



## OpenAIR@RGU

### The Open Access Institutional Repository at Robert Gordon University

<http://openair.rgu.ac.uk>

This is an author produced version of a paper published in

Journal of Chemical Technology and Biotechnology (ISSN 0268-2575,  
eISSN 1097-4660)

This version may not include final proof corrections and does not include  
published layout or pagination.

#### Citation Details

##### Citation for the version of the work held in 'OpenAIR@RGU':

MCCULLAGH, C., SKILLEN, N., ADAMS, M. and ROBERTSON, P. K. J.,  
2011. Photocatalytic reactors for environmental remediation: a  
review. Available from *OpenAIR@RGU*. [online]. Available from:  
<http://openair.rgu.ac.uk>

##### Citation for the publisher's version:

MCCULLAGH, C., SKILLEN, N., ADAMS, M. and ROBERTSON, P. K. J.,  
2011. Photocatalytic reactors for environmental remediation: a  
review. *Journal of Chemical Technology and Biotechnology*, 86 (8),  
pp. 1002-1017.

#### Copyright

Items in 'OpenAIR@RGU', Robert Gordon University Open Access Institutional Repository,  
are protected by copyright and intellectual property law. If you believe that any material  
held in 'OpenAIR@RGU' infringes copyright, please contact [openair-help@rgu.ac.uk](mailto:openair-help@rgu.ac.uk) with  
details. The item will be removed from the repository while the claim is investigated.

This is the accepted version of the following article: MCCULLAGH, C., SKILLEN, N., ADAMS, M. and ROBERTSON, P. K. J., 2011. Photocatalytic reactors for environmental remediation: a review. *Journal of Chemical Technology and Biotechnology*, 86 (8), pp. 1002-1017, which has been published in final form at <http://dx.doi.org/10.1002/jctb.2650>

1 **Photocatalytic reactors for Environmental Remediation: A review.**

2

3 Cathy McCullagh, Nathan Skillen, Morgan Adams and Peter K.J. Robertson\*

4 IDeaS, Innovation, Design and Sustainability Research Institute, Robert

5 Gordon University, Schoolhill, Aberdeen, AB10 1FR, UK

6 \*Corresponding Author:

7 Tel: +44 1224 263750

8 Fax: +44 1224 262759

9 [peter.robertson@rgu.ac.uk](mailto:peter.robertson@rgu.ac.uk)

10

11

12

1 **Abstract**

2 **OVERVIEW**

3 Research in the field of photocatalytic reactors in the past three decades has  
4 been an area of extensive and diverse activity with an extensive range of  
5 suspended and fixed film photocatalyst configurations being reported. The key  
6 considerations for photocatalytic reactors, however, remain the same; effective  
7 mass transfer of pollutants to the photocatalyst surface and effective  
8 deployments and illumination of the photocatalyst.

9 **IMPACT**

10 Photocatalytic reactors have the potential versatility to be applied to the  
11 remediation of a range of water and gaseous effluents. Furthermore they have  
12 also been applied to the treatment of potable waters.

13 **APPLICATIONS**

14 Photocatalytic reactors are being scaled-up for consideration within waste and  
15 potable water treatment plants. Furthermore systems for the reduction of  
16 carbon dioxide to fuel products have also been reported.

17 **Keywords:** Photocatalyst, reactor, fluidised bed, immobilised film, suspended  
18 catalyst mass transport, rate control.

19

## 1 INTRODUCTION

2 The application of semiconductor photocatalysis in the fields of engineering and  
3 science is an important area of research which has grown significantly in the last  
4 three decades with increasing numbers of publications appearing every year <sup>1-6</sup>.  
5 Semiconductor photocatalysis has been applied to a diverse array of  
6 environmental problems including air, potable and wastewater treatment. This  
7 versatile process has also been utilised for the destruction of micro-organisms  
8 such as bacteria<sup>6, 7</sup>, viruses <sup>8</sup> and for the inactivation of cancer cells<sup>9, 10</sup>.  
9 Semiconductor photocatalysis has also been applied to the photo-splitting of  
10 water to produce hydrogen gas <sup>11-14</sup>, nitrogen fixation <sup>15-18</sup> and for the remediation  
11 of oil spills <sup>19-21</sup>.

12 Heterogeneous photocatalysis for the remediation of polluted water streams falls  
13 into two distinct areas. Firstly the focus is on basic chemical transformations on  
14 semiconductor photocatalyst materials. These investigations have concentrated  
15 on the examination of basic photocatalytic processes such as including  
16 photocatalytic material science, surface interactions on photocatalysts, reaction  
17 mechanisms and kinetics that impact on the processes on a molecular level <sup>1-5</sup>.

18

19 In order to demonstrate the viability of semiconductor photocatalysis for  
20 environmental remediation, reactor design is an equally critical factor. Effective  
21 reactor design research and development aims to scale up lab bench scale  
22 processes to industrially feasible applications. Scaling up photocatalytic reactors  
23 is, however a complex process with many factors needing consideration to yield  
24 a technically and economically efficient process. These factors include  
25 distribution of pollutant and photocatalyst, pollutant mass transfer, reaction

1 kinetics and irradiation characteristics. The issue of effective photocatalyst  
2 illumination is particularly important as this essentially determines the amount of  
3 water that may be treated per effective unit area of deployed photocatalyst. A  
4 wide range of photocatalytic reactors have been developed and used in both  
5 basic research and pilot scale studies. The central problem of scale-up of  
6 photocatalytic reactors is the provision of sufficient high specific surface area of  
7 catalyst and the uniform distribution of illumination across this area.

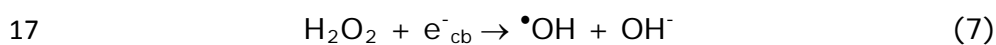
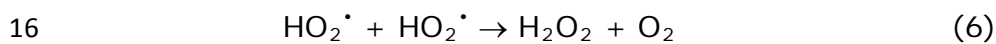
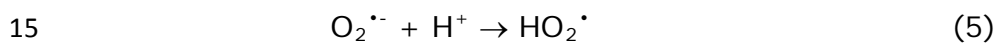
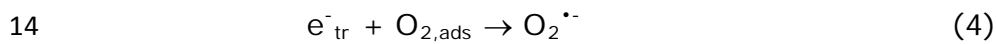
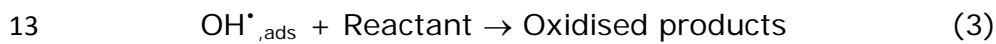
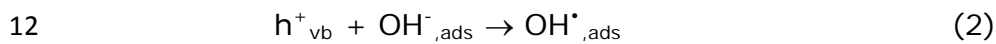
8

### 9 **Mechanism of heterogeneous photocatalysis**

10 As a process for water purification, photocatalysis has vast advantages over  
11 many existing technologies. The technique can result in the mineralisation of  
12 pollutants rather than transferring them to an alternative phase, such as is the  
13 case with activated carbon adsorption. Furthermore photocatalysis does not  
14 require the use of hazardous materials such as hypochlorite, peroxide or ozone<sup>4</sup>.  
15 Titanium dioxide is the catalyst of choice as it is inexpensive, non-toxic,  
16 chemically stable and is highly photocatalytically active<sup>2</sup>.  $\text{TiO}_2$  acts as a  
17 photocatalyst due to its electronic structure, characterised by an electronically  
18 filled valence band and empty conduction band<sup>22</sup> separated by a band gap. If a  
19 photon of energy greater than or equal to the bandgap energy,  $E_g$ , is absorbed  
20 by  $\text{TiO}_2$ , an electron is promoted from the valence band to the conduction band.  
21 This generates a reducing electron in the conductance band and an oxidising  
22 hole in the valence band (figure 1). The excited conduction band electrons may  
23 recombine with the valence band holes generating heat energy. Alternatively  
24 they may be trapped in surface states, undergo reactions with electron donating  
25 or accepting species that are adsorbed on the  $\text{TiO}_2$  surface. The electron and

1 holes formed are highly charged and result in redox reactions, which can  
 2 ultimately result in the mineralisation of aqueous pollutants. Hydroxyl radicals  
 3 are believed to be generated on the surface of TiO<sub>2</sub> through a reaction of the  
 4 valence band holes with adsorbed water, hydroxide or surface titanol groups (Eq  
 5 2). The photogenerated conduction band electrons react with electron  
 6 acceptors such as oxygen which generates superoxide (O<sub>2</sub><sup>-</sup>) (Eq 4).  
 7 Thermodynamically the redox potential of the TiO<sub>2</sub> electron/hole pair should  
 8 enable the production of hydrogen peroxide, primarily via the reduction of  
 9 adsorbed oxygen<sup>3, 23, 24</sup>(Eqn 1-7).

10



18

19 Figure 1 illustrates the basic processes involved in photocatalysis.

20

21 **Figure 1:**

22

23

24

## 1 TYPES OF REACTOR CONFIGURATIONS

2 A wide variety of reactor configurations have been reported in the literature over  
3 the past 30 years including:

- 4 • annular photoreactor <sup>25, 26</sup>,
- 5 • packed bed photoreactor <sup>27</sup>,
- 6 • photocatalytic Taylor vortex reactor <sup>28-31</sup>,
- 7 • fluidised bed reactor <sup>32, 33</sup>,
- 8 • coated fibre optic cable reactor <sup>34</sup>,
- 9 • falling film reactor <sup>35</sup>,
- 10 • thin film fixed bed sloping plate reactor <sup>36</sup>,
- 11 • swirl flow reactor <sup>37</sup>,
- 12 • corrugated plate reactor <sup>38</sup>.

13

14 Table 1 shows a comprehensive yet not exhaustive overview of photoreactor  
15 type, reactant phase, experimental targets, catalyst employed and industrial  
16 applications.

### 17 **Table 1.**

18 Examples of these various reactors will be considered in the following sections.

## 19 **Suspended Liquid Reactors**

### 20 **Slurry/Suspension systems**

21 TiO<sub>2</sub> catalysts have been investigated in both slurry/suspension systems and  
22 immobilised systems. The main advantage of using photocatalyst slurries is the  
23 larger surface area compared to the immobilised system. The separation of the



1 nanometre catalyst particles is, however, expensive and is a major drawback in  
2 the commercialisation of this type of system<sup>39</sup>. Nan Chong *et al.*<sup>40</sup> have reported  
3 the use of H-titanate nanofibres in an annular slurry photoreactor. They studied  
4 congo red as a model compound but one of the key advantages was the settling  
5 velocity of the catalyst. Using Kynch's theory batch settling trials revealed that  
6 the H-titanate nanofibre photocatalysts resulted in a settling velocity of  $8.38 \times$   
7  $10^{-4} \text{ m s}^{-1}$ . The authors proposed that these novel nanoparticles could deliver a  
8 true engineering solution to catalyst separation on an industrial scale. It was  
9 also reported that it was difficult to effect irradiation of all the photocatalyst  
10 particles in the slurry in the unit due to shielding from the light source of the  
11 particles in the body of the unit from particles closer to the reactor walls..  
12 Therefore depth of light penetration into the slurry reactor was restricted.

13

14 It has been reported the ratio between backward reflected and incident photon  
15 flow is strongly influenced by the geometry within the reactor. The Apparent  
16 Napierian Extinctance (ANE) coefficient for slurries is a parameter which is  
17 related to the photon absorption rate<sup>41</sup>. ANE depends on the particle size and is  
18 inversely proportional to the particle size where:

19 
$$(\text{ANE})_{\lambda} = -\ln(I/I_o) \quad (8)$$

20  $I$  is the outgoing light intensity after interaction with catalyst suspension and  $I_o$   
21 represents the light intensity obtained in a blank setup. The absorption of light  
22 within slurry systems cannot be separated from scattering, which makes kinetic  
23 analysis of experiments more challenging<sup>42</sup>.

1 An alternative approach is to attach the catalyst to a transparent stationary  
2 support over which the contaminated water passes. In such a system it is  
3 possible achieve effective illumination of all the photocatalyst deployed in the  
4 reactor <sup>43, 44</sup>. There are, however, also problems associated with immobilised  
5 systems which include the dependence on mass transfer of pollutant to the  
6 photocatalyst surface and the ensuring effective access to the photocatalyst  
7 surface of both activating photons and reacting molecules to the photocatalyst  
8 surface<sup>45</sup>. The photocatalyst film thickness may affect the internal mass transfer  
9 as it may not be possible to access the photocatalyst material in the proximity of  
10 the support-catalyst interface. Each of these factors will result in a reduction in  
11 the rates of decomposition in immobilised film photocatalyst units when  
12 compared to slurry processes.

13

14 An early slurry based reactor effected the complete mineralisation of chloroform  
15 to chloride and CO<sub>2</sub> <sup>46</sup>. This study was further supported by Kormann *et al* <sup>47</sup> who  
16 demonstrated the complete dehalogenation of chloroform but also illustrated an  
17 increase in [Cl<sup>-</sup>] as a function of time. Pramauro *et al.* <sup>48</sup> reported the complete  
18 degradation of monuron (figure 2a), a persistent herbicide, within one hours  
19 photocatalysis. The reaction conditions investigated were simulated solar  
20 irradiation, TiO<sub>2</sub> slurry and a batch reactor. Several intermediate products were  
21 determined after 30 minutes irradiation; these were not detected at the end of  
22 the irradiation time. The complete degradation and dechlorination of 3,4-  
23 dichloropropionamide (figure 2b), another persistent herbicide was reported by  
24 Pathirana and Maithreepala <sup>49</sup> using a TiO<sub>2</sub> suspension photocatalytic reactor. It  
25 was reported that the dechlorination of the herbicide was dependent on TiO<sub>2</sub>

1 concentration, and higher photocatalyst levels caused TiO<sub>2</sub> a shielding effect of  
2 the incident light. Li Puma and Yue<sup>50</sup> investigated the kinetics of degradation of  
3 single component and multi-component systems of chlorophenols. They studied  
4 the simultaneous use of short, medium and long wavelength ultra-violet light to  
5 investigate the integration of simultaneous photocatalysis and photolysis. The  
6 optimal loading of photocatalyst for the geometry of their batch reactor via the  
7 oxidation of 2-CP to CO<sub>2</sub> was determined as part of this investigation with  
8 mineralization rates achieving a 3-fold increase up to a maximum photocatalyst  
9 loading of 0.5 mg L<sup>-1</sup>. In the single component experiments the kinetics were too  
10 complex to allow the authors to determine whether the photolytic pathway  
11 would be preferential to the photocatalytic pathway or vice versa. During the  
12 multi-component experiments they observed that the overall oxidation kinetics  
13 were controlled by the reactant in excess, as the substrate present in smaller  
14 concentrations was found to degrade at a much slower rate than that in the  
15 single-component experiments. Overall the authors demonstrated that the entire  
16 course of photocatalytic oxidation of single-component and multi-component  
17 systems of chlorophenols can be predicted satisfactorily using simple kinetic  
18 models that could be useful in the design and modelling of large scale  
19 photocatalytic reactors. This work was followed by studies of a pilot-scale  
20 continuous-flow laminar falling film slurry photocatalytic reactor (LFFSIW)<sup>51</sup>,  
21 which utilised commercially available UV lamps. A comprehensive investigation  
22 of this unit was reported looking at a range of parameters including; the  
23 wavelength and intensity of the incident irradiation source, the influence of  
24 additional oxidising reagents, the concentrations of both reactants and  
25 photocatalyst and the irradiation time. Li Puma and Yu also studied six different  
26 photon based processes, and determined that the UVC photocatalysis/UVC

1 photolysis/UVC peroxidation process was superior to UVA photocatalysis, UVC  
2 photocatalysis/UVC photolysis, UVA photocatalysis/UVA peroxidation, UVC  
3 photolysis/UVC peroxidation and UVC photolysis systems. The results of the  
4 experiments conducted at different incident radiation intensities clearly indicated  
5 that low-wattage UVC lamps were preferable to high-wattage UVC lamps  
6 because enhancements of reactor conversions due to higher lamp power were  
7 offset by an increase in electricity costs.

## 8 **FIGURE 2**

9 Further work by Li Puma and Yue <sup>35</sup> compared the effectiveness of a range of  
10 photooxidation processes in a falling film pilot reactor. They investigated UVA-  
11 photocatalysis, UVA-photocatalysis-peroxidation, UVC-photolysis, UVC-  
12 photolysis-peroxidation, UVC-photocatalysis-photolysis and UVC-photocatalysis-  
13 photolysis-peroxidation. They selected salicylic acid as a model compound for all  
14 experiments. The highest conversion of salicylic acid was 58%, which was  
15 obtained with the UVC-photolysis-peroxidation process. Conversely the highest  
16 conversion to CO<sub>2</sub> was obtained with the UVC-photocatalysis-photolysis-  
17 peroxidation process (28%) and the lowest value was with the UVA-  
18 photocatalysis process (1.7%). The mineralisation efficiency of the photocatalytic  
19 and the photolytic-photocatalytic processes was in the range 58-65%, far above  
20 that of the photolytic process which was in the range of 14-18%.

21

22 San *et al.* <sup>52</sup> investigated the photodegradation of 3-aminophenol (3-AP) in a TiO<sub>2</sub>  
23 batch suspension photoreactor. They reported that this process followed  
24 pseudo-first-order kinetics, with the apparent rate constant depending on the

1 initial 3-AP concentration. Furthermore the addition of electron acceptors  
2 enhanced the reaction rate significantly.

3

4 A study of the adsorption and photocatalytic degradation of the safira HEXL dye,  
5 has been reported using a TiO<sub>2</sub> slurry reactor<sup>53</sup>. The authors concluded that the  
6 dye adsorption to the photocatalyst surface was critical for the efficient  
7 photocatalytic degradation. The process was also pH dependent with an  
8 improved degradation rate observed near the point of zero charge of TiO<sub>2</sub>. The  
9 slurry TiO<sub>2</sub> catalyst has also been employed within an internally circulating  
10 bubble column reactor for the degradation of trichloroethylene with a removal  
11 efficiency of 97% reported<sup>54</sup>. Employing a photocatalyst filtration with a bubble  
12 column reactor and TiO<sub>2</sub> slurry may be efficient process for the destruction of  
13 phenoxyacetic acid<sup>55</sup>. This overcomes one of the major drawbacks of slurry  
14 systems, separation of catalyst from remediated waste stream.

15

16 Adams *et al*<sup>56</sup> reported the comparison of a standard flat plate reactor with a  
17 novel drum reactor<sup>57</sup> for the removal of oil and gas hydrocarbon contaminants  
18 from water. The flat plate design was constructed from polymethylmethacrylate  
19 (PMMA) and was as a small scale lab unit, however, a 'concertina' designed  
20 reactor was proposed for the scale up. The drum reactor concept was a single  
21 pass continuous flow system for the treatment of waste water/effluents which  
22 utilised rotating paddles to ensure an even distribution of catalyst<sup>56</sup> (figure 3).

23 **FIGURE 3**

1 The reactor set up utilised three connecting drum reactors. The paddles  
2 positioned on the inside of the reactor drum were placed as to allow the removal  
3 of pellets from the reaction solution allowing for exposure to UV illumination  
4 before returning to the main stock<sup>56, 57</sup>. Dependant on the level of contaminant  
5 present the effluent would pass from one 'drum' to another for prolonged  
6 treatment. On average the sample would reside in each drum for approximately  
7 3 min with the overall reaction time at approximately 10 min. In the event of the  
8 final sample still being of a high hydrocarbon concentration the effluent would be  
9 recirculated into the system. As an attempt to develop a reactor deemed more  
10 environmentally efficient, pelletised catalyst was used to reduce downstream  
11 processing restrictions associated with the filtration of powdered catalyst. While  
12 both designs proved effective in the removal of hydrocarbons the drum reactor  
13 achieved 90 % removal in less than 10 min. This high level of reduction over a  
14 short period of time is attributed to samples passing through the 3 consecutive  
15 drums each with 200 g of TiO<sub>2</sub> catalyst present, therefore utilising 600 g for the  
16 entire system. The flat plate design was investigated in regards to optimising  
17 conditions to provide maximum destruction. It was concluded that a lower  
18 angled plate increases retention times of compounds and thus a chance of  
19 successful catalyst-pollutant interface<sup>56</sup>. The addition of air and hydrogen  
20 peroxide to increase destruction proved effective with 80% degradation over 135  
21 min recorded for H<sub>2</sub>O<sub>2</sub> compared to 40 % for air alone.

22 In a subsequent paper McCullagh *et al.*<sup>58</sup> described the degradation of  
23 methylene blue (MB) in a slurry continuous flow drum reactor (figure 4). A 98 %  
24 degradation of MB over 60 min of illumination utilising a high loading weight of  
25 30 g L<sup>-1</sup> of TiO<sub>2</sub> pellets was reported. The investigation of different loading  
26 weights concluded that a maximum weight of 180 g of catalyst was attainable.

1 While 98% degradation was recorded for 30 g catalyst (Pellet form) over 60 min,  
2 the use of Degussa P25 as a photocatalyst was significantly more efficient with  
3 90% of the MB be decomposed within 20 min photocatalysis.

#### 4 **FIGURE 4**

5 Further evaluation of the drum reactor has been carried out in the remediation  
6 of oily waste water (OWW) from an interceptor tank<sup>59</sup>. This study showed the  
7 unit achieved a 50% reduction in abundance for both decane and dodecane after  
8 90 min UV photocatalysis along with a >50 % abundance reduction in  
9 tetradecane. Following a further 90 min illumination period of the OWW in the  
10 reactor all volatile organic compounds (VOC) initially identified were almost  
11 completely removed. Additionally total organic carbon TOC was investigated  
12 showing a 35 % reduction in TOC for OWW samples passed twice through the  
13 reactor. The results highlighted demonstrate a high efficiency for a novel  
14 photocatalytic reactor with large scale applications as a polishing technique to be  
15 coupled with current water treatment processes<sup>59</sup>.

#### 16 **Mass Transfer Limitations within TiO<sub>2</sub> slurry reactors**

17 Mass transfer within slurry reactors has not received a great deal of attention  
18 primarily as such problems have not been recognised as being major  
19 impediments to the application of a slurry reactor. Chen and Ray<sup>60</sup> did not  
20 observe either intra or extra particle diffusive limitations when working with a  
21 suspended solid reactor. Considering the optical thickness of the suspension  
22 Martin *et al.*<sup>61</sup> found a loss in reactor efficiency when the optical thickness of the  
23 suspension was greater than an calculated optimum level. For large catalyst  
24 loading restrictions with mass and radiation transport inside the catalytic particle  
25 has been reported by Mehrotra *et al.*<sup>62</sup>. Peralta Muniz Moreira<sup>63</sup> found no

1 diffusion limitations within their reactor when pseudo 1<sup>st</sup> order kinetics were  
2 applied along with the Weiz and Prater criterion. How mass transport influences  
3 the rate of reaction within slurry reactors is very important in order to obtain  
4 relevant kinetic information about the reaction but also for photoreactor design.  
5 Ballari *et al.*<sup>64</sup> investigated the effects of photocatalyst irradiation, photocatalyst  
6 loadings, flow rates, total suspension volume and changes in illumination length  
7 of the reactor. They found significant concentration gradients that were likely to  
8 cause limitations in mass transport resulted from the non-uniformity of the  
9 irradiation area. The authors state that these concentration gradients are difficult  
10 to avoid but could be eradicated if a fully developed turbulent flow operated  
11 within the reactor. They concluded that mass transport problems could be  
12 overcome using a 1 g/L catalyst loading, irradiation rates of  $1 \times 10^{-7}$  Einsteins  
13  $\text{cm}^{-1} \text{s}^{-1}$  and effective mixing.

14

## 15 **Fluidised bed reactors**

16 The application of fluidised bed reactors have been extensively reported with  
17 round 1000 published on this topic over the past 6 years<sup>32, 64-74</sup>. Fluidised bed  
18 reactors are capable of utilising an upward stream of fluid (gas or liquid) to allow  
19 particles in a stationary phase to be brought to a suspended or 'fluidised' state  
20 allowing for photocatalytic transformations to occur (figure 5).

## 21 **FIGURE 5**

22 The advantages of this style of reactors include:

- 23 • a low pressure drop,
- 24 • high throughput and



- 1 • high photocatalyst surface area which consecutively allows for increased  
2 catalyst-reactant interaction.

3

4 In 1992 Dibble and Raupp<sup>66</sup> used a flat plate fluidised bed reactor (figure 6) to  
5 photoxidise trichloroethylene (TCE). A quantum efficiency range of 2-13 % was  
6 achieved with a reaction rate peaking at  $0.8\mu\text{mol TCE (g of catalyst)}^{-1} \text{ min}^{-1}$  [ $2$   
7  $\mu\text{mol}/(\text{g of TiO}_2)^{-1} \text{ min}^{-1}$ ]. These results are significant in that they are  
8 comparable to results produced in a liquid-solid slurry system for the oxidation  
9 of TCE, specifically demonstrating an order of magnitude increase. TCE is a  
10 standard evaluation test carried out by many including more recently Lim and  
11 Kim who investigated a circulated fluidised bed reactor (CFDB) <sup>67</sup>. The design of  
12 this reactor used a loop seal which allowed particles that were carried up by the  
13 air stream to flow back down and re-enter the system. Several other factors  
14 were investigated in this paper including UV wavelength, initial TCE  
15 concentration, circulation rate and O<sub>2</sub> and H<sub>2</sub>O concentrations.

## 16 **FIGURE 6**

17 A two-dimensional fluidised bed reactor was used by Lim *et al.*<sup>75</sup> in order to  
18 decompose NO. Reported in the research was efficient contact between the  
19 catalyst (P-25) and reactant gas (NO) which, when coupled with good UV-light  
20 transmission allowed for increased NO decomposition in comparison to an  
21 annular flow-type photoreactor. A series of altering conditions were tested as a  
22 means to increase photocatalytic efficiency including; initial gas concentration,  
23 residence time of gas, reaction temperature and irradiation intensity. A  
24 particular point of interest in this publication surrounded the superficial gas  
25 velocity and light transmission interaction. Results displayed in figure 7 show

1 increasing superficial gas velocity increased light transmission with light intensity  
2 significantly increasing at approx.  $1.3 U_{mf}$  (minimum fluidisation velocity) which  
3 equated to a sharp increase in NO conversion at  $1.3 U_{mf}$ . Below this value the UV  
4 light was not capable of transmitting through the catalyst bed. An increasing  
5 trend is observed until the  $U_{mf}$  reaches 2.5, where a 70 % photocatalytic  
6 decomposition of NO was recorded. Conversion recorded after this point was  
7 regarded as a result of NO bypassing through bubbles and the reduction of gas  
8 residence time in the catalysts bed<sup>69, 75</sup>. Based upon their results Lim *et al.*<sup>75</sup>  
9 concluded that NO photocatalytic decomposition required adequate residence  
10 time and an effective NO gas velocity to enable the appropriate bubble size  
11 formation enabling appropriate contact between UV light and the TiO<sub>2</sub>-NO  
12 system.

### 13 **FIGURE 7**

14 Son *et al.*<sup>70</sup> investigated the use of combined TiO<sub>2</sub> particles with Al<sub>2</sub>O<sub>3</sub> in an  
15 attempt to overcome the drawbacks associated with fluidised reactors for  
16 photocatalysis. Paz<sup>65</sup> reported that fluidisation of small particles such as P-25  
17 was challenging due to 'drifting' from the primary operation area in the unit.  
18 Combining the catalyst, however, with larger particles such as Al<sub>2</sub>O<sub>3</sub> could  
19 eliminate this problem and as such many researchers use an Al-TiO<sub>2</sub> catalyst.  
20 Son<sup>70</sup> focused on the decomposition of acetic acid and ammonia utilising a three  
21 phase photocatalytic system. Decomposition during the research was enhanced  
22 when carried out in the three-phase fluidised bed reactor, showed significant  
23 improvement over use in a steady reactor. Production of N<sub>2</sub> and CO<sub>2</sub> were  
24 monitored as means of measuring acetic acid and ammonia decomposition. In  
25 terms of acetic acid decomposition the fluidised bed showed increased efficiency

1 over that of a conventional steady reactor along with increased efficiency by the  
2 addition of Al-TiO<sub>2</sub> instead of solely TiO<sub>2</sub>; the conversion of acetic acid to CO<sub>2</sub>  
3 reached approximately 90 % after 600 min with 10 mol% Al-TiO<sub>2</sub>. A similar  
4 trend was observed for ammonia decomposition with a conversion rate of >95 %  
5 being reached with 10 mol% Al-TiO<sub>2</sub> compared to that of 70 % in the steady  
6 reactor. A point of interest is the suppression of the more undesirable products  
7 of NO<sub>2</sub> and NO<sub>3</sub> with the use of Al-TiO<sub>2</sub> when compared to increased levels with  
8 pure TiO<sub>2</sub>. In a previous study which used a FeTiO<sub>2</sub> material it was reported that  
9 the anatase structure of the catalyst transformed into a rutile structure after  
10 methanol destruction<sup>71</sup>. Son *et al.*<sup>70</sup>, however, found replacing the Fe with Al  
11 produces a catalyst with increased stability, thus providing enhancing ammonia  
12 decomposition. They concluded from their results that the removal of VOC  
13 efficiency is increased by both the use of Al-TiO<sub>2</sub> combined particles and a  
14 fluidised reactor.

15

16 Nelson *et al.*<sup>72</sup> reported the comparison of a fluidised TiO<sub>2</sub> system for methanol  
17 oxidation with a packed bed reactor. They concluded that fluidisation resulted in  
18 faster rates of photocatalytic decomposition than achieved on the packed bed  
19 unit. A rate of  $2.0 \times 10^{-7}$  mol/cm<sup>3</sup> cat/min for CO<sub>2</sub> production was achieved for  
20 the fluidised reactor compared to a CO<sub>2</sub> production rate of  $1.0 \times 10^{-7}$  mol/cm<sup>3</sup>  
21 cat/min obtained with the packed bed reactor. It was reported that the use of  
22 both static mixing and vibration in the process to reduce photocatalyst  
23 separation rates was, however, only effective with Degussa P25 and not TiO<sub>2</sub>-  
24 Al<sub>2</sub>O<sub>3</sub>. Overall TiO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> was found to be an effective photocatalyst which is in  
25 agreement with the results obtained by Paz<sup>65</sup>.

1

2 The drawbacks to fluidising pure  $\text{TiO}_2$  has led to an increased number of papers  
3 reporting results using combined catalyst, demonstrated by Kuo *et al.*<sup>32</sup> who  
4 investigated the removal of toluene vapours from a continuous gas stream. The  
5 specific limitations include the 'loss or trapping' of powder within a photoreactor  
6 due to the fine structure of  $\text{TiO}_2$ . These problems can often result in the  
7 powdered catalyst 'drifting' away from the main area of operation. The  
8 investigation used activated carbon (AC) particles with a  $\text{TiO}_2$  coating to  
9 overcome any fluidisation problem and promote good evenly distributed catalyst.  
10 The research investigated the impact of altered relative humidity (RH), varied  
11  $\text{TiO}_2$  loading weights, and the use of glass beads (GB) in replacement of AC  
12 along with and without the use of a reflector. The use of activated carbon solely  
13 is effective in the removal of toluene, however, upon saturation of the AC  
14 particles toluene removal decreased significantly. It was established that 30%  
15 RH was optimal for efficient toluene removal. Interestingly increasing RH did not  
16 result in increased rates of toluene removal suggesting there is competitive  
17 adsorption between water and toluene molecules at higher levels of RH. The  
18 degradation of toluene vapours was accredited to both the use of activated  
19 carbon and photocatalysis, however, saturation resulted in certain restrictions  
20 when solely using AC. The results displayed that the combination of AC removal  
21 of toluene and the photocatalytic removal could significantly extend toluene  
22 removal duration. In comparing the effectiveness of the two catalyst types it was  
23 found that the GB/ $\text{TiO}_2$  was half as effective as AC/ $\text{TiO}_2$  catalyst for toluene  
24 removal.

25

1 Voronstov *et al.*<sup>74</sup> also investigated the use of vibration to improve fluidisation of  
2 granular photocatalysts for the decomposition of gaseous acetone. A variety of  
3 fixed bed constructions were investigated together with the vibrofluidised bed  
4 system to enable an efficient comparison. The vibrofluidised bed system was the  
5 most effective with an 8.7 % in quantum efficiency being achieved. The high  
6 efficiency of the vibrofluidised bed was, interestingly, attributed to the external  
7 vibrations used together with the 'periodic light phenomenon'. This phenomenon  
8 resulted from the eccentric movement photocatalyst movement within the  
9 reactor which consequently enabled increased absorption of scattered light.

10

11 A fluidised bed system utilising an upward stream of air which brought TiO<sub>2</sub>  
12 pellets to a fluidised state has also been reported<sup>76</sup>. The reactor consisted of a  
13 reaction chamber which contained a foraminated member supporting a bed of  
14 mobile photocatalysts along with an aeration device to allow for agitation of  
15 photocatalytic particles. The aeration device generated gas bubbles through a  
16 perforated shelf allowing agitating of photocatalysts. The reactor configuration  
17 was utilised for the treatment of waste water in a flow through style process.  
18 The reactor concept was designed to reduce or completely remove the need for  
19 moving parts, thus allowing for a reactor concept with a reduced foot print  
20 suggesting a more energy efficient design. To ensure the constant agitation of  
21 the particles the terminal settling velocity of the particles must not exceed the  
22 velocity of any upward flow of the liquid through the perforated shelf by more  
23 than 10 ms. The reactor retained the advantage of the use of pellets which both  
24 allow for reduced downstream processing and via agitation present a number of  
25 faces capable of excitation by illumination.

## 1 Immobilised Liquid Reactors

### 2 Fixed bed

3 Al-Ekabi and Serpone <sup>77</sup> investigated TiO<sub>2</sub> supported on a glass matrix for the  
4 photo-decomposition of phenol, 4-chlorophenol, 2,4-dichlorophenol, and 2,4,5-  
5 trichlorophenol. The degradation followed first order kinetics with the reaction  
6 occurring on the surface of the semiconductor. The irradiation source was an  
7 AM-1 filter simulating solar irradiation. A fixed bed reactor system employing a  
8 fibre-optic cable (OFR) was reported by Pill and Hoffmann <sup>34</sup>. The system was  
9 conceived to allow for remote light distribution to photocatalysts, to effectively  
10 determine of quantum yields through effective light flux measurement.  
11 Furthermore OFR allowed for reactor reuse to assess different coatings and light  
12 input angles, and to minimise potential heating and photocatalyst delamination.  
13 They anchored TiO<sub>2</sub> particles onto quartz fibres and light was transmitted to the  
14 TiO<sub>2</sub> particles via radial refraction of light out of the fibre. A maximum quantum  
15 efficiency of  $\phi = 0.011$  for the oxidation of 4-chlorophenol was achieved. This  
16 can be compared to a maximum quantum efficiency of  $\phi = 0.0065$  for 4-  
17 chlorophenol oxidation in a TiO<sub>2</sub> slurry reactor.

18 This study was followed with an investigation into the application of the OFR  
19 system towards the photocatalytic degradation of pentachlorophenol, oxalate  
20 and dichloroacetate <sup>34</sup>. Relatively high apparent quantum efficiencies of  $\phi =$   
21 0.010, 0.17, and 0.08 were achieved for PCP, OX and DCA respectively, with  
22 complete mineralisation reported. It was concluded that the OFR system had the  
23 advantages of a fixed-bed unit together with the kinetic efficiency of a slurry  
24 reactor. The OFR configuration enhanced the not only the distribution but also  
25 the uniformity of activated photocatalyst within a particular reaction volume

1 compared to standard fixed-bed designs. These characteristics reduced mass  
2 transport limitations for photochemical conversion efficiency and allowed higher  
3 processing capacities. Furthermore, potential light loss via absorption or  
4 scattering by the reaction medium was minimised. The OFR system could be  
5 used in batch or continuous flow operation for both liquid and or gas phase  
6 reactions. The transmission cable also allowed for remote light delivery to the  
7 photocatalyst.

8

9 Nogueira and Jardim <sup>78</sup> reported the photodegradation of methylene blue using  
10 solar irradiation on a fixed bed reactor with TiO<sub>2</sub> immobilised on a flat glass plate  
11 as a support. They investigated the slope of the plate and found that it  
12 influenced the methylene blue photodegradation because of 2 factors:

- 13 i. the fluid thickness film which flowed over the plate and
- 14 ii. the light intensity that reached the system.

15

16 They reported a limited range of slopes 22° – 25° and found that 95.8 % of the  
17 model compound was degraded at 22° slope while 89 % was degraded at 25°  
18 angle. Ray and Beenackers<sup>78</sup> proposed a distributive type fixed bed reactor  
19 system that employed hollow glass tubes as of light conductors for distribution  
20 to photocatalyst particles. The reactor configuration increased the surface to  
21 volume ratio while eliminating the potential light loss through absorption and  
22 scattering by reaction matrix. This configuration facilitated a large surface of  
23 photocatalyst to be deployed within a relatively small reactor volume. Between  
24 70 and 100 fold increase in surface area per m<sup>3</sup> of reactor volume was achieved  
25 compared to a conventional annular reactor configuration. The photo-

1 degradation of special brilliant blue, a model dye pollutant, was investigated and  
2 a 90% photocatalytic destruction of the dye achieved after 100 min irradiation.  
3 This study was followed up with the development of a tube light reactor which  
4 had a 100-150 fold increase in surface area per unit volume of fluid being  
5 treated compared to a conventional annular reactor design and a 10-20 fold  
6 increase contrasted with an immersion reactor. In a study of a reactor volume of  
7  $3.65 \times 10^{-4} \text{ m}^3$  containing 21 U-shaped lamps of diameter 0.45 cm coated with  
8 the a P25 photocatalyst, a 695% increase reactor efficiency was achieved  
9 compared with an annular photocatalytic reactor. Furthermore a 259 % increase  
10 in efficiency was obtained for the new unite compared with a slurry reactor.

11

12 Feitz *et al.*<sup>79</sup> investigated two fixed bed photocatalytic reactors, a packed bed  
13 reactor and a coated mesh reactor, using solar illumination. They assessed the  
14 processing rate for 2 mg l<sup>-1</sup> phenol solutions, and calculated a rate of 140 mg m<sup>-2</sup>  
15 h<sup>-1</sup> for the packed bed reactor with a rate of 20 mg m<sup>-2</sup> h<sup>-1</sup> for the coated mesh  
16 reactor. The lower activity obtained with the coated mesh reactor was believed  
17 to be due to insufficient photocatalyst surface contact, low levels of available  
18 attached TiO<sub>2</sub> and a small reactor to tank volume ratio. Photonic efficiencies for  
19 the decomposition of 100 mg l<sup>-1</sup> dichloroacetic acid solutions using the packed bed  
20 unit were only 40% lower than suspension systems. They therefore proposed  
21 that this system was particularly effective for treating contaminated water.

22 Dionysiou *et al.*<sup>80</sup> developed a TiO<sub>2</sub> rotating disk reactor for the decomposition  
23 of organic pollutants in water (figure 8). They used a commercial TiO<sub>2</sub> composite  
24 ceramic ball photocatalyst material. LiCl tracer studies performed under different  
25 disk angular velocities, between 5 and 20 rpm, demonstrated that mixing in the



1 rotating disk photocatalytic reactor RDPR was similar to that of a continuous  
2 stirred tank reactor. They reported the destruction of >90 % 4-Chlorobenzoic  
3 acid after 6 hours irradiation. The light intensity distribution within the reactor  
4 was also determined and found to vary from about 30 to 1500  $\mu\text{W}/\text{cm}^2$  within  
5 the reactor. The RDPR has a number of advantages it eliminates the need for  
6 effluent filtration as the catalyst is immobilised, the 3-D nature of the flow  
7 created enabled effective mixing, whilst the formation of the thin film allowed  
8 more effective oxygen transport from the gas phase to the photocatalyst  
9 surface. The photonic efficiency calculated for the experiment at 4 rpm was 2.7  
10 %. The authors anticipate this value to improve with process optimisation<sup>81,82</sup>. A  
11 similar study by Hamill *et al.*<sup>82</sup> reported the use of a sealed rotating  
12 photocatalytic reactor (RPC) of similar configuration to the Dionysiou RDPR<sup>80, 81,</sup>  
13<sup>83</sup>. They investigated the photo-degradation of dichlorobutene and examined the  
14 effects of mass transfer and combinations of pollutants. They reported that the  
15 RPC could effectively degrade a range of substrates and that the degradation  
16 rate was dependent on rotation speed. They also reported that this configuration  
17 could be applied to both volatile and non-volatile pollutants.

## 18 **FIGURE 8**

19 Mehrvar *et al.*<sup>27</sup> reported the use of a photoreactor with  $\text{TiO}_2$  coated tellerette  
20 packing. The tellerette packings were constructed from stainless steel welding  
21 wire which was 'roughened' to promote the adhesion of  $\text{TiO}_2$  to the surface. The  
22 wires were then wound in a spiral or 'spring like' structure to be formed. The  
23 wound wire was cut into smaller lengths and each individual 'spring' adjusted to  
24 allow the ends to be brought together to form the tellerete shape. The tellerette  
25 type packings were selected and manufactured on the basis that they would

1 permit sufficient light dispersion into the interior of the bed to maintain effective  
2 photocatalytic reaction rates with no significant mass transfer limitations. The  
3 photoreactor allowed substantial UV light penetration throughout its interior, and  
4 had no significant mass transfer limitations during the photocatalytic degradation  
5 of 1,2-dioxane. It was concluded that the range of attenuation coefficients of  
6 interest, the reaction rate at various radial positions will not be significantly  
7 mass transfer limited unless the photocatalyst activity is increased by at least  
8 one order of magnitude.

9

10 A novel photocatalytic reaction system, composed of solution and gas spaces  
11 that were divided by a thin Teflon film and TiO<sub>2</sub> coated mesh or cloth has been  
12 reported<sup>84</sup>. The activity of TiO<sub>2</sub> immobilised on a stainless steel mesh and on a  
13 fibre-glass cloth using isopropanol as a model compound was investigated.  
14 Although both support materials yielded comparable photocatalytic activities the  
15 fibre glass cloth was the most stable. The Teflon membrane enhanced the O<sub>2</sub>  
16 levels in the reaction solution which increased the photocatalytic activity for the  
17 destruction of organic compounds in water. The benefit of this system for the  
18 photo-degradation of aqueous volatile organic carbons was that it did not require  
19 air bubbling, which resulted in volatilization of the contaminants to the  
20 atmosphere.

21

22 Lim *et al.*<sup>68</sup> reported the use of an external lamp, annular photocatalytic reactor  
23 with TiO<sub>2</sub> adsorbed on a quartz tube for the degradation of phenanthrene and  
24 pyrene from a dilute water stream. They reported that above a feed velocity of 7

1 cm min<sup>-1</sup> the process was rate controlled and not influenced by mass transfer  
2 limitations.

3 A tubular photocatalytic reactor for water purification using a ceramic cylindrical  
4 tube with a Pt-loaded TiO<sub>2</sub> film coated on the inner surface of the tube has been  
5 developed by Zhang *et al*<sup>85</sup>. Phenol, trichloroethylene and bisphenol A were  
6 used as model pollutants to examine the effectiveness of the photoreactor. The  
7 complete degradation of each pollutant within 2 hours reaction time was  
8 observed with the authors concluding that the performance of the reactor was  
9 dependent on the aeration of the system.

10

11 McMurray *et al.*<sup>86</sup> reported the use of a stirred tank reactor with immobilised  
12 Degussa P25 TiO<sub>2</sub> for the degradation of oxalic acid and formic acid. The rate of  
13 degradation of both acids was not mass transfer limited with propeller speeds  
14 greater than 1000 rpm. They reported apparent quantum yields of 5 % for oxalic  
15 acid and 10 % for formic acid.

16

17 A sol-gel prepared TiO<sub>2</sub> coating on a tubular photocatalytic reactor with re-  
18 circulation mode and a batch photocatalytic reactor was investigated by Lin *at*  
19 *al*<sup>87</sup> for the degradation of methylene blue and phenol. The sol-gel film  
20 synthesised demonstrated an effective photocatalytic activity for the  
21 decomposition of organic compounds in water and the authors proposed the use  
22 of this reactor for water purification. During a 180 minute photoreaction of  
23 phenanthrene, 67.6 % destruction was observed with a 40.1 % conversion to  
24 CO<sub>2</sub>.

1

2 Zhang *et al.*<sup>38</sup> reported the use of a corrugated plate reactor configuration which  
3 was developed and assessed using 4-chlorophenol as model pollutant. They  
4 compared the new configuration with a flat plate reactor and a slurry reactor.  
5 The corrugated plate reactor was reported to be 150 % faster with mass  
6 transfer rates 600 % higher than that of a flat plate reactor. The authors  
7 suggested that the enhanced performance of the corrugated configuration was a  
8 result of the relatively larger illuminated photocatalyst surface area per unit  
9 volume, coupled with an effective delivery of both photons and reactants to the  
10 photocatalyst surface.

11

12 The photocatalytic oxidation of a non ionic surfactant was carried out in a  
13 labyrinth flow reactor with an immobilised photocatalyst bed<sup>88</sup>. The work focused  
14 on the effects of flow-rate for the decomposition of the non-ionic surfactant. The  
15 authors concluded that the optimum photodegradation of the surfactant was  
16 observed with a flow-rate of 11.98 dm<sup>3</sup>/h. They further studied the remediation  
17 of Acid Red 18, an azo dye <sup>89</sup>. Long reaction times were investigated for the  
18 photo-degradation of Acid Red using an immobilised Aeroxide Degussa P25  
19 catalyst. Slower flow rates affected the efficiency of the system, with  
20 mineralisation times varying from 35 hours to 60 hours depending on flow rate.

21

22 An annular photocatalytic reactor, assimilated to a plug flow reactor, with a fixed  
23 bed of Degussa P25 immobilised onto a fibre glass support was reported for the  
24 remediation of gaseous acetone <sup>26</sup> (figure 9). There was no limitation on mass

1 transfer observed either internally or externally under the experimental  
2 conditions investigated.

### 3 **FIGURE 9**

4 An internally illuminated monolith reactor (IIMR) has been reported in the  
5 investigation of multi-phase photocatalysis<sup>90, 91</sup>. The IIMR had side light emitting  
6 fibres incorporated within the channels of a ceramic monolith containing a TiO<sub>2</sub>  
7 photocatalyst coated individual channel walls. Photonic efficiencies obtained with  
8 this reactor were below those obtained for a slurry reactor but greater than that  
9 reported for an annular photoreactor and a reactor configuration with side light  
10 fibres immersed in a TiO<sub>2</sub> slurry. The authors reported that the IIMR had a larger  
11 area of catalyst exposed to illumination and this is its key advantage over other  
12 photocatalytic reactors.

13

### 14 **Immobilised Gas Reactors**

15 The use of the Optical Fibre Reactor (OFR) for the photoreduction of CO<sub>2</sub> to fuels  
16 by a visible light activated catalyst has been reported by Nguyen and Wu<sup>92</sup>. The  
17 optical fibres were coated with a gel-derived TiO<sub>2</sub>-SiO<sub>2</sub> mixed photocatalyst. The  
18 OFR reactor operated on the principle of incident light being split in two beams  
19 when first in contact with the surface of the fibre. Part of the light penetrated the  
20 layer of catalyst on the fibre and creates excitation, while the other beam shall  
21 reflected off the fibre and transmit along the length of the optical fibre. This  
22 allowed the light to gradually spread through the length of the reactor. Two  
23 photocatalysts were utilised in this investigation; Cu-Fe/TiO<sub>2</sub> along with Cu-  
24 Fe/TiO<sub>2</sub>-SiO<sub>2</sub>. The products obtained in this study included ethylene and

1 methane, along with traces of ethane and methanol. The product formation  
2 under UVA illumination demonstrated selectivity towards the photocatalyst used,  
3 with the exception of methane which was evolved with each of the  
4 photocatalysts. The highest rate of methane production, 1.860  $\mu\text{mol/g-cat h}$  was  
5 found when the catalyst Cu(0.5 wt.%) -Fe(0.5 wt.%) /TiO<sub>2</sub>-SiO<sub>2</sub>-acetyl acetone  
6 (acac) on optical fibres was used. Ethylene evolution was selective in production  
7 and was only seen over Fe and Cu containing catalysts; the highest production  
8 rate of 0.575  $\mu\text{mol/g-cat h}$  was found when the catalyst Cu(0.5 wt.%) -Fe(0.5  
9 wt.%) /TiO<sub>2</sub> on optical fibres was used. In comparison natural sunlight (from a  
10 solar concentrator) produced a production rate of 0.279  $\mu\text{mol/g-cat h}$  with the  
11 catalyst Cu(0.5 wt.%) -Fe(0.5 wt.%) /TiO<sub>2</sub>-SiO<sub>2</sub>-acac on optical fibres,  
12 significantly higher than the rate for the TiO<sub>2</sub>-SiO<sub>2</sub> catalyst. These results  
13 achieved were attributed to the increased surface area from the use of TiO<sub>2</sub>-SiO<sub>2</sub>  
14 in comparison to pure TiO<sub>2</sub> and the presence of Cu and Fe metals to shifted the  
15 light absorption into the visible spectrum. The benefits of this system included  
16 uniform light distribution throughout the reactor, a feature not seen in traditional  
17 packed bed designs and the visible light driven catalyst which the authors  
18 concluded enhanced the applicability as a commercial and industrial application.

19

## 20 **FIGURE 10**

### 21 **Immobilised Vapour Reactors**

22 The development of a cost effective and ease-of-use catalyst support for an  
23 immobilised system was investigated by Haijiesmaili *et al*<sup>93</sup>. This investigation  
24 focused on TiO<sub>2</sub> being supported upon a 3-dimensional carbon foam for the  
25 oxidation of gaseous methanol in a vapour phase flow-through photoreactor. An

1 impregnation technique was adopted for the production of the TiO<sub>2</sub> supported  
2 carbon foam; carbon foam was immersed into a TiO<sub>2</sub> (P25)-containing  
3 water/ethanol (1:1) solution followed by drying. The photoreactor was  
4 established by packing the carbon foam supported TiO<sub>2</sub> into a Pyrex tube with  
5 internal illumination from an 8 W central UV-A light. Results were based upon  
6 methanol conversion and CO<sub>2</sub> and formaldehyde selectivity. The methanol  
7 conversion with the carbon foam supported TiO<sub>2</sub> was dependent on TiO<sub>2</sub> loading  
8 where TiO<sub>2</sub> loadings of 7.7, 9.4, 16.3 and 28.5 wt% achieved methanol  
9 conversions of 52, 57, 61 and 75 % respectively. A maximum methanol  
10 conversion of 81% was achieved for a 666 wt% TiO<sub>2</sub> loading. These results were  
11 impressive compared to those obtained in the photoreactor where the TiO<sub>2</sub>  
12 photocatalyst was simply coated on the inside of the reactor wall. In this case  
13 the highest methanol conversion reached was 22 %, where after any increase in  
14 TiO<sub>2</sub> loading resulted in a screening effect of excess particles and resulted in no  
15 further increase in methanol conversion. CO<sub>2</sub> and formaldehyde selectivity  
16 further supported the findings in the paper with CO<sub>2</sub> reaching 44 % compared to  
17 7 % wall coated reactor. The efficiency of the carbon foam supported TiO<sub>2</sub>  
18 photoreactor was attributed to the ability to increase the exposed surface of the  
19 carbon foam and hence increase the surface to reactor ratio which allowed for  
20 increasing the TiO<sub>2</sub> content within the reactor. The authors concluded the  
21 reactors air to surface ratio and ability to perform at very low pressure drops  
22 allow its use in practical applications.

23

24 **Suspension vs. Immobilised Reactors: A Comparison**

1 Dutta and Ray<sup>28</sup> developed a Taylor vortex photocatalytic reactor that created  
2 unsteady Taylor-Couette flow between two co-axial cylinders through re-  
3 circulation of fluids from the body of the reactor to the inner cylinder wall, coated  
4 with TiO<sub>2</sub>, (figure 11 a and b). The Taylor-Couette flow is the movement of  
5 viscous fluids in between a pair of coaxial cylinders which experience inner  
6 centrifugal instability when the inner cylinder rotates differentially with respect  
7 to the outer cylinder. They investigated the effect of the Reynolds number  
8 (figure 11 c) and catalyst loading on the photodegradation and compared the  
9 results with that of a slurry reactor. They noted that increasing the Reynolds  
10 number increased the rate of photodegradation of their model pollutant orange  
11 II demonstrating that external mass transfer controlled the overall reaction rate,  
12 figure 12.

### 13 **FIGURE 11**

14 The performance of a suspended catalyst system, an immobilised catalyst where  
15 the wall of the reactor was coated and an immobilised system packed with  
16 coated glass beads has been compared for the photodegradation of formic  
17 acid<sup>94</sup>. Mass transfer limitations were observed in the immobilised system with  
18 the catalyst coated on the reactor wall. However aerating the system overcame  
19 this mass transfer problem. The performance of the packed bed reactor was  
20 investigated for two different sized beads (d= 1.3 and 2 mm) to gain an  
21 understanding of the photocatalytic activity. The authors concluded that large  
22 beads enhanced the photocatalytic activity.

23

24 A pilot reactor utilising a TiO<sub>2</sub> coated 15 pores-per-inch alumina reticulated foam  
25 monolith incorporated in the space between a centrally deployed UV lamp and



1 the internal wall of the reactor has been compared with a Degussa P25 TiO<sub>2</sub>  
2 slurry system<sup>95</sup>. Results for the degradation of 1,8-diazobicyclo[5.4.0]undec-7-  
3 ene (DBU) indicated that the foam monolith immobilised photocatalyst system  
4 was more efficient compared to the slurry photocatalyst reactor suggesting that  
5 the immobilised system could be scaled up for water purification.

6

7 Three different reactor configurations, slurry reactor, wall reactor and fixed-bed  
8 reactor, were compared for the photocatalytic disinfection of *Escherichia coli*  
9 aqueous suspensions and methylene blue photodegradation (figure 12). Titania  
10 was in suspension for the slurry reactor, was immobilised on the reactor wall for  
11 the wall reactor and was immobilised on the packing in the fixed bed reactor<sup>96</sup>.  
12 The authors investigated the effect of increasing catalyst layer thickness and  
13 compared it to increasing concentration of catalyst in the slurry system. The  
14 results for methylene blue photo-oxidation were in good agreement for both  
15 slurry and immobilised system. For the photocatalytic disinfection, however, this  
16 was not the case. The increased density of TiO<sub>2</sub> film caused by the heat  
17 treatment reduced the surface area of catalyst available for the micro-organisms  
18 and therefore reduced the photocatalytic activity was observed. When the  
19 immobilised reactor was investigated for effluents sampled from a wastewater  
20 treatment plant, however, they required comparable irradiation times to the  
21 slurry system to reach the bacterial detection limit.

22 **FIGURE 12**

23

24 **Industrial Applications**

1 The potential industrial applications for semiconductor photocatalysis are wide  
2 and diverse ranging from treating oil and gas effluent to potable water. The  
3 reality of research based photoreactor designs are, however, that very few  
4 laboratory scale test reactors are ultimately feasible in terms of industrial scale  
5 up. In an industrial environment the volume and rate of waste effluent  
6 production is in the order of hundreds of cubic meters i.e. millions of litres per  
7 day. Typically laboratory photoreactors have a capacity of between 1 ml and 1  
8 L, with a UV illumination source between 36 W and 500 W. Transforming a  
9 even 1 L capacity reactor to a 1 m<sup>3</sup> capacity unit is not a simple transformation,  
10 with the relationship between materials, volume, catalyst loading, turbidity and  
11 UV penetration presenting complex challenges.

12 Examples where industrial scale photoreactors have been employed on a large  
13 scale are those of a solar activated designs. Solar photoreactors have the  
14 advantage of not requiring artificial light sources but do require huge amounts of  
15 space and are also depend on the solar insolation<sup>97-99</sup>.

16

## 17 **Pilot Scale Studies**

18 Imoberdorf *et al*<sup>100</sup> propose a scaled up multi-annular photocatalytic reactor for  
19 the remediation of air pollution. This consists of four concentric cylindrical  
20 borosilicate glass tubes (figure 13). The illumination source, Philips TL18W UV  
21 lamp, was placed on the central axis of the system. With the available reaction  
22 length of 177 cm the total surface area available for radical production was 5209  
23 cm<sup>2</sup>, a significant increase compared to the 81 cm<sup>2</sup> of the laboratory test reactor.  
24 Using a sol gel process a thin layer of TiO<sub>2</sub> was coated on two walls of the

1 reactor which were in contact with the gas flowing through each annulus of the  
2 unit.

### 3 **FIGURE 13**

4 In this study Imoberdorf *et al.* <sup>100</sup> examined the mechanisms of radiative  
5 transfer, rate kinetics and mass transfer. They proposed that the process  
6 should be free from mass transfer limitations with a reactor operated under the  
7 kinetic control. It was found that it was possible to make accurate predictions of  
8 reactor behaviour, based purely on the chemical reaction fundamentals, reactor  
9 engineering and radiation transport theory. This took no account of the  
10 adjustable or unknown parameters of photoreactor design.

11 Shu and Chang <sup>101</sup> have reported the investigation of a pilot scale thin gap  
12 annular plug flow (TGAPF) and photoreactor and recirculated batch reactor for  
13 the degradation of azo dye waste water using H<sub>2</sub>O<sub>2</sub> instead of a semiconductor  
14 photocatalyst. The TGAPF system was tested using acid orange 10 at a  
15 concentration of 20 mg/L<sup>-1</sup> with a simulated waste water prepared in a tank  
16 reservoir at a volume of 100 L. The TGAPF reactor having a capacity of 3000 ml  
17 and the dye was pumped through the TGAPF reactor at a rate of 1.5 L/min<sup>-1</sup> to  
18 6.5 L/min<sup>-1</sup>, which gave a through put of 2.35 – 9.32 m<sup>3</sup> day<sup>-1</sup>. The UV lamp  
19 source was a 5000 W medium pressure mercury at 253.7 nm and was positioned  
20 centrally in the quartz housing. This configuration allowed a azo dye  
21 degradation rate of 0.26 min per L of dye to 99% of original concentration, e.g.  
22 100 L in 26.9 min. When compared to the degradation rate of the recirculated  
23 batch reactor the TGAPF reactor was 233 times more efficient, with the batch  
24 reactor taking 6267 min to degrade the same volume and concentration of azo  
25 dye.

1

2 These two examples of scaled up pilot photoreactors show that there is the  
3 potential for increasing capacity, although it is worth noting that the Shu and  
4 Chang TGAPF reactor does require a 5000 W UV source which is both expensive  
5 and has significant associated operating costs.

6

7 The double skin sheet reactor (DSSR) comprises a flat transparent box  
8 framework constructed from PLEXIGLAS<sup>®</sup> <sup>98, 102</sup>(figure 14). The photocatalyst  
9 was deployed as a suspension in the waste water and the slurry was then  
10 pumped through the channels of the unit. After the degradation process the  
11 photocatalyst had to be removed from the suspension either by filtering or by  
12 sedimentation. The DSSR has been demonstrated to utilize both direct and the  
13 diffuse portion of solar radiation. A pilot scale DSSR, has been investigated for  
14 the treatment of industrial wastewater effluent in Wolfsburg, Germany<sup>99</sup>. 50 %  
15 of the organic pollutants in the waste water were degraded within between eight  
16 and eleven hours irradiation. The efficiency of the photocatalytic process was  
17 found to be dependent on the initial pollutant concentration, the time of  
18 illumination, and, not surprisingly, the solar UV light flux density.

19 **FIGURE 14**

20 **Conclusions**

21 The research detailed in this review highlight the diversity in photocatalytic  
22 reactor design along with their potential applications. Suspended Liquid  
23 Reactors, Immobilised Liquid Reactors, Immobilised Gas Reactors and

1 Immobilised Vapour were considered and, where appropriate, compared in an  
2 attempt to address the advantages and disadvantages of individual designs.

3 The following conclusions could be drawn from considering the current state of  
4 the art in this field;

5 1. Slurry and suspended systems offer the advantage of increased surface  
6 area allowing increased photocatalyst and reactant interaction and has  
7 proved effective in the treatment of wastewater. Limited light penetration  
8 and downstream processing procedures, particularly with respect to  
9 catalyst separation, however, restrict these concepts for  
10 commercialisation and scale up.

11

12 2. Fluidised bed reactors also present excellent photocatalyst surface area to  
13 pollutant ratios for photocatalytic transformations. Research has  
14 demonstrated that these systems have been effective for both gas and  
15 liquid phase photocatalysis. This has enabled the use of highly efficient  
16 powders such as Degussa P25 which benefit from the 'periodic light  
17 phenomenon' created by fluidised systems. Drawbacks to the system  
18 include the loss or 'drifting' of particles within the system and downstream  
19 processing restrictions.

20

21 3. Fixed bed designs utilise immobilised catalysts which have the advantages  
22 of no downstream processing restrictions such as separation and filtration  
23 and allow operation on both a batch and continuous flow phase. There are  
24 restrictions regarding this system including difficulty in the illumination of  
25 the entire support containing the catalyst and mass transfer limitations

1 affected by catalyst thickness. Furthermore in order to achieve and  
2 effective surface area of photocatalyst relative to the effluent being  
3 treated, scaled up units require a significant "footprint".

4  
5 4. Comparisons of the systems demonstrate that given the correct  
6 parameters the individual concepts are effective. While many  
7 comparisons of systems describe the immobilised catalyst reactor set-up  
8 as having decreased efficiency or restrictions due to mass transfer  
9 limitations, it was reported that if such restrictions were overcome results  
10 were comparable to that slurry and fluidisation systems.

11  
12 As has been demonstrated here, a vast array of photoreactor concepts have  
13 been reported, all displaying varying engineering characteristics in terms of  
14 efficiency for pollutant transport, photocatalyst deployment and activation.  
15 These characteristics are all critical however the future of photoreactor  
16 technology does not solely rely in the design of the reactor itself, but in the  
17 development of more effective photocatalysts, particularly in rate limited  
18 systems. The development of photocatalysts that can achieve greater  
19 conversion efficiencies at lower irradiation energies, and ultimately visible light  
20 absorbing materials will be a critical component in ensuring wide scale adoption  
21 of this versatile technology. For industrial applications photoreactors need to  
22 meet the challenge of capacity, ruggedness, reliability and ease of use.  
23 Currently the only design of photoreactor which is capable of processing the  
24 level of waste water required is that of suspension reactors, but as noted in this  
25 review they are far from infallible. Ultimately however, the application of

- 1 semiconductor photocatalysis for remediation is has real scope for impacting on
- 2 water pollution and hence global water scarcity.
- 3

## 1 References.

2

- 3 1. Ollis DF, Al-Ekhabi, H., Photocatalytic purification and treatment of water  
4 and air. *Elsevier* (1993).
- 5 2. Mills A and Le Hunte S, An overview of semiconductor photocatalysis.  
6 *Journal of Photochemistry and Photobiology A: Chemistry* **108**: 1-35  
7 (1997).
- 8 3. Hoffman AJ, Carraway ER and Hoffmann MR, Photocatalytic Production of  
9 H<sub>2</sub>O<sub>2</sub> and Organic Peroxides on Quantum-Sized Semiconductor Colloids.  
10 *Environmental Science & Technology* **28**: 776-785 (1994).
- 11 4. Fox MA and Dulay MT, Heterogeneous photocatalysis. *Chemical Reviews*  
12 **93**: 341-357 (1993).
- 13 5. Fujishima A, Rao TN and Tryk DA, Titanium dioxide photocatalysis. *Journal*  
14 *of Photochemistry and Photobiology C: Photochemistry Reviews* **1**: 1-21  
15 (2000).
- 16 6. Ireland JC, Klostermann P, Rice EW and R M Clark RM, Inactivation of  
17 *Escherichia coli* by titanium dioxide photocatalytic oxidation. *Applied*  
18 *Environmental Microbiology* **59**: 1668-1670 (1993).
- 19 7. McCullagh C, Robertson J, Bahnemann D and Robertson P, The application  
20 of TiO<sub>2</sub> photocatalysis for disinfection of water contaminated with  
21 pathogenic micro-organisms: a review. *Research on Chemical*  
22 *Intermediates* **33**: 359-375 (2007).
- 23 8. Sjogren JC, and Sierka, R.A., Inactivation of phage ms2 by iron-aided  
24 titanium dioxide photocatalysis. *Applied and Environmental Microbiology*  
25 **60**: 344-347 (1994).
- 26 9. Cai R, Hashimoto, K. Kubota, Y. and Fujishima, A. , Increment of  
27 Photocatalytic Killing of Cancer Cells Using TiO<sub>2</sub> with the Aid of  
28 Superoxide Dismutase. *Chemistry Lett* (1992).
- 29 10. Cai R, Kuboata, Y., Shuin, T., Hashimoto, K., and Fujishima, A., Induction  
30 of Cytotoxicity by Photo-excited TiO<sub>2</sub> Particles. *Cancer Res*: 2346-2348  
31 (1992).
- 32 11. Fujishima A and Honda K, Electrochemical Photolysis of Water at a  
33 Semiconductor Electrode. *Nature* **238**: 37-38 (1972).
- 34 12. Karakitsou KE and Verykios XE, Effects of altrivalent cation doping of  
35 titania on its performance as a photocatalyst for water cleavage. *J Phys*  
36 *Chem* **97**: 1184-1189. (1993).
- 37 13. Suzuki in, Photocatalytic purification and treatment of water and air. *Eds*  
38 *Ollis, David F, Al-Ekhabi, H Elsevier* (1993).
- 39 14. Borgarello E, Kiwi J, Pelizzetti E, Visca M and Gratzel M, Photochemical  
40 cleavage of water by photocatalysis. *Nature* **289**: 158-160 (1981).
- 41 15. Taqui Khan MM and Nageswara Rao N, Stepwise reduction of coordinated  
42 dinitrogen to ammonia via diazinido and hydrazido intermediates on a  
43 visible light irradiated Pt/CdS · Ag<sub>2</sub>S/RuO<sub>2</sub> particulate system suspended  
44 in an aqueous solution of K[Ru(EDTA-H)Cl]2H<sub>2</sub>O. *Journal of*  
45 *Photochemistry and Photobiology A: Chemistry* **56**: 101-111 (1991).
- 46 16. Schiavello M, Some working principles of heterogeneous photocatalysis by  
47 semiconductors. *Electrochimica Acta* **38**: 11-14 (1993).
- 48 17. Taqui Khan MM, Chatterjee D and Bala M, Photocatalytic reduction of N<sub>2</sub>  
49 to NH<sub>3</sub> sensitized by the [Ru(III)-ethylenediaminetetraacetate-2,2'-



- 1 bipyridyl]- complex in a Pt-TiO<sub>2</sub> semiconductor particulate system. *Journal*  
2 *of Photochemistry and Photobiology A: Chemistry* **67**: 349-352 (1992).
- 3 18. Khan MMT, Chatterjee D, Krishnaratnam M and Bala M, Photosensitized  
4 reduction of N<sub>2</sub> by RuII(bipy)<sub>3</sub><sup>2+</sup> adsorbed on the surface of Pt/TiO<sub>2</sub>/RuO<sub>2</sub>  
5 semiconductor particulate system containing RuIII-EDTA complex and L-  
6 ascorbic acid. *Journal of Molecular Catalysis* **72**: 13-18 (1992).
- 7 19. Gerischer H, Heller, A. , Photocatalytic oxidation of organic molecules at  
8 titanium dioxide particles by sunlight in aerated water. *J Electrochem Soc*  
9 **139** (1992).
- 10 20. Jackson NB, Wang, C. M., Luo, Z., Schwitzgebel, J., Ekerdt, J. G., Brock,  
11 J. R., Heller, A., Attachment of TiO<sub>2</sub> Powders to Hollow Glass Microbeads:  
12 Activity of the TiO<sub>2</sub>-Coated Beads in the Photoassisted Oxidation of  
13 Ethanol to Acetaldehyde. *J Electrochem Soc* **138**: 3660-3664 (1992).
- 14 21. Nair M, Luo Z and Heller A, Rates of photocatalytic oxidation of crude oil  
15 on salt water on buoyant, cenosphere-attached titanium dioxide.  
16 *Industrial & Engineering Chemistry Research* **32**: 2318-2323 (1993).
- 17 22. Boer KW, Survey of semiconductor physics. *Van Nostrand, New York*:  
18 1472 (1990).
- 19 23. Kormann C, Bahnemann DW and Hoffmann MR, Photocatalytic production  
20 of hydrogen peroxides and organic peroxides in aqueous suspensions of  
21 titanium dioxide, zinc oxide, and desert sand. *Environmental Science &*  
22 *Technology* **22**: 798-806 (1988).
- 23 24. Carraway ER, Hoffman AJ and Hoffmann MR, Photocatalytic Oxidation of  
24 Organic Acids on Quantum-Sized Semiconductor Colloids. *Environmental*  
25 *Science & Technology* **28**: 786-793 (1994).
- 26 25. Mohseni M, Gas phase trichloroethylene (TCE) photooxidation and  
27 byproduct formation: photolysis vs. titania/silica based photocatalysis.  
28 *Chemosphere* **59**: 335-342 (2005).
- 29 26. Vincent G, Marquaire PM and Zahraa O, Abatement of volatile organic  
30 compounds using an annular photocatalytic reactor: Study of gaseous  
31 acetone. *Journal of Photochemistry and Photobiology A: Chemistry* **197**:  
32 177-189 (2008).
- 33 27. Mehrvar M, Anderson WA and Moo-Young M, Preliminary analysis of a  
34 tellerette packed-bed photocatalytic reactor. *Advances in Environmental*  
35 *Research* **6**: 411-418 (2002).
- 36 28. Dutta PK and Ray AK, Experimental investigation of Taylor vortex  
37 photocatalytic reactor for water purification. *Chemical Engineering Science*  
38 **59**: 5249-5259 (2004).
- 39 29. Sczechowski JG, Koval CA and Noble RD, A Taylor vortex reactor for  
40 heterogeneous photocatalysis. *Chemical Engineering Science* **50**: 3163-  
41 3173 (1995).
- 42 30. Giordano RC, Giordano RLC, Prazeres DMF and Cooney CL, Analysis of a  
43 Taylor-Poiseuille vortex flow reactor--I: Flow patterns and mass transfer  
44 characteristics. *Chemical Engineering Science* **53**: 3635-3652 (1998).
- 45 31. Richter O, Hoffmann H and Kraushaar-Czarnetzki B, Effect of the rotor  
46 shape on the mixing characteristics of a continuous flow Taylor-vortex  
47 reactor. *Chemical Engineering Science* **63**: 3504-3513 (2008).
- 48 32. Kuo HP, Wu CT and Hsu RC, Continuous reduction of toluene vapours from  
49 the contaminated gas stream in a fluidised bed photoreactor. *Powder*  
50 *Technology* **195**: 50-56 (2009).

- 1 33. Lee D-K, Kim S-C, Cho I-C, Kim S-J and Kim S-W, Photocatalytic oxidation of microcystin-LR in a fluidized bed reactor having TiO<sub>2</sub>-coated activated carbon. *Separation and Purification Technology* **34**: 59-66 (2004).
- 2  
3  
4 34. Peill NJ and Hoffmann MR, Development and Optimization of a TiO<sub>2</sub>-Coated Fiber-Optic Cable Reactor: Photocatalytic Degradation of 4-Chlorophenol. *Environmental Science & Technology* **29**: 2974-2981 (1995).
- 5  
6  
7  
8 35. Li Puma G and Yue PL, Comparison of the Effectiveness of Photon-Based Oxidation Processes in a Pilot Falling Film Photoreactor. *Environmental Science & Technology* **33**: 3210-3216 (1999).
- 9  
10  
11 36. D. Bockelmann RG, D. Weichgrebe, and D. Bahnemann, Solar detoxification of polluted water: comparing the efficiencies of a parabolic through reactor and a novel thin-film fixed-bed reactor. *In Photocatalytic Purification and Treatment of Water and Air, D F Ollis and H Al-Ekabi, Eds, Elsevier Science, Amsterdam, The Netherlands pp. 771-776 (1993).*
- 12  
13  
14  
15  
16 37. Sahle-Demessie E, Bekele S and Pillai UR, Residence time distribution of fluids in stirred annular photoreactor. *Catalysis Today* **88**: 61-72 (2003).
- 17  
18 38. Zhang Z, Anderson WA and Moo-Young M, Experimental analysis of a corrugated plate photocatalytic reactor. *Chemical Engineering Journal* **99**: 145-152 (2004).
- 19  
20  
21 39. Dijkstra MFJ, Michorius A, Buwalda H, Panneman HJ, Winkelman JGM and Beenackers AACM, Comparison of the efficiency of immobilized and suspended systems in photocatalytic degradation. *Catalysis Today* **66**: 487-494 (2001).
- 22  
23  
24  
25 40. Chong MN, Jin B, Zhu HY, Chow CWK and Saint C, Application of H-titanate nanofibers for degradation of Congo Red in an annular slurry photoreactor. *Chemical Engineering Journal* **150**: 49-54 (2009).
- 26  
27  
28 41. Augugliaro V, Loddo V, Palmisano L and Schiavello M, Performance of Heterogeneous Photocatalytic Systems: Influence of Operational Variables on Photoactivity of Aqueous Suspension of TiO<sub>2</sub>. *Journal of Catalysis* **153**: 32-40 (1995).
- 29  
30  
31  
32 42. Minero C and Vione D, A quantitative evaluation of the photocatalytic performance of TiO<sub>2</sub> slurries. *Applied Catalysis B: Environmental* **67**: 257-269 (2006).
- 33  
34  
35 43. Ollis DF, Pelizzetti, E., Serpone, N., Photocatalyzed destruction of water contaminants. *Environmental Science & Technology* **25**: 1522-1529 (1991).
- 36  
37  
38 44. Ray AK and Beenackers AACM, Novel swirl-flow reactor for kinetic studies of semiconductor photocatalysis. *AIChE Journal* **43**: 2571-2578 (1997).
- 39  
40 45. Chen D, Li F and Ray AK, External and internal mass transfer effect on photocatalytic degradation. *Catalysis Today* **66**: 475-485 (2001).
- 41  
42 46. Pruden AL and Ollis DF, Degradation of chloroform by photoassisted heterogeneous catalysis in dilute aqueous suspensions of titanium dioxide. *Environmental Science & Technology* **17**: 628-631 (1983).
- 43  
44  
45 47. Kormann C, Bahnemann DW and Hoffmann MR, Photolysis of chloroform and other organic molecules in aqueous titanium dioxide suspensions. *Environmental Science & Technology* **25**: 494-500 (1991).
- 46  
47  
48 48. Pramauro E, Vincenti M, Augugliaro V and Palmisano L, Photocatalytic degradation of Monuron in aqueous titanium dioxide dispersions. *Environmental Science & Technology* **27**: 1790-1795 (1993).
- 49  
50

- 1 49. Pathirana HMKK and Maithreepala RA, Photodegradation of 3,4-  
2 dichloropropionamide in aqueous TiO<sub>2</sub> suspensions. *Journal of*  
3 *Photochemistry and Photobiology A: Chemistry* **102**: 273-277 (1997).
- 4 50. Li Puma G and Yue PL, Photocatalytic Oxidation of Chlorophenols in  
5 Single-Component and Multicomponent Systems. *Industrial & Engineering*  
6 *Chemistry Research* **38**: 3238-3245 (1999).
- 7 51. Li Puma G and Yue PL, Enhanced Photocatalysis in a Pilot Laminar Falling  
8 Film Slurry Reactor. *Industrial & Engineering Chemistry Research* **38**:  
9 3246-3254 (1999).
- 10 52. San N, Hatipoglu A, Koçtürk G and Çİnar Z, Prediction of primary  
11 intermediates and the photodegradation kinetics of 3-aminophenol in  
12 aqueous TiO<sub>2</sub> suspensions. *Journal of Photochemistry and Photobiology A:*  
13 *Chemistry* **139**: 225-232 (2001).
- 14 53. Jeon J, Kim S, Lim T and Lee D, Degradation of trichloroethylene by  
15 photocatalysis in an internally circulating slurry bubble column reactor.  
16 *Chemosphere* **60**: 1162-1168 (2005).
- 17 54. Sauer T, Cesconeto Neto G, José HJ and Moreira RFPM, Kinetics of  
18 photocatalytic degradation of reactive dyes in a TiO<sub>2</sub> slurry reactor.  
19 *Journal of Photochemistry and Photobiology A: Chemistry* **149**: 147-154  
20 (2002).
- 21 55. Kamble SP, Sawant SB and Pangarkar VG, Photocatalytic Mineralization of  
22 Phenoxyacetic Acid Using Concentrated Solar Radiation and Titanium  
23 Dioxide in Slurry Photoreactor. *Chemical Engineering Research and Design*  
24 **84**: 355-362 (2006).
- 25 56. Adams M, Campbell, I., Robertson, P.K.J., Novel Photocatalytic Reactor  
26 Development for Removal of Hydrocarbons from Water. *International*  
27 *Journal of Photoenergy* **Volume 2008** 7 pages (2008).
- 28 57. Author, Apparatus and method for treating a fluid by means of a  
29 transparent container (2006).
- 30 58. McCullagh C, Robertson PKJ, Adams M, Pollard PM and Mohammed A,  
31 Development of a slurry continuous flow reactor for photocatalytic  
32 treatment of industrial waste water. *Journal of Photochemistry and*  
33 *Photobiology A: Chemistry* **211**: 42-46 (2010).
- 34 59. Salu OA, Adams, M., Robertson, P.K.J., Wong, L.S., and McCullagh, C.,  
35 Remediation of oily wastewater from an interceptor tank using a novel  
36 photocatalytic drum reactor  
37 *Desalination and Water Treatment In Press* (2010).
- 38 60. Chen D and Ray AK, Photocatalytic kinetics of phenol and its derivatives  
39 over UV irradiated TiO<sub>2</sub>. *Applied Catalysis B: Environmental* **23**: 143-157  
40 (1999).
- 41 61. Martin ST, Herrmann, Hartmut, Choi, Wonyong, Hoffmann, Michael R.,  
42 Time-resolved microwave conductivity. Part 1.-TiO<sub>2</sub> photoreactivity and  
43 size quantization. *Journal of the Chemical Society, Faraday Transactions*  
44 **90**: 3315-3322 (1994).
- 45 62. Mehrotra K, Yablonsky GS and Ray AK, Kinetic Studies of Photocatalytic  
46 Degradation in a TiO<sub>2</sub> Slurry System: Distinguishing Working  
47 Regimes and Determining Rate Dependences. *Industrial & Engineering*  
48 *Chemistry Research* **42**: 2273-2281 (2003).
- 49 63. Moreira R, Sauer T, Casaril L and Humeres E, Mass transfer and  
50 photocatalytic degradation of leather dye using TiO<sub>2</sub>/UV. *Journal of Applied*  
51 *Electrochemistry* **35**: 821-829 (2005).

- 1 64. Ballari MdIM, Alfano OM and Cassano AE, Mass transfer limitations in  
2 slurry photocatalytic reactors: Experimental validation. *Chemical*  
3 *Engineering Science* **65**: 4931-4942 (2010).
- 4 65. Paz Y, Application of TiO<sub>2</sub> photocatalysis for air treatment: Patents'  
5 overview. *Applied Catalysis B: Environmental* **99**: 448-460 (2010).
- 6 66. Dibble LA and Raupp GB, Fluidized-bed photocatalytic oxidation of  
7 trichloroethylene in contaminated air streams. *Environmental Science &*  
8 *Technology* **26**: 492-495 (1992).
- 9 67. Lim TH and Kim SD, Trichloroethylene degradation by photocatalysis in  
10 annular flow and annulus fluidized bed photoreactors. *Chemosphere* **54**:  
11 305-312 (2004).
- 12 68. Lim TH and Kim SD, Photocatalytic degradation of trichloroethylene (TCE)  
13 over TiO<sub>2</sub>/silica gel in a circulating fluidized bed (CFB) photoreactor.  
14 *Chemical Engineering and Processing* **44**: 327-334 (2005).
- 15 69. Coates NH and Rice RL, Sulfur dioxide reduction by combustion of coal in  
16 fluidized bed of limestone. *AIChE Symp Ser* **70** (1974).
- 17 70. Son Y-H, Jeon, M-K., Ban, J-Y., Kang, M and Choung, S-J, Application of  
18 three-phase fluidized photocatalytic system to decompositions of acetic  
19 acid and ammonia. *J Ind Eng Chem* **11**: 938-944 (2005).
- 20 71. Kang M, Synthesis of Fe/TiO<sub>2</sub> photocatalyst with nanometer size by  
21 solvothermal method and the effect of H<sub>2</sub>O addition on structural stability  
22 and photodecomposition of methanol. *Journal of Molecular Catalysis A:*  
23 *Chemical* **197**: 173-183 (2003).
- 24 72. Nelson RJ, Flakker CL and Muggli DS, Photocatalytic oxidation of methanol  
25 using titania-based fluidized beds. *Applied Catalysis B: Environmental* **69**:  
26 189-195 (2007).
- 27 73. Yao G-h, Wang F, Wang X-b and Gui K-t, Magnetic field effects on  
28 selective catalytic reduction of NO by NH<sub>3</sub> over Fe<sub>2</sub>O<sub>3</sub> catalyst in a  
29 magnetically fluidized bed. *Energy* **35**: 2295-2300 (2010).
- 30 74. Vorontsov AV, N. Savinov E and Smirniotis PG, Vibrofluidized- and fixed-  
31 bed photocatalytic reactors: case of gaseous acetone photooxidation.  
32 *Chemical Engineering Science* **55**: 5089-5098 (2000).
- 33 75. Lim TH, Jeong SM, Kim SD and Gyenis J, Photocatalytic decomposition of  
34 NO by TiO<sub>2</sub> particles. *Journal of Photochemistry and Photobiology A:*  
35 *Chemistry* **134**: 209-217 (2000).
- 36 76. Foster NR, Bassiti, K., Photocatalytic Reactor. *European Patent -*  
37 *EP2089328 (A1), World Patent - WO2008050119(1998)* (2008).
- 38 77. Al-Ekabi H and Serpone N, Kinetics studies in heterogeneous  
39 photocatalysis. I. Photocatalytic degradation of chlorinated phenols in  
40 aerated aqueous solutions over titania supported on a glass matrix. *The*  
41 *Journal of Physical Chemistry* **92**: 5726-5731 (1988).
- 42 78. Nogueira RFP and Jardim WF, TiO<sub>2</sub>-fixed-bed reactor for water  
43 decontamination using solar light. *Solar Energy* **56**: 471-477 (1996).
- 44 79. Feitz AJ, Boyden BH and Waite TD, Evaluation of two solar pilot scale  
45 fixed-bed photocatalytic reactors. *Water Research* **34**: 3927-3932 (2000).
- 46 80. Dionysiou DD, Balasubramanian G, Suidan MT, Khodadoust AP, Baudin I  
47 and Laine J-M, Rotating disk photocatalytic reactor: development,  
48 characterization, and evaluation for the destruction of organic pollutants in  
49 water. *Water Research* **34**: 2927-2940 (2000).
- 50 81. Dionysiou DD, Khodadoust AP, Kern AM, Suidan MT, Baudin I and Laine J-  
51 M, Continuous-mode photocatalytic degradation of chlorinated phenols

- 1 and pesticides in water using a bench-scale TiO<sub>2</sub> rotating disk reactor.  
2 *Applied Catalysis B: Environmental* **24**: 139-155 (2000).
- 3 82. Hamill NA, Weatherley LR and Hardacre C, Use of a batch rotating  
4 photocatalytic contactor for the degradation of organic pollutants in  
5 wastewater. *Applied Catalysis B: Environmental* **30**: 49-60 (2001).
- 6 83. Dionysiou DD, Suidan MT, Baudin I and Laïné J-M, Oxidation of organic  
7 contaminants in a rotating disk photocatalytic reactor: reaction kinetics in  
8 the liquid phase and the role of mass transfer based on the dimensionless  
9 Damköhler number. *Applied Catalysis B: Environmental* **38**: 1-16 (2002).
- 10 84. Villacres R, Ikeda S, Torimoto T and Ohtani B, Development of a novel  
11 photocatalytic reaction system for oxidative decomposition of volatile  
12 organic compounds in water with enhanced aeration. *Journal of*  
13 *Photochemistry and Photobiology A: Chemistry* **160**: 121-126 (2003).
- 14 85. Zhang L, Kanki T, Sano N and Toyoda A, Development of TiO<sub>2</sub>  
15 photocatalyst reaction for water purification. *Separation and Purification*  
16 *Technology* **31**: 105-110 (2003).
- 17 86. McMurray TA, Byrne JA, Dunlop PSM, Winkelman JGM, Eggins BR and  
18 McAdams ET, Intrinsic kinetics of photocatalytic oxidation of formic and  
19 oxalic acid on immobilised TiO<sub>2</sub> films. *Applied Catalysis A: General* **262**:  
20 105-110 (2004).
- 21 87. Lin HF and Valsaraj KT, A titania thin film annular photocatalytic reactor  
22 for the degradation of polycyclic aromatic hydrocarbons in dilute water  
23 streams. *Journal of Hazardous Materials* **99**: 203-219 (2003).
- 24 88. Mozia S, Tomaszewska M and Morawski AW, Photocatalytic degradation of  
25 azo-dye Acid Red 18. *Desalination* **185**: 449-456 (2005).
- 26 89. Mozia S, Tomaszewska M and Morawski AW, Photodegradation of azo dye  
27 Acid Red 18 in a quartz labyrinth flow reactor with immobilized TiO<sub>2</sub> bed.  
28 *Dyes and Pigments* **75**: 60-66 (2007).
- 29 90. Du P, Carneiro JT, Moulijn JA and Mul G, A novel photocatalytic monolith  
30 reactor for multiphase heterogeneous photocatalysis. *Applied Catalysis A:*  
31 *General* **334**: 119-128 (2008).
- 32 91. Carneiro JT, Berger R, Moulijn JA and Mul G, An internally illuminated  
33 monolith reactor: Pros and cons relative to a slurry reactor. *Catalysis*  
34 *Today* **147**: S324-S329 (2009).
- 35 92. Nguyen T-V and Wu JCS, Photoreduction of CO<sub>2</sub> to fuels under sunlight  
36 using optical-fiber reactor. *Solar Energy Materials and Solar Cells* **92**:  
37 864-872 (2008).
- 38 93. Hajiesmaili S, Josset S, Bégin D, Pham-Huu C, Keller N and Keller V, 3D  
39 solid carbon foam-based photocatalytic materials for vapor phase flow-  
40 through structured photoreactors. *Applied Catalysis A: General* **382**: 122-  
41 130.
- 42 94. Salaices M, Serrano B and de Lasa HI, Experimental evaluation of photon  
43 absorption in an aqueous TiO<sub>2</sub> slurry reactor. *Chemical Engineering*  
44 *Journal* **90**: 219-229 (2002).
- 45 95. Ochuma IJ, Osibo OO, Fishwick RP, Pollington S, Wagland A, Wood J and  
46 Winterbottom JM, Three-phase photocatalysis using suspended titania and  
47 titania supported on a reticulated foam monolith for water purification.  
48 *Catalysis Today* **128**: 100-107 (2007).
- 49 96. van Grieken R, Marugán J, Sordo C and Pablos C, Comparison of the  
50 photocatalytic disinfection of E. coli suspensions in slurry, wall and fixed-  
51 bed reactors. *Catalysis Today* **144**: 48-54 (2009).

- 1 97. Dillert R, Cassano AE, Goslich R and Bahnemann D, Large scale studies in  
2 solar catalytic wastewater treatment. *Catalysis Today* **54**: 267-282  
3 (1999).
- 4 98. Benz V, Müller, M., Bahnemann, D.W., Weichgrebe, D., Brehm, M.,  
5 Reaktoren für die photokatalytische Abwasserreinigung mit  
6 Stegmehrfachplatten als Solarelemente. *Deutsche Offenlegungsschrift DE*  
7 **195 14 372 A1** (1996).
- 8 99. Dillert R, Vollmer, S., Gross, E., Schober, M., Bahnemann, D., Wienefeld,  
9 J., Pahlmann, K., Schmedding, T., Arntz, H.-J., Sager, G., Solar-catalytic  
10 treatment of an industrial wastewater. *Zeitschrift für Physikalische Chemie*  
11 **213**: 141-147 (1999).
- 12 100. Imoberdorf GE, Cassano AE, Irazoqui HA and Alfano OM, Optimal design  
13 and modeling of annular photocatalytic wall reactors. *Catalysis Today*  
14 **129**: 118-126 (2007).
- 15 101. Shu H-Y and Chang M-C, Pilot scale annular plug flow photoreactor by  
16 UV/H<sub>2</sub>O<sub>2</sub> for the decolorization of azo dye wastewater. *Journal of*  
17 *Hazardous Materials* **125**: 244-251 (2005).
- 18 102. van Well M, Dillert RHG, Bahnemann DW, Benz VW and Mueller MA, A  
19 Novel Nonconcentrating Reactor for Solar Water Detoxification. *Journal of*  
20 *Solar Energy Engineering* **119**: 114-119 (1997).
- 21 103. Mozia S, Morawski AW, Toyoda M and Inagaki M, Effectiveness of  
22 photodecomposition of an azo dye on a novel anatase-phase TiO<sub>2</sub> and two  
23 commercial photocatalysts in a photocatalytic membrane reactor (PMR).  
24 *Separation and Purification Technology* **63**: 386-391 (2008).
- 25 104. Alexiadis A and Mazzarino I, Design guidelines for fixed-bed photocatalytic  
26 reactors. *Chemical Engineering and Processing* **44**: 453-459 (2005).
- 27 105. Cernigoj U, Stangar UL and Trebse P, Evaluation of a novel Carberry type  
28 photoreactor for the degradation of organic pollutants in water. *Journal of*  
29 *Photochemistry and Photobiology A: Chemistry* **188**: 169-176 (2007).
- 30 106. Lo C-C, Hung C-H, Yuan C-S and Wu J-F, Photoreduction of carbon dioxide  
31 with H<sub>2</sub> and H<sub>2</sub>O over TiO<sub>2</sub> and ZrO<sub>2</sub> in a circulated photocatalytic  
32 reactor. *Solar Energy Materials and Solar Cells* **91**: 1765-1774 (2007).
- 33 107. Fu J, Ji M, Zhao Y and Wang L, Kinetics of aqueous photocatalytic  
34 oxidation of fulvic acids in a photocatalysis-ultrafiltration reactor (PUR).  
35 *Separation and Purification Technology* **50**: 107-113 (2006).
- 36 108. Li Puma G and Lock Yue P, The modeling of a fountain photocatalytic  
37 reactor with a parabolic profile. *Chemical Engineering Science* **56**: 721-  
38 726 (2001).
- 39 109. Agarwal A and Bhaskarwar AN, Comparative simulation of falling-film and  
40 parallel-film reactors for photocatalytic production of hydrogen.  
41 *International Journal of Hydrogen Energy* **32**: 2764-2775 (2007).
- 42 110. Ochuma IJ, Fishwick RP, Wood J and Winterbottom JM, Photocatalytic  
43 oxidation of 2,4,6-trichlorophenol in water using a cocurrent downflow  
44 contactor reactor (CDCR). *Journal of Hazardous Materials* **144**: 627-633  
45 (2007).
- 46 111. Chin SS, Lim TM, Chiang K and Fane AG, Hybrid low-pressure submerged  
47 membrane photoreactor for the removal of bisphenol A. *Desalination* **202**:  
48 253-261 (2007).
- 49 112. Orozco SL, Arancibia-Bulnes CA and Suárez-Parra R, Radiation absorption  
50 and degradation of an azo dye in a hybrid photocatalytic reactor. *Chemical*  
51 *Engineering Science* **64**: 2173-2185 (2009).

- 1 113. Erdei L, Arecrachakul N and Vigneswaran S, A combined photocatalytic  
2 slurry reactor-immersed membrane module system for advanced  
3 wastewater treatment. *Separation and Purification Technology* **62**: 382-  
4 388 (2008).
- 5 114. Hao X-g, Li H-h, Zhang Z-l, Fan C-m, Liu S-b and Sun Y-p, Modeling and  
6 experimentation of a novel labyrinth bubble photoreactor for degradation  
7 of organic pollutant. *Chemical Engineering Research and Design* **87**:  
8 1604-1611 (2009).
- 9 115. Pourahmad A, Sohrabnezhad S and Kashefian E, AgBr/nanoAIMCM-41  
10 visible light photocatalyst for degradation of methylene blue dye.  
11 *Spectrochimica Acta Part A: Molecular and Biomolecular Spectroscopy* **77**:  
12 1108-1114.
- 13 116. Shifu C, Wei Z, Wei L, Huaye Z, Xiaoling Y and Yinghao C, Preparation,  
14 characterization and activity evaluation of p-n junction photocatalyst p-  
15 CaFe<sub>2</sub>O<sub>4</sub>/n-Ag<sub>3</sub>VO<sub>4</sub> under visible light irradiation. *Journal of Hazardous*  
16 *Materials* **172**: 1415-1423 (2009).
- 17 117. Liu W, Ji M and Chen S, Preparation, characterization and activity  
18 evaluation of Ag<sub>2</sub>Mo<sub>4</sub>O<sub>13</sub> photocatalyst. *Journal of Hazardous Materials*  
19 **186**: 2001-2008.
- 20 118. Kanade KG, Baeg J-O, Kale BB, Mi Lee S, Moon S-J and Kong K-j, Rose-  
21 red color oxynitride Nb<sub>2</sub>Zr<sub>6</sub>O<sub>17</sub>-xNx: A visible light photocatalyst to  
22 hydrogen production. *International Journal of Hydrogen Energy* **32**: 4678-  
23 4684 (2007).
- 24 119. Jing D, Liu M, Chen Q and Guo L, Efficient photocatalytic hydrogen  
25 production under visible light over a novel W-based ternary chalcogenide  
26 photocatalyst prepared by a hydrothermal process. *International Journal*  
27 *of Hydrogen Energy* **35**: 8521-8527.
- 28 120. Shen S, Zhao L and Guo L, Cetyltrimethylammoniumbromide (CTAB)-  
29 assisted hydrothermal synthesis of ZnIn<sub>2</sub>S<sub>4</sub> as an efficient visible-light-  
30 driven photocatalyst for hydrogen production. *International Journal of*  
31 *Hydrogen Energy* **33**: 4501-4510 (2008).
- 32  
33

Reactor Type	Reactant Phase	Reactor Name	Experimentation	Catalyst Used	Industrial Application/ Comments	Reference
Immobilised	Liquid	Photocatalytic Membrane Reactor (PMR)	Azo dye degradation	Anatase-phase TiO <sub>2</sub>		103
		Fixed Bed	Waste water treatment	TiO <sub>2</sub>		104
		Corrugated plate	4-chlorophenol degradation	TiO <sub>2</sub> (P25)		38
		Rotating Disc	4-Chlorobenzoic acid degradation	TiO <sub>2</sub> (coated commercial ceramic and glass balls)		80
		Carberry photoreactor	4-chlorophenol degradation	TiO <sub>2</sub> (P25) on sodium glass support.		105
		Optical Fibre Reactor (OFR)	Degradation of 4-chlorophenol	TiO <sub>2</sub> on quartz fiber cores	Water treatment	34
	Gas	Annular Wall	PCE degradation	TiO <sub>2</sub>	Air purification	100
		Circulated system	CO <sub>2</sub> reduction	TiO <sub>2</sub> (P25), ZrO <sub>2</sub>		106
		Monolith	Cyclohexane	TiO <sub>2</sub> (Hombikat UV100)	Air purification	90
	Vapour	Carbon foam-based photoreactor	Gaseous methanol oxidation	TiO <sub>2</sub> (P25) supported on carbon foam		93



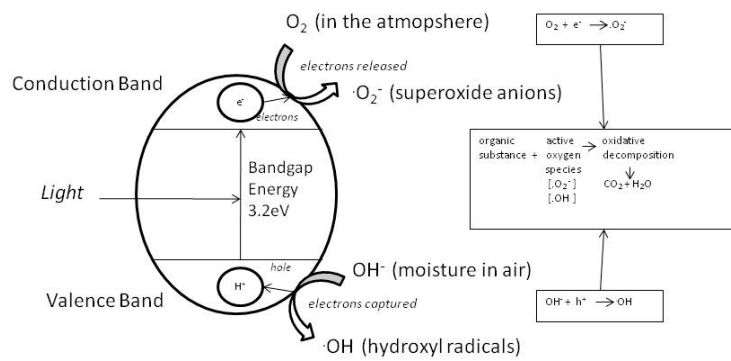
Reactor Type	Reactant phase	Reactor Name	Experimentation	Catalyst Used	Industrial Application/ comments	Reference	
Suspended	Liquid	Photocatalysis-Ultrafiltration Reactor (PUR)	Fulvic Acid	TiO <sub>2</sub> (P25)		<sup>107</sup>	
		Rotating Drum Reactor	Hydrocarbons	TiO <sub>2</sub> (P25)	Waterwater Treatment. Drinking water disinfection.	<sup>56</sup>	
		Taylor Vortex	Formate acid	TiO <sub>2</sub> (P25)		<sup>29</sup>	
		Fountain	Indigo carmine oxidation	TiO <sub>2</sub>		<sup>108</sup>	
		Falling film slurry	Salicylic acid oxidation	TiO <sub>2</sub> (P25)		<sup>51</sup>	
			Hydrogen production	CdS		<sup>109</sup>	
		Internally circulating slurry bubble column reactor	TCE	TiO <sub>2</sub> (P25)		<sup>53</sup>	
		Cocurrent downflow contactor reactor (CDCR)	2,4,6-trichlorophenol (2,3,6-TCP)	TiO <sub>2</sub> (VP Aeroperl P25/20)		<sup>110</sup>	
		Hybrid low-pressure submerged membrane photoreactor	Removal of bisphenol A	TiO <sub>2</sub> (P25)		<sup>111</sup>	
		Hybrid photoreactor	Azo dye – reactive blue 69	TiO <sub>2</sub> (P25)		Wastewater treatment. Uses Solar irradiation	<sup>112</sup>
		Slurry reactor-immersed membrane	Synthetic wastewater	TiO <sub>2</sub> (P25)		Wastewater Treatment	<sup>113</sup>
		Novel labyrinth bubble photocatalytic reactor	Methyl orange degradation	TiO <sub>2</sub> immobilised on quartz glass. (Used in suspension)		Wastewater treatment. Built on a commercial scale.	<sup>114</sup>
		Fluidised bed photoreactor	MC-LR destruction	TiO <sub>2</sub> coated activated carbon			<sup>33</sup>
		Standard slurry photoreactor	Methylene blue degradation	AgBr/nanoAlMCM-41		Visible light activated.	<sup>115</sup>
				p-CaFe <sub>2</sub> O <sub>4</sub> /n-Ag <sub>3</sub> VO <sub>4</sub>		Visible light activated.	<sup>116</sup>

		Methyl orange oxidation	$\text{Ag}_2\text{Mo}_4\text{O}_{13}$		<sup>117</sup>
		Hydrogen production	$\text{Nb}_2\text{Zr}_6\text{O}_{17-x}\text{N}_x$	Visible light activated	<sup>118</sup>
	$\text{Cu}_2\text{WS}_4$		Visible light activated	<sup>119</sup>	
	$\text{ZnIn}_2\text{S}_4$		Visible light activated	<sup>120</sup>	

Table 1: Overview of photoreactor type, reactant phase, experimental targets, catalyst employed and industrial applications.

1 Figure 1:

2



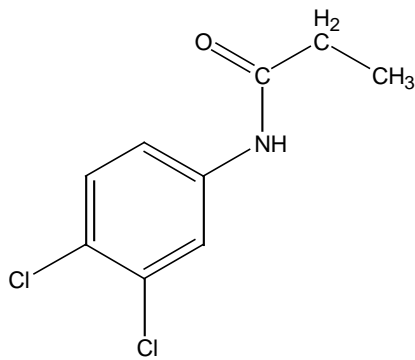
3

4

5 Figure 2:

6

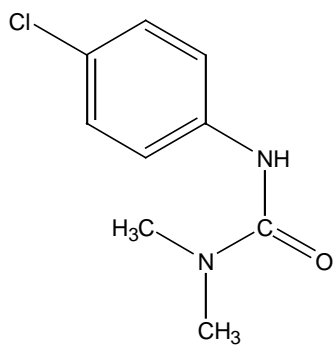
7 (A)



8

9 (B)

10

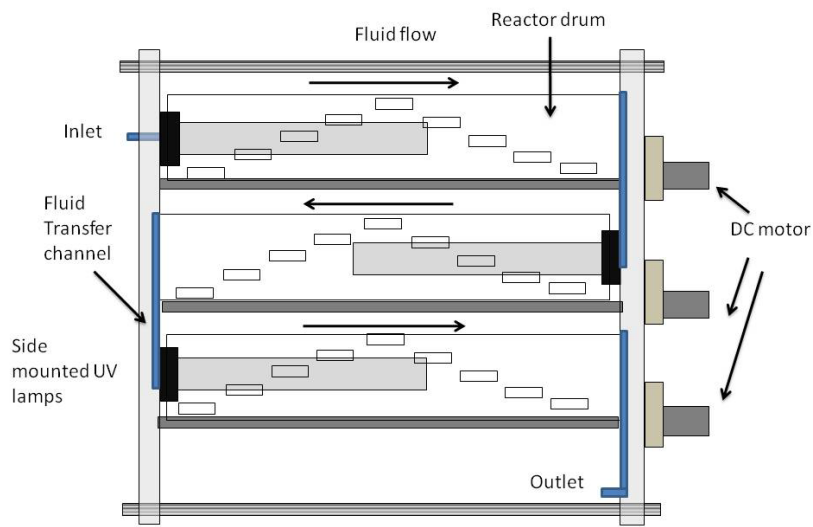


11

12

13

1 Figure 3:

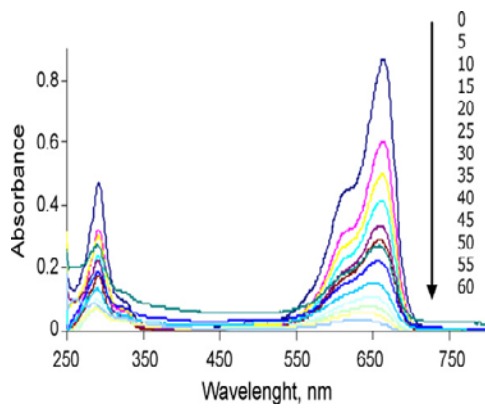


2

3

4 Figure 4:

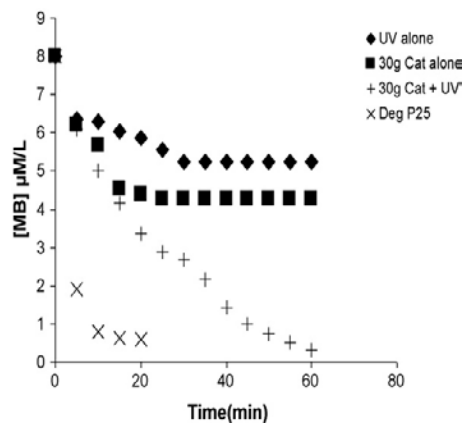
5 (A)



6

7 (B)

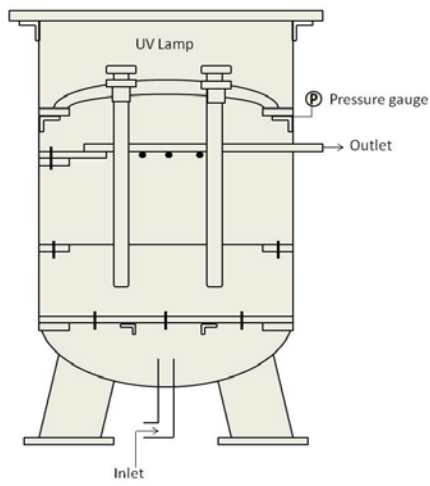
8



9

1 Figure 5:

2

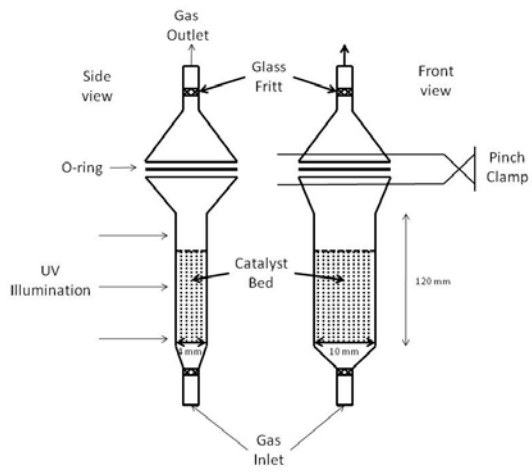


3

4

5 Figure 6:

6



7

8

9

10

11

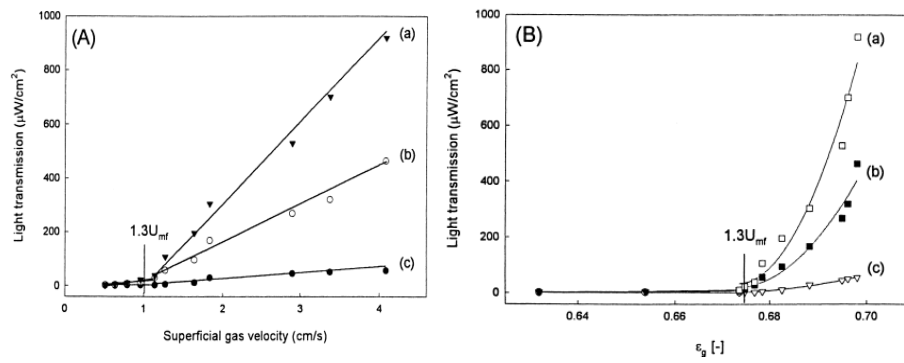
12

13

14

1 Figure 7:

2

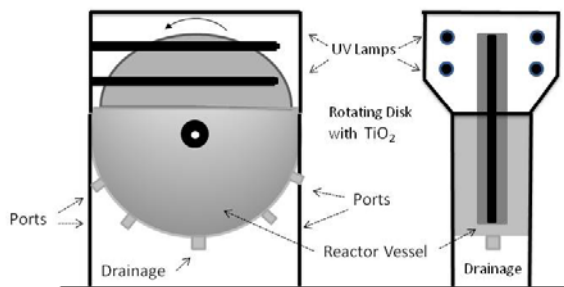


3

4

5 Figure 8:

6



7

8

9

10

11

12

13

14

15

16

17

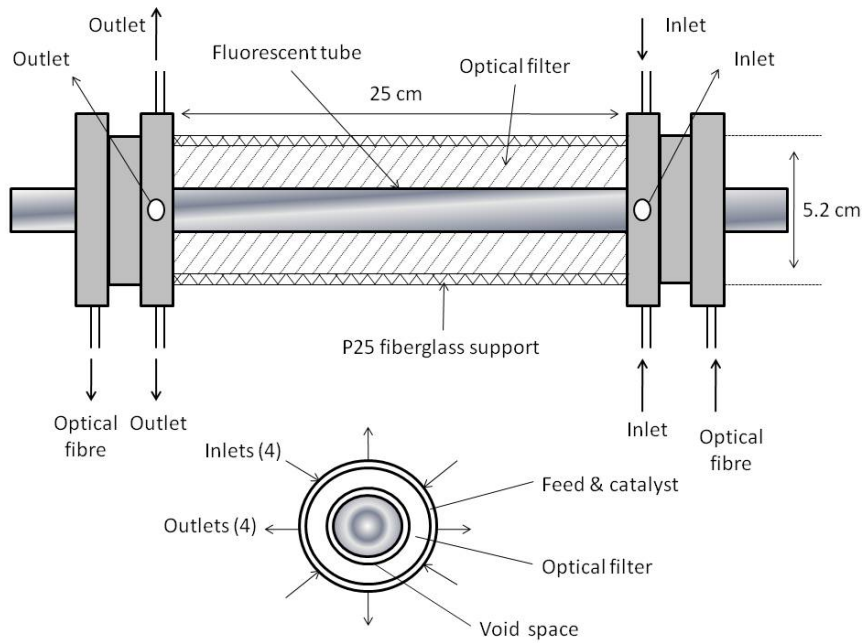
18

19

1 Figure 9:

2

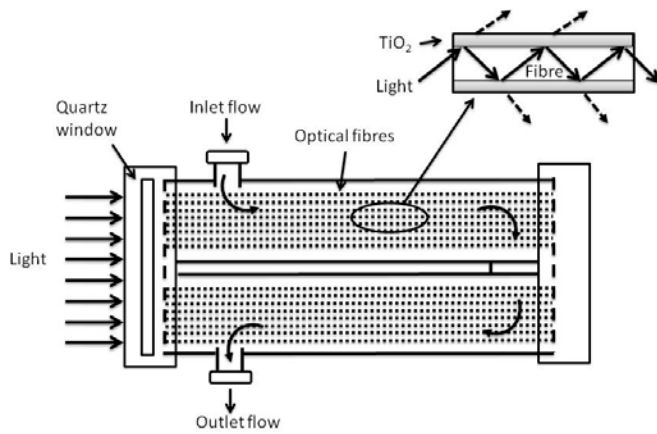
3



4

5 Figure 10:

6



7

8

9

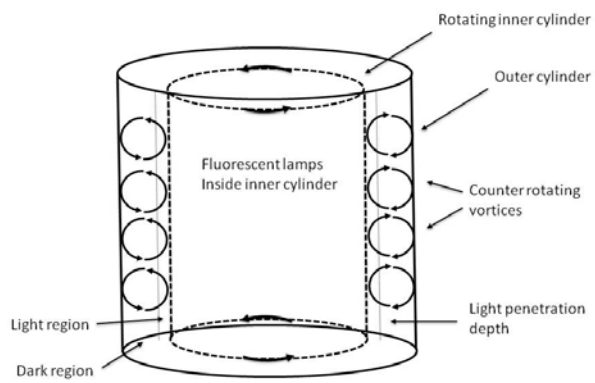
10

11

12

1 Figure 11:

2 (A)



3

4

5 (B)

6

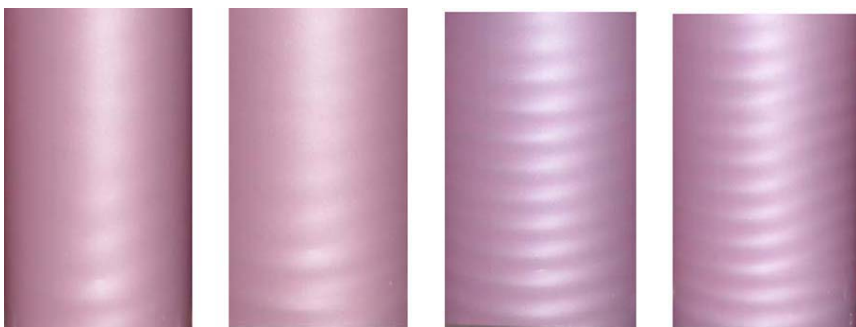


7

8

9 (C)

10



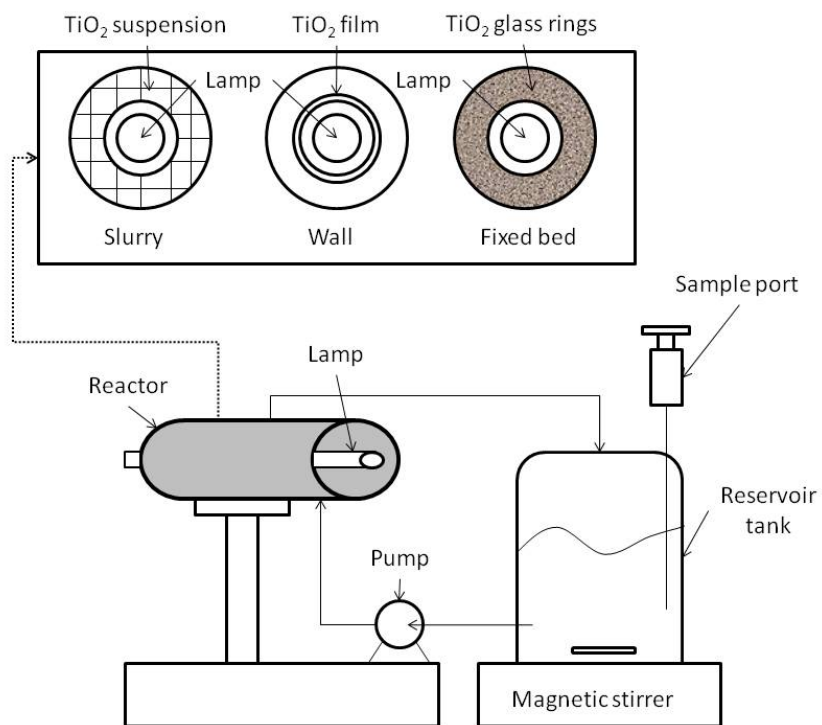
11

12



1 Figure 12:

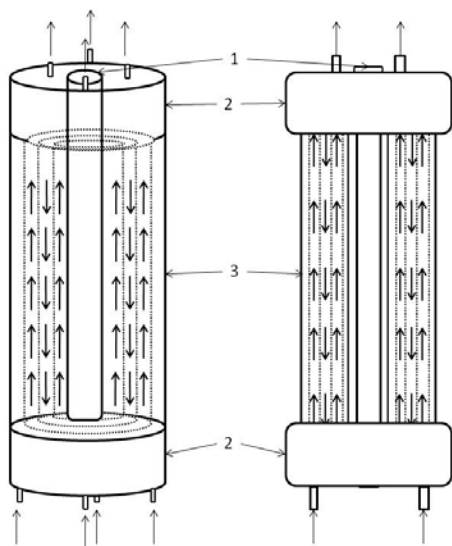
2



3

4 Figure 13:

5



6

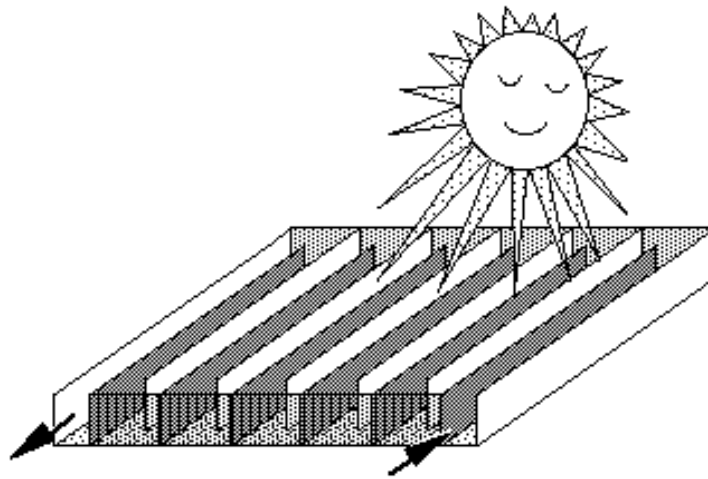
7

8

9

1 Figure 14:

2



3

4

5

6

7

1 **Captions for Figures**

2

3 Figure 1: Page 5

4 Processes that occur on photo-excitation of  $\text{TiO}_2$

5

6 Figure 2: Page 10

7 Chemical structure of the herbicides Monuron and 3,4-dichloropropionamide.

8

9 Figure 3: Page 11

10 Schematic of photocatalytic drum reactor

11

12 Figure 4: Page 13

13 (A) Temporal absorption spectral pattern displaying the degradation of MB over  
14 a 60 min time period and (B) effect of UV only, 30g catalyst (pellet form) only,  
15 UV combined with 30g catalyst and Degussa P25 (powder) effect on MB  
16 degradation.

17

18 Figure 5: Page 14

19 Schematic of fluidised bed reactor utilised for the destruction of microcystin-  
20 LR<sup>33</sup>.

21

22 Figure 6: Page 15

23 Schematic of flat plate fluidised bed reactor displaying the illumination direction  
24 and catalyst bed location<sup>66</sup>.

25

26 Figure 7: Page 16

27 Results observed for the effect of (A) superficial gas velocity and (B) voidage on  
28 light transmission with measuring light at (a) 96mm, (b) 53mm and (c) 10mm<sup>75</sup>.

29

30

1 Figure 8: Page 23  
2 Schematic representation of Rotating Disc Photo Reactor<sup>80</sup>.  
3  
4 Figure 9: Page 27  
5 Schematic representation (A) and sectional drawing (B) of the annular  
6 photoreactor<sup>26</sup>.  
7  
8 Figure 10: Page 28  
9 Schematic of Optical Fibre Photo Reactor displaying light transmission along  
10 coated fibres<sup>92</sup>.  
11  
12 Figure 11: Page 30  
13 (A) Schematic of Taylor vortex reactor<sup>29</sup> and (B) Image of Taylor vortex  
14 photocatalytic reactor. (C) Progress of time-dependent Taylor vortex flow  
15 around critical Reynolds number,  $Rec = 111$ <sup>28</sup>.  
16  
17 Figure 12: Page 32  
18 (A) A slurry reactor, using suspensions of Degussa P25 TiO<sub>2</sub>. (B) A wall reactor,  
19 immobilizing Degussa P25 TiO<sub>2</sub> onto the 15-cm long glass tube that constitutes  
20 the inner-tube wall of the reactor. (C) A fixed-bed reactor, immobilizing Degussa  
21 P25 TiO<sub>2</sub> onto 6mm\_ 6 mm glass Raschig rings placed into the annular reactor  
22 volume<sup>96</sup>  
23  
24 Figure 13: Page 33  
25 Schematic representation of a pilot-scale multi-annular photocatalytic  
26 reactor, (A) UV lamp, (B) distribution heads, and (C) borosilicate glass tubes<sup>100</sup>  
27  
28 Figure 14: Page 35  
29 Schematic View of a DSSR reactor<sup>102</sup> showing the inner structure of the transpa-  
30 rent structured box made of PLEXIGLAS<sup>®</sup> (reproduced from xx with permission  
31 from Elsevier BV).