Synthesis, Characterization, and Activity of Tin Oxide Nanoparticles: Influence of Solvothermal Time on Photocatalytic Degradation of Rhodamine B

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ABSTRACT
The SnO2 spheres-like nanoparticles have been successfully synthesized by a microwave solvothermal method, in which SnCl2·2H2O, poly(vinylpyrrolidone) PVP, H2O2 and NaOH as raw materials. The as-synthesized products have been characterized by X-ray diffraction, scanning electron microscope, and UV/Vis/NIR spectrophotometer. Photocatalytic activities of the samples have been evaluated by the degradation of rhodamine B (RhB) under UV-light illumination. Results showed that these products with diameter about 1 - 2 µm, and when the reaction time prolong, the surface of the SnO2 spheres will change to rough and then smooth when the time even longer. The product with nanorods on its surface shows the higher photocatalytic activity and red shift in the UV-vis absorption, which are relative to the unique structure. At last we studied the electron transfer reactions during photo-oxidation of RhB.

Keywords: SnO2; Nanoparticles; Photocatalytic; Solvothermal

1. Introduction
Environmental problems, especially, the sustained pollutions of water by various organic and metallic ion contaminants have been one of the most serious problems. And many efforts are dedicated to the remediation of environmental pollution [1-3]. For instance, photodegradation of organic compounds provides an available way to deal with the water pollution. Nanostructured semiconductors (such as TiO2, ZnO, SnO2 and so on) are proved to be an excellent photocatalyst which can degrade many kinds of persistent organic pollution [4-10].

Tin oxide (SnO2), as one of the most important semiconductor oxides, has been used as photocatalyst for photodegradation of organic compounds. The results indicate that SnO2 has exhibited photoactivity toward degradation of dye and other organic compounds [11,12]. However, just like other transition metal oxides photocatalysts such as TiO2 and ZnO, SnO2 suffer from low photocatalytic efficiency because of its wide-bandgap (energy of the band gap is about 3.6 eV) [13] and high recombination rates of photogenerated electron-hole pairs. This defects hinder SnO2 photocatalyst using widely and practically in the environmental application [14]. To overcome this problem, the fabrication of nanostructures provides an effective way.

The synthesis of TiO2-based photocatalysts has been reported in our previous papers [15-18]. Recently, SnO2-based photocatalysts were also synthesized by various approaches and have exhibited attractive performances [19-22]. In this work, we describe the synthesis of SnO2 nanoparticles via a microwave solvothermal method and discuss the effect of solvothermal time on the morphologies and nanostructures. The as-synthesized SnO2 nanoparticles were well characterised and their photocatalytic activities were evaluated by the photodegradation of Rhodamine B (RhB).

2. Experimental
2.1. Preparation
All the reagents used in this experiment were analytical grade and were used without further purification. In a typical preparation procedure, first, 1.353 g of SnCl2·2H2O was dissolved into 40 mL distilled water under conti-
nuous magnetic stirring to form white slurry. Then 1.44 g of NaOH and 5 mL of 30% H₂O₂ were introduced to the well-stirred mixture at room temperature with simultaneous vigorous agitation until NaOH dissolved completely. When the solution became transparent, 1.2 g of poly (vinylpyrrolidone) PVP (MW 30,000) was introduced. After several minutes of stirring, Subsequently, the obtained solution was transferred into five 100 mL teflon autoclaves, which was treated in a MDS-8 microwave hydrothermal system (manufactured by Shanghai Sineo Microwave Chemistry Technology Co. Ltd.) at 180°C for 30 min, 60 min, 90 min, respectively, and allowed to cool to room temperature naturally. The resulting white powder was collected from the bottom of the Teflon container after decanting the supernatant, washed several times with absolute ethanol and distilled water. Subsequently, the products were dried in vacuum at 60°C for 12 h for further characterization.

2.2. Characterization

Morphologies of the samples were observed by using a high-resolution field emission environmental scanning electron microscope (JSM-6700). All the images were obtained under high vacuum mode without sputter coating. X-Ray Diffraction (D/max-2200, Diffractometer with Cu Ka radiation) was used to verify crystal phase and estimate the crystal sizes of as-synthesized SnO₂ nanoparticles. Absorption spectrum was measured on a UV/Vis/NIR spectrophotometer (LAMBDA-950) in the wavelength range of 200 - 800 nm.

2.3. Photocatalytic Activity Measurement

The photocatalytic activity of as-synthesized SnO₂ nanoparticles was evaluated through the degradation of 5 mg/L Rhodamine B (RhB) in a BL-GHX-V multifunctional photochemical reactor (Shanghai Bilon Experiment Equipment Co. Ltd., Shanghai, China). The volume of the reaction solution was 250 mL (8 test tubes of 30 mL) into which 25 mg of photocatalyst was added and stirred for 30 min. The solution was dispersed by sonication, and then transferred to test tubes. Irradiation was provided by a medium-pressure Hg lamp (300 W), and the reaction temperature was kept about 25°C. Stirring was performed at all the times during the reaction. Sampling was also performed at regular intervals (every 15 min). The residual concentration of RhB was determined by measuring its absorbance at 554 nm using an UV/Vis/NIR spectrophotometer (LAMBDA-950).

3. Results and Discussion

3.1. SEM and XRD Analysis

XRD pattern of the as-prepared product is shown in Figure 1. All the diffraction peaks are quite similar to those of SnO₂, which can be indexed as the tetragonal rutile structure of SnO₂ with lattice constants of a = 4.738 Å and c = 3.187Å, which is in good agreement with the JCPDS file of SnO₂ (JCPDS 41 - 1445) [22]. No impurity diffraction peaks are observed, indicating the high purity of the final products. In additionally, when treated for 30 min, the diffraction peaks were broaden and weaken, due to the relative lower crystallinity and size-quantization effect of nanomaterials (shown in Figure 1(a)). The increase of crystallinity was corresponding to the reaction times during the solvothermal process, it clearly show when reacted 90 min, the sample has the most high crystallinity among these samples. According to the Scherrer equation, \[ D = \frac{K\lambda}{\beta\cos\theta}, \] the average crystal size of SnO₂ calculated from the main diffraction peak are about 9, 16 and 17 nm, respectively.

Scanning electron microscopy (SEM) image of the as-grown product were shown in Figure 2, the nanoparticles were seem to spheres, and the diameters about 1 - 2 μm are clearly observed. Higher magnification SEM image shown in Figure 2(b), the observed SnO₂ spheres are seem to be soft and have some holes, due to the effect of PVP. So we can indicate that the process of formation of the sphere is not completed and this is just a middle product. When the time prolong to 60 min, the spheres made up of numerous one-dimensional tetragonal prism nanorods with an average diameter of about 100 nm were clearly seen in Figure 2(d). Interestingly, when continue prolong the time to 90 min, the surface become smooth, but the nanorods also can be seen (shown in Figure 2(f)). I think the nano-sized particles (as shown in Figure 2(d)) around the sphere will gather on the surface of the sphere, and close the hole between nanorods. This process was simply shown in Figure 3, the nanoparticles will show this process in the solvothermal process, and when this will con-
Figure 2. FESEM images of the SnO$_2$ nanoparticles synthesized by microwave solvothermal method with different reaction time: (a) (b) 30 min, (c) (d) 60 min, (e) (f) 90 min.

Figure 3. Schematic diagram of the microwave solvothermal process with time increase.

continue occur when the time even longer. And this will also lead to the size of nanocrystals and diameter of the sphere increase. The BET surface areas of the sample were obtained from N$_2$ adsorption/desorption isotherms determined at liquid nitrogen temperature on an automatic analyzer (Micromeritics, ASAP 2010), and the BET surface area of the samples are 45.1, 56.5, 50.9 m$^2$/g, respectively.

3.2. UV-Vis Analysis

Figure 4 shows the absorption spectra of the samples, which are all nearly identical, indicating that their optical band gaps were also almost the same. The fundamental absorption edge of SnO$_2$ located in the UV region at about 325 nm. But they also show some differences, as shown in Figure 4 inset curves. The fundamental absorption edges were 327.5, 328.5 and 352.0 respectively. The products prepared during 60 min were shown a little red shift due to the unique nanostructures and creation of oxygen vacancies [15]. It will narrow the band gap and enhance the UV absorption.

Figure 4. UV-visible absorption spectra of the SnO$_2$ nanoparticles synthesized by microwave solvothermal method with different reaction time: (a) 30 min; (b) 60 min; (c) 90 min. Inset was show the absorption at the range of 250 - 450 nm.

3.3. Evaluation of Photocatalytic Activities

Photocatalytic activity of the synthesized SnO$_2$ products was evaluated by monitoring the change in optical absorption of an RhB solution at ~554 nm during its photocatalytic decomposition process. The kinetics of this reaction can be monitored by UV-vis spectroscopy as seen from UV-vis spectra measured at different times shown in Figures 5(a)-(c). It demonstrates RhB shows a strong absorption band at 554 nm and the addition of SnO$_2$ products leads to a decrease of the absorption band with time. The color of the dispersion disappeared, indicating that the chromophoric structure of the dye was destroyed. The order rate kinetics with respect to the RhB concentration could be used to evaluate the photocatalytic rate as done previously. As it clearly demonstrated the SnO$_2$ photocatalysts show higher photocatalytic activity. Figure 5(d) shows a comparison of the photocatalytic activities among SnO$_2$ photocatalyst treated in different times. Additional experiments in the absence of photocatalyst, was also present in Figure 5(d). It can be noted that a trend of the photocatalytic activity is as follow order: b > a > c. The photocatalytic can be attributed to UV absorption, and the products prepared during 60 min shows a red shift and shows the higher photocatalytic activity. The existing of the nanorods also increases the surface area and there is more reaction place during photocatalytic process [18].

At last, we study the mechanism for photocatalytic degradation of RhB, and the electron transfer reactions involved in the selective photooxidation of RhB with oxygen are proposed in Figure 6. When the SnO$_2$ photocatalyst is excited under light irradiation with greater energy than its band gap energy, it will cause the formation of the hole-electron pair in the SnO$_2$. Subsequently,
Figure 5. Photocatalytic degradation of RhB solutions under UV irradiation by the presence of SnO₂ nanoparticles synthesized in different time: (a) 30 min, (b) 60 min, (c) 90 min. The concentration of the reactants was as follow: [RhB] = 5 mg/L, [SnO₂] = 100 mg/L.

Hole (h⁺) with high activity may react with H₂O or hydroxyl groups adsorbed on the surface of the SnO₂, the formed hydroxyl radicals also have strong oxidizing activity Hole (h⁺) and Electron (e⁻) can react with the dye molecule in favor of its degradation directly and following mineralization. In the process, the RhB can interact with the photogenerated holes in the valence band (VB), and provides a direct chemical reaction between the dye and the photocatalyst [23].

4. Conclusion

In summary, we studied the SnO₂ photocatalyst prepared via microwave solvothermal process. When the reaction time prolong, the surface of the SnO₂ spheres...
will change to rough and then smooth when the time even longer. We also propose the modeling to explain this interesting phenomenon. The product with nanorods on its surface shows the higher photocatalytic activity and red shift in the Uv-vis absorption, which are relative to the unique structure.

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