## Contiguous Platinum Monolayer Oxygen Reduction Electrocatalysts on High-Stability-Low-Cost Supports

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Project FC009





a passion for discovery





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## **Overview**

## Timeline

Project start date: July 2009 Project end date: September 2013 Percent complete: Approx. 40%

## **Barriers**

Performance: Catalyst activity; ≥ 0.44 A/mg<sub>PGM</sub> Cost: PGM loading; ≤ 0.3 mg PGM /cm<sup>2</sup> Durability: < 40% loss in activity under potential cycling

## Budget in \$K

Total project funding:3,529Funding in FY10:267Funding in FY 11:860

#### **Partners**

Massachusetts Institute of Technology (MIT) Johnson Matthey Fuel Cells (JMFC)

#### Collaborations

UTC Power; Toyota M. C.,3M Corporation; U. Wisconsin, U. Stony Brook, U. Santiago de Compostela, CFN-BNL

BNL coworkers: Yun Cai, Stoyan Bliznakov, Kuanping Gong, Kurian Kuttiyiel



## Relevance

#### **Objectives:**

1. Developing high performance fuel cell electrocatalysts for the oxygen reduction reaction (ORR) comprising contiguous Pt monolayer (ML) on stable, inexpensive metal or alloy:

- nanoparticles (NPs)
- nanowires (NWs), nanorods (NRs),
- hollow NPs
- carbon nanotubes (CNT),
- 2. Increasing stability of cores and supports
- 3. Scale-up syntheses of selected catalysts,
- 4. MEA-testing, stack- testing





## Approach

Our approach is based on improved understanding of factors affecting the ORR from the studies of:

- 1. Oxygen binding energy
- 2. Surface contraction effects (size-induced, hollow NPs),
- 3. Surface segregation effects;
- 4. Core-shell structures 2<sup>nd</sup> generation (interlayer; core sub-surface layer)
- 5. Computations and modeling



- Reduced Oxygen binding energy
- Low number of low-coordination atoms
- Compressed top layer of atoms
- Moderately compressed (111) facets

Cores have a key role in achieving these requirements

Additional improvements with Pt as a contiguous monolayer

\*J.X. Wang, H. Inada, L.Wu, Y. Zhu, Y. Choi, P. Liu, W.P. Zhou, R.R. Adzic, J. Am. Chem. Soc, 131 (2009) 17298, JACS Select #8



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4

#### Synthesis of Pd nanowires and nanorods by electrodeposition on carbon



Deposition of Pd on carbon surfaces in 0.1M  $HClO_4$  with 1mM Pd<sup>2+</sup>.

The growth mechanism: Hupd in Pd acts as reducing promotor at terraces, while chlorides adsorb at lowcoordination sites and block growth in that direction. The type of deposit: NPs, NWs or NRs, depend on the potential and Pd ion concentration



FFT of the TEM image showing (111) pattern of Pd(111)

Scale-up: Cell for  $5 \text{ cm}^2$  electrode was constructed. It appears that a uniform deposition of NWs at electrodes of any practical surface area will be possible. Thus, this synthesis will be compatible with MEA fabrication.





High activity and excellent stability due to: (i) the smooth surface; (ii) the reduced number of defects (iii) low number of low-coordinated sites.



Pt/PdAu Mass activity: **3.5** A/mg<sub>Pt</sub> Specific activity: **1.5mAcm<sup>-2</sup>** 





Mass activity: **1.07** A/mg<sub>Pt</sub> Spec activity: **0.45mAcm<sup>-2</sup>** 

Additional Pd NWs from surfactant-based synthesis have similar activities ( with S, Wong, C. Koenigsmann) 6

1-Dimensional Interconnected Pt NPs on Amine-Functionalized MWCNTs for the ORR



(a) Polyol synthesis of 1-D Pt NPs/MWNTs, TEM image of (b) as prepared 1-D sample (c) after 6 M HCl treated (b) as prepared Pt NPs/MWNTs and (d) after heat at 220 °C.

Junhyung Kim, Seung Woo Lee, Christopher Carlton and Yang Shao-Horn, MIT



#### **1-D Interconnected Pt Nanoparticles on Amine-Functionalized MWCNTs**



Intrinsic ORR activity comparison. (a) CVs from 4 samples, (b) ORR at RDE at 1600 rpm ; 10 mV/s, (c) specific activities at 0.9 V, iR corrected and (d) specific activity of TKK and heat treated Pt NPs/MWNTs.



#### Syntheses of Pt wires, tests of TiC and WC supports



wire / solvent

suspension

Pt wires synthesized and transferred to CNPs successively

- further optimization required
- Pt was blocked by strong surfactant

New supporting materials developed:

1.TiC2.WC3.Pt wires on cores to thrift PGM

Initial tests are being conducted



40w% Pt (5nm) on Commercial TiC (100nm)





Rachel O'Malley, Laura Ashfield, Geoff Spikes



#### **Pt monolayer on hollow Pd nanoparticles**



TEM images of pulse-deposited Ni nanoparticles



(a), Pt(ML)-Pd<sub>20</sub>Au hollow particles fabricated using Ni nanoparticles as templates (b).





High activity is due to smooth surface morphology, and hollowinduced lattice contraction.

Scale-up synthesis is being developed using:

- The cell for electrodeposition 1. of Pd NWs
- 2. The microemulsion method.



#### Pt monolayer on Pd tetrahedral nanoparticles: Synthesis



Tetrahedral (TH) Pd NPs were prepared using a hydrothermal route using PVP as surfactant. PVP was removed from the NPs by CO adsorption-induced desorption.

# CO adsorption-induced surfactants desorption

A versatile, noninvasive method for removing surfactants - from NPs surfaces



#### Technical Accomplishments and Progress Pt monolayer on Pd tetrahedral nanoparticles



## The weaker BE-O of TH $Pt_{ML}Pd$ may cause its higher activity than that of $Pt_{ML}Pd$ .

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TH consists of much smaller number of atoms than a sphere-like (SP) model  $\rightarrow$  Lower Pd, Pt loading

	Pt <sub>ML</sub> /TH Pd/C
Half-wave potential (V)	891
ECSA (m <sup>2</sup> /mg <sub>Pt</sub> )	19
Pt specific activity (mA/cm <sup>2</sup> <sub>Pt</sub> )	0.64
Pt mass activity (A/mg <sub>Pt</sub> )	0.92
PGM mass act. (A/mg <sub>Pt, Pd</sub> )	0.14
12	BROOKH

**Technical Accomplishments and Progress** Improving cores by alloying: Pt/IrNi/C



**HAADF-STEM** image of a PtIrNi nanoparticle



**EELS intensities for the Pt and Ir M-edge** and Ni L-edge along the scanned line

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Thermal treatment induces

Ir segregation to surfaces



**Exceptional stability of cores alloys of Ir** with non-noble metals: e.g. IrNi, IrFe



**Pt Mass Activity:** 1.4 A/mg<sub>Pt</sub> **Spec. Activity:** 0.60 mA/cm<sup>2</sup> **PGM Mass Activity:** 0.78 A/mg<sub>Pt+lr</sub>

#### Improving cores by alloying: Pt/IrNi/C



DFT: Adding Ni to Ir makes BE-O weaker as a function of strain  $\rightarrow$  Higher activity



EELS intensities after 50,000 cycles; Ni is clearly observed



Fuel cell test of Pt<sub>ML</sub>/Pd<sub>9</sub>Au/C (a) and its improved version (b) Stability tests: potential cycling (0.6 -1.0 V; 50 mV/s,



SEM image and EDS line analysis of Pt/Pd<sub>9</sub>Au after 200K (a) and 100K cycles (b).

Pd "band" forms from dissolved Pd (a) sample, but not for a more stable Pd<sub>9</sub>Au<sub>1</sub> (b).

No Pt dissolution is observed

#### MEA tests at Toyota MC

Brookhaven Science Associates Sasaki et al. Angew. Chem. Int. Ed., 49 (2010) 8602.

#### New mechanism of stability of core-shell electrocatalysts: shell protected by the core

- 1. PtOH formation shifted positively.
- 2. Contraction of Pt and Pd lattices induced by some loss of Pd (hollow may form).
- 3. Cathodic protection effect.



Data supported by EXAFS, XANES, EELS, EDS, RDE, DFT results.



The structure of Pt shells was almost retained after the tests



Fuel cell performance stability test of Pt<sub>ML</sub>/Pd<sub>9</sub>Au/C and Pt<sub>ML</sub>/Pd/C electrocatalysts No change in performance in 200,000 potential cycles with Pt<sub>ML</sub>/Pd/C



TEM images and EDS line analysis of Pt/PdAu/C catalyst, before (a) and after (b) cycling test number of potential cycles



## **Collaborations**

- **1. Massachusetts Institute of Technology (MIT)** (University) **Yang Shao-Horn, Co-PI of the project**
- 2. Johnson Matthey Fuel Cells (JMFC) (Industry) Rachel O'Malley, Co-PI of the project, Sarah Ball, Graham Hard
- **3. Toyota Motor Company Hideo Naohara,** MEA test, catalysts scale-up
- 4. UTC Power (Industry) Collaboration on MEAs making, stack building and testing.
- 5. U. Wisconsin (University)Manos Mavrikakis, collaboration on theoretical calculations-
- 6. Center for Functional Nanomaterials, BNL Ping Liu, YongMan Choi, DFT calculations; Eli Sutter ad Yimei Zhu, TEM, STEM
- 7. 3M Corporation (Industry) Radoslav Atanasoski, Andrew Haug, Greg Haugen
- 8. University Santiago de Compostela, Spain David Bucheta, microemulsions-based syntheses of core-shell nanoparticles

17

## **Proposed Future Work**

#### FY11

- 1. Scale-up synthesis of Pd nanowires by electrodeposition at electrodes of 5 cm<sup>2</sup> and 25 cm<sup>2</sup>. Further study of the synthesis involving chemical route with surfactants. (BNL)
- 2. Further studies of hollow Pd and PdAu nanoparticles with a scale-up using:
- i) Electrodeposited Ni templates that are galvanically displaced by Pd
- ii) Developing the microemulsion method to synthesize hollow Pd nanoparticles. (BNL, JMFC, MIT)
  Scale-up of selected synthesis to produce 20 grams of the catalyst (JMFC), or the electrodes of 25 cm<sup>2</sup> compatible with MEA manufacturing.

#### FY12

- 1. Improving synthesis of Pd tetrahedra, scale-up to 5 grams
- 2. Improving synthesis of Pt/Ir-Ni/C ; Pt/Pd/Ru/C. (BNL, MIT, JMFC)
- 3. Improve metallization and catalyzation of CNTs, oxides, nitrides. (JMFC, MIT, BNL)
- 4. Further the work on Pd-W, Pd-N and Pd-V NPs. (BNL, MIT)
- 5. MEA fabrication and tests. Go/No go for these catalysts based on MEA tests. (JMFC, UTC)



## **Summary**

Pt monolayer electrocatalysts supported on several types of cores, which can have several shapes, are ready for scale-up syntheses and MEA testing.

Nanowires, nanorods, hollow NPs and tetrahedral NPs provide excellent supports-cores.

Pd, Pd alloys, Ir - non-noble metal alloys provide excellent cores.

Using refractory metal alloys in cores and metallization of CNTs require further studies.

Sub-surface ML modification of cores opens up possibilities for additional improvements.

New mechanism of stability of core-shell electrocatalysts, in which shell is protected by the core, has been proposed.

Pt<sub>ML</sub>/Pd<sub>9</sub>Au/C and Pt<sub>ML</sub>/Pd/C are practical electrocatalysts. Only 10 grams of Pt, and about 15-20 grams of Pd is needed for a FC of 100 KW. Currently, catalytic converters use 5-10 g of Pt per vehicle. No new Pt is needed for converting to fuel cells.

