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Dichloramine Hydrolysis in Membrane Desalination Permeate: Mechanistic Insights and Implications for Oxidative Capacity in **Potable Reuse Applications**

Liang Wu, Sitao Liu, and Haizhou Liu*

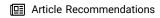


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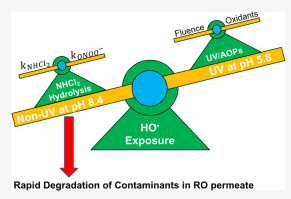
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ABSTRACT: Dichloramine (NHCl₂) naturally exists in reverse osmosis (RO) permeate due to its application as an antifouling chemical in membrane-based potable reuse treatment. This study investigated mechanisms of background NHCl2 hydrolysis associated with the generation of oxidative radical species in RO permeate, established a kinetic model to predict the oxidative capacity, and examined its removal efficiency on trace organic contaminants in potable reuse. Results showed that NHCl₂ hydrolysis generated transient peroxynitrite (ONOO⁻) and subsequently dissociated into hydroxyl radical (HO*). The maximal HO* exposure was observed at an RO permeate pH of 8.4, higher than that from typical ultraviolet (UV)-based advanced oxidation processes. The HO^o exposure during NHCl₂ hydrolysis also peaked at a NH₂Cl-to-NHCl₂ molar ratio of 1:1. The oxidative capacity rapidly degraded 1,4-dioxane,



carbamazepine, atenolol, and sulfamethoxazole in RO permeate. Furthermore, background elevated carbonate in fresh RO permeate can convert HO[•] to carbonate radical (CO₃•-). Aeration of the RO permeate removed total carbonate, significantly increased HO exposure, and enhanced the degradation kinetics of trace organic contaminants. The kinetic model of NHCl2 hydrolysis predicted well the degradation of contaminants in RO permeate. This study provides new mechanistic insights into NHCl₂ hydrolysis that contributes to the oxidative degradation of trace organic contaminants in potable reuse systems.

KEYWORDS: dichloramine, hydrolysis, peroxynitrite, HO[•] exposure, 1,4-dioxane, potable reuse

INTRODUCTION

Growing population, expanding urbanization, and frequent drought severely limit freshwater availability worldwide. 1,2 With the rapid development of advanced water treatment technologies, potable reuse of wastewater effluent becomes invaluable to augment drinking water supplies.3 Typically, municipal wastewater effluent is fed into a membrane-based water reuse treatment train that sequentially consists of microfiltration (MF), reverse osmosis (RO), and an ultraviolet light-based advanced oxidation process (UV/AOP).4 Because uncharged and low-molecular-weight trace organic contaminants can pass through RO membranes, and other wastewater effluent-derived organic contaminants can pass through aged RO membranes, 5,6 UV/AOP is employed post-RO to eliminate trace organic contaminants from RO permeate. This process harnesses the generation of reactive radical species from photo-oxidants in a photochemical system with an electrical energy input, facilitating an oxidative degradation of trace organic contaminants. 7,8 Different photo-oxidants including hydrogen peroxide (H_2O_2) , persulfate $(S_2O_8^{2-})$, free chlorine (HOCl), and chloramines (NH2Cl and NHCl2) are applied or proposed. 9-13

Monochloramine (NH2Cl), formed by HOCl addition to the typical ammonium ion (NH₄⁺)-containing wastewater effluent, is an antifouling chemical widely used in membranebased potable reuse treatment. Because NH₂Cl passes through RO membranes and subsequently undergoes photolysis, 14-16 the photolysis of NH₂Cl can be harnessed via a UV step to degrade micropollutants in the RO permeate. 11,17,18 Meanwhile, NH2Cl disproportionates to generate dichloramine (NHCl₂) in acidic conditions. ¹⁹ The formation of NHCl₂ is under a favorable condition in the RO permeate because of its acidic pH (~6) and the absence of NH₄⁺ due to the high rejection rate of positively charged ions by the RO membrane (Reaction 1) 20,21

$$2NH_2Cl + H^+ \rightarrow NHCl_2 + NH_4^+$$
 (R1)

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Prior studies suggest that the decomposition of NHCl₂ can lead to an oxidative capacity. For example, the addition of HOCl into NH₄⁺ or NH₂Cl was reported to rapidly degrade 1,4-dioxane (1,4-D), an indicator compound regulated for potable reuse in California. NHCl₂ decomposition also results in the formation of *N*-nitrosodimethylamine (NDMA) when subjected to high radical exposure in the presence of NDMA precursors. Study has investigated NHCl₂ decomposition with a focus on NDMA formation and other significant yet unidentified end products through the nitrogen species pathway. In particular, peroxynitrite (ONOO⁻), a nitrosating agent primarily responsible for NDMA formation in the presence of NDMA precursors, was identified as a critical intermediate during NHCl₂ decomposition.

However, the mechanisms of NHCl₂ hydrolysis on the production of reactive radical species through oxidative reactions need better understanding. ONOO- can trigger HO generation due to the self-decomposition of its protonated form, peroxynitrous acid (ONOOH). 27,28 Additionally, the oxidative capacity induced by background preexisting NHCl₂ hydrolysis in RO permeate for trace organic degradation under potable reuse conditions has yet to be quantified. Strategies including the upstream removal of precursors (e.g., membrane-based treatments for the removal of large molecules of NDMA precursors) and the posttreatment adjustment of water quality parameters (e.g., increasing pH to destabilize NHCl₂) can effectively minimize NDMA formation. 21,29 Moreover, the generated HO• has been proven effective in degrading NDMA. 30 Hence, the hydrolysis of pre-existing NHCl2 in RO permeate could contribute as a complementary non-UV AOP to conventional UV/AOPs, thereby reducing the electric energy consumption of traditional UV processes.

The pH of RO permeate can significantly impact the oxidative efficiency of NHCl₂ hydrolysis in several ways. First, the pH affects the speciation of the chloramines and carbonate, which are photo-oxidants and strong HO $^{\bullet}$ scavengers, respectively. Second, the pH of RO permeate affects the equilibrium between NH₂Cl and NHCl₂, which can ultimately change the NH₂Cl-to-NHCl₂ molar ratio. In addition, the pH affects the kinetics of chloramine decomposition and the equilibrium of the ONOOH/ONOO pair (p $K_a = 6.6$). Therefore, the effects of the RO permeate pH on the NHCl₂-initiated oxidative reactions need to be systematically quantified, and the optimal NH₂Cl-to-NHCl₂ ratio for HO $^{\bullet}$ generation needs more exploration.

Furthermore, it is important to examine the effects of total carbonate (TOTCO₃) on the oxidative capacity of the NHCl₂ hydrolysis because carbonate species can alter radical speciation and contribution.³⁸ RO desalination is highly pressurized, which increases the solubility of atmosphere carbon dioxide (CO₂) and therefore renders an elevated level of TOTCO3 in RO permeate. This increases the concentrations of dissolved CO₂ and bicarbonate (HCO₃⁻) and substantially suppresses the oxidative capacity of NHCl₂ hydrolysis. 23 For instance, CO2 can strongly scavenge ONOO-, preferentially forms selective carbonate radical (CO3 •-) via a two-step reaction instead of generating extremely reactive, nonselective HO[•]. ³⁹ HCO₃ can also scavenge HO $^{\bullet}$ to shift the radical speciation toward CO $_3$ $^{\bullet-}$ ($k_{\rm HO\bullet,HCO3}^-=8.6\times10^6~{\rm M}^{-1}~{\rm s}^{-1}$). Therefore, it is crucial to investigate the effects of TOTCO3 on NHCl2 hydrolysis in RO permeate.

Additionally, the efficiency of the NHCl $_2$ hydrolysis on the degradation of indicating trace organic contaminants in potable reuse needs investigation. For example, 1,4-D, a probable human carcinogen, has been widely used as a solvent stabilizer in industry; it passes through RO membranes and exists in RO permeate. Pharmaceuticals including carbamazepine (CBZ), atenolol (ATN), and sulfamethoxazole (SMZ) have been detected in RO permeate, mainly when membrane aged and separation efficiency decreases. Consequently, it is important to evaluate the performance of the NHCl $_2$ hydrolysis in potable reuse systems with respect to the removal of these indicating contaminants.

Accordingly, this study aimed to investigate the mechanism of NHCl₂ hydrolysis in RO permeate chemical conditions by understanding the reaction pathways leading to the generation of oxidative capacity, quantify the effects of RO permeate chemical parameters (i.e., pH, NH₂Cl-to-NHCl₂ ratio, and TOTCO₃) on radical yield and speciation, and identify the optimal condition for trace organic removal. 1,4-D, CBZ, ATN, and SMZ were chosen as the indicator organic contaminants to assess the treatment efficiency of the NHCl₂ hydrolysis.

■ MATERIALS AND METHODS

Materials and Reagents. All chemicals were ACS reagentgrade or higher from Fisher Scientific and Sigma-Aldrich. Ultrapure MQ water (>18.2 M Ω ·cm, Millipore) was utilized to prepare all solutions except for experiments using RO permeate. The chloramine synthesis followed well-established procedures.²³ Fresh NH₂Cl working solution with negligible ammonia was synthesized by titrating NaOCl to (NH₄)₂SO₄ at a nitrogen-to-chlorine molar ratio of 1.2 at pH 8.8 and buffered with 4 mM borate and equilibrated for 2 h before use. Fresh NHCl₂ stock solutions were prepared by acidifying the NH₂Cl working solution with 10 M HClO₄ to pH 3.5. A 100 mM persulfate stock solution was freshly prepared using Na₂S₂O₈. Phosphate buffers to control pHs between 7 and 9 were made using a predetermined combination of NaH2PO4 and Na₂HPO₄. Borate buffers to control pH values above 9 consisted of a proper combination of H₃BO₃ and Na₂B₄O₇. ONOO stock solutions were synthesized by rapidly adding HCl to the mixed solution containing H₂O₂ and NaNO₂ with instant quench using NaOH.46 Details on the preparation of the ONOO solutions are provided in Text S1 in the Supporting Information (SI). Other working solutions of photo-oxidants (i.e., H₂O₂, HOCl, and S₂O₈²⁻) were prepared

Synthetic fresh RO permeate was prepared by combining 1 μ g/L trace organic contaminants (specifically, 1,4-D, CBZ, ATN, or SMZ), 3 mg/L NHCl₂, and 2 mM NaHCO₃ at pH 5.8. The concentrations of these chemical components represent realistic RO permeate conditions. The synthetic aerated RO permeate sample was prepared the same way with the addition of 0.05 mM NaHCO₃. These carefully chosen levels of TOTCO₃ represent the background carbonate levels in fresh RO permeate and post-treatment after aeration, respectively.

Fresh RO permeate was collected in situ from a full-scale municipal potable reuse facility at Orange County Water District (OCWD) in Fountain Valley, California. Fresh RO permeate was collected from the outlet of the full-scale RO permeate module and preserved in a 4 L amble bottle without headspace. Aerated RO permeate was prepared by purging air

to the fresh RO permeate in a 4 L beaker with constant agitation for 2 h on-site to remove TOTCO₃.

Bench-Scale Experiments on NHCl₂ Hydrolysis and Oxidative Reactions. Bench-scale NHCl₂ hydrolysis experiments were conducted in a 50 mL Petri dish reactor in triplicates at 25 °C. The predetermined buffer volume at the targeted pH and nitrobenzene (NB) were added to the Petri dish reactor. Following that, an appropriate amount of fresh NHCl₂ stock solution was added to the reactor to start the reaction, creating an initial solution containing 3 mM NHCl₂, 5 μ M NB, and 60 mM buffer under constant rapid agitation. The initial NHCl₂ concentration of 3 mM was chosen to allow for accurate radical quantification. Samples were withdrawn from the reactor at 0 and 15 s of the hydrolysis reaction for immediate chemical analysis. The reaction time of 15 s was selected based on the rapid kinetics observed in preliminary experiments.

To investigate the effects of pH on NHCl₂ hydrolysis, buffers were introduced to the reactor with pHs ranging from 7 to 12. This wide pH range allows for a mechanistic investigation of the reaction system. Solution pH was stable during the experiments. To explore the effects of the NH₂Clto-NHCl2 ratio on NHCl2 hydrolysis, varying levels of fresh NH₂Cl stock solutions were added to the reactor in separate experiments to obtain a NH₂Cl-to-NHCl₂ molar ratio between 0 and 3, which covers relevant conditions in real-world potable reuse scenarios. Additional experiments to investigate carbonate effects were conducted by adding NaHCO3 to obtain a TOTCO₃ level between 0.1 and 20 mM at pH 8.4. HCO₃ was utilized to simulate scenarios in real-world RO permeate, as the highly pressurized RO process increases the solubility of CO₂ in acidic RO permeate, and it is transformed to HCO_3^- when the RO permeate is neutralized to a higher pH as final treated water. Twenty μ M N,N-dimethylaniline (DMA) was added as the $CO_3^{\bullet-}$ exposure probe, and 5 μ M NB was added as the HO exposure probe in carbonate experiments. Furthermore, bench-scale NHCl₂ hydrolysis experiments were conducted using synthetic RO permeates at a pH of 8.4, employing 0.8 μM NB as a probe compound to quantify the exposure to HO[•].

Experiments on NHCl₂ Hydrolysis Using RO Permeate at a Potable Reuse Facility. To assess the effectiveness of the proposed NHCl₂ hydrolysis as a novel non-UV AOP for degrading trace-level organic contaminants, field experiments were conducted at OCWD using fresh RO permeate from their full-scale municipal potable reuse facility. Fresh RO permeate was collected first. In addition, to evaluate the effects of aeration of RO permeate on oxidative capacity, additional freshly collected RO permeate was bubbled with air at ambient temperature for up to 2 h to degas and produce the aerated RO permeate. The concentrations of pre-existing NH₂Cl and NHCl₂, pH, and TOTCO₃ in both fresh and aerated RO permeate samples were measured, and the RO permeate was subsequently transferred to a 500 mL reactor. Background NHCl₂ was already present in the fresh RO permeate, and additional freshly prepared NHCl2 stock was added to obtain an initial NHCl₂ concentration of 3 mg/L in the RO permeate for the field experiment. This targeted level of NHCl₂ falls within the typical background concentration found in realistic RO permeate. Additionally, the solution was spiked with 1 μ g/L of 1,4-D and 0.8 μ M NB. After that, the RO permeate was adjusted to a pH of 8.4 to start hydrolysis reactions, and samples were taken after 15 s of reaction for analysis.

Decomposition of NHCl₂ and ONOO⁻ and UV Photolysis Experiments. Experiments on NHCl₂ and ONOO decay kinetics were conducted in a 3-mL quartz cuvette reactor at pHs from 7 to 12. Details of the decomposition experiments are provided in Text S2 and Figures S1 and S2. Control UV/AOP experiments with different photo-oxidants (i.e., H_2O_2 , HOCl, $S_2O_8^{2-}$, NH_2Cl , NHCl₂, NH₂Cl with H₂O₂, and NHCl₂ with H₂O₂) were conducted to compare with the oxidative capacity of the NHCl₂-initiated non-UV hydrolysis AOP. UV/AOP control experiments were conducted at pH 5.8 (typical RO permeate pH) in a 50-mL Petri dish reactor under constant agitation with a 254 nm low-pressure mercury lamp (Ultra Sum Technologies). The UV fluence of the UV system was determined by atrazine actinometry to confirm the targeted value of 850 mJ/cm², which is the typical UV fluence received by RO permeate at the full-scale UV/AOP treatment.⁴⁷ Details on the UV/AOP control experiments are provided in Text S3.

Analytical Methods. Each sample taken from the NHCl₂ hydrolysis experiments was analyzed for concentrations of NH₂Cl and NHCl₂ using the standard N,N-diethyl-p-phenylenediamine (DPD) method. 48 Alkalinity was measured according to the standard titration method.⁴⁹ Probe compounds of NB and DMA were quantified by utilizing highperformance liquid chromatography coupled with a diode array detector (HPLC-DAD, Agilent). Trace concentrations of 1,4-D (1 μ g/L or less) were determined according to EPA Method 522 by gas chromatography coupled with mass spectrometry (GC-MS, Agilent). Trace concentrations of CBZ, ATN, and SMZ were analyzed by ultrahigh-performance liquid chromatography coupled with high-resolution tandem mass spectrometry (HPLC-HRMS/MS, Q Exactive, Thermo Fisher Scientific). Details of analytical methods for 1,4-D and all indicator compounds are provided in Text S4. The concentration of H₂O₂ was quantified by the titanium oxalate method.⁵⁰ Concentration of S₂O₈²⁻ was determined using the KI titration method.⁵¹ The stock solution of ONOO⁻ was determined via a direct absorbance method at 302 nm.⁴⁶

To determine the cumulative HO^{\bullet} exposure from $NHCl_2$ hydrolysis and UV/AOP control, NB was used as a selective HO^{\bullet} probe (details regarding the calculation of cumulative HO^{\bullet} exposure are provided in Text S5 and Table S1). To quantify the contribution of radicals during $NHCl_2$ hydrolysis, NB and DMA were utilized as probe compounds for HO^{\bullet} and $CO_3^{\bullet-}$, respectively. A competition kinetic method was used for calculating the steady-state concentrations for the eventual calculation of the cumulative exposure of HO^{\bullet} and $CO_3^{\bullet-}$ (Text S5).

Mathematical Model to Predict the Oxidative Capacity of NHCl₂ Hydrolysis. A mathematical model was established to predict the oxidative capacity of the NHCl₂ hydrolysis-initiated AOP based on reaction pathways and mechanisms discovered in this study. The optimization of the model was carried out using a training data set composed of experimental data on the hydrolysis of NHCl₂ and ONOO⁻ to fit the HO⁴ reaction efficiency coefficient α in the model. A comprehensive global fitting procedure was employed to fit the experimentally observed mean values of HO⁴ exposure to model-predicted outcomes by optimizing the value of an intrinsic model parameter, γ , which is the NH₂Cl effect factor. During the model fitting to correlate the experimentally observed and model-predicted HO⁴ exposure data, the slope was constrained to be within 1 ± 0.25, the intercept ≤2 ×

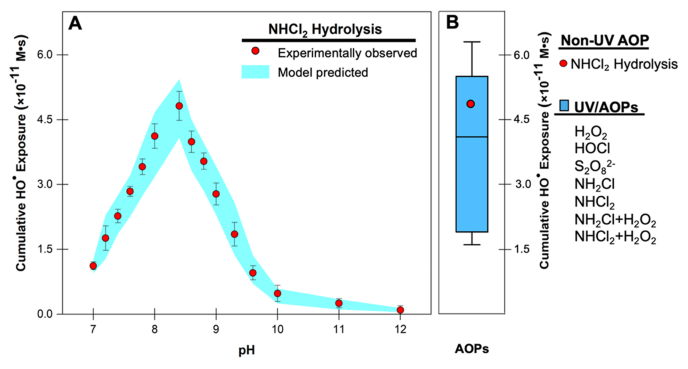
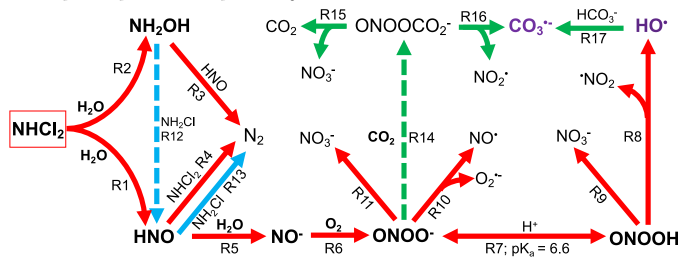


Figure 1. Cumulative hydroxyl radical exposure of non-UV NHCl₂ hydrolysis vs traditional UV/AOPs. (A) NHCl₂ hydrolysis at pH 7–12. Buffer = 60 mM for all pHs, [NHCl₂]₀ = 3 mM, [NB]₀ = 5 μM, and reaction time = 15 s. (B) UV photolysis of H₂O₂, HOCl, S₂O₈²⁻, NH₂Cl, and NHCl₂ at pH 5.8; in the single-oxidant UV/AOP system: [oxidant]₀ = 1 mM; in the combined-oxidant UV/AOP system, [chloramine]₀ = 1 mM, [H₂O₂]₀ = 2 mM; [NB]₀ = 100 μM, UV fluence = 850 mJ/cm². The solid line in the box plot is the median value, and the lower and upper box edges are the 25th and 75th percentiles. Whisker bars represent the minimum and maximum values. Details of the HO[•] exposure of UV/AOPs are provided in Table S6. Mathematical model fitting represents one standard deviation of the prediction: $\alpha = 0.25 \pm 0.05$, $\gamma = 1$ (details of the model fitting are provided in Text S6). The predicted shaded area represents a 68% confidence interval. Error bars represent the standard deviation of experimental triplicates.

Scheme 1. Reaction Scheme for the NHCl₂ Hydrolysis System, with Proposed Reaction Pathways for the Generation of Reactive Species Responsible for Micropollutant Degradation^a



 a Major reaction pathways (red), generated radicals for micropollutant removal (purple), NH $_{2}$ Cl-related reactions (blue), and scavenging reactions (green).

 10^{-5} , and the $R^2 \ge 0.85$. The detailed model optimization procedure is provided in Text S6. The fully optimized model was tested to predict the removal of trace organic contaminants under both synthetic and RO permeate conditions.

■ RESULTS AND DISCUSSION

Mechanisms of pH Effects on NHCl₂ Hydrolysis and Cumulative HO[•] Exposure. The oxidative capacity of the NHCl₂ hydrolysis system was examined as a function of the solution pH. The degradation of NB was observed during the NHCl₂ hydrolysis within 15 s at pHs ranging from 7 to 12 (Figure 1A). The results demonstrated that HO[•] was

generated through the hydrolytic decay of NHCl₂ at pH levels where NHCl₂ is unstable, consistent with observations from our previous study.²³ In the NHCl₂-initiated hydrolysis system, when NHCl₂ destabilizes at certain pH levels, it undergoes hydrolytic decay, producing nitroxyl (HNO) and hydroxylamine (NH2OH) via R1 and R2, respectively (R1 and R2 in Scheme 1; the following reactions all refer to Scheme 1). 26,54 NH₂OH further reacts with HNO to form nitrogen gas (N₂) via R3.55 HNO reacts with NHCl₂ as a side reaction to produce N2 (R4), but more significantly, HNO deprotonates to form nitric oxide anion (NO⁻) via R5, which is further rapidly oxidized by dissolved oxygen in the solution to generate the critical intermediate peroxynitrite (ONOO-; R6; $k = 2.7 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$. Upon protonation, ONOO becomes peroxynitrous acid (ONOOH; R7; $pK_a = 6.6$).³⁷ Most importantly, ONOOH directly decays into HO and nitrogen dioxide radical ($^{\circ}NO_2$) via R8 ($k = 3.5 \times 10^{-1} \text{ s}^{-1}$) or produces NO_3^- as a stable product via R9 ($k = 9.0 \times 10^{-1}$ s⁻¹).⁵⁸ Meanwhile, ONOO undergoes self-decomposition into NO $^{\bullet}$ and O₂ $^{\bullet-}$ (R10; $k = 2.3 \times 10^{-2} \text{ s}^{-1}$) or yields nitrate (NO₃ $^{-}$) by R11 ($k = 9.0 \times 10^{-6} \text{ s}^{-1}$). S4,58 The generated HO $^{\bullet}$ was not scavenged by the maximum 3 mM chloride (Cl⁻), which was introduced by the NaOCl solution utilized for NH₂Cl preparation (detailed calculations are provided in Text S5). This is because the first-order reaction rate of the forward reaction (HO $^{\bullet}$ + Cl $^{-}$ \rightarrow ClOH $^{\bullet-}$; $k \times [Cl^{-}] = 1.3 \times 10^{7} \text{ s}^{-1}$) is much slower than the backward reaction (ClOH $^{\bullet-} \rightarrow HO^{\bullet} +$ Cl^- ; $k = 6.1 \times 10^9 \text{ s}^{-1}$.

The maximal HO[•] exposure peaked at pH 8.4 and decreased either toward a neutral pH of 7 or at higher alkaline pHs up to 12, with suppressing percentages of 77 and 98% (Figure 1A), respectively. HO generated from NHCl2 hydrolysis can greatly benefit micropollutant abatement due to its high cumulative exposure within a short time.²² At the optimal pH of 8.4, the cumulative HO[•] exposure within 15 s of NHCl₂ hydrolysis was 4.8×10^{-11} M·s, a level that stands out as a very high value when compared with the HO exposure levels generated by existing UV/AOPs commonly utilized or suggested for implementation in potable reuse systems (Figure 1B). HO• exposure from UV/AOPs is between 1.6 and 6.3 × 10⁻¹¹ M·s. The fact that the optimal oxidative capacity of the NHCl₂ hydrolysis is at the higher end of UV irradiation-based AOPs for water reuse is very promising, suggesting an alternative non-UV-based AOP with significantly less energy footprint due to the omission of UV lamps and photon energy.

Prior literature proposed a two-step decay pathway that leads to HO generation, via first NHCl2 undergoing a hydrolytic decay to form ONOO (Reactions R1-R6), and subsequently the generated ONOO yields HO (R7-R8). 22 The decay of NHCl₂ and ONOO was quantified through the decomposition experiments and data showed that the NHCl2 hydrolysis rate increased by more than 1 order of magnitude when the solution pH increased from 7 to 12, increasing from 1.4×10^{-2} to 1.9×10^{-1} s⁻¹ (Figure 2). This is because NHCl₂ hydrolysis favors alkaline conditions and becomes unstable when the pH is above neutral. 60 In contrast, the decomposition rate of ONOO increased by nearly 2 orders of magnitude when the pH decreased from 12 to 7. ONOO remians stable as an anion at alkaline pH levels. However, as the pH decreases and protonation occurs, ONOO rapidly decomposes into HO• and •NO₂ (R7-R8). The opposite trends in NHCl₂ and ONOO decay rates (Figure 2) created a combined effect on

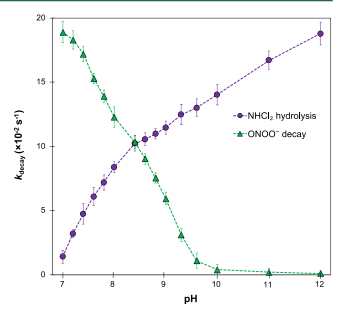


Figure 2. Experimentally observed *pseudo*-first-order rates of NHCl₂ hydrolysis and ONOO⁻ decay at different pHs. Buffer = 60 mM at all pHs, $[\mathrm{NHCl_2}]_0 = 42~\mu\mathrm{M}$, $[\mathrm{ONOO^-}]_0 = 850~\mu\mathrm{M}$, and reaction time = 15 s. Error bars represent the standard deviation of experimental triplicates.

HO• formation that peaked at an intermediate pH level between 7 and 12 (i.e., at pH 8.4 as observed in Figure 1A).

Mechanisms of NH₂Cl-to-NHCl₂ Molar Ratio Effects on NHCl₂ Hydrolysis and HO^o Exposure. The NH₂Cl-to-NHCl₂ molar ratio greatly affected the oxidative capacity of the NHCl₂ hydrolysis system (Figure 3A). The HO[•] exposure was the highest at a NH₂Cl-to-NHCl₂ molar ratio of 1 and declined when the NH₂Cl-to-NHCl₂ molar ratio deviated from 1. The HO exposure was suppressed by 19 and 47% when shifting the NH₂Cl-to-NHCl₂ molar ratio from 1 to 0 and 3, respectively. These trends of molar ratio effects resulted from two competing effects of NH2Cl on HO exposure during NHCl₂ hydrolysis. When NH₂Cl is introduced to the NHCl₂initiated AOP system, the generated NH2OH reacts with NH₂Cl to form HNO (R12 in Scheme 1).⁶¹ The formation of HNO resulted in higher HO^o exposure, as HNO is the primary precursor for the subsequent generation of ONOO-/ ONOOH, leading to HO^o production (R5-R8). However, as the concentration of NH2Cl further increases, it also scavenges HNO to produce N₂ (R13).⁵⁶ As the NH₂Cl-to-NHCl₂ molar ratio increases beyond the optimal level, R13 becomes more dominant than R12, consequently leading to a more substantial scavenging effect rather than a promoting effect on HO[•] exposure. The trade-off effects led to an optimal NH₂Cl-to-NHCl₂ molar ratio at 1 during NHCl₂ hydrolysis on HO[•] exposure (Figure 3A).

At the optimal $\mathrm{NH_2Cl}$ -to- $\mathrm{NHCl_2}$ molar ratio of 1, the highest HO^{\bullet} exposure of 5.9×10^{-11} M·s was observed at pH 8.4, consistent with the pH effects in the absence of $\mathrm{NH_2Cl}$ (Figure 1A). It showed an approximately 23% enhancement compared with the system without $\mathrm{NH_2Cl}$ across all pHs (Figures 3B vs 1A). The results suggested that $\mathrm{NH_2Cl}$ promoted HO^{\bullet} exposure across a wide solution pH during $\mathrm{NHCl_2}$ hydrolysis.

Mechanisms of Carbonate Effects on NHCl₂ Hydrolysis and Radical Speciation. Fresh RO permeate is elevated in dissolved CO₂ and therefore TOTCO₃ due to enhanced

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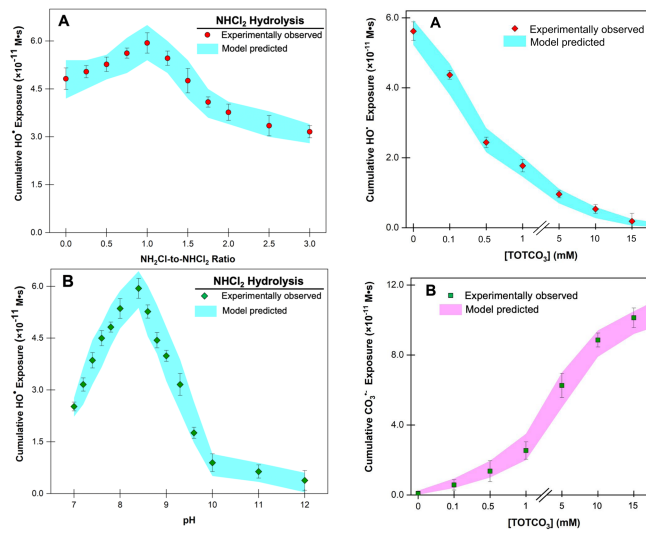


Figure 3. Cumulative hydroxyl radical exposure via NHCl₂ hydrolysis. Buffer = 60 mM for all pHs, [NHCl₂]₀ = 3 mM, [NB]₀ = 5 μ M, and reaction time = 15 s. (A) NH₂Cl-to-NHCl₂ molar ratio effects at pH 8.4; (B) pH effects at the NH₂Cl-to-NHCl₂ molar ratio of 1. Mathematical model fitting represents one standard deviation of the prediction: α = 0.25 \pm 0.05, γ = 1 (details of the model fitting are provided in Text S6). The predicted shaded area represents a 68% confidence interval. Error bars represent the standard deviation of experimental triplicates.

dissolution of the gas from pressurized air during RO operation and the fact that RO permeate is acidic. Harnessing the oxidative capacity of the NHCl₂ hydrolysis to its maximum needs to consider the TOTCO₃ effects. At acidic pHs, ONOO⁻ is scavenged by the dissolved CO₂ to form ONOOCO₂⁻ (R14 in Scheme 1; $k = 2.9 \times 10^4 \, \text{M}^{-1} \, \text{s}^{-1}$), which then self-decomposes into CO₂ and NO₃⁻ (R15) or decays into CO₃•- and NO₂• (R16). ^{39,62} Additionally, at neutral-to-alkaline pHs, HO• is further converted to CO₃•- by HCO₃⁻ (R17; $k = 8.6 \times 10^6 \, \text{M}^{-1} \, \text{s}^{-1}$). Therefore, the HO• exposure is suppressed and converted to CO₃•- exposure in the presence of TOTCO₃.

Experimental results showed that the cumulative HO[•] exposure was reduced by 30% in the presence of 0.1 mM TOTCO₃, further reduced by 70% when the TOTCO₃ level reached 1 mM, and suppressed to a negligible level in the presence of 20 mM TOTCO₃ (Figure 4A). The scavenging of

Figure 4. Cumulative radical exposure during NHCl₂ hydrolysis under pH 8.4 at varying TOTCO₃ concentrations. (A) Cumulative HO• exposure; (B) cumulative CO₃•- exposure. [NHCl₂]₀ = 3 mM, [NB]₀ = 5 μ M, [DMA]₀ = 20 μ M, and reaction time = 15 s. Mathematical model fitting represents one standard deviation of the prediction: α = 0.25 \pm 0.05, γ = 1 (details of the model fitting are provided in Text S6). The predicted shaded area represents a 68% confidence interval. Error bars represent the standard deviation of experimental triplicates.

HO[•] was accompanied by an increase in the level of CO₃^{•–} exposure. The CO₃^{•–} exposure was enhanced by nearly 19-fold when the TOTCO₃ was shifted from 0.1 to 20 mM (Figure 4B). Because the standard RO process is pressurized and elevates the dissolution of CO₂, the high TOTCO₃ levels in fresh RO permeate can compromise the HO[•] exposure. Consequently, the HO[•]-driven oxidative capacity of the NHCl₂-based AOP favors a low TOTCO₃, which can be achieved by an aeration step to decarbonize the fresh RO permeate in the water reuse facilities.

NHCl₂ Hydrolysis AOP in Synthetic RO Permeate Conditions. NHCl₂ hydrolysis-based AOP significantly degraded the four relevant trace organic contaminants in the synthetic RO permeate (Figure 5A). Under normal RO operation, the fresh RO permeate contains high levels of TOTCO₃ that can be up to 2 mM (details are provided in Text S7). The experimentally observed *pseudo-*first-order degrada-

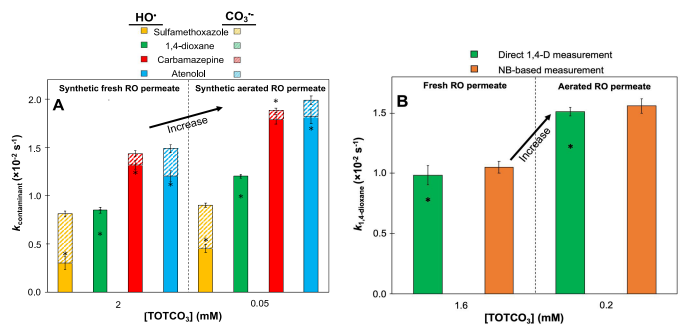


Figure 5. Effects of TOTCO₃ on *pseudo*-first-order degradation rates of trace contaminants via NHCl₂ hydrolysis at pH 8.4 under synthetic RO permeate vs RO permeate. The sign of "*" represents the predicted *pseudo*-first-order rates of trace contaminants by the mathematical model (details of the model prediction are provided in Text S6). (A) Synthetic fresh RO permeate: $[TOTCO_3]_0 = 2$ mM; synthetic aerated RO permeate: $[TOTCO_3]_0 = 0.05$ mM, buffer = 60 mM, $[NHCl_2]_0 = 3$ mg/L, $[1,4-D]_0 = 1$ μ g/L, $[CBZ]_0 = 1$ μ g/L, $[ATN]_0 = 1$ μ g/L, $[SMZ]_0 = 1$ μ g/L, $[NB]_0 = 0.8$ μ M; (B) fresh RO permeate: $[TOTCO_3]_0 = 1.6$ mM; aerated RO permeate: $[TOTCO_3]_0 = 0.2$ mM, $[NHCl_2]_0 = 3$ mg/L, $[NH_2Cl]_0 = 2.1$ mg/L, $[1,4-D]_0 = 1$ μ g/L, $[NB]_0 = 0.8$ μ M. Reaction time = 15 s. Error bars represent the standard deviation of the experimental triplicate.

tion rates (denoted as $k_{\rm contaminant}$) of SMZ, 1,4-D, CBA, and ATN were 8.1×10^{-3} , 8.5×10^{-3} , 1.4×10^{-2} , and 1.5×10^{-2} s⁻¹, respectively (left panel in Figure 5A). When the RO permeate undergoes an aeration process to decarbonate, the TOTCO₃ level in the RO permeate can be reduced to 0.05 mM (details are provided in Text S7). In this scenario, the experimentally observed degradation rates for all selected compounds were significantly enhanced (by up to 140%) in comparison to the normal RO operation conditions (right panel in Figure 5A). In both scenarios, the contribution of HO $^{\bullet}$ to the degradation of trace organic contaminants is predominant.

Specifically, when the RO permeate is aerated to decarbonate, the degradation rate of SMZ increased by 11% because the rate constants of SMZ with HO[•] and CO₃^{•-} are comparable (rate constants between the four selected contaminants with HO• and CO₃•- are provided in Table S2).^{63,64} 1,4-D degradation rate was enhanced by 42% because it reacts with HO but is not reactive with CO₃ -65,66 Therefore, the 1,4-D degradation favors a low TOTCO3 level. Similarly, the observed degradation rates of CBZ and ATN were greatly enhanced by 1.4 times from 2 to 0.05 mM TOTCO3 because CBZ and ATN react with HO with the same order of magnitude as 1,4-D,63,67,68 while CBZ and ATN have a lower reactivity with CO₃ in comparison to SMZ. 69,70 1,4-D was degraded by HO^o regardless of the TOTCO₃ level, while both HO• and CO₃• oxidized the other three pharmaceuticals. The sensitivity of TOTCO3 for the four selected compounds toward their oxidative capacity follows the order of 1,4-D > CBZ ≈ATN > SMZ, which is reversely correlated with their reactivities with $CO_3^{\bullet -}$.

Model Prediction of Cumulative HO* Exposure during NHCl₂ Hydrolysis. To further gain a fundamental understanding of HO* generation initiated by NHCl₂

hydrolysis, a mathematical model was established to predict the HO^{\bullet} exposure during $NHCl_2$ hydrolysis under different solution chemical conditions at pHs between 7 and 12, as shown in eq 1

$$\int_{0}^{t} [HO^{\bullet}]_{t} dt = \alpha \gamma \left(\frac{k_{\text{NHCl}_{2}} k_{\text{ONOO}^{-}}}{k_{\text{NHCl}_{2}} + k_{\text{ONOO}^{-}}} \right)$$

$$\left(\frac{k_{\text{cont}, \text{HO}^{\bullet}} [\text{cont}]}{k_{\text{HCO}_{3}^{-}, \text{HO}^{\bullet}} [\text{HCO}_{3}^{-}] + k_{\text{cont}, \text{HO}^{\bullet}} [\text{cont}]} \right) \frac{t}{k_{\text{cont}, \text{HO}^{\bullet}}}$$
(1)

where $\int_0^t [HO^{\bullet}]_t dt$ is the cumulative HO^{\bullet} exposure via NHCl₂. hydrolysis-based AOP; α is the HO $^{\bullet}$ reaction efficiency coefficient, which represents the combined reaction efficiencies of three sequential pathways to generate HO*, including NHCl₂ decays into ONOO (R1-R6), ONOO decomposition to HO[•] (R7-R8), and HO[•] reaction with chemical of concerns; γ is the NH₂Cl effect factor and is a function of the NH_2Cl -to- $NHCl_2$ molar ratio; k_{NHCl_2} is the pseudo-first-order decay rate of NHCl2 at a specific pH, experimentally determined through decomposition experiments conducted at various pH levels by measuring the decrease in NHCl₂ concentration over a reaction period of 15 s; k_{ONOO} - represents the pseudo-first-order decay rate of ONOO at a specific pH, also experimentally determined through decomposition experiments conducted at various pH levels by measuring the decrease in ONOO⁻ concentration over a decomposition time of 15 s, t is the reaction time of NHCl₂ hydrolysis, which equals to 15 s; $k_{\text{HCO}_3^-,\text{HO}}$ is the second-order rate constant of the reaction between HO^{\bullet} and HCO_3^{-} (8.6 × 10^6 M⁻¹ s⁻¹), 40 $k_{\text{cont. HO}}$ is the second-order rate constant of the reaction between HO and a chemical of concern (e.g., with NB, $k_{\text{NB, HO}} = 4.7 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$, and [HCO₃] is the HCO₃

concentration at a specific pH; [cont] represents the concentration of chemical of concern (for model fitting, the chemical of concern is NB, which was utilized to quantify the cumulative HO $^{\bullet}$ exposure). The term $\frac{k_{\mathrm{NHCl}_2}k_{\mathrm{ONOO}^-}}{k_{\mathrm{NHCl}_2}+k_{\mathrm{ONOO}^-}}$ is a

standard kinetic expression to depict coupled sequential reactions, which represents the combined kinetics resulting from sequential reactions of NHCl₂ hydrolysis and ONOO⁻ decomposition for the eventual HO[•] production. The term

 $\frac{k_{\text{cont,HO}}[\text{cont}]}{k_{\text{HCO}_3^-,\text{HO}}[\text{HCO}_3^-] + k_{\text{cont,HO}}[\text{cont}]} \text{ is the branching ratio of HO}^{\bullet}$

reacting with the chemical of concern vs reacting with HCO $_3$ ⁻. Model fitting using NB decay experimental data at different pHs (Figure 1A) obtained the value of α as 0.25 \pm 0.05, which indicated the overall molar efficiency from NHCl $_2$ hydrolysis to HO $^{\bullet}$ production averaged at 25% (details on model fitting are provided in Text S6). Model fitting using NB decay experimental data at different NH $_2$ Cl-to-NHCl $_2$ molar ratios (Figure 3) obtained the value of γ as a function of the NH $_2$ Cl-to-NHCl $_2$ molar ratio. Specifically, the value of γ equals to 1 when the NH $_2$ Cl-to-NHCl $_2$ molar ratio $(\frac{[NH_2Cl]}{[NHCl<math>_2]})$ is zero;

 $\gamma = 0.23 \frac{[\mathrm{NH_2Cl}]}{[\mathrm{NHCl_2}]} + 1$ when the molar ratio is ≤ 1 ; $\gamma = -0.3 \frac{[\mathrm{NH_2Cl}]}{[\mathrm{NHCl_2}]} + 1.4$ when the molar ratio is >1 (detailed calculations are provided in Text S6).

The model predicted that NB degradation rates matched very well with the experimentally measured values (Figure S3), indicating the robustness of the mathematical model. The established mathematical model predicted well the experimentally measured cumulative HO^{\bullet} exposure during $NHCl_2$ hydrolysis across all pHs (blue shaded area in Figure 1A), in the presence of NH_2Cl (blue shaded area in Figure 3A,B) and HCO_3^- (blue shaded area in Figure 4A). The model confirmed that the decay of $NHCl_2$ and $ONOO^-$ governs HO^{\bullet} generation during $NHCl_2$ hydrolysis.

In the presence of varying TOTCO₃, the cumulative CO₃•-exposure generated via NHCl₂ hydrolysis AOP at pH 8.4 is mathematically modeled in eq 2

$$\int_{0}^{t} \left[\text{CO}_{3}^{\bullet -} \right]_{t} \, \mathrm{d}t = \alpha \gamma \left(\frac{k_{\text{NHCl}_{2}} k_{\text{ONOO}^{-}}}{k_{\text{NHCl}_{2}} + k_{\text{ONOO}^{-}}} \right)$$

$$\left(\frac{k_{\text{HCO}_{3}^{-}, \text{HO}^{\bullet}} [\text{HCO}_{3}^{-}]}{k_{\text{HCO}_{3}^{-}, \text{HO}^{\bullet}} [\text{HCO}_{3}^{-}] + k_{\text{cont}, \text{HO}^{\bullet}} [\text{cont}]} \right) \frac{t}{k_{\text{cont}, \text{CO}_{3}^{\bullet -}}}$$
(2)

where $\int_0^t [\mathrm{CO_3}^{\bullet-}]_t \mathrm{d}t$ is the cumulative $\mathrm{CO_3}^{\bullet-}$ exposure via NHCl₂ hydrolysis-based AOP; α , $\gamma \left(\frac{k_{\mathrm{NHCl_2}}k_{\mathrm{ONOO}^-}}{k_{\mathrm{NHCl_2}}+k_{\mathrm{ONOO}^-}}\right)$, and t are the same as in eq 1; the term $\frac{k_{\mathrm{HCO_3^-,HO^0}}[\mathrm{HCO_3^-}]}{k_{\mathrm{HCO_3^-,HO^0}}[\mathrm{HCO_3^-}]+k_{\mathrm{cont,HO^0}}[\mathrm{cont}]}$ is the branching ratio of HO $^{\bullet}$ reacting with HCO $_3^-$ vs reacting with the chemical of concern; $k_{\mathrm{cont,CO_3^{\bullet-}}}$ is the second-order rate constant of the reaction between $\mathrm{CO_3^{\bullet-}}$ and a chemical of concern (e.g., with DMA, $k_{\mathrm{DMA,CO_3^{\bullet-}}}=1.8\times10^9~\mathrm{M^{-1}~s^{-1}});^{72}$ [cont] represents the concentration of chemical of concern (for model fitting, the chemical of concern is DMA, which was utilized to quantify cumulative $\mathrm{CO_3^{\bullet-}}$ exposure). The established mathematical model predicted well the experimentally measured cumulative $\mathrm{CO_3^{\bullet-}}$ exposure during NHCl₂

hydrolysis across all TOTCO₃ levels at the optimal pH of 8.4 (purple-shaded area in Figure 4B).

The optimized model on cumulative HO• and CO₃•-exposure was further utilized to predict the degradation of trace organic contaminants in synthetic RO permeate samples (detailed calculations are provided in Text S5). The model prediction matched well with the experimentally measured pseudo-first-order degradation rates (represented by star signs vs bars in Figure 5A). The only exception was a slight underprediction of SMZ degradation, possibly due to an underestimate of the rate constant between CO₃•- and DMA.

NHCl₂ Hydrolysis AOP in RO Permeate Condition. The freshly collected RO permeate comprised background NH₂Cl and NHCl₂ alongside an oversaturated TOTCO₃ level (1.6 mM) with respect to atmospheric CO₂ equilibrium. Conversely, the aerated RO permeate exhibited the same background NH₂Cl and NHCl₂ concentrations but a notably reduced TOTCO3 level of 0.2 mM. The NHCl2 hydrolysis AOP demonstrated a significant degradation of trace-level 1,4-D in RO permeate, as depicted in Figure 5B. During standard RO operation, the experimentally observed pseudo-first-order degradation rates for 1,4-D were $1.0-1.1 \times 10^{-2} \text{ s}^{-1}$, as determined via direct measurement and the nitrobenzenebased (NB-based) indirect measurement (left panel in Figure 5B). The direct 1,4-D measurement represents the quantification of trace-level 1,4-D degradation occurring during NHCl2 hydrolysis; the NB-based indirect measurement employs NB as a probe for HO[•], confirming its exposure and subsequent degradation of 1,4-D. The strong agreement of 1,4-D degradation rates between the direct 1,4-D measurement and the NB-based indirect measurement (Figure 5B) further supported the substantial oxidative capacity of NHCl2 hydrolysis AOP under RO permeate conditions. Aeration of fresh RO permeate resulted in a substantial reduction of TOTCO₃ by nearly 90% (Table S3), concurrently shifting the pH from a slightly acidic condition toward neutrality. This adjustment brings the system closer to optimal conditions conducive to NHCl₂ hydrolysis. Consequently, the pseudo-firstorder 1,4-D degradation rate was enhanced by approximately 50% to $1.5-1.6 \times 10^{-2} \text{ s}^{-1}$ in aerated RO permeate due to the significant reduction in TOTCO₃ (right panel in Figure 5B).

In the RO permeate experimental conditions (Text S8 and Table S4), NH₂Cl and NHCl₂ coexist with a NH₂Cl-to-NHCl₂ molar ratio of 0.7. As depicted in Figure 3A, cumulative HO[•] exposure increased with an ascending NH2Cl-to-NHCl2 molar ratio from 0 to 1. This implies that the oxidative capacity of the NHCl2-driven non-UV AOP is augmented when subjected to realistic NH₂Cl and NHCl₂ levels present in the RO permeate. Moreover, prediction from the optimized kinetics model well predicted 1,4-D degradation in the fresh RO permeate and aerated RO permeate from experiments conducted at a potable reuse facility (represented by star signs vs bars in Figure 5B). The consistency between the model-predicted and experimentally observed 1,4-D degradation rates, obtained from experiments employing fresh and aerated RO permeate with varying water chemical parameters (including pH, NH2Cl, NHCl₂, and TOTCO₃) from an actual water reuse facility, supported the proposed mathematical model and validated the efficacy of the newly introduced NHCl2 hydrolysis AOP for potable reuse applications. This highlights the applicability of our model to diverse water treatment scenarios, enhancing its potential as a valuable tool for optimizing water reuse processes.

Engineering Implications. The findings of the study provide fundamental mechanistic insights into the NHCl₂ hydrolysis-initiated AOP and develop strategies for optimizing RO permeate conditions to maximize treatment efficiency with a significantly reduced energy footprint in water reuse applications (calculation is shown in Text S9). The NHCl₂initiated hydrolysis can be harnessed from the background preexisting NHCl₂ as a complementary non-UV AOP for the rapid degradation of trace organic contaminants in potable reuse. More importantly, this oxidative pathway does not require UV photon energy and efficiently destroys trace organic contaminants in the RO permeate. The trace-level organic contaminants in the RO permeate may differ when the source water shifts. Therefore, water reuse facilities can optimize the extent of aeration and the decarbonation level to maximize removal efficiency, based on the reactivities of HO• and CO3•- with a particular contaminant, since the radical speciation between HO and CO3 in the RO permeate can be controlled accordingly (detailed calculations are provided in Text S10 and Table S5). In addition, if the target contaminant has high reactivity with both CO3 •- and HO*, an aeration step to decarbonate the RO permeate may not be needed, as both radical exposures equally contribute to the degradation of the particular contaminant. The NH₂Cl-to-NHCl₂ molar ratio can also be adjusted to maximize radical exposure by controlling the pH and the NH2Cl disproportionation reaction in the RO feedwater.

Meanwhile, the risk of NDMA formation can be mitigated by implementing a pretreatment step, including membrane-based filtration, to eliminate its precursors. Additionally, employing a post-treatment procedure involving pH adjustment is recommended when utilizing the NHCl₂-initiated AOP. Specifically, raising the pH to its maximum level of 8.4 enhances the oxidative capacity of the NHCl₂-initiated AOP to its fullest extent. At this optimal pH of 8.4, the formation of NDMA can also be significantly minimized.²¹ In essence, while the NHCl₂-initiated AOP demonstrates its maximum oxidative capacity, it also proves to be beneficial for NDMA control simultaneously.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.est.4c04547.

Experimental solution preparation, UV photolysis experiments, mathematical model prediction, radical quantification, and analytical methods (PDF)

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Notes

The authors declare no competing financial interest.

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