# 가시광 감응 산화티탄(TiO2)

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# Visible Light Responsive Titanium Dioxide (TiO2)

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산화티탄은 가장 많이 연구된 반도체 산화물로 환경 정화와 에너지 생산에 응용이 크게 기대되고 있다. 공기와 물 속의유해 유기물을 제거하고 물분해를 통한 수소 생산은 대표적인 응용 분야이다. 산화티탄의 저렴한 가격, 낮은 독성, 화학적및 열적 안정성은 잘 알려진 장점이다. 그러나, 산화티탄의 단점은 가시광 영역에서 광촉매 활성이 낮다는 점이다. 이러한문제점을 해결하기 위하여, 귀금속, 금속, 양이온, 음이온 도핑 방법으로 산화티탄의 표면과 전기적 구조를 변형시켜 가시광 영역에서 광촉매 활성을 높이기 위한 연구가 많이 진행되고 있다. 이번 총설에서는 산화티탄의 가시광 감응을 유도하는 방법에 대한 광범위한 정보를 정리하였다.

Titanium dioxide (TiO<sub>2</sub>) is one of the most researched semiconductor oxides that has revolutionised technologies in the field of environmental purification and energy generation. It has found extensive applications in heterogenous photocatalysis for removing organic pollutants from air and water and also in hydrogen production from photocatalytic water-splitting. Its use is popular because of its low cost, low toxicity, high chemical and thermal stability. But one of the critical limitations of TiO<sub>2</sub> as photocatalyst is its poor response to visible light. Several attempts have been made to modify the surface and electronic structures of TiO<sub>2</sub> to enhance its activity in the visible light region such as noble metal deposition, metal ion loading, cationic and anionic doping and sensitisation. Most of the results improved photocatalytic performance under visible light irradiation. This paper attempts to review and update some of the information on the TiO<sub>2</sub> photocatalytic technology and its accomplishment towards visible light region.

Keywords: visible light photocatalyst, titanium dioxide, semiconductor, TiO2 modification

#### 1. Introduction

TiO<sub>2</sub> semiconductor is the most popular photocatalyst because of its excellent optical and electronic properties, low cost, non toxicity, chemical and thermal stability[1]. TiO<sub>2</sub> has been used for environmental remediation purposes such as in the purification of water and air and also in the solar water splitting[2-7]. Photocatalysis is by far one of the most superior technologies in the environmental purification because unlike many other technologies, photocatalysis does not serve as a mere phase transfer but completely degrades the organic pollutants by converting to innocuous substances such as CO<sub>2</sub> and H<sub>2</sub>O. Photocatalytic water-splitting using TiO<sub>2</sub> for hydrogen pro-

duction provides a promising method to obtain clean, low-cost and environmentally friendly fuel by solar energy. Therefore the improvement and optimization of  $\text{TiO}_2$  is an important task for heterogeneous photocatalysis in the future[8]. Many researches have been carried out not only to improve the photoactivity of  $\text{TiO}_2$  under UV light but also to improve its activity under visible light region.

A number of scientific reviews have been published on visible light responsive (VLR) TiO<sub>2</sub> photocatalyst[9-15]. The above review papers mainly cover the recent developments and progress of VLR TiO<sub>2</sub> in terms of photocatalytic water-splitting and degradation of organic pollutants. It also includes TiO<sub>2</sub> modification to produce VLR TiO<sub>2</sub> such as chemical additives (electron donors and suppression of backward reaction), noble metal loading, ion doping, sensitization and

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Table 1. History and Breakthrough of TiO2

Year	Comment
1791	- Discovery of the titanium element by Willam Gregor in England
1800's	- Fundamental chemical reactions of titanium
1821	- Discovery of TiO <sub>2</sub>
1916	- Commercial preparation of white pigment TiO2 by the sulfate method in Norway
1972	- Honda-Fujishima effect (photocatalytic reaction) - photo oxidation was used to chemically split water over $TiO_2$ to form $H_2$ and $O_2$ .
1980	- Photocatalysis on TiO <sub>2</sub> and Pt/TiO <sub>2</sub> powder (short circuit photoelectron chemical cell)
1985	<ul> <li>Photocatalysis on highly dispersed TiO<sub>2</sub> anchored on various supports</li> <li>Photocatalysis in the zeolite (cavities and frameworks)</li> </ul>
1987	<ul> <li>Ti/Si and Ti/Al binary oxide photocatalysts</li> <li>Size quantization effect on the photocatalysis of TiO<sub>2</sub></li> <li>Nanoscale TiO<sub>2</sub> particles</li> </ul>
1990	- TiO <sub>2</sub> coating on ceramic surface
1991	<ul> <li>Idea of sensitized photocatalysis (Grätzel solar cell)</li> <li>Purification and detoxification of polluted water</li> </ul>
1993	- Applications of TiO <sub>2</sub> photocatalysis in the purification of polluted water and air
1996	- Inactivation of pathogenic microorganism in air and water
1997	- New function of TiO <sub>2</sub> thin film photocatalyst (superhydrophilic phenomena)
1998	- Second generation TiO2 in powder and thin film forms (ion-implantation and RF magnetron sputtering method)
2000	- Applications of TiO <sub>2</sub> photocatalysts on a global scale
2001	- Photocatalytic water splitting under visible light irradiation
2006	- Preparation of TiO <sub>2</sub> from wastewater sludge for massive, environmental-friendly and functional TiO <sub>2</sub> production

metal ion-implantation. In this paper, we have made an attempt to review detailed effects of each ion and its different modification techniques to harvest visible light response TiO<sub>2</sub>.

# 2. Background

# 2.1. History of TiO<sub>2</sub>

Ever since the element titanium was discovered, TiO2 nanoparticles have drawn increasing interests as shown in Table 1[16-19]. Titanium element was first discovered by William Gregor in 1791, which he called menachanite but was named titanium four years later by Klaproth and the fundamental chemical reactions on which the present titanium industry is based were known before 1800, although it was not available in the markets until 1916[16]. Titanium dioxide (TiO<sub>2</sub>) was discovered in 1821 but it was not until 1916 that modern technology had progressed to the point where it could be massively produced. Fujishima and Honda[20] invented photo-electrochemical splitting of water over TiO2 to form H2 and O2 which opened up greater possibilities of solar energy conversion by semiconductors. The extensive knowledge that was gained during the development of semiconductor photo electrochemistry during the 1970 and 1980's has greatly assisted the development of photocatalysis and its application in the environmental purification. In 1995 TiO2 was used as a coating on ceramic surface by TOTO Company in Japan[21] while its application for inactivation of pathogenic micro-organisms in air and water was found in 1996. Since then, a number of photocatalyst applications have significantly developed.

#### 2.2. Properties of TiO<sub>2</sub>

Titanium is widely distributed over the surface of the earth and is the ninth most abundant element in the earth's crust comprising an estimated 0.62% of the earth crust[22]. The naturally occurring titanium ores are ilmenite, mineral rutile and brookite. Ilmenite is a black sand or rock and has the formula FeTiO<sub>3</sub>. The iron is partially oxidized to the trivalent state and ore also contains some impurities of silicon compounds. The TiO<sub>2</sub> concentration in the ore varies from 45 to 60% depending upon the origin. Rutile is naturally occurring TiO<sub>2</sub> and its colour vary from brown to reddish black, and its major impurities are iron compounds. The TiO<sub>2</sub> content in this ore normally varies from 90 to 95%.

Table 2 shows some of the physical and structural properties of  $TiO_2[23]$ .  $TiO_2$  exists in three different phases: anatase, rutile and brookite.  $TiO_2$  is chemically inert and thermally stable, non flammable and non toxic.

Electronic properties such as band gap play an important role for a semiconductor photocatalyst. Anatase phase  $TiO_2$  has higher band gap (3.2 eV) compared with other two phases of rutile (3.0 eV) and brookite. Anatase  $TiO_2$  is more popular as photocatalyst although rutile has been found to be effective under certain specific circumstances[7].

Figure 1 shows the photocatalytic mechanism of water-splitting to produce hydrogen and oxygen and the formation of the precursor species of hydroxyl radical and superoxide ions when illuminated under UV light. The energy level between Valence Band (VB) and Conduction Band (CB) is called band gap. At normal state, electrons and the protons remain in the VB but when  $\text{TiO}_2$  is excited by photon

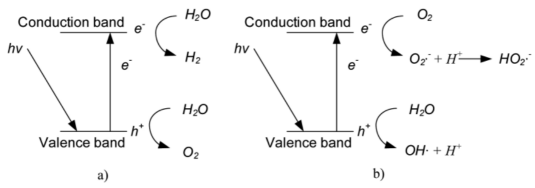


Figure 1. Mechanism of a) photocatalytic water-splitting from hydrogen production and b) formation of reacting species (hydroxyl radicals and superoxide ions) when illuminated under UV light.

Table 2. Physical and Structural Properties of Anatase and Rutile Structure of TiO<sub>2</sub>

Property	Anatase	Rutile	
Molecular weight (g/mol)	79.88	79.88	
Melting point (°C)	1825	1825	
Boiling point (°C)	$2500 \sim 3000$	$2500 \sim 3000$	
Specific gravity	3.9	4.0	
Light absorption (nm)	$\lambda \leq 385 \text{ nm}$	$\lambda \leq 415 \text{ nm}$	
Mohr's Hardness	5.5	6.5 to 7	
Refractive index	2.55	2.75	
Dielectric constant	31	114	
Crystal structure	Tetragonal	Tetragonal	
I -44: C(A0)	a = 3.784	a = 4.5936	
Lattice Constants(A <sup>o</sup> )	c = 9.515	c = 2.9587	
Density (g/cm <sup>3</sup> )	3.79	4.13	
Ti O hand langth (A9)	1.937 (4)	1.949 (4)	
Ti-O bond length (A°)	1.965 (2)	1.980 (2)	

with energy greater than its band gap energy level, electrons in the VB jump to the CB, creating e-/h+ pair. Because of its high band gap of 3.2 eV (anatase), only UV light source has been found to be effective in the excitation of the electrons in TiO<sub>2</sub>.

To produce hydrogen, water should be efficiently broken into hydrogen and oxygen by hydrolysis. When  ${\rm TiO_2}$  is irradiated by UV light,  ${\rm e^{\cdot}/h^{+}}$  pair is formed and when it comes into contact with absorbed water, it gets oxidized by positive holes which in the process forms hydroxyl radicals (OH $^{\cdot}$ ), having strong oxidative decomposing power and therefore reacts with organic matter. In the presence of oxygen, the intermediate radicals in the organic compounds and oxygen molecules undergo radical chain reactions and consume oxygen to finally form carbon dioxide and water as shown below:

$$TiO_2 (h^+) + H_2O_{ads} \rightarrow TiO_2 + OH_{ads} + H^+$$
 (1)

$$TiO_2 (h^+) + OH_{ads} \rightarrow TiO_2 + OH_{ads}$$
 (2)

$$OH_{ads}$$
 + organics  $\rightarrow CO_2 + H_2O$  (3)

Under certain conditions, organic compounds directly react with the

positive holes ( $h^+$ ) resulting in oxidative decomposition. In the presence of air, oxygen reduction takes place instead of hydrogen generation and forms superoxide anions ( $O_2$ ) which attach to the intermediate products in the oxidative reaction, forming peroxide or changing to hydrogen peroxide and then later to water.

$$TiO_2 (e^{-}) + O_{2,ads} \rightarrow TiO_2 + O_2^{--}$$
 (4)

$$TiO_2 (e^-) + H_2O_{2,ads} \rightarrow TiO_2 + OH^- + OH_{ads}$$
 (5)

#### 2.3. Preparation of TiO<sub>2</sub>

TiO<sub>2</sub> is produced using a variety of methods such as: i) sulfate method, ii) chloride method (vapor method), iii) alkoxide method, iv) specific method and v) sludge recycling method by Ti-salt flocculation in wastewater[18,19]. Among the methods, the sulphate process and the chlorine process are mainly used to produce TiO<sub>2</sub> powders. World production of TiO<sub>2</sub> is shared almost equally between these two production routes[24]. The chlorine process will continue to displace the sulphate process in the future due to the economic, environmental and market forces. Environmental problems with waste disposal costs associated with the large amount of acid and iron salt wastes which are produced during the sulphate and chorine process have resulted in the closure of certain U.S based plants[25]. To resolve the above problems, Shon et al.[19] developed a new solution to the sludge problem through a process which recovered economically valuable TiO<sub>2</sub> from wastewater. The work uses titanium tetrachloride (TiCl<sub>4</sub>) as an alternative coagulant instead of more commonly used salts of iron (FeCl<sub>3</sub>) and aluminum (Al<sub>2</sub>(SO<sub>4</sub>)<sub>3</sub>) to remove particulate and dissolved organic matter including persistent organic pollutant which is not removed in the conventional sewage treatment plants. TiCl4 coagulant successfully removed organic matter to the same extent as Fe and Al salts and moreover the floc size formed using titanium salt was bigger than that of Fe and Al salts which led to faster and effective settling. This technique is a novel method in solving the environmental problems associated with large quantity of sludge disposal using conventional coagulant. The settled floc was incinerated to produce functional TiO2 doped with carbon (C) and phosphorus (P) (TiO<sub>1.42</sub>C<sub>0.44</sub>P<sub>0.14</sub>) and up to 40 mg-TiO<sub>2</sub> per litre of synthetic wastewater was produced following TiCl4 flocculation.

Alkoxide can be used for the production of high purity TiO2 and thin film TiO2. Compared with different TiO2 preparation methods, TiO<sub>2</sub> by the alkoxide method is free from counter anions so that the effect of the anions on photocatalytic activity is marginal[18]. The alkoxide method is mainly used for TiO<sub>2</sub> coating. When the TiO<sub>2</sub> sol is coated onto a substrate, thin films are obtained which are used for many applications. Specific methods to produce thin film TiO2 consist of i) thermal decomposition of titanium isopropoxide vapour (amorphous TiO<sub>2</sub> of large surface area), ii) molten salt reaction of a mixture of titanium oxysulfate and alkaline nitrate (microcrystalline anatase TiO2), iii) laser induced decomposition of alkoxide (TiO2 microcystal), iv) hydrothermal oxidation of titanium metal in an autoclave (rutile TiO2), v) thermal reaction of titanium oxyacetylacetonate in organic media (microcrystalline anatase), v) high temperature hydrolysis of alkoxide with water dissolved in organic solvent (nanocystalline anatase and higher photoactivity), vi) ultrasonic technique, vii) microemulsion or reverse micelles and viii) thermal decomposition of alkoxide in inert organic solvents and high termperature hydrolysis of alkoxide with water liberated from solvent alcohols (nanocystalline anatase and higher photoactivity)[26].

#### 2.4. Applications of TiO<sub>2</sub>

TiO<sub>2</sub> is the most widely used metal oxide for environmental applications, cosmetics, paints, electronic paper and solar cells. As a pigment, TiO<sub>2</sub> has a remarkably high refractive index and an exceedingly high reflectance. It offers maximum opacity or hiding power as well as imparting whiteness and brightness. The above properties allow TiO<sub>2</sub> as a pigment in paints and coatings (including glazes and enamels), plastics, paper, inks, fibers, food and cosmetics. The material is used as an opacifier in glass and porcelain enamels, sunscreens and cement. As photocatalyst, TiO2 is used as photosensitiser for photovoltaic cells, and electrode coating in photoelectrolysis cells can enhance the efficiency of electrolytic splitting of water into hydrogen and oxygen. Photocatalytic activity of TiO2 results in thin coatings of the material exhibiting self cleaning and disinfecting properties under exposure to UV radiation. These properties make the material a candidate for applications such as medical devices, food preparation surfaces, air conditioning filters, and sanitary ware surfaces. As oxygen sensor, even in mildly reducing atmospheres TiO2 tends to lose oxygen and become sub stoichiometric. In this form the material becomes a semiconductor and the electrical resistivity of the material can be correlated to the oxygen content of the atmosphere to which it is exposed. Hence TiO2 can be used to sense the amount of oxygen (or reducing species) present in an atmosphere. The followings are the main different applications of TiO<sub>2</sub>.

#### 2.4.1. Energy Applications of TiO<sub>2</sub>

Hydrogen is considered as an ideal fuel for the future. However renewable energy contributes only about 5% of the commercial hydrogen production via water electrolysis, while other 95% hydrogen is mainly from fossil fuels. Nano-sized  $TiO_2$  photocatalytic water-split-

ting technology has great potential for low-cost, stable, non corrosive, environmentally friendly solar-hydrogen production to support the future hydrogen economy. The solar-to-hydrogen energy conversion efficiency is too low for the technology to be economically sound. The main barriers are the rapid recombination of photo-generated electron/hole pairs as well as backward reaction and the poor activation of TiO<sub>2</sub> by visible light.

In response to these deficiencies, many investigators have been conducting research with an emphasis on effective remediation methods. Some investigators studied the effects of addition of sacrificial reagents to prohibit rapid recombination of electron/hole pairs and backward reactions. The organic compounds act as sacrificial electron donors by consuming photo generated holes and/or oxygen, decreasing the rates of electron-hole recombination and increasing H<sub>2</sub>-production rates. EDTA, methanol, ethanol, CN, lactic acid and formaldehyde have been tested and proved to be effective to enhance hydrogen production [27-30]. Moreover, the decomposition of these hydrocarbons could be also contributed to a higher hydrogen yield since hydrogen is one of their decomposed products. Li *et al.*[27] reported that the addition of cyanide has improved the amount of hydrogen evolution over NiO/TiO<sub>2</sub> which is directly proportional to the cyanide concentration.

When complete liberalization of the sacrificial agent is achieved, the H<sub>2</sub>-production rate drops to steady-state values[31]. Photocatalytic decomposition of gaseous methanol on Pt/TiO2 nano-film can be carried out in a continuous-flow reactor for hydrogen production[32]. The pollutants can be acted as electron donors, so the photocatalytic production of hydrogen and decomposition of these pollutants happened at the same time. The present of carbonate salts was very efficient for hydrogen production using Pt loaded TiO2 semiconductor photo catalysts[33]. According to Sayama et al.[34] these carbonates salts can prevent backward reaction and thereby enhance the hydrogen production rate. According to Hashimoto et al.[35] the rate of hydrogen production from mixture of water and several aliphatic hydrocarbons when alkaline is added to the solution becomes 2~4 times lager than those without alkaline. Several aromatic compounds also serve as electron donors for the reaction of the present photocatalytic hydrogen production with Pt/TiO<sub>2</sub>. The rate of hydrogen production is about 5~10 times larger than those without alkaline. Other researchers focused on the enhancement of photo catalysis by modification of TiO2 by means of metal loading, metal ion doping, dye sensitization, composite semiconductor, anion doping and metal ion-implantation.

TiO<sub>2</sub> semiconductor photocatalyst loaded with noble metals like Pt can inhibit charge recombination and increase hydrogen production rate[36]. Since Pt is very expensive, low cost metals have been identified. Wu and Lee[28] deposited Cu particles on TiO<sub>2</sub> for hydrogen production from aqueous methanol solution. They reported that the maximum hydrogen production rate was ten times higher than that of blank TiO<sub>2</sub> and 4.5 times higher than P-25 TiO<sub>2</sub> for 1.2 wt% Cu. Anion doping, such as nitrogen doping and sulfur doping, is more effective the photocatalytic activity than metal ion doping[37,38]. Dye sensitization and composite semiconductor are very efficient methods

Photocatalyst	Time (h)	Amount of hydrogen evolved (micromoles)
·	1	370
Aqueous 1 mmol KCN and 1.5 M NaOH with NiO/TiO <sub>2</sub> catalyst	2	480
•		1000
	1	3000
Deposition TiO <sub>2</sub> with Cu particles from aqueous methanol solution	2	4200
	4	6200
100 mL of 4.9 $\times$ 10 <sup>-3</sup> M oxalic acid solution; 0.05 g 0.5 wt.% Pt-TiO <sub>2</sub>	1	60
	2	115
	4	220
	1	1200
Ar-calcined $TiO_2$ from water-methanol solution (vol.ratio = 1.4/1)		1400
	4	1450
N damed TiO (0.1 a) in 0.2 M No CO calution from 1990 and titanium totmobleside	1	270
N-doped TiO <sub>2</sub> (0.1 g) in 0.2 M Na <sub>2</sub> SO <sub>3</sub> solution from urea and titanium tetrachloride	2	450
(mole ratio $Urea/TiO_2 = 5$ )		1080

Table 3. Approximately Amount of Hydrogen Evolved after Different Times for Different Photocatalysts

to expand light response of  $TiO_2$  to visible region. Metal ion implantation with wavelength up to 600 nm is a promising modification technique for red shift of  $TiO_2$  photocatalyst.  $TiO_2$  is bombarded with high energy ions which are injected into the lattice and interact with  $TiO_2$ . The qualitative effectiveness of red shift observed to be the following order: Cr > Mn > Fe > Ni[39,40].

The hydrogen evolution rate still remains very low as can be seen from Table 3[27,28,41-43], the apparent energy conversion efficiency defined as the ratio of the combustion heat energy of hydrogen to the radiation energy from the Xe lamp to the reactor. Kida *et al.*[30] pointed out that according to some estimates an efficiency of  $10 \sim 15\%$  might be economical for hydrogen production from water using solar energy.

Based on the studies reported in the literature, metal ion-implantation and dye sensitization are very effective methods to extend the activating spectrum to the visible range[15]. They play an important role in the development of efficient photocatalytic hydrogen production to more than  $10 \sim 15\%$  under sunlight.

#### 2.4.2. TiO<sub>2</sub> Applications to Air Purification

The applications of photocatalysis have been expanded, and it has become more attractive. An air treatment system for ethylene removal has been placed in grocery stores to remove the naturally occurring ethylene that causes fruits and vegetables to spoil. TiO2 photocatalytic reactor traps and chemically oxidizes volatile organic compounds converting them primarily to carbon dioxide and water. Photocatalytic reactors are also modular and can be scaled to suit a wide variety of air quality applications. They operate at room temperature and with negligible pressure drop and therefore may be readily integrated into new and existing heating, ventilation, and air conditioning systems[44]. TiO<sub>2</sub> catalyst which is excited by plasma-induced high-energy particles (electrons, photons), resulting in an enhanced pollutant air removal cleaning system for such as tobacco smoke found in indoor environments[45]. Thin films of organic contaminants can be photo catalytically oxidized on TiO2-coated surfaces. The thickness of the organic layer oxidized per day is 1~5 pm, a rate sufficient for maintaining the cleanliness of a surface when the flux of contaminant is not excessive[46]. In the process of treating air streams,  $TiO_2$  must be suspended on some sort of surface to allow the gas to pass over it and react. This is usually some sort of matrix with a high surface area and UV light.

The main purpose of photocatalytic coatings is to maintain clean windows or windshields, strip finger marks off walls and to reduce the density of colonies of microorganisms on hospital walls. Toluene and formaldehyde are malodorous and cause indoor pollution. Ichiura et al.[47] prepared a composite sheets of TiO<sub>2</sub>-zeolite using a papermaking technique to decompose formaldehyde and toluene under UV irradiation. It was shown that these sheets are potentially applicable as highly functional materials to be placed on walls and ceilings of houses.

TiO<sub>2</sub> can be used to coat building materials, such as glass, tile, and cement[48]. Once coated, these sidings, roofs, or roads could remove outdoor air pollutants. These pollutants are converted into carbon dioxide and water vapor. Coating roadway systems are being considered for air pollution mitigation[49], and has already been used on several building structures in Italy[50,51]. Most of the exterior walls of buildings become soiled from automotive exhaust fumes, which contain oily components. When the original building materials are coated with a photocatalyst, a protective film of titanium provides the self-cleaning building by becoming antistatic, super oxidative, and hydrophilic. The hydrocarbon from automotive exhaust is oxidized and the dirt on the walls washes away with rainfall, keeping the building exterior clean at all times[52].

The glass covers on highway tunnel lighting fixtures darken from automobile exhaust, when TiO<sub>2</sub> coated lamp covers are used, the glass surface remains cleaner longer, and the number of required cleanings is greatly reduced[21]. The Japanese company TOTO developed TiO<sub>2</sub>-coated surfaces tiles for operating rooms, hospital applications and public restrooms on the basis of joint research with Fujishima, Hashimoto and their colleagues at the University of Tokyo[46]. Colonies of pathogenic microorganisms do not persist on them, even under low level irradiance from common fluorescent lamps.

TiO<sub>2</sub> coatings on sintered borosilicate glass were prepared employing a synthetic route based on the hydrolysis of titanium oxysulfate

(TiOSO<sub>4</sub>)[53]. The stability and lifetime of the coatings were tested for more than 4000 h of continuous operation. They have not noticed any appreciable degradation of the coatings was observed.

Air purifier with TiO2 can prevent smoke, soil, pollen, bacteria, virus and harmful gas as well as seize the free bacteria in the air by filtering percentage of 99.9% with the help of the highly oxidizing effect of photocatalyst TiO<sub>2</sub>[54]. It was powerful for the sterilization of floating germs. They noticed that the number of germs after seven months operation were almost the same as obtained in the first operation. Ao and Lee[55] have used TiO2 immobilized on activated carbon (TiO<sub>2</sub>/AC) filter for removing indoor air pollutant which was installed in an air cleaner. They found that the higher removal efficiency of using TiO2/AC filter is owing to the large adsorption capacity provided by the activated carbon. A 25% higher of nitrogen oxides (NOx) was achieved using TiO2/AC filter compared to the TiO<sub>2</sub>/AC only. Purification ability of TiO<sub>2</sub>/AC was found to be higher than single purification techniques in decomposing toluene, whether in high concentration or low concentration[56]. They found that the compound of TiO2 photocatalyst and active carbon has the advantage of active carbon absorption and TiO2 photocatalyst for decomposition.

#### 2.4.3. Water Applications of TiO<sub>2</sub>

Photocatalyst coupled with UV lights can oxidize organic pollutants into nontoxic materials, such as CO<sub>2</sub> and water and can disinfect certain bacteria in water. This technology is very effective in removing further hazardous organic compounds and at killing a variety of bacteria and some viruses in the secondary wastewater treatment. Heterogeneous photocatalysis is an alternative treatment method for decontamination of organic compounds and disinfection of microorganisms to purify potable water[57].

Disinfecting microorganisms such as E.coli is a great concern in drinking water supply facilities around the world. An annular type fluidized bed reactor has been constructed and tested to disinfect E.coli bacteria[58]. They prepared specially catalyst of (TiO<sub>2</sub> + glass bead + zeolite). The efficiency of E.coli removal was 99.99% by using the proposed technology. It has been found that the pilot scale is capable of degrading 84% of E.coli within 80 minutes compared with 77.3% by using UV only which has capacity of handling 46 litre/min. E.coli degradation in larger scale is being tested for the first time and its result compared with only UV destruction is promising. They recommended using this method on a continuous reactor to increases the potential for scale up. Peyton and DeBerry[59] used three different semiconductors (TiO2, ZnO, and Fe2O3) to decompose pollutant compounds in wastewater by using solar energy. They concluded that Fe<sub>2</sub>O<sub>3</sub> was found to be ineffective, both ZnO and TiO<sub>2</sub> catalyzed the removal of all compounds.

Yamashita *et al.*[60] has used the metal ion-implantation method to improve the electronic properties of the TiO<sub>2</sub> photocatalyst to realize the utilization of visible light. They investigated the properties of TiO<sub>2</sub> photocatalyst for the purification of water. The UV-VIS absorption spectra of these metal ion-implanted TiO<sub>2</sub> photocatalysts were found to shift toward visible light regions depending on the amount and the

kind of metal ions implanted. They were found to exhibit an effective photocatalytic reactivity for the liquid-phase degradation of 2-propanol diluted in water at 295 K under visible light ( $\lambda$  > 450 nm) irradiation. Zhanghai *et al.*[61] fabricated a composite nano-ZnO/TiO<sub>2</sub> photocatalyst with vacuum vaporized and solgel methods. They indicated that the nano-ZnO/TiO<sub>2</sub> film improved the separate efficiency of the charge and extended the range of spectrum, which showed a higher efficiency of photocatalytic than the pure nano-TiO<sub>2</sub> and nano-ZnO film. Under the optimal operation conditions, chemical oxygen demand (COD) values with the linear range of  $0.3 \sim 10.0$  mgl<sup>-1</sup> were achieved and these results were in good agreement with those from the conventional COD methods. Photocatalysis aided by TiO<sub>2</sub> nanoparticles is used in removing the organic chemicals which occur as pollutants in wastewater effluents from industrial and domestic sources.

In general, the main advantage of photocatalysis is that there is no further requirement for secondary disposal methods. Other treatment methods such as adsorption by activated carbon and air stripping merely concentrate the chemicals present by transferring them to the adsorbent or air and they do not convert them to non toxic wastes. Also as compared to other oxidation technologies, expensive oxidation methods are not required as ambient oxygen is used.

# 2.4.4. Pigment Applications of TiO<sub>2</sub>

The most important function of  $TiO_2$  however is in powder form as a pigment for providing whiteness and opacity to such products such as paints and coatings (including glazes and enamels), plastics, paper, inks, fibers, food and cosmetics.  $TiO_2$  is by far the most widely used white pigment.  $TiO_2$  very white and has a very high refractive index-surpassed only by diamond. The refractive index determines the opacity that the material confers to the matrix in which the pigment is housed. Hence, with its high refractive index, relatively low levels of  $TiO_2$  pigment are required to achieve a white opaque coating.

The high refractive index and bright white colour of TiO<sub>2</sub> make it an effective opacifier for pigments. The material is used as an opacifier in glass and porcelain enamels, cosmetics, sunscreens, paper, and paints. One of the major advantages of the material for exposed applications is its resistance to discoloration under UV light. Even in mildly reducing atmospheres TiO<sub>2</sub> tends to lose oxygen and become sub stoichiometric. In this form the material becomes a semiconductor and the electrical resistivity of the material can be correlated to the oxygen content of the atmosphere to which it is exposed. Hence TiO<sub>2</sub> can be used to sense the amount of oxygen (or reducing species) present in an atmosphere.

#### 2.4.5. Applications of TiO2 in Construction Materials

 $TiO_2$  photocatalyst have high application potential in the construction industry. Many applications of this material already exist, such as photocatalytic ceramics, self cleaning glass, photocatalytic cements and paints.  $TiO_2$  coating makes the material surface super hydrophilic and self-cleansing. Thin coating not only makes the surface wetting and rinsing more uniform so that water slides easily without

forming a droplets and float away organic surface contaminants. Moreover the organic dust that may come in contact with the surface will be removed by photocatalytic action in presence of sun light. This technology has allowed the creation of many exterior and interior facades and self-cleansing rooftops from ceramic and glass tiles[62]. More applications could be found in the future for building applications such as self cleansing pavements, walls, tunnels, curbs so on to remove NOx and SOx from the air produced by vehicular emissions.

Photocatalyst-modified cements are used for various applications in buildings, self-cleaning surfaces and solar-powered remediation devices for polluted waters[63]. They used a model reaction of photocatalytic atrazine degradation by white Portland cement samples modified with different semiconducting oxides TiO<sub>2</sub> to investigate a possible application of modified cements for the degradation of pollutants on building surfaces.

#### 2.4.6. Applications of TiO2 in Waste Treatment

The most common water treatment technologies for hazardous organic pollutants is using adsorption on granulated activated carbon and air stripping to destroy the contaminants. Its implementation on a large scale was prevented due to negative public perception and the potential hazard of incineration of organic toxic compounds. Hofstadlert and Bauer[64] recommended that one of the most effective approaches to resolve this problem is the mineralization of organic contaminants by using photocatalytic semiconductors (SrTiO<sub>2</sub>). They indicated that the potential is high enough to destroy most organic compounds to carbon dioxide and mineral acids.

Photocatalytic degradation using ultraviolet-irradiated  $TiO_2$  suspension has been investigated for destroying both free and complex cyanide with a concurrent removal of copper metal[65]. In contrast to conventional cyanide waste treatment processes, the photocatalytic processes convert both free and complex cyanide species into carbon dioxide and nitrogen with no residual harmful chemicals remaining. Results revealed that about 78% of free cyanide ( $10^{-3}$  M) was removed after illumination for 4 h in the presence of 1 g/L  $TiO_2$  at pH 11. Free copper ( $10^{-2}$  M) was completely removed in a shorter time of 3 h.

# 2.4.7. Oil Spills Applications of TiO2

Oil spills in the sea and ocean is one of the major environmental disasters in the marine environment. Researchers have been trying to study the effect of photodegradation in cleaning up the marine environment. The application of  $TiO_2$  photocatalysis has therefore been reported for the remediation of marine environment[66-68].

Photodegradation remains one of the least studied related to the crude oil and petroleum products spilled into the ocean or coastal waters. In fact, all the weathering process occurred simultaneously in the environment, and it is difficult to distinguish the changes induced by photodegradation from the changes induced by other process as biodegradation. Furthermore, the complex composition of crude petroleum and oil fractions makes the photodegradation process complicated. A multitude of photoinduced reactions could take place, pro-

ducing also a multitude of photodegradation products. Polycyclic aromatic hydrocarbons (PAHs) frequently appear in oil spills, petroleum fractions and coal. The possible applications in the cleaning up of marine oil spills have been considered[69]. The initial photoreactivity of petroleum distillates was found to depend greatly upon the aromatic fraction. PAHs are a class of persistent organic pollutants of special concern since they are carcinogenic and mutagenic[70]. PAHs were identified as the photosensitizing species responsible for most of the initiation reactions. An aromatic rich distillate photogrades about ten times faster than a homologue distillate with low aromatic content[71].

The PAHs reached a high degree of photodegradation in the water-soluble fraction, but the heavier PAHs have fewer tendencies to photodegradation. However, the organic fractions remained almost unaffected. The alkyl derivatives of PAHs show lower photodegradation rates than parent compounds (those without radical substituents). In general, the photodegradation rate lowers with the increasing substitution[68].

Photodegradation is in some cases as important as biodegradation, since these two processes act upon different components of oil: alkanes are biodegraded while aromatics tend to be photodegraded, and this fact leads to synergistic reactions which enhance the degradation of oil[72]. However, the *n*-alkanes with 10 or more carbon atoms and the isoprenoids such as prystane and phytane are very resistant to photodegradation[73], while in crude oil the secondary and the tertiary alkanes are easier to photogrades. *N*-alkanes were easily degraded by marine bacteria, while branched and cyclic alkanes were assumed to be less biodegradable[74]. Photoinduced toxicity of oil and PAHs has been confirmed due to the transient photodegradation intermediates which showed higher toxicity and solubility than the initial compounds, but were subsequently destroyed in the photocatalytic process [75].

Photocatalytic oxidations of dodecane and toluene have been considered in pure water and in synthetic sea water on different  $TiO_2$  powders under simulated solar radiation. Minero *et al.*[67] indicated that diluted hydrocarbons disappear in pure water in the presence of  $TiO_2$  and simulated solar radiation. The current properties of pure  $TiO_2$  have been found to be effective only under UV irradiation and hence there are still more challenges for in-situ application of  $TiO_2$  in open sea or water bodies.

# 2.4.8. Disinfections Applications of TiO2

Photocatalyst does not only kill bacteria cells, but also decompose the cell itself. The TiO<sub>2</sub> photocatalyst has been found to be more effective than any other antibacterial agent, because the photocatalytic reaction works even when there are cells covering the surface and while the bacteria are actively propagating[76]. The end toxin produced at the death of cell is also expected to be decomposed by photocatalytic action. TiO<sub>2</sub> does not deteriorate and it shows a long-term anti-bacterial effect. Generally speaking, disinfection by TiO<sub>2</sub> is three times stronger than chlorine, and 1.5 times stronger than ozone. Bacteria and viruses are decomposed on the tile surface due to the

strong oxidizing properties of TiO<sub>2</sub>. If you cover the walls, ceiling and floor with photocatalytic tiles, bacteria floating in the air in an operating room are also killed as they come in contact with the TiO<sub>2</sub> surface[21].

#### 2.4.9. Soil application of TiO<sub>2</sub>

One of the other applications of TiO<sub>2</sub> photocatalysis is the remediation of organically contaminated soil and sludge. Both pure and composite photocatalyst has been tried for the removal of organic pollutants from soil. Xie *et al.*[77] investigated the enhancement of photo degradation of organics on soil surfaces using TiO<sub>2</sub> induced by UV-light. They concluded that the photodegradation rate increased with the increase of the soil pH and photon flux. Higrashi and Jardim [78] used heterogeneous photocatalytic of TiO<sub>2</sub> to degrade pesticide (diuron) under laboratory conditions to evaluate the potential use of this technology for in situ remediation. They reported that the photocatalytic treatment combined with solar light to be very efficient in the destruction of diuron in the top 4 cm of contaminated soil, with the degradation rate markedly dependent on the irradiation intensity.

Pelizzetti *et al.*[79] mixed contaminated soils of 2-chlorophenol, 2,7-dichlorodibenzodioxin and atrazine with a photocatalyst (TiO<sub>2</sub>) in a aqueous slurry and exposed them to simulated solar radiation in the laboratory. They noticed that the organic contaminants were destroyed in relatively short time and reported that the photocatalytic processes could be effective chemical detoxification methods for contaminated soils. According to Hamerski *et al.*[80] the most active photocatalyst for soil purification was TiO<sub>2</sub> modified by calcium.

The common challenge on the above TiO<sub>2</sub> applications is to use its photoactivity under visible light irradiation. Thus, the following sections focus on the modification of VLR TiO<sub>2</sub>.

# 3. Modification of TiO<sub>2</sub> to Harvest Visible Light

Heterogeneous photocatalysis using semiconductor TiO2, is a fast growing field of basic and applied research for the degradation of organic pollutants in water and in air and photocatalytic water-splitting for hydrogen production[78,81-83]. Oxidation of organic pollutants occurs because of the electron-hole pairs of the semiconductor after being excited by UV light with energy equal to or greater than the band gap energy (Eg) of semiconductor. TiO2 is considered the noble photocatalyst because of its many intrinsic properties, but it has its own limitations which are critical to photocatalytic technology[84] such as: i) recombination of photo-generated electron/hole pairs, ii) fast backward reaction and iii) inefficient visible light utilization. The other significant drawback of TiO2 as photocatalyst is its wide bad gap because it is found effective only under UV irradiation and not under solar light that contain only about 4% of UV rays[5,7]. The use of UV as light source has many disadvantages such as requirement of external energy and also has to be conducted in a closed and protected space. Therefore in order to deal with these drawbacks, techniques such as i) addition of electron donors (hole scavengers), ii) addition of carbonate salts, iii) noble metal loading, metal ion doping, iv) anion doping, v) dye sensitization, vi) composite semiconductors and vi) metal ion-implantation have extensively been investigated[15].

Several attempts have been made to modify UV responsive TiO<sub>2</sub> to VLR TiO<sub>2</sub> and one of the methods is by doping of TiO<sub>2</sub> with other substances. Sol-gel method is one of the most popular methods used for doping TiO<sub>2</sub>. Mechanical alloying is also an effective process for refining grain size down to the nano-sized range and has the ability to alloy immiscible elements. In hydrothermal method the products prepared have well crystalline phase and have high thermal stability of the nano-sized materials. Flame spray pyrolysis is the coupling of high temperature and rapid quenching has been found to generate homogeneous doping even at a concentration level as high as 30 wt%. Ion-implementation technique is used for doping at higher concentrations of dopants up to 60% and it gives better properties of powders than sol-gel method.

Doping of  $TiO_2$  using noble metals such as gold[85,86], silver[6,87,88], platinum[5,89] transition metal ions such as Co, Cr, Cu, Fe, Mo, V and W[1,90,91], non-metals such as C, N, P and S[7,37, 83,92], carbon, coupling with organic dye-sensitizers[93-95] have been reported and also shown a promising result under visible light irradiation though the mechanism under which this process occurs still remains controversial. The followings are some of the attempts made to modify the electronic structures of  $TiO_2$  to improve its effectiveness under visible light.

#### 3.1. Noble Metal Loading (Pt, Au, Pd, Rh, Ni, Cu, Sn, Ag)

Noble metals deposited or doped on TiO<sub>2</sub> (Pt, Au, Pd, Rh, Ni, Cu, Sn and Ag) show various effects on the photocatalytic activity of TiO<sub>2</sub> by different mechanisms[88]. These noble metals act separately or simultaneously depending on the photoreaction conditions and they may (i) enhance the electron-hole separation by acting as electron traps, (ii) extend the light absorption into the visible range and enhance surface electron excitation by plasmon resonances excited by visible light and (iii) modify the surface properties of photocatalyst. While some authors have reported an increased photocatalytic activity under visible light due to certain metal ion doping, others have reported a decreased activity. But generally the photocatalytic property of metal ion doped TiO<sub>2</sub> has been reported to be quite higher than the undoped TiO<sub>2</sub> and therefore is seen to be of very high research value.

The effect of platinum (Pt) loading on TiO<sub>2</sub> was mainly studied for hydrogen generation from water but it was also observed that it increases the degradation of organic pollutants. Pt ion doped TiO<sub>2</sub> has been reported to have not only high photocatalytic activity under UV irradiation but also effective under visible light irradiation. Kim *et al.*[5] successfully demonstrated the visible light photocatalytic degradation of chlorinated organic compounds using Pt doped TiO<sub>2</sub>. It had lower band gap than that of undoped TiO<sub>2</sub> by about 0.2 eV and it showed higher photocatalytic activity than undoped TiO<sub>2</sub>. Analyses showed that Pt ions were substituted in the TiO<sub>2</sub> lattice and the visible light absorption is due to electronic transition between the band gap edge (CB or VB). The defect redox states of Pt ions substituted in the TiO<sub>2</sub> lattice. It was also found that the visible light activity of Pt-TiO<sub>2</sub>

Table 4. Characteristics of Noble Metal Loading on TiO2

Noble metal	Remarks
Pt	- Pt loading reduced the amount of Ti <sup>3+</sup> (evidence of the occurrence of electron transfer from TiO <sub>2</sub> to Pt)[97].
	- Pt showed better than Au loading in terms of hydrogen generation[36].
	- Anatase size (at 600 °C) decreased from 37 nm to 19 nm. Specific surface area increased from 45 to 63 m <sup>2</sup> /g. Photocatalytic activity
	increased by 18%. Phase transfer appeared from anatase to rutile at 700 °C and phase transformation was completed at 800 °C
Ag	[98].
Ag	- Ag/TiO <sub>2</sub> catalyst was evaluated for the photocatalytic degradation of reactive yellow-17 under UV and visible light irradiations.
	The enhanced degradation of reactive yellow-17 was obtained with Ag-deposited TiO2 due to trapping of conduction band electrons
	[99].
	- Optical characterization by UV-vis spectrophotometer showed a shift in optical absorption wavelength to visible region which may
	be due to the incorporation of gold nanoparticles ( $1 \sim 2\%$ ) into TiO <sub>2</sub> structure. The kinetic study showed that the rate of
Au	decomposition of phenol using Au/TiO <sub>2</sub> photocatalyst was improved by 2~2.3 times than undoped TiO <sub>2</sub> [100].
	- Au/TiO2 (0.45 wt%) in the oxidation of glucose resulted in high selectivity and high catalytic activities were made in the optimum
	conditions at a reaction temperature (40~60 °C) and a pH value[101].
	- Loadings of Pt and Au were better than that of Pd due to suitable electron affinity[102].
Pd	- Pd/TiO2 anatase exhibited higher acetylene conversion and ethylene selectivity than rutile TiO2 supported ones. Ag addition to
	Pd/TiO <sub>2</sub> anatase suppressed the advantageous effect of the Ti <sup>3+</sup> sites during selective acetylene hydrogenation[103].
	- Cu loading was almost comparable to Pt loading for hydrogen production.
Cu	- The optimum contents for Cu (1.5 wt%), Pd (1 wt%), and Au (2 wt%) loadings produced the H2 evolution rate of 360, 420, and
	557 µmol/h, respectively, suggesting that the H <sub>2</sub> evolution resulted showed that the photocatalytic capability of the cocatalysts was
	in the order of $Au > Pd > Cu[104]$ .
Ni	- Ni and Ag loadings among different noble metals were found to be effective for photocatalytic activity[105].
Sn	- At 500 °C a mixture of anatase and rutile phases was observed with rutile.

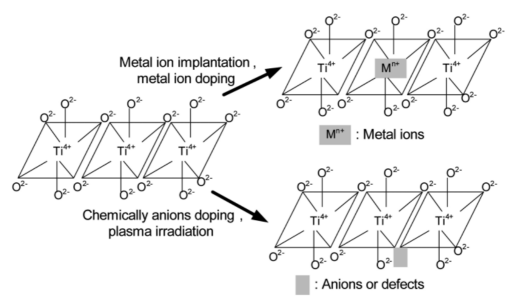


Figure 2. Electronic structure of TiO<sub>2</sub> by the substitution of metal ions or anions to harvest visible light response[10].

was observed to be strongly affected by the calcination temperature. Nano-Ag doped  $TiO_2$  was investigated[88] and observed that the Ag clusters give rise to localized energy levels in the band gap of  $TiO_2$  and the VB electrons of  $TiO_2$  are excited at wavelength longer than 370 nm. The photonic efficiency increases with an increase in the metal loading up to an optimum level while above the optimum level, the dopants behave as electron/hole recombination centres. Under UV irradiation, Ag deposits exhibit the effect only as electron traps, thus leading to the enhancement in the Ag- $TiO_2$  photocatalytic activity.

The substitution of Sn for Ti in the lattice structure is expected to aid the formation of solid solutions, alter the structure and contribute to the thermal stability of  $\text{TiO}_2[96]$ . More details of the noble metal loading are shown in Table 4.

# 3.2. Ion Doping

Cationic and anionic elements have been used as either dopants or simply as deposited with TiO<sub>2</sub> to increase its photocatalytic activity under visible light region. The Ti ions in TiO<sub>2</sub> has been substituted

with different metal ions by metal ion implantation or metal doping as shown in Figure 2 for increasing its photocatalytic properties. On the other hand, the substitution of O ions in  $TiO_2$  with N, S, C, F and P ions has also been reported to enhance its visible light response (Figure 2) but the effectiveness of doped  $TiO_2$  as photo anode or as photocatalyst critically depends on the preparation and doping methods[106].

# 3.2.1. Cationic (Metal) Doping (Fe, V, Mo, Ru, Os, Re, V, Rh, Mn, Ni, Co, Cr, La, Ce, Er, Pr, Gd, Nd, Sm, Zn)

The majority of the metal-ion-doped TiO<sub>2</sub> photocatalysts studied were prepared using the coprecipitation, incipient wet impregnation method and solgel method[107]. Transitional metal and rare earth metal ions have been tried as dopants to improve the photocatalytic efficiency of TiO<sub>2</sub> in the visible light region. As metal ions are doped into TiO<sub>2</sub>, impurity energy levels in the band gap are formed. This leads to the alteration of electron hole recombination. Transitional metals are either deposited or doped on the TiO<sub>2</sub> surfaces as metallic nanoparticles or the metals are doped as ionic dopants. A common method consists of doping of TiO<sub>2</sub> with transition metal cations while maintaining a good control of the primary particle size to achieve nanoscale configurations of the catalysts. The doping elements usually are Cr, Fe, V, Nb, Sb, Sn, P, Si, and Al[108].

Iron has been used to dope TiO2 and its photocatalytic activity was superior than the commercial Degussa P-25 under visible light irradiation[91]. Teoh et al.[109] also reported that Fe-doped TiO2 was found to have very high photocatalytic activity under visible light irradiation than Degussa P-25. Fe<sup>3+</sup> cations acted as shallow traps in the TiO<sub>2</sub> lattice. Optimum photocatalytic properties were achieved upon doping at a relatively weak level. This was closely related to the dynamics of the recombination process which was linked to the distance between dopant cations in the TiO2 lattice[110]. Fe ions trapped not only electrons but also holes, which leads to increase of photoactivity[111]. The maximum photoactivity appreared with 0.5 wt% of Fe<sup>3+</sup> due to decrease in the density of the surface active centers[14]. The added Fe atoms dissolved in TiO2 phase was found to have a rutile structure with average grain size less than 10 nm. Fe-doped powder had a higher absorption threshold in the range of 427~496 nm than the commercial P-25 powder (406 nm). Fe-doped powder color changed from white to bright yellow, Fe content more than 4.57 wt% decreased the UV-Vis absorption turning the bright vellow to dark yellow.

Vanadium doped TiO<sub>2</sub> and its visible light activities have been reported[2,112-118]. A vanadium-doped TiO<sub>2</sub> is reported to have shown quite high photocatalytic property under visible light irradiation. It is believed that under visible light irradiation the excited vanadium centres donate electron to the TiO<sub>2</sub> CB, which allows the oxidation of surface adsorbed molecules[2]. Vanadium is thus a rather interesting candidate to obtain valuable TiO<sub>2</sub>-doped photocatalysts. The photocatalytic activity of Eu3q-, La3q-, Nd3q- and Pr3q-doped TiO<sub>2</sub> was investigated[119] and observed that these doped nanoparticles showed high photocatalytic activity in comparison to undoped TiO<sub>2</sub> and their

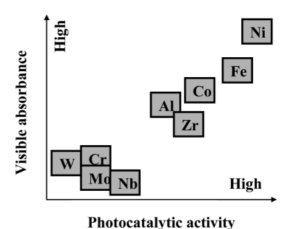


Figure 3. Effect of cationic doping compounds in terms of visible light absorbance and photocatalytic activity.

analysis indicated that this was attributed to the increased charge separation in the doped systems. Various transition-metal cation dopants have been extensively investigated. However the disadvantage of cationic dopants is that they can result in localized d-levels deep in the band gap of TiO<sub>2</sub>, which often serve as recombination centers for photogenerated charge carriers[3]. Figure 3 shows effect of different metals in TiO<sub>2</sub> to absorb visible light and to enhance photocatalytic activity[56,120,121]. More details of the cation doping TiO<sub>2</sub> are shown in Table 5.

## 3.2.2. Anionic (non-metal) Doping (C, N, S, P, F)

Anionic or non-metal dopants, namly, carbon (C), sulphur (S), nitrogen (N), phosphorus (P) and fluorine (F), are being investigated to extend the photocatalytic activity into the visible light region. It is said that the related impurity states are supposed to be close to the valence band maximum. Furthermore, the position of the conduction band minimum, which must be kept at the level of the H<sub>2</sub>/H<sub>2</sub>O potential, when TiO<sub>2</sub> is used for the photoelectrolysis of water into hydrogen and oxygen, is not affected[3].

N-doped TiO2 can be made by i) heating of titanium hydroxide and urea, ii) reactive magnetron sputtering, iii) nitriding of anatase TiO2 with alkylammonium salts and iv) treating TiO2 powder in NH3 (80%)/Ar gas flow at 550 °C[15]. Sato[128] firstly reported the visible light activity of TiO2-based products from the calcinations of Ti(OH)4 and ammonium salts and ascribed this beneficial doping effect to NO<sub>x</sub> impurities on the TiO2 lattice. Recently many groups have concluded that the visible light photoactivity was due to oxygen vacancies which gave rise to donor states located below the conduction bands, while the substitutional nitrogen acted as an inhibitor for electron-hole pair recombination. Asahi et al.[37] recently suggested that when N (2p) states and O (2p) states were mixed, the resultant band gap of nitrogen doped TiO2 was reduced below intrinsic band gap edge and the photoactivity under visible light due to substitutional nitrogen atoms. Irie et al.[128] concluded that nitrogen-induced effect allowing sub-band gap excitation was due to an isolated N (2p) states rather than band gap narrowing. Optical absorption identified the substitu-

Table 5. Characteristics of Cationic Doped TiO2

Cation	Remark
	- Fe <sup>3+</sup> cations acted as shallow traps in the TiO <sub>2</sub> lattice. Optimum photocatalytic properties were achieved upon doping at a relatively weak level. This was closely related to the dynamics of the recombination process which was linked to the distance between dopant cations in the TiO <sub>2</sub> lattice[110].
	- Fe and Cu ions trapped not only electrons but also holes, which led to increase of photoactivity[111].
Fe	- The maximum photoactivity appreared with 0.5% of Fe <sup>3+</sup> (weight) due to decrease in the density of the surface active centers [14].
	- The added Fe atoms dissolved in $TiO_2$ phase was found to have a rutile structure with average grain size less than 10 nm. Fe-doped powder had a higher absorption threshold in the range of $427 \sim 496$ nm than the commercial P-25 powder (406 nm). Fe-doped powder color changed from white to bright yellow, Fe content more than 4.57 wt% decreased the UV-Vis absorption turning the bright yellow to dark yellow.
Cr, Fe, V, Nb, Si	- Anatase-to-rutile transformation was accelerated by the mmol% content of Nb, Cr, Si, and Fe in TiO <sub>2</sub> . Interaction of coprecipitated or impregnated cations was found critical in the phase transformation process. Nb retarded the crystal growth during calcinations[108].
Al	- Aluminium doping was used on TiO <sub>2</sub> for a potential application in thermal shock due to its stable thermal expansion coefficient and physical property[122]. Al <sub>2</sub> O <sub>3</sub> and Al <sub>2</sub> TiO <sub>5</sub> were observed at AlCl <sub>3</sub> /TiCl <sub>4</sub> ratios higher than 1.1 at 1400 °C. They found that a new structure connected with Al-O-Ti framework was generated. For Al/TiO <sub>2</sub> , the anatase structure was stable after calcination at 800 °C, while pure TiO <sub>2</sub> was easily transferred to the rutile phase after calcination above 700 °C. The optical property of Al/TiO <sub>2</sub> prepared by a thermal plasma method responded visible light[123]. They also found that the size of synthesized powder decreased with increase of the amount of Al because Al species inhibited the particle growth. Al/TiO <sub>2</sub> was applied to gas sensor, which needed high conductivity of TiO <sub>2</sub> [124]. It has been reported that the conductivity of the Al-doped TiO <sub>2</sub> is higher than that of pure TiO <sub>2</sub> in a temperature (600~900 °C)
V	- V doping made in the sol-gel method provided a promising method to enhance the photoactivity of TiO <sub>2</sub> under visible light [125].
Li	<ul> <li>- TiO<sub>2</sub> doped with Li ions resulted in significant improvement in the rate of phenol degradation. On the other hand, Co, Cr, Mn, Ce, Al, and Fe doped TiO<sub>2</sub> (5 mol%) was inactive in removing phenol[14].</li> <li>- The crystalline size increased from 20 to 27 nm. The photocatalytic activity decreased.</li> </ul>
Na, K	<ul> <li>Na-doped TiO<sub>2</sub> decreased the nanoparticle size from 20 to 11 nm. The photocatalytic activity decreased.</li> <li>K-doped TiO<sub>2</sub> decreased the size from 22.4 to 7 nm. The photocatalytic activity decreased.</li> </ul>
Cu, Mn	- Cu, Mn and Fe ions trapped electrons and holes and these ions worked better than Cr, Co, and Ni ions[42].
Со	- Among the 21 metal ions doped, Fe, Mo, Ru, Os, Re, V, and Rh ions increased photocatalytic activity, whereas Co and Al ions caused detrimental effect[112].
Gd	- Gd, La, Ce, Er, Pr, and Nd doped with TiO <sub>2</sub> changed to red shift. Among them, Gd ions showed effective photoactivity and transfer of charge carriers[126].

tional-doping and localized N-states in the TiO2 lattice and the status of N to be anion-like (N-) and the chemical environment of N was in N-Ti-O in the TiO2 lattice[83]. Absorption below 500 nm was mainly due to nitrogen states located above the valence bands, whereas absorption above 500 nm was mainly caused by oxygen vacancies[128]. Valentin et al.[130] observed that nitrogen caused a significant change in the absorption spectra of the TiO2 and showed that the N 2p orbitals were localized above the top of the O 2p valence bands, even for relatively large values of doping. The N-doping effects on the electronic and optical properties of TiO2 rutile crystal have been studied using density functional theory[131]. Their calculations of several possible N-doped structures showed that band gaps had little reduction but some N 2p states lay within the band gap in the substitutional N to O structure and interstitial N-doped rutile supercell, which resulted in the reduction of the photon-transition energy and absorption of visible light. In contrast, substitutional N to Ti doped model had a significant band-gap narrowing.

Carbon-doped TiO2 thin films had hydrophilic property under visi-

ble light irradiation[132]. The localized C (2p) formed above the valence band should be the origin of visible light sensitivity, which resulted in an inferior hydrophilic property when irradiating with visible light compared with UV light. Park *et al.*[85] reported the preparation of vertically grown carbon-doped TiO<sub>2</sub> (TiO<sub>2</sub>-<sub>x</sub>C<sub>x</sub>) nanotube arrays with high aspect ratios for maximizing the photocleavage of water under white-light irradiation. The synthesized TiO<sub>2</sub>-<sub>x</sub>C<sub>x</sub> nanotube arrays showed much higher photocurrent densities and more efficient water splitting under visible-light illumination (> 420 nm) than pure TiO<sub>2</sub> nanotube arrays. The total photocurrent was 20 times higher than that with a P-25 nanoparticulate film under white-light illumination.

Wushu *et al.*[133] prepared S-doped nano-TiO<sub>2</sub> catalysts which showed photocatalytic activity under visible light. They observed that S-doped TiO<sub>2</sub> greatly increased grain distribution, a uniform dispersion of nanocrystals fabrication; In addition, S-doped TiO<sub>2</sub> in the treatment effectively inhibited the process of transformation from anatase to rutile. The replacement of some part of sulfur into the Ti<sup>4+</sup> TiO<sub>2</sub> lattice distortion and led to the lack of oxygen, improving the

Table 6. Characteristics of Anion Doped TiO2

Anion	Remark
N	- N-doped TiO2 caused a VB upward shift (narrow band gap and less oxidating holes). N-doped TiO2 for hydrogen generation was
	due to electrons for reducing protons and the ability of oxidation did not affect the performance because of more positive of VB
	level than hydrogen production[136].
	- N-doped TiO <sub>2</sub> was effective for methylene blue degradation under visible light[37].
	- C-doped TiO2 showed photoactivity on degrading 2-propanol under visible light. However, quantum efficiency (0.2%) of C-doped
С	TiO <sub>2</sub> was found inferior than that (2.25%) of N-doped TiO <sub>2</sub> for high photocatlaytic activity due to lower quantum efficiency[137].
C	- Carbon substitution (0.32%) was not effective for the shifting of upper level of the VB towards negative potential during band
	gap narrowing[138].
	- Ionic radius of S was too large to be incorporated into the lattice of TiO2 although S-doped TiO2 resulted in a similar band gap
	narrowing[37].
S	- The S ions were replaced with some of the Ti atoms in the form of S <sup>4+</sup> and S-doped TiO <sub>2</sub> showed better degradation of 2-propanol
	and methylene blue under visible light compared with undoped TiO <sub>2</sub> [134].
	- S doping restrained the transformation from anatase to rutile. The band gap shifted from 3 to 2.2 eV[137].
	- Dopants (P and C atoms) was less effective as the states were so deep that charge carriers were difficult to be transferred to
D	the surface of $TiO_2[37]$ .
P	- P-doped TiO2 in an anatase phase colored yellow exhibited a narrower band gap than pure TiO2 and an absorption tail in the
	visible range. This resulted in its effective photocatalytic degradation of 4-chlorophenol under visible-light irradiation[129].
	- The disorder layer was recovered by the surface migration of vacancy-type defects and implanted F atoms diffused to the outer
F	surface. The concentration increased with decreasing depths into the surface. The F doping gave rise to a modification of the
	electronic structure around the CB edge of TiO <sub>2</sub> [135].

Table 7. Characteristics of Metal Ion Implantation on TiO<sub>2</sub>

Metal	Remark	
Cr	- Cr deposited TiO <sub>2</sub> performed very effectively in the decomposition of NO under visible light irradiation[139].	
Aa Ma	- Metal ion-implantation with V, Cr, Mn, Fe, and Ni was possible to shift the absorption band toward visible light regions.	
Ag, Mg	However, Ag, Mg, or Ti ion-implanted TiO <sub>2</sub> showed no shift[18].	
	- The effectiveness of the red shift was in the order of $V > Cr > Mn > Fe > Ni[97]$ .	
V, Cr, Mn, Fe, Ni	- Metal ion implanted TiO2 (V, Mn, and Fe) exhibited high decomposition of propanol to CO2 under visible light irradiation	
	[60].	

visible catalyst activity[133]. Ohno *et al.*[134] synthesized S-doped  $TiO_2$  which had photo absorption greater than C-doped and N-doped  $TiO_2$  and their absorption depended on the calcination temperature. Their results indicated that S-atoms were incorporated into bulk phase of  $TiO_2$  although but it was not clear whether the  $S^{4+}$  species were introduced interstitially or at the lattice sites. However, the distortion of the local lattice was an important factor for showing the absorption in the visible region and for the shift of the onset of the absorption edge near 400 nm.

F-doped  $TiO_2$  was investigated and observed high photocatalytic property for the decomposition of acetaldehyde gas under both UV and visible light irradiation[135]. They concluded that the high photocatalytic activity was attributed to several beneficial effects produced by F-doping: enhancement of surface acidity, creation of oxygen vacancies, and increase of active sites. Table 6 shows detailed characteristics of anion-doped  $TiO_2$ .

# 3.3. Metal Ion Implantation

Several attempts have been made to deposit metal ions such as Cr, V, Mn, Fe, Ni, and Ar in the TiO<sub>2</sub> semiconductor to improve its photocatalytic activity in the visible light region[97,139-141]. High en-

ergy transitional metal ions are injected into the lattice of  $TiO_2$  using high energy bombardment to modify the electronic structure of the  $TiO_2$  to make it responsive to the visible light energy. Such modified  $TiO_2$  semiconductor are believed to be one of the most effective visible light photocatalysts and generally referred as second generation photocatalyst[15]. Detailed characteristics of metal ion implantation are shown in Table 7.

#### 3.4. Sensitization

Sensitization is classified into two methods: dye sensitization and composite semiconductor. Dye sensitization is widely applied to utilize visible light for energy conversion and textile wastewater. Table 8 shows commonly used dyes. The dyes have wavelength in the range from 442 to 665 nm. Some dyes which have redox property and visible light sensitivity are used in solar cells and photocatalytic reaction. When the dye absorbs visible light, the dye is excited. The dye in the excited state includes lower redox potential than the corresponding ground state. If the redox potential is lower than the CB of TiO<sub>2</sub>, an electron is injected from the excited state into the CB. This leads to initiation of photocatalytic reactions. Some dyes (safranine, O/EDTA and T/EDTA) absorb visible light and generate electrons as reducing

Table 8. Dye Classification[136]

Class	Dye
Thiazines	Thionine, methylene blue, new methylene blue, azure A, azure B, azure C
Hiazines	Toluidine blue
Phenazines	Phenosafranin, safranin-O, safranin-T, neutral red
Xanthenes	Fluorescein, erythrosin, erythrosin B, rhodamin B, rose Bengal, pyronine Y, eosin, rhodamine 6G
Acridines	Acridine orange, proflavine, acridine yellow
Triphenyl methane derivatives	Fusion, crystal violet, malachite green, methylviolet

Table 9. Characteristics of Dye Sensitization on TiO2

Dye	Remark
Erosin blue	- Enhancement of hydrogen production rate by different dyes was in the order of erosin blue > rose bengal > Ru(bpy) <sub>3</sub> <sup>2+</sup>
Erosiii biue	$>$ rhodamine B $\approx$ acriflavin $>$ fluorecein[143].
A	- In aqueous TiO <sub>2</sub> suspension, azo dyes were effectively degraded by photocatalytic degradation under visible light[144].
Azo, naphthol blue black	- Naphthol blue black dyes were decomposed by photosensitized oxidation[145].

agents to produce hydrogen[142]. Table 9 shows characteristics of dye sensitization.

The approach of semiconductor composition is that when a high band gap semiconductor (TiO2) is coupled with a low band gap semiconductor (CdS) with more negative CB level, CB electrons can be injected from the low band gap semiconductor to the high band gap semiconductor. Highly dispersed TiO<sub>2</sub> species prepared within zeolite frameworks as well as SiO2, B2O3 and Al2O3 matrices showed higher and unique photocatalytic performance as compared with bulk TiO2 photocatalysts[13]. Particularly, TiO<sub>2</sub> size of less than 10 nm showed significant improvement in photocatalytic reactivity attributed to the quantum size effect. Bo et al.[146] demonstrated that the photocatalytic activity of TiO2 towards the decomposition of gaseous benzene in a batch reactor was greatly improved by loading TiO2 on the surface of CeO2-ZrO2. The research investigated the effects of three metals doping into Ce<sub>0.5</sub>Zr<sub>0.5</sub>O<sub>2</sub> on photocatalytic activity of  $TiO_2/Ce_{0.45}Zr_{0.45}M_{0.1}OX$  (M = Y, La, Mn). Doping of metal in  $TiO_2$ increases more crystal defects and oxygen cavities on the surface which captures electron excited from VB to CB thereby impeding the electron-hole pair recombination. Photocatalytic process mainly occurs on the surface of the photocatalyst. Synthesis, characterization and photocatalytic activity of porous manganese oxide doped TiO2 for the decomposition of toluene reported by [82] which showed better thermal stability and regeneration activity, improved surface area than bulk TiO2 and require lower catalyst loading.

# **Conclusions**

 $TiO_2$  is one of the most promising semiconductor photocatalyst in the future for environmental applications because of its many excellent properties. Although the main limitation include its poor response to visible natural solar light nevertheless, with more research efforts worldwide, it wont be far when we could one day convert  $TiO_2$  photocatalyst into a very effective photocatalyst for use easily under natural solar light thereby solving many of the photocatalytic

limitations. Doping of  $TiO_2$  with other substances is a very promising method to achieve visible light response  $TiO_2$ . The use of  $TiCl_4$  as coagulant in the wastewater treatment serves the dual purpose of sludge management and in producing doped  $TiO_2$  which could open up greater possibilities for next generation of visible light response  $TiO_2$  photocatalyst.

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