IMPROVEMENT POSSIBILITIES OF HETEROGENEOUS PHOTOCATALYSIS WITH THE AIM OF IN-FIELD USE

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Heterogeneous photocatalysis can be successfully applied for the degradation of organic pollutants, although the efficiency of this method is insufficient. It can be increased by modification of the catalyst with precious metals (e.g. silver) or heterogeneous catalysis can be combined with other oxidative procedures such as ozonation. Another obstacle to the in-field use of the method is that separating the catalyst from the liquid phase is difficult. This problem can be eliminated by the immobilization of the catalyst. The poly(vinyl alcohol)-TiO₂ immobilized catalyst has been prepared for this purpose. During TiO₂-based photocatalysis, active oxygen species such as the hydroxyl radical, superoxide radical and hydrogen peroxide are produced. The formation rate of the reactive •OH radicals was also determined in the case of the previously mentioned techniques. As a scavenger of this radical, coumarin was added.

Keywords: TiO₂-based photocatalysis; Triton X-100; Benzenesulfonic acid; Silver-deposited TiO₂; Immobilization; Thermal decomposition of PVA

1. Introduction

Nowadays, a great proportion of anthropogenic pollutants cannot be mineralized by traditional biological and physicochemical wastewater treatment procedures. Therefore, important developments in chemical wastewater treatment technologies and in their utilization are required. For economic, technological and healthcare reasons, in terms of both type and quantity, it is important to use as few chemical additives as possible. Their applicability against remarkably different pollutants using minimal levels of energy consumption is even more important. Advanced Oxidation Processes satisfy these challenges [1]. A common characteristic of these processes is the generation of oxidative radicals, predominantly hydroxyl radicals, using solar radiation or other kinds of energy. Hydroxyl radicals are able to oxidize a great variety of organic compounds, therefore, heterogeneous photocatalysis has become an intensively studied field of research.

In the pharmaceutical industry, benzenesulfonic acid is mostly used for producing other specialty chemicals. A variety of pharmaceutical drugs are prepared as salts of benzenesulfonic acid, known as besilates. Triton X-100 with an average n ≈ 9.5 is one of the most widely applied man-made non-ionic surfactants. Non-ionic surfactants are more stable than ionic tensides and not sensitive to pH and electrolytes. Besides the hydrophilic polyethoxy chain, it also contains a hydrophobic octylphenyl group. Both compounds can hardly be degraded by biological treatments [2–4]. Hence, a more efficient degradation method like heterogeneous photocatalysis is indispensable for the total mineralization of these pollutants.

1.1. Photodegradation of model compounds

The titanium dioxide-mediated photocatalyzed degradation of benzenesulfonic acid (later abbreviated as BS) was investigated by monitoring the absorption and emission spectral changes, sulfate concentration, pH, as well as the total organic carbon (later abbreviated as TOC) content in both argon-saturated and aerated systems.

During the degradation of benzenesulfonic acid, a characteristic change in the π→π* transition (typical of aromatic systems) in the absorption spectrum of BS (λ_{max}=262 nm, ε=439 M⁻¹ cm⁻¹) can be observed. At first, there is a slight increase in the 260-270 nm range, while two new shoulders appear at around 280 and 300 nm and as the period of irradiation progresses the level of absorbance decreases. The reason for the increase in absorbance observed in the spectrum of this aromatic surfactant is that the absorbances of the intermediates
formed in the initial period of the process are higher than that of the starting compound. On the basis of the characteristic molar absorbances of 4-hydroxybenzenesulfonic (4-HBS) ($\lambda_{\text{max}}=277$ nm, $\varepsilon=868$ $\text{M}^{-1}\text{cm}^{-1}$) and 2,5-dihydroxybenzenesulfonic acid ($\lambda_{\text{max}}=302$ nm, $\varepsilon=3300$ $\text{M}^{-1}\text{cm}^{-1}$) as commercially available standards, the appearance of a shoulder at 280-300 nm suggests the formation of hydroxylated intermediates. The results indicate that the initial step of the degradation is hydroxylation of the starting surfactant, resulting in the production of hydroxy- and dihydroxybenzenesulfonates [5–7]. Therefore, liquid chromatography-mass spectrometry analysis was utilized for the detection of intermediates.

The sulfonate group of the substrate to be degraded can easily be oxidized by oxygen-containing reactants generated in these systems. Sulfate ions were detected in the irradiated reaction mixtures, which means that the hydroxylation reactions were accompanied by desulfonation. The initial rate of this process was lower in each case than that for the transformation (i.e. hydroxylation) of the starting compound (BS). These results indicate that the first step of mineralization is hydroxylation; desulfonation takes place afterwards.

During the photocatalytic oxidation of BS, especially in the first 180 min, a continuous decrease of pH was observed in both the argon-saturated and the aerated systems, i.e. strong acidification occurred. Probably, there is a direct correlation between the formations of hydrogen and sulfate ions, due to the following reaction (Eq. (1)), which can take place between the sulfonate group and hydroxyl radical, the main oxidizing agent in these systems:

$$\text{RSO}_3^- + \bullet OH \rightarrow \text{HSO}_4^- + \text{R} \bullet \rightarrow \text{H}^+ + \text{SO}_4^{2-} + \text{R} \bullet \quad (1)$$

The hydroxy species did not decay during the irradiation in the absence of dissolved oxygen. In the aerated system desulfonation and hydroxylation was much more efficient, moreover, a significant decrease in TOC took place during the initial stage. Further hydroxylation resulted in the cleavage of the aromatic system, through the formation of polyhydroxy derivatives, followed by ring-opening, which led to the production of aldehydes and carboxylic acids. Total mineralization was realized by the end of the photocatalysis. It has been proved that in this photocatalytic procedure the presence of dissolved oxygen is indispensable for the cleavage of the aromatic ring because hydroxyl radicals photochemically formed in the deaerated system alone are not able to break the C-C bond [5].

The degradation mechanism of Triton X-100 was also investigated [8]. In the case of this surfactant, hydroxyl radicals can attack in three ways; on the ethoxy chain, on the aromatic ring or on the alkyl chain. According to the absorption spectrum (no shift in the longer wavelength (275 nm) band and no new shoulders), no hydroxylation of the aromatic ring took place during the photocatalytic degradation. After the total disappearance of the starting surfactant (after approx. 60 min), an appreciable degree of absorbance arose at 275 nm, which indicates that aromatic intermediates were formed during the first hour. In liquid chromatographic experiments, Triton X-100 produced a multiple-peaked chromatogram. The retention time of these non-ionic surfactant components is in strong correlation with the length of the ethoxy chain. Intermediates formed from various lengths of ethoxy chains and others with aromatic rings were detected in the reaction mixture after irradiation started. The concentration of the shorter-chain molecules increased during the initial stage (in the first 10 min) of the irradiation, indicating that the fragmentation of the longer polyethoxy chains of the starting components initially increased the concentration of those with shorter ones, which was followed by a gradual decay. The component with a relative molecular weight of 206 Dalton could be clearly identified. Based on its structure, it was identified as the alkyl phenol part of the original Triton X-100. Its concentration peaked at around the 50th minute of irradiation. This is the result of the total cleavage of the polyethoxy chains without any oxidation of the rest of the original tenside molecules. Based on the results so far, it can be stated that hydroxyl radicals attack the ethoxy side-chain and subsequently the alkyl group, afterwards, the ring opens.

### 1.2. Challenges of heterogeneous photocatalysis

The obstacle to the practical application of heterogeneous photocatalysis is that its efficiency is relatively low. Fortunately there are several ways to improve its mineralization effect: by the combination of heterogeneous catalysis with other oxidative procedures such as ozonation [6, 9-10] or by the modification of the catalyst with precious metals. The latter solution helps promote charge separation by accumulating the electrons on the surface of the silver particles [11-12].

The other obstacle is that the catalyst is in a suspended form, therefore, its separation from the liquid phase is difficult and makes the technology more expensive. This problem can be eliminated if the catalyst is immobilized. There are several methods for immobilization, for example, the catalyst particles can be fixed directly into polymers. Between a polymer and the catalyst usually a physical contact is formed, but in some cases, such as with poly(vinyl alcohol), a chemical bond can be formed. Unfortunately, this method is not free of disadvantages. The fact that immobilization decreases the active surface area of the catalyst has to be taken into consideration, thus, it will become less efficient [13–16].

Irradiation in the near UV range generates electron-hole pairs in the TiO$_2$ nanoparticles. Under aerobic conditions the electrons in the conduction band are scavenged by adsorbed oxygen molecules with a relatively high quantum yield, producing O$_2$•$^-$ ions, which are readily protonated in acidic media. Beides,
the holes, in the absence of reducing species, oxidize the surface-adsorbed water or hydroxide ions to hydroxyl radicals. The \( \cdot \text{OH} \) radical can also be formed by thermal reactions following the primary electron transfer processes that occur at the surface of semiconductor particles or in the liquid phase [13,17].

The hydroxyl radical is often assumed to be the major reactant responsible for the photooxidation of various organic molecules. The first step of the mineralization of benzenesulfonic acid is hydroxylation of the aromatic ring, while the degradation of Triton X-100 is initialized by hydroxyl radicals, which attack the ethoxyl side chain (Fig. 1).

Due to the high reactivity and short lifetime of \( \cdot \text{OH} \), the direct detection of this species is difficult. Appropriate methods such as steady-state fluorescence were applied to measure the quantity of scavenged hydroxyl radicals. Recent studies have proved that several non- or weakly luminescent test molecules, such as coumarin, produce strongly luminescent compounds \( (7\text{-hydroxycoumarin}) \) with hydroxyl radicals. Thus, these molecules can be utilized for detecting and measuring \( \cdot \text{OH} \) radicals produced by UV excitation of TiO\(_2\) particles in heterogeneous systems [18-19].

The main goal of our work was to investigate the effect of silver doping and immobilization of the catalyst applied for photocatalytic degradations. In this work, coumarin was used as a scavenger to determine the formation rate of hydroxyl radicals under various circumstances.

2. Experimental section

2.1. Materials

For the experiments, the following materials were used: benzenesulfonic acid \( (\text{C}_6\text{H}_4\text{OS}) \) - Alfa Aesar, 4-hydroxybenzenesulfonic acid \( (\text{C}_6\text{H}_4\text{OHOS}) \) - Alfa Aesar, silver nitrate \( (\text{AgNO}_3) \) - Aldrich, Degussa P25 TiO\(_2\), 25±5% rutile, 75±5% anatase with a specific surface area of 50 m\(^2\) g\(^{-1}\) (now called Evonik AEROSID® TiO\(_2\) P25), coumarin \( (\text{C}_9\text{H}_8\text{O}_4\text{N}) \) - VWR, Triton X-100 \( (\text{C}_{15}\text{H}_{25}\text{O}(\text{C}_2\text{H}_4\text{O})_{n,5}) \) - Alfa Aesar, and poly(vinyl alcohol) \( (\text{PVA}) \) (later abbreviated as PVA) M=146000-186000 g mol\(^{-1}\), hydrolysis: 99±% - Aldrich). The materials used for the HPLC and IC measurements were the following: methanol \( (\text{CH}_3\text{OH}) \) - Sigma-Aldrich, tetra-n-butyllammonium bromide \( ((\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_3)\text{NBr}) \) - Sigma-Aldrich, sodium sulfate \( (\text{Na}_2\text{SO}_4) \) - VWR, 37% cc. hydrochloric acid \( (\text{HCl}) \) - VWR and sodium hydroxide \( (\text{NaOH}) \) - VWR. High purity water, used in this study as a solvent, was double distilled and then purified using a Milli-Q system.

2.2. Analytics

For the analysis, 4 cm\(^3\) samples were taken, the solid phase was removed by filtration using Millipore Millex-LCR PTFE 0.45 μm filters. The absorption spectral changes of the reaction mixtures irradiated were followed using a PerkinElmer Lambda 25 UV-Vis spectrophotometer, while the emission spectra were recorded using a PerkinElmer LS 50B spectrofluorometer. Both measurements were carried out in 1 cm quartz cuvettes. Mineralization was followed by measuring the total organic carbon (TOC) concentration, utilizing a Thermo Electron TOC TN 1200 apparatus. The pH of the aqueous phase of the reaction mixture was determined by a SP10T electrode connected to a Consort C561 instrument.

HPLC analyses of the benzenesulfonic acid samples were carried out using an Agilent 1100 instrument. A water:methanol (in a volume ratio of 95:5) solvent mixture containing 0.1 % (v/v) formic acid was the mobile phase with a flow rate of 1 cm\(^3\) min\(^{-1}\). During the experiments a 100×2 mm Synergi Hydro-RP C18 (Phenomenex, Torrance, CA, USA) column packed with 2.5 μm particles was used. The column was thermostated at 30°C. The sample volume injected was 1 μl, photometric detection was applied at wavelengths of 262, 277 and 302 nm.

Degradation of Triton X-100 was followed by the 1290 Infinity UHPLC system (Agilent Technologies Inc., Santa Clara, CA, USA). The mobile phase was 65:35 methanol:water for five minutes of analysis, then it was gradually increased to 75:25 over the following five minutes. The signal was detected with a DAD detector. The column used during the experiments was a 100×2 mm Synergi Hydro-RP C18 (Phenomenex, Torrance, CA, USA) column packed with 2.5 μm particles. The column was thermostated at 50 °C. The flow rate of the eluent was 1 cm\(^3\) min\(^{-1}\).

During the study of the degradation mechanism an Agilent 6890N gas chromatograph with an Agilent DB-5ms Ultra Inert column (30 m×0.25 mm×250 μm) was used for the separation. As a detector, an Agilent 5973 N-type mass spectrometer was used.

The surface of immobilized catalysts was tested using a Philips/FEI XL30 scanning electron microscope at an accelerating voltage of 20 kV, using backscattered electron imaging. The composition was measured by an EDAX Genesis energy dispersive X-ray analyzer. Prior to testing, the samples were sputter coated by a 10 nm
thick layer of gold. Elements of the qualitative composition do not overlap with the gold, therefore, the gold coating did not disturb the analysis.

2.3. Experimental circumstances

The concentration of benzenesulfonic acid to be degraded was 10^{-3} M in each experiment, while that of the catalyst was 1 g dm^{-3} in all cases. For homogenization, the suspension was stirred for 20 minutes in the reactor before irradiation. The silver-deposited TiO_2 was prepared by photoreduction from a mixture initially containing 10^{-3} M AgNO_3 and 1 g dm^{-3} TiO_2 as a photocatalyst. After 1 hour of photoreduction the catalyst, brown in color, was separated by vacuum filtration then dried at room temperature. To re-suspend the catalyst powder before each experiment an ultrasonic bath was applied in order to achieve a dispersion of particles sufficiently small in size. During subsequent ultrasonic treatment, the substrate to be degraded was added to the system only after a period of time, so that the *OH radicals generated during the sonication could recombine and not tamper with the experimental results.

A detailed description of the production of immobilized catalyst can be found in our previous article [20]. The PVA-TiO_2 catalyst was prepared by using a PVA with a molecular weight of 146 000-186 000 and a 99+% degree of hydrolysis and Degussa P-25 TiO_2. Immobilization was carried out using a solution-casting method [21]. The PVA-TiO_2 mixture was stirred at 90 then 60°C, subsequently a viscous solution was obtained, which was then poured into a petri dish and dried. The average weight of the immobilized catalyst was 0.47 g with a PVA content of 40%. Before its use, it was subjected to a 2 hour-long thermal treatment in an argon atmosphere to increase its stability. The Triton X-100 concentration was 2 \times 10^{-3} M with 1.5 g dm^{-3} TiO_2. It is important to note that when comparing the efficiencies of the catalyst in various forms, the amount of TiO_2 immobilized in PVA (0.3 g) and used as a suspension in 200 cm^3 of reaction mixture were both 1.5 g dm^{-3}.

Photochemical experiments were carried out using two different reactors and light-source setups.

One was a laboratory-scale reactor with an effective volume of 2.5 dm^3, equipped with a 40 W light tube located in the middle of the reactor. It emitted most of its energy at wavelengths exceeding 300 nm (\lambda_{\text{max}}=350 \text{ nm}, i.e. within the UVA range) [5]. The photon flux of the light source was I_0 = 4.3 \times 10^5 \text{ mol photon dm}^{-3} \text{ s}^{-1}, which was measured by the actinometry of trisoxalatoferrate(III). The heterogeneous reaction mixture was circulated by using a peristaltic pump through the reactor and a buffer vessel. Compressed air was bubbled through the reaction mixtures from gas bottles, to facilitate stirring and (with its O_2 content) as an electron acceptor. O_2 was produced by a LAB2B ozone generator, and introduced into the same airstream. The ozone concentration was determined by iodometry, using sodium iodide as a reagent and sodium thiosulfate for the titration of the iodine formed. The ozone dosage was estimated to be 0.35 mM min^{-1}.

The other setup consisted of one small reactor made of borosilicate glass (its volume was 250 cm^3) [20]. The catalyst film was placed on a perforated glass platform (foil holder) standing on a glass frit, through which compressed air was bubbled from a gas bottle at a flow rate of 10 dm^3 h^{-1}, ensuring a constant oxygen concentration. The 200 cm^3 reaction mixture (Triton X-100 solution or distilled water in the case of pre-treatment) was circulated by a magnetic stirrer. It was irradiated from above using an Oriel LCS-100 solar simulator resulting in a light intensity of 72 mW cm^{-2} on the surface of the reaction mixture.

Fig.2 shows the change in TOC during the photodegradation of benzenesulfonic acid under various experimental conditions. The band-gap energy of TiO_2 is 3.2 eV, indicating that the catalyst can be excited by light of 360 nm in wavelength. The difference between the two light sources used is clearly visible. The radiation intensity of the solar simulator in the UV range is very low, so the total organic carbon content of the initial solution decreased by only 10 mg over 360 minutes, while total mineralization was achieved using the UV lamp within this period.

3. Results

3.1. Increase in photocatalytic efficiency using various methods

As mentioned previously, the efficiency of heterogeneous photocatalysis can be increased in multiple ways: by combination with other oxidative procedures such as ozonation and by modification of the catalyst surface using noble metals.

Since the first reaction in the mechanism of photocatalytic oxidative degradation of benzenesulfonic
acid is hydroxylation of the aromatic ring, the determination of the rates of formation of hydroxyl radicals as the main oxidant in these systems was indispensable as far as the elucidation of the mechanism under various circumstances was concerned. Coumarin was used as an \*OH scavenger.

An intensive band at 278 nm and a shoulder at around 300 nm is visible in the absorption spectrum of coumarin. The level of absorbance quickly decreased under irradiation (Fig.S1), while the concentration of the fluorescent 7-hydroxycoumarin rapidly increased during the initial period, and subsequently decreased gradually, i.e. degraded (Fig.S2). By depicting the intensity of the emission line at 453 nm as functions of the irradiation time, the initial rate of formation of 7-hydroxycoumarin, which is proportional to that of the hydroxyl radicals formed, can be determined from the initial gradients of the increasing sections, so it is easy to compare the ability of each process to generate hydroxyl radicals [18].

Before measuring the formation of 7-hydroxycoumarin in the photocatalytic oxidative systems that apply ozone, blind probes were used (Fig.3). The reaction between coumarin and ozone in the dark (at 10^{-4} M coumarin and 0.35 mM min^{-1} ozone concentrations) yielded a negligible spectral change after 2 hours. Although ozone itself is a strong oxidant, a very low rate of formation of 7-hydroxycoumarin (0.32 INT min^{-1}) was observed under these experimental conditions. Upon UV irradiation of this probe, the rate of formation slightly increased (to 1.84 INT min^{-1}).

Ozone can be absorbed over a wide range, from infrared to vacuum UV, but is optimal at 254 nm. The light source applied in this work emits most of its radiation in the 300-380 nm range, where the molar absorbances of O_3 are rather low. Thus, it can poorly promote the dissociation of ozone via the homolytic cleavage of a bond.

Contrary to the blind probe with ozone, significant spectral changes were observed in irradiated reaction mixtures containing catalysts. The surface of the catalyst was also modified with silver and combined with ozonation, then the difference between the effects of the two methods of improvement was investigated by determining the rate of formation of oxidative \*OH radicals (Fig.4).

The combination of heterogeneous photocatalysis with ozonation resulted in a 2.5 fold increase in the rate of formation (from 7.42 INT min^{-1} to 19.4 INT min^{-1}). A further increase was achieved by the application of surface-modified (silver-deposited) titanium dioxide (to 21.9 INT min^{-1}). These results are in accordance with the fact that silver deposited on the surface of the catalyst diminishes the probability of the recombination of the photogenerated electron-hole pair, i.e. promotes charge separation [11].

3.1.1. Degradation of benzenesulfonic acid

Our research group has been studying the combination
of heterogeneous photocatalysis and ozonation in detail. It has been established that a joint application of these procedures results in synergic effects [6].

Another possibility of enhancing the efficiency of photocatalysis is the modification of the catalyst surface by the deposition of silver. Our results in this work are compared with previous observations [17].

The spectral changes during the irradiations of suspensions containing Ag-TiO₂ agree with earlier results. The degree of light absorption increases during the initial period (until the 50th minute), then it decreases (Fig. S3). The fine-band structure gradually disappears, while new bands arise at longer wavelengths (277 and 302 nm), suggesting the formation of isomers of hydroxy- and dihydroxybenzenesulfonic acid [6].

Characteristic changes can also be observed in the emission spectra of the samples in these experiments: an increase in the intensity of luminescence for 60 minutes, then a decrease accompanied by a red-shift of the 290 nm band (characteristic of the starting compound) and the appearance of a new emission band at about 330 nm (Fig. S4). The band-shift becomes considerable (46.5 nm) after the first hour of irradiation: from 286 nm to 333 nm. This finding also confirms the formation of hydroxy and dihydroxy intermediates with λ_{em,max} ≈ 305 nm and 352 nm, respectively.

The changes in the concentrations of benzenesulfonic acid and the hydroxylated intermediates were followed by HPLC analyses (Fig. 5). Of the three possible hydroxy isomers, the absolute concentration could only be determined for the commercially available 4-hydroxybenzenesulfonic acid (4-HBS, rt=0.86 min).

Although the concentration of the substrate gradually decreases, the content of the benzenesulfonic acid in the reaction mixture is practically zero by the 180th minute of irradiation. On the basis of the concentration vs. time plot, the rate of decay of the model compound was determined by polynomial fit: 2.1·10⁻² mM min⁻¹. Regarding the heterogeneous catalytic experiments (untreated TiO₂ catalyst), 8·10⁻³ mM min⁻¹ was the degradation rate of the starting surfactant, while 1.4·10⁻² mM min⁻¹ was measured in the case of combination with ozonation (Table 1) [6].

Mineralization of the model compound was followed by TOC (total organic carbon content) measurements. Taking the difference between the TOC value of the reaction mixture (TOC of the solution) and that of the benzenesulfonic acid (TOC of BS, calculated from its actual concentration), the TOC value regarding the intermediates in the solution can be obtained (Fig. 6). This figure also displays the TOC values belonging to 4-HBS (calculated from its actual concentrations, see Fig. 5).

The concentration of the starting material and its TOC value rapidly decrease in the heterogeneous photocatalytic experiments involving silver-modified TiO₂ while the amount of intermediates (predominantly hydroxy and dihydroxy derivatives) gradually increases. The initial rate of decrease in TOC for the reaction mixture was 1.58·10⁻¹ mg dm⁻³ min⁻¹.

The first step of the degradation of the model compound is hydroxylation. The highest initial rate of decay of BS was observed in the case of the system containing the silver-modified catalyst (Fig. 7, Table 1). These results are in good agreement with those obtained for the experiments using coumarin, an efficient scavenger of •OH. The highest initial rate of formation of hydroxyl radicals was obtained on the Ag-TiO₂ catalyst.

### Table 1. Comparison of the initial rates

<table>
<thead>
<tr>
<th>Condition</th>
<th>TOC</th>
<th>Benzenesulfonic acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>air+Ag-TiO₂+UV</td>
<td>1.58·10⁻¹</td>
<td>2.3·10⁻²</td>
</tr>
<tr>
<td>air+TiO₂+UV</td>
<td>2.24·10⁻¹</td>
<td>6.5·10⁻²</td>
</tr>
<tr>
<td>O₂+TiO₂+UV</td>
<td>5.92·10⁻¹</td>
<td>1.3·10⁻²</td>
</tr>
</tbody>
</table>

The error of the rate determinations is ±10%.

Figure 5. Transformation of the initial compound and change in concentration of the 4-HBS intermediate. c(BS)_0=10⁻³ M, air: 40 dm³ h⁻¹, c(Ag-TiO₂)=1 g dm⁻³

Figure 6. Change in TOC contents during the treatment. c(BS)_0=10⁻³ M, air: 40 dm³ h⁻¹, c(Ag-TiO₂)=1 g dm⁻³

- ▲ – TOC of the solution (measured)
- ▲ – TOC of BS (calculated)
- ▲ – TOC of intermediates (calculated)
- ○ – TOC of 4-HBS (calculated)
The cleavage of the aromatic ring is indispensable in terms of the mineralization of the model compound, besides the fragmentation of the hydrocarbon chain. Szabó-Bárdos et al. rendered it probable that ring-opening takes place in the reaction of hydroxylated intermediates with reactive oxygen-containing species (O₂−/HO₂−, O₂(Δg)) [5]. Deviating from the decrease in the BS concentration, the lowest rate of the decrease in TOC, an indicator of mineralization, was observed in the case of the silver-modified catalyst, while the highest value (3.7 times higher than the previous one) was obtained for the combined procedure (Fig. 7, Table 1).

Based on our results, the application of heterogeneous photocatalysis with silver-modified TiO₂ only accelerates the formation of hydroxyl radicals, while its combination with ozonation also increases the amount of other oxidative radicals (O₂−/HO₂−, O₂(Δg)).

3.2. Immobilization of the catalyst with poly(vinyl alcohol)

The thermal treatment of the immobilized catalysts, the preparation of which was thoroughly described in Section 2.3, resulted in significant physical and chemical changes; on the one hand, their thickness varied (Fig. 8), on the other hand, their color turned from their original white to brown (Fig. 9). Song et al. described similar results [22].

The thickness of the foils prepared was tested by scanning electron microscopy prior to and after the thermal treatment. Following the treatment, the average thickness decreased from about 75.6 µm to about 69.6 µm (Fig. 8). The shrinkage of about 6 µm can be attributed to the evaporation of water content.

It has been established in our earlier studies that during the thermal treatment the polymer underwent dehydroxylation and cracking processes; compounds containing double and triple bonds as well as aromatic rings appeared on the surface of the foil, increasing its degree of light absorption [20]. Irradiation of the immobilized catalysts in the absence of the model compound led to the dissolution of the side-products of the cracking processes, and they also underwent heterogeneous photocatalytic degradation due to the presence of TiO₂. Accordingly, stearic acid and its derivatives containing double bonds such as linoleic acid (9,12-octadecadienoic acid (Z,Z) and oleic acid (9-octadecenoic acid) were detected by GC-MS analysis of

![Figure 7](image)

*Figure 7. Change in concentrations of BS (a) and TOC (b) under various circumstances.*

![Figure 8](image)

*Figure 8. Cross sections of scanning electron micrographs of PVA-TiO₂ foils before (a) and after (b) thermal treatment.*
the liquid phase. During the photocatalytic degradation, organic acids with shorter hydrocarbon chains were also produced, e.g. tetradecanoic acid (C14), caprylic acid (C8) and malic acid (C4) [20]. Although thermal treatment improves the stability of the foil, it restricts its efficiency because a considerable proportion of the oxidative radicals formed during photocatalysis are consumed in the reactions with the pollutants produced during the thermal treatment. Taking all these factors into consideration, how the temperature of the thermal treatment affects the stability of the immobilized catalyst was also studied.

The foils heated at different temperatures were submersed in 200 cm$^3$ of distilled water, then irradiated for 8 hours. The change in TOC of the liquid phase is shown in Fig. 10/a, while the reductions in the foil masses are displayed in Fig. 10/b. For the interpretation of the results, the increase in the temperature of the solution due to irradiation also has to be taken into account, because it enhances the solubility of poly(vinyl alcohol), i.e. the amount of the dissolved PVA, which increases the actual TOC value. The organic materials (PVA and the products of the thermal treatment) dissolved off the foil which also underwent photocatalytic degradation in the presence of TiO$_2$. Both processes determine the change in TOC in the liquid phase, along with the mass reduction of the foil. The stability of the foils treated at lower temperatures (e.g. at 60$^\circ$C) is rather modest, the PVA is dissolved in water, the TOC value of the liquid phase increased to 130 mg dm$^{-3}$ by the end of the experiment, and the mass of the foil decreases considerably. The stability of the foils treated at higher temperatures (e.g. at 140$^\circ$C or 200$^\circ$C) significantly increased, while the TOC value of the liquid phase only grew slightly (to 34 mg dm$^{-3}$ at 140$^\circ$C, to 18 mg dm$^{-3}$ at 200$^\circ$C), in a similar fashion to the weight loss of the foils (140$^\circ$C - $\Delta$m=93 mg, 200$^\circ$C - $\Delta$m=24 mg).

The results clearly indicate that by increasing the temperature of the thermal treatment, the stability of the foil is gradually improved, due to efficient crosslinking [21]. The thermally treated immobilized catalysts cannot be applied for photocatalytic purposes, because they undergo degradation themselves during the irradiation, increasing the TOC value of the solution phase. Besides, the brownish organic compounds formed in the cracking processes diminish the catalytic efficiency because they occupy the active sites on the catalyst surface. Hence, a compromise has to be made between stability and efficiency. Efforts were made to mineralize the products of cracking during a pretreatment. Before the photocatalytic experiments, the thermally treated foils were cleaned by three sequential 8-hour-long pre-treatments, to remove the pollutants formed during the cracking processes. Since the pollutants formed whilst heated at the higher temperature (200$^\circ$C) were rather difficult to remove, 140$^\circ$C was applied for the thermal treatment of the foils in further experiments of ours [20].

3.2.1. Efficiency of •OH production during pre-treatments of immobilized catalysts

To detect the hydroxyl radicals formed during the irradiation of TiO$_2$ particles embedded in the polymer, 10$^{-4}$ M of coumarin as a scavenger was applied. Following the thermal treatment to enhance the stability of the foils, the undesirable products of this procedure have to be removed. In the first cycle of this process, when the foil is yet brown, the immobilized photocatalyst starts to work upon irradiation, but oxidative radicals formed in this process are mostly involved in the degradation of the products of cracking. The initial rate for the formation of 7-hydroxycoumarin just slightly exceeds the value observed in blind experiments (in the absence of a catalyst); 3·10$^{-2}$ INT min$^{-1}$ (blind), 4·10$^{-2}$ INT min$^{-1}$ (1st cycle) (Fig. 11).

Figure 10. Change in the TOC content of the liquid phase after various heat treatments (a) and the weight loss of foils after 8 hours of irradiation (b).
200 cm$^3$ distilled water, air: 10 dm$^3$ h$^{-1}$, c(TiO$_2$)=1.5 g dm$^{-3}$
The rate of formation of the hydroxyl radicals significantly increases during the 2nd cycle; the emission intensity at the end of this cycle ($v_0=10^{-4}$ INT min$^{-1}$) is more than twice as high as in the previous one. The results obtained during the 3rd cycle are similar to those during the 2nd one. Practically, during the three sequential 8-hour-long periods of irradiation, the polluting compounds that originate from the thermal treatment are totally degraded, hence, the active sites on the surface of the catalyst become more accessible, resulting in a much higher rate for the formation of oxidative radicals (in the 4th cycle $v_0=2.5-2.8\cdot10^{-4}$ INT min$^{-1}$, see Fig.11). In other applications, the radical producing efficiency of the catalyst does not change significantly, i.e. the foil can be applied at constant efficacy. The lower value determined during the 7th cycle suggests a slow rate of degradation of the immobilized catalyst, which has to be taken into consideration during its application.

3.2.2. Degradation of the model compound

The photocatalytic degradation of the model compound was investigated following the cleaning procedure. The average degree of degradation of Triton X-100 in three sequential cycles was 62%; the deviations might originate from measurement errors (Fig.12). This value is significantly lower than that observed in the case of the suspended catalyst (90%). Although the catalyst concentrations with regard to the volume of solution were equal (1.5 g dm$^{-3}$), immobilization considerably diminished the specific surface of the catalyst, leading to a substantial decrease in its efficiency. Mineralization of Triton X-100 takes place via cleavage-producing intermediates. The cleavage of the alkyl-phenyl part of the starting components of Triton X-100 took place from the beginning of the irradiation period. As complete intermediates originated from such a cleavage, ethylene glycol and dioxolane derivatives were detected by GC-MS. At the beginning of the irradiation the edge of the starting molecules or the initially generated intermediates mineralize to CO$_2$. Therefore, the decrease in TOC is much less than expected from the concentration change. The concentration changes of Triton X-100 components and the typical intermediates are in accordance with our previous results using a similar system that applied a suspended catalyst (see section 1.1). Remarkably, GC-MS analyses did not detect any intermediates which might derive from the PVA support. This suggested that three pre-treatment cycles of irradiation and rinsing were sufficient for a satisfactory level of removal of the derivatives formed during the thermal treatment.

3.2.3. Surface morphology of foils - SEM analysis

The behavior of the active surface of immobilized photocatalysts after repeated use raises an important issue. The surfaces of titanium dioxide containing PVA before and after thermal treatment are shown in Figs.13/a and 13/b, respectively. Some TiO$_2$-rich islands (Fig.14) are not fully covered by the PVA film, these are visible on the surface. Otherwise, no significant changes are observed after the thermal treatment. However, simulated solar radiation (repeated usage) induces visible changes in the morphology of the

Figure 11. Change in the intensity of luminescence of 7-hydroxycoumarin formed.

$c$(coumarin)$_0=10^{-5}$ M, air: 10 dm$^3$ h$^{-1}$, $\lambda_{exc}=332$ nm

Figure 12. Change in concentration change of Triton X-100 during the irradiation.

$c$(TX-100)$_0=2\cdot10^{-4}$ M, air: 10 dm$^3$ h$^{-1}$, $c$(TiO$_2$)$_0=1.5$ g dm$^{-3}$
surface. Such a period of radiation partially or totally removes the PVA layer covering the TiO$_2$ agglomerates. This process is enhanced after prolonged use: after four cycles the homogeneously distributed particles are already visible (Fig.13/c). Extended use - seven cycles - will enhance grain boundaries by removing more of the PVA matrix, indeed some of the particles become loosely bounded to the rest of the agglomerate (Fig.13/d). The observed changes in thickness strongly correlate with the degradation of films (Fig.15). After four cycles, a significant decrease in thickness can be observed: the foil thickness decreases to about 58.5 µm (from 69.6 µm). This corresponds to a decrease of about 2.7 µm / cycle. As for the thickness measured after seven cycles, the last three cycles only represent a decrease of about 0.9 µm / cycle. These observations imply a more intense degree of degradation during the initial stage.

4. Conclusion

Heterogeneous photocatalytic procedures are widely applied for the degradation of organic pollutants. The degradation of benzenesulfonic acid and Triton X-100 by heterogeneous photocatalysis was investigated under various circumstances. The formation of hydroxyl radicals as one of the major reactants responsible for the photooxidation of these molecules was also measured in the case of both setups of the catalyst.

Silver deposition on the surface of the TiO$_2$ catalyst significantly increased the initial degradation (hydroxylation) rate of BS, due to the hindered recombination of the photogenerated electron-hole pairs, increasing the rate of •OH production. However, mineralization of the intermediates was not accelerated by this method, deviating from the combination of the TiO$_2$-based photocatalysis with ozonation. The degradation efficiency in the latter procedure indicated a synergic effect on the mineralization rate as a consequence of the enhanced production of oxidative agents other than •OH radicals [6]. These reactive oxidants play key roles in the ring-opening processes needed for the mineralization of aromatic intermediates. Quantitative determinations of such reactants (e.g. O$_2$• and O$_2$($^1$Δg)) to gain a deeper understanding of the mineralization mechanism are in progress. Our results indicate that the methods applied for the enhancement of the degradation efficiency in heterogeneous photocatalysis affect the generation of the oxidizing agents in different ways, which ought to be taken into consideration in further studies.

In general, suspensions of the photocatalyst are used in these procedures. Thus, at the end of these processes the semiconductor particles have to be separated from the cleaned solution. This operation can be taken out by immobilization of the catalyst, which was achieved by using poly(vinyl alcohol) in this work.
The stability of the PVA support proved to be too low for practical use. The stability of the PVA-based foils could be considerably improved by thermal treatment. However, before photocatalytic application the water-soluble products of the heating process had to be removed from the surface of the PVA-TiO₂ foil. According to the efficiency of •OH formation, three cycles of irradiation and rinsing proved to be sufficient for this purpose. The foil pre-treated by this method could be applied for the photocatalytic degradation of a frequently used non-ionic detergent, Triton X-100. When using the immobilized catalyst, it should be noted that the foil catalyst is degraded, its thickness decreases and the catalyst particles become more and more accessible on the surface.

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Figure 15. Scanning electron micrographs of cross sections of film. a) after 4 cycles of irradiation b) after 7 cycles of irradiation.


