Impact of three different TiO2 morphologies on hydrogen evolution by methanol assisted water splitting: Nanoparticles, nanotubes and aerogels

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Abstract
Increasing the activity of a photocatalyst goes through the improvement of both its absorption (light) and adsorption (reactant) properties. For a given semiconducting material, the charge carrier separation is also a very important step. Properly combining chosen phases is one option to improve this separation (example of the commercial P25) and depositing platinum on the surface of the catalyst, another one. In some cases, coupling both may nevertheless lead to a decrease of photoactivity or at least limit the potentiality of the catalyst. A third option, consisting in modifying the morphology of the photoactive phase, has shown very promising results.

In this study, we have elaborated, characterized and evaluated the hydrogen evolution potentiality (through methanol assisted water splitting) of different TiO\textsubscript{2} morphologies: nanoparticles, nanotubes and aerogels. These materials have shown different behaviours depending on both their composition and morphology. Different types of separation processes have been claimed to account for the observed different photoactivities, with more or less pronounced synergetic effects, due to: the use of Pt as a co-catalyst, the mixture of different TiO\textsubscript{2} phases (anatase and TiO\textsubscript{2}(B) or rutile) and the specific morphology of the samples (nanotubes or aerogels). Among all the tested samples, the TiO\textsubscript{2} aerogel supported Pt one exhibited very promising performances, three times as active as P25 supported Pt, which is already much more active than pure P25 in our testing conditions.

Keywords: TiO\textsubscript{2}; photocatalysis; water splitting; morphologies; nanotubes; aerogels; hydrogen
**1. Introduction**

The energetic challenge of our century is to provide enough energy to a growing population with a minor impact on the environment. In the general context of the forthcoming energy demand increase, energy resources shortage and global warming, hydrogen may be a significant part of the future energetic landscape. Indeed, this need for new energy resources, non generating greenhouse gases, could be partially fulfilled with hydrogen, provided that the production processes are environmentally friendly.

The concept of direct solar energy conversion through water splitting (water photoelectrolysis) has already been validated on numerous semiconducting materials [1-4]. A lot of work has already been done on titanium dioxide (TiO$_2$) [5], beginning with the pioneer results obtained by Fujishima and Honda [6]. Both the optoelectronic properties and the chemical characteristics of the materials tested so far, as mediator in the hv / H$_2$ conversion process, did not allow to find a satisfactory solution to obtain, under natural irradiation (sun), both a high conversion efficiency and a satisfactory material lifetime. In general, in the standard utilization conditions, an improvement of efficiency is concomitant with a diminution of the material stability. It is notably the case when titanium dioxide is replaced with sulphur-based compounds such as cadmium sulfide (CdS).

We believe that there is still room to improve the TiO$_2$ activity, especially by playing on its morphology [7]. As an example, the morphology of nanoparticles has a direct influence on the photoactivity by affecting the specific surface area, the hydrophilic character or the availability of charge carrier by modifying the internal electric field close to the surface (space charge effects). Moreover, a great anisotropy of particles induces distinct behaviors by giving more importance to some crystalline planes in contact with the surrounding environment. It should then be possible, if the grain growth can be controlled, to develop the crystalline faces corresponding to the higher photocatalytic activity.

The purpose of the work presented here is to check the influence of the morphology of different nanostructured TiO$_2$ samples. Three different types of material have been synthesized, following different routes, in the scope to obtain a representative panel of samples with various morphologies: individual nanoparticles, nanotubes and aerogels. Among the structurally and morphologically characterized samples, the most promising ones have been tested for water splitting, with methanol as a hole scavenger. Within this study, we have confirmed some common sense assumptions and demonstrated the soundness of utilizing aerogels as photocatalysts for the foreseen application. They revealed to be more active than Degussa P25, whatever the selected operating conditions. Some hypotheses have been proposed to discuss such an encouraging result.

**2. Experimental**

**2.1 Materials**

Two batches of individual nanoparticles, three of nanotubes and four of aerogels have been prepared and characterized for this study. The selected samples have then been tested and their performances compared with those of commercial samples, namely Degussa P25 and Alfa Aesar anatase (ref. 036199).
2.2 Elaboration

2.2.1 Nanoparticles

TiO$_2$ nanoparticles have been synthesized by precipitation of titanium chloride (TiCl$_4$, *Fluka 89545*) in aqueous sodium hydroxide solution (NaOH, *Sigma Aldrich S8045*) followed by crystallisation in autoclave (hydrothermal transformation):

- a 250 ml TiCl$_4$ solution (0.4 M in water, containing 10 ml of HCl) and a 250 ml NaOH solution (2 M) are simultaneously added to 500 ml of distilled water, under mechanical stirring for one hour, at atmospheric pressure, maintaining the pH at 2 with 1.6 M NaOH and the temperature at 25 °C;
- the obtained amorphous precipitate is then washed three times with distilled water and added to distilled water to prepare a suspension at pH = 10 (adjusted with 1.6M NaOH);
- finally, the crystallisation is performed with 150 ml of the prepared suspensions introduced in a 300 ml Teflon lined Hastelloy autoclave, heated at 150 °C or 250 °C for 18 h under gentle magnetic stirring.

Starting from the native as-precipitated nanoparticles, two types of samples have thus been prepared depending on the temperature of the hydrothermal treatment: NP150 [pH = 10; T = 150 °C] and NP250 [pH = 10; T = 250 °C].

2.2.2 Nanotubes

Nanotubes have been prepared following the hydrothermal transformation route developed by Kasuga [8] and already used in a former study [9]:

- 8 g of commercial TiO$_2$ anatase (99.9% *Alfa Aesar 036199*) added to 100 ml of 10M NaOH (*Sigma Aldrich S8045*) are introduced in a 300 ml Teflon lined Hastelloy autoclave, heated at 150 °C for 72 hours under stirring;
- the sodium titanates thus obtained are then washed at room temperature, three times with distilled water;
- hydrogen titanates are obtained by ion exchange, realized three times with 2 M hydrochloric acid (HCl *Fluka 84415*) : adjustment of the final suspension pH to 2, stirring 4 h at room temperature, centrifugation and washing three times with distilled water at room temperature;
- hydrogen titanates are then dried and calcined in static air for 2 h at 400, 500 and 600 °C (2 °C/min).

Three types of samples have thus been prepared: NT400, NT500 and NT600.

2.2.3 Aerogels

Aerogels have been prepared following an acid-catalysed sol-gel route followed by a drying process under supercritical conditions [10] and a calcinations step.

- 100 ml of sol are prepared by slowly adding an aqueous solution of nitric acid (2 M HNO$_3$, *Riedel-de Haën 35278*), partially diluted in isopropanol (iPrOH purum > 99.0%, *Fluka*...
59310) (2.3 ml HNO₃, 0.9 ml H₂O, 58.2 ml iPrOH), to an alcoholic solution of titanium tert-butoxyde (Ti(OC₄H₉)₄, Fluka 86910) (19.2 ml Ti(OC₄H₉)₄, 19.4 ml iPrOH), under continuous mechanical stirring. These proportions correspond to the following molar ratio, defined by Boudjey et al. [11] in their early works: Ti(OC₄H₉)₄ / iPrOH / HNO₃ (2 M) / H₂O = 1 / 18 / 0.08 / 3;

- the gel, obtained after 30 min in room conditions, is covered with iPrOH and aged 4 days at room temperature before being thoroughly washed with iPrOH, in order to eliminate traces of water;
- aerogels are then obtained after supercritical (SC) drying with CO₂, in three steps: i) solvent exchange (iPrOH with SC CO₂ / 80 bars, 37 °C, 5 kg/CO₂/h, 4 h), ii) isothermal depressurization to atmospheric pressure (0.05 bar/min, 37 °C) and iii) inertial cooling down to room temperature;
- finally, the so-called aerogels are milled in an agate mortar and calcined for 2 h in air between 400 and 700 °C (2 °C/min).

Two batches of five aerogel samples have thus been prepared: AG (not calcined), AG400, AG500, AG600 and AG700

2.2.4 Platinum deposition
Pt nanoparticles have been synthesized at the surface of the materials by classical wet impregnation of TiO₂ under magnetic stirring at room temperature with an hexachloroplatinic acidic aqueous solution (H₂PtCl₆·x H₂O Aldrich 254029, 99.995%) so as to obtain a ratio Pt/TiO₂ = 0.3 wt% (5 ml water / g TiO₂). After evaporation of the solvent and drying at 110 °C for 1 h in air, the materials were calcined at 350 °C for 1 h in flowing air and finally reduced under flowing H₂ at 350 °C for 1 h to get metallic nanoparticles.

2.3 Characterizations
All the samples have been characterized so as to determine their structure, morphology and optical properties.

2.3.1 X-Ray Diffraction
The crystallographic structure has been determined by X-Ray Diffraction using a Bruker D8 diffractometer [(θ -2θ mode, cobalt source (Co Kα = 1.79 Å), counting time = 0.43 s, step = 0.0144°, range = 10-90°].

2.3.2 High Resolution Transmission Electron Microscopy
A Tecnai F20 ST HR-TEM with a point to point resolution of 0.24 nm has been used to observe the particles: shape, size and local crystallographic structure from the lattice planes, matrix defects. The samples are deposited on Agar Scientific analysis grids: Holey Carbon Film 300 Mesh Cu for nanoparticles and Formvar/Carbon 300 Mesh Cu for nanotubes.
2.3.3 **Nitrogen Sorption Analysis**

The specific surface area has been determined based on the BET model, using a Micromeritics ASAP 2010 to collect the N\textsubscript{2} sorption isotherms. The samples are first outgassed overnight at 60 °C under vacuum, before being analysed, at 77 K.

2.3.4 **Diffuse Reflectance UV-Vis Spectroscopy**

In order to determine the light absorption range of powder samples, Diffuse Reflectance spectroscopy has been used in the UV-Visible region, with the help of an integration sphere. These measurements have been carried out on a CARY 500 UV-VIS-NIR spectrophotometer, between 300 and 600 nm. Black paper and Teflon powder have been used as references respectively for 0% and 100% reflectance. The results have essentially been used to determine the band gap of the analysed materials as follows [12, 13].

Making the assumption that the Kubelka-Munk function F(R) is proportional to the absorption coefficient, we have:

- \([F(R). \text{hv}]^{0.5} \propto (\text{hv} - \text{Eg})\) for an indirect band gap
- \([F(R). \text{hv}]^2 \propto (\text{hv} - \text{Eg})\) for a direct band gap

where \(\text{h}\) is the Planck’s constant, \(\nu\) the frequency (s\(^{-1}\)) and \(\text{Eg}\) the band gap energy value (eV).

If the absorption edge gives a rough estimation of the band gap, plotting \((F(R)\text{hv})^n\) as a function of \(\text{hv}\), allows to more precisely determine the value of the band gap that is equal the \(x\)-intercept of the extrapolated linear part of the plot.

2.3.5 **Raman spectroscopy**

Measurements have been carried out on a LabRAM HR Raman spectrometer using 514.5 nm excitation wavelength, an objective 100X and a 1800 line mm\(^{-1}\) grating. The resulting power on the sample is about 2 mW on a diffraction limited area about 1\(\mu\)m\(^2\). This characterization technique has been used to quantify the respective amounts of anatase and TiO\textsubscript{2}(B) in some of our samples, following the method proposed by Beuvier et al. [14]. \(I_\text{A}\), the area of the main peak of anatase (144 cm\(^{-1}\)), and \(I_\text{B}\), that of the main peak of TiO\textsubscript{2}(B) (123 cm\(^{-1}\)), are proportional to \(m_\text{A}\) and \(m_\text{B}\), respective weights of anatase and TiO\textsubscript{2}(B) in the sample: \(I_\text{A} = m_\text{A} \cdot \sigma_\text{A} \text{ and } I_\text{B} = m_\text{B} \cdot \sigma_\text{B}\), where \(\sigma\) is the scattering factor of the corresponding peak. Drawing \(x_\text{A} = m_\text{A}/(m_\text{A} + m_\text{B})\) as a function of \(y = I_\text{A}/(I_\text{A} + I_\text{B})\), for several mixtures of known composition, allowed Beuvier et al. to set-up the following formula: \(x_\text{A} = y / [k_\text{R} + (1 - k_\text{R}) \cdot y]\) where \(k_\text{R}\), the anatase Raman sensibility factor (\(k_\text{R} = \sigma_\text{A} / \sigma_\text{B}\)), equals 6.2.

2.4 **Photocatalytic activity**

Once characterized, the materials have been tested for hydrogen evolution with or without Pt deposit. Two cylindrical photolysis reactors (one in Strasbourg-France [15] and one in Kyoto-Japan [16]) have been used, in the center of which is placed an irradiation source, water cooled thanks to a surrounding double walled quartz jacket.
A 1 litre suspension of the catalyst is prepared with deionized water and MeOH (10 vol%) as a sacrificial agent (to favour the hydrogen evolution [17, 18]) and irradiated under magnetic stirring. Before irradiation, the suspension is outgassed under inert gas bubbling, for 90 min, in order to evacuate the oxygen from both the suspension and the free space of the reactor. The hydrogen evolved under irradiation is analyzed by a micro-Gas Phase Chromatograph, calibrated with different H\textsubscript{2}/N\textsubscript{2} standards.

The use of the two reactors mentioned above is justified by their peculiar specificities.

In the Laboratoire des Matériaux, Surfaces et Procédés pour la Catalyse (LMSPC, Strasbourg):
- the irradiation source is a 150 W metal halide lamp (BLV-Licht, C-HIT, 150 W) whose spectrum is more or less centered on the visible region (figure 1a);
- the hydrogen analysis is made under continuous nitrogen flow so that a hydrogen flow rate is actually measured.

In the Institute of Advanced Energy (IAE, Kyoto University):
- the irradiation source is a 450 W mercury vapour lamp (Ushio, UM-452, 450 W), more intense in the UV region than the metal halide lamp used in the previous reactor (figure 1b);
- the argon bubbling is stopped once the suspension has been outgassed, so the hydrogen evolved is accumulating in the free space of the reactor. Hence, here it is the total amount of hydrogen which is measured at the time of analysis.

The main difference stands in the type of lamp used, which will allow us to check the activity of our materials in different irradiation conditions (more or less energetic and powerful).

![Fig. 1. Irradiation spectrum of the sources used at a) the LMSPC, Strasbourg, France () and b) the Institute of Advanced Energy, Kyoto, Japan (UM-452 450 W)](image)

### 3. Results and discussion

#### 3.1 Characterizations

#### 3.1.1 Nanoparticles (NP)

The main structural and morphological features of the NP particles are reported in table 1.
Table 1. Structural and morphological features of commercial Degussa P25 and Alfa Aesar anatase, nanoparticles (NP), nanotubes (NT) and aerogels (AG) samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Calcination Temperature (°C)</th>
<th>Crystalline structure</th>
<th>Particle size* (nm)</th>
<th>Specific Surface Area $S_{BET}$ (m²/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degussa P25</td>
<td>-</td>
<td>A/R</td>
<td>30</td>
<td>46</td>
</tr>
<tr>
<td>Alfa Aesar Anatase</td>
<td>-</td>
<td>A/R</td>
<td>80/20 (wt%)</td>
<td>7.4</td>
</tr>
<tr>
<td>(ref. 036199)</td>
<td></td>
<td>A</td>
<td>180</td>
<td></td>
</tr>
<tr>
<td>NP150</td>
<td>150</td>
<td>A</td>
<td>10</td>
<td>143</td>
</tr>
<tr>
<td>NP250</td>
<td>250</td>
<td>A</td>
<td>90</td>
<td>17</td>
</tr>
<tr>
<td>NT300</td>
<td>300</td>
<td>B</td>
<td>-</td>
<td>246</td>
</tr>
<tr>
<td>NT400</td>
<td>400</td>
<td>A/B</td>
<td>-</td>
<td>117</td>
</tr>
<tr>
<td>NT500</td>
<td>500</td>
<td>A / traces of B</td>
<td>-</td>
<td>90</td>
</tr>
<tr>
<td>AG</td>
<td>As-prepared</td>
<td>amorphous</td>
<td>-</td>
<td>569 (a) – 600 (b)</td>
</tr>
<tr>
<td>AG400</td>
<td>400</td>
<td>A</td>
<td>10</td>
<td>159 (a) – 150 (b)</td>
</tr>
<tr>
<td>AG500</td>
<td>500</td>
<td>A</td>
<td>15</td>
<td>95 (a) – 85 (b)</td>
</tr>
<tr>
<td>AG600</td>
<td>600</td>
<td>A/R</td>
<td>10/20</td>
<td>59 (a) – 60 (b)</td>
</tr>
<tr>
<td>AG700</td>
<td>700</td>
<td>R</td>
<td>200</td>
<td>6 (a)</td>
</tr>
</tbody>
</table>

$A =$ anatase, $R =$ rutile, $B =$ TiO$_2$(B)

$a =$ batch #1, $b =$ batch #2

* from TEM observations

** determined by Raman spectroscopy

All the prepared samples are composed of anatase TiO$_2$ (figure 2). As expected, the higher the autoclave treatment temperature, the better the crystallinity, the bigger the particles and the lower the specific surface area.

So considering the foreseen application, we will focus on the low temperature treated nanoparticles NP150 (figure 3), which is composed of the smallest pure anatase nanoparticles responsible of a reasonable specific surface area (143 m²/g). This sample has then been optically characterized.

The band gap of NP150 has not surprisingly been estimated to 3.2 eV after reflectance data treatment, assuming an indirect band gap [19-22] (figure 4a). The assumption of a direct band gap, reported by some authors for nanoparticles [23] or crystals [24], would give, with this determination protocol on our materials, a value of 3.45 eV which is inconsistent with literature data reported for anatase particles [25, 26]. Such a large value could have been considered, due to the very small size of the particles. However, the assumption of an indirect bandgap is confirmed by the determination of that of NP250 (figure 4b), made up of larger particles (3.2 eV for an indirect bandgap also). Tolbert et al reported that the quantum behaviour does not depend on the nature of the bandgap (direct or indirect)
Moreover, according to Monticone et al. [22], it seems that the quantum size effect is rather limited on anatase, due to a very small exciton Bohr radius. No increase of the bandgap has been observed before diameter lower than 1.5 nm. Finally, Nozik et al. reported 3.45 eV and 3.59 eV as values for the first allowed direct transition (respectively corresponding to perpendicular and parallel light transitions) [28].

**Fig. 2.** XRD pattern of the nanoparticles NP150 and NP250, calcined at $T = 150 \, ^{\circ}C$ and $250 \, ^{\circ}C$ respectively, from the alkaline suspension (pH=10).

**Fig. 3.** TEM picture of the nanoparticles NP 150, calcined at $T = 150 \, ^{\circ}C$, from the alkaline suspension (pH=10).
Fig. 4. Band gap estimation of the nanoparticles: a) NP150 (calcined at T = 150 °C) and b) NP250, (calcined at T = 250 °C)

3.1.2 Nanotubes (NT)

The main structural and morphological features of the NT samples are reported in table 1. Figure 5 shows the XRD patterns of the NT samples for different calcination temperatures.

The TiO$_2$(B) phase appears as the only detected one after calcinations at 300 °C as already reported in the literature [29]. Neither XRD nor HR-TEM allowed us to find any trace of anatase.

Anatase appears as the main phase after calcination at 400 °C (NT400). A significative amount of TiO$_2$(B) still remains in this sample. Raman spectroscopy (figure 6) allowed us to identify both anatase [30] and TiO$_2$(B) [31] characteristic vibration modes. After subtraction of the background and decomposition of the two TiO$_2$ contributions, the quantification method presented in 2.3.5 allowed us to estimate the amount of anatase in NT400 to 80 wt%, the remaining phase being TiO$_2$(B) (20 wt%).

Finally, both phases are still present after calcination at 500 °C, in NT500, which is particularly well crystallised. TiO$_2$(B), a very minor component in NT500, has been identified on TEM pictures (figure 7).

From a morphological point of view the samples are tube-like, with, at least for NT300 and NT400, typical length and diameter of 50-100 nm and 10 nm respectively (as evaluated from TEM observations). From 400 °C and above, they tend to decompose into nanoparticles after calcination. The higher the temperature of calcination, the lower the specific surface area and the more pronounced the degradation into nanoparticles (figure 8). Cross-checking TEM observations and XRD results, it seems highly probable that, in our samples, nanotubes are made of TiO$_2$(B) while only particles issued from their decomposition are made of anatase.

These samples have been optically characterized in order to determine their band gap value. It decreases with increasing calcination temperature, from 3.35 eV for NT300 (TiO$_2$(B)) down to 3.25 eV for NT500 (anatase).
Fig. 5. XRD patterns of the nanotubes NT300, NT400 and NT500 (NT calcined at 300, 400 and 500 °C).

Fig. 6. Raman spectrum of the nanotubes NT400 (NT calcined at 400 °C), A=anatase, B=TiO$_2$(B).
Fig. 7. TEM picture of the nanotubes NT500 (NT calcined at 500 °C) evidencing the presence TiO$_2$-B (FFT realized on the white square).

Fig. 8. Modification of the nanotubes morphology with the calcination temperature, a) NT300, b) NT400, c) NT500.
3.1.3 Aerogels (AG)

The main structural and morphological features of the aerogels samples (AG) are reported in table 1. Calcined at 400 °C, the samples are made of anatase. The anatase-rutile phase transition has been observed for a calcination temperature between 500 and 600 °C (figure 9). At 700 °C, the only detected phase corresponds to rutile. The calcination temperature of our aerogel samples has thus been selected at 500 °C, in order to reach both a sufficient crystallinity and an acceptable specific surface area (97 m²/g) while avoiding the presence of rutile.

This sample (AG500), whose TEM picture is shown on figure 10, has then been optically characterized and its indirect bandgap value evaluated at 3.25 eV (the assumption of a direct bandgap would give a value of 3.6 eV).

Fig. 9. Evolution of the XRD pattern of TiO₂ aerogels with the calcination temperature.

Fig. 10. TEM picture of TiO₂ aerogels calcined at 500 °C (AG500).
3.2 *Photocatalytic activity*

The different selected materials prepared within the framework of this study have been tested for \( \text{H}_2 \) evolution by methanol assisted water photolysis and their behavior compared to that of reference materials, namely commercial anatase TiO\(_2\) from Alfa Aesar (*ref. 036199*) and P25 from Degussa. The main structural and morphological features of the reference samples are also reported in table 1.

3.2.1 *Nanoparticles (NP 10-150)*

NP150 nanoparticles revealed a photocatalytic activity (0.31 \( \text{µmol H}_2/\text{min} \)) between that of the selected commercial anatase particles Alfa Aesar (0.1 \( \text{µmol H}_2/\text{min} \)) and Degussa P25 (0.37 \( \text{µmol H}_2/\text{min} \)), the latter exhibiting the highest activity (figure 11). This can be ascribed to both the composition (crystal phase) and the morphology of the samples. On one hand, our nanoparticles, made of pure anatase, have a higher specific surface area than the selected commercial anatase ones, mainly due to smaller particles. The photocatalytic activity being strongly related to the available surface area in contact with water, the observed result is consistent with the fact that the higher the specific surface area, the higher the activity. On the other hand, despite a lower specific surface area, the slightly better behavior of P25 may be due to its particular composition, a mixture of rutile and anatase TiO\(_2\), favorable to an enhanced charge separation [32]. So the specific surface area, obviously being a major feature of photocatalysts, is probably not as important as the charge separation enhancement. These effects will be studied hereafter through the evaluation of TiO\(_2\) nanotubes and aerogels that exhibit very particular morphological features, which should also act on the charge separation.

![Fig. 11. H\(_2\) evolution on TiO\(_2\) nanoparticles NP150 (pH = 10, \( T_{\text{calc.}} = 150 \) °C) compared to P25 (Degussa) and anatase nanoparticles (Alfa Aesar) - 150 W metal halide lamp, 0.9 l water + 0.1 l MeOH, 0.7 g samples, room temperature – Strasbourg, France.](image-url)
### 3.2.2 Nanotubes (NT)

The different nanotube samples show diverse behaviors depending on their calcination temperature (figure 12).

**Fig. 12.** $\text{H}_2$ evolution on TiO$_2$ nanotubes compared to P25 (Degussa) and anatase nanoparticles (Alfa Aesar) -150 W metal halide lamp, 0.9 l water + 0.1 l MeOH, 0.7 g samples, room temperature – Strasbourg, France.

First of all, NT300, calcined at 300 °C, is the less active of the tested materials, even if it has the highest specific surface area. It is actually composed of TiO$_2$(B), whose activity has already been observed to be lower than that of anatase [33-35]. Moreover a 3.35 eV bandgap has been measured, slightly higher than that of anatase, what can partially account for the lower photocatalytic activity in the testing conditions.

The two others, NT400 and NT500, are more active than the selected commercial anatase particles. They are mainly composed of anatase (80 wt% for NT400 and almost 100 wt% for NT500), accounting for their measured bandgap, close to 3.25 eV.

The higher the calcination temperature, the lower the amount of TiO$_2$(B) and the better the cristallinity (as evidenced by Raman spectroscopy and XRD results). Both evolutions are consistent with the observed photocatalytic activities. NT400 (0.27 $\mu$mol$\text{H}_2$/min) is indeed much more active than NT300 (0.04 $\mu$mol$\text{H}_2$/min). As regards NT500, it is notably more active than NT400 during the first 15 min of the experiment, before its activity begins to decline, to end around that of the selected commercial anatase particles. TEM observations gave evidence that the higher the calcination temperature the less stable the nanotubes. They actually tend to decompose into nanoparticles. A TEM observation of NT500 after test (figure 13), seems to reveal a more pronounced decomposition than before test. The observed behavior is thus not that surprising, and may partially be ascribed to the loss of the particular
1D morphology or the weakest interaction of the nanoparticles with the remaining TiO$_2$(B) nanotubes. Finally all the samples are less active than P25. Neither their specific surface area, nor the tubular morphology of the remaining TiO$_2$(B), was it assumed to improve the charge separation thanks to an increased conductivity along the tube axis, seems to be a strong enough lever to drastically improve their performances in comparison with P25, at least in the operated testing conditions. The crystallographic phases in presence have here to be discussed. The comparison between P25 (80 wt% A, 20 wt% R) and NT400 (80 wt% A, 20 wt% TiO$_2$(B)) is particularly interesting since they are both made up of two phases, with one in common and in similar proportion: anatase. In the considered irradiation conditions, we can expect the following order of light absorption efficiency: rutile (3 eV) > anatase (3.25 eV) > TiO$_2$(B) (3.35 eV), which may partly explain the better activity of P25 (A+R) compared to that of NT400 (A+B). Moreover, the charge separation process due to the contact between the two TiO$_2$ phases seems to be more efficient with P25 than with NT400, which actually has not been specifically designed for such a purpose. Considering the composition of NT400 and P25, the respective positions of the conduction bands of rutile, anatase and TiO$_2$(B) may also partly account for the better activity of P25 in the considered irradiation conditions.

Fig. 13. TEM picture of NT500 (NT calcined at 500 °C) before (left) and after (right) H$_2$ evolution test.

It is noteworthy that when irradiated with a mercury vapor lamp, instead of a metal halide one, NT400 revealed to be more active than P25 (figure 14). In such conditions, the incident photons are more energetic and NT400 is more efficient, probably due to the higher activity of the TiO$_2$(B) part, leading to an enhanced charge separation (compared to former irradiation conditions). Compared to P25, this charge separation process may not anymore be as a determining step. Another process, related to the charge conductivity along the tube axis may be evoked to account for an additional charge separation effect and hence an increased activity of NT400 compared to P25, in such irradiation conditions.
Fig. 14. H₂ evolution on TiO₂ nanotubes calcined at 400 °C compared to P25 (Degussa) - 450 W Hg lamp, 0.72 l water + 0.08 l MeOH, 0.8 g samples, 30 °C – Kyoto, Japan.

3.2.3 Aerogels (AG)

The sample referenced AG500 has been selected based on structural and morphological characterization results. Compared to P25, a slightly better photocatalytic activity has been observed (figure 15): 16 % relative increase from 0.37 μmolH₂/min for P25 to 0.43 μmolH₂/min for AG500. This value has been confirmed in a complementary experiment carried out in the japanese photoreactor (figure 16).

At least, four parameters can be referred to, in order to explain such a behavior, i) the specific surface area, ii) the surface chemistry, iii) the connectivity of the particles and iv) the crystallinity.

AG500 unfolds a significantly higher specific surface area than P25 (97 m²/g versus 46 m²/g), what could account for part of its good activity, in correlation with better adsorption properties as already mentionned by Dagan and Tomkiewicz [36] for the photoassisted oxidation of salicylic acid for instance. However, as shown earlier, this is probably not a completely satisfaying justification. All the more so that this sample is composed only of anatase whereas P25 is a mixture of anatase and rutile TiO₂ and should see its activity enhanced compared to that of AG500, because of a better charge carrier separation.

The surface chemistry of the particles may also in this case play a significant role. As a matter of fact, aerogels are prepared following a wet chemistry route, involving water and alkoxides, all the process being realized at a relatively low temperature. Hence, we can expect a higher surface concentration of titanol groups (≡ Ti–OH) for AG500 than for P25, responsible for a supposed higher hydrophilicity [37]. Znaidi et al. reported that the number of hydroxyl groups on the surface of TiO₂ aerogels only
slightly decreases up to 500 °C, compared to the as-prepared amorphous material [38]. Since the water splitting activity is closely related to the water adsorption efficiency at the surface of the catalyst, such surface chemical states are beneficial to the process.

Finally, the higher activity of the aerogel sample may partially be explained by its very particular texture. Indeed, whereas P25 is made up of individual particles forming agglomerates through electrostatic forces (Van der Waals bonds), in aerogels materials, the particles are strongly linked through real covalent bonds. With a reduced number of interfaces, such a texture may induce a better conductivity, beneficial to the charge carrier separation, as what could be expected for nanotubes.

The enhanced electronic conductivity mentioned above, is also strongly related to the cristallinity of the material, which is improved at higher calcination temperature. The higher the calcination temperature of the aerogel samples, the lower their specific surface area but the higher their photocatalytic activity (figure 16). This is also, by the way, another illustration of the relative importance of the specific surface area. Note that the activity of the sample calcined at 500 °C is comparable to that obtained in the previous test.

![Graph](image)

**Fig. 15.** H$_2$ evolution on TiO$_2$ aerogel calcined at 500 °C (AG500) compared to P25 (Degussa) - 150 W metal halide lamp, 0.9 l water + 0.1 l MeOH, 0.7 g samples, room temperature – Strasbourg, France.
Finally the promising AG500 sample has been loaded with platinum (0.3 wt%) in order to increase its activity [39-43]. STEM-High Angular Annular Dark Field (STEM-HAADF) microscopic observations evidenced that 2.5 nm Pt particles have been deposited on the surface of AG500 (figure 17), without any particular agglomeration. In comparison, TEM observation of Pt-P25 showed a similar Pt particles distribution on the surface, with a slightly lower average particle size (1.5 nm in diameter) (figure 18).

**Fig. 16.** H₂ evolution on TiO₂ aerogels calcined at different temperatures - 450 W Hg lamp, 0.9 l water + 0.1 l MeOH, 1 g samples, 30 °C – Kyoto, Japan.

**Fig. 17.** STEM-HAADF pictures of AG500 supported Pt (0.3 wt%). White spots corresponds to Pt particles.
Fig. 18. TEM-HAADF picture of P25 supported Pt (0.3 wt%) (Black spots corresponds to Pt particles) and Pt particles size distribution (statistics on 656 particles).

Both Pt loaded samples (Pt-AG500 and Pt-P25) are much more active than bare TiO$_2$ photocatalysts, what can be ascribed to the beneficial effect of Pt nanoparticles in the charge carrier separation process [44]. Pt particles act as electron sinks one hand and, more important, catalyze the (dark) reduction of protons one the other hand. If this effect has already been well reported in the literature, more surprising is the behavior of the aerogel supported Pt sample. It remains more active than Pt-P25 but the difference of performance is significantly increased (figure 19).

Fig. 19. H$_2$ evolution on TiO$_2$ Pt loaded samples: aerogel calcined at 500 °C and P25 - 150 W metal halide lamp, 0.9 l water + 0.1 l MeOH, 0.7 g samples, 0.3 wt% Pt/TiO$_2$, room temperature – Strasbourg, France.

This may be due to a synergetic effect of the two charge separation processes, ascribed to the peculiar aerogel morphology on one hand and to the co-catalyst on another hand. If we consider that the proton reduction on Pt particles is similar on both photocatalysts, the Pt particles seem to act as electron sinks
much more efficient on Pt-AG500 than on Pt-P25. The minor increase of activity observed on Pt-P25 may also be analysed with respect to the conclusion of Emilio et al. [45]. These authors claimed that the deposit of Pt particles on the surface of P25 has a negative effect on the phenol photodegradation whereas it is beneficial for pure anatase TiO$_2$, thus confirming the results of Sun et al. [41]. This conclusion is based on the assumption that the charge carrier separation process induced by Pt on pure anatase TiO$_2$ has no influence on P25 since the charge carrier separation process induced by the connection between anatase and rutile is already very efficient. Hence they claimed that negative effects of Pt, such as light absorption or the introduction of recombination sites, are prevalent to the beneficial ones. As a consequence, if the efficiency on anatase TiO$_2$ is increased after Pt deposit, that of P25 is decreased. Regarding our results, what is claimed for phenol photodegradation cannot strictly apply to hydrogen evolution. This may nevertheless be used to analyze the difference observed. After a closer look at the results presented by Emilio et al., it is noteworthy that the photoactivity of anatase supported platinum is similar to that of pure P25 (50 min are necessary to completely degrade phenol in both cases). This seems to indicate that the beneficial effect of Pt on anatase is comparable to that of mixing anatase and rutile, at least as far as phenol photodegradation is concerned (figure 20).

![Diagram comparing photoactivity of materials](image)

**Fig. 20.** Proposed diagram to compare and analyze the different photoactivities of our materials.

A similar observation can be made comparing our results on P25 and AG500. Provided that the specific morphology of AG500 may be responsible of its good photocatalytic activity, we have, here also, two different charge carrier separation processes leading to similar photoactivity. On the contrary to what Emilio et al. reported for phenol photodegradation, we have nevertheless observed an increase of activity, concerning H$_2$ evolution, for both catalysts supported Pt (figure 20). The water degradation kinetics might not be as a limiting step as phenol degradation kinetics, what could partially accounts for the additional activity induced by the Pt deposit. In both cases, two different charge carrier
separation processes are coupled: Pt deposit, plus the anatase-rutile interface for P25 or the aerogel morphology for AG500. The former interaction (Pt + anatase-rutile interface) is here again less efficient than the latter (Pt + aerogel morphology). This could partially account for a less impressive increase of photoactivity of Pt-P25 compared to that of Pt-AG500.

Pt-AG500, more than three times more active than Pt-P25 in the selected operating conditions, appears to be a good candidate for methanol assisted water splitting applications. The mesoporous texture of TiO₂ has recently been evidenced as a major feature to increase its photocatalytic activity [41], thus confirming the interest of aerogel materials in this area.

4. Conclusion
The different TiO₂ samples tested in this study have shown different photocatalytic behaviors, depending on both their composition and morphologies. The nanoparticles we have tested revealed to be more active for H₂ evolution than the selected commercial anatase ones because of a higher specific surface area. Nevertheless, despite a higher specific surface area than P25, they were less active, stressing the point on the importance of a good charge carrier separation. The two other type of materials, namely nanotubes and aerogels, have been chosen because of the possible role of their specific morphology in the aforementioned separation process. Our nanotubes exhibited promising performances but tend to decompose in nanoparticles. Made of TiO₂(B), they showed that, under UV irradiation, the use of 1D morphology combined with the mixture of two phases (TiO₂(B) and anatase as in NT400) could lead to a better activity than P25, while under “visible” irradiation the performances are only slightly below that of P25.

TiO₂ aerogels are very promising materials since they exhibit very good performances. Without Pt co-catalyst, they can reach the level activity of P25 for hydrogen evolution. The addition of 0.3 wt% Pt nanoparticles has even allowed to largely surpass that of Pt-P25, whose activity is already ten folds that of unsupported P25. We suppose that such a good behavior may be ascribed to the beneficial interaction of two charge separation processes: one due to the use of Pt, already well illustrated in the literature, added to a process linked to the specific morphology of aerogels.

Aerogels photocatalysts will be more deeply investigated in our group, in order to clarify the origin of such good performances, that we think resides in their specific morphology, probably responsible of efficient charge carrier separation.

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References


