Low Pt Loading Fuel Cell Electrocatalysts

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(This presentation does not contain any proprietary or confidential information.) Project ID # FC09

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Overview

Timeline

- Project start date: 06.02.
- Project end date: Multi-year
- Percent complete

Budget

- Total project funding: \$1314K
- DOE share: \$1314K
- Funding received in FY05: \$330K
- Funding for FY06: \$360K

Barriers addressed

- 1. Precious metal loading
- 2. Electrocatalysts' Activity
- 3. Electrocatalysts' Durability

Target	2010:	0.3 g/kW
Target	2010:	5000hr

Collaborations

- Los Alamos National Laboratory (Fuel cell tests F. Uribe, P. Zelenay)
- Battelle Memorial Institute (CRADA Scale-up of synthesis J. Sayre and A. Kawczak)
- 3M (PdCo catalyst, exploratory activities R. Atanasoski)
- Plug Power Test of the PtRu₂₀ anode catalyst: 2390 hr showed a small loss in activity (B. Do).
- General Motors Co. planning stage (F. Wagner)

Objectives

--- Comprehensive

To assist the DOE in the development of fuel cell technologies by providing low-platinum-loading electrocatalysts.

--- During the current year

- To demonstrate the stability of Pt mixed metal monolayer (Pt₈₀Ir₂₀/Pd/C) and Pt/Au/Ni/C electrocatalysts in fuel cell tests.
- To explore a novel class of electrocatalysts for O₂ reduction consisting of **Pt monolayer on** *noble metal -non-noble metal core-shell* **nanoparticles**.
- Further studies of the electrocatalysts with **low**, or **no Pt**.
- Scale up of synthesis based on displacement of a Cu monolayer.
- Addressing the problem of Pt dissolution under potential cycling regimes.
- Testing the effects of Au clusters on Pt stability.

APPROACH

Pt monolayer electrocatalysts

Why Pt monolayer catalysts?

- Complete Pt utilization (of all atoms that are not blocked by Nafion[®])
- Ultimate reduction of Pt loading
- Increased activity
- (Six patents pending)

For 3-5nm nanoparticles ~25% of atoms are on the surface, ~75% are not available for catalysis. Three types of Pt electrocatalysts:

1. <u>Pt on Pd nanoparticles</u>

Improved activity

Improved durability

2. <u>Mixed-metal Pt on Pd nanoparticles</u>

Even higher activity Durability tests at LANL

and Battelle



Pt

Pd

3. <u>Pt on noble/non-noble core-shell nanoparticles</u>

Yet higher activity per mass of noble metal

Durability to be tested





Long-term test (one segment) of the anode PtRu₂₀ electrocatalyst at PLUG POWER



Compared to commercial MEA baselines (0.6 mg Pt/Ru/cm²): Higher than neat H₂ baseline; lower than reformate baseline. Total run time: 2394 hours

New class of catalysts: Pt monolayer on *noble metal - non-noble metal core-shell* nanoparticles



New class of electrocatalysts that facilitate a **further decrease of noble metal content** while possessing a very high activity.



Pt O₂ Reduction Kinetics and Mass Activities of Pt_{ML}/PtCo₅/C and Pt_{ML}/PdCo₅/C



Pt mass activity

Total noble metal mass activity



2- to 5-fold increase in noble metal mass activity with Pt on core-shell nanoparticles



O₂ Reduction Kinetics and Mass Activities of Pt_{ML}/NbO₂/C and Pt/C/C

Pt/C



Pt_{ML}/NbO₂/C has 3 times higher mass activity than the commercial Pt/C

Pt/NbO₂/C

Pt/C

Formation of Au clusters on Pt electrodes



E / V RHE



Effect of submonolayer of Au clusters on O₂ reduction on Pt(111) and Pt/C



Small inhibition of O₂ reduction by Au, not in proportion to Au coverage



Effect of potential cycling on the activity of Pt/C in the ORR

30,000 cycles from 0.6 to 1.1V at 25°C in O₂-saturated solution



A negative shift of 40mV in $E_{1/2}$ and a loss of surface area of 45%

Stabilization of Pt/C by a submonolayer of Au clusters under a potential cycling regime

30,000 cycles from 0.6 to 1.1V at 25°C in O₂-saturated solution



A negligible shift in $E_{1/2}$ and a loss of surface area



Stabilization of Pt_{ML}/Pd/C by a submonolayer of Au nanoparticles under a potential cycling regime

30,000 cycles from 0.6 to 1.1V at 25°C in O₂-saturated solution



A negative shift of 26mV in E_{1/2}

A negligible shift in E_{1/2}



Pd₃Fe alloy O₂ reduction electrocatalyst



The activity of Pd_3Fe/C is as high as that of commercial Pt/C.

Compressive strain-induced downshift of the d - band center decreases the activity of Pd and the blocking effect of OH, O_2 , O_2^- , H_2O_2 .



The optimal Pd-Pd distance is 0.273 nm.



Fine tuning of the Pt-substrate interaction: Very high activity of Pt_{ML}/Pd₃Fe/C



Pt_{ML}/Pd3</sub>Fe/C has 0.14g/kW, which meets 0.15g/kW,Total noble mi.e., 1/2 of 0.3g/kW, the DOE's target for 20105 times that of

Total noble metal mass activity is about 5 times that of the commercial Pt catalyst.

Methanol tolerance of the Pd₂Co electrocatalyst

Pd₂Co

XANES data indicates pronounced electronic effects in Pd-Co and an increased stability of Pd, its activity being close to that of Pt(10%)/C.

Fuel cell methanol tolerance test by P. Zelenay et al., LANL



BNL#101-103 are from one sample obtained by different acid wash procedure.



Dual-Pathway Kinetic Equation for the Hydrogen Oxidation Reaction on Pt Electrodes

The Butler-Volmer equation, $j_k = j_0 \quad (e^{2.3\eta/b} - e^{-2.3\eta/b})$, appears to be inappropriate for describing the kinetics of the H₂ oxidation reaction (HOR) on a Pt electrode (1,2).

- 1. Gasteiger, et al., Power Sources, 127 (2004), 162.
- 2. Chen, Kucernak, J. Phys. Chem. B 108 (2004) 13984.
- 3. J. X. Wang, T. E. Springer, R. R. Adzic, J. Electrochem. Soc. Submitted.

Based on the Tafel-Heyrovsky-Volmer mechanism, we derived a new equation, ref. 3, to describe the HOR on Pt over the entire potential region:

$$j_{k} = j_{0T} (1 - e^{-F\eta/\beta RT}) + j_{0H} (e^{F\eta/2RT} - e^{-F\eta/(2+\beta)RT})$$

A fast, inverse exponential rising of kinetic current at small overpotentials through the Tafel-Volmer pathway, j_{TV} , and a gradual rise at $\eta > 50$ mV through the Heyrovsky-Volmer pathway, j_{HV} .

The anode behavior in a PEM fuel cell can now be well understood.

New basis for fuel cell modeling, optimization, and diagnosis.



Measured (symbols, from ref [2]) and fitted (solid lines using the Eq.) polarization curves for the HOR on Pt microelectrodes



Calculated kinetic current (solid line), components, jTV (dashed line) and jHV (dotted line), the kinetic current from the Butler-Volmer equation (dotted-dashed line).



Future Work

Remainder of FY2006

- Pt_{ML}/Pd/C electrocatalyst
 - Post fuel cell tests Z-contrast TEM, XANES.
- Pd₂Co electrocatalyst
 - Stability studies; and fuel cell tests.
- Mixed-metal Pt monolayer electrocatalysts
 - Stability studies. Effects of Au clusters in MEA's under potential cycling regimes.

FY 2007

- Stabilization effects of Au clusters:
 - RDE and MEA studies under potential cycling conditions with Pt/Pd/C; Pt/PdCo₅/C.
- Pt/AuNi/C electrocatalyst
 - Segregation of Pt, Au; Stability tests; Fuel cell tests.
- Further reduction of Pd content: Pt monolayers on core-shell nanoparticles
 - Basic in situ surface science and electrochemical studies; stability and fuel cell tests.

Summary: Pt monolayer electrocatalysts

BNL ANODE PtRu₂₀/C electrocatalyst:

Durability and CO tolerance confirmed in a 2400h test at Plug Power. The performance of **0.063** g_{Pt}/kW meets the 2010 target of **0.15g/kW**, i.e., 1/2 of 0.5g/kW (Ru not counted).

BNL CATHODE electrocatalysts:

- Novel class of electrocatalysts developed: Pt monolayer on *noble metal - non-noble metal core-shell* nanoparticles. The performance of 0.14g/kW meets the target for 2010 of 0.15g/kW.
- 2. MEA test of Pt/Au/Ni/C failed, attributable to inadequate synthesis.
- 3. Scale up of catalysts synthesis to 1g/batch accomplished.

4. Au cluster submonolayer has a pronounced stabilization effect on Pt under potential cycling regime. It is a highly promising solution to that problem.

5. The Pd_2Co and Pd_3Fe have activities similar to that of Pt; Methanol tolerance of Pd_2Co confirmed in a fuel cell test at LANL.

6. NbO₂ is a promising support for Pt monolayer catalysts.

Responses to Previous Year Reviewers' Comments

- Include repeated potential cycling tests as part of durability screening of promising candidates... --- The tests have been started.
- 2. Do small-scale development of the processes needed to make 50cm² MEA and, eventually, short-stack quantities of catalysts through the UPD-replacement process. ---The synthesis has been scaled up to make 1 g of the catalyst per day.
- 3. Should try to reduce the amount of Pd sooner in timeline (2 reviewers) -- Pt monolayer on noble metal -non-noble metal core-shell nanoparticles provide such a possibility.
- 4. A more appropriate industry collaboration would be with companies that make catalysts,--- BNL is open for collaboration.

Publications

- J. Zhang, F.H.B. Lima, M. H. Shao, K. Sasaki, J.X. Wang, J. Hanson, R. R. Adzic, Platinum monolayer on non-noble metal - noble metal core-shell nanoparticles electrocatalysts for O2 reduction, *J. Phys. Chem. B*, 109 (2005) 22701-22704.
- 2. M.H. Shao, P. Liu, R.R. Adzic, Superoxide is the intermediate in the oxygen reduction reaction on platinum electrode. *J. Am. Chem. Soc.* (submitted).
- 3. J. Zhang, M. B Vukmirovic, K. Sasaki, F. Uribe, R. R. Adzic, Platinum monolayer electrocatalysts for oxygen reduction: effect of substrates, and long-term stability, *J. Serb. Chem. Soc.*, 70 (2005) 513-525 (75th Anniversary issue).
- J. X. Wang, F. A. Uribe, and R. R. Adzic, Parameterizing H₂/air-PEMFC polarization curves, and quantifying cell performance and major voltage losses", *Electrochem. Solid-State Lett*. (submitted).
- 5. M.H. Shao, K. Sasaki, R.R. Adzic, Pd-Fe nanoparticles as electrocatalysts for oxygen reduction. *J. Am. Chem. Soc.* 128 (2006) 3536.

Presentations

Five papers at national and three at international meetings,

Critical Assumptions and Issues

- Stability of Au clusters on Pt surfaces
- Stability of surface segregation induced by annealing
- The extent of mixing in bimetallic systems at 80-100°C
- Electronic effects vs. strain effects