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TITLE TRITIUM EXPERIMENTS ON COMPONENTS FOR FUSION FUEL PROCESSING AT THE TRITIUM SYSTEMS TEST ASSEMBLY

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TRITIUM EXPERIMENTS ON COMPONENTS FOR FUSION FUEL PROCESSING

AT THE TRITIUM SYSTEMS TEST ASSEMBLY

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ABSTRACT

Under a collaborative agreement between US and Japan, two tritium processing components, a palladium diffuser and a ceramic electrolysis cell have been tested with tritium for application to a Fuel Cleanup System(FCU) for plasma exhaust processing at the Los Alamos National Laboratory. The fundamental claracteristics, compatibility with tritium, impurities effects with tritium, and long-term behavior of the components, were studied over a three year period.

Based on these studies, an integrated process loop, "JAERI Fuel Cleanup System" equipped with above components was installed at the TSTA for full scale demonstration of the plasma exhaust reprocessing.

I. INTRODUCTION

The Japan Atomic Energy Research Institute (JAERI) and the United States Department of Energy (DOE) agreed a collaborative program for the development of the components, the palladium diffuser and the ceramic electrolysis cell for the Fuel Cleanup System(FCU), which would be applicable to the next geogration of fusion experimental devices. Both designed and manufactured by JAERI and shipped to the Tritium Systems Test Assembly(TSTA) at the Los Alamos National Laboratory for the testing with tritium.

A palladium diffuser separates hydrogen isotopes from all other impurity species and produces a pure hydrogen stream that will be handled in the isotope separation system in the fusion fuel koop. The ceramic electrolysis cell decomposes tritiated water vapo, without generating solid waste or having a large tritium inventory. Since 1984, preliminary tests on both components with tritium have successfully been performed at the TSTA under an early agreement is inter JAERT collaborations. Both components proved to be feasible for tritium service and seemed attractive for application in the processes if the Eucl Cleanup^{1,4}. Thus further studies were suggested for development of practical dependents.

TT. EXEMPTS

A Fallation toffiser

the jailatium diffuser contains 45 fine palladoum ally tibes that selectively permeate hydrogen csst pest. Ecquire 1 shows the schematic R. V. Carlson, K. E. Binning, J. R. Bartlit and J. L. Anderson, Los Alamos National Laboratory, Los Alamos, NM 87545, 505-667-3651

of the diffuser. A mixture of hydrogen isotopes and impurity is separated into a pure hydrogen stream and a bleed stream that contains mostly impurity and some residual unpermeated hydrogen. The diffuser is contained in a double jacket in order to recover tritium permeated from the primary containment that is heated up to 450°C by a heater wire wound on it. The flow diagram of the experiment is shown in Figure 2. One or two metal bellows pumps are used to evacuate the inside of the palladium tubes and recirculate the permeated hydrogen into the feed stream. Impurities such as methane and carbon monoxide were added in order to investigate the chemical effect as well as measure the separation characteristics. Gas samples were occasionally taken from the feed



Fig. 1. Schematic of the Palladium Diffuser

in bleed for Panan or mass spectroscopy. Flow rates and pressures were measured at the feed and permeated sides of the diffuser. Tritium was supplied to the experiment from the ZrCo red. Approximately 2500Ci were used. The experiment was independent of the TSTA main loop and was placed in a separate glovebox.



Fig. 2. Flow Diagram of the Palladium Diffuser Experiment.

B Ceramic Electrolysis Cell

The electrolysis cell contains 10 sintered stabilized zirconia tubes that has calcined platinum electrodes at the inside and outside. Figure 3 shows the structure of the cell. Water vapor in the feed gas stream is decomposed at the surface of the inner electrode to thim hydrogen/tritium while pure oxygen generates at the outer surface of the ceramic tube.

The rest loop consisted of a tritium source, a catalytic reactor filled with hypcalite, a LNP freezer, the CEC, a metal relives pump, and a ZrCo tritium storage bed as clown in Figure 4. Fure or high level tritiated water was formed by exidation of tritium gas at the hopeslite bed. The water vapor was then collected in a freezer so that regeneration of the freezer in the FCU system was simulated. In some rests, vapor was continuously sent to the cell for decomposition followed by recombination with exygen from the exygen side of the cell. There gas and deuterium was used as carrier gas in the loop. Effects of imputities such as CO and CDP were investigated. The experiment was located in a gl vebox in an auxiliary laboratory at the TGTA

1. Zardenaum Cohalt Bed

the Assessmentalist schemetalist mpound is a new material developed by JABPT as a substitute for unaryum for tritium recovery and storage⁵., and some beds for tritium service are developed⁶.



Fig. 3. Schematic of the Ceramic Electrolysis Cell.



Fig. 4. Flow Dingram of the CEC Experiments.

Both experiments were equipped with ZrCo beds for recovery, storage and supply of tritium as shown in the figures. Some features of the ZrCo teds were tested in practical tritium service.

III RESULTS

A. Palladium Diffuser Experiment 1 Fermeability Measurement

The permeation flow rates of pure H₂, D₂ and T₂ were measured as the functions of pressure at various temperatures. Examples at 200°C are summarized in Figure 5. Flow rates are expressed in cm^2 per minutes at 0°C, 103kPa. Linear relations are observed between flow rates and differential square root of pressures across the membrane although some deviation at both high and low pressures appeared. Permeability of tritium was approximately one half of that for hydrogen.



Fig. 5. Fermeability of Pure Hydrogen Isotopes through the Diffuser.

2 Separation Characteristics

A mixture of hydrogen and 10% methane was introduced into the system to measure the reparation characteristics of the diffuser. Seed and permeation flow rates were measured as a function of the concentration of hydrogen in the blood stream. The result indicates a low oncentration of hydrogen is the blood from a feed intaining major amount of hydrogen, theory the permeation flow rate is much smaller than in the case of pure hydrogen feed. This is aured by a high partial pressure of impurity in the feed wide of the diffuset. Numerical "alculation suggests that an increased diffuser length was desirable for achieving low hydrogen concentration in the output impurity stream. The system was left in a continuous operation mode for months and little change in the separation characteristics was observed.

3 Carbon Monoxide Testing

A high concentration of CO in the stream may be encountered in some applications of the diffuser in tritium processing loops. The diffuser was operated for an extended period with a T2-CO mixture. Approximately a 10% loss of permeability occurred in a 6 month operation. Oxidation treatment of the membrane followed by hydrogen reduction regenerated the permeability as in Fig.6. This result suggests that the reduction of the permeability might have been caused by surface contamination of the palladium alloy membrane. A possible deposit of carbon caused by the radiolysis of CO was suspected.



Fig. 6 Regeneration of the diffuser after longterm exposure to Tp=CO.

4 Long term Reliability

The diffuser was operated at relatively low temperature (150°C = 300°C) to covertigate the effect of ⁴He formed in the palladium metal liftic degradation of the permeability was observed in more than 1 year of operation

9 Ceramic Electrolysis Cell Experiment 1 Decomposition Characteristics

Decomposition characteristics of the electrolysis cell was measured with H_2O-N_2 , T_2O-N_2 N2, T20-D2 systems under the conditions simulating application for a plasma exhaust process. In the Fuel Cleanup Systems in TSTA or TPL, DTO captured at the DTO freezers should be regenerated and decomposed to recover tritium. Carrier gases such as No, He or Do may be used for regeneration of the freezers. Carrier flow rate was 400 - 1000cc min, where no effect of the flow rate was observed in the characteristics of the cell. Figure 7 shows the conversion efficiency of water vapor to hydrogen obtained with the cell operated at 600°C. Conversion ratio was determined from the ratio of the inlet and the outlet humidities. The Irfree voltage is measured between the two electrodes on the cell that composes an open circuit where no current is applied. This value indicates the electrochemical potential across the cell generated by the difference of oxygen potential at the 0_2 side and the T_2/T_2O side. The observed conversion efficiency was around These values are lower than expected, 958. probably due to the error caused by the residual humidity at the outlet of the cell.A small isotopic difference was observed between T20 and HpC. The theoretical conversion efficiencies for each system are also shown on the figure. The results show similar trends although marked deviations are observed.



iq 2. Conversion efficiency from vapor to hydrogen for H2 o N2, T2 (62) and T20012 systems at the electrolysis coll Theoretical relationship also shown.

Teuterium was tested as carrier gas for the tritiated water. Use of the D_2 for regenerating the freezer is advantageous because only gaseous hydrogen isotopes (DT)2 is expected to be obtained as product stream from the cell. The experiment proves that T_2O can be decomposed at high efficiency at 1.4V, that is higher than the case with inert carrier as predicted by the valculation.

2 Impurity Testing

Systems of CO_2-N_2 and $CO_2-D_2O-N_2$ were tested to investigate the effect of CO_2 that might be electrolyzed by the cell while water is decomposed. A thermochemical calculation was made for the electrochemical equilibria of the CO_2 , CO, H_2O , H_2 , C and O_2 systems in the cell.

Figure 8 shows the relationship between conversion ratios for the reactions discussed above and Ir-free potential at 600°C. For the reactions $CG \leftrightarrow \frac{1}{2}O_2 + C$, and $CO_2 \leftrightarrow C + O_2$, initial concentrations of CO or CO₂ are assumed to be 0.1. Each line shows how the reaction can proceed at the given Ir-free voltage in the cell. As seen in the figure, the decomposition of water is the easiest to occur in the cell, but all the reactions are possible at the potential of 1.3V to 1.4V where the cell is usually operated. Either CO₂ or CO can be decomposed to form carbon as the result of the reaction.



Fig. H Theoretical equilibrium conversion of H₂C and CO, at the electrolysis cell at 600°C. Feed concentration was assumed to be 0.1 for CO and CO₂.

Figure 3 shows the result of the tests with Coy. The electrolysis of Coy starts at about 2.97, that is close to the calculater voltage for reaction Coy $\epsilon + \frac{1}{2} \gamma \gamma + C \sigma$, however

the conversion ratio was less than $3\times$. It is inderstood that the percus platinum electronelies not have catalytic activity for the decomposition of CO₂, while it does for the decomposition of water. Electrolysis of D₂O -CO₂ - N₂ mixture shown in the figure indicates no effect on the electrolysis performance of water. These results suggest that CO₂ impurity in the cell does not have any undesirable effect on the decomposition of water.



Fig. 9. Electrolysis of $CO_2 - N_2$ and $D_2O - CO_2 - N_2$.

3 Batch Operation

One of the major practical applications of the CEC is the batch processing of a tritiated water gas mixture. In the test, a mixture of tritiated water and carrier was processed with the cold trap, electrolysis cell and a ZrCo bed to convert tritiated water and recover tritium in a closed loop. More than 99.99% of the water was decomposed and tritium was trapped at the bed. It was proved that the CEC is suitable to process relatively small amounts of high level initiated water in a batch operation.

4 Long-term Reliability

The electrolysis experiment was operated in a closed loop mode in which oxygen generated at the anode is recycled to the catalytic reactor inlet for recombining with electrolyzed tritium. A test with T2O-CO2-He was performed as long as one year to evaluate the long term reliability of the cell with tritium and impurity. The cell has successfully worked for the test period, however marked embrittlement of the atabilized zirconia material was observed when the test was completed.

C.f.r.s.num-Cobalt Bed 1 Four Dibrium Pressure of Tritium

Pressions imposition isotherms of the Ty-Zife system was measured with the Zife Led installed in the CEC experiment. Preliminary results indicine that the equilibrium pressure of Ty-culture Trick is higher than that of hydrogen by a factor of less than 2. It is suggested that the isotopic difference was negligible in the practical use of 2rCc for tritium service. Detailed measurement was done by the program and apparatus under the Annex IV agreement.

2 Practical Application

Both palladium diffuser and electrolysis cell experiments were equipped with ZrCo beds of 5 liter of hydrogen in capacity. The beds were used for storage and supply of pure and mixed tritium. The recovery of tritium was performed both by absorption of pure isotopes and recycling of the mixture with inert through the bed. These practical experiences of the beds verified that a ZrCo bed is a suitable substitute for uranium beds.

IV. CONCLUSION

The experimental program on the "process ready components" under Annex III was completed and all of the objectives were achieved experimentally in the tests performed in these three years. Through the tests, it is concluded that the palladium diffuser is applicable to the processing of plasma exhaust to produce pure hydrogen isotopes for as long as 3 years without any maintenance. Use with carbon monoxide was not a problem. The ceramic electrolysis cell was verified as an attractive component for the decomposition of tritiated water in various processes. Carbon dioxide affected it little. Thus, both process-ready components, the palladium diffuser and the ceramic electrolysis cell, were proved to be suitable for fusion fuel processing. Long term reliability and compatibility of the components with tritium and impurity was verified.

Based on the results, an integrated process loop, "JAERI Fuel Clearup System" that utilizes both components was developed and designed by JAERI for full scale demonstration of the plasma exhaust reprocessing. The system will be tested with simulated fusion fuel in the TSTA main loop in the near future.

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