Ultrasound/chlorine sono-hybrid-advanced oxidation process: Impact of dissolved organic matter and mineral constituents

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ABSTRACT

In this work, after exploring the first report on the synergism of combining ultrasound (US: 600 kHz) and chlorine toward the degradation of Allura Red AC (ARAC) textile dye, as a contaminant model, the impact of various mineral water constituents (Cl−, SO4 2−, NO3 −, HCO3 − and NO2 −) and natural organic matter, i.e., humic acid (HA), on the performance of the US/chlorine sono-hybrid process was assessed for the first time. Additionally, the process effectiveness was evaluated in a real natural mineral water (NMW) of a known composition. Firstly, it was found that the combination of ultrasound and chlorine (0.25 mM) at pH 5.5 in cylindrical standing wave ultrasonic reactor (f = 600 kHz and P a = 120 W, equivalent to P A ~ 2.3 atm) enhanced in a drastic manner the degradation rate of ARAC; the removal rate being 320% much higher than the arithmetic sum of the two separated processes. The source of the synergistic effect was attributed to the effective implication of reactive chlorine species (RCS: Cl2 , ClO2 and ClO3−) in the degradation process. Radical probe technique using nitrobenzene (NB) as a specific quencher of the acoustically generated hydroxyl radical confirmed the dominant implication of RCS in the overall degradation rate of ARAC by US/chlorine system. Overall, the presence of humic acid and mineral anions decreased the efficiency of the sono-hybrid process; however, the inhibition degrees depend on the type and the concentration of the selected additives. The reaction of these additives with the generated RCS is presumably the reason for the finding results. The inhibiting effect of Cl−, SO4 2−, NO3 − and NO2 − was more pronounced in US/chlorine process as compared to US alone, whereas the inverse scenario was remarked for the effect of HA. These outcomes were associated to the difference in the reactivity of HA and mineral anions toward RCS and ‘OH oxidizing species, in addition to the more selective character of RCS than hydroxyl radical. The displacement of the reaction zone with increasing the additive concentration may also be another influencing factor that favors competition reactions, which subsequently reduce the available reactive species in the reacting medium. The NMW exerted reductions of 43% and 10% in the process efficiency at pH 5.5 and 8, respectively, thereby confirming the RCS-quenching mechanism by the water matrix constituents. Hence, this work provided a precise understanding of the overall mechanism of chlorine activation by ultrasound to promote organic compounds degradation in water.

1. Introduction

Azo dyes are broadly used in dyeing, weaving, tanning and paper industries. This class of dyes represent over 70% of the total production of synthetic dyes [1]. They are aromatic compounds with large structural diversity that always provide a high degree of photolytic, biological and chemical stability; in simply words, they resist to degradation [2]. A numerous of these substances are toxic, carcinogens and mutagenic, making them potential risk for human health and aquatic environments [3,4]. Accordingly, industrial wastewaters containing azo dyes cannot be treated with conventional methods and, consequently, an exigent challenge is devoted in evolving more appropriate...
technologies for the complete removal of these persistent chemicals before discharging their effluents in environmental water basins.

Advanced oxidation processes (AOPs) based on hydroxyl or sulfate radicals have often gained recognition as a promising approach towards removing recalcitrant textile dyes from aqueous effluent in water [5]. UV/H₂O₂ (or SO₂O₅⁻), Fe(II)/H₂O₂ (or SO₂O₅⁻), UV/O₃, photocatalysis, sonolysis and other innovative AOPs like UV/IO₄⁻ and H₂O₂/IO₄⁻ have shown efficient degradation of textile dyes, even in very complex matrices [6–11]. Thanks to the high oxidation potentials of OH and SO₄•⁻ (E₀ = 2.8 and 2.6 V/SHE, respectively), the degradation of micropollutants may be forced for up to mineralization [5].

The basic event of sonication in aqueous solution is the acoustic cavitation phenomenon, which is defined as the formation, growth and collapse of short-lived tiny bubbles in water [12]. The quasi-adiabatic collapse of these bubbles compresses intensively the gas and water vapor trapped therein up to achieving dramatic thermodynamic conditions, i.e. temperature and pressure of the order of thousand kelvin and hundred atmospheres [13]. Hydroxyl and hydrogen radicals have been then generated from the homolytic cleavage of water vapor molecules (H₂O → OH + H), and with other gases present, i.e. O₂ and air, further reactive species like HO₂• and atomic oxygen (O) can be formed during the radical-driven chain reactions inside the cavity at the collapse [14–17]. Of all, ‘OH radical is the most recognized oxidizing agent, as identified by EPR-spin trapping [18,19], chemical dosimetry [20,21] and chemical probe-techniques [22,23]. ‘OH can drive oxidation reaction in the gas phase for volatile substrates, at the bubble/solution interface for hydrophobic compounds and in the bulk solution for hydrophilic substances [24]. However, the highest concentration of these transient species is located at the bubble/solution interface, and only about 10% of this quantity can achieve the bulk of the solution [25].

Chlorine is generally the most used chemical agent for drinking water disinfection [26]. Despite its low activity on microorganisms in biofilms, chlorine can lead to a significant removal of the majority of planktonic bacteria [26]. Lately, ultraviolet (UV) combined with chlorine (i.e., the UV/chlorine process) is being one promising AOP for the degradation of persistent micropollutants [27,28]. The synergy resulted from applying UV/chlorine process for the degradation of diverse organic pollutants was attributed to the implication of ‘OH and reactive chlorine species RCS (Cl•, ClO• and ClO₂•) in the degradation process [29–33]. Primarily ‘OH and chlorine (Cl•) radicals and secondary ClO• and ClO₂• are formed in the system through UV photolysis of HClO/ClO⁻ and their subsequent chain-reaction [28]. A similar reaction mechanism has also been reported for the catalytic activation of chlorine with iron (II and III), which has been successfully applied for fastly degrading of textile dyes [34,35]. Redox potentials of RCS are of 2.43 V/SHE for Cl•, 2.13 V/SHE for ClO• and 1.5–1.8 V/SHE for ClO₂•, and they react with organic matter with second order rate constants varies from ~10⁹–10¹¹ M⁻¹s⁻¹ for Cl• to ~10⁷–10⁸ M⁻¹s⁻¹ for ClO• and 10⁵–10⁶ M⁻¹s⁻¹ for ClO₂• [28,36]. In such multiple-free radical systems, ‘OH and RCS work together to facilitated the degradation of micropolllutants. While ‘OH is non-selective oxidant, RCS are selective and preferentially react with contaminants containing electron rich moieties [37,38], with similar reaction mechanisms as for ‘OH (i.e. electron transfer, H-atom abstraction and addition in unsaturated band). In UV/chlorine system, RCS were found to be the key species implicated in the destruction of trimethoprim [39] and benzoic acid [40], whereas ‘OH dominates for the degradation of rodinazole [41] and ibuprofen [42].

The present work will primarily provide the first report on chlorine activation by high frequency ultrasound (US: 600 kHz) at near neutral pH. The US/chlorine process has been applied for the degradation of Allura Red AC (ARAC) synthetic dye, as a contaminant model. ARAC is a very persistent textile dye of established carcinogenic and toxic effects [43–45] and, therefore, any presence of this compound could have a detrimental effect in aquatic life. After exploring the synergism of combining US and chlorine towards the degradation of ARAC, the impact of various mineral water constituents (Cl⁻, SO₄²⁻, NO₃⁻, HCO₃⁻ and NO₂⁻) and natural organic matter, i.e., humic acid (HA), on the performance of the US/chlorine sono-hybrid process has been assessed for the first time. Additionally, degradation experiments have been conducted in a natural mineral water to check the limit of applicability of this new sono-hybrid process for water treatment.

2. Materials and methods

Throughout the study, ultrapure water was used for solutions and samples preparation. Sodium hypochlorite solution (~16% available active chlorine basis) and Allura Red AC (abbreviation: ARAC; CAS number: 25956–17–6; chemical formula: C₁₁H₄₂N₂Na₂O₅S₂; molecular weight: 496,42 g mol⁻¹) were supplied by Sigma-Aldrich. The molecular structure of ARAC is given in Fig. S1 (supplementary data). All other reagents (NaOH, H₂SO₄, KI, ((NH₄)₂MoO₄·4H₂O), nitrobenzene, humic acid, Na₂SO₄, NaCl, NaHCO₃, NaNO₃ and NaNO₂) were commercial products of the purest grade available (Sigma-Aldrich).

Sonolytic runs were conducted using 150 mL of open to air-solution in the cylindrical water-jacketed glass reactor presented in Fig. S2 (supplementary-data). The sonication system operates at a single frequency (600 kHz) and electric power (120 W) and emits irradiation waves in continuous mode from a piezoelectric disc (diameter of 4 cm) pasted on stainless steel plate (diameter 5 cm) fixed at the bottom of the reactor. The temperature of the irradiating liquid was controlled through the cooling jacket and displayed by a thermocouple (Hanna Instruments) immersed in the solution. The acoustic energy dissipated in the solution (~23 W, corresponding to an acoustic intensity I₀ = 1.83 W/cm²) was estimated by the calorimetric method.

Stock solutions of chlorine (100 mM, pH 5) and ARAC (500 mg L⁻¹, pH ~ 5.5) were prepared and stored in the dark at 4 °C. Experiments were carried out under different conditions at pH 5.5 and ambient temperature (25 ± 1 °C). The pH of the solution was adjusted using NaOH or H₂SO₄ (0.1 M). Quantitative analysis of the dye concentration was performed using a Biochrom WPA Lightwave II UV–vis spectrophotometer at λmax = 504 nm. Hydrogen peroxide concentrations were quantified according to the iodometric method. To ensure reproducibility of the results, all runs were performed in triplicate and results were presented as averages. Error bars, plotted in relevant data, represent the deviation of means.

In tests of chlorination alone (without sonication), runs were conducted in a cylindrical water-jacketed glass cell of 200 mL under a fixed magnetic stirring (300 rpm). When US is applied, no mechanical stirring is applied; the stirring is efficiently ensured by sonication. Sonication through the acoustic cavitation is known to create an efficient micro-stirring of the reaction system (please see [46,47], for example, where the advancing role of stirring by ultrasound is shown on the extraction of arsenic(V) and copper (II) ions for the aqueous solution by an ELM ‘Emulsion Liquid Membrane’ system).

3. Results and discussion

3.1. Characteristics of ARAC chlorination

The effect of pH in the range of 3 to 10 and chlorine dosage (0.05–0.3 mM) on direct chlorination of ARAC (C₀ = 5 mg L⁻¹) at 25 °C is shown in Fig. 1(a) and (b). In all these runs, magnetic agitation was performed at 300 rpm. After a long contact time, i.e., 40 min, low removals of ~ 10–15% were obtained at pH 3–6, whereas a relatively faster rate was observed at pH 8 and 10, where ~ 69% and 90% of ARAC was eliminated after 40 min of treatment, respectively (Fig. 1(a)).

In aqueous solution, the main free chlorine species are Cl₂O, HOCl and OCI⁻ [28,36]. At 25 °C, the calculated distribution of chlorine species vs. solution pH is shown in Fig. S3 for a chlorine concentration of 0.5 mM. Cl₂ is only present at low pH values (pH < 3). HOCl is the predominant free chlorine species at pH < 7.5, and ClO⁻ at pH > 7.5. >98% of free chlorine is present as HOCl in the pH range of 2.5–6, and as ClO⁻ at pH
Fig. 1. ARAC chlorination kinetics for different solution pH (a) and diverse chlorine dosages (b) (conditions: \(C_0 = 5 \text{ mg L}^{-1} (10 \mu \text{M})\), \([\text{chlorine}]_0 = 0.25 \text{ mM for (a) and 0.05–0.3 mM for (b), pH 3–10 for (a) and pH 5.5 for (b), V = 150 \text{ mL}, temperature: 25 \pm 1 ^\circ \text{C, stirring speed: 300 rpm}}\).
> 9. At pH 8, chlorine exists as mixture of 26% HOCl and 74% of OCl⁻. Therefore, under typical water treatment conditions in the pH range 6–9, hypochlorous acid and hypochlorite are the main chlorine species. It is important to note that in addition to these major chlorine species, other chlorine intermediates such as Cl₂ and Cl₂O can also be formed [36], but their concentrations are very low to be considered in chlorination processes [36]. In addition, given that the pKₐ value of ARAC is 11.4 [48], the dye could not change its structure between pH 3–8. Therefore, ARAC reacts with OCl⁻ (pH 8 and 10), whereas the dye molecules showed a strong persistence toward the reaction with HOCl (pH 3–6), even at varying chlorine dosages as stated in Fig. 1(b).

As a recapitulation, at pH 8–10, the breakdown of ARAC by chlorination alone is ensured via its reaction with OCl⁻ present in a large amount (Fig. S3 of the Supplementary Data) whereas, in the pH's range 3–6, the reaction of ARAC with HOCl (present in a large quantity in this range of pH according to Fig. S3) is very slow even at varying chlorine dosages as stated in Fig. 1(b).

3.2. Degradation synergism upon US/chlorine sono-hybrid treatment

Degradation experiments were carried out at pH 5.5 for different reaction systems including chlorination, sonication (US: 600 kHz, 120 W) as well as US/chlorine combination for C₀ = 5 mg L⁻¹ (10 µM) and [chlorine]₀ = 0.25 mM. As seen from Fig. 2(a), after 20 min of treatment, 96.2% of ARAC was eliminated by US/chlorine against 10% for chlorine sole, 57% for sonication alone and 68.5% for the sum of two separated processes. This higher synergism was maintained for pH 3–6 where the sole chlorination did not affect significantly the degradation of the dye (Fig. 1(a)). At pH 8–10, ARAC chlorination happened at appreciable initial rates of 0.41 mg L⁻¹ min⁻¹ (Fig. 1(a)). Besides, Fig. S4(a) of the Supplementary Material showed that the sonolytic degradation of ARAC in basic medium (pH 8–10) is as higher as that ensured by the sole chlorination (0.35 mg L⁻¹ min⁻¹). Consequently, at pH 8 and 10, the effect of applying US/chlorine is additive as the synergistic index is equal to 1 (i.e. r₀,US/Chlorine ~ 0.8 mg L⁻¹ min⁻¹, r₀,US ~ 0.35 mg L⁻¹ min⁻¹, r₀,Cl ~ 0.41 mg L⁻¹ min⁻¹). Therefore, the synergism of applying US/Chlorine treatment was could be only obtained at pH 4–6 where HOCl is the sole chlorine species (Fig. S3).

For the purpose of process improvement, the ARAC degradation by the US/chlorine system has been evaluated through assisting the sonicated system by a mechanical agitation. The obtained results are reported in Fig. S5 of the Supplementary Material. As seen, solution stirring by 100 to 300 rpm has practically no effect on the degradation performance of ARAC. This confirms the performing action of sonication as a perfect stirring tool. Above 500 rpm, the process performance is reduced, mainly to the negative impact of stirring on the wave propagation, as a great part of the acoustic energy dissipated into the solution could be lost by the wave diffraction at the surface of the propeller, in addition to the fact that higher stirring speed could accelerate the coalescence of bubbles which could reduce the number of active bubbles, and then the radicals generation rate in the sonicated solution. Therefore, the conclusion was that there is no need to assist the sonochemical treatment by mechanical stirring; the strategy which has been adopted in all following sections. Also, pH adjustment with acids other than H₂SO₄ can affect the process performance; however, this scenario is not posed as the substitution of H₂SO₄ with HCl did not show any difference.

![Fig. 2. ARAC degradation kinetics (a), and corresponding pseudo-first order rate constant (b), obtained under different oxidation systems, i.e., chlorine, ultrasound (US), US/chlorine, US/NB, and US/chlorine/NB (conditions: frequency: 600 kHz, power: 120 W, C₀ = 5 mg L⁻¹ (10 µM), [chlorine]₀ = 0.25 mM, [NB]₀ = 1 mM, 25 ± 1 °C, pH 5.5). NB: nitrobenzene.](image-url)
(Fig. S6 of the Supplementary Material).

Although it is impossible to make an exact comparison with other advanced oxidation process (AOPs) due to the disparity in the experimental conditions and the substrate pollutant, we have tried to conduct a qualitative comparison with three chlorine-based AOPs (i.e., UV/chlorine, Fe(II)/chlorine and Fe(III)/chlorine) where the degradation process is similar to that of US/chlorine (i.e., 'OH and RCS pathways). The comparison is only based on the removal yield of the different pollutants at given degradation times. The comparison data was added in Table S1 of the Supplementary Data. As shown in this Table, the US/chlorine provides degradation performance as higher as other ones, particularly after a long irradiation time (10 min in this case). At the initial stage, the UV/chlorine provided them most advancement; however, this is because of the lower power supply of our ultrasonic device. If higher power is applied, the process performance of the US/chlorine will be much preferable, as you can see in [49]. Besides, our discovered system is more efficient than the Fe(III)/chlorine process either at the initial or at the final stage of the treatment. On the other hand, despite the fact that the US/chlorine started slowly than the Fe(II)/chlorine system (35% for the former against 80% for the latter), the sono-chlorination overlapped the Fe(II)/chlorine at the last stage of reaction (10 min), where 92% of removal was achieved against 84% for the Fe(II)/chlorine. Furthermore, the performance of the US/chlorine is incomparable with those of US/persulfate and US/periodate. These lasts induced a rather limited enhancement in the degradation of dyes [50,51] as compared to the drastic improvement caused by the US/chlorine process (of this study). This is in addition to the fact that we do not need to remove the residual chlorine after the treatment, while this operation is necessary in the case of persulfate and periodate. Besides, all iron-based AOPs require supplementary steps for removing iron; the problem which is not posed by the developed sono-chlorination system.

3.3. Source of the synergism

ARAC is a highly water soluble solute (solubility: 225 g L\(^{-1}\), log \(K_{ow}\) = – 0.55 [52]) of negligible volatility. This compound cannot enter the bubbles to be pyrolyzed but it could be oxidized outside the bubble by 'OH radical ejected from the bubbles inside at the collapse. We confirmed this mechanism by adding nitrobenzene (NB, a nonvolatile substrate) as a specific scavenger of 'OH (\(k_{NB,OH} = 3.9 \times 10^9\) [36]). NB at 1 mM inhibited the sonolytic degradation of ARAC by >91% as stated in Fig. 1(a) and (b), i.e., the \(k\) value decreased from 0.059 to 0.005 min\(^{-1}\) due NB addition. It seems that the ARAC degradation occurred preferentially at the interfacial region, as it was confirmed by \(H_2O_2\) analysis (Fig. 3). \(H_2O_2\) could mainly be formed at the bubble/solution interface via 2'OH → \(H_2O_2\) (\(k = 5.5 \times 10^9\) M\(^{-1}\) s\(^{-1}\)) [20,53]. The rate of hydrogen peroxide formation decreased from 5.6 \(\mu\)M min\(^{-1}\) in pure water to 4.17 \(\mu\)M min\(^{-1}\) in ARAC aqueous solution (5 mg L\(^{-1}\)), meaning that ARAC molecules scavenge an appreciable portion of hydroxyl radicals located at the reactive interfacial region. Consequently, the sono-degradation of ARAC mainly takes place at the bubble/solution interface via 'OH reaction pathway, but some non-negligible degradation reactions can take place in the bulk solution as NB did not quench completely the dye removal (~90% of ARAC stayed not degraded after a long irradiation time in the presence of NB).

On the other hand, at pH 5.5, hypochlorous acid (HOCl) is the main species of chlorine in aqueous solution (Fig. S3). This species is highly reactive toward the acoustically generated reactive species ('OH, \(H^+\), HO\(^{\cdot}\) and \(H_2O_2\)) and the susceptible reactions between them can generate several reactive chlorine species (RCS: Cl\(^{\cdot}\), ClO\(^{\cdot}\), HOC\(^{\cdot}\) and Cl\(_2\)\(^{\cdot}\)) as can be summarized in the following reaction scheme [2,28,36,54,55]:

\[
\text{'OH} + \text{HOCl} \rightarrow \text{ClO}^{\cdot} + \text{H}_2\text{O} \quad k_1 = 2 \times 10^8 \text{ M}^{-1} \text{s}^{-1} \quad [28] (1)
\]
3.4. Effect of NOM on the performance of the US/chlorine sono-hybrid process

Natural organic matter (NOM) is comprised of humic and fulvic acids, found in almost all surface waters, and are due to breakdown of vegetation [60]. Surface waters contain NOM in the concentration range of 0.1–20 mg L⁻¹ [60]. Humic acids (HA) have molecular weights ranging from several hundred to several thousand and are comprised primarily of aromatic compounds (i.e., with benzene-rings in various forms) [60]. NOM has a negative charge, i.e., hydrophilic compounds, which stay relatively far from the reactive interfacial zone of the cavitation bubbles and being hardly degraded by ultrasound [60]. To look at the effect of HA on the performance of US/chlorine sono-hybrid process, ARAC (5 mg L⁻¹) degradation runs were conducted in the presence of 5, 10 and 15 mg L⁻¹ of HA at pH 5.5. The results for US/chlorine process are shown in Fig. 5a and the corresponding effect of HA concentration on the pseudo-first order rate constant (k) for US and US/chlorine treatment of ARAC are presented in Fig. 5b.

For the sono-chlorination process, HA at 5 mg L⁻¹ did not affect the degradation rate whereas 8% and 15% of reduction in the dye removal at 20 min were recorded for 10 and 15 mg L⁻¹ of HA, respectively (Fig. 5a). The corresponding decrease in the rate constant (k) were 32% and 47% for [HA]₀ = 10 and 15 mg L⁻¹, respectively (Fig. 5b). Conversely, a nearly constant reduction of about 17 ± 2% in the dye elimination at 20 min and ~ 50% in the rate constant k were obtained for the three HA concentrations upon the sole sonication of ARAC (Fig. 5b). This latter observation confirms that HA can compete for hydroxyl radical reaction (i.e., k_{HA-OH} = 2.5 × 10⁸ M⁻¹ s⁻¹) [61] in the bulk solution only since no further reductions in the removal efficiency was obtained for [HA]₀ > 5 mg L⁻¹. Hamdaoui and Merouani [23] showed an insignificant effect of HA (5–40 mg L⁻¹) on the sonochemical degradation of acid orange 7 (20 mg L⁻¹, 57.09 μM) at 600 kHz and 120 W. Similar outcome has also been reported by Neppolian et al. [62] for the oxidation of Arsenic(III) to Arsenic(V) using an initial concentration of 10.10 μM (0.7567 mg L⁻¹) by 20 kHz sonication. The difference between our findings and those of Hamdaoui and Merouani [23] and Neppolian et al. [62] resides in the difference of initial pollutant concentration (C₀). For lower C₀, like in our case, degradation may occur in both the bubble/solution interface and the bulk of the solution whereas, for higher C₀, the bubbler/solution interface is the preferable reaction zone for pollutant oxidation [63]. Thus, addition of HA for cases where higher C₀ is used could not affect the degradation rate as the pollutant could be degraded at the interfacial reactive zone where the availability of HA is unattended due to its too high hydrophilic nature [23]. However, a competition could take place between the contaminant and HA to react with hydroxyl radical if the degradation reaction happens also in the bulk solution, i.e., when lower C₀ is used.

For the system, US/chlorine, the appreciable decrease in the rate constant (k) when [HA]₀ increased from 10 to 15 mg L⁻¹, i.e., 32% and 47%, respectively, was attributed to competition between ARAC molecules and HA to react with both 'OH and R₂C₂O₂⁻ (i.e., k_{HA-ClO₂⁻} = 4.5 × 10⁷ M⁻¹ s⁻¹) [64]; whose able to drive reaction in the bulk solution due to their higher lifetime than 'OH (i.e., 5 μs for Cl₂⁻ [65] and fractions of milliseconds for Cl₂⁺ [66], against ~ 1 ns for 'OH). HA may also react directly with HOCI (k = 0.7–9 M⁻¹ s⁻¹) [66] leading to an instant consumption of available chlorine [67]. Fang et al. [40] have showed that the k value for benzoic acid degradation in UV/chlorine process decreased by about the half when the concentration of NOM increased from 0 to 10 mg/L. Recently, Dong et al. [68] have showed that the removal yield of ciprofloxacin in UV/chlorine system decreased from 98.5% to 62.2% in the presence of only 0.4 mg L⁻¹ of HA. With the continuous increase of HA to 4 mg L⁻¹ [61], the degradation efficiency further decreased to 43.2% [68]. In their detailed study on the degradation of 34 pharmaceuticals and personal care products (PPCPs), Guo et al. [37] have recorded a significant reduction in the degradation rate of some PPCPs, while the degradation of some others where less affected by NOM. The calculated 'OH-degradation rate constant (k'OH) decreased by ~ 10%, while that of R₂C₂O₂⁻ (k'Cl₂⁻) decreased by 19.3% to 90% [37]. Moreover, the steady-state concentration of 'OH, Cl₂⁻ and Cl₂O₂⁻ decreased by < 20% [37]. The authors concluded that R₂C₂O₂⁻ is more sensitive toward NOM than 'OH [37].

Besides, the molecular weight of humic acid (Sigma-product) ranges of 2,000–500,000 g/mol, against only 496.42 g/mol for ARAC. Thus, if we consider only 2000 g/mol for HA, the 5 mg/L of HA corresponds to...
2.5 μM whereas the 5 mg/L of ARAC corresponds to 10 μM. Thus, for 5 mg/L for each one (ARAC and HA), the molar concentration of HA in the solution is much lower than that of ARAC, by about 4-fold (10/2.5 = 4). This justifies the non-significance of HA addition at 5 mg/L on the degradation rate of ARAC substrates are quite similar. It is also seen that even with the use of 15 mg/L of HA, there is no much decrease in the degradation rate of ARAC. However, as the concentration of HA increases to 10 and 15 mg/L (i.e., 5 and 7.5 μM), this could create a competition with the ARAC (5 mg/L or 10 μM) because the molar concentrations of both substrates are quite similar. It is also seen that even with the use of 15 mg/L of HA, there is no much decrease in the degradation rate of ARAC either in the US or in the US/chlorine systems (Fig. 4(a)). However, as the concentration of HA increases to 10 and 15 mg/L (i.e., 5 and 7.5 μM), this could create a competition with the ARAC (5 mg/L or 10 μM) because the molar concentrations of both substrates are quite similar. It is also seen that even with the use of 15 mg/L of HA, there is no much decrease in the degradation rate of ARAC either in the US or in the US/chlorine systems (Fig. 4(b)).

3.5. Effect of water mineral anions on the performance of the US/chlorine sono-hybrid process

Inorganic ions constitute a major part of wastewater. The effect of various mineral water constitutes, i.e., SO₄²⁻, Cl⁻, NO₃⁻, HCO₃⁻ and NO₂⁻ at 1 and 10 mM on the degradation rate constant (k) of ARAC (5 mg L⁻¹) upon US and US/chlorine sono-hybrid system is shown in Fig. 5 for [chlorine]₀ = 0.25 mM and pH 5.5, except for HCO₃⁻ where the initial pH was 8 to ensure maximum quantity of bicarbonate in the reacting medium [69]. Practically, all mineral anions act as radical scavengers, thereby decreasing the degradation rate of the dye in both US and US/chlorine system. However, the degree of reduction differs from one anion species to another and it sometimes may depend on the applied process.

The degradation rate of the dye upon the sole sonication was decreased by ~15% and 20% with adding 1 and 10 mM of Cl⁻, respectively, whereas more inhibiting effect of 18% and 34% was recorded for the sono-hybrid system for 1 and 10 mM of chloride, respectively. In the US system, Cl⁻ can react with ‘OH (Eq. 9), thereby decreasing the degradation rate through the generation of the less reactive radical anion HClO⁻ [70]. The competition between ARAC and Cl⁻ to react with •OH may take place in the bulk solution at low Cl⁻ dosage, but it was also probable at the bubble/solution interface in the
presence of a high quantity of chloride, i.e., 10 mM. The increase in competition with increasing Cl\textsuperscript{−} concentration could result in lowering the degradation rate of the dye. However, even though HOCl\textsuperscript• can simultaneously be decomposed via several routes to generate Cl\textsuperscript{•−}, Cl\textsubscript{2}\textsuperscript{•−} and also OH\textsuperscript• (Eqs. 13–15), no enhancing-degradation rate was observed, which may be attributed to the fact that the consequent radicals from Eqs. 13–15 could also react with Cl\textsuperscript{−} itself which is still in excess as compared to the dye molecules (10 μM), thereby decreasing the overall yield of radicals in the solution. The same trend of chloride effect was observed for several available sonochemical treatments [22,71,72].

In the light of the foregoing, the detrimental effect exerted by Cl\textsuperscript{−} in the US/chlorine system, as compared to the sole sonolysis, can be associated to (i) the scavenging of the acoustically generated ‘OH by the excess of Cl\textsuperscript{−} like in the case of sonolysis alone and (ii) the consumption of HOCl by chloride ions according to the reaction (24), which reduced the available concentration of chloride in the solution (i.e., the yield of RCS could then be lower than that in the absence of Cl\textsuperscript{−}).

\[
\text{HOCl} + \text{Cl}^{-} \rightarrow \text{Cl}_{2}\text{OH}^{-} \quad k_{24} = 1.5 \times 10^{9} \text{ M}^{-1} \text{s}^{-1} [54] (24)
\]

On the other hand, no significant effect was observed for sulfate ions on the sonoletic degradation of ARAC, while remarkable losses of 19% and 34% in the degradation rate constant (k) were obtained for the US/chlorine treatment in the presence of 1 and 10 mM of sulfate, respectively. Sulfate is generally inert toward sonolysis [22,71,72]. Also, no effect of sulfate ions was reported for the degradation of reactive green 12 (RG12) dye by UV/chlorine process, where HOCl\textsuperscript• reacts rapidly with the sulfate ions [34,35]. However, sulfate ions can react efficiently with 'OH with a second-order rate constant of 1 × 10\textsuperscript{10} M\textsuperscript{−1} s\textsuperscript{−1} (Eq. 25) [2,35]. Therefore, the dye removal could be quenched even at a low concentration of NO\textsubscript{2}\textsuperscript{−}. In the US/chlorine system, NO\textsubscript{2}\textsuperscript{−} not only quenches 'OH but also consume RCS as well as free chlorine (HOCI) (Eqs. 26 and 27) [2], resulting in a more inhibiting effect of the degradation rate as compared to the sonication in the absence of HOCI (Fig. 5).

\[
\text{NO}_{2}^{-} + \text{OH} \rightarrow \text{OH}^{-} + \text{NO}_{2} \quad k_{25} = 1 \times 10^{9} \text{ M}^{-1} \text{s}^{-1} (25)
\]

\[
\text{NO}_{2}^{-} + \text{Cl}_{2}\text{O} \rightarrow 2\text{Cl}^{-} + \text{NO}_{2} \quad k_{26} = 2.5 \times 10^{9} \text{ M}^{-1} \text{s}^{-1} (26)
\]

Contrarily to nitrite, NO\textsubscript{3}\textsuperscript{−} moderately inhibited the degradation of ARAC by US and US/chlorine processes (Fig. 5). Losses of 17% and 30% in the k value were calculated for ARAC degradation in US/chlorine system in the presence of 1 and 10 mM of nitrite, respectively. Accordingly, 35% and 48% of losses in the k value were recorded in the US reaction system. It is therefore noted that the decelerating effect of NO\textsubscript{3}\textsuperscript{−} vis-à-vis ARAC degradation was more pronounced for US than US/chlorine system. In the US system, NO\textsubscript{3}\textsuperscript{−} can react rapidly with the acoustically generated ‘OH and H\textsuperscript{\•} according to reactions (28) and (29), thereby producing the lesser reactive NO\textsubscript{2}\textsuperscript{•} and NO\textsubscript{3}\textsuperscript{•} radicals [74]. For the system, US/chlorine, RCS are the main driver of the pollutant degradation. Based on the obtained results, it seems that NO\textsubscript{3}\textsuperscript{−} is less reactive toward RCS as for ‘OH and H\textsuperscript{•}. Unfortunately, there is no available data of second-order rate constants to confirm this suggestion. The one found reaction is that of NO\textsubscript{3}\textsuperscript{−} with Cl\textsuperscript{•−}, given in reaction (30).

\[
\text{NO}_{3}^{-} + \text{OH} \rightarrow \text{OH}^{-} + \text{NO}_{2} \quad k_{28} = 1 \times 10^{9} \text{ M}^{-1} \text{s}^{-1} (28)
\]

\[
\text{NO}_{3}^{-} + \text{H}^{\bullet} \rightarrow \text{NO}_{2}^{-} + \text{OH}^{-} \quad k_{29} = 4.4 \times 10^{8} \text{ M}^{-1} \text{s}^{-1} (29)
\]

\[
\text{NO}_{3}^{-} + \text{Cl}^{-} \rightarrow \text{NO}_{2}^{-} + \text{Cl}^{-} \quad k_{30} = 1 \times 10^{8} \text{ M}^{-1} \text{s}^{-1} (30)
\]

Bicarbonate can react at high second-order rate constants with ‘OH, Cl\textsuperscript{•−} and Cl\textsubscript{2}\textsuperscript{•−} (Eqs. 31–33), while ClO\textsuperscript{•} is not reactive toward HCO\textsubscript{3}\textsuperscript{−} ions (k < 600 M\textsuperscript{−1} s\textsuperscript{−1}) [64,69].

\[
\text{HCO}_{3}^{-} + \text{OH} \rightarrow \text{CO}_{3}^{2-} + \text{H}_{2}O \quad k_{31} = 8.5 \times 10^{8} \text{ M}^{-1} \text{s}^{-1} [64] (31)
\]

\[
\text{HCO}_{3}^{-} + \text{Cl}^{-} \rightarrow \text{CO}_{3}^{2-} + \text{Cl}^{-} + \text{H}^{\bullet} \quad k_{32} = 2.2 \times 10^{9} \text{ M}^{-1} \text{s}^{-1} [64] (32)
\]

\[
\text{HCO}_{3}^{-} + \text{Cl}_{2}\text{O} \rightarrow \text{CO}_{3}^{2-} + 2\text{Cl}^{-} + \text{H}^{\bullet} \quad k_{33} = 8 \times 10^{7} \text{ M}^{-1} \text{s}^{-1} [64] (33)
\]

As a result, the concentration of ‘OH, Cl\textsuperscript{•−} and Cl\textsubscript{2}\textsuperscript{•−} could be decreased in the presence of HCO\textsubscript{3}\textsuperscript{−}, while that of CO\textsubscript{3}\textsuperscript{2−} increased. The oxidation potential of carbonate radical (1.78 and 1.59 V/SHE at pH 7 and pH 12.5, respectively [69]) is less than those of Cl\textsuperscript{•−} (2.43 V/SHE) and Cl\textsubscript{2}\textsuperscript{•−} (2.13 V/SHE) [28,34]. This radical, even less reactive than ‘OH and RCS, has shown a rate enhancement of the sonochemical degradation of
several water contaminants when the pollutant concentration is low and the HCO$_3^-$ loading is relatively high [69,72,75,76]. The mechanism of enhancement was usually attributed to the selectivity and the long lifetime of CO$_3^{2-}$ as compared to OH. However, this scenario was valid only where the CO$_3^{2-}$ radical is reactive toward the target pollutant [2]. In fact, the production of carbonate radical in the UV/chlorine oxidation system has decreased the degradation of some pollutants like naproxen, nalidixic acid, metronidazole, ibuprofen, diethyltoluamide, Propranolol, flumequine, N,N-diethyl-m-toluamide, and caffeine [37,38,77]. However, the degradation of other ones were practically not affected (e.g. erythromycin, azithromycin, roxithromycin, venlafaxine, and Salbutamol) or even increased like the case of Ractopamine and Clenbuterol [37]. In our case, bicarbonate addition decreased the sonolytic degradation of ARAC by about 40% (Fig. 5), indicating that ARAC is presumably not reactive with CO$_3^{2-}$. Thus, the substitution of a certain amount of OH by CO$_3^{2-}$ could reduce the degradation via decreasing the available concentration of hydroxyl radical. Accordingly, Fig. 5 showed that bicarbonate has an insignificant effect on ARAC degradation by US/chlorine process, perhaps for the same reason as reported early for the US system. However, the huge generation of RCS may offset the remarkable reducing effect of bicarbonate on the sonolytic reaction system.

3.6. Process efficiency in natural mineral water

Complementary degradation experiments using US/chlorine process for ARAC degradation were performed in natural mineral water (NMW) whose the characteristics are: Ca$^{2+}$ = 81 mg L$^{-1}$, Mg$^{2+}$ = 24 mg/L, Na$^+$ = 15.8 mg L$^{-1}$, Cl$^-$ = 72 mg L$^{-1}$ (2 mM), SO$_4^{2-}$ = 53 mg L$^{-1}$ (0.55 mM), HCO$_3^-$ = 265 mg L$^{-1}$ (4.34 mM). The achieved results are presented in Fig. 6 for pH 5.5 and 8. For the US/chlorine system, the degradation rate constant in the NMW at pH 5.5 was 0.125 min$^{-1}$, compared to 0.221 min$^{-1}$ in deionized water, thus resulting in 43% of decrease in the value of k. A comparable decrease of 39% was provoked by the NMW for the US/chlorine performance in the NMW compared to the DW matrix is probably owing to the fact that the inhibition of HOCl (through the different anions) is offset by the action of ClO$^-$ on ARAC. However, as it was indicated previously, the efficacy of US/chlorine process is clearly shown in the pH's range 5.5–8 (Fig. 4). However, the decrease in the k values at pH 8 due to NMW is much lower for US/chlorine against US alone (i.e., 10% against 18%, respectively), meaning that the process is very effective at the near neutral pH. Thus, the US/chlorine process is a promising technique for ARAC removal in real natural mineral waters.

3.7. Energetic cost of US/chlorine process

It should be noted that the efficiency of US process is certainly improved by the addition of chlorine to the medium as an oxidation enhancer. This is corroborated through the different findings obtained in Fig. 2, 4, 5 and 6. The technological cost of our system (US/chlorine process) may be determined through the energy consumed (kWh) per 1 g (or mol) of ARAC converted (under the action of OH radicals and RCS), in addition to the amount of chlorine added to the sonoreactor. Considering the maximal elimination of ARAC (Fig. 2) where 96.2% (5 mg/L) is decomposed in 20 min, this gives a conversion of 4.008 × 10$^{-2}$ g L$^{-1}$ s$^{-1}$. On the other hand, the power delivered to the sonicated medium (150 mL) is equal to 120 W, which gives a power density of 800 W/L. As a result, our energetic consumption for ARAC conversion (5 mg/L) is equal to 14.43 g/kWh (2.907 × 10$^{-2}$ mol/kWh) at an ultrasound frequency of 600 kHz. Regarding the consumption of chlorine (cheap compound), if we suppose that the total chlorine (0.25 mM) is activated (via US) during the conversion of ARAC (10 μM), a chloride consumption of 2.083 × 10$^{-7}$ mol L$^{-1}$ s$^{-1}$ is obtained for the same acoustic conditions (600 kHz and 120 W). Regarding the challenges of the US application in the proposed process (US/chlorine AOP), this issue is resumed in the following points:

(i) Reduction of energetic consumption of sonoreactor through the optimization of ultrasound frequency, acoustic intensity and liquid temperature in accordance with the amount of chlorine added to the sonicated solution.

![Fig. 6. Effect of water matrices on ARAC degradation rate constant (k) for US and US/chlorine treatments (conditions: frequency: 600 kHz, power: 120 W, C$_0$ = 5 mg L$^{-1}$ (10 μM), [chlorine]$_0$ = 0.25 mM, 25 ± 1 °C, pH 5.5 and 8). DW: deionized water, NMW: natural mineral water.](image-url)
4. Conclusion

US/chlorine is being discovered as one promising advanced oxidation process which can alternate for US/O₂ toward the degradation of persistent organic contaminants. Chlorine activation with ultrasound can promote the degradation of pollutants in similar manner as for UV/chlorine AOP, in which both ·OH and RCS may participate in the oxidation process. The process was found efficiently synergistic toward the degradation of ARAC. Overall, the presence of NOM and mineral anions decreased the efficiency of the sono-hybrid process; however, the degree of inhibition depends on the type and the concentration of the selected additives.

This study could open effective perspectives for the application of US/chlorine process as alternative AOP for water treatment. Nevertheless, the viability of the process in treating real wastewater treatment is still depending on by-products analysis and evaluating the TOC abatement and the toxicity of the residual effluent. Even though the objective of the present work was to provide initial results on the effect of HA and mineral anion in term of pollutant removal, further complementarily in-depth studies are under realization to check the effective applicability of the US/chlorine process, by which degradation products, TOC evolution and toxicity of the residual effluent will be analyzed.

Finally, with simple comparison to US/O₂ process, the most examined hybrid treatment, the US/chlorine process may present several initial advantages, like:

(i) US/chlorine is facile to handling with more safety as chlorine is less harmful as ozone. Additionally, the liquid phase of chlorine facilitates its employment instead of O₃ which is used in a gas phase (toxic element).

(ii) Chlorine is available and less expensive than ozone, i.e., the latter requires in situ generation via a specific expensive apparatus. Thus, the cost of US/chlorine treatment could be lower than that of US/O₂ for the same experimental operation.

(iii) The synergy resulted from US/chlorine is more important than that reported for US/O₃ process. Therefore, the US/chlorine system allows achieving higher removal yield in a short treatment time than US/O₂.

(iv) The US/chlorine process does not require removal of the residual chlorine as it is originally used as a disinfectant, whereas the elimination of residual O₂ after the sono-ozonation is obligatory.

**CRediT authorship contribution statement**

Oualid Hamdaoui: Investigation, Conceptualization, Methodology, Formal analysis, Project administration, Supervision, Data curation, Funding acquisition, Resources, Visualization, Writing – review & editing. Slimane Merouani: Conceptualization, Methodology, Formal analysis, Writing – original draft, Writing – review & editing. Meriem Alt Idir: Investigation, Visualization, Writing – review & editing. Hadjer C. Benmahmoud: Investigation, Visualization, Writing – review & editing. Aissa Debane: Validation, Visualization, Writing – review & editing. Abdalaziz Alghyamah: Validation, Visualization, Writing – review & editing.

**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.

**Appendix A. Supplementary data**

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ultsonch.2022.105918.

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