Theoretical Calculation on the Reaction Mechanisms, Kinetics and Toxicity of Acetaminophen Degradation Initiated by Hydroxyl and Sulfate Radicals in the Aqueous Phase

Mengmeng Xu 1, Junfang Yao 1, Simei Sun 2, Suding Yan 3 and Jingyu Sun 1,*

1 Hubei Key Laboratory of Pollutant Analysis & Reuse Technology, College of Chemistry and Chemical Engineering, Hubei Normal University, Cihu Road 11, Huangshi 435002, China; xumm@stu.hbnu.edu.cn (M.X.); yaojf@stu.hbnu.edu.cn (J.Y.)
2 Huangshi Key Laboratory of Photoelectric Technology and Materials, College of Physics and Electronic Science, Hubei Normal University, Huangshi 435002, China; simeisun@hbnu.edu.cn
3 College of Urban and Environmental Sciences, Hubei Normal University, Huangshi 435002, China; yansd@hbnu.edu.cn
* Correspondence: sunjy@hbnu.edu.cn; Tel./Fax: +86-0714-6515602

Abstract: The •OH and SO₄²⁻ play a vital role in degrading pharmaceutical contaminants in water. In this paper, theoretical calculations have been used to discuss the degradation mechanisms, kinetics and ecotoxicity of acetaminophen (AAP) initiated by •OH and SO₄²⁻. Two significant reaction mechanisms of radical adduct formation (RAF) and formal hydrogen atom transfer (FHAT) were investigated deeply. The results showed that the RAF takes precedence over FHAT in both •OH and SO₄²⁻ with AAP reactions. The whole and branched rate constants were calculated in a suitable temperature range of 198–338 K and 1 atm by using the KiSThelP program. At 298 K and 1 atm, the total rate constants of •OH and SO₄²⁻ with AAP reactions were 3.23 × 10⁹ M⁻¹ s⁻¹ and 4.60 × 10¹⁰ M⁻¹ s⁻¹, respectively, considering the diffusion-limited effect. The chronic toxicity showed that the main degradation intermediates were harmless to three aquatic organism, namely, fish, daphnia, and green algae. From point of view of the acute toxicity, some degradation intermediates were still at harmful or toxic level. These results provide theoretical guidance on the practical degradation of AAP in the water.

Keywords: acetaminophen (AAP); density functional theory; degradation mechanisms; rate constants; acute toxicity; chronic toxicity

1. Introduction

The problem of water pollution caused by the drug residues have been paid much more attention. Even though the content of these drugs in the water environment is very low, they brings potential dangers to human health and ecological environment due to its strong persistence, bioaccumulation and slow biodegradation [1,2]. Acetaminophen (AAP), as one kind of antipyretic analgesics, enters to water environment by the excretion of humans and animals. The concentration of AAP rose to 6 µg/L in European STP effluents [3]. Its concentration up to 10 µg/L was detected in natural waters in the United States [4], and over 65 µg/L concentration was measured in the Tyne river in the United Kingdom [5]. The removal of micropollution is challenging for water treatment technology. Researches on this area were also relatively limited.

Advanced oxidation processes (AOPs) are highly efficient engineering technologies in the elimination of water micropollutants. The active free radicals (i.e., •OH and SO₄²⁻) generating in AOPs can decompose these contaminants. The ultraviolet/hydrogen peroxide (UV/H₂O₂), Fenton (H₂O₂/Fe²⁺) and Photo-Fenton (UV/H₂O₂/Fe²⁺) processes can contribute to •OH, which is shown in the following equation:
\[ \text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \cdot\text{OH} + \text{HO}^- \]

\( \cdot\text{OH} \) could be generated from hydrogen peroxide activated by electrochemical process with \( \text{Fe}^{2+}/\text{Fe}^{3+} \), which can degrade pentachlorophenol (PCP) \([6]\). The removal of carbamazepine (CBZ) was attributed to \( \cdot\text{OH} \) formed by coupling \( \text{H}_2\text{O}_2 \) with UV and \( \text{Fe}^{2+}/\text{Fe}^{3+} \) \([7]\). The removal efficiency of three AOP systems has been compared and found the order of \( \text{O}_3/\text{H}_2\text{O}_2/\text{Fe}^{2+} > \text{UV/\text{H}_2\text{O}_2}/\text{Fe}^{2+} > \text{H}_2\text{O}_2/\text{Fe}^{2+} \) \([8]\). Certainly, other processes such as photocatalysis and photoelectrocatalysis are able to produce \( \cdot\text{OH} \) \([9]\).

Meanwhile, the ultraviolet/persulfate (UV/S\text{O}_2\text{O}_5\text{2}^-) can generate \( \text{SO}_4^{*+} \), which is described by the following equation \([10]\):

\[ \text{S}_2\text{O}_8^{2-} \xrightarrow{h\nu} \text{SO}_4^{*+} + \text{SO}_4^{*}^- \]

Surely, \( \cdot\text{OH} \) can be produced when \( \text{SO}_4^{*+} \) reacts with \( \text{H}_2\text{O} \), which is presented by the following equation \([11]\):

\[ \text{SO}_4^{*+} + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + \cdot\text{OH} + \text{H}^+ \]

Thus, \( \cdot\text{OH} \)-mediated and \( \text{SO}_4^{*+} \)-mediated degradation of contaminants were available. The redox potential and rate constants are summarized in Table 1. It was reported that the \( \cdot\text{OH} \)-initiated degradation rate constant was about \( 10^{8}–10^{10} \text{ M}^{-1} \text{ s}^{-1} \). The \( \text{SO}_4^{*+} \)-initiated rate constant was about \( 10^{2}–10^{10} \text{ M}^{-1} \text{ s}^{-1} \) \([12]\). The degradation rates of two reactive radicals are nearly equivalent, which is consistent with their high redox potential (2.5–3.1 V for \( \text{SO}_4^{*+} \) versus 1.8–2.7 V for \( \cdot\text{OH} \)) \([13–15]\). The degradation processes of contaminants triggered by \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) were investigated in recent years. For example, Tong et al. determined the rate constants of syringic acid reactions with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) in aqueous phase by laser flash photolysis. They found that \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) possessed similar reaction rate at the same pH \([16]\). Gao et al. measured the rate constants of neutral sulfamethoxazole with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) were \( (7.27 \pm 0.43) \times 10^9 \) and \( (2.98 \pm 0.32) \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \) in the systems of \( \text{UV/H}_2\text{O}_2 \) and \( \text{UV/PS} \), respectively \([17]\). Similarly, Wang et al. detected the rate constants for AAP with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) reactions were \( (3.26 \pm 0.41) \times 10^9 \) and \( (1.80 \pm 0.17) \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \) in the \( \text{Fe}^{2+}/\text{persulfate} \) system, respectively \([18]\). The second-order rate constants of \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) were conformed as \( 5.15 \times 10^9 \) and \( 7.66 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \), respectively, using the ultraviolet light emitting diode (UV-LED)-based method by Li et al. \([19]\). However, the study of degradation mechanisms of \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) with the target contaminants still faced with great challenge. At atom level, quantum chemistry calculation was a powerful tool to gain a in-depth understanding for mechanisms and kinetics of \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) reacting with some pollutants \([20–23]\).

| Radicals (M) | Redox Potential \( ^{a} \) (V) | Rate Constants \( ^{b} \) (M\(^{-1}\) s\(^{-1}\)) | The Second-Order Rate Constants of Neutral Sulfamethoxazole \( ^{c} \) (M\(^{-1}\) s\(^{-1}\)) | The Second-Order Rate Constants of Acetaminophen (M\(^{-1}\) s\(^{-1}\)) |
|-------------|-------------------------------|---------------------------------|------------------------------------------------|---------------------------------|
| \( \cdot\text{OH} \) | 1.8–2.7 | \( 10^{8}–10^{10} \) | \( (7.27 \pm 0.43) \times 10^9 \) | \( (3.26 \pm 0.41) \times 10^9 \) |
| \( \text{SO}_4^{*+} \) | 2.5–3.1 | \( 10^{7}–10^{10} \) | \( (2.98 \pm 0.32) \times 10^9 \) | \( (1.80 \pm 0.17) \times 10^9 \) |

\(^{a}\) (Xiao, et al., 2020; Devi, et al., 2016; Ghanbari, et al., 2017); \(^{b}\) (Li, et al., 2020, \[12]\); \(^{c}\) (Gao, et al., 2020); \(^{d}\) (Wang, et al., 2019); \(^{e}\) (Li, et al., 2020, \[19]\).

Theoretical studies are essential for discussing the degradation processes of AAP with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \). Therefore, the reaction mechanisms and kinetics of the AAP with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) have been studied by using quantum chemistry calculations. Rate constants of every possible pathways for AAP with \( \cdot\text{OH} \) and \( \text{SO}_4^{*+} \) reactions were calculated. Even more importantly, the ecotoxicity of AAP and its degradation products has been evaluated in order to know their risk.
2. Computational Methods

2.1. Mechanism Calculation

Usually, reaction mechanisms are investigated by Density functional theory (DFT). M06-2X functional can solve noncovalent interactions for some complexes better than other density functional such as B3LYP [24]. The functional ratio of exchange correction of M06-2X is 54% which will obtain more accurate energies [25]. Furthermore, M06-2X method of DFT was selected in the reactions of AAP with \( \text{-}\bullet \text{OH and SO}_4^{2-} \) without hesitation, because satisfactory results were acquired on the degradation of some micropollutants [27–30]. For example, the thermodynamic and kinetic data for ibuprofen reactions with hydroxyl and sulfate radicals reported by Yang et al. were calculated with M06-2X method [27]. All electronic structures and energy calculations were performed using Gaussian 09 software [31]. The reactants (R), transition states (TS) and intermediates (IM) were optimized at the M06-2X/6-31+G(d,p) level. IM (all positive frequencies) and TS (only one imaginary frequency) are primarily identified by harmonic vibration frequency analysis. Moreover, the method of intrinsic reaction coordinates (IRC) was applied to determine every right transition state [32]. The water solvent effect was taken into account by a universal solvation model (SMD) [33] when these structures were optimized in the aqueous phase. Based on right structures, the single point energies were calculated accurately at high level of M06-2X/6-311++G(3df, 2p). The Gibbs free energy barrier of activation (\( \Delta G^\ddagger \)) and free energy of reaction (\( \Delta G \)) are calculated as follows:

\[
\Delta G^\ddagger = G_{\text{TS}} - G_{\text{R}} \\
\Delta G = G_{\text{IM}} - G_{\text{R}}
\]

2.2. Kinetic Calculation

The conventional Transition State Theory (TST) was used to calculate the rate constants implemented by KiSThelP program [34] that has obtained accurate results for contaminants with free radicals reactions [35–39]. The calculation formula is employed in KiSThelP:

\[
k = \kappa \sigma \frac{k_b T}{h} \left( \frac{R T}{P_0} \right)^{\Delta n} e^{-\frac{\Delta G^\ddagger}{R T}}
\]

Some parameters need to be explained. \( \kappa \) is tunneling correction factor of Wigner approach [34]. \( \sigma, k_b \) and \( h \) are the reaction path degeneracy, Boltzmann’s constant and Planck’s constant, respectively. \( \Delta G^\ddagger \) is the standard Gibbs free energy of activation. \( RT/P_0 \) has the unit of the inverse of a concentration. For bimolecular reactions, \( \Delta n \) is equal to 1.

The diffusion-limited effect was considered to obtain the apparent rate constants (\( k_{\text{app}} \)) of aqueous phase based on Collins-Kimball theory [40].

\[
k_{\text{app}} = \frac{k_{aq} k_D}{k_{aq} + k_D}
\]

where, \( k_{aq} \) is calculated by TST as aqueous rate constant. \( k_D \) is calculated by the Smoluchowski equation as the diffusion-limited rate constants:

\[
k_D = 4 \pi R_{AB} D_{AB} N_A
\]

\( R_{AB} \) means the reaction distance, and \( N_A \) is Avogadro’s number, \( D_{AB} \) represents the sum of diffusion coefficient of the reactants A (AAP) and B (\( \bullet \text{OH or SO}_4^{2-} \)). The calculations of \( D_A \) and \( D_B \) are realized by using the Stokes–Einstein approach [41]:

\[
D = \frac{k_b T}{6 \pi a \eta}
\]

where \( k_b, T, \eta, \) and \( a \) are the Boltzmann constant, temperature, viscosity of the solvent, and radius of the solute, respectively. For water, \( \eta = 8.9 \times 10^{-4} \text{ Pa s} \).

2.3. Ecotoxicity Calculation

The aquatic toxicity of AAP and its degradation products was evaluated by using the Structure Activity Relationship (SAR) method with the ECOSAR program [42], which has
been successfully used to evaluate the acute and chronic toxicity [43–46]. Three aquatic organisms of green algae, daphnia and fish were considered to assess the acute and chronic toxicities. Acute toxicity of the target compounds was estimated by median lethal concentration (LC50) and median effect concentration (EC50). LC50 is defined 50% lethal concentration for fish and daphnia in 96 and 48 h, respectively. EC50 is 50% effective concentration for green algae in 96 h. The chronic toxicity was described by the chronic toxicity value (ChV) for green algae, daphnia and fish.

3. Results and Discussion
3.1. Degradation Mechanisms

The degradation mechanisms of AAP initiated by •OH and SO$_4$•$^-$ mainly include radical adduct formation (RAF) and formal hydrogen atom transfer (FHAT). Similarities and differences of mechanisms about two reactions were adequately investigated. The Gibbs free energy of reaction (Δ$G$) and Gibbs free energy barrier of activation (Δ$G^\ddagger$) of the initial reaction of AAP with •OH and SO$_4$•$^-$ were calculated and discussed. The binding distances and angles of AAP, •OH and SO$_4$•$^-$ are shown in Figure 1. All structures of transition states are plotted in Figure S1 (Supplementary Materials).

![Figure 1. The structures of AAP, •OH and SO$_4$•$^-$ calculated at the M06-2X/6-31 + G(d,p) level. The bond distance (Å) and angles (°) are listed. Here, ⋅ = H, ⋅ = C, ⋅ = O, ⋅ = S, ⋅ = N.](image-url)
3.1.1. Radical Adduct Formation

RAF pathways of AAP with •OH and SO₄⁻⁻ reactions are displayed in Figure 2. •OH-initiated and SO₄⁻⁻-initiated RAF channels consist of addition on the benzene ring and the acetamide group. It is uniform for the RAF mechanisms of AAP with •OH and SO₄⁻⁻ reactions. The acetamide group addition has no advantage over that of the benzene ring because the free energy barriers are 15.23 and 30.86 kcal/mol for acetamide group addition of •OH and SO₄⁻⁻, respectively. However, the free energy barriers were 5.95–9.26 kcal/mol and 2.66–8.74 kcal/mol for •OH and SO₄⁻⁻ addition to six carbon atoms of benzene ring, respectively. Generally, •OH-triggered reactions are higher exothermic than that of SO₄⁻⁻. Based on the values of ΔĜ and ΔG̃, C₆ addition (path 6 for •OH-triggered reactions versus path 13 for SO₄⁻⁻-triggered reactions) is the most favorable channels because their barriers are only 5.95 and 2.66 kcal/mol, respectively. Recently, the similar addition results were proved by Li et al. [47]. Figure 3 shows the comparison of potential energies for RAF mechanisms of two radicals reactions. SO₄⁻⁻-initiated reactions have the lower free energy barriers than that of •OH. In SO₄⁻⁻-initiated reactions, TS13, C₆ addition transition state, has stronger hydrogen bond intermolecular interaction, namely, hydrogen atom of phenolic hydroxyl group of AAP with oxygen atom of SO₄⁻⁻. IRC intuitively shows hydrogen atom of phenolic hydroxyl group of AAP tends to be close to oxygen atom of SO₄⁻⁻. The interaction will greatly decrease reaction barrier. However, SO₄⁻⁻-initiated reactions have less reaction heats compared with •OH-initiated reactions. For example, the energy barrier of path 6 is higher 3.29 kcal/mol than path 13, but path 6 is more exothermic than 6.95 kcal/mol.

Figure 2. The radical adduct formation channels of AAP with •OH and SO₄⁻⁻ reactions with the Gibbs free energy of reaction (ΔĜ) and Gibbs free energy barrier of activation (ΔG̃) (unit: kcal/mol).
Figure 2. The radical adduct formation channels of AAP with •OH and SO₄⁻⁻ reactions with the Gibbs free energy of reaction (ΔG) and Gibbs free energy barrier of activation (ΔG≠) (unit: kcal/mol).

Figure 3. The free energy diagram of RAF pathways initiated by two radicals.

3.1.2. Formal Hydrogen Atom Transfer

Ten hydrogen abstraction pathways from benzene ring and methyl group are found and shown in Figure 4. Hydrogen abstractions from C2, C3, and C5 of benzene ring experience TS15, TS16, TS17 with free energy barriers of 19.13, 18.75 and 19.75 kcal/mol for AAP with •OH reactions, respectively. For SO₄⁻⁻-initiated reactions, the free energy barriers of hydrogen abstractions from C2, C3, C5, and C6 of benzene ring are 30.58, 19.60, 19.30 and 30.43 kcal/mol, respectively. The hydrogen atom can be abstracted from C6 of benzene ring and the methyl group via 14.27 and 14.41 kcal/mol barriers for •OH-initiated path 18 and path 19. Moreover, the corresponding products are exothermic 7.02 and 11.37 kcal/mol, respectively. The results indicate that hydrogen abstractions from C6 of benzene ring and methyl group are two important channels for •OH with AAP reaction. However, methyl group hydrogen abstraction is the most important channel for SO₄⁻⁻ with AAP reaction due to the lowest energy barrier of 10.91 kcal/mol. Figure 5 describes the comparison of free energies for FHAT mechanisms. As shown in the Figure 5, the free energy barriers for SO₄⁻⁻ abstracting hydrogen atom from benzene ring are higher than that of •OH, and the corresponding paths (path 20–path 23) initiated by SO₄⁻⁻ are less exothermic than path 15–path 18 initiated by •OH. However, the free energy barrier of SO₄⁻⁻-initiated path 24 is lower 3.5 kcal/mol than •OH-initiated path 19, and path 24 is more exothermic 1.76 kcal/mol than path 19.

The comparison of FHAT and RAF mechanisms is shown in Figure 6. It is implied that RAF has an advantage over FHAT for both reactions. The free energy barrier for the most important RAF channel is lower 8.32 and 8.25 kcal/mol than the most favorable FHAT channel for •OH-initiated and SO₄⁻⁻-initiated reactions, respectively.

3.2. Kinetics

The rate constants involved free radicals reactions are of great value for predicting the degradation rate. However, the measurement of such data is difficult due to these rapid reactions. The theoretical calculations play an important role in attaining rate constants for these radical-participating reactions.
3.1.2. Formal Hydrogen Atom Transfer

Ten hydrogen abstraction pathways from benzene ring and methyl group are found and shown in Figure 4. Hydrogen abstractions from C2, C3, and C5 of benzene ring experience TS15, TS16, TS17 with free energy barriers of 19.13, 18.75 and 19.75 kcal/mol for •OH reactions, respectively. For SO4•−-initiated reactions, the free energy barriers of hydrogen abstractions from C2, C3, C5, and C6 of benzene ring are 30.58, 19.60, 19.30 and 30.43 kcal/mol, respectively. The hydrogen atom can be abstracted from C6 of benzene ring and the methyl group via 14.27 and 14.41 kcal/mol barriers for •OH-initiated path 18 and path 19. Moreover, the corresponding products are exothermic 7.02 and 11.37 kcal/mol, respectively. The results indicate that hydrogen abstractions from C6 of benzene ring and methyl group are two important channels for •OH with AAP reaction. However, methyl group hydrogen abstraction is the most important channel for SO4•− with AAP reaction due to the lowest energy barrier of 10.91 kcal/mol. Figure 5 describes the comparison of free energies for FHAT mechanisms. As shown in the Figure 5, the free energy barriers for SO4•− abstracting hydrogen atom from benzene ring are higher than that of •OH, and the corresponding paths (path 20–path 23) initiated by SO4•− are less exothermic than path 15–path 18 initiated by •OH. However, the free energy barrier of SO4•−-initiated path 24 is lower 3.5 kcal/mol than •OH-initiated path 19, and path 24 is more exothermic 1.76 kcal/mol than path 19.

The comparison of FHAT and RAF mechanisms is shown in Figure 6. It is implied that RAF has an advantage over FHAT for both reactions. The free energy barrier for the most important RAF channel is lower 8.32 and 8.25 kcal/mol than the most favorable FHAT channel for •OH-initiated and SO4•−-initiated reactions, respectively.
The rate constants for AAP with •OH reactions are given in Table 2. The apparent rate constant of •OH reaction with AAP ($k_{app}$) is $3.23 \times 10^9$ M$^{-1}$ s$^{-1}$ at 298 K. The calculated rate constant is consistent with experimental results of $(3.26 \pm 0.41) \times 10^9$ and $5.15 \times 10^9$ M$^{-1}$ s$^{-1}$ [18,19]. The C$_6$ site addition (path 6) has the largest rate constant of $3.56 \times 10^9$ M$^{-1}$ s$^{-1}$ with the 84.8% branching ratio. The other RAF and FHAT pathways contribute the total reactions weakly. The rate constants for AAP with SO$_4$•$^-$ reactions are depicted in Table 3. The apparent rate constant of SO$_4$•$^-$ reaction with AAP ($k'_{app}$) is $4.60 \times 10^{10}$ M$^{-1}$ s$^{-1}$ at 298 K, which is higher six times than experimental value of $7.66 \times 10^9$ M$^{-1}$ s$^{-1}$ [19]. The possible reason is that the lower barrier leads to higher reaction rate, which agrees with discussion of mechanisms. Theoretical model and method will lead to some deviations, but the accuracy of experiment is affected by some factors such
as equipment, reagent, and operation. Theoretical calculations can predict and explain some results. Consequently, the benefits of theoretical calculations cannot be underestimated. The C₆ site addition (path 13) is dominant channel with the largest rate constant of 8.65 × 10¹³ M⁻¹ s⁻¹ that possesses the 92.8% branching ratio. The other RAF and FHAT pathways have a little contribution for AAP with SO₄²⁻ reaction. As shown in Table 4, C₆ of benzene ring and methyl group hydrogen abstractions are dominant channels for •OH with AAP reaction with the branching ratio of 50.42% and 49.58%, respectively. For AAP with SO₄²⁻ reaction, methyl group hydrogen abstraction contributes 100% to FHAT channels.

Table 2. The calculated rate constants (kₐq), steady-state rate constant (kₐ), apparent rate constant (kₐp) and branching ratio (Rₐq) for the AAP with •OH reaction in the aqueous phase at 298 K.

| Paths                          | kₐq (M⁻¹ s⁻¹) | Rₐq (%) | kₐ (M⁻¹ s⁻¹) | kₐp (M⁻¹ s⁻¹) |
|-------------------------------|---------------|---------|---------------|---------------|
| APP + •OH → IM1 (k₁)          | 8.04 × 10⁷    | 1.9     | 9.80 × 10⁹    | 7.97 × 10⁷    |
| APP + •OH → IM2 (k₂)          | 1.87 × 10⁸    | 4.5     | 9.80 × 10⁹    | 1.83 × 10⁸    |
| APP + •OH → IM3 (k₃)          | 1.51 × 10⁷    | 0.4     | 9.80 × 10⁹    | 1.51 × 10⁷    |
| APP + •OH → IM4 (k₄)          | 3.22 × 10⁸    | 7.6     | 9.80 × 10⁹    | 3.12 × 10⁸    |
| APP + •OH → IM5 (k₅)          | 3.33 × 10⁷    | 0.8     | 9.80 × 10⁹    | 3.32 × 10⁷    |
| APP + •OH → IM6 (k₆)          | 3.56 × 10⁵    | 84.8    | 9.80 × 10⁹    | 2.61 × 10⁵    |
| APP + •OH → IM7 (k₇)          | 6.75 × 10⁵    | 0       | 9.80 × 10⁹    | 6.75 × 10⁵    |
| APP + •OH → IM15 (k₁₅)        | 4.75          | 0       | 9.80 × 10⁹    | 4.75          |
| APP + •OH → IM16 (k₁₆)        | 9.75          | 0       | 9.80 × 10⁹    | 9.75          |
| APP + •OH → IM17 (k₁₇)        | 1.91          | 0       | 9.80 × 10⁹    | 1.91          |
| APP + •OH → IM18 (k₁₈)        | 1.15 × 10⁴    | 0       | 9.80 × 10⁹    | 1.15 × 10⁴    |
| APP + •OH → IM19 (k₁₉)        | 1.13 × 10⁴    | 0       | 9.80 × 10⁹    | 1.13 × 10⁴    |
| APP + •OH → Product (k₉)      | 4.20 × 10⁵    | 100     |               | 3.23 × 10⁵    |

Table 3. The calculated rate constants (kₐq), steady-state rate constant (kₐ), apparent rate constant (kₐp) and branching ratio (Rₐq) for the AAP with SO₄²⁻ reaction in the aqueous phase at 298 K.

| Paths                          | kₐq (M⁻¹ s⁻¹) | Rₐq (%) | kₐ (M⁻¹ s⁻¹) | kₐp (M⁻¹ s⁻¹) |
|-------------------------------|---------------|---------|---------------|---------------|
| APP + SO₄²⁻ → IM8 (k₈)        | 6.00 × 10¹²   | 6.4     | 8.05 × 10⁹    | 8.04 × 10⁹    |
| APP + SO₄²⁻ → IM9 (k₉)        | 1.61 × 10¹¹   | 0.2     | 8.05 × 10⁹    | 7.67 × 10⁹    |
| APP + SO₄²⁻ → IM10 (k₁₀)      | 2.60 × 10¹¹   | 0.3     | 8.05 × 10⁹    | 7.81 × 10⁹    |
| APP + SO₄²⁻ → IM11 (k₁₁)      | 3.28 × 10¹⁰   | 0.01    | 8.05 × 10⁹    | 6.46 × 10⁹    |
| APP + SO₄²⁻ → IM12 (k₁₂)      | 2.52 × 10¹¹   | 0.3     | 8.05 × 10⁹    | 7.80 × 10⁹    |
| APP + SO₄²⁻ → IM13 (k₁₃)      | 8.65 × 10¹³   | 92.8    | 8.05 × 10⁹    | 8.05 × 10⁹    |
| APP + SO₄²⁻ → IM14 (k₁₄)      | 1.77 × 10⁻⁶   | 0       | 8.05 × 10⁹    | 1.77 × 10⁻⁶   |
| APP + SO₄²⁻ → IM20 (k₂₀)      | 14.3          | 0       | 8.05 × 10⁹    | 14.3          |
| APP + SO₄²⁻ → IM21 (k₂₁)      | 1.11 × 10²    | 0       | 8.05 × 10⁹    | 1.11 × 10²    |
| APP + SO₄²⁻ → IM22 (k₂₂)      | 1.88 × 10²    | 0       | 8.05 × 10⁹    | 1.88 × 10²    |
| APP + SO₄²⁻ → IM23 (k₂₃)      | 5.55          | 0       | 8.05 × 10⁹    | 5.55          |
| APP + SO₄²⁻ → IM24 (k₂₄)      | 1.33 × 10⁸    | 0       | 8.05 × 10⁹    | 1.33 × 10⁸    |
| APP + SO₄²⁻ → Product (k₉₅)   | 9.32 × 10¹³   | 100     |               | 4.60 × 10¹⁰   |

Table 4. The calculated rate constants (kₐq, kₐq) and branching ratio (Rₐq, Rₐq) for the formal hydrogen atom transfer channels in the aqueous phase at 298 K.

| Paths                          | kₐq (M⁻¹ s⁻¹) | Rₐq (%) | Paths                          | kₐq (M⁻¹ s⁻¹) | Rₐq (%) |
|-------------------------------|---------------|---------|-------------------------------|---------------|---------|
| APP + •OH (FHAT)              | 2.28 × 10⁴    | 100     | APP + SO₄²⁻ (FHAT)            | 1.33 × 10⁸    | 100     |
| APP + •OH → IM15 (k₁₅)        | 4.75          | 0       | APP + SO₄²⁻ → IM20 (k₂₀)      | 14.3          | 0       |
| APP + •OH → IM16 (k₁₆)        | 9.75          | 0       | APP + SO₄²⁻ → IM21 (k₂₁)      | 1.11 × 10²    | 0       |
| APP + •OH → IM17 (k₁₇)        | 1.91          | 0       | APP + SO₄²⁻ → IM22 (k₂₂)      | 1.88 × 10²    | 0       |
| APP + •OH → IM18 (k₁₈)        | 1.15 × 10⁴    | 50.42   | APP + SO₄²⁻ → IM23 (k₂₃)      | 5.55          | 0       |
| APP + •OH → IM19 (k₁₉)        | 1.13 × 10⁸    | 49.58   | APP + SO₄²⁻ → IM24 (k₂₄)      | 1.33 × 10⁸    | 100     |
The temperature dependence of rate constants is shown in Figure 7 at the temperatures from 198 to 338 K and 1 atm, and the corresponding data are listed in Tables S1 and S2 (Supplementary Materials). The total rate constants have weakly negative temperature dependence for •OH-initiated reaction. However, SO$_4^{•-}$-initiated reactions have distinctly negative temperature dependence.

![Figure 7. Temperature dependence of the calculated rate constants.](image)

3.3. The Aquatic Toxicities of AAP and Its Degradation Intermediates

The acute and chronic toxicities of AAP and the important degradation intermediates are assessed in three different aquatic organisms, which is drawn in Figure 8. Four types are classified and listed in Table S3 (Supplementary Materials). The toxic values of AAP and the important degradation intermediates are shown in Table S4 (Supplementary Materials).

3.3.1. Toxicity of AAP

The acute toxicity value of AAP is calculated as 323 mg/L of LC50 for fish, 63.1 mg/L of LC50 for daphnia and 26.3 mg/L of EC50 for green algae, respectively. These results indicate that AAP is harmful to daphnia and green algae, but not harmful to fish. The calculated ChV of AAP is 26.3 mg/L for fish, 5.13 mg/L for daphnia, and 37.2 mg/L for green algae. AAP is not harmful to fish and green algae at chronic level. However, it is harmful to daphnia chronically.

3.3.2. Toxicities of the Degradation Products

The most important intermediate (IM6) is harmful to three aquatic organisms in acute toxicity, but is harmless to three aquatic organisms in chronic toxicity. For other degradation intermediates, IM1 is acutely toxic for fish and green algae, and harmful to daphnia. The chronic toxicity of IM1 is harmful for three aquatic organisms. IM13 and IM8 are not harmful for three aquatic organisms chronically. Moreover, IM13 and IM8 are not acutely harmful for fish and daphnia, but pose a severe threat for green algae. In brief, the most important degradation intermediate (IM6) from •OH-initiated reaction is still harmful to aquatic organisms. IM13 from SO$_4^{•-}$-initiated reaction is harmless to fish and daphnia, but is very toxic to green algae. Thus, the toxicity of these compounds should be concerned.
3.3.1. Toxicity of AAP

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3.3.2. Toxicities of the Degradation Products

The most important intermediate (IM6) is harmful to three aquatic organisms in acute toxicity, but is harmless to three aquatic organisms in chronic toxicity. For other degradation intermediates, IM1 is acutely toxic for fish and green algae, and harmful to daphnia. The chronic toxicity of IM1 is harmful for three aquatic organisms. IM13 and IM8 are not harmful for three aquatic organisms chronically. Moreover, IM13 and IM8 are not acutely harmful for fish and daphnia, but pose a severe threat for green algae. In brief, the most important degradation intermediate (IM6) from •OH-initiated reaction is still harmful to aquatic organisms. IM13 from SO4•−-initiated reaction is harmless to fish and daphnia, but is very toxic to green algae. Thus, the toxicity of these compounds should be concerned.

4. Conclusions

In this work, the reaction mechanisms and rate of AAP with •OH and SO4•− have been explored theoretically in aqueous phase. The toxicity of AAP and its transformation intermediates to three aquatic organisms have been assessed. The novelty are summarized as below:

1. M06-2X/6-311++G (3df, 2p)//M06-2X/6-31+G (d, p) has been used to study the •OH-initiated and SO4•−-initiated transformation mechanism of AAP. •OH and SO4•− with AAP reactions have the same reaction sites, even reaction mechanisms. The results implied that the C6 addition is prominent pathway in RAF mechanisms and hydrogen abstraction of methyl group is dominant pathway for both reactions in FHAT mechanism. RAF takes precedence over FHAT.

2. At 298 K, the total apparent rate constant of AAP with SO4•− is larger than that of •OH. The calculated rate constants basically matched with experimental values. Theoretical calculations predicted the kinetic data at 198 K–338 K.

3. Toxic assessment shows that some representative degradation intermediates present an acute threat to the target organisms. Thus, subsequent degradation should be implemented until they are degraded into non-toxic substances.

In brief, this work explains the degradation processes of AAP initiated by •OH and SO4•− from microscopic points, and solves the problem of structures of intermediates and products which are associated with reactivity. The calculation of eco-toxicity plays an important role on assessing toxicity of degradation process. Finally, these results can apply to the practical degradation of AAP in AOPs.
Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/toxics9100234/s1, Figure S1: Optimized geometries involving the transition states of AAP with \( \cdot \mathrm{OH} \) and \( \mathrm{SO}_4^{\cdot-} \) at the M06-2X/6-31+G(d,p) level, Table S1: Calculated rate constants of AAP with \( \cdot \mathrm{OH} \) from 198 to 338 K and 1 atm, Table S2: Calculated rate constants of AAP with \( \mathrm{SO}_4^{\cdot-} \) from 198 to 338 K and 1 atm, Table S3: The acute and chronic toxicity class (mg L\(^{-1}\)), Table S4: Eco-toxicity values of AAP and its transformation intermediates to aquatic organisms (mg L\(^{-1}\))

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