

PEM REGENERATIVE FUEL CELLS

Larry L. Swette, Anthony B. LaConti, Stephen A. McCatty
Giner, Inc.
14 Spring Street
Waltham, MA 02154-4497

ABSTRACT

This paper will update the progress in developing electrocatalyst systems and electrode structures primarily for the positive electrode of single-unit solid polymer proton exchange membrane (PEM) regenerative fuel cells. The work was done with DuPont Nafion 117 in complete fuel cells (40 cm² electrodes). The cells were operated alternately in fuel cell mode and electrolysis mode at 80°C. In fuel cell mode, humidified hydrogen and oxygen were supplied at 207 kPa (30 psi); in electrolysis mode, water was pumped over the positive electrode and the gases were evolved at ambient pressure. Cycling data will be presented for Pt-Ir catalysts and limited bifunctional data will be presented for Pt, Ir, Ru, Rh and Na_xPt₃O₄ catalysts as well as for electrode structure variations.

INTRODUCTION

In prior work [1, 2, 3], a large number of candidate bifunctional positive electrode catalysts were evaluated for chemical and electrochemical stability and for catalytic activity in 30% KOH at 80°C. As a result of this work, two potentially bifunctional catalyst systems were identified: 1) Na_xPt₃O₄ [4] and 2) metal/metal oxide combinations of Rh, Pt and Ir [3]. More recently the scope of the program was expanded to include development of bifunctional positive electrode catalysts for the regenerative PEM fuel cell, focusing on many of the same catalysts.

In order to alternate between oxygen reduction and oxygen evolution on the same electrode, in addition to bifunctional catalysis, it is necessary to have an electrode structure that can perform in both of these modes. Some of the effort on this program was directed to the development of such structures. The approach taken was to develop catalyst/binder compositions optimized separately at the particulate level for either oxygen reduction (more hydrophobic) or oxygen evolution (more hydrophilic) and then combine these at an optimal ratio in a single electrode, referred to as an "integrated dual-character" (IDC) electrode. In this approach the catalyst for each function can be the same material if it shows bifunctional activity (e.g., Na_xPt₃O₄), or two different monofunctional catalysts (e.g., Pt for O₂ reduction and IrO₂ for O₂ evolution). This approach, developed for the alkaline system, was extended to the PEMRFC by introducing a two-layer structure to achieve an appropriate interface to the more confined two-dimensional character of the proton exchange membrane.

EXPERIMENTAL

Catalyst Materials

Pt black, used in both the positive and negative electrode structures, was fuel cell grade material obtained from Engelhard Industries. Metal oxide catalysts were prepared by proprietary modifications of Adams-type fusions of the metal salts in a nitrate flux [5]; metal/metal oxide catalysts (Pt-MO_x) were prepared by co-fusion of the metal salts in a similar manner followed by electrochemical reduction of the PtO₂ to Pt, or were blends of separately prepared materials (Pt-Na_xPt₃O₄).

Na_xPt₃O₄ was typically prepared by firing mixtures of Na₂CO₃ and PtO₂ under oxygen at 650°C or higher for several hours. In most cases it was necessary to use a proprietary activation procedure to develop significant catalytic activity and acceptable conductivity.

The surface areas of all materials were determined by the BET nitrogen adsorption method using a Micromeritics Flowsorb II 2300 instrument. The electrical conductivities of materials were estimated by a 2-point method measuring the resistance of the powders under moderate compression [2]. The compositions of the metal oxides prepared by nitrate fusion were determined in earlier work by X-ray diffraction analysis. Iridium chloride fusions yield IrO_2 , which becomes slightly oxygen deficient IrO_{2-x} ($x \approx 0.05$) if electrochemically reduced. Rhodium chloride fusions yield RhO_2 . The ruthenium salt fusions yield RuO_2 , which also becomes oxygen deficient RuO_{2-x} ($x \approx 0.1$) if electrochemically reduced. The $\text{Na}_x\text{Pt}_3\text{O}_4$ preparations were also analyzed by XRD for chemical characterization and determination of phase purity. The sodium content in $\text{Na}_x\text{Pt}_3\text{O}_4$ can be determined from the unit cell parameter, a_0 :

$$x = (a_0 - 5.59)/0.11$$

Recent preparations of $\text{Na}_x\text{Pt}_3\text{O}_4$ have exhibited moderate surface areas and marginally acceptable conductivities without any post-synthesis treatment and have been tested in that form. Materials characterization data is summarized in Table I.

PEM Regenerative Fuel Cell Testing

A rechargeable fuel cell requires 1) a negative electrode that is active for both H_2 oxidation and H_2 evolution and 2) a positive electrode that is active for both O_2 reduction and O_2 evolution. The general approach to achieve bifunctional electrode structures was to partially distribute the charge and discharge functions to separate catalyst layers of a composite electrode each of which is optimized primarily for one function. A hydrophilic PEM-bonded layer supports the gas evolution functions; for oxygen evolution the selected catalyst, at a loading of 1-3 mg/cm^2 , was blended with 10-20 wt% of the ionomer and pressed onto the surface of the membrane. The gas-consumption discharge functions are supported by a separate more hydrophobic catalyst layer which was physically pressed against the PEM-bonded layer by the mechanical compression of the cell assembly.

Platinum is a good catalyst for the negative electrode reactions in both alkaline and acid electrolytes. We used the composite electrode structure for most of the PEM cell configurations. For the negative electrode the combination consisted of a platinum layer bonded directly to the PEM (optimized for hydrogen evolution) and a separate IDC electrode (optimized for hydrogen oxidation). The "free-standing" component of the composite electrode consisted of an IDC platinum composition bonded to wetproofed carbon paper.

The requirements for the positive electrode are more difficult to meet because of the scarcity of efficient bifunctional oxygen catalysts. In addition, a titanium current collector is required because carbon support materials are not stable under oxygen evolution conditions. Platinum is an excellent O_2 -reduction catalyst for the fuel-cell mode, but is relatively poor for oxygen evolution. IrO_2 is an excellent O_2 -evolution catalyst, but a poor O_2 -reduction catalyst. For electrolysis cells it is mixed or alloyed with Pt to improve the conductivity of the IrO_2 . The general approach was to use Pt black for the oxygen reduction function in the free-standing component and to incorporate the more favorable oxygen evolution catalyst in the PEM-bonded layer, with Pt added to improve conductivity. For baseline performance data we used Pt or Pt- IrO_2 as the oxygen electrode catalyst. Three baseline cell types were fabricated and tested: a Pt/Pt fuel cell (also tested as an electrolyzer), a Pt/Pt- IrO_2 electrolyzer, and several Pt/Pt- IrO_2 bifunctional cell configurations; subsequently, Pt/Pt- $\text{Na}_x\text{Pt}_3\text{O}_4$ bifunctional cells of similar construction were also tested [6]. More recently, as the oxygen evolution catalyst, we have evaluated Pt- RuO_x , RuO_x - IrO_x and RhO_2 , and we examined the performance of Pt- IrO_x and $\text{Na}_x\text{Pt}_3\text{O}_4$ with repeated cycling.

The baseline Pt/Pt fuel cell and the experimental bifunctional cells were fabricated with a Nafion 117 PEM and utilized composite electrodes (catalyst layer bonded to the PEM plus a free-standing IDC electrode) on both sides. The baseline fuel cell and all of the experimental bifunctional cells had the same negative (H_2) electrode configuration. On the oxygen side, the PEM-bonded electrode was Pt, Pt- IrO_2 , Pt- RuO_x , RuO_x - IrO_x , RhO_2 or Pt- $\text{Na}_x\text{Pt}_3\text{O}_4$, and the free-standing electrode was Pt or Pt- $\text{Na}_x\text{Pt}_3\text{O}_4$ on titanium mesh (Pt-plated). The baseline electrolysis cell had only the PEM-bonded electrode layers, optimized for gas evolution only (without ionomer

added). In electrolysis mode, the cells were run at ambient pressure with heated water pumped through the positive electrode chamber; in fuel-cell mode the gases were supplied at 207 kPa and the H₂ stream was humidified at 90-120°C. The majority of testing was done at a cell temperature of 80°C.

RESULTS

As described above, three different types of PEM cells were constructed and tested to establish a performance baseline for single-function cells (fuel cell or electrolyzer) and a bifunctional cell. Subsequently, bifunctional PEM cells with Pt-Na_xPt₃O₄ positive electrodes were fabricated and tested to determine the relative performance of this potential catalyst in acid electrolyte. More recently we have evaluated Pt-RuO_x, RhO₂ and RuO_x-IrO_x as the oxygen evolution components in bifunctional cells and investigated the performance of Pt-IrO_x and Pt-Na_xPt₃O₄ under limited cycling conditions.

Baseline Cells

Baseline Pt/Pt Fuel Cell. - In fuel-cell mode, this cell ran at 0.874 V at 100 mA/cm² and at 0.714 V at 500 mA/cm² at 80°C. As an electrolyzer, performance was poor, as expected, primarily because Pt is not a very good catalyst for O₂ evolution; 1.77 V at 100 mA/cm², 2.00 V at 500 mA/cm² at 80°C. This cell was provided with the same positive electrode structure as used in the bifunctional cells (an IDC platinum composition at a loading of 8-10 mg/cm² on a Ti current collector mesh) so that the electrolyzer performance of Pt could also be measured. More recently a non-reversible Pt/Pt cell, optimized for fuel cell operation, was built and tested. In this cell the anode and cathode were of the same composition and configuration; the PEM-bonded layers consisted of Pt at a loading of 1 mg/cm² with 10 wt% ionomer, and the free-standing electrodes consisted of an IDC composition of Pt at a loading of 4 mg/cm² on wetproofed carbon paper (Toray HO-30). The fuel cell performance of these two cells is compared in Figure 1. It can be seen that the cell optimized for fuel cell operation gave slightly better performance.

Baseline Pt/40Pt-60IrO₂ Electrolyzer. - This cell ran at 1.50 V at 100 mA/cm² and at 1.671 V at 500 mA/cm². This is fairly typical performance for a dedicated electrolyzer cell with Nafion and Pt/Pt-IrO₂ catalysts. This cell was not tested in fuel-cell mode. The performance is presented in Figure 2 in comparison to the baseline Pt/Pt fuel cell and the baseline Pt/Pt-IrO₂ bifunctional cell.

Baseline Bifunctional Pt/Pt-IrO₂ Cell. - Several Pt/Pt-IrO₂ cells with varying IrO₂ contents and electrode structures were evaluated initially to optimize bifunctional performance [6]. A Pt/40Pt-60IrO₂ cell (#5 in Tables 1 & 2) cell exhibited the best performance in both modes. At 500 mA/cm², this cell ran at 1.587 V in electrolysis mode and at 0.723 V in fuel-cell (#5) mode. The comparable voltages for the baseline cells are 1.671 V for the electrolyzer and 0.714 V for the fuel cell. The performance is presented in Figure 2 in comparison to the baseline Pt/Pt fuel cell and the baseline Pt/Pt-IrO₂ electrolyzer.

Catalyst Evaluation

Bifunctional Pt/Pt-Na_xPt₃O₄ Cells. - Two different bifunctional cells were constructed for the initial evaluation of Na_xPt₃O₄ as a component of the oxygen electrode catalyst. The first cell (#6, Na_xPt₃O₄ prep.#21, see Tables 1 & 2) used a two-layer oxygen catalyst structure bonded to the PEM consisting of very hydrophilic Na_xPt₃O₄ (10% ionomer) and a less hydrophilic Pt transition layer. The balance of the composite electrode was a free-standing IDC electrode consisting of a 33/67 mix of Na_xPt₃O₄ and Pt. This cell gave excellent fuel cell performance (0.74 V at 500 mA/cm²), as shown in Figure 2 and Table 2, better than the Pt/Pt cell (0.714 V) and the best Pt/40Pt-60IrO₂ cell (0.723 V). The cell performance in electrolysis mode was 1.697 V at 500 mA/cm², better than Pt/Pt (2.00 V), but not as good as the best Pt/40Pt-60IrO₂ cell (1.587 V).

The second Na_xPt₃O₄-containing bifunctional cell (#7, Na_xPt₃O₄ prep.#34A3) was constructed using a single layer of Na_xPt₃O₄ with 10% ionomer bonded to the PEM and the standard IDC Pt free-standing IDC electrode as the composite oxygen electrode. This cell showed reduced performance in both modes compared to

the first $\text{Na}_x\text{Pt}_3\text{O}_4$ cell (Figure 2 and Table 2). In the fuel-cell mode, the voltage at 500 mA/cm^2 was 0.657 V . In electrolysis mode, the voltage was 1.766 V at 500 mA/cm^2 , which is still a significant improvement over a pure Pt/Pt cell. The differences in performance between these two cells may be partly due to differences in catalytic activity since the materials, although very similar in physical characteristics, are from two separate and somewhat different preparations (see Table 1); a larger part of the differences, however, may be attributed to the changes in electrode structure in Cell #7 resulting in a higher resistance interface.

Pt/Pt-IrO₂. - Cell #8 was an attempt to repeat the build of the baseline bifunctional Pt/Pt-IrO₂ cell (#5) but with two slight differences: slightly more ionomer in the PEM-bonded layer and the introduction of a thin Pt transition layer between the bonded layer and the free-standing electrode. The cell had somewhat higher resistance and did not perform quite as well as Cell #5; the performance curves are shown in Figure 3.

RuO_x-IrO_x was used as the PEM-bonded layer for oxygen evolution in Cell #9 and the Pt transition layer was included for comparison to Cell #8. The electrolysis performance was better than for Cell #8 but the fuel cell performance was not as good (see Figure 3). It is anticipated that a structural reconfiguration of the cell duplicating Cell #5 would significantly upgrade fuel cell performance with this catalyst.

Pt-RuO_x was used as the PEM-bonded layer for oxygen evolution in Cell #10 and the Pt transition layer was again included for comparison. The performance was quite similar to the performance observed for RuO_x-IrO_x (see Figure 3).

RhO_x was used as the PEM-bonded layer for oxygen evolution in Cell #11 but the Pt transition layer was eliminated. This material was difficult to work with because of the very high surface area and more development would be required for a fair evaluation. Two performance points are shown in Table 2, but the cell was not very stable in either mode.

Repetitive Cycling Tests

Cells #12 and #13 were attempts to reproduce the electrode structures and performance of the baseline bifunctional Pt/40Pt-60IrO₂ cell (#5) for the purpose of investigating tolerance to repetitive cycling. The first build (#12) had somewhat higher than typical internal resistance and showed lower performance than Cell #5 (Table 2). The cell was subjected to six charge/discharge cycles at 500 mA/cm^2 , shown in Figure 4, after which it developed a leak and testing was terminated.

Cell #13 came somewhat closer in performance to Cell #5; in fuel cell mode it was not quite as good as Cell #5 but in electrolysis mode it was somewhat better (see Table 2). This build was subjected to fifteen charge/discharge cycles, at which point it also developed a leak. The performance at 500 mA/cm^2 , shown in Figure 5, remained fairly constant for both modes of operation over the fifteen cycles.

Cells #14, #15 and #16 were attempts to fabricate bifunctional positive electrodes similar to that used in Cell #5 but based on $\text{Na}_x\text{Pt}_3\text{O}_4$, also for the purpose of investigating tolerance to repetitive cycling. The first two builds (#14 & #15) did not give very stable performance and were not cycled (see Table 2). Cell #16 did not show any significant improvement in performance but was somewhat more stable (see Figure 6) and was subjected to five charge/discharge cycles. Neither the performance of Cell #5 nor the performance of the earlier $\text{Na}_x\text{Pt}_3\text{O}_4$ build (#6) was reproduced, but the cyclic performance was stable and better than Pt alone.

CONCLUSIONS

Regenerative PEM fuel cells showing efficient bifunctional performance can be fabricated with traditional Pt and Pt-Ir catalysts if the electrode structures are properly designed. This investigation has also identified RuO_x as a catalyst with good potential for improving the oxygen evolution component of a bifunctional positive electrode. $\text{Na}_x\text{Pt}_3\text{O}_4$ was identified in our earlier work as a catalyst with potential for bifunctional operation in

both alkaline and acid electrolytes; the current work has confirmed this with limited charge/discharge cycling but additional development will be required to achieve more reproducible catalyst properties and performance.

ACKNOWLEDGEMENTS

This phase of the work was supported by NASA Lewis Research Center under the direction of Dr. Patricia Loyselle on Contract No. NAS3-24635. We would also like to acknowledge the assistance of Prof. B.L. Chamberland of the University of Connecticut in the selection and identification of numerous materials tested in this program.

REFERENCES

1. L. Swette and J. Giner, *J. of Power Sources*, 22 (1988) 399-408.
2. L. Swette and N. Kackley, *J. of Power Sources*, 29 (1990) 423-436.
3. L. Swette, N. Kackley and S.A. McCatty, *J. of Power Sources*, 36 (1991) 323-339.
4. L. Swette and N. Kackley, "Bifunctional Alkaline Oxygen Electrodes," IECEC Conf. Proceedings, Boston, MA, Aug. 4-9, 1991.
5. R. Adams and R.L. Shriner, *J. Am. Chem. Soc.*, 45 (1923) 2171-2179.
6. L. Swette, N.D. Kackley and A.B. LaConti, "Regenerative Fuel Cells," Presented at the 27th Intersociety Energy Conversion Engineering Conference, San Diego, CA, Aug. 3-7, 1992

TABLE 1

PEMRFc positive electrode: composition and characteristics of PEM-bonded layer

Cell #	Catalyst	Ratio (wt %)	Loading (mg/cm ²)	Ionomer (wt %)	Surface Area (m ² /g)
5	Pt-IrO _x	40-60	1	10	104
6	Pt overlay Na _x Pt ₃ O ₄ #21*	33 67	1 2	0 10	28 12
7	Na _x Pt ₃ O ₄ #34A3*	100	2	10	7
8	Pt-IrO _x	40-60	1	15	117
9	RuO-IrO _x	50-50	2	25	114
10	Pt-RuO _x	40-60	1	15	69
11	RhO ₂	100	1	18	224
12	Pt-IrO _x	40-60	1	10	117
13	Pt-IrO _x	40-60	1	10	117
14	Pt black + Na _x Pt ₃ O ₄ #41*	33 67	1 2	10 10	29 18
15	Pt black + Na _x Pt ₃ O ₄ #43*	33 67	1 2	10 10	29 35
16	Pt black + Na _x Pt ₃ O ₄ #43*	33 67	1 2	10 10	29 35
* Na _x Pt ₃ O ₄ Characteristics					
Prep No.	Surface Area (m ² /g)	Conductivity (ohm-cm ⁻¹)	Unit Cell Parameter	Na _x -value	
#21	12	33	5.671(3)	0.74	
#34A3	7	30	5.674(1)	0.76 ("activated")	
#41	18	0.3	5.675(3)	0.77	
#43	35	3.4	5.688(1)	0.89	

TABLE 2

Cell voltage in fuel-cell and electrolysis modes for baseline cells and bifunctional PEM cells. 80°C, electrolysis at ambient pressure, fuel cell at 207 kPa.

Cell Performance in --->	Electrolysis Mode		Fuel-Cell Mode	
	100 mA/cm ² (V)	500 mA/cm ² (V)	100 mA/cm ² (V)	500 mA/cm ² (V)
Baseline Fuel Cells				
Pt/Pt (reversible)	1.770	2.000	0.874	0.714
Pt/Pt (fuel cell)	Not Tested	Not Tested	0.901	0.733
Baseline Electrolyzer				
Pt/40Pt-60IrO ₂	1.500	1.671	Not Tested	Not Tested
Bifunctional Cells				
#5 Pt/40Pt-60IrO ₂	1.456	1.587	0.901	0.723
#6 Pt/Na _x Pt ₃ O ₄ -Pt	1.545	1.697	0.903	0.740
#7 Pt/Na _x Pt ₃ O ₄ -Pt	1.621	1.766	0.839	0.657
#8 Pt/40Pt-60IrO _x	1.477	1.624	0.888	0.704
#9 Pt/50RuO _x -50IrO _x	1.406	1.627	0.778	0.565
#10 Pt/40Pt-60RuO _x	1.417	1.589	0.784	0.566
#11 Pt/RhO ₂	----	1.675	----	0.460
#12 Pt/40Pt-60IrO _x	1.489	1.620	0.858	0.640
#13 Pt/40Pt-60IrO _x	1.455	1.550	0.870	0.714
#14 Pt/33Pt-67Na _x Pt ₃ O ₄	----	1.873	0.873	0.698
#15 Pt/33Pt-67Na _x Pt ₃ O ₄	1.599	1.797	0.866	0.704
#16 Pt/33Pt-67Na _x Pt ₃ O ₄	1.618	1.799	0.838	0.597

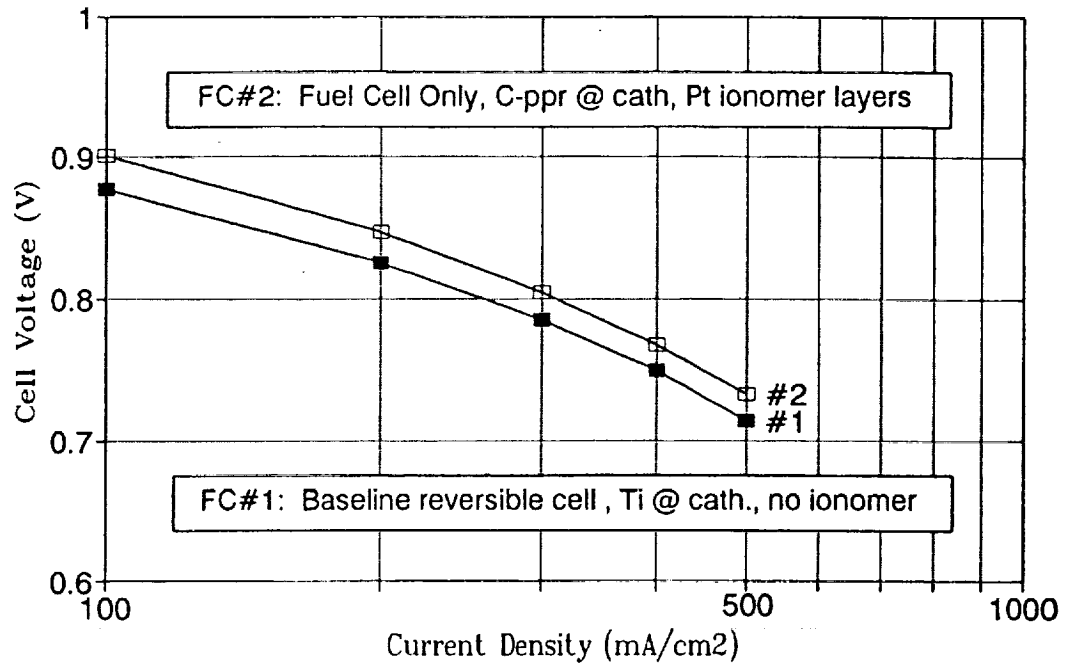


Fig. 1.—PEM Pt/Pt fuel cell performance (fuel cell mode only; 80°C, 30 psi).

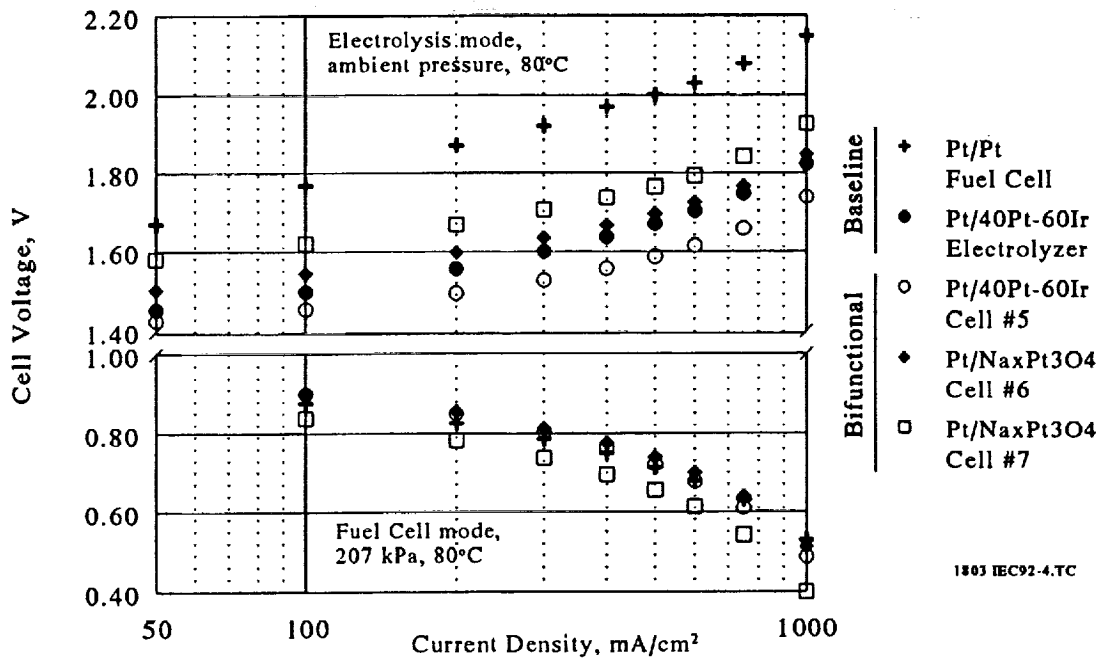


Fig. 2.—Performance of bifunctional Pt/Na_xPt₃O₄ PEM cells.

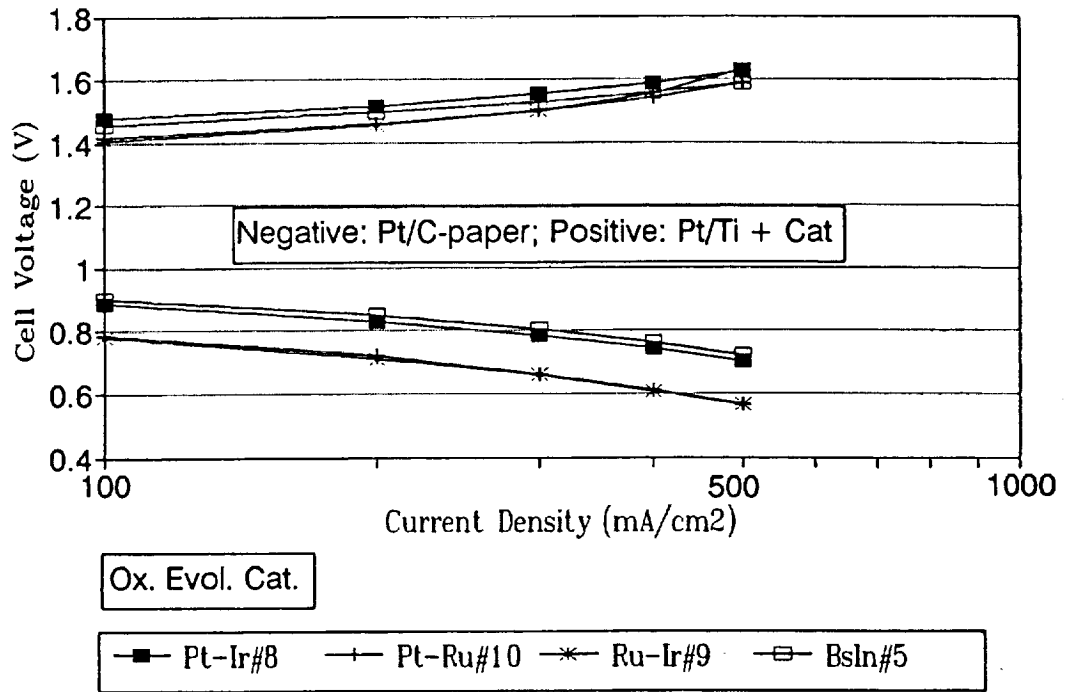


Fig. 3.—Bifunctional oxygen catalyst evaluation (Nafion 117, 80°C, 30 psi).

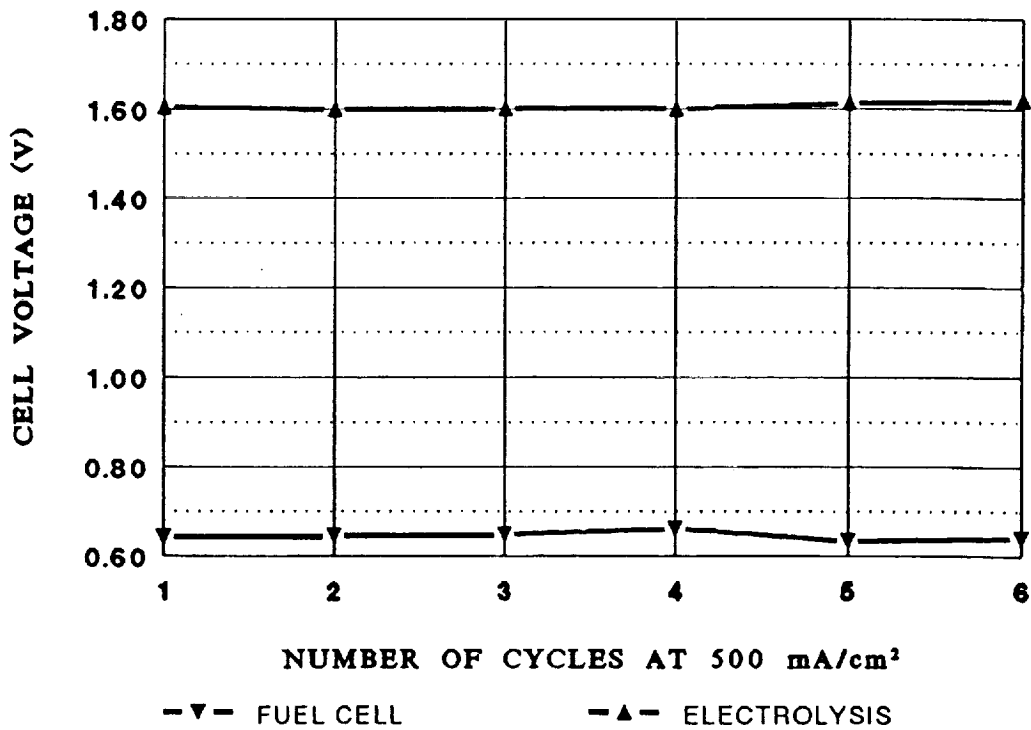


Fig. 4.—Cyclic performance of Cell #12 (Pt/Pt-IrO_x, 80°C).

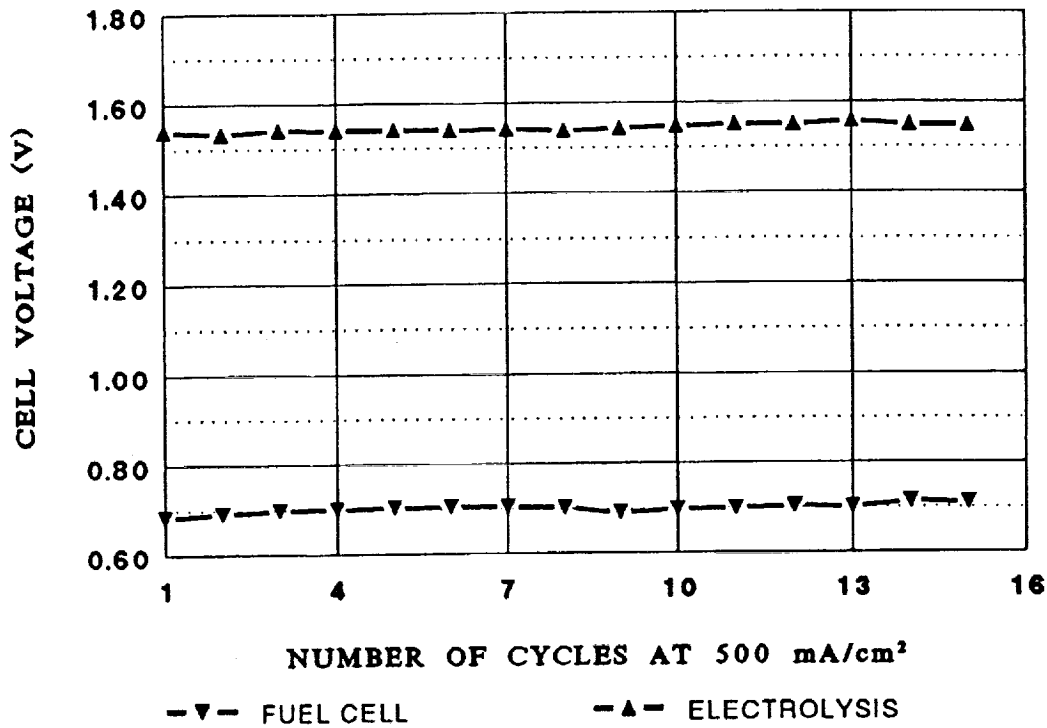


Fig. 5.—Cyclic performance of Cell #13 (Pt/Pt-IrO₂, 80°C).

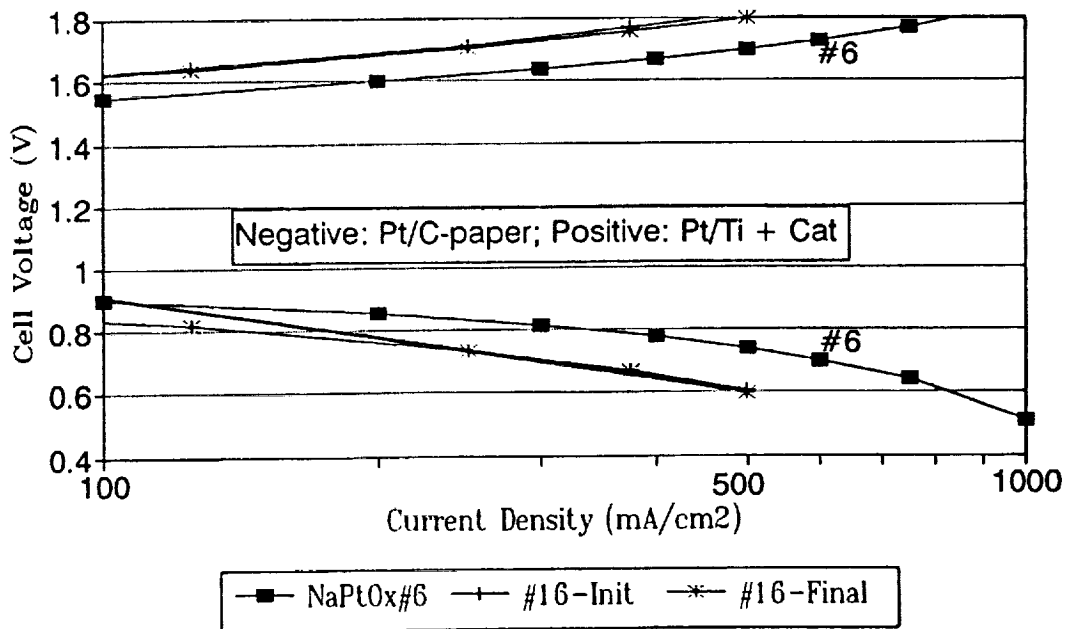


Fig. 6.—Cell #16 before and after five cycles (Nafion 117, 80°C, 30 psi).