Modification of the Surface Properties of Core-Shell Semiconductors and Their Effects on the Photodecolorization Activity and Adsorption

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Abstract
Semiconductor-based photocatalysts have been extensively studied for oxidative photodestruction of organic pollutants in wastewaters, releasing non-toxic substances such as Azo dyes. Various synthesized catalyst specimens were characterized to determine the correlation between preparation conditions (catalyst type, dopant, microstructure, preparation routes, optical and physicochemical properties) on the photocatalytic activity. Some researchers focused on the process parameters to optimize them to reach higher photoactivity. The specific surface areas, crystalline size, charge and pretreatment of the surface have significant effects on the physical and photocatalytic properties of the semiconductors. The surface sites of catalyst (TiO₂) were modified by doping ZnS nanoparticles in the form of Core-Shell structure and the photocatalytic activities were determined by using color degradation and hydrogen production tests. The dye adsorption isotherms of photocatalyst were determined using UV-Vis spectroscopy. The specific surface properties were determined from BET, Zeta meter and Particle size analyzer tests. Photocatalytic decolorization of AR and water splitting test were applied to understand the relation between the surface properties and the photocatalytic activity. The result indicated that core-shell prepared samples had different surface suitable sites to cooperate in photocatalytic reaction.

Keywords
ZnS, TiO₂, Nanocomposite, Core-Shell, Surface Properties, Photocatalysis

1. Introduction
Photocatalysis has recently become a common green nano material. Its applica-
tion in advanced oxidation processes was rapidly increased, because of world’s main problems, energy and fuel. The control of organic, inorganic and bacterial pollutions in water and air is an important measure in environmental protection [1] [2] [3].

The kinetic of heterogeneous catalysts like photocatalysts is included five steps. These are: 1) diffusion of reactants to the surface; 2) adsorption of reactants onto the surface; 3) reaction on the surface. This step in photocatalytic reaction proceed mainly by the contributions of active oxygen species, such as hydroxyl radical, OH−, superoxide radical, O2− and hydrogen peroxide, H2O2; 4) desorption of products from the surface; and 5) diffusion of products from the surface [1].

In a photocatalysis system, photo irradiation inducing molecular transformations or reactions takes place at the surface of a heterogeneous catalyst; therefore, the surface properties and sites are so important [4].

The composite materials are engineered or naturally occurring materials made from two or more constituent materials with significantly interface between matrix and reinforcements [5]. The interest in nano composite materials stems from the fact that their totally properties are a function of their size, composition, interface and microstructures. Therefore, main strategies are required to prepare suitable nano composites reliably and predictably in order to optimize properties by nanotechnology. In this way, many nano composite structures were developed, consisting of nano structured particles or bulks. When two compounds were in the nano size scale, different composites were developed like bimetals, core-shells or multi layers. Similar to other nano materials, many processes were used to prepare these composites [6], like co precipitation, sol gel, solvothermal, microemulsion and hybrid methods and some self assembled techniques. Composite powders are utilized in the areas of coatings, electronics, catalysis, separations and drugs. The beginning of core-shell particles is also of interest from a fundamental and academic viewpoint, especially in the areas of composite and surface science. The core-shell structure is a composite in nano scale which both substrate and thin layer or individual particles on it are in nano size. They are applicable as model systems to investigate factors governing colloidal interactions and stabilization and to find valuable information on the properties of composite dispersions as nano particle engineering.

Controlled properties of surface in hetero catalysts are often accomplished by coating or encapsulating them within a shell of a preferred material, or from the other viewpoint, controlling the morphology and size of the coating by using templates is done for some applications. For example, the shell could alter the charge of the surface, functionality, and reactivity of the surface sites, and could enhance the stability and dispersibility of the colloidal powders. Magnetical, optical, or catalytical functions may be readily imparted to the dispersed colloidal matter depending on the properties of the coating. Encasing nano materials in a shell may also protect the core from extraneous environmental changes in drug delivery.
Core-shell composites often show functionalized specific properties over their single-component counterparts, and hence are potentially useful in a wide range of applications [5].

Optimization of the specific characteristics of the surface of nano materials through coating processes is the goal of many researchers. Polymer coating, inorganic, composite coating and Biomacromolecule Layer are used as a shell in synthesized core-shell particles in the nanometers size to multilayer of thin films. In the field of catalytic materials, applications of onion shape materials were very interests. Optimization of catalyst properties by controlling the specific surface area, the band gap position by quantum dot shells, the stability of catalyst in different conditions, and the reaction between electron and hole in the semiconductor powders were the goal of these materials. ZnO, CdSe, CdTd, CdS, ZnTe and similar compounds [7] [8] [9] were applied as a core shell system with different groups.

Photocatalytic decomposition of dye near the surface of the photocatalyst nanoparticles is related to their band gap energy and recombination sites. In this paper, the nano particle size effect on the optical activity studied and demonstrated significant increase in the resulting photocatalysis with decreasing the size of such immobilized nanoparticles for the first time. The higher levels of photocatalytic activity seen in the case of nano composites in compare with the cases of only TiO₂ with the same total metal-oxide nano particle concentration.

2. Materials and Methods

In the powder preparation procedure, nano size TiO₂ were used by hydrothermally prepared anatase powders according to kolenko method [10]. Then, the mixtures of ethanol (C₆H₅OH) and triethanolamine (TEA, C₆H₁₂NO₃) and TiO₂ powders were prepared. Thiocarbamide, (CH₃COO)₂Zn·2H₂O and water as the raw materials employed to produce ZnS shell’s according to Xu method [11]. Zinc acetate dehydrate [Zn(CH₃COO)₂·2H₂O] crystals were added to the initial solution under 400 rpm agitation until the pH of the solution was reached to about 9. To increase the pH of solution up to 11, sodium hydroxide (NaOH) solution was added. The resultant solution was then transferred into an autoclave at 140°C for 5 h. After the reaction was completed the autoclave was cooled down to room temperature naturally. To increase the precipitation rate and remove unwanted impurities, the precipitates were filtered, sonicated and washed with deionized water for several times and dried at 60°C in an oven. The obtained powders with (0, 50, 80 and 100 mole % TiO₂) were prepared.

The nano composites with (0.5% Pt) prepared by wet impregnation methods used as photocatalysis reaction. Other chemicals used in the experiments were analytical grade. X ray diffraction (XRD; SIEMENS D500; 1.784 nm CuKα), used to identify the crystalline size by means of the Scherrer’s equation, the ZnS, rutile and anatase phases was determined by internal standards The average specific surface area and open pore size distribution of powder was determined by analyzing the N₂ adsorption isotherm and calculated with the BET and BJH equa-
utions. The surface and particle morphology characterizations were also done by using transmission electron microscopy (TEM, Jcol Co., 1200EX, Japan). The surface charge was characterized by (Malvern, Zetasizer 3000HS).

The photocatalytic activities and adsorption isotherms of the samples were calculated in the photo degradation of aqueous Acid Red 14 Azar Rang co. (Iran), and water splitting in Fujishima’s reactor [3] which was used directly without further purification. In a typical procedure, the investigated photocatalyst sample was added into a 500 mL reactor which contains Azo dye. The concentration of the photocatalyst was estimated by UV-Vis spectrophotometer (UV-Vis 2500, Shimadzu, Japan). The pH value of the reaction batch reactor was about seven. The suspension was then transferred into the transparent photoreactor and irradiated with a high-pressure UV-C lamp. Samples were obtained at an interval of 10 min and the absorbencies of the samples were checked after paper filtering. Hydrogen production test was carrying out in a sealed batch pyrex reactor with monomer and septum link for sampling. Gas chromatograph (HP 4890, GC) applied to calculate hydrogen production rates.

3. Results and Discussion

The surface properties of powders are controlled by the preparation conditions. The crystalline size, phase composition (derived from XRD) and specific surface area (derived from BET) of nano size powders are given in Table 1. These powders applied to characterize photocatalytic activities with various phases’ ratios, crystalline size, and reaction time. The phase ratios did not change by secondary heat treatment and Shell preparation process adjust to reach similar crystalline size and surface area [4].

Figure 1 illustrates TEM images of powders synthesized at different conditions with different precursors. The images show that as-synthesized powders have different particle morphology. The ZnS and TiO₂ powders consist of cubic like and spherical particles, respectively but composite powders seem to be similar to core shapes. Those images clearly show that with adjusting components ratio, the microstructure and surface of powders are controlled.

Table 2 shows the effect of particles on the rates of photodegradation of AR

<table>
<thead>
<tr>
<th>Reference</th>
<th>Synthesis</th>
<th>Phase composition</th>
<th>Particle size nm (XRD)</th>
<th>BET, m²·g⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>TiCl₄ solution, HCl 160C, 24 h, TEA [8]</td>
<td>Anatase 100%</td>
<td>20</td>
<td>80</td>
</tr>
<tr>
<td>A20Z</td>
<td>A + 20% Zn(Ac) hydrothermal</td>
<td>Anatase 80% + ZnS 20%</td>
<td>~20</td>
<td>85</td>
</tr>
<tr>
<td>A50Z</td>
<td>A + 50% Zn(Ac) hydrothermal</td>
<td>Anatase 50% + ZnS 50%</td>
<td>~20</td>
<td>70</td>
</tr>
<tr>
<td>Z</td>
<td>Zn(Ac) hydrothermal</td>
<td>ZnS 100%</td>
<td>20</td>
<td>50</td>
</tr>
</tbody>
</table>
Table 2. Decolorization and hydrogen production rates, adsorption wavelength and surface charge of composite powders synthesized by hydrothermal method.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Decolorization rate (min⁻¹)</th>
<th>Hydrogen production rate (micro mol H₂/gr cat·h)</th>
<th>Adsorption wavelengths (nm)</th>
<th>Surface change (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.23</td>
<td>532</td>
<td>361</td>
<td>2.7</td>
</tr>
<tr>
<td>A20Z</td>
<td>0.49</td>
<td>609</td>
<td>320</td>
<td>12.4</td>
</tr>
<tr>
<td>A50Z</td>
<td>0.37</td>
<td>281</td>
<td>332</td>
<td>24.1</td>
</tr>
<tr>
<td>Z</td>
<td>0.20</td>
<td>0</td>
<td>355</td>
<td>29.5</td>
</tr>
</tbody>
</table>

Figure 1. TEM images of powders synthesized at different conditions with different precursors according to Table 1: (a) A, (b) Z, (c) A20Z and (d) A50Z.

It can be seen from the chart that the order of photocatalytic activity of powders is A20Z > A50Z > A > Z respectively. Many papers on AOP claim that the photocatalytic efficiency of TiO₂ depends on its phases [12], and crystalline size [13]. This variability has been attributed to difference in the levels of semiconductor, the rate of electron-hole recombination [14] [15], and adsorption and desorption behavior of surface sits [15]. The mobility of charge-carriers and electron hole pair’s recombination rate are critical steps in the reaction rate. the band gap of ZnS is about 3.54 ev that could not contributed in this irradiation, and the Fermi level of ZnS is higher than anatase [1]. In mixed ZnS and anatase, sample (A) generally enhances the photocatalytic activity as seen in Table 2. This activity derived from the results of dye decolonization and water splitting tests. Thus, the surface of catalyst is the most effective structural parameter of kinetic in five step of reaction, because photocatalysis is an interfacial reaction. Thus, a higher specific surface area induces a higher number of accessible active sites. It has other effects on the e⁻/h⁺ recombination time [15] band gap energy, penetration of light, consequently, photocatalyst reactivity. The band gaps of powders are related to their ledge of adsorption which is indicated in Table 2.
The composite samples have shorter adsorption wavelength that is against the core and shell's band gap. Smaller 2D crystalline size of ZnS could show quantum size effect and broadening of adsorption range [16] but conductance and valance levels of two semiconductor indicated that electron and hole could transfer between these levels to decrease the band gap and recombination rate. Also, the surface charge of samples is collected in Table 2. The charge value for composite samples gradually exchanged from pure anatase to pure ZnS. This shows that ZnS and TiO₂ present on the surface area of composite. Increased reaction sites on the surface of the catalyst enhanced the reactivity.

The coupling or capping of two semiconductors provides a more efficient electron/hole separation, delayed charge recombination, and enhanced interfacial suitable sites [17] as shown in composite results. The summary table shows that this sample has good photodecolorization, it can be related to: 1) electron-hole transfer process (IPET) [16]; 2) increased surface sits.

Dye adsorption isotherms of powders show in Figure 2. The accumulation of the dye on the surface of ZnS is very low and on the anatase is limited, but on the composite samples is increased and not limited. The graphs for composite samples have some steps which indicate different mechanisms cooperated on the adsorption of dye on the surface sites.

4. Conclusion

In this study, photocatalytic activity of TiO₂/ZnS nano composites has been investigated using two systems. The reaction rate was related to the surface area, phase's ratio, particle size, adsorption mechanism of reactants, concentration of dye and catalyst. The surface reaction sites and photo reactivity are higher in
composite powders with similar particle size and band gap. The concentration of reactants (dye or water, H₂, O₂) is important because they react in suitable surface sites. The suitable sites are determined by size of compounds and surface charges. Additional to the surface properties, effectively charge separation increased the photo reactivity of composites.

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