Preparation and Gas Sensing Properties of Orderly Porous Tin Dioxide

Junliang Liua, Yiyang Zhange, Chengwu Zhoue, Haiyan Chena,c, Ming Zhanga*

a School of Chemistry and Chemical Engineering, Yangzhou University, Yangzhou, PR China, 225002
b School of System Informatics, Kobe University, Kobe, Japan, 657-8501
c Yangzhou Polytechnic College, Yangzhou, PR China, 225127

*Corresponding Author: lxyzhangm@yzu.edu.cn

Abstract

The orderly porous tin dioxide materials have been fabricated by a templated method with using mono-dispersed polystyrene microspheres as the templates. The as-synthesized porous tin dioxide materials showed a good sensing property with a low optimum working temperature of 310 °C and a wide sensor response of alcohol, a fast response time of 27 s and a short recovery time of 31 s (as the alcohol concentration was 200 ppm), which was much better than those of the sol-gel derived tin dioxide ultrafine powders.

Keywords: Tin dioxide, porous, gas sensing.

1. Introduction

Tin dioxide as one kind of wide bandgap semiconductors is characterized of excellent gas sensing properties(1-2), which gives it a broad application in commercially manufactured sensors for toxic and combustible gases. To enhance its sensitivity and decrease its working temperature, the design of porous order structures is believed to be one of the feasible ways(3-7). The designed structures with higher specific surface and porosity could improve sensitivity, and reduce response and recovery times(3-7). Focusing on enhancing gas sensing properties of tin dioxide materials, many efforts have been made and the results are encouraging. In this paper, a polymer templated method has been designed and used to fabricate orderly porous tin dioxide materials, which showed a good alcohol-sensing property.

2. Experimental

2.1 Preparation

The porous tin dioxide was synthesized by a templated method, which was briefly described as follows: 1) Stannous oxalate as the tin source was dissolved in the citric acid aqueous solution, and then triethanolamine was employed to adjust the pH value to 6-7. After stirring and reacting for 8h, a transparent yellow sol was obtained. 2) Mono-dispersed polystyrene micro-spheres with the particle size about 400 nm were fabricated via a dispersion polymerization(8) and dispersed in the above yellow sol. The mixture was centrifugated and the precursor was obtained. After re-dispersed in alcohol, the polystyrene microspheres combining with tin sol were assembled under the gravity force with evaporation of alcohol at 35 °C. 3) templates with tin sol were transferred into a crucible and place in a muffle furnace. The polystyrene templates were removed after heat-treatment at 500 °C for 2 h and the porous tin dioxide powders were collected. For comparison, the conventional tin dioxide powders were fabricated by a sol-gel method with stannous oxalate as the starting material and citric acid as the chelating agent.

2.2 Characterization

The phase compositions of the synthesized tin dioxide powders were identified by the X-ray diffraction measurement with Cu Kα (Bruker, D8 Advance). Their morphology was analyzed by scanning electronic microscope (Hatachi, S4800). Their gas sensing properties were characterized by a WS-30A gas sensitivity instrument (Zhengzhou Winsen Electronics Co., Ltd., Henan, China). The pastes containing tin dioxide powders were uniformly coated on a ceramic tube with electrodes. Prior to the test, the assembled samples were aged at 300 °C for about a week. The alcohol was heated to form the gas source, injected into a glass chamber and mixed with ambient air. The gas sensing properties were determined according to the testing principle of gas sensor(5, 9). The sensor response (S) was determined by
the following formula:

\[ S = \frac{R_a}{R_g} \]  

(1)

Where \( R_g \) and \( R_a \) were calculated by:

\[ \frac{R_g}{R_a} = \frac{R_L (V_c - V_g)}{V_c} \]  

(2)

\[ \frac{R_a}{R_g} = \frac{R_L (V_c - V_a)}{V_a} \]  

(3)

\( R_L \) is the load resistance with a certain value, \( V_c \) is the supplied circuit voltage. The output voltage \( (V_{out}) \) reflects the immediate voltage of the load resistor in the entire testing process, and it is described as \( V_a \) and \( V_g \) when the sensor is loaded in ambient air and alcohol gas.

3. Results and discussion

Fig.1 gave the XRD patterns of the as-synthesized porous tin dioxide and sol-gel derived tin powders. All the diffraction peaks were indexed and ascribed to tetragonal rutile tin dioxide and the relative intensities of the diffraction peaks were very similar to JCPDS No. 41-1445, indicating the successful formation of single phase tin dioxide without obvious impurity phases. There was no carbonizing product of polystyrene remaining in the synthesized porous tin oxide. Also, the sol-gel derived powders consisted of single phase tin dioxide.

Fig.1.  The XRD patterns of the synthesized porous tin oxide(up) and sol-gel derived powders (down).

Referring to the morphology of the two kinds of tin dioxide. As shown in Fig.2, the as-synthesized tin dioxide via the designed template method demonstrated a porous structure with the homogenous pores of about 200 nm in it. The pores were of high orderly arrangement owing to the self-assembly of the polystyrene micro-spheres. The thickness of the pore walls was in the nano scale about 40 nm. Such a highly order and porous structure may increase the contacting area between tested gas and SnO2 surface and provide abundant diffusion channels for gas molecules, which was expected to have a good gas sensing property. For comparison, the sol-gel derived tin dioxide ultrafine powders were of a broad size distribution in the range of 200 nm to 1 \( \mu \)m and with a severe agglomeration.

Fig. 2.  The SEM images of the synthesized porous tin oxide(up) and sol-gel derived powders (down).

Fig. 3.  The output voltage as a function of testing time as loading alcohol gas with various concentrations.

Fig.3 demonstrated the output voltage-testing time curves with increasing alcohol concentration, indicating the gas sensor with the as-synthesized porous tin dioxide approached a stable sensing ability in a wide concentration.
range from 10 to 200 ppm and the response and recovery time appeared to be very short. The sensor response as a function of alcohol concentration was shown in Fig. 4. As the alcohol concentration increased, the sensor response grew up continuously and approached a near saturation value as the alcohol concentration was over 1000 ppm, which certified it could be utilized as a good gas sensor for alcohol.

Moreover, the sensor response-working temperature curves were reflected in Fig.6. It was found that: 1) the gas sensor fabricated by the as-synthesized porous tin dioxide had a higher sensor response all through the working temperature range from 160 to 420 °C, comparing to the gas sensor fabricated by the sol-gel derived tin dioxide. 2) The optimum working temperature decreased from 370 °C to 310 °C as the successful formation of the high orderly and porous structure.

4. Conclusions

In conclusion, high orderly and porous tin dioxide with pore size of 200 nm and wall thickness of 40 nm has been successfully fabricated by using the polystyrene microspheres as the templates. The formation of the high orderly and porous structure provided a larger contacting area and more abundant diffusion channels, which endowed the gas sensor with the as-synthesized porous tin dioxide a good sensing property to alcohol: a low optimum working temperature of 310 °C, a wide sensor response of alcohol, and a faster response about 27 s as well as a short recovery time of 31s.

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