



Developing Novel Platinum Group Metal-Free Catalysts for Alkaline Hydrogen and Oxygen Evolution Reactions (DE-EE0008082)

PI: Prof. Sanjeev Mukerjee

Department of Chemistry and Chemical Biology,

Northeastern University

360 Huntington Ave, Boston, MA 02115

April 30th, 2019

Project ID: P156

This presentation does not contain any proprietary, confidential, or otherwise restricted information









Lawrence Livermore National Laboratory





Project Overview

Project	Partners
---------	----------

PI: Sanjeev Mukerjee - Northeastern University Partner organizations:

- University of Delaware Prof. Yushan Yan
- Advent North America Dr. Emory De Castro

Project Vision

This project will develop stable, high-conductivity, and high-strength AEMs, stable and active PGM-free catalysts for hydrogen and oxygen evolution reaction (HER/OER), and high-performance electrode architectures.

Project Impact

The materials being developed under this project have the potential to unlock the cost advantages of AEM electrolyzers, enabling a clear pathway for achieving <\$2/KgH2 (on scale) with efficiency of 43 kWh/kg H2 via anion exchange membrane (AEM)-based electrolysis.

*this amount does not cover support for HydroGEN resources leveraged by the project (which is provided separately by DOE)









Approach - Summary

Project Motivation

The focus of this project is on durable, highperformance materials advanced water splitting, enabling a pathway for achieving <\$2/Kg_{H2} with efficiency of 43 kWh/kg H₂ via anion exchange membrane (AEM)-based electrolysis. We aim to advance these goals via an understanding of both hydrogen and oxygen evolution reactions leading to novel platinum group metal (PGM)-free catalyst materials in conjunction with improvements in membrane and ionomers and gas evolution electrodes with corresponding characterization and testing.

Barriers

Current state-of-the-industry anionic exchange membranes are subject to degradation above 60 °C. This project aims to develop a new anionic exchange membrane capable of operating at elevated to temperatures, thereby increasing the kinetics of electrolysis.

Key Metrics

	Units	Year 1	Year 2
Cell voltage	V	1.92	1.82
Current density	A/cm ²	0.8	1
Operating temp	°C	80	90
Membrane ASR	$\Omega{ m cm^2}$	0.15	0.12
HER overvoltage @500 mA/cm ² , 25 ^o C	mV	300	200
OER overvoltage @500 mA/cm ² , 25 ^o C	mV	150	125
Model voltage	V	1.74	1.69
Unexplained voltage loss	V	0.18	0.13



Approach - Summary

Partnerships and Roles

- Northeastern Univ., (Prime) Boston, MA: S. Mukerjee (P.I):
 - Novel HER catalysts:
 - Metal-Metal Oxides
 - Functionalized mono-metallic
 - Embedded Metal-N-C Systems
 - Novel OER Catalysts:
 - Double layer metal oxides on Raney Ni (strategies for increasing active site density)
 - Half and Single Cell Tests (NEU)
 - In situ Raman and Synchrotron X-ray Spectroscopy
- University of Delaware (Sub-awardee), Newark, DE: Prof. Y. Yan
 - Novel Membranes and lonomers
 - Design of multi-cation Poly(aryl-piperidinium) Tri-quaternary-ammonium (PAP-TQN) membranes and ionomers for high temperature operation (90°C).
- Advent North America (Sub-awardee), Cambridge, MA: Dr. Emory De Castro
 - **Novel Electrodes** designed for gas evolution with and without liquid flow
- <u>National Lab Partners</u>:
 - NREL MEA Prep and testing
 - LBNL Small Angle Scattering and Transport Modeling
 - Sandia National Labs Interfacial modeling



Approach - Innovation

NUCRET will leverage its expertise in catalyst development to synthesize and characterize catalysts for both OER and HER.

- HER catalysts
 - Metal/metal oxide NiCr/C synthesized through borohydride reduction
 - Metal nitride Ni²⁺ was chelated using cupferron, a nitrogenous compound. The complex was reduced via heat treatment in argon.
 - Metal organic framework Ni and N precursors were incorporated into the ZIF-8 template.
- OER catalyst Metal/metal oxide
 - NiFe/Raney Ni has been shown to be an effective non-PGM catalyst for OER

BP1 activities focused on identifying the most promising HER catalysts, validating the OER catalyst, and establishing baseline performance of an electrolysis equipped with these materials.

The University of Delaware has developed a polyaryl piperidine-based AEM with robust physical properties capable of operating at elevated temperatures (60 $^{\circ}C$ <). BP1 activities related to AEM development included:

- Thorough characterization of physical properties
- Quantifying thermal stability
- Increasing the ion-exchange capacity (IEC) through the introduction of an additional active site

Relevance and Impact



- This program aims to develop thermally sustainable anionic membranes that will enable the use of cost-efficient transition metal catalysts.
- Our group will be able to leverage the capabilities of the ENM to generate multiscale modeling and molecular dynamics generated simulations of the membrane catalyst interface. In addition, its expertise with advanced ionomer materials, the EMN will provide durability and validation protocols of electrolyzer materials
- These activities can be applied to other areas of electrolysis research to characterize ionomer stability and electrolyte ion flux.







Approach – HER in Alkaline

Volmer: M + H₂O + $e^- \rightarrow M$ -H_{ads} + OH⁻

Heyrovsky:

 $M-H_{ads} + H_2O + e^- \rightarrow H_2 + OH^-$

Tafel: 2 M-H_{ads} \rightarrow H₂



Northeastern University Center for Renewable Energy Technology

Approach - Materials Design Strategy: Non-PGM HER Electrocatalysts

Strategy 1: State of the art Metal-Metal Oxide electrocatalysts



Strategy 3: Embedded metal nanoparticle electrocatalysts









 Hydride formation did not go past sub-surface in the case of Ni-cup/C, evidenced by stable HER's performance of initial and post 1h hold @ -0.3 V.

Northeastern University

Center for Renewable Energy Technology

- In the case of both Ni/C and NiCr/C, hydride formation to go beyond surface and change the Ni lattice structure, resulting in lower HER performance
- Cycling the Ni-cup/C at high oxidation voltage did not oxidize the catalyst
- Opposite trend observed in the case of Ni/C and NiCr/C



Irreversible process for Ni metal.





Approach – Surface Protection

Ni reduced /K600 has no graphite.





Ni-Cup



848

Ni-cup/ K600

- Multi SAD patterns show graphene plane (002)
- Each Ni particle has about 5 graphite layers or more (some have 10 layers)

XPS

- Ni/C-No Metal-Surface comprised of a mixture of oxides
- Ni-cup/C-Surface Ni metal-Reduced
 amounts of surface oxides





- At anodic potentials (0.1V and 0.7V vs. RHE), the Ni-cup/K600 remains at Ni(0) state, where the Ni/ K600 is oxidized even at 0.1 V vs. RHE.
- Hydrogen gas bubbles formed at cathodic potentials and interfered with the XAS acquisition.

Ni-Cup/C remains metallic (more noble) while Ni/C is oxidized



Approach - Mechanistic Aspects

Materials Strategy $\Delta \mu$ shift $\propto E$ in situ XAS Precipitation 12 Annealing Reduction Ni K-edge Ni(NO₃)₂ Ni-O_v/C Co(NO₃)₂ 11 Ni-Co-O_v/C Raney Fe(NO₃)₃ Raney Raney Ni-Fe-Co/Raney - Ni-Fe-O_v/C Ni NaBH₄ Ni NH₂ Argon 10 2. Ni-Fe-Co-O_v/C dropwise 400°C ΔE (eV) vs. addition 30min 9 Metal/Aniline complex Ni/Fe Alloy Shift of 8 Ni^{2+/3+} Strategy 1: Improve surface area of ** Peaks Ni support Strategy 2: Improve active site 6 density 1.5 1.0 1.1 1.2 1.3 1.4 1.6 Strategy 3: Optimize Fe/Ni Ratio E (V vs. RHE) ΔE (eV, XAS) vs •••• appi Fe stabilizes Ni in lower oxidation state 4 OH-O₂ + 2H₂O + 4e⁻ Co facilitates Ni oxidation Yeager Mechanism (validated for base-metal oxides) $2S^{z} + 2OH^{-}$ \rightarrow 2S^z-OH + 2e-(Fast adsorption of OH⁻) $2S^{z+1}$ -OH + 2e- (Oxidation of metal - rds) 2S^z-OH \rightarrow 2 S^{z+1}-OH + 2 OH⁻ \rightarrow 2S^z + O₂ + H₂O (Fast Acid-Base reaction) DFT results: Fe³⁺ sites exhibit optimized binding of -OH & -OOH intermediates

Proton-transfer from Ni³⁺ redox shuttles

Bell et al. JACS 2015

y-NiOOH:Fe

Northeastern University Center for Renewable Energy Technology

Approach – Ionomer Synthesis



Milestone #	Project Milestone	Progress
2.1	Synthesis of PAP-TP polymers	Completed
2.2	Synthesis of two- ammonium-cations iodo- alky side-chain (DQN-I)	Completed
2.3	Synthesis of PAP-TQN polymers:	Completed (MQN-X side chain was found to be more suitable for synthesis rout and membrane properties)



NREL Characterization



PAP-TP-MQN	I (UD-PAP)	24.3	No strip; melted
PAP-TP-85 (1) UD 39	17.6	16.5
PAP-TP-85 (2) (UD 45)	13	8
Milestone #	Project	Milestone	Progress
Go/No Go	Achieve AE ASR of (M membrane $0.15 \ \Omega \ { m cm^2}$	Completed



- IEC_{titration}: 2.9 mmol/g (CI) IEC_{theory}: 3.1 mmol/g (CI)
- Almost no change over 1000 h at 90 ^oC in 1M KOH solution observed by ¹H-NMR
- After 1000 hour in KOH: 0.025 ÷ 1 ÷ (85 %)
 ~ 2 % (decomposition yield)



Temp: 60°C, HOR: Pt/C (TKK 47.2%), Anode: Humidified H₂; Cathode: Humidified N₂.



Milestone #	Project Milestone	Progress
1.1	Deliver three 5 gm batch samples of Ni-MOx/C HER catalysts for RDE and Single Cell tests with η of 300 mV @ 500 mA/cm2 and 2 A/cm3 (HER)	Completed (did not meet overvoltage target)
1.2	Deliver one 5 gm batch of Ni-Nx/C HER catalysts for RDE and Single cell studies η of 300 mV @ 500 mA/cm2 and 2 A/cm3 (HER)	Completed
1.3	Deliver one 5 gm batch of M-N(S)-C MOF HER catalysts for RDE and Single cell studies η of 300 mV @ 500 mA/cm2 and 2 A/cm3 (HER)	Catalyst was synthesized but <u>did not meet</u> performance milestone.
1.4	Deliver three 5 gm batches of MMOx OER catalysts for RDE and single cell tests	Completed
Go/No Go	OER Electrocatalysts: Using PGM-free OER electrocatalsyts in an AEM MEA operating under electrolysis conditions, at a current density of 500 mA/cm2 achieve 150 mV or lower over-voltage and 6 A/cm3 (VCD).	Completed

Water splitting cell equipped with at non-PGM OER electrode (NiFe/Raney) out performed a cell with an iridium oxide electrode. 1.8 V constitutes ~ 150 mV (η), Temp: 50°C. Northeastern University Center for Renewable Energy Technology



- NiFe/Raney Ni was used as the OER catalyst
- NiCr/C and Ni-cup/C required 1.92 and 1.95 V, respectively, to achieve a current density of 800 mA/cm²
- High temperature stability of University of Delaware's ionomer materials enable this performance.
- Ni-cup/C and NiCr/C both showed similar stability in a water electrolysis cell.
- H₂ studies and RDE data suggests that over time, NiCr/C is less stable.

Milestone #	Project Milestone	Progress
Go/No Go		



Collaboration – SNL modeling

Potential as a function of surface charge





• Simulated the effects of charging the electrode on the ionomer and mobile species to tie local pH and other resolved quantities to water splitting efficiency

Water surface coverage for increase oxide coverage



(a) bare Ni Ni: gray, N+: blue, O: red, H: white

(c) partial NiO coverage Quarter circle centered on lower left corner



Collaboration – LBNL modeling



- Model fit to DI water-fed electrolyzer data from NEU
- Carbonate feed at the anode performs significantly better *despite* the predicted Nernstian voltage loss
 - Observed difference between feeding water and carbonate at anode equivalent to 1,000x increase in product of *active surface area* and *exchange current density*
 - Carbonate increases ionic pathways (decreased tortuosity) and maybe impacts kinetics



AEM does not a strong phase-separation and structural features

Northeastern University

Center for Renewable Energy Technology

Do not change with hydration

SAXS Profiles of UD-AEM (PAP-TP-85)

•

- Do not change with <u>temperature</u>
 - AEM does not respond to environment the way other PEMs (e.g., Nafion) do. Indicates a set morphology that swells macroscopically without affecting nanophase separation

- Conductivity in liquid water correlates with hydration (upon anion-exchange)
 - In OH form, conductivity is slightly lower than that of Nafion in proton form (~ 60-80 mS/cm)
 - OH form showing the highest uptake and conductivity
 - Reveals critical role of anion form and water content despite lack of a phase-separated nanostructure
 - Large shift in conductivity in OH form is in accord with the mixed, gel-like morphology facilitating hopping





Catalyst development:

- HER Electrocatalysts: An overvoltage of 300 mV or lower was achieved at a current density of 500 mA/cm² using PGM-free HER electrocatalysts in an AEM MEA operating under electrolysis conditions.
- OER Electrocatalysts: At a current density of 500 mA/cm2 achieved at 150 mV using PGM-free OER electrocatalysts in an AEM MEA operating under electrolysis conditions,
- AEM MEA: Demonstrated a PGM-free AEM MEA electrolyzer performance of 0.8 A/cm2 at ≤1.92 V.

Membrane development:

- Reliable methods for preparing single and multi-cation polyaryl piperidinyl triphenyl AEM membranes were devised.
 - Physical properties of these materials such as ion-exchange capacity, physical stress, water uptake, swelling, and conductivity were obtained.
- University of Delaware materials showed an area-specific resistance of 0.048 Ω ·cm²
- Materials showed a loss of <2% of ion-exchange capacity after a 1000 hour treatment in 95 °C 1M KOH

Additional Activities

- Mathematical modelling of AEM electrolysis systems showed a decreased pH at the electrode/solution interface resulting in voltage loss. (LBNL)
- Modelling of the electrode/ionomer interface demonstrate the importance of the presence of surface metal oxides in facilitating the adsorption of water molecules. (SNL)
- SAXS profiles elucidate the microstructure of the UDel baseline membrane, showing its nanoscopic features remain consistent under various hydration conditions. (LBNL)



- 1. Materials Discovery: (NEU & Udel) Q5-Q8
 - HER Catalysts: Ni-MOx-Nx catalyst optimization, stability tests, structure function determination.
 - OER catalyst: Enhance active site density of Ni-Fe-Co/Raney Ni.
 - Reinforced PAP-MQN membranes leading to PAP-TQN system for FY3.
 - Membrane and Ionomer: Extensive durability tests, Full Cell Testing at elevated temperatures
 - Electrodes and Ink Composition: New electrode design with novel ink compositions
- 2. Cell Tests: (NEU) Q6-Q8
 - Accurate determination of OER overvoltage
 - Optimization of Full Cell for long duration tests
- 3. Detailed modeling of transport processes for deciphering carbonate effect (LBNL) Q6-Q8
- **4. SAXS measurement** for membrane structure and morphology as a function of temperature, ion exchange and humidity (LBNL) Q6-Q8
- 5. Interfacial modeling to understand ionomer effects (Sandia) Q6-Q8
- 6. Durability measurements on PAP-MAC membranes (NREL) Q7-Q8
- 7. MEA optimization and testing (NREL) Q5-Q8



Publications:

- Unifying Alkaline Hydrogen Evolution/Oxidation Reaction Kinetics by Identifying the Role of Hydroxy-Water-Cation Adducts. Ershuhai Liu, Li Jiao, Huong Doan, Zeyan Liu, Yu Huang, Kuzhikalail M. Abraham, and Sanjeev Mukerjee, *J. Amer. Chem. Soc.* 2019, 141, 3232-3239.
- Current Understanding of Sluggish Kinetics of Hydrogen Evolution and Oxidation reactions in Base. Qingying Jia, Ershuhai Liu, Li Jiao, and Sanjeev Mukerjee, *Current Opinion in Electrochemistry*. **2018**, *12*, 209-217.

Presentations:

- Mukerjee, Sanjeev, and Qingying Jia. "Fundamental aspects of regenerative hydrogen electrocatalysis in alkaline pH." In ABSTRACTS OF PAPERS OF THE AMERICAN CHEMICAL SOCIETY, vol. 256. 1155 16TH ST, NW, WASHINGTON, DC 20036 USA: AMER CHEMICAL SOC, 2018.
- Mukerjee, Sanjeev, Jingkun Li, and Qingying Jia. "Current Understandings of the Slow Kinetics of the Hydrogen Evolution Reaction in Alkaline Media." In *Meeting Abstracts*, no. 29, pp. 1703-1703. The Electrochemical Society, 2018.
- Qingying Jia, Jingkun Li and Sanjeev Mukerjee. "Understanding the Improved Kinetics of the Hydrigen Evolution/Oxidation Reactions on Pt-Oxophilic Metal Systems in Alkaline Media." In Meeting Abstracts, no 29, pp 1710-1710. The Electrochemical Society, 2018.
- Ian Kendrick, Michael Bates, Qingying Jia, Huong Doan, Wentao Liang and Sanjeev Mukerjee. "Tuning Ni Surfaces for Enhanced Oxygen Evolution Reaction." In *Meeting Abstracts*, no 29, pp 1702-1702. The Electrochemical Society, 2018.



Technical Backup Slides





Model physics

Thermodynamics

Standard cell potential Equilibrium H₂O content membrane, liquid, vapor

Kinetics

Butler-Volmer for HER and OER H_2O phase change

Transport

Stefan-Maxwell diffusion for gas-phase components Darcy's law for liquid, gas phases Ohm's law for e⁻ current Modified Ohm's law for ion current H₂O transport by ion drag H₂O diffusion in membrane Enthalpy balance for energy **Conserved** quantities

Mass; Charge; Energy

Constitutive relations

Faraday's law Ideal-gas law

Properties

Function of Tand H₂O content

Equations (12): 7 2nd-order PDEs; 5 Algebraic equations



- PAP-TP-85: HCO₃⁻ form possesses slightly greater κ than CO₃²⁻ counter ion for given hydration number
- PAP-TP-85-MQN exhibits greater
 water uptake than PAP-TP-85 for a
 given RH, which could be due its
 higher IEC value (3.0 VS. 2.1 mmol/g)

MQN has also higher conductivity





Anion Conductivity: Temperature Impact

Optional slide on conductivity-temperature of new chemistry



- The anion conductivity of PAP-TP-85-MQN increases with temperature
- Activation energy of anion conduction decreased from 56.5 to 15.5 kJ/mol, when increasing the RH from 20 to 98%
 - ✤ The plotting of Ea as a function of RH is in progress





Notes and Comments

Not for AMR – this slide provides a status update and comments

• Slide -1: Summary (self-contained)

Solution New results showing temperature-**in**dependent morphology in water

• Slide -2: Comparison of New (MQN) AEM

 $\boldsymbol{\boldsymbol{\boldsymbol{\forall}}}$ It exhibits higher uptake and conductivity than the previous set

• Key overall findings:

- Lack of a strong phase-separated morphology indicates a gel-like structure with strong ion concentration which cause a big shift in OH conductivity due to effective facilitation of hopping
- Sor the same reason activation energy for conductivity shows a strong dependence on hydration (RH)
- In hydrated state, lack of morphological changes with heating agrees with the low activation energy for conductivity

• Our data analysis will continue this year to draft a paper summarizing these results



Collaboration – LBNL modeling



0

7

pH

14

 [1] C. C. Pavel *et al.*, *Angew. Chemie - Int. Ed.*, vol. 53, no. 5, pp. 1378–1381, 2014.