The influence of ferrous ions on the efficiency of aqueous photocatalytic oxidation of 2-ethoxy ethanol

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Abstract. The complex influence of ferrous ions on the efficiency of aqueous photocatalytic oxidation (PCO) of 2-ethoxyethanol (2-EE) was examined. A drastic efficiency increase at lower concentrations of ferrous ions was observed to change to a sharp decrease at higher concentrations. An explanation was proposed for the observed phenomena based on the low sensitivity of the pollutant towards radical-oxidation reactions and the competitive adsorption of metallic ions and 2-EE on the TiO2 surface.

1. INTRODUCTION

2-Ethoxyethanol (2-EE, CH3-CH2-O-CH2-CH2-OH) is used as a solvent, a de-icing agent for runways and aircrafts, and an antifreeze jet fuel additive. This substance is often present in groundwater as a result of leakage, accidental spillages and uncontrolled disposal of fuels and de-icing liquids. It has negative effects on the kidneys, respiratory, and reproductive systems, and also acts as a depressant of the central nervous system [1]. The de-icing agents have been proven by earlier studies to be refractory against biodegradation and thus they accumulate and remain intact in groundwater for long times [2].

The photocatalytic oxidation (PCO) of organic pollutants is based on the action of positively charged holes on the semiconductor surface [3]. However, the holes may recombine rapidly with conduction band electrons, decreasing the PCO efficiency. Prolongation of the holes' lifetime should, therefore, increase the efficiency of PCO. For this purpose, multivalent metal ions may be added to the treated solutions, or, more precisely, to the photocatalyst surface, to scavenge electrons at the surface of the titanium dioxide, thus preventing electron-hole recombination and improving the oxidation performance.

This paper analyses the influence of ferrous ion additives on the PCO of 2-EE. Several previous studies [4-6] have reported higher PCO rates following the addition of small amounts of ferrous ions. Sucrose, carboxylic acids and textile azo dye were the organic pollutants in these studies. However, higher concentrations of ferrous ion reduced the decomposition rate significantly. A similar trend was observed with other multivalent metals influencing PCO of phenol and sucrose [7, 8]. Our previous studies [9] showed the positive role of ferrous ions at low concentrations in the efficiency of PCO of methyl tert-butyl ether (MTBE), although the PCO experiments with 2-EE, carried out in the same range of concentrations of Fe2+, indicated only a negative effect of iron ions. The results of more detailed investigation, given here, extend the knowledge of the influence of iron ions on pollutants of various types and show the complex character of the dependence of the PCO efficiency on the concentration of iron ions.

2. EXPERIMENTAL SECTION

Two 200-mL simple batch reactors with inner diameter 100 mm (evaporation dishes), aperture 40 m2 m-3, thermostatted at 20 ± 1 °C and mechanically agitated with magnetic stirrers were used in the PCO experiments: the reactor used for the PCO was called “active” and the other, containing no photocatalyst, was called “reference”. Both reactors were exposed to identical experimental conditions. The samples from the active reactor were compared to the reference samples to avoid complications caused by water evaporation. A UV-light source, Phillips TLD 15 W/05 low-pressure luminescent mercury UV-lamp with the emission maximum at 360-nm, was positioned horizontally over the reactors, providing irradiance of about 0.7 mW cm-2 measured by the optical radiometer UVX at a distance corresponding to the level of the free surface of the reactor.

The experiments were conducted with synthetic solutions at a concentration of 300 mg L-1 of 2-EE. The concentration of 2-EE was chosen to be consistent with its presence in polluted groundwater and experimental conditions applied previously [9]. All experiments were carried out at pH 3.0, adjusted with sulphuric acid. The treatment time, 24 h, was chosen to reduce...
the concentration of 2-EE below 50% of the residual concentration and was used in calculations of the process efficiency $E$ (see eq. (1)). All the experiments were carried out three times under identical conditions. The average deviation of data in parallel experiments did not exceed 5%.

The experiments were performed using titanium dioxide as Degussa P25. Shurry at 1 g L$^{-1}$ was used as a suspended catalyst option. The supported catalyst option was TiO$_2$ attached to buoyant hollow glass microspheres and to the surface of the glass plates. The hollow glass micro-spheres used in this study had an average diameter of 60 to 70 µm and a density 0.27 g cm$^{-3}$ (the product of LP-ImEx, Estonia). Titanium dioxide was attached to the surface of the micro-beads by the thermal method [10]: equal volumes of dry micro-spheres and aqueous suspension of titanium dioxide with a concentration of 1 g L$^{-1}$ were mixed by stirring and sonication for 30 min. The micro-spheres were then separated from the mixture by filtration with a membrane filter, heated to dryness at 120 °C and calcined at a temperature of 300 °C for 4 h. This procedure was repeated six times, which was found to be the optimum number of attachment operations [11]. Analogously, TiO$_2$ was attached to the surface of the glass plates (one side) in approximately equal amounts either by multiple submerging of the plates in the TiO$_2$ suspension with subsequent drying after each submersion, or by spraying the TiO$_2$ suspension over the surface of the plates and drying. TiO$_2$ attached to glass plates, submerged horizontally in the solution to be treated at a depth from 5 to 10 mm, was used in the PCO experiments. Since 2-EE does not absorb UV light at 360 nm, the depth of the submersion was not important within the indicated limits. This was confirmed by specially conducted experiments.

The decrease in the pollutant concentration was determined for 2-EE from the decrease in chemical oxygen demand (COD), measured by a standard method [12].

Adsorption experiments for oxygenated hydrocarbons and ferrous ion, introduced as sulphate, on the TiO$_2$ surface were conducted in the dark in closed flasks equipped with magnetic stirrers, thermostatted at 20 ± 1 °C and adjusted to pH 3.0. The amount of substances adsorbed was derived from the batch mass balance: the concentration of the dissolved substance was determined before and after adsorption. The points of the isotherms are the average of three experimental sets. The concentrations of iron ions were determined by a spectrophotometric phenantrolin method described in [12].

The PCO by-products of 2-EE, mainly carbonic acids, were qualitatively determined by the methods described in [13]. 200 mg of resorcinol was added to 5 ml of the sample in a test-tube. After the resorcinol had dissolved, 10 ml of concentrated sulphuric acid was carefully inserted into the bottom of the test-tube. The coloured rings corresponding to carbonic acids appeared in the following sequence: the blue ring indicating oxalic acid appeared at the liquids’ interface, the red ring below the blue one indicated glycolic acid, the orange ring above the blue one corresponds to formic acid. The rings were observed distinctively and repeatedly with both synthetic solutions of the acids and the samples of PCO treated 2-EE solutions. Oxalic acid was also separated from the sample as calcium oxalate and quantitatively determined by means of titration with potassium permanganate. Acetic acid was qualitatively determined as red crystals of iron (III) acetate precipitated as a result of addition of iron (III) chloride to the sample. These identifications were available only in the absence of iron ions in the treated samples, i.e. when the samples were treated without iron ions added.

3. RESULTS AND DISCUSSION

The performance of PCO with artificial radiation sources was characterised by the process efficiency $E$. The efficiency $E$ is defined as the decrease in the amount of pollutant divided by the amount of energy reaching the surface of the treated sample (eq. (1)). The efficiency values were calculated after a period of time equal to the treatment time (24 h)

$$E = \frac{\Delta c \cdot V \cdot 1000}{I \cdot s \cdot t}$$

where $E$—PCO efficiency, mg W$^{-1}$ h$^{-1}$; $\Delta c$—the decrease in the concentration of the pollutant, mg O$_2$ L$^{-1}$ for COD of 2-EE; $V$—the volume of the sample to be treated, L; $I$—irradiance, mW cm$^{-2}$; $s$—irradiated area, cm$^2$; $t$—treatment time, h.

Both ferrous and ferric ions were tested in the experiments. Sulphate was chosen as the counter-ion due to its low inhibitive effect on PCO efficiency, observed with 2-EE in [9]. The results of the experiments are shown in Figure 1. One can see a similar trend in the dependence of PCO performance on the concentrations of both ionic species. Ferric ions were seen to have the same effect as ferrous ions, which may be explained with a dynamic equilibrium established between ferrous and ferric ions at the UV-irradiated TiO$_2$ surface and, probably, in its closest vicinity, as described in [14].

Ferrous and ferric ions appeared to play a significant role in PCO of 2-EE. The addition of small amounts of Fe$^{2+/3+}$-ion, up to 0.09 mM, resulted in a drastic, up to 60%, increase in the process efficiency (Figure 1). With further increase in the concentration of Fe$^{2+/3+}$, the PCO efficiency decreased dramatically. Therefore, the experiments were conducted with ferrous ions the characterisation of which at $t = 0$ and in aqueous solution was far simpler. During the PCO process, the dynamic equilibrium Fe$^{2+}$/Fe$^{3+}$ took place very fast.

A similar dependency pattern, in which the PCO efficiency of 2-EE had its maximum with a Fe$^{2+/3+}$
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Figure 1. The efficiency of PCO of 2-ethoxyethanol in TiO₂ slurry vs. concentration of Fe²⁺ (Δ) and Fe³⁺ (•) ions at pH 3.0.

Figure 2. Efficiency of PCO of 2-ethoxyethanol at 0.09 mM of Fe²⁺ with TiO₂ application modes: 1-TiO₂ Degussa P25 suspension (1 g L⁻¹); 2-TiO₂ sprayed on the glass plate, 0.6 mg cm⁻²; 3-TiO₂ settled at the glass plate, 0.7 mg cm⁻²; 4-TiO₂ attached to hollow glass micro-spheres, the amount of buoyant catalyst in the reactor 50 g m⁻²; 5-same as 4 at 25 g m⁻².

Figure 3. The efficiency of PCO of MTBE in TiO₂ slurry vs. concentration of Fe²⁺-ions at pH 3.0.

of Fe²⁺-ions was followed by a decrease with further addition of ferrous ions (Figure 3) [9]. However, the maximum efficiency for MTBE was observed at the Fe²⁺ concentration of 1 mM, which is about ten times bigger than that for 2-EE. A further increase in the concentration of Fe²⁺-ions resulted in a gradual increase of PCO efficiency, which was not observed with 2-EE. This may be explained by the difference in the PCO mechanism of these two substances and their different adsorption properties at the surface of titanium dioxide.

The difference in the behaviour of 2-EE and MTBE can be explained by a dual mechanism [3]: (1) the adsorption of the pollutant by the surface of the titanium dioxide is followed by a direct subtraction of the pollutant’s electrons, i.e. oxidation, with positively charged holes; (2) oxidation with hydroxyl radicals takes place at the catalyst surface or in its vicinity. Both reactions proceed simultaneously, although positively charged holes have an oxidation potential about 1.25 times bigger than OH-radicals [15]. Which mechanism dominates in PCO depends on the chemical and adsorption properties of the pollutant.

The promotion of the OH-radical was less effective in oxidation of ethylene glycol derivatives [16, 17]. The increasing PCO efficiency for 2-EE with the addition of ferrous ions in small, below 0.09 mM, concentrations may then be explained by the partial occupation of adsorption sites with Fe²⁺-ions, overpowered, however, with the electron scavenging by iron ions, extending the lifetime and thus the oxidation performance of the positively charged holes. A further increase in the concentration of metallic ions results in the blockage of adsorption sites with a resultant drastic decrease in PCO efficiency. The formation of hydroxyl radicals has little effect on the overall oxidation rate.

Methyl tert-butyl ether is less resistant towards oxidation with a hydroxyl radical. For example, oxidation methods, such as ozonation, exhibited poor
performance in the abatement of MTBE, although introduction of OH-radical promoters significantly enhanced oxidation [18]. This makes both PCO mechanisms effective in PCO of MTBE. Small concentrations of ferrous ions therefore enhance the direct oxidation of MTBE with positively charged holes, and also the formation of OH-radicals. Radical oxidation reactions contribute to the overall oxidation rate successfully until a certain concentration of metallic ions, 1 mM, is achieved (Figure 3). Above this concentration, the formation of OH-radicals is probably obstructed by a "short-circuit" phenomenon, described in [4]: the blockage of the TiO2 adsorption sites with ferrous ions results in reduced generation of OH-radicals. The increase of the PCO efficiency with further increasing ferrous ion concentration may be explained by increased OH-radicals formation in the bulk solution initiated by the UV-irradiated Fe2+ ions. It was also observed that MTBE oxidized to some extent under UV-irradiation in aqueous solutions containing Fe2+ ions with no TiO2 present, although this reaction was beyond the scope of the present paper. In the TiO2-free experiments the oxidation rate increased with increasing concentration of Fe2+ ions. The authors presume that oxidation of MTBE with UV-irradiated Fe2+ may be enhanced in the presence of TiO2. However, further experimental studies are necessary to confirm this statement.

To support the hypothesis concerning the role of ferrous ions in PCO performance determined by the adsorption properties and the sensitivity of the target compounds towards radical attacks, the impact of the Fe2+ ions on the adsorption of a pollutant by TiO2 was studied in a series of adsorption experiments. The ferrous ions content dependency of adsorption of 2-EE from a solution with an equilibrium concentration of 2-EE 300 mg L−1 by the TiO2 surface was determined experimentally. As one can see from Figure 4, the surface concentration of 2-EE steadily decreased within the Fe2+ ions equilibrium concentration range attributable to the peak zone of the PCO efficiency (Figure 1).

An analogous dependence of the equilibrium surface concentration of MTBE was also observed: the MTBE surface concentration in equilibrium with an MTBE solution of 100 mg L−1 decreased from 0.9 mmol g−1 at zero concentration of Fe2+ ions to 0.5 mmol g−1 at 1 mM and 0.1 mmol g−1 at 10 mM of Fe2+ ions (Figure 5). This explains the difference between the critical concentration values of Fe2+ ions for 2-EE and MTBE: the adsorption of MTBE was less affected by the presence of Fe2+ ions than 2-EE.

The last observation may be explained by differences in adsorption of 2-EE and MTBE. The carbonic acids determined qualitatively as PCO products of 2-EE include oxalic, acetic, formic and glycolic acids. The possible route of the reactions is outlined in Figure 6. The presence of acetic and especially glycolic acids among the oxidation by-products indicate the adsorption of 2-EE with its etheric oxygen and, therefore, electrophilic attack of positively charged holes to the etheric bond. This results in fracture of the 2-EE molecule into two fragments, the acetic and glycolic acids (reaction A). The formation of oxalic acid appears to be possible only when both oxygen atoms in the 2-EE molecules are adsorbed on the TiO2 surface and are under simultaneous attack of positively charged holes (reaction B). A further possible precursor of oxalic acid is glycolic acid, when the alcoholic group is oxidised (reaction C). Hypothetically, acetic acid may also be a product of 2-EE stepwise oxidation starting from the alcohol group only (reaction D) with formic acid as a product of the last reaction. The ethoxy carbonic acids were not identified and were only shown as hypothetical by-products in rectangular brackets. Further oxidation of reactive by-products, such as formic acid also observed in samples, may proceed with OH-radicals formed at the TiO2 surface, although their formation may be

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**Figure 4.** The surface concentration of 2-EE on TiO2 versus Fe2+ ions equilibrium concentration at pH 3.0 (20 °C).

**Figure 5.** The surface concentration of MTBE on TiO2 versus Fe2+ ions concentration at pH 3.0 (20 °C).
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Figure 6. Outline of aqueous photocatalytic oxidation reaction of 2-ethoxy ethanol on TiO₂.

The adsorption of ferrous/ferric ion on the surface of titanium dioxide was also studied. The results shown in Figure 7 are consistent with the adsorption equilibrium of MTBE: the equilibrium aqueous concentration of Fe²⁺ for the minimum adsorption of MTBE approximately coincides with the maximum adsorption of Fe²⁺, which shows a direct connection between the occupation of adsorption TiO₂ sites with iron ions and the performance of PCO—the minimum PCO efficiency for MTBE was observed around this concentration of Fe²⁺ [9].

4. CONCLUSIONS

The complicated influence of the addition of multivalent metallic ions to aqueous solutions of groundwater pollutant, 2-ethoxy ethanol (2-EE), was studied. An optimum concentration range of ferrous ions was observed.

The hypothesis was supported that the optima are the result of a dual mechanism of oxidation at the surface of the TiO₂ photocatalyst, including the direct suppression in acidic media. The OH-radical, although less reactive with 2-EE, may also participate in oxidation reactions initiated by positively charged holes at the TiO₂ surface.
oxidation of adsorbed pollutants with positively charged holes and the radical oxidation: the PCO of 2-EE resistant to radical oxidation practically stopped when the positively charged holes at the TiO₂ surface were blocked by ferrous/ferric ions. The effective PCO of MTBE at high concentrations of Fe²⁺-ions observed in our previous work may be explained by the attack by hydroxyl radicals more efficient with MTBE.

ACKNOWLEDGMENTS

The authors express their gratitude to the Estonian Science Foundation (grant 5899), Centre for International Mobility (CIMO), Finland, and the Academy of Finland (Project 208134) for financial support.

REFERENCES
