Hydrogen, Fuel Cells & Infrastructure Technologies Program, 2006 Annual Review Arlington, Virginia – May 16-19, 2006

Non-Platinum Cathode Catalysts

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Project ID FC14

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Project Overview

<u>Timeline</u>

- Start date Jan 2004
- End date TBD

Budget

- **FY05 Funding** \$350K
- **FY06 Funding** \$500K

Technical Barriers

- A. Durability
 - catalyst
 - electrode layer
- B. Cost
 - catalyst
 - -MEA
- C. Electrode Performance
 - ORR overpotential
 - O₂ mass transport



Partners

- University of Illinois, Urbana-Champaign (Andrzej Wieckowski)
- Université de Poitiers, Poitiers, France (Nicolas Alonso-Vante)
- University of New Mexico, Albuquerque (Plamen Atanassov)
- University of California, Riverside (Yushan Yan)
- Mesoscopic Devices LLC, Denver (Jerry Martin)

Targets

DOE Targets: Electrocatalysts for Transportation Applications (Stack)				
Characteristics (Units)	2004 Status	2010	2015	
PGM Total Content (g/kW)	1.3	0.5	0.4	
PGM Total Loading (mg/cm ²)	0.8	0.3	0.2	
Cost (\$/kW _e)	20	8	8	
Durability with cycling @ T \leq 80°C (h)	1,000	5,000	5,000	
Activity (µA/cm ² @ 0.9 V _{iR-free})	180	720	720	
Non-Pt Catalyst Activity (A/cm ³ @ 0.8 V _{iR-free})	< 8	> 130	300	

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Objectives

Primary Objective:

Develop low-cost non-platinum oxygen reduction reaction (ORR) catalysts for the polymer electrolyte fuel cell (PEFC) cathode, with similar activity and performance durability to the currently used Pt-based cathode catalysts

Individual Objectives:

- Identify and/or synthesize new cathode catalysts
- Determine performance of these catalysts, including ORR kinetics, low-pH stability, performance durability, etc.
- Establish the ORR mechanism, in particular, identify ORR active sites
- Design and optimize the performance of membrane-electrode assemblies (MEAs) with new catalyst used at the cathode
- Test and optimize catalyst performance durability
- Collaborate with the fuel cell industry on efficient integration of new catalysts into MEAs and facilitate catalyst technology transfer



Approach

- Develop advanced non-platinum cathode catalysts with either minimum precious metal content or entirely free of precious metals
 - Non-precious metal / heteroatomic polymer nanocomposites
 - "Chalcogenide-type" ORR catalysts by surface modification
 - *a.* Organic solvent route *b.* Aqueous route *c.* PGM loading reduction via core-shell approach
- Through experimentation develop understanding of the ORR mechanism and factors impacting catalyst performance and durability
- Maximize performance, utilization of PGM-based catalysts and achievable loading of non-precious metal catalysts by designing novel ("open frame") cathodes



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Non-Precious Metal/Heteroatomic Polymer Nanocomposites

- *Hypothesis:* CoN2 (CoN4) sites claimed to act as ORR active sites (e.g. in pyrolized Co porphyrins)
- **Objective:** Generate ORR active sites <u>without</u> destroying ordered structure of the catalyst
- Approach: Heteroatomic polymer as a matrix for entrapping and stabilizing non-precious metal
- Choice: Cobalt-polypyrrole-carbon composite (Co-PPY-XC72)







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Synthesis & Basic Properties of the Co-PPY-XC72 Nanocomposite



Composite	Surface Area (m ² /g)
XC72	240
PPY-XC72 PPY-XC72 (reduced)	100 100
Co-PPY-XC72	124
Co-PPY-XC72 (pyrolized)	140



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Fuel Cell Testing of the Co-PPY-XC72 Nanocomposite



• OCV in excess of 0.85 V in both O₂ and air operation

• PEFC cathode performance of the Co-PPY-XC72 nanocomposite close to that of the best performing pyrolized porphyrin catalysts

A new class of non-precious metal ORR catalysts identified!



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Does the "By-Design" Structure Matter?



Pyrolysis: 2 h @ 800°C; N₂ (T-change: 10°C/min)

 Cell performance of Co-PPY-XC72 composite ~10× better at 0.4 V than that of carbon-supported Co catalyst

 Cell performance of non-pyrolized Co composite ~7× better at 0.4 V than that of pyrolized Co-composite

Co-N site crucial to ORR activity



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Co-PPY-XC72 Nanocomposite: Performance Durability



- Stable operation of the Co nanocomposite catalyst for 110 hours
- Major improvement in stability over pyrolized-porphyrin catalysts

Unique performance durability for a non-precious metal catalyst at low pH



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Structural Characterization by XANES and EXAFS



- <u>XANES</u>: Both Co(0) present in as-synthesized catalyst; Co(II)-to-Co(0) ratio increasing during catalyst break-in
- <u>EXAFS</u>: Most of Co(0) transforming to Co(II)-O/N (i.e. O or N) states (not CoO)
- <u>EXAFS</u>: Good stability in time-resolved experiments (0th, 24th, and 190th hour)



Co-PPY-XC72 Nanocomposite: 4e⁻ vs. 2e⁻ Process (RRDE Study)





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Co-PPY-XC72 Nanocomposite: ORR Kinetics



High performance, matching or exceeding that of 'state-of-the-art' non-precious catalysts, achieved in the first year of nanocomposite research!

Performance milestone for non-precious metal catalysts achieved

(cf. Supporting Information)

Characteristics (Units)	Value	
ORR activation energy, E _a ° (kJ mol ⁻¹)	40.6	
ORR activity in H ₂ -O ₂ fuel cell at specified voltage, 80°C (A cm ⁻³)	0.80 V	4.9
	0.70 V	24.3
O ₂ turnover at RDE, specified potential <i>vs.</i> NHE, 25°C (10 ⁻⁵ site ⁻¹ s ⁻¹)	0.80 V	5.5
	0.70 V	25.0
O ₂ turnover in H ₂ -O ₂ fuel cell at specified voltage, 80°C (10 ⁻³ site ⁻¹ s ⁻¹)	0.80 V	3.2
	0.70 V	14.0



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"Decorated" Ru Nanoparticles for Oxygen Reduction





Project in close collaboration with

- University of Illinois, Urbana-Champaign
- Université de Poitiers, France



- Reduced Ru a very good ORR catalyst but prone to oxidation
- Ru decoration with selenium:
 - Se + $Ru \rightarrow$ Se/Ru
- New, and different than Ru, chalcogenide-type, oxidationresistant catalyst synthesized

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Se/Ru Catalyst: Structure by XRD and EDX



Se/Ru Catalyst: Fuel-Cell Evaluation of Performance



Electrocatalysts for Transportation Applications (Stack)				
Characteristics (Units)	Se/Ru Organic Path	Se/Ru Aqueous Path	2010 target	
Non-Pt Catalyst Activity (A/cm ³ @ 0.8 V _{iR-free})	19	43	> 130	



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Se/Ru Catalysts: 4e⁻ vs. 2e⁻ Process (RRDE Study)



Number of electrons: $n = (4j_{disk}) / (j_{disk} + j_{ring}/\varepsilon)$ (ε - ring collection efficiency)Peroxide yield: $\% H_2O_2 = 100 (4 - n) / 2$

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Se/Ru Catalysts: RRDE Evaluation of ORR Kinetics

RRDE Kinetic Evaluation of Se/Ru Catalysts			
Characteristics (Units)	Value		
Kinetic current at RDE, specified potential <i>vs.</i> NHE, 25°C (10 ⁻⁵ A cm ⁻²) (Surface area estimated by spheres)	0.75 V	0.8	
	0.70 V	2.0	
	0.65 V	4.0	
Apparent rate constant at RDE, specified potential <i>vs.</i> NHE, 25°C (cm ⁴ mol ⁻¹ s ⁻¹)	0.75 V	0.1	
	0.70 V	0.3	
	0.65 V	0.6	
O ₂ turnover at RDE, specified potential <i>vs.</i> NHE, 25°C (10 ⁻³ site ⁻¹ s ⁻¹)	0.75 V	9	
	0.70 V	20	
	0.65 V	40	

- <u>RRDE</u>: Significantly higher O₂ turnovers for Se/Ru than nanocomposites
- <u>RRDE & fuel cells</u>: Mass transport limitations much less pronounced at surface chalcogenides than nanocomposites
- *Fuel cells:* Aqueous path yielding Se/Ru with higher activity than organic path
- <u>Fuel cells</u>: "Volumetric" activity of Se/Ru at ~ <u>30%</u> of the DOE 2010 target



Surface Chalcogenide Catalysts: Durability and Lowering Ru Content



Summary

- Demonstrated <u>a new class</u> of non-precious metal/heteroatomic polymer nanocomposite catalysts, such as Co and Fe composites, with promising ORR activity and unique for non-precious metal catalysts performance durability
- Identified a dual role of the heteroatomic polymer: (i) formation of the active ORR site and (ii) stabilization of the non-precious metal center via the formation of a bond with the heteroatom
- Confirmed by XANES/EXAFS the key role of the Co-N (or Co-O)
- Developed two new synthesis paths for fabricating surface chalcogenide Se/Ru catalysts via "decoration" of Ru nanoparticles by Se
- Demonstrated very high fuel cell activity (<u>30% of the DOE's 2010 activity target</u>) and respectable performance stability of Se/Ru catalysts
- In RDE/RRDE experiments, determined ORR kinetic parameters on nanocomposites and chalcogenides and identified inefficient O₂ mass-transport in the catalyst layer as a limiting factor for the scale-up of the nanocomposite performance with catalyst loading
- Experimentally verified validity of the core-shell approach for reducing Ru loading via addition of a non-precious metal (Fe)



Future Work

Remainder of FY06:

- Complete detailed characterization of oxygen reduction on Co nanocomposite
- Finish, already advanced, RDE/RRDE study of ORR kinetics on the cobalt nanocomposite and two surface chalcogenides
- Initiate development of methods for reducing O₂ concentration overpotential in the PEFC cathode

FY07:

- Develop novel mesoporous "open-frame" structures for improved mass transport of O₂ in catalyst layers to (i) maximize catalyst utilization (surface chalcogenide) and (ii) allow for higher catalyst loading (nanocomposites)
- Synthesize and characterize composites based on other heteroatomic polymers, e.g. polyaniline, poly(vinyl) pyridine, poly(ethylene dioxy) thiophene, and non-precious metals other than cobalt, e.g. Ni, Fe, W, Mo, and their alloys
- Establish collaborative effort with the university partners for lowering Ru content in surface chalcogenides (alloying with non-precious metals; "coreshell" catalysts) and further increasing activity ("ORR activators")
- Perform structure-mechanism correlations; propose approaches to lowering ORR overpotential on non-Pt catalysts



Non-Platinum Cathode Catalysts

Supporting Information

Responses to Reviewers' Comments

"CoTPP types of catalyst are just not sufficiently stable for consideration."

Based on the durability data acquired in the first year of the project (2004-2005), metalloporphyrins have been deleted from the scope and replaced by much more stable non-precious metal/heteroatomic polymer nanocomposites.

"Further emphasis on understanding mechanisms rather than demonstrating the highest fuel cell performance should be considered."

Catalysts in both studied groups have become subject of a detailed mechanistic analysis. The focus of that analysis has been on the identification of the active reaction site and ORR kinetics. XANES/EXAFS, SEM EDX, and XRD/RRDE have been used extensively as complementary techniques to fuel cell testing.

"Unclear if on track to meeting key DOE technical barriers/targets."

Performance of all catalysts has been referred to the DOE targets for non-Pt electrocatalysts, in particular to the 2010 "volumetric" activity target (130 A/cm³ at 0.8 V, iR-corrected). Se/Ru catalyst has already shown activity greater than 30% of that target.

"Program requires reliable, reproducible catalyst sources."

This project focuses on entirely new materials, none of them available commercially. Catalysts from external sources as well as synthesized at LANL have been carefully evaluated for purity, noble-metal contamination, and reproducibility (cf. batch-tobatch consistency data for the Co nanocomposite in "Supporting Information").



Selected Project-Relevant Publications and Presentations

- 1. "New Class of Non-precious Metal Composite Catalysts for Fuel Cells," B. Rajesh, P. Zelenay, Nature, submitted
- 2. University of California Riverside, Department of Chemical and Environmental Engineering, Riverside, California, October 14, 2005. Title: "Polymer Electrolyte Fuel Cells: Highlights from the Fundamental and Applied Research at Los Alamos," P. Zelenay (invited lecture)
- 3. 207th Meeting of the Electrochemical Society, Los Angeles, California, October 16-21, 2005. Title: "A New Non-Precious Metal Catalyst for Oxygen reduction," B. Rajesh,* P. Zelenay
- 4. NANO Commerce SEMI NanoForum, Chicago, Illinois, November 1-3, 2005. Distributed Energy Systems–Panel; moderated by David Forman, Small Times Magazine, P. Zelenay* (invited panelist)
- 5. 2005 Fuel Cell Seminar, Palm Springs, California, November 14-18, 2005. Title: "Compact, Portable, and Robust DMFC System Using Mixed-reactants" P. Zelenay, V. Hovland, A. Kulprathipanja,* J. Martin
- 6. International Battery Association & Hawaii Battery Conference, Waikoloa, Hawaii, January 9-12, 2006. Title: "Non-Platinum Electrocatalysis – A Major Challenge for Polymer Electrolyte Fuel Cells," R. Bashyam, J.-H. Choi and P. Zelenay* (invited lecture)
- 7. LANL Materials Science and Technology Division Review Meeting, April 4-6, 2006. Title: "New class of non-precious metal nanocomposite electrocatalysts for fuel cells,"
 B. Rajesh,* R. Mukundan, E. Brosha, F. Garzon, S. Conradson, P. Zelenay



- Nanocomposite-type catalysts: Development of an open-frame, highly O₂-permeable cathode structure allowing for a significant increase in the catalyst loading without introducing a mass-transfer hindrance
- Surface chalcogenides: Reduction in the Ru content through partial replacement by a non-precious metal, such as Fe ("coreshell" approach); also, further enhancement in the O₂ turnover, for example, via an introduction of ORR activators
- All non-platinum catalysts: Sufficient stability in the very acidic, high-temperature and oxidizing environment of the polymer electrolyte fuel cell cathode



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- Identify the active ORR site of a selected well-performing nanocomposite*. – Close to completion
- Determine the kinetic parameters and turnover rates on nanocomposite and surface chalcogenide catalysts. – Close to completion
- Demonstrate a Co-composite catalyst with performance and performance stability comparable to the state-of-the-art nonprecious metal cathode catalysts. – Achieved (with developed in-house a new class of catalysts)



^{*}Non-precious metal/heteroatomic polymer composites replace less promising metalloporphyrins

Loading Effect and Batch-to-Batch Consistency



Fe-based Nanocomposite



• Similar performance of Fe and Co nanocomposites at low current densities

• Significant mass-transfer limitations observed with Fe-PPY-XC72 below 0.65 V

A <u>class</u> of non-precious metal/heteroatomic polymer composites demonstrated



"Aqueous" Se/Ru Catalyst: DMFC Performance



 Very good methanol tolerance of the "aqueous-path" Se/Ru catalyst in cells with up to ~17 M methanol concentration in the anode feed stream

 Delivered maximum power of 65 mW cm⁻² and 43 mW cm⁻² with 0.5 M and 17.0 M methanol, respectively, ~2× better than cells with "organic-path" Se/Ru catalyst



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