Environmental Safety and Health (ESH) Impacts of Emerging Nanoparticles and Byproducts from Semiconductor Manufacturing

Tasks 425.023 and 425.024

#### **Research Team**

#### **PIs:**

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#### **Other Researchers:**

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#### **Cost Share (other than core ERC funding):**

• \$80k from UA Water Sustainability Program

### **Overall Objectives**

- Characterize toxicity of current and emerging nanoparticles (NP) & NP byproducts
- Develop new rapid methodologies for assessing and predicting toxicity

## **ESH Metrics and Impact**

1. Reduction in the use or replacement of ESH-problematic materials

This project will evaluate the toxicity of various types of nanoparticles utilized or considered for application in semiconductor manufacturing, and the impact of manufacturing steps on their toxicity. This information can assist in selecting materials which are candidates for replacement or use reduction.

2. *Reduction in emission of ESH-problematic material to environment* 

The knowledge gained can be utilized to modify the manufacture of nanoparticles so that they have a lowered toxicity and thus a lowered environmental impact.

## **Surface Physical Characterization**

Hypothesis: The size and size distribution of nanoparticles intrinsically makes them more adsorptive to external chemicals, and these surface molecules lead to the observed toxic effects of nanoparticles on cells.

## **Surface Physical Characterization**

- Particle size distribution (dynamic light scattering)
- Specific area (area/volume or area/mass of NP)
- > Active site density; site energetics
- > Physical adsorption vs chemical adsorption
- Ability of the surface to concentrate bulk contaminants (selective adsorption)
- > Retention of contaminants

### **Surface Physical Characterization**

<u>Objective:</u> determine surface ability to concentrate and retain bulk contaminants. Key parameters are specific area, active site density, and surface energetics for selective adsorption



## **Experimental Method & Typical Results**



- Physical adsorption of inert adsorbent (similar to BET isotherm) for area measurement
- Chemical adsorption of reactive adsorbent for measuring site density



• Temperature-Programmed Interaction (TPI) for measuring site energetics

# <u>Comparison of Surface Activity of</u> <u>Different NP Materials</u>



Contamination retention is compound dependent: highest for CeO<sub>2</sub> and lowest for HfO<sub>2</sub>; adsorption on CeO<sub>2</sub> seems to be strong chemisorption

# **NPs Retention of Contaminants**

#### **Dynamics of Moisture Desorption**



**Contamination retention of NPs is size dependent (smaller NPs show higher retention)** 

## **Surface Characterization**

	Particle Size	Adsorption Rate Coeff.	Desorption Rate Coeff.	Active Site Density	Adsorption Capacity
	d <sub>p</sub> (nm)	k <sub>a</sub> (cm <sup>3</sup> mol <sup>-1</sup> s <sup>-1</sup> )	k <sub>d</sub> (s <sup>-1</sup> )	S <sub>0</sub> (mol/cm <sup>2</sup> )	C <sub>s0</sub> (mol/cm <sup>2</sup> )
HfO <sub>2</sub>	20	3.30E+08	2.4	7.00E-10	6.56E-10
HfO <sub>2</sub>	100	8.00E+08	0.8	2.50E-10	2.48E-10
SiO <sub>2</sub>	20	5.30E+08	360	2.00E-08	2.74E-09
CeO <sub>2</sub>	20	<b>3.00E+08</b>	1	8.75E-10	8.49E-10

• Small HfO<sub>2</sub> particles adsorbed contaminants more energetically than larger particles (*higher activation energy*)

• Small particles have higher *capacity* for adsorption and retention of secondary contaminants

## **Fractionation of CeO<sub>2</sub> by Centrifugation**

Fractioning CeO<sub>2</sub> 2g/L Eppendorf Centrifuge 4500 rpm



## **Role Surfactant Conc. on CeO<sub>2</sub> NP Size**



## **Impact Biological Media on NP Dispersions**

#### Intensity Based Particle Size Averages in Water (DLS) (units = nm)

MEDIUM	MA	TERIAL	Comment	
	HfO <sub>2</sub>	CeO <sub>2</sub>		
MQ Water	359 ± 12	1741 ± 275		
MQ Water + dispex	138 ± 2	209 ± 25		
МТТ	284 ± 2		MTT = mitochondrion toxicity test medium	
HBSS	3242 ± 270		HBSS = Hanks' Balanced Salt Solutions	
DMEM	593 ± 252		DMEM = Dulbecco's Modified Eagle Medium (+25KBS, no HCO3)	
Microtox	901 ± 406	236 ± 21		

## **Surface Chemical Characterization**

The University of Washington has a strong campus resource facility permitting us to perform state-of-the-art nanoparticle surface analysis. Instrumentation that is available for this purpose includes:

- Electron spectroscopy for chemical analysis (ESCA)
- Secondary ion mass spectrometry (SIMS)
- Surface plasmon resonance (SPR)
- Atomic force microscopy (AFM)
- Sum Frequency Generation (SFG)
- Attenuated Total Reflectance IR (ATR-IR)

# <u>Secondary Ion Mass Spectrometry (SIMS)</u> <u>Time-of-flight (ToF) SIMS; Static SIMS</u>



- Probably the most informationrich of the modern surface analysis methods
- Various organic/inorganic contaminants detected on the surface of HfO<sub>2</sub> NPs





 Positive and negative spectra can be used to identify impurities including metals from fabrication or organics from unidentified sources

# <u>Nanoparticle Impurities – ToF SIMS</u>

**Positive Spectra Impurities** 

mass	ID	Ref Micron	NP1 20 nm	NP2 1-2 nm	NP3 100 nm
27	ΔΙ	+	+		+
20					
20		Ŧ	TT		TT
30	$CH_4N$	+	+		+
40	Ca	++			+
45	$C_2H_5O$	++		++	+
46	$C_2H_6O$	+		+	+
52	$C_3H_2N$		+		+
55	Fe	+			+
58	Ni		+		
78	$C_2H_6O_3$		+		
90	Zr	++	+		+
118	$C_5H_{12}NO_2$	+		+	+
135	$C_9H_{11}O$	++		++	+
161	$C_{11}H_{13}O$	++		+++	+

"+" represents presence of listed fragment. "++" and "+++" are used to indicate relative amounts of listed fragments within row and cannot be used to compare rows one to another.

## **Nanoparticle Impurities – ToF SIMS**

Negative	Spectra	Impurities
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mass	ID	Ref	NP1	NP2	NP3
		Micron	20 nm	1-2 nm	100 nm
19	F	+++	+	++	++
26	CN	+	++		+
31	Р		+		
35	CI	+++	+	+	++
47	PO	+	++		
51	CIO	+			
59	$C_2H_3O_2$	+		++	
78	$C_3H_7OF$		+		
78.96	PO <sub>3</sub>		+		
78.92	<sup>79</sup> Br	+	++		
81	<sup>81</sup> Br	+	++		
104	$C_3H_8N_2O_2$		+		
127	I	+			
205	C <sub>13</sub> H <sub>19</sub> NO			+	

"+" represents presence of listed fragment. "++" and "+++" are used to indicate relative amounts of listed fragments within row and cannot be used to compare rows one to another.

## <u>Surface Characterization</u> <u>Summary/Preliminary Conclusions</u>

### **SIMS Analysis**

Impurity	Ref Micro	NP1 20 nm	NP2 1-2 nm
Light Organics (<100 MW)	+	+	+
Heavy Organics (>100 MW)			+
Silicon	+		+
Chlorine	+	+	
Bromine		+	
Rare Earth Metals	+	+	÷

### **Catechol Treated CeO<sub>2</sub> XPS Spectra**



#### Sodium, carbon, cerium and oxygen were observed in the spectrum

### **<u>CeO<sub>2</sub></u>** Core Level XPS Spectra Comparison



Ce 3d core level photoemission spectra

from (A)  $CeO_2$  (111), (B) Ce (III) oxide\*

\* D.R. Mullins, S.H. Overbury, D. R. Huntley. Surface Science 409 (1998) 307-319.

#### **Surface Chemistry Results and Future Plans**

- Our central hypothesis about the presence of surface contaminant species and high surface adsorptiveness of these nanoparticles is supported by our data.
- Comparison between Ce3d photoemission spectra of catechol treated CeO<sub>2</sub> and literature suggests that sample is in Ce4+ state.
- It has been shown in the literature that X-ray emission might have an effect on the oxidative state of the sample.
- In order to find the oxidative state of a pristine sample, the effect of x-ray on CeO<sub>2</sub> nanoparticles should be investigated.

## **Toxicity Assessment and Prediction**

#### **Objectives**

- Establish role for reactive oxygen species (ROS) and oxidative stress as a potential marker for NP toxicity assessment
- Develop predictable models of toxicity based on physicochemical properties elucidated by advanced surface analysis techniques
- Validate toxicity assessments and predictions with organ skin cultures (and advanced lung cultures)

## **Materials**

• Nanoparticles

Hafnium Oxide (HfO<sub>2</sub>), immersion lithography

Silica Oxide (SiO<sub>2</sub>), CMP

Ceria Oxide (CeO<sub>2</sub>), CMP



Others (Al<sub>2</sub>O<sub>3</sub>, carbon and germanium- nanotubes, quantum dots etc)

• Biological targets

Human skin cell line (HaCat)

Human lung epithelial cell line (16HBE14o-)

- Human foreskin rafted organ culture (ROC)
- Bacterium (Vibrio fischeri) Microtox test



Others (methanogens, bacterial cultures, yeast etc)

## **Methods**

#### • Main Toxicity Tests Utilized

Live/Dead Assay with HaCat Skin Cell Line (HaCat)

Mitochondrial Toxicity test (MTT) (ureter cells)

Microtox (Vibrio fischeri)

Methanogenic Activity





3-(4,5-Dimethylthiazol-2-yl)-2,5-

diphenyltetrazolium bromide

HfO, 3 ppm

Live: calcein AM)

YELLOW

HfO, 300 ppm

mitochondri reductase



Dead: ethidium homodimer-1

HfO, 3000 ppm

purple formazan

• Chemical: Reactive Oxygen Species (ROS) Production



## **Results on HfO**<sub>2</sub>

Four distinct batches of hafnium oxide tested. Example Live/Dead test (HaCat skin cells)



## **Results on CeO**<sub>2</sub>

#### Cerium oxide (MTI, "20 nm"). Example Microtox Test



## **Results on CeO**<sub>2</sub>

Cerium oxide (MTI, "20 nm"). Example Live/Dead with Dispex



## **Results on Mn<sub>2</sub>O<sub>3</sub>**

Manganese Oxide (SSNano, "40-60 nm"). Example Microtox with Dispex



### **Hypothesis ROS**



#### **Chemical Production ROS**

CeO2 (MTI, "20 nm")



Results indicate that the oxidation of L Dopa by CeO2 NP produces ROS. Direct reaction of CeO2 with dissolved oxygen and water does not produce ROS

#### **Chemical Production ROS**

Mn2O3 SSNano "40-60 nm").



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### **<u>Correspondence ROS Versus Inhibition</u>**



## **Development New Techniques**

New dye-free techniques that are less prone to interferences

• xCELLigence based on measuring impedance





• Wound healing assay, based on time to close scrape wound



## **Development New Techniques**

New dye-free techniques that are less prone to interferences (continued)

• O2 uptake assay for yeast and bacterial cells

Inhibition of Yeast, Saccharomyces cerevisiae, by CeO2



## **Preliminary Conclusions**

- Microtox Methanog L/D • HfO2, ZrO2 and CeO2 50% inhib 50% inhib 50% death NPs mild to no toxicity. ----- mg/L -----Higher Toxicity of Batch 1 HfO2 HfO2 >2000 3000 >2500 may be due to chemical >300\*\* contamination (from synthesis) CeO<sub>2</sub> **2500**<sup>\*\*</sup> >1000 >1000\*\* ZrO2 \*batch3 \*\* with dispersant
- NPs producing ROS directly in water most toxic. Chemical ROS production indicative of NP toxicity
  - $Mn_2O_3$  50% IC microtox = 70 mg/L
  - $Fe_2O_3$  50% IC microtox  $\approx$  500 mg/L
  - Fe<sup>0</sup> 50% IC microtox  $\approx$  500 mg/L

<u>Industrial Interactions and</u> <u>Technology Transfer</u>

- ISMI-Sematech (Steve Trammell, Laurie Beu)
- AMD (Reed Content)
- IBM (Arthur T. Fong)
- Intel (Steve W. Brown, Paul Zimmerman, Mansour Moinpour)

## **Future Plans**

#### **Next Year Plans**

- Fractionation of CeO2 for toxicity study size fractions
- Biochemical indicators of oxidative stress
- Complete development of new non-dye based techniques

#### **Long-Term Plans**

- Rapid screening protocols of for assessing NP toxicity
- Toxicity to organ models

## **Publications, Presentations**

- Brownbag presentation: Nanoparticle Interaction with Biological Wastewater Treatment Processes, Water Sustainability Program, Phoenix, Arizona Jan 20<sup>th</sup>, 2010 at Arizona Cooperative Extension
- Sierra-Alvarez, R. 2009. Toxicity characterization of HfO<sub>2</sub> nanoparticles. SRC/Sematech Engineering Research Center for Environmentally Benign Semiconductor Manufacturing Teleseminar Series. August 6.
- Boitano, S. 2009. Measuring cytotoxicity of nanoparticles in human cells. SRC/Sematech Engineering Research Center for Environmentally Benign Semiconductor Manufacturing Teleseminar Series. Sept. 17.
- Ratner, B. 2009. Static SIMS: A Powerful Tool to Investigate Nanoparticles and Biology. SRC/Sematech Engineering Research Center for Environmentally Benign Semiconductor Manufacturing Teleseminar Series. May 14.