V.D.3 Rationally Designed Catalyst Layers for PEMFC Performance Optimization

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Overall Objectives

- Realize the oxygen reduction reaction (ORR) mass activity benefits of advanced Pt-based cathode electrocatalysts in membrane electrode assemblies (MEAs) and stacks operating at high current densities and on air and at low precious group metal (PGM) loading (≤0.1 mgPt/cm² on cathode).
- Determine the source(s) of performance limitations of de-alloyed PtNi (d-PtNi)-containing MEAs at high current densities (>1,000 mA/cm²) when operating on air.
- Design and develop an electrode layer composition and/or structure, based on in-cell diagnostics, advanced characterization, and performance modeling to exceed the technical targets for MEAs for transportation applications.

Fiscal Year (FY) 2016 Objectives

- Determine effect of the equivalent weight (EW) of the ionomer in the cathode catalyst layer (CCL) on cell performance and performance durability.
- Diagnose source of performance loss with Ni²⁺ contamination in CCL utilizing X-ray scattering, water uptake measurements, and doping Ni²⁺ into Pt-based CCLs.
- Diagnose source(s) of activity losses of d-PtNi with functionalized support.
- Calculate porosities and diffusivities of Pt/C and d-PtNi/C electrodes utilizing X-ray tomography and transmission electron microscopy images, image analysis, and image reconstructions.
- Achieve a hydrogen-air current density of 1,250 mA/cm² at 0.675 V, under differential cell conditions (high stoichiometries), using cells with 0.125 mg-Pt/cm² fabricated using the best compositions and preparation procedures developed throughout the project.
- Determine the durability of cells with d-PtNi cathode catalysts and total cell Pt loadings of <0.125 mg-Pt/cm² using DOE-recommended protocols and alternative protocols. Performance and durability goals are 6.5 kW/g-Pt at peak power and <10% drop in power after application of the U.S. DRIVE Fuel Cell Tech Team Cell (FCTT) component accelerated stress tests (ASTs).

Technical Barriers

This project addresses the following technical barriers from the Fuel Cell section of the Fuel Cell Technologies Program Multi-Year Research, Development and Demonstration Plan.
(C) Performance
(B) Cost
(A) Durability

Technical Targets

The technical targets for this project are listed in Table 1.

FY 2016 Accomplishments

- Achieved a hydrogen-air current density of 1,259 mA/cm² at 0.675 V with cathode loading of 0.112 mg-Pt/cm² and under differential conditions. Achieved 1,060 mA/cm² at 0.675 V with a total cell...
loading of 0.1107 mg-Pt/cm² when using a modified FCTT protocol with higher cell operating temperature, allowing cell to meet Q/ΔT targets. (Achieved 890 mA/cm² at 0.675V using standard FCTT protocol).

- MEA with d-PtNi/C cathode at a loading of 0.1159 mg-Pt/cm² and a Pt/C anode at a loading of 0.018 mg-Pt/cm² achieved 7.7 kW/g-Pt at peak power under differential conditions and 6.2 kW/g-Pt under standard DOE FCTT-recommended polarization curve protocol.

- Achieved a 7% loss in peak power after 30,000 triangle cycles (0.6–0.925 V, 50 mV/s) and 22% loss in peak power after 30,000 trapezoid cycles (0.6–0.95 V, 700 mV/s, 6 s/cycle).

### INTRODUCTION

One of the major contributors to the cost of PEMFC systems for automotive and stationary power applications is the PGM cathode electrocatalyst [1]. The high cost of the cathode electrocatalyst results from the high loadings of catalyst necessary to overcome the limitations of low ORR activity, low utilization of PGM, and loss of activity with operating time. Alloying platinum with base metals (e.g., cobalt, iron, and nickel) is well known to improve its intrinsic ORR activity [2]. While ORR mass activities exceeding the DOE 2020 targets (>0.44 A/mg PGM and 720 µA/cm² at 900 mV) have been demonstrated for numerous Pt alloy and core-shell nanoparticle catalysts [3], the full high current density performance of CCLs based on low loadings of these promising catalysts has yet to be achieved in MEAs. There are several possible reasons the full potentials of these advanced catalysts have not been realized in MEAs, such as inadequate transport of protons and oxygen to the catalytic sites arising from less than optimum spatial distribution of ionomer, catalyst, and pores. The goal of this project is to optimize the composition, structure, and materials properties of cathodes based on an advanced cathode catalyst to translate the catalyst’s intrinsically high ORR mass activity into high performance at peak power. The advanced catalyst chosen for this project is dealloyed PtNi (d-PtNi) developed by Johnson Matthey Fuel Cells within a DOE-supported General Motors project [3].

### APPROACH

The overall approach of the project is to:

- Determine the properties of advanced alloy-based catalysts and/or cathode catalyst layers that limit the high current density/air performance using:
  - *In-cell* diagnostics of d-PtNi/C versus high surface area Pt/C and Pt/C of comparable electrochemically active surface area.
  - A suite of in situ and ex situ techniques, such as transmission electron microscopy, cryogenic transmission electron microscopy, dynamic light scattering, ultra-small angle X-ray scattering, X-ray absorption spectroscopy, X-ray tomography, and porosimetry.

- Design the catalyst layer composition and structure and support functionality to mitigate the performance limitations, guided by computational modeling, by:
  - Studying the dispersion of d-PtNi/C catalyst aggregates and the ionomer particles in liquid media and in electrodes and compare them to Pt/C-based inks and electrodes.
  - Developing an ink composition and/or ink processing (e.g., solvent removal process) that result in optimum agglomerate structure in d-PtNi/C-based electrodes.
Developing the catalyst support surface functionality to increase the performance of the catalyst and cathode – to decouple proton conductivity from ionomer content.

The d-PtNi catalysts were synthesized by Johnson Matthey Fuel Cells using a catalyst precursor comprised of 5.4 nm mean diameter PtNi, alloy particles deposited on Ketjen black carbon, via a commercially-scalable method, followed by annealing to drive alloy formation. The resulting catalyst precursor is dealloyed via an acid treatment step to leach Ni to form a catalyst with an approximate composition of Pt$_3$Ni$_2$, and a Pt loading on the carbon support of approximately 30 wt%. As shown in Table 1, the ORR mass activity of this catalyst in an MEA far exceeds the DOE 2020 target.

**RESULTS**

The effects of ionomer EW on the agglomerate structure in catalyst-ionomer-solvent inks, agglomerate structure in electrodes, and CCL performance and performance durability under a variety of oxygen partial pressures and relative humidities (RHs) were determined for the d-PtNi/C catalyst and, for comparison, for a Pt/C catalyst with a particle size distribution comparable to that of the d-PtNi/C (An-Pt/C). The following summarizes the results of these characterization and performance studies.

- Lower EW ionomer in inks:
  - In general causes greater break-up of agglomerate structure than the standard 1100 EW ionomer, effect is dependent on I/C (Figure 1)
  - Best overall cell performance under high and low RHs observed for lowest EW ionomer and intermediate I/C (1.0) (Figure 2)

- Functionalization of carbon support with –SO$_3$H groups improves low RH performance, but decreases catalyst mass activity.
  - Lower mass activity due to adsorption of sulfur species on Pt surface and to loss of both Pt and Ni from catalyst during functionalization

- Performance Status (cells with ~0.1 mg-Pt/cm$^2$ anode loading):
  - Differential conditions, organic ink, I/C = 0.8, acid-washed CCM: 1,300 mA/cm$^2$ at 0.675 V

- Performance Status (cells with ~0.025 mg-Pt/cm$^2$ anode loading):
  - Differential conditions, organic ink, I/C = 1.0, 850 EW, acid-washed CCM: 1,259 mA/cm$^2$ at 0.675 V

- FCTT polarization curve:
  - 1,060 mA/cm$^2$ at 0.675 V with a total cell loading of 0.1107 mg-Pt/cm$^2$ at 90°C
  - 890 mA/cm$^2$ at 0.675 V under standard FCTT conditions (80°C)

- Durability status after 30,000 AST cycles (cells with ~0.025 mg-Pt/cm$^2$ anode loading) (Figure 3)
  - Mass activity loss: 34%; electrochemical surface area (ECA) loss: 49%, (0.6–1.0 V, 50 mV/s, triangle, limited diagnostics)
  - Mass activity loss: 26%; ECA loss: 26%; Voltage decrease at 1.5 A/cm$^2$: 33 mV (0.6–0.925 V, 50 mV/s, triangle, limited diagnostics)
  - Mass activity loss: 46%; ECA loss: 51%; Voltage decrease at 1.5 A/cm$^2$: 121 mV (0.6–0.95 V, 700 mV/s trapezoid, extensive diagnostics)
CONCLUSIONS AND FUTURE DIRECTIONS

Issues with the d-PtNi/C catalyst-based electrodes are drop-off in performance at >1,000 mA/cm² (i.e., not maintaining mass activity benefit relative to high-surface area Pt/C) under high and intermediate humidity conditions and a severe drop-off at low RHs. The sources of these issues and potential solutions to the issues are:

- Mass transport to low loadings of large particles.
  - Can be mitigated with smaller particles that are >~4 nm to insure stability against dissolution-related ECA loss.
- Sensitivity to low humidity conditions is unique to d-PtNi/C and can be partially mitigated with increased ionomer content, organic solvents, and lower EW ionomer which improve dispersion of d-PtNi/C particles, and can be greatly improved with acid treatment of CCM.

Results indicate that Ni²⁺ in ionomer, leached during ink and CCM fabrication is altering the structure of the ionomer, which alters the agglomerate structure of cathode catalyst layer, decreasing mass transport, especially at low RH.

- More aggressive leaching of Ni²⁺ prior to ink synthesis and CCM fabrication can improve d-PtNi/C cathode performance, especially under dry conditions.

Leaching of Ni from particles continues during ASTs, especially when upper potential of the AST is 1.0 V

- Mitigate through control of highest voltage encountered by cathode.
- Continue development of catalyst particles with more uniform morphology, intra-particle composition distribution, more uniform Pt shell.

d-PtNi/C shows higher ECA, mass activity, and high current density performance loss than An-Pt/C when subjected to catalyst ASTs.

Acid-washing d-PtNi/C-containing CCMs does not adversely affect ECA and mass activity decay with AST cycling and decreases low and high current density performance decay.

The decay rates with AST cycling were comparable for d-PtNi/C-containing electrodes containing either 850 or 1100 EW ionomer.
• Intra-AST-cycling diagnostics, such as CO stripping for ECA measurement, dramatically increases ECA, mass activity, and high and low current density performance loss of the d-PtNi/C-based CCL (Figure 3).

• New trapezoid AST causes more degradation of ECA, mass activity, and air performance than DOE 0.6–1.0 V, 50 mV/s triangle AST and General Motors-suggested 0.6–0.925 V triangle AST (Figure 3).

• d-PtNi/C can meet ECA and mass activity durability targets and approaches high current density performance loss durability target (30 mV) when limiting upper potential of triangle AST to 0.925 V (Figure 3).

FY 2016 PUBLICATIONS/PRESENTATIONS


REFERENCES

