Catalytic oxidation of organic sulfides by H$_2$O$_2$ in the presence of titanosilicate zeolites

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Catalytic oxidation of organic sulfides by H$_2$O$_2$ in the presence of titanosilicate zeolites (FER) and its delaminated form (ITQ-6), with various Si/Ti molar ratios, were synthesized and tested as catalysts for diphenyl sulfide (Ph$_2$S) and dimethyl sulfide (DMS) oxidation with H$_2$O$_2$. The zeolites were characterized with respect to their chemical composition (ICP-OES), structure (XRD, UV–vis DRS) and texture (low-temperature N$_2$ adsorption-desorption). Titanium in the FER and ITQ-6 samples was present mainly in the zeolite framework with a significant contribution of titanium in the extraframework positions. Titanosilicate zeolites of FER and ITQ-6 series were found to be active catalysts of diphenyl and dimethyl sulfides oxidation by H$_2$O$_2$ to sulfoxides (Ph$_2$SO/DMSO) and sulfones (Ph$_2$SO$_2$/DMSO$_2$). The efficiency of these reactions depends on the porous structure of the zeolite catalysts – conversion of larger molecules of diphenyl sulfide was significantly higher in the presence of delaminated zeolite Ti-ITQ-6 due to the possibility of the interlayer mesopores penetration by reactants. On the other side diphenyl sulfide molecules are too large to be accommodated into micropores of FER zeolite. The efficiency of dimethyl sulfide conversion, due to relatively small size of this molecule, was similar in the presence of Ti-FER and Ti-ITQ-6 zeolites. For all catalysts, the organic sulfide conversion was significantly intensified under UV irradiation. It was suggested that Ti cations in the zeolite framework, as well as in the extraframework, species play a role of the single site photocatalysts active in the formation of hydroxyl radicals, which are known to be effective oxidants of the organic sulfides.

1. Introduction

Zeolites have been known as very important materials for catalysis since their successful application in petrochemistry in the 60’s of 20th century [1]. Since that time many new zeolite topologies and their applications in industry have been developed. One of the most interesting synthetic zeolite is ferrierite (FER), based on the 5-membered rings (MR) with two types of perpendicularly intersecting channels (delimited by 8 and 10 MR) [2]. Precursors of ferrierite, PREFER, are characterized by the layered structure, in which the zeolite sheets are separated by surfactant molecules. During their calcination organic surfactants are removed from PREFER resulting in the condensation of silanol groups from the pinnately placed layers with the formation of 3D microporous structure of FER [2]. The specific layered structure of PREFER gives also an opportunity to obtain delaminated zeolitic materials, characterized by the hierarchical microporous and mesoporous structure [3]. Such delaminated zeolitic materials, called ITQ-6, and also microporous FER are very interesting as catalysts or catalytic supports for various chemical processes [4]. Their applicability is related not only to the porous structure but also to the presence of acid sites, as well as ion-exchange properties and therefore possibility of uniform deposition of catalytically active metal ions [4–7]. Moreover, a very important advantage represented by ITQ-6 is its delaminated structure consisting of larger pores located between chaotically arranged zeolite layers and mesopores inside zeolite layers. Such hierarchical porous structure was reported to be effective in the conversion of bulkier molecules due to reduced internal diffusion restrictions of reactants. Examples of this effect are comparative studies of Ti-FER and Ti-ITQ-6 based catalysts for epoxidation of 1-hexene with H$_2$O$_2$ [5,8] or styrene epoxidation with tertbutyl hydroperoxide [7]. Titanium substituted into the zeolite framework results in the modification of its acidic character. Titanosilicate zeolites have been reported to be more effective in binding and
activation of some organic molecules and therefore have been known to be efficient catalysts for various oxidation processes [9]. An example is application of Ti-zeolites – TS-1, TS-2, Ti-beta – as effective catalysts for the selective oxidation of diphenyl, methyl phenyl and dipropyl sulfides [10]. One of the main problems of bulky organic sulfides oxidation is related to the internal diffusion limitations of bulky reactants inside pores. To overcome this problem the zeolitic catalysts with the combined micro-mesoporous structure can be used [9–11]. Internal diffusion of bulky molecules in mesopores is much faster than in micropores and therefore the overall reaction rate in the case of the catalysts with the hierarchical micro-mesoporous structure should be much faster comparing to microporous catalysts. Many organic sulfides, e.g. dimethyl sulfides, diphenyl sulfides and products of their oxidation, organic sulfoxide and sulfones, are important chemicals for various applications, including pharmacy and medicine. They are used in production of various pharmaceuticals, such as vasodilators, phototropics, antiulcer and antihypertensive medicaments, as well as antibacterial and antifungal agents. Organic sulfoxide and sulfones can be produced by selective oxidation of suitable organic sulfides. Among various oxidizing agents hydrogen peroxide, \( \text{H}_2\text{O}_2 \), which is nontoxic, clean and produces only water as by-product, seems to be the most promising one [12,13]. Our previous studies have shown very promising results of diphenyl sulfides oxidation to diphenyl sulfoxide and sulfone by \( \text{H}_2\text{O}_2 \) in the presence of TiO\(_2\)-based catalysts [14]. These studies were extended for tetrainsolate zeolites, Ti-FER and Ti-ITQ-6, used as catalysts in the process of dimethyl and diphenyl sulfides oxidation using hydrogen peroxide as an oxidant with and without UV irradiation.

2. Experimental

2.1. Synthesis of catalysts

Two series of the zeolitic samples, Ti-FER and Ti-ITQ-6, with different Si/Ti molar ratios were prepared based on the recipe reported earlier [3]. Ti-PREFER materials were synthesized using fumed silica (Aerosil 200, silicon source), titanium (IV) ethoxide (TEOTi, titanium source), 4-amino-2,2,6,6-tetramethylpiperidine (R, structure directing agent), \( \text{NH}_4\text{F} \), HF and distilled water in the following molar ratios – 1 SiO\(_2\)-x TEOTi: 1 R: 1.5 NH\(_4\)F: 0.5 HF: 10H\(_2\)O, where \( x = 0.08, 0.04, 0.02 \) and 0.01 for the intended Si/Ti molar ratios equal to 12.5, 25, 50 and 100, respectively. The obtained gels were continuously stirred in autoclaves at 135 °C for 10 days. The resulting solid products were filtered, washed with distilled water and dried at 60 °C. The synthesis resulted in four Ti-PREFER samples with the attended Si/Ti molar ratios of 12.5, 25, 50 and 100, denoted as Ti-PREFER-12.5, Ti-PREFER-25, Ti-PREFER-50 and Ti-PREFER-100, respectively.

Each of the obtained Ti-PREFER samples was divided into two portions. The first one was calcined at 650 °C for 10 h resulting in the condensation of the ferrierite layers with the formation of three dimensional microporous ferrierite zeolites with the intended Si/Ti molar ratios of 12.5, 25, 50 and 100, denoted as Ti-FER-12.5, Ti-FER-25, Ti-FER-50 and Ti-FER-100, respectively. The second part of the Ti-PREFER samples was dispersed in a solution consisting 40 g of \( \text{H}_2\text{O} \), 200 g of cetyltrimethylammonium bromide (CTMABr, 25 wt%, 50% exchanged \( \text{Br}/\text{OH} \)) and 60 g of tetrapropylammonium bromide (TPABr, 40 wt%, 30% exchanged \( \text{Br}/\text{OH} \)) and refluxed at 80 °C for 16 h. Then, the slurries were sonicated in an ultrasound bath (50 W, 40 kHz) for 1 h to disperse the swollen ferrierite sheets. In the next step pH of slurries was decreased to about 3.0 with the use of concentrated HCl and then the solid samples were recovered by centrifugation and washed with distilled water. After drying at 60 °C and calcination at 650 °C for 10 h, a series of the Ti-ITQ-6 samples with the intended Si/Ti molar ratios of 12.5, 25, 50 and 100, denoted as Ti-ITQ-6-12.5, Ti-ITQ-6-25, Ti-ITQ-6-50 and Ti-ITQ-6-100, respectively, was obtained.

2.2. Characterisation of the zeolite samples

The obtained zeolite materials were characterized with respect to their chemical composition, structure and texture. The chemical composition of the samples was determined by ICP-OES method using an iCAP 7400 instrument (Thermo Science). The solid samples were dissolved in a mixture of hydrofluoric, hydrochloric and nitric acid solutions assisted by microwave radiation using Ethos Easy system (Milestone). X-ray diffractograms of the zeolite samples were obtained using Brucker D2 diffractometer. The measurements were performed with Cu-Kα radiation in the 2 Theta range of 2–45° with a step of 0.02° and a counting time of 1 s per step. Textural properties of the samples were determined by N\(_2\) adsorption-desorption measurements at –196 °C using a 3 Flex (Micrometrics) automated gas adsorption system. Prior to measurements, the samples were outgassed under vacuum at 350 °C for 24 h. The specific surface area value was determined using BET equation. Distributions of micropore sizes were determined using the Horvath-Kawazoe model, while for the mesopore range according to BJH model. The total pore volume was determined by means of the total amount of adsorbed nitrogen at \( p/p_0 = 0.98 \). Micropore volume was determined using the t-plot method. The UV-vis diffuse reflectance spectra of the samples were measured at room temperature using an Evolution 600 (Thermo Science) spectrophotometer. The spectra were recorded in the range of 190–900 nm with a resolution of 4 nm.

2.3. Catalytic tests

The zeolitic samples of Ti-FER and Ti-ITQ-6 series were tested as catalysts for oxidation of diphenyl (Ph\(_2\)S) and dimethyl (DMS) sulfides to sulfoxides (Ph\(_2\)SO/DMDSO) and sulfones (Ph\(_2\)SO\(_2\)/DMSO\(_2\)) in the presence of hydrogen peroxide (H\(_2\)O\(_2\)) as an oxidant. The reaction was performed in a 100 cm\(^3\) round-bottom flask equipped with stirrer, dropping funnel and thermometer. The reaction mixture consisted of 0.4 mmol of diphenyl sulfide (or dimethyl sulfide, DMS), 20 cm\(^3\) of acetonitrile used as a solvent, 0.1 mmol of bromobenzene used as an internal standard and 25 mg of the catalyst. The obtained mixture was stirred (1000 rpm) at 25 °C for 10 min and then 2 mmol of hydrogen peroxide (30% solution of H\(_2\)O\(_2\)) was added. The catalytic reaction was performed in the dark in order to avoid photocatalytic conversion of Ph\(_2\)S (conditions marked here as “DARK”). Moreover, the reaction was also performed under UV irradiation (marked as “LIGHT”). In this case a 150 W xenon short arc lamp was used as a UV light source (11.65 mW cm\(^{-2}\)). To avoid excitation of Ph\(_2\)S and its direct photooxidation a 320 nm cut off filter was applied, as well as a NIR and IR filter (10 cm optical path, 0.1 mol dm\(^{-3}\) solution of CuSO\(_4\)). The reaction progress was monitored by analysis of the reaction mixture by HPLC method. The mixture of acetonitrile/water with the volume ratio of 80:20 was used as the eluent. The samples of the reaction mixture were taken in regular intervals – every 10 min within the first hour and every 30 min afterwards, filtered through the 0.22 μm Nylon membrane filter and analysed at a Flexar chromatograph (PerkinElmer) equipped with the analytical C18 column (150 mm × 4.6 mm i.d., 5 μm pore size). The column was maintained at 25 °C throughout analysis and the UV detector was set at 254 nm for oxidation of Ph\(_2\)S or 210 nm for oxidation of DMS. Catalytic and photocatalytic tests were conducted with the over-stoichiometric excess of H\(_2\)O\(_2\) (H\(_2\)O\(_2\)/sulphide molar ratio of 5). In such conditions the reaction rate is not limited by the actual content of H\(_2\)O\(_2\) in the reaction mixture. The examples of the results of the photocatalytic tests conducted with different H\(_2\)O\(_2\)/Ph\(_2\)S molar ratios and the ratios of the H\(_2\)O\(_2\)/Ph\(_2\)S conversions in these reactions are presented in Supplementary materials.

Hydroxyl radicals generation was examined by testing the reaction of terephthalic acid (TA) hydroxylation. Studied materials (0.5 g dm\(^{-3}\)) suspended in 16 cm\(^3\) of the terephthalic acid solution (Aldrich, 98%; 3 × 10\(^{-3}\) mol dm\(^{-3}\) dissolved in 0.01 mol dm\(^{-3}\) NaOH, pH = 11) were irradiated with an XBO-150 xenon lamp (Instytut Fotonowy, 8.1 mW
cm$^2$). To avoid excitation of TA a 320 nm cut off filter was used as well as NIR and IR filter (10 cm optical path, 0.1 mol dm$^{-3}$ solution of CuSO$_4$). Samples of 2 cm$^2$ were collected during irradiation and then centrifuged to separate the photocatalyst powder. In the reaction of non-fluorescent TA with hydroxyl radicals hydroxysteraphelic acid (TAOH) is formed. Formation of TAOH was monitored by the emission spectroscopy. It shows a broad emission band at $\lambda_{\text{max}} = 425$ nm (when excited at $\lambda_{\text{exc}} = 315$ nm).

3. Results and discussion

3.1. Characterization of the samples

Chemical composition of the zeolite samples is presented in Table 1. It can be seen, that the real Si/Ti molar ratios are higher than intended values, indicating the lower titanium content in the samples then it was planned. The Si/Ti molar ratios in the analogous Ti-FER and Ti-ITQ-6 samples are slightly different. It is possible that part of titanium was removed from Ti-PREFER during its delamination. This effect is more distinct for the samples with the higher titanium content.

The X-ray diffraction patterns of the Ti-FER samples, presented in Fig. 1A, are typical of ferrierite zeolite [15]. An increase in titanium loading resulted in a decrease in intensity of the reflections, what is possibly related to decreased ordering of the zeolite framework and crystallinity of the samples with the larger Ti-content [4,16,17]. Delamination of the Ti-PREFER structure, resulting in the Ti-ITQ-6 series, decreased intensity of the reflections characteristic of ferrierite (Fig. 1B). It is related to the significantly limited long-distance ordering in delaminated structures. Similarly to the Ti-FER samples, an increase in titanium content resulted in decreased intensity of the reflections characteristic of ferrierite in the Ti-ITQ-6 series. No reflections characteristic of TiO$_2$ or any other titanium containing phases were identified in diffractograms of the zeolitic samples.

Nitrogen adsorption-desorption isotherms of the studied samples are presented in Fig. 2, while their textural parameters are compared in Table 2. Isotherms of the Ti-FER samples can be qualified as isotherms of the type I characteristic of microporous materials (Fig. 2A). This type of isotherm shows a steep adsorption at low relative pressure, which is assigned to nitrogen condensation in micropores [18]. Comparison of textural parameters (Table 2) of the Ti-FER samples shows only small decrease in the BET surface area from 400 m$^2$ g$^{-1}$ for zeolite with the lowest titanium content (Ti-FER-100) to 378 m$^2$ g$^{-1}$ for the sample with the highest titanium loading (Ti-FER-12.5). The changes in micropore volume ($V_{\text{MLO}}$) follow the same tendency. Pore size distributions, determined in the range of micropores and mesopores for a series of the Ti-FER samples, are presented in Fig. 3A. The maximum of pore size distribution in a series of the Ti-FER samples is located at about 0.53–0.57 nm, what is in full agreement with the diameter of 10 MR channels in ferrierite [19]. No peaks in the mesopore range were found in the pore size distribution profiles of the Ti-FER samples.

The nitrogen adsorption-desorption isotherms recorded for the series of the Ti-ITQ-6 samples, presented in Fig. 2B, is the type IV, characteristic for mesoporous materials. Moreover, an increase in adsorbed volume of nitrogen observed at very low relative nitrogen pressure indicates also a significant contribution of micropores in this series of the samples. Micropores are located in the zeolitic layers, while mesopores are the spaces between chaotically oriented zeolite layers. The hysteresis loops are the H3 type, characteristic of non-rigid aggregates of plate like particles [20], typical of the ITQ-6 structure [21,22]. Profiles of pore size distributions in the micropore and mesopore ranges for a series of the Ti-ITQ-6 are presented in Fig. 3B. In the micropore range the maximum of pore size distribution is located at about 0.53–0.58 nm with a broad tail from the side of larger pores. Thus, the location of this maximum fits very well to the diameter of 10 MR channels in ferrierite [19]. The intensity of this maximum is significantly reduced comparing to the Ti-FER samples. In the mesopore range the maximum of pore size distribution is centered in the range of 3.7–5.1 nm. In the case of the sample with the lowest titanium content, Ti-ITQ-6-100, a sharp maximum is located at 3.7 nm with a broad tail from the side of larger pores. For other samples of this series much broader peak of mesopore size distribution was observed. Textural parameters, presented in Table 2, show significantly higher BET surface areas of the Ti-ITQ-6 samples compared to the Ti-FER series, especially in the case of zeolite with the highest titanium content (Ti-ITQ-6-12.5). Moreover, delaminated zeolites are characterized by the total pore volume of about 4–6 times larger and micropore volume significantly reduced compared to the Ti-FER samples. These results clearly show the successful delamination of Ti-PREFER resulting in Ti-ITQ-6 zeolites.

UV-vis-DRS method was used for determination the form and aggregation of titanium species introduced into zeolites. The original UV-vis-DR spectra and sub-bands obtained by their deconvolution are presented in Fig. 4. For the Ti-FER samples the intensive bands at about 220 nm, attributed to tetrahedrally coordinated Ti$^{4+}$ cations incorporated into the zeolite framework, are present (Fig. 4, left side). These bands result from the ligand-to-metal charge transfer within tetrahedral TiO$_4$ and O$_2$TiOH moieties incorporated into the zeolite framework [23–25]. Moreover, the less intensive bands above 220 nm can be attributed to extraframework titanium species, such as isolated Ti$^{4+}$ cations in the octahedral coordination (about 230–250 nm) and partially polymerized hexacoordinated Ti-species containing Ti–O–Ti bridges (about 260–320 nm) [5,15,26,27]. For the sample with the lowest titanium content, Ti-FER-100, the second band, at about 250 nm, is attributed to monomeric extraframework Ti$^{4+}$ cations in the octahedral coordination and partially polymerized hexacoordinated Ti-species containing Ti–O–Ti bridges, located in the extraframework positions [23–25]. The contribution of this subband to 290, 296 and 303 nm for Ti-FER-50, Ti-FER-25 and Ti-FER-12.5, respectively. This interesting effect is related to the formation of extraframework, partially polymerized, hexacoordinated Ti-species, in which the polymerization degree increased with an increase in titanium content.

In the case of the Ti-ITQ-6 samples the original bands were deconvoluted into two subbands (Fig. 4, right side), similar to the Ti-FER samples. The first subband at about 225 nm is assigned to tetrahedrally coordinated Ti$^{4+}$ cations incorporated into the zeolite framework, while the second band, at 255–285 nm, is related to a partially polymerized hexacoordinated Ti-species containing Ti–O–Ti bridges, located in the extraframework positions [5,15,26,27]. The contribution of this subband in spectra increased with an increase in titanium content. Moreover, the shift of this maximum from 255 nm for Ti-ITQ-6-100 to 285 nm for Ti-ITQ-6-12.5 indicates the tendency to the formation of polymerized titanium species in the samples with the higher titanium content.

3.2. Catalytic studies

Zeolites of the Ti-FER and Ti-ITQ-6 series were studied as catalysts for oxidation of diphenyl sulfide (Ph$_2$S) to diphenyl sulfoxide (Ph$_2$SO) and diphenyl sulfone (Ph$_2$SO$_2$) in the presence of hydrogen peroxide (H$_2$O$_2$) as an oxidant. Apart from Ph$_2$SO and Ph$_2$SO$_2$ no other reaction

| Sample       | Si/wt% | Ti/wt% | Si/Ti | Sample       | Si/wt% | Ti/wt% | Si/Ti |
|--------------|--------|--------|-------|--------------|--------|--------|-------|
| Ti-FER-12.5  | 43.6   | 4.0    | 26    | Ti-FER-25    | 45.3   | 1.8    | 60    |
| Ti-FER-100   | 46.1   | 0.8    | 137   | Ti-ITQ-6-12.5| 44.6   | 2.8    | 38    |
| Ti-ITQ-6-12.5| 46.3   | 1.8    | 61    | Ti-ITQ-6-100 | 46.6   | 0.2    | 257   |
products were detected. Moreover, zeolitic samples were tested as catalysts for dimethyl sulfide (DMS) oxidation by H$_2$O$_2$. Dimethyl sulfoxide (DMSO) and dimethyl sulfone (DMSO$_2$) were the only detected reaction products. As it was shown in our previous paper the oxidation of Ph$_2$S in the absence of H$_2$O$_2$ was not effective [28].

Fig. 5 shows the results of the Ph$_2$S oxidation in the presence of the Ti-FER and Ti-ITQ-6 catalysts without (DARK) as well as with UV irradiation (LIGHT). As mentioned, Ph$_2$SO and Ph$_2$SO$_2$ were the only detected reaction products, thus the selectivity to Ph$_2$SO$_2$ can be determined by subtraction of the selectivity towards Ph$_2$SO from 100%. Conversion of Ph$_2$S depended on titanium content in the Ti-FER catalysts. In the case of the tests conducted without UV irradiation for the most effective catalyst of this series, Ti-FER-12.5, the Ph$_2$S conversion of about 80% was achieved after 4 h of the catalytic reaction. Other catalysts of this series were less active. The selectivities to Ph$_2$SO obtained for all catalysts of this series were above 94%. The oxidation of Ph$_2$S in the presence of the Ti-FER catalysts was significantly improved under UV irradiation (Fig. 5). The efficiency of diphenyl sulfide oxidation, similarly to catalytic tests without UV irradiation, increased with an increase in titanium content in the Ti-FER samples. Ph$_2$SO was the only reaction product during the first hour of the tests and afterwards the formation of small amounts of Ph$_2$SO$_2$ was detected.

The Ti-ITQ-6 samples were found to be much more effective catalysts of Ph$_2$S oxidation than the catalysts of the Ti-FER series, both in the tests conducted without and with UV irradiation. Similarly to the Ti-FER series, effectiveness of Ph$_2$S oxidation increased with an increase of the titanium content in the Ti-ITQ-6 catalysts. In the case of the catalytic tests conducted without UV irradiation 100% of diphenyl sulfide conversion was obtained only for the sample with the highest titanium content after 2 h of the catalytic reaction. Other catalysts of this series were less catalytically active than Ti-ITQ-6-12.5, however presented significantly higher activity than the analogous catalysts of the Ti-FER series. The selectivity to Ph$_2$SO and Ph$_2$SO$_2$ depended on titanium content in the Ti-ITQ-6 catalysts. For the Ti-ITQ-6-12.5 catalyst, Ph$_2$SO$_2$ was the only product of Ph$_2$S oxidation after 3 h of the catalytic reaction. Also for other catalysts of this series the selectivity to Ph$_2$SO$_2$ was significantly higher compared to the analogous Ti-FER samples. Efficiency of Ph$_2$S oxidation in the presence of the Ti-ITQ-6 catalysts was significantly improved under UV irradiation (Fig. 5). In this case the complete Ph$_2$S conversion was obtained for the Ti-ITQ-6-25 and Ti-ITQ-6-12.5 catalysts during less than 1 h of the catalytic reaction.

### Table 2

| Sample          | $S_{BET}$/m$^2$/g | $V_{MIC}$/cm$^3$/g | $V_{TOT}$/cm$^3$/g |
|-----------------|------------------|-------------------|-------------------|
| Ti-FER-12.5     | 378              | 0.136             | 0.228             |
| Ti-FER-25       | 384              | 0.139             | 0.222             |
| Ti-FER-50       | 380              | 0.139             | 0.220             |
| Ti-FER-100      | 400              | 0.149             | 0.185             |
| Ti-ITQ-6-12.5   | 940              | 0.046             | 1.266             |
| Ti-ITQ-6-25     | 642              | 0.036             | 0.823             |
| Ti-ITQ-6-50     | 721              | 0.056             | 1.092             |
| Ti-ITQ-6-100    | 729              | 0.046             | 1.069             |

Fig. 1. X-ray diffraction patterns of Ti-substituted Ti-FER (A) and Ti-ITQ-6 (B) zeolites with different Si/Ti ratio.

Fig. 2. N$_2$ adsorption-desorption isotherms of Ti-FER (intervals of 25 cm$^3$ g$^{-1}$) and Ti-ITQ-6 (intervals of 200 cm$^3$ g$^{-1}$) zeolites with different Si/Ti ratios.
titanium content and their catalytic activity is still present. The selectivity to Ph$_2$SO decreased, while selectivity to Ph$_2$SO$_2$ increased during the catalytic tests. This effect is more distinct for the catalysts with the higher titanium content, however 100% selectivity to Ph$_2$S was obtained in the presence of the samples with the lowest Ti-content – Ti-FER-100 and Ti-ITQ-6-100. Efficiency of DMS oxidation in the presence of both series of the catalysts was significantly improved under UV irradiation (Fig. 6). For all catalysts of both series the complete DMS conversion was obtained during 30 min of the catalytic test. In contrast to the DMS oxidation in DARK conditions, for the reaction conducted under UV irradiation there is a correlation between titanium content in the catalysts and their catalytic activity. The selectivity to DMS decreased by about 5% under UV irradiation comparing to the reaction conducted in DARK conditions (Fig. 6).

The efficiency in DMS conversion is significantly higher comparing to Ph$_2$S. The size of dimethyl sulfide molecules is significantly smaller compared to diphenyl sulfide and therefore DMS molecules can penetrate not only mesopores but also micropores of the Ti-FER samples. A slightly higher efficiency of the DMS conversion, observed for Ti-ITQ-6 catalysts, is possibly related to the faster rate of the internal diffusion of reactants in the hierarchical meso- and microporous structure of this series of the samples comparing to the slower internal diffusion in micropores in Ti-FER catalysts.

Comparing the results of the catalytic oxidation of Ph$_2$S (Fig. 5) and DMS (Fig. 6) obtained for the Ti-ITQ-6 catalysts, it can be seen that the selectivity of diphenyl sulfide oxidation to Ph$_2$SO$_2$ decreased, while selectivity to Ph$_2$SO$_2$ increased with the reaction time. On the other hand, the selectivities to DMSO and DMSO$_2$ were nearly the same during the catalytic tests. This interesting effect could be explained by different reaction mechanisms of Ph$_2$S and DMS oxidation. It seems that the diphenyl sulfide conversion in the presence of the Ti-ITQ-6 catalysts is a sequence of two consecutive oxidation steps: Ph$_2$S $\rightarrow$ Ph$_2$SO $\rightarrow$ Ph$_2$SO$_2$, while oxidation of DMS occurs in parallel directly to DMSO and DMSO$_2$.

Turnover frequency (TOF) values determined for the reactions of Ph$_2$S and DMS oxidation, conducted with and without UV irradiation, are compared in Table 3. It was assumed that all titanium cations play a role of catalytically active sites. TOF values were determined for the initial period of 30 min of the reactions. In general, TOF values increase with a decrease in titanium content in the samples (the only exception is Ti-ITQ-6-100). This effect could be related to the higher reactivity of Ti$^{4+}$ cations incorporated into the zeolite framework comparing to the extramolecular titanium species. Moreover, it was assumed that all titanium cations are accessible for the reacting molecules but in the case of extramolecular, more aggregated species this may not be met. In general, TOF values determined for the conversion of DMS are higher than for the Ph$_2$S conversion, what is possibly related to different internal diffusion rates of smaller DME and larger Ph$_2$S molecules. This same trend is observed for the conversion of large Ph$_2$S molecules in the presence of microporous Ti-FER zeolites (lower TOF values) and micromesoporous Ti-ITQ-6 samples (higher TOF values). Moreover, in the case of the reactions conducted under UV irradiation a significant increase in TOF values was observed.

As it was shown, the organic sulfides oxidation with H$_2$O$_2$ is possible without UV irradiation, however it is significantly less effective comparing to the process conducted under UV irradiation. The catalytic oxidation of various organic compounds over zeolites containing titanium has been reported in literature [29-33]. Ravinder et al. [29] postulated the formation of Ti-hydroperoxide complexes (=Ti-O–O–H) as a result of H$_2$O$_2$ interaction with titanium cations in titanium silicate molecular sieves. Similar results were reported for H$_2$O$_2$ interacting with Ti$^{4+}$ cations, present in the Ti-silicate framework, by Bordiga et al. [30] and Tozola et al. [31]. On the other hand, the formation of such reactive Ti-hydroperoxide complexes was reported not only for monomeric Ti$^{4+}$ cations in the zeolite framework but also for =Ti-O–Ti= pairs in...
TiAlPO-5 by Novara et al. [32]. Thus, possibly also small aggregates of TiO$_2$, present in the extraframework positions of zeolites, can participate in the catalytic oxidation of organic molecules. Chen et al. [33] showed that the white TS-1 powder became light yellow when TS-1 was immersed in aqueous solution of H$_2$O$_2$, indicating the formation of titanium-hydroperoxide complexes by the direct interaction of TS-1 with H$_2$O$_2$. A similar effect was also observed in this work for the samples of Ti-FER and Ti-ITQ-6 series, thus the formation of titanium-hydroperoxide complexes is postulated also for these catalysts.

Theoretical studies of the TS-1 catalyzed epoxidation of ethylene with H$_2$O$_2$ resulted in a conclusion that the O–O bond length in titanium-hydroperoxide complexes ($\equiv$Ti–O–O–H) is 1.521 Å, which represents a remarkable activation of the O–O bond compared to H$_2$O$_2$ molecule [34]. A high oxidation reactivity of titanium-hydroperoxide complexes was proven for sulfoxidation of thioethers [29], epoxidation of ethylene [34], oxidation of dibenzothiophene [35] and other organic compounds. Thus, oxidation of organic sulfides by H$_2$O$_2$ over the Ti-FER and Ti-ITQ-6 catalysts possibly includes the formation of...

Fig. 4. UV–vis DR spectra of the Ti-FER (left) and Ti-ITQ-6 (right) zeolites with different Si/Ti ratios.
highly reactive titanium-hydroperoxide complexes (\(\approx Ti-O-O-H\)), which effectively oxidize organic sulfides to sulfoxide and sulfones.

A comparison of the results of the catalytic tests conducted with and without UV radiation (Figs. 5 and 6) shows a significant intensification of organic sulfides oxidation by UV irradiation. Juan et al. [35] postulated two possible pathways of thioether oxidation by \(H_2O_2\) over TS-1 catalysts under UV radiation. The first mechanism involves the formation of titanium-hydroperoxide complexes (\(\approx Ti-O-O-H\)), which under UV radiation decompose to hydroxyl radicals (\(HO\)), reactive in the oxidation of organic sulfides. Karlsen and Schöffel [34] as well as Dae Lee et al. [36] postulated that \(HO\) radical can be formed much easier from titanium-hydroperoxide complexes than from \(H_2O_2\). The second mechanism is related to the presence of small oligomeric Ti–O–Ti–O–Ti species [35, 37]. Such small aggregated species were identified in the Ti-FER and Ti-ITQ-6 samples by UV-vis-DRS analysis (Fig. 4). Howe and Krisnandi [37] reported that electron transfer may occur between the Ti–O–Ti–O–Ti chains and guest species in the pores of Ti-containing zeolite, resulting in Ti\(^{3+}\) cations in such oligomeric species. It was reported that such species play a role of a single-site photocatalyst active in the formation of free radicals involved in polymerization of ethylene [37]. A similar activity in the formation of \(HO\) radicals from the reduction of \(H_2O_2\) (resulting in \(HO^+\) and \(OH^-\)) cannot be excluded. In order to verify the possible formation of hydroxyl radicals under UV irradiation, tests of terephthalic acid (TA) to hydroxyterephthalic acid (TAOH) oxidation by \(HO\) radicals were done in the presence of the most active catalysts of the Ti-ITQ-6 series. The reaction rate is a measure of the efficiency of hydroxyl radicals generation in the reaction mixture. Results of these studies, presented in Fig. 7, clearly show that \(HO\) radicals are intensively formed only under UV irradiation. Therefore, enhanced oxidation of organic sulfides in LIGHT conditions as a result of the hydroxyl radicals formation, according to one or both described mechanisms, is postulated. Moreover, \(HO\) radicals are well known as highly reactive, often regarded as the most effective, oxidants involved in photocatalytic processes. Because of the involvement of non-selective \(HO\) radicals, a decreased selectivity of sulfoxides formation under irradiation was observed.

4. Conclusions

Titanosilicate ferrierite (Ti-FER) and its delaminated form (Ti-ITQ-6), with the various Si/Ti molar ratios, were synthesized and tested as catalysts for diphenyl sulfide (Ph\(_2\)S) and dimethyl sulfide (DMS) oxidation with \(H_2O_2\) without and with UV irradiation. The main conclusions of these studies are:

1. Activity of the zeolitic catalysts in oxidation of Ph\(_2\)S and DMS, both without and with UV irradiation, increased with an increase in titanium content;
2. Conversion of Ph\(_2\)S was more effective in the presence of delaminated Ti-ITQ-6 catalysts than microporous Ti-FER. It is related to the possible internal diffusion of bulky Ph\(_2\)S molecules and products of its oxidation in mesopores of Ti-ITQ-6 catalysts. In the case of Ti-FER catalysts, due to their microporous structure, oxidation of Ph\(_2\)S possibly occurred only on the external surface of the zeolite grains;
3. Conversion of DMS was significantly more effective than Ph\(_2\)S, for both series of zeolitic catalysts, due to easy accessibility of micropores for small DMS molecule;
4. The conversion of organic sulfides was significantly intensified under UV irradiation, what was related to the UV induced decomposition of...
H$_2$O$_2$ on titanium centers, resulting in the formation of reactive hydroxyl radicals (HO$^\cdot$);
5. Based on the results of the catalytic studies it is postulated that conversion of Ph$_2$S in the presence of Ti-ITQ-6 catalysts is a sequence of two consecutive oxidation steps: Ph$_2$S $\rightarrow$ Ph$_2$SO $\rightarrow$ Ph$_2$SO$_2$, while DMS oxidation occurs in parallel directly to DMSO and DMSO$_2$.

**Declaration of competing interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

**Table 3**

| Sample        | TOF/h$^{-1}$ | Ph$_2$S (Dark) | DMS (Dark) | Ph$_2$S (Light) | DMS (Light) |
|---------------|--------------|----------------|------------|----------------|-------------|
| Ti-FER-12.5   | 13.8         | 34.1           | 14.2       | 38.3           |
| Ti-FER-25     | 16.2         | 76.6           | 17.0       | 85.1           |
| Ti-FER-50     | 21.1         | 176.2          | 34.5       | 191.5          |
| Ti-FER-100    | 23.0         | 452.0          | 53.6       | 766.1          |
| Ti-ITQ-6-12.5 | 49.8         | 50.3           | 54.5       | 54.7           |
| Ti-ITQ-6-25   | 50.2         | 77.0           | 82.6       | 85.1           |
| Ti-ITQ-6-50   | 85.8         | 269.7          | 104.2      | 306.4          |
| Ti-ITQ-6-100  | 30.6         | 536.3          | 61.3       | 766.1          |

**Fig. 6.** The results of catalytic oxidation of DMS by H$_2$O$_2$ conducted with (LIGHT) and without UV irradiation (DARK) in the presence of Ti-FER (left) and Ti-ITQ-6 (right) zeolites.

**Fig. 7.** Tests of terephthalic acid (TA) to hydroxyterephthalic acid (TAOH) conversion by $\cdot$OH radicals with (LIGHT) and without (DARK) UV irradiation ($\lambda > 320$ nm, 8.1 mW/cm$^2$) in the presence of the Ti-ITQ-6-12.5 and Ti-ITQ-6-25 catalysts.

**CRediT authorship contribution statement**

Marcelina Radko: Formal analysis. Małgorzata Rutkowska: Investigation, Methodology. Andrzej Kowalczyk: Formal analysis.
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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.micromeso.2020.110219.

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