Comparison of radical-driven technologies applied for paraben mixture degradation: mechanism, biodegradability, toxicity and cost assessment

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Abstract
Parabens (esters of p-hydroxybenzoic acid) are xenobiotics belonging to endocrine disruptors and commonly used as a preservative in cosmetics, food, pharmaceutical, and personal care products. Their wide use is leading to their appearance in water and wastewater in the range from ng/L to mg/L. In fact, the toxicity of benzylparaben is comparable to bisphenol A. Therefore, it is important to find not only effective but also ecofriendly methods for their removal from aqueous environment since the traditional wastewater treatment approaches are ineffective. Herein, for the first time, such extended comparison of several radical-driven technologies for paraben mixture degradation is presented. The detailed evaluation included (1) comparison of ozone and hydroxyl peroxide processes; (2) comparison of catalytic and photocatalytic processes (including photocatalytic ozonation); (3) characterisation of catalysts using SEM, XRD, DRS, XPS techniques and BET isotherm; (4) mineralisation, biodegradability and toxicity assessment; and (5) cost assessment. O₃, H₂O₂/Fe²⁺, H₂O₂/UVC, O₃/H₂O₂, O₃/UV A, O₃/H₂O₂/UV A, UVA/catalyst, O₃/catalyst and O₃/UV A/catalyst were selected from advanced oxidation processes to degrade parabens as well as to decrease its toxicity towards Aliivibrio fischeri, Corbicula fluminea and Lepidium sativum. Research was focused on the photocatalytic process involving visible light (UVA and natural sunlight) and TiO₂ catalysts modified by different metals (Ag, Pt, Pd, Au). Photocatalytic oxidation showed the lowest efficiency, while in combining ozone with catalysis and photocatalysis process, degradation efficiency and toxicity removal were improved. Photocatalytic ozonation slightly improved degradation efficiency but appreciably decreased transferred ozone dose (TOD). Results indicate that the degradation pathway is different, or different transformation products (TPs) could be formed, despite that the hydroxyl radicals are the main oxidant.

Highlights
• O₃, H₂O₂/Fe²⁺, H₂O₂/UVC, O₃/H₂O₂, O₃/UV A, O₃/H₂O₂/UV A, UVA/catalyst, O₃/Cat and O₃/UV A/Cat were compared.
• Noble metal catalysts cause higher effectiveness under natural sunlight than under UVA.
• O₃/Cat supported by UVA increases degradation and decreases ozone dose requirements.
• Lower toxicity on different bioindicators after all advanced oxidation processes were observed.
• Combining ozone with catalysis and photocatalysis process improved toxicity removal.

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Introduction

Xenobiotics are organic compounds foreign to living organisms, which do not occur in the form of neither primary nor secondary metabolites. While some xenobiotics are natural compounds, most of them are from anthropogenic source (Hashmi et al. 2017; Yang et al. 2017). They are included in personal protective equipment or pharmaceuticals, referred as pharmaceuticals and personal care products (PPCPs), whose presence in water reservoirs is a serious environmental problem (Yang et al. 2017). The compounds belonging to the PPCP group (e.g. parabens) have different physicochemical properties, which make them difficult to detect and remove from wastewater and natural environment (Nakada et al. 2007; Yang et al. 2017; Wang et al. 2018). Many of xenobiotics have been classified as compounds that disrupt the hormonal balance of the body (xenoestrogenic or endocrine compounds) called endocrine-disrupting compounds (EDCs) (Nakada et al. 2007; Esplugas et al. 2007; Giulivo et al. 2016; Wang and Wang 2016). Numerous studies have confirmed the harmful effect of xenobiotics on disturbing the hormonal balance of organisms, hepatotoxic effects damaging liver cells and the possibility of producing free radicals which cause oxidative stress (Vilela et al. 2018).

Xenobiotics are an interesting research object since they are a group of compounds with significant toxic effects. In addition, in trace amounts, they occur in the aquatic environment. However, even at very low concentrations (ppb or ppm), these compounds are dangerous for the environment, and contaminated water causes a threat to people and also affects the condition of animal populations and plants (Gu et al. 2002). Unfortunately, the still growing consumerism means that the presence of xenobiotics in the lives of people, animals and the environment is inevitable. It is known that the removal of EDCs in wastewater treatment plants using conventional methods is insufficient (Nakada et al. 2007; Esplugas et al. 2007; Wang and Wang 2016; Gmurek et al. 2017; Yang et al. 2017; Wang et al. 2018). Poor efficiency is probably related to physical and chemical properties as well as its toxicity. Considering the product cycle, these compounds in unmetabolised form were found in wastewater, from where they get mainly to water reservoirs (e.g. used pesticides as a result of runoff from agricultural areas get into surface waters). Consequently, they are detected in drinking water from which they can be absorbed into the body.

Parabens (esters of \(\beta\)-hydroxybenzoic acid) represent xenobiotics, PPCPs and EDC groups commonly used as preservative in cosmetics and personal care products, food, pharmaceuticals, etc. Although it was found that butylparaben can be synthesised by marine bacteria (Peng et al. 2006), only produced synthetically commercial parabens are used for industrial purpose (Juliano et al. 2017). Furthermore, for the last 10 years, the EU regulation about paraben content in cosmetic products has been restricted. According to Regulation (EU) No. 358/2014 (Juliano et al. 2017), isopropylparaben, isobutylparaben, phenylparaben, benzylparaben and pentylenparaben are prohibited in cosmetic products. As reported by Regulation (EU) No. 1004/2014 (Juliano et al. 2017), propylparaben and butylparaben (also their isoforms and salts) were banned from cosmetic products for children under 3 years of age, based on those substances’ potential endocrine activity. Additionally, the maximum concentration allowed for these two parabens was reduced in the cosmetic products for adults and kids above 3 years of age (Juliano et al. 2017). Parabens can be considered as endocrine disruptors and carcinogens (Giulivo et al. 2016; Juliano et al. 2017). Moreover, the toxicity and oestrogenicity of benzylparaben is comparable to that of bisphenol A (Martins et al. 2016). Parabens were detected in influents from WWTPs (even at concentrations of mg/L), and after the treatment in effluents, their concentrations were still high (including BzP) (Canosa et al. 2006; Haman et al. 2015; Marta-Sanchez et al. 2018; Gomes et al. 2018a). The presence of paraben was recorded in natural water in Japan, the UK, Portugal, Switzerland, Belgium, China, Spain and the USA in the range of 0.1–676 ng/L (Jonkers et al. 2010; Haman et al. 2015). Some of them were detected in drinking water in the concentration of ng/L in Germany, Spain or the USA (Haman et al. 2015; Marta-Sanchez et al. 2018). Swimming pools are also contaminated by parabens as well as sediments (Haman et al. 2015; Lempart et al. 2018). Due to the characteristics of parabens, which have a negative impact on human and animal populations, a method of their degradation is sought, because nature also fails on removing them through biological process. Moreover, the complex structure causes their insufficient removal in the wastewater treatment process, and as a result of which, they accumulate in living organisms. Until now, commonly applied methods of wastewater treatment or water treatment are ineffective, and a significant amount of these compounds is presented in water. Therefore, it is important to develop efficient methods for eliminating them from wastewater.

Based on literature data, the most effective method of removing them is chemical oxidation using reactive oxygen species generated using ultraviolet (UV) or visible (Vis) radiation or ozone (Nakada et al. 2007; Gottschalk et al. 2010; Nawrocki and Kasprzyk-Hordern 2010; Tay et al. 2010; Gmurek et al. 2015, 2017; Wang and Wang 2016; Pipolo et al. 2017; Gomes et al. 2017a; Wang et al. 2018;
Foszpańczyk et al. 2018). These methods often allow to obtain products more susceptible to biodegradation. Studies focusing on photodegradation processes of compounds contaminating the aquatic environment are mostly aimed at intensifying the degradation process and obtaining non-toxic phototransformation products. The preferred direction of the development of these methods is the possibility of using renewable energy in the form of solar radiation. Many of the known photochemical methods belong to the group of advanced oxidation processes (AOPs). Lately, AOPs are the most applicable treatment methods for removing hazardous water contaminants, despite that lots of them have a limitation due to optimal pH parameter. O₃ has two pathways of degradation: in acidic pH, direct oxidation via O₃, while in alkaline pH, indirect oxidation via OH (Glaze 1986; Glaze et al. 1987; Lucas et al. 2010). H₂O₂/Fe²⁺ has an optimum effectivity at pH ≈ 3, particularly due to the precipitation of ferric oxyhydroxide at higher pH value (Ikhlaq et al. 2018, 2019). However, alkaline environment is the favorable pH for most AOPs (Glaze et al. 1987; Lucas et al. 2010; Xiao et al. 2015; Boczkałj and Fernandes 2017; Shahzad Munir et al. 2019; Asgari et al. 2019; Ikhlaq et al. 2019). In fact, the AOPs are aimed to be applied as tertiary systems for the removal of hazardous water contaminants from municipal wastewater treatment plants (Gmurek et al. 2017; Wang et al. 2018; Miklos et al. 2018; Soriano-Molina et al. 2018). Usually, these effluents are treated in secondary reactors based in biologic systems that require neutral pH conditions for proper operation. Therefore, it is interesting that the following treatment technologies also operate under those conditions (further pH adjustment is not necessary) which reduce operating costs, even in case of Fenton process, however, with the disadvantage of iron precipitation (Soriano-Molina et al. 2018). The second important issue in case of AOP is the light source. Unfortunately, the visible light can be applied as an effective light source only for few AOPs. Mostly, they focus on the application of UV radiation with various combinations of other oxidants (H₂O₂, O₃, Fenton reagent, TiO₂). Even though processes using UV radiation are energy-consuming, these are still widely used due to their high efficiency and rate of degradation process. The use of UV radiation also allows for process control (constant radiation conditions). However, it should be borne in mind that the use of additional oxidants may cause further environmental problems (for example, the resulting sediment during the photo-Fenton process). Thus, processes such as photolysis using UV, UV/H₂O₂, TiO₂/UV, O₃/UV and TiO₂O₃/UV can form the foundations of research works using photochemical processes, allowing determination of quantum yields or kinetic constants. Nowadays, the best catalyst is sought; for example, the addition of a second semiconductor will shift the spectrum absorbed by the semiconductor towards the visible light. Moreover, photocatalyst surface modifications by metals are affected by changing the distribution of electrons on the properties of the semiconductor, which increases the photodegradation activity (Nawrocki and Kasprzyk-Hordern 2010; Grabowska et al. 2016; Gmurek et al. 2017; Gomes et al. 2017b; Foszpańczyk et al. 2018).

Bearing in mind the problem associated with parabens, it is important to find not only effective but also ecofriendly methods for their removal from aqueous environment. From all well-known water and wastewater treatment technologies, AOPs are not only important, efficient and ecofriendly but also promising methods that allow to remove persistent aqueous contaminants.

To the best of our knowledge, this is the first time that such extended comparison of several AOPs for the mixture of water pollutant degradation has been presented. The detailed evaluation included (1) comparison of ozone and hydroxyl peroxide processes; (2) comparison of catalytic and photocatalytic processes (including photocatalytic ozonation); (3) characterisation of catalysts using SEM, X-ray diffraction (XRD), DRX, XPS techniques and BET isotherm; (4) mineralisation, biodegradability and toxicity assessment; and (5) cost assessment. H₂O₂/Fe²⁺, H₂O₂/UV, O₃, O₃/H₂O₂, O₃/UV, O₃/H₂O₂/UV, UVA/catalyst (UVA/Cat), O₃/Cat and O₃/UV/catalyst (O₃/UV/Cat) were selected from AOPs to degrade parabens from aqueous environment. As photocatalysts, noble metals (Pt, Pd, Au, Ag) that modified TiO₂ were applied and compared with pure TiO₂. For photocatalytic oxidation, the comparison of UVA light and natural sunlight was made. The toxicity assessment before and after treatment was evaluated by Vibrio fischeri luminescence inhibition and the freshwater Asian clam Corbicula fluminea mortality. Moreover, the phytotoxicity test using Lepidium sativum was performed.

Materials and methods

Chemicals and reagents

A solution of five parabens [methylparaben (MP), ethylparaben (EP), propylparaben (PP), butylparaben (BuP) and benzylparaben (BeP)] was used as a model effluent. The pollutant solutions were prepared with Milli-Q (MQ) water using 10 mg/L of each compound. A commercial form of TiO₂ (P25, crystalline composition of 80% anatase and 20% rutile, with a surface area of 50 m²/g) was obtained from Evonik (Germany). 2-Propanol (for analysis) was purchased at Merck (Germany). All solvents [HPLC water (VWR), acetonitrile and formic acid (both from Merck)] were analytical grade (≥99%). Experimental solutions were prepared using ultrapure water.
Preparation of photocatalysts

Two methods of TiO₂ modification were used. TiO₂-Pt, TiO₂-Pd and TiO₂-Ag were prepared by UV reduction of Pt⁴⁺, Pd²⁺ and Ag⁺⁺ used as chloride salts, in the TiO₂ (P25) suspension, while TiO₂-Au was prepared by the sol–gel method, according to Gomes et al. (2017a). In photodeposition method, an aqueous solution of isopropanol containing H₂PtCl₆ (0.5 E/Ls; 31.8 W/m²) was degassed with nitrogen and irradiated by UV–Vis light (1000 W Xe lamp) for 6 h. For TiO₂-Ag preparation, an aqueous solution of ethanol containing AgNO₃ (0.5 wt%) was degassed with nitrogen and irradiated by UV–Vis light (1000 W Xe lamp) for 100 min. All catalysts prepared by the photodeposition method were separated by centrifugation and dried at 65–120 °C for 12 h. In the sol–gel method, titanium(IV) isopropoxide (TIP), known as the titanium source for the anatase-type TiO₂, was used for TiO₂-Au preparation. TIP was mixed with AuCl₄⁻ (0.5 wt%), methanol and distilled water. The solution was stirred at room temperature for 2 h, followed by a 24-h thermal treatment (45 °C) and calcinations at 400 °C for 2 h.

Characterisation of photocatalysts

The microstructural analysis of the surfaces was performed by TESCAN VEGA3 SBH - Easy Probe Scanning Electron Microscopy (SEM) with a tungsten-heated cathode. The SEM images were acquired with a working tension of 5 kV, using the secondary electron detector. The optical properties of the photocatalysts were characterised by an ultraviolet/visible diffuse reflectance spectrophotometer (Evolution 220; Thermo Scientific), in which BaSO₄ was employed as the internal reflectance standard. The band gaps for all photocatalysts were determined using the Tauc plot method. The surface areas of catalysts Brunauer–Emmett–Teller were determined using nitrogen (7–196 °C) with an accelerated surface area and porosimetry analyser (ASAP 2000, Micrometrics). The crystal structure of photocatalysts was determined from XRD pattern measured in the range of 2θ = 20–80° using an X-ray diffractometer (X’Pert PRO-MPD, Philips) with Cu target (λ = 1.542 Å). The lattice parameters were estimated by the Le Bail method using FullProf package.

Light sources

Irradiation experiments were performed using a UVC (254-nm) lamps, UVA (365-nm) lamps and natural sunlight (in Poland as well as in Portugal). Six low-pressure mercury lamps (USHIO, model G8T5 Hg; 7.2 W each) with a maximum emission of 254 nm (88.6%) were used as a UVC source (E₀ = 5.75 × 10⁻⁷ E/Ls; 8.9 W/m²). Three lamps made of blacklight blue glass, which transmits UVA radiation (Philips TL 6 W BLB; tube diameter of 16 mm), with the maximum emission of 365 nm were used as UVA source (E₀ = 5.75 × 10⁻⁷ E/Ls; 8.9 W/m²). The natural sunlight was applied, and the experiments were conducted in the sunny days (in July 2016 and May 2018) with the average irradiance of 340 W/m², corresponding to the photon flux rate E₀ = 2.60 × 10⁻⁴ E/Ls.

The experiments with UVC lamps were carried out in quartz test tubes (10 mL and an average optical path length of 0.85 cm) placed between two exposure panels using a merry-go-round device.

The experiments with a UVA lamps were carried out in a 2-L reactor with internal lamp source. The experiments with sunlight were carried out in a cylindrical borosilicate reactor (0.5 L) equipped with a compound parabolic concentrator (CPC) collector. All reactors set up are presented in Fig. S1.

During the experiments, the light exposition was measured with an Oceans Optics USB 4000 fibre optic spectrometer with an approximate resolution of 0.4 nm. All the experiments were performed twice. Results are presented as a mean value of single experiments.

Experimental procedure

H₂O₂/Fe²⁺ process

Reaction was carried out in a batch-stirred glass reactor. Ferrous ion (Fe²⁺) was added in the form of iron sulphate, and the reaction started when the desired amount of H₂O₂ was injected. pH was left to run freely (at the beginning, 6 ± 0.7, and after 120 min of reaction, the final pH value falls to 3.2 ± 0.3). A detailed procedure and an experimental set-up were previously published in the study of Martins et al. (2016).

H₂O₂/UVC

The reaction set-up was composed by quartz test tubes (10 mL) placed between two exposure panels using a merry-go-round device at a distance of 15 cm from its centre. Each panel was composed by three low-pressure mercury lamps (7.2 W each) with a maximum emission for 254 nm. The pH of the reaction medium was 7, adjusted by a phosphate buffer. A detailed procedure and an experimental set-up were previously published in the study of Gmurek et al. (2015).

O₃-based technology

Two kinds of equipment were used for ozone-based technology. The first one was a semi-batch reactor of 500 mL which was used to perform ozonation experiments (single ozonation (O₃) and O₃/H₂O₂). Ozone was produced in situ by an ozone generator that was fed by a pure oxygen stream (99.9%) supplied by Praxair. Ozone concentration in the gas entering ([O₃]in) and leaving ([O₃]out) the reactor was measured by a gas ozone meter. To ensure chemical regime, the reactor was
agitated at the approximate speed of 700 rpm by means of a magnetic stirrer. The pH of the reaction medium was 7, adjusted by a phosphate buffer. A detailed procedure and an experimental set-up were previously published in the study of Pipolo et al. (2017). The second one was a 2-L glass reactor with the possibility of using 3 UVA lamps which was used for O₃, O₃/UV A, UVA/Cat and O₃/Cat O₃/UV A/Cat experiments. The stirring speed was previously optimised at 700 rpm to ensure chemical regime. Along the reaction, samples were withdrawn and immediately centrifuged at 3500 rpm to remove the photocatalyst. pH was left to run freely (at the beginning, 5.7). Ozone was produced from pure oxygen stream (99.9%), and the inlet ([O₃]ᵢⁿ) and outlet ([O₃]ₒᵘᵗ) ozone concentrations were measured by ozone analysers. A detailed procedure and an experimental set-up were previously published (Gomes et al. 2017a, b, c, d).

The photocatalytic experiments were carried out with 0.07 g/L of the photocatalyst. To test the adsorption capacity of photocatalyst, it was placed in the reactor with the solution of parabens 5 min before ozone being fed and turning on the light irradiation. During the degradation experiments, the reactor was bubbled by pure oxygen stream (99.9%, Praxair, 0.2 L/min).

All process parameters are collected in Table S1.

**Analytical methods**

The concentration of paraben in the mixture was followed by HPLC (UFLC, Shimadzu) with a C18 (SiliaChrom) column at 40 °C. The detection wavelength was 255 nm. The injection volume was 20 μL, and the mobile phase consisting a mixture of 50:50 methanol:acidic water (0.1% orthophosphoric acid) was used with the flow rate of 0.5 mL/min.

Chemical oxygen demand (COD) was measured according to the standard method 5220D using a calibration curve obtained for potassium hydrogen phthalate (Panreac). The absorbance of the samples was measured at 445 nm (WTW photolab S6 photometer) after 2 h of digestion at 150 °C (ECO25; VELP Scientifica).

The absorption spectra of parabens were measured in a 1-cm quartz cell on a Unicam spectrophotometer.

**Toxicity assessment**

The toxicity assessment was investigated by luminescence inhibition tests with *A. fischeri* using a LUMISTox 300 apparatus. The bioassays with C. fluminea were conducted using adult clams. Before clams were used as a bioindicator, they were maintained at least 1 week under continuous aeration at constant temperature and photoperiod (16-h light and 8-h dark) prior to use. The laboratory culture water was fully renewed once a week. Mortality tests were conducted under static conditions (Gomes et al. 2014). Blank controls with dechlorinated municipal water were applied along with a dilution series of untreated and treated model effluents with exposure lasting 72 h. *L. sativum* was considered as a bioindicator in the phytotoxicity test. Toxic effects, expressed as germination index (GI), were calculated according to Gomes et al. (2017a). Moreover, the level of the toxic impact on *L. sativum* was assessed by the method suggested by Wang (1992).

**Results**

**Preliminary study**

UV radiation is very often used in photodegradation processes. However, the use of UV radiation alone in purification or treatment processes is uneconomical (Gmurek et al. 2017). Most often, monochrome radiation with a wavelength of 254 nm is used, which, in the case of most organic substances, overlaps with the maximum of their absorbance (Fig. 1).

Unfortunately, UVC radiation in this range does not occur in...
the spectrum of solar radiation at the Earth’s surface. Thus, studies based on UVC (254 nm) techniques cannot be directly translated into processes occurring in the natural water environment. Nevertheless, these technologies are still very popular, especially in disinfection systems. Photochemical degradation techniques are also conducted using polychromatic UVA radiation, which falls within the solar spectrum (Fig. 1). Most often, UV radiation is combined with other oxidants, which leads to the generation of hydroxyl radicals (•OH) (Gmurek et al. 2017). It is known that the use of •OH generated as an oxidant in AOPs leads to very rapid degradation of compounds and high mineralisation. Such high degradation efficiency is obtained as a result of the low selectivity of these oxidants. However, to generate hydroxyl radicals, a lot of energy and an additional oxidiser are needed. On the other hand, the use of less energetic visible radiation is associated with a decidedly lower reaction rate, i.e. a longer photodegradation time. As it was mentioned above, the modification of photocatalysts can improve their photoreactivity in the range of visible light. As can be seen in Fig. 1, all investigated photocatalysts exhibit very strong absorption below 400 nm that can be explained by the property of titanium to absorb UV light. Above 400 nm in the visible light region, the best absorption properties have been shown by TiO_2-Au and TiO_2-Pd, while the worst absorption properties have been shown by TiO_2-Ag. However, all modified photocatalyst are able to absorb visible light, which can be attributed to the surface plasmon resonance of noble metal nanoparticles (Leong et al. 2015; Grabowska et al. 2016).

To evaluate the applicability of photodegradation processes to decontamination of parabens, the direct photolysis of their mixture under UVC and UVA lamps as well as sunlight has been compared. As it was expected, according to the results depicted in Fig. 1, almost complete removal after 1 h has been achieved when a UVC lamp was used (Fig. S2). Besides, negligible degradation was observed under UVA light, while natural sunlight has not let any degradation at all (Fig. S1).

**Comparison of ozone and H_2O_2 technologies**

It is well known that the capable action of ozone, H_2O_2 and light improves the effectivity of decontamination. However, it should be noticed that the process parameters such as hydrodynamic conditions are significant. The synergetic effects are not always observed; moreover, a scavenging effect can be also expected, while some oxidants are added in extent. Furthermore, the decrease of initial concentrations is useful for the understanding of reaction mechanisms, but it is not sufficient for real effluents containing a mixture of organic compounds and no information is given about degradation of by-products. More detailed information is given by COD removal since the global organic matter removal is followed. As can be seen in Fig. 2, the process parameters are extremely important. Although all processes led to complete removal of initial concentration of parabens after 2 h (mostly resulted in 80% concentration reduction of parabens after 1 h), the COD removal is totally different for all of them.

The efficiency of ozone-based technology is highly dependent on pH, which has the ability to change the ozone oxidation pathway and dissolve ozone concentration (Hoigné and Bader 1976; Olak-Kucharczyk and Ledakowicz 2016; Pipolo et al. 2017; Asgari et al. 2019). In an acidic condition, molecular ozone pathway occurs; in neutral and alkaline environments, the hydroxyl radical pathway is promoted. Furthermore, O_3 solubility increased with a decrease in pH (Biń 2006), affecting ozone concentration in liquid. It is well known that the ozone saturation is dependent not only on temperature but also on pH which can be effected by liquid constituents or impurities such as salinity and ionic strength (Biń 2006). If the AOP treatment is taken into consideration, two possible situations can occur: it can be supplemented for secondary effluents after biological treatment (pH close to neutral) or before biological treatment (effluents do not have to be buffered). Therefore, three reaction conditions during single ozonation have been investigated. At buffered neutral condition (pH ~ 7) in small volume reactor (500 mL), two ozone inlet gas concentrations were used (45 mg O_3/L and
20 mg O₃/L). While in non-buffered solution (2-L reactor), 8 mg O₃/L was applied. In both cases, the mass transfer resistance can be neglected (Gomes et al. 2017; Pipolo et al. 2017). It is well known that hydroxyl radicals are more reactive than molecular ozone. Second-order rate constants for reaction with •OH are 10²–10¹⁰ M⁻¹/s order of magnitude (Tay et al. 2010; Gmurek et al. 2015), while the second-order rate constants for reaction of organic molecules with ozone are lower ~ 10² order rate constants for reaction of organic molecules with ozone are lower ~ 10²–10⁴ M⁻¹/s or ~ 10²–10⁶ M⁻¹/s in acidic or neutral condition, respectively (Tay et al. 2010; Olak-Kucharczyk and Ledakowicz 2016). Ozone dose is an important key parameter in determining the residual ozone concentration. However, the ozone decomposition increases with increasing pH. Therefore, at low pH, less ozone concentration is required because ozone lifetime is much longer than in neutral or alkaline environment (Bin 2006; Hansen et al. 2016).

As can be seen in Fig. 2, by single ozonation in buffered solution when the reaction occurs via •OH, the COD removal was much higher than in not buffered solution. At the buffered solution, it was observed that with increasing ozone dosage, the reaction was accelerated. However, in the non-buffered solution, the ozone concentration was much lower, and the increase of it would be meaningless. The ozone pathways will always be slower than •OH. In case when the pH was not adjusted (initial pH = 5.7 then decreased during the reaction), even when H₂O₂ or UV A was used, the effectiveness has not increased significantly (Fig. 2). From integration of ozonation with the biological system point of view, if the initial concentration of contaminants was degraded, higher COD removal is not required, because biological treatment will be able to handle it.

However, application of single ozonation led to the problem of residual O₃ in treated wastewater that does not allow for the immediate use of biological treatment after it (the remaining dissolved ozone must be let to be decomposed first before the biological reactor). This disadvantage can be overcome by the merging action of oxidants. It was mentioned that combining O₃ with UV or H₂O₂ to enhance •OH generation is often more effective than O₃ alone. The best results have been obtained for O₃/H₂O₂ (0.5 L) treatment when the •OH radical pathways occurred (pH = 7). Even the application of H₂O₂/ UVC and Fenton reagent did not give better results (Fig. 2). It should be noticed that in the case of the Fenton process after the addition of iron sulphate and hydrogen peroxide, pH decreased from neutral to pH 3.6 [optimal condition according to Neyens and Baeyens 2003, Martins et al. 2016 and Ikhlaq et al. 2018]. The synergetic effect of H₂O₂ and O₃ has been also observed in the case of non-buffered solution (Fig. 2).

The generation of •OH in O₃/UV is initiated by the photolysis of O₃ by UV (Gottschalk et al. 2010). The application of UVC (254 nm) enhances the oxidation potential due to the high molar extinction coefficient of O₃ (ε₂₅₄ = 3300 M⁻¹/cm) (Gottschalk et al. 2010) (Fig. 1b). However, the major drawback of the ozone-based technology especially UV/O₃ is generating high operational costs (the energy requirement of O₃ production and UVC lamp equipment). The cheaper UVA lamp (365 nm) is not effective as a UVC source, because the ozone molecule is not absorbed in this region, so the •OH is not effectively produced. Therefore, the COD removal is even lower than in the case of single ozonation (Fig. 2). A better COD removal is achieved when O₃/H₂O₂ was applied, especially when effectiveness has been considered from the transferred ozone dose (TOD) point of view (Fig. S3). The action of O₃ with H₂O₂ and UVA led to worse results than O₃/H₂O₂ and lower COD removal and required more TOD (Fig. 2, Fig. S3). However, the O₃/H₂O₂/UV A system exhibited higher COD removal in comparison to single ozonation and O₃/ UVA and required less TOD (Fig. 2, Fig. S3). Based on the results, it was shown that the effectiveness of the process with respect to time as well as TOD must be considered.

### Photocatalytic processes

As shown in Fig. 1b, the photocatalysts with noble metals like Pd, Ag, Pt and Au effectively absorbed UVA as well as visible light. Moreover, it was proved that both of this light do not cause the photodegradation of parabens directly via photolysis (Fig. S1). TiO₂-Pd and TiO₂-Au exhibit the best absorption characteristics in the visible light range than TiO₂-Pt and TiO₂-Ag, while TiO₂ has the worst ability to absorb visible light (Fig. 1b). That can be attributed to the width of the band gaps as shown in Table 1 and Fig. S4. Indirect/direct band gaps of pure TiO₂ (3.27 eV/3.38 eV) were reduced to 3.00 eV/3.22 eV, 2.74 eV/3.23 eV, 2.59 eV/3.19 eV and 2.5 eV/ 3.13 eV for TiO₂-Ag, TiO₂-Pt, TiO₂-Au and TiO₂-Pd (all 0.5 wt%), respectively (Table 1). Since TiO₂-Pd and TiO₂-Au have the narrowest and similar band gaps (both direct and indirect), both perform in the Vis light range better, increasing the efficiency of the photocatalysis. The improvement in visible range absorption ability may be attributed to the surface plasmon resonance of noble metal nanoparticles. In Fig. 1b, in visible range, the maximum absorption peak due to localised surface plasmon resonance (LSPR) for TiO₂-Au and TiO₂-Ag can be observed (positioned in the range of 525–600 nm and 450–500 nm, respectively). It is well known that the photocactivity has also influenced the particle size, shape and particle distribution. Moreover, the sol–gel method gives ability to synthesize nanosized titania with high purity at low temperature (Xiao et al. 2015), while the photodeposition method allows a better particle size distribution control (Wenderich and Mul 2016). SEM images present the successfully deposited noble metals (Fig. S5). The modification by noble metal resulted in the homogeneous distribution of the Pd, Ag as well as Au particles on the catalyst surface. However, in the case of Pt and Au, particles were incorporated on the TiO₂ surface in an irregular manner presenting higher agglomeration of particles or the size of particles. XRD
measurement confirmed crystallographic forms after modification and showed that the anatase phase is predominant (Table S2). The results proved that surface modification with noble metals did not change significantly the basic structure of TiO₂ (P25). Moreover, results indicated that titania, for the TiO₂-Au nanoparticles, prepared by TIP, was present in the pure anatase structure. The size of anatase crystallites was the biggest for TiO₂-Pd while the smallest for TiO₂-Au. The BET analysis shows that the specific surface area was not significantly changed after modification (Table S2).

As can be seen in Fig. 3a, the photocatalytic oxidation under UVA with TiO₂ led to negligible paraben decontamination after 3 h. Unfortunately, the photocatalytic oxidation with TiO₂ modified by noble metals has not caused the complete degradation of paraben concentration; therefore, the COD reduction exceeded 20%. In comparison to TiO₂, only a slight improvement has been obtained when TiO₂-Au was applied. The best results, but still not enough satisfactory, were obtained for TiO₂-Pd and TiO₂-Ag (Fig. 3a). The highest degradation was achieved for BeP (73% and 53% of BeP initial concentration for TiO₂-Pd and TiO₂-Ag, respectively). However, for the rest of parabens, the removal was below 50% of their concentration, while when COD removal is taken into account, only 19% and 17% were obtained after 3 h of the treatment for TiO₂-Pd and TiO₂-Ag, respectively. It was surprising that despite the low initial paraben degradation during TiO₂-Pt photocatalytic treatment, 18% of COD removal was achieved. In case of TiO₂ as well as TiO₂-Au, the COD removal was around 10%.

Much better results have been obtained when natural sunlight was applied. Indeed, significant improvement was observed for all catalysts. Moreover, the use of TiO₂-Pd led to total paraben degradation (Fig. 3b). Also, higher degradation was achieved in the case of TiO₂-Pt rather than for TiO₂-Ag, indicating that this photocatalyst has better visible light photocatalytic ability (Foszpańczyk et al. 2018). However, the highest improvement has been observed for TiO₂-Au, proving that the modification by Au increases the visible light ability, which is in agreement with absorption spectra. Despite higher degradation under sunlight, the effectivity of photocatalytic processes was much below expectation. Therefore, the catalytic ozonation as well as the photocatalytic ozonation have been investigated (Fig. 4). As it was expected, the catalytic ozonation as well as the photocatalytic ozonation increased effectivity. The parabens were totally removed during the first 2 h of both treatments, leading to higher mineralisation than for O₃ or O₃/UVA. It is well known that the catalytic ozonation allows for the effective formation of hydroxyl radicals also at a low pH in contrast to single ozonation (Nawrocki and Kasprzyk-Hordern 2010). Furthermore, as expected, the photocatalytic ozonation led to higher COD removal. In case of 2 h of O₃/Cat treatment, TiO₂ and TiO₂-Au exhibited similar effectivity (28% of COD removal). While the application of TiO₂-Pt, TiO₂-Ag and TiO₂-Pd photocatalysts caused

### Table 1

| Band gap | TiO₂ (eV) | TiO₂-Ag (eV) | TiO₂-Pt (eV) | TiO₂-Au (eV) | TiO₂-Pd (eV) |
|----------|-----------|--------------|--------------|--------------|--------------|
| Direct   | 3.38      | 3.22         | 3.23         | 3.19         | 3.13         |
| Indirect | 3.27      | 3.00         | 2.74         | 2.59         | 2.5          |

![Fig. 3](https://example.com/fig3.png)

**Fig. 3** Photocatalytic oxidation of paraben mixture under UVA (a) and natural sunlight (b) during 15 min, 30 min, 60 min, 120 min and 180 min
around 35% of COD removal. After 2 h of O3/UV A/Cat treatment, 38% of COD removal was achieved when TiO2 and Au-TiO2 were used, while 41%, 43% and 49% of COD were removed when TiO2-Pd, TiO2-Ag and TiO2-Pt, respectively, were applied. It was surprising that O3/H2O2 as well as O3/H2O2/UV A gave better COD removal (both above 33%) efficiency than O3/Cat and O3/UV A/Cat [after 1 h, 27% (TiO2-Pd) and 28% (TiO2-Pt), respectively] (Figs. 2 and 4).

However, when TOD is taken into account, the requirements are much lower when catalytic ozonation and photocatalytic ozonation were considered (Fig. S3, Fig. 4b). As can be seen in Fig. 4b, the orders of the highest ozone-demanding catalysts are TiO2-Au < TiO2-Ag < TiO2 < TiO2-Pt < TiO2-Pd and TiO2-Pd < TiO2 < TiO2-Au < TiO2-Ag < TiO2-Pt during O3/Cat and O3/UV A/Cat, respectively. The differences can be related with ozone adsorption on the surface of the catalyst and whether this process leads to ozone decomposition followed by the formation of surface-bound or free radicals (Nawrocki and Kasprzyk-Hordern 2010) and with absorption of light ability. In fact, the presence of UVA with these catalysts photogenerates electrons that can be trapped by ozone reducing to ozonide radical which, in acidic conditions, can be adjuvant on hydroxyl radical production (Mehrjouei et al. 2015; Gomes et al. 2017d, 2018b).

Kinetic studies

Kinetic studies for applied AOPs were performed for the degradation of each paraben in admixture assuming pseudo-first-order kinetics. The pseudo-first-order kinetic model fitted well with the experimental results with coefficients of determination ($R^2$) ranging from 0.790 to 0.998. Pseudo-first-order rate constants ($k'$) of the processes are collected in Tables 2, 3 and 4. Firstly, the ozone and hydrogen peroxide process will be discussed (Table 2). The calculated values of pseudo-first-order rate constants for the H2O2/UVC process for all parabens are 1 order of magnitude higher than those for other processes. It must be mentioned that for this process, the complete removal of each paraben was achieved after just 30 min., while for the H2O2/O3/UVA process, 90 min was required. The rest of the processes (O3, H2O2/UVC, H2O2/O3, UVA/O3) needed 2 h or 3 h to remove the paraben concentration at the satisfactory levels. Despite maintaining the same order of magnitude of $k'$, the slight increase in the following order (average values between brackets): O3 (0.02475 ± 0.0102 l/min), H2O2/O3 (0.02839 ± 0.0026 L/min), UVA/O3.

Table 2  Pseudo-first-order rate constants for paraben mixture phototransformation during O3, H2O2/UVC, H2O2/O3, UVA/O3 and H2O2/O3/UVA processes

| Process                  | Paraben | $k'$ (1/min) |
|--------------------------|---------|--------------|
| O3 (8 mg O3/L)           | MP      | 0.01737      |
|                          | EP      | 0.02295      |
|                          | PP      | 0.0245       |
|                          | BuP     | 0.01699      |
|                          | BeP     | 0.04192      |
| H2O2/UVC                 | MP      | 0.62102      |
|                          | EP      | 0.75672      |
|                          | PP      | 0.65232      |
|                          | BuP     | 0.56989      |
|                          | BeP     | 0.88343      |
| H2O2/O3 (70 mg H2O2/L, 8 mg O3/L) | MP | 0.02847      |
|                          | EP      | 0.02483      |
|                          | PP      | 0.02701      |
|                          | BuP     | 0.03038      |
|                          | BeP     | 0.03128      |
| UVA/O3 (8 mg O3/L)       | MP      | 0.03111      |
|                          | EP      | 0.02938      |
|                          | PP      | 0.03151      |
|                          | BuP     | 0.02776      |
|                          | BeP     | 0.03264      |
| H2O2/O3/UVA (13.5 mg H2O2/L) | MP | 0.036       |
|                          | EP      | 0.0363       |
|                          | PP      | 0.04206      |
|                          | BuP     | 0.03402      |
|                          | BeP     | 0.03585      |
When the photocatalytic oxidation process is considered (Table 3), it can be seen that the calculated values of pseudo-first-order rate constants for the processes under sunlight irradiation are 1 order of magnitude higher than those for the process under UVA conditions. The exceptions are TiO$_2$-Ag and TiO$_2$-Au. For TiO$_2$-Ag, the $k'$ is, on average, twice as high using the natural sun in comparison with UVA (0.0076 ± 0.00017 L/min and 0.0036 ± 0.00032 L/min (average values), respectively), while for TiO$_2$-Au, the $k'$ determined for photocatalytic oxidation under sunlight condition is 2 orders of magnitude higher (0.01126 ± 0.0003 L/min) than that in UVA condition (0.00061 ± 0.00016 L/min). However, the highest $k'$ values were obtained for Pd-TiO$_2$ (0.00407 ± 0.00196 L/min and 0.03672 ± 0.0034 L/min (average values), respectively), UV A/Cat and Sun/Cat processes.

However, when the catalytic ozonation is compared with photocatalytic ozonation, the pseudo-first-order kinetic constants are similar with a slight advantage in favour of the UVA/Cat/O$_3$. But when COD removal is taken into account, the implementation of UVA is not reasonable.

In case of all processes, the pseudo-first-order kinetic constants for methylparaben were the lowest while the highest values were obtained for benzylparaben degradation.

**Proposed mechanism**

It is known that generally AOPs enable a high degree of degradation of hazardous aqueous contaminants in a short time. Its wide application is connected with $\cdot$OH generation in near ambient condition (temperature and pressure). The $\cdot$OH is one of the most powerful oxidants and the most commonly used in water treatment processes, due to its non-selectivity (Glaze et al. 1987). As presented above, many processes are qualified as AOP. The well known are processes involving combinations of O$_3$, H$_2$O$_2$, UV, TiO$_2$ and Fenton reagent. The mechanism of those processes is widely investigated. In the H$_2$O$_2$/UVC system, the cleavage of H$_2$O$_2$ with UVC light produces a quantum yield of two $\cdot$OH radicals per unit of radiation absorbed (Eq. (1)). However, the $\varepsilon_{254}$ value of H$_2$O$_2$ is very low (19.6 M$^{-1}$cm$^{-1}$) (Glaze et al. 1987), which determines its very high concentration to produce sufficient $\cdot$OH. Moreover, as can be seen in Fig. 1b, H$_2$O$_2$ is able to absorb photons below 300 nm, which excludes the UVA lamps.

H$_2$O$_2$ + h$v$→2$\cdot$OH

(1)

The Fenton process overcomes these drawbacks because the oxidation processes utilise the activation of H$_2$O$_2$ by iron salts which allows the generation of $\cdot$OH (Eq. (2)). Moreover, Fe$^{3+}$ can react with H$_2$O$_2$ and hydroperoxyl radical (so-called ‘Fenton-like reaction’), which leads to Fe$^{2+}$ regeneration (Eq. (3))

Fe$^{3+}$ + H$_2$O$_2$→HO$^-$ + $\cdot$OH + Fe$^{2+}$

(2)
UV A/Cat/O3 MP Pd-TiO2 0.04346  
O3/Cat MP Pd-TiO2 0.03345  

Process Paraben Type of catalyst k (1/min)  

O3/Cat  

MP Pd-TiO2 0.03346  
Pt-TiO2 0.03507  
Ag-TiO2 0.0204  
Au-TiO2 0.01817  
TiO2 0.02372  

EP Pd-TiO2 0.03071  
Pt-TiO2 0.03395  
Ag-TiO2 0.0187  
Au-TiO2 0.01641  
TiO2 0.02136  

PP Pd-TiO2 0.03267  
Pt-TiO2 0.03409  
Ag-TiO2 0.02043  
Au-TiO2 0.01801  
TiO2 0.02342  

BuP Pd-TiO2 0.02383  
Pt-TiO2 0.02019  
Ag-TiO2 0.01782  
Au-TiO2 0.01811  

BeP Pd-TiO2 0.05988  
Pt-TiO2 0.03812  
Ag-TiO2 0.03639  
Au-TiO2 0.03453  
TiO2 0.04889  

UVA/Cat/O3  

MP Pd-TiO2 0.04346  
Pt-TiO2 0.03698  
Ag-TiO2 0.03925  
Au-TiO2 0.02933  
TiO2 0.03886  

EP Pd-TiO2 0.04258  
Pt-TiO2 0.03443  
Ag-TiO2 0.03946  
Au-TiO2 0.02716  
TiO2 0.03816  

PP Pd-TiO2 0.04542  
Pt-TiO2 0.03725  
Ag-TiO2 0.04022  
Au-TiO2 0.02918  
TiO2 0.03506  

BuP Pd-TiO2 0.03978  
Pt-TiO2 0.03912  
Ag-TiO2 0.03673  
Au-TiO2 0.02427  
TiO2 0.02993  

BeP Pd-TiO2 0.05755  
Pt-TiO2 0.03777  
Ag-TiO2 0.05482  
Au-TiO2 0.03312  


Fe^{3+} + H_2O_2 → HO_2^- + H^+ + Fe^{2+}  

The reaction based on O_3 is more complicated. Firstly, the mechanism of its action is strongly dependent on pH (as discussed above): at high pH values, ozone acts not just by direct reaction but also by formation and reaction of ‘OH. The decomposition of ozone in a cyclic chain process can be indicated by hydroxide ion (Fig. 5a), H_2O_2 supplementation to the system or photolysis of ozone (Fig. 5b). Moreover, despite the two pathways, both the formations of hydrogen peroxide can occur, which can initiate chain decomposition of ozone, resulting in ‘OH formation (Glaze 1986). Therefore, even reactions that are performed below or near neutral pH may have a component of radical character, because peroxide is often a by-product of the ozonolysis (Glaze 1986). Moreover, the stoichiometric yield of ‘OH is greater from the photolysis of H_2O_2 (only 0.5 mol H_2O_2 is consumed for 1 mol of ‘OH production) in comparison to the photolysis of ozone (1.5 mol O_3 is consumed for 1 mol of ‘OH production) in comparison to the photoactivity of ozone (1.5 mol O_3 is consumed for 1 mol of ‘OH production) in comparison to the photoactivity of ozone (1.5 mol O_3 is consumed for 1 mol of ‘OH production) in comparison to the photoactivity of ozone (1.5 mol O_3 is consumed for 1 mol of ‘OH production) in comparison to the photoactivity of ozone (1.5 mol O_3 is consumed for 1 mol of ‘OH production). When O_3/H_2O_2 is considered, only 1 mol O_3 and 0.5 mol H_2O_2 are required to form 1 mol ‘OH (Glaze et al. 1987). However, taking into account the absorption coefficients, the higher amount of ‘OH is formed during O_3 photolysis (2 radicals per incident photon) than H_2O_2 photolysis (0.09 radicals per incident photon). Therefore, O_3/H_2O_2 and O_3/H_2O_2/UV have higher effectivity and are more cost-effective (Lucas et al. 2010).

It is known that in the TiO_2 photocatalytic process, the photocatalytic activity is exhibited by photogenerated charge carriers which is a consequence of the TiO_2 activation by radiation with energy higher than its band gap energy. Positive holes (h^+) are responsible for oxidising organic compounds due to reactions with a water molecule or OH ions and ‘OH radicals (Gmurek et al. 2017), while negative electrons (e^-) for reducing molecular oxygen to superoxide radical anions (Fig. 5c). Moreover, the generation of H_2O_2 from the electroreduction of oxygen occurred that may also lead to the production of ‘OH radicals (Salvador 2007; Xiao et al. 2015), while modification of TiO_2 by a noble metal changed the mechanism of electron transfer. The photons are absorbed by semiconductor (TiO_2) and/or also by metal nanoparticles through their LSPR excitation (mostly in case of Ag and Au) (Bumajdad and Madkour 2014; Zielińska-Jurek 2014; Wysocka et al. 2018); see Fig. 5d-f. The enhancement in photoactivity is attributed to Schottky barriers under UV range radiation and to LSPR that induces a collective coherent oscillation of the conducting band electrons of the metal nanoparticles. However, it should be mentioned that under UV irradiation (where energy is much higher than the surface plasmon resonance (SPR), the improvement in photoactivity is not related to SPR (Bumajdad and Madkour 2014). However, under visible range, the enhancement is strongly related with LSPR and Schottky barrier (Wysocka et al. 2018; Borowska et al. 2019). Moreover, the electron transfer through the noble metal/TiO_2 interface under a wide range (UVA and visible) occurred (Hu and Bürgi 2012). The differences between ozone-based or hydrogen peroxide–based AOP and photocatalysis with semiconductors modified by noble metals indicate that photodegradation can be attributed not only to ‘OH radicals but also to other forms of reactive oxygen species (ROS) O_2^-, 1O_2 and H_2O_2 or e_{cb}^- and h^+ (Wysocka et al. 2018).

The addition of ozone significantly enhances the mineralisation prior to heterogeneous photocatalysis alone or
even in case of catalysis. This synergetic effect is due to ready hydroxylation of the semiconductor catalyst in aqueous solution (Xiao et al. 2015). Heterogeneous photocatalytic ozonation is a very complicated system that involves chemical, catalytic and photocatalytic reactions. The improvement of the mineralisation during simultaneously occurring photocatalysis and ozonation synergistically intensified the formation of free radicals (Fig. 5g).

Based on the experiments of scavengers, the main degradation pathways via $\cdot OH$ were proved for $H_2O_2/Fe^{2+}$, $H_2O_2/UV C$, $O_3/H_2O_2$, $O_3/UV A$, $O_3/H_2O_2/UV A$, UVA/Cat, $O_3/Cat$ and $O_3/UV A/Cat$. It is known that oxidation of aqueous water contaminants can be proceeded via direct oxidation (the reaction of free holes/
electrons with adsorbed contaminates) or indirect oxidation (through oxidation by ROS). However, in photocatalytic processes in which the hydroxyl radicals were confirmed as the main oxidants, the other ROS present in the system (such as ozone, ozonide and superoxide radicals) have a lower effect on paraben degradation and by-product formation and further decay. As a main transformation by-product, $p$-hydroxybenzoic acid, 2,4-dihydroxybenzoic acid and 3,4-dihydroxybenzoic acid as well as monohydroxylated parabens were identified and confirmed the degradation pathway during AOP treatment accounted predominantly via hydroxylation (Fig. S6).

**Mineralisation, biodegradability and toxicity assessment**

For the tested AOPs, the degree of mineralisation was assessed. As discussed above, the mineralisation was expressed by COD. However, for a deeper analysis, total organic carbon (TOC) removal (after 2 h) will be discussed as well (Table S3). Between 80% and 90% COD reduction and 50% TOC removal were observed after 60 min of treatment. As discussed above, the highest COD removal was obtained when $\text{H}_2\text{O}_2$/UVC was used, which also corresponded with the highest TOC removal. Very good COD reduction was observed after $\text{O}_3$/H$_2$O$_2$ (70%), while TOC removal was equal to 26%. For all catalytic processes, the highest TOC reduction was achieved when TiO$_2$-Pd was used: UVA/Cat (25%), Sun/Cat (34%), $\text{O}_3$/Cat (18%) and UV/A/O$_3$/Cat (38%). In case of UVA/Cat, Sun/Cat and $\text{O}_3$/Cat, there were no significant differences between the other photocatalysts. The application of UVA/O$_3$/Cat process caused the highest COD as well as TOC removal (Table S3). The TOC and COD analysis showed that the total mineralisation was not achieved in any of the processes. It can be suspected that organic transformation by-products are still present in solution after treatments. Therefore, it is necessary to investigate the biodegradability as well as the toxicity of the solution after the treatment. The biodegradability study was based on the investigation of changes in the degree of oxidation, which is an indicator of the oxidation degree of complex solutions, providing indirect information on their probability of biodegradation. The COD/TOC values and average oxidation state (AOS) were used for biodegradability evaluation (Al Momani et al. 2004). The results are shown in Fig. 6.

![Fig. 6 Biodegradability assessment of COD/TOC and AOS B values obtained for the decomposition of paraben mixture after several AOPs](image-url)
transformation by-products decreased, despite high differences between COD and TOC removal, while when the catalytic ozonation was proceeded, the COD/TOC is higher than that for initial solution. Moreover, for TiO₂ and TiO₂-Ag, the values of that parameter were similar to single ozonation.

It is known that AOS is a valuable parameter that can be used as a general oxidation measure of complex mixtures containing various oxidation products. Theoretically, AOS can take values between +4 for CO₂ (the most oxidised state of C) and −4 for CH₄ (the most reduced state of C). For example, the AOS for benzene is −1, that for formaldehyde and acetic acids is 0 while that for formic acid and oxalic acid is +2 and +3, respectively (Al Momani et al. 2004). As can be seen in Fig. 6, almost all of the tested AOPs resulted in significant increases in the AOS values (initial AOS = −2.27). The AOS values after H₂O₂/UV, O₃/UVA, photocatalytic oxidation (both UVA and Sun) and photocatalytic ozonation confirmed that more oxidised by-products are obtained. However, it is still not as simple as carboxylic acids. The lowest oxidation degree was achieved in the case of single ozonation or catalytic oxidation when TiO₂ and TiO₂-Ag were used, confirming results from COD/TOC parameter.

As can be seen in Table 5 and as discussed above, almost all processes led to complete degradation of paraben concentration after 120 min. Moreover, when some processes have been applied, the removal of each paraben was achieved much faster. Only during photocatalytic oxidation the degradation after 3 h was incredibly low, and the decontamination has been influenced by the photocatalysts. For all effluents (containing paraben mixture) before treatment, the initial COD value as well as Alivibrio fischeri luminescence inhibition (LI) after 15 min of exposure were 102 ± 7.5 mg O₂/L and 95.57 ± 0.04%, respectively.

The best results of COD removal have been obtained when H₂O₂/UVC and H₂O₂/Fe²⁺ were applied, in spite of the lowest time of treatment (1 h). Contrarily, the worst COD abatement was obtained by photocatalytic oxidation (UVA) when the treatment time was the longest one (3 h). It should be noticed that during this process, no more than 50% removal of each paraben concentration was achieved, while during the other processes, the degradation of parabens was obtained before 120 min.

The initial paraben mixture was highly toxic to Alivibrio fischeri. During the reaction, the toxicity has been reduced. According to the literature, the sample can be considered non-toxic when LI is below 30% (Miralles-Cuevas et al. 2017). Taking this value into account, for the Fenton process and H₂O₂/UVC, non-toxic effluents have been achieved. Similar effect has been obtained for O₃/H₂O₂ and O₃/UVA (irrespective to COD removal). When the processes with photocatalysts were considered, large differences have been noticed. It was found

### Table 5: Maximum paraben concentration, COD removal, A. fischeri luminescence inhibition (15 min of exposure (LIₑ₅ mₜ)), C. fluminea mortality and L. sativum germination index (GI) after 120 min of treatment

| Process                | Max. C. parabens | COD (%) | LIₑ₅ mₜ | Mortality | GI (%) |
|------------------------|------------------|---------|---------|-----------|--------|
| H₂O₂/Fe²⁺              | 100%             | 40      | Nd⁰      | 37%       |
| H₂O₂/UVC               | 100%             | 41      | Nd⁰      |           |
| UVA/TiO₂               | 12% BuP          | 10      | > 80%    | ab/p      | (43 ± 0) |
| UVA/TiO₂-Pt            | 42% BuP          | 18      | > 80%    | 55%       | (57 ± 23) |
| UVA/TiO₂-Pd            | 73% BuP          | 19      | > 80%    | 29%       | (70 ± 15) |
| UVA/TiO₂-Ag            | 50% Al          | 17      | > 80%    | 21%       | (61 ± 9) |
| UVA/TiO₂-Au            | 16% BuP          | 8       | > 80%    | ab/p      | (44 ± 5) |
| O₃/H₂O₂                | 100%             | 70      | 57.1 ± 2.8 | 29%    |
| O₃/UVA                 | 100%             | 27      | 59.9 ± 1.8 | 0%     | (93 ± 11) |
| O₃/TiO₂                | 100%             | 28      | 26.5 ± 0.5 | 21%     | (71 ± 10) |
| O₃/TiO₂-Pt             | 100%             | 35      | 32.9 ± 1.0 | 0%      | (86 ± 4) |
| O₃/TiO₂-Pd             | 100%             | 35      | 33.7 ± 1.1 | 15%     | (90 ± 3) |
| O₃/TiO₂-Ag             | 100%             | 35      | 35.2 ± 3.9 | 0%      | (80 ± 1) |
| O₃/TiO₂-Au             | 100%             | 28      | 31.4 ± 1.4 | 0%      | (71 ± 5) |
| O₃/UVA/TiO₂            | 100%             | 38      | 36.3 ± 0.0 | 0%      | (90 ± 0) |
| O₃/UVA/TiO₂-Pt         | 100%             | 49      | 61.4 ± 1.2 | 0%      | (107 ± 11) |
| O₃/UVA/TiO₂-Pd         | 100%             | 41      | 44.2 ± 1.4 | 0%      | (108 ± 7) |
| O₃/UVA/TiO₂-Ag         | 100%             | 43      | 43.4 ± 1.4 | 0%      | (112 ± 1) |
| O₃/UVA/TiO₂-Au         | 100%             | 38      | 55.4 ± 1.0 | 0%      | (107 ± 5) |

For photocatalytic oxidation, the maximum removal is indicated (the paraben considered is given as superscript) ab/p abnormal behaviour and/or paralysis of C. fluminea, Nd not determined

*After 60 min of treatment

⁰The toxicity after the treatment time was too low to inhibit luminescence

⁰After 180 min of treatment
that the photocatalytic oxidation gave the worst results (LI15 min > 80%). According to the literature when LI is higher than 80%, the bioassay test is not sensitive to changes, which indicates still very high toxicity towards A. fischeri (Miralles-Cuevas et al. 2017). Catalytic ozonation gives much better toxicity results than photocatalytic ozonation, despite lower COD removal. This can be related with the by-product production. In fact, Gomes et al. (2019) verified that the presence of hydroxyl radicals for the same paraben mixture in a general way shows more toxicity than the single ozone due to the generation of different by-products.

In the case of Asian clams, for the same conditions, for the initial mixture of parabens, 100% mortality was achieved after 72 h of contact. It has to be mentioned that C. fluminea is intolerant at the concentration above 7.5 mg/L of each, and the LC50 of the paraben mixture solution is equal to 56% (Gomes et al. 2017a). Therefore, for all treatment processes (photolysis, UVA/TiO2 and UVA/TiO2-Au) where the degradation was not efficient enough to cause higher concentration, abnormal foot extension and/or paralysis was observed. However, for TiO2-Pt, TiO2-Ag and TiO2-Pd, the reduction of paraben toxicity towards C. fluminea after the treatment was observed. While after all catalytic ozonation experiments (except for TiO2-Pd- and Pd-TiO2), no mortality was observed. For the samples obtained after photolytic ozonation and photocatalytic ozonation with different photocatalysts, no mortality was verified for all concentrations tested after 72 h of exposure. The solution after the Fenton process exhibits higher mortality than O3/H2O2 and even after single ozonation (29%).

The high toxic impact of paraben mixture (GI = (42 ± 11)%) towards Lepidium sativum was investigated. According to Trautmann and Krasný (1997), this germination index proves strong or severe inhibition, while according to Wang (1992), the inhibition of the root growth of L. sativum can be classified as very strong or strong. Almost all treatments increased the phytotoxicity. The photocatalytic oxidation (UVA/TiO2, UVA/TiO2-Au, UVA/TiO2-Pt) slightly reduced toxicity; however, the inhibition is still strong. After single ozonation (GI = (65 ± 5)%) with UVA/TiO2-Pd and UVA/TiO2-Ag and catalytic ozonation with TiO2, TiO2-Au and TiO2-Ag (O3/TiO2 and O3/TiO2-Ag), just mild inhibition is observed, which means moderate phytotoxicity. While after O3/H2O2 and photocatalytic ozonation with all photocatalysts, no inhibition of plant growth was achieved.

**UV dose and cost-effectiveness**

Kinetics was also investigated from the UV fluence point of view (Fig. 7). Following these, the pseudo-first-order degradation rate constants for UV processes were also determined for the curves ln(C/C0) (as an average for all paraben degradations) vs. fluence (not shown), revealing linear correlation coefficients of 0.92–0.98.

It can be seen that the H2O2/UVC process which was the most effective in paraben decomposition is also the least time- and energy-consuming process. The application of UVA (8.9 W/m²) instead UVC (31.8 W/m²), despite the lower lamp power, decreased dramatically kf. Even additional oxidant (O3) did not cause the comparable degradation values, proving that H2O2/O3/UVA and O3/UVA are much more energy-consuming. The highest value was obtained for TiO2-Pd [(7.63 ± 3.69) × 10−6 cm²/J and (8.96 ± 0.34) × 10−5 cm²/J], for photocatalytic oxidation processes and photocatalytic ozonation, respectively. Also for both processes, TiO2-Ag was very effective [(6.80 ± 0.59) × 10−6 cm²/J for UVA/TiO2-Ag and (8.56 ± 0.27) × 10−5 cm²/J for UVA/TiO2-Ag/O3]. For the rest of photocatalysts, the behaviour was different for the case of UV A/Cat and UV A/Cat/O3. When the photocatalytic oxidation was performed, the kf value decreases in the following order: TiO2-Pt ((3.58 ± 1.63) × 10−6 cm²/J) > TiO2-Au ((1.14 ± 0.23) × 10−6 cm²/J) > TiO2 ((0.72 ± 0.07) × 10−6 cm²/J), while for that system where O3 was supplemented, the UV fluence–based pseudo-first-order rate constant for TiO2 ((8.35 ± 0.49) × 10−5 cm²/J) was higher than that for TiO2-Pt ((7.40 ± 0.23) × 10−5 cm²/J) > TiO2-Au ((5.84 ± 0.34) × 10−5 cm²/J).
These results indicate that UVA/Cat processes are the most energy-consuming process. Moreover, it was shown that the application of UVA not only increases the time of treatment decreasing mineralisation, which leads to higher energy consumption which determines the treatment costs.

If the application of these methods is to be considered in actual wastewater treatment, the energy cost as well as the operational cost must be known. In this regard, specific energy consumption (SEC) which defines the amount of electrical energy consumption (kWh) per unit mass of COD was calculated (Saien et al. 2014; Shen et al. 2017). When the process was based on the photochemical reaction, Eq. (4) was used (Saien et al. 2014), while in O₃-based processes, SEC was calculated according to Eq. (5) (Shen et al. 2017).

\[
\text{SEC} = \frac{P \times t}{V \times (\text{COD}_0 - \text{COD}_t)} \quad (4)
\]

\[
\text{SEC}_{O_3} = \frac{P \times t + r \times \text{TOD}}{V \times (\text{COD}_0 - \text{COD}_t)} \quad (5)
\]

where \( P \) is the nominal electric power (kW) of the photochemical system, \( t \) is the reaction time (h), \( r \) is the energy requirement for O₃ production (15 kW h/kg O₃) (Katsoyiannis et al. 2011) and \( V \) is the volume (L) of the solution in the reactor.

The cost of a treatment from the electricity point of view was found by multiplying the cost of electricity with the SEC value. In the total operational costs of a treatment, several factors were included: oxidants costs, operating costs for ozone production (1.97 €/kg O₃ (Ried et al. 2009) and costs of catalyst (synthesis and the metal modification).

It can be seen that the lowest SEC was calculated for O₃, O₃/H₂O₂ and Cat/O₃ due to the lack of requirement of light source application, while for processes UVA/Cat/O₃, O₃/H₂O₂ and O₃/UVA, SEC is 1 order of magnitude lower than that for UVA/Cat and H₂O₂/UVC (it is obvious that SEC for Sun/Cat is equal to 0). Despite the UVA energy consumption for the decomposition, the UVC treatment is much higher in energy consumption.
cost. Besides the cost of electricity, there are also expenses for hydrogen peroxide and ozone production. However, the costs of photocatalyst synthesis (especially the cost of noble metals) are predominant in the catalytic process. Despite the highest efficiency, TiO$_2$-Pd is characterised by one of the highest operational costs, due to the Pd price, while for TiO$_2$-Ag, the costs are positioned around 197 €/g COD. In case of those catalysts, the energy consumption is not too high if we consider it from the TOC and COD decrease point of view. The results indicate that the use of 0.42 kWh/g COD causes 37% and 43% of TOC and COD decrease, respectively (Fig. 8, Table S3), which provides a significant and attractive advantage.

**Conclusions**

The comparison of several radical-driven technologies for mixture degradation of five parabens was presented. Several AOP methods were examined to degrade parabens from aqueous environment as well as to decrease its toxicity. The ozonation in different process parameters was discussed, bearing in mind that combining ozone with H$_2$O$_2$, UVA as well as noble metal modified TiO$_2$. The pseudo-first-order kinetic constants (based on time and on UV fluence) were determined. The biodegradability and toxicity assessment has been presented, considering the toxic effect over several species.

For all applied processes, the same trend was observed. The BuP was the fastest degraded paraben, then followed in the order BuP > PP > EP > MeP. The treatments reduced not only COD but also toxicity towards *V. fischeri, C. fluminea* and *L. sativum*, despite the fact that the total COD removal was not achieved. This indicates that the refractory by-products produced are less toxic than parabens.

These results can indicate that the degradation pathway is different, despite the hydroxyl radicals are the main oxidant. The next explanation is that the same TPs formed at the last stage of degradation are in higher concentration. This finding can be with agreement that lower COD removal gives better toxicity results. However, it should be also noticed that the application of pure TiO$_2$ leads to lower toxicity and lower COD removal in comparison to photocatalysts modified by noble metals. Furthermore, from all metals used for TiO$_2$ modification, platinum seems to be the best. Moreover, it is evident that the application of ozone not only improves detoxification, but also higher mineralisation is observed after O$_3$-modification, despite the fact that the total COD removal was not achieved. This indicates that the refractory by-products produced are less toxic than parabens.

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The results reveal that the treatment efficiency as well as the costs largely depend on the implemented AOP. Nevertheless, the application of hydrogen peroxide, ozone with second oxidant (UVA or H$_2$O$_2$) and TiO$_2$ modified by a noble metal led to much short depletion times when compared with photolysis. Moreover, the synergetic effect not only improved mineralisation and biodegradability but also reduced toxicity and energetic costs.

However, the application of natural sunlight for the photocatalytic oxidation process reduced the cost of lamp equipment as well as electricity, but taking into account the highest mineralisation and biodegradability, the most appropriate cost-effective treatment seems to be photocatalytic ozonation with a UVA lamp. Moreover, detailed characterisation of photocatalysts showed that surface modification of TiO$_2$ with noble metals significantly improved the absorption properties of the catalyst resulting in higher COD abatement from aqueous solutions compared to pure TiO$_2$. This statement was also supported by the width of the band gap of the photocatalysts. The most promising photocatalysts seem to be TiO$_2$ modified by Pd nanoparticles, due to the highest band gap and highest mineralisation. However, when the total operating cost will be taken into account, more reasonable is to use the TiO$_2$-Ag that gives satisfactory mineralisation.

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