Research Article

Synthesis, Characterization, and Photocatalysis of Fe-Doped TiO$_2$: A Combined Experimental and Theoretical Study

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Fe-doped TiO$_2$ was prepared by hydrothermal treating Ti peroxide sol with different amount of iron nitrate. Fe ions can enter TiO$_2$ lattice by substituting Ti$^{4+}$ ions, which significantly affect the crystallinity and morphology of TiO$_2$ nanoparticles. Fe doping also influences the UV-Vis absorption and photoluminescence of TiO$_2$, due to the change of electronic structure. It is shown that Fe ions are more easily doped on TiO$_2$ surface than in bulk. The theoretical computation based on the density functional theory (DFT) shows that the Fe ions in TiO$_2$ bulk are localized and mainly act as the recombination centers of photoinduced electrons and holes. Some results support that the Fe$^{3+}$ ions on surface can form intermediate interfacial transfer pathway for electrons and holes, which is beneficial for increasing the photocatalytic activity of TiO$_2$. The photocatalytic activity first increases and then decreases as the Fe concentration increases, which is affected by the bulk-doped and surface-doped Fe ions.

1. Introduction

Since the publishing of photocatalytic water splitting [1], photocatalysis has drawn much attention in the world for many years [2–4]. Compared with other photocatalysts, TiO$_2$ is extensively studied because of low-cost, non-toxicity, and high chemical-stability. Generally, pure TiO$_2$ suffers from problems of large band gap ($E_g$) [5, 6] and low quantum yield. Doping is proved to be effective to narrow $E_g$ and increases photocatalytic activity [7–14]. Transition-metal-doped TiO$_2$ have been widely studied due to their d electronic configuration [15–18]. Fe is the most studied because it has the radium identical to Ti$^{4+}$ and the half-filled d electronic configuration [19–21]. It was reported that the photocatalytic activity of Fe-doped titanate nanotubes is 2 times higher than P25 under visible light illumination [19], and the Fe-doped Q (quantum)-TiO$_2$ particles can remove chloroform under UV light illumination due to the trapping of photoinduced electrons and holes [20].

It is always reported that the photocatalytic activity under UV light illumination will firstly increase and then decrease with the increase of doped Fe concentration [17, 19, 20]. It is considered that the Fe ions can trap holes or electrons for low-level doping, while they will change to recombination centers for high-level doping [17, 19, 22]. However, the physical reason that the doped Fe$^{3+}$ ions mainly act as recombination centers is not well known. Although tunneling recombination is said to become significant for high doping level, no experimental proofs can support this conclusion. The research 20 devoted to studying the photocatalytic mechanism of transition-metal-doped Q-TiO$_2$ particles of 3-4 nm. In this case, Fe ions are mainly on surface due to the small particle size, which can form effective pathways for charge interfacial transfer. For example, Wang et al. reported that the Fe$^{3+}$ ions can combine with the hydroxyl groups on TiO$_2$ nanoparticle surface and form (OH)Fe$^{3+}$ complexes, which is a good way for electron
In this work, we aim to discuss the di-dopant on photocatalysis is not systematically investigated. Compared with the Q-particles, the specific surface area decreases for large particles (>15 nm), and it is possible for the Fe ions to enter TiO$_2$ bulk beside surface. Since the carrier exciton Bohr radius in TiO$_2$ is ca. 2 nm [20], the holes or electrons trapped at Fe ions in bulk are difficult to migrate to surface before going to recombination. It is also reported that the optimal Fe-doped concentration is dependent on the particle size [26], and the increase of particle sizes will result in the decrease of optimal concentration, which may be due to that more Fe ions that enter the bulk for larger particles can increase the recombination of electrons and holes. It predicts that the Fe ions on surface may increase the photocatalytic activity, while the Fe ions in bulk will have negative effect.

However, the different effect of bulk dopant and surface dopant on photocatalysis is not systematically investigated. In this work, we aim to discuss the different functions of Fe ions on surface and in bulk during photocatalysis under UV light illumination. It is found that the surface doping is dominant for low-level doping, and more Fe ions will enter TiO$_2$ bulk with the increase of doping concentration. The ab initio computation based on density functional theory (DFT) shows that the Fe ions in TiO$_2$ bulk are localized, indicating that they mainly act as the recombination centers of electrons and holes. It is suggested that the role change of Fe ions from trapping to recombination centers as the doped Fe concentration increases is related to the transformation from surface to bulk doping, and this transformation explains the photocatalytic behavior of Fe-doped TiO$_2$ observed by us and other researchers.

2. Experimental

2.1. Sample Preparation. The TiO$_2$ samples were synthesized using hydrothermal method. TiCl$_4$ and Fe(NO$_3$)$_3$ were used as the precursors of TiO$_2$ and Fe ions. 4.0 mL TiCl$_4$ was added to 400 mL ice distilled water under vigorous stirring, and then, the resulted solution was stirred for 24 h at room temperature. Subsequently, ammonia aqueous solution (1:9) was added to adjust the solution pH to ca. 7 to get Ti(OH)$_4$ precipitation. The precipitation was filtered and cleaned using distilled water for many times. Afterwards, the Ti(OH)$_4$ precipitation was dispersed in ice-distilled water. 30% H$_2$O$_2$ was dropped until it became yellow and transparent. A designed amount of Fe(NO$_3$)$_3$ was added under stirring, and then, the solutions were heated in enclosed autoclave for 12 h at 160°C to get Fe-doped TiO$_2$ hydrosols. A series of Fe-doped TiO$_2$ samples were prepared with the Fe: Ti molar ratio being 0, 0.1, 0.3, 0.5, 0.8, 1.0, 2.0, and 5.0 at. %, respectively. Finally, the hydrosols were dried at 40°C to obtain the powders.

2.2. Characterization. The X-ray diffraction (XRD) patterns were collected in the range of 20–80° (2θ) with a HZG4-PC diffractometer using CuKα radiation with an accelerating voltage of 40 kV and current of 40 mA ($\lambda_\alpha = 1.5406$ Å).

The surface composition and element chemical states of Fe-doped samples were characterized by an X-ray photoelectron spectroscopy (XPS) in an ESCA system with monochromatic Mg kα source and a charge neutralizer. The particle morphologies were observed with a JEM-2001 transmission electron microscope (TEM) at an accelerating voltage of 200 kV. Diffused reflectance spectra were recorded on a Shimadzu UV2550-UV spectrophotometer equipped with an integrating sphere assembly in the range of 190–700 nm, with BaSO$_4$ used as the reference. The photoluminescence (PL) spectra were measured at room temperature with Shimadzu RF-5301 fluorospectrophotometer by using 320 nm line of Xe lamp as excitation source. The energy-dispersive X-ray (EDX) equipped on the JSM-7500F field emission scanning electron microscope was used to detect the real element concentrations in the Fe-doped TiO$_2$ samples.

2.3. Computational Detail. The computational calculation has been performed within DFT plane wave pseudo potential method. The general gradient approximation (GGA) with the Perdew-Burke-Ernzerhof (PBE) functional and ultrasoft pseudopotentials was used to describe the exchange-correlation effects and electron ion interactions. The kinetic energetic cutoffs of 24 and 288 Ry for the smooth part of electronic wave functions and augmented electron density were used. Quantum-ESPRESSO code, Pwscf package [27], was used to perform ab initio quantum calculation. Monkhorst-Pack (M-P) k-space sampling was adopted for geometry optimization and density of the electronic states (DOS) calculation. The convergence threshold for the self-consistency energy calculation was set to $10^{-8}$ Ry, and all atomic relaxations were carried out until all components of the residual forces were less than $10^{-3}$ Ry/Bohr.

4.3. Photocatalytic Activity Evaluation. The photocatalytic activities of the Fe-doped TiO$_2$ samples under UV were evaluated on the basis of the degradation rate of methyl orange (MO) in a 100 × 20 self-designed quartz reactor containing 0.3 g of catalyst and 30 mL of 10 mg/L MO aqueous solution. Prior to photocatalytic reactions, the suspension was allowed to reach adsorption equilibrium by stirring in dark about 60 min. The irradiation was performed with a 365 nm 1300 μWcm$^{-2}$. 
UV fluorescent lamp. In addition, the blank experiment without using UV illumination and TiO₂ photocatalyst was also conducted for comparison. Generally, the photocatalytic degradation follows a Langmuir-Hinshelwood (LH) mechanism for the low concentration [28, 29], and the reaction rate equation can be expressed as ln(C₀/C) = kᵗ, where C₀ and C are the initial concentration and reaction concentration, respectively.

3. Results and Discussion

3.1. Characterization. Figure 1(a) shows the XRD patterns of the Fe-doped TiO₂ samples. All of the as-prepared samples are mainly composed of anatase TiO₂. There is a little brookite, and its content becomes less and less with the increase of Fe concentration, indicating that Fe doping can restrain the brookite formation to some extent. The average anatase crystallite sizes decrease with the increase of Fe concentration because the doping restrains the grain growth [30, 31]. It can be seen from Figure 1(b) that the anatase (101) peak shifts to low 2θ direction as the Fe concentration increases, which indicates the (101) plane spacing becomes larger and larger when more and more Fe ions are doped. Because the radius of Fe³⁺ is a little smaller than that of Ti⁴⁺, the widening of (101) plane spacing indicates that it is possible for some Fe³⁺ ions to enter the interstitial voids of TiO₂ lattice.

Figure 2 shows the TEM photos of pure (A-1), 2.0 at. % (B-1), and 5.0 at. % Fe-doped TiO₂ (C-1), with A-2, B-2, C-2 being their high-resolved photos. The pure sample is mainly composed of needles, sheets, and sticks. The TEM photos clearly show that the nanoparticles become smaller and smaller as the doped Fe content increases, which is in accordance with the XRD analysis. This is easy to be understood because doping foreign ions in TiO₂ lattice can actually prohibit the crystal growth. In addition, it can also be seen that the TiO₂ nanoparticles change to leaf-like morphology after Fe doping, indicating that Fe doping may also affect the particle growth. As shown in Figure 2, some white arrows are labeled to show the [001] crystal direction of anatase nanoparticles. The nanoneedles and sticks of the pure TiO₂ are long in [001] direction. After doping Fe, the length of [001] becomes shorter and shorter as the Fe-doped content increases. This result indicates that Fe doping may prohibit the nanoparticle growth in [001] direction.

Figure 3 shows the O₁s, Ti₂p, and Fe₂p core spectra of the pure and Fe-doped samples (2.0 and 5.0 at. %). Ti₂p peaks contain Ti₂p₁/₂ and Ti₂p₃/₂ subpeaks due to the spin-orbital coupling effect [32, 33]. The Fe doping almost has no effect on the Ti₂p peak, indicating that Fe doping does not change the coordination and chemical state of Ti⁴⁺ ions. Compared with the symmetrical O₁s peak of pure TiO₂ sample, a new broad peak at ca. 531.5 eV after Fe doping shows the absorption of hydroxyl groups on TiO₂ surface, which may indicate the formation of Fe³⁺(OH) groups. It is reported that the Ti₂p peak of N-doped TiO₂ shifts to low binding energy (BE) due to the bond weakening between nitrogen and titanium, implying that the nitrogen enters TiO₂ matrix by replacing oxygen ions [34]. In present research, the shifting of O₁s peaks to high BE shows that the Fe doping changes the chemical coordination of oxygen. In addition, the Fe₂p spectra show a positive shift as compared to those in Fe₂O₃, probably indicative of more positively charged surface Fe³⁺ [22]. The slight enhancement of Fe₂p BE is due to the Fe³⁺ diffusion into TiO₂ lattice and the formation of Fe–O–Ti bond.
The EDX area scanning technique was used to detect the average Fe on surface and in bulk. The 2.0 at. % and 5.0 at. % Fe-doped TiO₂ samples contain ca. 1.9 and ca. 4.9 at. % Fe ions. It can be seen that the Fe concentrations are very close to the designed values. The Fe content on surface checked using XPS is 8.2 at. % and 14.1 at. %, respectively. Therefore, it can be seen that the Fe ions are not uniformly distributed in but enrich on TiO₂ surface, in agreement with the study [35]. Since XPS is a detecting technique of surface chemical information of one and two atomic layers, it can be known that the Fe ions are enriched on TiO₂ surface. The ratio between the surface Fe concentrations and the designed values is ca. 4.31 and 2.82 for the 2.0 at. % and 5.0 at. % samples. Therefore, the surface doping becomes difficult gradually with the increase of doping concentration, which will make more Fe ions enter TiO₂ bulk.

Figure 4 shows the DRS spectra of the Fe-doped TiO₂ samples. The onset of the absorption edge for pure TiO₂ is at ca. 390 nm [36]. Those spectra show an obvious red shift after Fe doping, which is induced by the electron transition from Fe3d orbitals to TiO₂ conduction band (CB). The absorption peak at 480 nm is from the d→d transition (T₂g → A₁g) [37], and it becomes clearer and clearer with the increase of Fe concentration. Figure 5 shows the normalized PL spectra of Fe-doped TiO₂ samples. The peak at 3.43 eV is from the CB→valence band (VB) recombination [38], and its density decreases gradually with the increase of Fe concentration due to the increase of non-irradiative recombination. The electrons and holes trapped on surface contribute to the PL band from 3.0 to 3.2 eV [39]. This band intensity decreases with the increases of Fe concentration, which shows that that Fe doping has a significant effect on surface electronic structure. Since the PL signal is from the recombination, lower PL intensity may indicate lower recombination [40, 41], which indicates that the surface Fe ions may increase the probability of interfacial transfer of photoinduced electrons and holes. The PL peaks within 2.6 and 2.75 eV are suggested to be from the indirect recombination via bulk defects [42]. The PL shoulders from 2.2 to 2.45 eV are from the deactivation of the photoinduced electrons in [TiO₆] groups on TiO₂ surface [39], which also show that the Fe doping has little effect on the Ti–O chemical environment, in agreement with the XPS result.

3.2. Electronic Structure Investigation. Based on the experiments, DFT computations were performed. The projected density of states (PDOSs) is shown in Figure 6. It can be seen that the VB and CB of TiO₂ are mainly composed of O2p orbitals and Ti3d orbitals, respectively [42, 43]. For pure TiO₂ (Figure 6(a)), the Fermi level (Eₚ) locates in the middle of the forbidden band, indicating that VB is full filled, while CB is empty. The Eₚ of pure TiO₂ is ca. 1.9 eV, which is underestimated due to the well-known shortcoming of
exchange-correction functional in describing excited states. The theoretical study indicates that oxygen vacancy can be easily introduced by Fe doping [44], so the electronic structure of reduced anatase was also investigated, which is shown Figure 6(b). The $E_F$ shifts to CB because some $\text{Ti}^{4+}$ ions are reduced as $\text{Ti}^{3+}$, and some energy levels related to oxygen defects appeared at the CB bottom (labeled as the arrow). The $E_g$ is narrowed by downshifting the CB bottom ca. 0.2 eV, which is in agreement with other reports [45, 46]. For the supercell containing a $\text{Fe}^{3+}$ ion, the Fe3d orbitals split into two bands (Figure 6(c)). The upper hybrid band with CB is mainly composed of $d_{x^2}$ and $d_{x^2-y^2}$ (A2g), and the inter-gap band consists of $d_{xy}$, $d_{xz}$, and $d_{yz}$ (T2g), $\sim 0.5$ eV above the VB top. The T2g-A2g distance is about 1.8 eV. Because $E_F$ is in the middle of T2g band, the electrons mainly occupy T2g band, and the A2g band is empty. Therefore, T2g $\rightarrow$ A2g transition is possible, which is the d-d transition in the UV-Vis DRS spectra. In addition, the hybrid A2g band partly contributes to the DRS red shift. When an oxygen vacancy is introduced in the Fe-doped anatase (Figure 6(d)), the Fe3d orbitals are still composed of two subbands, one midgap band (T2g) and one hybrid band (A2g). However, the T2g band is ca. 1.2 eV above the VB top, and the T2g-A2g distance is reduced to 0.9 eV due to the weakening of the crystal field effect. The $E_F$ is almost at the top of A2g band because the electrons in oxygen vacancy are shared by the two adjacent Fe ions, which may be the reason that no $\text{Ti}^{3+}$ ions are detected from XPS spectra. Consequently, DFT computation gives a clear result that the Fe doping will induce localized bandgap states, namely, T2g band.

Figure 3: The O1s, Ti2p, and Fe2p core XPS spectra of 0.0, 2.0, and 5.0% Fe-doped TiO$_2$ samples.
3.3. Photocatalytic Performance Evaluation. Figure 7(a) shows the dependence of $\ln(C_0/C)$ on time of the Fe-doped TiO$_2$ samples photodecomposing MO aqueous solution. Figure 7(b) shows the UV-Vis absorption spectra of MO aqueous solution for different periods of UV light illumination for 0.3 at. % doped TiO$_2$. The blank experiment shows that the MO cannot be degraded in the absence of either TiO$_2$ or UV light. The Fe doping obviously influences the photocatalytic activity under UV light illumination. The 0.3 at. % Fe-doped sample has the best activity, better than the photocatalytic activity of P25. An optimal Fe concentration for the Fe-doped TiO$_2$ to present the best photocatalytic activity was observed, which is not a surprise because this has been seen for many times. Although it is stated that the Fe ions mainly form trapping centers for holes and electrons for low doping level, while they will become recombination centers for high doping [47], the mechanism of this trapping to recombination transformation is still unclear and in debate.

As shown by DFT calculation, the Fe$^{3+}$ ions in bulk mainly form the recombination centers of electrons and holes; more Fe ions entering TiO$_2$ bulk cannot contribute to the photocatalysis. Compared with the Fe$^{3+}$ ions in bulk, the Fe$^{3+}$ ions on surface have different coordination environment. They can directly combine with the oxygen molecule (Fe–O$_2$) or the hydroxyl groups (Fe–OH), so the electrons or holes trapped on surface Fe$^{3+}$ ions are possible to transfer to O$_2$ and –OH, resulting in the formation of O$_2^−$ and •OH, which has also been reported by others. Therefore, the Fe ions on surface can form intermediate pathways for the interfacial transfer. Because the surface recombination of the trapped electrons and holes is generally much slower than the interfacial transfer [48], the presence of Fe$^{3+}$ ions on surface is useful to improve photocatalytic speed.

For the present experiment, the Fe$^{3+}$ ions are not uniformly distributed in TiO$_2$ nanoparticles. In the case of low-level doping, they are mainly on TiO$_2$ surface. According to the above discussion, it can be known that the Fe ions mainly form the pathway for the electron-hole transfer for low doping concentration, so the photocatalytic activity will increase with the increase of doping concentration. For medium-level doping, many Fe$^{3+}$ ions can be doped in bulk beside those on the surface, so the photocatalytic activity will increase with the increase of doping concentration. For high-level doping, many Fe$^{3+}$ ions can be doped in bulk beside those on the surface, so the photocatalytic activity will increase with the increase of Fe content. It is widely accepted that the doped Fe$^{3+}$ ions will change from the trapping centers to recombination centers as the doped concentration increases, how this change takes place is still not clear. It is claimed that the tunnel recombination through Fe ions is the reason of this trapping to recombination changing. We give new explanation of the trapping to recombination changing with the increase of Fe concentration, which is related to surface doping and bulk doping, as illustrated above.

4. Conclusion

The Fe-doped TiO$_2$ nanoparticles are prepared using hydrothermal method. It is confirmed that Fe$^{3+}$ ions are...
mainly doped on TiO$_2$ surface for low doping concentration, and some Fe ions will enter TiO$_2$ bulk for high doping concentration. The experiments and DFT calculation show that the Fe$^{3+}$ ions in TiO$_2$ bulk are localized, so they mainly form the recombination centers. The Fe$^{3+}$ ions on surface can form bridges for the transfer of electrons and holes, so surface doping is beneficial to the photocatalysis. It is observed that the photocatalytic activity of as-prepared Fe-doped TiO$_2$ firstly increases and then decreases with the increase of doping concentration, which is due to the trapping to recombination changing of doped Fe$^{3+}$ ions. It is confirmed that this changing is related to the increase of bulk doping as the doped concentration increases.

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