Degradation of nitro-based pharmaceuticals by UV photolysis: Kinetics and simultaneous reduction on halonitromethanes formation potential

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This study investigated the degradation kinetics and halonitromethanes formation potential (HNMsFP) of two nitro-based pharmaceuticals (i.e., ranitidine (RNTD) and nizatidine (NZTD)) during ultraviolet (UV) photolysis. It was found that the degradation kinetics of RNTD and NZTD exhibited pH-dependent trends, in accordance with their deprotonation equilibria. The neutral species of RNTD and NZTD were more photo-reactive than their corresponding deprotonated species, with their specificfluence-based first-order rate constants varying in the range of 5.64–31.90 m2 E−1. Both the RNTD and NZTD were prone precursors of HNMs (with molar yields of 5.6±0.3% and 4.7±0.4%, respectively at pH 7.0). Acidic and neutral circumstances facilitated the HNMs formation. The UV photolysis of RNTD and NZTD could reduce their HNMsFP simultaneously. Positive linear relationships between residual RNTD or NZTD concentration and HNMsFP were observed and the denitration during the UV photolysis accounted for the HNMsFP reduction. With the mandatory UV disinfection fluences in China (i.e. 20–80 mJ cm−2), the effective abatement of RNTD and NZTD and their HNMsFP could not be fully achieved, highlighting the necessity of increasing UV fluence or developing UV-based advanced oxidation process in future.

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1. Introduction

Formation of toxic nitrogenous disinfection by-products (N-DBPs) is a growing concern for water utilities using impaired source waters with wastewater effluent discharges or algal blooms due to elevated concentrations of dissolved organic nitrogen as N-DBP precursors (Fang et al., 2010; Chu et al., 2014). HNMs are one class of emerging N-DBPs in drinking water, which have drawn public concern due to their high toxicity (Bond et al., 2012). Among 50 high priority DBPs monitored in a 2000–2002 U.S. nationwide survey at drinking water treatment plants (DWTPs), the cytotoxicity of HNMs with in vitro mammalian cell tests could be several orders of magnitude higher than currently regulated trihalomethanes (THMs) and haloacetic acids (HAAs) (Weinberg et al., 2002). Generally, the chemical structure of HNMs contains a nitro-group and different halogens on the α-carbon including mono-, di- and tri-halogenated species (Plewa et al., 2004).

Continued efforts have been made to identify the major HNM precursors and control HNMs formation by optimized pretreatment and disinfection processes for water and wastewater. After analyzing the HNM concentrations in 20 drinking water supply systems in the United Kingdoms, Bond et al. (2015) found that the summed concentration of HNMs was not correlated with that of THMs or HAAs. Yang et al. (2012) observed that the tri-chloronitromethane (TCNM) formation was higher during chlorination than chloramination, when spiked with 31 organic nitrogen compounds. During chloramination, both organic nitrogen and nitrate contributed to the nitrogen source of HNMs, while monochloramine did not contribute to the nitrogen source of TCNM. Nitro-based and amino-based compounds are considered as the direct and indirect precursors of HNMs, respectively (Bond et al., 2012; Deng et al., 2014).

Belonging to the group of histamine-2 blockers, ranitidine (RNTD) and nizatidine (NZTD) are prescribed worldwide as gastrointestinal drugs, which can reduce the amount of acid producers in stomach (Wu et al., 2001). As RNTD and NZTD cannot be
effectively removed in biological wastewater treatment process, both have been frequently detected in wastewater treatment plant (WWTP) effluents and receiving waters (Castiglioni et al., 2006; Radjenovic et al., 2009). The detected RNTD concentrations were reported in the range of 70–540 ng L$^{-1}$ in primary effluents of WWTPs (Radjenovic et al., 2009) and around 10 ng L$^{-1}$ in several surface waters (Kolpin et al., 2002). Apart from inducing adverse ecological consequences to receiving waters, the nitro-group of RNTD and NZTD may become the precursor of HNMs during disinfection. Effective strategy for reducing the HNMs formation is removing the main nitrogen-containing precursors before chlorination. As conventional drinking water treatment processes (e.g., coagulation/sedimentation and filtration) are ineffective in reducing low-molecular-weight organic nitrogen precursors (Lee and Westerhoff, 2006; Chu et al., 2011), pre-oxidation with chlorine, ozone, and some other oxidants has been evaluated to decrease the HNMsFP by transforming some key structural moieties of HNM precursors (Wang et al., 2016). Unfortunately, the HNMs formation may be promoted after the attack of oxidant and/or radicals. For precursors containing an amine-group, the combined ultraviolet (UV)/chlorine treatment could oxidize the amine-group to a nitro-group, which is always a rate-limiting step for the formation of HNMs (Deng et al., 2014). Ozonation also led to a promoted TCNM formation and primary and secondary alkyl amines are the predominant precursors (McCorry et al., 2016). UV irradiation has been recognized as a promising disinfection means because it neither requires additional chemical(s) nor impairs the aesthetic quality (e.g., taste and odor) of treated water and wastewater. However, little is known so far about the degradation of RNTD and NZTD by UV photolysis and the subsequent impact on the HNMsFP during chlorination is also largely unknown.

The objectives of this study were to: (1) determine the kinetic parameters of RNTD and NZTD degradation under UV (254 nm) irradiation; (2) clarify the effects of pH and chlorine concentration on the HNMsFP of RNTD and NZTD; and (3) determine the effect of UV photolysis on the subsequent HNMsFP of RNTD and NZTD during chlorination. It is expected to illustrate the behavior and fate of nitro-based pharmaceuticals during the disinfection of water and wastewater.

2. Materials and methods

2.1. Materials

Individual HNMs, including monochloronitromethane (MCNM), dichloronitromethane (DCNM) and TCNM, were all purchased from CanSyn Chem. Co. (BC, Canada). An internal standard, 1,2-dibromopropane, was purchased from J&K Chemical Co. (Beijing, China). RNTD and NZTD were purchased from Tokyo Chemical Industry Co. (Tokyo, Japan). High performance liquid chromatography (HPLC) grade methanol and acetonitrile were obtained from Thermo Fisher Scientific (NJ, USA), and formic acid (≥99.0%) from DKma Technologies (CA, USA). All other chemicals, of at least analytical grade, were provided by Beijing Chemical Reagents Company (Beijing, China). All solutions were prepared in ultrapure water (18.2 MΩ cm) produced by a Milli-Q system (Advantage A10, Millipore, MA, USA).

2.2. Experimental procedures

2.2.1. UV photolysis system

A cylindrical glass reactor (inner diameter (i.d.) = 10.0 cm, height = 25.0 cm) was used as the UV photolysis reactor with a working volume of 1.0 L (Fig. S1). The temperature of the UV system was controlled at 25 °C by circulating the water in the bath. A quartz tube (i.d. = 4.5 cm) and one low pressure (LP) Hg UV lamps (Heraeus, GPH 212TL5/4, 10 W) were fixed in the center of the UV reactor, which was magnetic mixed at the bottom of the reactor. The UV lamp was initially warmed up for about 30 min to ensure a relatively stable output. The photo flux ($I_0$) was determined to be 0.75 mE s$^{-1}$ for kinetic tests via iodide-iodate chemical actinometry (Bolton and Linden, 2003). The effective path length (L) was determined to be 7.2 cm by measuring the photolysis rate of H$_2$O$_2$ (Garoma and Gurol, 2005). The average UV fluence rate was determined to be 2.1 mW cm$^{-2}$ by atrazine actinometer and the UV photolysis experiments were conducted in triplicate.

The quantum yield ($\Phi$) is a fundamental parameter that quantifies the photon efficiency of a photochemical reaction (Shu et al., 2013). For the analysis of $\Phi$ during the UV photolysis at 254 nm, the aqueous solution to be irradiated was buffered with 10 mM phosphate buffer. To verify the UV photolysis of RNTD and NZTD in real waters, three real waters including filtered water (FW) from a DWTP, river water (RW) and WWTP effluent water (WE) were sampled and filtered (0.45 µm) prior to the UV photolysis. The addition of the stock solutions of RNTD and NZTD to the real water samples did not cause any substantial dilution (<1%) of the water matrix components. A UV–vis spectrophotometer (DR5000, Hach, CA, USA) was used to measure the apparent molar absorptivity of RNTD and NZTD at 254 nm. RNTD and NZTD concentrations ranged from 0 to 10 µM, and pH was controlled using 10 mM phosphate buffer.

2.2.2. HNMsFP test

HNM precursors are diverse and may be amenable to different treatment processes based on their chemical structures. To evaluate the effectiveness of treatment techniques, HNM precursors can be measured using a HNMsFP test, which was conducted by chlorinating water containing precursors and measuring the amount of HNMs formed over 24 h. The HNMsFP reduction is then quantified as the reduction in formation potential before and after the UV photolysis. HNMsFP tests were conducted in triplicate in the dark using 50 mL headspace-free screw-cap amber glass vials with PTFE-lined septa at 25 °C for 24 h.

2.3. Analysis

RNTD and NZTD concentrations were measured with an Agilent 1200 series HPLC (CA, USA) equipped with an Atlantis C18 column (4.6 × 250 mm, 5.0 µm; Waters, MA, USA) at an UV wavelength of 254 and 230 nm, respectively. The mobile phase was a mixture of methanol/water (65:35, v/v). The flow rate was set at 0.6 mL min$^{-1}$ and the injection volume was 50 µL. The temperature of the column was set at 25 °C. The formation of HNMs was analyzed after samples being extracted with methyl tert-butyl ether using a gas chromatograph (7890, Agilent, CA, USA) equipped with an HP 5 capillary column (30 m × 0.25 mm, 0.25 µm, J&W, USA) and an electronic capture detector using the modified USEPA method 551.1 (Munch and Hautman, 1995). NO$_3$ and NO$_2$ were analyzed by ion chromatography (IC) (ICS–2000, Dionex, CA, USA) with an AS19 column (4 × 250 mm) and an eluent of 30 mM KOH at a flow rate of 1 mL min$^{-1}$. Water samples were filtered (0.45 µm) prior to chromatographic analysis and the injection volume was 20 µL. Chlorine concentration was determined using DPD colorimetry.

The transformation by-products (TBPs) of RNTD and NZTD were identified by ultra-performance liquid chromatography–tandem mass spectrometry (UPLC–MS/MS). Agilent 1290 Infinity LC, Agilent 6420 Triple Quad LC/MS detector, CA, USA) coupled with an Acquity UPLC™ C18 column (2.1 × 100 mm, 1.7 µm) (Waters, USA). The mobile phase consisted of (A) formic acid/methanol (1:1000, v/v/...
v) (1:1000, v/v) and (B) formic acid/Milli-Q water (1:1000, v/v) pumped at a flow rate of 0.3 mL min\(^{-1}\). Elution gradient was programmed as follows (% of A, min): (5, 5), (30, 25), (30, 30), (90, 40), (90, 42), and (0, 60). The injection volume was 20 \(\mu\)L. All samples were analyzed with both total ion chromatogram and selected ion monitoring modes.

3. Results and discussion

3.1. Determination of molar absorption coefficients at 254 nm

Molar extinction coefficient (\(\varepsilon\)) has been widely acknowledged as a significant parameter which strongly affects the UV photolysis rate. As shown in Fig. S2, the absorption spectrum of RNTD exhibits the maximum band at 229 nm, and a weak band with a smooth maximum at 312 nm, while the absorption spectrum of NZTD exhibits the maximum band with a maximum at 311 nm, and a weak band with a smooth maximum at 254 nm. Fig. 1 shows that in the studied pH range of 5.0–9.0, the \(\varepsilon\) values of RNTD and NZTD decreased with the increase of the solution pH. This result reflects that the different species of an RNTD and NZTD should have different \(\varepsilon\) values at 254 nm. To quantitatively analyze the pH dependence of UV photolysis of RNTD and NZTD, the \(\varepsilon\) was modeled as a function of pH by assigning each protonated/deprotonated species a specific molar absorptivity (\(\varepsilon_0\) and \(\varepsilon_1\)), with the apparent absorptivity being the summation of the absorptivity contributed by each species (Eqs. (1)–(3)) (Canonica et al., 2008; Bahnmuller et al., 2015). The specific molar absorptivity for each RNTD and NZTD species was fit by minimizing the sum of differences squared between observed and modeled apparent molar absorptivity.

\[
\varepsilon = \varepsilon_0 \alpha_0 + \varepsilon_1 \alpha_1
\]  
\[
\alpha_0 = \frac{1}{1 + 10^{\beta pK_a - pH}} 
\]  
\[
\alpha_1 = \frac{1}{1 + 10^{\beta pK_a - pH}}
\]

Results indicated that the overall \(\varepsilon\) values experimentally measured could be well fitted by Eq. (1), and the specific \(\varepsilon\) values were 575 and 820 m\(^2\) mol\(^{-1}\) for neutral species and 491 and 752 m\(^2\) mol\(^{-1}\) for anionic species of RNTD and NZTD, respectively. For water and wastewater treatment, the typical pH usually ranges from 6.5 to 9.5, where the neutral and anionic species predominate in RNTD and NZTD solutions.

3.2. UV photolysis of RNTD and NZTD

The UV photolysis kinetics at pH 5.0–9.0 of RNTD and NZTD were further investigated. As shown in Figs. S3 and S4, the decrease

| Compound | \(\varepsilon_0\) (m\(^2\) mol\(^{-1}\)) | \(\varepsilon_1\) (m\(^2\) mol\(^{-1}\)) | \(k_0\) (m\(^{-1}\) E \(\text{m}^2\) \(\text{s}^{-1}\)) | \(k_1\) (m\(^{-1}\) E \(\text{m}^2\) \(\text{s}^{-1}\)) | \(\Phi_0\) | \(\Phi_1\) |
|---------|-----------------|-----------------|---------------------|---------------------|------|------|
| RNTD    | 575             | 491             | 31.90               | 18.53               | 0.022| 0.012|
| NZTD    | 820             | 752             | 9.49                | 5.64                | 0.0046| 0.0032|
of RNTD and NZTD concentrations as a function of time during the UV photolysis could be described by first-order kinetics, enabling calculation of pseudo-first-order photo degradation rate constants ($R^2 > 0.99$). Observed pseudo-first-order rate constants ($k$, s$^{-1}$), for RNTD and NZTD were obtained by linear regression of its logarithmic relative residual concentration over photolysis time $t$, according to Eq. (4).

$$-\ln \frac{C}{C_0} = kt$$

To allow a comparison of this study with those using other chemical pollutants by UV photolysis, the fluence-based first-order rate constant ($k_E$, m$^2$ J$^{-1}$) was calculated through Eq. (5) (Canonica et al., 2008; Bahnmuller et al., 2015).

$$k_E = \frac{k}{E}$$

where $E$ (einstein m$^{-2}$ s$^{-1}$) is the photon fluence rate, determined by chemical actinometry.

From Fig. 2, it is clear that the $k_E$ decreases from pH 5.0 to 9.0. These apparent rate constants were deconvoluted to calculate the specific fluence-based rate constants (Table 1) associated with the neutral and anionic species of the RNTD and NZTD during the UV photolysis with Eq. (6). After these fit parameters had been determined, the $\Phi$ values for the neutral and deprotonated species were determined using Eq. (7). Both the decreasing reactivity of deprotonated RNTD and NZTD with UV is due, in part, to the decreased molar absorptivity of the anionic compared to the neutral form.

This means that both the neutral forms of RNTD and NZTD, which dominate at pH below the $pK_a$, are more photo-reactive than the deprotonated form. The specific $k_E$ values of RNTD and NZTD vary by one order of magnitude, with a minimum for the anionic form of NZTD (5.64 m$^2$ E$^{-1}$) and a maximum for the neutral form of RNTD (31.90 m$^2$ E$^{-1}$). The high $\Phi$ and $k_E$ values at acidic and neutral pHs indicate that the photo efficiency of the transformation reaction, as measured by $\Phi$ at 254 nm, may be lower under alkaline circumstances.

$$k_E = k_{E0} + k_{E1}$$

$\Phi = \frac{k_E}{2.303\epsilon}$

The $\Phi$ values of RNTD and NZTD follow the same trend, decreased with the increase of pH. As shown in Table 1, the $\Phi$ values for neutral and anionic species of RNTD were 0.022 and 0.012, respectively (3.8 and 2.8 fold higher than those of NZTD, respectively), accounting for the faster degradation during the UV photolysis. The pH dependence of $\Phi$ might result from the acid-base speciation of excited-state species or from acid-catalyzed deactivation of photolysis intermediates. Meanwhile, the $\Phi$ values of RNTD and NZTD were higher than those of carbamazepine (Wols and Hofman-Caris, 2012), naproxen (Pereira et al., 2007), and 2,4-dichlorophenoxyacetic acid (Chu, 2001), while lower than those of diclofenac (Shu et al., 2013) and sulfamethoxazole (Lian et al., 2015), indicating a moderate UV photolysis rate of RNTD and NZTD. With these rate constants, the apparent transformation kinetics of the RNTD and NZTD can be determined.
for any water/wastewater pH, providing an estimate of baseline removal by UV photolysis. Note that the mineralization of RNTD and NZTD during the UV photolysis may be rather limited as no oxidant was added during the photolysis. Thus, the HNMsFP may increase with the cleavage of chemical bond and TBPs formation, which needs a further HNMsFP evaluation before and after the UV photolysis.

3.3. HNMsFP of RNTD and NZTD

The effect of initial chlorine concentration on the HNMs formation from RNTD and NZTD was firstly investigated. As shown in Fig. 3, the initial chlorine concentration impacts not only the formed HNMs species, but also the HNMs formation amounts. DCNM was the dominated HNMs species formed when the molar ratio of [Chlorine]/[RNTD or NZTD] ≤ 10. However, TCNM became the dominated formed HNMs species with the increase of chlorine concentration. With the chlorine concentration increased from 0.05 to 1.0 mM, the TCNM concentration increased from 4.6 ± 0.33 to 92.0 ± 3.1 µg L⁻¹ for RNTD and NZTD, respectively. Thus, a 0.5 mM chlorine concentration was adopted in the HNMsFP test based on these results and previous studies (Bond et al., 2014; Wang et al., 2016).

pH is another important factor governing the formation of DBPs by influencing on HOCl/OCl⁻ species, halogenation and hydrolysis reactions during chlorination. As shown in Fig. 4, the solution pH significantly affected the HNMs formation amounts. The highest TCNM formation was found under acidic and neutral circumstances, which is different from the previous results (Joo and Mitch, 2007; Chiang et al., 2010). There are no obvious differences among the HNM yields at pH 5.0–7.0, while the HNMsFP formation amounts decreased in alkaline circumstance with the minimum value of 9.2 and 4.0 µg L⁻¹ at pH 9.0 for RNTD and NZTD, respectively. The pKa of RNTD, NZTD and HOCl were 8.7, 6.8, and 7.5, respectively (Degim et al., 2001; Tian et al., 2014). As RNTD and NZTD are prone precursors of HNMs during chlorination, it is speculated that the both the neutral and anionic species of RNTD and NZTD contributed to the HNMs formation. However, the fraction of active chlorine species at different pH seems to determine the HNMs formation amount (Hansen et al., 2012). Meanwhile, the stability of HNMs, are subject to be influenced by pH. Joo and Mitch (2007) reported that TCNM was stable at pH 5.0, whereas it decomposed rapidly at pH 9.0. Moreover, the precursors of HNMs were also unstable under alkaline circumstances, which may restrict HNMsFP formation during chlorination (Yang et al., 2010).

Based on the results above, a HNMsFP test method was developed (i.e., [chlorine]₀ = 0.5 mM, pH 7.0, 25 °C, and after 24 h) to investigate the variation of HNMsFP during the UV photolysis of RNTD and NZTD. It was found that the molar yields of HNMs from RNTD and NZTD at pH 7.0 were 5.6 ± 0.3% and 4.7 ± 0.4%, respectively under this method. Wang et al. (2016) reported a comparable result for the conversion of RNTD and NZTD to HNMs (5.8 ± 0.6% for RNTD and 5.0 ± 1.1% for NZTD). Although both the chemical structures of RNTD and NZTD contain two secondary amine groups (i.e., one primary amine group and one nitro group), the molar yield of HNMs from the chlorination of amine group was rather low (<0.04%, 0.01%), probably due to the weak oxidation ability of chlorine and the slow transformation of amine group into nitro group (Bond et al., 2014). Thus, the nitro group in RNTD and NZTD is speculated to the prone functional group for HNMs formation during the chlorination.
3.4. Effect of UV photolysis on HNMsFP

The role of UV photolysis on the HNMsFPs of RNTD and NZTD were further investigated by measuring the residual HNMsFP during their degradation at pH 7.0. As shown in Fig. 5, the sum of HNMsFP including MCNM, DCNM and TCNM decreased linearly with the decreasing RNTD and NZTD concentrations ($R^2 > 0.97$). The HNMsFP from RNTD and NZTD were significantly reduced by UV photolysis and a more effective reduction of HNMsFP from RNTD was observed than those of NZTD under the same UV fluence. For example, the HNMsFP from RNTD were reduced to $<5\%$, while the remained HNMsFP from NZTD remained 40% of the initial formation after 30 min, which was consistent with the degradation kinetics of RNTD and NZTD during the UV photolysis.

The regression results indicated that the mother structure of RNTD and NZTD played an important role for the HNMs formation during chlorination. Once the mother structure of RNTD and NZTD was destroyed by UV photolysis, the HNMsFP decreased simultaneously. As mentioned above, the nitrogen source of HNMs during chlorination of RNTD and NZTD was inferred to the nitro group of RNTD and NZTD. Thus, it was speculated that the attack on the nitro group of RNTD and NZTD by UV photolysis induced the reduction of HNMsFP and other denitrated TBPs have negligible contributions to HNMs formation compared with RNTD and NZTD. To verify this speculation, the formed anions during the UV photolysis of RNTD and NZTD were investigated and obvious NO$_2^-$ and NO$_3^-$ formations were observed. As shown in Fig. 6, the NO$_2^-$ and NO$_3^-$ concentrations increased quickly following the degradation of RNTD and NZTD in the initial stages of photolysis (0–10 min) and remained at constant level at later phase. During the photolysis of...
UV photolysis could be produced from C (McMurry, 2008). Therefore, the main TBPs of RNTD and NZTD by significantly lower than that of NO2. The third pathway may involve the cleavage of C–S bond. Then the TBP R4 and TBP N4 may undergo the similar homolytic and heterolytic denitration pathways, with the TBP R6, TBP N6, NO3, NO2 formations subsequently (Fang et al., 2013). Note that the inter-transformation between NO3 and NO2 may occur during the UV photolysis. Complementary experiments with the tested UV fluence in this study were conducted and it was found that both the concentrations of NO3 and NO2 kept stable during the UV photolysis (data not shown). Overall, the denitration occurred among all the three proposed pathways, which contributed to the simultaneous reduction of HNMsFP during the photolysis of RNTD and NZTD.

3.5. Degradation pathways of RNTD and NZTD during UV photolysis

The UV photolysis could induce the transformation of compound via photoelimination, photoreduction, photooxidation, and photodehydrogenation (Fang et al., 2013). The dissociation energies of C–N and C–S bonds are 306 and 272 kJ mol$^{-1}$, respectively, significantly lower than those of C–H, N–H and C–C bonds (McMurry, 2008). Therefore, the main TBPs of RNTD and NZTD by UV photolysis could be produced from C–N and C–S bond fissions (Table S1). Combined with TBPs analysis, three photolytic pathways of RNTD and NZTD are proposed in Fig. 7A and Fig. 7B, respectively. The first pathway was the homolytic cleavage of C–N bond without deprotonation, which generated the methyl radical and NO2 radical. Then the intermediate underwent acid-catalyzed hydrolysis to form TBP R1 and TBP N1 and the NO2 radical was released as NO3 (Phillips et al., 2005; Cole et al., 2007). The second pathway was the heterolytic cleavage of C–N bond, which produced TBP R2 and TBP N2 and NO3 ion as the result of a nucleophile attack in aqueous solution, which was facilitated by the nucleophilic attack of a water molecule in the nitro group. As mentioned above, the NO3 formation derived from the oxidation of NO2 even if the formation amount is not obvious. Stefan and Bolton (2002) also observed NO3 was generated along with the NO2 during the UV photolysis of N-Nitrosodimethylamine, but at a lower level than NO2. The third pathway may involve the cleavage of C–S bond. Then the TBP R4 and TBP N4 may undergo the similar homolytic and heterolytic denitration pathways, with the TBP R6, TBP N6, NO3, NO2 formations subsequently (Fang et al., 2013). Note that the inter-transformation between NO3 and NO2 may occur during the UV photolysis.

3.6. Effect of water matrix

To investigate the effect of UV photolysis on the HNMsFP of RNTD and NZTD in real waters, three real water samples (including FW, RW and WE with their water qualities shown in Table S2) were spiked with 10 $\mu$M RNTD or NZTD. Prior to irradiation, the pH of the samples was adjusted to 7.0 to allow direct comparison with control group (ultrapure water). As expected, the degradation of RNTD was much faster than that of NZTD in the real water. Also, all the degradation rate constants of the RNTD and NZTD in the real water samples were lower than those in the ultrapure water followed the descending order of FW > RW > WE (data not shown). The reduced degradation rates of RNTD and NZTD in the real water samples should be primarily attributable to the presence of natural organic matter via acting as an inner filter to reduce the effective fluence rate, and consequently the rate of any photochemical reaction (Latch et al., 2003; Wammer et al., 2013). As shown in Fig. 8, the HNMsFP reductions in real waters by UV photolysis were also lower to those in the ultrapure water with a reduction of the HNMsFP by 55.6–86.0% (FW > RW > WE).

UV-based processes are increasingly adopted not only for disinfection of water and wastewater, but also for the treatment of emerging pollutants which can be further configured to generate reactive species through photo chemical and photo catalytic reactions. This study determined the fundamental parameters for the UV photolysis of RNTD and NZTD (including $\epsilon$, $k$, and $\Phi$), which can be extended to other UV systems to predict degradation kinetics. In China, the UV fluences for disinfection of wastewater, drinking water and reclaimed water are regulated as 20, 40 and 80 $\text{mJ cm}^{-2}$, respectively (GB/T 19837–2005, UV Disinfection Equipment for Municipal Water and Wastewater Treatment). At a typical pH of 7.0, the removal efficiencies of RNTD and NZTD in these three waters during UV disinfection were estimated to be $<70\%$. Obviously, all the mandatory disinfection UV fluences in different waters cannot achieve the effective abatement of RNTD, NZTD and their HNMsFP, highlighting the necessity of increasing UV fluence or developing UV-based advanced oxidation process in future.

4. Conclusions

Based on the experimental results, the following conclusions can be drawn:

- Both the RNTD and NZTD are photo-reactive under UV irradiation and exhibit pH-dependent degradation rates at pH 5.0–9.0, in accordance to their deprotonation equilibria.
The specific $k_4$ were obtained with a minimum for the anionic NZTD (5.64 m$^2$ E$^{-1}$) and a maximum for the neutral NZTD (31.90 m$^2$ E$^{-1}$).

- Both the RNTD and NZTD are prone precursors of HNMs with molar yields of 5.6±0.3 and 4.7±0.4%, respectively at pH 7.0 and acidic and neutral circumstances facilitated the HNMs formation.

- The UV photolysis of RNTD and NZTD could reduce their HNMsFP simultaneously and denitrification accounted for this reduction.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.watres.2017.04.049.

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