Catalytic Membrane Ozonation

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Definition: Catalytic membrane ozonation is a hybrid process that combines membrane filtration and catalytic ozonation. The membrane deposited with an appropriate solid material acts as catalyst. As a consequence, the catalytic membrane contactor can act simultaneously as contactor (i.e., improving the transfer/dissolution of gaseous ozone into the liquid phase), as well as reactor (i.e., oxidizing the organic compounds). It can be used in water and wastewater treatment limiting the disadvantages of membrane filtration (i.e., lower removal rates of emerging contaminants or fouling occurrence) and ozonation (i.e., selective oxidation, low mineralization rates, or bromate (BrO₃⁻) formation). The catalytic membrane ozonation process can enhance the removal of micropollutants and bacteria, inhibit or decrease the BrO₃⁻ formation and additionally, restrict the membrane fouling (i.e., the major/common problem of membranes’ use). Nevertheless, the higher operational cost is the main drawback of these processes.

Keywords: membrane catalytic ozonation; ozone; membrane filtration; hybrid process; fouling control; micropollutants; bromate formation; cost analysis; water treatment

1. Introduction

Clean water is very essential for the sustainable human future in planet Earth. However, water can be extensively contaminated and the development of innovate processes for its effective treatment and use constitutes a constantly evolving field [1]. Among the most recently detected pollution issues is considered the occurrence of emerging/persisting contaminants/chemicals in the aquatic ecosystems and especially, the presence of micropollutants. Micropollutants are termed the compounds of mostly anthropogenic origin that occurring usually in the aqueous environment in concentrations lower than mg/L (i.e., in the range of µg/L, or even ng/L). The evolution of analytical science and techniques especially during the recent years has permitted the accurate detection of them [1,2]. They can be divided into the following six main categories: (1) perfluorinated compounds (e.g., perfluorooctanesulfonate, perfluorooctanoic acid), (2) disinfection by-products (e.g., halomethanes, nitrosamines, hydroxyl acids), (3) gasoline additives (e.g., tert-butyl ether, benzene, 1,3-butadiene), (4) manufactured nanomaterials (e.g., carbon nanotubes, silicon dioxide, titanium oxide), (5) human and veterinary pharmaceuticals (e.g., estradiol, 17α-ethinylestradiol, acetaminophen), and (6) sunscreens/ultraviolet filters (e.g., benzophenone-3, homosalate, 4-aminobenzoic acid) [2]. Micropollutants are omnipresent and generally contribute to improving the quality of human life, but they are consisting also an issue of major concern for the water and wastewater treatment plants. Micropolllutants are mostly persist in the treated effluents, because the conventional systems are not specifically designed to remove them effectively [3]. The optimization of these treatment plants maybe a sufficient strategy for the efficient treatment of these problematic contaminants. Due to the respective developments the relevant scientific literature, regarding the treatment of emerging/persistent contaminants, has highly increased during the past few years, as shown in Figure 1.
Figure 1. Yearly published scientific articles considering the emerging contaminants treatment from Science Direct [4].

The decontamination technologies generally have to be effective, cheap and eco-friendly in order to be widely applicable [1]. The Advanced Oxidation Processes (AOPs) present relevant advantages; among them belongs the membrane catalytic ozonation, a hybrid technique that can combine the processes of catalytic ozonation and of membrane filtration [4].

2. Catalytic Ozonation

Catalytic ozonation is a highly studied advanced oxidation process (AOP) and it is based on the acceleration of hydroxyl radicals’ production (HO•) through the addition of appropriate materials (solids or dissolved), acting as catalysts. Ozone can be self-decomposed into hydroxyl radicals via a complex mechanism that can be summarized in the following reaction [5]:

$$3\text{O}_3 + \text{HO}^- + \text{H}^+ \rightarrow 2\text{HO}^* + 4\text{O}_2$$

(1)

However, the addition of an appropriate catalyst can further increase the respective reaction rate and also, can decrease the relevant energy demands, which are considered as the main disadvantage of single ozonation process. Two main types of catalysts can be used in the catalytic ozonation technique, i.e., either several transition metals in dissolved form (leading to the “homogeneous” catalytic ozonation), or several solid materials, acting through their surface as potential catalysts (leading to the “heterogeneous” catalytic ozonation) [5]. Recently, the combination of membrane filtration with ozone has been studied extensively and this hybrid process can be performed via three modes, in which the ozonation can takes place before or after the membrane filtration, or both processes can operate simultaneously. In the catalytic ozonation the last (combined) mode is usually applied, while there are rather few studies, regarding the application of the other two alternative techniques [6–8].

When a solid material is used as catalyst, catalytic ozonation can be also divided in two sub-categories. Whether the catalyst is a solid material, suspended in the aqueous phase (usually in powdered form), the process is termed as “single heterogeneous catalytic ozonation”; whereas the process is commonly referred as “membrane catalytic ozonation”, whether the solid material is appropriately immobilized onto the surface of membrane [4]. In this hybrid ozonation process the membrane can act simultaneously as catalyst for the acceleration of the hydroxyl radicals’ production, as well as a membrane contactor for the improvement of transfer and diffusion of ozone gas to the aqueous phase during treatment [9].
3. Catalytic Membrane Contactors

Membrane contactors (of polymeric or ceramic nature) nowadays are used in the gas-liquid processes, permitting the effective transfer of a constituent (usually gaseous) between a liquid and a gas phase [10]. In the case of ozonation process the ozone gas is diffused into the liquid phase most commonly by the application of injectors, bubble diffusers, or static mixers [11]. However, these techniques produce bubbles of rather larger diameters that are decreasing the respective mass-transfer rates, leading to un consumed/unreacted ozone losses, and therefore to higher operational costs [12]. Alternatively, the application of membrane contactors is considered as a more eco-friendly technology [13] that can decrease the operational costs and increase the efficiency of the ozonation process, although the initial capital costs are higher, than the more conventional ones [10]. When solid materials are being used in ozonation processes (e.g., as catalysts) the nature of them should be carefully considered. Ozone is a strong oxidant agent that reacts and decomposes most organic compounds, including the polymeric materials [2]. As a consequence the use of common polymeric membrane contactors are quite limited [12]. It seems that only polytetrafluoroethylene (PTFE), polyvinylidene fluoride (PVDF), and polydimethylsiloxane (PDMS) polymeric membranes are presenting sufficient resistance, when contacting with ozone [4,14]. Alternatively, ceramic membranes can be often selected [12]. Considering the removal of micropollutants, the nano-filtration membranes are generally more efficient, mainly due to their smaller pore size and lower pressure operation, than, e.g., the reverse osmosis membranes [15].

Nevertheless, the applied contactors are mainly membranes and therefore, fouling may commonly occur, being among the main drawbacks of membrane application process. The fouling appears when organic particles/colloids and/or microorganisms are deposited on the surface of the membrane, increasing the trans membrane pressure (TMP). This incident acts as an additional resistance and decreases the efficiency of the process. It can be divided into four types: (a) complete pore blocking, (b) intermediate pore blocking, (c) standard pore blocking, and (d) surface cake layer. The formation of a cake layer in the surface of the membrane is usually observed [15].

The virgin membranes ordinarily do not present any particular catalytic activity; appropriate metal oxide particles have to be deposited (usually) onto them in order to accelerate the production of hydroxyl radicals. However, some materials, such as Al₂O₃, TiO₂, and ZrO₂, may present an inherent catalytic activity and can act as catalytic ceramic membrane contactors without further modification [16,17]. The most frequently used oxides are the iron and manganese oxides. Park et al. [18] coated a γ-Al₂O₃ membrane with iron oxide nanoparticles and Chen et al. [19] coated a relevant membrane with a TiO₂ and a Ti-Mn layer. In both cases the membrane fouling was significantly reduced through the increase of hydroxyl radicals’ production during the oxidation process. The same was observed also by Corneal et al. [20], when the manganese oxide (α-Mn₂O₃) was used as catalytic layer. The removal of micropollutants is based upon the same radicals’ production mechanism. Zhu et al. [21] deposited CeO₂ on (inorganic) titania membrane to examine the removal of tetracycline by the application of catalytic ozonation. The concentration of micropollutant was reduced by 85%, due to the oxidation process, while the adsorption on the membrane surface contributed also to the removal. Furthermore, Guo et al. [22] deposited CuMn₂O₄ particles in a ceramic membrane, and observed that superoxide ions (O₂•⁻) and hydrogen peroxide (H₂O₂) were produced during the oxidation process, which have the ability to accelerate further the production of hydroxyl radicals [5].

In most relevant ozonation cases the ceramic membranes have been commonly applied in the catalytic ozonation systems, because they are fully ozone resistant and the substrate material is simpler to be used as coating substrate. However, Yu et al. [23] who have coated a PVDF membrane with MnO₂ nanoparticles; they reported that the presence of these oxide particles lowered the fouling rate and the presence of ozone has limited the bio-accumulation of several components onto the membranes’ surface, hence restricting the fouling problem. Furthermore, Sun et al. [24] have doped a similar membrane with TiO₂
nanoparticles to examine the removal of nitrobenzene. Under acidic pH values the removal efficiency of this pollutant was rather low, but it was further increased with the increase of pH values up to 10, due to the higher production of hydroxyl radicals. Nevertheless, a further increase of pH value leaded to a quick decompose of hydroxyl radicals into simple ions, hence reducing the removal rate. Zhu et al. [25] used for the ozonation treatment of municipal wastewater the same catalytic membrane and observed that only the larger molecular weight compounds were degraded into smaller ones. The produced micromolecules could not be removed effectively by this membrane, and thus the overall COD value of the wastewater was reduced only by 27.6%. Figure 2 shows schematically the fundamental mechanism of catalytic membrane ozonation process.

![Figure 2. Fundamental mechanism of catalytic membrane ozonation process (adapted from [4]).](image)

In order the catalytic effect to be employed the ozone dosing must be continuous. When only a single ozone dose at the beginning of the process is applied, then the oxidation takes place at the first min of the reaction and the ozone concentration in the system is quickly substantially decreasing; as a result membrane filtration takes over and the membrane fouling can occur [16,26].

There are five main factors that influence the catalytic performance of a membrane: (1) surface charge; (2) coating times; (3) pore size; (4) preparation method regarding the metal oxide deposition; and (5) adsorption capacity [15,18,27]. More specific information about how these factors are influencing the catalytic membrane ozonation can be found in the respective reference of Psaltou and Zouboulis [4].

Almost all researchers, studying this process, have observed that the catalytic membrane ozonation follows a mechanism based on the production of radicals. The prominent radical species in most cases were the hydroxyl ones, noting also that in some cases both hydroxyl and superoxide radicals were found to participate in the oxidation process [27]. However, Scarati et al. [28] have noticed that when CuO was used as (catalytic) coating for membranes, then the primary radical species were the superoxide radicals, which however did not lead to the further production of hydroxyl radicals. Although, the superoxide radicals are part of the ozone decomposition mechanism towards the formation of hydroxyl radicals [29,30], it seems that in the aforementioned case the decomposition pathway was different. However, the obtained results were not very promising and the catalytic ozonation with the use of this coated membrane may not be sufficiently effective, when applied for the removal/treatment of micropollutants.

Except for the removal of micropollutants and the prevention of membrane fouling some researchers have used the membrane catalytic ozonation process also for the removal of bacteria. The relevant studies have shown that bacteria adhere to the metal oxide surface,
increasing their retention time at the membrane and subsequently, their contact with ozone will be prolonged, improving the efficiency of oxidation. The inactivation of bacteria occurs by the presence of radical species and especially by the hydroxyl radicals, similar with the case of micropollutants removal. Therefore, the radical mechanism can enhance also the disinfection behavior of this treatment system [31,32]. All the aforementioned studies are summarized in Table 1.

Table 1. Summary of studies regarding wastewater treatment by the application of membrane catalytic ozonation.

| Catalytic Membrane | Target Parameter/Pollutant | Efficiency | Reference |
|--------------------|---------------------------|------------|-----------|
| α-Al₂O₃            | Color TOC (WWTP effluent)  | 68%        | [16]      |
| TiO₂               |                           | 21%        |           |
|                    |                           | 88%        |           |
|                    |                           | 43%        |           |
| Cement             | p-chloro-nitro-benzene    | 90%        | [17]      |
| IONs * / α-Al₂O₃   | p-chlorobenzoic acid (River water) | 56% | [18] |
| Ti-Mn/TiO₂/Al₂O₃  | NOM (aquaculture wastewater) | 52.1% | [19] |
| α-Mn₂O₃/CéRAM     | Flux (Lake water) (flux recovery) | 95% | [20] |
| CeO₂-TiO₂/α-Al₂O₃ | Tetracycline              | >80%       | [21]      |
| CuMn₂O₄/ZrO₂/α-Al₂O₃ | Benzophenone              | 76.6%      | [22]      |
| MnO₂/PVDF         | Model raw water Prevention of membrane fouling | | [23] |
| Nano-TiO₂/PVDF    | Nitrobenzene              | 59.5%      | [24]      |
| Nano-TiO₂/PVDF    | COD (Municipal wastewater after primary treatment) | 27.6% | [25] |
| TiO₂/α-Al₂O₃      | Color TOC (Municipal wastewater treatment) | 88.5% | [26] |
|                   |                           | 48.7%      |           |
| CuO/α-Al₂O₃-ZrO₂  | 1,4-dioxane               | 65%        | [28]      |
| Iron Oxide/CéRAM  | E. Coli (Lake water) >99%(mortality) | | [31] |

* IONs: Iron Oxide Nanoparticles.

4. Important Issues of Consideration, When Applying the Ozonation Process

4.1. Bromate Formation

Although, ozone has been applied widely in drinking water and in wastewater treatment, its use may cause secondary problems, when the water to be treated contains bromide (Br⁻) at concentrations higher than 50 µg/L. The Br⁻ concentrations in natural waters may usually range between 4 and 1000 µg/L [33]. Due to salt intrusion, Br⁻ is detected in surface waters, as well as in the influent of wastewater treatment plants at concentrations up to 400 µg/L [34]. The main by-product of the reaction between ozone and Br⁻ is bromate (BrO₃⁻). BrO₃⁻ once formed, it is neither biodegradable nor removable by the application of conventional treatment [33]. It is considered as a potential carcinogen (Group 2B) by the International Agency for Research on Cancer (IARC) [34]. The maximum permissible contaminant level for BrO₃⁻ is 10 µg/L, according to USEPA and Drinking Water Commission of Europe [35]. Due to that the minimization or the prevention of its production in water treatment is crucial [33]. Figure 3 shows the formation of BrO₃⁻ via
the direct ozonation and through the radical mechanism, noting that BrO$_3^-$ is mostly produced by the 2nd (radical) pathway. It is reported that in Milli-Q water experiments, almost 70% of BrO$_3^-$ formation occurs through the hydroxyl radicals oxidation reactions and the remaining 30% depends on the molecular ozone reactions [36]. More specific details about the reaction pathways, regarding the production of BrO$_3^-$ in ozonation processes can be found in Fischbacher et al. 2015 [37].

![Oxidation Reaction Diagram](image)

**Figure 3.** Comparison of the molecular ozone mechanism and the $^*$OH mechanism in the presence of bromide anions (Br$^-$) for the production of bromates (adapted from [35]).

There are four main factors, affecting BrO$_3^-$ formation, in ozonation processes [33]:

1. **pH value:** In alkaline pH region the BrO$_3^-$ formation is higher. The reaction rate constants of Br$^-$/OBr$^-$ with molecular ozone and with hydroxyl radicals are 160 M$^{-1}$s$^{-1}$/330 M$^{-1}$s$^{-1}$ and $1.1 \times 10^9$ M$^{-1}$s$^{-1}$/4.5 $\times 10^9$ M$^{-1}$s$^{-1}$, respectively. When the solution pH increased, more hydroxyl radicals are formed, leading to more Br$^-$ and hypobromide ions (OBr$^-$), which are subsequently oxidized to form BrO$_3^-$ [34].

2. **Initial bromide concentration:** The production of BrO$_3^-$ increase with the increase of Br$^-$ concentration.

3. **Ozone dose:** The BrO$_3^-$ concentration increases with an increased ozone dose. In membrane ozonation there is a strong linear relationship between the BrO$_3^-$ concentration in the permeate and the dissolved ozone concentration [36].

4. **Temperature:** The BrO$_3^-$ concentration increases with increasing temperature up to 25 °C, because at relatively higher temperature (>25 °C) the ozone stability also becomes a major factor. Additionally, according, to Henry’s law, the efficiency of ozone solubility in an ozonation system decreases with the increase of temperature. Under these conditions the ozone concentration could be even below the respective threshold limit required for BrO$_3^-$ formation [38]. Table 2 shows the ozone escape to the gas phase at the third min of ozonation process, as well as the respective half-lives at various temperatures.

**Table 2.** Ozone escape to the gas phase at the 3rd min of ozonation process and its typical half-life at various temperatures [39,40].

| Temperature (°C) | Ozone Escape (%) | Half-Life (min) |
|-----------------|------------------|-----------------|
| 15              | 7.7              | 30              |
| 25              | 9.4              | 15              |
| 35              | 11.1             | 8               |
In terms of \( \text{BrO}_3^- \) formation control, the application of heating process is financially unprofitable. Cooling down the ozonation system is preferable, if there is already an adequate cooling device in the treatment system [38]. The reduction of pH/temperature or the addition of carbonate alkalinity are more effective ways to control the \( \text{BrO}_3^- \) production; however, it is neither cost effective, nor practical considering full-scale applications [34,36]. Furthermore, the control of these parameters may present also negative effects regarding the micropollutants removal, e.g., by slowing down the production of hydroxyl radicals [39]. Therefore, a key challenge is to develop a process that can effectively degrades micropollutants and simultaneously minimizes the formation of \( \text{BrO}_3^- \) [36]. To overcome these drawbacks, the heterogeneous catalytic ozonation process was introduced as a potential solution [34]. However, catalytic ozonation can enhance the production of hydroxyl radicals and because the hydroxyl radicals are mainly responsible for the \( \text{BrO}_3^- \) formation, its application must be carefully studied [41].

Liu et al. [38] applied the catalytic ozonation process with the use of different oxides to control \( \text{BrO}_3^- \) formation. The addition of magnesium (MgO) and titanium oxide (TiO\(_2\)) promoted the formation of \( \text{BrO}_3^- \), while the presence of KMnO\(_4\) inhibited that production. More particularly, by increasing the KMnO\(_4\) dose the noted inhibition was further enhanced. Increased production of hydroxyl radicals was observed in the MgO/O\(_3\) and TiO\(_2\)/O\(_3\) systems. On the contrary, the ozone decomposition, but not towards hydroxyl radicals, was promoted by the KMnO\(_4\) addition and as a consequence, the oxidation ability of the treatment system was reduced.

The most applicable catalyst, as reported in the literature for the elimination of \( \text{BrO}_3^- \) formation, is CeO\(_2\). Wang et al. [42] showed that CeO\(_2\) can be a good inhibitor for the \( \text{BrO}_3^- \) production. However, the organic micropollutants removal efficiency by using the O\(_3\)/CeO\(_2\) process was rather small. The same results were reported by Zhang et al. [43], who observed that the inhibition effect was optimal at slightly acidic conditions, while at the neutral pH value the efficiency of process was decreased considerably. This limitation significantly restricts the application of this catalyst in drinking water treatment processes.

Ye et al. [41] showed that the \( \text{BrO}_3^- \) formation rate by using graphene oxide (GO) as catalyst was almost twice, than the case where the single ozonation was applied. Moreover, by adding Tert-Butyl-Alcohol (TBA), the \( \text{BrO}_3^- \) formation was decreased to 25%, because the hydroxyl radicals’ production was limited. However, when the 0.36CeO\(_2\)/RGO (reduced graphene oxide) material was used as catalyst, the degradation efficiency of DEET (N,N-Diethyl-meta-toluamide) was enhanced during the ozonation process, but adsorption experiments were not conducted, aiming to eliminate the possibility this micropollutant (DEET) to be simply adsorbed by GO. Alongside, Ye et al. were compared the formation of \( \text{BrO}_3^- \) by the addition of various metal oxides in the ozonation system. After 10 min, the \( \text{BrO}_3^- \) production during the O\(_3\), O\(_3\)/CeO\(_2\), O\(_3\)/FeOOH, O\(_3\)/Fe\(_2\)O\(_3\), or O\(_3\)/ZnO processes were found 1.41, 0.59, 1.34, 0.92, and 1.74 \( \mu \text{M} \), respectively. All these concentrations were lower than the \( \text{BrO}_3^- \) produced during the application of single ozonation except for the O\(_3\)/ZnO process, in which 1.74 \( \mu \text{M} \) of \( \text{BrO}_3^- \) was formed. CeO\(_2\) achieved an inhibition efficiency of 58% for \( \text{BrO}_3^- \) formation; ozone adsorbed to CeO\(_2\) active sites and the decomposition of ozone to hydroxyl radicals was decreased. Furthermore, as Shen et al. [44] observed, the use of a relevant catalyst was found to decreased the degradation efficiency of p-CNB (p-chloro-nitro-benzene), as this compound can be practically removed only by hydroxyl radicals’ mechanism.

The material LaCoO\(_3\) was also examined by Zhang et al. [34], as catalyst for the elimination of \( \text{BrO}_3^- \) formation and for the removal of organic compounds during the application of catalytic ozonation. When the concentration of benzotriazole (BEN) was 0.148 mg/L, the formed \( \text{BrO}_3^- \) reached the highest concentration and the Br\(^-\) conversion rate was 89.4%. However, when the BEN concentration increased, then the formed \( \text{BrO}_3^- \) decreased significantly. This might be due to the competition reactions taken place between BEN and Br\(^-\) with the ozone molecules and the hydroxyl radicals. The formed \( \text{BrO}_3^- \) in catalytic ozonation was lower than that formed during the application of single ozonation
at each pH value, suggesting that catalytic ozonation is more efficient process for the reduction of both BrO$_3^-$ and micropollutant (BEN). At the pH value 7.22 (i.e., a typical value for a drinking water treatment effluent) the degradation of BEN was reduced simultaneously [34]. However, this study did not show whether the BEN reduction can be considered enough for a micropollutant treatment process. In general, as von Gunten et al. [35] reported, BrO$_3^-$ formation can be lower in an advanced oxidation processes, e.g., when compared to single ozonation, if the ozone dosage is kept constant.

Another way (and probably more effective) to lower the BrO$_3^-$ formation is the use of membranes. Moslemi et al. [36] observed that under constant operating pressure, as the molecular weight cut-off (MWCO) of the membrane increased, the permeate flux also increased, while the residence time in the treatment system decreased. Therefore, for the membranes presenting higher MWCOs, the ozone exposure (i.e., the retention time multiplied by the dissolved ozone concentration) was reduced. Therefore, in such hybrid systems there is a another (6th) factor that affects the BrO$_3^-$ formation, which is the contact (residence) time of ozone with Br$^-$ in the membrane and the entire treatment system.

Hamid et al. [45] studied also the formation of BrO$_3^-$ using a hybrid filtration-ozonation system. The results suggested that the BrO$_3^-$ formation can be minimized when using ceramic membranes. The BrO$_3^-$ formed by the application of membrane ozonation was 50% lower, than that of the ozonation using a conventional diffuser. The reduced BrO$_3^-$ formation observed in this study may be due to the lower ozone exposure with Br$^-$, due to the catalytic formation of hydroxyl radicals (even in small extent) at the surface of the membrane. The reduction of BrO$_3^-$ production could also explained by the surface properties of ceramic membrane that can cause the attraction or repulsion of ions. When a charged membrane is in contact with an electrolyte solution, the electrostatic repulsion will repel ions with the same charge, because the concentration of these ions will be generally lower near the membrane surface and within the membrane pores; inversely, the membrane surface will attract ions having the opposite charge, resulting in higher concentrations (locally) of them. The surface properties of ceramic membranes can play an important role and are worthy to be explored further as an additional mechanism for BrO$_3^-$ control.

Merle et al. [46] combined homogeneous catalytic ozonation with membranes in the so-called MEMBRO$_3$X process, noting also that for the efficient removal of p-chlorobenzoic acid (p-CBA) are required large H$_2$O$_2$ dosages (5.67 mg/L). At this dose and for an ozone concentration 1.0 g/Nm$^{-3}$ around 98% of p-CBA was degraded and about 6 µg/L of BrO$_3^-$ was formed, whereas the BrO$_3^-$ formation were about 79% higher in comparison with the conventional peroxone reaction. The key feature of the MEMBRO$_3$X process is the addition of ozone over multiple injection points. In ozone-based processes for BrO$_3^-$ formation can be limited by keeping the O$_3$ concentration low enough, which is achieved by dosing it in small quantities over the membrane contactor device and transforming it efficiently to hydroxyl radicals, which can further react with the micropollutants [35].

Until now there is no study conducted about the control of BrO$_3^-$ formation during the catalytic membrane ozonation process. This process, due to the dual role played by the membrane (i.e., acting as a diffuser and as a reactor), has the potential to remove efficiently the micropollutants, as well as to minimize the BrO$_3^-$ formation and simultaneously, to disinfect the treated water; therefore, presenting certain significant advantages. Table 3 summarizes the studies about bromate formation during catalytic ozonation process.
Table 3. Bromate formation during the application of catalytic ozonation process.

| Used Catalyst | Experimental Conditions | Bromate Formation (µg/L) | Reference |
|---------------|-------------------------|--------------------------|-----------|
| LaCoO$_3$     | [Br$^-$] = 100 µg/L,  [O$_3$] = 2 mg/L, pH = 6.41 | 84.4 | [34] |
| MgO           | [Br$^-$] = 50 mg/L,  [catalyst] ≈ 4 mg/L | ≈ 80 | [38] |
| TiO$_2$       | [O$_3$] = 0.4 mg/L,  pH = 7 | ≈ 135 | |
| KMnO$_4$      | [catalyst] ≈ 0.25 g/L, pH = 6.41 | ≈ 35 | |
| CeO$_2$       | [Br$^-$] = 1 mg/L,  [catalyst] = 0.5 g/L, pH = 7 | 75.5 | |
| FeOOH         | [O$_3$] = 10 mg/L/min, time = 10 min | 171.4 | |
| Fe$_2$O$_3$   | [catalyst] = 0.5 g/L, pH = 7, T = 18 °C, Time = 30 min | 117.7 | [41] |
| ZnO           |  | 222.5 | |
| CeO$_2$       | [Br$^-$] = 1.8 mg/L,  [catalyst] = 0.1 g/L, pH = 6.30, T = 18 °C, Time = 30 min | 205.58 | |
| MgO           | [O$_3$] = 5.21 mg/L,  [catalyst] = 0.1 g/L, pH = 6.20, T = 15 °C, Time = 30 min | 471.01 | |
| FeOOH         |  | 351.53 | [42] |
| CeO$_2$       | [Br$^-$] = 2 mg/L,  pH = 6.20, T = 15 °C, Time = 30 min | < 300 | |
| α-FeOOH       | [O$_3$] = 1.41 mg/L,  [catalyst] = 0.1 g/L, pH = 6.20 | ≈ 350 | |
| γ-FeOOH       |  | < 350 | |
| α-Fe$_2$O$_3$ | [catalyst] = 0.5 g/L, pH = 8.1, T = 20 °C, time = 25 min | ≈ 450 | |
| MEMBRO$_3$X   | [Br$^-$] = 180 ± 4 µg/L,  [O$_3$] = 5 g/m$^3$,  [catalyst] = 5.67 mg/L, pH = 8.1, T = 20 °C, time = 25 min | ≈ 35 | [46] |

4.2. Operational Cost Evaluation

Except for the BrO$_3^-$ formation another major drawback of ozonation is the relatively high energy consumption that leads to higher operation costs. In general there are two main parameters that can make a technology suitable for the industry: (1) Technical, and (2) Economical feasibility [47]. High capital, operation and maintenance costs reflect negatively on any applicable method [48]. High energy consumption leads to a costly unit with a larger footprint treatment [49] and influences the operational cost. In the single ozonation process ozone dose is the main factor that influences in large extent the relevant cost. Table 4 shows, the operational costs, according to different ozone doses, as well as the micropollutant degradation efficiency obtained for each dose. Increasing ozone dosages from 2 to 6 mg/L leads to 64% cost increase.

Table 4. Electrical Energy and Operating Cost required for the ozonation process (adapted from [50]).

| Ozone Concentration (mg/L) | Electrical Energy (kWh m$^{-3}$) | No. of PPCPs Removed by ≥ 90% Efficiency from a Total Number 37 | Operational Cost (Yen m$^{-3}$) |
|---------------------------|-----------------------------------|---------------------------------------------------------------|---------------------------------|
| 2                         | 0.03                              | 24                                                            | 0.5                             |
| 4                         | 0.06                              | 32                                                            | 0.9                             |
| 6                         | 0.09                              | 35                                                            | 1.4                             |

The composition of the influent to be treated also affects the operational cost of the process, although indirectly, through the determination of the necessarily applied
ozone dosage, because some pollutants are less degradable than others and they need higher oxidant concentrations to be efficiently treated. Furthermore, the applied ozone concentration is related to the initial micropollutant concentration [48]. Table 5 shows the removal efficiency of different micropollutants with the use of different ozone dosages and the respective operational cost of the treatment system. As becomes obvious, phenol having 100 times higher initial concentration, needs 3 times lower ozone dose to be effectively removed, than the tri-chloro-ethylene (TCE). TCE with reaction rate constant with ozone \((k_{O3})\) equal to 17 M\(^{-1}\)·s\(^{-1}\) [51] is more difficult to be oxidized, than phenol presenting much higher \(k_{O3}\), equal to \(1.3 \times 10^3\) M\(^{-1}\)·s\(^{-1}\) [52].

**Table 5.** Cost estimation for the removal of different pollutants by the application of ozonation (adapted from [47]).

| Pollutant          | Ozone Concentration (mg/L) | Pollutant Concentration (mg/L) | Efficiency (%) | Operational Cost ($/1000 gal) |
|--------------------|-----------------------------|--------------------------------|----------------|-------------------------------|
| Phenol             | 2                           | 235.28                         | 90             | 1.2023                        |
| Reactive azo dye   | 12.4                        | 20                             | 90             | 4.0839                        |
| TCE                | 6                           | 2.2                            | 40.9           | 2.3549                        |

Advanced oxidation processes (AOPs) are generally more complex to handle (e.g., for ozone generation, light sources, pumps, etc.) and are presenting usually higher energy costs. For that reason, they are still categorized as rather expensive processes [53]. However, among the various AOPs the application costs differs widely. Catalytic and photocatalytic ozonation are two of the main processes in this category. Table 6 shows the energy consumption and the treatment cost for the removal of oxalic acid from aqueous solutions by the application of three different AOPs. All these processes in the study were performed with the use of TiO\(_2\) as catalyst for comparison reasons. The TiO\(_2\) proved to be inefficient catalyst for the catalytic ozonation process and the removal of the micropollutant was quite small, noting also that the treatment cost of catalytic ozonation may have been decreased with the use of an optimum catalyst. Furthermore, considering the oxidation of oxalic acid, 25 mg/L and 70 mg/L of O\(_3\) was applied in the catalytic and photocatalytic ozonation processes respectively (in the gas phase), which means that the ozone concentration in the liquid phase was about 10 mg/L and 25 mg/L [53]. Thus, the photocatalytic ozonation seems more cost efficient treatment process because the relevant economic parameters were calculated based on the cost per micropollutant removal and not per treated volume.

**Table 6.** Specific energy consumption and treatment costs of photocatalytic oxidation, photocatalytic ozonation and catalytic ozonation for the removal of oxalic acid (adapted from [53]).

| AOPs               | Specific Energy Consumption (kWh/mM) | Treatment Costs (euro/m\(^3\)) |
|--------------------|--------------------------------------|-------------------------------|
| Catalytic ozonation| 0.017                                | 3.65                          |
| Photocatalytic oxidation | 0.063                              | 13.55                         |
| Photocatalytic ozonation | 0.007                              | 1.51                          |

Another parameter that can affect the cost of catalytic ozonation process is the specific type of the catalyst and its potential reusability [54]. For example, Biard et al., [55] select ruthenium among other metals (Co, Pd, Pt, and Rh) not only for its better reactivity, but also for its lower cost (about 30 euros/g). From the same perspective Martins et al. [54] examined three low-cost materials (i.e., volcanic rock, sepiolite, and iron shavings) as potential catalysts for the treatment of olive mill wastewater by the application of heterogeneous catalytic ozonation. Iron shavings proved to be the most efficient catalyst among the others, when the same ozone dose was applied; therefore the ozonation system can achieve the lower operational cost.
5. Limitations and Future Perspectives

Although the catalytic membrane ozonation seems to be a quite promising technology, the stability of applied coatings may be considered as an important issue of concern. In several cases the initially deposited metals can be leached back into the aqueous solution during the reaction procedure, leading to a secondary pollution problem. Even for the metals that specific regulation limits have not been yet established (e.g., for the case of Sn) their secondary occurrence in drinking water supply may raise health issues for the living organisms that should be carefully addressed [56, 57]. Furthermore, the formation of by-products during the oxidation process must be studied in depth, as it possibly constitutes an additional pollution source. These by-products can be produced due to the incomplete mineralization of organic pollutants and their acute toxicity in some cases is even higher than the initially oxidized compounds [58]. Gomes et al., [58] studied the ecotoxicity of a five parabens mixture (methyl-, ethyl-, propyl-, benzyl-, and butylparaben) and their by-products. They found that although catalytic ozonation could degrade efficiently the parabens, the treated sample was more toxic than the initial one. It presented higher toxicity over D. Magna, R. subcapitata and L. savitum. The most toxic by-products were hydroquinone and 1,4-benzoquinone. Hydroquinone is also one of the by-products of quaiacol oxidation, a well-known chemical intermediate for food materials and perfumery products [59]. Therefore, the conduct of toxicity assays is essential.

In terms of materials science, the coating techniques should be further developed to facilitate the further use of catalytic membranes. New solid materials, deposited into the membranes surface to convert them into catalytic ones, should be also examined. Additionally, ozone-resistant polymeric membranes or coated materials that make them more stable against the contact with ozone should be developed and accompanied with new techniques that make easier the deposition of catalytic materials. These developments are expected to lead to an increase of catalyst ranges/options, raising further the effective/economic applications of catalytic membrane ozonation process.

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