



University at Buffalo
The State University of New York



2018 DOE H₂ and Fuel Cell Annual Merit Review Meeting

Advanced Catalysts and MEAs for Reversible Alkaline Membrane Fuel Cells

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Newton, MA

June 12, 2018

Project #
FC 129

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Project Overview

Timeline

- Project Start Date: June 1, 2015
- Project End Date: June 30, 2018

Budget

- Total \$1,200,496
 - DOE share \$959,334
 - Contractors share \$241,162
- Spent \$1, 182,565 (by April. 2018)

Principal Researchers

Shuai Zhao and Tom McCallum

Subcontractors

- SUNY-Buffalo: Prof. Gang Wu and Shiva Gupta
- NREL: Drs. Bryan Pivovar and Shaun Alia, Andrew Park

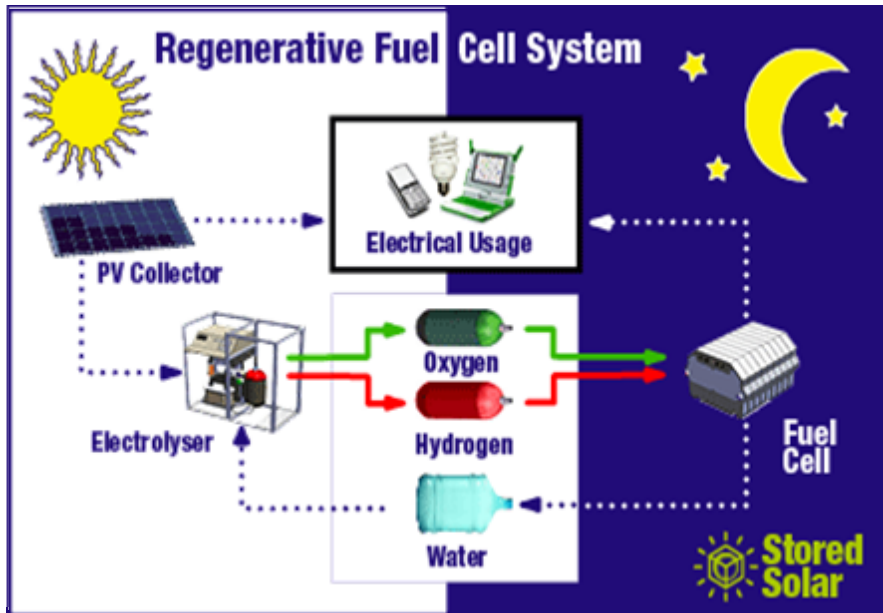
Barriers Addressed

- Activity (catalyst; MEA)
- Durability (catalyst; MEA)
- Cost (catalyst; MEA)

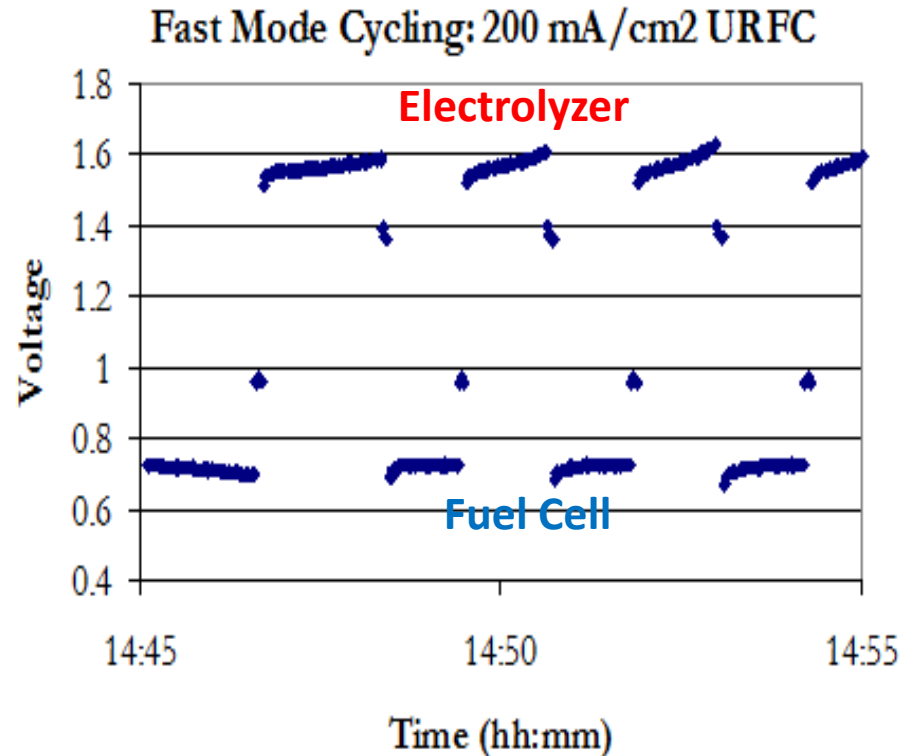
Technical Targets

- Design and develop ORR/OER bi-functional oxide catalysts
- Integrate ORR/OER bifunctional oxide catalysts and alkaline membranes to develop highly efficient reversible alkaline membrane fuel cells (AMFCs) for stationary energy storage

Reversible Fuel Cells

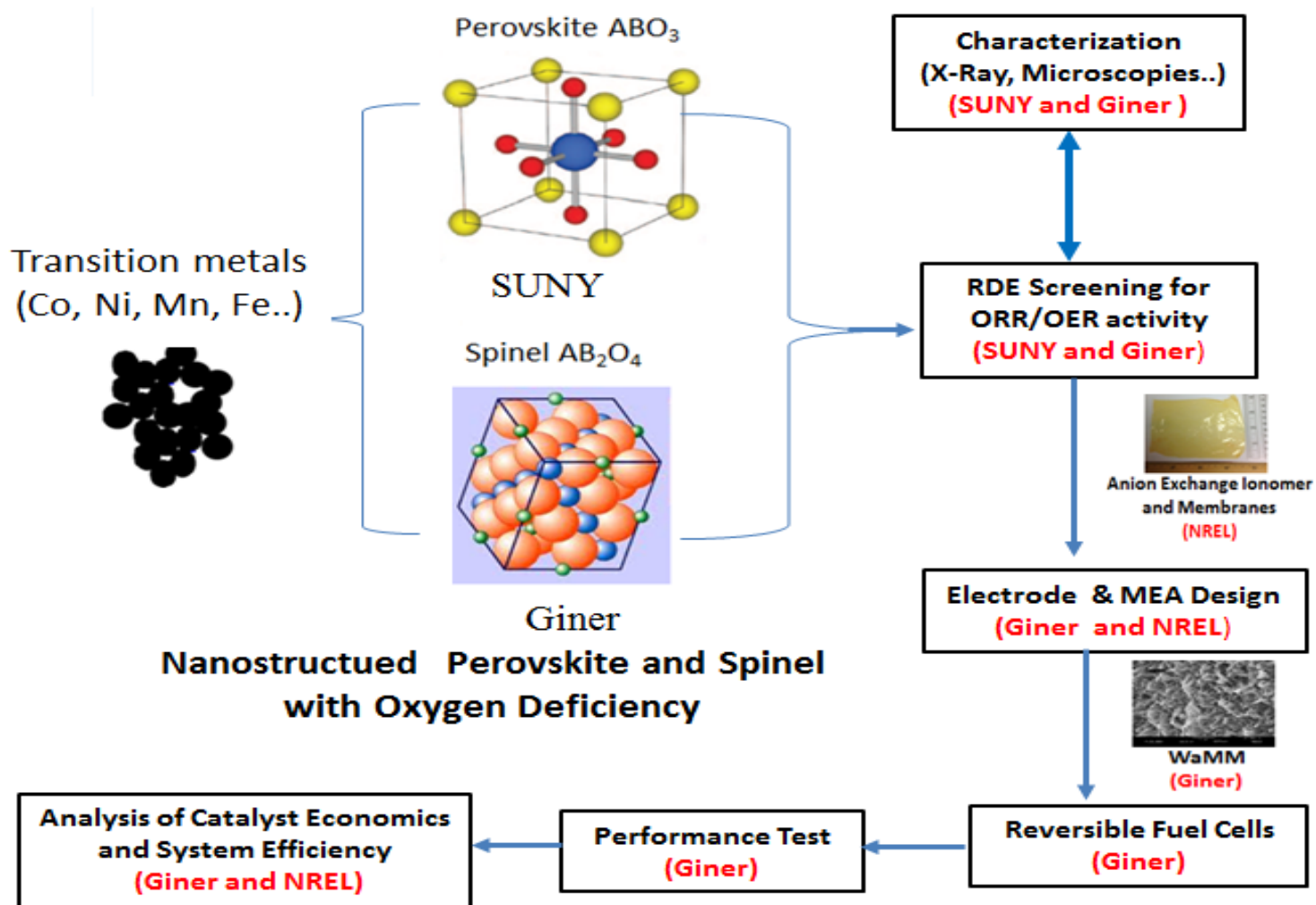


Giner unitized reversible PEM fuel cell



- Water electrolyzer is an ideal device to store energy from wind mills and solar farms, where surplus (off peak) energy is nearly free
- Stored H₂ can be used for fuel cells to generate electricity in peak time

Technical Approaches



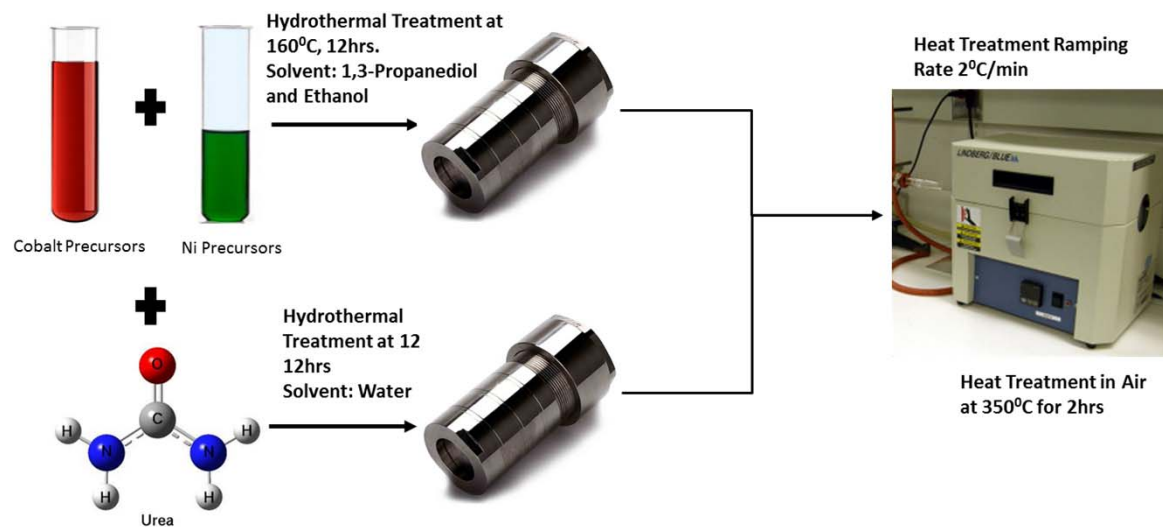
- Catalyst Long-term Stability;
- MEA Fabrication Technology

Technical Milestones

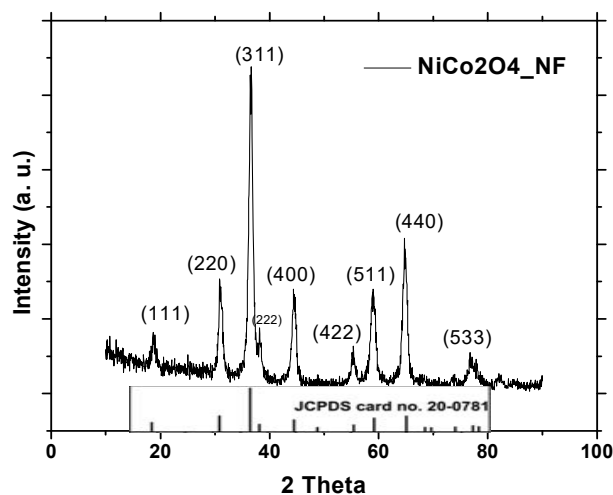
Time	Milestone Description	Completion
Q1	Synthesize BaTiO _{3-x} perovskites with 3 different oxygen vacancy concentrations	100%
Q2	Prepare 3 other oxygen-deficient AA'BB'O _{3-x} multiple perovskite catalysts (e.g., BaSrCoFeO _{3-x} or BaSrMnCrO _{3-x}) with optimized defect structures	100%
Q3	Reduce perovskite particle size to nanoscale (<10 nm) with much increased surface areas (>20 m ² /g)	100%
Q3	Prepare 3 A _x B _{1-x} C ₂ O ₄ spinel catalysts (A, B and C represent Co, Mn, Fe or other Metals) with particle size <10nm	100%
Q4 (go/no-go point)	In RDE, demonstrate ORR activity > 1 mA/mg oxide at IR-free 0.9 V; and OER activity > 15 mA/mg oxide at IR free 1.6 V.	100%
Q4	Provide 20g of PF AEM material in membrane/ ionomer form Membrane conductivity >0.05 S/cm at 60°C and 100% RH; H ₂ permeability: 10 ⁻¹² mol/(kPa.s.cm)	100%
Q5	3 AEI ionomer categories and 5 ionomer loadings will be evaluated to identify the best electrode composition	100%
Q6	Achieve RFC performance 0.55V for fuel cell and 2.0V for electrolyzer, both at 600mA/cm ²	100%
Q7	Achieve fuel cell and electrolyzer life of 500 hours with less than 10% performance decay	60%
Q8	Generate a full report of catalyst and reversible fuel cell economics	50%

Accomplishment 1: Carbon-free Metal Oxide Catalysts (UB)

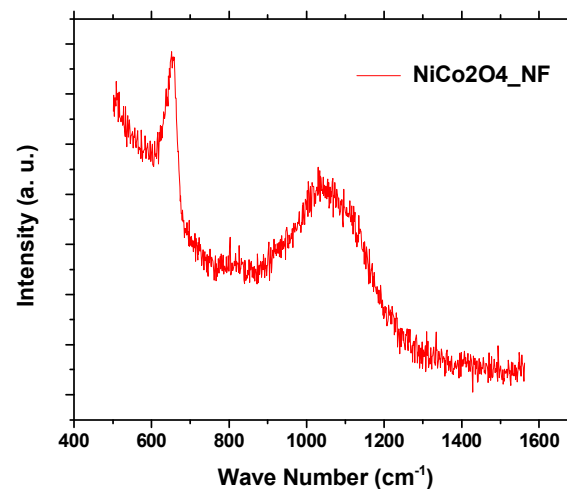
Low-cost and scalable hydrothermal synthesis



XRD Patterns

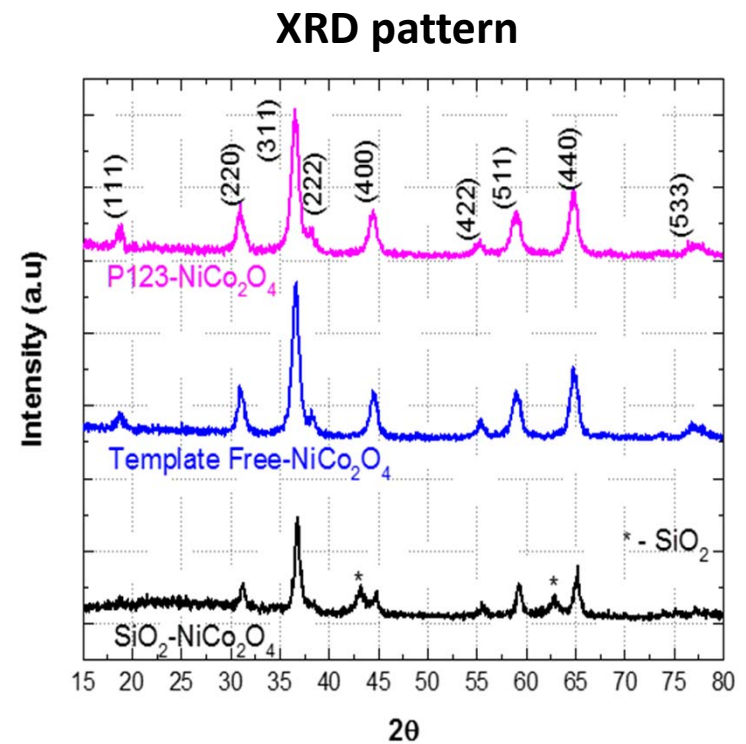
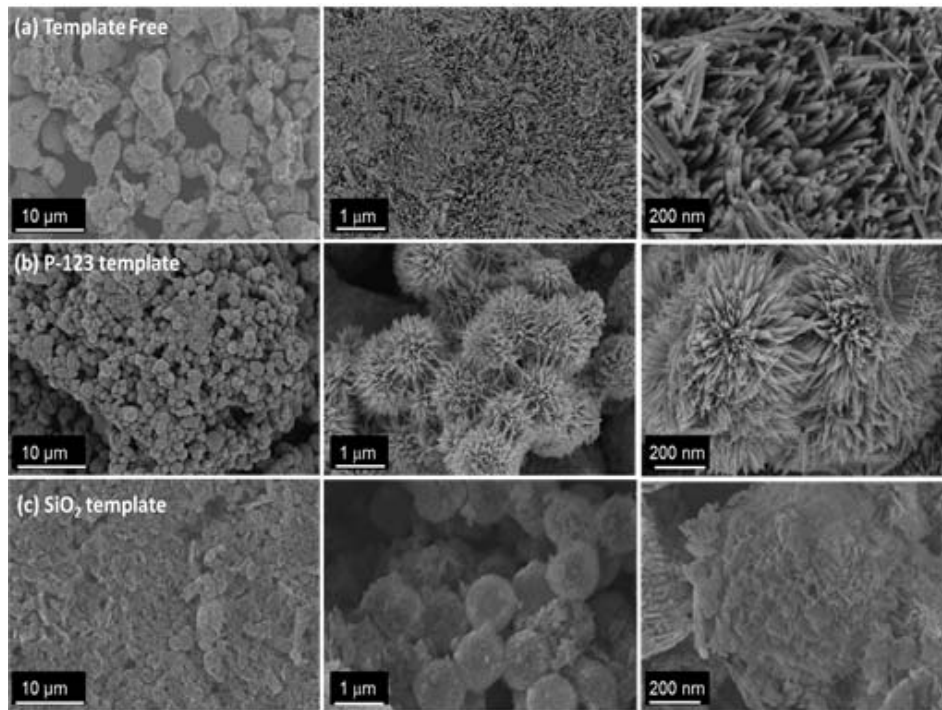


Raman spectroscopy



The XRD and Raman confirmed NiCo₂O₄ phase and peaks for Nickel-Cobalt Oxide are well in agreement confirmation the formation of NiCo₂O₄ phase.

Accomplishment 2: Carbon-free Metal Oxide Catalysts (UB)

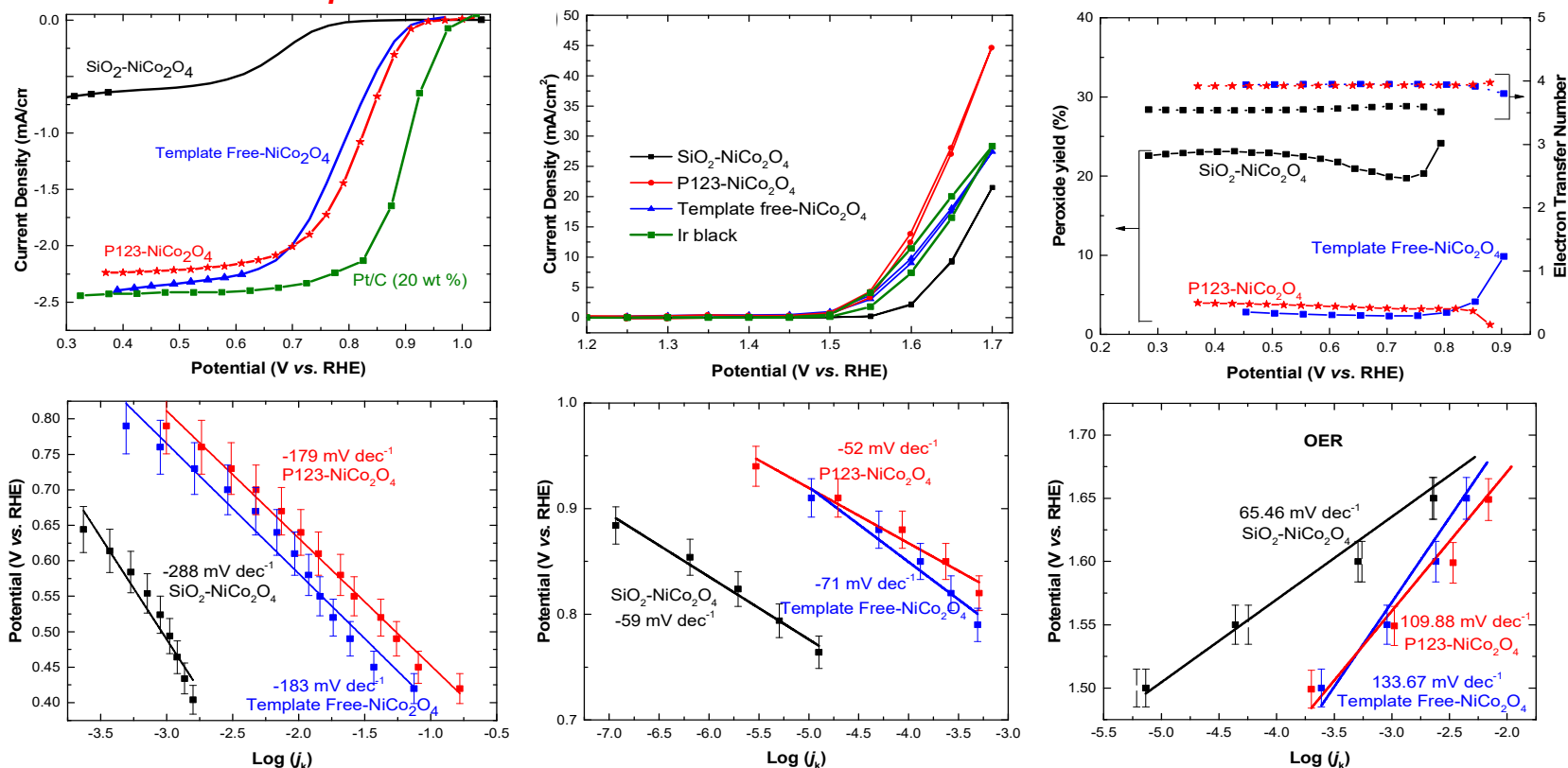


- ❑ Different morphologies of NiCo₂O₄ catalysts were achieved with different synthesis templates: (a) template-free, (b) P-123 soft template, and (c) SiO₂ hard template.
- ❑ The XRD pattern for template free and soft template assisted NiCo₂O₄ were well in agreement with the spinel-like NiCo₂O₄.

RDE Activity and Mechanistic Evaluation of NiCo_2O_4 Catalysts

Comparison between different morphologies

Steady state polarization plots for the ORR and the OER of various NiCo_2O_4 catalysts recorded in O_2 saturated 1.0 M NaOH at 900 and 1600rpm

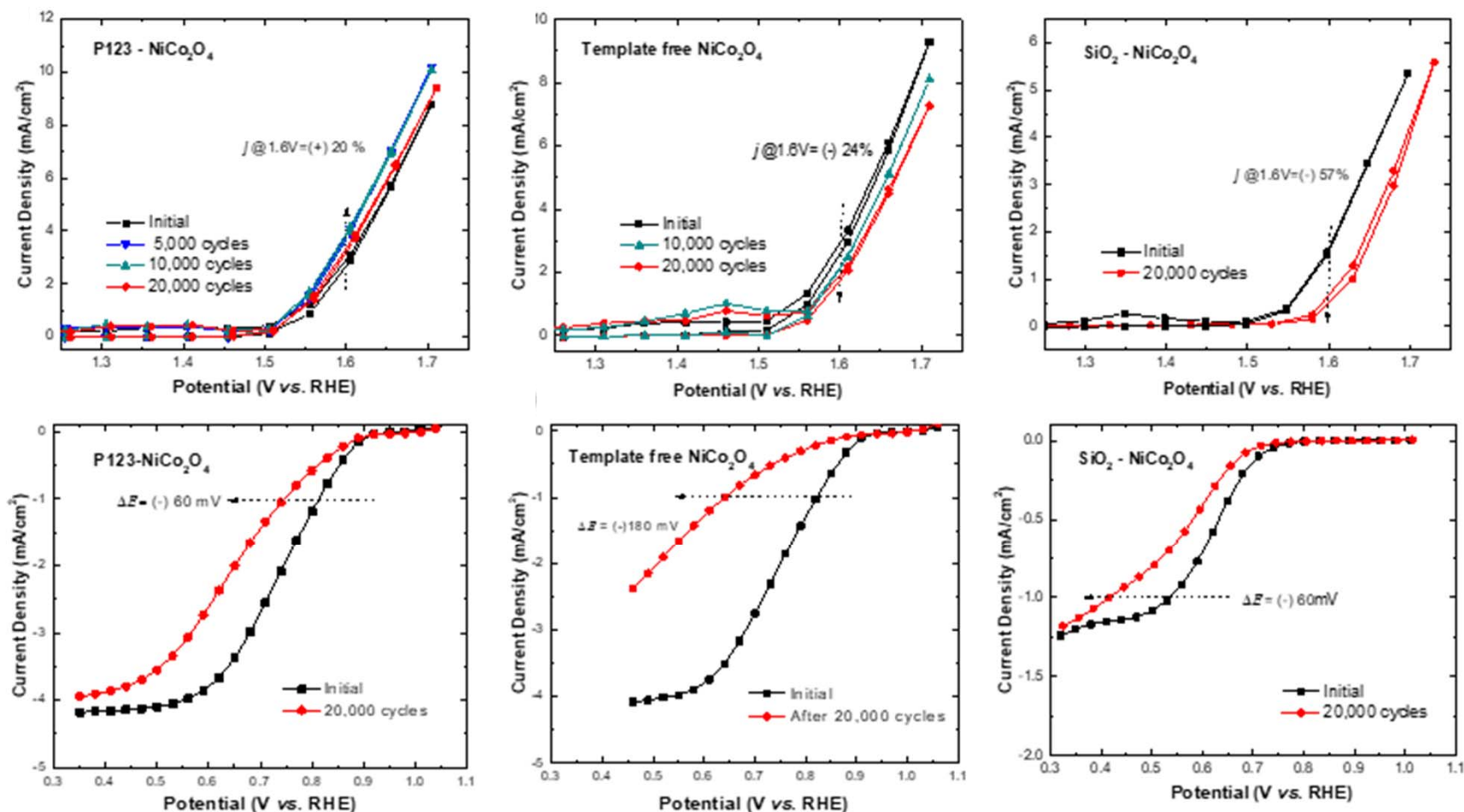


- ❑ The best activity was observed with the soft-templated NiCo_2O_4 catalyst, achieving onset (E_{onset}) and half-wave potentials ($E_{1/2}$) of 0.94 V and 0.82 V, representing one of best oxide ORR catalysts
- ❑ The formation of peroxide HO_2^- was monitored using ring currents measured from RRDE tests for selectivity study
- ❑ Tafel slopes for the ORR and OER were further determined to elucidate the overall reaction mechanisms.

RDE Stability Evaluation of NiCo₂O₄ Catalysts

Comparison between different morphologies

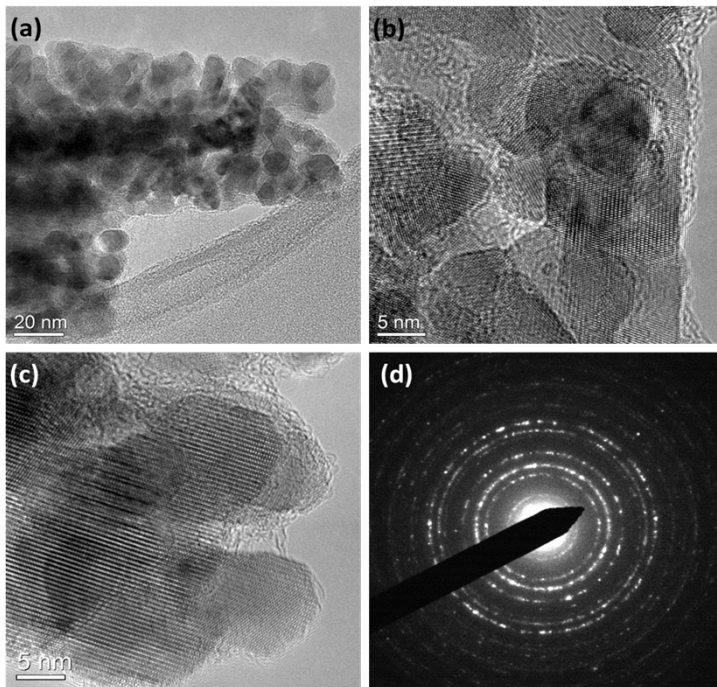
0.0 to 1.9 V in O₂ saturated 0.1 M NaOH, 25°C, 500 mV/s



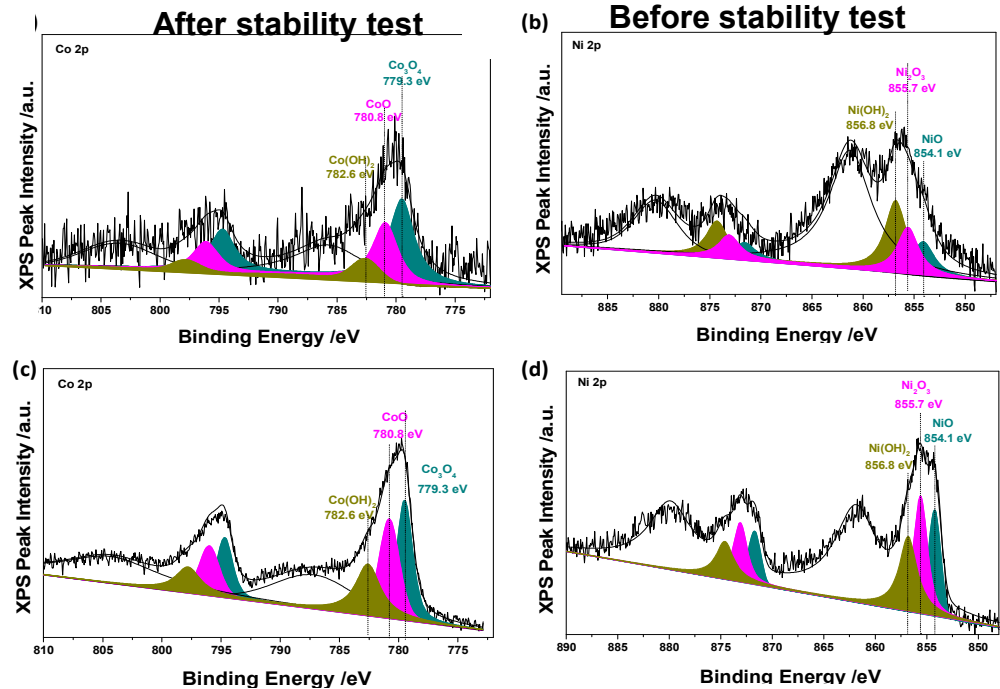
- ❑ Best performing soft-templated NiCo₂O₄ was also highly stable up to 20,000 cycles even with a 20% increase in current density generated at 1.6 V
- ❑ Much larger ORR degradation was observed with template-free NiCo₂O₄ catalyst with a loss of 220 mV in E_{1/2}

Mechanistic Evaluation of the Stability Enhancement

HR-TEM images

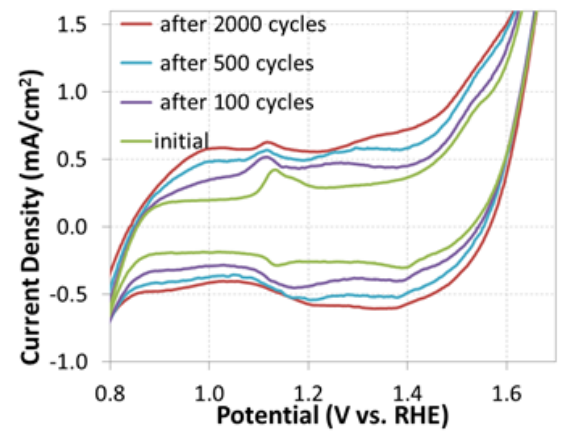
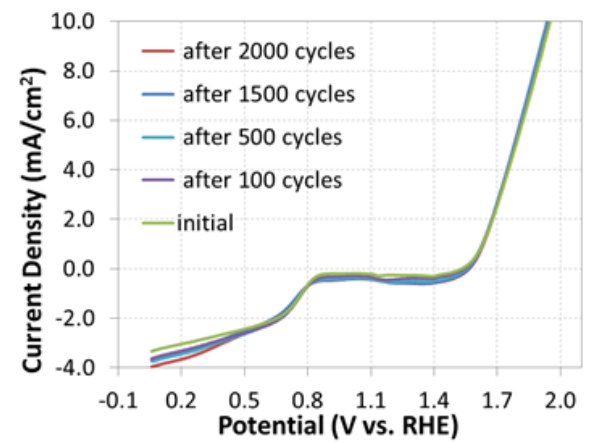
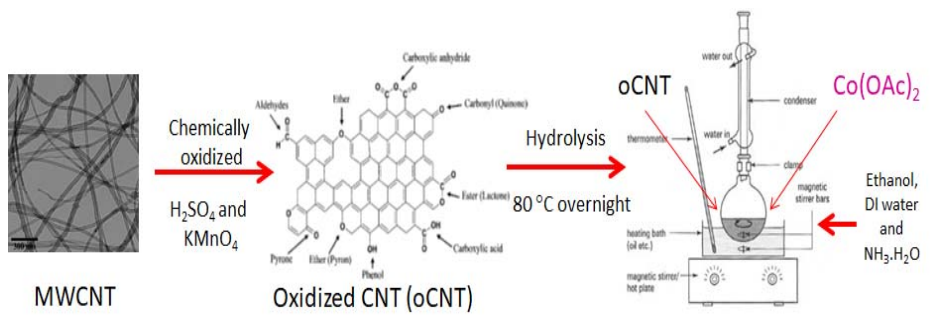


XPS Spectra



- ❑ The crystallinity of NiCo₂O₄ particles was well retained after 20,000 potential cycles from 0-1.9 V.
- ❑ After 20,000 potential cycles, Co species and their ratio at the catalyst surface remained nearly the same, the content of CoO was slightly increased after cycling, which could promote the OER.
- ❑ Ni species was not stable during the AST. Ni(OH)₂ became dominant over Ni₂O₃ and NiO.
- ❑ The formation of additional CoO and Ni(OH)₂ was likely responsible for the ORR activity loss.

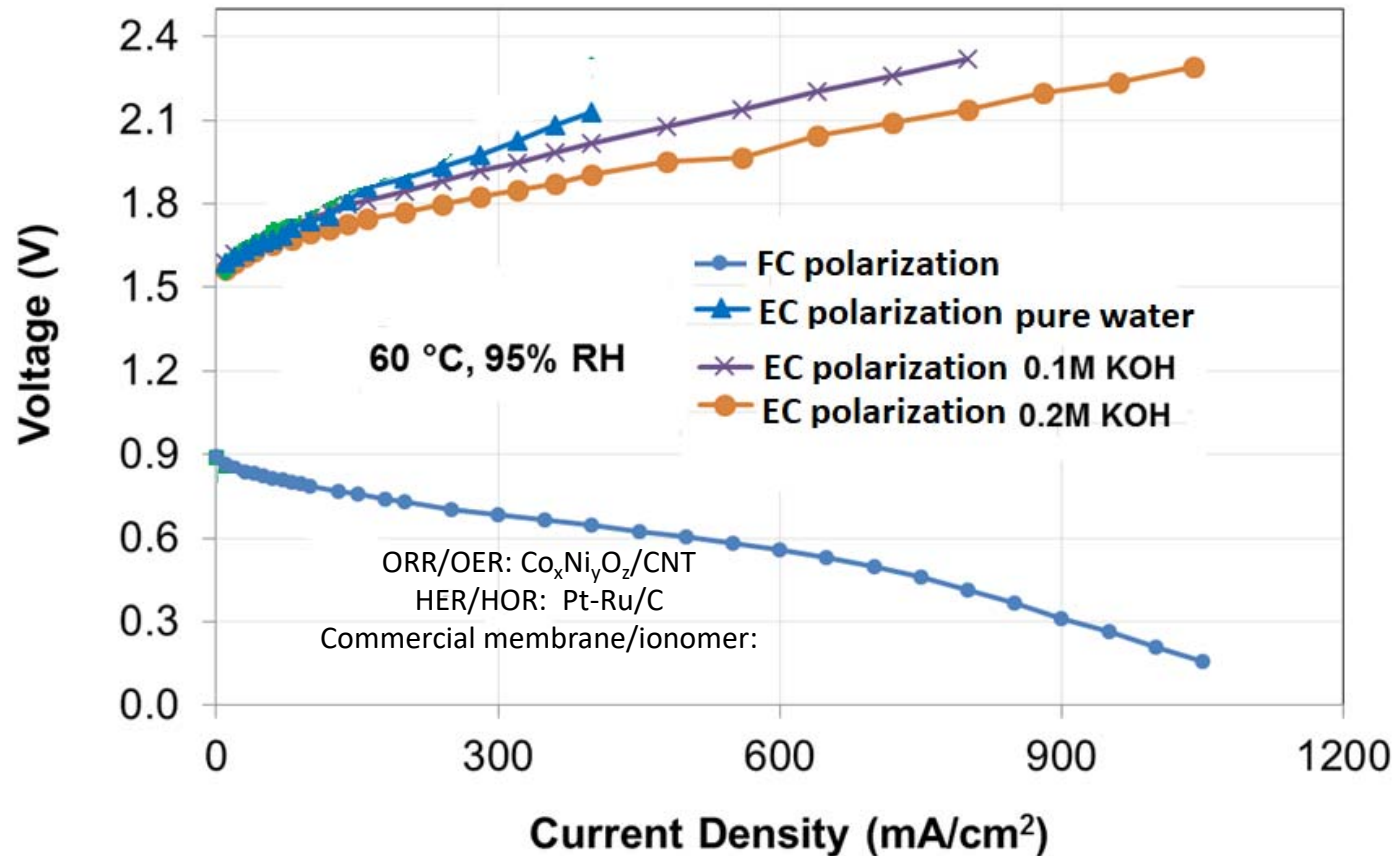
Accomplishment 2: Scale-up of $\text{Co}_3\text{O}_4/\text{CNT}$ Catalyst Synthesis (Giner)



5 grams of catalyst has been synthesized, with reproducible RDE results

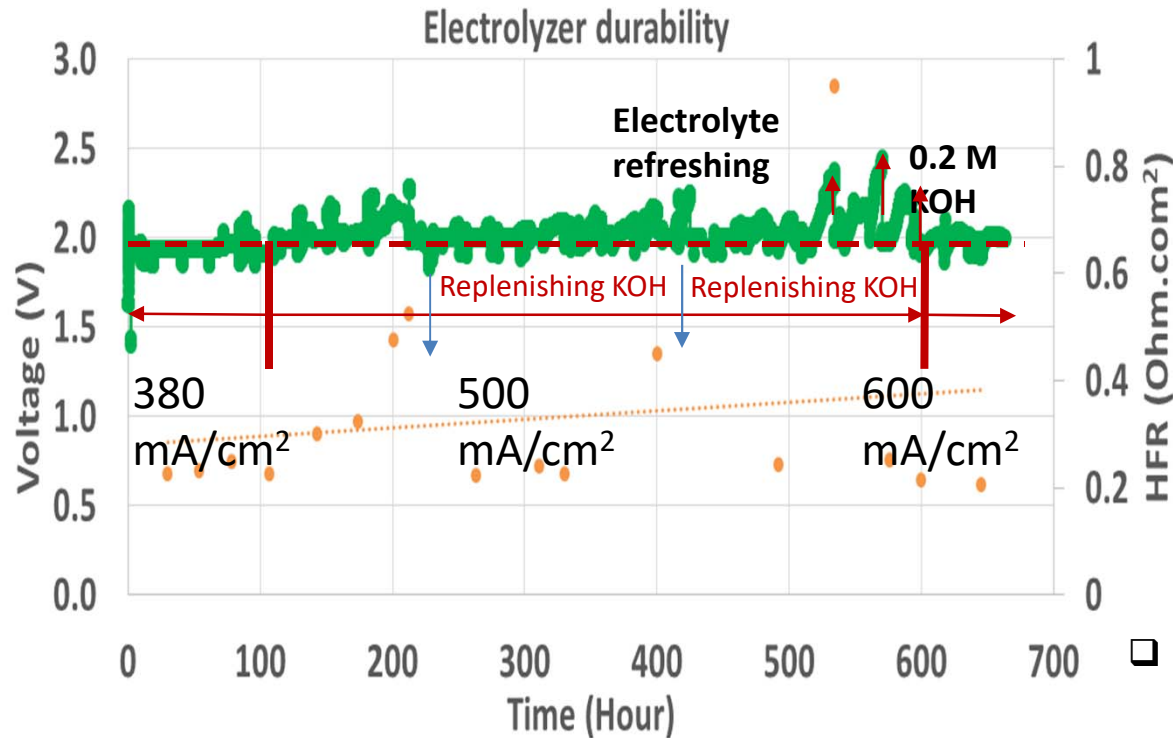
☐ SUNY NiCo_2O_4 will be mixed with Giner $\text{Co}_3\text{O}_4/\text{CNT}$ → $\text{Co}_x\text{Ni}_y\text{O}_z/\text{CNT}$

Accomplishment 5: MEA Performance Evolution (Giner)



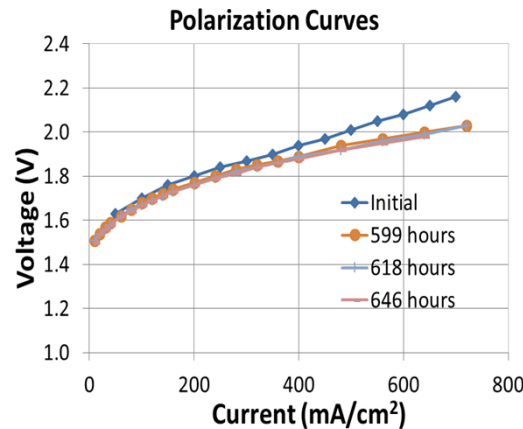
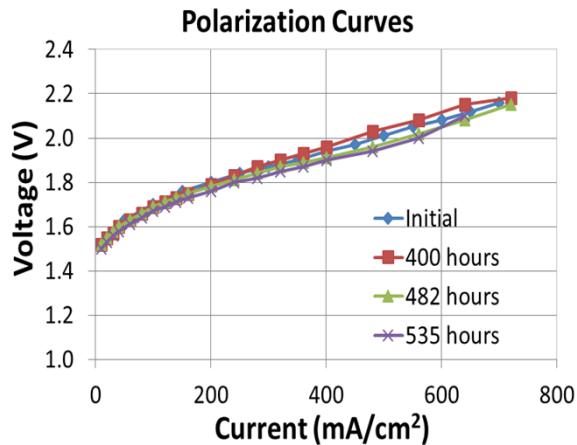
- ❑ The $\text{Co}_3\text{O}_4/\text{CNT}$ demonstrates good reversibility between FC and EC
- ❑ Poor EC performance w/o water can be improved by adding diluted KOH

Electrolyzer Cell Durability Tests



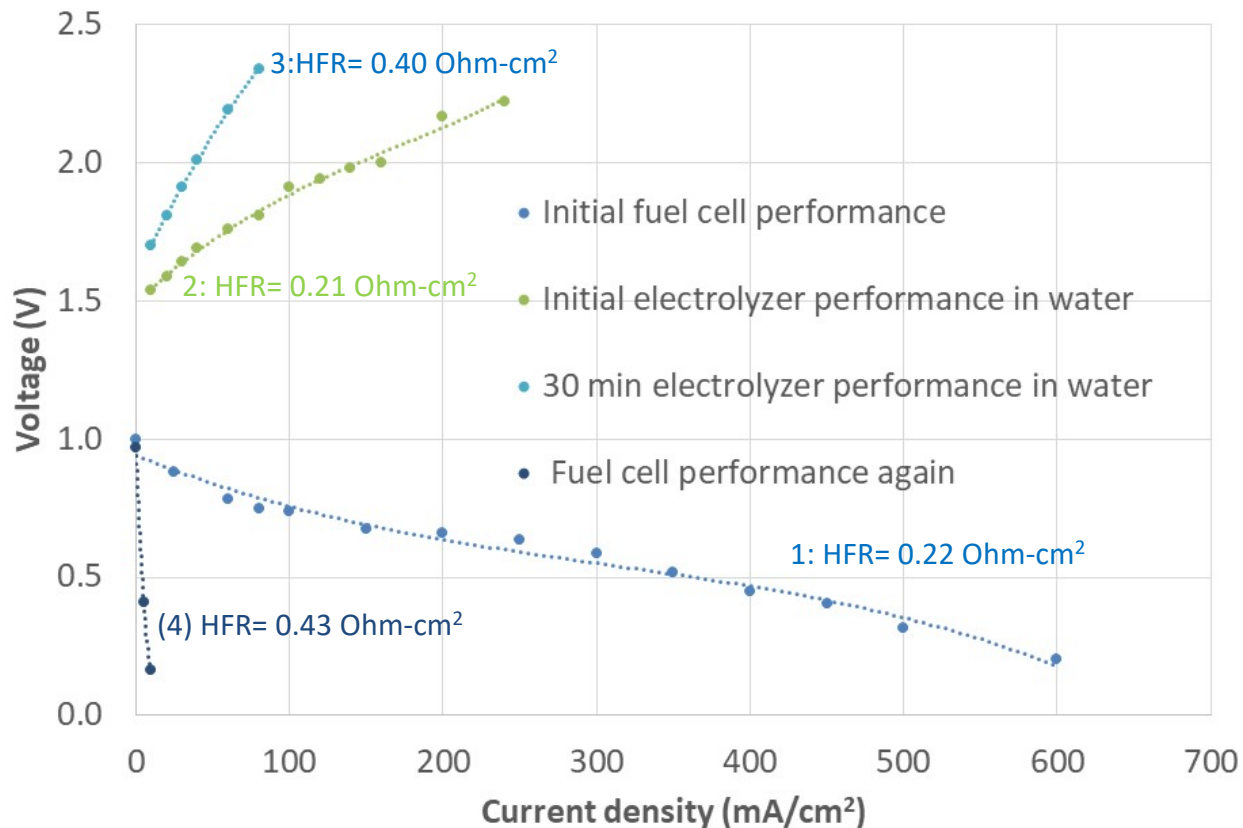
EC test conditions

Cell T=60 °C,
 (0.1 M KOH, 0.2 M KOH after 500 h)
Cathode: Co_xNi_yO_z/CNT
 (2 mg/cm², ionomer=20 w.t %)
Anode: PtRu/C
 (0.7 mg/cm², Ionomer=26 w.t. %).



- EC Durability test for 600 hours showed no performance decay, surpassing millstone:
 - 2.0V @ 600 mA/cm²
 - >10% decay after 500 hours
- Feeding diluted KOH solution improved EC performance
- HFR increased as the KOH was depleted and replenishing KOH lower the HFR and improve the EC performance

Reversible Operation with Water in EC



MEA

Cathode: $\text{Co}_x\text{Ni}_y\text{O}_z/\text{CNT}$
 (2 mg/cm², ionomer=20 w.t.%)
 Anode: PtRu/C
 (0.7 mg/cm², ionomer=26%).

FC Conditions

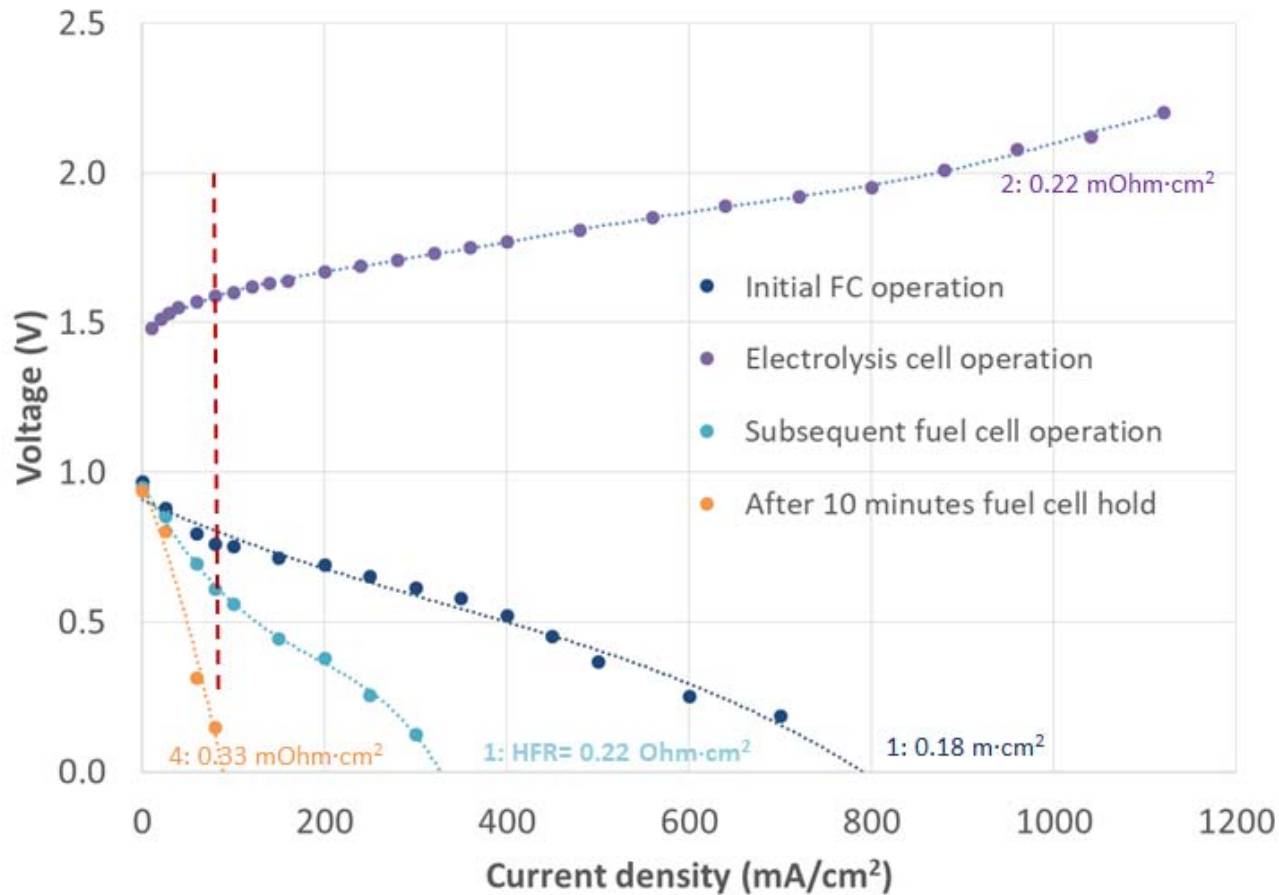
Cell T= 60 °C,
 Anode/cathode T: 57/60 °C
 H₂/O₂ backpressure: 35 psia.

EC test conditions

Cell T= 60 °C,
 DI water feeding

- EC performed poorly with pure water feeding
 - Performance degraded even after 30 minutes
- FC performance cannot be recovered after EC test
- HFR increased after 30-minute EC test, indicating alkaline membrane

Reversible Operation with KOH Solution in EC



MEA

Cathode: Co_xNi_yO_z/CNT
(2 mg/cm², ionomer=20 w.t.%)
Anode: PtRu/C
(0.7 mg/cm², ionomer=26%).

FC Conditions

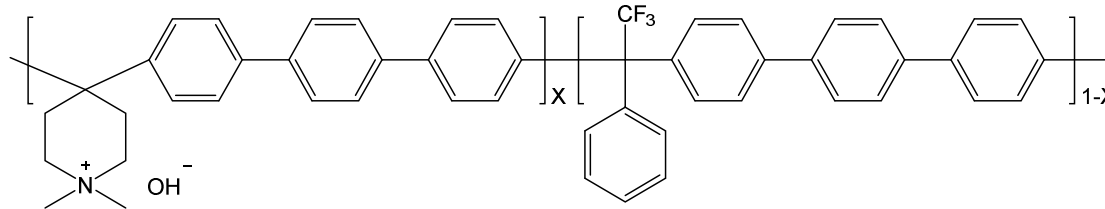
Cell T= 60 °C,
Anode/cathode T: 57/60 °C
H₂/O₂ backpressure: 35 psia.

EC test conditions

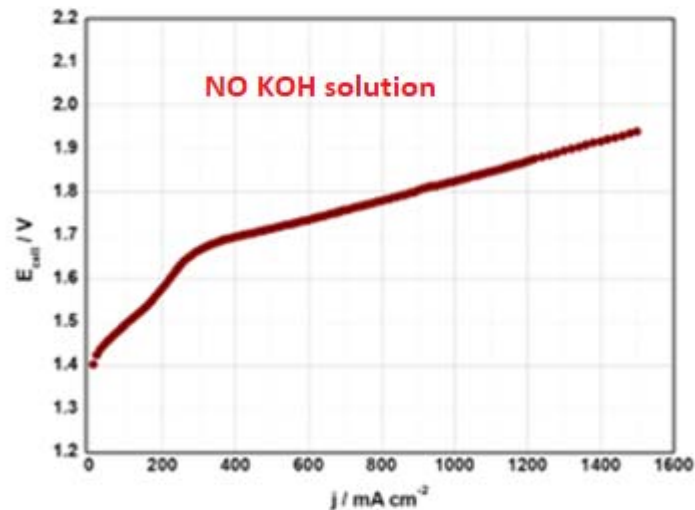
Cell T= 60 °C,
DI water feeding

- ❑ Introduction of KOH introduction in the water tremendously improve the EC performance
-Due to improved OH⁻ transport in the electrode
- ❑ FC performance cannot be recovered due to likely degraded membrane and ionomer under EC operation, evidenced by increased HFR

New Membrane and Ionomer: Enabled Electrolysis Operation without KOH



#1 HEMEL Performance of TP-100 ionomer with water-fed @ 80 °C



Data provided by Prof.
Yushan Yan at University
of Delaware

- Polarization curve obtained by applying current steps (30 sec/step),
- Operation Conditions: $T = 80\text{ }^{\circ}\text{C}$; DI water fed @ 3 ml/min;
- HEMEL Performance: 900 mA/cm² @ 1.8 V.
- AC impedance after the cycles is 135 - 180 mΩ·cm².

- ❑ New membrane and ionomer developed by University of Delaware enabled EC operation without KOH: stable under high potential or oxidation-resistant

Summary

- ❑ Bifunctional OER/ORR catalysts NiCo_2O_4 and $\text{Co}_3\text{O}_4/\text{CNTs}$ have been optimized with improved activity and durability:
 - Efforts focused more on the non-carbon catalyst development
- ❑ Bifunctional electrodes and MEAs have been optimized for durability tests and reversible operations:
 - Commercial alkaline membrane and ionomer are unstable under electrolyzer cell (EC) operation, likely due to high-potential (oxidation) caused materials degradation
 - Introduction of diluted KOH tremendously improved the EC operation but did not improved fuel cell (FC) operation without KOH solution
 - MEA under EC operation demonstrated 600-hour durability barely showing any performance decay, demonstrating remarkable catalyst stability
- ❑ Oxidation-resistant alkaline membrane and ionomer are needed to enable EC operation and reversible fuel cell operation without introducing KOH solution

Collaborations

Institutions	Roles
<u>Giner Inc. (Giner)</u> Lead: Hui Xu (PI), Shuai Zhao, Tom McCallum	Oversees the project; metal oxide supported on CNTs; bi-functional MEA fabrication and optimization; reversible fuel cell design; cost analysis
<u>SUNY -Buffalo(SUNY)</u> Sub: Gang Wu and Shiva Gupta	Graphene tube and metal oxide based OER/ORR bi-functional catalyst development; MEA fabrication
<u>National Renewable Energy Laboratory (NREL)</u> Sub: Bryan Pivovar, Shaun Alia, and Andrew Park	AEM development; HOR/HER catalyst development
<u>University of Delaware</u> Collaborator: Yushan Yan	Novel alkaline membrane and ionomer development

Future Plans

- ❑ Improve fuel cell/electrolysis operational reversibility using oxidation-resistant alkaline membrane and ionomer
- ❑ Complete dual operation durability test
- ❑ Perform techno-economical analysis

Any proposed future work is subject to change based on funding levels

Acknowledgments

- ❑ **Financial support from DOE EERE Fuel Cell Technologies Office**, Incubator Program Award # DE-EE0006960
- ❑ **DOE Program Managers**
 - Dr. David Peterson
 - Donna Ho
- ❑ **Giner**: Shuai Zhao, Tom McCallum, Corky Mittelsteadt
- ❑ **SUNY**: Prof. Gang Wu, Shiva Gupta, Mengjie Chen, Surya Devaguptapu
- ❑ **NREL**: Drs. Bryan Pivovar, Shaun Alia and Andrew Park
- ❑ **U. Delaware**: Prof. Yushan Yan