The Aging Behavior of Polyvinyl Chloride Microplastics by UV/Sodium Percarbonate Oxidation: Efficiency and Mechanism

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Abstract: The aging behavior of microplastics (MPs) in the environment has garnered significant attention, yet the exact aging process undergone by microplastics during advanced oxidation still remains unclear. This study emphasized elucidating the oxidation and dechlorination mechanisms of polyvinyl chloride microplastics (PVC MPs) within the UV-activated sodium percarbonate system (UV/SPC). After 35 h of UV/SPC (0.05 M) treatment, it exhibited effective dechlorination with a Cl\(^{-}\) release of 100.62 ± 13.17 mg/L. And this treatment resulted in the fragmentation and release of fragments from the PVC MPs, thereby contributing to their further aging and dechlorination. In addition, an increase in oxygen-containing functional groups was identified. Quenching experiments confirm that \(\cdot\)OH and \(\cdot\)CO\(^3\)- are the primary active species present in the system. In summary, the aging of PVC MPs results in a process of fragmentation, releasing fragments that are more fragile and susceptible to mineralization compared to the original PVC MPs. The release of chloride ions originates from the mineralization of these fragments released from PVC MPs by \(\cdot\)OH and \(\cdot\)CO\(^3\). This study has provided valuable insights into the aging mechanisms of MPs in practical wastewater treatment.

Keywords: advanced oxidation processes; microplastics; UV/Sodium percarbonate; microplastics; aging behavior

1. Introduction

Over the past 50 years, global plastics production has skyrocketed, increasing from 1.7 million tons in 1950 to 335 million tons in 2016, a near-two-hundred-fold jump [1]. This rampant growth, however, has come with a cost: the fragmentation of discarded plastics in the environment results in the release of microplastics, varying in size from the nanometer scale up to 5 mm [2]. Microplastics are widely distributed and of various types detected in diverse environments, encompassing soil [3], oceans [4], rivers [5], bottled water [6], sediments [7], urban wastewater [8], and even the air we breathe [9]. Microplastics, which can adsorb harmful substances [10–12] and are toxic to organisms [13], have been found in various biological bodies [14–16], including humans [17]. Among these microplastics, polyvinyl chloride (PVC) stands out as one of the most commonly encountered types [18–20]. The aging of microplastics is divided into two categories: natural aging and artificial aging. Since the natural aging process of microplastics often takes an exceptionally long time, the artificial aging of microplastics has garnered considerable attention from scholars [21]. The aging of microplastics leads to changes in their surface properties, such as specific surface area [22,23] as well as in their physical characteristics, including crystallinity [24], ultimately affecting their behavior within the environment [25,26]. In the methods of artificial aging, advanced oxidation techniques (AOPs) excel due to their stability and performance across diverse conditions, coupled with lower operational complexities and costs compared to other methods [27].
Among the various radicals generated by advanced oxidation, hydroxyl radicals have proven effective in degrading microplastics [28]. Fenton oxidation, a classic technique utilizing hydroxyl radicals to remove pollutants, faces challenges due to the strict transportation and storage requirements of H₂O₂, resulting in high supply costs and limiting its practical application in wastewater treatment [29]. Sodium percarbonate (Na₂CO₃·1.5H₂O₂, SPC), known as solid hydrogen peroxide, has emerged as an ideal alternative due to its stability, low cost, convenience and safety to transport [30]. SPC can also be activated by transition metals like Fe(II)/Fe(III) to achieve a Fenton-like reaction in water treatment [31], but the presence of metal ions relies on acidic pH, leading to secondary contamination from metal precipitates [32]. However, UV-activated SPC (UV/SPC) has no such limitations. UV/SPC has been applied for the removal of a variety of pollutants, including aniline [33], natural organic matter (NOM) [34], tetracycline (TC) [35], and bisphenol A (BPA) [36], demonstrating significant potential for application in water treatment.

Since Na₂CO₃·1.5H₂O₂ is an adduct formed by the connection of Na₂CO₃ and H₂O₂ through hydrogen bonds, it easily decomposes and releases Na₂CO₃ and H₂O₂ when dissolved in water [37]. The generation of radicals in the UV/SPC system can be divided into two aspects: (1) the production of ·OH by the decomposition of SPC releasing H₂O₂ under UV irradiation; and (2) the reaction of ·OH with coexisting HCO₃⁻ and CO₃²⁻ to produce ·CO₃⁻. As shown in Equations (1)–(4), the production of ·CO₃⁻ consumes ·OH, the redox potential of ·OH (2.32 V vs. SHE at pH 7) is higher than that of ·CO₃⁻ (1.78 V vs. SHE at pH 7) [36], so the presence of HCO₃⁻ and CO₃²⁻ reduces the efficiency in degrading some pollutants such as aniline [33]. The presence of ·CO₃⁻, however, allows the degradation efficiency of certain pollutants to remain unchanged or even be enhanced, such as BPA and TC [35,36]. Gao et al. concluded that ·CO₃⁻ contributes significantly to the degradation and transformation of electron-rich organic pollutants, which can react with electron-rich compounds by electron transfer or hydrogen extraction [36]. Furthermore, the high steady-state concentration of ·CO₃⁻ can offset its limited reactivity towards target reactants and the consumption of ·OH [38], potentially enhancing the degradation rate of pollutants. In conclusion, ·CO₃⁻ has the potential to efficiently degrade organic pollutants. However, to date, there have been few studies investigating the mechanism of how microplastics (MPs) undergo aging under UV/SPC conditions.

\[
\begin{align*}
\text{H}_2\text{O}_2 & \overset{\text{UV}}{\rightarrow} 2\text{.OH} & (1) \\
\text{CO}_3^{2-} + \cdot\text{OH} & \rightarrow \cdot\text{CO}_3^- + \text{OH}^- & (2) \\
\text{HCO}_3^- + \text{H}_2\text{O}_2 & \rightarrow \text{HCO}_4^- + \text{H}_2\text{O} & (3) \\
\text{HCO}_4^- & \overset{\text{UV}}{\rightarrow} \cdot\text{OH} + \cdot\text{CO}_3^- & (4)
\end{align*}
\]

In this study, PVC was selected as the target MPs because its aging process releases chloride ions [38], providing a more intuitive demonstration of the aging extent. The main objectives of this study are as follows: (1) to investigate the dechlorination and morphological changes in PVC MPs during the reaction process; (2) to explore the active species that play a major role in the aging of PVC MPs; (3) to delve into the aging mechanism of PVC MPs in UV/SPC, and (4) to assess the aging effect of PVC MPs under UV/SPC treatment in practical wastewater treatment.

2. Materials and Methods

2.1. Materials

PVC MPs were purchased from Aladdin Co., Ltd. (Shanghai, China). Sodium percarbonate (Na₂CO₃·1.5H₂O₂, 13–14% active oxygen) was purchased from Macklin Co., Ltd. (Shanghai, China). Hydrogen peroxide (H₂O₂, ≥90%) was purchased from Kelong Co., Ltd. (Chengdu, China). Detailed information on PVC and handling before use is shown in Text S1. Other chemicals used in the experiment were included in Text S2.
2.2. Experimental Procedures

A low-pressure mercury lamp (with a primary emission wavelength of 254 nm, 10 W) was used as the irradiation source for the photosresponse experiments. A 200 mL aqueous solution, 0.5 g of PVC and the required amount of SPC were filled in a 250 mL tall beaker and irradiated. The beaker was wrapped in tinfoil to prevent UV from escaping. The system was set in a thermostatic water bath at 30 °C with sufficient agitation so that the PVC and SPC were evenly dispersed in the solution. Samples were withdrawn at specified intervals. The samples to be measured were filtered through a 0.45 µm membrane and the concentration of Cl⁻ released from PVC was subsequently measured by IC (IC, Dionex AQ-1100, Thermo Fisher Scientific, Waltham, MA, USA). Before testing, the treated PVC needed to be freeze-dried. All experimental procedures were standardized to ensure the accuracy of the results, including the effects of different SPC dosages and the control experiments using only SPC or UV. Error analysis was carried out in all experiments.

2.3. Analytical Methods

2.3.1. Characterization Methods

The Fourier Transform Infrared Spectrometer (FTIR, Nicolet 6700, Thermo Fisher Scientific, MA, USA) is utilized to analyze the functional group composition and changes in PVC MPs before and after reactions, with a detection wavelength ranging from 400 to 4000 cm⁻¹. A Scanning Electron Microscope (SEM, JSM-7500F, JEOL, Tokyo, Japan) is employed to investigate the surface morphological changes in the PVC MPs. Prior to analysis, the PVC MPs are coated with a conductive layer to enhance their surface conductivity and then scanned in secondary electron mode with an appropriate resolution of 5 µm at 5.0 KV. An X-ray diffractometer (XRD, EMPYREAN, PANalytical B.V., The Netherlands) is used to study the crystallinity changes in PVC MPs before and after reactions. The crystal structure images of the PVC MPs are obtained using the Cu target χ radiation mode. The scanning range is set from 5 to 60°, with a step size of 0.025°, and the scanning mode is set to continuous. An X-ray photoelectron spectrometer (XPS, ESCALAB 250Xi, Thermo Fisher Scientific, MA, USA) is employed to characterize the surface elemental content changes in PVC MPs before and after reactions. The full-spectrum power is set at 15 kV, 5 mA, while the sub-spectrum power is 15 kV, 10 mA. All spectra are calibrated using C 1s = 284.6 eV. The specific surface area of the PVC MPs is determined using a rapid surface area and pore size analyzer (ASAP2020, Micromeritics, Shanghai, China), employing the Brunauer–Emmett–Teller (BET) method. The final degassing temperature is set at 50 °C, and the ultimate vacuum value is 50 µm Hg. Gas chromatography-mass spectrometry (GC-MS, TSQ9000, Thermo Fisher Scientific, MA, USA) was used to detect the reaction by-products, and the filtered reaction solution was first subjected to solid phase extraction before detection.

2.3.2. Release of Chloride Ions

The Ritger and Peppas model is employed to investigate the release mechanism of Cl⁻ and fit the release curve:

\[ \frac{M_t}{M_\infty} = kt^n \]  

Generally, where \( M_t \) and \( M_\infty \) are the amount of Cl⁻ released at time t and equilibrium, respectively, k is the release rate constant, and n is the release index indicating the nature of the release mechanism. In this study, \( M_\infty \) represents the release of Cl⁻ at the end (35 h) of the reaction.

The Ritger and Peppas model was originally used to describe diffusion mechanisms [39] and now mostly used to describe drug release mechanisms. Due to the special skeleton structure of PVC microplastics and the different forms of dechlorination and release in the UV/SPC system, which were similar to certain photosensitizing drug carriers for drug release. Therefore, the Ritger and Peppas model was used to describe the mechanism of Cl⁻ release from PVC MPs in the photocatalytic system.
When \( n < 0.45 \), the chloride ion release mechanism is Fick diffusion, when \( 0.45 < n < 0.89 \), the chloride ion release mechanism is non-Fick diffusion and when \( n > 0.89 \), the chloride ion release mechanism is skeleton erosion [40]. The detailed meaning is mentioned later.

### 2.3.3. Carbonyl Index

The FTIR spectra obtained for PVC MPs samples were used to estimate the Carbonyl Index (CI), since the FTIR technique allows for the measurement of several by-products containing carbonyl functional groups formed on the particle surface as a result of the oxidation of the material [41]. The CI is determined by calculating the ratio of the carbonyl peak to the reference peak [42]. Therefore, this work used CI as a measure of PVC oxidation, which enabled the quantification of several modifications to the material’s chemical structure. The PVC MPs selected in this study were easily identified as having a newly generated absorption peak at wave number 1710–1730 cm\(^{-1}\) after processing. The intensity levels observed varied depending on the aging process and exposure time as a result of the C=O stretching of the carbonyl group generated by the reaction, which carries a strong and characteristic infrared absorption peak at 1715 cm\(^{-1}\) [43].

According to Mariana et al.’s statistics [41], there is no universal method for calculating CI. Some researchers have chosen to calculate the CI as the ratio of the absorption areas of the two peaks or the maximum absorption heights of the two peaks. In addition, the selection of reference peaks for each microplastic varied in different studies, and researchers also used different methods based on the transmittance or absorbance patterns of the FTIR spectra. In our study, the FTIR measurements were performed in transmission mode, and the maximum height of the selected peaks was transformed from transmittance (T) to absorbance (A) by Equation (6). After screening various peaks as reference peaks, we selected the peak formed by the symmetric stretching vibration of the methylene (-CH\(_2\)-) group, which exhibited little variation under all the different system treatments mentioned in this study, as the reference peak for PVC MPs. The CI was calculated as shown in Equation (7).

\[
A = \log_{10} \frac{1}{T(\text{maximum height})} \quad (6)
\]

\[
CI = \frac{A_{\text{carbonyl group}(1710–1730)}}{A_{\text{methylene group}(2910 \text{ for PVC})}} \quad (7)
\]

### 3. Results and Discussion

#### 3.1. The Dechlorination of PVC

To explore the aging effect of PVC MPs in UV/SPC systems and to validate the synergistic effects of UV and SPC, we conducted a controlled variable study, investigating the dechlorination effects of PVC in systems where UV or SPC was applied individually. As shown in Figure 1a, the release of Cl\(^-\) from PVC MPs in the system with SPC alone proceeded at a slow pace throughout the entire treatment process. After 35 h of treatment, the concentration of Cl\(^-\) released into the solution was 15.16 ± 1.81 mg/L. The Cl\(^-\) release from PVC MPs in the system with UV irradiation alone reached 54.05 ± 0.52 mg/L. This finding is consistent with the research conducted by Ouyang et al., who demonstrated that UV can age PVC MPs in water and promote the release of Cl\(^-\) [44]. After the same treatment time, the chlorine release in the UV/SPC system reached 100.62 ± 13.17 mg/L, which was 1.86 times higher than that in the UV-alone system, highlighting the synergistic effect of combining UV with SPC. The enhanced dechlorination effect can be attributed to the oxidative radicals, such as \( \cdot \text{OH} \) and \( \cdot \text{CO}_3^- \) generated by the activation of SPC by UV (Equations (1)–(4)). These oxidative radicals effectively oxidize and age microplastics [30], and Cl\(^-\) is released during this process [38]. The dechlorination ability of other photocatalytic, biodegradation and hydrothermal carbonization (HTC) technologies for PVC MPs was summarized as shown in Table S1, and it can be concluded that the dechlorination ability of UV/SPC is higher than that of UV/PS [44] and photocatalytic LMWOA-Fe(III) [38]. Biodegradation is inherently different from advanced oxidation technology in terms of
degradation efficiency [45], while HTC has a good dechlorination efficiency in a shorter period of time but consumes additional energy [46].

\[
\cdot \text{OH} + \cdot \text{OH} \rightarrow 0.5 \text{O}_2 + \text{H}_2\text{O} \quad (8)
\]

\[
\cdot \text{CO}_3^- + \cdot \text{CO}_3^- \rightarrow \text{O}_2 + \text{C}_2\text{O}_4^{2-} \quad (9)
\]

Figure 1. The concentration of released \( \text{Cl}^- \) in PVC MPs is compared between (a) UV and SPC and (b) Different concentrations of SPC and UV only. Conditions: [PVC] = 2.5 g/L.

In addition, experiments were conducted to investigate the initial dosage of SPC. In the UV/SPC system, SPC is irradiated by UV to produce a large number of oxidative radicals. Therefore, the initial dosage is an important factor that affects the aging and dechlorination of PVC MPs, necessitating further investigation. As shown in Figure 1b, the dechlorination effect of PVC MPs increases with an increase in the initial dosage of SPC. However, SPC contains a small amount of NaCl impurities, and the chlorine ions released from NaCl can interfere with the measurements, leading to increased errors. The larger the initial dosage of SPC, the greater the error. Therefore, the dosage of SPC was limited to 0.05 M. With the increase in initial SPC concentration (0.005–0.05 M), the release of chlorine ions from PVC rapidly rose from 63.00 ± 0.26 mg/L to 100.62 ± 13.17 mg/L under UV irradiation for 35 h. As evident from the graph, the dechlorination rate was not constant, it accelerated as the treatment time progressed. Additionally, the timing of this acceleration varied across systems with different initial concentrations. Specifically, a significant inflection point was observed at 15 h for the system with an initial concentration of 0.005 M, while the inflection point occurred at 25 h for the 0.05 M system. This phenomenon could be attributed to the self-quenching of high concentrations of radicals. When SPC dissolves in water, it generates a significant amount of \( \cdot \text{OH} \) and \( \cdot \text{CO}_3^- \). At high concentrations, these reactive species compete with pollutants for reactions (Equations (8) and (9)) [36], thereby affecting the efficiency of oxidation and dechlorination. In conclusion, the dechlorination of PVC MPs increased with the initial SPC dosage in the range of 0–0.05 M dosage.

The Ritger Peppas model was used to fit the nonlinearities of the UV alone and UV/SPC systems, and the fitting results are shown in Figure S1. Based on the analysis of the fitted data, the UV and UV/SPC models both presented patterns of skeleton erosion, which means the destruction of the PVC MPs’ structure and carbon chain, a phenomenon that occurs during the oxidation of MPs [30]. However, oxidizing radicals are weakly dechlorinating [47], and the release of \( \text{Cl}^- \) is often accompanied by a high degree of oxidation, suggesting that \( \text{Cl}^- \) are released during the mineralization of PVC MPs by oxidizing radicals. This result may also explain the accelerated release of \( \text{Cl}^- \) in the UV/SPC system. Oxidation is a cumulative process, and researchers have found that MPs produce fragile intermediate products that detach from their skeletons during oxidation [48].
The fragile intermediate is derived from the main chain breakage of PVC MPs, which are essentially oxidized chlorinated hydrocarbons. Because they are fragile, they are more susceptible to mineralization and Cl\(^-\) is released during this process. With increasing treatment time, shedding intermediates were accumulated, so the rate of Cl\(^-\) release became faster.

3.2. Morphology Characterization of PVC

3.2.1. Morphologies

SEM was used to observe the change in surface morphology of PVC MPs before and after treatment. Figure 2a shows the initial sample without UV/SPC treatment, and it can be observed that the surface is relatively intact and dense with only some small pits. And as shown in a–f, after 35 h of UV/SPC treatment with different initial dosages, varying sizes of holes or cracks appeared on the surfaces, which would expose the interior of PVC MPs to the attack of free radicals and favor further aging. UV easily induces photochemical reactions in the chromophores of PVC MPs [49] and causes particle surface aging [50]. While UV aging of PVC MPs accelerates the aging process in the presence of oxidants, Dong et al. showed that direct photolysis of UV dominated microplastics at low concentrations of oxidants or in the early stages of aging and that as the concentration of oxidants increased or as the degree of aging increased, indirect photolysis gradually dominated [51]. From the SEM images, it can be clearly observed that when the initial SPC dosage was 50 mM, the surface of PVC MPs was severely damaged and the internal structure was completely exposed, which indicates that UV/SPC can effectively age PVC MPs. In addition, it can be observed that the surface of the treated PVC MPs has generated some fragments that seem to be ready to peel off at any time, and it can be assumed that it is the peeling of these fragments that causes holes or cracks to appear on the surface. As the fragments gradually peel off, the holes or cracks expand in size, until the internal structure is completely exposed, thus resulting in a new round of peeling. As mentioned earlier, these flaked fragments continue to be attacked by radicals, and in the process, chloride ions are released. It is worth noting that after the addition of a sufficient amount of PhOH to quench the major radicals in the system, the surface of PVC MPs was still damaged, and the extent of the damage was similar to that of the initial dosage of 5 mM, which may be caused by the presence of other minor active species in the system, like O\(^{•-}\) or the direct photolysis by UV. Due to its low redox potential (E\(_{0}\) = -0.33 V), which contributes less to the direct degradation of the target pollutant, O\(^{•-}\) is often ignored [52]. However, when the main active substance in the system is quenched, the role of O\(^{•-}\) may be revealed.

The appearance of surface holes and cracks is often accompanied by an increase in the specific surface area, and for the further study of the morphological changes in PVC MPs, the Brunauer–Emmett–Teller (BET) method was used to characterize the changes in the specific surface area before and after the treatment. Specific data are shown in Table S2. The specific surface area of the treated PVC MPs is 1.68 times higher than that before the treatment, and there is also a substantial increase in the specific surface area after quenching experiments, which supports the above conjecture.

To further investigate the microstructural changes in PVC MPs before and after the treatment, XRD was used to analyze the changes in crystallinity (Figure 3). Ouyang et al. found that two peaks appear between 10° and 30° for PVC MPs, and the peaks become sharp, probably due to the change in the structure of PVC MPs itself and the destruction of C-Cl bonds, as the crystallinity of PVC MPs increases [44]. In agreement with the research, it can be observed that the peak near 20° becomes sharp after UV/SPC treatment. In addition, the increase in crystallinity means that the plastic becomes more fragile and thus more prone to producing smaller particles [53], which coincides with the appearance of fragments on the surface of the treated PVC MPs observed in the SEM images.
Figure 2. SEM of (a) pristine and (b–f) treated PVC MPs. (b–e) 5 mM, 10 mM, 20 mM and 50 mM SPC with UV, (f) 50 mM SPC + 50 mM PhOH with UV.

Figure 3. XRD image of PVC MPs before and after PVC MPs treatment. Conditions: [PVC] = 2.5 g/L, [SPC] = 50 mM.

3.2.2. Functional Groups

In order to explore the oxidation mechanism and further investigate the dechlorination mechanism of PVC MPs in the UV/SPC system, the functional groups on the surface of PVC MPs were analyzed by FTIR. As shown in Figure 4a, there are two characteristic peaks associated with chlorine on the infrared spectrum of PVC MPs, which appear at 605 cm\(^{-1}\) and 1250 cm\(^{-1}\), respectively, and they are generated by the stretching vibrations of the C-Cl group and the CH-Cl group [44,54]. Another characteristic peak appearing at 2909 cm\(^{-1}\) represents the stretching vibrational signal of methylene (-CH\(_2\)) [54]. A new characteristic peak at 1715 cm\(^{-1}\) emerged on the spectrum of UV/SPC-treated PVC MPs, attributable...
to the stretching vibration of the C=O group [43], and the corresponding carbonyl index in Figure 4b demonstrates a significant increase for all treated PVC MPs, indicating that the active species present in the system can oxidize PVC MPs. It can be clearly observed that the signal intensity belonging to the C=O group is the strongest when the initial SPC dosage is 50 mM, and the concentration of chloride ions released from PVC MPs is also the highest at the same condition, which suggests that there exists a relationship between the concentration of released Cl\(^{-}\) and the degree of oxidation. However, the characteristic peak signals of CH-Cl and C-Cl groups related to chlorine both showed different degrees of enhancement after treatment, which may be related to the inability of oxidizing radicals to attack the C-Cl bond directly. Evidence for this speculation is the very low contribution of oxidizing radicals to dechlorination found by Miao et al., who found 1% dechlorination of PVC MPs in an anode chamber where only oxidizing radicals were present (8–10% in a cathode chamber) [47]. The enhanced signals representing chlorine are due to the increased specific surface area of the treated PVC MPs, resulting in more chlorine-containing groups being detected. In conclusion, the enhancement of the signals at 1250 cm\(^{-1}\) and 605 cm\(^{-1}\) illustrates the inability of oxidizing radicals to release Cl\(^{-}\) by selectively breaking the C-Cl bond.

Figure 4. (a) The FTIR spectra and (b) CI values of pristine and treated PVC MPs with different SPC initial dosages.

3.2.3. Chemical Composition and Intermediates of PVC

As shown in Figure 5, the XPS analysis exhibits the changes in surface elemental content and functional groups before and after the treatment of PVC MPs. Figure 5a shows the XPS total spectrum; the total amount of carbon on the surface of PVC MPs after UV/SPC treatment decreased. Combined with Figure 2e, it can be concluded that the main reason is that the MPs surface was severely damaged by UV/SPC at the initial dosage of 50 mM to form oversized voids and lose a large number of carbon chains, which was also seen in other studies [30]. And there may be two reasons for the decrease in total oxygen content: the first is also because the surface of PVC MPs is severely damaged to expose the internal surface that has not yet been oxidized, and the second is that on the surface of PVC MPs in the oxidation process, there will be a lot of fragments containing oxidation intermediates flaking, and these fragments of flaking take away part of the oxygen element. The high amount of oxygen on the surface of the initial PVC MPs may indicate that PVC is also easily oxidized in the environment. A similar observation has been made in other studies, where the oxygen content of untreated PVC reached 10.42% [44]. But in combination with Figure 2a, it can be found that the slight degree of oxidization remains only on the surface. In addition, the fluorine that appeared on the surface after the treatment came from the PTFE stirrer, and it is possible that the surface of the stirrer was broken, resulting in debris mixing into the PVC MPs sample.
As shown in Figure 5b, the XPS spectrum of C 1s is divided into four peaks, including C=C/C-C/C-H, C-Cl, C-O, and C=O, with binding energies centered at 284.6, 285.7, 286.1, and 288.2 eV, respectively [55,56]. The specific proportion of each peak is shown in Table S3, which shows that the proportion of the peaks representing C-Cl after treatment did not change significantly, which further indicates that the active species within the UV/SPC system cannot directly break the C-Cl bond to release Cl\(^-\). As shown in Figure 5c, the XPS spectrum of O 1s is divided into two peaks, including C=O (532.6 eV) and C-O (531.7 eV) [55], and the percentage of the C=O peak is slightly increased after the treatment, but it is not much different from the original PVC MPs, for reasons that are consistent with the total spectrum. In addition, the XPS spectrum of Cl 2p (Figure 5d) is decomposed into two peaks at 201.0 and 199.4 eV [55], which belong to Cl 2p\(_{3/2}\) and Cl 2p\(_{1/2}\), respectively. The reason for the decrease in the total amount of chlorine after the treatment is in line with the reason for the decrease in the total amount of carbon.

In addition, the filter solution of UV/SPC-treated PVC MPs was analyzed by GC-MS. As shown in Table S4, seven major products were detected, including ketone and ester organics. These products originate from oxidative processes with oxygenated functional groups (-C=O, -COOH). They detach from the surface of the MPs and exist in solution, which corresponds to taking away the elemental oxygen from the surface, which is consistent with the XPS total spectrum.

### 3.3. Oxidation and Dechlorination Mechanism of UV/SPC System for PVC MPs

#### 3.3.1. Reactive Species Contribution to Dechlorination

According to the relevant studies, the main reactive species present in the UV/SPC system are -CO\(_3^-\) and ·OH [30,36,57], and in order to further determine the contribution of the different radicals for dechlorination, ethanol (EtOH) was used for -OH scavenging...
(\(k_{\text{EtOH-HO}} = 2.8 \times 10^9 \text{ M}^{-1}\text{s}^{-1}\)), and phenol (PhOH) was used for quenching of \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) (\(k_{\text{PhOH-HO}} = 2.8 \times 10^9 \text{ M}^{-1}\text{s}^{-1}\), \(k_{\text{PhOH-CO}_3}\) = \(1.4 \times 10^{10} \text{ M}^{-1}\text{s}^{-1}\)) [36]. As shown in Figure 6, the addition of 75 mM EtOH resulted in a 38.47\% decrease in chloride release from 100.62 ± 13.17 mg/L to 61.91 ± 15.23 mg/L in the UV/SPC system. The addition of 50 mM PhOH led to a 93.10\% decrease in chloride release to 6.94 ± 0.51 mg/L, indicating that \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) are the main active species in this system. From the results of the quenching experiments, it can be concluded that the contribution of \(\cdot\text{CO}_3\) to the dechlorination of PVC MPs in the UV/SPC systems is greater than that of \(\cdot\text{OH}\). However, this result does not mean that the dechlorination capacity of \(\cdot\text{CO}_3\) is higher than that of \(\cdot\text{OH}\), because the generation of \(\cdot\text{CO}_3\) consumes part of \(\cdot\text{OH}\) (Equations (2)–(4)). The formation of \(\cdot\text{CO}_3\) in the UV/SPC system occurs in two ways. The first one is to consume \(\cdot\text{OH}\) directly to produce the same amount of \(\cdot\text{CO}_3\) (Equation (2)), and the second one is to react with H\(_2\)O\(_2\) to form an intermediate, and the intermediate will continue to produce the same amount of \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) (Equations (3) and (4)), and the original H\(_2\)O\(_2\) activated by the UV can produce twice the amount of \(\cdot\text{OH}\) as its own (Equation (1)), so the production of \(\cdot\text{CO}_3\) can be theoretically regarded as the consumption of the same amount of \(\cdot\text{OH}\).

![Figure 6. Concentration of released Cl\(^-\) of PVC MPs in quenching experiment. Conditions: [PVC] = 2.5 g/L, [SPC] = 50 mM.](image)

In order to further investigate the dechlorination ability of \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\), experiments with H\(_2\)O\(_2\) and Na\(_2\)CO\(_3\) instead of SPC were designed, and the results of the experiments are shown in Figure 7a. According to the experimental results, the releases of Cl\(^-\) in the two systems of UV/H\(_2\)O\(_2\) and UV/H\(_2\)O\(_2\)/Na\(_2\)CO\(_3\) were 72.86 ± 18.85 mg/L and 75.60 ± 2.07 mg/L, respectively, with almost no difference, which indicated that the dechlorination ability of the active species in the two systems was not much different, and the main active species in the UV/H\(_2\)O\(_2\) system was \(\cdot\text{OH}\), while both \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) existed in the UV/H\(_2\)O\(_2\)/Na\(_2\)CO\(_3\) system. However, \(\cdot\text{CO}_3\) was generated by consuming an equal amount of \(\cdot\text{OH}\) in the UV/H\(_2\)O\(_2\)/Na\(_2\)CO\(_3\) system, but the fitted reaction rate constants \(a\), release constants \(n\), and the Ritger and Peppas model curves were similar (Figure 7b), which proved that the dechlorination ability of \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) was comparable for PVC MPs. While both UV/H\(_2\)O\(_2\) and UV/H\(_2\)O\(_2\)/Na\(_2\)CO\(_3\) systems were not as effective as the UV/SPC system for dechlorination, this could be due to the excessive initial dosage of oxidant leading to the self-quenching of \(\cdot\text{CO}_3\) and \(\cdot\text{OH}\) (Equations (8) and (9)) [36]. However, the H\(_2\)O\(_2\) in SPC is required to be produced by SPC decomposition, which may contribute to the slow release of H\(_2\)O\(_2\), especially advantageous in degrading PVC MPs, which is a long-term treatment process.
This discovery holds significant value for the application of the UV/SPC system in the aging of PVC MPs, as the removal efficiency of $\cdot$CO$_3^-$ does not always reach the same level as that of $\cdot$OH for all pollutants. For instance, when degrading aniline [33], the reaction rate constant of the UV/$\text{H}_2\text{O}_2$ system is nearly twice as high as that of the UV/SPC system. However, some pollutants, $\cdot$CO$_3^-$ showed a notable oxidizing ability. For example, in the study of treating TC, the removal efficiency of the UV/SPC system reached 89.1% under the same treatment conditions, while the removal efficiency of UV/$\text{H}_2\text{O}_2$ for TC was only 45.1%, which was significantly lower than that of the UV/SPC system [35]. When removing BPA, the removal efficiency of UV/$\text{H}_2\text{O}_2$, UV/SPC and UV/$\text{H}_2\text{O}_2$/Na$_2$CO$_3$ for BPA was basically the same under the same conditions [36]. In the removal of PS MPs, quenching experiments revealed that $\cdot$CO$_3^-$ plays a primary role while $\cdot$OH plays a secondary role [30]. However, unlike PVC MPs, TC, BPA, and PS MPs all have a benzene ring and are electron-donating groups with the benzene ring, making them electron-rich pollutants, which is consistent with the electrophilic nature of $\cdot$CO$_3^-$. The high capacity exhibited by $\cdot$CO$_3^-$ for PVC MPs aging can be attributed to the higher steady-state concentration of $\cdot$CO$_3^-$ compared to $\cdot$OH, which needs to be further explored.

In addition, in a study that also investigated the aging of PVC MPs by UV-catalyzed oxidants [4], the dechlorination efficiency of UV/PS for PVC MPs was lower than that of ours, and the major radicals in their system were SO$_4^-$ and $\cdot$OH, whereas the major radicals in our system, UV/SPC, were $\cdot$OH and $\cdot$CO$_3^-$, which indicates that $\cdot$CO$_3^-$ has better performance for PVC dechlorination than SO$_4^-$. 

### 3.3.2. Reactive Species Contribution to Oxidation

To further investigate the ability of $\cdot$CO$_3^-$ and $\cdot$OH to oxidize PVC MPs, samples of PVC MPs treated by quenching experiments were taken to analyze the functional groups by FTIR. As shown in Figure 8a, observing the C=O group representing the degree of oxidation in the FTIR band, it was found that the intensity of the peak at 1715 cm$^{-1}$ of the PVC MPs samples produced by the UV/SPC system with the addition of PhOH did not change significantly, while the intensity with the addition of EtOH increased significantly, this indicates that the quenching of $\cdot$CO$_3^-$ and $\cdot$OH by PhOH completely inhibited the generation of oxidizing groups on the surface of PVC MPs, which means the oxidation by the UV/SPC system was inhibited, whereas the quenching of $\cdot$OH by EtOH still generated some oxidizing groups on the surface of PVC MPs, which proves that even though $\cdot$OH was quenched, $\cdot$CO$_3^-$ was still able to oxidize the PVC MPs in the UV/SPC system. Combined with Figure 8b, the CI of PVC MPs samples produced by the UV/SPC system with the
addition of EtOH is 0.42, and the CI value of the UV/SPC system is 0.49, which suggests that the \( \cdot \text{CO}_3 \) can oxidize the PVC MPs efficiently and plays a major role in the UV/SPC system.

![Figure 8. (a) FTIR spectrum and (b) CI value of PVC MPs before and after quenching treatment. Conditions: [PVC] = 2.5 g/L, [SPC] = 50 mM, [EtOH] = 75 mM, [PhOH] = 50 mM.](image)

However, the FTIR spectra of the PVC MPs samples produced by the UV/SPC system with PhOH showed the same characteristics as the other oxidized PVC MPs samples, which means the intensity of the peak representing the CH-Cl group at 1250 cm\(^{-1}\) was significantly enhanced compared to the pristine PVC MPs sample. This is also caused by the increased specific surface area (Table S2). In addition, the reason for the decreased CI value may be due to the previously mentioned \( \text{O}_2^- \) (\( E_0 = -0.33 \text{ V} \)), because it is not an oxidizing radical, it breaks the chain on the microplastic surface, taking some of the oxygen with it. This explains both the surface destruction and aging (Figure 2f and Table S2) of the microplastics and why the CI decreases instead. Although the surface of PVC MPs had been damaged and the specific surface area increased accordingly, only 6.94 ± 0.51 mg/L of CI\(^-\) were released, suggesting that the dechlorination was asynchronous with the oxidation of PVC MPs and that dechlorination was accompanied by a higher degree of aging.

3.3.3. Oxidation and Dechlorination Mechanism of PVC MPs in UV/SPC

Based on the dechlorination effect, the characterization results and the radical quenching experiments, the oxidation and dechlorination mechanisms of PVC MPs in the UV/SPC system were proposed. Dechlorination and oxidation of PVC MPs do not occur together; in the early stages of oxidation, direct photolysis and \( \cdot \text{CO}_3 \) and \( \cdot \text{OH} \) from UV/SPC oxidize PVC MPs, generating oxidation intermediates (fragments) that can be detached from the polyvinyl chloride surface. These intermediates are formed when active species break down the chains on the surface of PVC MPs, whose chains originally contained a large amount of chlorine. With the increase in reaction time, the oxidation intermediates are gradually mineralized; in this process, the CI\(^-\) are released and in the whole process, the main role is played by \( \cdot \text{CO}_3 \), and \( \cdot \text{OH} \) plays an auxiliary role.

3.4. Application of UV/SPC in Actual Water Bodies

The aging performance of UV/SPC for PVC MPs in two real water bodies was investigated. The river water samples were taken from Jiangan River (River J, Chengdu, China), and the secondary effluent of the sewage plant (SE) was taken from a sewage treatment plant in Shuangliu, Chengdu, China. The indicators of the two actual water bodies are shown in Table 1. As shown in Figure 9, the dechlorination effect of PVC MPs was inhibited in both actual water bodies. Similarly, when Gao et al. applied UV/SPC for BPA removal in secondary effluent from a wastewater plant, the same inhibition was observed. The explanation was due to the scavenging effect of complexes contained in the effluent,
which led to a significant reduction in the concentration of radicals and turbidity, thereby decreasing the UV transmittance [36]. Additionally, it is also possible that coexisting anions and organic pollutants in the water body are competing with the PVC MPs for the reaction, depleting ·OH and subsequently hampering the generation of ·CO$_3$ $^-$ [58]. Thus, it seems that UC/SPC may be more suitable for the treatment of reuse water [36]. Currently, there are scant reports on advanced oxidation techniques effectively removing microplastics from actual water bodies, a domain that deserves further exploration and research.

Table 1. Various indicators of actual water bodies.

<table>
<thead>
<tr>
<th>Original pH</th>
<th>UV-Vis Absorbance @ 254 nm</th>
<th>Cl$^-$ (mg/L)</th>
<th>NO$_3^-$ (mg/L)</th>
<th>SO$_4^{2-}$ (mg/L)</th>
<th>PO$_4^{3-}$ (mg/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>River J</td>
<td>8.56</td>
<td>0.315</td>
<td>107.38</td>
<td>11.70</td>
<td>64.85</td>
</tr>
<tr>
<td>SE</td>
<td>7.12</td>
<td>0.061</td>
<td>22.64</td>
<td>2.27</td>
<td>58.27</td>
</tr>
</tbody>
</table>

Figure 9. Dechlorination effect of PVC MPs in different water qualities. Conditions: [PVC] = 2.5 g/L, [SPC] = 50 mM.

4. Conclusions

In this study, the aging and dechlorination mechanisms of PVC MPs in UV/SPC were investigated. The aging and dechlorination of PVC MPs increased with the increase in SPC dosage when the initial dosage of SPC was 0.005–0.05 M and the maximum chloride release concentration was 100.62 ± 13.17 mg/L in 35 h. The characterization results revealed that the surface of treated PVC MPs exhibited fragments, and the peeling off of the fragments resulted in the formation of cracks and holes, accompanied by an augmentation in the specific surface area. This confirmed that the aging of PVC MPs involved a fragmentation process, releasing fragile fragments that were more susceptible to mineralization compared to the original PVC MPs. And the release of chloride originates from the mineralization of fragments by ·CO$_3$ $^-$ and ·OH. Additionally, this study has provided valuable insights into the aging mechanisms of MPs in practical wastewater treatment, yet further research is still needed to deepen our understanding, such as by experimenting with a wider range of MPs samples.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/w16111529/s1, Text S1: The information of PVC. Text S2: Materials and reagents. Figure S1: Release kinetic model of PVC MPs in each system. A: only UV; B: UV+5 mM SPC; C: UV+10 mM SPC; D: UV+20 mM SPC; E: UV+50 mM SPC Conditions: [PVC] = 2.5 g/L. Table S1: Comparison of dechlorination efficiency of various PVC MPs aging technologies. Table S2: Specific surface area of PVC MPs before and after reaction. Table S3: The area proportion of each peak in the XPS spectrum. Table S4: The major products of PVC MPs under UV-activated SPC treatment.
Author Contributions: Conceptualization, Z.L., Y.W. and Q.W.; Methodology, L.S.; Investigation, L.S.; Resources, Z.L., Y.W. and Q.W.; Data curation, J.H. and L.S.; Writing—original draft, L.S.; Supervision, Q.W.; Funding acquisition, Q.W. All authors have read and agreed to the published version of the manuscript.

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