TITLE: COST-EFFECTIVE METHOD FOR PRODUCING SELF SUPPORTED PALLADIUM ALLOY MEMBRANES FOR USE IN EFFICIENT PRODUCTION OF COAL DERIVED HYDROGEN

QUARTERLY TECHNICAL PROGRESS REPORT

REPORTING PERIOD START DATE: 9/09/03 (PROGRAM START)

REPORTING PERIOD END DATE: 01/31/05

PRINCIPLE AUTHOR(S): B. LANNING, J. ARPS

DATE REPORT WAS ISSUED: 01/28/05

DOE AWARD NUMBER: DE-FC26-03NT41849

SUBMITTING ORGANIZATION: SOUTHWEST RESEARCH INSTITUTE
6220 CULEBRA ROAD (78238-5166)
P.O. BOX 28510 (78228-0510)
SAN ANTONIO, TEXAS

OTHER TEAM MEMBERS: COLORADO SCHOOL OF MINES
IDATECH

SUBMIT TO: NETL AAD DOCUMENT CONTROL
BLDG. 921 U.S. DEPARTMENT OF ENERGY
NATIONAL ENERGY TECHNOLOGY LABORATORY
P.O. BOX 10940
PITTSBURGH, PA 15236-0940

APPROVED:

[Signature]

James H. Arps, Manager
Surface Engineering Section
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 or any information, apparatus, product, or process disclose, 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.”
ABSTRACT

Over the last quarter, we developed procedures for producing free-standing, defect free films using rigid silicon and glass substrates over areas up to 12 square inches. Since formation of contiguous Pd-Cu films in the 2-3 \(\mu\)m-thick range is ultimately governed by the size of the particle contamination on the supporting substrate surface, we have adopted techniques utilized by the semiconductor industry to reduce and eventually eliminate particle contamination. We have found these techniques to be much more effective on rigid substrates and have made a down select decision on removal methods (a key milestone) based on these results and the performance of membranes fabricated by this technique. The path to fabricating even larger membranes is straightforward and will be demonstrated in the coming months.

Hydrogen permeation tests were also conducted this quarter on as-deposited, Pd-Cu membranes, between 6-14 \(\mu\)m-thick. In the case of a 6 \(\mu\)m-thick film, the pure hydrogen flux at 20 psig and ~260 °C was 36 cm\(^3\) (STP)\(/cm^2\) min. This flux corresponds to a pure hydrogen permeability of \(7.4\times10^{-5} \text{ cm}^3 \text{ cm}^{-2} \text{ s}^{-1} \text{ cm Hg}^{1/2}\) at 250 °C. This value is within 20% of the pure hydrogen permeability at 250 °C reported in the McKinley patent. In the case of a 14 \(\mu\)m-thick membrane tested at 350 °C, the pure hydrogen flux, measured before initiating a pinhole-size leak, was \(2.1\times10^{-5} \text{ cm}^3 \text{(STP) cm/cm}^2 \text{ s cm Hg}^{0.5}\). This value is considerably lower than the expected permeability of Pd\(_{60}\)Cu\(_{40}\) materials at 400 °C.

To date, essentially all of the sputtered deposited Pd-Cu thin film membranes have had palladium compositions that were as much as 3% greater than the ideal 60 weight percent composition (this is a direct consequence of sputtering from a 60/40, Pd/Cu alloy target). As the concentration of Pd is increased beyond the optimum 60% value, a less desirable two-phase structure forms at the higher temperatures (in this case, above 260 –280 °C). As we continue development of procedures for producing thinner Pd-Cu films next quarter, we will also be optimizing alloy composition and corresponding hydrogen permeation flux as well.
# Table of Contents

DISCLAIMER.................................................................................................................III

ABSTRACT ..................................................................................................................... IV

TABLE OF CONTENTS ...................................................................................................V

1.0 EXECUTIVE SUMMARY ........................................................................................ 1

2.0 EXPERIMENTAL..................................................................................................... 1

3.0 RESULTS AND DISCUSSION ................................................................................ 1

   3.1 PROGRESS ........................................................................................................... 1

   3.2 PROBLEMS ENCOUNTERED: ............................................................................. 8

   3.3 PLANS FOR NEXT REPORTING PERIOD: ......................................................... 8

4.0 CONCLUSION ......................................................................................................... 8

5.0 REFERENCES.......................................................................................................... 9
1.0 EXECUTIVE SUMMARY

Refer to abstract.

2.0 EXPERIMENTAL

Pd-Cu alloy Vacuum Deposition

Flexible substrates – As part of a near term effort to reduce through-thickness defects in as-deposited membranes less than 5µm, techniques were investigated to modify and optimize the surface properties of the sacrificial polymer backing material(s). Since surface energy (chemical functionalization) and roughness of the polymer surface can have a dramatic effect on the agglomerization and wetting characteristics of a vapor deposited film, two approaches were selected to optimize surface properties of the polymer. First, testing of other polymers such as Kapton (polyimide), and PET were included in addition to PS and PVA (Solublon), to expand the range of polymer surface energies. Next surface pretreatments (modification) with an Ar/O₂ (95/5) plasma were investigated. Argon creates long-lived carbon radicals while oxygen creates functional groups, such as hydroxyl, carbonyl, or carboxyl that can reduce particulate adhesion and promote uniform deposition.

Both patches and continuous segments (11” wide and up to 48” in length) of the polymer films were treated/coated in the vacuum web coater (controlled drum temperatures between 0-60 °C) using either co-evaporation of Pd and Cu or magnetron sputtering from a composite target. Plasma treatments were conducted in-line with the Pd-Cu deposition steps.

Rigid substrates – Based on procedures reported last quarter (10/30/04), for depositing and releasing Pd-Cu films from rigid silicon substrates, we extended this development to include larger silicon and glass substrates. In brief, processing parameters were optimized to produce films with poor adhesion and minimal electrostatic interaction forces using magnetron sputtering from an alloy target.

H₂ Testing – A membrane foil is first sandwiched between two circular supports, such as alumina paper, and then sealed with either a Kalrez O-ring (max. use to 315 °C) or Grafoil packing material (allowing a 650 °C upper use temperature in oxygen-free environments) in the 25 mm Millipore membrane cell. The membrane is then checked with helium to confirm a tight seal and that the membrane is defect (pinhole) free. Subsequently, the membrane is heated to operating temperature to begin permeation testing.

3.0 RESULTS AND DISCUSSION

3.1 Progress

3.1.1 Optimization of Pd-Cu Membrane Formation

Flexible Polymer Substrates

Although procedures have been developed and routinely employed to deposit and release Pd-Cu films, 2-5 µm-thick and >75 in² in area from PS and PVA polymer backing materials, we
observe appreciable through-thickness defects (pinholes) for film thicknesses less than 6 µm. Though a small percentage of the defects are related to the deposition process (and therefore potentially eliminated), the majority of defects are due to characteristics of the sacrificial substrate.

In the case of flexible polymer substrates, the low surface energy (related to surface functional groups), roughness, and surface charge (electrostatic) can promote defect formation by methods such as surface particles/contamination, or agglomeration/clustering of incident atoms (as opposed to spreading (wetting) into a contiguous layer). Whether a particle or a cluster on the surface, at some point in the deposition, the incident atoms must bridge across these points to form a contiguous film. Although this “abridged” film is essentially contiguous, the coating will contain “through-thickness” defects (porosity) that compromise the mechanical integrity of the film; i.e., with only minor tensile stress, the reduced cross-section induced at the defect will break and form a pinhole. Hence, to form a <5 µm-thick, free-standing, gas impermeable membrane, the characteristics of the substrate must be such that vacuum deposited films must spread at thicknesses <<1 µm-thick.

Two treatment methods were investigated to promote spreading of the Pd-Cu films on the polymer backing materials; i.e. Ar/O₂ plasma and deposition of a precursor layer, such as SiOₓ with a more favorable surface energy. Although argon plasma treatment (with 5% O₂) was shown to decrease water contact angles (increase film wettability) on all plastics tested (PS, PVA, Kapton, and PET), we were not able to completely eliminate defects in films less than 5 µm. Subsequently, based on results presented in the previous quarter for SiO₂ on rigid silicon substrates, an evaporated silica (SiOₓ) layer was deposited on plasma treated polymer substrates. Unlike the case for rigid silicon substrates, the SiOₓ-coated polymer substrates did not yield films with a lower defect density. Therefore, irrespective of the surface treatment options tested (plasma or seed layer), we were not able to produce defect-free Pd-Cu films on flexible polymer substrates over large areas at thicknesses <5 µm (It is important to note however that defect-free films have been produced at thicknesses >6 µm).

**Rigid Substrates (silicon/glass)**

Since the primary cause of through-thickness defect formation in thin-films is due to surface particle contamination prior to deposition, we investigated methods, established by the semiconductor industry, to minimize and essentially eliminate effects from particle contamination. We continued using rigid silicon, in addition to including glass, in order to demonstrate the effect of particle contamination on the formation of defects in the films less than 5 µm-thick. During last quarter, we established vacuum deposition parameters and surface pretreatments of silicon and glass that have enabled us to produce defect-free, 5 µm-thick Pd-Cu films on 4” diameter silicon and 1” x 3” glass substrates. Two released Pd-Cu films are shown in Figure 1 below. These deposition procedures are currently being further optimized to produce defect-free films <3 µm - thick. Membrane samples on larger area rigid substrates up to 75 in² will also be prepared in the coming months.
Figure 1 Free-standing Pd-Cu Foils, 5 µm-thick; a) from rigid, 4” silicon wafer and b) glass slide.

3.1.2 H₂ Permeation Testing

Results from H₂ Permeation Testing (Sputtered films on silicon)

Permeation testing was completed on 6 and 14 µm-thick Pd-Cu films. Details from one of each of the two thicknesses are presented below.

Membrane 080304r1p4 (thickness ~ 6 µm) was sealed in the cell using a Kalrez® o-ring. This membrane exhibited a small helium leak that was attributed to Knudsen diffusion (Pure hydrogen flux vs. driving force measurements showed a pressure dependence of $\Delta P^n$ where $n$ was approximately 0.6), indicating that there is likely a pinhole in the membrane. All of the following hydrogen fluxes are corrected for the observed leak. The membrane was heated to 257 °C under helium, and hydrogen flow to the membrane was initiated after remaining at 257 °C for 12 hours. The hydrogen flow remained on, and the pure hydrogen flux at 20 psig was measured. The flux at these conditions slowly decreased from about 28 cm³/cm² min to about 20 cm³/cm² min over a period of 5 days. At this point, the flow to the membrane was switched from hydrogen to helium and left overnight with a feed pressure of 0 psig. After remaining under helium for approximately 15 hours, the membrane was tested again for pure hydrogen permeation at 20 psig. The pure hydrogen flux jumped to 36 cm³(STP)/cm² min. The pure hydrogen flux was measured again at 36 cm³/cm² min the following day after the membrane was exposed to helium again overnight. This flux corresponds to a pure hydrogen permeability of $7.4 \times 10^{-5}$ cm³ cm cm⁻¹ s⁻¹ cm Hg⁻¹/² at 250 °C. This value is within 20% of the pure hydrogen permeability at 250 °C reported in the McKinley patent.
The furnace temperature was then ramped to 275 °C. A temperature of 281 °C was measured using a thermocouple probe placed inside the cell, on the feed side of the membrane. A continuous flow of pure hydrogen was used on the feed side of the membrane for several days and the pure hydrogen flux at 20 psig decreased steadily to about 14 cm³/cm² min over this time period. The temperature was then ramped down again to 257 °C and the pure hydrogen flux did not recover. It dipped slightly lower to 13 cm³/cm² min. Repeating the procedure of leaving the membrane under helium at 0 psig overnight did result in a sharp increase in pure hydrogen flux back to 36 cm³/cm² min, which subsequently began to decrease with time.

One hypothesis for the above behavior is that oxygen had been diffusing back through the permeate lines of the system and oxidizing the surface of the foil. To test this hypothesis, the feed side of the membrane was exposed to helium at 20 psig for 15 hours and then hydrogen flux was measured. The pinhole leak ensured that there was helium flow in the permeate lines when the feed side was pressurized to 20 psig. This would limit oxygen from diffusing back through these lines. Figure 2 below shows the behavior of pure hydrogen flux at 20 psig for these conditions.

![Figure 2. Pure hydrogen flux at 20 psig after exposing to helium at 0 psig](image-url)

Figure 2. Pure hydrogen flux at 20 psig for membrane 083004r1p4 at 250 °C.
The above figure clearly shows the peak in hydrogen flux after exposing the membrane to helium at 0 psig and the decreasing trend from 35.9 cm$^3$ cm$^{-2}$ min$^{-1}$ to around 16.5 cm$^3$ cm$^{-2}$ min$^{-1}$, where the flux levels off. The enhanced hydrogen flux was not observed when the membrane was left under helium at 20 psig, instead, the flux decreased significantly. This supports our hypothesis that the enhanced hydrogen flux is a result of air oxidation of the foil. The flux recovered back to 33.7 cm$^3$ cm$^{-2}$ min$^{-1}$ when the membrane was exposed to helium at 0 psig for 15 hours, showing that this effect is reproducible.

Membrane 083104r1p2a (thickness ~14 µm) was placed into the Millipore fixture over a ceramic paper support and sealed using a graphite seal. The ceramic support ensured that there would be no intermetallic diffusion between the foil and the material that makes up the Millipore fixture. Furthermore, the graphite seal gave us the freedom to test at elevated temperatures.

The membrane was leak-free at room temperature and thus deemed acceptable for testing at high temperature. The system was brought to 250 °C and left under hydrogen. Pure-gas flux measurements were made with a hydrogen feed pressure of 20 psig. The hydrogen permeate flow rate showed a gradual decreasing trend from 22.3 cm$^3$/min to 11.4 cm$^3$/min over a period of 210 hours. The temperature was then raised to 350 °C to increase the rate at which any sort of phase change could be occurring. From this point on, the pure hydrogen flux began to increase. The system seemed to be approaching a steady state, with a pure hydrogen flux of 4.7 cm$^3$/cm$^2$ min and then the flux suddenly increased. Tests with helium gas revealed that a pinhole might have formed in the membrane, as there was a measurable helium flux of 0.7 cm$^3$/cm$^2$ min. Figure 3 shows the behavior of pure hydrogen flow-rate with respect to time at the two temperatures. The pure hydrogen flux measured before the leak corresponds to a hydrogen flux of $2.1 \times 10^{-5}$ cm$^3$(STP)$\cdot$cm$^2$•s•cm Hg$^{0.5}$. This value is considerably lower than the expected permeability of Pd$_{60}$Cu$_{40}$ materials at 400 °C.

![Figure 3. Influence of time on hydrogen permeation of membrane 083104r1p2a with a hydrogen feed pressure of 20 psig.](image-url)
A 25 µm thick palladium-copper foil known to be 60 wt.% Pd was supplied by Idatech and is being used as a standard for compositional analysis of subsequent membranes. A 25 mm disc of this material was tested for hydrogen permeation. The material was tested as received, with no prior activation. A steady-state measurement of pure hydrogen flux was performed and the hydrogen permeability at 400 °C was calculated to be $7.2 \times 10^{-5} \text{ cm}^3/(\text{STP}) \cdot \text{cm}^2 \cdot \text{s} \cdot \text{cm Hg}^{0.5}$. This value is nearly 50% of the maximum permeability reported in the literature for Pd$_{60}$Cu$_{40}$ materials. Upon examining the foil after testing, it appeared that the surface had significant carbon contamination that may have come from the graphite seal used in the membrane holder. See Figure 4 for an image of the surface of the membrane. This apparent carbon contamination could explain why the permeability observed is lower than that reported in the literature.

![Image](image.jpg)

Figure 4. Image of Pd$_{60}$Cu$_{40}$ standard foil showing the apparent carbon contamination on the surface that appeared after testing.

**Effect of Membrane foil Pre-treatment (25 µm-thick foil)**

An experiment was performed to determine the crystal structure of membrane 090104r1p2. The foil was annealed in nitrogen for 7 days at 450 °C, and then rapidly quenched in an ice bath. The surface of the foil appeared slightly oxidized. Figure 5 shows the XRD pattern of this foil after annealing. This pattern suggests that the crystal structure is fcc and the foil is wholly in the α-phase. This is not consistent with the Pd-Cu phase diagram for a composition near 60 wt. % Pd. We expected to see peaks corresponding to both the α and β phases. EDS analysis of this sample showed a composition of 68 wt. % Pd, which is more consistent with the observed XRD pattern.

The composition of sample 083004r1p4 was determined, using EDS, to be 63 wt.% Pd and 37 wt.% Cu. Table 1 below shows the EDS analysis of this membrane as well as a 60 wt.% Pd and 40 wt. % Cu standard. The results at different spots were averaged and then normalized against the standard.
Figure 5. XRD pattern of membrane 090104r2p4 annealed at 450 °C.

Table 1. EDS analysis of membrane 083004r1p4 and Pd$_{60}$Cu$_{40}$ standard from Idatech.

<table>
<thead>
<tr>
<th>EDS Compositional Analysis</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Spot</td>
<td>Pd wt. %</td>
<td>Cu wt. %</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>60.71</td>
<td>39.29</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>60.54</td>
<td>39.46</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>61.97</td>
<td>38.03</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>59.19</td>
<td>40.81</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>59.32</td>
<td>40.68</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>61.60</td>
<td>38.40</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>61.14</td>
<td>38.86</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>59.40</td>
<td>40.60</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>59.37</td>
<td>40.63</td>
<td></td>
</tr>
<tr>
<td>Pd$<em>{60}$Cu$</em>{40}$ Standard</td>
<td>62.45</td>
<td>37.45</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>63.14</td>
<td>36.86</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>65.27</td>
<td>34.73</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>63.33</td>
<td>36.67</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>62.30</td>
<td>37.70</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>63.22</td>
<td>36.78</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>63.57</td>
<td>36.43</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>63.57</td>
<td>36.43</td>
<td></td>
</tr>
<tr>
<td>083004r1p4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>63.33</td>
<td>36.66</td>
<td>99.99</td>
</tr>
<tr>
<td>5</td>
<td>62.95</td>
<td>36.99</td>
<td>99.94</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Average Pd</th>
<th>Average Cu</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard</td>
<td>60.36</td>
<td>39.64</td>
</tr>
<tr>
<td>083004r1p4</td>
<td>63.33</td>
<td>36.66</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Standardized Pd</th>
<th>Standardized Cu</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard</td>
<td>60.00</td>
<td>40.00</td>
</tr>
<tr>
<td>083004r1p4</td>
<td><strong>62.95</strong></td>
<td><strong>36.99</strong></td>
</tr>
</tbody>
</table>
The composition determined using EDS is reasonable as it corresponds well to the measured pure hydrogen flux. The steady state, pure hydrogen flux of membrane 083004r1p4 at 250 °C was determined to be $3.3 \times 10^{-5} \text{cm}^3(\text{STP})\text{cm}^2\text{s}^{-1}\text{cm Hg}^{0.5}$. The McKinley patent (USP 3,439,474) reports the pure hydrogen flux at 350 °C for a membrane that is 62.5 wt.% Pd and 37.5 wt.% Cu as well as pure hydrogen fluxes for a 60 wt.% Pd foil at various temperatures. Using data from the patent and assuming that the activation energy for a 62.5 wt% Pd foil is the same as that of a 60 wt.% foil, we can estimate the flux for the 62.5 wt.% foil at 250 °C to be approximately $5.2 \times 10^{-5} \text{cm}^3(\text{STP})\text{cm}^2\text{s}^{-1}\text{cm Hg}^{0.5}$. This value compares reasonably well to the hydrogen permeability of membrane 083004, particularly considering that the composition is even further from the 60 wt.% Pd target so a lower permeability is expected.

3.2 Problems Encountered:

Although we have established procedures and demonstrated the ability, using these procedures, to produce “free-standing” Pd-Cu alloy films over large areas, we have not been able to produce pinhole-free films less than 6 µm-thick using flexible polymer substrates. With our current efforts to address surface particle contamination and demonstrated ability to produce thinner membranes using flat silicon and glass substrates, we plan to apply these results to larger area rigid substrates as well as a limited series of polymer-type substrates.

3.3 Plans for Next Reporting Period:

- Fabrication and permeation testing of “pinhole-free” membranes less than 5µm in thickness,
- Preparation of membranes on rigid substrates up to 75 in$^2$ in area,
- Optimization of membrane alloy composition and structure (i.e., effects of single versus two phase structures, optimum Pd/Cu ratio, and specific alloy additions),
- Continued efforts to improve methods for handling and sealing membranes less than 5 µm in thickness

4.0 CONCLUSION

Although efforts were conducted to eliminate pinholes in as-deposited Pd-Cu films less than 5 µm-thick on flexible polymer substrates, employing alternative plastic materials such as Kapton and PET, Ar/O$_2$ plasma treatment for increased wettability, and SiO$_x$ seed layers, we still observed sufficient defects over large areas. In parallel with this effort, however, we developed procedures for producing free-standing, defect free films using rigid silicon and glass substrates and have demonstrated the process to prepare membranes up to 12 square inches in area. Since formation of contiguous Pd-Cu films in the 2-3 µm-thick range is ultimately governed by the size of the particle contamination on the supporting substrate surface, we have adopted techniques utilized by the semiconductor industry to reduce and eventually eliminate particle contamination. A down-select decision has been made to focus our further development on rigid substrates, achieving a stated milestone for the project.
Hydrogen permeation tests were also conducted this quarter on as-deposited, Pd-Cu membranes, between 6-14 µm-thick. In the case of a 6 µm-thick film, the pure hydrogen flux at 20 psig and ~260 °C was 36 cm³(STP)/cm² min. This flux corresponds to a pure hydrogen permeability of $7.4 \times 10^{-5}$ cm³ cm⁻¹ s⁻¹ cm Hg⁻¹/2 at 250 °C. This value is within 20% of the pure hydrogen permeability at 250 °C reported in the McKinley patent. In the case of a 14 µm-thick membrane tested at 350 °C, the pure hydrogen flux, measured before initiating a pinhole-size leak, was $2.1 \times 10^{-5}$ cm³(STP) cm/cm² s cm Hg⁰.⁵. This value is considerably lower than the expected permeability of Pd₆₀Cu₄₀ materials at 400 °C.

To date, essentially all of the vacuum deposited Pd-Cu thin film membranes have had palladium compositions that were as much as 3% greater than the ideal 60 weight percent composition (this is a direct consequence of sputtering from a 60/40, Pd/Cu alloy target). As the concentration of Pd is increased beyond the optimum 60% value, a less desirable two-phase structure forms at the higher temperatures (in this case, above 260 - 280 °C). As we continue to produce thinner Pd-Cu films next quarter, we will also be optimizing the composition and corresponding hydrogen permeation flux as well.

5.0 REFERENCES