Generation of hydroxyl radical during chlorination of hydroxyphenols and natural organic matter extracts

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A R T I C L E   I N F O

Article history:
Received 23 January 2020
Received in revised form 2 March 2020
Accepted 3 March 2020
Available online 6 March 2020

Keywords:
Chlorine
Hydroxyphenols
Electron transfer
Quinones
Semiquinones
Superoxide radical
Hydroxyl radical

A B S T R A C T

The generation of hydroxyl radicals (·OH) during the chlorination of air saturated solutions of different hydroxyphenols (hydroquinone, resorcinol, catechol, gallic and tannic acids) at pH 7 has been determined by the formation of phenol (in presence of benzene in excess) or 2-hydroxyterephthalic acid (in presence of terephthalic acid). Formation of ·OH was only detected during the chlorination of o- or p-hydroxyphenols, compounds that react with chlorine by electron transfer forming the corresponding semiquinones/quinones. In aerated solutions, oxygen is reduced by the semiquinone to the superoxide radical, O2−, which reacts with HOCl to ·OH. Compared to the studied o-hydroxyphenols, the lower reactivity of hydroquinone towards chlorine favours the reaction between chlorine and O2−, and its ·OH formation potential is ~50 times higher. The extent of ·OH generated increased with the concentration of the hydroxyphenol and chlorine, but the ·OH yield (moles formed per mole of hydroxyphenol eliminated), decreased due to the formation of the quinone, that acts as O2− scavenger. The yield was almost not affected by the pH (6 ≤ pH ≤ 7.5), whereas a strong impact of dissolved O2 was observed. The ·OH production was null in absence of O2 and 2.5–3 times higher at oxygen saturated conditions compared to air-saturated. Contrary to chlorination, during bromination of hydroquinone ·OH was not formed, which can be attributable to a much faster consumption of the oxidant, with no chance for O2− to react with bromine.

Formation of ·OH during the chlorination of different NOM extracts (SRHA, SRFA, PLFA and Nordic Lake NOM) and water from Lake Greifensee (Switzerland) was also studied using terephthalic acid as ·OH scavenger. For SRHA, SRFA and Nordic Lake NOM (all of allochthonous origin and presenting high electron-donating capacity, EDC), ·OH yields expressed as moles formed per mole of DOCn (%) were between 1.1 and 2.0, similar to that of hydroquinone (~1.5). For PLFA and Lake Greifensee water (autochthonous, lower EDC) much lower ·OH yields were observed (0.1–0.3). Both chlorination rate and EDC, the later favouring the formation/stabilization of O2−, seem to be key factors involved in ·OH generation during the chlorination of NOM. A mechanism for these findings is proposed based on kinetic simulations of hydroquinone chlorination at pH 7.

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1. Introduction

Dissolved organic matter (DOM) is typically the main sink for oxidants commonly applied in drinking water treatment (Wenk et al., 2013; von Gunten, 2018). The oxidants react with electron-rich functional groups such as phenolic moieties, amines, olefines, etc., partially leading to the formation of low molecular weight organic disinfection byproducts (DBPs) (Lee et al., 2007; Bond et al., 2012; von Sonntag and von Gunten, 2012; Le Roux et al., 2016). Among the different structures present in DOM, phenolic moieties are highly reactive towards many different oxidants. The content of aromatic moieties in DOM has been usually measured in terms of the specific ultraviolet absorbance at 254 nm, SUVA254 (Weishaar et al., 2012; von Sonntag and von Gunten, 2012).
et al., 2003), however, recently more specific measurements such as the electron-donating-capacity (EDC) are being used as a proxy of the phenolic content (Aeschbacher et al., 2012; Walpen et al., 2016; Önnby et al., 2018). Although, in general, phenolic structures react with chlorine by electrophilic aromatic substitution (EAS) forming chlorophenols, recent studies have demonstrated that chlorination of ortho and para hydroxyphenols occurs via an electron transfer (ET) with the formation of the corresponding quinones. These reactions are likely responsible for the high initial hypohalous acid consumption by DOM through ET reactions (Criquet et al., 2015).

The chemistry of o- and p-hydroxyphenols in water is complex. According to Eyer (1991), hydroquinone (HQ) can be oxidized by dissolved oxygen in aqueous solution through the following mechanism, based on the hydroquinone/semiquinone/benzoquinone (HQ/SQ+/BQ) equilibria:

\[
\begin{align*}
HQ + O_2 & \quad \overset{k_1}{\rightleftharpoons} O_2^- + SQ^+ + 2H^+ \quad (1) \\
SQ^+ + O_2 & \quad \overset{k_2}{\rightleftharpoons} BQ + O_2^- \quad (2) \\
HQ + BQ & \quad \overset{k_3}{\rightleftharpoons} 2SQ^+ + 2H^+ \quad (3) \\
HO_2^* & \quad \overset{k_4}{\rightarrow} O_2^- + H^+ \quad (4) \\
2H^+ + 2O_2^- & \quad \overset{k_5}{\rightarrow} H_2O_2 + O_2 \quad (5) \\
HO_2^* + O_2^- & \quad \overset{k_6}{\rightarrow} HO_2 + O_2 \quad (6)
\end{align*}
\]

At pH \(\leq 7\) equilibrium 1 is almost completely displaced to the left (\(K_1 = 10^{-14}\) at pH 7, room temperature and air saturated, Eyer, 1991), and the oxidation of HQ to SQ+ by O2, with generation of superoxide radical (O2- \(^\bullet\)) is minimal. If SQ+ were formed, it would be quickly oxidized by O2 to BQ (reaction 2, \(k_2 = 5 \times 10^4 \text{M}^{-1}\text{s}^{-1}\) at pH 7 and room temperature, Eyer, 1991), under formation of more O2- \(^\bullet\). BQ could then react with HQ (\(k_3 = 58 \text{M}^{-1}\text{s}^{-1}\), \(k_3 = 8 \times 10^7 \text{M}^{-1}\text{s}^{-1}\) at pH 7, Yamazaki and Ohnishi, 1966), and also with O2- \(^\bullet\) (\(k_2 = 10^8 \text{M}^{-1}\text{s}^{-1}\) at pH 7, Eyer, 1991), both reactions leading to SQ+. With a pKa of 4.8 for equilibrium 4 (Bielski et al., 1985), at pH 7 the formation of hydroperoxyl radical (HO2-) from superoxide protonation is expected to be very low, whereas O2- \(^\bullet\) disproportionation to H2O2 (reaction 5) is slow leading to a certain stability of O2- \(^\bullet\) in solution under these conditions (Sheng et al., 2014). Although the reaction between HO2- and O2- \(^\bullet\) to HO2- has a high second-order rate constant (\(k_6 = 9.7 \times 10^7 \text{M}^{-1}\text{s}^{-1}\)), at pH 7 this reaction is not favored due to the low concentration of HO2- (0.6% of [O2- \(^\bullet\)])

In the above mechanism the possible reaction between HQ and O2- \(^\bullet\) was not considered. For this reaction, a second-order rate constant of \(1.7 \times 10^4 \text{M}^{-1}\text{s}^{-1}\) was reported by Rao and Hayon (1975), in disagreement with the findings of Nadezhdin and Dunford (1979), among others, who determined a second-order rate constant of \(1.7 \times 10^4 \text{M}^{-1}\text{s}^{-1}\) for the HQ/HO2- reaction. Since HO2- is a stronger oxidant than O2-, the rate constant of HQ-O2- reaction is expected to be \(< 1.7 \times 10^4 \text{M}^{-1}\text{s}^{-1}\).

According to the mechanism provided in reactions 1–6, superoxide radical will not be formed in absence of dissolved O2 and its steady state concentration will be very low in an aerated medium at pH \(\leq 7\).

The presence of an agent that can oxidize HQ to BQ could promote the formation of SQ+ through reaction 3 and hence the generation of O2- \(^\bullet\) (reaction 2). As stated before, Criquet et al. (2015) determined that chlorine and bromine react with o- and p-hydroxyphenols, HQ among them, via ET, with formation of the corresponding quinones. In the case of HQ, the apparent second-order rate constant of its reaction with free available chlorine (FAC, sum of [HOCI] and [ClO-]) at pH 7 (reaction 7; \(k_7 = 2.16 \times 10^5 \text{M}^{-1}\text{s}^{-1}\) measured in terms of BQ formation) is comparable to that of reaction 3. Hence, depending on the experimental conditions as BQ is formed, some SQ+ can be generated (reaction 3), thereby promoting the formation of O2- \(^\bullet\). Also, the oxidation of
SQ\textsuperscript{*} by HOCI, with formation of HOCI\textsuperscript{+} (reaction 8), cannot be ruled out.

\[ \text{HOCI} + \text{HQ} \xrightarrow{k_f} \text{BQ} + \text{Cl}^- \] (7)

\[ \text{HOCI} + \text{SQ}^* \xrightarrow{k_f} \text{BQ} + \text{HOCI}^* \] (8)

According to Candeias et al. (1993), superoxide reacts with HOCI to form *OH (reaction 9; \( k_9 = 7.5 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \), determined by monitoring the decay of O\text{\textsubscript{2}}\textsuperscript{*}; Long and Bielski, 1980).

\[ \text{HOCI} + \text{O}_2^* \xrightarrow{k_f} \text{**OH + Cl}^- + \text{O}_2 \] (9)

Therefore, if during HQ chlorination SQ\textsuperscript{*} (and hence, O\text{\textsubscript{2}}\textsuperscript{*}) is generated, hydroxyl radicals could be formed through reaction 9. Also, if reaction 8 develops, HOCI\textsuperscript{+} would lead to *OH formation through reaction 10 (\( k_{10} = 6.1 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \) and \( k_{10} = 4.3 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \), Jayson et al., 1973).

\[ \text{HOCI}^+ \xrightarrow{k_{10}} \text{**OH + Cl}^- \] (10)

A previous study revealed that an initial bromination of natural organic matter (NOM) mainly occurs through ET instead of EAS (Criquet et al., 2015). Hence, with the presence of phenols being structural moieties of NOM and in analogy to bromination, formation of SQ\textsuperscript{*} during chlorination of NOM, with a concomitant formation of HOCI\textsuperscript{+} and/or O\text{\textsubscript{2}}\textsuperscript{*} and subsequently *OH, cannot be ruled out.

Because of the short lifetime of hydroxyl radicals, it is common to use scavengers to determine their formation (Flyunt et al., 2003; Rodriguez et al., 2015). Moreover, if the product from the reaction between *OH and a scavenger is known and can be measured, the extent of the formation of *OH can be estimated. In addition, it is also important to exclude reactions of the scavenger and/or the reaction products with other substances present in the reaction medium (reagents and/or intermediates and/or products). Therefore, the following criteria need to be considered for the selection of a scavenger to quantify *OH formation: (i) the reactivity of the scavenger, the organic surrogate (hydroxyphenols or NOM in this case) and the oxidant (chlorine in this study) towards the radical species; (ii) the nature of the products formed from the reaction between the scavenger and the radical (stability, reactivity towards oxidants, etc.); (iii) easy detection and quantification of the products; and (iv) the reactivity of the scavenger, the transformation products and the organic surrogate towards the oxidant. Furthermore, the scavenger has to be present in a sufficiently high concentration, so that a fraction of >95% of the *OH is scavenged. Even though the product from the reaction between the scavenger and *OH is an indicator of the generation of this species, there are some uncertainties on its quantification due to possible effects of the reaction conditions on the product yield (moles of product formed per mole of *OH generated).

In this study, to investigate the formation of *OH, dimethyl sulfoxide (DMSO), tert-butyl alcohol (TBA), benzene (BZ) and ter-ephthalic acid (TPA) were initially selected as scavengers. However, DMSO, that reacts with *OH to form methane sulfinic acid (MSI) (\( k_{*OH:DMSO} = 4.5 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \); Bardouki et al., 2002), also has a relatively high reactivity with FAC (\( k_{FAC:DMSO} = 315 \text{ M}^{-1} \text{s}^{-1} \) at pH 7, 20 °C; Amels et al., 1997), wherefore its use is impossible for this system. The second order rate constants for the reactions of the different *OH scavengers towards *OH and FAC (pH 7 and 25 °C) and their reaction products are provided in Table 1.

The main products from the reaction between *OH and TBA (\( k_{*OH:TBA} = 6 \times 10^8 \text{ M}^{-1} \text{s}^{-1} \); Buxton et al., 1988) in presence of oxygen are formaldehyde (FAL; ~25% of *OH; Flyunt et al., 2003), acetone, 2-hydroxy-2-methylpropanal and 2-methyl-2-hydroxypropanal, as well as considerable amounts of HO\text{\textsubscript{2}}/O\text{\textsubscript{2}} (Schuchmann and von Sonntag, 1979; Cederbaum et al., 1983; Flyunt et al., 2003). Neither TBA nor FAC reacts with HOCl. The reactivity/stability of other reaction products is unknown.

In aqueous solution and in presence of dissolved oxygen the reaction between *OH (generated radiolytically) and BZ (\( k_{*OH:BZ} = 7.8 \times 10^9 \text{ M}^{-1} \text{s}^{-1} \); Buxton et al., 1988) proceeds through the formation of a hydroxycyclohexadienyl radical that further reacts with O\text{\textsubscript{2}}, to mainly phenol (Pan et al., 1993). The molar phenol yields were found in the range of 53–93% depending on the pH, and O\text{\textsubscript{2}}/HO\text{\textsubscript{2}}, small amounts of various aldehydes and formic acid were also formed. By photolysis of aerated NO\text{\textsubscript{3}} to produce *OH at pH ~6, the molar phenol yield was 95% (Deister et al., 1990). An average value of 85% has been adopted by different authors (Dong and Rosario-Ortiz, 2012). Contrary to BZ, which does not react with FAC, phenol (PHEN) reacts with an apparent second order rate constant \( k_{FAC-PHEN} = 18 \text{ M}^{-1} \text{s}^{-1} \), at pH 7 and 25 °C (Gallard and von Gunten, 2002).

The reaction of *OH with TPA in aqueous solution (\( k_{*OH:TPA} = (3.3-4.4) \times 10^9 \); Fang et al., 1996; Page et al., 2010; Charbouillot et al., 2011) results in the formation of the corresponding hydroxyterephtalic-type radical intermediate with a yield of about 85%. In presence of dissolved oxygen as the only oxidant, this radical intermediate reacts with O\text{\textsubscript{2}} generating 2-hydroxyterephthalic acid (hTPA) and *OH (Fang et al., 1996). The molar yield of hTPA based on moles of *OH produced is ~30% at pH 7 and 25 °C and slightly increasing with increasing pH and temperature (Charbouillot et al., 2011). The reactivity of the hydroxyterephtalic-type radical formed from the TPA-*OH reaction with O\text{\textsubscript{2}} is ~30 times lower than that of the radical formed from BZ-*OH reaction (Fang et al., 1995). Because of that, in presence of a stronger oxidant such as ICl\textsuperscript{2}, the hTPA yield can increase to up to 85% without formation of O\text{\textsubscript{2}}/HO\text{\textsubscript{2}} (Fang et al., 1996). Since there are no data in literature about the reactivity of TPA nor hTPA towards FAC, the corresponding apparent second-order rate constants (pH 7 and 25 °C) have been experimentally determined in this study.

The above mechanism (reactions 1–10) suggests that *OH might be formed during the chlorination of hydroxy-phenolic compounds. However, so far, there is no experimental evidence for this process. The main objective of this study was to determine whether chlorination of NOM leads to *OH formation and identify the main NOM structures involved. To this end, different o-, m- and p-hydroxyphenols (hydroquinone, catechol, resorcinol, gallic and tannic acid) and NOM extracts (humic and fulvic acids) were chlorinated, and the formation of *OH and the role of O\text{\textsubscript{2}}\textsuperscript{*} were investigated with the aid of multiple radical scavengers. The influences of hydroxyphenol, NOM, FAC and dissolved oxygen concentrations as well as pH on *OH production were also studied. From the results obtained a kinetic model for *OH formation during hydroquinone chlorination at pH 7 was developed to support the proposed mechanism.

2. Materials and methods

2.1. Chemicals

With the exception of the sodium hypochlorite solution (10–15% active chlorine, Sigma-Aldrich, reagent grade), all organic and inorganic compounds (purchased from Sigma-Aldrich, Fluka Analytical, Merck or Carlo Erba), were of analytical grade or higher and used without further purification. Stock solutions of the
selected hydroxyphenols (15 mM for hydroquinone (HQ), catechol (CAT), resorcinol (RES) and gallic acid (GAL); 2.5 mM for tannic acid (TAN)), were freshly prepared in Milli-Q ultrapure water. Their structures are presented in Table 2 together with the corresponding second order rate constants for the reactions with FAC, $^\bullet$OH and $O_2^\bullet$. The NOM extracts were purchased from the International Humic Substances Society (IHSS) and used as received: Suwannee River humic acid II (SRHA, catalogue number: 2S101H), Suwannee River fulvic acid II (SRFA, catalogue number: 2S101F) and Nordic Lake NOM (NL-NOM, catalogue number: 1R108N), representing organic materials from allochthonous origin; and Pony Lake fulvic acid (PLFA, catalogue number: 1R109F), as a representative of autochthonous NOM in aquatic systems. Some of their characteristics are summarised in Table S1 (supplementary information, SI). A given amount of each extract was dissolved in pH 7 phosphate buffer (PBS, 1 mM) to obtain stock solutions of 20 mg L$^{-1}$. Chlorination of a filtered (0.45 μm nylon membrane, VWR) water from Lake Greifensee (LG) (Canton of Zurich, Switzerland), was also carried out. The main characteristics of LG water were: DOC 3.3 mg L$^{-1}$, alkalinity 355 mg L$^{-1}$ CaCO$_3$, and pH 8.37. Stock NOM solutions and LG water were stored at 4°C until used.

Table 1
Second order rate constants for the reactions of different scavengers with hydroxyl radical ($^\bullet$OH) and HOCl and the corresponding reaction products.

<table>
<thead>
<tr>
<th>Scavenger</th>
<th>Second order rate constant for the reaction with $^\bullet$OH $k_{^\bullet$OH,M}$M^{-1}s^{-1}$</th>
<th>Selected product from the reaction with $^\bullet$OH</th>
<th>Second order rate constant for the reaction with FAC $k_{FAC,M}$ $M^{-1}s^{-1}$</th>
<th>Second order rate constant for the reaction of the product with FAC $k_{^*$FAC,M}$ $M^{-1}s^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimethyl sulfoxide (DMSO)</td>
<td>$4.5 \times 10^8$ (Bardouki et al., 2002)</td>
<td>Methanesulfonic acid (MSI)</td>
<td>$k_{FAC,DMSO} = 315$, pH 7, 20 °C (Amels et al., 1997)</td>
<td>?</td>
</tr>
<tr>
<td>tert-butyl alcohol (TBA)</td>
<td>$6 \times 10^8$ (Buxton et al., 1988)</td>
<td>Formaldehyde (FAL) $^a$</td>
<td>NR</td>
<td>NR</td>
</tr>
<tr>
<td>Benzene (BZ)</td>
<td>$7.8 \times 10^8$ (Buxton et al., 1988)</td>
<td>Phenol (PHEN) $^a$</td>
<td>NR</td>
<td>$k_{FAC-PHEN} = 18$, pH 7, 25 °C (Gallard and von Gunten, 2002)</td>
</tr>
<tr>
<td>Terephthalic acid (TPA)</td>
<td>$3.3 \times 10^9$ (Fang et al., 1996)</td>
<td>2-hydroxy-TPA (hTPA) $^a$</td>
<td>$k_{FAC-hTPA} &lt; 0.003$, pH 7, 25 °C (this study)</td>
<td>$k_{FAC-hTPA} = 0.147$, pH 7, 25 °C (this study)</td>
</tr>
</tbody>
</table>

FAC: Free available chlorine = [HOC] + [ClO$^-_\bullet$].  
$^a$ Is the main product. 
$^b$ Is not the main product; NR: No Reaction.

Table 2
Structures of the selected hydroxy-phenols and the apparent second order rate constants for their reactions with FAC, $^\bullet$OH and $O_2^\bullet$ at pH 7.

<table>
<thead>
<tr>
<th>Compound</th>
<th>Structure</th>
<th>$k_{FAC}$ (pH 7, 25 °C), $M^{-1}s^{-1}$</th>
<th>$k_{^\bullet$OH,M}$M^{-1}s^{-1}$</th>
<th>$k_{O_2^\bullet,M}$ $M^{-1}s^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydroquinone (HQ)</td>
<td>$C_6H_6O_2$; MW 110.11</td>
<td>21.6 (Criquet et al., 2015)</td>
<td>$1.2 \times 10^{10}$ (Gurol and Vatistas, 1987)</td>
<td>$&lt;1.7 \times 10^6$ (Nadezhdin and Dunford, 1979)</td>
</tr>
<tr>
<td>p-benzoquinone (BQ)</td>
<td>$C_6H_4O_2$; MW 108.09</td>
<td>$\sim 3$ (This study)</td>
<td>$6.6 \times 10^9$ (Schuchmann et al., 1998)</td>
<td>$10^9$ (Eyer, 1991)</td>
</tr>
<tr>
<td>Catechol (CAT)</td>
<td>$C_6H_6O_2$; MW 110.11</td>
<td>217.5 (Criquet et al., 2015)</td>
<td>$1.1 \times 10^{10}$ (Buxton et al., 1988)</td>
<td>$2.7 \times 10^8$ (Bielski et al., 1985)</td>
</tr>
<tr>
<td>Resorcinol (RES)</td>
<td>$C_6H_6O_2$; MW 110.11</td>
<td>$\sim 4000$ (Rebenne et al., 1996)</td>
<td>$1.2 \times 10^{10}$ (Buxton et al., 1988)</td>
<td>2 (Bielski et al., 1985)</td>
</tr>
<tr>
<td>Gallic acid (GAL)</td>
<td>$C_7H_6O_5$; MW 170.12</td>
<td>1600 (Criquet et al., 2015)</td>
<td>$10^{10}$ (Dwibedy et al., 1999)</td>
<td>$2.3 \times 10^4$ (Huisman et al., 2004)</td>
</tr>
<tr>
<td>Tannic acid (TAN)</td>
<td>$C_{76}H_{52}O_{46}$; MW 1701.2</td>
<td>$1.9 \times 10^8$ (Criquet et al., 2015)</td>
<td>$3 \times 10^{10}$ (Bors and Michel, 1999)</td>
<td></td>
</tr>
</tbody>
</table>

A FAC stock solution (~50 mM) was prepared every week by dilution of a commercial hypochlorite solution with ultrapure water and stored at 4 °C. HOBr was obtained from the reaction between 1 mM Br⁻ and 1.1 mM ozone at pH 4 (5 mM PBS) during 30 min, purging any residual ozone with nitrogen for 1–2 h (Heeb et al., 2017). The lower reaction time applied compared to that of Heeb et al. (2017) is probably the reason why the final HOBr concentration was lower than expected (~0.7 mM instead of ~1 mM).

Stock solutions of the different scavengers were prepared in ultrapure water: tert-butyl alcohol (TBA) 3.5 M; saturated aqueous benzene (BZ) – 23 mM, stirred overnight and kept in a fume hood; terephthalic acid (TPA) 50 mM, adjusted to pH 6.5 by adding some drops of concentrated NaOH; and MnCl₂·4H₂O 20 mM, the latter used as a superoxide scavenger. When tetrani-tromethane (TNM) was used as superoxide scavenger, 4 µL of TNM 8.2 M (pure compound) was added to 20 mL of the reaction medium (final [TNM] = 1.65 mM) in the fume hood.

2.2. Analytical methods

Stock solutions of chlorine and bromine were photometrically standardized at pH 11 before use (εCl₂ - at 292 nm = 350 M⁻¹ cm⁻¹; Hand and Margerum, 1983; εBr₂ - at 330 nm = 327 M⁻¹ cm⁻¹; Heeb et al., 2017). Residual concentrations of FAC were determined using a colorimetric method based on diethyl-p-phenylene diamine (DPD), measuring the absorbance at 515 nm (Rodier et al., 2009). The selected hydroxyphenols (HQ, CAT, RES, GAL and TAN), p-benzoquinone (BQ), formaldehyde (FAL), phenol (PHEN), and 2-hydroxyterephthalic acid (hTPA), were measured by a high-performance liquid chromatography system (HPLC Dionex Ultimate 3000) equipped with a Nucleosil 100-5 C18 column (125 × 3 mm, 5 µm, 100 A, Macherey-Nagel) with UV diode array and fluorescence detectors. The column temperature and flow were set as 30 °C and 0.8 mL min⁻¹, respectively, using 0.1% H₃PO₄ in ultrapure water (A) and methanol (B) to prepare the mobile phases. In all cases, 100 µL samples were injected except for HTPA (20 µL).

Formaldehyde (FAL) was firstly derivatized with 2,4-dinitrophenylhydrazine (DNPH), to form the corresponding hydrazone, and then analysed by HPLC-DAD at 360 nm (U.S. EPA, 1996). Acetone and BO, if present, were also quantified by this method.

A description of the preparation of reagents, conditions of the HPLC analyses, measuring ranges and limits of detection (LOD) and quantification (LOQ) are provided in Text S1 (SI).

The nitroform anion (NF⁻) formed from the tetrani-tromethane (TNM) reduction by O₂⁻ was determined spectrophotometrically at 350 nm (ε₃50nm = 14600 M⁻¹ cm⁻¹; Rabani et al., 1965).

The dissolved organic carbon (DOC) and inorganic carbon (IC) content in LG water were measured using a Shimadzu total organic carbon (TOC) analyzer. Initial DOC concentration (DOC₀) of the hydroxyphenols and NOM extract solutions was calculated from their molecular structures and chemical composition, respectively.

2.3. Experimental procedures

2.3.1. Chlorination of phenolic surrogates, NOM extracts and LG water in the presence of *OH scavengers

To determine the formation of *OH during chlorination, 0.4 mL phosphate buffer (PBS) 0.5 M (pH 6–7.5), and fixed volumes of hydroxyphenol solutions (or NOM solutions), radical scavenger stock solutions and ultrapure water were added, to obtain the desired concentrations in a final volume of 20 mL (22 mL amber glass vials). In the case of LG water, only the scavenger solution was added, decreasing the pH from 8.37 to 7.85.

After mixing, different volumes of a FAC stock solution were added to each vial while stirring and the vials were closed. A control sample was always prepared under the same conditions but without adding FAC. The mixtures were allowed to react at room temperature for 1–72 h (depending on the nature of the surrogate and the concentration of reactants) and then analysed by HPLC. In some cases bromine instead of chlorine was added.

To establish the initial concentrations of all reagents, the reactivity of the *OH scavenger, the organic surrogates and FAC towards *OH was taken into account to ensure that more than 95% of the generated *OH ([f(*OH)] >0.95), is captured by the *OH scavenger, according to the following equation:

\[
f(*OH) = \frac{k_{*OH-scavenger} [OH scavenger]}{k_{*OH-scavenger} [OH scavenger] + k_{*OH-hydroxyphenol} [hydroxyphenol] + k_{*OH-HOCl} [HOCl] + k_{*OH-ClO^-} [ClO^-]}
\]

where \( k_{*OH-scavenger} \) represents the second order rate constant for the reaction between *OH and the selected scavenger (Table 1); \( k_{*OH-hydroxyphenol} \) the second order rate constant for the reaction between *OH and the phenolic surrogate (Table 2); \( k_{*OH-HOCl} \) (1.21 × 10⁹ M⁻¹ s⁻¹; Bulman et al., 2019) and \( k_{*OH-ClO^-} \) (6.37 × 10⁸ M⁻¹ s⁻¹; Bulman et al., 2019), are the second order rate constants for the reactions of *OH with HOCl and ClO⁻, respectively. The [OH scavenger], [hydroxyphenol], [HOCl] and [ClO⁻] are the initial molar concentrations of each reactant, the last two determined taking into account the pH of the medium, equilibrium 12, and the concentration of FAC ([FAC] = [HOCl] + [ClO⁻]). Since the second order rate constants for the reactions between *OH and H₂PO₄⁻/HPO₄²⁻ are extremely low (2 × 10⁴ and 1.5 × 10⁵ M⁻¹ s⁻¹, respectively; Buxton et al., 1988), at the tested conditions (PBS 10 mM) they can be neglected as *OH scavengers.

\[
HOCl + \text{pK}_a - 7.49 \rightarrow \text{ClO}^- + H^+
\]

Thus, for [hydroxyphenol]₀ ≤ 62.5 mM, molar [FAC]₀/[hydroxyphenol]₀ ratios ≤ 2.5 and pH ≤ 7.5, 50 mM TBA, 10 mM BZ and 4 mM TPA were used to fulfill a >95% scavenging criteria for \( f(*OH) \) according to equation (11) (see Table S2 in SI for calculated values).

To investigate the participation of O₂⁻, a similar procedure was applied by adding different amounts of a 15 mM BQ stock solution, pure TNM or 20 mM Mn(II) (as MnCl₂·4H₂O), in the latter case filtering the samples (0.45 µm nylon filter) before the HPLC analysis.

When needed, the dissolved O₂ concentration in the reaction medium was increased or reduced by purging with pure O₂ or N₂, respectively. When BZ was used, bubbling was performed before adding the scavenger to avoid losses by stripping.

\[
\text{HOCI} + \text{pK}_a - 7.49 \rightarrow \text{ClO}^- + H^+
\]
2.3.2. Determination of chlorine demand (72 h) of NOM extracts and LG water

In a series of 22 mL amber glass vials, 0.4 mL of 0.5 M PBS pH 7 and a fixed volume of NOM extract stock solutions were added, thereafter diluted to 20 mL with ultrapurified water. After mixing, to obtain different [FAC]₀/[DOC]₀ molar ratios a predetermined volume of a FAC stock solution was added to each vial while stirring and the vials were closed. The mixtures were allowed to react at room temperature in the dark, and after 72 h the residual FAC was analysed. For LG water, the procedure was the same but without adding PBS or ultrapurified water. Once the chlorine demand (μM) of each NOM solution was determined, the specific chlorine demand was calculated by dividing this value by the DOC₀ (μM) content.

2.3.3. Determination of the apparent second order rate constants for the reactions between chlorine and TPA, hTPA or BQ at pH 7 (10 mM PBS) and 25 °C

Apparent second order rate constants for the reactions of TPA, hTPA or BQ with chlorine at pH 7 (10 mM PBS, 25 °C), were determined in excess of FAC (pseudo-first order conditions; see experimental conditions in Fig. S1, SI), by measuring the evolution of the parent compounds (TPA and hTPA by HPLC-DAD/fluorescence; BQ by the decrease of the absorbance at 247 nm). In all cases but BQ residual chlorine was quenched before the analysis by adding a few μL of a concentrated solution of Na₂S₂O₃ (0.5 M). For TPA, experiments in excess of TPA (4 mM) were also performed measuring the evolution of residual FAC ([FAC]₀ = 45 μM). For BQ, a competition kinetics method was also applied using phenol as competitor \( k_{\text{FAC-TPA}} = 18 \text{ M}^{-1} \text{s}^{-1} \) at pH 7, 25 °C; \cite{Gallard and von Gunten, 2002}. In this case, vials containing BQ (50 μM) and PHEN (5–10 μM) (10 mM PBS, pH 7 at 25 °C), were dosed with different volumes of a FAC stock solution to obtain \([\text{FAC}]_0\) in the range of 0–50 μM. The samples were mixed, and after a reaction time of 3 h, the final concentrations of BQ and PHEN were determined by HPLC. In all cases, control samples were prepared in absence of chlorine. All the experiments were carried out at least in duplicate.

2.3.4. Modelling of *OH generation during chlorination of HQ

The kinetics of the formation of *OH during HQ chlorination at pH 7, 25 °C and air saturated conditions (concentration of dissolved oxygen assumed to be 3 × 10⁻⁴ M), was modeled by the Tenua kinetic simulator (http://bililite.com/tenua/).

3. Results and discussion

3.1. Determination of *OH formation from hydroquinone chlorination

3.1.1. Influence of the nature of the selected *OH scavengers on HQ chlorination

In a first series of experiments, chlorination of an air-saturated HQ solution 62.5 μM (DOC 4.5 mg L⁻¹) at pH 7 (PBS 10 mM) and room temperature (22 ± 2 °C) was carried out, using different molar [FAC]₀/[HQ]₀ ratios, in absence/presence of the selected *OH scavengers: TBA 50 mM, BZ 10 mM or TPA 4 mM (Fig. 1).

Under these experimental conditions, and as expected due to their low reactivity towards FAC (see Table 1 and Fig. S1, SI) the scavengers had practically no effect on the extent of HQ oxidation. Thus, for a given [FAC]₀/[HQ]₀ ratio, the extent of HQ abatement after 1.5 h contact time (Fig. 1a) was very similar regardless of the presence of a scavenger. Furthermore, the HQ abatement (solid symbols) corresponded well with the BQ formed (open symbols), i.e., the BQ yields were close to 1 mole per mole of HQ oxidized. For [FAC]₀/[HQ]₀ ratios > 1.5, a decrease in the final BQ concentration is observed, most probably due to its reaction with FAC \( (k_{\text{FAC-BQ}} = 3 \text{ M}^{-1} \text{s}^{-1} \text{ at pH 7, 25 °C, obtained in this study; Fig. S1, SI}) \). Also, as the dose of FAC increased, the BQ concentration was higher in presence of a scavenger, which could indicate that under these conditions some BQ abatement is prevented. The apparent global stoichiometry of the chlorine-HQ reaction was close to 2 mol of chlorine per mole of HQ consumed (Fig. 1b), which is consistent with previous observations \cite{Criquet et al., 2015}.

3.1.2. Influence of the nature of the scavenger on *OH quantification

In Fig. 2a, the concentrations of FAL (from TBA), PHEN (from BZ) or hTPA (from TPA) formed from chlorination (1.5 h contact time) of air-saturated HQ 62.5 μM solution (pH 7), for different [FAC]₀/[HQ]₀ molar ratios are shown. In absence of FAC (control), FAL, PHEN or hTPA were not detected. The FAC concentration was much lower than PHEN or hTPA, results that cannot be attributable to the low FAL yield of the TBA-*OH reaction (−25%; \cite{Flyunt et al., 2003}). Moreover, the FAC concentration initially increased with increasing molar [FAC]₀/[HQ]₀ ratios and decreased for ratios > 0.5. These results indicate that for the tested conditions, FAL and acetone (not shown) are not stable in solution due to unknown secondary reactions. Therefore, in the further experiments the formation of FAL from TBA was not considered to determine the formation of *OH from HQ chlorination.

For molar [FAC]₀/[HQ]₀ ratios ≤ 0.5, the PHEN and hTPA concentrations were similar despite the different yields of PHEN (from BZ + *OH) and hTPA (from TPA + *OH) reported in air saturated and neutral pH conditions (−85% and 30%, respectively). As it will be discussed in the kinetics section, this could indicate that in presence of chlorine, similarly to what was observed in presence of IrC³⁺\(^{(3)}\) \cite{Fang et al., 1996}, the yield of hTPA is higher than when using O₂ as the only oxidant. For [FAC]₀/[HQ]₀ > 1 the extent of PHEN formation was clearly lower than for hTPA and it was negligible for [FAC]₀/[HQ]₀ > 2. Undoubtedly, these results are related to the reactivity of PHEN towards HOCl, which is about 100 times higher than for hTPA (\( k_{\text{FAC-hTPA}} = 0.15 \text{ M}^{-1} \text{s}^{-1} \text{ at pH 7 and 25 °C; Table 2 and Fig. S1, SI} \)).

The yield of PHEN or hTPA as moles formed per mole of HQ eliminated (%), is shown in Fig. 2b. For the lowest molar [FAC]₀/[HQ]₀ ratio, yields of PHEN and hTPA were −20% and −15%, respectively. As the [FAC]₀/[HQ]₀ ratios increased, both yields decreased, in case of PHEN to < 5% for molar ratios ≥ 1.5, potentially due to the reaction of PHEN with chlorine. For hTPA, as will be discussed later, the decrease in the yield is probably related to a superoxide scavenging effect of the formed BQ, and remained almost constant at about 9% for molar [FAC]₀/[HQ]₀ ratios ≥ 0.5.

For the conditions tested, the compliance with equation (11) \( (f^{(\ast OH)} = 0.95) \) was also experimentally corroborated as shown in Fig. S2 (SI) as an example, where the influence of the TPA concentration on hTPA formation from HQ chlorination (\([\text{HQ}]_0 \text{ 62.5 μM; } \text{[FAC]}_0 \text{ 93.7 μM}, \text{ at pH 7 and room temperature, is presented. As observed, a constant hTPA concentration was determined when TPA > 3 mM was added.} \)

The fact that both PHEN and hTPA are detected implies that under the applied experimental conditions, chlorination of HQ leads to the formation of *OH. Moreover, to suppress the *OH -TPA reaction, the formation of hTPA was also tested in presence of high TBA concentrations. The results (Fig. S3, SI) indicate that TBA drastically inhibited the formation of hTPA. Therefore, it is very likely that *OH is the main species involved in hTPA formation and the same can be assumed for PHEN when BZ was used as a scavenger.
3.1.3. Influence of the initial concentration of HQ

The influence of the initial HQ concentration on *OH formation at pH 7 for TPA as *OH scavenger is shown in Fig. 3. As presented in Fig. 3a, for [HQ]₀ < 15 μM, the concentration of hTPA increased almost linearly as [FAC]₀/[HQ]₀ increased, whereas for higher [HQ]₀ a linearity can only be observed for low [FAC]₀/[HQ]₀ ratios, with hTPA concentrations approaching a plateau for higher values. The molar hTPA yields, based on moles of HQ abated (Fig. 3b), decrease with increasing [HQ]₀, approaching constant values at higher molar [FAC]₀/[HQ]₀ ratios. Similar results have been obtained for the PHEN yields when BZ was used as scavenger (Fig. S4, SI).

These observations could be at least partly explained by the nature of the intermediates and/or products formed during HQ chlorination. BQ, which reacts very fast with O₂⁻ (k₂ = 10⁹ M⁻¹ s⁻¹), could compete with HOCl (k₉ = 7.5 × 10⁸ M⁻¹ s⁻¹) for O₂⁻, thereby lowering the *OH yield. To test this, BQ (0–60 μM) was added to samples containing 62.5 μM HQ and 10 mM BZ. Under these conditions, BQ did not impact the hTPA yield (Table S2, SI). The mixtures were chlorinated and PHEN was measured. Fig. 4 shows the evolution of the PHEN yields as a function of [FAC]₀/[HQ]₀ for experiments performed with/without initial BQ (see also Fig. S5, SI). For low molar [FAC]₀/[HQ]₀ ratios (and therefore low BQ formation from HQ chlorination), the addition of BQ clearly reduced the PHEN yield. For increasing molar [FAC]₀/[HQ]₀ ratios, this effect was compensated by the BQ formed from HQ oxidation. Hence, it can be hypothesized that BQ can become a superoxide scavenger (reaction 2), lowering the *OH yield.

In absence of FAC (control samples) the concentration of HQ varied very slightly during the reaction time considered (between 1.5 and 4 h), indicating that a contribution of reaction 1 (HQ autooxidation) to O₂⁻ and SQ⁺ formation was likely negligible. This means that BQ is mainly formed from HQ chlorination (reaction 7), allowing the formation of SQ⁺ through reaction 3. Depending on the experimental conditions, the SQ⁺ formed could be further oxidized to BQ by O₂ (reaction 2), with concomitant generation of O₅⁻, and/or HOCl (reaction 8) to HOCl⁺, which could also
initial HQ concentrations on (a) hTPA formation and (b) hTPA yields. Symbols: , [HQ]₀ = 3.9 μM; : [HQ]₀ = 7.8 μM; : [HQ]₀ = 15.6 μM; : [HQ]₀ = 31.2 μM; : [HQ]₀ = 62.5 μM. Experimental conditions: TPA = 4 mM; air saturated PBS (10 mM), pH 7; Tₑxₜ = 22 ± 2 °C; reaction time = 1.5–4 h. The lines in Fig. 3a are shown to guide the eye.

contribute to *OH generation through reaction 10.

To test the potential role of superoxide radicals, experiments were performed with TNM or Mn(II) as O₂⁻⁺ scavengers. Unfortunately, due to the reactivity of TNM and Mn(II) towards HQ, that at the conditions tested was oxidized to BQ by these reagents in absence of chlorine, their use as O₂⁻⁺ scavenger seems to be impossible for this system (for further information see Text S2, S1).

3.1.4. Influence of pH

To test the influence of speciation of chlorine on *OH production from HQ chlorination, experiments were performed in the pH range 6–7.5 (PBS 10 mM, [HQ]₀ 62.5 μM, air saturated), using TPA as *OH scavenger. At pH > 7.5, HQ is not stable in solution due to its fast oxidation by O₂ (James et al., 1938; WHO, 1994; Criquet et al., 2015). At pH 7.5, the formation of a BQ ~1.5 times higher than the control sample (absence of chlorine), attributable to partial HQ autooxidation by O₂ through reactions 1–2. However, since the extent of HQ oxidized per mole of FAC also slightly increased with increasing pH, the differences in terms of hTPA yield are negligible (Fig. 5b). In presence of oxygen as the only oxidant, an increase of pH causes a slight increase in the yield of hTPA formed from TPA-·OH (from −28% at pH 6 to ~33% at pH 7.5, at 25 °C) (Charbouillot et al., 2011). Although k₆ also increases with increasing pH (58 M⁻¹ s⁻¹ at pH 7, Yamazaki and Ohnishi, 1966; 570 M⁻¹ s⁻¹ at pH 7.4, Eyer, 1991), which could favor the formation of O₂⁻⁺, an increase in k₇ is also expected counteracting this effect.

3.1.5. Influence of dissolved oxygen concentration

In contrast to HOC⁺⁺ (reaction 8), the formation of O₂⁻⁺ requires the presence of oxygen (reaction 2). Hence, if this species is mainly responsible for *OH generation during HQ chlorination, the formation of hTPA or PHEN must strongly depend on the O₂ concentration. In addition, it is important to note that, in absence of other oxidants, the mechanisms of hTPA and PHEN formation (from the TPA-·OH or BZ-·OH reactions) also require the presence of O₂ (Mathews, 1980; Fang et al., 1996; Pan et al., 1993). The reactions of the radical intermediates with O₂ leading to hTPA or PHEN could also produce molar equivalents of O₂⁻⁺, which could contribute to the *OH formation by reaction 9. In absence of O₂, products different than hTPA or PHEN could be formed. All these aspects will be discussed in more detail in the section on reaction kinetics.

To investigate the effect of O₂, air saturated ([O₂] = 300 μM), oxygen saturated ([O₂] = 1200 μM) and N₂ saturated (O₂ purged by stripping with N₂) solutions containing 62.5 μM HQ were chlorinated (pH 7 and room temperature), using TPA as *OH scavenger. The results in Fig. 6, show that hTPA is not formed in absence of O₂, both hTPA concentrations (Fig. 6a) and yields (Fig. 6b) increased with increasing O₂ levels. For oxygen saturated conditions (the O₂ concentration increased four fold compared to air saturation), hTPA concentrations were 2–3 times higher and the yield increased ~3–4 times, the latter due to the lower efficiency of HQ abatement for increasing O₂ concentrations (Fig. 6c). Overall, an increase in O₂ favors the formation of O₂⁻⁺ through reaction 2 and, at
the same time, the chlorine consumption by reaction 9. From these results, it seems quite evident that \( \text{O}_2/\text{C}_{15}/\text{C}_0 \) participates in the generation of \( \text{C}_{15} \text{OH} \). Moreover, the strong influence of \( \text{O}_2 \) levels on hTPA and PHEN generation would indicate that a contribution of HOCl/\( \text{C}_{15}/\text{C}_0 \) to \( \text{C}_{15} \text{OH} \) production through reaction 8 is less important.

Similar results were obtained for the influence of \( \text{O}_2 \) on PHEN formation and yields when BZ was used as \( \text{C}_{15} \text{OH} \) scavenger. However, in this case, to avoid BZ volatilization, the level of \( \text{O}_2 \) in the samples could only be partially reduced or increased (Fig. S6, SI).

The possible contribution to \( \text{C}_{15} \text{OH} \) formation from superoxide that can be generated from BZ-\( \text{C}_{15} \text{OH} \) or TPA-\( \text{C}_{15} \text{OH} \) reactions will also be discussed in the kinetics section.

### 3.1.6. Bromination of hydroquinone

Similar to chlorine, bromine oxidizes HQ to BQ, with a significantly higher second order rate constant for this reaction at pH 7 compared to FAC (\( k_{\text{HOBr-HQ}} = 6.4 \times 10^4 \text{ M}^{-1} \text{ s}^{-1} \) vs \( k_{\text{FAC-HQ}} = 2.16 \times 10^3 \text{ M}^{-1} \text{ s}^{-1} \); Criquet et al., 2015). The second order rate constant for the reaction between HOBr and \( \text{O}_2^\bullet \) (reaction 13) is also about three orders of magnitude higher than for HOCI (reaction 9) (\( k_{13} = 3.5 \times 10^3 \text{ M}^{-1} \text{ s}^{-1} \); Schwarz and Bielski, 1986), and leads to HOBr\(^*\) that further evolves to \( \text{C}_{15} \text{OH} \) through reaction 14

\[
\begin{align*}
\text{HOBr} + \text{O}_2^\bullet & \rightarrow \text{HOBr}^* + \text{O}_2 \\
\text{HOBr}^* & \rightarrow k_{14} k_{14} \quad \text{OH} + \text{Br}^- \\
\text{HOBr}^* & \rightarrow k_{15} k_{15} \quad \text{Br} + \text{OH}^- 
\end{align*}
\]

The experiments were performed at pH 7 (PBS, 10 mM, air saturated) with 62.5 \( \mu \text{M} \) [HQ], using 4 mM TPA as a scavenger and molar [HOBr]/[HQ] ratios between 0 and 1.5. The apparent stoichiometry of HOBr:HQ reaction was \( \sim 1:1 \) (Fig. S7, SI), in agreement with Criquet et al. (2015). In these experiments hTPA was not detected. A plausible explanation for this observation could be the
much higher value of $k_{\text{HOBr-HQ}}$ compared to that of $k_{\text{FAC-HQ}}$, wherefore, HOBr would be quickly and totally consumed by HQ with no chance to react with $O_2^-$ so neither HOBr$^-$ nor $O_2^-$ would be generated in this reaction system.

### 3.2. *OH formation from chlorination of different hydroxyphenols at pH 7

The behavior of other hydroxyphenols (CAT, RES, GAL and TAN) was investigated to explore the applicability of our findings to a wider range of compounds. Their structures and reactivities towards different species (FAC, $*\text{OH}$ and $O_2^-$) are summarised in Table 2. At pH 7, all of these compounds except RES lead to the formation of quinones upon chlorination and have higher FAC reactivities than HQ (Criquet et al., 2015).

The experimental procedure was the same as for HQ, i.e., chlorination in 10 mM PBS air saturated at pH 7, using 4 mM TPA as scavenger with [CAT]$_0$, [RES]$_0$, or [GAL]$_0$, being 62.5 μM and [TAN]$_0$= 62.5 μM. Hence, DOC$_0$ as hydroquinone was 375 μM for CAT and RES, and 475 μM for GAL and TAN. Relative hTPA yields normalized to DOC$_0$ (%) are shown in Fig. 7a. For comparative purposes, hTPA yields corresponding to HQ$_0$ 62.5 μM (DOC$_0$ = 375 μM) are also included, and were much higher than for the rest of the tested compounds (squares, right Y-axis in Fig. 7a). In absence of FAC (control runs), formation of hTPA was negligible.

As expected, hTPA was not formed when RES was chlorinated (stars in Fig. 7a). Unlike the other selected hydroxyphenols, due to the m-position of the two OH substituents, RES chlorination proceeds through EAS, with chlorine addition, instead of ET (Criquet et al., 2015). Since there is no quinone/semiquinone formation, neither $O_2^-$ nor HOCl$^-$ can be generated. In contrast, for CAT, GAL and TAN, formation of hTPA was detected, although the yields were significantly lower than for HQ. Again, in line to what was proposed in the case of HQ bromination, a possible explanation could be the higher reactivity of these o-hydroxyphenols with FAC compared to HQ (Table 2). Since FAC is consumed more quickly, its reaction with $*\text{OH}$ can take place to a much lower extent.

Previously, it was shown that the presence of borate buffer decreased the reactivity of o-hydroxyphenols towards chlorine through the formation of a borate complex (Criquet et al., 2015). To test if the reactivity of these compounds with chlorine is a key parameter for $O_2^-$ formation, solutions containing 60 μM CAT were chlorinated at pH 7 in 10 mM PBS, pH 8 in 25 mM PBS and pH 8 in 50 mM borate buffer with 4 mM TPA as $*\text{OH}$ scavenger. Fig. 7b shows that the presence of borate leads to a much higher normalized hTPA yield. This supports the hypothesis, that the rate of reaction 7 is important for generation of $*\text{OH}$ through reaction 9.

Taking into account the complexity of these reaction systems, the reactivity between FAC and the hydroquinol (reaction 7) is probably not the only factor affecting $*\text{OH}$ formation. For example, both the relative position of the OH groups and the presence of electron donating/withdrawing groups on the aromatic ring can affect the standard reduction potentials ($E^0$) of the quinone/semiquinone pairs and, therefore, reaction 2 and the stability of $O_2^-$ (Song and Buettner, 2010). To test this, methyldihydroquinone (MeHQ), with higher electron donating properties than HQ due to the presence of the methyl group, was chlorinated at pH 7 alone (MeHQ$_0$ 60 μM) or in combination with HQ ([MeHQ]$_0$ = [HQ]$_0$ = 30 μM), and the concentration of HQ, MeHQ and hTPA were determined. Although the reactivity of MeHQ towards FAC was higher than for HQ as expected (see Fig. 58, SI), the hTPA yields were very similar and close to [HQ]$_0$ 60 μM (Fig. 58, SI). These results can be explained by the influence of the methyl group on $E^0$ (quinone/semiquinone) and hence the ratio of $k_2$/k$_3$. For MeHQ, $k_2$ = $1.1 \times 10^6$ M$^{-1}$s$^{-1}$ and $k_3$/k$_2$ = $1.4 \times 10^{-3}$, whereas for HQ $k_2$ = $5 \times 10^4$ M$^{-1}$s$^{-1}$ and $k_3$/k$_2$ = $5 \times 10^{-5}$ (Song and Buettner, 2010). This means that the presence of the methyl group favors the formation/stabilization of $O_2^-$ countering the negative effect of its higher reactivity towards FAC.

### 3.3. $*\text{OH}$ formation from chlorination of different NOM extracts at pH 7

The formation of $*\text{OH}$ during the chlorination of NOM extracts (SRHA, SRFA, PLFA and NL-NOM; Table S1, SI) was investigated in presence of TPA, and the formation of hTPA was measured after 72 h. The [DOC]$_0$ of the NOM extracts was 375 μM (SRHA), 392 μM (SRFA), 442 μM (PLFA) and 248 μM (NL-NOM). The specific chlorine demand of each extract (pH 7; 72 h contact time) was determined, and resulted to be 0.49, 0.34, 0.38 and 0.36 moles of FAC per mole of DOC$_0$ for SRHA, SRFA, PLFA and NL-NOM, respectively (Fig. 59, SI).

The DOC$_0$ normalized yields of hTPA (mol/mol, %) are shown in Fig. 5 as a function of the specific chlorine dose. The maximum hTPA yields followed the order PLFA (0.3) < SRFA (1.1) < SRHA (1.6) < NL-NOM (2.0). They were achieved for specific chlorine doses slightly higher than the corresponding specific chlorine
demands, which is due to some FAC consumption by TPA under these conditions. After reaching these maxima, higher chlorine doses led to lower hTPA yields probably due to its reaction with FAC (kFAC-hTPA ≈ 0.15 M/C01 s/C01 at pH 7 and 25 °C; this study). It has to be mentioned that hTPA was also formed in very low concentrations in absence of FAC (control runs). This means that the NOM extracts can generate *OH in aqueous solution in the dark. This effect was also observed previously during aeration of different anoxic lake waters and attributed to the development of Fenton-like reactions that involve HQ/SQ and/or Fe(II) (Minella et al., 2015). After 72 h, the hTPA yields (mol hTPA/mol DOCo, %) of the control samples were about 2 orders of magnitude smaller, but following the same sequence as for chlorination (0.002 (PLFA) < 0.006 (SRFA) < 0.02 (SRHA) < 0.03 (NL-NOM)).

Among the selected extracts, PLFA (autochthonous NOM) presents the lowest SUVA254, EDC, aromatic C and phenolic content (Table S1 in SI, and references therein). PLFA has the lowest *OH yield, which seems to indicate that phenolic structures in NOM are involved in *OH formation during chlorination of NOM. With the exception of PLFA, the DOC-normalized hTPA yields for the selected NOMs were close or even higher than for HQ (Fig. 7a). Thus, in case hydroxyphenols in NOM are the main source of *OH during chlorination, HQ moieties with higher *OH formation potential than HQ must be present. This points towards the presence of HQ-like structures in NOM with lower reactivity with FAC than HQ, thus allowing FAC to react with O2*. Alternatively, this could be explained by the presence or formation of quinone structures with higher k2 and k2/k-2 values than HQ (that is, structures with low E0 quinone/semiquinone) favoring the formation/stabilization of O2*. Also, NOM moieties different from phenols could be responsible for *OH formation upon chlorination. Based on the information in Table S1 (SI), N-containing structures are probably not involved, since PLFA has the lowest *OH formation potential, but contains 4–10 times more N than the other extracts. And finally, the higher the iron content, the higher the amount of *OH produced. To this end, a study by Santana-Casiano et al. (2010) carried out in seawater demonstrated that Fe(III)-CAT complexes are formed, followed by Fe(III) reduction to Fe(II) and CAT oxidation to SQ*, with generation of O2*. Hence, the enhancement of *OH formation due to iron complexes in NOM cannot be disregarded.

Water from Lake Greifensee (LG, Switzerland) was also chlorinated (DOC0 250 μM; pH 7.85 after the addition of TPA; specific chlorine demand 0.22 moles of FAC per mole of DOC), and the formation of hTPA was measured. In absence of FAC (control runs), after 72 h contact time, the molar DOC-normalized hTPA yield (%) was ~0.003. The results obtained (see Fig. S10, SI), indicate that DOM in LG water was capable of producing *OH when chlorinated. However, the maximum DOC-normalized hTPA formation was low, ~0.1% mol (mol DOC0)−1, similar to PLFA (also of autochthonous origin; Wenk et al., 2011), and more than ten times lower than for HQ and the other investigated NOM extracts.
3.4. Modelling of the kinetics of *OH formation during HQ chlorination at pH 7

In the last part of the study, the formation of *OH during HQ chlorination at pH 7 has been modeled based on the experimental results and the kinetic data available. All the reactions considered and their rate constants are compiled in Table S3 in SI. The kinetic model is based on reactions (1)–(3) and (7) assuming values of $10^{-7}$ and $10^2$ M$^{-1}$ s$^{-1}$ for $k_1$ and $k_{17}$, respectively ($K_1 = 10^{-14}$; Eyer, 1991).

At pH 7, $k_5$ is very low (Sheng et al., 2014), and from equilibrium 4 the contribution of reaction 6 can be disregarded (see below). Also, due to the strong effect of O$_2$ concentration on hTPA and PHEN formation (see section 3.1.5), keeping in mind that reactions 8 and 10 are oxygen-independent, the contributions of these reactions to *OH formation is expected to be minimal and they have not been considered. For reaction 7, according to the experimental results a molar stoichiometry 2:1 FAC:HQ has been assumed (see Fig. 1b; and also Fig. S11, SI).

According to the pKa of FAC (7.49, equilibrium 12), at pH 7, the molar fractions of HOCl and ClO$^-$ are 0.756 and 0.244, respectively. Since ClO$^-$ does not react with O$_2$ (Long and Bielski, 1980), reaction 9 can be rewritten in terms of FAC with an apparent second order rate constant $k_{9b} = 5.6 \times 10^6$ M$^{-1}$ s$^{-1}$ at pH 7 (Long and Bielski, 1980):

$$FAC + O_2^{1-} \rightarrow k_{9b} \quad Cl^{-} + *OH + O_2$$

(9b)

In presence of BZ as *OH scavenger, the following reactions have been considered (Pan et al., 1993; Fang et al., 1995):

$$*OH + BZ \rightarrow k_{18} \quad *CHDt$$

(16)

$$*CHDt + O_2 \rightarrow k_{17,22} \quad *CHDHpt$$

(17)

$$*CHDHpt \rightarrow k_{18} \quad PHEN + O_2^{1-}$$

(18)

$$*CHDHpt \rightarrow k_{19} \quad P1BZ$$

(19)

where *CHDt is a cyclohexadienyl-type radical intermediate ($k_{16} = 7.8 \times 10^8$ M$^{-1}$ s$^{-1}$; Buxton et al., 1988), *CHDHpt is the cyclohexadienylhydroperoxide-type radical ($k_{17} = 3.1 \times 10^9$ M$^{-1}$ s$^{-1}$; $k_{17} = 1.2 \times 10^9$ s$^{-1}$; Fang et al., 1995), and P1BZ another final product other than PHEN. Since the rate constant of *CHDHpt decrease is 800 s$^{-1}$ (Fang et al., 1995), by assuming a yield of 85% for PHEN and 15% for P1BZ (Dong and Rosario-Ortiz, 2012), $k_{18}$ and $k_{19}$ values have been assumed to be 680 s$^{-1}$ and 120 s$^{-1}$, respectively.

In presence of TPA as *OH scavenger, the following reactions have been considered (Fang et al., 1996):

$$*OH + TPA \rightarrow k_{20} \quad P1TPA$$

(20)

$$*OH + TPA \rightarrow k_{21} \quad *CHDt$$

(21)

$$*CHDt + O_2 \rightarrow k_{22a,22b} \quad *CHDHpt$$

(22a)

$$FAC + 2*CHDt \rightarrow k_{22b} \quad 2hTPA + Cl^{-}$$

(22b)

*CHDHpt $\rightarrow k_{24}$ hTPA + O$_2^{1-}$

(23)

*CHDHpt $\rightarrow k_{24}$ P2TPA

(24)

where P1TPA and P2TPA represent products different from hTPA and *CHDt the cyclohexadienyl-type radical formed when *OH was added to the ortho position of TPA. From the overall rate constant of the TPA - *OH reaction (4.4 $\times$ 10$^9$ M$^{-1}$ s$^{-1}$; Page et al., 2010) and the corresponding yields (85% for *CHDt and 15% for P1TPA; Fang et al., 1996), values of 0.66 $\times$ 10$^9$ M$^{-1}$ s$^{-1}$ and 3.74 $\times$ 10$^3$ M$^{-1}$ s$^{-1}$ have been assigned for $k_{20}$ and $k_{21}$, respectively. According to Fang et al. (1995), $k_{22a} = 1.6 \times 10^7$ M$^{-1}$ s$^{-1}$ and $k_{22a} = 3.4 \times 10^5$ s$^{-1}$, whereas the rate constant of *CHDHpt decomposition (reactions 23 and 24) is 390 s$^{-1}$. By considering a global yield of 30% for hTPA and 55% for P2TPA, $k_{24}$ and $k_{24}$ values were assumed to be 138 s$^{-1}$ and 252 s$^{-1}$, respectively.

Similarly to what was observed by Fang et al. (1995) during the TPA - *OH reaction in presence of IrCl$_6^{3-}$, another possible scenario in which FAC can also oxidize *CHDt directly and irreversibly to hTPA (without formation of *CHDHpt or O$_2^{1-}$) has been considered (reaction 22b), initially assuming the same rate constant for this reaction as for reaction 21, that is, $k_{22b} = 3.74 \times 10^9$ M$^{-1}$ s$^{-1}$.
Finally, the reactions of FAC and *OH with other compounds have also been introduced in the mechanism (reactions 25–28 and 29–34, respectively. See Table S3, SI). However, since at the experimental conditions applied in this study \( f(\cdot \text{OH}) \) > 0.95 for TPA and BZ (Table S1, SI), reactions 29–34 are expected to be negligible.

A comparison between experimental and modeled final concentrations of PHEN and hTPA for chlorination of HQ 62.5 μM for different [FAC]₀/[HQ]₀ molar ratios is shown in Fig. 9a (symbols: experimental results, lines: model calculations). When TPA was used as *OH scavenger the two possible scenarios (that is, without/ with participation of 22b) were considered, in the latter case initially assuming \( k_{22b} = k_{21} \) as indicated before. In Fig. 9a, a good agreement between experimental and modeled data is observed for PHEN, whereas in the case of hTPA its formation through the \( \text{O}_2^- \) dependent mechanism (reactions 22a and 23, without participation of 22b), clearly underestimate the experimental values (Fig. 9a, dotted line). In contrast, when reaction 22b is also considered, the experimental/calculated hTPA agree within the experimental errors (Fig. 9a, solid line).

To further investigate the role of reaction 22b, the two scenarios were tested for lower HQ₀ and FAC₀ conditions, again initially assuming \( k_{22b} = k_{21} \). From the evolution of experimental and modeled hTPA as a function of the FAC concentration (see Fig. S12, SI) it seems that for \( [\text{FAC}]_0 > 1 \text{ mg L}^{-1} \) reaction 22b (solid lines) is the main source of hTPA, and hence \( \sim 0.85 \) mols of hTPA are formed per mole of *OH generated. At lower FAC doses (<1 mg/L) the participation of the \( \text{O}_2^- \)-based mechanism (reactions 22a and 23) increases, with hTPA yields in the range of 30%–85%. The fact that in presence of FAC the yield of hTPA formation from the TPA -*OH reaction is higher than 30% would explain why the experimental concentrations and yields of PHEN and hTPA were similar for [FAC]₀/[HQ]₀ molar ratios \( \leq 1 \) (see Fig. 2). The value for \( k_{22b} \) was optimized by various model calculations to minimize the differences between experimental and calculated hTPA for different initial HQ and FAC concentrations. The optimized value is \( k_{22b} = 10^8 \text{ M}^{-1} \text{s}^{-1} \) (see Fig. S13, SI) which was then applied for all model calculations.

In any case, the yields of both hTPA (from the TPA-*OH reaction) and PHEN (from the BZ-*OH reaction) are lower than 100%. This implies that the total formation potential of *OH (*OHFP), is higher than the measured PHEN or hTPA concentrations. The *OHFPs of HQ 62.5 μM as a function of [FAC]₀/[HQ]₀ at pH 7 in presence of BZ 10 mM or TPA 4 mM were modeled. As shown in Fig. 9b, the total production of *OH in presence of BZ (dashed line) is higher than for TPA (solid line), attributable to the \( \text{O}_2^- \) generated through reaction 18. The calculated relative contribution (%) of \( \text{O}_2^- \) formed from reactions 18 or 23 to *OHFP, is also indicated in Fig. 9b (reaction 18: dashed line; reaction 23: solid line). For TPA this contribution was minimal (lower than 1%), whereas for BZ it decreased from ~22% to 10% as the FAC dose increased.

At the conditions tested and as expected, the introduction of equilibrium 4 and reaction 6 in the mechanism did not cause any change. The presence of FAC and/or BQ in the reaction medium together with the low concentration of \( \text{O}_2^- \) and \( \text{HO}_2^- \) in solution (~10⁻¹¹ and 10⁻¹² M, respectively) prevent the formation of HO₂ through 6.

The effect of the initial HQ concentration on the experimental hTPA yield (that is, for a given [FAC]₀/[HQ]₀ the hTPA yield decreases with increasing [HQ]₀, see Fig. 3) is also well predicted by the proposed mechanisms (Fig. S14, SI).

The negative effect of BQ on *OHFP experimentally observed (see Fig. 4), is also corroborated by this mechanism. As an example, Fig. S15 (SI) shows the modeled reduction of the *OHFP (%) during the chlorination of HQ (62.5 μM, pH 7, air saturated) in presence of different concentrations of BQ, with 4 mM TPA or 10 mM BZ as scavengers. As observed, the higher the initial BQ concentration, the higher the reduction of the *OHFP. For a given [BQ]₀, this effect is attenuated by the BQ formed from HQ chlorination as the [FAC]₀ increases. Thus, the formation of BQ from the HQ - chlorine reaction seems to be the main reason why, for a given [HQ]₀, *OHFP (and hence hTPA or PHEN yields) decreases as the dose of FAC increases, BQ acting as \( \text{O}_2^- \) scavenger.

Finally, the role of dissolved \( \text{O}_2 \) on *OH generation and hence hTPA formation is also well predicted by the proposed mechanisms, as shown in Fig. S16 (SI) for the experiments performed using TPA as *OH scavenger. Fig. S16a (SI) shows that for \( \text{O}_2^- \)-saturated conditions ([O₂] = 1200 μM) reaction 22b seems to be the main source of hTPA, although in this case the model for the optimized \( k_{22b} \) value (dashed line) slightly underestimates the actual hTPA concentrations. However, the model fit is still very good considering the significant number of reactions. Fig. S16b (SI) shows the *OHFP predicted at air-saturated conditions ([*OHFP]ₐir sat., being [O₂]ₐir sat = 300 μM) for HQ 62.5 μM chlorination as a function of FAC dose, together with the evolution of the predicted [*OHFP]/[*OHFP]ₐir sat. ratio for different \( \text{O}_2 \) concentrations. As observed, for dissolved oxygen concentrations 50, 100 and 200 μM, that is, [O₂]/[O₂]ₐir sat values 0.17, 0.33 and 0.67, respectively, the [*OHFP]/[*OHFP]ₐir sat. is ~ 0.21, 0.39 and 0.71, whereas at \( \text{O}_2 \) saturation conditions ([O₂]...
When the aid of different scavengers the generation of hydroxyl radical (‘OH) during the chlorination of α- and p-hydroxyphenols at neutral pH conditions has been demonstrated.

The mechanism of ‘OH generation requires the formation of semiquinones capable of reducing dissolved oxygen to superoxide radical, which further reacts with HOCl to ‘OH.

For p-hydroxyphenols the ‘OH yield (in terms of moles formed per mole of DOC) was clearly higher and comparable to that of SRHA, SRFA and NL NOM, all these extracts presenting high electron donating capacities.

According to the results obtained, a low reactivity towards chloride and a high presence/formation of quinone structures with low E0 favor the formation/stabilization of O2H, thereby increasing the potential of ‘OH formation during the chlorination of NOM.

From the combination of experimental results with kinetic modelling, a mechanism for hydroquinone chlorination at pH 7 has been proposed, which can explain the findings well.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Acknowledgments

Dr Eva M. Rodríguez thanks Ecole Politechnique Fédéral de Lausanne (EPFL, Switzerland) for funding (contract number 49770) and the Consejería de Educación y Cultura of Junta de Extremadura (Spain) for the financial support provided through the mobility program (grant MOV15033).

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.watres.2020.115691.

References


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