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Kinetic Study on Degradation of Micro-organics by Different UV-based advanced oxidation processes in EfOM Matrix

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Abstract:

Effluent Organic Matter (EfOM) contains a large number of substances that are harmful to both the environment and human health. To avoid the negative effects of organic matter in EfOM, advanced treatment of organic matter is an urgent task. Four typical oxidants (H₂O₂, PS, PMS, NaClO) and UV-combined treatments were used to treat micro-contaminants in the presence or absence of effluent organic matter (EfOM), because the active radical species produced in these UV-AOPs are highly reactive with organic contaminants. However, the removal efficiency of trace contaminants was greatly affected by the presence of EfOM. The degradation kinetics of two representative micro-contaminants (benzoic acid (BA) and para-chlorobenzoic acid(p-CBA)) was significantly reduced in the presence of EfOM, compared to the degradation kinetics in its absence. Using the method of competitive kinetics, with BA, p-CBA and 1,4-dimethoxybenzene (DMOB) as probes, the radicals (HO·, SO₄²⁻, ClO·) proved to be the key to reaction species in advanced oxidation processes. UV irradiation on EfOM was not primarily responsible for the degradation of micro-contaminants. The second-order rate constants of the EfOM with radicals were determined to be (5.027±0.643)×10⁻² (SO₄²⁻), (3.192±0.153)×10⁴ (HO·) and 1.35×10⁶ (ClO·) (mg-C/L)⁻¹·s⁻¹. In addition, this study evaluated the production of three radicals based on the concept of R_{ct}, which can better analyze its reaction mechanism.

Key words: Effluent organic matter (EfOM); advanced oxidation processes (AOPs); second-order reaction rate constants; sulfate radical; hydroxyl radical
Urban sewage contains a large amount of organic matter of various and complex types. Although more than 90% of this organic matter can be effectively removed with conventional biological treatment, some refractory organic residue inevitably remains. These refractory dissolved organic substances remaining in sewage after primary and secondary biochemical treatment are collectively referred to as Effluent Organic Matter (EfOM). Vigneswaran, 2006) While the concentration of emerging contaminants (ECs) in EfOM is extremely low in the environment, (Rosal et al., 2010) their high stability in wastewater makes them difficult to degrade with conventional biological treatment. (Taoufik et al., 2021) Conventional wastewater treatment processes including coagulation, sedimentation, filtration, and disinfection—can remove only a limited amount of EfOM, and sometimes highly toxic intermediate products are produced during treatment. Advanced Oxidation Process (AOPs) is a new type of high-efficiency pollutant control technology developed in the 1980s. (Hisaindee et al., 2013) Because AOP has strong oxidizing ability and low selectivity to pollutants, and can remove trace amounts of harmful chemicals and refractory organics, it has been widely used in the treatment of contaminated groundwater, especially for the removal of some special trace pollutants in water. With the development of these advanced oxidation processes, in addition to the initial application of OH, other highly reactive free radicals (such as SO$_{4}^-$·, O$_{2}^-$·, Cl·, etc.) can undergo electron transfer, or hydrogen addition or substitution, to react with refractory organic matter, (Khan and Adewuyi, 2010) thereby causing chemical bond breakage of the organic matter. It is even possible to directly mineralize the organic matter into carbon dioxide and water.

UV/H$_2$O$_2$ is a conventional advanced oxidation process (AOP), based on the production of a hydroxyl radical (HO·) (E$_0$ = 2.8 V) via UV/H$_2$O$_2$. The major water constituents known to scavenge HO· are EfOM and inorganic species such as carbonate, bicarbonate, nitrite, and bromide ions. (Keen et al., 2014; Wols and Hofman-Caris, 2012) HO· water matrix demand is commonly calculated based on measured concentrations of these compounds and the respective second-order rate constants ($k_{OH, P, M}$) for their reaction with HO·. The second-order rate constants for the reaction between HO· and EfOM have been reported as 1.0-4.5×10$^7$ MC$^{-1}$s$^{-1}$ (And and Fulkersonbrekken†, 1998; Donham et al., 2014; Reisz et al., 2003), and these vary depend on the origin, characteristics, and composition of the EfOM.

An advanced oxidation process based on SO$_4^-$· (E$_0$ = 2.6 V) could be applied as an alternative to those based on a hydroxyl radical (HO·) for the remediation of organic pollutants in surface water, groundwater or wastewater. (Hori et al., 2005; Yang et al., 2014) SO$_4^-$· is generated via the activation of peroxymonosulfate (HSO$_5^-$, PMS) or persulfate (S$_2$O$_8^{2-}$, PS) by UV, heat or transition metals. (Matta et al., 2011; Zhou et al., 2013) (Milh et al., 2021) UV/persulfate possesses several advantages, including stability of the precursors (PMS or PS), ease of storage and transportation, high water solubility, versatile activation strategies and a wide operating pH range. (And and Dionysiou, 2004; Das, 2017) (Giannakis et al., 2021)

UV/Cl is an emerging AOP alternative to the UV/H$_2$O$_2$ process, as it produces HO· and reactive chlorine species (RCS). The quantum yields of HOCI and OCI$^-$ by UV photolysis and their absorptivity are reported to be higher than those of H$_2$O$_2$. (Feng et al., 2007; Watts and Linden, 2007) Compared to HO·, RCS such as Cl·, Cl$_2^-$ and ClO$^-$ are powerful oxidants, with oxidation potentials of 2.47 V, 2.0 V and 1.5-1.8 V, respectively. (Alfassi et al., 1988; Beitz et al., 1998)

Previous studies have demonstrated the feasibility of adopting advanced oxidation processes to treat micro pollutants. (Cong et al., 2015) (He et al., 2020) However, data on this topic are still scarce,
especially for UV/oxidant methods for municipal wastewater treatment. In the present work, benzoic acid (BA), para-chlorobenzoic acid (p-CBA) and 1,4-dimethoxybenzene (DMOB) were chosen as model compounds to investigate the degradation of micro pollutants in UV/H₂O₂, UV/PS, UV/PMS and UV/Cl₂ processes. A kinetic model of UV-based AOPs was established for the degradation of micro pollutants in wastewater, and the second-order rate constants of EfOM with radicals were evaluated.

2. Materials and methods

2.1 Samples and chemicals

Secondary wastewater effluent was obtained from a wastewater treatment plant in Beijing with a capacity of 1,000,000 m³/d. The municipal sewage was purified by screens, aerated grit chambers and primary settling, and an A²/O process (anaerobic, anoxic and oxic conditions) and secondary clarification were carried out. The water parameters of the WWTP effluent are listed in Table 1.

Oxone was purchased from Alfa Aesar; H₂O₂ solution (30%), sodium peroxydisulfate (>99%), Sodium hypochlorite (NaClO, ≥8%), and 1,4-dimethoxybenzene (DMOB), benzoic acid (BA; 99%) and tert-butyl alcohol (t-BuOH>99.5%) were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China); para-chlorobenzoic acid (p-CBA; 98%) was purchased from J&K (Beijing, China); and methanol (>99.9%) and acetonitrile (>99.9%) were purchased from J. T. Baker. Ultrapure deionized water (resistivity >18.0 MΩ*cm) was used in all experiments.

2.2. UV-based advanced oxidation experiment

The photo reactor used for the AOPs was equipped with a Xenon lamp peaking at 254 nm (CEL-HXUV300, Zhongjiao Jinyuan). The average UV fluence rate (E₀) was 1.217 mW cm⁻². A 200-mL test solution containing EfOM and 1.0 μM BA, p-CBA, or 5.0 μM DMOB was dosed with the oxidant stock solution (PS, PMS, H₂O₂ or NaClO) and simultaneously exposed to UV irradiation at 25±0.2°C. The oxidant dosages of the reaction were 0.588, 1.176, 2.352 and 5.880 mM, respectively. Samples were collected at 10-min intervals for an hour, for further analysis. Reactions of UV/PS, UV/PMS and UV/H₂O₂ were quenched with 100 mM sulfite and UV/Cl₂ reactions were quenched with ascorbic acid at a molar ratio of [ascorbic acid]/[chlorine]=1.5:1. All tests were conducted at least twice. All data plots represent the average of the experimental data of the duplicated test results.

2.3 Analytical methods

High-performance liquid chromatography (HPLC; Waters 2695, USA) was used to determine [BA], [p-CBA] and [DMOB] (separation conditions given in Table S1). The column used in the liquid chromatographic analysis of BA, p-CBA and DMOB was a Waters Acquity UPLC BEH C18 (1.7 μm, 2.1×100 mm). The dissolved organic matter in the EfOM was determined using a TOC analyzer (TOC; Shimadzu, Japan), and UV absorbance at 254 nm (UV₂₅₄) was measured using a UV/Vis (Evolution 300, Thermo Scientific, USA). Ion chromatography (ICS3000, Dionex Corp., USA) was used to determine [Cl⁻], [NO₃⁻], and [HPO₄²⁻]. The separation was finished in an IonPac AS11 column with a
constant gradient mode. The mobile phase eluent was NaOH solution (30.0 mM) and the flow rate was 1.0 mL/min. Before the separation, 25.0 μL of the sample was injected by an autosampler. The pH was measured using a pH meter (S210 Seven Compact, Mettler Toledo).

2.4 Kinetic model of UV-based AOPs

2.4.1 Pseudo-first-order dynamics model

Because UV irradiation showed no effect on the reference compound degradation (shown in Fig.1), and considering that the degradation reaction of the reference compound (R) may be related to the oxidizing properties of the oxidants (PS, PMS and H$_2$O$_2$), SO$_4^-$ and HO·, the degradation of R can be assumed to follow second-order kinetics:

$$\frac{d[R]}{dt} = k_1[PS][R] + k_2[SO_4^-][R] + k_3[HO^·][R]$$  (1)

$$\frac{d[R]}{dt} = k_4[PMS][R] + k_5[SO_4^-][R] + k_6[[H O^·]][R]$$  (2)

$$\frac{d[R]}{dt} = k_7[H_2O_2][R] + k_8[[HO^·]][R]$$  (3)

where $k_i$ is the second-order rate constant of the reaction of PS, PMS, H$_2$O$_2$, SO$_4^-$ and HO· with R. It is known that the minimum concentration of PS, PMS or H$_2$O$_2$ is 0.588 mM, and it can be assumed that the BA and p-CBA concentration is 1.0 μM [Oxidant] >> [R]. In order to simplify Eqs. (1)-(3), $k_{app}$ is introduced:

$$\frac{d[R]}{dt} = k_{app}[R]$$  (4)

Then, the integral is transformed to

$$-\ln \left( \frac{[R]}{[R]_0} \right) = k_{app}t$$  (5)

where $k_{app}$ is the apparent reaction rate constant (s$^{-1}$), and [R]$_0$ (mM) and [R] (mM) represent the concentrations of R at the reaction times at $\theta$ and $t$, respectively.

2.4.2 Determination of second-order rate constants of HO· and EfOM

To quantify the reactivity of EfOM with HO·, the second-order rate constants ($k_{HO\cdot EfOM}$, M$^{-1}$s$^{-1}$) between HO· and EfOM were determined based on the competition kinetics method using BA, p-CBA and MeOH (or t-BuOH) in UV/H$_2$O$_2$, since the reaction rate constants of BA, p-CBA and MeOH (or t-BuOH) with HO· were known.

$$k_{app}^{R} = k_{HO\cdot R} \times [HO^·]_{ss} = k_{HO\cdot R} \times \frac{\alpha_{HO\cdot}}{k_{HO\cdot EfOM} + k_{HO\cdot MeOH/t-BuOH} + k_{HO\cdot R} + k_{HO\cdot H_2O_2} + k_{HO\cdot H_2O_2}}$$  (6)

Then,

$$\frac{1}{k_{app}^{R}} = \frac{k_{HO\cdot MeOH/t-BuOH}}{k_{HO\cdot R} \times \alpha_{HO\cdot}} + \frac{k_{HO\cdot EfOM} + k_{HO\cdot R} + k_{HO\cdot H_2O_2} + k_{HO\cdot H_2O_2}}{k_{HO\cdot R} \times \alpha_{HO\cdot}}$$  (7)

where $k_{app}^{R}$ is the apparent degradation rate constant of R (s$^{-1}$), and $k_{HO\cdot R}$ is the second-order rate constant for the reaction between HO· and compound R (M$^{-1}$s$^{-1}$). [HO·]$_{ss}$ is the steady-state concentration of HO· (M), and $\alpha_{HO\cdot}$ is the formation rate of HO· (M s$^{-1}$).
In order to eliminate the capture of hydroxyl radicals by inorganic ions in the EfOM and determine
the concentrations of Cl\(^{-}\), NO\(_{3}^{-}\) and HPO\(_{4}^{2-}\), we introduced the parameter \(\beta\):

\[
\beta = k_9 \times [\text{Cl}^{-}] + k_{10} \times [\text{NO}_3^{-}] + k_{11} \times [\text{HPO}_4^{2-}] 
\]

(8)

where \(k_9\), \(k_{10}\), \(k_{11}\) are the second-order rate constants of HO\(^{·}\) with Cl\(^{-}\), NO\(_{3}^{-}\) and HPO\(_{4}^{2-}\),
respectively. (Alfassi et al., 1988; Buxton et al., 1988; Herrmann et al., 1999)

Bringing \(\beta\) into Eq. 7 gives

\[
\frac{1}{k_{\text{app}}} = \frac{k_{\text{HO}^{·}, \text{MeOH/t-BuOH}}}{k_{\text{HO}^{·}, \text{R}}} [\text{MeOH/t-BuOH}] + \frac{k_{\text{HO}^{·}, \text{EfOM}} + k_{\text{HO}^{·}, \text{R}} + k_{\text{HO}^{·}, \text{H}_2\text{O}_2} + k_{\text{HO}^{·}, \text{R}}} {k_{\text{HO}^{·}, \text{R}}} \beta \]

(9)

2.4.3 Determination of second-order rate constants of SO\(_{4}^{−}\) and EfOM

Similar to calculating the secondary reaction rate of HO\(^{·}\) with EfOM, the second-order rate constant
\((k_{\text{SO}_4^{−}, \text{EfOM}}, \text{MC}^{-1} \cdot \text{s}^{-1})\) of the reaction between EfOM and SO\(_{4}^{−}\) is the same as above, using BA, \(p\)-CBA
and MeOH (or \(t\)-BuOH) in the UV/PS system and UV/PMS system.

\[
\gamma = k_{12} \times [\text{Cl}^{-}] + k_{13} \times [\text{NO}_3^{-}] + k_{14} \times [\text{HPO}_4^{2−}] 
\]

(10)

\[
\frac{1}{k_{\text{app}}} = \frac{k_{\text{SO}_4^{−}, \text{MeOH/t-BuOH}}}{k_{\text{SO}_4^{−}, \text{R}}} [\text{MeOH/t-BuOH}] + \frac{k_{\text{SO}_4^{−}, \text{EfOM}} + k_{\text{SO}_4^{−}, \text{R}} + k_{\text{SO}_4^{−}, \text{PS/PMS}} [\text{PS/PMS}] + \gamma} {k_{\text{SO}_4^{−}, \text{R}}} \]

\[
(11)
\]

where \(k_{12}\), \(k_{13}\) and \(k_{14}\) are the second-order rate constants of SO\(_{4}^{−}\) with Cl\(^{-}\), NO\(_{3}^{-}\) and HPO\(_{4}^{2−}\), respectively;
\(k_{\text{app}}\) is the apparent degradation rate constant of R(s\(^{-1}\)); \(k_{\text{SO}_4^{−}, \text{R}}\) is the second-order rate constant for
the reaction between SO\(_{4}^{−}\) and compound “R;” and \(\alpha_{\text{SO}_4^{−}}\) is the formation rate of SO\(_{4}^{−}\) (M s\(^{-1}\)).

2.4.4 Determination of second-order rate constants of ClO\(^{·}\) and EfOM

The second-order rate constants \((k_{\text{ClO}^{·}})\) for the reaction of ClO\(^{·}\) with EfOM were determined by
competition kinetics between EfOM and a reference compound of 1,4-dimethoxybenzene (DMOB),
which was selected to be the reference compound because of its available \(k\) value with ClO\(^{·}\) of \(2.1 \times 10^9\)
M\(^{-1}\) s\(^{-1}\). The pseudo-first order rate constant \((k')\) of DMOB varied depending on the presence or absence
of EfOM, as shown in Eq 12:

\[
k_{R-0}' = \frac{k_{R-i}'}{k_{R-i}'} = 1 + \frac{k_{\text{ClO}^{·}, \text{EfOM}}}{k_{\text{ClO}^{·}, \text{R}}} \times \frac{[\text{EfOM}]_0}{[\text{R}]_0} 
\]

(12)

where \([\text{EfOM}]_0\) and \([\text{R}]_0\) represent the initial concentrations of the EfOM and the DMOB respectively;
\(k_{R-0}'\) and \(k_{R-i}'\) represent the \(k'\) of the reference compound (DMOB) in the absence and presence of
EfOM, respectively; and \(k_{\text{ClO}^{·}, \text{EfOM}}\) and \(k_{\text{ClO}^{·}, \text{R}}\) represent the second-order rate constants of
ClO\(^{·}\) reacting with the EfOM and the DMOB, respectively. (Guo et al., 2018)

3. Results and discussion

3.1 Removal efficiencies of reference compounds in EfOM by UV/ oxidants

Fig. 1 Degradation profile of probes (BA and \(p\)-CBA) in UV-based AOP: (A) UV/ \(\text{H}_2\text{O}_2\), (B) UV/PS, (C)
UV/PMS, (D) UV/Cl; 1-BA, 2-\(p\)-CBA.

Reaction conditions: [BA or \(p\)-CBA] =1.0 \(\mu\)M, \(E_0= 1.217\) mW cm\(^{-2}\), solution pH= 7.9
Fig. 1 compares the degradation of BA (or \( p\)-CBA) by the UV/H\(_2\)O\(_2\), UV/PS, UV/PMS and UV/Cl systems at different oxidant concentrations in EfOM and ultrapure water. BA and \( p\)-CBA are commonly used as radical probe compounds in UV-based AOPs because they have high reactivity with radicals, especially with HO\( \cdot \) and SO\(_4\)\( \cdot \). On the whole, probe compounds have the best degradation effect in ultrapure water, as the concentration of oxidants is 0.588 mM background in different oxidation systems.

The degradation rates of BA in the UV/H\(_2\)O\(_2\), UV/PS, UV/PMS, and UV/Cl systems were 86.07%, 79.05%, 56.94% and 54.91%, respectively, and the degradation rates of \( p\)-CBA in the UV/H\(_2\)O\(_2\), UV/PS, UV/PMS and UV/Cl systems were 69.42%, 50.17%, 44.77% and 60.02%, respectively. It can be concluded that the AOPs had the potential to degradate the contaminants. The pseudo-first-order reaction rate constants for BA were 4.90\( \times \)10\(^{-4}\), 3.54\( \times \)10\(^{-4}\), 2.31\( \times \)10\(^{-4}\) and 2.14\( \times \)10\(^{-4}\) cm\(^2\) m\(^{-1}\) in the UV/H\(_2\)O\(_2\), UV/PS, UV/PMS and UV/Cl systems, respectively, and the pseudo-first-order reaction rate constants for \( p\)-CBA were 2.73\( \times \)10\(^{-4}\), 1.55\( \times \)10\(^{-4}\), 1.60\( \times \)10\(^{-4}\) and 2.58\( \times \)10\(^{-4}\) cm\(^2\) m\(^{-1}\) in the UV/H\(_2\)O\(_2\), UV/PS, UV/PMS and UV/Cl systems, respectively (as shown in Table S2). It can be seen that the reaction rates of BA were higher than those of \( p\)-CBA in the UV/H\(_2\)O\(_2\), UV/PS and UV/PMS systems, indicating that the reaction mechanisms between the two probes and the radicals (HO\( \cdot \) and SO\(_4\)\( \cdot \)) differed. The reaction rate was opposite in the UV/Cl system, where the degradation of \( p\)-CBA in ultrapure water was significantly faster than the degradation of BA, while the opposite results were obtained in the EfOM background.

The degradation of BA and \( p\)-CBA in the secondary effluent organic matter by UV alone had little effect. The efficiency of the degradation rate of the two probes under UV irradiation showed a significant relationship with the molar absorption coefficient (\( \varepsilon \)) and quantum yield (\( \Phi \))(Kwon et al., 2015). As the concentration of oxidants increased, the degradation efficiency of the probe compounds also increased, indicating that the concentration of radicals increased. But this increase was not continuous. It can be seen that when the concentration reached 5.880 mM, the degradation efficiency was basically the same as for the concentration of 2.352 mM. For example, the pseudo-first-order reaction rate constant for BA was 1.13\( \times \)10\(^{-4}\) cm\(^2\) m\(^{-1}\) with a PS concentration of 2.235 mM, and the reaction rate constant for BA was 1.15\( \times \)10\(^{-4}\) cm\(^2\) m\(^{-1}\) with a concentration of 5.880 mM in the UV/PS system. The reason for this difference is that the concentration of the micro-contaminant was limited, in the wastewater. When the concentration of free radicals reaches a certain value, the reaction is saturated. Therefore, an oxidant concentration of 2.352 mM was selected for the next experiment.

The degradation rate of the UV/H\(_2\)O\(_2\) system was higher than for other processes. For example, the pseudo-first-order reaction rate constants for BA were 1.64\( \times \)10\(^{-4}\), 1.13\( \times \)10\(^{-4}\), 1.56\( \times \)10\(^{-4}\), and 1.57\( \times \)10\(^{-4}\) cm\(^2\) mJ\(^{-1}\) in UV/H\(_2\)O\(_2\), UV/PS, UV/PMS and UV/Cl systems, respectively, with a 2.352-mM concentration of oxidants in the presence of EfOM. In the UV/H\(_2\)O\(_2\) system, the major oxidant was the hydroxyl radical (HO\( \cdot \)); the reaction is shown below (Reaction 1). H\(_2\)O\(_2\) is decomposed to generate a powerful oxidant HO\( \cdot \) under the irradiation of ultraviolet light and to trigger free radical chain reactions. PS is stable at room temperature, and the UV led to the cleavage of the O-O bond of PS and generated two SO\(_4\)\( \cdot \) molecules (Reaction 2), which was efficient at degrading the probes. Compared to PDS, PMS has a shorter bond, and more energy is required to cleave the peroxide bond and generate HO\( \cdot \) and SO\(_4\)\( \cdot \) (Reaction 3). The redox potential of SO\(_4\)\( \cdot \) is equal to or even better than HO\( \cdot \) (Ghauch and Tuqan, 2012) but SO\(_4\)\( \cdot \) is more selective than HO\( \cdot \) in degrading the contaminants, which may have led to the lower degradation rate in the UV/PMS and UV/PDS than in the UV/H\(_2\)O\(_2\) system. In the UV/Cl system, there are several kinds of radicals, such as OH\( \cdot \), Cl\( ^{-} \), ClO\(^{-} \), and Cl\(_2\)\( \cdot \), that are responsible for degrading the probes in the effluent organic matter (Reactions 5-10).
In particular, a large amount of SO$_4^{2-}$ is transformed to HO• under basic conditions (Reaction 4). In the presence of p-CBA, the conditions are more acidic compared to those in the presence of BA, indicating that the degradation rate of BA is higher than that of p-CBA. Besides, the main reaction of BA with HO• is the direct attack of the HO• on the aromatic ring to form a hydroxy-substituted compound. (Singla et al., 2004) However, the reaction of SO$_4^{2-}$ with BA first leads to the formation of a radical cation followed by hydrolysis, to form 4-hydroxybenzoic acid (HBA). (Ying-Hong et al., 2011a) This is why the BA degrades faster in UV/H$_2$O$_2$ than in the UV/PS or UV/PMS systems. 

Table 2 Reactions involved in the different UV-based advanced oxidation processes

In the UV/PS system, the degradation efficiencies of BA and p-CBA were 40.66% and 26.85%, respectively, at the concentration of 5.880 mM in the presence of EfOM. Under the same conditions, the degradation efficiencies under UV/PMS were 44.04% and 28.35%, respectively. Mahdi-Ahmed (Mahdi-Ahmed and Chiron, 2014) and Minhwan Kwon (Lee et al., 2018) also found that the removal rate of the probe compound in the UV/PS process is higher than that of the UV/PMS process in ultrapure water, whereas the UV/PMS process has a higher removal rate of the probe compound than the UV/PS process in wastewater from a sewage treatment plant. Guan’s research results proved that UV/PMS significantly enhances the degradation of BA in the pH range of 9-11, while the concentration of PMS has little effect. (Ying-Hong et al., 2011b) By comparing the degradation results in ultrapure water and EfOM, we observed that the degradation of the probe compound by different oxidant concentrations in EfOM is not as effective as in ultrapure water. It can be reasonably inferred that this is most likely because the organic matter contained in EfOM has a trapping effect on the radicals, which leads to different results in the degradation of the probe compound by different oxidation systems. Therefore, we elected to use the competition kinetics method to calculate the second-order rate constants of EfOM and radicals. 

3.2 Contribution of different radicals to contaminant degradation in EfOM

To demonstrate the HO• and SO$_4^{2-}$ reactivity with the EfOM, the second-order rate constants between radicals and the EfOM were determined based on the competition kinetics method using probes (BA, p-CBA) and inhibitors (MeOH, t-BuOH) in the UV/PS, UV/PMS and UV/H$_2$O$_2$ systems. The initial concentration of MeOH (or t-BuOH) was varied from 0 mM to 0.10 mM, and the initial concentrations of probes and oxidants were fixed at 1.0 μM and 2.352 mM, respectively. The introduction of MeOH and t-BuOH significantly inhibited the degradation of probes in UV/PS, UV/PMS and UV/H$_2$O$_2$ compared to results in the absence of quencher, indicating that HO• and SO$_4^{2-}$ are the main reactive oxidizing species. In the sulfate radical systems, when 0.1 mM of radical scavenger was applied, the removal efficiency of BA and p-CBA was reduced by about 30% in the presence of MeOH (Fig. 2); while almost no BA or p-CBA decrease was observed with the addition of t-BuOH (Table S3 and Fig. 3). These results indicated that SO$_4^{2-}$ was the predominant reactive species in the UV/PS and UV/PMS systems, a result consistent with a study by Osburn. (Osburn et al., 2009) Fig. 4 and Fig. 5 show the experimental results of the competition kinetics for the calculation of the second-order rate constants. Due to the reactivity of ClO•, the second-order rate constant for the reaction between ClO• and EfOM was determined using DMOB as a reference compound with varying EfOM concentrations, which react with ClO• at the second-order rate constant of 2.1×10$^8$ M$^{-1}$ s$^{-1}$ (Alfassi et al., 1988) (Fig. 6).
Reaction conditions: \([\text{BA}]_0 = [\text{p-CBA}]_0 = 1.0 \, \mu\text{M}, \; [\text{PS}]_0 = [\text{PMS}]_0 = [\text{H}_2\text{O}_2]_0 = 2.352 \, \text{mM}\)
\([\text{MeOH}]_0 = 0.10, \; 20, \; 50 \; \text{and} \; 100 \, \text{mM}\)

**Fig. 2** Pseudo primary degradation of added MeOH in EfOM (includes BA and p-CBA)

**Fig. 3.** Pseudo primary degradation of added t-BuOH in EfOM (includes BA and p-CBA)

**Fig. 4** Reciprocal of the apparent rate constants for BA and p-CBA vs. MeOH concentrations: (a) UV/H\(_2\)O\(_2\) system; (b) UV/PS system; (c) UV/PMS system
 Reaction conditions: \([\text{f-BuOH}]_0 = 1.0 \, \mu\text{M}, \; [\text{MeOH}]= 2.352 \, \text{mM}, \; \text{pH}=7.9\)

**Fig. 5** Competitive dynamics of BA and p-CBA at different tert-butanol concentrations
 Reaction conditions: \([\text{BA}]_0 = [\text{p-CBA}]_0 = 1.0 \, \mu\text{M}, \; [\text{PS}]_0 = [\text{PMS}]_0 = [\text{H}_2\text{O}_2]_0 = 2.352 \, \text{mM} \; [\text{t-BuOH}]_0 = 0, 10, 20, 50 \; \text{and} \; 100 \, \mu\text{M}\)

**Fig. 6** Competition kinetic plot for ClO\(^-\) reaction with EfOM using 1,4-dimethoxybenzene (DMOB) as a reference compound.
 Reaction conditions: \([\text{chlorine}]= 2.353 \, \text{mM}, \; [\text{DMOB}]_0 = 5.0 \, \mu\text{M}, \; [\text{EfOM}]= 0, 1444, 3344, 5244, 6384 \; \mu\text{L}^{-1}\)

Table S6 lists the second-order rate constants determined for the reactions between the EfOM and radicals. To verify the method and rate constants determined in this study, both probes and inhibitors were measured in each of the different oxidation systems. The second-order rate constant of EfOM and HO\(^-\) was determined to be \((3.192 \pm 0.153) \times 10^4 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\), which is within the commonly reported range of second-order rate constants. These rate constants are similar to the \(10^{3}-10^{11} \, \text{M}^{-1} \text{s}^{-1}\) range of second-order rate constants for the reaction of organics with HO\(^-\) presented in the literature: these are compared with other studies in Table S5. The results of Yang and NAGARNAIK P M—\(3.3 \times 10^4 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\) and \((7.1 \pm 0.81) \times 10^4 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\), respectively (Nagarnaik and Boulanger, 2011; Yang et al., 2016)—are very similar to our findings. In the sulfate radical-mediated oxidation system, the error of the determined rate constant \(k_{\text{SO}_4^- - \text{EfOM}} = (5.027 \pm 0.643) \times 10^2 \, ((\text{mg-C}/\text{L}) \times \text{s}^{-1})\) in this study was small, indicating the reliability of the measurement method. And compared with the other studies shown in Table S4, Yang’s samples of EfOM were isolated from RO Brine A by solid-phase extraction, and contain a large amount of chloride ions, exceeding the probe compounds concentration by 1300- to 2300-fold. They observed that SO\(_4^-\) can be converted to more selective halogen and carbonate radicals, resulting in a wider range of degradation efficiencies among the contaminants (Yang et al., 2016) Zhou measured the absolute rate constants of the reaction of SO\(_4^-\) with four types of organic matter: two fulvic acids and two types of lake organic matter, and their results were close to those in our research (Zhou et al., 2017) The differences in the organic matter contained in the effluent are the main cause of the difference in research results, and are also related to the choice of secondary biochemical reaction process.

Because the UV/Cl process is an emerging advanced oxidation process (AOP) used for the degradation of micropollutants, there has been little research on the secondary reaction rate constant of ClO\(^-\) reacting with different substances in aqueous systems. The second-order kinetic rate constant for EfOM with a chlorooxyl radical was calculated to be \(1.35 \times 10^6 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\) in this research. This value is two orders of magnitude higher than that from EfOM from the Tai Cang wastewater treatment plant in Shanghai—\(1.83 \times 10^4 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\). (Guo et al., 2018) Guo (Guo et al., 2017) researched simulated drinking water prepared by spiking NOM in pure water (1mg L\(^{-1}\)), and the k value of the organic matter with ClO\(^-\) was \(4.52 \times 10^4 \, (\text{mg-C}/\text{L}) \times \text{s}^{-1}\). The difference in k values between the organic matter of different sources and...
ClO \textsuperscript{•} may be due to the different components of the wastewater.

### 3.3 Calculating the radical production rate in EfOM

It is well known that a wastewater matrix such as carbonate species and EfOM affects the removal efficiency of a reference compound (R) in a radical-mediated system (Rosenfeld and Linden, 2007; Yuan et al., 2011). Von Gunten and Linden introduced the concept of R\text{st} to model these complex matrix effects in different AOPs (Elovitz et al., 2000b; Rosenfeld and Linden, 2007). The R\text{st} concept, defined as the experimentally determined radical exposure per UV fluence for a given water matrix and initial oxidants concentration, can characterize the effectiveness of the UV/oxidant AOPs within a specific water matrix (Elovitz et al., 2000a). To quantify the scavenging effect of EfOM, \( R_{\text{HO} \cdot \text{UV}}, \) and \( R_{\text{SO}_4^{\cdot} \cdot \text{UV}} \) were calculated with Eqs. (12) and (13), respectively:

\[
R_{\text{SO}_4^{\cdot} \cdot \text{UV}} = \frac{k_{R,\text{UV}/\text{Oxids},\text{D,EfOM}} - k_{R,\text{UV}/\text{D,EfOM}}} {k_{R,\text{SO}_4^{\cdot} \cdot \text{D,EfOM}}} \int [\text{SO}_4^{\cdot} \cdot \text{UV}]_t \text{d}t
\]  
(13)

\[
R_{\text{HO} \cdot \text{UV}} = \frac{k_{R,\text{UV}/\text{H}_2\text{O}_2/\text{PMS} \cdot \text{D,EfOM}}} {k_{R,\text{HO}}} \int [\text{HO} \cdot \text{UV}]_t \text{d}t
\]  
(14)

where \( k_{R,\text{UV}/\text{D,EfOM}} \) is the apparent first-order rate constant (s\textsuperscript{-1}) of R destruction in EfOM under UV conditions, and \( k_{R,\text{UV}/\text{Oxids},\text{D,EfOM}} \) is the apparent first-order rate constant (s\textsuperscript{-1}) of R degradation in the UV/oxidants system in the presence of EfOM. The superscript “D” indicates that the value is fluence-based. UV fluence (H in the unit of mJ cm\textsuperscript{-2}) is simply the product of \( E_0 \) and t. The detailed calculation method refers to the study of Gao et al. (Gao et al., 2019).

Figure 7 displays the first-order degradation kinetics of probe compounds as a function of applied UV fluence at EfOM in the different UV/oxidant systems. The decay is first order with UV fluence, with only one kinetic regime throughout. Through Figure 7, we can use Formulas (13) and (14) to calculate the data in Table S7, which illustrates the R\text{st} values of SO\textsuperscript{4-} and HO \textsuperscript{•} to BA or p-CBA degradation in a UV/PS, UV/PMS or UV/H\textsubscript{2}O\textsubscript{2} system. When BA is used as a probe, UV/PS produces an SO\textsuperscript{4-} value of \( 9.42 \times 10^{-14} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}, and UV/PMS produces two kinds of radicals —HO and SO\textsuperscript{4-} — \( 2.64 \times 10^{-14} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1} and \( 1.30 \times 10^{-13} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}, of which the one mainly producing SO\textsuperscript{4-} accounted for 83.1%. The HO produced by UV/H\textsubscript{2}O\textsubscript{2} was \( 2.78 \times 10^{-14} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}. And using p-CBA as a probe, UV/PS produced SO\textsuperscript{4-} of \( 1.94 \times 10^{-13} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}, while HO and SO\textsuperscript{4-} in UV/PMS produced fluxes of \( 1.78 \times 10^{-14} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1} and \( 2.48 \times 10^{-13} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}, and the proportion of SO\textsuperscript{4-} was 93.3%. The HO produced by UV/H\textsubscript{2}O\textsubscript{2} was \( 2.66 \times 10^{-15} \) M s cm\textsuperscript{2} mJ\textsuperscript{-1}. By comparing the scavenging effects of different oxidation systems on the two probes, it can be observed that the SO\textsuperscript{4-} produced by UV/PS and UV/PMS was more effective. Although the degradation of EfOM is more rapid in UV/H\textsubscript{2}O\textsubscript{2}, SO\textsuperscript{4-} is more reactive with specific contaminants than HO \textsuperscript{•}, in the degradation of specific pollutants.

In general, the most useful application of the R\text{st} parameter may be in calculating the scavenging properties of the wastewater matrix (Rosenfeld and Linden, 2007) and assessing the potential of SO\textsuperscript{4-} based AOPs in wastewater treatment systems. We used a method of experimentally assessing the exposures of radicals in UV/oxidant AOPs. This parameter can be used to compare water-matrix effects on such UV/oxidant AOPs, to compare the model oxidation of environmental pollutants of concern during AOP treatments in different types of water. This parameter can also be useful in comparing efficiencies among several AOPs.
Fig. 7 Pseudo-first-order degradation of BA (or p-CBA) by different oxidant concentrations in EfOM and ultrapure water

Reaction conditions: [BA or p-CBA] = 1.0 μM, $E_0 = 1.217$ mW cm$^{-2}$, solution pH = 7.9

Conclusions

The effect of UV-based advanced oxidation processes (UV/H$_2$O$_2$, UV/PS, UV/PMS, UV/Cl) in degrading probes (BA, p-CBA, DMOB) in the absence or presence of EfOM was investigated. Direct UV photolysis is not effective because of the low quantum yield. With the addition of oxidants (H$_2$O$_2$, PS, PMS, NaClO), the removal rate was largely improved due to the formation of HO·, SO$_4$· and ClO·. As the concentration of oxidants increases, the degradation efficiency of the probe compounds also increases. The oxidant concentration of 2.352 mM was selected to degrade the probes. Due to its different degradation mechanism, the reaction rate of BA was higher than p-CBA in the UV/H$_2$O$_2$, UV/PS and UV/PMS systems. The degradation rates of BA decreased in the order UV/H$_2$O$_2$ > UV/PS > UV/PMS > UV/Cl, whereas the order of the p-CBA degradation rates was UV/H$_2$O$_2$ > UV/Cl > UV/PS > UV/PMS in ultrapure water. With the competition kinetics method using a probe (BA, p-CBA) and inhibitors (MeOH, t-BuOH) in the UV/PS, UV/PMS and UV/H$_2$O$_2$ systems, results indicated that HO· and SO$_4$· are the principal radicals. The second-order rate constants of the EfOM with radicals were determined to be $(3.192±0.153)×10^4$ M s$^{-1}$ mJ$^{-1}$ (HO·), $(5.027±0.643)×10^2$ M s$^{-1}$ mJ$^{-1}$ (SO$_4$·) and $1.35×10^6$ M s$^{-1}$ mJ$^{-1}$ (ClO·). With the introduction of $R_{ct}$ to the wastewater, results indicated that SO$_4$· was the principal radical in the UV/PMS and UV/PS systems, and HO· is the principal radical in UV/H$_2$O$_2$.

Declarations

Ethics approval and consent to participate

Not applicable

Consent for publication

Not applicable

Availability of data and materials

All data generated or analyzed during this study are included in this published article and its supplementary information files.

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

DY: Writing-Editing & Review, Funding acquisition; GL: Data curation, Writing-Original draft preparation; FQ: Writing-Editing & Review, Conceptualization; JW: Writing-Review, Visualization; YK: Writing-Editing & Review, Funding acquisition; YC: Investigation; MB: Visualization; XL: Methodology

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Figures

Figure 1

Degradation profile of probes (BA and p-CBA) in UV-based AOP: (A) UV/ H2O2, (B) UV/PS, (C) UV/PMS, (D) UV/Cl; 1-BA, 2-pCBA.
Figure 2

Pseudo primary degradation of added MeOH in EfOM (includes BA and p-CBA) Reaction conditions: [BA] 0 = [p-CBA] 0 = 1.0 μM, [PS] 0 = [PMS] 0 = [H2O2] 0 = 2.352 mM, [MeOH] 0 = 0, 10, 20, 50, 100 μM
Figure 3

Pseudo primary degradation of added t-BuOH in EfOM (includes BA and p-CBA) Reaction conditions: [BA]₀ = [p-CBA]₀ = 1.0 μM [PS]₀ = [PMS]₀ = [H₂O₂]₀ = 2.352 mM t-BuOH₀ = 0, 10, 20, 50, 100 μM
Figure 4

Reciprocal of the apparent rate constants for BA and pCBA vs. MeOH concentrations: (a) UV/H2O2 system; (b) UV/PS system; (c) UV/PMS system Reaction conditions: \([BA \text{ or } p-CBA]_0 = 1.0 \mu \text{M}, [\text{oxidant}]_0 = 2.352 \text{ mM}, E_0 = 1.217 \text{ mW cm}^{-2}, \text{solution pH}=7.9\)
Figure 5

Competitive dynamics of BA and p-CBA at different tert-butanol concentrations Reaction conditions $[\text{BA}]_0 = [\text{p-CBA}]_0 = 1.0 \ \mu\text{M}, [\text{PS}]_0 = [\text{PMS}]_0 = [\text{H}_2\text{O}_2]_0 = 2.352 \ \text{mM}, [\text{t-BuOH}]_0 = 0, 10, 20, 50, 100 \ \mu\text{M}$
Figure 6

Competition kinetic plot for ClO• reaction with EfOM using 1,4-dimethoxybenzene (DMOB) as a reference compound. Reaction conditions: [chlorine]₀ = 2.353 mM, [DMOB]₀ = 5.0 μM, [EfOM]₀ = 0, 1444, 3344, 5244, 6384 μg L⁻¹
Figure 7

Pseudo-first-order degradation of BA (or p-CBA) by different oxidant concentrations in EfOM and ultrapure water Reaction conditions: [BA or p-CBA] = 1.0 μM, E0 = 1.217 mW cm⁻², solution pH = 7.9

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