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TEST EVALUATION OF

FUEL CELL CATALYSTS

bу

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Final Report

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#### SUMMARY

The objective of this work was to test materials prepared by the Pittsburgh Coal Research Center for NASA as anodic catalysts for ammonia in alkaline electrolytes and carbon monoxide in acidic and aqueons carbonate fuel cell electrolytes.

A total of fifty-four samples consisting of carbides, nitrides, borides, nitrocarbides, carbonitrides and Raney alloys of iron, cobalt, and nickel were tested. Corrosion potentials were run to determine usable electrolyte and catalyst combinations. Surface areas measured by double layer capacitance ranged from 27.1  $\text{Cm}^2/\text{gm}$  to 117,000  $\text{Cm}^2/\text{gm}$ . In general the areas were found to be: carbides > nitrocarbides > metals > borides and for the metallic moiety: silver > nickel  $\simeq$  cobalt >> iron.

Voltammograms at 25°C under approximately steady-state conditions were run in electrolytes compatible with the materials. Results showed the best materials were less than 2% of the activity obtainable with platinum for the same fuel. When ammonia was the fuel, activity followed the trend: carbides  $\cong$  nitrocarbides >> metals and nickel >> cobalt  $\cong$  silver >> iron. Carbon monoxide activity was generally nitrocarbides >> metals > carbides >> borides with silver > nickel > cobalt >> iron. With only one exception the sixteen best materials of the twenty-nine tested in acetate buffer contained nickel.

This work was carried out for the National Aeronautics and Space Administration with Mr. E. M. Cohn as Technical Monitor. Principal investigators were T. Webb and J. R. Moser.

#### I. INTRODUCTION

The high cost and scarcity of the noble metals prompted a search for cheaper, more abundant fuel cell catalysts. Initial results with Hagg carbide (Fe<sub>2</sub>C) indicated that iron group interstitial carbides might be the answer. Consequently, a total of fifty-four carbides, nitrides, nitrocarbides, carbonitrides, borides and Raney alloys synthesized by Bureau of Mines were tested in this laboratory. Nineteen iron compounds including four carbides, six nitrides, five carbonitrides and four nitrocarbides were tested for corrosion at 25°C in 30% KOH, saturated K<sub>2</sub>CO<sub>3</sub>, acetate buffer, 2N H<sub>2</sub>SO<sub>4</sub> and 85% H<sub>3</sub>PO<sub>4</sub>. Thirty-five cobalt and nickel materials consisting of nine metallic alloys, eleven carbides, ten nitrocarbides, and five borides were tested for corrosion at 25°C in 30% KOH, acetate buffer and 2N H<sub>2</sub>SO<sub>4</sub>. The surface area of each material was determined by double layer capacitance measurements (1).

Compatible catalyst-electrolyte combinations were further tested for anodic activity toward the oxidation of either ammonia or carbon monoxide. Equipment, testing procedures, treatment of catalysts and floating electrode preparation have all been previously described (2-4). All materials were tested during the first three quarters of this contract. No additional materials were received during the fourth quarter.

### II. RESULTS

Surface area measurements and the identity of each material tested are given in Table I. Voltammetric data for the oxidation of ammonia are shown in Tables II and III. Results for carbon monoxide oxidation are listed in Tables IV through VII.

#### III. DISCUSSION

### III.-1 Surface Areas

Double layer capacitance measurements on the 54 materials tested gave values for surface areas from 27.1  $\rm Cm^2/gm$  to 117,000  $\rm Cm^2/gm$ . Our platinum black measured was 210,000  $\rm Cm^2/gm$ .

As a group, the iron compounds exhibited the lowest area. The remaining materials which contained silver generally showed the largest areas. Surface areas according to type of non-ferrous material were usually in the order carbides> nitrocarbides> metals> borides.

#### III.-2 Ammonia Oxidation

Voltammograms for the oxidation of ammonia were run in 30% KOH for all materials stable in that electrolyte. Additionally the iron compounds which were stable in saturated  ${\rm K_2^{CO}_3}$  were tested in that electrolyte.

In 30% KOH the half-cell potential for the reaction was between 0.54 and 0.58 V vs DHF for most catalysts. This corresponds to the calculated value. Five of those tested, however, had lower half-cell potentials; one was higher. All six of these anomalies were due to corrosion of the electrode upon addition of ammonia. This was evidenced by the discolored electrolyte accompanying high exchange currents. The tafel slope for the oxidation with platinum is 0.04 volt while that for most of the other materials is around 0.12 volt. This may be explained by the mechanism of Owen and Salomon (5) who calculated Tafel slopes for the four steps in the reaction:

#### III. DISCUSSION (Cont'd)

				Calculated Tafel Slope
(i)	$NH_3 + M + OH$	<b>→</b>	$M-NH_2 + H_20 + e^{-}$	0.118 volt
(ii)	M-NH <sub>2</sub> to H		$M=NH + H_2^0 + e^{-}$	0.039 volt
(iii)	M=NH + OH	-	$M=N + H_2^0 + e^-$	0.024 volt
(iv)	2 M=N	<b>-</b>	$2M + N_2$	0.010 volt or ∞

Reaction (ii) is the rate determining step for platinum but reaction (i), the initial dissociative adsorption, seems to be the rate determining step for the other materials.

When saturated potassium carbonate was the electrolyte much lower currents, high Tafel slopes and variable half-cell potentials were obtained. These results were not readily interpreted. This electrolyte was abandoned after the first group of materials was tested.

In general for the oxidation of ammonia all of the materials tested were greatly inferior to platinum. However, those interstitial compounds containing carbon and/or nitrogen were much better catalysts than the pure metals or alloys. The order found for the catalysts activity of metallic constituents was Nickel >> Cobalt == Silver >> Iron.

#### III. DISCUSSION (Cont'd)

### III.-3 Carbon Monoxide Oxidation

Four electrolytes were used in this study. Currents were very low in 85%  ${\rm H_3PO_4}$  and saturated  ${\rm K_2CO_3}$ ; consequently, these systems were abandoned after the first group of materials was tested. The maximum current was obtained with platinum in  $2{\rm N}$   ${\rm H_2SO_4}$ . All other materials gave currents less than 2% of this value.

The oxidation reaction mechanism for carbon monoxide has not been well established. It is known that for platinum a reaction occurs around 0.26 volt versus DHE during which the electrode is poisoned by a reaction product. The major reaction occurs around 0.96 volt vs DHE where the product is carbon dioxide. For most of the catalysts tested the first reaction is predominant in 85%  ${\rm H_3PO_4}$  and acetate buffer while the second reaction is the major one in  ${\rm 2N}$   ${\rm H_2SO_4}$ . In  ${\rm K_2CO_3}$  the potentials are well scattered. Three Tafel slopes were observed in the acidic electrolytes: 0.040, 0.080 and 0.140 volt. However, the second was absent in  ${\rm H_3PO_4}$  while the third was absent in  ${\rm H_2SO_4}$ . The overall reactions seem to be:

(v) 
$$M + CO + OH^{-} \rightarrow MCO_2^{H} + e^{-}$$

(vi) 
$$MCO_2H + OH_{d1} \rightarrow M + CO_2 + H_2O + e^-$$

(vii) 
$$co_{ad} + 20H_{d1} \rightarrow co_2 + H_20 + 2e^{-}$$

#### III. DISCUSSION (Cont'd)

### III.-3 Carbon Monoxide Oxidation (Cont'd)

The nitrocarbides were much better catalysts than the pure metals or alloys while lowest in activity were the carbides and finally the borides.

The trend in catalytic activity was Silver > Nickel > Cobalt > Iron. Silver is known to strongly adsorb hydroxy ions; unoxidized nickel forms a carbonyl at room temperature -- nickel must catalyze reaction (v) since in acetate buffer with only one exception the sixteen best materials of the twenty-nine tested contained nickel. Two containing nickel and silver were actually better than platinum in this electrolyte although much lower than platinum in  $2N H_2 SO_4$ .

#### IV. CONCLUSIONS

The catalytic activity of all the materials tested was greatly inferior to the best results that could be obtained with platinum. For ammonia and carbon monoxide oxidation none of these proved to be a good platinum substitute. The results indicate that catalytic activity was limited by failure of these materials to adsorb the fuels. The reason is in some cases undoubtedly due to the presence of surface oxide (passivation) which also prevents corrosion of the metal by the electrolyte.

#### V. RECOMMENDATIONS

Since all of these materials were originally prepared for possible use as oxygen electrodes in alkaline electrolytes it would be fortuitous if any were found to be suitable as anodes in acid solutions. A thorough understanding of the fuel cell reaction mechanisms would be an extremely desirable asset in selecting potential catalysts. With this knowledge and a systematic approach as used by Bond (6), it should be possible to arrive at the optimum catalyst for a particular reaction under a fixed set of conditions.

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## TABLE I

		Surface Are	eas (cm²/gm)		
			Metals .	•	
RC-1 RN-2 RAL-6 RAL-8	Cobalt Nickel 3Ni/lAg 1Co/lAg	5,300 5,300 26,900 69,800	RAL-9 RAL-10 RAL-11	3Co/lNi 1Ni/lCo/lAg 1Ni/lCo/lAu	3,090 6,200 9,700
		Reduced Hy	droxides		
<b>5</b> 3R	lNi/lAg	32,500	61R	3Ni/1Co	121
		Carbi	ldes		
10C 12C 15C 23C 27C 29C 33C 35C	θ-Fe <sub>3</sub> C,α-Fe ε-Fe <sub>2</sub> C,α-Fe θ-Fe <sub>3</sub> C ε-Fe <sub>3</sub> C,α-Fe 3Ni/ICo 1Ni/3Ag Cobalt Nickel	33.5 32.9 30.0 27.1 7,060 117,000 4,920 54,400	39C 42C 43C 46C 53C 60C 61C	1Co/lAg 1Ni/lCo 1Ni/lAg 3Ni/lAg 1Ni/lAg 1Ni/lAg 1Ni/lAg/lAu 3Ni/lCo	20,200 10,400 13,650 4,050 1,870 3,000 403
		Carboni	itrides		
1CN 4CN 5CN	$\begin{array}{c} \varepsilon & -Fe \\ \varepsilon & -Fe \\ \varepsilon & -Fe \\ \end{array} \begin{array}{c} x \\ \varepsilon & -Fe \\ \end{array} \begin{array}{c} x \\ \end{array}$	46.0 156 54.2	7CN 8CN	€ -Fe <sub>2</sub> X, Ag € -Fe <sub>2</sub> X, 3Ag	90.4 83.8
	•	Nitri	ides		
1N 10N 18N	$\varepsilon$ -Fe <sub>2</sub> N, $\gamma$ -Fe <sub>4</sub> N $\varepsilon$ -Fe <sub>3</sub> N $\zeta$ -Fe <sub>2</sub> N, $\varepsilon$ -Fe <sub>3</sub> N	51.0 27.1 82.6	19N 20N 21N	$\xi$ -Fe <sub>2</sub> N, -Fe <sub>3</sub> N $\varepsilon$ -Fe <sub>3</sub> N $\gamma$ -Fe <sub>4</sub> N, -Fe <sub>3</sub> N	33.8 72.0 74.0
		Nitroca	arbides		
1NC 2NC 3NC 4NC 15NC 21NC 22NC	€ -Fe <sub>2</sub> X, Fe <sub>3</sub> 0 <sub>4</sub> X -Fe <sub>2</sub> X, € -Fe <sub>2</sub> X € -Fe <sub>2</sub> X € -Fe <sub>2</sub> X 3Ni/ICo Cobalt 3Co/1Ag	97.5 85.6 103 27.6 6,900 2,700 36,300	23NC 26NC 28NC 32NC 38NC 44NC 45NC	Nickel 1Co/lAg 1Ni/lCo 3Ni/lAg 1Ni/lAg 1Ni/lAg 1Ni/lAg 3Ni/lCo	19,800 45,500 29,200 78,400 37,400 48,000 4,900
		Воз	rides		
$   \begin{array}{c}     B6 \\     B7 \\     B9 \\     \hline     X = (C, \cdot)   \end{array} $	Nickel Cobalt 1Ni/1Co	565 700 1,100	B18 B20	1Ni/3Co 3Ni/1Co	6,900 2,650

TABLE II

Activity of Materials Toward NH $_3$  Oxidation in 30% KOH at 25 $^{\rm o}$ C

	io ma/gm	Tafel Slope <u>Volt</u>	E <sub>r</sub> Volt		io <u>Ma/gm</u>	Tafel Slope Volt	E <sub>r</sub>
Pt	883	0.04	0.58	RAL-6	1.78	0.14	0.54
33C*	106	0.04	0.22	20N	1.66	0.13	0.56
43C*	60.1	0.08	0.68	RAL-8	1.57	0.13	0.54
46C*	42.4	0.08	0.24	RN-2	1.42	0.12	0.54
28NC*	28.3	0.10	0.32	1NC	1.37	0.12	0.54
27C*	14.1	0.12	0.26	RC-1	1.35	0.12	0.54
12C	3.54	0.17	0.55	RAL-11	1.28	0.14	0.54
15NC	3.53	0.23	0.56	5CN	1.13	0.26	0.55
B6*	3.50	0.03	0.33	RAL-9	1.07	0.13	0.54
21N	3.15	0.12	0.56	4CN	0.99	0.11	0.56
61C	2.80	0.11	0.54	RAL-10	0.71	0.07	0.56
18N	2.47	0.12	0.56				

<sup>\*</sup> Dissolves in NH<sub>3</sub>

TABLE III

Activity of Materials Toward NH $_3$  Oxidation in Saturated  $\rm K_2^{CO}_3$  at  $\rm 25^{\circ}C$ 

	io ma/gm	Tafel Slope <u>Volt</u>	Er <u>Volt</u>		io ma/gm	Tafel Slope <u>Volt</u>	Er <u>Volt</u>
Pt	205	0.20	0.79	19N	1.20	0.26	0.09
1NC	2.40	0.56	0.16	4NC	0.885	0.85	0.13
2NC	1.87	0.27	0.20	21N	0.813	0.17	0.09
<b>1</b> N	1.66	0.70	0.12	5CN	0.777	1.40	0.20
1CN	1.55	0.36	0.17	12C	0.177	0.16	0.14
3NC	1.52	0.65	0.32				

	io ma/gm	Tafel Slope Volt	E <sub>r</sub> Volt		io ma/gm	Tafel Slope Volt	E <sub>r</sub> Volt
Pt	17.7	0.02	1.00	10C	0.254	0.49	0.04
2NC	1.59	0.11	0.23	15C	0.177	0.07	0.26
3NC	1.34	0.22	0.22	19N	0.177	0.11	0.25
10N	1.17	0.04	0.04	23C	0.152	0.14	0.26
21N	0.849	0.13	0.25	4NC	0.120	0.05	0.29
1N	0.814	0.49	0.98	12C	0.035	0.04	0.29
1CN	0.283	0.50	0.15				

	io ma/gm	Tafel Slope Volt	Er <u>Volt</u>		io ma/gm	Tafel Slope Volt	E <sub>r</sub> Volt
Pt	1.980	0.11	0.89	61C	3.5	0.04	0.05
1CN	36.4	2.20	0.50	B20	2.5	0.07	1.09
45NC	15.0	0.04	0.00	4NC	2.12	0.13	0.67
19N	12.7	0.22	0.96	RN-2	2.12	0.08	0.99
<b>53</b> C	8.8	0.07	0.42	60C	2.1	0.05	0.96
В9	5.7	0.15	1.09	12C	1.87	0.08	0.64
44NC	5.0	0.08	0.85	В7	1.8	0.07	1.02
10N	4.95	0.22	0.88	В18	1.4	0.03	1.07
100	4.60	0.60	0.58	1NC	1.31	0.30	0.88
61R	4.2	0.10	0.03	В6	1.0	0.05	1.06
20N	3.71	0.49	0.92				

.  $\underline{ TABLE \ VI}$  Activity of Materials Toward CO Oxidation in Acetate Buffer at  $25^{\circ}\text{C}$ 

	io ma/gm	Tafel Slope Volt	E <sub>r</sub> Volt		io ma/gm	Tafel Slope Volt	Er Volt
32NC	97.2	0.30	0.67	61R	2.8	0.54	0.46
RAL-10	95.4	0.25	0.58	60C	2.6	0.04	1.03
Pt	47.6	0.17	0.21	В7	2.0	0.05	1.02
RAL-11	25.4	0.70	0.48	18N	1.77	0.12	1.00
23NC	24.7	0.12	0.46	23C	1.76	0.09	0.36
43C	20.5	0.27	0.72	В6	1.2	0.08	1.06
В9	14.	0.02	0.42	1NC	1.13	0.18	0.98
10C	12.4	0.14	0.31	21N	1.13	0.16	1.03
<b>3</b> 5C	7.42	0.14	0.48	RC-1	1.13	0.12	1.06
46C	6.36	0.07	0.50	В18	1.1	0.07	1.03
61C	4.6	0.15	0.84	B20	0.88	0.04	1.04
RAL-9	4.24	0.16	0.73	4NC	0.85	0.18	1.04
RN-2	3.89	0.08	1.08	4CN	0.81	0.14	1.05
53C	3.5	0.03	0.48	12C	0.56	0.14	1.06
15NC	3.00	0.17	0.46	5CN	0.42	0.15	0.41

	io ma/gm	Tafel Slope <u>Volt</u>	Er <u>Volt</u>		io <u>Ma/g</u> m	Tafel Slope <u>Volt</u>	E <sub>r</sub> Volt
Pt	40.7	0.04	0.63	1NC	0.233	0.90	0.30
21N	0.530	0.14	0.40	12C	0.223	0.20	0.40
2NC	0.282	0.11	0.40	19N	0.212	0.33	0.18
1N	0.257	0.29	0.23	4NC	0.191	0.45	0.52
1CN	0.247	0.17	0.30	3NC	0.141	0.13	0.48
5CN	0.247	0.90	0.19				

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