### [S-3]

### Impact of Nanoparticulates on Respiratory Health Effects

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Published pulmonary toxicology studies in rats have demonstrated thatultrafine or nanoparticles (generally defined as particles in the size range < 100nm) administered to the lung cause a greater inflammatory response when compared to larger particles of identical chemical composition at equivalent mass concentrations. However, this common perception that all nanoparticles are more toxic than fine-sized particles is based upon a systematic comparison of only 3 particle-types (titanium dioxide particles, carbon black particles and diesel exhaust particles). Additional factors, other than particle size, may play more important roles in modifying pulmonary toxicity of nanoparticles. These include: surface coatings of particles; the tendency of aerosolized particles to aggregate/disaggregate; whether the particle was generated in the gas or liquid phase (i.e., furned vs. colloidal/precipitated); and surface charge. Results of pulmonary bioassay hazard studies will be presented demonstrating that fine-sized quartz particles (1.6µm) may produce greater pulmonary toxicity in rats when compared to nanoscale quartz particles (50nm) but not when compared to smaller nanoquartz sizes (e.g., < 30nm). In addition, other studies have demonstrated no difference in pulmonary toxicity between fine-sized TiO<sub>2</sub> particles (300nm) and TiO<sub>2</sub> nanodots (25nm) and nanorods. Finally, studies will be presented which demonstrate that surface coatings on particles can modify lung inflammatory effects. In summary, these are the most important conclusions:

- 1) Risk is a product of Hazard and Exposure;
- 2) one cannot assume that nanomaterials have the same toxicity as their microscale or macroscale counterparts (i.e., either greater than or less than);
- 3) therefore, each particle-type should be tested on a case-by-case basis.

## Impact of Nanoparticulates on Respiratory Health Effects

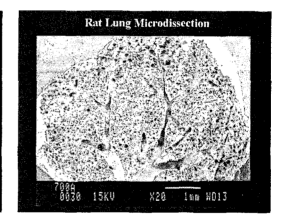
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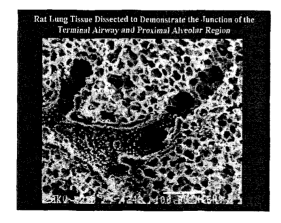
### Outline

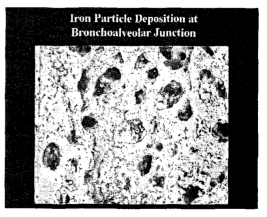
- · Lung structure and particle deposition
- Pulmonary bioassay as a measure of lung toxicity- Hazard Assessment
- Pulmonary bioassay with Fine/Nanoscale TiO<sub>2</sub> dots and rods; Fine/Nanoscale Quartz particles, and Fine/Nanoscale ZnO particles
- Impacts of Particle Surface Coatings
- Summary

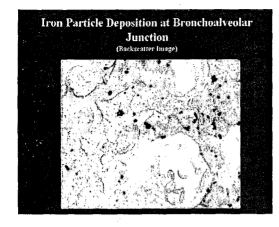
### **Definitions- Particle Size**

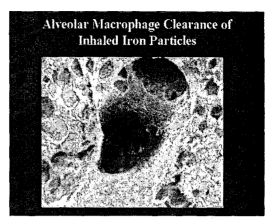
- Nano = Ultrafine = < 100 nm
- Fine = 100 nm 3  $\mu$ m
- Respirable (rat) =  $< 3 \mu m \text{ (max = 5 } \mu m)$
- Respirable (human) =  $< 5 \mu m \text{ (max = } 10 \mu m)$
- Inhalable (human) =  $\sim 10 100 \mu m$

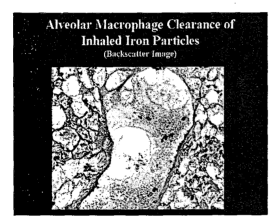


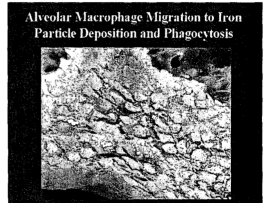


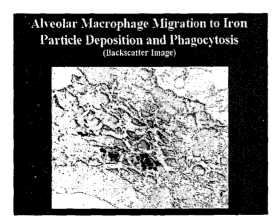


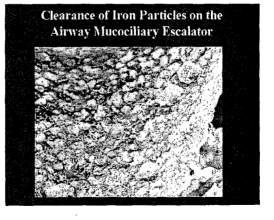


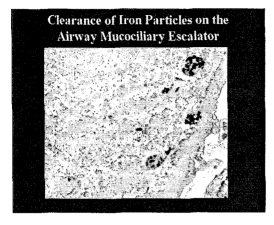


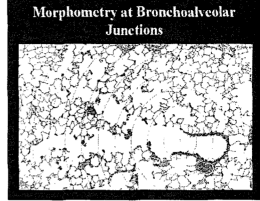












### Common Perceptions on Pulmonary Toxicity of Nanoparticles

- Nanoparticles are more toxic (inflammogenic, tumorigenic) than finesized particles of identical composition.
- Concept generally based on 3 particle-types:
  - Titanium Dioxide particles
  - Carbon Black particles
  - Diesel Particles

## Complications related to the Dogma of Nanoparticulate Toxicology

- · Not all Nanoparticles are more toxic
- · Surface coatings of particles
  - Coatings passivated or dispersion
- Species Differences in Lung Responses

   Rat is the most sensitive species
- Particle aggregation/disaggregation potential
- · Fumed vs. precipitated Nanoparticles
- · Surface charge of particles

The Key Issue: Risk

Health Risk is a product of

· Hazard and Exposure

Studies to Assess Pulmonary Hazards to Nanoparticulates

### Pulmonary Bioassay Studies

- · Working hypothesis
- · Four factors influence the development of pulmonary fibrosis
  - 1) inhaled materials which cause cell/hung injury
  - inhaled materials which promote ongoing inflammation
  - 3) inhaled materials which reduce alveolar macrophage
  - 4) inhaled materials which persist in the lung

### **Pulmonary Bioassay Components**

### Bronchoalveolar Lavage Assessments

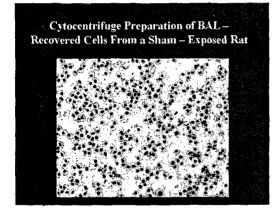
ung Inflammation & Cytotoxicity
Cell Differential Analysis

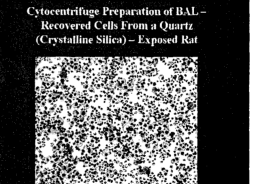
- Cen Directions analysis
  BAL Fluid Lactate Dehydrogenase (cytotoxicity)
  BAL Fluid Alkaline Phosphatase (cpithelial cell toxicity)
  BAL Fluid Protein (lung permeability)

### Lung Tissue Analysis

- Lung Weights
  Lung Cell Proliferation (BrdU)

  > Parenchymal





# Cytocentrifuge Preparation of BAL - Recovered Cells From a Carbonyl Iron - Exposed Rat

Use of Bronchoalveolar Lavage, Cell Proliferation, and Histopathology to Assess the Lung Toxicity of Particulate samples

### **Parameter**

#### Indicator

(BALF = Bronchoalveolar Lavage Fluid Analysis)

BALF Cells and Differentials BALF Luctate Dehydrogenase BALF Alkaline Phospharase BALF Protein

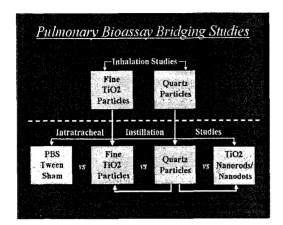
Lung Weights Macrophage phagocytosis Cell Proliferation

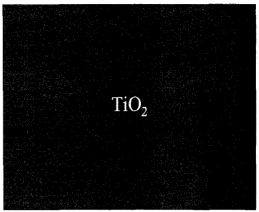
Histopathology

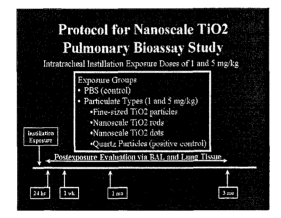
Lung Inflammation Non-specific cytotoxicity Type 2 cell epithelial toxicity Permeability of alveolar capillary barrier Pulmonary edema or fibrosis

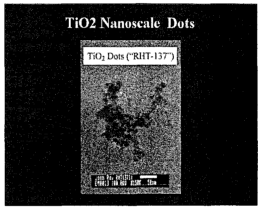
Lung clearance functions Inflammation lung fibrosis and tumor potential

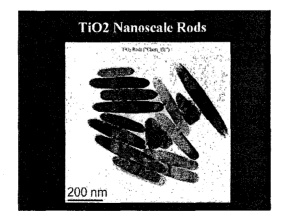
Evaluation of lung tissue responses

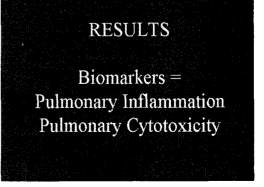


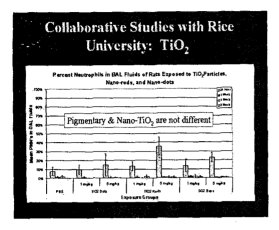


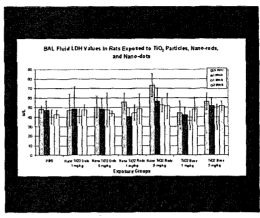










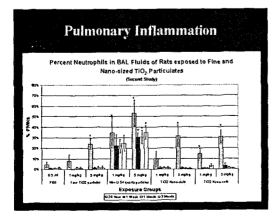


### Characterization of Nanoscale TiO<sub>2</sub> Particles

XRD particle size Surface Area

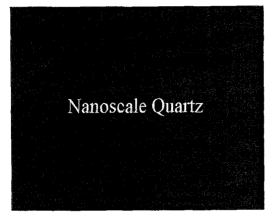
- Fine  $TiO_2$  rutile  $d_{50} = 300 \text{ nm}$  6 m<sup>2</sup>/g
- TiO<sub>2</sub> Nanorods anatase length= 90 233 nm width = 20 35 nm 26.5 m<sup>2</sup>/g
- $TiO_2$  Nanodots anatase  $d_{50} = 6$  nm 169.4 m<sup>2</sup>/g

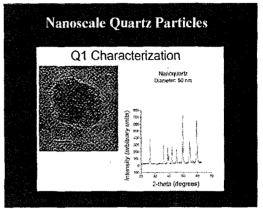
Second Nanoscale TiO2 Study

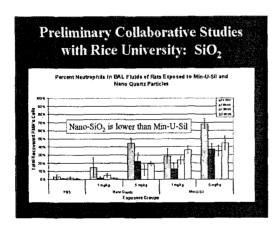


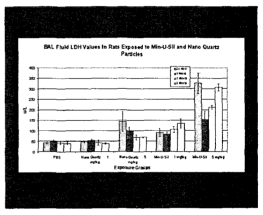
### Hypothesis and a Question

- Hypothesis: At similar doses Ultrafine (Nano) particles have greater pulmonary toxicity than fine-sized particles of identical composition.
- Question generally this dogma applies to low toxicity particulates. However, in considering a cytotoxic particle such as crystalline silica – would nanoquartz particles be even more toxic than finesized Min-U-Sil quartz particles?

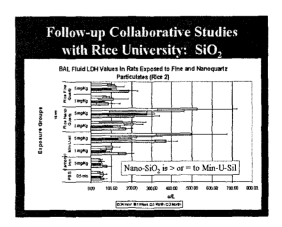


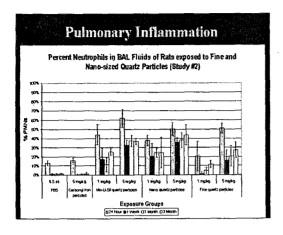


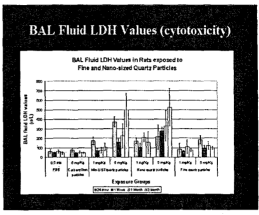


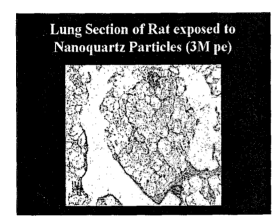


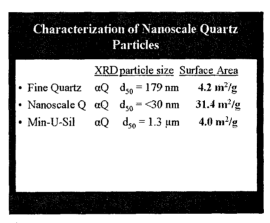
Second Nanoscale Quartz Study

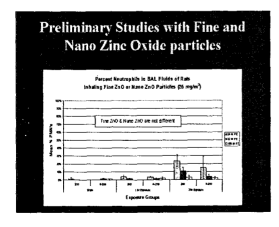


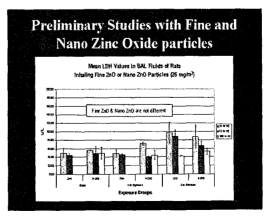










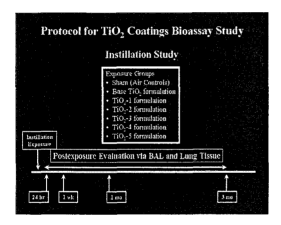


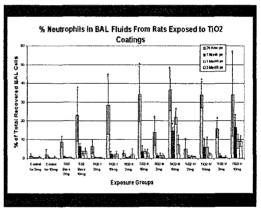
## Impact of Surface Treatments/Coatings on TiO<sub>2</sub> Particles

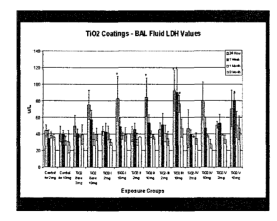
- · Inhalation Studies
- Pulmonary Bioassay Intratracheal Instillation Studies

### TiO<sub>2</sub> Coatings Formulations

- TiO2 base 99% TiO2 1% alumina
- TiO2 I 99% TiO2 1% alumina + organic grinding aid
- · TiO2 II 96% TiO2 4% alumina
- TiO2 III 83% TiO2 6% alumina 11% amorphous silica
- TiO2 IV 91% TiO2 3% alumina 6% amorphous silica
- TiO2 V 94% TiO2 3% alumina 3% amorphous silica







### **Important Particle Characteristics**

- · Primary particle size
- Particle shape (SEM)
- · Surface area
- · Surface charge
- · Composition- e.g crystalline vs.amorphous
- Surface Coatings
- · Aggregation status
- · Particle number
- Method of synthesis (gas vs. liquid phase)

### Summary

- Risk is a product of Hazard and Exposure
- Cannot assume that nanomaterials are the same as their bulk counterpart
- Each particle-type should be tested on a case-by-case basis

### Acknowledgments

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- Tom Webb and Ken Reed provided the pulmonary toxicology technical expertise for the study. Denise Hoban, Elizabeth Wilkinson and Rachel Cushwa conducted the BAL fluid biomarker assessments. Carolyn Lloyd, Lisa Lewis, John Barr prepared lung tissue sections and conducted the BrdU cell proliferation staining methods. Don Hildabrandt provided animal resource care.