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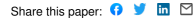
Sammy W. Verbruggen, Sammy W. Verbruggen

Institutions: Katholieke Universiteit Leuven, University of Antwerp

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TiO₂ photocatalysis for the degradation of pollutants in gas phase: from morphological design to plasmonic enhancement

Sammy W. Verbruggen ^{a,b*}

^a Sustainable Energy, Air & Water Technology (DuEL), Department of Bioscience Engineering, University of Antwerp, Groenenborgerlaan 171, 2020 Antwerp, Belgium

^b Center for Surface Chemistry and Catalysis, Department of Microbial and Molecular Systems, KU Leuven, Kasteelpark Arenberg 23, 3001 Heverlee (Leuven), Belgium

* Corresponding author: <u>Sammy.verbruggen@uantwerpen.be</u>

Tel: +32 3 265 32 60, Fax: +32 3 265 32 25

ABSTRACT

TiO₂-based photocatalysis has become a viable technology in various application fields such as (waste)water purification, photovoltaics/artificial photosynthesis, environmentally friendly organic synthesis and remediation of air pollution. Because of the increasing impact of bad air quality worldwide, this review focuses on the use and optimization of TiO₂-based photocatalysts for gas phase applications. Over the past years various specific aspects of TiO₂ photocatalysis have been reviewed individually. The intent of this review is to offer a broad tutorial on (recent) trends in TiO₂ photocatalyst modification for the intensification of photocatalyst modification is discussed both on a morphological as well as an electronic level from the perspective of gas phase applications. The main focus is laid on recent developments, but also possible opportunities to the field. This review is intended as a solid introduction for researchers new to the field, as well as a summarizing update for established investigators.

Keywords: Photocatalysis; titanium dioxide; gas phase; modification; morphology; plasmon

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1 INTRODUCTION: IMPORTANCE OF AIR CLEANING

Robert Orben, an American comedy writer once said: "There's so much pollution in the air now, that if it weren't for our lungs there'd be no place to put it all". Unfortunately, some truth can be found in his playful words. As an illustration: for 2008 the World Health Organization (WHO) published a world map indicating the number of deaths caused by poor outdoor air quality (Figure 1). In a recent press release they claimed that in 2012 one out of eight deaths worldwide can be linked to air pollution.[1] Also the European Environment Agency (EEA) has published data on the statistical life expectancy reduction due to high levels of airborne pollutants. In their 2012 air quality report it is stated that human life expectancy is reduced by more than eight months on average and by more than two years in the most polluted cities and regions.[2] Though this is a controversial statement, it clearly indicates that air pollution is a serious problem that calls for urgent action from policy makers (e.g. by developing concise guidelines and enforcing realistic exposure levels), as well as from scientists by investigating sustainable environmental solutions.

Indoor air quality is often even worse that outdoor air quality. Indoor air pollution levels are roughly the same as those outdoors *plus* additional contaminants from indoor sources. Preventing pollution by source oriented measures is obviously the most convenient way of improving air quality levels, but it is utopia to believe that this alone will solve the problem. In case these primary measures are not sufficient, secondary measures need to be considered. For instance increased ventilation of confined spaces is frequently used to reduce pollutant levels indoors, but this merely transfers the contaminants to another location.[3,4] Purifiers based on filters or adsorbents can successfully remove harmful

components from the air, but the net effect is a mere transfer from one phase to another.[3,5] Therefore direct abatement of airborne contaminants is one of the more interesting options. A number of advanced oxidation processes (AOPs) have been proposed to serve as viable air cleaning technologies. AOPs are chemical oxidation reactions that exploit the high reactivity of oxygen (more specifically hydroxyl ('OH)) radicals.[6] Air-based AOPs include ozonation, often combined with H_2O_2 , a catalyst or UV irradiation,[7,8] and photocatalytic oxidation (PCO). PCO offers the advantage that there is no need for additional chemicals such as H_2O_2 and there are basically no waste streams.

The first accounts on gas-solid heterogeneous photocatalysis can be traced back to the 1970's with the work of Teichner's and Stone's groups.[9–11] Since then the use of photocatalysis in gaseous systems has been investigated with exponential interest, especially during the last two decades (Figure 2). In these studies, titanium dioxide (TiO_2) is the most encountered photocatalyst, because it is considered inexpensive, (photo)stable, non-toxic, and easily activated by (near-)UV light.[12]

This review focuses on the use and optimization of TiO_2 -based photocatalysts for gas phase applications. At this point it is important to stress that although this review is written from the perspective of air purification, photocatalysis is also a promising technology in the fields of (waste)water purification,[13–15] photovoltaics/artificial photosynthesis[16,17] and environmentally friendly organic synthesis.[18–20] Given the vast amount of information available on the topic, it is impossible to include *all* contributions to the field. In addition, many different aspects of TiO_2 photocatalysis and catalyst modification have been reviewed individually. The intent of this review is to offer a broad overview of trends in TiO_2 photocatalyst modification, both on a morphologic as well as electronic level, for the intensification of photocatalytic applications in the gas phase. The text is illustrated with validated results from literature and some recent results of our group. Where applicable, more detailed (review) articles will be suggested to the reader for further reference.

2 TARGETING THE POLLUTANTS

Against expectation, pollutant levels indoors are often higher than outside (EPA, 2013), due to recent building techniques where better insulation is not always compensated by sufficient ventilation. This led to new symptomatic diseases such as "sick building syndrome", characterized by headaches, nausea and irritation of skin, eyes, nose and throat.[21] Fortunately, many of the airborne contaminants can be destroyed by PCO. The most important classes of outdoor and indoor gaseous (in)organic contaminants amenable to PCO by TiO₂-based photocatalysts, are nitric oxides (NO_x), sulfur oxides (SO_x), volatile organic compounds (VOCs) and dioxins, as illustrated in Table 1. With regard to this table, note that the booming field of gas phase photocatalysis has led to the development of a myriad of photoreactors and testing protocols (various flow rates, concentrations, catalyst loadings and spatial organizations, lamps and corresponding intensities,...), resulting in many different styles of reporting, making it very difficult to mutually compare results. In that regard several institutions are collaborating on the development of standardized photocatalytic test protocols.

Next to actual gaseous pollutants, PCO of organic solids can be considered as a border case of gas phase photocatalysis, as the reactions take place at the solid-gas interface, for instance after deposition of suspended solids in aerosols. Furthermore, the mass transfer laws governing the supply of O_2 or the carry-off of CO_2 are similar to those of gas phase reactions. Such reactions include PCO of polymer films such as polyethylene[56] and polystyrene[57], but also solid acids. The degradation of stearic acid films is a typical example of the latter. Stearic acid is considered as a good model compound for organic fouling on, for instance, glass windows and is therefore often the pollutant of choice for studies involving self-cleaning surfaces.[58–61] Recently, Mills and Krysa collaborated on the development of another quick qualitative test for evaluating photocatalytic performance based on color changes of solid resazurin ink films.[62] A final, socially relevant application of solid-gas heterogeneous PCO is the degradation of soot (black carbon) deposits. In this context Smits *et al.* have shown that it is possible to photocatalytically degrade considerable amounts of carbon black (ca. 22 μ g cm⁻²) using TiO₂-coated mortar substrates.[63] Within the framework of air pollution, the role of soot is becoming more and more prominent, which is why several groups' recent activities are concentrated on this emerging topic.[63–66]

3 FUNDAMENTALS OF PHOTOCATALYSIS

The previous sections underline the stringent need for air purification and illustrate that PCO could offer a viable solution. In first instance this chapter briefly reviews the fundamentals of photocatalytic reactions and points out the benefits and limitations. The second section of this chapter investigates the driving properties of gas phase photocatalysis in particular. Although this review focuses on 'classic' TiO₂-based photocatalysis, it has to be highlighted that in recent years a lot of work has been devoted to the development of alternative photocatalysts, other than TiO₂. These are often semiconductors based on d⁰ transition metal cations such as Ta⁵⁺ and Nb⁵⁺, or nitrides and oxides of Ga³⁺, In³⁺ and Bi³⁺ (d¹⁰ configuration). Metal-organic frameworks (MOFs) make up another special class of alternative photocatalysts. Photoactivity has been demonstrated for several members of this material class such as MOF-5, UiO-66 and UiO-66(NH₂).[67–69] Hernández-Alonso *et al.* have reviewed a great number of these alternative catalyst materials, so we refer the interested reader to this review paper for more information.[70] The rest of this literature overview is restricted to TiO₂-based photocatalysts.

3.1 GENERALITIES OF TIO₂ PHOTOCATALYSIS

Anatase, rutile, brookite and TiO_2 (B) monoclinic are the four TiO_2 polymorphs that are encountered in nature, of which anatase is the most photoactive phase and rutile thermodynamically the most stable.[71] Both the anatase and rutile crystal lattices are composed of chains of TiO_6 octahedra, but with different connectivity. In the anatase lattice the octahedra share four edges (Figure 3a), whereas in rutile only two edges connect the octahedra with their nearest neighbors (Figure 3b). According to Pauling's third rule, structures tend to become less stable as the constituting polyhedra are connected by an increasing number of edges or faces due to increasing repulsion of the cationic cores. This explains why rutile is thermodynamically more stable than anatase.

As any semiconductor, TiO_2 is characterized by a band structure in which the energy levels of the valence band (highest occupied level) and conduction band (lowest unoccupied level) are separated by a band gap (E_g) of ca. 3.2 eV in the case of anatase.[72] A photocatalytic event is initiated when the catalyst is activated by light with an energy content that is sufficient to overcome the band gap.[73] The incident light wavelength is related to its energy content according to (eq. 1).

$$E = h\nu = \frac{hc}{\lambda} \tag{eq. 1}$$

with E = energy content h = Planck's constant (6.63 x 10^{-34} J s) v = frequency of incident light c = speed of light (3 x 10^8 m s⁻¹) λ = incident light wavelength

In the case of TiO_2 anatase the band gap energy corresponds to UV light with a wavelength of 388 nm or shorter to bridge the band gap. This enlightens one of the biggest limitations of TiO_2 photocatalysis, as UV light only contributes to ca. 5% of the entire solar spectrum. It would therefore be of tremendous interest to extend the photocatalytic activity towards the visible light region (400-700 nm) that contributes to an approximate additional 40% of the spectrum.[74]

After absorbing light of sufficient energy, an electron (e^{-}) is promoted from the valence band to the conduction band, leaving behind a positively charged hole (h^+) ((eq. 2) and Figure 4)

$$TiO_2 + h\nu \to e_{CB}^- + h_{VB}^+ \tag{eq. 2}$$

The photogenerated charge carriers can in turn initiate the reduction/oxidation of species adsorbed on the catalyst surface, depending on the relative positions of their redox potentials. At neutral pH, the redox potential of the hole on TiO_2 is +2.53 V, whereas that of the excited electron is -0.52 V (both versus standard hydrogen electrode (SHE)).[75] Hence, reductions can only occur when the redox potential of the e⁻ is negative enough to reduce the oxidant, while oxidations only take place when the redox potential of the h⁺ is more positive than that of the reductant. As an illustration, consider the application of H₂ gas evolution from H₂O (i.e. water splitting) (Figure 4). Two half reactions need to take place: reduction of H₂O to H₂ (-0.413 V vs. SHE) on the one hand and the oxidation of H₂O to O₂ (+0.83 V vs. SHE) on the other hand. Using UV-activated TiO₂, this reaction is allowed since the photogenerated electrons and holes are strong enough reducing agents (-0.52 V < -0.413 V) and oxidizing agents (2.53 V > 0.83 V), respectively.

With regard to the degradation of organic contaminants, the generally accepted reaction mechanism involves a radical pathway. The radical chain reaction is typically initiated by the formation of hydroxyl radicals (°OH) from H₂O (+2.27 V vs. SHE) and superoxide anion radicals (O^{2-•}) from O₂ (-0.28 V vs. SHE) with the aid of photogenerated h⁺ and e⁻, respectively (equations (eq. 3) and (eq. 4)).[75,76] This is justified in view of the relative positions of the redox potentials (Figure 4). Subsequent reactions of O^{2-•} with H⁺ can yield other reactive oxygen species such as hydroperoxyl radicals (•OOH), H₂O₂ and finally •OH (eq. 5-7) that may participate in the further photocatalytic process.[71,74]

Singlet oxygen (${}^{1}O_{2}$) is another specific oxygen species that can be formed through photo-oxidation of $O_{2}^{-\bullet}$ on the TiO₂ surface, often in sensitized reactions.[77–80] Direct oxidation of adsorbed organic species by photogenerated holes has also been reported, especially at low water content or high surface coverage of pollutant.[71,75,81]

$$H_2O + h_{VB}^+ \rightarrow {}^{\bullet}OH + H^+ \tag{eq. 3}$$

$$O_2 + e_{CB}^- \to O_2^{-\bullet} \tag{eq. 4}$$

$$O_2^{-\bullet} + H^+ \to {}^{\bullet}OOH \tag{eq. 5}$$

$$^{\bullet}OOH + ^{\bullet}OOH \to H_2O_2 + O_2$$
 (eq. 6)

$$H_2 O_2 + e_{CB}^- \to {}^{\bullet} OH + OH^-$$
 (eq. 7)

Apart from initiating a radical chain reaction, the possibility also exists for photogenerated charge carriers to neutralize one another (eq. 8). This process is known as recombination.[82,83] It is facilitated by lattice defects, crystal imperfections and impurities.[84] According to Hoffmann *et al.*,[85] charge recombination is a semi-fast process occurring on a time scale of 10-100 ns, whereas the initial charge generation is very fast (fs time scale) and interfacial charge transfer is rather slow (100 ns - ms time scale). Of course, recombination should be avoided as much as possible because it results in drastic efficiency decrease. Several strategies have been -and are being- considered in order

to reduce charge recombination by improving separation of $e^{-}h^{+}$ pairs (cfr. Section 5), thus enhancing the photocatalytic efficiency.

$$e_{CB}^- + h_{VB}^+ \rightarrow energy$$
 (eq. 8)

3.2 DRIVING PHOTOCATALYTIC PROPERTIES IN GAS PHASE

After these general considerations, it is useful to try defining the driving parameters of gas phase photocatalysis and how they differ from liquid phase PCO. An insightful case-study is offered by comparing the often used P25 (Evonik) and PC500 (Crystal Global) TiO₂-based photocatalysts.[41] In liquid phase PCO, P25 outperforms PC500.[86–89] In the gas phase PCO of acetaldehyde, however, the extreme opposite is observed. [41,90] Apparently, both catalysts possess characteristics that render them particularly more (un)suited for either gas or liquid phase applications. For more insight, an overview of the relative magnitude of several important photocatalytic and material properties of PC500 with respect to P25 is given in Table 2. From this table it is clear that the superior activity of PC500 in the gas phase PCO of acetaldehyde can be attributed to its larger specific surface area, absolute amount of surface OH-groups and adsorption capacity. However, considering adsorption properties alone, the turnover frequency (TOF) on PC500 should exceed the TOF on P25 by a factor of at least three. This is plainly not the case (factor 1.3), which is attributed to the unfavorable bulk properties of PC500, as evidenced by the low photocurrent ratio (<0.01). This is indicative of a lower photon utilization efficiency of the PC500 photocatalyst. It has to be added that in aqueous phase the low performance of the PC500 catalysts is in part expected to be due to adsorbed and bulk water in the small, intricate pore system (mean size around 4-5 nm) that will imply a serious mass transfer resistance. In addition, the pH of the solution in liquid phase PCO also plays an important role in governing the attraction/repulsion between the contaminants and the catalyst surface.[85]

Summarizing, good morphological properties can dominate the photocatalytic efficiency over poor electronic properties and is thus the main driving force for gas phase PCO. It is in fact useful to exploit the low mass transfer resistance encountered when working in gas phase by developing photoactive materials with high surface area, which in most cases will be quite accessible. On the other hand, the ideal gas phase photocatalyst should not only possess a large specific surface area and adsorption capacity, but also a high photon utilization efficiency. This implies that (i) light has to be able to penetrate the material and (ii) the incident photons should be absorbed and effectively induce the generation of free charge carriers. This means that two main issues should be addressed: the limited activity window of TiO₂ (restricted to UV light) and the tendency for free charge carriers to recombine. At this point it is evident that both morphological design (to the first degree) as well as electronic

modifications (to a second extent) can add substantially to the overall photocatalytic performance in gas phase. These two main types of photocatalyst modification will be the subject of the next two chapters.

4 MORPHOLOGICAL TIO₂ DESIGN AND SELECTED GAS PHASE EXAMPLES

As already indicated above, TiO₂ photocatalysis can benefit from morphological design in several ways. Nakata and Fujishima have proposed a classification of TiO₂ photocatalysts in four categories, based on their dimensionality (Figure 5).[91] In the following discussion this classification will be followed and extended with additional examples on gas phase applications. It is not the intention to discuss into detail all different synthesis strategies. The main goal is to give an overview of available TiO₂ catalyst architectures with the associated benefits and drawbacks and illustrate them with gas phase PCO studies. For an in-depth overview on photocatalyst design with more detail on synthesis procedures, the reader is referred to the excellent review article by Nakata and Fujishima.[91] Note that some structural modifications will also affect the electronic properties and vice versa.

4.1 NANOPARTICLES

As photocatalytic reactions take place at active sites on the TiO₂ surface, it is useful to maximize exposure of the catalyst surface to the gaseous environment and the light source.[41] By reducing the primary particle size, a very high surface-to-volume ratio can be obtained. A second advantage of using nanoparticles is offered by the short charge carrier diffusion distances involved, reducing e⁻h⁺ recombination.[92] By decreasing the particle size, an overall increase of photocatalytic performance has been shown for gas phase PCO of toluene, [93] ethanol and acetaldehyde, [94] amongst others. Also industrially, fabrication of photocatalyst powders with small particle sizes is a well-known strategy. For instance, Evonik's P25 Aeroxide, which is one of the most widely used commercial photocatalysts and the benchmark for most comparative studies, is a nano-sized powder with an average primary particle size of approximately 21 nm. Maira et al. demonstrated that 7 nm is the optimal size of TiO_2 nanoparticles for PCO of gaseous trichloroethylene.[39] Larger individual particles are deemed to be less efficient because less of the incident light is absorbed due to the larger scattered fraction, but this can be circumvented by the ultimate catalyst and/or reactor design. Multiple scattering can eventually lead to improved light utilization and associated high efficiencies.[34] On the other hand the activity also drops when decreasing the nanoparticle size below 7 nm. This is due to the fact that the TiO_2 semiconductor displays discretization of its band structure for such small particles. As a consequence, the band gap experiences a blue shift, leading to less efficient utilization of incoming photons. [95,96] This phenomenon is better known as the "quantum size effect", that limits the exploitation of the net effect of increasing surface area by shrinking the particle size.

Fabrication of hollow nanospheres[97] or sphere-in-sphere structures[98] are more advanced variations on the theme of nanoparticle design. Such architectures show the additional advantage of multiple reflections of incident light inside the particles, leading to improved photon utilization.

4.2 NANOTUBES AND (HOLLOW) FIBERS

TiO₂ nanotubes (Figure 6a), nanofibers (Figure 6b), nanowires and other related one-dimensional structures possess similar advantages to zero dimensional nanoparticles.[99–101] They too benefit from high surface-to-volume ratios, as well as fast interfacial charge transfer, leading to low recombination rates, especially in the case of thin-walled hollow structures.[91] Furthermore, an assembly of fibrous or wire-like structures can serve as a mat that is well suited for practical gas phase applications such as gas sensing.[102,103] Creating additional surface area through the introduction of (multiple) hollow channels (Figure 6c) further enhances the photocatalytic activity of TiO₂ fibers. Gas phase PCO of acetaldehyde was shown to progress more efficiently, as the number of channels in hollow fibers increases. This result was attributed to a combined effect of better molecule trapping, as well as the occurrence of multiple light reflections in the cavities.[104] A similar effect was observed for long nanotube arrays that were also used in the PCO of acetaldehyde.[105]

4.3 NANOSHEETS AND THIN FILMS

TiO₂ nanosheets are commonly employed in applications involving self-cleaning surfaces, i.e. photocatalytic reactions at the solid-gas (or solid-liquid) interface. This is because of their specific properties. Nanosheets are flake-shaped structures with a thickness of a few nanometers and cross-sectional dimensions up to a few microns.[106] Generally, nanosheets adhere well onto various substrates and their surface is very smooth. That property is advantageous towards anti-fouling applications as it is more difficult for pollutants to stick to a smooth surface compared to rough surfaces. Thin, long and narrow belt-like structures can be considered as a special kind of nanosheets, called nanoribbons.[107,108]

TiO₂ thin films can be regarded to cross the bridge between two dimensional and three dimensional architectures. Very thin films (with a thickness of a few nanometers) can still be approximated as two dimensional. These films often consist of hard TiO₂ glazes prepared by various techniques, including vapor-type depositions (including atomic layer deposition, ALD),[61,109,110] sputtering,[111] sol-gel coating[112,113] and thermal methods.[114] A commercially available material that fits this category and is often used in comparative tests with self-cleaning surfaces, is Pilkington Activ[™] glass,[115]

consisting of a 15 nm thick nanocrystalline TiO_2 layer.[116] Thicker films (sub-micron to several hundreds of microns) are usually prepared with the intention of introducing a certain degree of porosity or structural ordering. These can no longer be regarded as being two dimensional and will therefore be dealt with in the next section.

4.4 POROUS, INTERCONNECTED AND SUPPORTED STRUCTURES

Although quite some benefits are associated with the use of nanoparticles, a big disadvantage is that they have high tendency to agglomerate into larger aggregates. Generally this is at the expense of photocatalytic activity, but in particular cases it can also lead to an improved efficiency. Bahnemann's group have proposed the 'antenna mechanism', which involves the transfer of photon energy from one nanoparticle to another when they are arranged as an interconnected network along a fixed crystallographic orientation.[117] This could be an important effect in many TiO₂ powders consisting of large secondary agglomerates of closely interconnected crystalline nanoparticles. In many other cases, however, TiO₂ nanoparticles are incorporated into (porous) substrates or alternatively, (ordered) interconnected 3D structures of pure TiO₂ are synthesized.

4.4.1 TiO₂ supported on porous materials

Aggregation of individual nanoparticles is often prevented by their incorporation into porous substrates such as zeolites and (ordered) mesoporous silica.[118–120] Zeolites are micro- and mesoporous materials with uniform pore sizes and very good adsorption properties, which makes them interesting supports for environmental remediation when combined with TiO₂.[121–123] Gas phase PCO over TiO₂-zeolite composites has for instance been applied for the degradation of acetaldehyde,[124] pyridine[125] and propionaldehyde.[126] Another peculiar 3D support is of biological origin. Van Eynde *et al.* have used the silica frustule of diatoms (i.e. siliceous algae skeletons) as porous substrates with a multimodal pore size distribution for the fixation of TiO₂ in photocatalytic air purification applications (Figure 7a).[127,128] Apart from the selected examples mentioned above, many other ordered and unordered (bulk) (meso)porous supports are available and have been reviewed in literature.[120,129]

Next to accommodating or preventing agglomeration of nanoparticles, some supports can play an active role in the catalytic reaction. Especially carbonaceous supports often exert such a function, which has been thoroughly reviewed by Leary and Westwood.[130] For instance, TiO_2 can be deposited onto activated carbon. It is known that activated carbon is an excellent adsorbent. The combination with TiO_2 offers the very attractive feature that pollutants can be efficiently 'caught' from the surroundings and the reaction is less affected by excessive humidity levels. Thereafter they are photocatalytically degraded while surface concentrations are kept high.[131,132] Another example of

a carbonaceous substrate participating in the PCO reaction is the use of carbon nanotubes (CNTs). CNTs are well known for their excellent electron conductive properties. When TiO₂ is brought into electrochemical contact with CNTs, photogenerated electrons can be transported through the CNT from the spot of photo-excitation to the site where the reduction reactions take place. That way they are physically separated from the holes so the rate of charge carrier recombination is reduced. TiO₂-CNT composites have for instance proven to be quite efficient in the PCO of gaseous acetone and propene for this reason.[133,134] We have also exploited this effect for the swift electron transfer in a stand-alone photo-fuel cell that produces hydrogen gas from ambient humid air.[135] Alternatively, Faria's group has shown that CNTs may also act as sensitizers (cfr. Section 5.3.1) by generating e⁻-h⁺-pairs on the tubes and subsequently transferring electrons to TiO₂.[136] Combination of TiO₂ with graphene (and fullerene) is also frequently encountered. The beneficial effect is again based on efficient conductivity of photogenerated electrons in order to reduce charge recombination, as shown for gas phase PCO of benzene.[137] Note that these are all examples of how design aspects simultaneously influence the electronic properties.

4.4.2 Porous and interconnected TiO₂ structures

It would be very convenient to have a strategy available that results in self-supporting, porous, macroscopic TiO₂ networks that can be readily incorporated in gas phase PCO reactors. A foam-like structure is a suitable candidate to this end because of its large exposed surface area and its open structure that keeps mass transfer resistances low and still allows sufficient light penetration. Keller's group made nice advancements in this domain by using foam-like supports for various gas phase applications, including disinfection of airborne bacteria.[139–141] An alternative strategy has been explored by Tytgat *et al.*, who fabricated self-supporting TiO₂ foams using a direct foaming technique (Figure 7b).[142] It was shown that these TiO₂ foams were active towards PCO of ethylene in the gas phase.

Usually a templating strategy is followed in order to fabricate structured TiO₂ networks. The sacrificial template technique involves deposition of TiO₂ onto, often ordered, foreign substrates. In the next step the foreign template is removed by chemical or thermal treatment, leaving an interconnected TiO₂ network with a structure commensurate to the original template. A high degree of porosity can be introduced due to pores and channels caused by the disappearance of the template. An illustration of this principle is given by some of our recent work in which carbon nanotubes and carbon nanosheets were used as sacrificial carbonaceous templates for atomic layer deposition (ALD) of TiO₂. A thin, conformal layer of TiO₂ was deposited on the surface of these carbonaceous structures. ALD was used as the deposition method since it ensures complete coverage of nanostructures with high aspect ratios, without obstructing the pores.[143] Calcination of the samples led to removal of the template,

crystallization into the photoactive anatase phase, and a morphology change into an interconnected network of TiO₂ nanoparticles with an overall morphology commensurate to the template (Figure 8). The obtained films thus present a very large surface area enhancement and were shown to be more active compared to commercially available reference materials in the gas phase PCO of acetaldehyde.[144–146]

A particular case of the templating method that deserves special attention is the synthesis of inverse opals as photonic structures.[147,148] Photonic structures are ordered materials with void sizes corresponding to the incident wavelength. Light that enters the crystal is trapped in those voids by multiple internal reflections and diffractions, increasing the effective optical path length through the catalyst (Figure 9).[149] This way the probability of light absorption and subsequent charge generation is enhanced. That is the reason why inverse opals are preferably designed to match light wavelengths corresponding to the onset of the semiconductor band gap, that normally have low probability of leading to e^-h^+ -pair formation.[150] The study of Ren and Valsaraj on the use of TiO₂ inverse opals for the enhanced PCO of 1,2-dichlorobenzene is one of the few accounts in literature in which photonic structures are employed to improve gas phase photocatalysis.[151] This application again illustrates how morphologic engineering and electronic properties are intertwined.

5 MODIFYING TIO₂ on the electronic level: promising strategies and selected gas phase examples

A lot of effort has been devoted to improve the photo-electronic properties of TiO_2 in order to circumvent one of the two major limitations of TiO_2 photocatalysis:

- (i) Tendency for photogenerated charge carriers to recombine
- (ii) Photoactivity window restricted to UV illumination

In this section an overview is given on current strategies to bypass one or both of these shortcomings. The focus is laid on the most recent or promising developments.

5.1 DOPING

Semiconductor doping is the most encountered and facile way of changing the electronic properties of TiO_2 by altering the band gap. There are three main doping categories: self-doping, cation doping and anion doping. Co-doping with several elements simultaneously is a variation on the same theme.

5.1.1 Self-doping

The term "self-doping" refers to the fact that the TiO_2 semiconductor is changed without the introduction of other metals or non-metals. It is achieved by disrupting the stoichiometric oxygen

balance by generating vacancies in the lattice, giving rise to Ti³⁺ sites.[153] Oxygen vacancies are indeed known to be good e⁻ traps. These trapping states are located just below the conduction band. After visible light absorption they are promoted to the surface where they can participate in surface reactions, which has for instance been employed in the gas phase PCO of NO.[154,155] Mul's group reported the one-step synthesis of blue colored, Ti³⁺-rich TiO₂ that is active under visible light illumination,[156] but such materials can suffer from stability issues. On the other hand, Pillai's group recently described the visible light activity of oxygen rich TiO₂. This was attributed to an upward shift of the valence band after thermal decomposition of peroxo-TiO₂ complexes.[157] The material remained stable up to 900°C due to an increased Ti-O-Ti bond strength.

5.1.2 Cation doping

Increasing visible light absorption through cation doping (with Fe, Cu, Mn, Mg, Al, Zn, Cr, Ag, Ni, Pb, Ca, etc.) has been the focus of many research efforts for more than two decades. Unfortunately, this strategy did not lead to big breakthroughs. Nonetheless there is a vast amount of literature available on the subject. It is impossible to strive for completeness within the limited frame of this literature overview. The interested reader is therefore referred to an extensive review by Litter on the effects of transition metal doping in photocatalysis and references therein.[158] The following discussion will thus be limited to general considerations and recent trends.

In general, the goal of cation doping is to introduce additional states in the TiO_2 band gap.[92] These impurity levels result in more absorption in the visible light region of the spectrum because energy transitions are enabled that require less energy (Figure 10). The resulting energy levels of the dopant states and resulting photocatalytic activity strongly depend on several factors: the chemical nature of the dopant, its concentration and its chemical and structural environment. It is obvious that every metal will exhibit a different effect on the activity and even for a given metal cation, the resulting effect may differ depending on the synthesis or reaction conditions.[159] With regard to dopant concentration, there is a consensus that at low amounts, the dopants act as trapping sites that can improve charge separation and introduce intermediate energy levels in the band gap that promote electron excitation at longer (visible light) wavelengths. At higher concentrations they can become recombination sites.[160] Hence, there often exists an optimum amount of dopant, which is typically in the order of 1 wt% or lower (TiO₂ basis). The influence of the structural environment of the dopant is illustrated by Sun et al., who did not observe visible light activity when the cations were present as surface complexes.[161] The requirement for ion incorporation in the bulk lattice is also confirmed by Kang et al.[162] Metal ions that are not well incorporated in the lattice may block active sites at the surface and lead to lower activities.

One recent trend that emerged the last five to ten years is doping with metals from the lanthanide group of the periodic table. Especially doping with Ce is a promising approach. It has a high thermal stability, as well as adsorption selectivity that may improve adsorption of pollutants at the catalyst surface, as has been shown in the case of nitrite adsorption on 0.7wt%Ce-TiO₂.[163] Another special feature of Ce is that it acts as a good structural promoter. Ce doping leads to well-defined mesoporosity and an increased thermal stability of the anatase phase,[164] which is beneficial for gas phase applications that benefit strongly from increased surface area. Several researchers have observed a retardation in the crystal growth and a suppression of the thermal anatase to rutile recrystallization.[164,165] This structural stabilizing effect is attributed to the formation of Ti-O-Ce species at the surface, as a consequence of the large atomic radius that prevents Ce from being incorporated in the TiO₂ lattice.[159,163] A final aspect of Ce doping is the increased visible light absorption. The Ce³⁺/Ce⁴⁺ redox potential (1.8 eV) lies within the TiO₂ band gap. Consequently, a red shift is observed in the range of 400-500 nm after doping with Ce.[166,167]

5.1.3 Anion Doping

Increasing visible light absorption by doping with anions has been considered since the end of the 1980's. Carbon, fluorine, sulfur, boron, phosphor and halides are all known examples, but doping with nitrogen has attracted most attention since the first discovery of Sato *et al.* in 1986.[168] It is easily incorporated in the TiO₂ lattice due to its low ionization potential and comparable atomic radius to oxygen.[169] On the other hand, the precise chemical nature and location of the species that account for the enhanced visible light absorption of N-doped TiO₂ is still under debate: NO_x, NH_x, N²⁻, NO⁻, NO₂⁻ and NO₃⁻ have all been proposed.[170] There is a general hypothesis that incorporation of anions with lower electronegativity than oxygen in the TiO₂ lattice, results in an upward shift of the 2p-levels of the valence band and thus narrows the band gap, in contrast to cation doping that results in additional energy states within the band gap itself (Figure 10). At this point it is not 100% clear whether the N-related states are localized or intrinsically part of the band gap.[84]

An interesting application of anion doping for air purification is demonstrated by Li *et al.*[171] They studied N-F-co-doped TiO₂ for the degradation of gaseous acetaldehyde, toluene and trichloroethylene in a simulated a commercial blue LED air cleaner. Co-doping of nitrogen and fluorine showed a synergetic effect. N-doping led to an increased visible light absorption, while F-doping led to the formation of new active sites through the introduction of oxygen vacancies that give rise to [•]OH and $O_2^{-•}$ radicals.

As a final remark, it is important to realize that although the main idea of ion doping is to increase visible light absorption, high visible light *absorption* does not necessarily mean high visible light photocatalytic *activity*!

5.2 COMPOSITE SEMICONDUCTORS

By bringing two independent semiconductors into close electrochemical contact, photogenerated charge carriers can be transferred from one component to the other. The result is an improved charge separation, less recombination, with expected higher photocatalytic activity. SnO_2/TiO_2 and WO_3/TiO_2 are two representative examples of such composites.[172–175] In both cases the TiO_2 conduction band position is more negative and the TiO_2 valence band position is less positive than the conduction and valence bands of the other oxide respectively. This enables a vectorial charge transfer after photo-excitation that results in a physical separation of electrons and holes (Figure 11).

Interestingly, this type of charge separation can also occur in polymorphic TiO₂ (i.e. co-presence of anatase and rutile). It has been suggested that the good photocatalytic performance of commercial P25 (Evonik) is not only attributable to its nanocrystalline nature, but also to its polymorphic composition (roughly 80% anatase, 20% rutile). After the initial work of Bickley *et al.*[177] and several other researchers trying to unravel the synergy of anatase and rutile, [178–181] Ohtani *et al.* concluded in 2010 that the different fractions of P25 behave independently, without any interaction.[182] However, one year later Su *et al.* contradicted this finding in a convincing study that demonstrates the synergetic effect for anatase-rutile composites with an anatase content of at least 40% and rutile fraction of at least 20%.[176] Their study was unique in that sense that different composite samples were kept constant. Since morphological parameters are ruled out of the equation, it can be expected that the synergy is an intrinsic catalytic property and thus applies to both liquid and gas phase reactions. A maximal synergetic effect was observed in their experiments for 60% anatase and 40% rutile, which was 50% higher compared to pure anatase (Figure 12). This result is confirmed by surface photovoltage measurements of TiO₂ nanopowders with variable rutile content by our group.[90]

5.3 SENSITIZED AND PLASMONIC TIO2

This final part on modification of the electronic properties of TiO_2 deals with the use of nanostructures in contact with the TiO_2 surface in order to induce or enhance visible light activity. Two main categories are considered: sensitizers and nanoparticles displaying surface plasmon resonance.

5.3.1 Sensitizers

The general principle of a photosensitizer in electrochemical contact with a substrate (e.g. TiO_2) is that it absorbs visible light energy and transforms it into chemical energy that in turn can be transferred to the otherwise inactive oxide (Figure 13a).[183] Photosensitized photochemistry has mainly been employed in application fields such as dye-sensitized solar cells (DSSC) or Grätzel cells,[184] semiconductor-sensitized solar cells (SSSC),[185] water splitting[186,187] and water purification.[188,189] In order to restore the sensitizer to its ground state, an electron donor (e.g. EDTA) is added to the solution.[190] This technical requirement is the main reason why photosensitization is not a common approach in gas phase photocatalysis, where sensitizer depletion is observed.[191]

Photosensitizers can be classified into three major categories: inorganic sensitizers (quantum dots), organic dyes, and organo-metal complexes. Inorganic sensitizers are in fact narrow band gap semiconductors such as CdS, CdSe, PbS and ZnS.[193–195] They have a more negative conduction band compared to TiO₂ thus enabling the transfer of visible photogenerated electrons from the narrow band gap semiconductor to TiO₂. A unique property of quantum dots is that, due to their nanometer size, they show quantum confinement effects that allow to tune their absorption spectrum by varying the particle size.[196] They also show high extinction coefficients, so a thin layer of quantum dots (Figure 13b) already leads to appreciable visible light absorption. Plenty of organic dyes have been employed as sensitizers. Examples include eosin Y,[191] rose bengal,[188] and cresyl violet.[197] The large drawback of dyes is their lack of stability. Dissolution as well as photocatalytic degradation of the dyes has been reported.[190] Finally, organo-metal complexes can also be applied as sensitizers. They often possess chromophoric centers that mimic nature's light harvesting systems. Known examples include Ru complexes,[198] Os porphyrines[199] and Pt complexes[200] anchored to the TiO₂ surface.

5.3.2 Plasmonic photocatalysis

Surface plasmon resonance (SPR) is a unique optical property of metallic nanoparticles that arises from a photo-induced collective oscillation of conduction band electrons.[201] SPR enables concentrating and manipulating light at the nanoscale. This led to a myriad of applications ranging from sensors,[202] over energy production,[203] biochemical applications,[204,205] surface enhanced Raman spectroscopy (SERS),[206,207] to environmental remediation through photocatalysis.[208,209] The interested reader is also encouraged to read the recent review paper by Zhang *et al.* that includes more photo-physical aspects of plasmonic photocatalysis, also on materials other than TiO₂.[210]

Although the use of SPR in nanotechnology seems a recent development, exploitation of the unique optical properties of noble metal nanoparticles dates back to thousands of years ago.[211,212] For instance, red glassware from the late Bronze era (1200-1000 BC) was colored by means of copper nanoparticles. The *Lycurgus* Cup from the 4th century (AD) has the peculiar property that when the glass of the cup is illuminated, its color changes from yellowish-green to bright ruby-red, which is attributed to the presence of gold-silver alloy nanoparticles (Figure 14). A lot of the still existing stained glass frames in medieval churches receive their color from the presence of metallic nanoparticles displaying SPR. It is amazing that only in 1908, thanks to the work of Gustav Mie, people actually started to realize that they were practicing nanotechnology all along![213]

An important physical feature of plasmonic nanoparticles is their extinction cross section, equal to the sum of the absorption and scattering cross sections.[214] For particles much smaller than the incident light wavelength, the contribution of scattering is negligible and extinction is mainly determined by absorption.[214,215] Plasmonic nanoparticles can have absorption cross sections several times larger than their actual size.[216,217] Tuning of the SPR maximum to a specific wavelength is an important tool for achieving efficient utilization of incident light in various applications. Therefore the different factors that exert an influence on the SPR wavelength need to be taken into consideration. The first, quite obvious parameter that influences SPR, is the kind of metal that is used. The most encountered metals are those with pronounced SPR in the visible range of the spectrum, such as gold, silver and copper (Figure 15a). An alternative way of SPR wavelength tuning is the fabrication of metal alloy nanoparticles. This strategy enables to cover a wide range of SPR wavelengths by merely altering the alloy composition (Figure 16).[218] A second parameter that governs the SPR wavelength is particle size. Although a small red-shift of the SPR maximum occurs with increasing particle size, the dominant effects are broadening and increasing the intensity of the absorption band (Figure 15b).[217] A third influencing parameter is the shape of the particles. Not only spherical particles, but also a lot of other nanosized architectures such as rods, [219,220] cubes, [221] wires, [222] and many other (organized) geometries display SPR effects. Since the restoring force for the electron oscillation in plasmonic structures depends on the charge accumulation on the surface, it will be affected by the geometry. This is quite clear in the case of plasmonic nanorods. The charge accumulation, proportional to the restoring force, is small along the rod axis and bigger in the perpendicular direction. Therefore SPR occurs at larger wavelengths (lower energy) for oscillations along the rod axis (longitudinal plasmons) and at shorter wavelengths for oscillations along the radial direction (transversal plasmons). The larger the aspect ratio of the rod, the larger the red-shift of longitudinal plasmons. This is a general trend: the larger the deviation from sphericity, the more SPR shifts towards longer wavelengths.[220] A final main parameter affecting SPR is the dielectric environment. The charge accumulation at the particle surface induces an electric field in the near surrounding environment. This induces polarization of the medium which partially balances the charge accumulation at the surface, reducing the restoring force and consequently shifting SPR towards longer wavelengths.[216] The higher the dielectric constant of the medium (ε_m), the more pronounced this effect will be (Figure 15c).

The main concern within the context of this review is how these plasmonic structures can be used to improve photocatalytic performance under visible light illumination. In recent years, quite some studies have been published on the subject, mainly with regard to photocatalytic water splitting.[209,224,225] Despite all efforts, the exact mechanism of plasmon mediated photocatalysis is still under debate. A number of possible mechanisms have been proposed and tentatively proven, but

there seems to be no overall consensus, except for the fact that the highest activities on plasmonic photocatalysts are obtained when the incident light wavelength matches the SPR wavelength. Linic *et al.* describe the three prevailing explanations, that can be regarded as non-mutually exclusive mechanisms: (i) direct charge injection, (ii) near-field enhancement and (iii) extended optical path length by plasmon induced scattering (Figure 17).[226]

The direct charge injection mechanism is quite similar to the mechanism of photosensitization. Light absorption at the SPR wavelength leads to the formation of excited, energetic electrons. Under the right conditions and with intimate electrochemical contact between the plasmonic particle and the semiconductor, an electron is transferred from the excited plasmon state to the TiO₂ conduction band (Figure 17a). For instance, the studies by Silva *et al.*,[225] Tsukamoto *et al.*,[227] Tian and Tatsuma,[228,229] and Kowalska *et al.*[208] are all based on plasmon-induced charge transfer as the working hypothesis in relation to their observations.

The enhanced near-field hypothesis assumes radiative energy transfer between metal and semiconductor. The electrical field associated with charge accumulation on the metal surface can be quite intense. The field intensity drops exponentially with increasing distance from the nanoparticle, but it can still be significant at a distance of several nanometers (Figure 17b). Since the amount of e⁻ h^+ formation in a photocatalyst is proportional to the intensity of the surrounding electromagnetic field, it is to be expected that more photogenerated charge carriers are formed in the close vicinity of plasmonic nanoparticles. Because the field intensity strongly decreases with distance, more charge carriers are generated close to the semiconductor surface, rather than in the bulk.[226] Consequently short charge carrier diffusion distances apply that minimize charge recombination, and potentially lead to higher photocatalytic efficiency. Awazu and coworkers provided strong evidence for this near-field mechanism.[230] They covered plasmonic silver nanoparticles with layers of SiO₂ of increasing thickness to prevent direct contact with TiO_2 . The photocatalytic activity decreased with increasing thickness of the SiO₂ barrier, which is in accordance with the proposed mechanism. Liu *et al.*, [224] Kumar et al. [231], Amrollahi et al. [232] and Christopher et al. [233] use this enhanced near-field mechanism to explain their findings. As a last point of attention, one should be aware that field enhancement can only be successful if the photocatalytic semiconductor already shows at least modest activity at the incident light wavelength used to generate SPR.[231,233] Indeed, the field *intensity* is increased by SPR, but not its *energy* level to overcome the band gap.

The final mechanistic hypothesis is mainly important for large plasmonic nanoparticles, i.e. with diameters of the same order of magnitude as the incident light wavelength. In contrast to small nanoparticles, light extinction for large nanoparticles is mainly due to scattering.[234] This property is

exploited in order to increase the effective optical path length of incident light inside the metalsemiconductor system, as illustrated in Figure 17c.

Apart from the three prevailing mechanisms discussed above, local heating of plasmonic nanoparticles and direct photocatalysis on plasmonic nanostructures have also been proposed.[235,236]

Although the current discussion on plasmonic photocatalysis is extensive, the number of actual applications with regard to TiO_2 photocatalysis in the gas phase are scarce to non-existent. With this we hope that this fascinating research domain will also prove its potential in photocatalytic environmental remediation. A first step in this direction is provided by our group by studying the degradation of stearic acid at the solid-gas interface. Gold-silver plasmonic alloy nanoparticles were used to tune the photocatalytic activity of TiO_2 to the solar spectrum.[223] Very recently it has been confirmed by means of action spectra and photocurrent measurements that such gold-silver alloy- TiO_2 photocatalysts can be used for solar-driven applications.[237]

As a final remark it is important to distinguish between visible light and UV illumination of metal-TiO₂ nanostructures. Under visible light illumination, the goal is to exploit the plasmon response on the metal in order to induce visible light photocatalytic activity of TiO₂. Under UV illumination on the other hand, TiO₂ is intrinsically able to generate charge carriers and the metals do not necessarily display SPR at the incoming wavelength. Still, in these conditions they can serve as co-catalysts that offer active sites with lowered activation barriers. Furthermore they can act as electron sinks, ensuring efficient e^- -h⁺ separation.[238–240]

6 TECHNO-ECONOMIC CONSIDERATIONS CONCERNING PHOTOCATALYTIC GAS PHASE APPLICATIONS

The success of a photocatalytic reaction is not solely determined by the catalyst properties, but also by reactor design and operating conditions. A key aspect to take into account in photoreactor development is the immobilization of the catalyst. Especially in gas phase applications involving high air flow rates, it is crucial to contain nanosized particles so they do not become airborne and pose a risk to human health. Process parameters such as light intensity, pollutant concentration, amount of catalyst present, flow rate and temperature obviously all affect the PCO efficiency.[241,242] Another very important parameter is gas humidity. It is commonly observed that PCO is impeded in the absence of water vapor.[42,243] On the other hand, an excess of water vapor can also inhibit PCO because water molecules competitively adsorb on the active sites.[244] Another important process condition is the composition of the gas feed. Sometimes the presence of certain VOCs can cause cross-sensitivity of PCO for other VOCs.[245] Conversely, when partial degradation of certain compounds yields radicals with a high oxidation potential (e.g. •Cl), the rate of PCO might be enhanced for other pollutants present.[243,246]. From all this it is clear that careful control over all different process parameters is crucial in photocatalytic research and applications, but this is outside the scope of this overview.

From a socio-economic perspective, photocatalytic air purification still offers great opportunities. Already 15 years ago, Fisk estimated that the productivity gains that may be obtained by improving indoor air quality in the United States amount to 6-14 billion USD by reducing respiratory diseases, 1-4 billion USD by reducing allergies and asthma, 10-30 billion USD from decreased Sick Building Syndrome (SBS) symptoms and 20-160 billion USD dollars by improving general worker performance.[247] In that regard it is surprising that a BCC Research market survey from 2010 reported that only 0.5% of the total photocatalyst product sector is directly related to environmental applications. The largest share (87.4%) is accounted for by the construction sector (photocatalysts in building materials).[248] It has to be noted that the latter can also indirectly exert a beneficial influence on the environment, mainly through passive systems (e.g. coated surfaces exposed to the surrounding environment). Given these appealing prospects and the ample room for growth in the photocatalyst sector for environmental applications, it is wise not only to focus on improving the performance of photocatalysts through all kinds of high-end modification methods, but also keep taking into account the resulting costs, especially with the eye on commercialization.[249]

7 GENERAL CONCLUSIONS AND OUTLOOK

Photocatalysis has proven to be a viable technology for environmental air remediation. Its use has been demonstrated towards the abatement of various simple and complex, common and less encountered pollutants from indoor as well as outdoor sources. One particular compound that has not received much attention so far, is (airborne) particulate matter (consisting of soot and other poly aromatic organic and inorganic components). Employing photocatalysis in order to reduce soot concentrations in air could be a promising new area of application.

With regard to catalyst modification a lot has been achieved over the last years. The synthesis and use of nanoparticles has been studied quite extensively. Nanotubes, interconnected structures and (ordered) porous materials remain attractive building blocks for new and improved photocatalysts. The number of possible architectures that can be created is endless. In combination with properly conceived reactor design, this could mean a big step forward in photocatalytic efficiency. Photoreactor design could benefit a lot from accurate models, for instance based on computational fluid dynamic calculations, that take into account the entire photocatalytic process (flow pattern, mass transfer,...) and not only overall reaction kinetics.[250]

Tuning of electronic properties of TiO₂ photocatalysts has evolved a lot over the past few decades. Improving charge separation through formation of composite structures is a very efficient approach. Inducing visible light activity by cation doping seems to become less popular and many alternative approaches have been proposed. Particularly (dye) sensitized and plasmon-mediated visible light active photocatalysis have recently become hot research topics. Applications of these materials in the field of air purification are, however, extremely scarce, which opens up a whole new world of opportunities.

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Figure Captions

Figure 1. Deaths (per 100,000) attributable to outdoor air pollution (2008). ©World Health Organization ref. [1]

Figure 2. Number of publications on gas phase photocatalysis as a function of the year of publication according to an ISI Web of Knowledge search with the keyword combinations photocat* AND air (black bars), and photocat* AND gas* (grey bars).

Figure 3. Crystal structure of a) TiO₂ anatase and b) TiO₂ rutile. (color version online)

Figure 4. Schematic representation of the TiO_2 band structure with indication of the redox potentials of photogenerated holes and electrons (vs. SHE) at neutral pH. As an example, the required redox potentials are indicated for the water splitting reaction (grey) or the formation of reactive oxygen radicals for photocatalytic degradation reactions (dark orange). Based on the values reported in ref.[75]. (color version online)

Figure 5. Morphological classification of photocatalyst materials based on their dimensionality. Adapted from ref.[91] ©2012 with permission from Elsevier. (color version online)

Figure 6. Electron microscopy pictures of a) aligned TiO_2 nanotube arrays, b) TiO_2 nanofibers and c) hollow TiO_2 nanofibers with no, one, two and three channels.

a) reprinted with permission from ref.[99] ©2006 American Chemical Society, b) reprinted with permission from ref.[100] ©2012 Elsevier and c) reprinted with permission from ref. [104]©2010 The Royal Society of Chemistry.

Figure 7. a) SiO₂ diatom frustule support and b) macroscopic self-supporting TiO₂ foams. a) reproduced from ref.[127] ©2012 with permission of The Royal Society of Chemistry. b) reproduced from ref.[138] with permission from the author. (color version online)

Figure 8. Picture of 10 μ m CNT template with increasing number of TiO₂ deposition cycles by ALD after annealing (a) and increasing number of TiO₂ deposition cycles by ALD on CNS template after annealing (b), cross section SEM image of TiO₂ PC500 reference film (Crystal Global) (c), SEM image of 200 TiO₂ ALD cycles on CNS template as deposited (d), after calcination (e) and TEM details of the latter (f). Cross section SEM image of 100 TiO₂ ALD cycles on CNT template as deposited (g), after calcination (h) and TEM details of the latter (i). Reprinted with permission from ref.[145] ©2014 Elsevier. (color version online)

Figure 9. SEM image of an ordered TiO₂ inverse opal film with a schematic illustration of the increase in effective light path length in the photonic structure. Adapted with permission from ref.[152] ©2007 Elsevier. (color version online)

Figure 10. Schematic illustration of the effect of doping with cations (dopant energy level arbitrarily chosen) and anions on the band gap. (color version online)

Figure 11. Schematic representation of the vectorial charge transfer in a TiO₂/SnO₂ composite semiconductor

Figure 12. Demonstration of the synergetic effect of co-presence of anatase and rutile. Adapted with permission from ref.[176] ©2011 American Chemical Society. (color version online)

Figure 13. a) Schematic representation of electron flow and mechanism of photosensitization. b) TiO₂ nanorod decorated with CdS quantum dots. Panel b reprinted with permission from ref.[192] ©2010 American Chemical Society. (color version online)

Figure 14. Lycurgus Cup dating from the 4th century (AD) as it is (left) and under illumination (right). Picture: British Museum Images. (color version online)

Figure 15. Simulated extinction spectra of a) 10 nm silver and gold nanoparticles in air, b) silver nanoparticles in air of different sizes, and c) 10 nm Ag nanoparticles in air and water. simulated using MiePlot v4300. (color version online)

Figure 16. Photographs of $Au_xAg_{(1-x)}$ with x from left to right 0.2, 0.4, 0.6, 0.8 and 1.0 of a) colloidal alloy nanoparticle solutions and b) plasmonic photocatalyst powders obtained after photodeposition of the colloidal nanoparticles on TiO₂. Corresponding UV-VIS absorption spectra of c) colloidal $Au_xAg_{(1-x)}$ nanoparticle solutions and d) plasmonic $Au_xAg_{(1-x)}$ -TiO₂ photocatalysts (the gray curve corresponds to crude TiO₂). STEM-EDX maps of e) 100% gold nanoparticles on TiO₂ (Au: red, Ti: green) and f) and $Au_{0.6}Ag_{0.4}$ alloy nanoparticle on TiO₂. Reprinted with permission from ref. [223] ©2014 Elsevier. (color version online)

Figure 17. Potential mechanisms of plasmon-mediated photocatalysis. a) Direct charge injection of an electron from a plasmon excited state on the metal into the semiconductor's conduction band. b) Near-field enhancement close to the

plasmonic metal surface. c) Extension of the effective optical path length by means of multiple scattering. Panel b reprinted with permission from ref.[224] ©2011 American Chemical Society. (color version online)

Figure 1 (double column)

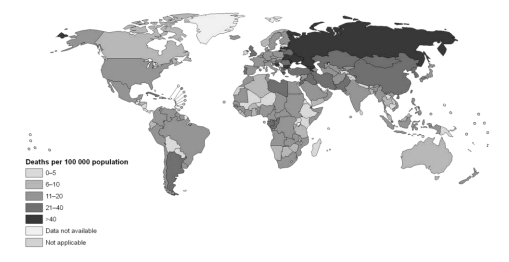


Figure 2 (single column)

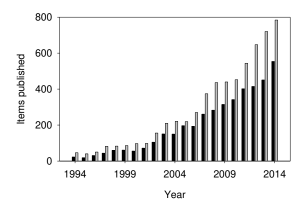


Figure 3 (single column, color online)

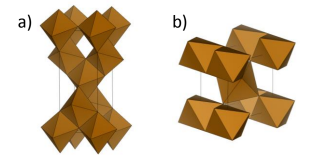


Figure 4 (single column, color online)

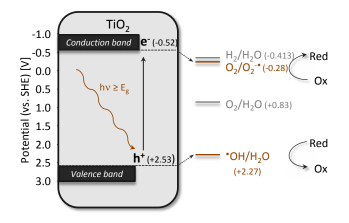


Figure 5 (single column, color online)

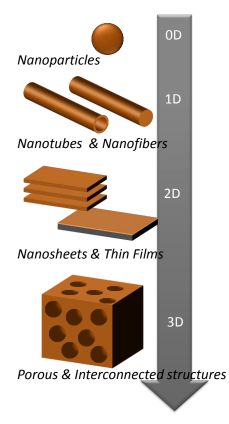


Figure 6 (single column)

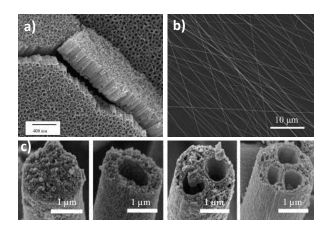


Figure 7 (single column, color online)

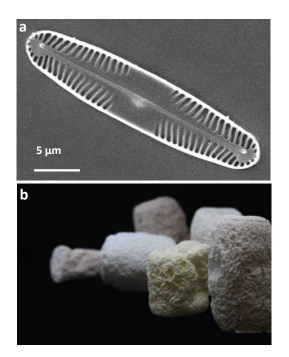


Figure 8 (double column, color online)

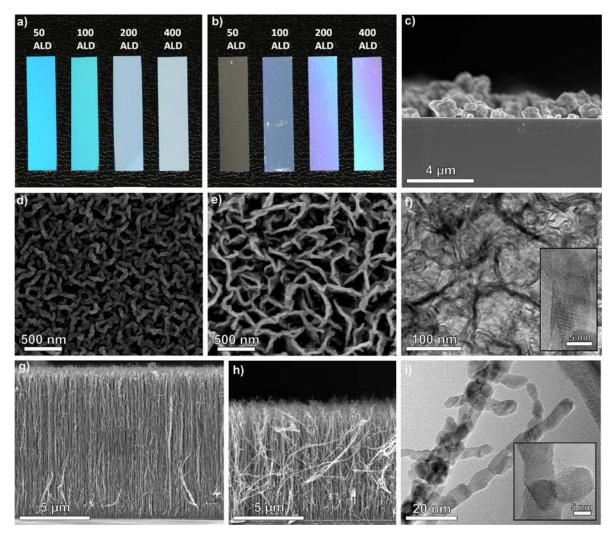


Figure 9 (single column, color online)

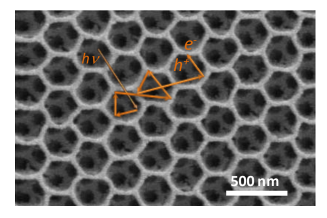
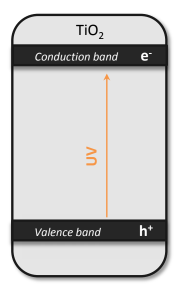
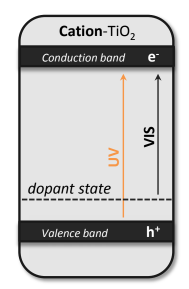


Figure 10 (double column, color online)





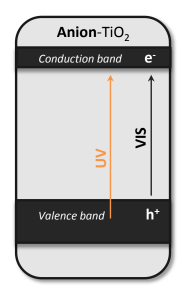
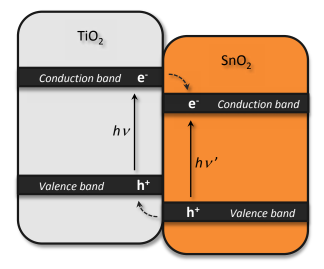
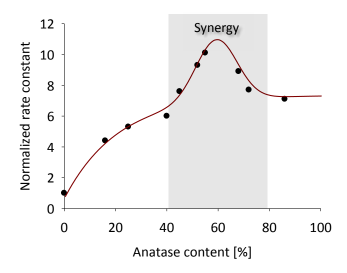


Figure 11 (single column, color online)







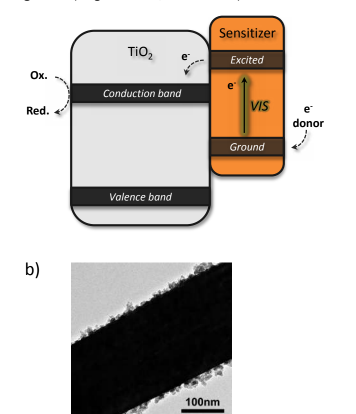
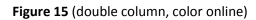


Figure 13 (single column, color online)

Figure 14 (single column, color online)





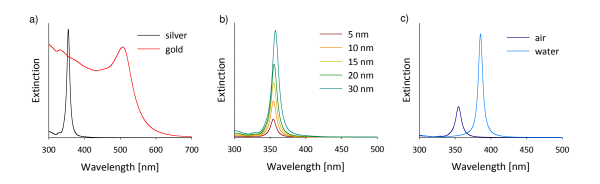
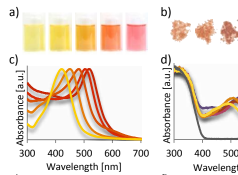
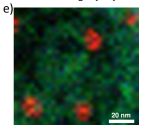
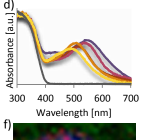
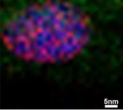


Figure 16 (single column, color online)









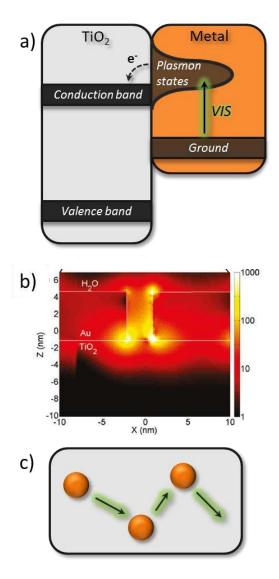


Figure 17 (single column, color online)

Pollutant class	Compound	Catalyst ^a	Concentration ^b	Absolute intensity ^b	(Best) achievement ^c	Reference
NOx	NO	P25	5 ppm	0.2 - 0.8 mW/cm ²	70% conversion	Devahasdin [22]
		TiO ₂ -NH ₃	200 ppb	750 μW/cm²	76.6% conversion	Li [23]
		Aerolyst 7710	5000 ppm	not specified	Mechanistic elucidation	Hauchecorne [24]
	NO/NO ₂	TiO ₂ , not specified	10-100 ppm	not specified	Surface reaction by XPS	Dalton [25]
		23% TiO ₂ /Al ₂ O ₃	48 ppm	Only distribution given	320 µmol/g catalyst	Shelimov [26]
SO _x	SO ₂	TiO ₂ on asphalt	175 ppb	2 mW/cm ²	23% conversion	Hassan [27]
		Synthesized TiO ₂	2529 mg/m³	not specified	98% conversion	Zhao [28]
VOCs						
Alkanes	Isooctane	P25	400 ppmv	7.56 x 10 ⁻⁹ E/cm ² /s	98.9% conversion	Alberici [29]
	Cyclohexane	P25	250 ppm	not specified	66% → 57% conversion	Einaga [30]
Halo-alkanes	Dichloromethane	$0.005 \text{ mol}\% \text{ Fe-TiO}_2$	96 ppm	0.83 mW/cm ²	87% conversion	Hung [31]
	Chloroform	Synthesized TiO ₂	162.6 ppmv	1.7 x 10 ⁻⁷ E/s	90%, fast deactivation	Yamazaki [32]
Alkenes	Ethylene	Aerolyst 7710	100 ppmv	640 μW/cm² @ 254 nm	100% conversion	Tytgat [33]
		Aerolyst on beads	75 ppmv	1.3 mW/cm² @ 254 nm	97% conversion	Verbruggen [34]
	Propene	P25	100 ppmv	not specified	100% conversion	Bouazza [35]
	1-butene	Synthesized TiO ₂	2.75 ppm	0.70 mW/cm ²	0.4 μmol/cm²/h	Cao [36]
	1,3-butadiene	P25	~ 40 ppmv	0.009 W/cm ²	0.65 μmol/cm²/h	Obee [37]
Halo-alkenes	1,2-cis-dichloroethylene	TiO ₂ fiberglass cloth	1000 ppmv	1 mW/cm²	96% conversion to CO_2	Ozaki [38]
	Trichloroethylene	Synthesized TiO ₂	240 ppm	not specified	1.7 μmol/s/g	Maira [39]
	Tetrachloroethylene	TiO ₂ , not specified	16.5 g/m³	32.6 mW/cm ²	56.7% conversion	Hager [40]
Aldehydes	Formaldehyde	P25	~ 45 ppmv	0.0093 W/cm ²	10 μmol/cm²/h	Obee [37]
	Acetaldehyde	PC500	170 ppmv	205 μW/cm² @ 365 nm	99% conversion	Verbruggen [41]
	Butyraldehyde	P25	118 mg/m³	5 x 10 ⁻⁷ E/cm²/min	1.5 μg/cm²	Peral [42]
Alcohols	Methanol	Synthesized TiO ₂	300 ppm	2.095 mW/cm ²	~1.7 x 10 ³ mol/m ³ /min	Kim [43]

Table 1. Selected examples of pollutants degraded by PCO on TiO₂-based photocatalysts in gaseous environment

	Ethanol	Synthesized TiO ₂	Not specified, spike vapors	0.12 mW/cm ²	Mechanistic elucidation	Coronado [44]
	2-propanol	PC500	400 ppbv	3.2 mW/cm ²	96% mineralization	Vildozo [45]
	1-butanol	Hombikat UV100	145 ppm	<0.65 mW/cm² (UV part)	>99% mineralization	Kirchnerova [46]
Ketones	Acetone	Synthesized TiO_2	450 ppmv	2.7 mW/cm ²	100% conversion	Coronado [47]
Aromatics	Benzene	P25	<5 ppm	5 mW/cm²	100% conversion	Jacoby [48]
		P25	100 ppmv	5.7 mW/cm ² UVC	100% conversion	Bouazza [49]
	Toluene	P25	0.45 g/m³	3.05 mW/cm ² UVC	3.9 g/m²/h	Keshmiri [50]
		TiO ₂ , not specified	1043 mg/m³	not specified, UVC	100% conversion	Bouzaza [51]
	m-xylene	P25	not specified trace	5 x 10 ⁻⁷ E/cm²/min	0.2 μg/cm²/min	Peral [42]
	Phenol	P25	not specified; adsorbed vapor	not specified for gas experiment	Full conversion + mechanistic elucidation	Palmisano [52]
Dioxins ^d	1,2-dichlorobenzene	P25	3.9 mmol/Nm ³	14.8 mW/cm ²	~100% destruction and removal efficiency	Lu [53]
	2,3,7,8-tetrachlorodibenzo- p-dioxin	P25	500 ng/cm² on solid-gas interface	not specified	99.86% conversion	Wu [54]
	1,2,3,6,7,8- hexachlorodibenzo-p-dioxin	P25	12.5 ng/cm ² on solid-gas interface	6.33 mW/cm ²	100% conversion	Wu [55]

^a all catalysts are TiO₂-based. This column mainly indicates if commercially available, or lab-made catalysts were used.

^b given in the same units or terminology as reported in (or directly derived from) the corresponding reference

^c either the conversion of the pollutant, the oxidation rate or a mechanistic insight, in the same units or terminology as reported in (or directly derived from) the corresponding reference

^d including model compounds for dioxins

Table 2. Dimensionless ratio of several material and catalytic properties of PC500 and P25 for the gas phase PCO of acetaldehyde.^a

Property	Ratio PC500/P25		
Turnover frequency (TOF)	1.3		
Specific surface area	6.0		
Acetaldehyde adsorbed	3.5		
Absolute amount OH groups	3.75		
Photocurrent	<0.01		

^a adapted from ref. [41] based on the original data