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Activation of persulfate by irradiated laterite for removal of fluoroquinolones in multi-component systems

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Graphical Abstract
HIGHLIGHTS

- Laterite/PS/UVA is efficient to simultaneously remove three FQs.
- Degradation rate constants can be ranked as follows: CIP > NOR > FLU.
- Degradation and mineralization extents decrease in binary/ternary systems.
- CIP oxidation in wastewater is less affected by FLU and NOR.
- An excellent catalytic stability of laterite in mixture systems.

Abstract

Although several emerging contaminants (e.g. fluoroquinolones (FQs)) have been simultaneously detected in environmental systems, there is very limited information on their elimination from contaminated waters in multi-component systems. In this study, removal of three FQs including flumequine (FLU), ciprofloxacin (CIP) and norfloxacin (NOR) were investigated in single and mixture systems, using natural laterite soil and persulfate (PS) under UVA irradiation. Both sorption and oxidation reactions contribute to the removal of FQs from aqueous phase, whereas quenching experiments showed that $SO_4^{\cdot-}$ is mainly responsible for the FQs oxidation. The kinetic rate constants can be ranked as follows: CIP > NOR > FLU, regardless of whether the compound was alone or in mixture. The higher degradation rate constant of CIP relative to those of NOR and FLU could be explained by the high reactivity of $SO_4^{\cdot-}$ radical with cyclopropane-ring containing compounds. Fall in oxidation performance was observed in synthetic wastewater, probably due to sulfate radical scavenging by wastewater components. However, degradation rate constants of CIP in wastewater remains unchanged in mixture systems as compared to single ones. This environmentally friendly remediation technology may appear as a promising way for the removal of fluoroquinolone antibiotics from multi-contaminated waters.

Keywords: fluoroquinolones; persulfate; laterite; oxidation, multi-components.
1. Introduction

Because of increasing multiple contaminations in aquatic ecosystems worldwide, surface waters and groundwater often contain multiple contaminants including antibiotics (e.g. (fluoro)quinolones, sulfonamides and macrolides) [1, 2]. As a result of their releases from human and intensive farming activities, different compounds belonging to fluoroquinolones have been simultaneously detected in sediments and aquatic environments [3, 4]. Although affected environmental systems often contain contaminant mixtures, elimination mechanisms and competition effects in multi-component systems have been scarcely investigated. Indeed, most of previous works have mainly focused on removal processes in mono-component systems, and few works have been dedicated to develop cost-effective methods for the elimination of environmental contaminant mixtures [5, 6].

Activated persulfate oxidation processes (PS-AOPs) have received a growing attention to treat a wide range of contaminants in water [7, 8]. In these processes, persulfate anion (S$_2$O$_8^{2-}$) is usually activated by thermal [9], alkaline [10], UVC [11] or transition metal [12-14] to form sulfate radical (SO$_4^{•-}$), which has high oxidation-reduction potential (SO$_4^{•-}$/SO$_4^{2-}$, E° = 2.6-3.2 V vs NHE). One of the most common activators of persulfate (PS) includes Fe under different forms, e.g. dissolved, colloidal or supported [13, 15].

To overcome the drawbacks of homogeneous reaction, heterogeneous activation using Fe-solids may offer several advantages such as recovery/re-use of catalyst, no sludge formation, applicability for a wide range of pH, etc. [14,15]. Different heterogeneous iron-containing catalysts have been used in oxidation processes, including synthetic and natural iron-bearing solids [11-16].

Laterite is an abundant porous and coarse soil, which has diverse minerals in oxide forms, and can be found in several parts of earth including Africa [16, 17]. Although some studies were
dedicated to the use of laterite in Fenton reactions [16-19], much remains to be learned on the sorption/oxidation activity of this mineral and its catalytic stability.

In this present work, the ability of laterite to activate PS under UVA irradiation and then effectively remove mixture of emerging contaminants from water was assessed. Here, flumequine (FLU), norfloxacin (NOR) and ciprofloxacin (CIP), emerging contaminants belonging to fluoro(quinolone) (FQs) group, were used as target contaminants, which co-exist and persist in aquatic systems [20]. It was reported that these FQs concentrations ranged from 3 ng L⁻¹ to 240 µg L⁻¹ in hospital wastewaters and from 0.5 ng L⁻¹ to 6.5 mg L⁻¹ in the fresh surface water, which depend on location and sampling time [2]. Higher concentrations of FQs were also detected in the wastewater of drug producers ranging from 6 ng L⁻¹ to 31 mg L⁻¹ [21]. Removal of FQs from contaminated waters using PS-based oxidation processes have been recently investigated [22-27]. However, the use of laterite as sorbent/catalyst in the removal of FQs has never been investigated to date. Furthermore, most UV sources applied in PS activation studies are in the UVC range (λex ~ 254 nm) [7, 8, 11], while UVA has recently received great attention [11, 22].

First, the influence of laterite loading and PS concentration on the removal efficiency rate was investigated. The effect of water matrices was then evaluated by performing oxidation tests in synthetic wastewaters (SWW). Competition effects were also investigated by determining kinetic rate constants and degradation percentages in single, binary and ternary systems. Finally, quenching experiments were conducted to identify the main reactive species in PS/UVA/laterite system. Moreover, catalytic stability of laterite was assessed by conducting sequential oxidation cycles in PS/UVA/Laterite system. The use of iron-containing soil may offer a cost-effective alternative for large-scale applications of water treatment, particularly in the developing countries.
2. Experimental

2.1. Materials

Flumequine (FLU, 99% purity), norfloxacin (NOR, 98% purity) and ciprofloxacin (CIP, 98% purity) were purchased from Sigma–Aldrich (St. Louis, USA). Potassium persulfate (PS) (K₂S₂O₈, > 99.5% purity), 1,10-phenanthroline (> 99% purity), sodium acetate (> 99% purity), and ferrous ammonium sulfate hexahydrate ((NH₄)₂Fe(SO₄)₂·6H₂O) were obtained from Sigma–Aldrich. Hydrochloric acid (HCl, 37% v/v), Sodium hydroxide (NaOH, 98% purity), Isopropanol (Isopr, C₃H₈O), Tert-butanol (Tert-b, C₄H₁₀O) and 1,4-Benzoquinone (BQ, C₆H₄O₂) were also provided from Sigma–Aldrich. Solutions were prepared with high-purity water obtained from a Millipore Milli-Q system.

Laterite soil was collected from Ivory Coast, West Africa. The sample was ground and passed through 250 µm sieve, and then characterized by X-ray powder diffraction (D8 Bruker diffractometer). X-ray powder diffraction data revealed the presence of quartz, goethite, hematite, kaolinite and gibbsite (Fig. S1). However, the most abundant phases in our sample are quartz, goethite and hematite, which is in agreement with a previous work [17]. Consistently, the TEM-EDX indicated a composition of Fe, Al and O (i.e. under oxide forms), together with two major elements Ca and Mg (Fig. S1). More details are given in the supplementary material. To determine the metal contents in laterite, elements were analyzed using an inductively coupled plasma atomic emission spectrometer (ICP-AES, Jobin-Yvon JY 70 Type HORIBA) after acid digestion of the sample (Table S1). According to elemental composition analysis, the most abundant elements excluding Si are: Fe (151 g kg⁻¹), Mn (184 mg kg⁻¹), Cr (170 mg kg⁻¹) and Al (83 mg kg⁻¹).

Potentiometric titrations of the laterite were conducted in thermostated double walled pyrex cell at 293 K in 0.001, 0.01 and 0.1 M NaCl solutions. The pH value of the suspensions was
adjusted with titrant solutions (HCl or NaOH). The Point Zero Charge (PZC) of laterite soil lied at 6.5 ± 0.3 (Fig. S2).

Based on the N₂ adsorption/desorption isotherms (Fig. S2), the surface adsorption properties such as B.E.T. specific surface area, micro pore volume and total pore volume were determined as 23 m² g⁻¹, 0.06 m³ g⁻¹ and 0.07 m³ g⁻¹, respectively.

2.2. Photoreactor and Photodegradation experiments

All experiments were performed in a 500 mL capacity batch photoreactor (made of borosilicate glass) at room temperature. The reactor was designed in a column shaped in 34 cm high and 3.8 cm diameter. This setup has an enclosed chamber comprising a reactor; an UVA lamp 24 W (Philips PL-L) placed in the center of the glass cell emitting in a wavelength region 320–400 nm with emission peak centered at λmax = 360 nm, yielding a irradiation intensity of 16 mW. cm⁻² as detected with a UVA Radiometer (VLX- 3W equipped with a sensor CX 365, ALYS Technologies, Switzerland). The solution with catalyst was continuously stirred with a magnetic bar at 180 rpm. The pH and temperature of suspension were checked along the experiments.

Two different experiments were carried out beforehand at room temperature. In a first experiment, FQ solution and laterite were stirred overnight in the dark to reach the adsorption equilibrium without UVA irradiation and PS. In the second test, FQ solution, PS, and laterite were mixed simultaneously under UVA irradiation.

Photodegradation experiments were carried out for FLU, NOR and CIP in single, binary and ternary systems. Water samples were spiked with each molecule separately (single-component experiments at 77 µM), or with mixture of two or three molecules in equimolar concentrations (mixture experiments).
Aqueous concentrations of FLU, CIP and NOR were determined using a high performance liquid chromatography (Waters 600 Controller) equipped with a reversed-phase C18 column (250 mm×4.6 mm i.d., 5 μm) and UV detector (Waters 2489). The detector was set to 246, 275, 277 nm for FLU, CIP and NOR, respectively. The mobile phase was a mixture of acetonitrile/water (30/70 v/v) contained 0.1% formic acid. The flow rate was set at 1 mL min⁻¹ in isocratic mode. Under these conditions, the retention times of FLU, NOR and CIP were 6.5, 8 and 5 min, respectively. Total Organic Carbon (TOC) was determined using a TOC-meter (Shimadzu TOC-VCSH). Total dissolved iron concentrations were measured by the 1,10-phenanthroline method at 510 nm [28]. The scavenging experiments were performed using 100 mM of Isopr, Tert-b and BQ each, to determine the contributions of SO₄²⁻, ·OH and HO₂/ O₂⁻ in the degradation of NOR. The optimum concentration of scavengers (i.e. 100 mM) was obtained, according to a preliminary study (Fig. S3). The possible release of trace elements such as Cr, Mn, Cu, Co, Ni and Zn from laterite surfaces was also checked by ICP/AES, which indicated that these elements were under quantification limits (i.e. 0.1 μg L⁻¹) under our experimental conditions. Residual PS concentration was monitored by a UV-vis spectrophotometer (Varian, 50 Probe) at 352 nm according to the modified method of Liang et al. [29]. All experimental runs were performed in triplicates within a temperature of 20 ± 1°C. All results were expressed as a mean value of the 3 experiments.

3. Results and discussion

3.1. Removal efficiencies of single components

The kinetics of FLU, NOR and CIP removal using different oxidation processes in presence and absence of laterite are shown in Figure 1. First, the direct photolysis is less than 12%. The observed degradation in PS/UVA may result from the photochemical activation of PS generating sulfate radicals (i.e. S₂O₈²⁻ + hv → 2SO₄²⁻), though PS activation is supposed to be
low in UVA. Indeed, PS activation decreases as photolytic wavelength increases, as the extinction coefficients of UV photolytic persulfate activation at 248, 308, and 351 nm are 27.5, 1.18, and 0.25 mol⁻¹ cm⁻¹ [30].

The heterogeneous process (i.e. PS/UVA/laterite) showed the best removal performance, with 63%, 72% and 79% for respectively FLU, NOR and CIP, suggesting synergetic effect of laterite and UVA for PS activation. To account for the sorption on laterite, desorption tests (with addition NaOH to pH 11) were carried out and total amounts of FLU, NOR and CIP in aqueous phase were plotted versus time (see Fig. 1). Indeed, the yields of degradation of FLU, NOR and CIP were 53%, 63% and 73% at pH 6.5 ± 0.1, respectively. The amounts of degraded compounds were found slightly lower than the removal amounts (as illustrated by [FLU]tot vs [FLU]aq in Fig. 1), whereas the adsorbed amounts of FLU, NOR and CIP lied at 10%, 9% and 6%, respectively. An overnight pre-equilibration of compound with laterite suspension in the dark before oxidation did not significantly change the kinetic behavior (see e.g. Fig. S4 for FLU). TOC measurements confirmed the oxidative degradation in PS/UVA/laterite system, with 30%, 43% and 55% of mineralization achieved for FLU, NOR and CIP, respectively (Fig. S5a). The Fe leaching of laterite surfaces in PS/UVA/laterite system was very low (i.e, less than 0.03 µmol L⁻¹) (Fig. S5b), as expected from the Fe-solubility equilibrium at pH 6.5 ± 0.1.

Therefore, the efficiency of laterite combined with UVA irradiation could be attributed to the higher production of reactive oxygen species (e.g. SO₄²⁻, 'OH and HO₂⁻/O₂⁻). These radicals could be directly generated from PS under the assistance of the redox cycle of ≡Fe³⁺/≡Fe²⁺ on laterite surfaces promoted by UVA irradiation, as previously reported [31, 32]:

\[ ≡Fe^{III}OH^2+ + h\nu \rightarrow ≡Fe^{II} + 'OH \]  (1)

Indeed, Fe(II) could be generated via photo-reduction of Fe(III)-sites on laterite surfaces, which in turn reacts with PS to generate SO₄²⁻ following:
\[ \text{Fe}^{II} + \text{S}_2\text{O}_8^{2-} \rightarrow \text{Fe}^{III} + \text{SO}_4^{2-} + \text{SO}_4^{2-} \quad k = 2.7 \times 10^1 \text{M}^{-1} \text{s}^{-1} \quad [31, 32] \quad (2) \]

Then, \( \cdot \text{OH} \) could be also generated through the reactions between the \( \text{SO}_4^{2-} \) and \( \text{H}_2\text{O}/\text{HO}^+ \):

\[ \text{SO}_4^{2-} + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + \cdot \text{OH} + \text{H}^+ \quad k \quad [\text{H}_2\text{O}] = 1.3 \times 10^3 \text{ s}^{-1} \quad [33, 34] \quad (3) \]

\[ \text{SO}_4^{2-} + \text{OH}^- \rightarrow \text{SO}_4^{2-} + \cdot \text{OH} \quad (\text{pH} > 8.5) \quad k = 6.5 \pm 1 \times 10^7 \text{ M}^{-1} \text{s}^{-1} \quad [35] \quad (4) \]

Moreover, \( \text{SO}_4^{2-} \) and \( \cdot \text{OH} \) could be also formed through chain reactions dominated by the superoxide anion radical [36]:

\[ \text{Fe}^{II} + \text{O}_2 \rightarrow \text{Fe}^{III} + \text{O}_2^{\cdot} \quad (5) \]

\[ \text{S}_2\text{O}_8^{2-} + \text{O}_2^{\cdot} \rightarrow \text{SO}_4^{2-} + \text{O}_2 + \text{SO}_4^{2-} \quad (6) \]

\[ 2\text{O}_2^{\cdot} + 2\text{H}^+ \rightarrow \text{H}_2\text{O}_2 + \text{O}_2 \quad (7) \]

\[ \text{Fe}^{II} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{III} + \cdot \text{OH} + \text{OH}^- \quad (8) \]

To get further insights into the reaction mechanism, quenching studies using scavengers were performed to identify the main radical species involved in PS/UVAlaterite system. First, isopropanol was supposed to quench efficiently both generated \( \text{SO}_4^{2-} \) and \( \cdot \text{OH} \) considering the second order rate constants of \( k_{\text{Isopr}, \text{SO}_4^{2-}} = 7.42 \times 10^7 \text{ M}^{-1}\text{s}^{-1} \) and \( k_{\text{Isopr}, \cdot \text{OH}} = 1.9 \times 10^9 \text{ M}^{-1}\text{s}^{-1} \) (very high concentration of scavengers (100 mM) were used according to preliminary optimization tests, Fig. S6), while Tert-b can be considered to be more selective toward \( \cdot \text{OH} \) \((k_{\text{Tert-b}, \cdot \text{OH}} = 6.0 \times 10^8 \text{ M}^{-1}\text{s}^{-1})\) than \( \text{SO}_4^{2-} \) \((k_{\text{Tert-b}, \text{SO}_4^{2-}} = 8.31 \times 10^5 \text{ M}^{-1}\text{s}^{-1})\) [37, 38]. It is then possible to estimate the radical scavenging percentage by using the aqueous concentration of each chemical and the reactivity of radicals with NOR \((k_{\text{NOR, SO}_4^{2-}} = 10^7\cdot 10^{10} \text{ M}^{-1}\text{s}^{-1})\) and \( k_{\text{NOR, \cdot OH}} = 1.0 - 6.2 \times 10^9 \text{ M}^{-1}\text{s}^{-1} \) [40, 41]) (more detail is given in supplementary material).

With \( k_{\text{NOR, SO}_4^{2-}} = 10^7 \text{ M}^{-1}\text{s}^{-1}, \) more than 99\% of \( \text{SO}_4^{2-} \) would be quenched with Isopr or Tert-b. With \( k_{\text{NOR, SO}_4^{2-}} = 10^{10} \text{ M}^{-1}\text{s}^{-1}, \) about 91\% and 10\% of \( \text{SO}_4^{2-} \) could be scavenged by Isopr and Tert-b, respectively (Fig.S6). It is worth noting that the bimolecular reaction rate
constants with radicals were mostly determined at pH buffered to 7 (pKa₁ < pH < pKa₂), where the zwitterionic form of NOR is the predominant species (See Fig. S7 for distribution of NOR species at various pH values). In the present work, the oxidation reactions were conducted at pH 6.5 ± 0.1 where both protonated and zwitterionic (or neutral) forms may co-exist.

In the presence of Isopr, about 14% of degradation of NOR was still observed, whereas 49% of NOR degradation was inhibited with respect to the degradation without scavengers. 14% of inhibition was observed using Tert-b, and there was still 49% of NOR degradation (Fig. 2a). We can suppose that the difference observed in NOR degradation (i.e. 35%) when the scavengers were used separately should correspond to the contribution of SO₄⁻⁻.

Addition of BQ to Isopr in PS/UVA/laterite system inhibited completely the degradation of NOR (difference between Isopr + BQ and Isopr = 4%). It is well known that BQ (k_{BQ, \text{HO}_2^*/\text{O}_2^-} = 9.6 \times 10^8 \text{ M}^{-1}\text{s}^{-1}) is an electron acceptor able to interrupt dissolved oxygen accepting electrons, and so acts as a very effective trap to avoid the formation of radical couple (hydroperoxyle radical /superoxide radical anion, \text{HO}_2^*/\text{O}_2^-) [11, 42]. It is then possible that \text{O}_2^- can be formed in our experimental conditions (pH = 6.5 > pKa (\text{HO}_2^*/\text{O}_2^-)), as shown in eq.5. Since 35%, 14% and 14% of degradation of NOR were due to SO₄⁻⁻, \text{OH} and \text{O}_2^-, respectively, the relative contributions of SO₄⁻⁻, \text{OH} and \text{O}_2^- for NOR degradation (63%) were estimated as 56%, 22% and 22%, respectively (Fig. 2b).

3.2. Effects of PS concentration and laterite loading

Removal kinetics of FLU and NOR were determined for a range of PS concentration (0.4 - 2 mM) and laterite loading (0.5 - 4 g L⁻¹) at pH 6.5 ± 0.1. Assuming that FLU or NOR was mainly degraded by sulfate radical species, the degradation kinetic can be described as a second-order reaction:
where \([SO_4^-]\) is steady-state concentration of sulfate radical, \([FLU]\) is concentration of FLU in water, \(k\) is the second-order rate constant, and \(t\) is the reaction time. By assuming that \(SO_4^-\) instantaneous concentration is constant, the kinetics of FLU degradation in water can be described according to the pseudo-first-order equation as given below:

\[
[FLU]_t = [FLU]_0 \exp(-k_{app}t)
\]

where \(k_{app}\) is the pseudo-first-order apparent rate constant (min\(^{-1}\)). \(k_{app}\) (min\(^{-1}\)) obtained by linear regression of \(\ln ([FLU]_t/[FLU]_0)\) versus time \(t\), was plotted versus PS concentration or laterite loading (Fig. 3).

Firstly, kinetic rate constant increased with increasing amount of PS from 0.4 to 0.7 mM, reached an optimum value and then decreased at higher PS concentrations (Fig.3a), probably due to the scavenging effect of \(SO_4^-\) radical by PS and/or recombination of radicals \([43]\) as following:

\[
SO_4^- + S_2O_8^{2-} \longrightarrow SO_4^{2-} + S_2O_8^{4-} \quad k = 6.62 \times 10^5 \text{ M}^{-1}\text{s}^{-1}
\]

\[
SO_4^- + SO_4^- \longrightarrow S_2O_8^{2-} \quad k = 8.1 \times 10^8 \text{ M}^{-1}\text{s}^{-1}
\]

Total consumption of PS was achieved for all PS concentrations, except for the highest concentration \((i.e. \ 2 \text{ mM})\). For the latter, about 50% of initial PS concentration was detected at the end of reaction time (See Fig. S8). Here, we note that the optimal PS concentration for the degradation rate of FLU and NOR is around 0.7 mM, where PS has been totally disappeared after 500 min of reaction time. At this concentration of PS, the degradation rate constant of NOR \((2.2 \times 10^{-3} \text{ min}^{-1})\) is slightly higher than that for FLU \((1.8 \times 10^{-3} \text{ min}^{-1})\) (Fig. 3a). Likewise, kinetic rate constants first increased with laterite loading and then decreased (Fig. 3b). When the laterite concentration varied from 0 to 1 g L\(^{-1}\), the first-order rate constants for
the removal efficiency increased from $7.10^{-4}$ to $1.8.10^{-3}$ min$^{-1}$ for FLU and $6.10^{-4}$ to $2.2.10^{-3}$ min$^{-1}$ for NOR. The fall observed at higher laterite loadings can be explained by screening effects occurring at high solid loading in aqueous suspension and/or scavenging effects of involved radicals by laterite [16, 44]. For instance, excessive Fe-sites can act as SO$_4^{•−}$ radical scavenger as follows [45]:

\[
\text{SO}_4^{•−} + \equiv \text{Fe}^{II} \rightarrow \text{SO}_4^{2−} + \equiv \text{Fe}^{III} \quad k = 3 \times 10^8 \text{M}^{−1} \text{s}^{−1}
\]  

(13)

As shown in Fig.3b, the optimal laterite loading is around 1 g L$^{-1}$, regardless of the target compound.

### 3.3. Degradation of fluoroquinolones (FQs) in mixture systems

In order to evaluate the effectiveness of PS/UVA/laterite system for removal of FQs in multi-component systems, pseudo-first-order apparent rate constants were determined for FLU, NOR and CIP in single, binary and ternary systems using the optimum conditions (0.7 mM of PS and 1 g L$^{-1}$ of laterite) (Fig.4).

The kinetic rate constants can be ranked as follows: CIP > NOR > FLU, regardless of the investigated system (single, binary or ternary). While both SO$_4^{•−}$ and ‘OH can be produced simultaneously in such process, SO$_4^{•−}$ preferentially reacts with target compounds via electron transfer mechanism and ‘OH through hydrogen abstraction or addition reactions [36]. The higher degradation rate constant of CIP relative to those of NOR and FLU could be explained by the high reactivity of SO$_4^{•−}$ radical with the cyclopropane ring of CIP [23]. Indeed, Jiang et al. [23] have compared the different transformation pathways and oxidation byproducts of CIP and enrofloxacin (ENR), two compounds containing cyclopropane ring, with those of norfloxacin and ofloxacin (cyclopropane ring free). First, the most important degradation pathway is initiated by piperazine ring cleavage followed by stepwise oxidation caused by SO$_4^{•−}$ attack. Indeed, SO$_4^{•−}$ radical preferentially reacts as an electron acceptor with the N
atoms on the piperazine ring containing lone pair electrons [46]. This concerns CIP and NOR, but not FLU that lacks the characteristic piperazine ring. Second, an additional degradation pathway can be initiated by the cleavage of the cyclopropane moiety (e.g. in case of CIP) via loss of one CH₂ unit [23] or two CH₂ units [24]. Consistently, Sturini et al. [47] reported that degradation rates of three compounds containing cyclopropane ring (i.e., CIP, ENR and danofloxacin) were greater than those without cyclopropane ring (i.e., levofloxacin, marbofloxacin and moxifloxacin).

In binary systems, the kinetic rate constants of CIP decreased from 17.10⁻³ min⁻¹ to 12.10⁻³ min⁻¹ in CIP/FLU mixture and to 10.10⁻³ min⁻¹ in CIP/NOR mixture. Similar conclusion can be drawn for the two other compounds, i.e. the kinetic rate constants decreased in binary as compared to single systems (Fig. 4a). In the ternary mixture (CIP/NOR/FLU), the kinetic rate constants of CIP, NOR and FLU fall down to 9.9.10⁻³ min⁻¹, 4.4.10⁻³ min⁻¹ and 3.1.10⁻³ min⁻¹, respectively (Fig. 4a). Consistently, the degradation percentages decreased in mixture systems, as compared to those of single compounds (Fig. 4b).

3.4. Effect of water matrices
To assess the ability of PS/UVa/laterite system to remove FQs under environmentally relevant conditions, pseudo-first-order apparent rate constants were determined for FLU, NOR and CIP in single, binary and ternary systems as previously explained, but in synthetic wastewaters (SWW). Synthetic wastewater (SWW) were prepared by adding 400 mg L⁻¹ of NaCl, 50 mg L⁻¹ of citric acid, 30 mg L⁻¹ of ascorbic acid, 100 mg L⁻¹ of saccharose and 230 mg L⁻¹ Na₂HPO₄ to tap water (conductivity 457 µS cm⁻¹, Cl⁻ 0.15 mg L⁻¹, SO₄²⁻ 7.31 mg L⁻¹, NO₃⁻ 34.7 mg L⁻¹, DOC 0.254 mg L⁻¹) and adjusting pH to 7.25 ± 0.1. This composition resembles that of pharmaceutical industry wastewater [48].
As previously observed in pure water, degradation rate constants as well as degradation percentages decreased in binary or ternary systems with respect to those observed for individual compounds (Fig. 5). However, strong inhibition of CIP degradation was observed in SWW, as compared to those measured in pure water regardless of the investigated system (single or mixture). Indeed, 4-fold decrease in degradation rate constant (from $17.10^{-3}$ min$^{-1}$ to $4.1.10^{-3}$ min$^{-1}$) was observed for CIP, while less than 2-fold decrease was observed for NOR or FLU (Figs. 4a and 5a). This may result from the higher reactivity of CIP with sulfate radicals, which make it more subject for scavenging effects than NOR or FLU. In fact, SWW components may compete with target compounds for reaction with radical species and particularly sulfate radical. For instance, chloride ion (400 mg L$^{-1}$) participates to the quenching of SO$_4$* ($k_{SO_4^*,cr} = 2.0 \times 10^8$ M$^{-1}$ s$^{-1}$ [24]) to generate Cl$^*$ which rapidly combines with another chloride ion in water forming dichloride radical anion (Cl$_2$*) $k_{Cl^*,Cl^-} = 0.8 - 2.1 \times 10^{10}$ M$^{-1}$ s$^{-1}$ [46]. Other SWW components, such as phosphates, saccharose or citrate could also actively contribute to sulfate radical scavenging. It is worth noting that, unlike NOR or FLU, no significant decrease in kinetic rate constants or degradation percentages was observed for CIP in mixture systems as compared to single ones (Fig. 5). In conclusion, these results show that CIP oxidation was more affected by SWW components than by the presence of co-occurring FQs (i.e. FLU and NOR).

### 3.5. Reusability of laterite

The reusability of laterite has been evaluated over three successive oxidation cycles, with the above-mentioned optimum conditions, at pH 6.5 ± 0.1 and in the ternary system (Fig. 6). At the end of the oxidation process, the solid was easily removed from the reactor, washed with ultra-pure, and dried at 50°C overnight, and then used for next experiment. Thus, the degradation efficiencies using the recovered solid of FLU, NOR and CIP, remained stable.
during the three oxidation cycles with 52 ± 3%, 64 ± 3%, and 73 ± 3% respectively. Mineralization extent remained also unchanged, with 32 ± 4%, 44 ± 4%, and 55 ± 4% for FLU, NOR, and CIP respectively. Therefore, the good stability of the catalytic activity of laterite could be attributed to the very low iron leaching during oxidation cycles and to the structural stability of the solid. The latter was checked by recording XRD diffractogram at the end of oxidation reaction, which exhibits no change with that obtained before reaction (Fig.S1).

4. Conclusion

This paper demonstrated that laterite soil under UVA irradiation can activate PS to mainly generate $\text{SO}_4^{2-}$ and effectively remove FQs such as FLU, NOR and CIP. Quenching experiments showed implications of $\text{SO}_4^{2-}$, $\cdot\text{OH}$ and couple $\text{HO}_2^\cdot/\text{O}_2\cdot^\cdot$ in the degradation of FQs, but $\text{SO}_4^{2-}$ was the most involved radical in FQs degradation. The kinetic rate constants can be ranked as follows: CIP > NOR > FLU, regardless of the investigated system (single, binary or ternary). The higher degradation rate constant of CIP relative to those of NOR and FLU could be explained by the high reactivity of $\text{SO}_4^{2-}$ radical with cyclopropane ring. In pure water, kinetic rate constants were found lower in binary/ternary systems with respect to those of individual compounds. Unlike FLU and NOR, strong inhibition of CIP degradation was observed in wastewater, probably due to sulfate radical scavenging. However, no significant decrease in kinetic rate constants was observed for CIP in mixture systems as compared to single ones. Collectively, these results showed that CIP oxidation was less impacted by the presence of FLU and NOR, particularly in wastewater. The laterite can be re-used for several oxidation cycles without structural changes or deactivation of surface sites, but water washing and drying processes were applied before re-use in a new oxidation cycle. This new PS-
activation process may promote the development of a cost-effective technology of water remediation, for the removal of emerging compounds in multi-contaminated systems.

Acknowledgements

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Supplementary data

Supplementary data associated with this article can be found, in the online version.
References


**Capture Figures**

**Fig.1.** Kinetic removals of FLU (a), NOR (b) and CIP (c) for various oxidation processes at pH 6.5 ± 0.1: \([\text{FLU}]_0 = [\text{NOR}]_0 = [\text{CIP}]_0 = 77 \mu\text{M}; \ [\text{PS}]_0 = 0.7 \text{ mM}; \ [\text{Laterite}]_0 = 1 \text{ g L}^{-1}; \ \text{UVA irradiation}; \) \([ ]\text{aq} = \text{concentration of pollutant removal in the liquid phase}; \ [ ]\text{tot} = \text{total concentration of pollutant representing both aqueous (residual) concentration and adsorbed concentration obtained after desorption.}\

**Fig.2.** Effect of Isopr, Tert-b and BQ (a) and radical species involved (b) in PS/UVA/Laterite system after 540 min at pH 6.5 ± 0.1: \([\text{Laterite}]_0 = 1 \text{ g L}^{-1}; \ [\text{PS}]_0 = 0.7 \text{ mM}; \ [\text{NOR}]_0 = 77\mu\text{M}; \ [\text{Isopr}]_0 = [\text{Tert-b}]_0 = [\text{BQ}]_0 = 100 \text{ mM}; \ \text{UVA irradiation.}\

**Fig.3.** Effect of PS concentration in presence of 1g/L laterite (a), and laterite loading in presence of 0.7 mM PS (b) on FLU and NOR removals by PS/UVA/Laterite system at pH 6.5 ± 0.1: \([\text{FLU}]_0 = [\text{NOR}]_0 = 77\mu\text{M}; \ \text{UVA irradiation}; \ \text{reaction time} = 540 \text{ min.}\

**Fig.4.** Degradation rate constants \((k_{\text{app}})\) (a) and percentages (b) of FLU, NOR and CIP in single, binary and ternary systems at pH 6.5 ± 0.1: \([\text{FLU}]_0 = [\text{NOR}]_0 = [\text{CIP}]_0 = 19 \mu\text{M}; \ [\text{laterite}]_0 = 1 \text{ g L}^{-1}; \ [\text{PS}]_0 = 0.7 \text{ mM}; \ \text{UVA irradiation}; \ \text{reaction time} = 300 \text{ min.}\

**Fig.5.** Degradation rate constants \((k_{\text{app}})\) (a) and percentages (b) of FLU, NOR and CIP in single, binary and ternary systems in synthetic wastewater (SWW) at pH 6.5 ± 0.1: \([\text{FLU}]_0 = [\text{NOR}]_0 = [\text{CIP}]_0 = 19 \mu\text{M}; \ [\text{laterite}]_0 = 1 \text{ g L}^{-1}; \ [\text{PS}]_0 = 0.7 \text{ mM}; \ \text{UVA irradiation}; \ \text{reaction time} = 300 \text{ min.}\

**Fig.6.** Degradation of FLU, NOR and CIP in three successive oxidation cycles of PS/UVA/Laterite system at pH 6.5 ± 0.1: \([\text{FLU}]_0 = [\text{NOR}]_0 = 77 \mu\text{M}; \ [\text{PS}]_0 = 0.7 \text{ mM}; \ [\text{Laterite}]_0 = 1 \text{ g L}^{-1}; \ \text{UVA irradiation}; \ \text{reaction time} = 540 \text{ min.}
Fig. 1
Fig. 2

(a) Remaining concentration of NOR (%) for different scavengers:
- No scavenger
- Isopr
- Tet-b
- Isopr + BQ

(b) Implication in the degradation (%):
- $\text{SO}_4^{\cdot-}$
- $\text{OH}^{\cdot}$
- $\text{O}_2^{\cdot-}$

Fig. 2
Fig. 3
Fig. 5

### a) $k_{app}$ (min$^{-1}$)

- **CIP**
- **FLU**
- **NOR**

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### b) Degradation percentage (%)

- **CIP**
- **FLU**
- **NOR**

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Fig. 6