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# Automotive fleet repair facility wastewater treatment using air/ZVI and air/ZVI/H<sub>2</sub>O<sub>2</sub> processes

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**Abstract:** Advanced automotive fleet repair facility wastewater treatment was investigated with Zero-Valent Iron/Hydrogen Peroxide (Air/ZVI/H<sub>2</sub>O<sub>2</sub>) process for different process parameters: ZVI and H<sub>2</sub>O<sub>2</sub> doses, time, pH. The highest Chemical Oxygen Demand (COD) removal efficiency, 76%, was achieved for ZVI/H<sub>2</sub>O<sub>2</sub> doses 4000/1900 mg/L, 120 min process time, pH 3.0. COD decreased from 933 to 227 mg/L. In optimal process conditions odor and color were also completely removed. COD removal efficiency was increasing with ZVI dose. Change pH value below and over 3.0 causes a rapid decrease in the treatment effectiveness. The Air/ZVI/H<sub>2</sub>O<sub>2</sub> process kinetics can be described as d[COD]/dt = -a [COD]t<sup>m</sup>, where 't' corresponds with time and 'a' and 'm' are constants that depend on the initial reagent concentrations. H<sub>2</sub>O<sub>2</sub> influence on process effect was assessed. COD removal could be up to 40% (560 mg/L) for Air/ZVI process. The FeCl<sub>3</sub> coagulation effect was also evaluated. The best coagulation results were obtained for 700 mg/L Fe<sup>3+</sup> dose, that was slightly higher than dissolved Fe used in ZVI/H<sub>2</sub>O<sub>2</sub> process. COD was decreased to 509 mg/L.

#### Introduction

Automotive fleet repair facility wastewater contains many persistent and toxic petroleum organic compounds:wasteoils, petrol components and heavy metals (Makowska and Mazurkiewicz 2016, Piecuch et al. 2015, US EPA 1991). They are refractory to biodegradation. Thus, conventional biological treatment may no longer be considered as satisfactory treatment process.

Advanced oxidation processes (AOP), such as Fenton process, are promising treatment methods for wastewater containing hardly biodegradable and potentially toxic organic compounds. They consist in the effective production of free radicals which oxidize organic pollutants. Equation 1 is recognized as Fenton reaction and implies the oxidation of Fe<sup>2+</sup> to Fe<sup>3+</sup> ions to decompose H<sub>2</sub>O<sub>2</sub>into 'OH. It is usually considered as the core of the Fenton chemistry. Furthermore, other reactions must be considered to understand the whole process(Babuponnusami and Muthukumar 2014).

$$Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH^- + OH - k_1 = -40 - 80 L/(mol \cdot s)$$
 (1)

$$Fe^{3+} + H_2O_2 \rightarrow Fe^{2+} + O_2H + H^+ \qquad k_2 = -9.1 - 10^{-7} L/(\text{mol} \cdot \text{s})$$
 (2)

$$Fe^{2+} + OH \rightarrow Fe^{3+} + OH - k_s = -2.5 - 5.10^8 L/(mol \cdot s)$$
 (3)

$$Fe^{2+} + O_2H \rightarrow Fe^{3+} + OH_2$$
,  $k_A = 0.72 - 1.5 \cdot 10^6 L/(mol \cdot s)$  (4)

$$Fe^{3+} + {}^{\star}O_{2}H \rightarrow Fe^{2+} + O_{2} + H^{+}$$
  $k_{5} = 0.33 - 2.1 \cdot 10^{6} L/(mol \cdot s)$  (5)

As shown in (Eq. 2), the generated  $Fe^{3+}$ ions can be reduced by reaction with excess of  $H_2O_2$  to form again  $Fe^{2+}$ ion. This reaction is called Fenton-like reaction, it is slower than Fenton reaction, and allows for  $Fe^{2+}$  regeneration in an effective cyclic mechanism. Equations (2–5), represent the limiting steps in the Fenton chemistry since  $H_2O_2$  is consumed and  $Fe^{2+}$  ionsare regenerated from  $Fe^{3+}$  ion through these reactions.

The application of homogeneous Fenton process to treat large quantity of water may produce large amount of sludge in the final neutralization/coagulation step. Usually Fe<sup>2+</sup> ions are introduced to wastewater as FeSO<sub>4</sub> solution, which increases salinity usually above the standard 1000ml/L and 500 mg/L of chlorides and sulfates (VI) respectively (Directive of the Minister of the Environment, Journal of Laws of 18 November 2014, Item 1800). In order to avoid these disadvantages, zero valent iron (ZVI) has appeared as an effective heterogeneous catalyst for wastewater treatment. ZVI and other iron based catalysts were used inter alia for the treatment of dye (Chang et al. 2006, Devi et al. 2009, Greic et al. 2012, Moon et al. 2011, Weng et al. 2013, Zhang et al. 2014, Yang et al. 2015a, Yang et al. 2015b), nitrobenzene (Dong et al. 2010), 2,4-dinitroanisole (Shen et al. 2013), pharmaceutical compounds (Bautitz et al. 2012, Segura et al. 2013), TNT wastewater (Barreto-Rodrigues et al. 2009), phenols and chlorophenols (Kim et al. 2011, Shimizu et al. 2012), pesticides (Fjordboge et al. 2013), BPA (Xi et al. 2014) or coking wastewater (Lai et al. 2007), palm oil mill effluent (Taha and Ibrahim 2014), nitrites reduction (Fatemina et al. 2013, Suzuki et al. 2012), oil sands reclamation (Pourrezaei et al. 2014), and surfactants removal (Martins et al. 2014).

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Ferrous and ferric saltsare low cost material widely used in wastewater treatment as coagulant or as a component of Fenton reagent. Adding ZVI in aqueous phase has proved to be a fairly fast process to reduce various organic compounds according to the dissolution of Fe<sup>2+</sup> ions and electrons transfer to react with the organic molecules on the iron surface (Cao et al. 1999, Chen et al. 2001). In heterogeneous Fenton reaction, oxidation of ZVI provides an alternative means of inducing Fenton oxidation as shown in (Eq. 6–11):

$$Fe^0 + O_2 + 2H^+ \rightarrow Fe^{2+} + H_2O_2$$
 (6)

$$Fe^0 + 2H^+ \rightarrow Fe^{2+} + H_2$$
 (7)

$$Fe^{3+} + Fe^0 \rightarrow 3Fe^{2+}$$
 (8)

$$Fe^0 + H_2O_2 + 2H^+ \rightarrow Fe^{2+} + 2H_2O$$
 (9)

$$Fe^0 \to Fe^{2+} + 2e \tag{10}$$

Organic pollutant + Fe $^0$   $\rightarrow$  Reduced organic pollutant + Fe $^{2+}$  (11)

The objective of this study was to provide additional knowledge regarding the treatment of automotive fleet repair facilities wastewater. The studies were especially aimed at the estimation of the efficiency of (i) Air/ZVI,Air/ZVI/H $_2$ O $_2$  and coagulation process, (ii) estimation of the stirring method and pH influence on efficiency of Air/ZVI/H $_2$ O $_2$  process, and (iii) description of its kinetics.

#### Materials and methods

The raw wastewater was collected in April 2014 from automotive fleet repair facility, located in Warsaw, Poland. All types of wastewater produced in facility are collected in equalization tank and thentransferred to emulsion separator. Samples were collected after emulsion separation in the separator. All studies concerning the wastewater treatment were carried out within 72 h after collection of samples. After wastewater collection, the samples were acidified to pH 3.0 and refrigerated at 4°C until analysis to prevent biochemical decomposition processes.

The following parameters were determined according to the EN or ISO standards: COD, COD after 30 min sedimentation (COD $_{sed}$ ) and COD dissolved (for sample filtered through 0.45  $\mu$ m filters) COD $_{dis}$ (ISO 6060), total suspended solids (TSS) (EN 872), pH (EN ISO 10523), turbidity (EN ISO 7027) and specific conductivity (EN 27888).

Wastewater after 30 min of sedimentation, for easily settleable solids (ESS) removal, were treated by Air/ZVI, Air/ZVI/H<sub>2</sub>O<sub>2</sub> or coagulation process.

Air/ZVI and Air/ZVI/H<sub>2</sub>O<sub>2</sub> experiments were carried out in 0.7L cylindrical reactor filled with 0.5L of sample. Four doses of ZVI were used (1000, 2000, 3000, 4000 mg/L). 30% H<sub>2</sub>O<sub>2</sub> solution was added to wastewater in ZVI/H<sub>2</sub>O<sub>2</sub> process. H<sub>2</sub>O<sub>2</sub>/COD mass ratio was equal to 0.5, 1, 1.5. Based on density and percentage concentration data, H<sub>2</sub>O<sub>2</sub> volumes added were 1.2, 2.4 and 3.6 mL respectively. All samples were stirred by magnetic or ultrasonic stirrer. Magnetic stirring was very intensive. Stirring speed was set as 1000rpm (Heidolph MR3000). Studies on ultrasonic stirring were performed in ultrasonic bath SONIC 5 630 W (Polsonic). Ultrasound power

used was  $2\times320\mathrm{W}$  and 40 kHz.The pH was adjusted to 3.0. During studies on pH influence, the pH was adjusted to 2.0, 4.0, 5.0 or 6.0. After preselected times (15, 30, 60, 120 min) the process was stopped by increasing the pH to 9.0 with 1.1 M NaOH. The sample was left overnight for sedimentation of iron precipitate and  $\mathrm{H_2O_2}$  decomposition. Iodometric test on  $\mathrm{H_2O_2}$  removal was performed, and after that COD was determined.

Studies on coagulation were performed in 0.7L reactor with magnetic stirrer (Heidolph MR3000). The reactor was filled with 0.5 L of sample. The wastewater was coagulated with FeCl<sub>3</sub> at pH 9.0. The coagulant doses wereinwide range (50 to 700 mg Fe<sup>3+</sup>/L). The samples were stirred for 5 min for fast, followed by 10 min for the slow stirring. The sample was left overnight for sedimentation of iron precipitate. After that COD was determined.

All reagents used in this study were manufactured by POCH and were of puriss p.a. grade.

For ZVI/ $\rm H_2O_2$  process, the efficiency of  $\rm H_2O_2$  was evaluated according to the Kang and Hwang equation (Eq. 12) (Kang and Hwang 2000):

$$\eta (\%) = \{\Delta COD/(0.4706 \cdot [H_2O_2])\} \cdot 100\%$$
 (12)

where  $\Delta COD$  is the decrease of the COD value (mg/L), 0.4706 is a constant factor for the amount of COD theoretically removed (mg) by 1 mg of  $H_2O_2$ , and  $[H_2O_2]$  is the hydrogen peroxide dose (mg/L).

#### Results and disscusion

#### Raw wastewater

It must be pointed out that all experiments in this work were carried out with real wastewater, without any dilution or other pre-treatment. The parameters of wastewater are shown in Table 1. All parameters except COD were determined after sedimentation. 30 min sedimentation for ESS removal allows to remove 9.9% of initial COD. Suspended solids left after sedimentation contributed to a 185 mg/L COD value. All treatment processes were performed on wastewater after initial sedimentation and Easily Settleable Solids (ESS) removal.

Raw wastewater wascharacterized by a moderate COD value (in comparison with other industrial wastewater sources, that could generate wastewater with COD even over 100 000 mg/L), but unacceptable color and odor. Based on a known source of wastewater and the smell, it could be suspected that the compounds contained in wastewater are, among others, components of fuels. Crucial for this type of wastewater treatment is to ensurethe color and odorremoval to an acceptable level.

Table 1. Raw wastewater parameters

Parameter	Unit	Value
COD	mg/L	1035
COD <sub>sed</sub>	mg/L	933
COD <sub>dis</sub>	mg/L	748
TSS	mg/L	640
рН	_	7.8
Conductivity	mS/cm	1.241
Colour	_	Dark green
Odor	_	Oily, strong

#### Air/ZVI and Air/ZVI/H2O2process

Results of Air/ZVI and Air/ZVI/H<sub>2</sub>O<sub>2</sub>processes are shown in Figures 1–4.

The COD removal was increasing with iron dose in a twostage pattern. At first sharp decline, followed by a further small decrease in time, was observed. COD during ZVI process rapidly decreased in 15 min from 933 mg/L to 733-600 mg/L. The decrease was significant in the case of 4000 mg/Lcompared with 1000 mg/L ZVI dose. The COD value increased slowly at time 15 to 60 min for different amount of ZVI. Finally, the second decrease at time 60 min to 120 min was observed. After 15 min COD minimally decreased to 773 mg/L (17%), while after 120 reached the maximal value of 560 mg/L (40% decreasing) for the highest dose. It might be due to some compounds transformation. On the one hand, radical oxidation leads to organic compounds rearrangements and partial decomposition but, on the other hand, coagulation persistent, more polar, compounds could be created. As a result of those two opposite processes, no significant COD removal effect was observed. For longer process times coagulation persistent compounds were decomposed and new ones were not created. Process effect increased with increasing ZVI dose, which was consistent with the studies of Lai et al. (Lai et al. 2007). The oxidation-reduction ability of Air/ZVI could transform the structures and characteristics of some compounds. Air/ZVI is more efficient in eliminating low molecular weight compounds, but there were still some residual contaminants which could

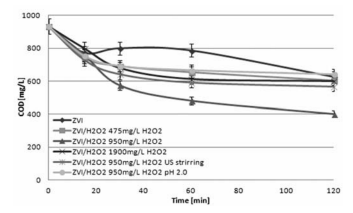


Fig. 1. COD changes for Air/ZVI and Air/ZVI/H $_2$ O $_2$  processes for 1000 mg/LZVI dose

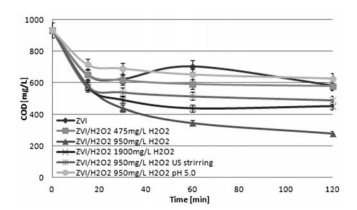


Fig. 3. COD changes for Air/ZVI and Air/ZVI/H<sub>2</sub>O<sub>2</sub> processes for 3000 mg/LZVI dose

not be removed by this process (Lai et al. 2007, Cao et al. 1999, Chen et al. 2001).

ZVI addition declined the color and odor significantly. It was observed that color and odor removal increased after the addition of  $\rm H_2O_2$ . The color and odor removal efficiencies incremented promptly to the first 15 min then increasing more with the passage of time to 120 min. In this study, the optimal deodorization and decolorizationwere consistent with some other studies (Cao et al. 1999, Chen et al. 2001). The Air/ZVI process could be used for the decolorization of wastewater, but the COD efficiency was too low to justify its use. To overcome this, Air/ZVI/ $\rm H_2O_2$  process, that was similar to a Fenton process, was examined.

Air/ZVI/ $\mathrm{H_2O_2}$  system was more effective than Air/ZVI one. There must be a balance between the concentration of  $\mathrm{H_2O_2}$  and ZVI. It was found that the result for Air/ZVI/ $\mathrm{H_2O_2}$  COD/ZVI ratio 1:1 were better than 1:0.5 and 1:2 (Figures 1–4) regardless or ZVI dose. Figures 1–4 show COD rapid decrease at time 0 min to 30 min. The amount of decrease was significant in the case of 4000 mg/L compared with 1000 mg/L ZVI dose. For longer times COD was still decreasing, but the rate was lower. After 120 min COD reached the value of 227 mg/L (76% decreasing) for the highest ZVI dose.

In comparison with other studies (Deng and Engelhardt 2006, Gogate and Pandit 2004, Lau et al. 2001, Zhang et al. 2005) sludge settling characteristic was excellent at the COD/H<sub>2</sub>O<sub>2</sub> ratio 1:1, but became poor for 1:2 ratio. It seemed

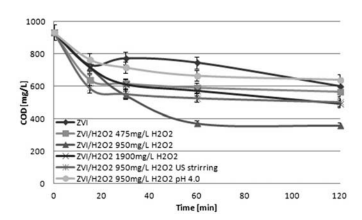


Fig. 2. COD changes for Air/ZVI and Air/ZVI/H $_2$ O $_2$  processes for 2000 mg/LZVI dose

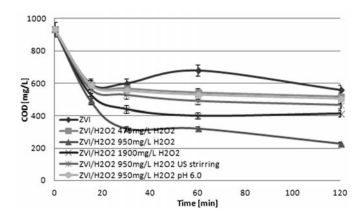


Fig. 4. COD changes for Air/ZVI and Air/ZVI/H $_2$ O $_2$  processes for 4000 mg/LZVI dose

that excessive  $\mathrm{H_2O_2}$ , when decomposed, would produce oxygen bubbles that made sludge settling difficult (Lau et al. 2001, Zhang et al. 2005). In addition, residual  $\mathrm{H_2O_2}$  might inhibit downstream chemical treatment. When  $\mathrm{COD/H_2O_2}$  ratio was 1:0.5 residual ferrous iron was observed, with the bluish green effluent (probably originated by  $\mathrm{Fe^{2^+}}$  compounds such as  $\mathrm{Fe(OH)_2}$ ). Iron sludge required further treatment (Deng and Engelhardt 2006, Gogate and Pandit 2004). Therefore  $\mathrm{COD/H_2O_2}$  ratio 1:1 was chosen for further experiments.

### Comparison between dissolving of oxygen by ultrasound and magnetic stirring in Air/ZVI/H<sub>2</sub>O<sub>2</sub> processes

Ultrasonic treatment involves the generation of 'OH by acoustic cavitation. This phenomenon is produced through compression and expansion cycles of US waves ()))), which result in the growth and collapse of the gas bubbles and the production of local areas of high energy and pressure. These conditions are so extreme that they are capable of breaking up the entrapped molecules within the bubbles, such as vaporized water or dissolved gases (e.g.), to produce radicals, mainly 'OH and 'OH, (Eq. 13–17).

$$H_2O + ))) \rightarrow H + OH$$
 (13)

$$O_2 + ))) \rightarrow 2O \tag{14}$$

$$H_2O + O \rightarrow 2 \cdot OH$$
 (15)

$$H + O_2 \rightarrow O + OH$$
 (16)

$$2 \cdot H + O_2 \rightarrow O + \cdot OH_2 \tag{17}$$

These radicals are capable of initiating or promoting many fast oxidation reactions, leading to the degradation of more hydrophobic compounds at the bubble–bulk solution interface and to a lesser extent, of more hydrophilic species via 'OH diffused or ejected into the bulk solution(Rubio-Clemente et al. 2014).

However, the magnetic stirring is more efficient in wastewater treatment than ultrasonic, as shown inthe Figures 1–4. Oxygen is paramagnetic and can be effectively introduced to wastewater by a magnetic field(Kim et al. 2011, Zhang et al. 2005). Reactions of oxygen with dissolving Fe, creation of H<sub>2</sub>O<sub>2</sub> are the most important comparing differences in COD. Where the consumption of dissolved oxygen by ZVI reaction (Eq. 6) was also enhanced in the presence of magnetic field, the competing reaction of H<sub>2</sub> production (Eq. 7) from proton reduction was retarded. Since the ZVI induced oxidation is mainly driven by the in situ generated 'OH (Kim et al. 2011), ZVI constantly consumes dissolved oxygen (Eq. 6). Under condition of weak reaeration (ultrasonic stirring) ZVI reduces protons according to (Eq. 7).

Shimizu et al. (2012) confirmed that in oxygen saturated wastewater 'OH generation is the same with and without aeration. Afterward the 'OH generation without aeration was significantly suppressed due to the shortage of dissolved oxygen. It may be concluded that 'OH generation was significantly related to the dissolved oxygen concentration. Because of that oxygen introduction mechanism is crucial for the purification effect. Ultrasonic stirring is less effective than magnetic stirring, so even though the reactions (13–17)

take place, they have less effect on the process then dissolved oxygen (Eq. 6).

Moreover Taha and Ibrahim (2014) found that for palm oil mill effluent treated withZVI,sonification intensity and duration time have a weak correlation with Fe<sup>2+</sup> production and COD decrease. They found by referring to shell – core concept, that at pH 3.0 and 4.0 only a small amount of Fe<sup>2+</sup> was in aqueous solution resulting from partially dissolved shell.

#### Effect of pH

Experiments were also performed to assess the generation of 'OH for previously foundedoptimal ZVI/ $H_2O_2$  doses. The slower rate of degradation at lower pH is also due to the excess  $H^+$  ions present in the solution acting as 'OH scavenger according to reaction (Eq. 18) (Devi et al. 2009).

$$H^+ + {}^{\bullet}OH + e^- \rightarrow H_2O$$
 (18)

While the fresh surfaces were kept, the adsorption ability and reactivity of the ZVI might be high. At the initial stage, pollutant molecules could easily transport and be oxidized by 'OH formed near Fe<sup>o</sup> surface and/or adsorb to Fe<sup>o</sup> surface.

At lower pH,  $Fe^0$  oxidizes to  $Fe^{2+}$  in aqueous acidic conditions due to the dissolution. present in the solution decomposes formed by the follow reaction (Eq. 6, 1) to generate OH.

Therefore, at lower pH the Fenton reaction generating 'OH may occur near the ZVI powder surface and dominate the overall removal of color and odor (Shimizu et al. 2012).

In this study, it was found that the optimal value for COD was reached at pH 3.0. It means the increasing value of pH effect slightly for decreasing COD, which was consistent with other studies (Devi et al. 2009, Fan et al. 2009). They discovered that although the cumulative 'OH production at pH 2.0 was about a half of that at pH 3.0, the contribution of the Fenton degradation at pH 2.0 was only 0.1%. The excess acidity retarded the degradation rate as excess H<sup>+</sup> ions act as 'OH scavenger shown in reaction (20) (Devi et al. 2009, Fan et al. 2009).

The pH influence is a very important operating parameter which affects color and odor removal efficiency in  ${\rm Air/ZVI/H_2O_2}$  and  ${\rm Air/ZVI}$  oxidation process. The decrease of ZVIdecolorization rate by raising pH to the alkaline region is mentioned by Chen et al. (Chen et al. 2001). Fe<sup>2+</sup> ions, from the iron surface and hydroxyl ions in the alkaline solution precipitate ferrous hydroxide on the surface of iron occupying the reactive sites thus hindering the reaction (Chang et al. 2006).

#### Coagulation

Figure 5 shows that COD decreased even for low Fe<sup>3+</sup> doses. The decrease was significant in the case of concentration of Fe<sup>3+</sup> 50 mg/L up to 737 mg/L (21%). For increasing Fe<sup>3+</sup> dose, COD was still decreasing, but the rate was lower for 700 mg/L, it reached the value of 509 mg/L (45% decrease).

The main mechanisms for COD decreasing in coagulation could be pollutants adsorption-followed precipitation. For the removal of organic pollutants in sample, however, the oxidation – reduction mechanism must be considered in addition to the coagulation and precipitation mechanism. On the other hand, Air/ZVI and Air/ZVI/H<sub>2</sub>O<sub>2</sub> could be improved coagulation process, since the contaminants removal was enhanced by oxidation – reduction reactions in the system (Lai et al. 2007).

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#### **Kinetics**

The rate of the ZVI process is difficult to describe. The following equations were used to describe the investigated processes:

- pseudo-first-order reaction with respect to the COD value

$$d[COD] / dt = -k_{1st order}[COD];$$
 (19)

- pseudo-second-order reaction with respect to the COD value

$$d[COD] / dt = -k_{2nd \text{ order}}[COD]^2;$$
 (20)

 empirical equation that accounts for changes in the process rate due to other factors that influence changes with process time (the Balcerzak's equation, Eq. 21) (Naumczyk et al., 2012):

$$d[COD]/dt = -a [COD]t^{m}.$$
 (21)

Where 't' corresponds with time and 'a' and 'm' are constants that depend on the initial reagent concentrations.

After integration and application of the double logarithm, the following formula was obtained (Eq. 22):

$$\ln \ln[COD]_0/[COD] = (m+1) \ln t + \ln a - \ln (m+1)$$
 (22)

where [COD]<sub>0</sub> is the initial COD.

Calculations made using the pseudo-first- and pseudo second order reaction equation resulted in too low correlation

coefficients for all three investigated processes. Much better correlation coefficients were obtained for (Eq. 21). Usually ZVI processes kinetics is described as first – order one [6]. This simple approach has not worked in the case of automotive fleet repair facility wastewater.

The abovementioned equation has been used to describe the kinetics of the Fenton and photo-Fenton reactions in landfill leachate and cosmetic wastewater treatment(Naumczyk et al. 2012, Marcinowski et al. 2014).

The result of kinetic calculations and  ${\rm H_2O_2}$  efficiency are presented in Tables 2–5.

It was found that in the case of Air/ZVI/H<sub>2</sub>O<sub>2</sub> process, the Balcerzak's equation (Eq. 21) is characterized by a high compatibility (R<sup>2</sup> is usually over 0.9, especially for larger ZVI doses). But for ZVI it failed to satisfactorily describe the kinetics (very low R<sup>2</sup> values, decreasing with ZVI dose). In the case of this wastewater ZVI process seems to be too complex for simple approach, but models built on this approach are very complicated (Grcic et al. 2012).

#### H<sub>2</sub>O<sub>2</sub> efficiency

The oxidant  $(H_2O_2)$  efficiency is shown in Tables 2–5.The highest oxidant efficiency, up to 157.9% was observed in Air/  $ZVI/H_2O_2$  process, for  $ZVI/H_2O_2$  doses 4000/950 mg/L, under magnetic stirring at pH 3. Oxidant efficiency over 100% could be a proof for a chain radical reaction presence. More radicals than just hydroxyl one are used in oxidation process. These conditions are also optimal in the case of COD removal. The most important factor indicating  $H_2O_2$  efficiency is pH. If

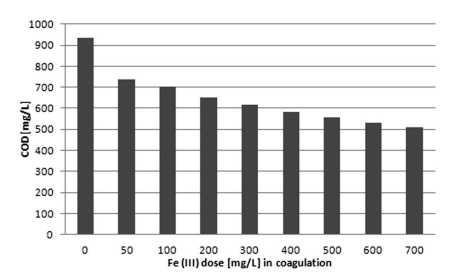


Fig. 5. Effect of coagulation

Table 2. Kinetics of Air/ZVI the process

Doses [mg/L]	Air/ZVI (1000 mg/L)	Air/ZVI (2000 mg/L)	Air/ZVI (3000 mg/L)	Air/ZVI (4000 mg/L)
а	0.0200	0.0253	0.0181	0.0058
m	-0.6603	-0.7133	-0.9377	-0.9853
n	0.3397	0.2867	0.0623	0.0147
k	0.0590	0.0883	0.2914	0.3989
R <sup>2</sup>	0.4954	0.4759	0.0669	0.0042

**Table 3a.** Kinetics of Air/ZVI/ $H_2O_2$  process

Doses [mg/L]	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (1000/450)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (2000/450)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (3000/450)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (4000/450)
а	0.0312	0.0357	0.0353	0.0409
m	-0.6973	-0.8615	-0.8579	-0.8742
n	0.3027	0.1385	0.1421	0.1258
k	0.1031	0.2582	0.2486	0.3252
R <sup>2</sup>	0.9917	0.9994	0.9426	0.9997
η	73.5	82.0	79.1	93.2

**Table 3b.** Kinetics of Air/ZVI/ $\mathrm{H_2O_2}$  process

Doses [mg/L]	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (1000/950mg)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (2000/950mg)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (3000/950mg)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (4000/950mg)
а	0.0272	0.0343	0.0667	0.0954
m	-0.3420	-0.3596	-0.5479	-0.6549
n	0.6580	0.6404	0.4521	0.3451
k	0.0413	0.0535	0.1475	0.2766
R <sup>2</sup>	0.9058	0.8898	0.9583	0.8613
η	119.2	128.1	146.0	157.9

**Table 3c.** Kinetics of Air/ZVI/ $H_2O_2$  process

Doses [mg/L]	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (1000/1900)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (2000/1900)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (3000/1900)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> (4000/1900)
а	0.0238	0.0378	0.0600	0.1343
m	-0.5036	-0.5895	-0.806	-0.6412
n	0.4964	0.4105	0.194	0.3588
k	0.0480	0.0921	0.3097	0.3744
R <sup>2</sup>	0.8386	0.9496	0.7873	0.7343
η	74.4	98.4	107.3	116.3

Table 4. Kinetics of Air/ZVI/ $H_2O_2$ /US process

Doses [mg/L]	Air/ZVI/H <sub>2</sub> O <sub>2</sub> /US (1000/950)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> /US (2000/950)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> /US (3000/950)	Air/ZVI/H <sub>2</sub> O <sub>2</sub> /US (4000/950)
а	0.0363	0.0450	0.0466	0.0409
m	-0.6649	-0.8568	-0.8595	-0.8742
n	0.3351	0.1432	0.1405	0.1258
k	0.1085	0.3145	0.3316	0.3252
R <sup>2</sup>	0.9066	0.9840	0.9843	0.9998
η	82.0	96.1	98.8	104.6

**Table 5.** Kinetics of Air/ZVI/ $\mathrm{H_2O_2}$  process, pH influence

Doses [mg/L]	pH 2.0, Air/ZVI/	pH 4.0, Air/ZVI/	pH 5.0, Air/ZVI/	pH 6.0, Air/ZVI/
	H <sub>2</sub> O <sub>2</sub> (1000/950)	H <sub>2</sub> O <sub>2</sub> (2000/950)	H <sub>2</sub> O <sub>2</sub> (3000/950)	H <sub>2</sub> O <sub>2</sub> (4000/950)
а	0.0293	0.0274	0.0306	0.0469
m	-0.7371	-0.6837	-0.8006	-0.8568
n	0.2629	0.3163	0.1994	0.1432
k	0.1116	0.0867	0.1535	0.3276
R <sup>2</sup>	0.9110	0.9628	0.9922	0.9840
η	65.3	65.3	68.0	96.1

system operates in not optimal pH conditions,  $\rm H_2O_2$  efficiency rapid decrease is observed even up to 65.3%. Another factor affecting the efficiency is a stirring method (the amount of dissolved oxygen available) but the influence is much smaller-oxidant efficiency decrease only to 82–104.6%, comparing to 65.3–96.1% in case of pH. It is possible that the difference in efficiency, compared with optimum efficiency 157.9%, is related to the  $\rm H_2O_2$  produced by the reaction (Eq. 6).

#### Conclusion

All processes (Air/ZVI, Air/ZVI/H<sub>2</sub>O<sub>2</sub>, coagulation) are suitable for automotive fleet repair facility wastewater treatment. Effectiveness of the processes was Air/ZVI/H<sub>2</sub>O<sub>2</sub>>Air/ZVI > coagulation. In Air/ZVI/H<sub>2</sub>O<sub>2</sub> process, the amount of H<sub>2</sub>O<sub>2</sub> is crucial for the efficiency of the process and must be close to COD value.In Air/ZVI and Air/ZVI/H2O2 processes higher ZVI dose causes better effectiveness. For the same amount of dissolved Fe, Air/ZVI is better than coagulation through the production of H<sub>2</sub>O<sub>2</sub> and oxidation of organic compounds. The optimal pH for Air/ZVI/H<sub>2</sub>O<sub>2</sub> process is 3.0. Change of pH value causes a rapid decrease in the effectiveness. Magnetic stirring is better than ultrasonic, because more oxygen is introduced to wastewater. Economically Air/ZVI and Air/ZVI/H2O2 are better than coagulation because the ZVI powder is very cheap (steel shavings). The highest COD removal efficiency, 76%, was achieved for Air/ZVI/H<sub>2</sub>O<sub>2</sub> process and ZVI/H<sub>2</sub>O<sub>2</sub>doses 4000/1900 mg/L, 120 min process time. COD decreased from 933 to 227 mg/L at pH 3.0. Odor and color were also completely removed.

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## Oczyszczanie ścieków z zakładów naprawczych floty samochodowej z wykorzystaniem procesów powietrze/ZVI i powietrze/ZVI/H<sub>2</sub>O<sub>2</sub>

**Streszczenie:** Ścieki z zakładu naprawczego floty samochodowej poddano oczyszczaniu z wykorzystaniem żelaza metalicznego i nadtlenku wodoru (Air/ZVI/H<sub>2</sub>O<sub>2</sub>). Badano wpływ dawki żelaza i nadtlenku wodoru, czasu i pH na efektywność procesu. Największy stopień usunięcia ChZT, 76%, uzyskano dla dawek ZVI/H<sub>2</sub>O<sub>2</sub> 4000/1900 mg/L, czasu 120 min i pH 3.0. ChZT zmniejszono z 933 do 227 mg/L. Dodatkowo uzyskano całkowite usunięcie barwy i zapachu. Skuteczność usunięcia ChZT rosła wraz ze wzrostem zastosowanej dawki ZVI. Zmiana pH na inne niż 3, powoduje gwałtowne zmniejszenie efektywności procesu.

Kinetyka procesu może zostać opisana z wykorzystaniem równania d[COD]/dt = -a [COD]t<sup>m</sup>, gdzie 't' oznacza czas a 'a' i 'm' są stałymi zależnymi od początkowego stężenia reagentów.

Badano także wpływ H<sub>2</sub>O<sub>2</sub> na efektywność procesu. Skuteczność usunięcia ChZT wynosi 40% (560 mg/L) w przypadku zastosowania ZVI bez dodatku H<sub>2</sub>O<sub>2</sub>. Określono także skuteczność koagulacji z wykorzystaniem FeCl<sub>3</sub>. Najlepsze rezultaty uzyskano dla dawki Fe<sup>3+</sup> 700 mg/L, zmniejszając ChZT do 509 mg/L.