Comparison Study Between Hydrogen Palladium Sensor and Hydrogen Manganese Dioxide Sensor

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ABSTRACT

This research work oriented to compares between Palladium (Pd) thin films sensor that have been prepared on glass substrates and hydrogen sensor using optical fiber, coated with Manganese dioxide (MnO₂) nanostructures. Palladium thin films prepared by vaporization deposition technique with annealing temperature around 600°C. Where, manganese dioxide, prepared on Optical Multimode Fiber (MMF) of 125m in diameter as the transducing platform was tapered to 20m to enhance the evanescent field of the light propagates in the fiber core.

The optical characteristics of the prepared films showed that they are highly sensitive, but their properties considerably vary when the measurements conducted in vacuum or in air. The response-recovery time of Pd materials to hydrogen gas is characterized to be extremely short.

The tapered optical fiber coated with Pd/MnO₂ nanograins was found to be sensitive towards hydrogen with different concentrations in synthetic air at 240°C operating temperature. The annealed sensor showed higher response and sensitivity compared with the as-prepared sensors when measured in the visible to near infrared optical wavelength range. The absorbance response of the annealed Pd/MnO₂ on fiber has increased to 65% compared to 20% for the as-prepared Pd/MnO₂ upon exposure to 1% Hydrogen in synthetic air.

Keywords: Manganese dioxide, Palladium, Semiconductor, Spectrometer, Sensor, Thin-Film

I. INTRODUCTION

Hydrogen gas occupies about more than 90% of the atmosphere. It is a highly flammable gas that burn at concentrations as low as 4% in air. It has a larger window of flammability in comparison to other natural gases such as gasoline, propane, ethane, methane, propylene, etc. The flammability limit of hydrogen is even times wider than methane. It is therefore, critical for a hydrogen sensor to have a wider measurement range for safety applications than most common fuels. Hydrogen is the lightest of elements and the smallest molecule; it, therefore, has the greatest tendency to leak[1].

The use of hydrogen as a clean fuel in various applications requires practical and robust sensors to minimize risk of explosion associated with its volatile properties[2]. Thus, for a process safety application, a hydrogen leak can be more dangerous and its detection becomes more challenging than other flammable gases[1].

II. HYDROGEN SENSOR TECHNOLOGY

Different of approaches that used to sense and detect hydrogen. Number of which are used in industry include the typical Gas Chromatography (GC), Mass Spectrometry (MS), Catalytic Bead (CB), and thermal conductivity. Semiconducting metal oxide and CB sensors are popular solid-state technologies, which employ heated catalysts to sense hydrogen. These
Electrochemical sensors are based on known electrolytic reactions of hydrogen. Sensors based on catalytic combustion are generally nonspecific, electrochemical hydrogen sensors with liquid or solid type electrolytes having leakage issues. The hydrogen sensors based on thermal conductivity, CB, metal oxide, and electrochemical technologies require the presence of oxygen for sensor operation. Oxygen plays a crucial role in promoting the grain boundary formation in metal oxide sensors and electron transfer reactions in electrochemical sensors. The most promising solid-state technology is based on a hydrogen-specific material, palladium, which does not require oxygen for operation. Palladium-based sensors are gaining wide popularity in industry due to their reliability and high specificity to hydrogen. Any hydrogen sensor technology needs to satisfy the three basic requirements sensitivity, selectivity, and specificity.

Several types of palladium-based hydrogen sensors have been reported in the literature. The most notable ones are based on Pd thin-film resistors, FETs, Pd nanowires, Pd nanoparticle networks, Pd nanoclusters, and Pd nanotubes. Palladium Field-Effect Sensors: “Hydrogen sensors based on the “field effect” of palladium have been investigated extensively in the literature”[3]. “The field effect results due to the rapid dissolution of hydrogen in the palladium surface arranged in a Pd–SiO2–Si configuration. The sensor relies on an electric field resulting from the charge transfer between palladium and hydrogen on its surface. The FETs”[4] and “metal–insulator–semiconductor (MIS)”[5] are the two major types of device structures that have been studied for palladium-based hydrogen sensing. Palladium is catalytically active, permeable to hydrogen, and can be readily used in FET and MIS devices.

A wide variety of solid-state sensors based on hydrogen-specific palladium, “Metal Oxide Semiconductor (MOS), CB, electrochemical, and Surface Acoustic Wave (SAW) technology are used in the industry for several years. Microelectromechanical systems (MEMS) and nanotechnology-based devices for the measurement of hydrogen are the recent developments. These developments are mainly driven by the demands of the fuel cell industry. Solid-state approaches are gaining rapid popularity within the industry due to their low cost, low maintenance, replacements, and flexibility of multiple installations with minimal labor[1].

Number of traditional approaches can be used to detect Hydrogen, these may include:

i. Thermal Conductivity (TC): is the most widely applied measuring principle for the determination of hydrogen. “The measuring principle is based on the differences in thermal conductivity of the gases to be measured. A Thermal Conductivity Detector (TCD) measures the concentration of a gas in a binary gas mixture by measuring the thermal conductivity of the sample gas and comparing it to the thermal conductivity of a selected reference gas”[1].

ii. Gas Chromatography (GC): is also another widely applied measuring principle for hydrogen detection. “The disadvantages of GC are long response times (minutes) due to the chromatography, time-intensive sample preparation, consumable (carrier and calibration gases), and labor-intensive handling procedures. An advantage, however, is the ability to measure other gases such as nitrogen, oxygen, and carbon dioxide in the presence of hydrogen. But, this adds time to the total analysis”[1].

A wide variety of solid-state sensors require heating to about 300°C to enable surface reactions that promote hydrogen sensing. Electrochemical sensors are based on known electrolytic reactions of hydrogen. Sensors based on catalytic combustion are generally nonspecific, electrochemical hydrogen sensors with liquid or solid type electrolytes having leakage issues. The hydrogen sensors based on thermal conductivity, CB, metal oxide, and electrochemical technologies require the presence of oxygen for sensor operation. Oxygen plays a crucial role in promoting the grain boundary formation in metal oxide sensors and electron transfer reactions in electrochemical sensors. The most promising solid-state technology is based on a hydrogen-specific material, palladium, which does not require oxygen for operation. Palladium-based sensors are gaining wide popularity in industry due to their reliability and high specificity to hydrogen. Any hydrogen sensor technology needs to satisfy the three basic requirements sensitivity, selectivity, and specificity[1].

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ii. Palladium-Coated Fiber Optic Sensors: A fiber optic hydrogen sensor consists of a palladium coating at the end of an optical fiber that senses the presence of hydrogen in air. "When the coating reacts with the hydrogen, its optical properties are changed. Light from a central electro-optic control unit is projected down the optical fiber where it is either reflected from the sensor coating back to central optical detector or is transmitted to another fiber leading to the central optical detector. A change in the reflected or transmitted intensity indicates the presence of hydrogen. The fiber optic detector offers inherent safety by removing all electrical power from the test sites and reduces signal-processing problems by minimizing electromagnetic interference. The fiber optic hydrogen sensors can be fabricated using a palladium-coated single-mode tapered optical fiber" [10]. The attenuation change of the fiber-mode when the device was exposed to hydrogen is used to detect and measure hydrogen concentration in gaseous atmospheres.

III. HYDROGEN PALLADIUM SENSOR COMPONENTS

Hydrogen palladium thin films sensor consists basically on seven components include; chamber, white photo diode, spectrometer, vacuum unit, gauge pressure, cylinder of hydrogen gas, and computer. Figure (1) and figure (2) below illustrate sensor component and sensor setup successively. A sample of palladium coated glass slide sensor should be prepared first to detect hydrogen, and should be tested by X-ray to check its crystallization[11].

The main sensor components are;

a. The coated glass substrate (Palladium thin films): should be placed inside the developed polymer square test chamber of 50 mm square base and of 75 mm height with the top removable cover. The effective volume of the chamber is 187500 mm³; it has an inlet to allow the test gas to flow in.[11]

b. White photo diode: is the light source used in the sensor.

c. CCS Spectrometer: used to measure transmitted.

d. Vacuum unit: to evacuate the test chamber.

e. Gauge pressure: to measure the current pressure of the chamber.

f. Cylinder of hydrogen gas of a known concentration: to flow through the test chamber during measurement.

g. Computer: to process the recorded signal.

IV. SETUP OF HYDROGEN MANGANESE DIOXIDE SENSOR

The optical hydrogen gas sensing setup comprised of tungsten halogen light source modelled HL-2000, Ocean Optics USA which exhibit wavelength range of 360 nm to 2500 nm. Optical absorbance spectrum was monitored via spectrophotometer (USB4000 VIS-NIR, Ocean Optics USA) with spectral response ranged between 200nm to 1100 nm of wavelength. The light source was coupled into the MMF by using a standard SMA optical cable (600 nm in core diameter) for light transmission. The MMF termination is FC/PC type. Hence, an SMA-FC adapter was used to couple the SMA cable to the MMF. From the adapter data sheet, typical insertion loss for mating FC/PC terminated MMF to SMA terminated cable is 1.4 dB. The calculated coupling efficiency based on this insertion loss was 99.6%. The MnO₂-coated on tapered optical fiber sensor was
placed in the sealed gas chamber as demonstrated in Figure (3). The gas purging into the chamber was controlled by a computer-controlled mass flow controller which regulates gas flux at a flow rate of 200 sccm. In order to obtain the desired concentration of hydrogen gas, pure synthetic air was used to dilute the hydrogen gas to concentration varied from 0.125% to 1.0%. The absorbance measurement and dynamic response of cumulative absorbance were monitored while purging alternately hydrogen gas and pure synthetic air, respectively. The chamber is also equipped with a heater plate which can be varied by dc voltage supply as a source of operating temperature [12].

\[ \text{Fig: 3 Gas sensing experimental setup} \]

V. MEASUREMENTS AND RESULTS

Two tests were carried out to examine the palladium thin film sensor. In both tests, results were obtained through adopting the following steps:

Opening the test chamber to place the palladium thin film sensor on the sensor holder and close it. The necessary light source is then directed by optical fiber and allowed to pass through the sample to the spectrometer. The rotary pump is then switched on to evacuate the test chamber to about -0.7 bars. Next, the hydrogen gas of a known concentration allowed passing from the cylinder through the special inlet to the test chamber by opening the cylinder valve. Test chamber pressure should be measured by observing the gauge pressure. Spectrometer is then detects and analyzes transmitted signals and sends the data to the computer.

Numbers of measurements were carried out in different pressures. Six readings were observed for each sample test including pressures of -0.6, -0.5, -0.4, -0.3, -0.2 and -0.1 bar. Frequency against transmitted light intensity graphs were produced for each reading. The six graphs were then combined together in one diagram with a background graph representing -0.7 pressure without Hydrogen. Figure (4) and figure (5) below represent the resultant combined graphs for both sample tests [11].

\[ \text{Fig: 4. Combined graphs of the palladium thin film sample one (Pd 1) with a numbers of measurements were carried out indifferent pressures. Six readings were observed for each sample test including pressures of -0.6, -0.5, -0.4, -0.3, -0.2 and -0.1 bar.} \]

\[ \text{Fig: 5. Combined graphs of the palladium thin film sample two (Pd 2)} \]

From the figures obtained above it can be noted that by increase of concentration of hydrogen gas in the test chamber, the transmitted light is also increase which, means that the palladium thin film sample successfully detect hydrogen gas. And it can obviously note that the suitable wavelength that can be used to detect hydrogen gas is a visible band that located in the region of 500-650 nm.

In the Manganese dioxide sensor the gas absorbance response was carried out at operating temperature of 240°C. No response was recorded below this
operating temperature, possibly because of a slow chemical activation between adsorbed gas molecules and the sensing layer [13]. Figure (6) demonstrates the absorbance response of Pd/MnO$_2$ coated optical fiber sensor for as-prepared and annealed samples towards 1% H$_2$. The integration of Pd on MnO$_2$ layer is important to catalyze the gas dissociation and alter the layer optical properties when interact with H$_2$ [14]. Pd has the ability to highly absorb H$_2$ at almost 900 times of its own volume [15]. When both as-prepared and annealed samples were exposed to synthetic air and 1% of H$_2$ gas at elevated temperature of 240°C, a distinctive change of absorbance magnitude over the range of 500–800 nm was observed. However, the annealed sensor gives higher magnitude of absorbance as compared to the as-prepared when exposed to H$_2$. The absorbance change in the MnO$_2$ layer can be explained with the following mechanism. During H$_2$ adsorption onto the sensing layer, the dissociation of H$_2$ molecules onto Pd resulting in the generation of H$^+$ ions and electrons. Ions of H$^+$ were spilled over onto the MnO$_2$ nanograins layer which then reacted with chemisorbed oxygen (O$_2^-$ and O$^-$) resulting in production of H$_2$O molecules [13]. The generated electrons reduced Mn$^{4+}$ in the center of the MnO$_2$ crystal lattice to Mn$^{3+}$ [16]. Hence, the film refractive index and its absorbance changed. Upon exposure to synthetic air, the adsorbed oxygen as well as the desorbed H$^+$ ions in/out of the sensing layer returned the layer absorbance spectrum to its original baseline.

Fig. 6: Absorbance versus optical wavelength of Pd/MnO$_2$ sensor for as-prepared and annealed samples exposed to 1% concentration of H2 in synthetic air at 240 °C.

VI. CONCLUSION

This work is a comparison study between two developed hydrogen sensors. The first depended on the Palladium coated glass thin film and the other depended on optical fiber coated with manganese dioxide. From the measurements carried out and results obtained, it can be concluded with that both sensor efficiently detected hydrogen and electromagnetic interference free.

The Hydrogen palladium thin films sensor has rapid response to low hydrogen concentration, also it has a wide dynamic range specifically in 500-650nm range. Moreover, the sensor can be used in the room temperature.

On the other hand, Pd/MnO$_2$ coated optical fiber sensor is sensitive at elevated temperature of 240°C therefore, it can be used in the hot areas but, it is more expensive compared with Palladium coated glass thin film sensor.

REFERENCES


