### OFFICE OF NAVAL RESEARCH

### FINAL REPORT

for

### Grant Number: N00014-98-1-0278

### **PR Number 98PR03689-00**

### ONR POC: Dr. Richard Carlin, ONR-331 (703) 696-5075

### "Electrode Surface Modification for Cathode Catalysis in Semi-Fuel Cells"

by

### Russell R. Bessette, Ph.D.

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16 December, 1998

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### PART I

### OFFICE OF NAVAL RESEARCH PUBLICATIONS/PATENTS/PRESENTATIONS/HONORS REPORT

PR Number: 98-PR-03689-00

Contract/Grant Number: N00014-98-1-0278

Grant Title: "Electrode Surface Modification for Cathode Catalysis in Semi-Fuel Cells"

Principal Investigator(s): Russell R. Bessette, Ph.D.

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Phone Number: (508)999-8587 Fax Number: (508)999-9167 E-mail Address: RBessette@UMASSD.EDU http address:

- a. Number of papers submitted to refereed journals, but not published: <u>1</u>\_\_\_\_
- b. Number of papers published in refereed journals (for each, provide a complete citation): <u>0</u>
- c. Number of books or chapters submitted, but not yet published: \_0\_\_\_
- d. Number of books or chapters published (for each, provide a complete citation): <u>0</u>
- e. Number of printed technical reports/non-refereed papers (for each, provide a complete citation): \_0\_\_\_
- f. Number of patents filed: <u>0</u>
- g. Number of patents granted (for each, provide a complete citation): <u>0</u>
- h. Number of invited presentations (for each, provide a complete citation):
- i. Number of submitted presentations (for each, provide a complete citation): 1
- j. Honors/Awards/Prizes for contract/grant employees (list attached): \_0\_\_\_\_ (This might include Scientific Society Awards/Offices, Selection as Editors, Promotions, Faculty Awards/Offices, etc.)
- k. Total number of Full-time equivalent Graduate Students and Post-Doctoral associates supported during this period, under this PR number: <u>0.4</u>

Graduate Students: <u>0.4</u>

Post-Doctoral Associates: \_0\_\_\_

including the number of,

Female Graduate Students: \_\_\_\_\_

Female Post-Doctoral Associates: \_\_\_\_

the number of

Minority\* Graduate Students: \_\_\_\_\_

Minority\* Post-Doctoral Associates: <u>0</u>

and, the number of

1.

Asian Graduate Students: <u>0</u>

Asian Post-Doctoral Associates: 0

Other funding (list agency, grant title, amount received this year, total amount, period of performance and a brief statement regarding the relationship of that research to your ONR grant)

### a. Abstracts Submitted:

An abstract entitled "Electrode Surface Modification for Cathode Catalysis in Semi Fuel Cells" by R.R.Bessette, J.M. Cichon and D.W. Dischert has been submitted for review, acceptance and subsequent presentation at the Electrochemical Society (ECS) Meeting to be held in Boston, MA, November, 1998. A subsequent paper is planned for publication in the Journal of the Electrochemical Society.

### h. Invited Papers:

Aspects of this work and preliminary results were presented by R.R. Bessette at the AFOSR/ONR 6.1 Electrochemical Science and Technology Program Review, 15-16 January, 1998, Ventura, CA, entitled "Electrode Surface Modification for Cathode Catalysis in Semi Fuel Cells". Co-authors include J.M. Cichon and D.W. Dischert.

### i. Submitted Presentations:

Naval Undersea Warfare Center (NUWC) Division Newport - FY98 Independent Research (IR), PI: E.G. Dow, AI(s): R.R. Bessette, J.M. Cichon, "Surface Stabilization and Characterization of Electrochemical Catalysts for Use in the Aluminum Hydrogen Peroxide Semi-Fuel Cell Battery System". July, 1997.

### 1. Other Funding:

1) Received 6.1 funding of \$135K for FY98 from ONR-33's Competitive Technology Assistance Program (CTAP)- "Semi-Fuel Cell Power System S&T"

2) Naval Undersea Warfare Center (NUWC) Division Newport - FY98 Independent Research (IR), "Surface Stabilization and Characterization of Electrochemical Catalysts for Use in the Aluminum Hydrogen Peroxide Semi-Fuel Cell Battery System", \$30K, FY98.

3) Laboratory Participation Program(LPP) from NUWC-Newport Division, \$20K, FY98.

### PART II

- a. Principal Investigator: Russell R. Bessette
- b. Telephone Number: (508

c. ONR Program Officer:

(508) 999-8587

Dr. R. Carlin, ONR-331, 6.1 Electrochemical Science and Technology Program

- d. Program Objective(s):
  - 1. Identify the optimum conditions for fabricating a stable electrocatalytic cathode of palladium and iridium that will:
    - a. improve the cathode polarization potential
    - b. diminish the heterogeneous decomposition and direct chemical reactions
    - c. significantly improve (gain of 50%) the overall electrochemical cell efficiency

2. To evaluate the electrochemical performance of the catalyzed cathode

3. To characterize the structure and composition of the catalytic surface

### e. Significant Results:

Comparison of the cell voltages measured at  $250 \text{ mA/cm}^2$  for a baseline silver catalyst and for planar nickel, reticulated nickel and planar carbon substrates catalyzed with palladium and iridium show the catalyzed planar carbon yields the best cell voltage (1.37 V vs. 1.25 V for reticulated Ni) and thus the best power. Multiple discharges at 100mA/cm<sup>2</sup> for 45 minutes followed by the acquisition of a polarization curve show continued catalytic activity and stable electrochemical performance for catalyzed carbon and silver foil substrates only. The catalyzed carbon electrode exhibits the highest cell voltage between 50 and 300 mA/cm<sup>2</sup>.

Scanning Electron Microscopy (SEM) data of Pd and Ir catalyzed nickel and planar carbon substrates show non-uniform coverage of the nickel substrate but uniform coverage of the carbon substrate. Energy Dispersive Spectrometry (EDS) analysis of the surface confirms the SEM results.

X-Ray Photoelectron Spectroscopy (XPS) spectra of Pd alone, Ir alone and the Pd/Ir combination on nickel foil and carbon substrates have been obtained for samples prior to use in an  $Al-H_2O_2$  electrochemical cell. The data to date are very inconclusive. The presence of adventitious carbon and perhaps oxygen make interpretation of the spectra very difficult at this time. Additional experimentation is required. An X-Ray Diffraction (XRD) analysis of the Pd/Ir combination on a nickel substrate suggests the very interesting possibility of the formation of a Pd/Ir solid solution. Again, additional experimentation is needed.

Anodic and cathodic efficiency studies using silver foil and Pd/Ir on carbon as catalysts and EB50V aluminum alloy as the anode show an increase in the Al efficiency from 56% with silver foil to 64% with Pd/Ir on carbon. The  $H_2O_2$  efficiency increases from 32% with silver foil to 38% with Pd/Ir on carbon. The overall low efficiency numbers for the  $H_2O_2$  reduction is believed to be due to the small electrode area (6.45 cm<sup>2</sup>) used relative to the 100 mL electrode reservoir.

Other tests utilizing a 100 mL electrolyte reservoir in combination with a  $63 \text{ cm}^2$  electrode have yielded  $H_2O_2$  reduction efficiencies in the 60 to 70% range. Additional efficiency determinations using the larger electrode surface area while keeping the electrolyte volume at 100 mL are planned.

### f. Summary of Experimentation Plan

X-Ray Photoelectron Spectroscopy (XPS) and SEM/EDS analyses will continue to be used with addition of X-Ray Diffraction Spectroscopy (XRD) to characterize the structure of the catalyzed surface. Initial elemental distribution data have been obtained by XPS. Ultimately the chemical composition of the species on the surface needs to be determined.

Capillary gas chromatography (GC) will be used for evolved oxygen and hydrogen gas analyses. The  $O_2(g)$  analysis will quantify the  $H_2O_2$  decomposition process. The  $H_2(g)$  analysis will quantify the corrosion reaction of the aluminum anode with the hydroxide:

 $2AI + 2OH^{-} + 2H_2O - --> 2AIO_2^{-} + 3H_2(g)$ Quantification of the  $H_2O_2$  decomposition and the aluminum corrosion reactions will allow us to quantify the direct reaction of the Al with the  $H_2O_2$ . We will then be able to perform experiments to minimize this direct reaction.

Once a stable catalytic deposit is achieved, uniform catalysis of nanosized (extremely high surface area per unit volume upwards of 10,000 cm<sup>2</sup>/cm<sup>3</sup>) material of either carbon or nickel will be attempted. Successful completion of these experiments should lead to very high (>85%) efficiency for the reduction of H<sub>2</sub>O<sub>2</sub> catholyte or the newly proposed hypochlorite (ClO<sup>2</sup>) catholyte.

Experiments will be designed to quantify the low rate performance of the catholyte reduction ( $H_2O_2$  and ClO) and stability of the electrocatalysts as a function of temperature, flow rate, surface composition, substrate preparation and catholyte concentration.

Characterization of the fundamental chemical and electrochemical equations that describe these phenomena occurring at the electrode surfaces will be accomplished.

### g. Students Working on the Project

Mr. James M. Cichon, MS Chemistry candidate at UMASS-Dartmouth, began a position at Duracell, Inc. in Needham, MA on 29 June 1998. He will also be involved in the completion of his work on this project over the next 3 to 6 months.

Mr. Craig Deschenes, BS Chemistry candidate at UMASS-Dartmouth, began work on this project on 15 June 1998. He will work 24 hours per week throughout the summer and will continue to be actively involved with this project during academic year '98-'99. **Electrode Surface Modification for Cathode Catalysis** 

# in Semi-Fuel Cells

Bessette/Cichon, UMASS-Dartmouth

Technology Issues: Development of affordable, long endurance semifuel electric propulsion systems to meet the demands of current and future UUV applications.

**Objectives:** To identify the optimum conditions for fabricating a stable electrocatalytic cathode of palladium and iridium; to improve the cathode polarization potential; to diminish the heterogeneous decomposition and direct chemical reactions; to significantly improve (gain 50% relative) the overall electrochemical cell efficiency

### Approach:

- Utilize SEM/EDS and XRD to characterize the structure of the catalyzed surface
- Perform GC analyses of evolved gases (O<sub>2</sub> and H<sub>2</sub>) to quantify H<sub>2</sub>O<sub>2</sub> decomposition and Al reactions (corrosion/direct)
- Determine best method of forming a stable and reproducibly performing Pd/Ir catalytic surface
  Ouantify electrochemical

efficiencies

### Accomplishments:

- Pd/Ir catalyzed carbon substrate yielded best cell voltage, best power, continued catalytic activity and stable electrochemical performance
  - SEM and EDS data of Pd/Ir catalyzed on Ni is non-uniform and uniform on carbon
- •Al oxidation and H<sub>2</sub>O<sub>2</sub> reduction efficiencies increase for Pd/Ir catalyzed carbon vs. Ag foil catalyst



- Developments on a solution phase catholyte for Al aqueous batteries at reduced current densities indicate specific energies in excess of 200 Wh/lb are possible utilizing sodium hydroxide mixed with seawater as caustic electrolyte
  - Achievement of a more stable Pd/Ir catalyzed cathode will significantly improve power and catholyte electrochemical efficiency and allow application in a hypochlorite catholyte system

Electrode Surface Modification for Cathode Catalysis

# in Semi-Fuel Cells

Bessette/Cichon, UMASS-Dartmouth







REPORT DOCUMENTATION PAGE			⊢orm Аррго∨ва ОМВ No. 0704-0188		
Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0188), Washington, DC 20503.					
1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE 11 Jan 99	3. REPORT TYPE AND	DATES COV	ered Jan 98 - 30 Sept 98	
4. TITLE AND SUBTITLE	11 Jan 35			IG NUMBERS	
Electrode Surface Modification for Cathode Catalysis in Semi-Fuel Cells				G - N00014-98-1-0278	
6. AUTHOR(S) Russell R. Bessette, Ph. D.					
7. PERFORMING ORGANIZATION NAMES(S) AND ADDRESS(ES) University of Massachusetts Dartmouth Department of Chemistry and Biochemistry 285 Old Westport Road North Dartmouth, MA 02747				RMING ORGANIZATION T NUMBER	
9. SPONSORING / MONITORING AGENCY NAMES(S) AND ADDRESS(ES) Department of the Navy Office of Naval Research 800 North Quincy Street Arlington, VA 22217-5660			10. SPONSORING / MONITORING AGENCY REPORT NUMBER		
11. SUPPLEMENTARY NOTES     a. DISTRIBUTION / AVAILABILITY STATEMENT     12. DISTRIBUTION CODE					
Approved for Public Release; distribution is unlimited.					
13. ABSTRACT (Maximum 200 words) The characterization and use of a palladium and iridium catalyst combination in an aluminum-hydrogen peroxide semi-fuel cell is described. The Pd/Ir combination outperforms palladium alone or iridium alone on the same substrate. Scanning Electron Microscopy (SEM) data of Pd and Ir catalyzed nickel and planar carbon substrates show non-uniform coverage of the nickel substrate but uniform coverage of the carbon substrate. X-ray Photoelectron Spectroscopy (XPS) spectra of Pd alone, Ir alone and the Pd/Ir combination on nickel foil and carbon substrates are inconclusive.					
The presence of adventitious carbon and perhaps oxygen make interpretation of the spectra inconclusive at this time. Anodic and cathodic efficiency studies using silver foil and Pd/Ir on carbon as catalysts and EB50V aluminum alloy as the anode are reported upon. Operation of the AI-H2O2 semi-fuel cell incorporating a Pd/Ir on carbon cathode at low current densities yielded H2O2 reduction efficiencies in the 60 to 70% range					
14. SUBJECT TERMS Aluminum Electrochemical Cells; Hydrogen Peroxide Electrocher Cells;Semi-Fuel Cells; Catalysis				15. NUMBER OF PAGES 7 16. PRICE CODE	
17. SECURITY CLASSIFICATION OF REPORT UNCLASSIFIED	18. SECURITY CLASSIFICATION OF THIS PAGE UNCLASSIFIED	19. SECURITY CLASSIFICA OF ABSTRACT UNCLASSI		20. LIMITATION OF ABSTRACT	

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