Stannous Oxide Thick Film Nanosensors Design by Screen Printing Technology: Structural, Electrical Parameters and H₂S Gas Detection Study

UMESH JAGANNATH TUPE,¹* ARUN VITTHAL PATIL², MADHUKAR SARVOTTAM ZAMBARE³ and PRASHANT BHIMRAO KOLI⁴

¹,³Research Centre in Electronic Science, Department of Electronic Science, Fergusson College, Pune, Maharashtra, Affiliated to Savitribai Phule Pune University, Pune, India.
²Department of Physics, Arts, Science and Commerce, College, Surgana, Taluka- Surgana, District- Nashik, Maharashtra, India, Affiliated to Savitribai Phule Pune University, Pune, India.
⁴Karmaveer Abasaheb Alias and N.M Sonawane Arts, Commerce and Science College, Satana, Taluka-Baglan, District- Nashik, Maharashtra, India, Affiliated to Savitribai Phule Pune University, Pune, India.

Abstract
The present research deals with the fabrication of stannous oxide nanoparticles by conventional and cost effective co precipitation method. The thick film sensors of SnO₂ nanoparticles were prepared by standard screen-printing technique by photolithography. The prepared SnO₂ material was characterized by several techniques to confirm the structural properties. Initially, the prepared nanoparticles of SnO₂ were investigated by x-ray diffraction technique to confirm the synthesis of prepared material within nanoscale. From XRD data the average particle size of prepared thick films was found to be 21.87 nm calculated using Debye-Scherer formula. The material was further characterized by using scanning electron microscopy (SEM) to investigate the structural and surface characteristic of SnO₂. SEM data clearly indicates the heterogeneous surface, and some voids present over the surface of SnO₂ nanoparticles. The Fourier transfer infra red technique was employed to investigate the metal oxygen frequency of SnO₂ material. The prepared sensor was exclusively utilized to sense the hydrogen sulfide gas vapors at various concentrations. The prepared sensor was found to be highly sensitive to H₂S vapors nearly 63.8% sensitivity was recorded. The response and recovery study shows the response time of 9 seconds and recovery time of 19 seconds for hydrogen sulfide gas vapors. The SnO₂ sensor was further utilized for recycling performance to get the firm results of sensitivity in four turns with period of 15 days.

CONTACT Umesh Jagannath Tupe umeshtupe14@gmail.com Department of Electronic Science, Fergusson College, FC Road, Shivajinagar, Pune, Maharashtra, India.

© 2021 The Author(s). Published by Enviro Research Publishers.
This is an Open Access article licensed under a Creative Commons license: Attribution 4.0 International (CC-BY).
Doi: http://dx.doi.org/10.13005/msri/180108
Introduction

Material science recently becomes more promising field in the catalysis and sensors technology. Most of the researchers are working on the material technology to defeat the environmental and industrial problems. Due to rapid urbanization and industrialization the pollution related problems are enhanced with high percentage leading to fatal toxic effects on human and animals. Thus, everyone is looking to get perfect remedy to get rid off from these pollution related problems. Material science has great potential to defeat these problems. Material science has great potential to defeat these problems. Since, most of the material prepared as sensors are easily employed to sense these toxic gases at various concentrations. The main air pollutants such as carbon dioxide, carbon monoxide, sulphur dioxide and NO\textsubscript{2} etc. are responsible to make polluted environment.\textsuperscript{1,3} Hence, the similar gases must be detected with effective sensors. These gases are emerging as automobile exhaust and industrial by products and their desired concentration is highly toxic.

On the other hand there are several gases which are emerged during chemical reactions, as well as by products of several chemical processes and mining. The gases like hydrogen sulfide, VOC’s (volatile organic compounds), methane, and natural hydrocarbons etc. Out of these the hydrogen sulfide concentration at elevated level can be highly toxic and leads to fatality in many ways. According to research, nearly 700-800 ppm concentration of H\textsubscript{2}S leads to loss of consciousness, fatigue, breathing problem, cardiological arrest etc. Hence, the good sensors that can sense the hydrogen sulfide vapors will be very useful defeat H\textsubscript{2}S toxicity.\textsuperscript{3,4} Hydrogen sulfide (H\textsubscript{2}S) is colorless, more hazardous and toxic element. It can be identified by a characteristic rotten egg odor at small concentration. The major noticeable point of hydrogen sulfide is easily smelled perceptible at concentrations low concentration in air but if the concentration of gas is increases in the surrounding, smell is disabling. The presence of high concentration of hydrogen gas in the surrounding with no perceivable odor is extremely dangerous for human being, so it is necessary to control and monitor H\textsubscript{2}S gas in the environment.\textsuperscript{4,5} The gas is inherently found in manure, hot springs, volcanoes, several sewage, swamps, geysers, by-products of several chemical reactions, mining etc.

There are several methods to prepare the nanoparticles some of the popular methods can be listed as physical vapor deposition (PVD), chemical bath deposition (CBD), spray pyrolysis, green synthesis method, sol-gel method, combustion method and co-precipitation method. Depending upon the requirement the method can be selected to synthesis the desired nanomaterials. The stannous oxide nanoparticles can be cost effectively synthesized by co-precipitation and sol-gel route. Many of the research papers flaunting that metal oxides nanoparticles can be useful as gas sensors for the most of the gases.\textsuperscript{6,7} A great deal of efforts has been put into developing new sensing materials with improved sensor properties. The listed synthesis techniques can be easily employed to fabricate the various semiconducting nanosensors in low time and cost effective way.\textsuperscript{7,8}

The tin dioxide (SnO\textsubscript{2}) is a versatile metal oxide because of its two characteristics: variable valence state and existence of oxygen vacancy defects. It has wide band gap of ~3.6 eV and it shows n-type semiconductor behavior. Tin dioxide also referred as tin oxide and stannous oxide. SnO\textsubscript{2} has numerous special attentions in the field of material science due to their diverse applications like catalysis, transparent conducting oxides, battery materials, solar cell, optics and gas sensing etc. The stannous oxide sensors are effectively used to sense the various toxic gases as mentioned earlier as thin and thick film gas sensor.\textsuperscript{8,9}

In the present research we are reporting the cost effective synthesis of SnO\textsubscript{2} by co-precipitation method and its application for detection of H\textsubscript{2}S gas at various concentration and temperature. The structural and electrical properties of SnO\textsubscript{2} thick films are also reported in the present investigations. In the present investigations the SnO\textsubscript{2} thick film sensor was exclusively utilized to sense the hydrogen sulfide gas vapors. In addition to that the response and recovery curves and recycling properties are reported of SnO\textsubscript{2} sensor for hydrogen sulfide gas vapors.

Material and Methods

The chemicals required for synthesis of SnO\textsubscript{2} are sodium hydroxide, stannous chloride dihydrate, acetone, ammonia, and double distilled water. All the chemicals used were analytical grade and used without further purification. The screen printing
apparatus was utilized to prepare stannous oxide thick film sensors.

**Synthesis of SnO$_2$ Nanoparticles using Precipitation Method**

SnO$_2$ nanoparticles were synthesized by cost-effective co-precipitation method. In this method, stannous chloride dihydrate used as source of tin (Sn). To prepare SnO$_2$ nanoparticles initially, 0.01 moles of tin chloride dihydrate (SnCl$_2$·2H$_2$O) were dissolved in 100 ml of double distilled water. In a separate beaker 0.1 molar solution of ammonia was prepared. Then, 0.1 a molar ammonia solution was dropped into tin chloride solution drop wise with constant stirring. The stirring was continued till white precipitate of tin hydroxide was obtained. The, white precipitate was filtered off, dried and calcined under muffle furnace at 400$^\circ$C for three hours. The white colour tin dioxide nanoparticles were recovered from muffle furnace on next day and used for further process of research.

**Thick Films Preparation of SnO$_2$ Nanoparticles by Using Standard Screen Printing Technique**

The thick films of pure SnO$_2$ were prepared by using screen printing technique on glass substrate. The organic compounds used as binders and tin oxide nanoparticles were used in appropriate ratios as 30:70 respectively. The compounds used as binders over the glass substrates were 8% BCA (Butyl Carbitol Acetate) and 92% ethyl cellulose (EC). Where, inorganic composition used as tin oxide nanoparticles. Together, these both organic and inorganic compounds were mixed in mortar and pestle and mixed mechanically, the desired concentration of EC and BCA were added to get thixotropic paste. The prepared paste mass was then applied on glass substrate (previously cutted with 2.5×1.25 cm) by screen printing technology. Screen printing setup consists of nylon screen of 120 meshes to put paste on glass substrate. After complete coating of the films, the films were dried under IR lamp for 30 minutes. In these process IR radiations removes temporary organic volatile materials. Finally, films were annealed at 350$^\circ$C in the muffle furnace for 3 hours. The calcined films were used in further gas sensing and electrical study.

**Thickness Measurement of the Films**

The surface coating of the films was calculated using equation 1 as mentioned below. The thickness of the stannous oxide film sensor was calculated as 4.22 μm, (4220 nm). The thickness is observed in the thick region. The thickness of the SnO$_2$ film sensor was calculated using mass difference method.\(^{10, 11}\)

\[
t = \frac{\Delta M}{A \times \rho} \ldots(1)
\]

\[
\Delta M = \text{Mass difference of the film before and after deposition.}
\]

\[
\rho = \text{Composite density of stannous oxide film sensor SnO}_2
\]

\[
A = \text{Area of the film}
\]

**Fig.1: XRD spectrum of SnO$_2$ thick film**

**Results and Discussion**

**X-ray Diffraction Study**

The screen printed thick films of stannous oxide were investigated by means of x-ray diffraction technique model number D8 Bruker AXS (Germany), possessing Braggs scanning angle varying from 20-90$^\circ$. The instrument consisting CuKa (wave length 1.51 Å) used to generate x-rays.\(^{12, 13}\) The XRD spectrum of stannous oxide thick films is as depicted in figure 1. The XRD spectrum as depicted in figure 1, from which the Bragg’s reflection peaks can be assigned to the formation of SnO$_2$ material having tetragonal crystal lattice with space group P42/mnm.\(^{14, 15}\) The 2θ values of diffraction peaks gained from x-ray diffraction statistics for stannous oxide material are 26.18, 33.53, 38.26, 51.59, 54.10, 57.12, 61.14, 65.18, 71.40, and 79.07. The two theta values as mentioned above can be assigned to reflection of (110), (101), (200), (211), (202), (002), (310), (112), (202), (321) planes. The diffraction peaks mentioned above with respective hkl planes confirms the formation tetragonal SnO$_2$ crystal.
The average particle size was calculated from Debye-Scherer's formula represented in equation 2:

$$D = \frac{K \lambda}{\beta \cos \theta} \quad \text{(2)}$$

Where $D$ is average particle size, $K$ is constant (0.9 to 1), $\beta$ is full width half maxima (FWHM) of diffracted peak, $\theta$ is the angle of diffraction. The average particle size of stannous oxide nanoparticles calculated using equation 2 was found to be 51.10 nm. The match scan data of stannous oxide exhibits the fabrication of SnO$_2$ material with JCPDS cart number 00-001-0625.

From SEM images it can be seen that the nanoparticles with varied size and diameter having voids and cavities over the surface of SnO$_2$ nanoparticles. These voids are responsible for the adsorption of the gas molecules like H$_2$S. The specific surface area calculated by using SEM images was measured by using Image J software for spherical particles. The rate of adsorption and desorption increases with large surface area of the films and chemisorptions properties between adsorbent SnO$_2$ nanoparticles and adsorbate gas molecules. Using BET method, specific surface area of SnO$_2$ thick films was calculated by equation 3. Specific surface area of SnO$_2$ material found to be 1.4312 m$^2$/g.

$$S_w = \frac{6}{\rho \times d} \quad \text{(3)}$$

Where, $d$ is the diameter of the particles and $\rho$ is the density of the particles and $S_w$ is specific surface area.

**Energy-Dispersive X-ray Spectroscopy (EDAX)**

Energy dispersive x-ray spectroscopy of SnO$_2$ thick film samples fired at 350°C was performed. From EDAX spectrum of SnO$_2$ the elements tin and oxygen are well resolved. From EDAX spectrum it can be seen that there is no other impurity elements were found in SnO$_2$ thick films. From the EDAX scale of SnO$_2$ it can be observed that the elemental tin is well resolute at 3.4-4 KeV, while the elemental oxygen is resolute at 0.6-08 KeV. The EDAX results obtained for stannous oxide thick are in good agreement with reported results. Excess amount of oxygen observed in the E-DAX analysis, may be due to the adsorption of the oxygen during the sintering or firing stage of the SnO$_2$ thick films. Due to high electron affinity of surface molecules, oxygen gets attached to the molecules on the surface of the thick films.

**Scanning Electron Microscopy (SEM)**

Figure 3 shows SEM image of pure SnO$_2$ thick film. The 2D image with high magnification was used to study surface morphology of this film. SEM results indicate the formation of particles with different shapes and sizes. The image shows, larger particles or grain agglomerates. From SEM images it can be seen that the nanoparticles with varied size and diameter having voids and cavities over the surface of SnO$_2$ nanoparticles. These voids are responsible for the adsorption of the gas molecules like H$_2$S. The specific surface area calculated by using SEM images was measured by using Image J software for spherical particles. The rate of adsorption and desorption increases with large surface area of the films and chemisorptions properties between adsorbent SnO$_2$ nanoparticles and adsorbate gas molecules. Using BET method, specific surface area of SnO$_2$ thick films was calculated by equation 3. Specific surface area of SnO$_2$ material found to be 1.4312 m$^2$/g.

**Electrical Characterization**

The electrical characterization of prepared SnO$_2$ thick films studied in this section. The electrical characterization of sample performed by determination of resistivity ($\rho$), the activation energy ($\Delta E$) and the temperature coefficient of resistance (TCR).

Resistivity of SnO$_2$ Thick Film

Resistance determination of SnO$_2$ thick films was done using home built characterization system shown in figure 4. Resistance of SnO$_2$ thick films was determined by using half bridge method. In half bridge method the value of $R_{\text{ref}}$ = 10 MΩ and 30 VDC power supply were used. The resistance of the thick film samples was determined using following equation:

$$R_{\text{sample}} = R_{\text{ref}} \left[ \frac{(V_{\text{supply}} / V_{\text{ref}})}{1} \right]$$

To calculate resistivity of thick film, $R = 24 \times 10^8$ Ω, $L = 2.5$ cm, $b = 1.25$ cm and $t = 23$ µm were used.

$$\rho = \frac{R \cdot b \cdot t}{L}$$

Activation Energy of Stannous Oxide Thick Film Material

The activation energies of prepared SnO$_2$ thick films was measured at the low temperature and high temperature, the activation energy of SnO$_2$ film was calculated using Arrhenius plot. Figure 6 shows graph of $\log R$ in ohm versus reciprocal of temperature $(1/T)$ in °K for SnO$_2$ thick films. The Arrhenius plot indicates two distinct regions of temperature low and high temperature region respectively. The annealing temperature of films plays a very important role to decide the transition temperature from low region to high region. The Activation energy for SnO$_2$ thick film found 0.2767 eV and 0.5988 eV at low and at high temperature respectively. The energy of activation within the low temperature region is often but the energy within the high temperature region because material passes from one conduction mechanism to a different. In the low temperature region, the increase in conductivity is because of the mobility of charge carrier, which is rely upon the dislocation concentration. The conduction mechanism is
typically called the region of temperature conduction, during this region energy of activation decreases because at very low thermal energies quite sufficient for the activation of charge carriers to require part within the conduction process. Hence increase in conductivity within the lower temperature region may be attributed to the rise of charge mobility.

Gas Sensitivity Properties of SnO$_2$ at Elevated H$_2$S Gas Concentration

The H$_2$S gas sensing behavior of pure SnO$_2$ thick films was studied by using home-built static apparatus. Pure SnO$_2$ thick film resistance was measured by using half bridge method with temperature in the H$_2$S gas atmosphere. Time interval of operating temperature is 50ºC during the measurement of gas response. For measurement of H$_2$S gas sensitivity the temperature range varied from room temperature 35ºC to 250ºC. The gas sensitivity of the prepared SnO$_2$ thick film was calculated using equation 7.

\[
\text{% Sensitivity} = \frac{R_a - R_g}{R_a} \times 100 \quad \text{(7)}
\]

Where, $R_a$ - Resistance in air and $R_g$ - Resistance in H$_2$S gas atmosphere.

The concentration of H$_2$S gas in the current gas sensing study were taken in parts per million (ppm). Here four concentrations viz. 100 ppm, 200 ppm, 500 ppm and 1000 ppm of H$_2$S gas were used for gas sensing purpose. From Figure 7, SnO$_2$ thick film showed 63.8 % sensitivity to H$_2$S gas at operating temperature at 200ºC and gas concentration 500 ppm. H$_2$S gas also shows good response to 200 ppm at 200ºC. The response of H$_2$S gas slowly increases to 100 ppm concentration with
increase in temperature. It has been observed that when nanoparticles of SnO₂ were exposed to the H₂S gas, the resistance of thick film decreases because the nature of H₂S gas is reducing. At 500 ppm concentration of H₂S gas SnO₂ thick film show more sensitivity as compare to 100 and 200 ppm. It has been also found that at 1000 ppm of gas concentration sensitivity decreases.

**H₂S Gas Sensing Mechanism Tosno₂ Thick Film Sensor**
The SnO₂ thick film sensor shows good response to H₂S gas. SnO₂ is n-type semiconductor and the majority of charge carriers in n type semiconductors are electrons. H₂S is reducing gas, when a molecule of hydrogen sulfide gas comes in contact with the surface of SnO₂ thick film sensor the sample resistance decreases due to adsorption reaction and sensor gives response to H₂S gas.

**Response Time and Recovery Time**
The thick film sensor was kept at 200°C optimum temperature. Then, H₂S gas was injected using syringe in the glass dome of the gas sensing system for measurement of response time. To determine response time variation in resistance is measured with time. From figure 8 it was observed that response time is 08 seconds whereas recovery time was 20 seconds to SnO₂ thick film sensor for H₂S gas at concentration 500 ppm and at 200°C optimum temperature.

**Reusability of SnO₂ Thick Film Sensor for Hydrogen Sulphide Gas**
The SnO₂ thick film sensor reusability experiment was carried out in four cycles with a 15 days interval between each cycle. The main goal of the reusability study was to see whether the results of a hydrogen sulphide gas sensor could be repeated. The optimum response for H₂S provided by SnO₂ was 63.8 %, and the experiment looked into whether the thick film sensor could replicate the same results. In the second turn the optimum response for H₂S gas was 62.48%, in the third the response found to be 61.13% and in the final set, the optimum response was recorded as 60.65%. It may be due to slight variation in composition of the prepared thick film sensor of SnO₂, minimum decline in response for H₂S was observed after 15 days interval in four turns. Figure 9 depicts the reusability outcomes of the experiment.

**Conclusions**
From the obtained results, nanocrystalline powder of SnO₂ has been fabricated by using co-precipitation
method. SnO$_2$ thick films were prepared by standard screen printing technique on glass substrate. From XRD data it is observed that synthesized SnO$_2$ nanoparticles belongs to tetragonal lattice. The SnO$_2$ thick film sensor showed nearly 63.8 % sensitivity at 200°C temperature. The structural and electrical investigation supports SnO$_2$ is good semiconducting material with optimum activation energy. The reusability experiment performed for SnO$_2$ thick film sensor proved that the sensor is promising for hydrogen sulfide gas vapors for long time stability and reusability.

Acknowledgments
Authors are very thankful to Department of Electronic Science, L. V. H. College, Panchavati, Nasik and Department of Electronic Science, Fergusson College, Pune for providing necessary laboratory facilities. Authors are gratefully acknowledged to the Department of Physics, Savitribai Phule Pune University, Pune providing facilities for XRD,SEM analysis.

Funding
This research received no specific grant from any funding agency.

Conflict of Interest
The authors do not have any conflict of interest.

References


