristics which were measured using Keithley-4200.

TC-ThP3 Study on Multiple Stacked 6,13-bis(triisopropylsilyl)ethyl) (TIPS) Pentacene for Improved Organic Thin Film Transistor. S. Lee, Sungkyunkwan Univ. & Samsung Mobile Display, Republic of Korea, J.J. Han, K. Kim, I.J. Bae, I. Chung, Sungkyunkwan Univ., Republic of Korea

We found that the crystallization of TIPS pentacene thin film plays an important role in determining the electrical property of organic thin film transistor (OTFT). Ink-jetted TIPS pentacene film reveals 2 different types of grains namely due to the coffee strain effect. The better electrical properties were obtained from the OTFTs with bigger grains that had been possible due to the multiple stacked TIPS pentacene layer. Poly-4-vinylphenol (PVP) was used as a gate insulator, and Au electrode was evaporated using a shadow mask. The channel lengths of OTFTs with the bottom-gate structure were of 20-50 um. The physical properties of TIPS pentacene films were analyzed using optical microscope (OM), x-ray diffraction (XRD) and secondary electron microscopy (SEM). The electric characteristics of OTFT were obtained using Keithley-4200.

TC-ThP4 Catalyst-assisted Pulsed Laser Deposition of Tin (IV) Oxide on Si Substrates: Growth Evolution of Low-dimensional Nanostructures. K. T. Leung, University of Waterloo, Canada

Single-crystalline nanostructures of SnO₂ have been grown in the aid of size-controllable gold nanosized catalysts supported on a Si substrate by using the Pulsed Laser Deposition (PLD) method. By changing the gas atmosphere and manipulating the deposition at a relatively low substrate temperature (500-700°C), we produce faceted nanobricks, nanograss, and nanoribbons on oxidized Si and cubic nanoparticles on H-terminated Si. Scanning electron microscopy clearly shows the faceted morphology of these one-dimensional and zero-dimensional nanostructures and suggests a vapour-solid and a vapour-liquid-solid growth mechanisms for nanoparticles and nanobricks and for nanograss and nanoribbons, respectively. X-ray diffraction results reveal the tetragonal crystalline phase of the SnO₂ nanostructures, and the relative intensity ratios obtained for different peaks further show a preferred growth orientation of (110) for the nanoparticles and nanobricks, and of (200) for the nanograss and nanoribbons. For nanobricks and nanoribbons, transmission electron microscopy confirms the single-crystalline nature of these nanostructures, and the corresponding high-resolution and selected area electron diffraction data illustrate their different growth orientations that generally lead to the preferred growth directions as inferred from the corresponding X-ray diffraction data. We have also recently used Helium Ion Microscopy to elucidate not only the intricate surface details but also the growth evolution of these SnO₂ nanostructures. These results demonstrate the versatility of
the catalyst-assisted PLD technique in depositing a variety of SnO2 nanostructures, which can be easily optimized and/or modified by appropriate doping within the PLD method for producing desirable optoelectronic, magnetic, gas-sensing, and semiconducting properties for emerging applications.

**TC-ThP5** From Discrete and Hollow Nanocavities to the Formation of Continuous Indium-Filled Indium Oxide Nanotubes, M. Kumar, South Dakota State University, B.R. Mehta, J.P. Singh, Indian Institute of Technology-Delhi, India

The growth mechanism for single crystalline indium oxide nanostructures is being under debate in scientific community. The mechanism was proposed for the growth of nanotubes based on the growth parameters dependent morphological transformation from discrete and hollow nanocavities in nanowires to continuous indium-filled indium oxide nanotubes. The gas flow rate induced change in indium partial pressure and hence supersaturation. The nucleation of species has taken place on the different substrate, which led to the development of two different nanostructures. We discussed a unified growth mechanism based on vapor-solid growth followed by the out diffusion or in-diffusion of metal indium depending on it’s partial pressure during growth. The out diffusion of indium resulted discrete and hexagonal nanocavities enclosed with minimum surface energy planes, [111] while the higher partial pressure of indium support the merging of discreet nanocavities into continuous connecting and filled with In metal.

**Triology Focus Topic**

**Room: East Exhibit Hall - Session TR-ThP**

**TR-ThP1** Subsurface Characteristics of an Abraded Fe-0.4wt%Cr Martensitic Steel using Nanoindentation and Cross-Sectional TEM Techniques, F. Kataaki, Sumitomo Metal Industries, Limited, Japan

The present investigation of unidirectional abraded surfaces of a martensitic (0.4 wt%Cr) steel with silicon (1.5wt%Si), chromium (1.5wt%Cr) and molybdenum (1.9wt%Mn) addition elucidates the work hardening and the softening near the surface layer caused by abrasion, particularly its relation to the wear behavior. The abrasion testing was performed using a pin-abrasion apparatus in which a small pin of the specimen was ground on an abrasive paper at an applied load of 2.1N and sliding speed of 0.66m/s. Crushed silica particles (size: 15-67μm) were used as the abrasive medium.

The abraded surfaces were examined with a nanoindentation apparatus to evaluate the variation of nanohardness with sliding time on a nanometer scale. A cross sectional transmission electron microscope (TEM) technique was also employed to clarify the structural changes in the region close to the abraded surface. It has been found that abrasion induced work hardening with sliding time was observed in the case of chromium and molybdenum addition steels. A fine dispersion of molybdenum carbide (Mo2C) was observed in the surface of the molybdenum steel after abrasion. Mo2C was also employed to clarify the structural changes in the region close to the abraded surface. The present work was motivated to develop an abrasive paper at an applied load of 2.1N and sliding speed of 0.66m/s. Crushed silica particles (size: 15-67μm) were used as the abrasive medium. The abraded surfaces were examined with a nanoindentation apparatus to evaluate the variation of nanohardness with sliding time on a nanometer scale. A cross sectional transmission electron microscope (TEM) technique was also employed to clarify the structural changes in the region close to the abraded surface. The present work was motivated to develop a new technique to understand the wear behavior of steels. The present work was motivated to develop a new technique to understand the wear behavior of steels.


For vacuum applications, lubricants which have low vapor pressure and good tribological performance are required. Solid lubricants have been widely adopted for vacuum applications due to their low vapor pressure. Thin film lubricants and high-temperature solid lubricants are discussed as the latest alternatives to fluorine grease. These lubricants feature lower outgassing for clean and vacuum environments. One big drawback of solid lubrication is limited life. We investigated the frictional behavior of molybdenum disulfide (MoS2) films, characterized by high performance ultra-high vacuum (UHV) manipulation mechanism and high-temperature drive devices. We focused sputtering coating method and Molybdenum Disulfide (MoS2), since this material has been reported to be suitable in vacuum, but not well been investigated. As a first step, we have carried out basic study of MoS2 thin film to find out suitable sputtering parameter. It was deposited on martensite stainless steel (SUS440C) substrates, using radio frequency magnetron sputter. Their tribological characteristics in UHV and high-temperature was evaluated by vacuum friction measurement system, high-temperature friction measurement system based on Bowden-Leben type and abrasion measurements by Stylus Surface Profiler.

**TR-ThP3** Structure of Thin Diamond-Like Carbon Films and its Relationship to its Tribological Performance, F.J. Flores-Ruiz, F.J. Espinoza-Beltran, M.O. Vazquez-Lepe, A. Herrera-Gomez, CINVESTAV-Unidad Queretaro, Mexico

Diamond-like carbon (DLC) films are promising materials for dry-contact applications where resistance to surface damage or lubricating performance is required. In the present work, the structure of 3 and 10 nm Focus Cathodic Arc (FCA) carbon films and 3 and 10 nm plasma carbon films grown by filtered cathodic arc (FCA) are studied. The films were deposited on Si(OH)4 and Octadecyltrichlorosilane (OTS) Self-Assembled Monolayer (SAM) coatings. The gas flow rate induced change in ion partial pressure and hence supersaturation. The concentration of species is related to the distribution of sp2 and sp3 carbon within the film was assessed with methods described elsewhere [1]. Atomic force microscopy (AFM) with a silicon tip was used to study the friction coefficient μ on the nanoscopic scale. It is defined by the Amonton law, μ = Ff/Fn, where Ff and Fn are the normal and the lateral force applied to a probe [ii]. The calculation of the normal and lateral force applied during the process of sliding the tip over the sample was done according to the methodology proposed by Carpick [i].

The present investigation of unidirectional abraded surfaces of a martensitic (0.4 wt%Cr) steel with silicon (1.5wt%Si), chromium (1.5wt%Cr) and molybdenum (1.9wt%Mn) addition elucidates the work hardening and the softening near the surface layer caused by abrasion, particularly its relation to the wear behavior. The abrasion testing was performed using a pin-abrasion apparatus in which a small pin of the specimen was ground on an abrasive paper at an applied load of 2.1N and sliding speed of 0.66m/s. Crushed silica particles (size: 15-67μm) were used as the abrasive medium. The abraded surfaces were examined with a nanoindentation apparatus to evaluate the variation of nanohardness with sliding time on a nanometer scale. A cross sectional transmission electron microscope (TEM) technique was also employed to clarify the structural changes in the region close to the abraded surface. The present work was motivated to develop a new technique to understand the wear behavior of steels. The present work was motivated to develop a new technique to understand the wear behavior of steels.


**TR-ThP4** Tribological Investigations of Octadecylphosphonic Acid (ODP) and Octadecyltrichlorosilane (OTS) Self-Assembled Monolayers: A Comparative Study of MEMS-type Interfaces, N. Ansari, Auburn University, S. Barkley, Luther College, C. Doussin, M. Deram, N. Eigenfeld, Saint Olaf College, O. Mathews, Luther College, A. Poda, W.R. Ashurst, Auburn University, B.P. Borovsky, Saint Olaf College, E.D. Luther, Luther College

Microelectromechanical systems (MEMS) are critically-limited by interfacial phenomena such as friction and adhesion. One strategy to reduce friction between MEMS surfaces is to coat them with molecularly-thin self-assembled monolayer (SAM) coatings. Historically, silicon MEMS have been coated with silane-based SAMs, such as octadecyltrichlorosilane (OTS). However, continued progress in the development of MEMS may require new material systems to be employed. Therefore, in this study, we have investigated the frictional properties of octadecylphosphonic acid (ODP) monolayers deposited on aluminum oxide surfaces, across speed regimes. Measurements using an atomic force microscope (AFM) and separately using a nanoindenter-quartz crystal microbalance system were performed each with a microsphere-terminated probe. This allows for a comparative study between different velocity regimes using contacts with similar sizes, pressures, surface roughnesses, and interfacial chemistries. AFM colloidal probe friction measurements indicate that for a bare tip the friction is related to the adhesion force between tip and sample. However, the friction coefficient values do not show significant changes as the strength of adhesion varies, indicating that the value of the friction coefficient depend on the contributions of atomic bonds at the surface.
observed strong evidence of transfer of the ODP molecules to the tip when the tip is uncoated. The results presented in this study are significant contributions towards our goal of better understanding the frictional properties of phosphonate SAMs in pursuit of alternative MEMS materials.

**TR-ThP6 The Effect of Test Parameters on the Tribocorrosion Behavior of Multilayers, M. Flores, O. Irienes, J. Garcia, E. Rodriguez, Universidad de Guadalajara, Mexico, L. Huerta, Universidad Nacional Autonoma de Mexico**

The multilayer coatings can improve the corrosion and wear resistance of materials for biomedical applications. The tribocorrosion behavior of TiAlN and TiAlPₓN coatings and TiAlPₓN-TiAlPₓ multilayers immersed in a corrosive environment was investigated. The coatings were deposited on 316L stainless steel and Ti6Al4V alloys by magnetron sputtering. The period thickness of multilayers was 300 nanometers and the total thickness was 3.6 microns. In order to evaluate the influence of the environment, the corrosive test was studied using open circuit potential (OCP) measurements and potentiodynamic polarization techniques in saline and a Ringer’s solution. For the tribocorrosion test a counterbody of Alumina with 10 mm diameter was used. The loads used were from 1 to 5 N, the oscillating frequencies were 1Hz to 5 Hz. The electrochemical noise measurements were performed during, and after the sliding and scratch tests. The structure and composition of multilayers were studied by means of XRD, XPS and RBS techniques. It was found that the codeposition of Pt and TiAlN-TiAl multilayer can improve the wear-corrosion resistance of materials for biomedical applications. The tribocorrosion behavior results in terms of the coefficient of friction showed a dependence against the force and the sliding frequency.

**TR-ThP7 Tribological Properties of Electron Beam Excited Plasma Nitrided Surfaces, P. Abruja, J. Miyamoto, Meijo University, Japan**

Tribological properties play significant role on the performance of tool steel surfaces. Here, application of electron beam excited plasma nitriding and its effect on tribological properties is described. The technique eliminates the formation of the brittle and rough compound layer that is common in nitriding processes. The hardening process is done through diffusion of the plasma species in to the subsurface of the treated material without altering the initial surface finish. The applications of the process can be in areas of hard coating where adhesion of the coating material with the tool steel is of significant importance.

The experimental tool steel material is SKD 61 with a chemical composition of 0.36% C, 5.05% Cr, 1.21% Mo, 0.83% V, 0.92% Si, 0.43% Mn, 0.008% P, 0.001% S, Fe bal. The sample was heat treated, hardened and triple tempered to a hardness of 630 Hv. The sample was then treated in a nitrogen plasma produced by a beam current of 8 A under a working pressure of 0.4 Pa. The temperature was set at 500 degrees centigrade throughout the treatment time. The experimental set up includes bias terminals that reduce the ion density within the vicinity of the tool steel material. This is done to reduce nitriding due to ion and increase the chance of nitriding due to neutral species within the plasma. The cross sectional hardness distributions and wear measurements of the nitrided tool steels were examined to determine the mechanical and tribological surface properties. The surface has no trace of the compound layer that is usually observed in the ion nitriding processes. This is also confirmed from the X-ray Diffraction peaks, as there is no visible Fe,N and Fe,N peaks observed. These results are attractive as they open new areas of application especially in the coating industry where adhesion remains to be the limiting factor in lots of the hard coatings that protect cutting and forming tools against wear.

**TR-ThP8 Parameter Optimization of Ion Plated Nickel-Copper-Silver Lubrication, M. Danyuk, A. Dhingra, University of Wisconsin Milwaukee**

In this paper we present a connection between argon ion flux with ion mixing and rolling contact fatigue (RCF) life of a thin solid nickel-copper-silver film lubrication on ball bearings tested in high vacuum. Using a Langmuir probe we measure plasma properties and ion flux and then calculate plane stress within the film during deposition using a validated model found in the thin film science literature. Experiments reveal that there is an inverse relationship between ion flux and RCF life for most deposition voltage and pressure combinations tested, specifically, 15.5 to 18.5 mTorr and 1.5 to 3.5 kV. For voltages up to 2.5 kV, RCF life decreases as deposition voltage and ion flux increase. Experiments also confirm that as ion flux increases deposition rate decreases due to sputter removal from the ball surface. For voltages between 2.5 and 3.5 kV interlayer mixing and contamination of the 100 nm thick coating reduces RCF life even as ion flux decreases with decreasing process pressure within a constant power process. At ion energy greater than 2.5 kV and ion flux above 10¹³ cm⁻² s⁻¹, SRIM results suggest elemental mixing of copper and nickel at the interface and this is confirmed using Auger Electron Spectroscopy (AES) on steel and Si₃N₄ 5/16” diameter balls.

**TR-ThP9 Shear-induced Tribofilm Formation: Boron Containing Molecules on Copper, B.P. Miller, O.J. Furlong, W.T. Tysoe, University of Wisconsin Milwaukee**

The push for greener lubricants has steered focus away from compounds containing sulfur and phosphorus. The tribological chemistry can depend critically on the nature of the substrate so that a good lubricant additive for one type of surface may not be applicable to another. In particular, the lubrication of sliding copper-copper interfaces in electrical motors provides a challenge due to the requirement for a conducting interface. Boron containing molecules have been proposed as potential green lubricants since environmentally safe boric acid is a prominent decomposition product in the presence of water. The following investigates the chemistry and frictional properties of alkoxy dioxaborolane (borolane) on copper surfaces in ultrahigh vacuum (UHV) to determine the precursor at room temperature that can potentially form a tribofilm needed for the lubrication of sliding copper-copper contacts. Temperature programmed desorption (TPD) and X-ray photoelectron spectroscopy (XPS) experiments provide the background information for investigating the frictional properties of borolane. A sliding copper-copper interface is exposed to borolane under UHV conditions, and a significant reduction in friction is found from the clean-surface values. A lasting tribofilm persisted even after the borolane dosing was stopped, and carbon was found on and below the surface in the wear scar region by in-situ Auger spectroscopy. Because the interfacial temperature rise under the experimental conditions used to measure friction is <1 K, the tribofilm formation is shear- and not thermal-induced.
Spectroscopic Ellipsometry: Future Directions and New Techniques

Moderator: H. Wormeester, MESA, The Netherlands

8:20am EL+AS+EM+MS+PS+TF-FrM1 Current Trends and Future Outlook for Spectroscopic Ellipsometry, J.N. Hijfler, B. Joks, C.M. Herzinger, T.E. Twidal, J.A. Woollam Co., Inc.

This talk reviews the significant developments in spectroscopic ellipsometry (SE) in areas including extending spectral range, improving accuracy, and enhancing speed. Current SE applications o ws much to hard ware and software developments of the past. Thus, today’s research efforts may reach full potential for applications years or even decades from now. With this in mind, we point to the current state-of-the-art and what this may mean for future SE applications.

Three important areas will be explored. First, there has been a continual trend to expand SE wavelength range. This has included extensions to both shorter and longer wavelengths. For the latter, there is current development into the THz. More immediate benefit may come from smaller SE extensions from the ultraviolet to the near infrared. For example, further near-infrared extensions help to characterize modern transparent conductive oxides (TCOs), used in both inorganic and organic photovoltaic stacks.

Second, we look at the search for improved SE accuracy. Substantial improvements have come with the development of new ellipsometer technologies, progressing from rotating analyzer/polarizer to rotating compensator and now dual-rotating compensator ellipsometers. In addition to improved accuracy, this technology provides advanced measurements, including the complete Mueller-matrix. This will open SE characterization to new applications of anisotropic, nanostructured, and even patterned thin films. Accuracy enhancements must be compatible with the expanding SE spectral range. Infrared SE has overcome many non-ideal optical components to provide measurements competitive to standard FTIR measurements.

Third, we look at the quest for improved measurement speed. This development is constrained by the previous requirements. The benefits of a wide spectral range generally outweigh speed requirements; otherwise laser-based ellipsometry would still have a strong foot-hold. Thus, compromises are made depending on application. Current instrumentation typically utilizes detector arrays for multi-channel SE measurements.

To conclude, we will look at the SE outlook and how it may take advantage of wavelength range, accuracy, and speed. In-line and in-situ SE measurements show special promise. Significant improvements in instrumentation, computing speed, and software are now making these applications more feasible. In addition, there are novel ideas to provide sample access and overcome non-ideal measurement conditions for in-line and in-situ SE. Significant progress in many different areas promises to extend ellipsometry into new areas – many of which are being studied by researchers today.


Nanometer-scale dielectric thin films with controlled thickness and sheet carrier density are of great interest for the terahertz (THz) frequency region and beyond. The MIR measurements are performed for analysis of the heterostructure constituents’ layer thickness, phonon modes, and the existence of significant volume charge carrier concentrations could be excluded. NIR to VUV ellipsometry is used to determine the thickness of the thin top layers. From a line-shape analysis in the VUV spectral range the optical constants of the HfO2 passivation layer could be extracted. An amorphous structure of the HfO2 passivation layer could be confirmed by comparison with existing studies in literature.

THz dielectric anisotropy in the THz spectral range is performed for characterization of the 2DEG channel parameters. A classical Drude model for free charge carrier contribution to the dielectric function was applied to determine the sheet density, the carrier mobility, and the effective mass of the 2DEG electrons. The electron effective mass of (0.22±0.04) m was extracted here using OHE corroborates the values found in previous SdH and CR studies. The values for the high-frequency sheet density and carrier mobility obtained by the optical investigations in the THz spectral range are in excellent agreement with results from dc EHE measurements indicative within linear Boltzmann transport theory for frequency-independent carrier scattering mechanisms of the 2D carrier distribution.

9:40am EL+AS+EM+MS+PS+TF-FrM5 Vector-Magneto-Optical Generalized Ellipsometry on Sculptured Thin Films, D. Schmidt, C. Briley, E. Schubert, M. Schubert, University of Nebraska - Lincoln

Sculptured thin films are self-organized and self-assembled three-dimensional nanostructures with tunable geometries. These artificial nanostructured thin films exhibit highly anisotropic physical properties, which mainly depend on their specific geometry.

Slanted, highly-spatially coherent, columnar nanostructure samples were prepared by glancing angle electron-beam deposition. Glancing angle deposition is a bottom-up fabrication technique that employs a physical vapor deposition process at oblique angles where the trajectory of the incoming particle flux is not parallel to the substrate normal. The technique allows to engineer the columnar film structure and is today amongst the most promising self-organized fabrication processes in micro- and nanotechnology.

We present and discuss the novel approach of vector-magneto-optical generalized ellipsometry on ferromagnetic permalloy nanostructured thin films carried out at room temperature. Investigations have shown that the metal alloy thin films are highly transparent, reveal strong form-induced birefringence, and exhibit intriguing magneto-optical anisotropy. Spatial magnetization orientation hysteresis and magnetization magnitude hysteresis properties are studied using a three-dimensional Helmholtz coil arrangement. This particular octupole setup allows for arbitrary magnetic field directions at the sample position with field strengths up to 200 mT while optical access is granted for reflection and transmission-type ellipsometry measurements. Analysis of data obtained within this unique vector-magneto-optic setup reveals magnetization anisotropy of the permalloy slanted nanocolumns and gives insight into switching behavior of confined magnetic domains.

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targeted optical properties of STF in the THz range can be achieved by variation of slanting angle, lateral column density, and material.

Ellipsometric porosimetry (EP) is a non-contact, non-destructive technique that is cited as a reference technique for porous thin film analysis [1, 2]. As it is based on a spectroscopic ellipsometric measurement, the technique allows the precise determination of the refractive indices and thickness of the porous films. The advantage of these EP tools is that the combination of this well-established spectroscopic ellipsometric (SE) technique with a suitable adapted adsorption chamber permits access to all the information obtained by classic adsorption experiments (e.g. BET) on thin films with an excellent sensitivity. Information such as open and closed porosity, pore size distribution etc... can be thus obtained.

In addition the EP allows access to a multitude of information that the classic equipment does not. For example, Spectroscopic Ellipsometry allows to follow the variation of the sample thickness during the adsorption experiment, leading to the determination of the Young’s Modulus for the thin films. This will be presented. The technique is highly sensitive to the detection of interfaces; it is thus possible to detect a porosity gradient or to study a multilayer structure and thereafter simultaneously plot the two corresponding adsorption isotherms [3]. In the same manner, the instrument permits the use of different gases adsorptive in order to tailor the probe molecule to the morphology and to the chemistry of the porous layer at ambient temperature [4]. We thus obtain information on the chemistry of the pores within the layer, before, during and after the adsorption experiment. Recent developments include the implementation of the FTIR interferometer SE extension to the EP system. It allows a precise characterization of the chemistry of the pores within the layer. We thus obtain information on the chemical bonds present in the layers before, during and after the adsorption experiment. Preliminary results will be presented.

Specifically, this fundamental technique permits the thorough characterization of porous thin film samples. We will demonstrate some of the different features of the EP technique with regards to the morphological and chemical properties of the porous thin films. Additionally, we will illustrate the technique for various thin film applications such as solgel thin films, nanofilms for catalysis, photovoltaic cells, fuel cells, optical sensors, and bio-compatible materials to name but a few.

References:


or equal to 90°, the [211] orientated growth is supported. The [111] surface of the rock salt ErAs is typically a high-energy surface; however, the Ga-rich (1 -1 -1) plane provides a flux mediated epitaxial growth surface for the ErAs analogous to a vapor-liquid-solid type of growth. In-situ RHEED, LEED and STM surface studies will be presented along with a detailed growth model to explain differences in the growth process and in nanorod formation for different substrates and substrate orientations.

Supported by AFOSR FA9550-10-1-0119 and ARO W911NF-07-1-0547.


9:00am EM+SS-FrM3 Bulk Topological Insulators and Superconductors: Discovery and the Frontier, M.Z. Hasan, Princeton University

INVITED

While most known phases of matter are characterized by broken symmetries, the discovery of quantum Hall effects (1980s) revealed that there exists an organizational principle based on topology rather than broken symmetry. In the past few years, theory and experiments have suggested that new types of topological states of matter exist in certain insulators without any applied magnetic field. These topological insulators are characterized by a full band gap in their bulk and gapless conducting edge or surface states protected by time-reversal symmetry. Unlike the quantum Hall systems, the topological insulators can be doped into superconductors and magnets revealing the interplay between topological order and broken symmetry order. In this talk, I will briefly review the basic theory and highlight the experimental developments in topological insulators. I will then conclude by drawing connections between the emergent novel physics and their potential applications.

10:20am EM+SS-FrM7 Inter-band GaN/InGaN/GaN Tunnel Diodes, S. Krishnamoorthy, D.N. Nath, S. Bajaj, S. Rajan, Ohio State University

The III-Nitride material system has demonstrated its potential for a broad range of optoelectronic and electronic applications. However there are no reports of efficient III-Nitride tunnel junctions due to the large band gaps in this material system. In this work, we show that with unique properties such as the polarization, tunneling can be enhanced using band bending over smaller distances in nitride heterostructures, leading to record reverse and forward tunneling current density for the III-nitride material system.

We have designed and demonstrated GaN/InGaN/GaN tunnel junction with a record high current density of 118 A/cm² at a reverse bias of 1 V by utilizing a 6.4 nm thick In0.53Ga0.47N barrier material. N-polar p-GaN/n-GaN heterostructures were fabricated for tunneling studies. A zero bias was grown by plasma assisted molecular beam epitaxy by choosing the critical thickness of InGaN barrier appropriately. The tunnel junction sample shows five orders of magnitude higher current at a reverse bias of 1 V as compared to a standard p+nn+ GaN sample indicating efficient tunneling across the InGaN barrier. The tunneling turn-on close to zero voltage could be ascribed to the larger band offsets and high current density achieved in this work demonstrates the potential of polarization-engineered tunnel junctions.

Two distinct regimes of transport are identified based on the temperature dependent I-V measurements. At lower reverse bias, defect assisted tunneling with strong temperature dependence is found to dominate. In this regime, a plot of ln(I/V) vs V¹/² shows a linear behavior suggesting a Frenkel- Poole emission mechanism due to the high field in the InGaN quantum well. A direct band to band tunneling regime resulting in weak temperature dependence that arises from band gap variation with temperature is observed from a reverse bias of 1 V. A decrease in current density is observed with increase in temperature in the range of 77-150 K and this can be attributed to the presence of band tail states which has been observed previously in In face GaN.

We discuss the design of these quantum well tunnel junctions. Although higher indium compositions yield higher band to band tunneling probability, calculations using a simplified Kane model reveal that the wider depletion region in n GaN due to higher band offset considerably reduces the net tunneling probability. Calculations also reveal the need for very high doping in the n GaN layer so as to minimize the depletion region thickness in order to achieve very high current densities in such polarization charge assisted tunnel junctions. These calculations can guide future tunnel junctions with better performance characteristics.

10:40am EM+SS-FrM8 Probing Surface-Induced Fluctuations in Organic Materials using an Atomic Force Microscope, N.C. Hoepker, S. Lekkala, R.F. Loring, J.A. Marohn, Cornell University

The development of organic electronics calls for new tools to study organic thin films. By measuring the frequency noise experienced by a cantilever near a surface, we are able to microscopically probe organic materials. In previous work, we used an Atomic Force Microscope to measure frequency noise due to dielectric fluctuations as a function of cantilever height and voltage over a thin film of polyvinyl acetate. In parallel, we have developed a zero-free parameter linear-response theory of thermally induced dielectric fluctuations. Although this method successfully describes our observations, it does not account for carrier mobility. Dielectric fluctuations are characterized by zero bias, and it is possible that the fluctuations are enhanced by the presence of mobile carriers.

Having understood dielectric fluctuations, we are now investigating fluctuations induced by carrier motion in polymeric semiconductors. Charge transport in these devices is not well understood. Previous work indicates that the ratio of diffusion constant to mobility in these materials violates what is predicted by the Einstein relation. In addition, there is an ongoing controversy on the charge density and electric field dependence of mobility. While the correlated-disorder model correctly predicts the electric field dependence of mobility, models that predict a density dependence of mobility rely on uncorrelated site-to-site energies.

A resolution of these controversies calls for new tools to study carrier motion in organic semiconductors. By measuring the frequency fluctuations experienced by a cantilever near a surface, we are able to microscopically probe carrier motion in organic materials. Comparing our observations over a poly(3-hexylthiophene) transistor to a calculation based on free diffusion, we find that while theory overestimates the observed fluctuations, it predicts the correct spectral shape and distance dependence of the fluctuations. Even at high gate bias, the observed cantilever frequency fluctuations differ from what we expect based on free diffusion and on the measured carrier mobility. This discrepancy indicates a breakdown of the Einstein relation. Further we present a number of different charge hopping models. We find that the predicted cantilever frequency noise is very sensitive to the details of the model, indicating that frequency noise spectra are a vital tool for selecting appropriate charge transport models.


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11:00am EM+SS-FrM9 2011 AVS Albert Nerken Award Lecture - Electron Spectroscopy of Reconstructed Surfaces: From Silicon to Graphene, J.E. Rowe*, North Carolina State University

INVITED

Surface reconstruction of silicon (and other materials) refers to the process by which atoms at the surface of a crystal assume a different structure than that of the bulk and has been extensively discussed and reported at a number of AVS meetings from the early 1960’s until present time. In the 1970’s a number of electron spectroscopy methods were applied to study this effect and many atomic models were proposed. Early 1970’s experiments using electron energy loss spectroscopy and photoemission spectroscopy are described along with methods that led to the now accepted dimer model (later confirmed by STM) for the Si(100)(2x1) and most other reconstructed (100) semiconductor surfaces. Additional core-level synchrotron spectroscopy are described along with very recent studies which include adsorbate-induced surface reconstruction and the role of interface reconstruction of SiC(0001) used for the growth of graphene and studied by STM and STS. Spectroscopy has continued to play an important role even during the past 25 years after the discovery of atomic-scale imaging by STM of the Si(111)x7x7 reconstruction. Both early and more recent studies of reconstruction by the author are reviewed.

11:40am EM+SS-FrM11 Molecular Motion Confined to Self-Assembled Quantum Corrals, E. Vitambay, R.A. Rosenberg, N.P. Guisinger, Argonne National Laboratory

Engineering molecular superstructures on metals opens great possibilities for the control and exploration of complex nanosystems for technological applications. Of particular interest is the use of chiral molecules, such as alanine, to build self-assembled nanoscale structures for the trapping of the two-dimensional free electron gas of a metal. In the present work, molecules of D- or L-alanine were deposited on Cu(111). Scanning tunneling microscopy and spectroscopy revealed the formation of a uniform network of hexagonal pores of average diameter ~1.2 nm. Each pore acts as a quantum corral by confining the two-dimensional electron gas of the Cu(111) surface state. Furthermore, excess alanine molecules were trapped at the inner perimeter of the hexagonal pore, and were observed as rotating or immobile spatial states. This study demonstrates the engineering of one of the smallest quantum confined structure, and the dynamics of molecular motion within these potential wells.

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* Albert Nerken Award Winner
Energy Frontiers Focus Topic
Room: 203 - Session EN+AC-FrM

Materials Challenges for Nuclear Energy
Moderator: L. Petit, Daresbury Laboratory, UK

8:20am EN+AC-FrM1 Multi-Electron Correlation in UO2 from Soft X-ray Spectroscopy. J.G. Tobin, S.W. Yu, Lawrence Livermore National Laboratory

Resonant Inverse Photoelectron Spectroscopy (RIPES) and X-ray Emission Spectroscopy (XES) have been used to probe the electronic structure of Uranium Dioxide, UO2. From these variants of soft x-ray spectroscopy, the nature of the main and satellite features at the U4f/5 edge can be ascertained. This leads to important insights into the multi-electronic correlations underlying the ubiquitous satellite features of UO2 and implications for future experiments with Pu.

Lawrence Livermore National Laboratory is operated by Lawrence Livermore National Security, LLC, for the U.S. Department of Energy, National Nuclear Security Administration under Contract DE-AC52-07NA27344. This work is funded by the DOE Office of Science, Office of Basic Energy Science, Division of Materials Sciences and Engineering. The Advanced Light Source (ALS) is supported by the Director, Office of Science, Office of Basic Energy Sciences.

8:40am EN+AC-FrM2 New Results from the DOE EFRC on the Materials Science of Actinides. P.C. Burns, University of Notre Dame, G. Sigmon, EFRC INVITED

The Materials Science of Actinides EFRC has three major themes: (1) Complex actinide materials, with complexity arising from chemistry, structure, and properties; (2) Nanoscale control of actinides; and (3) Behavior of actinide materials in extreme environments of pressure, temperature and radiation fields (including coupled effects). This presentation will cover several recent results that span the range of themes in the EFRC, with emphasis on those that are most significant to the challenges of nuclear energy.


We have performed x-ray absorption experiments on uranium dioxide (UO2) at the O 1s, U 4d, and U 5f electron energies. After comprehensive energy calibrations for O 1s, U 4d, and U 5f spectra, we have used the U 4d and U 5f spectra to sort the energetic positions of the 5f and the 6d states in the unoccupied band unambiguously. This demonstrates conclusively that UO2 is an f-f Mott-Hubbard insulator, where the electronic repulsion between f electrons is responsible for the insulating state. Calculations performed within the U-corrected generalized gradient approximation of the optical response of UO2 permit direct comparison with the absorption spectra and confirm the experimental results.

Lawrence Livermore National Laboratory is operated by Lawrence Livermore National Security, LLC, for the U.S. Department of Energy, National Nuclear Security Administration under Contract DE-AC52-07NA27344. This work is funded in part by the DOE Office of Science, Office of Basic Energy Science, Division of Materials Sciences and Engineering. This work is also funded in part by Laboratory Directed Research and Development (LDRD) Program (10-SI-016) of Lawrence Livermore National Laboratory. The Advanced Light Source (ALS) is supported by the Director, Office of Science, Office of Basic Energy Sciences, of the U.S. Department of Energy under Contract No. DE-AC02-05CH11231. We would like to thank Ian Hutcheon, Patrick Allen, Anthony Van Buuren, Trevor Wiley, and Joseph Zaug for valuable discussions. POV would like to acknowledge CONACyT Mexico.

10:00am EN+AC-FrM6 Atomic Models for Actinide-Acetate Oxide Interfaces. S.M. Valone, Los Alamos National Laboratory INVITED

An important class of materials problems of great interest to nuclear energy production consists of composites of metals and metal oxides, and in particular, actinide metals and actinide oxides. Individually, either type of material, actinide or oxide, can involve strong electron correlation effects. At an interface, the situation becomes even more complex. In traversing a metal-oxide interface, a radical compositional change is encountered. Most atomic models address only metals, or only ceramics, but rarely both. In addition, the actinide oxides themselves exhibit multiple oxidation states, depending on the composition. Thus, in traversing an interface, the oxidation state need not change abruptly. As a result, these sorts of interfaces present new challenges that must be met in order to understand this important class of materials systems. To address these challenges, a new, “fragment” model Hamiltonian is constructed at the atomic level, as opposed to the one-electron model Hamiltonians that underlie tight-binding and density functional theory methods. The model encompasses both actinides and actinide oxides, and provision is made for transitioning gradually through multiple oxidation states. The extremes of the models, the dioxides and the metals, map closely to existing models for these materials. The model for metals conforms generally to a modified embedded atom method (MEAM), meaning that the embedding function (atomic site energy model) is analytical. The differences between the fragment Hamiltonian potential and the existing MEAM models appear in the explicit form of the embedding function and in the fact that there are two distinct terms in the embedding energy in the new model. The second term is critical to strongly-correlated-electron materials, as it is an atomic analogue to terms appearing in Hubbard models. The model also possesses a sense of electron hopping that imparts ways to both regulate the net charge on sites in the material and to change important energy gaps that determine metallic and insulating behavior.


For future generation nuclear fuel cycle, actinide co-conversion processes are one option for the co-management of actinides. Oxalic acid is a well-known reagent to recover actinides thanks to the very low solubility of Am(IV) and An(III) oxalate compounds in acidic solution. Therefore, considering mixed-oxide fuel or considering minor actinides incorporation in mixed-oxide fuels for transmutation, oxalic acid co-conversion is convenient to synthesize mixed oxalate compounds, precursors of oxide solid solutions.

Up to now, only U(IV)-Ln(III) oxalate solid solutions have been completely structurally described started from single-crystals. In these oxalate compounds, a mixed-crystallographic site which accommodates both uranium and lanthanide elements in spite of their different charges has been established. The extent of this study on powder compounds allowed to examine the influence of (AnIV) and (AnIII) nature. The switch of actinide’s nature causes unexpected structural modifications underlying the complexity of specific transuranium elements physical chemistry and the need to pursue studies on single crystal on these actinides.

As the existing oxalate single crystal syntheses are not adaptable to the actinide-oxalate system, several original crystal growth methods allowing the formation of mixed actinide oxalate crystals were first developed. Applied to the mixed actinide systems, they lead to the formation of the first mixed Am(IV)-An(III) oxalate single crystals. The results, including the different structural resolutions, are presented.

10:40am EN+AC-FrM8 Recent Work on Magnetism, Actinides and Defects at ORNL. G.M. Stocks, B.C. Larson, Oak Ridge National Laboratory INVITED

The extent to which the collective effects of defects can be manipulated and controlled yields the combination of structural materials properties – strength, toughness, and resistance to degradation in extreme chemical and radiation environments. In this presentation I shall outline the scope of studies of fundamental properties of defects such as formation energies and radiative induced defect displacement cascades being conducted within the Center for Defect Physics (CDP) at Oak Ridge National Laboratory and its partner institutions. Within the CDP, the focus is on the quantitative measurement and direct quantum simulation of defects at the level of unit dislocation and cascade events. For dislocation interactions, the focus is on high-spatial-resolution techniques such as X-ray microscopy and electron diffraction, which measure the local strains/stresses near isolated defects directly and thereby quantify the interactions and dynamics of defects in the bulk. For displacement cascades, the focus is on quantitative measurement of the formation and time evolution of energetic-ion-induced atomic displacement cascades using ultrahigh-resolution time-resolved X-ray diffuse scattering measurements made possible by the ultrahigh brilliance of femtosecond X-ray pulses produced at the Linac Coherent Light Source (LCLS) and the time-averaged brilliance of the Advanced Photon Source (APS). Theoretically, the focus is on developing high-fidelity models that treat spin and ion dynamics on an equal footing and to address system sizes and time scales commensurate with experiments. I will outline early progress with respect to addressing the feasibility of experimentally observing unit events and on developing ab initio electronic
structure based theories of combined atomistic and spin dynamics. For the latter, I will show preliminary results for Fe that address the importance of the disruption of the magnetic state of Fe caused by the introduction of defects such as dislocations and displacement cascades that are based on large scale (~10,000 atom) models and order-N electronic structure methods.

Work supported by the “Center for Defect Physics in Structural Materials” which is a Department of Energy, Office of Science, Energy Frontier Research Center (EFRC).

Graphene and Related Materials Focus Topic Room: 208 - Session GR+MS+EM-FrM

Graphene Device Physics and Applications

Moderator: M. Arnold, University of Wisconsin-Madison

8:20am GR+MS+EM-FrM1 Fabrication and Characterization of Graphene p-n Junction Devices, J.U. Lee, University at Albany-SUNY

INVITED

Graphene is a newly discovered material composed of two-dimensional array of hexagonal carbon atoms. It has a number of unique electronic properties, the most remarkable of which is the zero band-gap light-like linear electronic dispersion, giving rise to Dirac fermions. This feature can be used to make devices based on previously unexplored physical properties. For example, in analogy to optics, we describe new devices based on optics-like manipulation of electrons.

Our devices are based on graphene and bi-layer graphene p-n junctions doped using electrostatic doping techniques from buried split gates. In the present context, graphene p-n junctions do not rectify, i.e. behave as semiconduc tor diodes. Instead, in graphene p-n junctions, carriers launched from a point contact from one side of the junction are able to refocus back to a point on the other side of the junction. This behavior, known as the Veselago effect, can be the basis for new logic devices for replacing Si CMOS. In addition, using the same platform, we describe interconnect structures that can be reconfigured. Together, we envision a new circuit paradigm based on these components that seamlessly reconfigure between devices and interconnect components.

In this talk, we describe the details of graphene p-n junction fabrication and characterization, and circuits that are enabled by the p-n junction devices. The devices are fabricated at CNSE’s state-of-the-art 300nm Si wafer fabrication line using processing techniques that leave atomically flat top oxide surface above the patterned split gates. For characterizing the p-n junctions, we perform transport and SPM measurements.


Graphene has shown successful application in RF transistors and frequency doublers where its high mobility and high saturation velocity translate into operation at high frequencies while utilizing little power. However, a major detraction to graphene development for other device applications is that it does not have a band gap. The lack of a band gap means that graphene’s current cannot be turned off. Bilayer graphene is regarded as one possible solution to this problem, since bilayer graphene is capable of developing a band gap if the symmetry of the system can be broken. That said, bilayer graphene (from exfoliation or growth) forms a highly ordered A-B stack of the top graphene sheets resulting in little to no band gap, unless a high electric field can be applied.

In this presentation, we will demonstrate a novel method for creating bilayer graphene where a single layer of CVD graphene grown on Cu is bonded to a single layer of epitaxial graphene grown on Si-face SiC. This process results in a bilayer system that has a built-in asymmetry that yields unique physical and electrical properties not previously observed. For example, we demonstrate that the transfer of CVD graphene to epitaxial graphene results in a morphological transformation of the graphene to a SO2 and that bonding of CVD graphene to epitaxial graphene can avoid the damage caused by the drying step necessary in the poly (methyl methacrylate) transfer method. X-ray photoelectron spectroscopy and Raman microscopy demonstrate that the sheets are coupled together but strained differently, in contrast to a naturally formed bilayer. Electrical characterization of Hall devices fabricated on the novel bilayer show higher mobilities and lower carrier concentrations than the individual CVD graphene or epitaxial graphene sheets alone. Modeling of the electric field produced by opposite doping in the graphene sheets will also be presented, as CVD graphene is typically p-type and epitaxial graphene is typically n-type.

10:00am GR+MS+EM-FrM6 Rectification at Graphene / Semiconductor Junctions: Applications Beyond Silicon Based Devices, S. Tongay, X. Miau, K. Berke, M. Lemaître, B.R. Appleton, A.F. Hebard, University of Florida

Schottky barriers are crucial and necessary device components of metal-semiconductor field effect transistors (MESFETs) and high electron mobility transistors (HEMTs). Here, we report on the formation of Schottky barriers at graphene-multilayer graphene/semiconductor junction interfaces which have been characterized by current density vs. voltage (J-V) and capacitance vs. voltage (C-V) measurements. After graphene transfer onto various semiconductors such as Si, GaAs, GaN and SiC, we observe a strong rectification at the interface, i.e., high (low) resistance in the reverse (forward) bias directions. The J-V characteristics have been analyzed using theoretical transmission theory and the extracted barrier height values are consistent with the Schottky-Mott model. When capacitance is plotted as 1/C^2 vs V, a linear dependence is observed, which by extrapolation to the intercept identifies a built-in potential in that is consistent with the Schottky barrier height extracted from J-V measurements. Graphene's low Fermi energy together with its robust thermal, chemical, structural and physical properties provide numerous advantages when used to form Schottky barriers in device applications: namely, voltage tunability of the Schottky barrier height, stability to high temperatures, resistance to impurity diffusion across the interface, and the use of absorbates to chemically tune the Fermi energy and hence the Schottky barrier height.


It has been known and observed that there forms a Schottky junction between graphene and SiC in epitaxial graphene due to the work function difference and the charge transfer between them. As a result, it is viable to apply the electron beam induce current (EBIC) technique on the epitaxial graphene directly due to the fact that it needs a built-in field and ample electron generation volume to generate EBIC. EBIC is an important characterization technique which identifies electric impurities/defects, detects local built-in field, and measures minority carrier diffusion length. In this paper, we use a FEI SEM equipped with a current amplifier to investigate the spatial mapping of EBIC generation and collection in a two terminal geometry. The incident electron beam generates excited electron-hole pairs in SiC and the minority carriers are collected through the Schottky junction before flowing into graphene. EBIC imaging reveals mesoscopic domains of bright and dark contrast areas due to local EBIC polarity and magnitude, which is believed to be the result of spatial fluctuation in the carrier density in graphene. We also investigate the electron energy dependence, which modulates the EBIC magnitude. With an analytical drift-diffusion current model, we are able to extract the minority carrier diffusion length in the SiC, which is on the order of micro meter and agrees well with other published data.

10:40am GR+MS+EM-FrM8 Potassium-Ion Sensors Based on Valinomycin-modified Graphene Field-Effect Transistors, Y. Sofie, Y. Ohno, K. Machashi, K. Inoue, M. Matsumoto, The Institute of Scientific and Industrial Research, Osaka University, Japan

Highly sensitive ion sensors based on valinomycin-modified graphene field-effect transistors (VGFTs) have been developed to selectively detect K ions. VGFTs are an essential tool for biological applications including human life. Graphene single-layers were obtained by mechanical exfoliation. Graphene FETs were fabricated by conventional e-beam lithography and lift-off method on a thermally grown SiO2 layer. To demonstrate selective detection of K ions, the graphene channels were covered with ion selective membrane, which consisted of polyvinyl chloride and valinomycin. Transfer characteristics of VGFTs in a 100 mM Tris-HCl buffer solution with various KCl concentrations over the range from 10 nM to 1.0mM. With increasing K ion concentration, the solution-gated voltage at the Dirac point shifted toward negative direction. The shifts are due to the accumulation of positively charged K ions surrounded by valinomycin on the graphene surfaces. The electrostatic potential of graphene surfaces exhibit a rather linear dependence on log[K]. These results indicate that VGFTs effectively detected K ions with concentration from 10 nM to 1.0 mM. To investigate selectivity in VGFTs, Na-ion concentration dependence was also measured. The transfer characteristic in VGFTs remained almost constant over the Na ion concentration range between 10 nM and 1.0 mM. These results indicate that VGFT selectively detected K ions with high sensitivity.
11:00am GR-MS+EM-FrM9 Band-gap Generation by using Ionic-Liquid Gate in Bilayer Graphene. Y. Yamashiro, Y. Ohno, K. Maehashi, K. Inoue, K. Matsumoto, Osaka University, Japan

Electric fields were applied to a bilayer graphene to generate a band gap using an ionic-liquid gate instead of the general top-gate structures. The ionic-liquid gate can apply higher electric field than other type of the gates because of its large capacitance and electric strength. In this abstract, the graphene layers were extracted from kish graphite by a mechanical exfoliation and were put on highly doped Si substrates covered with a 300-nm-thick SiO2 layer. Side-gate electrodes were patterned approximately 20 mm away from the channels. An ionic liquid (DEME-TFSI) was put on the bilayer graphene and the side-gate electrode. Electrical characteristics at 300 K revealed that the electrical double layer in the ionic-liquid, which works as a very thin insulator, had 200 times larger capacitance than a 300-nm-thick SiO2 layer. The thickness of electrical double layer was estimated to be 3.75 nm. In electric field dependence measurements, an increase in a sheet resistance of the bilayer graphene channel was clearly observed with increasing the magnitude of electric field in bilayer graphene. On the other hand, the increase in the sheet resistance didn’t appear in the monolayer- and trilayer- graphene. That is why the increasing of the sheet resistance was caused by a band gap generated in ionic-liquid gated bilayer graphene by the electric field.


Graphene films grown by chemical vapor deposition on copper foils and hydrogenation using low kinetic energy were hydro-hydrogenated using low kinetic energy. Graphene films grown by chemical vapor deposition on copper foils and hydrogenation using low kinetic energy were hydro-hydrogenated using low kinetic energy. Baldwin (NRL) Electric fields were applied to a bilayer graphene to generate a band gap. K. Inoue, K. Matsumoto

Our bandstructure calculations on the ground state electronic properties of Graphene, hexagonal Boron Nitride (h-BN), Graphene and h-BN. Our results on doped Graphene indicate that upon electron (hole) doping, the Dirac-point in the electronic bandstructure shifts below (above) the Fermi level and a gap appears at the high-symmetric K-point. Upon co-doping of Graphene by both Boron and Nitrogen a small energy gap between the conduction and valence band appears at the Fermi level, making the CBN nanomaterial a narrow band semiconductor. The energy gap depends sensitively on the degree of doping and on the thickness of CBN layer. These results are in agreement with recent experimental measurements [1,2]. Our bandstructure calculations on the multilayers of Graphene and h-BN indicate that these nanostructured multilayers exhibit semiconducting behaviour with band gap in the range 60-600 meV depending on the relative orientation and thickness of the layers. (Author: Sugata Mukherjee, work done in collaboration with T.P. Kaloni)

1. X. wang et al, Science 324, 768 (2009)

MEMS and NEMS Group
Room: 105 - Session MN-FrM

Characterization of Materials and Structures at the Micro- and Nano-scale
Moderator: M. Metzler, Cornell University

8:20am MN-FrM1 Nanomechanics: Controlling Near-Field Interactions between Mechanical Systems. D. Lopez, Argonne National Laboratory

Metallic and dielectric objects are surrounded by fluctuating electromagnetic fields due to thermal and quantum fluctuations of the charge and current density at the surface of the bodies. Immediately outside the objects, this electromagnetic field exists partly in the form of propagating electromagnetic waves and partly in the form of evanescent waves that decay exponentially with distance away from the body’s surface. These fluctuating electromagnetic modes are responsible for a great variety of near-field phenomena such as the Van der Waals force, the Casimir force, near-field heat transfer, and non-contact friction forces. As devices evolve from micro- to nanoscale structures, these forces become relatively stronger, and their effect cannot be disregarded any further. For example, researchers working to develop NEMS devices need to consider the effects caused by Van der Waals and Casimir forces which can lead to compromise in the range of motion or in the voltages required for actuation. To improve our understanding of these near-field interactions and to develop mechanisms to control them is extremely important for a diversity of seemingly different fields, such as nanomechanics, quantum computing with trapped ions, measurements of gravitational forces at the nanometer scale, and detection of single spins for magnetic resonance force microscopy.

In this presentation I will describe the fundamentals of near-field forces, I will review recent scientific advances regarding manipulation of these interactions in the field of nanomechanics, and I will illustrate novel applications that could be enabled once we are capable of control these forces.

9:00am MN-FrM3 Pull-in Experiments on Electrostatically Actuated Microfabricated Meso Scale Beams. Y. Gerson, I. Sokolov, Tel Aviv University, Israel, T. Nachmias, RAFAEL LTD, Israel, S. Lulinsky, S. Krylov, Tel Aviv University, Israel

Meso scale (hundreds of micrometers to several millimeters) MEMS sensors and actuators are beneficial in applications where large displacements, manufacturability, and ease of integration with existing mechanical and packaging environments are required. In this work we report on the results of characterization and modeling of electrostatically actuated meso scale beams. The beams with clamped ends were 5000 μm long, 150 μm thick and 10, 12 and 15 μm wide and were operated by a parallel plate electrode located at the distance of 20 μm from the beam. The goal of the work was twofold. First, we demonstrate the feasibility of electrostatic actuation of the meso scale devices and ability to achieve relatively large displacement. Second, an electrostatically actuated double-clamped micro beam was viewed as a kind of benchmark problem and was intensively studied. However, the number of reported experimental results, which can serve for validation of models, is limited. We anticipate that our experimental results, obtained using larger meso scale structures and therefore relatively more accurate, could provide a reliable experimental reference for a double clamped beam actuated by a parallel-plate electrode.

The devices were fabricated by deep reactive ion etching (DRIE)-based process from highly doped Si using a silicon on insulator (SOI) wafer with (111) orientation and 150 μm thick device layer. The experimental approach based on the use of SOI wafers allows to fabricate devices with low residual stress and excellent mechanical properties of Si. The devices were operated in ambient air conditions. Linearly increasing (ramp) voltages were applied quasi-statically to the actuation electrode and easily visualized in-plane (parallel to the wafer surface) motion of the devices was registered using an optical microscope and a CCD camera. The response was video recorded, the movie was split into separate frames and the voltage-displacement dependence was built using customized edge detection image processing procedure implemented in Matlab. The critical pull-in voltage varied between 70 V in (nominally) 10 mm wide beam and up to 125 V in 15 mm wide beams. In addition, pull-in behavior of the beams was modeled using several approaches, starting from simplified reduced order models based on the Galerkin decomposition with linear eigenmodes as base functions and up to fully coupled nonlinear large deflection three-dimensional simulations. The actual dimensions of each beam, carefully measured using scanning electron microscope (SEM) were used in calculations. Excellent agreement between the results provided by the model and the experimental data was observed.

9:20am MN-FrM4 Absorption and Emission of Plasmonic Antenna Arrays. K.E. O'Brien, P.H. Holloway, M.R. Davidson, University of Florida

New and more portable means of generating narrow band radiation are of interest, especially in the terahertz (THz) range. One potential method for generating radiation involves photo-mixing over nano/micro scale plasmonic structures. The plasmonic structures can serve as antennas for absorbing incoming photons and conversely emit radiation of a lower frequency. Designs include 2-dimensional arrays of these resonant structures fabricated on Ag thin films using electron-beam lithography and
lift-off. Patterns vary from arrays of linear structures, “bowties,” and interlocking structures. We have shown emission of visible radiation from similar structures when excited by space charge from electrons. The absorption and emission of light by the structures has been measured for micron-scale and nano-scale antenna arrays and has exhibited a polarization dependent behavior. The effect of different antenna structures on the absorption and emission will be discussed.

10:00am MN-FrM6 Fabrication and Characterization of Structural and Electrical Properties of Ultrananocrystalline Diamond Nanowires. X. Wang, University of Puerto Rico, A.V. Sumant, Y. Joshi, L.E. Ocola, B. Kabius, D. Lopez, Argonne National Laboratory

Due to extraordinary mechanical, optical and electrical properties as predicted by theory, there has been tremendous amount of interest in making diamond nanowires (DNWs) and diamond nano-rods (DNRs). Synthesizing or fabricating these nanostructures is proving to be very challenging. To date, only a few attempts have been reported, either by etching single crystal diamond using focus ion beam (FIB) to produce diamond NRs or by coating Si nanowires with nanocrystalline diamond to produce diamond NWs. We report a top-down method based on e-beam lithography and reactive ion etching of ultrananocrystalline diamond produce diamond NWs. We report a top-down method based on e-beam lithography and reactive ion etching of ultrananocrystalline diamond (UNCD) to produce UNCD nanowires (UNCDNWs) with nanowire diameters as small as 30 nm. Since they are produced by lithographic approach (top-down), they can be fabricated at well-defined position with nanometer-scale precision. Compare to other fabrication techniques like FIB, our UNCDNWs maintain intrinsic diamond structure and properties without degradation after fabrication process, which has been confirmed by Raman spectroscopy (ultraviolet and visible), transmission electron microscope (TEM) and electron energy loss spectroscopy (EELS). Preliminary electrical measurement of UNCDNWs will be discussed. The ability to fabricate UNCDNWs provides an opportunity to study the fundamental mechanism of transport processes in UNCDNWs, which will enable new ideas and possibilities for the fabrication of new functional nanoelectronic devices.

10:20am MN-FrM7 Investigation of Heat Transfer Enhancement in Nanofluids with Molecular Dynamics Simulations – Role of Particle Charge and Fluid Polarity. J.D. Schall, Oakland University, A.S. Comfort, U.S. Army RDECOM-TARDEC

Thermal loads are increasing in military vehicles because of the greater use of microelectronics, higher power density engines, and restricted air flow from up-armor kits. Conventional methods to increase heat dissipation, such as increasing heat exchanger size produce an undesired increase in vehicle weight and packaging issues. One approach to mitigate these issues is the development of heat transfer fluids with improved thermal transport properties. Nanofluids are suspension of nanometer sized particles in solvent, and represent a potential method to increase the effective fluid thermal conductivity and heat transfer coefficient of coolants without creating the adverse effects found in larger particle suspensions, such as settling, clogging, and abrasion. Since their introduction by U.S. Choi in 1995, a great deal of uncertainty about the mechanisms of enhanced thermal conductivity of nanofluids continues to employ researchers and limits the development of optimized nanofluids in heat transfer applications. In this paper, molecular dynamics simulations are used to investigate heat conduction between model particle surfaces separated by a liquid layer. In particular, effects of base fluid charge, polarity, and nanoparticle surface charge on the solid-liquid interface liquid structure, thermal (i.e. Kapitza) resistance, and thermal conductivity are investigated. Results are compared with previous simulations from the literature which used simple monoatomic models interacting through Lennard-Jones potentials.

10:40am MN-FrM8 Novel CMOS MEMS Double Parallel Plate Capacitive Tactile Sensors For Blood Flow Monitoring. C.J. Hsieh, J.C. Liu, C.T. Sun, Y.C. Lin, W.-C. Tian, National Taiwan University

This research focuses on the developments and characterizations of non-invasive tactile blood flow sensors using CMOS MEMS technologies. The capacitive sensing structure consists of two parallel plate capacitors which can be connected in different configurations in cape with different measuring ranges. Sensor detection scope is set to be from 0 to 150 mmHg according to the estimated maximum human vessel pressure. The sensor is fabricated in commercial 2 polysilicon and 4 metal CMOS technology followed by self-developed post processes. The dimension of each sensor is 400μm in length with the membrane thickness of 1.45μm. An anisotropic inter metal dielectric layer etch step was utilized on CMOS chips to open wet metal etching holes. After this dielectric layer etch, a metal wet etching process was applied to release sensing structures. In order to protect the metal bonding pads in post CMOS MEMS processes, an Au layer was deposited on the pad areas. Based on the experiment results, the lateral metal sacrificial layer etching rate is 1.85μm per minute and the lateral etching rate underneath gold layer is 2.9μm per minute. We have successfully demonstrated the post CMOS MEMS processes for our sensors.

Initial finite element method analysis results showed that the sensitivities of two different designs are 6.7 and 2.2 fF per mmHg with a dynamic range of 75 and 200 mmHg. The sensor behavior measurement data will be presented.


The purpose of this study is to develop a sensitive gas sensor with engineered TiO2 nanostructures using semiconductor nanotechnologies for micro gas chromatography. Many TiO2 nanowires for gas sensing nowadays were fabricated by chemical synthesis methods, and the nanowires array are in irregular formats and the amount of sensing material may be varied chip to chip. The behavior of sensing repeatability of these TiO2 nanowires using conventional methods is hard to control. With the combination of E-beam lithography and the TiO2 thin film deposition, the TiO2 sensing nanowire arrays with well-controlled structures (100-300 nm wide with 1 μm period), were placed in between the Au interdigitated electrodes. A microheater were fabricated by deposition of the 350 nm thick Cr/Au films on the backside of the sensor. The great linear heating with increasing input power and uniform heating (329.3 °C in average, STDV of 9.3 °C, power of ~0.8 W) were obtained through an IR camera.

The performance of nanowire detector (100 nm wide, 183.5 MΩ) is compared to the microwire detector (20 μm wide, 24.6 MΩ) at various ethanol and benzene concentrations or at various operation temperatures. The measured resistance to the initial resistance ratio of the nanowire detector changed from 1 to 0.35 at 284 °C as 6.5% ethanol concentration. The effects of the rapid thermal annealing and an O2 plasma treatment to improve the sensor performances is investigated and will be presented.

11:20am MN-FrM10 Ultra-high Aspect Ratio High-speed Silicon Nanowire and Three-dimensional Formation Using a Hydrogen-assisted Deep Reactive Ion Etching. Z. Saeideh, S. Azimi, M. Pouidineh, S. Mohajerzadeh, A. Sandoughhsaz, University of Tehran, Iran

We report the formation of ultra-high aspect ratio and three dimensional features on silicon substrates using a novel low-density capacitive-coupled plasma reactive ion etching (13.56MHz). The etching process is based on using three gases of hydrogen/oxygen and SF6 in two sub-sequences called as passivation and etching sub-cycles. All three gases are used in the passivation step and SF6 in the etching step. Unlike Bosch process no polymer is used for passivation. By controlling the passivation sub-cycle, one is able to allow desired under-etching followed by “recovery” of the formerly under-etched features to make unique three-dimensional structures directly on silicon substrates [1].

Cleaned silicon samples are placed in an e-beam evaporation unit to deposit a 40nm chromium layer as the mask for the subsequent processing steps. The masking layer is patterned using precision projection lithography to achieve desired features between 100nm and 20um. For ultra-high aspect ratio and scallop-free etching while keeping the etch-rate of 1um/min, it is necessary to include trace values of H2/O2 during the etching step. Typical flows for H2/O2 and SF6 are 200/200 and 5 sccm in the passivation step while the etching is mainly practiced with SF6 (35 sccm). The plasma power is set at 250 W for the passivation and 130 W for the etching sub-cycle. By controlling these important parameters, we have realized three-dimensional features where the vertical structures have serpentine surfaces.
with desired recessions of 10um. Moreover, we have been able to realize arrays of nano-metric 3-D features using Si/SiO$_2$ structures with a diameter of 2-3um and features of the order of 100nm.

We realized 9-10um high and 90nm wide nano-wires where the mask undercut is 30nm and the surface of the wires is almost free of ‘scallop’. Scallop is side-effect of time-multiplexed processes where the periodic track of the etching step is seen on side-walls. To avoid this, while obtaining high-rates we have included H$_2$/O$_2$ gases during the etching sub-cycle. Normally H$_2$/O$_2$ gases act as the passivation layer, however the trace value of these gases does not affect the etching. Instead a slight passivation is formed on side-walls while the etching proceeds, prohibiting further lateral-etching of walls. We have studied the passivation layer using XPS and Ellipsometry. Thickness of the passivation layer is 2-3nm and it is mainly SiOF bonds (XPS). Field-emission SEM has been used to compare the results. Using this process we obtained high etch-rates of 0.8-1.1um/min for features around 100nm. The height of the nano-wires is around 10um with a side-wall angle of 70-80 and 90nm. This process uses low-density plasma with rapid steps and apart from MEMS/NEMS applications it can be used for “solar-cells” where nano-wires can significantly affect the efficiency and cost.


Plasma Science and Technology Division Room: 201 - Session PS-FrM

Plasma Modeling
Moderator: K. Bera, Applied Materials, Inc.

8:40am PS-FrM1 Delivering Activation Energy to Surfaces in Atmospheric Pressure Plasmas: Local and Remote, Z. Xiong, N.Y. Babaeva, M.J. Kushner, University of Michigan

Non-equilibrium atmospheric pressure plasmas (APPS) are efficient at producing reactive chemical environments where electron impact dissociation and ionization of feedstock gases. Other than generating UV photon fluxes, APPS are not thought to be sources of non-thermal activation energy in the form of energetic ions or hot atoms. Although mean free paths of ions may be less than 1 micron, the transient production of electric fields of 100s kV/cm to 1 MV/cm when the ionization fronts of streamers intersect with surfaces provide the possibility of accelerating ions to many to tens of eV. This high quality delivery of activation energy is a function of not only the properties of the streamer but also depends on the properties of the surface.

For example, delivery of high energy ions to the surface of a bulk polymer may differ from a layered polymer due to differences in their capacitive properties. These differences extend to organic materials as well – the deliver of energetic ions to cells and tissue will depend on their respective dielectric properties and those of the surrounding medium. Delivery of high quality activation energy in any form (photons or ions) to remote sites or locations is challenged by line-of-site issues and the charging of surrounding materials that may reduce ion energies. Being able to deliver activation energy to the crevices of rough surfaces may be important in the context of plasma sterilization. In this talk, results from modeling studies of atmospheric pressure plasma streamers and jets intersecting with dielectric surfaces will be discussed. The delivery of activation energy by ions and photons to rough surfaces will be discussed in the context of polymer modification and sterilization. Plasma sources will include direct applied dielectric barrier discharges and remote plasma source delivered by capillary tubes. * Work supported by the Department of Energy Office of Fusion Energy Sciences.

8:40am PS-FrM2 Kinetic Effects in Low Pressure Capacitatively Coupled Plasmas, A. Likhanski, P. Stolte, Tech-X Corp.

We present results of particle-in-cell/Monte Carlo collision simulations of kinetic effects in low pressure capacitively coupled plasma discharge.[1] In particular, we examine discharges of various gases (including Ar, Xe, and others) in the pressure range of 10s of mT and the frequency range of 10s of MHz. We track the formation of high energy electrons (e.g., at the ionization threshold or greater) as a marker for enhanced ionization, and look at the effects of elastic and inelastic collisions on the formation of these high energy electron bunches. [2,3] We show results for 2D and 3D simulations where we include density gradient effects, and results for plasma chemistry effects on the bulk electron energy distribution function and the ion energy distribution function at a plasma surface interface. We discuss the role of the bunches on electron heating in the plasma bulk and on their presence on how electron heating is treated in fluid simulations of plasma sources.


INVITED

The development of technologies for the plasma treatment of living tissue is in large part based on controlling plasma sources to deliver the desired fluxes of radicals and ions to surfaces. This process is complicated by scientific and technical issues. From a scientific standpoint, although it is generally accepted that reactive oxygen species (ROS) and reactive nitrogen species (RNS) are important in, for example, wound healing, sterilization and cancer treatment, it is not clear which species and in what proportions are optimum for each type of treatment. The situation becomes more complex when considering the UV photons, energetic ions and electric fields produced by the discharge which also interact with the tissue. From a technological viewpoint, the interaction between the tissue (and wounds in particular), the surrounding materials and the plasma can significantly affect the plasma. For example, the shape of the wound and the permittivity of the fluid in a wet wound can warp local electrical fields which then feed back to the plasma. Given this complexity and interdependencies, computer modeling of plasma-tissue interactions might provide insights to these interactions. In this talk, results from computer modeling of plasma-tissue interactions will be discussed. The modeling platform solves for charged particles, neutral and photon fluxes while also solving Poisson’s equation, and resolving spatial scales on reactor-to-cellular levels. Plasma transport through gases and liquids are included. Two types of plasma sources will be considered - dielectric barrier discharges (DBDs) where the plasma is in direct contact with the tissue and remote plasma jets, where dominantly neutral species and photons reach the tissue. We will discuss the treatment of wounds through a liquid layer covering exposed cells wherein the blood serum contains blood platelets. The characteristics of the plasma sources, and the interaction of plasma generated species and electric fields with the wound, fluid and underlying cells will be discussed.

* Work supported by the Department of Energy Office of Fusion Energy Sciences.


Low pressure magnetized capacitively coupled plasmas are extensively used for advanced microelectronics device fabrication. Due to the long mean free path of electrons in this regime, kinetic effects characterize the plasma dynamics in low pressure discharges. To take into account the kinetic effects, a hybrid 2-dimensional (2D) plasma modeling software has been developed that couples a particle-in-cell (PIC) model of charged particle species with a fluid method for neutral species. The electron motion due to electric and magnetic fields is incorporated in 3-dimensional velocity space using the Lorentz force law. The PIC model uses the Monte Carlo Collision (MCC) method to account for collision processes. The fluid model for neutral species takes into account species transport in the plasma, chemical reactions, and species depletion. Capacitively coupled plasmas in Ar have been computationally investigated for a 2D parallel plate plasma reactor in Cartesian co-ordinates. The inter-electrode gap is 5 cm (in y-direction). The bottom electrode is powered using a 60 MHz very high frequency (VHF) source, and the top electrode is grounded. The two electrodes are separated by quartz inserts. Ar plasma is simulated for a range of magnetic fields (25 - 100 Gauss), pressures (10 - 50 mTorr) and rf voltages (100 - 300 Volts). In this range of magnetic fields, the electrons are magnetized due to a small Larmor radius while the ions remain non-magnetized. For a symmetric reactor configuration without magnetic field, the plasma is symmetric, and the peak in plasma density occurs at the center plane between the top and bottom electrodes. The electron density increases with increase in pressure and rf voltage. With magnetic field in the x-direction (parallel to the electrodes), the plasma becomes more confined. When the magnetic field is increased, the plasma density increases, and the plasma becomes asymmetric. When the magnetic field direction is reversed, E x B drift reverses, therefore, the direction of plasma asymmetry reverses. The effect of magnetic field on plasma symmetry will be examined. In addition, results from the kinetic simulation will be compared to corresponding results from a fluid plasma model.

10:20am PS-FrM7 Simulations of SF$_6$ Plasma Etching in the GEC Reference Cell, S. Lopez-Lopez, Quanteml - University College London, UK, J.J. Mauco, D. Brown, Quanteml Ltd., UK, J. Tennyson, University College London, UK

Electrically driven plasmas containing halogens are very used in different material modification and surface cleaning processes. Sulfur Hexafluoride
(SF\textsubscript{6}) is used industry-wide in a range of processes for the dry etching of silicon or silicon dioxide for microelectronic feature definition, such as the Bosch process. However, the performance and efficiency of these processes and machines can vary widely, and the use of simulations can give us significant insight into the optimization problem and provide a low cost means for further development. That is especially relevant in the case of SF\textsubscript{6}, given its environmental impact, with a Greenhouse Warming Potential that is 22,000 times that of CO\textsubscript{2}. It is therefore vital to use SF\textsubscript{6} sparingly and efficiently in every process, and simulation here helps to find ways of remediating harmful waste gases and optimize the process for typical processing goals (e.g. etch rate, uniformity) as well as improving SF\textsubscript{6} consumption efficiency and other environmental measures.

A key aspect of the plasma processes here considered is that some type of work is done at the plasma / surface boundary layer, and realistic simulations must therefore incorporate the surface material and the etch product chemistry. This increases drastically the complexity of the problem but is the only way to represent all of the physics. Radical species from the surface entering the gas phase will take part in the phase and surface reactions that are associated with the parent gas, including negative ion formation and electron dissociation among others.

Here we present 2D simulations of an inductively driven SF\textsubscript{6} silicon etch process in the GEC Reference Cell [1], building upon previous calculations of SF\textsubscript{6} plasma chemistries using Quanternol-P [2]. Etch rate, pressure and power trends along with chamber wide contour plots of gas-phase species concentrations and potential contours are presented. Radical species from the surface are considered. We have found a good agreement with experimental results [3], which validates the underlying model and points to the important role of simulation-assisted plasma process development and optimization.

REFERENCES


10:40am PS-FrM8 Simulation of InP Etching under ICP Ar/Cl\textsubscript{2}/N\textsubscript{2} Plasma Discharge: Role of N\textsubscript{2} in the Sidewall Passivation. R. Chanson, A. Rhallabi, M.C. Fernandez, Ch. Cardinaud, J.P. Landsmeer, Institut des Matériaux Jean Rouxel (IMN), France. InP-based optoelectronic devices need reliable dry etching processes characterized by high etch rate, profile control and low damages. High density plasma etching, using inductively coupled plasma ICP reactors, has been found to be very important for the transfer of patterns from the mask to InP substrate and InP-based layers. In order to investigate the role of N\textsubscript{2} in the InP etching process under Cl\textsubscript{2}/Ar/N\textsubscript{2} plasma discharge, we have developed an ICP etching simulator permitting to determine the InP etch profile evolution through the mask as a function of the operating conditions and the initial mask geometry.

The InP etching simulator is divided in three modules: the global kinetic model of Cl\textsubscript{2}/Ar/N\textsubscript{2} ICP plasma discharge is based on 0D approach which allows to calculate the averaged densities of neutrals and ions as well as the electron density and electron temperature versus the machine parameters. The resolution of the differential equations associated to the mass balance of each considered species coupled to charge neutrality equation and the differential power balance equation from t=0 until the steady state allows to determine all reactive specie densities as well as their fluxes into the InP substrate. n\textsubscript{e} and T\textsubscript{e} calculated from the plasma global kinetic model are introduced in the sheath model to estimate the average sheath thickness. The Monte-Carlo technique is used to study the ion transport in the sheath. The calculation of energies and angles of positive ions impinging on the substrate allows determining the angular and energy distribution functions of positives ions. Such distribution functions with Cl, N and positives ions fluxes are introduced as input parameters into the etching model. The later is based on the cellular approach combined to the Monte-Carlo method which the considered domain (InP substrate and mask) is discretized on 2D uniform cells which each cell represents a real number of InP atoms. The fluxes of neutral species and positive ions are introduced as input parameters into the etching model. All the particle surface interaction processes like adsorption of atomic neutrals Cl and N on InCl\textsubscript{N} surface sites, desorption of InCl\textsubscript{N} sites, sputtering of both InCl\textsubscript{N} and mask by positive ions and redeposition of InCl\textsubscript{N} sites are described in probabilistic ways. Simulation results show the effect of the N\textsubscript{2} on the passivation of the lateral edges and in consequence the improvement of the etched profile anisotropy. However, a diminution of the etch rate by increasing the percentage of N\textsubscript{2} is observed. The simulated etch profiles are compared to those obtained by the experiments and the good agreements are obtained.

11:00am PS-FrM9 Three-Dimensional Modeling and Formation Mechanisms of Atomic-Scale Surface Roughness during Si Etching in Chlorine-Based Plasmas. H. Tsuda, Y. Takahashi, K. Erioguchi, K. Ono, Kyoto University, Japan.

Three-dimensional measurement and prediction of atomic-scale surface roughness on etched features become increasingly important for the fabrication of next-generation devices; however, the feature profiles are too small or too complex to measure the surface roughness on bottom surfaces and sidewalls of the etched features. To predict the surface roughness on atomic or nanometer-scale, we have developed our own three-dimensional atomic-scale cellular model (ASCeM-3D) [1] and feature profile simulation. Emphasis is placed on a better understanding of the formation mechanisms of atomic-scale surface roughness during Si etching in chlorine-based plasmas and the relationship between the ion incident energy and angle and etched feature profiles.

In the ASCeM-3D model, the simulation domain is divided into a number of small cubic cells of \( r_{\text{sc}} = 2.7 \times 10^{-5} \text{ cm} \), where the atomic density of Si substrates. Ions and neutrals are injected from the top of the simulation domain, and etch and/or sputter products are taken to be desorbed from etching surfaces into microstructural features, where two-body elastic collision processes between incident ions and substrate atoms are also taken into account to analyze the ion reflection on etched feature surfaces and penetration into substrates. The ASCeM-3D takes into account surface chemistries based on the Monte Carlo (MC) algorithm [2-4], including adsorption and reemission of neutrals, chemical etching, ion-enhanced etching, physical sputtering, and redeposition of etch and/or sputter products on feature surfaces.

Numerical results indicated that nanoscale convex features increase in size with increasing etching or plasma exposure time, and surface roughness increases with increasing incident ion energy. The ripple structures of etched surfaces were found to occur depending on incident angle of ions. Ion-enhanced etched surfaces show good results which affects the evolution of feature profiles and surface roughness on atomic scale.


The energy of ions bombarding the substrate is critical in plasma etching and deposition of thin films, especially when precise etching without damage is required. The ion energy distribution (IED) may be controlled by applying “tailored” bias voltages on the substrate, or on nearby electrodes immersed in the plasma. A Particle-in-Cell simulation with Monte Carlo Collision Model (P-ICMC) was used to investigate the application of DC voltage steps (and staircases) on an electrode, during the afterglow of a capacitively-coupled pulsed argon discharge, to control the energy of ions incident on the counter-electrode holding the wafer. Staircase voltage waveforms with selected amplitudes and durations resulted in ion energy distributions with distinct narrow peaks, having controlled peak energies and fraction of ions under each peak. A semi-analytical model was also employed to achieve “tailored” IEDs, i.e., distributions with a desired shape and energy spread (for example a nearly-nonmonoenergetic IED with given FWHM). This was again accomplished by applying judicious voltage waveforms on the substrate electrode. Predicted IEDs were compared with experimental data. Strategies to control the energy flux of bombarding ions or to distribute the total ion energy flux to different energies were identified.

Work supported by DoE: Plasma Science Center and NSF.

11:40am PS-FrM11 Molecular Dynamic Simulation for Selective Etching of Silicon Nitride over Silicon Oxide by Hydrofluorocarbon Ions. R. Shigekawa, M. Isohe, Osaka University, Japan, M. Fukiwasa, T. Tatsunami, Sony Corporation, Japan, S. Hamaguchi, Osaka University, Japan.

Selective etching of silicon oxide (SiO\textsubscript{2}) over silicon nitride (SiN) has been widely used in microelectronics fabrication processes such as contact hole etching in self-aligned processes, formation of a stress liner, and dual/triple hard mask (DHM/THM) etching processes of dual-damascene structures. Opposite selective etching of SiO\textsubscript{2} over SiN with high selectivity would be also desirable for various processes. In general, when a fluorocarbon gas is used for etching purposes, a carbon film tends to be accumulated on SiN surfaces which affects the evolution of feature profiles and surface roughness on atomic scale.

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etching rates and surface chemistry, especially focusing on effects of hydrogen on the process. The reactive interatomic potential functions for atomic systems of Si, O, F, C, N, and H were developed in-house for the MD simulations code, based on atomic interaction data of small molecules in ground states obtained from ab-initio calculations. Details of the atomic potential functions used in the simulations will be presented elsewhere. Simulations are typically performed on a small block of a model substrate that consists of several thousand atoms and is subject to bombardment by energetic particles such as CHxFy. In the simulations, we evaluate sputtering yields, surface modification during the process, and characteristics of sputtered products. From the simulations, it has been found that hydrogen of CHxFy ions tends to reduce F accumulation on SiN surface, forming volatile HF, and sometimes promotes formation of cyanides such as HCN. Detailed simulation results, including sputtering yields and surface chemical compositions, will be given in this presentation.

Late Breaking Session
Room: 109 - Session SS2-FrM

Surface Science Late Breaking Session
Moderator: C.R. Eddy Jr., U.S. Naval Research Laboratory, S.B. Sinnott, University of Florida

Surface Science Late Breaking Session

8:20am SS2-FrM1 CO Oxidation Facilitated by Robust Surface States on Au-Covered Topological Insulators, H. Chen, The University of Tennessee, Knoxville, W. Zhu, The University of Tennessee, Knoxville and ORNL, D. Xiao, Oak Ridge National Laboratory, Z. Zhang, Univ. of Sci. and Tech. of China and The University of Tennessee, Knoxville

Surface states—the electronic states emerging as a solid material terminates at a surface—are usually vulnerable to contaminations and defects. The robust topological surface state(s) (TSS) on the three-dimensional topological insulators (3DTI) provide a perfect platform for exploiting surface states in less stringent environments. Employing first-principles density functional theory calculations, we demonstrate that the TSS can play a vital role in facilitating surface reactions by serving as an effective electron bath. We use CO oxidation on gold-covered Bi2Se3 as a prototype example, and show that the robust TSS can significantly enhance the adsorption energy of both CO and O2 molecules, by promoting different directions of electron transfer. The concept of TSS as an electron bath may lead to new design principles beyond the conventional d-band theory of heterogeneous catalysis.

8:40am SS2-FrM2 A Theoretical Study of Methanol Synthesis from CO2 Hydrogenation on Metal-doped Cu(111) Surfaces, Y. Fang, State University of New York (SUNY) at Stony Brook, M.G. White, Stony Brook University and Brookhaven National Laboratory, P. Liu, Brookhaven National Laboratory

The synthesis of methanol (CH3OH) from CO2 hydrogenation (CO2 + 3H2 → CH3OH + H2O) has attracted considerable attention in the past decades. It is not only industrially important, but also of great environmental significance due to its application in the conversion of greenhouse gas, CO2. Commercially, the reaction is performed on a catalyst containing Cu, ZnO and Al2O3, on which the conversion of CO2 to CH3OH is kinetically limited to 15-25%. To improve the performance of the Cu catalysts, the effect of alloying on CH3OH synthesis was investigated in this study.

Density functional theory (DFT) calculations and Kinetic Monte Carlo (KMC) simulations were employed to investigate the CH3OH synthesis reaction from CO2 hydrogenation on metal-doped Cu(111) surfaces. Both the formate pathway and the reverse water gas shift (RWGS) reaction followed by CO hydrogenation pathway (RWGS + CO-Hydro) were considered. Our calculations showed that the overall CH3OH yield increased in the sequence: Au/Cu(111) < Cu(111) < Pd/Cu(111) < Rh/Cu(111) < Pt/Cu(111) < Ni/Cu(111). On Au/Cu(111) and Cu(111), the formate pathway dominates the CH3OH production. Doping Au does not help the CH3OH synthesis on Cu(111). Pd, Rh, Pt and Ni are able to promote the CH3OH production on Cu(111), where the conversion via the RWGS + CO-Hydro pathway is much faster than that via the formate pathway. Further kinetic analysis revealed that the CH3OH yield on Cu(111) was controlled by three factors: the dioxygenmethylene hydrogenation barrier, the CO hydrogenation step energy and the CO hydrogenation barrier. Accordingly, two possible descriptors are identified which can be used to describe the catalytic activity of Cu-based catalysts towards CH3OH synthesis. One is the activation barrier of dioxygenmethylene hydrogenation; the other is the CO binding energy. An ideal Cu-based catalyst for the CH3OH synthesis via CO2 hydrogenation should be able to hydrogenate dioxygenmethylene easily and bond CO moderately, being strong enough to prevent CO desorption, but weak enough to prevent CO poisoning. In this way, the CH3OH production via both the formate and the RWGS+CO-Hydro pathways can be facilitated.

9:00am SS2-FrM3 Synthesis and Characterization of Surface Oxide Films on CoGa(100), D.C. O’Connor, Stony Brook University, S. Axanda, W.-P. Zhou, Brookhaven National Laboratory, M.G. White, Stony Brook University and Brookhaven National Laboratory

Methanol is currently garnering a lot of attention as both a chemical feedstock for synthesis of organic substances and as a possible source of energy. It is currently produced industrially from a mix of syngas and hydrogen over a copper-zinc oxide catalyst. However, it has been shown that palladium-gallium oxide is a more active catalyst for methanol production, and uses a carbon dioxide-hydrogen feed, which is desirable for efficient CO conversion to CO2. Surface Science characterization of this catalyst poses a problem as a gallium oxide substrate is not conductive enough for use with X-ray photoelectron spectroscopy (XPS) or low energy electron diffraction (LEED). The most straightforward way to circumvent this limitation is to grow Ga2O3 on a conductive substrate. It has been shown that a Ga2O3 film forms on the surface of CoGa alloy crystals when exposed to oxygen. In this work we report the results of the characterization of Ga2O3 films on the CoGa(100) surface using XPS, LEED, and ion scattering spectroscopy (ISS). The films were synthesized using either O2 or NO2 as the oxidant at 300 K or in excess of 700K. ISS scans showed that cobalt was always present in the top surface layer regardless of oxidation conditions. XPS illustrated that depending on the oxidant and the temperature, the composition of the oxide films vary with some being nearly all Ga2O3 and ordered and others Co-Ga mixed oxides that gave no diffraction pattern.


9:20am SS2-FrM4 Apparent Ferroelectricity from Electrochemistry: Local Vacancy Diffusion in Oxides, A. Kumar, Y. Kim, A. Tslev, S.Y. Kalinin, A. Baddorff, Oak Ridge National Laboratory

Multiple reports of unexpected ferroelectricity in nanoscale systems have been made based on electrochemical hysteresis detected by piezoresistance force microscopy (PFM). These include thin films of ferroelectric materials below the critical thickness for ferroelectricity in the bulk, such as strontium titanate and manganites. However, recent studies of ionic systems, such as Li-ion conductors, suggest that local electrochemistry, usually ignored in these studies, may have an important role to play in the origin of the hysteresis loops. We report the role of local electrochemistry, including surface and bulk oxygen vacancy generation and dynamics on ionic conductors such as YSZ and lanthanum strontium cobaltes (LaSrCoO) and compare results with less understood systems including strained and unstrained SrTiO, and the lanthum aluminate - strontium titanate (LAO-STO) interface.

The role of oxygen vacancies in oxides, of obvious relevance in fuel cells, memristors, superconductors, etc., have traditionally been studied in macroscopic volumes and has been limited by high activation temperatures. We have developed an approach for spatially resolved local assessment of the oxygen dynamics and kinetic evolution in the gas phase oxidation of oxygen vacancies by utilizing volume variations of these oxides upon application of a concentrated electric field. In band excitation electrochemical strain microscopy (ESM), a strongly confined electric field at a tip is used to drive the oxygen vacancies in these oxide materials and the tip used to detect the resulting local electrochemical strain due to migration of oxygen vacancies. Vacancy diffusion is relatively slow, so that local strain hysteresis loops are open. Mapping the loop opening as a function of the final bias establishes the onset and kinetics of the diffusion process. Signal relaxation experiments locally characterize the diffusion dynamics of the vacancies. In mixed ionic-electronic oxide systems, current-voltage measurements provide complementary information on electronic transport.

Mapping of local oxygen vacancy diffusion with 30 nm resolution has been made based on a nanotip of oxides ranging from pure ionic conductors (YSZ) to mixed ionic electronic conductors (LaSrCoO), including ferroelectrics and resistive switching in TiO2 and SrTiO3 films. This approach was further applied to the LAO-STO interface, now a prototypical 2D electron gas with ferroelectric behavior proposed due to local hysteresis.
loops. While the role of static oxygen vacancies has been well discussed, we explore the local dynamic processes in the system. We observe two parallel dynamic processes that we tentatively ascribe to surface charge dynamics and bulk vacancy injection. This behavior is unique for LAO-STO and is not observed in YSZ and LaSrCoO$_3$. These observations both explain hysteresis without ferroelectricity and provide insight into the behavior of this system.

This research was conducted at the Center for Nanophase Materials Sciences, which is sponsored by Oak Ridge National Laboratory by the Office of Basic Energy Sciences, U.S. Department of Energy.

9:40am SS2-FrM5 Dopant-enhanced Neutralization of Low Energy Li$^+$ Scattered from Si(111). R.D. Guan, University of California, Riverside, Z. Sroubek, Czech Academy of Sciences, Czech Republic. J.A. Yarmoff, University of California, Riverside

The role of doping in semiconductor surface reactions is of fundamental scientific and technological importance, yet the effects of doping in atom-surface charge exchange have never been investigated directly. For example, in the dry processing of silicon, a dependence of oxidation, surface charge exchange have never been investigated directly. For scientific and technological importance, yet the effects of doping in atom-doping dependences. In addition to populating the bands, the band gap of Si excess majority carriers, which could have a large effect on electron tunneling rates between reactants and the surface, leading to the observed doping dependences. In addition to populating the bands, the band gap of Si narrows with increasing dopant density, which may also influence charge exchange.

This work presented here demonstrates that the charge exchange between scattered low-energy Li ions and a passivated Si surface depends strongly on doping. The neutralization of 3 keV Li$^+$ ions scattered from Si(111) is measured as a function of doping density, dopant type, and hydrogen coverage using time-of-flight spectroscopy. When the surfaces are saturated with hydrogen to unpin the Fermi level, the neutral fractions decrease for lightly doped samples, but become anomalously large for high-doped n-type Si. The neutralization does not correlate with the surface work function, indicating that the models used for metal surfaces are not directly applicable here.

A model is presented that includes the many-body band-gap narrowing effect, which predicts the neutralization to good accuracy using a tunneling mechanism similar to the free-electron gas jellium model normally employed for ion/metal interactions, but excluding levels in the gap. This work demonstrates that the surface of Si behaves, as far as electron transfer is concerned, like a jellium electron gas with states missing in the band gap region.

10:00am SS2-FrM6 Charge Transfer and Diabatic Dissipation of Surface Released Chemical Energy in GaP/Pt Schottky Nanostructures with a Resistively Heated Nanocathode Layer. E.G. Karpov, S.K. Dasari, A. Hashemin, University of Illinois

Recent observations of chemically induced hot electron flow over Schottky barriers in nanoscale nanostructures provides interesting possibilities for electrolyte-free conversion of chemical energy into electricity and novel sensor applications. The work aims to make advantage of the nonthermal nature of the diabatic dicromatics and explores the low-temperature (340-430K) chemovoltaic effect of hydrogen to water oxidation on Pt/GaP nanostructure surface. The chemoincurrent is also studied at high pressures of the oxygen/hydrogen mixture diluted with nitrogen, all the way to atmospheric pressures. Separation of the diabatic component to the total generated current is performed with an accurate method based on resistive nanofilm heating.

10:20am SS2-FrM7 Utilizations of Low Platinum Loading Pt-Co Bimetallic Alloy Catalyst for Proton Exchange Membrane Fuel Cells (PEM) in UHV Conditions. A.S. Ahsen, Gebze Institute of Technology, Turkey, O.K. Ozdemin, Yildiz Tekn. Univ., Turkey, O. Oturuk, Gebze Institute of Technology, Turkey

In this ongoing study it has been tailored surface structures and morphologies to increase chemical and structural stability for higher efficiency and improved utilization compared to presently available Pt-based catalysts used for fuel cell application. Pt-Co bi-metallic alloys is selected to explore the electronic and structural properties of tailored surface and interfaces of electrocatalysts, and it is intended to exploit surface chemical dynamics and interaction in fuel cell where chemical reaction pathways are influenced by the nature of the underlying support, the surfaces can be designed and modified appropriately to either promote or inhibit particular reaction. A series of Pt and Pt-Co bi-metallic nanocatalysts alloys films were deposited on glassy-carbon disks and on a commercial hydrophobic carbon paper substrate by dc magnetron sputtering. The electronic structure and chemical states of surface were investigated by Photoelectron Spectroscopy (XPS and UPS). Cyclic Voltammetry (CV) and Rotating Disc Electrode (RDE) methods were used to investigate Active Surface Area (ASA) and Oxygen Reduction Reaction (ORR) kinetics. Low loading values, 6-22 µg Pt/cm$^2$, were observed the kinetic properties of the nanocatalysts. The highest active surface area was observed with an optimum loading of 10 µg Pt/cm$^2$ value. Besides that the Pt-Co alloy catalysts showed significant improvement on catalytic activity against pure platinum catalysts. The effects of deposition temperature on alloy formation and reaction kinetics were also observed. A photoelectron spectroscopy study was also used for understanding electronic interaction between Pt and Co by the function of preparation temperature.

CV measurements indicate that carbon supported Pt-Co catalyst on gas diffusion electrode had electrochemical stability in the acidic environment. Also, it was observed that with increasing Co ratio metal oxidation and reduction current peaks rises. The enhancement on the ORR activity for the PtCo catalysts to form a Pt-enriched layer of the Pt-shell layer as revealed from the CV analysis, where the PtCo/C electrodes show a delayed formation of Pt-OH and faster reduction of the Pt-oxgenated containing species compared to Pt/C. It means that the oxophilicity of the PtCo/C catalysts are low compared to Pt/C. The hydrogen desorption peaks on the CV curve showed that the highest ESA value was obtained from Pt-Co(3:1) ratio. The capacitance region of PtCo catalysts increases with rising Co ratio. This is a strong evidence for that additional Co layers change the catalyst morphology. As a result, gas can reach to the reaction area easily and this reduces the ohmic resistance.

Surface Science Division
Room: 107 - Session SS-FrM

Surfacing Science on Graphene
Moderator: I.I. Oleynik, University of South Florida

8:20am SS-FrM1 Towards Controlled Growth of a Single-Layer of MoS$_2$. D. Sun, W. Lu, D. Kim, J. Mann, L. Bartels, University of California, Riverside

MoS$_2$ is a semiconducting material consisting of sulfur-molybdenum-sulfur tripledecker layers loose bound by van der Waals interactions. MoS$_2$ has been used technologically for a long time, for instance as lubricant, where similar to graphite its layered character was employed. Recently, its electronic characteristics have attained increased attention with the finding that it transitions from an indirect bandgap semiconductor at 1.6eV gap to a direct bandgap one at 1.9eV gap at the transition from multilayers to a single layer. A transistor has been constructed from a MoS$_2$ and shown appreciable properties. The increased bandgap and high fluorescence yield may also suggest applications of the material for photonic or photocatalytic applications.

MoS$_2$ can be exfoliated mechanically similar to graphene. While this method is simple, it is hard to control and not amendable to mass production of thin films. Solution-based processes have been proposed and may provide a scalable source of a mixture of single and multilayer material. Here we show an alternative avenue for the fabrication of MoS$_2$ monolayers: growth of MoS$_2$ on a sulfur-preloaded copper surface. In contrast to all other methods, this route has the potential of providing exclusively monolayer material, as the sulfur source is only available until the substrate is covered. Practically, this approach is related to the growth of graphene monolayers on copper or ruthenium films, where segregation of carbon to the surface is employed in aggregating a carbonaceous layer that transforms into graphene under the correct conditions.

Small MoS$_2$ triangles of a few nanometers in size have been grown previously on gold in a dilute H$_2$S atmosphere. Here we show significantly larger patches, tens of nanometers in size. In contrast to gold, copper forms a multilayer metal-sulfur surface coverages and also readily absorbs sulfur into the bulk. Thus, we can preload the substrate with a specific amount sulfur using an easy to handle liquid precursor, benzenethiol. In previous work we have shown that heating to below 400K removes the phenyl group of benzenethiol reliably from copper leaving sulfur coverages behind.

8:40am SS-FrM2 Oxygen Adsorption on Electronically Modified Graphite Surfaces Studied by Molecular Beam Scattering. J.P. Oh, T. Kondo, K. Arakawa, Y. Saito, J. Nakamura, University of Tsukuba, Japan

The graphite surface consists of π conjugated system. When the π conjugated system is broken, the non-bonding π electronic states are known to form on the surface. Recently, the non-bonding π electronic states at the Fermi level of the graphite-related materials are expected to the active sites
for the specific chemical reaction such as oxygen-reduction reaction in the fuel cell [1]. It is thus important to understand the interaction between an oxygen molecule and the graphite surface for the efficient usage of the graphite-related materials. We have reported previously that the defects induced by Ar\textsuperscript{+} ion bombardment on the graphite surface significantly affects the gas-graphite interaction based on the measurements of the angular intensity distributions of He and Ar beam scattered from the pristine and the defect induced graphite surfaces [2]. The difference in the gas- surface interaction has been ascribed to the local breaking of the conjugated system of graphite by defect formation. To further investigate the effect of the modification of the graphite electronic states on the gas- surface interaction, especially for the oxygen adsorption, we have measured angular intensity distributions of O\textsubscript{2} from electronically modified graphite surfaces, namely potassium intercalated graphite, nitrogen-doped graphite (graphite bombarded by N\textsubscript{2} \textsuperscript{+} ion) and defective graphite (graphite bombarded by Ar\textsuperscript{+} ion). The detail of scattering features as well as the effects of the electronic modification of graphite on the oxygen adsorption will be discussed in detail with our recent STM and STS results.

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### 10:00am SS-FrM6 Ripening Behavior of Pt Clusters on Monolayer Graphene Supported by Ru(0001) and the System’s Thermal Stability

**C.U. Lorenz, A.K. Engstfeld, Ulm University, Germany, H.E. Hoster, Technische Universität München, Germany, R.J. Behm, Ulm University, Germany**

The Moiré-type nm-scale patterns of graphene monolayers supported by metal single crystals were recently used for the fabrication of ordered arrays of metal nanoclusters by metal vapor deposition under ultrahigh vacuum UHV conditions [1-3]. The corrugation within the adsorption potential of the graphene layer result in virtually monodisperse clusters. These are important for model (electro-)catalysis studies investigating the size dependency of Pt clusters on carbon support.

In this study we analyze the ripening behavior of Pt clusters (formed at room temperature) on monolayer graphene supported by Ru(0001) at temperatures above 450 K. The STM images demonstrate that change of the clusters gives insight into the rate determining step and into the mechanism of the ripening process. Two different annealing step methods (i: a single sample was successively heated to higher temperatures and ii: individual samples reproducibly prepared in the same manner annealed to certain temperatures; both methods using the same heating period of 10 min) bring us to the conclusion that Pt cluster ripening between 450 K and 725 K occurs via a mechanism proposed by M. Smoluchowski. Above 725 K we observe indications for a change in the ripening mechanism, where Smoluchowski ripening is likely in competition with Ostwald ripening. Also above 725 K, we observed an adverse influence of the Pt on the stability of the monolayer graphene. Single defects in the otherwise well ordered graphene appeared, which were absent after annealing to lower temperatures.


### 10:20am SS-FrM7 Mechanisms of Graphene Growth on Metals.

**N.C. Bartelt, Sandia National Laboratories**

INVITED

Growth on metal substrates is a promising route for synthesizing high-quality graphene films. In addition, moving electronic applications into the real world requires understanding and controlling the properties of graphene in contact with metals. This talk will focus on the properties and growth mechanisms on several metals distinguished by varying binding strengths to graphene’s electronic properties (i.e., band structure and work function) result from changes in orientation. On Cu foils, graphene islands nucleate with a large range of orientation. Thus, the weak film-Cu interaction leads to a high defect density. Finally, the mechanism of bilayer graphene growth has been explored. Diffraction analysis reveals that the second graphene layer in Ir(111) grows next to the substrate, not on top of the first layer. This “underlayer” growth mechanism occurs when the carbon source is either segregation from the substrate or deposition on top of the first layer. How this unusual mechanism affects thickness uniformity will be discussed.

This work was supported by the Office of Basic Energy Sciences, Division of Materials and Engineering Sciences of the U.S. Department of Energy under Contract No. DE-AC04-94AL85000.
sizes and the hydrogen coverage and the opening of a band-gap in the graghene/H/Ru(0001) system for some of the Moiré/hydrogen adlayer coverages. This work was supported by the Nanoscale Science and Engineering Center for High-rate Nanomanufacturing (NSF NSEC-425826) and NSF DMR-1006863.

11:20am SS-FrM10 Modifying Ni(111)/Graphene Interfaces by Sn-Ni Interface Alloy Formation, R.Q. Addou, A. Dahal, L. Adamska, I.I. Oleynik, M. Batzill, University of South Florida

Graphene growth on metal surfaces (Ni, Pt, Ir, Rh and Cu) has been studied extensively [1]. Ni(111) is special among these metals because it is closely lattice matched with graphene (\(a_{\text{Ni(111)}} = 0.246 \text{ nm} \approx a_{\text{graphene}} = 0.249 \text{ nm}\) allowing the growth of graphene with a single domain and in registry with the substrate [2]. However, compared to most other metal substrates the interaction between Ni and graphene is rather large, resulting in a small metal-carbon distance and a large shift of the graphene \(n\)-band compared to freestanding graphene. In order to de-couple graphene from the Ni-substrate other weaker interacting metals such as Cu and Au have been successfully intercalated between the graphene and Ni-substrate [3]. These metals have, however, a different lattice parameter and consequently the registry between the substrate and graphene is lost. Here we demonstrate a new approach that weakens the metal-graphene interaction without destroying the lattice registry. By intercalating Sn-atoms an ordered \(\sqrt{3} \times \sqrt{3}\) R30° Sn-Ni alloy is formed. The intercalation process is characterized by Auger electron spectroscopy (AES) and low energy electron diffraction (LEED). In this alloy Sn substitutes for surface Ni atoms without changing the lattice parameter of the substrate and consequently the registry between the metal substrate and graphene is maintained. DFT simulations indicate that Sn alloying with Ni weakens the interaction of graphene with the metal substrate and consequently increasing the graphene-substrate distance and restoring the graphene \(n\)-band close to the position of free-standing graphene. Atomic-resolution scanning tunneling microscopy (STM) imaging reveals that the alloy periodicity is reproduced in the graphene layer, i.e. a \(\sqrt{3} \times \sqrt{3}\) R30° superstructure is imposed on the graphene by the alloy substrate. This indicates a variation of the local density of states for C-atoms located on top of Sn-substrate sites compared to Ni-sites. Further experimental and theoretical characterization of the influence of the substrate on the electronic and structural properties of graphene is ongoing.


11:40am SS-FrM11 Silicene Epitaxial Sheets: Silicon New Start, P. Vogt, Aix Marseille Univ, CNRS-CNAM, France and Technische Universität Berlin, Germany, P. De Padova, C. Quaresima, CNR-ISM, Italy. J. Avila, E. Frantzeskakis, M.C. Asensio, Synchrotron SOLEIL, France. B. Edel, Aix Marseille Univ, CNRS-CNAM, France. G. Le Lay, Aix Marseille Univ, CNRS-CNAM, France and CNR-ISM, Italy

We have just synthesized in Marseille silicene sheets [1], i.e., atom-thin graphene-like silicon layers with an in-plane Si-Si interatomic distance of \(a_{\text{graphene}} = 0.23 \text{ nm}\), upon in-situ epitaxial growth on silver (111) surfaces. The honeycomb atomic structure is revealed in Scanning Tunnelling Microscopy, while the long-range epitaxial order is confirmed by sharp 4x4 Low Energy Electron Diffraction patterns. Dirac cones at the K and K’ points of the silicene Brillouin zone, evidenced in High-Resolution Synchrotron Radiation Angle-Resolved PhotoElectron Spectroscopy measurements, point to massless relativistic fermions with a Fermi velocity of \(1.3 \times 10^6 \text{ m/s}\), as theoretically predicted [3], quite the same as graphene, and four times higher than previously obtained on a one-dimensional grating of silicene nano-ribbons [4]. Density Functional Theory calculations including the Ag(111) substrate confirm the stability of the epitaxial arrangement. The demonstration that silicon can form sheets of graphene with a single domain and in registry with the substrate and graphene is lost. Here we demonstrate a new approach that allows the growth of graphene with a single domain and in registry with the substrate [2]. However, compared to most other metal substrates the interaction between Ni and graphene is rather large, resulting in a small metal-carbon distance and a large shift of the graphene \(n\)-band compared to freestanding graphene. In order to de-couple graphene from the Ni-substrate other weaker interacting metals such as Cu and Au have been successfully intercalated between the graphene and Ni-substrate [3]. These metals have, however, a different lattice parameter and consequently the registry between the substrate and graphene is lost. Here we demonstrate a new approach that weakens the metal-graphene interaction without destroying the lattice registry. By intercalating Sn-atoms an ordered \(\sqrt{3} \times \sqrt{3}\) R30° Sn-Ni alloy is formed. The intercalation process is characterized by Auger electron spectroscopy (AES) and low energy electron diffraction (LEED). In this alloy Sn substitutes for surface Ni atoms without changing the lattice parameter of the substrate and consequently the registry between the metal substrate and graphene is maintained. DFT simulations indicate that Sn alloying with Ni weakens the interaction of graphene with the metal substrate and consequently increasing the graphene-substrate distance and restoring the graphene \(n\)-band close to the position of free-standing graphene. Atomic-resolution scanning tunneling microscopy (STM) imaging reveals that the alloy periodicity is reproduced in the graphene layer, i.e. a \(\sqrt{3} \times \sqrt{3}\) R30° superstructure is imposed on the graphene by the alloy substrate. This indicates a variation of the local density of states for C-atoms located on top of Sn-substrate sites compared to Ni-sites. Further experimental and theoretical characterization of the influence of the substrate on the electronic and structural properties of graphene is ongoing.


8:20am TF-FrM1 Low-temperature CVD with Growth Inhibitors to Afford Uniform, nm-thick Films in Structures with Aspect Ratio ~ 200, S. Babar, J. Abelos, University of Illinois at Urbana Champaign

In order to grow uniform, smooth and pinhole-free films of nm thickness, the nucleation step must occur with high areal density on the substrate and the nuclei must have a narrow size distribution. If the film must be deposited inside of a deep structure of very high aspect ratio, then the growth process must be strongly controlled by the nucleation rate of the precursor species rather than by the rate at which precursor is transported to the growth surface. Here, we present a method of control in chemical vapor deposition that simultaneously meets both of the above criteria. We previously reported that the steady-state film growth rate is reduced when a molecular species, called the inhibitor, is added to the CVD process. The inhibitor interacts transiently with the growth surface but does not decompose and incorporate its elements into the film, i.e., the inhibitor is not consumed. We also reported that the inhibitor greatly improves the film nucleation step, e.g. for the growth of HfB2, a high performance diffusion barrier, on SiO2 substrates. In the absence of the inhibitor the nucleation density is low and the size distribution is broad, such that some islands attain a height of > 10 nm before the film even coalesces. In the presence of the inhibitor, nuclei attain a height of ~ 1 nm, but then grow only very slowly. Additional nuclei continue to form and fill in the bare substrate, such that coalescence occurs at a thickness of ~ 2 nm with a rms surface roughness of < 0.5 nm.

The use of a growth inhibitor allows the nucleation regime of film growth to afford excellent nm-thick coatings in extremely high aspect ratio features. Note that when the film growth rate is low, so also is the consumption of precursor, such that a partial pressure of precursor species persists to the bottom of the feature. The inhibitor further assures that the nucleation process is greatly reduced. In features of aspect ratio >> 200, we show that a film is deposited on all surfaces with a rms roughness of 0.6 nm. We also discuss the mechanisms that can afford the observed results. We suggest that the ability to reduce (homogenize) the size distribution and increase the areal density of nuclei will greatly extend the useful range of CVD precursor-substrate combinations which can afford nm-thick coatings in very high aspect ratio features.

8:40am TF-FrM2 Understanding the Role of Hydrogen Impurity Scattering in Manganese Thin Films on the Si(001) Surface, A.J. Stollenhoff, B.J. Friend, University of Northern Iowa

Interest in energy production via hydrogen has been increasing due to the fact that it does not emit harmful greenhouse gases associated with fossil fuels. Unfortunately, hydrogen is an explosive gas that is both colorless and odorless. This will make hydrogen sensors an important piece of safety equipment in green energy-based systems. Diffusion of hydrogen to the Mn/Si interface is believed to have an amphoteric effect on the interface characteristics making it a possible hydrogen sensor. This motivates a need to better understand electron transport properties through Mnn/Si Schottky diodes. We performed ballistic electron emission microscopy (BEEM) on Mn/Si(001) Schottky diodes to study hot electron transport properties through this system. Samples were fabricated using electron-beam deposition of Mn onto n-type Si(001) with thickness ranging from 10 to 40 Å. Scattering in the Mn films as function of energy has been measured and compared to results obtained on Au/Si(001) Schottky diodes. Schottky heights on these samples were determined from the corresponding BEEM spectra using the Bell-Kaiser model. Future plans include introducing an atmosphere consisting of varying amounts of hydrogen and measuring the resulting effects on hot electron transport in the Mn film as well as the Schottky height. Should electron scattering or the Schottky height be affected by the presence of hydrogen, this system may prove useful as an inexpensive hydrogen sensor.

9:00am TF-FrM3 Elimination of Indium Surface Segregation in InGaN Grown Throughout the Miscibility Gap, M.W. Moseley, B. Gunning, J.E. Lodver, Georgia Institute of Technology, G. Namkoong, Old Dominion University, W.A. Doolittle, University of Alabama in Huntsville, Georgia Institute of Technology

InGaN alloys have great potential in optoelectronics due to the tunable bandgap which spans the visible spectrum. However, these alloys are difficult to obtain as a result of thermal decomposition, indium surface segregation, and spinodal decomposition. To solve these problems, low growth temperatures, fast growth rates, and in situ surface analysis must be
used. These limitations make Metal-Modulated Epitaxy (MME) a promising growth candidate. Using MME, smooth InGaN has been grown without phase separation throughout the miscibility gap. MME is a growth technique applied to molecular beam epitaxy of III-nitrides in which metal and dopant cell shutters are periodically opened and closed while active nitrogen flux remains constant. This technique uses metal-rich fluxes that would accumulate droplets in traditional MBE, taking advantage of the enhanced adatom mobility provided by excess metal. The periodic shuttering of the effusion cells allows the excess metal to be consumed, providing smooth, dry surfaces required for devices that demand abrupt interfaces.

In this study, MME is applied to the growth of InGaN and transient RHEED intensities are monitored for differing metal shutter open times (Fig. 1). It is found that these RHEED transients are the result of a RHEED oscillation associated with the buildup and consumption of the metal adlayer. This allows for observation and control of fractions of adsorbed metal layers. However, there is a drastic difference between the RHEED transients of low and high metal shutter open times. This difference is attributed to a growth regime which is characterized by a high metal concentration and by the generation of highly ionized deposition fluxes stemming from high electron (plasma) densities. Cathodic arc and pulsed laser deposition are examples of plasma deposition processes where such discharges are used to achieve high deposition rates and to facilitate the growth of metastable phases, nanostructures as well as micro-electro-mechanical systems and ultra-high-density magnetic storage. These materials have also been shown to possess a unique strain-induced chessboard cutoowell structure between their hard L10 and soft L12 magnetic phases that features exchange coupling effects. Within this class of materials Fe-Pd alloys possess a low order-disorder transition temperature making them a good candidate for ordered structure studies.

Nanoparticles and thin films of eutectoid (61.5 at% Pd) Fe-Pd were deposited using pulsed laser deposition. Additional ternary nanoparticles were deposited utilizing a novel technique involving the matrix assisted decomposition of metal-organic precursors. Nanoparticles and films were subsequently characterized by x-ray diffraction, vibrating sample magnetometry, high resolution transmission electron microscopy, and scanning electron microscopy.

9:20am TF-FrM4 Aging Effect on the Hydrophilicity of Metal Nanorod Arrays, N. Albarakati, D. Ye, Virginia Commonwealth University

Time-dependent wettability of nickel nanorod arrays were studied by measuring the water contact angles on the samples. Nickel nanorod arrays were deposited on silicon substrates by sputtering oblique angle deposition with an 85° incident angle. The substrates are rotated at a speed of 0.5 rps by a stepper motor. Samples with varying nanorod heights were prepared. Water contact angles were measured for each sample on a sequence of time up to three months. The water contact angles on the fresh nanorod arrays are less than 10°. However, the contact angle increases with time after the samples were exposed to air. We observed that the contact angles on the samples with short nanorods increase quickly to an angle close to hydrophobic and that on the samples with long nanorods increase slowly and remain in the hydrophilic region. X-ray photoelectron spectroscopy was employed to study the change of the surface composition due to oxidation and hydrocarbon contamination. We believe that the change of water contact angles is due to the accumulation of hydrocarbon on the nanorod surface.

9:40am TF-FrM5 Cubic ZnMgO and NiMgO for UV-C Applications, R.C. Boutwell, J.W. Mares, M. Wei, W.V. Schoenfeld, University of Central Florida

We will report on two new ternary cubic oxides offering close lattice matchings to MgO, ZnMgO and NiMgO, and compare the properties of films grown by both RF Plasma-Assisted MBE and a much more economical sol-gel deposition process. Using both methods we have successfully demonstrated band gap tuning in the UV-C spectral region, with ZnMgO films producing band gaps ranging from ~5 eV to 7.8 eV and NiMgO films having band gaps ranging from 3.5 eV to 7.8 eV. XRD revealed 20 FWHM values as narrow as 0.19°, and atomic force microscopy of ZnMgO films demonstrated surface roughness of 3.4 nm with NiMgO films having roughness below 1 Å. Optical, compositional, and morphological results from films produced using both synthesis methods will be presented. We will also report on initial photodetectors with 5 μm interdigitated fingers that produced peak responsivities of 12 mA/W at 250 nm.

10:00am TF-FrM6 Growth of Fe-Pd Nanoparticles and Thin Films via Pulsed Laser Deposition and the Matrix Assisted Pulsed Laser Evaporation of Metal Based Acetates, M.A. Steiner, J.M. Fitz-Gerald, University of Virginia

Nanoparticle and thin film magnetic alloys of 3d-4d/5d metals such as Fe-Pt, Co-Pt, and Fe-Pd are of technological interest due to their ordered L10 tetragonal phase which exhibits high magnetocrystalline anisotropy. Hard magnetic properties combined with ductility and corrosion resistance make these alloys ideal for applications including micro-electro-mechanical systems and ultra-high-density magnetic storage. These materials have also been shown to possess a unique strain-induced chessboard cutoowell structure between their hard L10 and soft L12 magnetic phases that features exchange coupling effects. Within this class of materials Fe-Pd alloys possess a low order-disorder transition temperature making them a good candidate for ordered structure studies.

Nanoparticles and thin films of eutectoid (61.5 at% Pd) Fe-Pd were deposited using pulsed laser deposition. Additional ternary nanoparticles were deposited utilizing a novel technique involving the matrix assisted decomposition of metal-organic precursors. Nanoparticles and films were subsequently characterized by x-ray diffraction, vibrating sample magnetometry, high resolution transmission electron microscopy, and scanning electron microscopy.

10:40am TF-FrM8 Surface Plasmon Resonance Excited in RuO2 Films Grown on Glass and on Crystalline (901) TiO2, L. Wang, E. Yang, C. Clavero, E. Crisman, V. Dinavahi, I. Novikova, R.A. Lukaszew, College of William and Mary

***PLEASE NOTE YOU MUST IDENTIFY A DIFFERENT PRESENTER FOR THIS ABSTRACT. YOU MAY PRESENT ONE PAPER ONLY (ORAL OR POSTER) AT THE CONFERENCE. YOU ARE ALREADY LISTED AS PRESENTER OF ABSTRACT #554: Surface Plasmon Resonance (SPR) excitation in metal oxide films and nanostructures has been widely applied to study biomolecules, absorption, bio-imaging, bio-sensing, and sensitivity enhanced Raman spectroscopy [1,2,3]. However, although the theoretical principles underlying SPR excitation are applicable to any conductive material, only metals, and in particular Au and Ag, have been considered for practical applications. Here we investigate SPR excitation in a conducting metal oxide, such as RuO2. Due to the large optical nonlinearities, the frequency doubling of SPR excitation is more intense in the infrared region for this material. The RuO2 thin films investigated were grown using reactive magnetron sputtering on two different substrates resulting in amorphous RuO2 films when deposited on glass and crystalline RuO2 films when epitaxially deposited on TiO2 (901) substrates. We have used Atomic Force Microscopy (AFM) and Reflection High-Energy Electron Diffraction (RHEED) to characterize the surface morphology and microstructure of these samples. Four-point probe was used to investigate the electrical conductivity properties and ellipsometry was used to characterize the optical properties of the films. We will show a comparison of the physical properties including the SPR excitation between these two kinds of thin film RuO2 samples. The investigation of SPR in conducting metal oxide materials provides a significant advancement for thin film characterizations as well as opens new venues for photonic and plasmonic applications. This work was financially supported by NSF (DMR-1006013).


11:00am TF-FrM9 Towards Achieving a High Degree of Carbon Ionization in Magnetron Sputtering Discharges, A. Alizaj, K. Sarakinos, D. Lundin, U. Helmersson, Linköping University, Sweden

Physical vapour deposition (PVD) methods, which are characterized by highly ionized deposition fluxes of the film forming species, provide added means for the fabrication of tailored materials. They can, for instance, facilitate the growth of meta-stable phases, nanostructures as well as selective deposition on complex-shaped substrates. In such methods, the generation of highly ionized deposition fluxes stems from high electron (plasma) densities. Cathodic arc and pulsed laser deposition are examples of such discharges where electron densities in the order of 1014 m−3 can be obtained. These techniques, while providing as high as 100% degree of ionization of the deposition flux, exhibit several drawbacks, such as macroparticle ejection from the target, lack of lateral film uniformity, and in some cases are difficult to scale up. Magnetron sputtering based techniques
are technologically more relevant, owing to their inherent advantages of conceptual simplicity, upscalability, and film uniformity. However, electron densities in magnetron discharges are significantly smaller, in the range of \(10^4-10^6\) m\(^{-3}\) and therefore generation of a highly ionized deposition flux is often difficult. This difficulty is overcome by high power impulse magnetron sputtering (HiPIMS), where plasma densities on the order of \(10^9\) m\(^{-3}\) are achieved. HiPIMS has been successful in enhancing the ionization for most common metals (Cu, Al, Ta, Ti), but it is challenged when nonmetals such as carbon is considered. Previous investigations have shown that C+/C ratio in HiPIMS does not exceed 5%, which does not provide efficient control over the physical properties and synthesis of carbon in various technologically relevant forms, e.g. tetrahedral amorphous carbon.

In the present study we address the low degree of ionization of carbon in magnetron discharges. We have developed a new HiPIMS based process, which provides a plasma characterized by high electron temperature and plasma density as determined by time-resolved Langmuir probe measurements. The C\(^+\) ion energy distribution functions (IEDFs) determined by time-averaged energy resolved mass spectrometry demonstrate an energetic C\(^+\) ion population and an overall five-fold increase of the C\(^+\) ion fraction as compared to standard HiPIMS methods. The enhanced ionized fraction of carbon facilitates the growth of carbon films with mass densities as high as approx. 2.8 g/cm\(^3\) as determined by high resolution x-ray reflectively measurements. Determination of the D-peak to G-peak ratio \((I(D)/I(G))\) and full width at half maximum of the G-peak in Raman spectra indicate that the films contain a large fraction of diamond-like bonded (sp\(^3\)) carbon.

11:20am TF-FrM10 Kinetics of Sputtered Metal Film Growth on Vertically Aligned Carbon Nanotube Arrays, C. Muratore, A. Reed, A. Waite, J. Bulman, J. Hu, T. Smith, A.A. Voevodin, Air Force Research Laboratory

Controlling the surface morphology of metallized arrays of vertically aligned nanotubes (VACNTs) is useful for diverse technological applications, such as interface materials for thermal management of high-heat flux electronics and electrical contacts for MEMS switches. We have observed variation of the morphology of metal films sputtered on carbon nanotubes, ranging from thin continuous films surrounding individual bundles of carbon nanotubes along their entire length for tubes up to 100 microns in length, to metal canopies covering the surface, with only minimal penetration (10-20 microns) into the nanotube forest. There is a strong link between flux of metal ions and their kinetic energy and the morphology of metal films of technological interest, including titanium, nickel, copper, gold and aluminum. To explore the mechanisms dictating metal film architectures on VACNT arrays, we systematically altered the flux and energy of incident metal and inert gas ions. These processing conditions were characterized using a mass spectrometer and energy analyzer adjacent to VACNT substrates. For conditions of interest, film growth was carried out for different times in an ultra high vacuum processing chamber, followed by electron microscopy of VACNT film cross sections, which were examined to observe different stages of growth and identify mechanisms of film nucleation and growth on nanotube sidewalls and tips. Comparison of growth kinetics of metal films on planar (002) graphite surfaces to that observed on VACNTs was used to identify effects of curvature on film growth. Switch performance was shown to be strongly dependent on morphology.

11:40am TF-FrM11 Crystallographic Orientation of Vanadium Dioxide Nano-Grains on Various Single-Crystal Sapphire Substrates, F. Rivera, Brigham Young University, J. Nag, R.F. Haglund Jr., Vanderbilt University, R. Davis, R. Vanfleeter, Brigham Young University

Vanadium dioxide (VO\(_2\)) is a material of particular interest due to the reversible semiconductor to metal phase transition that VO\(_2\) exhibits near room temperature (~ 68 °C) and the accompanied hysteresis. Recent studies suggest that external stresses applied to VO\(_2\) crystals have an effect on the transition temperature and hysteresis. Thin films of VO\(_2\) were deposited on three different cuts of sapphire by Pulsed Laser Deposition (PLD). Electron Back-Scattered Diffraction (EBSD) was used to study the orientation of the crystalline VO\(_2\) grains obtained and showed epitaxial relationships between the different single-crystal substrates. A predominant family of crystallographic relationships is present in all cuts of sapphire wherein the rutile VO\(_2\) {001} planes tend to lie parallel to the substrate's \{10-10\} and the rutile VO\(_2\) {100} planes lie parallel to the substrate's \{1-210\} and \{0001\}. This family accounts for the majority of the VO\(_2\) grains observed on all substrates with specific orientations (and stresses) depending upon the substrate's normal direction. Transmission Electron Diffraction patterns taken from cross-section of particles of the A and R cut sapphire substrates helped to lift the ambiguity present in the rutile \{100\} axes. EBSD showed additional families of relationships in the C and R cuts of sapphire, most related to the aforementioned predominant family. Variations in the transition temperatures of the deposited thin films are attributed to the stresses generated by the epitaxial relationships.
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