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The role of boundary conditions in the bromide-enhanced ozonation process for ammonia nitrogen removal and nitrate minimization / Ruffino, Barbara; Zanetti, Mariachiara. - In: JOURNAL OF ENVIRONMENTAL CHEMICAL ENGINEERING. - ISSN 2213-3437. - 8:(2020), p. 102933. [10.1016/j.jece.2019.102933]

Availability:

This version is available at: 11583/2850957 since: 2020-11-04T11:24:11Z

Publisher: Elsevier Ltd

Published

DOI:10.1016/j.jece.2019.102933

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1 The role of boundary conditions in the bromide-enhanced ozonation

2 process for ammonia nitrogen removal and nitrate minimization

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     Barbara Ruffino ^{1,2,*} and Mariachiara Zanetti ^1
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      <sup>1</sup>Department of Environment, Land and Infrastructure Engineering, Politecnico di Torino, corso Duca
 6
 7
      degli Abruzzi 24, 10129 Torino, Italy
     <sup>2</sup>CleanWaterCenter@PoliTo, Politecnico di Torino, corso Duca degli Abruzzi 24, 10129 Torino, Italy
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15
      *Corresponding author
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17
      Barbara RUFFINO
      DIATI, Department of Environment, Land and Infrastructure Engineering
18
      Politecnico di Torino
19
      Corso Duca degli Abruzzi, 24
20
21
      10129 Torino, ITALY
22
     Ph. +39.011.0907663
     Fax +39.011.0907699
23
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e-mail: barbara.ruffino@polito.it

Abstract

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In this study we investigated the effect of some operating conditions, such as the ratio between ammonia nitrogen (AN), ozone and bromide, the initial value of water pH, the amount of bicarbonate used as a buffering substance and the possible interference of organic matter, in a bromide-enhanced ozonation process for the removal of AN. One of the peculiarities of this process is that the oxidation of AN proceeds through the generation of reaction intermediates (bromoamines) and ends with molecular nitrogen (N₂) with none or limited generation of nitrate. The results of the ozonation tests carried out at a lab scale demonstrated that nitrate generation could be lowered to less than 10% of the theoretical value when three conditions coexist. Firstly, all the AN was present in the form of ammonium ion (NH₄⁺); this happened when the pH of the system was lower than 8. Secondly, the ratio between bromide and ozone species was optimized, so as to avoid the presence of free ozone and the consequent direct reaction with AN. Finally, the acidification induced by the oxidation of AN was controlled with a buffering substance (bicarbonate). The generation of inorganic bromoamines from the reaction between AN and bromide inhibits the oxidation of bromide to bromate. In fact, in the tests carried out in this study the formation of bromate was limited to values of less than 0.2% of the molar bromate yield. Organic substances in the form of glucose or potassium hydrogen phthalate (KHP) did not affect the removal rate of AN. Conversely, the presence of substances such as phenol and hydroquinone, that had a higher reactivity towards ozone and HOBr, determined a reduced AN removal rate and a bromate generation of one order of magnitude higher than that observed with KHP and glucose. Finally, all the

organic substances employed in the study seemed to have a positive effect in reducing the nitrate

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Keywords: ozone, pH, bicarbonate, bromate, organic matter

generation (-30% of the value obtained with sole AN).

Introduction

- Ammonia (NH₃) and ammonium ion (NH₄⁺), which can be transferred to each other by a pH depending
- equilibrium (shown in (1), pKa = 9.23), are collectively named as ammonia nitrogen (AN).
- 53 $NH_4^+ \leftrightarrow NH_3 + H^+ (1)$
- AN is present in almost all types of wastewater, from municipal, agricultural and industrial origin. High
- concentrations of ammonia are especially found in wastewaters generated from the industry sector,
- 56 including electronic, petrochemical, pharmaceutical, fertilizer, food and steel manufacturing industries
- 57 (Lin et al., 2018). AN is a source of nitrogen, which is a nutrient for algae and other forms of plant life
- 58 when released into the environment, and thus contributes to eutrophication of receiving water bodies. In
- 59 addition, AN in fresh waters causes acute toxicity to fish species, already at concentrations of less than
- 60 10 mg/l (Park et al., 2015). For these reasons, processes for removal of AN from ammonia-rich streams
- has gained increasing attention in recent years. Furthermore, because of the more and more stringent
- discharge limits for AN that have been steadily imposed on wastewater treatment plants (WWTPs)
- 63 worldwide, AN needs to be removed from wastewater to very low concentration values before discharge
- and preferably decomposed into harmless compounds.
- 65 The state-of-the-art treatment technologies for AN removal include traditional biological nitrification –
- denitrification, anaerobic ammonium oxidation (ANAMMOX) and, especially for wastewaters from
- 67 industrial origin, air stripping (Hsieh et al., 2013; Yuan et al., 2016), chemical precipitation (Cheng et
- al., 2016), eventually with the aim of recovering valuable compounds, ion exchange, adsorption, break-
- 69 point chlorination, reverse osmosis (Jin et al., 2013), electrodialysis and hollow fiber membrane contactor
- and membrane distillation (Lin et al., 2018). Each of these technologies has its own limitation. For
- 71 example, due to the toxic nature of AN in industrial wastewater, biological treatments are often not
- appropriate (Charmot-Charbonnel et al., 1999). Bacteria used in these processes are very sensitive and
- cannot withstand wide ranges of pH and temperature, halogen compounds, cyanides, and other heavy
- metals present in ammonia-containing water (Moussavi et al., 2009). Air-stripping creates additional air
- 75 pollution when AN is converted from liquid to gas phase. Chemical precipitation generates sludge, from
- 76 which the extraction of valuable products is not often possible, and leaves substantial amounts of AN in
- exhaust waters even after recovering it from concentrated streams in the form of (NH₄)₂SO₄ (Ulbricht et
- al., 2013). Ion exchange and reverse osmosis generates concentrated streams that have to be further
- 79 subjected to treatment.
- 80 In the last years, oxidation processes, where AN can be oxidized to molecular nitrogen, eventually
- 81 enhanced by using catalytic substances, have attracted increasing attention (Moussavi and

82 Mahdavianpour, 2016). Various metal oxide catalysts such as Co, Fe, Ni, Zn can improve the performances of a conventional oxidation of AN with ozone (Ichikawa et al., 2014). The most efficient 83 84 catalysts (Co₃O₄, MgO and NiO) could increase the rate of AN decomposition up to 3-5 times and, because of the high selectivity to gaseous products, including N₂ and N₂O, suppressed the formation of 85 soluble compounds such as nitrate. Recently, Chen and coauthors (2018) found that a catalyst with 8:2 86 MgO-Co₃O₄ molar ratio, calcined at 500°C for 3 hr, allowed to obtain an AN removal rate of 85.2% and 87 88 gaseous nitrogen selectivity of 44.8%. Among the catalytic oxidation processes, ozonation enhanced by bromide can be included. The main 89

90 peculiarities of the role of bromide in an ozonation process were firstly discovered in early Eighties (Haag et al., 1984). During ozonation, bromide is converted to hypobromous acid (HOBr). The mechanism of 91 92 the process is similar to that of ammonia breakpoint chlorination. The difference is that in an ozonation 93 process, HOBr can be continuously produced by the oxidation of bromide and bromide can be utilized 94 repeatedly. HOBr is known to be a stronger and more effective oxidant than the only ozone in the oxidation of AN (von Gunten, 2003). The oxidation process of AN proceeds through the generation of 95 96 reaction intermediates (bromoamine) and ends with molecular nitrogen (N₂) as in the series of reactions reported in Tanaka and Matsumura (2003): 97

- 98 $HOBr + NH_3 \rightarrow NH_2Br + H_2O$ (1a)
- 99 $HOBr + NH_2Br \rightarrow NHBr_2 + H_2O$ (1b)
- $100 \quad HOBr + NHBr_2 \rightarrow NBr_3 + H_2O \quad (1c)$
- 101 $2H_2O + NHBr_2 + NBr_3 \rightarrow N_2 + 3Br^- + 3H^+ + 2HOBr$ (1d)

Some patents, developed by Japanese inventors in the Nineties, used the basics of the bromide-enhanced 102 103 oxidation process for the treatment of wastewaters with a high content of AN deriving from the deionization of cooling waters of power stations (Nitto Kikai KK and Permelec Electrode LTD, 1994; 104 105 Japan Organo CO LTD, 1997). However, the above-mentioned patents did not report the effect of the operating parameters, that is both process parameters and the characteristics of the solution that 106 underwent the treatment, on the performance of the process in detail. Some years later, Yang and 107 108 coauthors (1999; 2000), by using water samples that reproduced the characteristics of the real 109 wastewaters from cooling towers (containing N-NH₄⁺, Na⁺, SO₄⁼ and very low amount of organic carbon, 1-4 mg/l), demonstrated that the formation of nitrate was mainly affected by the Br-/N ratio, N load, and 110 pH. The application of the bromide-enhanced ozonation process may be of great attractive for the 111 112 treatment of cooling waters, because aqueous bromine is often used as a primary biocide in cooling tower

disinfection applications. It is highly effective at inactivating bacteria responsible for forming biofilms, 113 such as Pseudomonas aeruginosa (Boal, 2016). In this case bromide should not be added for the 114 115 treatment. Later, Tanaka and Matsumura (2003) treated real samples of spent brine that already contained bromide 116 at the concentration of 100 mg/l. They proved that increasing concentrations of bromide had positive 117 effect on AN removal and that a pH decrease slowed down the reaction. Brines, other than ammonia, 118 119 contained small amounts of organic matter (COD = 50 mg/l). Ozone had also effect on COD removal but the competition between AN and organic carbon for the use of the oxidant (HOBr) was not 120 121 completely elucidated. More recently, Khuntia and coauthors (2013), using a batch apparatus equipped with a fine diffuser to sparge ozone-enriched oxygen microbubbles, observed that the rate of oxidation 122 123 of ammonia by ozone becomes faster in the presence of bromide ions. Samples used for the tests were on purpose made and contained only inorganic species such as NH₄Cl, (NH₄)₂SO₄, Na₂CO₃, HCl. The 124 125 authors concluded that the advantage of using bromide is that the nitrate formed by the oxidation of ammonia is ultimately converted to nitrogen. 126 127 As pointed out by the afore-mentioned experiences, the peculiarities of the bromide-enhanced ozonation process could be used for a fast and effective treatment of effluents from industrial origin that contain 128 from moderate to high AN loads. However, for the application of the process at a real scale, the effect of 129 operating conditions needs to be elucidated, such as the initial pH of the water, the presence of buffering 130 substances, the ratio between AN, ozone and bromide, and the possible interference of organic matter on 131 the transformation of the species of concern (AN, nitrate). This work wants to identify, and subsequently 132 tune, the operating conditions that determine the minimization of nitrate generation in a bromide-133 134 enhanced ozonation, with a regard to the problem of bromate formation. Secondly, it wants to provide preliminary insights concerning the competition between AN and organic carbon for the use of the 135

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2. Materials and Methods

oxidant (HOBr).

2.1 Experimental set-up and ozone quantification

Ozone was generated from oxygen gas (99.95% v/v) using an ozone generator (Ozone Lab TM, Ozone Services Division of Yanco Industries, Canada). Fluxes of oxygen and ozone-enriched oxygen (after ozone generation) of 200 ml/min were carried to and from the ozone generator through Teflon tubes. All tests were carried out in batch modality in a 300 ml glass gas washing bottle equipped with a filter disc

- for ozone diffusion. Volumes of solutions of 200 ml were subjected to ozonation treatments. All tests
- took place at a room temperature value of $20 \pm 2^{\circ}$ C.
- The dose of ozone distributed to the solution was quantified by using the iodometric wet-chemistry
- method (Rakness et al., 1996). The ozone enriched oxygen flow was bubbled through a 1% potassium
- iodide (KI) solution. Ozone oxidizes iodide to iodine, according to reaction (1)
- 149 $O_3 + 2H^+ + 2I^- \rightarrow O_2 + I_2 + H_2O$ (1)
- and the generated iodine was titrated with a 0.1 N solution of sodium thiosulfate ($Na_2S_2O_3 \cdot 5H_2O$).
- 151 The concentration of ozone in water was quantified by using the method described in Bader and Hoigné
- 152 (1981), that is based on the reaction of ozone with potassium indigo trisulfonate ($C_{16}H_7N_2O_{11}S_3K_3$) and
- the detection of the residual concentration of indigo at 600 nm.

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2.2 Experimental procedure and analytical parameters determination

- In this study four series of tests were carried out. The first series was aimed at investigating the effect of
- $[Br^-]/[O_3]$ and $[NH_4^+]/[Br^-]$ ratios on AN depletion and generation of nitrate. Solutions were prepared by
- dissolving NH₄Cl, NaHCO₃ (both of analytical grade and purchased by Sigma Aldrich) amounts in order
- to obtain final concentrations of 200 mg NH₄⁺/L (11.1 mM) and 1000 mg HCO₃⁻/L (16.4 mM). Aliquots
- of a 2000 mg/L solution of potassium bromide (KBr) were added in order to obtain bromide
- 161 concentrations that ranged from 44.3 to 282 mg Br-/L (0.55 3.53 mM). $[NH_4^+]/[Br^-]$ ratios were in the
- range 20.0 3.15 mM/mM. The generation of by-products (specifically bromate) was measured on the
- 163 system with $[Br^{-}] = 3.33 \text{ mM}.$
- The second series of tests was aimed at investigating the effect of the presence of sodium bicarbonate
- 165 (NaHCO₃), as a buffering agent, on the progress of AN oxidation and generation of nitrate. Solutions
- were prepared by keeping constant the concentrations of NH_4^+ (200 mg/L, 11.1 mM) and bromide (88.6
- mg/L, 1.11 mM), for a $[NH_4^+]/[Br_-] = 10.0$. Bicarbonate content was varied from 0 to 1000 (distributed
- in two aliquots, 500 + 500) mg HCO₃-/L.
- The third series of tests was used for investigating the effect of the initial pH value of the solution on the
- progress of the AN oxidation and generation of nitrate. Solutions were prepared by keeping constant the
- 171 concentrations of NH₄⁺ (200 mg/L, 11.1 mM), bicarbonate (1000 mg/L, 16.4 mM) and bromide (136
- mg/L, 1.70 mM), with a resulting [NH₄⁺]/[Br⁻] ratio of 6.51. The natural pH of the solution
- (approximately 8) was corrected to 9 by adding some drops of a 0.1 M NaOH solution.
- A fourth series of tests was used to obtain preliminary information concerning the possible interference
- of organic substances in the HOBr oxidation of AN. Five solutions containing AN (as NH₄Cl, 90.0 mg/L,

5.0 mM), bicarbonate (as NaHCO₃, 460 mg/L, 7.5 mM) and bromide (as KBr, 266 mg/L, 3.33 mM) were prepared. One solution was used as a reference. To each of the other four solutions an organic substance (glucose, potassium hydrogen phthalate (KHP), hydroquinone, phenol, all of analytical grade, purchased by Honeywell) was added to obtain a 0.5 M concentration. The compounds chosen for the test are representative of substances that are widely diffused into the environment. Specifically, phenolic compounds, like phenol and hydroquinone, have often been used as model compounds to describe substances that are naturally present in freshwater, as constituents of natural organic matter (NOM), or in wastewater, after conventional treatments, as effluent organic matter (EfOM) (Bond et al., 2009). Residual AN and alkalinity, nitrate and bromate were measured after a 30-min ozonation. Analytical parameters (AN, nitrate, pH, alkalinity, bromate) were determined according to Standards Methods (APHA, AWWA, WEF, 2005).

Table 1. Summary of the conditions used for the tests

Series of tests	N-NH ₄ ⁺ (mM)	HCO ₃ - (mM)	Br ⁻ (mM)	[N-NH ₄ ⁺ / Br ⁻]	рН	Organic (mM)
I	11.1	16.4	0.55 - 3.53	20.0 - 3.15	8	0
II	11.1	0 - 16.4	1.11	10.0	8	0
III	11.1	16.4	1.70	6.51	8-9	0
IV	5.0	7.5	3.33	1.50	NC	0.5

NC: not corrected

3. Results and Discussion

3.1 Effect of the [Br⁻]/[O₃] ratio on AN depletion and nitrate generation

193 It was verified, by using the iodometric wet-chemistry method, that the dose of ozone distributed to the 194 solution, through the ozone-enriched flow (200 mL/min) with the employed set-up, was of 26.6 mg/min. 195 The ozone concentration reached in the aqueous phase at steady-state, determined with potassium indigo 196 trisulfonate, was of approximately 0.20 mM.

Figure 1 shows the trend of AN depletion in four solutions with an identical initial concentration of AN, equal to 200 mg NH₄+/L (11.1 mM), due to an oxidation process where ozone species (i.e. ozone and HO* radicals) and bromide were contemporaneously present in the system. For the systems shown in Figure 1, the [Br-]/[O₃] ratio was of 5.54, 8.51, 11.1 and 16.6 (mM Br-/mM O₃) for bromide concentrations of 88.6, 136, 177 and 266 mg/L (1.11, 1.70, 2.22 and 3.33 mM), respectively. It can be seen that processes from approximately 70 to 120-minute long, depending on the bromide concentration, were required to reduce the concentration of AN from 200 mg/L to zero. Then, the resulting AN removal

- rates were of 1.80, 2.36, 2.62 and 2.79 mg $NH_4^+/L \cdot min (0.100, 0.131, 0.145 \text{ and } 0.155 \text{ mM/min})$ for [Br⁻
- 205]/[O₃] ratios of 5.54, 8.51, 11.1 and 16.6 (mM Br⁻/mM O₃), respectively. As expected, the increase in
- bromide concentration made the oxidation process faster. In fact, in an oxidation process with ozone and
- bromide, it must be assumed that the oxidation of AN due to HOBr, also known as "active bromine"
- 208 (Khuntia et al., 2013), that proceeds through the generation of the series of bromoamines (see Equations
- 209 1 a-d), overcomes the direct oxidation of AN due to ozone, as described in Equation (2)
- 210 $4O_3 + NH_3 \rightarrow NO_3^- + H^+ + H_2O + 4O_2$ (2)
- The rates of the two processes differ by several order of magnitude: $k_1 = 8 \cdot 10^7 \text{ M}^{-1} \text{s}^{-1}$ (Haag and Hoigné,
- 212 1983; Haag et al., 1984; Tanaka and Matsumura, 2003), in the case of HOBr, vs. $k_2 = 1 \text{ M}^{-1}\text{s}^{-1}$ (at pH=8)
- 213 (Haag et al., 1984), in the case of the sole ozone.
- 214 The AN removal rates observed in this study were in good agreement with the capacity of ozone and
- active bromine to oxidize AN found by Khuntia and coauthors (2013). They studied the effect of the
- 216 combination of bromide and ozone in the removal of AN and the consequent generation of nitrate. By
- using an ozone dose of 5.6·10⁻⁷ kg/s (equivalent to 33.6 mg/min, approximately 25% higher than that
- used in this study) and a molar ratio between AN and bromide of 18:1 (approximately 30 mg Br-/L) they
- obtained an AN removal rate in the order of 1.3 mg $NH_4^+/L \cdot min$.
- 220 It can be observed that the oxidation process followed a zero-order kinetic, as described by Equation (3)
- 221 $[AN]_t=-m\cdot t + [AN]_0$ (3)
- In a zero-order process the removal rate, m, is independent on the concentration of AN that remains in
- 223 the solution time after time. This result was in agreement with the findings of previous studies (Tanaka
- and Matsumura, 2002; Ruffino and Zanetti, 2011; 2012), that had demonstrated that a bromide-aided
- ozonation process, carried out on solutions containing concentrations of AN of more than 10 mg/l,
- followed a zero order kinetic. Then, the removal rate, m, could be assumed as a constant value for almost
- 227 all the duration of the process. The starting pH of the three systems was in the order of 8.0 8.1 and it
- decreased to 6.8-6.5 at the end of the process (data not shown).
- Figure 2 shows the effect of the [Br]/[O₃] ratio on the removal rate of AN (m, mg NH₄+/L·min). In the
- presence of $[Br^-]/[O_3]$ ratios from 2.8 to 18 $[mM Br^-]/[mM O_3]$, the observed removal rates of AN were
- in the range $1.4 3.0 \text{ mg NH}_4$ ⁺/L·min (0.078 0.166 mM/min). It is interesting to observe that the trend
- of the AN removal rates could be fitted well by a line in the left part of the diagram, that is for [Br⁻]/[O₃]
- ratios from 2.8 to approximately 11-12. Conversely, moving towards the right part of the diagram, that
- is for [Br]/[O₃] ratios of more than 11-12, the trend of the AN removal rates tended to an asymptote. The
- trend of the AN removal rates shown in Figure 2 suggested that, above a fixed value of the [Br⁻]/[O₃]

to form the "active bromine" (i.e. HOBr) and that subsequent increases in the bromide concentration were not able to further increase the oxidizing power of the solution. It can be stated/concluded that the oxidizing power of the solution was limited by the amount of ozone able to dissolve in the solution. With the employed set-up and oxygen flow (200 ml/min) and in the presence of the highest bromide concentration, i.e. when all the ozone that was dissolved into the solution had reacted with the available bromide to form active bromine, the specific consumption of ozone to remove AN was in the order of 60 mg O₃/mg NH₄⁺ (22.5 mmol O₃ / mmol NH₄⁺). Additionally, Figure 2 shows the relation between the [Br-]/[O₃] ratio and the capacity of the process of generating nitrate. For each test carried out at a different value of the [Br-]/[O₃] ratio, Figure 2 shows the percentage of generated nitrate, compared to the theoretical value. The theoretical value of nitrate concentration is the value that would be found if all the AN was oxidized to nitrate. As shown in Figure 2, an increase in bromide concentration, at a fixed dissolved ozone concentration value, determined a decrease in the actual nitrate generation. The generation of nitrate decreased to less than 10% of the theoretical value for [Br-]/[O₃] ratios higher than 6. As in the case of the removal rate of AN, even the trend of the nitrate generation showed a plateau for [Br-]/[O₃] ratios of more than 10-12. The observed trend can be justified because all the ozone species were involved in the formation of HOBr and could not react with AN to form nitrate. As reported in Figure 2, for a [Br-]/[O₃] ratio equal to 1.5, the generation of nitrate was approximately 40% of the theoretical value. This value was in good agreement with the findings of Berne and coauthors (2004) that observed concentrations of nitrate of 105 µM and 260 μM (for [NH₄⁺]/[Br⁻] ratios of 2 and 5, respectively), after a complete depletion of the AN, which were approximately 50% of the initial concentration of AN and, then, of the theoretical nitrate

ratio, all the ozone species (ozone and HO* radicals) available in the solution had reacted with bromide

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3.2 Effect of bicarbonate concentration on AN depletion and nitrate generation

As demonstrated from the results of Figure 2, even if nitrate was not formed (or was formed to a very small extent), the oxidation of AN determined a release of H⁺ ions in the solution and a consequent acidification. Previous studies demonstrated that the pH is a key parameter in the process of oxidation of AN by active bromine and that the inhibition of AN oxidation started from pH values around 5.0 (Tanaka and Matsumura, 2003). Consequently, in order to avoid a too fast acidification of the system, a buffering agent must be added before the sparging of ozone begins.

Figure 3 shows the effect of increasing concentrations of sodium bicarbonate, NaHCO₃, a buffering 267 substance, on both the performance of the AN oxidation process and the trend of the pH value in the 268 269 solution after treatments with durations from 15 to 120 minutes. This second series of tests was carried out at a [Br-]/[O₃] ratio of 5.54. From the results of Figure 3 it can be observed that, without the presence 270 of a buffering source, the pH of the solution prepared for the test naturally assumed the value of 6 (see 271 the light blue bar). pH dropped to the value of 3.5 fifteen minutes after the beginning of the test. Without 272 bicarbonate, the removal rate of AN decreased from approximately 1.5 mg NH₄+/L·min, in the first 273 fifteen minutes of the test, to values of less than 0.5 mg NH₄⁺/L·min in the remaining part of the test. 274 275 Amounts of bicarbonate in the order of 200 or 500 mg HCO₃-/L (i.e. 1 mg HCO₃-/mg NH₄+ or 2.5 mg HCO₃-/mg NH₄+) were not sufficient to make the oxidation of AN complete (to bring the oxidation to 276 277 completion) without determining a decrease in the pH to values of less than 4.0. From Figure 3 it can be 278 seen that the addition of a dose of bicarbonate of 200 mg HCO₃-/L had been able to buffer the release of 279 H⁺ for approximately 30 minutes (i.e. that dose could guarantee the depletion of approximately 25% of the total amount of AN), while a concentration of 500 mg/l guaranteed a steady development of the 280 281 process for approximately 60 minutes. In order to bring the removal of AN to completion without inducing an irreversible acidification of the 282 solution, the process described by the blue bars (Figure 3) was stopped after 60 minutes and a new amount 283 of bicarbonate was added to the solution. The dose added at minute 60th was such to bring the HCO₃⁻ 284 285 /NH₄⁺ ratio (by weight, mg/mg) to 5. Ideally, after the addition, the overall concentration of bicarbonate 286 in the solution was of 1000 mg/L (500 + 500 mg/L). The effect of the NaHCO₃ addition was the increase in the pH value observed for the blue bars from 60 to 90 minutes (Figure 3). 287 288 Consequently, it could be calculated an overall bicarbonate specific consumption in the order of 5 mg HCO₃-/mg NH₄⁺ to maintain the pH value in the range of neutrality for all the duration of the treatment. 289 290 These observations were in good agreements with the findings of a previous study that involved the 291 treatment of spent brine, where the initial concentrations of AN and bromide were of 13 mM and 106 (or 292 208) mg/l, respectively (Tanaka and Matsumura, 2003). The authors of this study observed that when all the alkalinity was depleted, ammonia was difficult to be oxidized and the removal rate decreased from 293 approximately 0.8 mM h⁻¹ to zero.

As shown in Figure 3, the decrease in the pH value had also a clear effect on nitrate generation. In the 295 296 systems where the oxidation of AN was not completely buffered (i.e 0; 200 and 500 mg NaHCO₃/l), the 297 generation of nitrate, expressed as the ratio between the actual and the theoretical concentration of nitrate, 298 increased to reach values in the order of 40% (see the series of light blue and red bars, Figure 3).

Conversely, where the pH was maintained at a nearly constant value for all the duration of the oxidation process, with the aid of the buffer substance (see the blue bars), nitrate was generated at an approximately constant rate that was in the order of 25% of the theoretical value.

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3.3 Effect of the initial pH value on AN depletion and nitrate generation

- Values of pH of more than 9.0 displaces the equilibrium between the forms of AN towards ammonia
- 305 (NH₃) and this species is removed preferentially by stripping (Luo et al., 2015). Figure 4 shows the effect
- of the starting pH values of 8.0 and 9.0 on AN removal and the pH value found at the end of the test.
- Tests were carried out in the presence of a [Br-]/[O₃] ratio of 8.51 (Br- concentration of 136 mg/l). As
- 308 expected, the progress of AN oxidation determined a decrease in the pH value in both systems. However,
- the rate of AN removal for the system with a starting pH of 8.0 was approximately 10% faster than that
- of the system with a starting pH of 9.0 (2.36 vs. 2.15 mg NH₄+/L·min). After 75 minutes, the residual
- AN in the system with a starting pH of 9.0 was 24.3%, in that with a starting pH of 8 was only of 17.2%.
- The increase in the pH at minute 60th, for the system with a starting pH of 8.0, was due to an addition of
- NaHCO₃, that was necessary because the residual amount of alkalinity was too low to guarantee the
- 314 completion of the process in the neutrality field.
- As shown in Figure 4, the generation of nitrate seemed to be greatly affected by the starting pH of the
- system. The generation of nitrate for the system with a starting pH of 8.0 was approximately 20% of the
- 317 theoretical value. Conversely, the more basic system registered a generation of nitrate in the order of 35-
- 318 37% of the theoretical value. The generation of nitrate between the two systems differed by
- approximately 100%.
- This result clarifies the role of the AN dissociation (NH₃ + H⁺ \leftrightarrow NH₄⁺) on nitrate generation in an
- oxidation process with both ozone and active bromine. At pH 9.0 approximately 60% of the AN is present
- in the form of ammonia (NH₃), while at pH 8.0 the amount of ammonia is only of 6% and decreases to
- less than 2% at pH 7.5. The observed results could be explained only if ammonia (NH₃) was directly
- oxidized to nitrate, while the ammonium ion was involved in the process of bromoamine generation that
- ends with the liberation of nitrogen gas (Rahmadi and Kim, 2014).

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3.4 Bromate generation potential

- 328 It is well known that bromide during ozonation may be converted to bromate (BrO₃-), classified as a
- probable or likely human carcinogen (WHO, 2005), and for which many countries have established the
- maximum allowable level in drinking water at 10 ug/L. Bromate is produced during ozonation through

- a multistep process that involves the oxidation of bromide (Br⁻), ensuing generation of HOBr and bromite
- 332 (BrO₂-) and their reactions with ozone (Fischbacher et al., 2015; von Gunten, 2003; von Gunten and
- Oliveras, 1998). Previous studies have revealed that many factors, including O₃ dosage, contact time,
- pH, and Br- and AN concentrations, influenced BrO₃- formation during ozonation (Antoniou and
- Andersen 2012; Lin et al., 2014; Pinkernell and von Gunten 2001).
- Figure 5 relates the residual concentration of AN, after a 30 or 60-minute ozonation process, to the molar
- bromate yield for the system containing $[NH_4^+] = 11.1 \text{ mM}$; $[HCO_3^-] = 16.4 \text{ mM}$ and $[Br^-] = 3.33 \text{ mM}$.
- 338 Molar bromate yield can be defined as the dimensionless ratio of molar bromate concentrations
- normalized by the initial bromide level, BrO₃/Br, mmol L⁻¹/mmol L⁻¹ (Li et al., 2017).
- Figure 5 shows that after a 30-minute process, the residual concentration of AN was in the order of 60%
- of the initial concentration and the molar bromate yield was 0.07%, with a bromate concentration of
- approximately 290 ug/L. After 60 minutes, the residual concentration of AN was of 16% and the molar
- bromate yield rose to 0.15%, with a bromate concentration of 644 ug/L. These results demonstrate that
- residual AN concentrations limited the generation of bromate to values much lower than 1% of the
- potential generation, even in the presence of very high initial bromide concentrations (i.e. in the order of
- 100 mg/l or more). In fact, AN inhibits bromate formation by reacting with HOBr/OBr⁻ and thus starting
- the series of reactions that leads to the formation of amines with different degree of bromination (mono-
- 348 , di-, tri-bromoamine, see Reactions 1a 1d). Bromoamines are cyclically formed, oxidized and
- destroyed to regenerate HOBr and Br- and release N₂. The participation of HOBr/OBr⁻ in the reactions
- of formation of bromoamines prevents it to be further oxidized to BrO₃- (Hofmann and Andrews 2001;
- Heeb et al. 2014). The results concerning the generation of bromate observed in this study are in good
- agreement with the findings of Berne and coauthors (2004) who observed delays in substantial bromate
- 353 generation that increased with the ratio between AN and bromide. Even in that case a residual presence
- of AN in the solution inhibited the generation of bromate.

360

- 355 However, notwithstanding the well-known role of AN in the suppression of bromate generation, small
- amounts of bromate have however been observed. This evidence suggests that radicals like CO₃• or
- 357 HCO₃• (coming from the bicarbonate introduced as a buffering agent) could be involved in the process
- of generation of bromate and their effect should be more in-depth investigated (Yang et al., 2019).

3.5 Interference of the presence of organic substances in AN depletion and nitrate generation

- As demonstrated in the previous sections, the bromide-enhanced ozonation process, if well regulated for
- 362 what concerns the initial pH value and the presence of buffer substances, could efficiently deplete AN

by keeping low the generation of nitrate. Moreover, the cyclical process of generation and destruction of 363 bromoamines, that involves HOBr/OBr- species, prevents the further oxidation of such species to 364 bromate. However, the extension of the applicability of the enhanced ozonation process to systems that 365 contain substances different from the sole AN, for example organic compounds, requires a more in-depth 366 comprehension of the interferences that can occur. 367 To obtain some preliminary insights into the possible interference due to the presence of organic matter 368 in the process of oxidation of AN with ozone and bromide, tests were performed in the condition of 369 maximization of the amount of dissolved bromide ($[Br-]/[O_3] > 12$). In this way, all the ozone species 370 (ozone and hydroxyl radicals) should have reacted with bromide to form "active bromine" and none (or 371 traces) of free oxidant (O₃ or HO*) was available. 372 373 Figure 6 shows some preliminary results that concern the effect of the presence of organic substances in the removal of AN in a bromide-enhanced ozonation process. It can be observed that, after 30 minutes, 374 375 the oxidant had consumed approximately 60% of the starting amount of AN in the system that did not contain any organic substance. From Figure 6 it can be seen that glucose and KPH did not affect the 376 377 removal rate of AN, in fact the concentration of AN after a 30-min oxidation, in the systems that contained the one or the other of these two substances, was of the same order of the AN concentration 378 found in the absence of organic substance. Conversely, the presence of hydroquinone and phenol slowed 379 down the AN removal. In fact, the residual amount of AN was of 60.1% (+50.2%, with respect to the 380 system containing the sole AN) and 63.6% (+58.8%) of the initial concentration for the systems that 381 contained hydroquinone and phenol, respectively. 382 The competition between the phenolic compounds and AN in the oxidation process takes probably place 383 384 in the very first stages of the process. It is well known that during oxidative processes (e.g., application of ozone), the phenolic functional groups found in NOM are the main sink for the added oxidants, being 385 386 susceptible to chemical modifications during exposure to oxidants (Onnby et al., 2018). The apparent second order rate constants for the reaction between phenol and ozone varies from 10³ to 10⁸ M⁻¹s⁻¹, in 387 the presence of pH values from pH 2 to 9 (Deborde and von Gunten, 2008). In the field of neutrality, 388 where the tests of the present study were carried out, the constant is in the order of 10⁶ M⁻¹ s⁻¹ and differs 389 for some orders of magnitude from that of the reaction between ozone and bromide $(1.60 - 2.58 \cdot 10^2 \, \text{M}^{-1})$ 390 s⁻¹, Liu et al., 2018). Phenol and compounds containing phenolic groups, like hydroquinone, are also 391 392 involved in reactions with HOBr and inorganic bromoamines. Recent studies demonstrated that HOBr and bromoamines are highly reactive towards compounds containing phenolic groups. Apparent second-393 order rate constant at pH = 7 in the order of $6.5 \cdot 10^4 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$, $2.1 \cdot 10^2 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$ and $2.3 \cdot 10^1 \,\mathrm{M}^{-1} \,\mathrm{s}^{-1}$ and were 394

measured for HOBr, monobromoamine and dibromoamine respectively (Criquet et al., 2015; Heeb et al., 2017).

As shown in Table 2 the compresence of AN and an organic substance decreased the nitrate production by approximately 30%. The presence of KHP or glucose seemed not to affect the generation of bromate that was maintained on yield values in the order of 0.04-0.07%. Conversely, the addition of substances such as phenol and hydroquinone determined a bromate generation of an order of magnitude higher than that observed with KHP and glucose. Results obtained in previous works (Chon et al., 2015; Li et al., 2017; Wu et al., 2018) proved that, at bromide concentrations in the order of few hundreds of micrograms per liter, reactions between the most reactive parts of NOM and ozone/HO* limited bromate formation. Conversely, the results found in this study demonstrate that, at high bromide concentration values, ready-degradable substances like phenolic compounds were not only unable to suppress the generation of bromate but seemed to be a cause for an enhanced generation. However, the series of mechanisms responsible for this enhanced bromate generation deserves a more detailed investigation and has still to be elucidated.

Table 2. Nitrate and bromate generation in systems containing ammonia (5 mM) and an organic substance (0.05 mM)

	Ammonia only	КНР	glucose	phenol	hydroquinone
NO ₃ -/theor NO ₃ - (%)	8.65±0.33	6.00±0.37	6.17±0.14	6.07±0.53	5.73±0.65
Variation (%)	-	- 30.6	- 28.7	- 29.8	- 33.8
BrO ₃ - (mg/l)	0.298	0.179	0.238	3.28	1.66
BrO ₃ - molar yield (%)	0.070	0.042	0.056	0.77	0.39

Conclusions

This study demonstrated that a bromide-enhanced ozonation process could effectively remove AN and minimize the generation of nitrate when all the ozone species reacted with bromide to form active bromine (HOBr). With the set-up and oxygen flow (200 ml/min) employed in this study, the condition of limiting ozone was realized for $[Br^-]/[O_3]$ ratios in the order of 12 or more. The generation of nitrate could decrease to 10% or less of the theoretical generation value.

The pH value played a key role in the process for two main reasons. On the one hand, it affected the equilibrium between ammonium ion and ammonia (NH₃ + H⁺ \leftrightarrow NH₄⁺). It was observed that, in the pH range where ammonia is the predominant form, the reaction of direct oxidation of ammonia, with

422 generation of nitrate, competed with the reaction that generated bromoamines and ends with the liberation 423 of nitrogen gas. On the other hand, the oxidation of AN determined the release of H⁺ ions and a consequent acidification of the solution. pH values below 5.0 resulted in a significant slowdown of the 424 reaction rate of AN oxidation and in an uncontrolled generation of nitrate. In order to avoid these 425 undesired effects, the addition of a buffering substance such as NaHCO₃ is compulsory. The 426 recommended ratio between HCO₃⁻/NH₄⁺ (by weight, g/g) is equal to 5. 427 428 The presence of AN inhibited the oxidation of bromide to bromate thanks to the generation of inorganic bromoamines. The generation of bromate was limited to values of less than 0.2% of molar bromate yield. 429 However, even when a residual concentration of AN was present, secondary reaction pathways that lead 430 to the generation of bromate could not completely be excluded. 431 432 Organic substances in the form of KHP or glucose did not affect the removal rate of AN. Conversely, the addition of substances such as phenol and hydroquinone, that have a higher reactivity towards ozone and 433

HOBr, determined a reduced AN removal rate and a bromate generation of one order of magnitude higher

than that observed with KHP and glucose. All the organic substances employed in this study seemed to

deplete the nitrate generation (-30% of the value obtained with sole AN). However, the role of organic substances in nitrate and bromate generation was not completely elucidated yet.

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Acknowledgements

The authors wish to thank Dr. M. Fungi from SMAT company for performing bromate analysis. This work was partially supported by SODAI Italia S.p.A. Some results are part of the research project entitled "Bromide enhanced ozonation of wastewater with high loads of ammonia nitrogen" funded by Compagnia di San Paolo through 2017 PoC Project.

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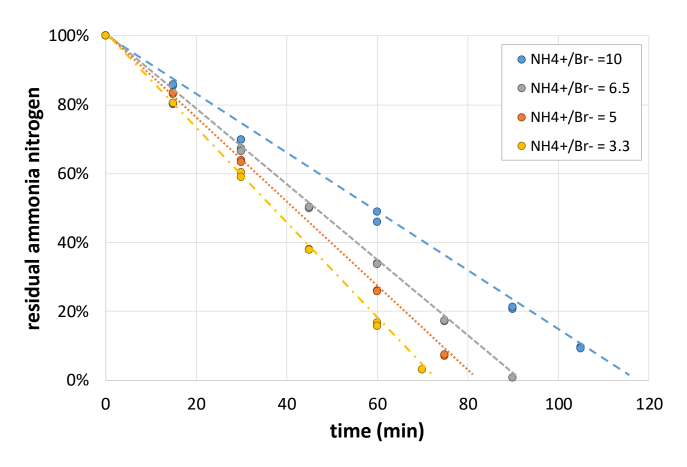


Figure 1. Trend of AN depletion in time, $[Br^-/O_3] = 5.54$; 8.51; 11.1 and 16.6 mM/mM (test I)

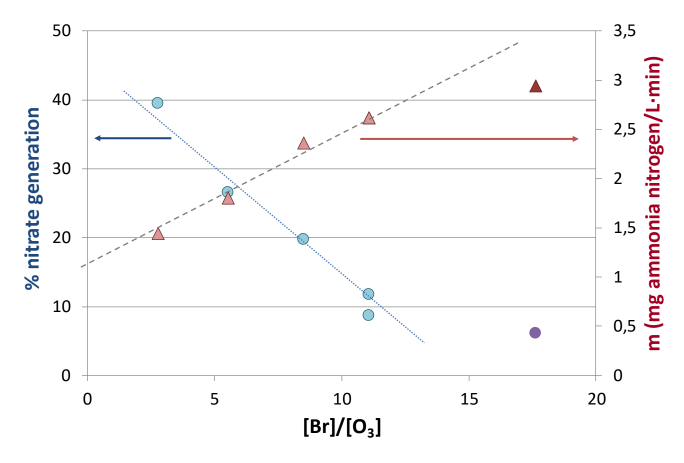
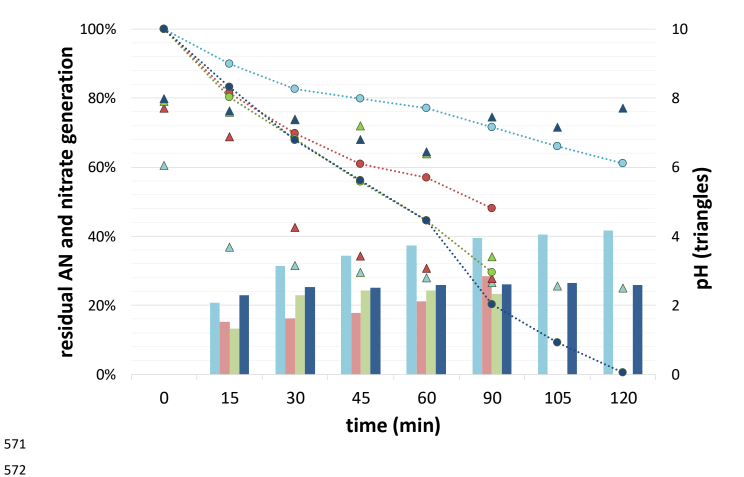


Figure 2. Effect of the $[Br-/O_3]$ ratio on AN removal rate (m, see Equation 2) and nitrate generation (test I)



0 mg HCO₃⁻/I
 200 mg HCO₃⁻/I
 500 mg HCO₃⁻/I
 500 mg HCO₃⁻/I

Figure 3. Effect of NaHCO₃ doses (0; 200; 500; 500+500 mg/L) on the progress of AN oxidation (dots and dotted line), nitrate generation (bars) and pH (triangles) after ozonation. Test conditions: $[NH_4^+] = 11.1 \text{ mM}$; $[Br^-] = 1.11 \text{ mM}$; $[NH_4^+]/[Br^-] = 10.0$, test II.

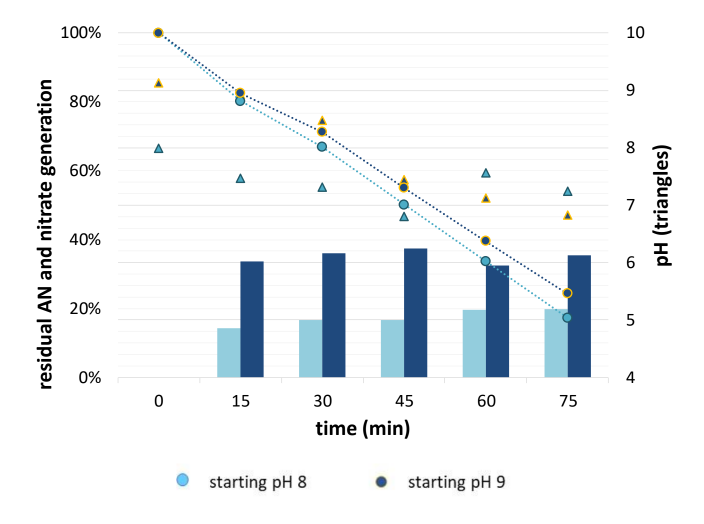


Figure 4. Effect of the starting pH (8; 9) on the progress of AN oxidation (dots and dotted line), nitrate generation (bars) and pH (triangles) after ozonation. Test conditions: $[NH_4^+] = 11.1 \text{ mM}$; $[HCO_3^-] = 16.4 \text{ mM}$; $[Br^-] = 1.70 \text{ mM}$; $[NH_4^+]/[Br^-] = 6.51$, test III.

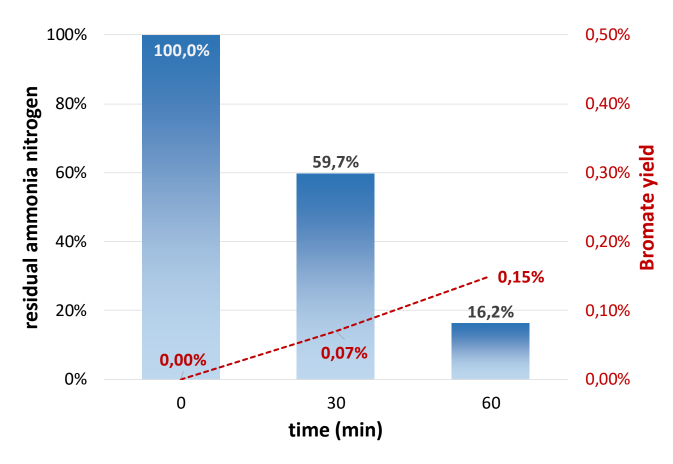


Figure 5. Trend of residual ammonia nitrogen and bromate yield ($[NH_4^+] = 11.1 \text{ mM}$; $[HCO_3^-] = 16.4 \text{ mM}$; $[Br^-] = 3.33 \text{ mM}$)



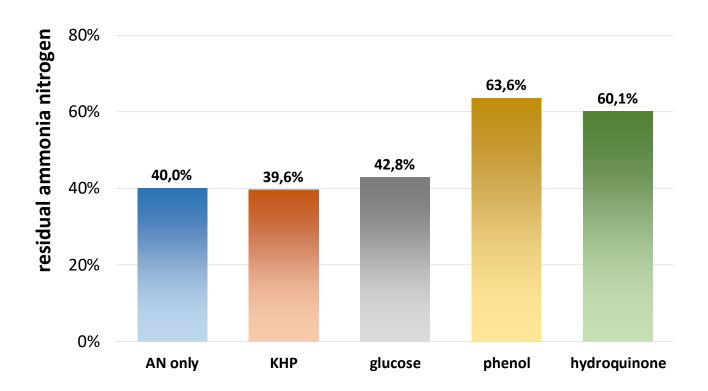


Figure 6. Residual ammonia nitrogen after a 30-minute ozonation in the presence of organic substances (test IV)