



Microstructural Characterization of PEM Fuel Cell MEAs

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Project ID FC3

Project Overview

Timeline

- Project initiated in FY2000
- Continuous fundamental research on microstructural characterization to improve MEA durability

Budget

- Total Funding to date ~\$1.6M
- Funding in FY06 \$375k (\$300k baseline project plus \$75k project with Arkema)
- Funding in FY07 \$300k

Barriers

- Fuel Cell Barriers Addressed
 - A: Durability
 - B: Cost
 - C: Performance

Partners

- Los Alamos National Laboratory
- Argonne National Laboratory
- PlugPower
- Honda Research Institute
- Rensselaer Polytechnic Institute
- Arkema



ORNL Research Objectives

- Identify high-resolution imaging and compositional/chemical analysis techniques for characterization of the material constituents comprising PEM fuel cell MEAs
 - $-\mu m$ to nm-scale using FEG-SEM
 - nm- to sub-angstrom scale imaging using TEM/STEM
 - Compositional analysis using TEM/STEM with EDS
 - Surface chemistry using XPS
- Apply these analytical and imaging techniques for the evaluation of microstructural and microchemical changes to MEA materials during life-testing
- Elucidate microstructure-related degradation mechanisms contributing to PEM fuel cell performance loss



Approach: Used Advanced Imaging and Compositional Analysis Techniques to Evaluate Atomic-Scale MEA Microstructures

- Develop innovative methodologies to prepare samples for microstructural analysis of MEA constituents
 - The use of "partial-embedding" for microtomy
 - Cryo-microtomy/cryo-transfer for membrane analysis in FEG-TEM
- Apply state-of-the-art electron microscopy techniques for the analysis of MEA materials
 - High-resolution FEG-SEM Hitachi S4800
 - High-resolution FEG-TEM imaging (nm-scale) Hitachi HF-2000
 - High-angle annular dark field (HAADF) imaging (Z-contrast imaging) in an aberration-corrected STEM (*sub-angstrom scale*) - JEOL 2200FS-AC
- Collaborate with industry, academia, and national laboratories to make these techniques (and expertise) available for MEA processing and/or life-testing studies



Technical Accomplishments and Progress ORNL/LANL Collaboration

- Primary focus has been on characterization of an E-TEK 20 wt% Pt-Co (3:1) bimetallic catalyst
- As-received powder Pt-Co on Vulcan XC-72
- Fresh MEA with Pt-Co/C cathode
- Same MEA aged to evaluate catalyst changes:
 - H₂/N₂ cycled 0.4-0.96 V @ 80°C, <u>100% RH</u>, ~35k cycles
 - H₂/N₂ cycled 0.4-0.96 V @ 80°C, <u>50% RH</u>, ~40k cycles
 - DOE drive cycle for 8905 cycles (results not complete)



Bimetallic Catalysts are Being Evaluated to Replace Pt-only Catalysts

- Pt is expensive and supply is limited
- Pt alloyed with transition metals (M=Co, Ni, Cr) have shown enhanced catalytic activity compared with Pt
 - Changes in Pt-Pt and Pt-M bond lengths
 - Changes in atomic coordination/nearest neighbors
 - Electron density of states in Pt 5d orbital
 - Nature of surface
- Atomic ordering of the Pt alloys will enhance the catalytic activity
- Specific crystallographic planes/facets can enhance activity

References: Stamenkovic et al., J. Phys. Chem. B 106 (2002) 11970.

Xiong and Manthiram, *J. Mater. Chem.* 14 (2004) 1454.
Mun et al., *J. Chem. Phys.* 123 (2005) 204717.
Gasteiger et al., *Appl. Cat. B* 56 (2005) 9.
Xiong and Manthiram, *J. Electrochem. Soc.* 152(4) (2005) A697.



Size and Size-Distribution Differences Between Pt and Pt-Co Catalyst Particles



Crystal Structure of E-TEK 20 wt% Pt-Co/XC-72



The Crystal Structure of Individual Pt-Co Particles Is Determined From HAADF-STEM Images



The Crystal Structure of Individual Pt-Co Particles Is Determined From HAADF-STEM Images



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JT-BATTE

During H₂/N₂ Cycling (0.4-0.96 V) at 80°C, Loss Of EASA Observed for Pt-Co Cathode Catalysts



- Pt-Co exhibits lower loss of surface area compared with Pt during potential cycling
- Higher %RH enhances particle growth

Borup et al., 2006 DOE Hydrogen Program Review, FC28 for Pt results.



During H₂/N₂ Cycling (0.1-0.96 V), Growth of Pt-Co Particles Is Quantified *f*(%RH) by TEM



Pt-Co particles in fresh MEA cathode



• 2<u>0 nm</u>

Pt-Co ~36k cycles @ 100% RH



There is a change in the form of the Pt-Co particle size distribution, which is more pronounced following cycling at 100% RH.



During H₂/N₂ Cycling (0.4-0.96 V), Growth of Pt-Co Particles Occurs Via Combined Mechanisms: Dissolution-Reprecipitation and Coalescence



Typical particle growth associated with an Ostwald Ripening process - large particles grow at the expense of small particles such that surface energy is reduced.

Size distribution shift from narrow distribution of small particles to wider distribution of larger particles

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Kinetic particle growth associated with combined mechanisms of dissolutionreprecipitation and particle coalescence.

More gradual size distribution change from a narrow distribution of small particles to a bimodal particle size distribution that includes small and large particles



What About Other Ordered Bimetallic Catalysts?



XRD data clearly shows superlattice reflections for Pt₃Cr particles and smaller average particle size

STEM shows differences in particle size distributions between Pt-Co and Pt₃Cr

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Pt₃Cr particles



Pt-Co particles

Unlike Pt-Co Catalyst, All Pt₃Cr Particles Ordered



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Many Pt-Based Bimetallic Particles Exhibit No Order



Pt-W "Assembles" Into Particles That Are Not Always Crystalline - Pt Atoms Cover Surface of KB Support



Note that individual, non-assembled Pt and/or M species on the carbon surface were not observed for highly crystalline, ordered bimetallic catalysts (Pt-Co or Pt-Cr systems)



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Technical Accomplishments and Progress ORNL/ANL Collaboration

10 wt% Pt/Vulcan imbibed with Nafion solution -supported on carbon fiber cloth

cycled 32 times (0.01-1.4V) in
0.6 M perchloric acid electrolyte

• cycled (as above) then held 72h at 1.1 V (vs. RHE)

cycled (as above) then held
72h at 1.2 V (vs. RHE)

Interested in Pt particle size changes





5 μm

TEM Used to Evaluate Pt Particle Size Changes



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TEM Used to Evaluate Pt Particle Size Changes



Future Work

- Apply the HAADF STEM imaging technique to fresh and electrochemically-aged MEAs in addition to new bimetallic catalysts. This technique will be especially useful for identifying species migration within the cathode and into the membrane.
- Characterize carbon support microstructural degradation (this has proved to be a difficult challenge because of the recast ionomer in electrodes!)
- Continue to establish collaborations with industries, universities, and national laboratories (including access to ORNL User Facilities) to facilitate "transfer" of unique capabilities.
- Support new DOE projects with microstructural characterization and technique development.



Summary

- The HAADF STEM (Z-contrast) imaging technique has been used to characterize the degree of atomic ordering in numerous Pt-based bimetallic catalyst systems:
 - Pt-Co catalyst shows mostly disordered Pt-Co alloy particles, with some ordered Pt₃Co particles and (Pt₃Co/Pt-Co) particles observed in the distribution. Pt-Co had a much larger particle size that Pt-only and a wider particle size distribution
 - Pt₃Cr catalyst particles are highly ordered (nearly 100% of particles) with a very uniform size distribution
 - Pt-W had a very narrow particle size distribution with particles <2.0 nm; much of the Pt-W remained atomically dispersed on the carbon support surface and did not assemble into crystalline particles.
 - Bimetallic catalyst systems also being evaluated include Pt-Ti and Pt-Ru
- Amount of %RH has a similar effect on EASA loss of Pt-Co as observed for Pt-only cathode catalysts, but to a much lesser degree
- Small Pt particle size differences observed for cycled/held ANL samples, but additional work should be done to image Pt within ionomer using HAADF imaging

