Nanocasting Synthesis of Ultrafine WO$_3$ Nanoparticles for Gas Sensing Applications

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Abstract Ultrafine WO$_3$ nanoparticles were synthesized by nanocasting route, using mesoporous SiO$_2$ as a template. BET measurements showed a specific surface area of 700 m$^2$/gr for synthesized SiO$_2$, while after impregnation and template removal, this area was reduced to 43 m$^2$/gr for WO$_3$ nanoparticles. HRTEM results showed single crystalline nanoparticles with average particle size of about 5 nm possessing a monoclinic structure, which is the favorite crystal structure for gas sensing applications. Gas sensor was fabricated by deposition of WO$_3$ nanoparticles between electrodes via low frequency AC electrophoretic deposition. Gas sensing measurements showed that this material has a high sensitivity to very low concentrations of NO$_2$ at 250°C and 300°C.

Keywords WO$_3$ · Nanocasting · Low frequency AC electrophoretic deposition · NO$_2$ gas sensor

Introduction

Semiconductor gas sensors offer good advantages with respect to other gas sensor devices due to their simple implementation, low cost and good reliability for real-time control systems [1–3]. Among them, ZnO, WO$_3$ and SnO$_2$ have widely been studied [4]. Tungsten oxide (WO$_3$) is a wide band gap, n-type semiconductor that has interesting physical and chemical properties [5–9]. WO$_3$ is a good photochromic, electrochromic, thermochromic and catalytic material [9–12].

Gas sensing is a surface effect. As gas adsorption occurs at surface level, an increase in the active surface area of the semiconductor oxide would enhance the properties of materials used as gas sensors [4, 13–15]. Nanostructured mesoporous materials have been widely studied in the development of gas sensing and catalytic systems, due to their large, controllable pore size and high surface area [1, 13, 16–18]. Nanocasting is a novel methodology for preparing nanoparticles by using mesoporous materials as a template to accommodate different oxides. In this synthesis pathway, the surface and voids of a preformed mesoporous solid are coated with desired precursors. The subsequent mineralization of these precursors and removal of the former solid templates may lead to mesostructures with other compositions and crystalline frameworks [1, 13, 19].

While using in chemical sensors, the sensing material should be aligned and connected between the electrode structures, so that the change of their properties is converted into electrical signal, when exposed to the measurand. Several approaches have been reported in the literature for placement of the nanomaterials between the electrodes, but most of them do not offer selective settlement of particles or need expensive apparatus and high vacuum [20]. Using electric field is one of the most versatile techniques known till date for selective and inexpensive assembly of nanoparticles [21].

In this research, we applied a nanocasting route for synthesis of ultrafine WO$_3$ nanoparticles. In the first step, mesoporous silica (SBA-15 structure) was prepared using a soft template, and in the second step, this material was used as a hard template in nanocasting of WO$_3$ nanoparticles. Also, alternating electric field was used to manipulate the
sensing material between electrodes, in order to fabricate gas sensor.

**Experimental**

Mesoporous SiO$_2$ was synthesized, using bottom-up method. Polyethylene oxide–polypropylene oxide block copolymer was used as soft template and tetraethyl orthosilicate (TEOS) as silicate source. In the first step, 6 gr of the copolymer, 180 cc distilled water and 30 gr HCl (35% wt) were mixed together, and the solution was stirred for 6 h at 35°C. In the second step, TEOS was added to this acidic solution and stirred for 24 h at the same temperature and then heated at 100°C for another 24 h as a hydrothermal treatment. Small volume of NH$_3$ was added in order to increase the PH up to 7. Then, the product was dried at room temperature in air atmosphere and calcined at 550°C for 4 h. The solid product of this process is mesoporous SiO$_2$, which will be used as nanotemplate in production of WO$_3$ nanoparticles.

The synthesis of WO$_3$ nanoparticles was carried out as follows: equal amounts of phosphotungstic acid hydrate and mesoporous SiO$_2$ were dissolved in ethanol, stirred vigorously for 1 h, dried at room temperature at air atmosphere and heated at 570°C for 5 h. The SiO$_2$ hard template was removed by HF (2% wt), and the product was collected by centrifugation. The product was washed both with water and ethanol for three times and dried at room temperature.

For sensor fabrication, WO$_3$ nanopowders were deposited on an alumina substrate between electrodes, via low frequency AC electrophoretic deposition [21–23]. For this purpose, a suspension of 1 gr/l WO$_3$ nanoparticles in acetone was prepared and ultrasonicated for 15 min to become stable and homogenous. Deposition was done by applying a voltage of 25 v and a frequency of 1 KHz. The sensor was calcined for 1 h at 58°C in an electric furnace and air atmosphere.

Synthesized powders were characterized by X-ray diffraction (XRD, Siemens D500, cu K$_x$ radiation) and transmission electron microscopy (HRTEM, JEOL2000 and TEM, Phillips). Nitrogen adsorption–desorption measurements were done using Gemini2375 instrument to determine the Brunauer–Emmett–Teller (BET) surface area. Response of the sensors toward various concentrations of NO$_2$ was measured using a normal set-up. The gas response was calculated as $R_a/R_g$, where $R_a$ and $R_g$ represent samples’ resistances without and with the presence of gas, respectively.

**Results and Discussion**

TEM image of mesoporous SiO$_2$ and HRTEM image of WO$_3$ nanoparticles are shown in Fig. 1a and b, respectively. The results revealed that WO$_3$ replica was constructed by hexagonally packed nanoparticle array in the two-dimensional SBA-15 structure of SiO$_2$. These nanoparticles were randomly oriented in the mesostructured framework and rather uniform in diameter (~5 nm) due to the confined growth in the channels of the mesoporous silica template. The d spacing has been measured to be 3.8 Å, which is in accordance with (001) planes in monoclinic WO$_3$ crystal structure.

![Fig. 1 a TEM image of mesoporous SiO$_2$; b HRTEM image of a single WO$_3$ nanoparticle](image1.jpg)

![Fig. 2 XRD pattern of synthesized WO$_3$ a before and b after removing the silicate template](image2.jpg)
BET measurements showed a surface area of about 700 m²/gr for mesoporous SiO₂ template that was reduced to 43 m²/gr for WO₃ nanoparticles.

XRD patterns before and after removing the SiO₂ template are shown in Fig. 2, indicating that WO₃ is fully crystalline with monoclinic crystal structure, which is in a good agreement with TEM results. The results show that SiO₂ peaks were removed after washing with HF and pure WO₃ has remained.

In AC electric field, the nanomaterials align in a highly uniform way and orientate along the electric field direction by the electrophoresis force. As it can be seen from the SEM images in Fig. 3a and b, nanoparticles assemble along the direction of electric field, connect end to end and bridge the electrode gap. Based on the SEM images, AC electrophoresis has the potential for the precise manipulation of nanoparticles, which would be useful in making micro/nano electronic devices.

Optical microscopy image of fabricated sensor via AC electrophoretic deposition is shown in Fig. 4. Obviously, WO₃ particles have connected the electrodes, by bridging over them as a result of AC electrophoresis forces.

The sensing properties of fabricated sensor exposed to dilute NO₂ were investigated in a dynamic mode switching from air to the specific concentration of the target gas and again back to the base air in each cycle. Figure 5a and b show the response transients and sensitivities of synthesized WO₃ nanoparticles at 200°C. Sharp response of the sensor to NO₂ (Fig. 5a) is an indication of sensor quality and is a direct consequence of extremely small particle size and high specific surface area. The sensor needed a rather long time to reach its original level after switching back to the air. Sensitivity values (defined as the ratio of sensor resistance in target gas over its resistance in base air) presented in Fig. 5b follow a line presenting the device as a trustable NO₂ sensor.
Sensing test results prove that the synthesized nanoparticles are sensitive to low concentrations of NO₂, which may be a consequence of ultrafine particle size that leads to higher active surface. Application of electric field for deposition of particles also improves the sensing properties of the fabricated gas sensor, thanks to precise placement of particles side by side and establishment of a good connection between the electrodes facilitating the charge transfer between electrodes.

Conclusions

Ultrafine WO₃ nanoparticles rather uniform in diameter (~5 nm) were synthesized efficiently by nanocasting route. AC electrophoresis was applied to deposit WO₃ nanoparticles between electrodes in order to fabricate the gas sensor. The sensing measurements showed that the synthesized particles are sensitive even to very low concentrations of NO₂ (100 ppb), which is a result of very small particle size combined with application of proper deposition method.

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