

Task 3.10
Gas Separation and Hot-Gas Cleanup

Topical Report

Michael L. Swanson

November 1995

Work Performed Under Contract No.: DE-FC21-93MC30097

For
U.S. Department of Energy
Office of Fossil Energy
Morgantown Energy Technology Center
Morgantown, West Virginia

By
University of North Dakota
Grand Forks, North Dakota

DISTRIBUTION OF THIS DOCUMENT IS UNLIMITED

kg

MASTER

DISCLAIMER

This report was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor any agency thereof, nor any of their employees, makes any warranty, express or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or any agency thereof. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or any agency thereof.

This report has been reproduced directly from the best available copy.

Available to DOE and DOE contractors from the Office of Scientific and Technical Information, 175 Oak Ridge Turnpike, Oak Ridge, TN 37831; prices available at (615) 576-8401.

Available to the public from the National Technical Information Service, U.S. Department of Commerce, 5285 Port Royal Road, Springfield, VA 22161; phone orders accepted at (703) 487-4650.

Task 3.10
Gas Separation and Hot-Gas Cleanup

Topical Report

Michael L. Swanson

Work Performed Under Contract No.: DE-FC21-93MC30097

For
U.S. Department of Energy
Office of Fossil Energy
Morgantown Energy Technology Center
P.O. Box 880
Morgantown, West Virginia 26507-0880

By
University of North Dakota
Energy and Environmental Research Center
P.O. Box 9018
Grand Forks, North Dakota 58202

November 1995

ACKNOWLEDGMENT

This semiannual was prepared with the support of the U.S. Department of Energy (DOE) Morgantown Energy Technology Center Cooperative Agreement No. DE-FC21-93MC30097. However; any opinions, findings, conclusions, or recommendations expressed herein are those of the author(s) and do not necessarily reflect the views of the DOE.

TABLE OF CONTENTS

LIST OF FIGURES	ii
LIST OF TABLES	ii
1.0 INTRODUCTION	1
2.0 OBJECTIVES	1
3.0 ACCOMPLISHMENTS	2
3.1 Equipment Description and Operating Procedures	2
3.2 Experimental Results	6
4.0 FUTURE WORK	10
5.0 REFERENCES	15

LIST OF FIGURES

1	4-lb/hr CFBR	2
2	Zinc oxide sorbent vessel	4
3	Piping modifications on the 4-lb/hr CFBR	5
4	Permeate flow rate versus run time for Membrane 1 tested at various conditions on the CFBR	8
5	Permeate flow rate versus run time for Membrane 2 tested at various conditions on the CFBR	8
6	Permeate flow rate versus run time for Membrane 3 tested at various conditions on the CFBR	9
7	Comparison of H ₂ permeability for bottled gas and coal gas tests in the CFBR using Pd-Ag membrane	11
8	H ₂ permeability for bottled gas and coal gas tests in the the CFBR using palladium membrane as compared to Pd-Ag membrane baseline	11
9	ESCA analysis of Pd-Ag membrane with deposited layer on the feed gas side of the membrane	12
10	Detailed ESCA analysis of Pd-Ag membrane identifying deposited layer as elemental sulfur	12

LIST OF TABLES

1	Proximate and Ultimate Analyses of Wyodak Subbituminous Coal from the Powder River Basin Mixed with CaCO ₃	6
2	Operating Conditions from Product Gas Tests on Pd-Ag Membrane	7
3	Operating Conditions from Product Gas Tests Using Palladium Membrane	7
4	Gas Chromatograph Data from Bottled Gas Tests on Pd-Ag Membrane Including Corrected Data from Permeate Gas Stream	13
5	Gas Chromatograph Data from Bottled Gas and Coal Gas Tests on Pd-Ag Membrane Including Corrected Data from Permeate Gas Stream	13
6	Gas Chromatograph Data from Bottled Gas and Coal Gas Tests on Palladium Membrane	14
7	Separation Factors and Permeabilities of Corrected Permeate Gases Through Pd-Ag Membranes	14

TASK 3.10 GAS SEPARATION AND HOT-GAS CLEANUP

1.0 INTRODUCTION

Catalytic gasification of coal to produce H₂-, CO-, and CH₄-rich mixtures of gases for consumption in molten carbonate fuel cells is currently under development; however, to optimize the fuel cell performance and extend its operating life, it is desired to separate as much of the inert components (i.e., CO₂ and N₂) and impurities (i.e., H₂S and NH₃) as possible from the fuel gas before it enters the fuel cell. In addition, the economics of the integrated gasification combined cycle (IGCC) can be improved by separating as much of the hydrogen as possible from the fuel, since hydrogen is a high-value product. Researchers at the Energy & Environmental Research Center (EERC) and Bend Research, Inc., investigated pressure-driven membranes as a method for accomplishing this gas separation and hot-gas cleanup. These membranes are operated at temperatures as high as 800°C and at pressures up to 300 psig. They have very small pore sizes that separate the undesirable gases by operating in the Knudsen diffusion region of mass transport (30–50 Å) or in the molecular sieving region of mass transport phenomena (<5Å). In addition, H₂ separation through a palladium metal membrane proceeds via a solution-diffusion mechanism for atomic hydrogen. This allows the membranes to exhibit extremely high selectivity for hydrogen separation.

Technological and economic barriers to be resolved before these membranes are commercially viable include improved gas separation efficiency, membrane optimization, sealing of membranes to metal vessels, high burst strength of the membrane, pore thermal stability, and material chemical stability. Hydrogen separation is dependent on the temperature, pressure, pressure ratio across the membrane, and surface area of the membrane. For gas separation under Knudsen diffusion, increasing feed pressure and pressure ratio across the membrane should increase the gas permeability, while decreasing the temperature and decreasing the permeate-to-total flow ratio should also increase the gas permeability. In the molecular sieving regime of mass transport, the inlet pressure and pressure ratio should have no effect on gas permeability, while increasing temperature should increase permeability.

2.0 OBJECTIVES

The selectivity of the ceramic membranes for removing undesirable gases while allowing the desired gases to be concentrated in the permeate stream was to be determined. Specific questions to be answered included:

- What are the effects of membrane properties (i.e., surface area, pore size, and coating thickness) on permeability and selectivity of the desired gases?
- What are the effects of operating conditions (i.e., temperature, pressure, and flow rate) on permeability and selectivity?
- What are the effects of impurities (i.e., small particulate, H₂S, HCl, NH₃, etc.) on membrane performance?

3.0 ACCOMPLISHMENTS

3.1 Equipment Description and Operating Procedures

The most cost-effective way to perform the desired tests on the membrane was to conduct both the baseline separation efficiency tests using bottled gas and the impurity degradation tests using actual coal-derived fuel gas in the same equipment. Originally, it was proposed to build a separate test stand to conduct the separation efficiency tests using bottled gases. However, it was decided that this method would waste too many valuable resources duplicating similar capabilities. Modification of the EERC's continuous fluid-bed reactor (CFBR) represented the most cost-effective alternative.

Figure 1 shows the 4-lb/hr CFBR used for gasification tests with various coals. The unit was originally designed as a pyrolysis unit for a U.S. Department of Energy (DOE) mild gasification program, but it has been used since for gasification, pyrolysis, and combustion on a variety of projects. Although predominantly operated as a fluid bed, the unit has been operated as a fixed bed by using lower gas flow rates or larger particle sizes in the bed. The unit was intentionally designed to be flexible.

Gas used for fluidization is mixed in a gas manifold. Bottled gas, house nitrogen, house air, and any liquid desired (such as steam) are first preheated, then mixed, and heated to reaction temperature in a superheater (20 ft of 3/8-in. tubing coiled into an 18-in. ceramic fiber heater). Two bottled gases in combination with either house air or house nitrogen and a liquid can be used at the present time. Using more bottled gases in the feed gas mix would involve additional mass flow controllers.

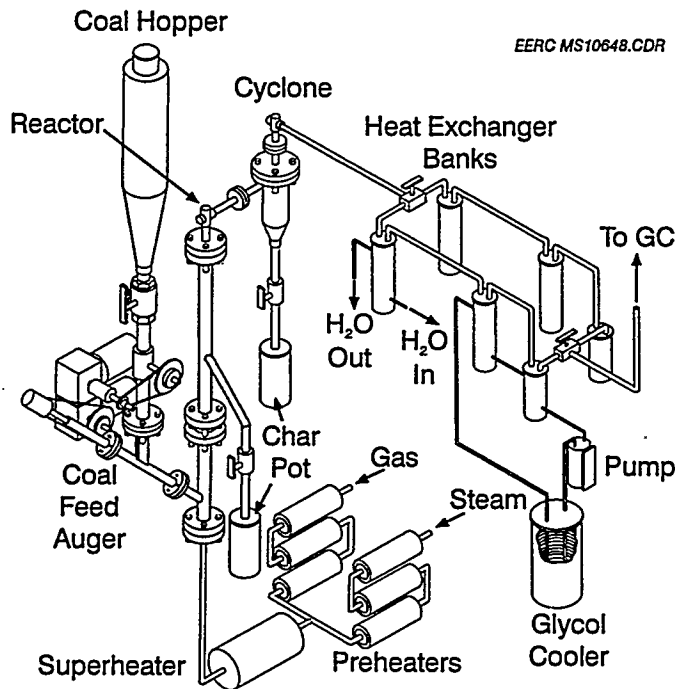


Figure 1. 4-lb/hr CFBR.

The reactor was constructed of 316H stainless steel Schedule 80 pipe. The first (bottom) section is made of 3-in. pipe and is 33 in. in length. The next (top) reactor section is made of 4-in. pipe, 18.75 in. in length. The two sections are connected with a 316H weld reducer. The unit was designed such that the top of the fluid bed lies 33 in. above the coal injection point. A solids offtake leg at the top of the bed is the primary means of solids removal from the reactor. A ball valve facilitates the collection of product while the system is operating.

The reactor has two ceramic fiber heaters to maintain the vessel's temperature and eliminate hot spots. Using external heaters allows the evaluation of internal and external heating methods for process development and scaleup. The reactor is capable of operating at a maximum of 155 psig and 840°C.

A 3-in.-diameter cyclone is used for solids removal from the gas stream. A ball valve allows the changing of the solids catch pot while the system is operating. The cyclone is heated with a ceramic fiber heater capable of operating at a temperature of 1650°F (900°C) and 200 psig. Figure 2 is a schematic of the 8-in.-long section of 2-in. 316H stainless steel Schedule 80 pipe constructed specifically for this project to be utilized as a pressure vessel to contain a fixed bed of approximately 1000 grams of zinc oxide sorbent to reduce the H₂S levels to less than 10 ppm and preferably to less than 1 ppm. Figure 3 is a drawing showing how the zinc oxide sorbent vessel, the H₂ separation membrane, a set of gas sampling lines, and ice-cooled knockout pots for removing moisture from the gas samples were installed on the CFBR.

Three 4-in.-diameter vessels are used to remove all condensables from the gas stream. Two separate trains were installed: one for mass balance sampling and the other for heatup, non-steady-state conditions, and cool down. The first condenser pot is indirectly cooled by water and typically cools the gas stream from 570°F (300°C) to 200°F (95°C). The next two condensers, also indirect, are glycol-cooled. The exit gas temperature is typically 50°F (10°C). A glass wool filter was used to capture aerosols passing through the condenser system.

A Genesis[®] software package is used for process control and data acquisition. Pressure drop across the bed is measured by two transmitters, and thermocouples throughout the unit measure temperatures. Temperature and pressure readings are recorded every 30 seconds, and these data are directly transferred to Lotus[®] spreadsheets.

Mass flow controllers for hydrogen, carbon monoxide, carbon dioxide, and nitrogen for supplying a clean gas stream from bottled gases were available for temporary use from other projects at the EERC. These mass flow controllers provided a clean gas stream to the CFBR. The CFBR was used as a mixing chamber and heater to generate hot gases for the initial bottled gas membrane tests. These shakedown tests were accomplished without the presence of impurities that would result from the gasification of coal in the CFBR. All gas analyses were performed on gas bag samples using a Hewlett-Packard 5880 gas chromatograph equipped with a refinery gas analysis package. The refinery gas analysis package provides an analysis equivalent to that specified by Universal Oil Products (UOP) Method 539-73. The gas chromatograph used thermal conductivity detectors and employs two carrier gas streams. Hydrogen and helium are separated and determined in an argon carrier gas stream. In an argon carrier gas, the response for hydrogen is linear over the entire concentration range from 0% to 100% hydrogen. The rest of the gas components are separated and determined in a high-purity helium carrier gas stream. The instrument is calibrated daily with

Gas Inlet

EERC MS11238.CDR

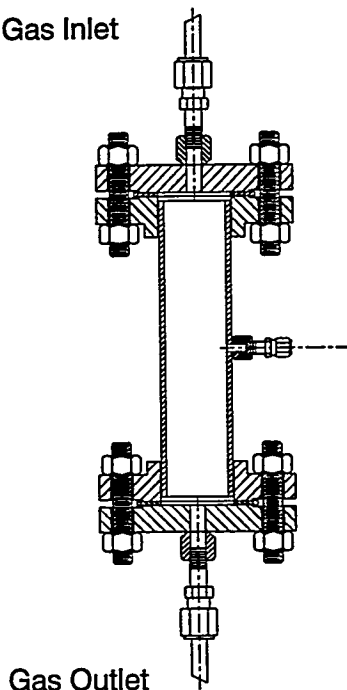
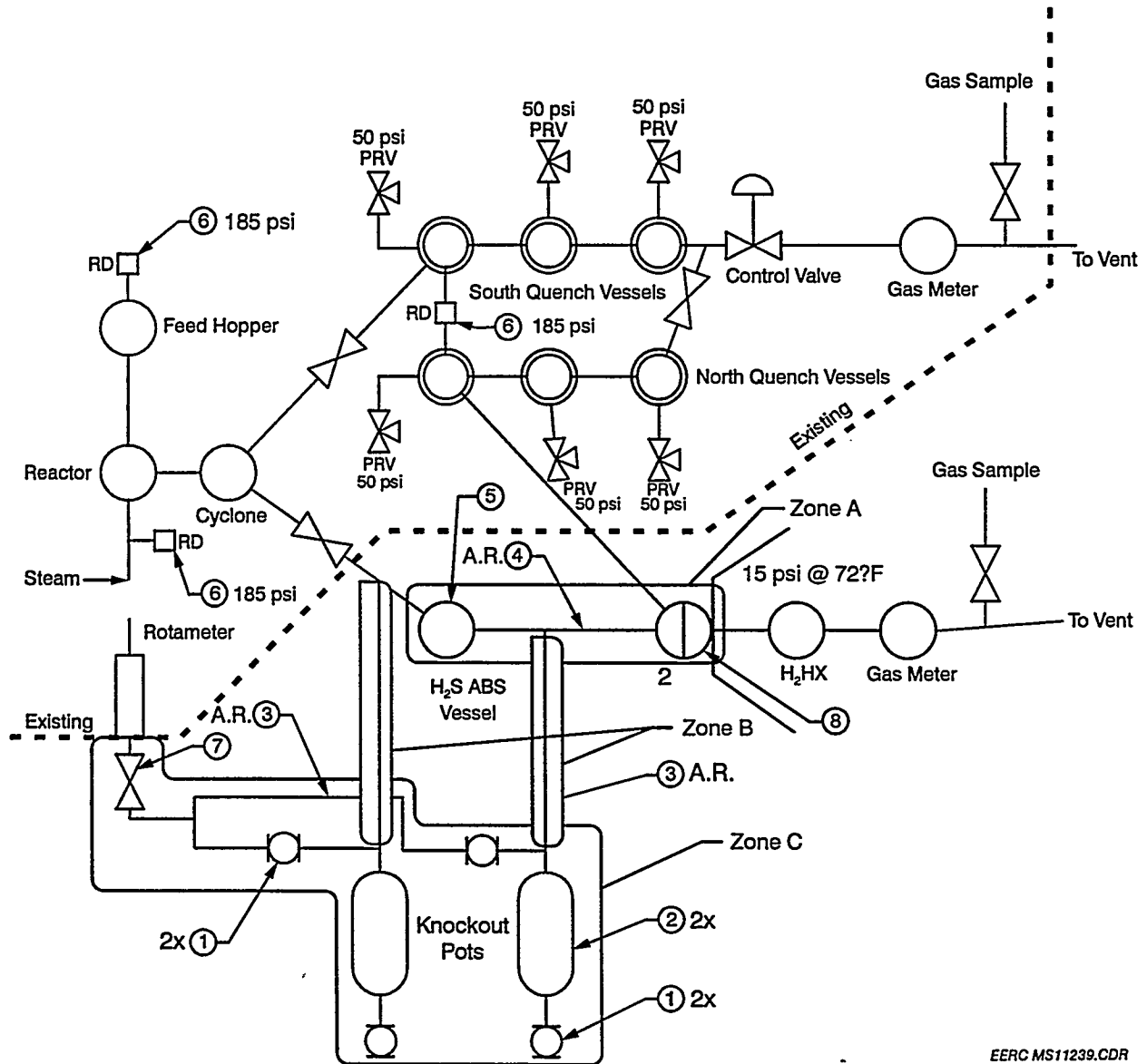


Figure 2. Zinc oxide sorbent vessel.

Certified Grade standard gas mixtures. The minimum detection limit (MDL) for the refinery gas analyzer package is 0.01 mole% except for CO_2 , CO , N_2 , O_2 , CH_4 , and H_2S , which all have a MDL of 0.02 mole%. The on-line gas analyzer available at the EERC was not capable of detecting H_2S (hydrogen sulfide) levels down to the low levels needed; however, a wet-chemistry method was used to determine the H_2S levels leaving the zinc oxide bed. This technique allows H_2S concentrations as low as 1 ppm to be measured, but requires a cubic foot of gas per hour to be bubbled through a solution that is then titrated to determine the concentration. This procedure results in a significant delay in obtaining H_2S concentration data.

Initial tests were performed to conduct short-term performance tests using the membrane to separate hydrogen from the mixed-gas stream. Subsequent tests then looked at the effects of H_2S impurities by injecting a known amount of H_2S into the gas stream. One operational problem encountered in performing the bottled gas tests was the formation of fine carbon particles from the reaction of carbon monoxide with itself to form carbon dioxide and carbon. This fine carbon blocked flow through the distributor plate of the CFBR and blocked the pressure taps on the CFBR, which resulted in shutting down the CFBR three times to either clear the distributor plate or the pressure taps. Further testing examined the effect of other impurities (such as H_2S , HCl , etc.) by operating the membrane using actual coal-derived fuel gases from the same CFBR equipment. The gas composition of the coal-derived gas stream was adjusted by adding some bottled hydrogen gas to the gas stream entering the reactor. Table 1 shows the results of proximate and ultimate analyses of the coal used to provide the coal-derived fuel gas for the testing of the membranes. The Wyodak subbituminous coal from the Powder River Basin was selected since its low sulfur and ash levels would allow the lower H_2S contaminant levels desired by the Bend Research, Inc., membrane. To



EERC MS11239.CDR

Figure 3. Piping modifications on the 4-lb/hr CFBR.

TABLE 1

Proximate and Ultimate Analyses of Wyodak Subbituminous Coal from the
Powder River Basin Mixed with CaCO₃

Wyodak Subbituminous	As-Run	Moisture-Free
Proximate Analysis		
Moisture	28.2	NA
Volatile Matter	31.22	43.48
Fixed Carbon	34.46	47.99
Ash	6.11	8.51
Ultimate Analysis		
Hydrogen	6.21	4.28
Carbon	47.08	65.57
Nitrogen	0.56	0.78
Sulfur	0.42	0.58
Oxygen, ind.	39.59	20.23
Ash	6.11	8.51
Heating Value, Btu/lb	7,953	11,077

reduce the H₂S levels further, solid CaCO₃ was added to the fuel at 3 wt% to capture as much of the sulfur as possible in the bed.

3.2 Experimental Results

Table 2 shows the operating conditions at which the bottled gas tests were performed on the 0.03-mm-thick Pd/Ag membrane. After the first membrane was exposed to H₂S, a new membrane was installed after Test M453 for the coal gas test.

Table 3 shows the operating conditions from product gas tests utilizing a 0.06-mm-thick palladium membrane using both a bottled gas baseline test and a coal-gas-derived baseline test, after which the membrane was exposed to 10 ppm H₂S to speed up the membrane deterioration process. Figures 4-6 show the measured permeate flow rates for the three membranes tested as a function of the total run time on each membrane. In these figures, every change in a run condition is indicated. From Figures 4 and 5, it appears that operating at higher temperatures increased the hydrogen permeability. Running at a higher partial pressure increased the permeate flow rate but did not increase the permeability of hydrogen, indicating that flow across the membrane is controlled by molecular sieving. The addition of H₂S (even at levels down to 1 ppm) to the bottled gas mixture significantly reduced the permeate flow rates. The initial gas flow rates used in these tests were comparable to those expected from a coal-derived gas stream. In an effort to save on bottled gas consumption and to determine the effect of product gas flow rate through the membrane, the product gas flow rate was reduced by approximately one-half after Test M451. The reduction of the flow rate of product gas through the membrane (Test 452) did not appear to change the hydrogen permeability from the 500° bottled gas baseline. However, it appears the possible shutdown of the product gas flow through the membrane for 3 hours to clear plugged pressure taps on the CFBR probably allowed the H₂S contaminant from Test M451 to be removed via some reversible equilibrium reaction. From these figures, it is apparent that as soon as the H₂S addition was stopped, the permeate flow rates would increase significantly.

TABLE 2

Operating Conditions from Product Gas Tests on Pd/Ag Membrane							
Run Number:	M449	M450	M451	M452	M453	M454	M455
Gas Source:	Bottle	Bottle	Bottle	Bottle	Bottle	Bottle	Coal Gas
Temperature, °C	400	500	500	500	500	500	500
H ₂ Part. Pres., psia	41.8	38.3	41.7	40.8	39.9	71.8	28.2
Inlet Gas Flow Rates, scfh							
H ₂	54	47.4	53	19.6	20.2	35	21
N ₂	74.2	71.2	72	27.8	29	17.5	132
CO ₂	21.2	21.2	21.2	8	8	6	NF ¹
CO	21.2	21.2	21.2	8	8	6	NF
H ₂ O	42.1	42.1	42.1	15.8	15.8	15.8	53.3
H ₂ S Conc., ppm	0	0	1	0	10	0	<5
Coal Feed Rate, lb/hr	0	0	0	0	0	0	1.5

¹Not fed.

TABLE 3

Operating Conditions from Product Gas Tests Using Palladium Membrane			
Run Number:	M456	M457A	M457B
Gas Source:	Bottle	Coal Gas	Coal Gas w/H ₂ S
Temperature, °C	500	500	500
H ₂ Part. Pres., psia	40.8	26.3	26.6
Inlet Gas Flow Rates, scfh			
H ₂	20.0	22	22
N ₂	29.0	135	135
CO ₂	8.0	0	0
CO	8.0	0	0
H ₂ O	15.8	49.4	49.4
H ₂ S Conc., ppm	0	<5	>10
Coal Feed Rate, lb/hr	0	1.5	1.5

While these flow rates would not return to their previous levels, they would increase by over a factor of two very rapidly, presumably as some of the H₂S contaminant on the surface of the membrane was reversibly removed via a reaction or just vaporized off as elemental sulfur. Approximately 5 hours into the coal-derived fuel gas test (Test M455) on the second palladium-silver (Pd-Ag) membrane, a couple of valves, which were used to take gas samples from either upstream or downstream of the zinc oxide sorbent bed, were accidentally left open at the same time, allowing some of the high-H₂S (\approx 280 ppm) fuel gas to bypass the zinc oxide bed. This reduced the permeate flow rate to near zero in a matter of a few minutes and it did not recover substantially on the coal gas after the valves were closed. Excluding the portion of the test where the zinc oxide bed was bypassed, the H₂S concentrations were determined to be less than 1 ppm during the early part of the test and had

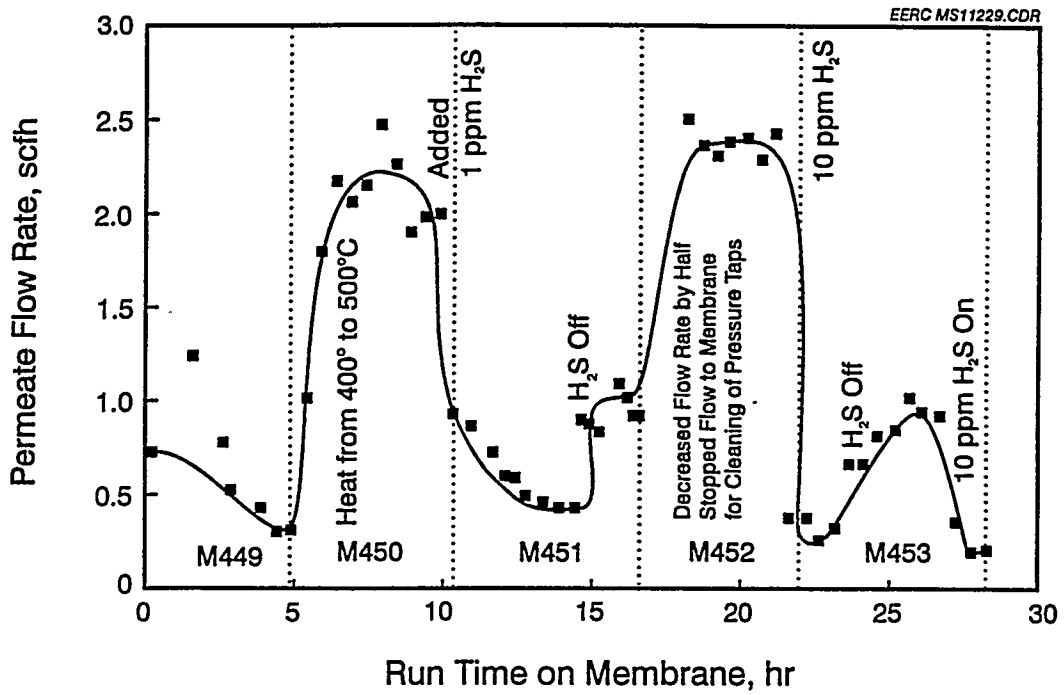


Figure 4. Permeate flow rate versus run time for Membrane 1 tested at various conditions on the CFBR.

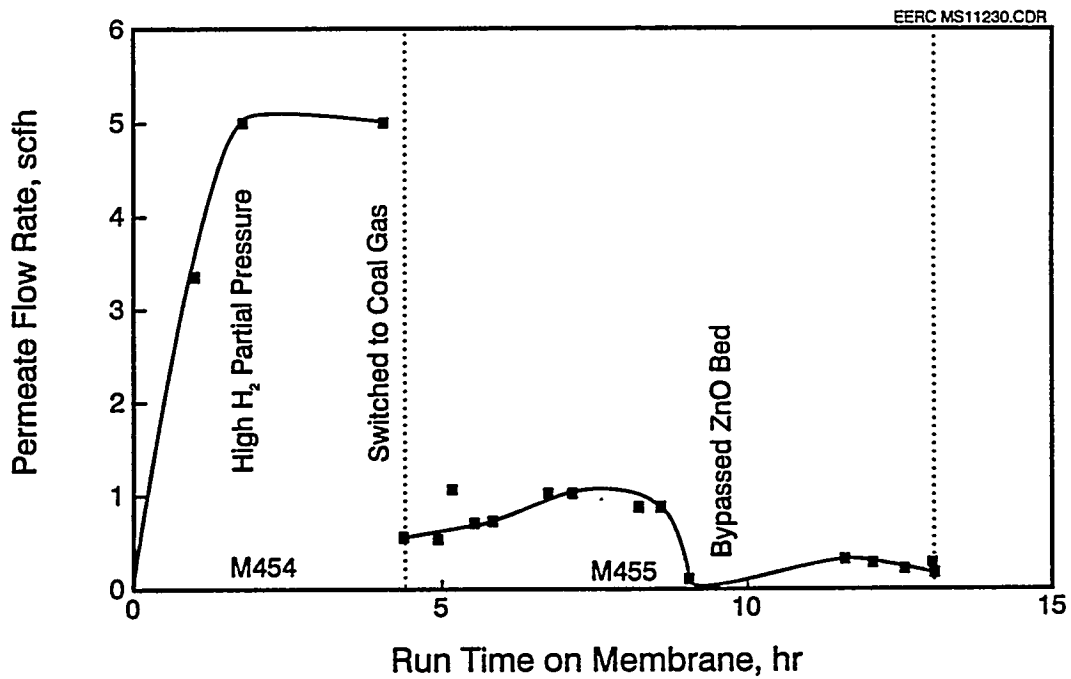


Figure 5. Permeate flow rate versus run time for Membrane 2 tested at various conditions on the CFBR.

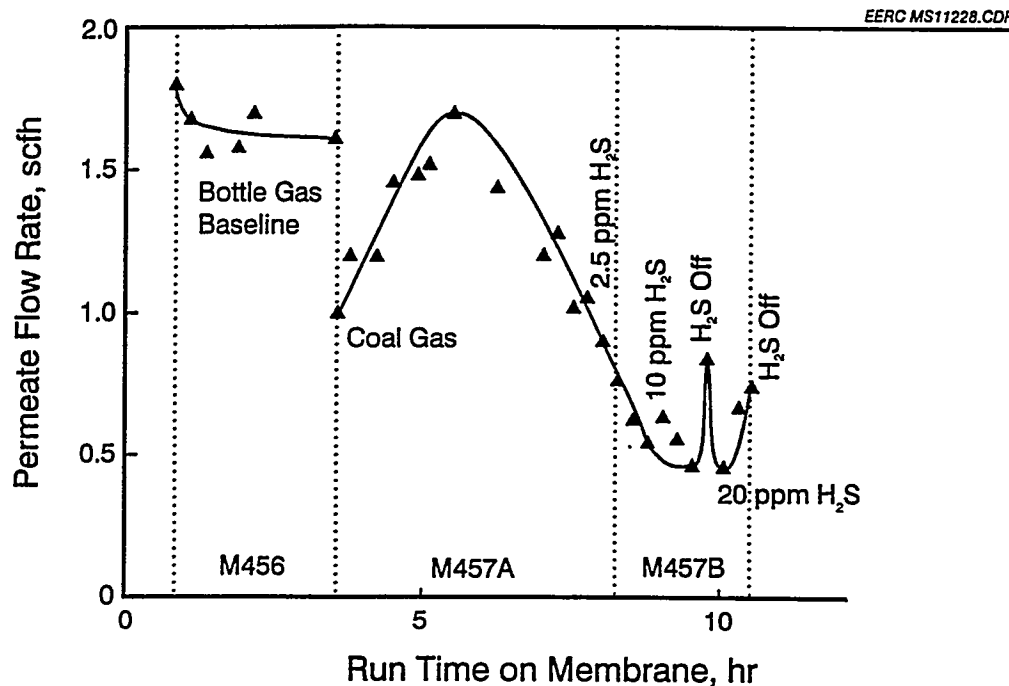


Figure 6. Permeate flow rate versus run time for Membrane 3 tested at various conditions on the CFBR.

increased to over 4 ppm by the end of the test on the membrane. The continued presence of H_2S in the fuel gas after the valves were shut probably limited the amount the membrane could recover.

The decrease in the baseline permeate flow rate for the third membrane is partially the result of the thickness (0.06 mm) of the all-palladium membrane as opposed to the two Pd-Ag membranes (0.03 mm). The Pd membrane was definitely affected by the presence of the H_2S in the product gas but did not appear to be as sensitive as the Pd-Ag membranes were.

Tables 4-6 show the gas analysis results of the feed gas, raffinate, and permeate streams taken for these tests. These tables also show the gas analysis results for the permeate gas stream after correcting for some dilution effects that resulted from the purging of the gas bags with nitrogen before they were evacuated with a vacuum pump. This correction assumed that the N_2 permeability would be equal to that of the CO (because of their similar molecular weights), and thus the ratio of N_2 to CO in the permeate gas stream should be the same as the ratio of N_2 to CO in the feed gas stream.

A common measure of membrane performance is its separation factor, which is defined in the following equation as (1):

$$\alpha_{ij} = \frac{Y_i/X_i}{Y_j/X_j} \quad [\text{Eq. 1}]$$

where α_{ij} = separation factor (dimensionless)
 Y_i = mole fraction of component i in permeate
 Y_j = mole fraction of component j in permeate
 X_i = mole fraction of component i in feed
 X_j = mole fraction of component j in feed

Separation factors for these product gas tests on the Bend Research, Inc., membranes are shown in Table 7. The permeability results have been reported as a permeate gas flow rate per unit component partial pressure drop across the membrane per unit membrane surface area (i.e., $\text{m}^3/\text{hr-atm-m}^2$). These results are also displayed in Table 7.

These values indicate that high separation factors can be achieved with these membranes, since the permeabilities of the undesired gases is very low. However, the permeability of hydrogen gas is also much lower than those reported for ceramic membranes (2).

Figures 7 and 8 show the hydrogen permeability for these tests as determined from the permeate flow rates and gas analyses. The hydrogen permeabilities again indicate how sensitive the palladium-based membranes are to hydrogen sulfide in the feed gas. The permeability also is significantly affected by membrane temperature or hydrogen partial pressure, but does not appear to be significantly affected by product gas feed rate through the membrane.

X-ray photoelectron spectroscopy was performed on a prototype hydrogen-permeable membrane produced by Bend Research, Inc. The membrane was exposed to hydrogen sulfide in the CFBR during a test to simulate coal gasification conditions. The objective of the analysis was to determine the composition of a thin white coating that was deposited on the retentate side of the membrane during the test run. In photoelectron spectroscopy, a sample is irradiated with photons of a known energy, which ionize the sample and result in the ejection of electrons. By measuring the kinetic energy of the electrons, we can determine their binding energies in the atoms from which they came. Since all elements have characteristic binding energies for electron shells, one can determine from which element an electron of a given energy came. Thus using this technique, we can determine the composition of an unknown. By looking at subtle shifts in binding energies produced in the chemical formation of compounds, we can also determine the chemical state of elements by this method.

For this analysis, a 1253.7 eV mg x-ray source was used at 10 kv and 350 watts power. A wide energy scan was performed using electron spectroscopy for chemical analyses (ESCA) to determine the elements present. Figure 9 shows this scan with the major elements identified. The elements found were palladium, silver, and sulfur. Since the membrane is a palladium-silver alloy, the deposit was assumed to be composed of a sulfur compound. A narrow energy scan, shown in Figure 10, was next performed to determine the state of the sulfur. The binding energies of the peaks were compared to those determined by Wagner and others (3). From these data, the deposit appears to be predominantly elemental sulfur emitting at 163.74 eV, with a small quantity of a sulfate-like compound emitting at 168.6 eV.

4.0 FUTURE WORK

This project will be continuing under the DOE Cooperative Agreement as Task 6.5 - Materials for Gas Separation and Hot-Gas Cleanup under the overall Task 6.0 - High-Temperature Materials program. The objective of this task will be to develop and test gas separation membranes to be developed with an EERC proprietary process. The remainder of this project was spent performing a literature review to ensure that this EERC process does not duplicate other techniques developed

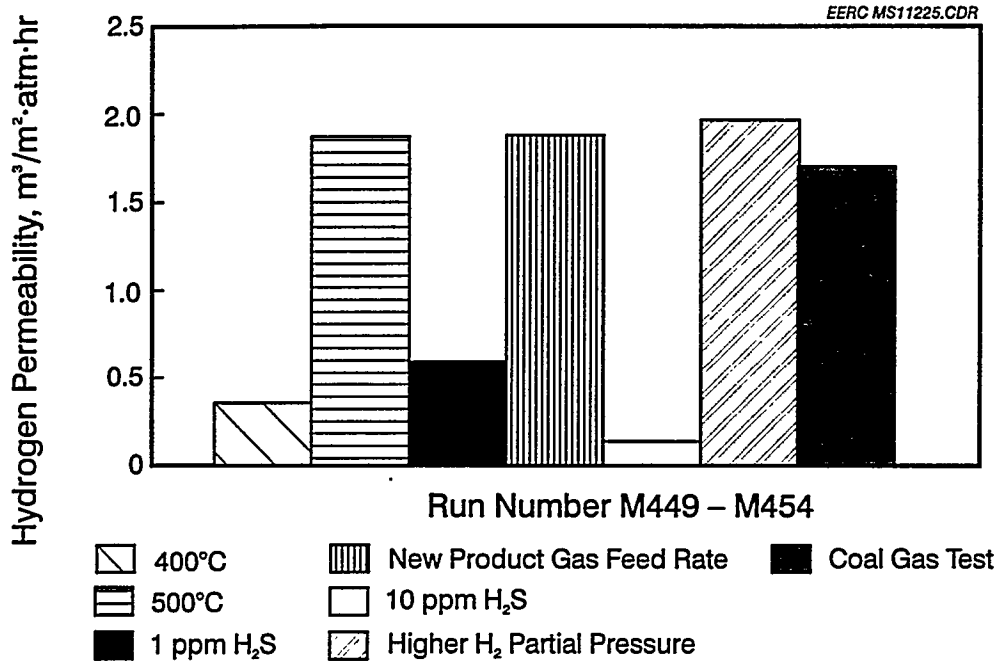


Figure 7. Comparison of H₂ permeability for bottled gas and coal gas tests in the CFBR using Pd-Ag membrane.

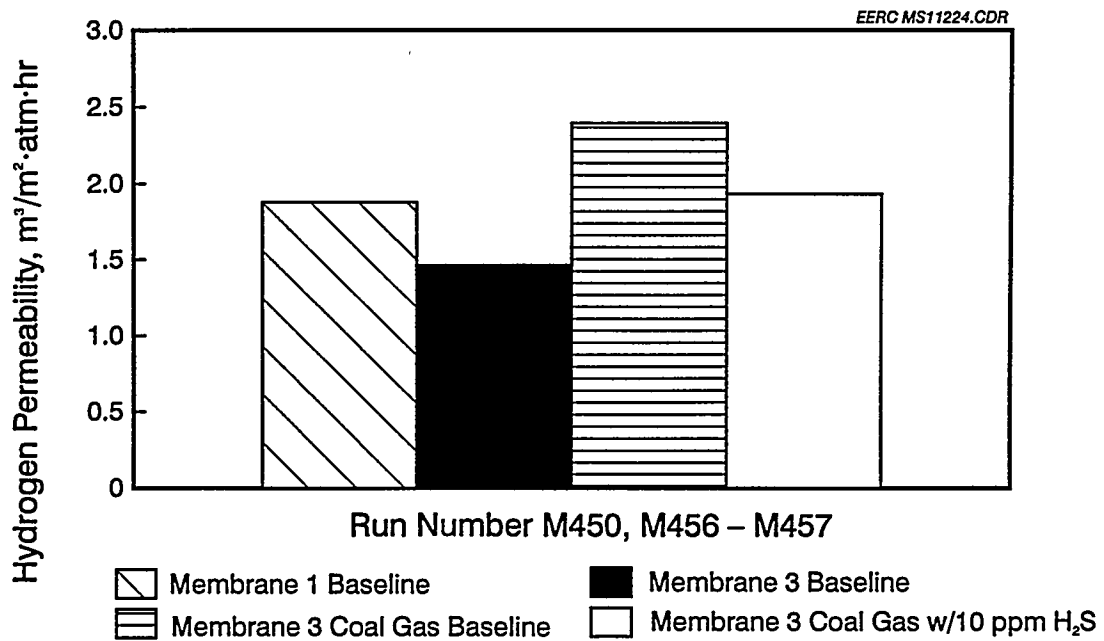


Figure 8. H₂ permeability for bottled gas and coal gas tests in the CFBR using palladium membrane as compared to Pd-Ag membrane baseline.

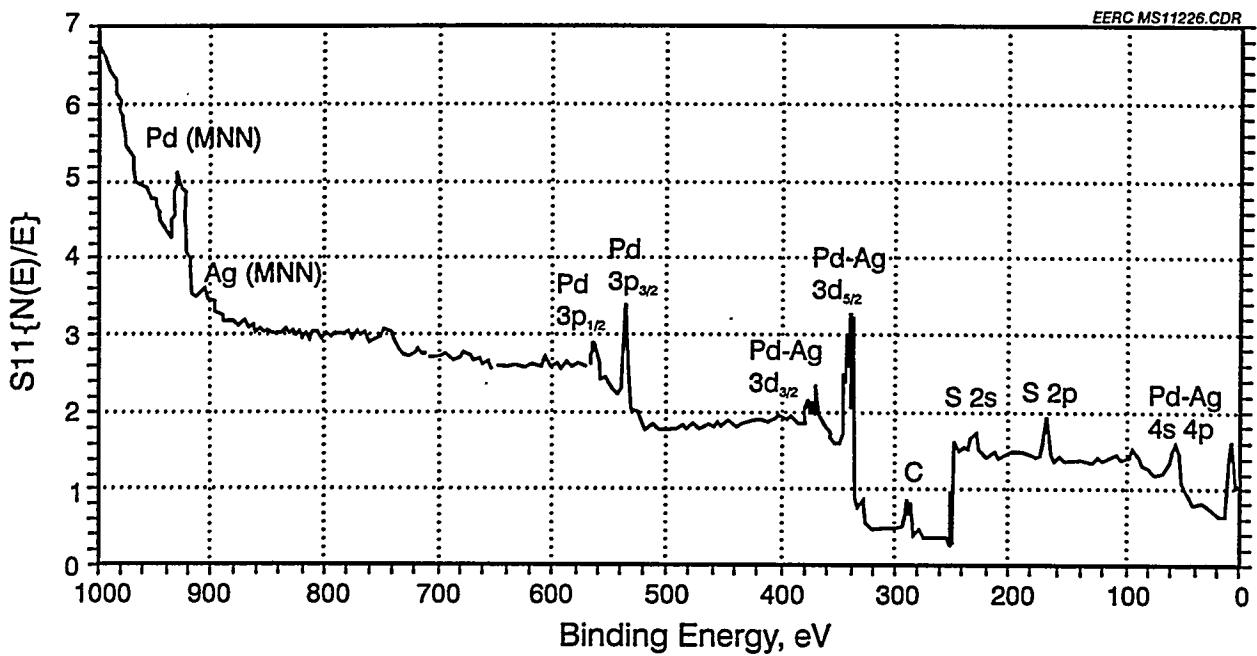


Figure 9. ESCA analysis of Pd-Ag membrane with deposited layer on the feed gas side of the membrane.

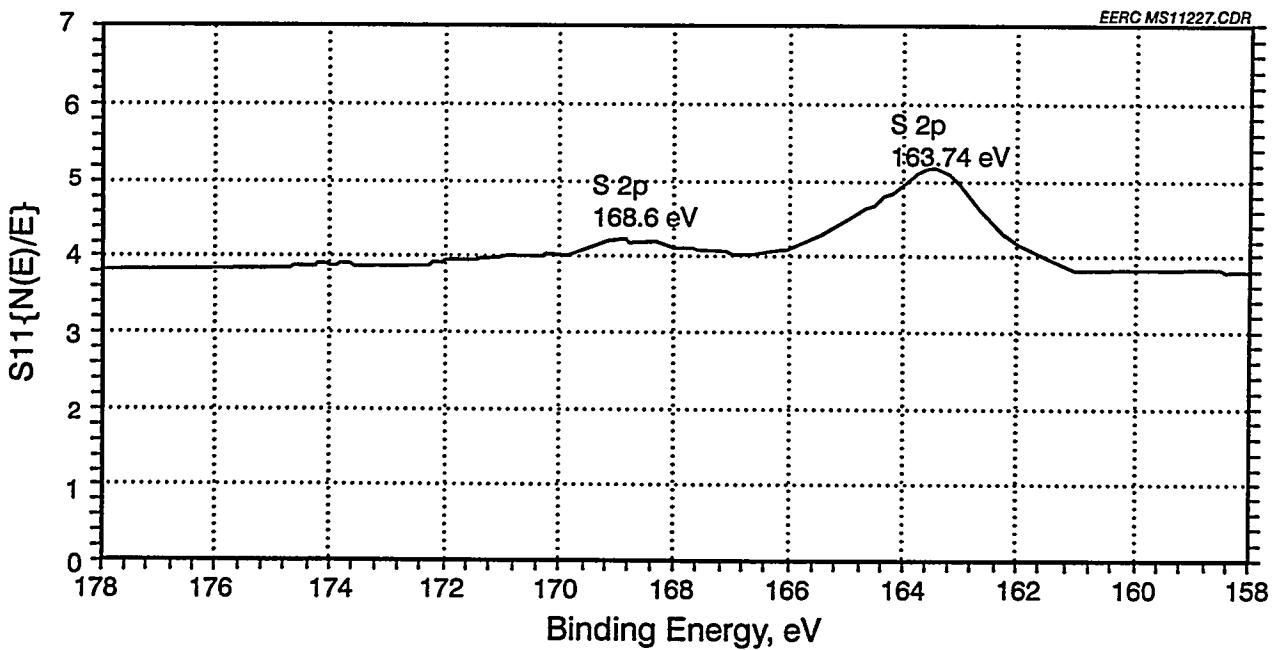


Figure 10. Detailed ESCA analysis of Pd-Ag membrane identifying deposited layer as elemental sulfur.

TABLE 4

Gas Chromatograph Data from Bottled Gas Tests on Pd-Ag Membrane Including Corrected Data from Permeate Gas Stream

Run Number Sample Type	M449		M450		M451		M452		M452		Cor.					
	Feed	Ref.	Feed	Ref.	Feed	Ref.	Feed	Ref.	Feed	Ref.	Per.	Cor.				
Gas Composition, mol%																
H ₂	32.6	31.4	84.6	86.3	32.6	30.3	98.7	99.7	32.0	32.1	97.4	98.8	30.1	30.8	90.2	94.0
N ₂	42.8	43.5	10.2	8.6	42.8	45.0	0.95	0.21	43.5	43.7	1.88	0.73	56.3	47.6	8.50	4.74
CO ₂	11.8	12.6	2.53	2.6	11.8	12.3	0.07	0.07	12.3	12.4	0.24	0.24	6.45	11.49	0.65	0.68
CO	12.4	12.1	2.44	2.5	12.4	12.0	0.06	0.06	12.0	11.7	0.20	0.20	6.93	.56	0.56	0.58
CH ₄	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.08	0.06	0.0	0.0
He	0.02	0.01	0.02	0.0	0.02	0.15	0.11	0.0	0.02	0.04	0.02	0.0	0.04	0.03	0.02	0.0
O ₂ /Ar	0.3	0.36	0.22	0.0	0.3	0.15	0.10	0.0	0.15	0.05	0.24	0.0	0.05	0.55	0.04	0.0
Total	100	100	99.9	100	100	99.9	100	100	100	100	99.9	100	99.9	100	100	100

TABLE 5

Gas Chromatograph Data from Bottled Gas and Coal Gas Tests on Pd-Ag Membrane Including Corrected Data from Permeate Gas Stream

Run Number Sample Type	M453A		M453B		M453B		M454		M454		M455	
	Ref.	Per.	Ref.	Per.	Ref.	Per.	Ref.	Per.	Ref.	Per.	Feed	Per.
Gas Composition, mol%												
H ₂	31.1	32.4	31.7	73.1	84.7	55.9	28.6	98.9	20.2	98.9	20.2	98.9
N ₂	44.9	49.1	48.8	20.2	9.26	27.6	44.8	1.09	73.0	1.09	73.0	1.02
CO ₂	12.0	9.77	10.8	2.90	3.36	9.37	16.3	0.0	4.40	0.0	4.40	0.0
CO	12.0	8.26	8.26	2.28	2.64	6.91	10.2	0.0	1.37	0.0	1.37	0.0
CH ₄	0.0	0.09	0.14	0.0	0.0	0.13	0.12	0.0	0.99	0.0	0.99	0.0
He	0.0	0.15	0.04	0.02	0.0	0.01	0.01	0.02	0.02	0.01	0.02	0.01
O ₂ /Ar	0.0	0.29	0.33	1.59	0.0	0.04	0.05	0.03	0.17	0.03	0.17	0.04
Total	100	100	100	100	100	100	100	100	100	100	100	100

Note: No correction was applied to M454 and M455 tests as no CO was detected in permeate gas stream.

TABLE 6

Gas Chromatograph Data from Bottled Gas and Coal Gas Tests on Palladium Membrane											
Run Number:	M456	M456	M456	M456	M457	M457A	M457A	M457A	M457B	M457B	M457B
Sample Type:	Feed	Raf.	Per.	Per.	Feed	Raf.	Per.	Per.	Raf.	Per.	Per.
Gas Composition, mol%											
H ₂	24.1	24.7	99.7	98.5	20.2	21.2	98.5	98.5	22.1	98.1	98.1
N ₂	54.5	48.9	0.21	1.25	73.0	71.8	1.25	1.25	69.3	1.39	1.39
CO ₂	7.23	9.54	0.0	0.01	4.20	4.44	0.01	0.01	4.77	0.07	0.07
CO	13.1	15.7	0.0	0.0	1.37	1.43	0.0	0.0	2.82	0.0	0.0
CH ₄	1.01	1.10	0.0	0.0	0.99	0.99	0.0	0.0	0.72	0.0	0.0
He	0.06	0.02	0.09	0.05	0.02	0.02	0.05	0.05	0.02	0.21	0.21
O ₂ :Ar	0.04	0.1	0.04	0.22	0.17	0.08	0.22	0.22	0.23	0.27	0.27
Total	100.0	100.1	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0

TABLE 7

Separation Factors and Permeabilities of Corrected Permeate Gases Through Pd-Ag and Pd Membranes											
Run Number:	M449	M450	M451	M452	M453A	M453B	M454	M455	M456	M457A	M457B
Sample Type:	Per.	Per.	Per.	Per.	Per.	Per.	Per.	Per.	Per.	Per.	Per.
Separation Factor ^a											
H ₂ to N ₂	13.2	742.9	182.5	37.1	14.2	15.2	45-∞	350-∞	1072-∞	284-∞	254-∞
H ₂ to CO ₂	12.1	630.7	156.4	29.7	11.2	10.3	∞	∞	∞	∞	∞
H ₂ to CO	13.2	735.8	182.5	39.1	13.2	13.1	∞	∞	∞	∞	∞
Permeability, m ³ /m ² -atm-hr											
H ₂	0.352	1.867	0.587	1.876	0.1716	0.135	1.940	1.656	1.456	2.392	1.932
N ₂	0.018	0.0016	0.0021	0.0446	0.0093	0.0073	0.034	0.0022	0.0014	0.0041	0.0037
CO ₂	0.019	0.0018	0.0024	0.0220	0.0110	0.0094				0.0006	0.0032
CO	0.018	0.0016	0.0020	0.0190	0.0093	0.0073					

^a Calculated from Eq. 1 in text where H₂ is component Y, and X is N₂, CO₂, or CO.

elsewhere. These membranes will be developed by using ceramic hot-gas particulate filters as a substrate on which to develop a thin-film membrane. After these membranes are developed and characterized, the two best membranes will be tested by mixing bottled gases in a manifold and feeding these gases to the membrane where gas composition and volumetric flow rates of the inlet, permeate, and raffinate gas streams will determine separation factors and permeabilities.

5.0 REFERENCES

1. Hackworth, J.H.; McIlvried, H.G. "Evaluation of the Use of Membranes in Power Generation Systems Involving Coal Gasification," final report for contract no. DE-AC21-90MC27346; Nov. 1992, p 195.
2. Lin, C.L.; Wu, J.C.S.; Gallaher, G.R.; Smith, G.W.; Flowers, D.L.; Gerdes, T.E.; Liu P.K.T. "Gas Separations Using Ceramic Membranes," report for U.S. Department of Energy Contract No. DOE/MC/25135-3341 (DE93000278), Feb. 1993, p 191.
3. Wagner, C.D. et al. *Handbook of X-Ray Photoelectron Spectroscopy*; Perkin Elmer, 1979; p 56.