Photooxidation of atrazine and its influence on disinfection byproducts formation during post-chlorination: effect of solution pH and mechanism

Yucan Liu1, Kai Zhu2*, Huayu Zhu3, Min Zhao2, Lihua Huang2, Bin Dong2 & Qianjin Liu2*

Partial photooxidation of micropollutants may lead to various degradation intermediates, obviously affecting disinfection byproducts (DBPs) formation during the post-chlorination process. The photooxidation of atrazine (ATZ) in aqueous solutions with low-pressure mercury UV lamps in UV, UV/H2O2 and UV/TiO2 treatment system and the formation of chlorinated disinfection byproducts (DBPs) during subsequent chlorination processes including dichloroacetic acid (DCAA), trichloroacetic acid (TCAA), 1,1,1-trichloro-2-propanone (TCP), trichloromethane (TCM) and chloropicrin (CHP) were investigated in this study. The effect of solution pH on the oxidation pathway of ATZ in three UV photooxidation treatment process and the impact of photooxidation on the DBPs formations were assessed. Based on UPLC-ESI–MS/MS analyses, identification of main oxidation intermediates was performed and the plausible degradation pathways of ATZ in photooxidation system were proposed, indicating that photooxidation of ATZ in UV/H2O2 and UV/TiO2 process system was significantly pH-dependent processes. Dichloroacetic acid (DCAA), trichloroacetic acid (TCAA), 1,1,1-trichloro-2-propanone (TCP), trichloromethane (TCM) and chloropicrin (CHP) were detected in photooxidized ATZ solutions. Compared to the other three DBPs, TCM and TCP were the main DBPs formed. The DBPs formations were greatly promoted in oxidized ATZ solutions. Solution pH and UV irradiation time exhibited obvious impact on the DBPs formation on the basis of DBP species. The variation tendency of DBPs observed relates to the combustion of ATZ in photooxidation system and the production oxidation intermediates.

Pesticides have been widely used for pests control and kill broadleaf weeds in agriculture, forestry and animal husbandry, owing to the proper biological toxicity and long half-life1. Due to the enormous application amounts and special molecular structures, pesticides can be transported to natural waters easily and has been detected in rivers, lakes, underground waters and sediments frequently2–4.

Atrazine (2-chloro-4-ethylamino-6-isopropylamino-s-triazine, ATZ), the most widely used pesticides for the control of weeds due to its excellent performance, is commonly detected in natural waters5. ATZ is also considered as an environmental hormone that induces the complete feminization, posing potential adverse effect to human health. Furthermore, trace of ATZ residue (μg/L or ng/L) in water can cause carcinogenic, teratogenic and mutagenic effect6. Because of the poor efficiency of conventional water treatment processes in removing the ATZ residue, this contaminant can react with chlorine during disinfection process forming various disinfection byproducts (DBPs)7. Therefore, ATZ would be an important precursor of DBPs during drinking water treatment and it is meaningful to assess the potential of ATZ to produce DBPs.

Recently, chemical pre-oxidation methods including photolysis and advanced oxidation processes (AOPs) have been applied to drinking water treatment with a lot of trace organic pollutants8. Apart from physical methods, chemical oxidation of organics includes the cleavage chemical bonds and damage of molecular structures.

1School of Civil Engineering, Yantai University, Yantai 264005, China. 2Shandong Provincial Key Laboratory of Water and Soil Conservation and Environmental Protection, College of Resources and Environment, Linyi University, Linyi 276000, China. 3School of Chemistry and Chemical Engineering, Linyi University, Linyi 276000, China. *email: zhukai@lyu.edu.cn; liuqianjin@lyu.edu.cn
Among these chemical pre-oxidation methods, photooxidation is considered as the most environmentally friendly and safe technology. The most popular photooxidation technologies are UV radiation, UV radiation with H₂O₂ (UV/H₂O₂) and UV radiation with TiO₂ (UV/TiO₂).9–12 The simplest photooxidation technology is UV radiation, the degradation of organics during sole-UV process is induced by the absorption of UV photons and direct photolysis.13 When UV radiation is conducted along with H₂O₂ or TiO₂, the processes are called UV-AOPs13,14. The UV-AOPs with H₂O₂ or TiO₂ in solution originate the production of hydroxyl radicals (·OH), the most important radical in AOPs, in the bulk. This free radical, with standard reduction potential of 2.8 V/SHE, is the second strongest oxidizing agent after fluorine.15 The generation of ·OH in solution promotes the oxidation rate of organic contaminants, producing large amounts of intermediates.

However, complete mineralization of organic contaminants to water and carbon dioxide generally demands relatively long reaction time and more chemicals consumption.16 It is unprocurable to obtain complete mineralization of organics in actual water treatment process owing to the limits of reaction time and water treatment cost. Hence, oxidation products may be present in water prior to the chlorination reaction. These oxidation intermediates may be easier to react with chlorine than the precursor themselves during chlorination process.17–20 Surprisingly, relevant papers focus on this topic are lacking, particularly regarding the role of solution pH and oxidation intermediates. Despite the level of individual organic contaminant is very low, the total contributions of contaminants to the formation of DBPs may not be ignorable.

The object of present study is to evaluate and analyze the effect of photolysis intermediates on the formation of DBPs from UV, UV/H₂O₂ or UV/TiO₂ oxidized ATZ aqueous solutions following chlorination. The formation of dichloroacetic acid (DCAA), trichloroacetic acid (TCAA), 1,1,1-trichloro-2-propanone (TCP), trichloromethane (TCM) and chloropicrin (CHP) during chlorination process is analyzed by ultra-performance liquid chromatography-electrospray ionization mode-triple quadruple mass spectrometry (UPLC-ESI–MS/MS) and a 7890A gas chromatograph fitted with a 7000A triple quadrupole mass spectrometer (GC-QqQ-MS/MS). The effects of solution pH during photooxidation processes on DBPs formation were also evaluated.

Materials and methods

Materials. ATZ (> 97%) was provided by TCI (Shanghai) Development Co., Ltd. (Shanghai, China) and used without further purification. Sulfuric acid (guarantee reagent), sodium hydroxide (analytical reagent), ascorbic acid (analytical reagent) and sodium hypochlorite (5% effective chlorine) were purchased from Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). The following standard solutions, HANs standard solution US EPA 511B (CHP and TCP, 2000 μg/mL of each in acetone), HAAs mixed standard solution (DCAA and TCAA, 2000 μg/mL of each in MTBE) and THMs mixed standard solution (TCM, 2000 μg/mL in acetone), were obtained from Sigma-Aldrich Corporation (Bellefonte, PA, USA). Hydrogen peroxide (30%, w/w) and TiO₂ (P-25, mainly in anatase form) used in photooxidation process were supplied by Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China) and Evonik Degussa Co. (Dusseldorf, Germany), respectively. Ultrapure water (18.2 MΩ·cm) from Elga Purelab Ultra system (Bucks, UK) was used to prepare all working solutions.

Photooxidation experiments. An annular vessel, which had been studied in our previous research21, was used as photochemical reactor. The photon flux into the working solution from the low-pressure mercury UV lamp was 1.18 × 10⁻⁷ Einstein/s, detected with an iodode–iodate chemical actinometer. Before the oxidation reaction, 300 mL ATZ solution (5 mg/L) was transfused into the photochemical reactor, using a magnetic stirrer to maintain reaction solution homogeneity. During photooxidation process, the solution temperature was kept at 20 ± 0.5 °C with a thermostatic water recirculation system. In order to acquire stable output, the UV lamp was ignited for 30 min before photooxidation experiments. UV/H₂O₂ oxidation was conducted with H₂O₂ added into the working solution at 5 mL/L, and UV/TiO₂ oxidation was performed with Degussa P25 TiO₂ powder added into the working solution at 5 mg/L. Solution pH was maintained at 4.0, 7.0 and 10.0, respectively, with 2 mM phosphate and/or borate buffers. Samples were withdrawn during photooxidation process at certain intervals to analyze the oxidation dynamics and mechanism of ATZ.

Chlorination processes. In chlorination experiments after photooxidation, the pH value of working solutions was first adjusted to 7.0 using 1 M sulfuric acid or 1 M sodium hydroxide. After pH adjustment, a certain amount of chlorine was added into solutions to obtain free chlorine residual concentration of 1.0 ± 0.5 mg/L after 24 h chlorination. Stirred for 30 s after addition of chlorine, 50 mL ATZ photooxidation solution was injected into an amber glass flask coupled with stopper, 40 mL ATZ photooxidation solution was transferred into head-space-free amber glass bottle fitted with caps and PTFE-lined septa. These flasks and bottles were kept in dark at 25 °C for 24 h. Ascorbic acid solution (100 g/L, 10 μL) was injected into the solution to annihilate residual chlorine to prevent further chlorination reaction. The chlorine demand was calculated from Eq. (1). All the experiments were conducted in triplicate, and average values and standard deviations were reported.

\[
\text{Chlorine demand} = \text{Cl}_0 - \text{Cl}_{24}
\]

where Cl₀ and Cl₂₄ were the chlorine concentrations at initial time and reaction 24 h, respectively.

Analytic methods. The concentration of ATZ and its oxidation products were detected using UPLC-ESI–MS/MS (Waters Corporation, Milford, MA, USA) equipped with an ACQUITY UPLC BEH C₁₈ column (2.1 mm × 100 mm, 1.7 μm particle). The detailed information of operation parameters is presented in Text S1 in the Supplementary materials. The extent of ATZ removal was calculated from Eq. (2).
where ATZ₀ and ATZₜ were the detected ATZ concentrations at initial time and reaction time t, respectively. The concentrations of DCAA and TCAA were measured with UPLC-ESI–MS/MS system based on our previous research with detailed information of operation parameters presented in Text S2 and Supplementary Table S1. The formations of TCP, TCM and CHP during chlorination process were detected by GC–QqQ-MS/MS (Agilent Technologies, Palo Alto, CA, USA) in the multiple reaction monitoring mode with specific operation parameters in Text S3 and Supplementary Table S2.

Results and discussion

Effects of solution pH on ATZ removal. Different initial pH values in the range of 4–10 were chosen to evaluate the impact of solution pH on ATZ removal. The photooxidation of ATZ at different solution pH values during UV, UV/H₂O₂, and UV/TiO₂ process is shown in Fig. 1.

After 240 min irradiation, the removal efficiency of ATZ during UV process reached 77.5%, 96.5% and 91.8% with solution pH values as 4.0, 7.0 and 10.0, respectively. Obviously, the removal efficiency of ATZ increased sharply with the increase of solution pH value from 4.0 to 7.0. Nevertheless, when the solution pH value further increased from 7.0 to 10.0, the removal efficiency of ATZ exhibited slight drop. The effect of solution pH on the photooxidation of ATZ is mainly due to the distribution of pH-dependent species, protonated and deprotonated ATZ species. As solution pH increased from 4.0 to 7.0, ATZ molecules mainly exist in negative forms, which increase the electron density of s-triazine ring and thus promoted the indirect photodegradation of ATZ. Further increased the solution pH to 10.0, the UV–Vis absorbance spectra of ATZ showed significant decreasing trend, blocking the absorption of photons. During UV/H₂O₂ process, the removal efficiency of ATZ exhibited analogous tendency, attaining 79.5%, 98.5% and 93.7% after 240 min treatment under solution pH values of 4.0, 7.0 and 10.0, respectively. The oxidation of ATZ in UV/H₂O₂ system was due to the synthetic action of ·OH generated by photolysis of H₂O₂ and direct photolysis. The oxidation power of ·OH decreased as the increase of solution pH, however, in neutral condition, the photolysis rate of ATZ is the highest. The results also indicated that addition of H₂O₂ during UV irradiation promoted the degradation of ATZ due to the formation of ·OH in the bulk.

During the initial stage of UV/TiO₂ process, the removal efficiency of ATZ decreased significantly as solution pH increased from 4.0 to 10.0. The pHₚz of TiO₂ is reported in the range of 6.3–6.9 by several papers. The surface of TiO₂ particle would be negatively charged when pH > pHₚz, positively charged when pH < pHₚz. Meanwhile, the distribution of protonated and deprotonated ATZ species varied with solution pH, and ATZ molecules had a negative charge in acidic condition. The ATZ molecules were easily adsorbed to the surface of TiO₂ particles in acidic condition in UV/TiO₂ process. In order to confirm that the decay of ATZ in UV/TiO₂ process was caused by the photooxidation only, adsorption experiment was carried out in dark using 5 mg/L TiO₂ as adsorbent in 300 mL solution with ATZ concentration at 5 mg/L. No significant signs of adsorption were observed, indicating that TiO₂ adsorption play no distinct role during photooxidation of ATZ in UV/TiO₂ process (see Supplementary Fig. S1).

Supplementary Figure S2 shows the semi-log graph of the ATZ versus UV photooxidation time with different solution pH. Supplementary Tables S3–S5 exhibits the relevant apparent rate constant k and correlation coefficient R². These figures and tables confirmed that the photooxidation of ATZ at different pH values during UV, UV/H₂O₂, and UV/TiO₂ process followed the pseudo-first-order kinetics. Solution pH had obvious significance on the k value, the k values were 0.00779, 0.01545 and 0.01308 min⁻¹ in sole-UV system with initial solution pH at 4, 7 and 10, respectively. The k values were 0.02042, 0.01765 and 0.01289 min⁻¹ in UV/H₂O₂ system with initial solution pH at 4, 7 and 10, respectively. The k values were 0.00842, 0.01703 and 0.01253 min⁻¹ in UV/TiO₂ system with initial solution pH at 4, 7 and 10, respectively. The above-mentioned tendency can be attributed to the oxidation effect of ·OH generated in UV/H₂O₂ and UV/TiO₂ process. With the increase of solution pH, ATZ molecules mainly exist in negative forms, which increase the electron density of s-triazine ring and thus promoted the indirect photodegradation of ATZ in solution. On the other hand, the generation of ·OH generated in UV/H₂O₂ and UV/TiO₂ process was strikingly affected by the solution pH.

Photooxidation intermediates and plausible mechanism. The primary intermediates of ATZ during photooxidation process were detected and identified by comparing the total ion chromatograms (TICs) of ATZ solutions before and after UV, UV/H₂O₂ and UV/TiO₂ process treatment.

Photooxidation intermediates and degradation pathway in UV process. The extracted ion chromatograms (EICs) of ATZ solutions after UV treatment are exhibited in Fig. 2. As can be seen, the influence of solution pH on ATZ degradation mechanism is negligible during UV treatment, since the EICs of ATZ solutions at pH of 4.0, 7.0 and 10.0 are almost the same. ATZ and its ten photolysis intermediates were detected, and for clear description, these peaks were labeled as P1, P2, P3, P4, P5, P6, P7, P8, P9 (ATZ), P10 and P11. MS and MS/MS spectra analyses under ESI⁺ and ESI⁻ mode were performed for identifying the molecular structures of these intermediates. The retention times and MS spectral information of the mentioned 11 intermediates are listed in Supplementary Table S6. Molecular weight (MW) of the 11 intermediates, deduced from the m/z, were used to analyze the molecular structures. Furthermore, in order to acquire further information of these intermediates, collision induced dissociation (CID) experiments were performed. Argon was used as collision gas and the collision energy of each intermediate was optimized. The precursor ions of each intermediate under daughter scan mode are shown in Supplementary Table S7, and the particular information of these fragments...
Figure 1. Effect of solution pH on ATZ removal: (a) UV process; (b) UV/H₂O₂ process; (c) UV/TiO₂ process.
during MS/MS analysis are shown in Supplementary Figs. S3–S12. Molecular structure of these intermediates were deduced on the basis of MS spectra and MS/MS spectra.

In the case of P1, only detected under ESI+ mode, the \( m/z \) of main fragment in MS spectra was 154. Under ESI+ mode, \( H^+ \) added to P1 produced the positive ion, indicating the MW of P1 was 153 Da. In addition, four fragment ions (\( m/z \) 112, 85, 70 and 68) were detected in the MS/MS spectra as shown in Supplementary Fig. S3. P1 (\( m/z \) 154) lost –CH(CH\(_3\))\(_2\) (\( m/z \) 42) produced the fragment ion of \( m/z \) 112, and fragment ion of \( m/z \) 85 lost –NH\(_2\) generated fragment ion of \( m/z \) 70. MS/MS spectra indicated the s-triazine ring of P1 attached –NHCH(CH\(_3\))\(_2\) and –NH\(_2\).

The MW of P2 (\( m/z \) 184), detected under ESI+ mode, was 183 Da. P2 lost –(CH\(_3\))\(_2\) generated fragment ion of \( m/z \) 156, lost –CH(CH\(_3\))\(_2\) generated fragment ion of \( m/z \) 142, and lost –CH(CH\(_3\))\(_2\), –NHCH\(_3\) and –OH generated fragment ion of \( m/z \) 97, indicated the molecular structure of P2 included –NHCH(CH\(_3\))\(_2\) and –NHCH\(_2\).

The precursor ion of P3 (\( m/z \) 196) produced five fragment ions (\( m/z \) 145, 97, 89, 71 and 65). Based on MS, MS/MS spectra and previous researches\(^{29,30}\), the possible molecular structure of P3 was concluded.

P4 was detected under both ESI+ and ESI− modes. P4 lost –CH(CH\(_3\))\(_2\) generated fragment ion of \( m/z \) 156, lost –CH(CH\(_3\))\(_2\) and –CH\(_2\)CH\(_3\) generated fragment ion of \( m/z \) 128, and lost –NHCH(CH\(_3\))\(_2\), –CH\(_2\)CH\(_3\) and –OH generated fragment ion of \( m/z \) 97 under ESI+ mode. Under ESI− mode, P4 lost –(CH\(_3\))\(_2\) generated fragment ion of \( m/z \) 168, lost –CH(CH\(_3\))\(_2\) generated fragment ion of \( m/z \) 154, lost –NHCH(CH\(_3\))\(_2\), and –CH\(_2\)CH\(_3\) generated fragment ion of \( m/z \) 125, and lost –NHCH(CH\(_3\))\(_2\) and –CH\(_2\)CH\(_3\) generated fragment ion of \( m/z \) 111. Based on above analysis, the molecular of P4 is shown in Supplementary Table S8.

The precursor ions of P5 and P6 were detected under both ESI+ and ESI− modes with the same \( m/z \). For P5, the fragment ion of \( m/z \) 156 was produced by the loss of –COCH\(_3\), the fragment ion of \( m/z \) 139 was produced by the loss of –NHCOCH\(_3\) and the fragment ion of \( m/z \) 113 was produced by the loss of –NHCOCH\(_3\) and –CH\(_2\)CH\(_3\). For P6, the fragment ion of \( m/z \) 156 was produced by the loss of –CHCH\(_2\) and –OH, the fragment ion of \( m/z \) 153 was produced by –CH\(_3\) instead of –NHCHOCH\(_3\) in this position, the fragment ion of \( m/z \) 127 was produced by the loss of –NHCHCH\(_2\), –CH\(_3\) and –OH, and the fragment ion of \( m/z \) 113 was produced by the loss of –NHCHCH\(_2\) and –CHOHCH\(_3\).

P7 and P8 had same molecular ions \( m/z \) 212 under ESI+ mode, indicating the MWs of P7 and P8 were 211. For P7, the fragment ion of \( m/z \) 170 was produced by the loss of –CH(CH\(_3\))\(_2\), the fragment ion of \( m/z \) 128 was produced by the loss of –CH(CH\(_3\))\(_2\), and –COCH\(_3\). The MS/MS spectra of P8 suggested the fragment ion of \( m/z \) 182 was produced by the loss of –OCH\(_3\), the fragment ion of \( m/z \) 170 was produced by the loss of –CH(CH\(_3\))\(_2\), the fragment ion of \( m/z \) 142 was produced by the loss of –CH(CH\(_3\))\(_2\) and –CH\(_2\)CH\(_3\), the fragment ion of \( m/z \) 128 was produced by the loss of –CH(CH\(_3\))\(_2\), –CH\(_2\)CH\(_3\) and –OH.

---

**Figure 2.** The extracted ion chromatograms (EICs) of ATZ solution after 90 min irradiation in UV process: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
P9 was identified as ATZ by the TIC of ATZ solution before UV oxidation. P10 was only detected under ESI+ mode and P11 was detected under both ESI+ and ESI− modes. For P10, two fragment ions (m/z 81 and 72) were detected in the MS/MS spectra as shown in Supplementary Fig. S12. For P11, under ESI+ mode, four fragment ions (198(M + H), 220(M + Na), 395(2 M + H) and 417(2 M + Na)) were detected in the MS/MS spectra, 196(M-H) was detected under ESI− mode. On the basis of the MS/MS spectra and previous study, the molecular structures of P10 and P11 were deduced and presented in Supplementary Table S8.

Based on these identified photooxidation intermediates, the main reactions of ATZ in UV irradiation treatment were proposed. The dominating reaction is dechlorination–hydroxylation, replacing the Cl atom with −OH. The product of dechlorination–hydroxylation reaction included P2, P3, P4, P5, P6 and P7. Dechlorination–dealkylation reaction, including the cleavage of C–Cl, C–C and C–N bonds, produced P2. Dealkylation reaction, depriving alkyl groups from ATZ generated P1 and P10. Deamination reaction, including the removal of lateral chains connected to the −NH2, produced P10. Alkylic-oxidation, resulting in the formation of carbon radical compound, produced P5 and P7. Dehydrogenation–olefination, abstracted H from lateral chains of ATZ by reactive radicals, produced P3 and P6. Dechlorination–hydrogenation, resemblance to dechlorination–hydroxylation, generated P1 and P10. Dechlorination–methoxylation, initiated by small number of methanol from reserving solution of ATZ, generated P8 and P11. Dehydroxylation, containing the removal of hydroxyl from s-triazine and photooxidation products, formed P1. The plausible degradation pathway of ATZ in aqueous solution during direct UV irradiation process was proposed based on above analysis as shown in Fig. 3.

Photooxidation intermediates and degradation pathway in UV/H2O2 process. The EICs of ATZ solutions after UV/H2O2 oxidation are exhibited in Fig. 4. The main photooxidation intermediates of ATZ were found to vary with solution pH. In the case of pH 4.0, ATZ and its fourteen oxidation intermediates were identified, including P1–P10, P11, P12, P14, P15 and P16. At pH 7.0, ATZ and fifteen oxidation intermediates were detected (P1–P16), including a new appeared peak (P13). As for pH 10.0, eighteen peaks were extracted, peaks of P14, P15 and P16 disappeared and five new peaks appeared (P17, P18, P19, P20 and P21). The precursor ions of P12–P21 under daughter scan mode are shown in Supplementary Table S9, and the particular information of these fragments during MS/MS analysis are shown in Supplementary Table S10 and Supplementary Figs. S13–S22. The proposed structure and chemical name of P12–P21 are shown in Supplementary Table S11.
For P12, detected under ESI+ mode, the MS spectra indicated its MW was 209 Da. Four fragment ions (m/z 168, 152, 123 and 115) were detected in the MS/MS spectra as shown in Supplementary Fig. S13. P12 (m/z 220) lost –COCH₃ produced the fragment ion of m/z 168, and fragment ion of m/z 168 lost –OH generated fragment ion of m/z 152. MS/MS spectra indicated the s-triazine ring of P12 attached –OH, –NH(C=CH₂)CH₃ and –NH₂COCH₃. P13 only detected under ESI+ mode, with MW of 138 Da. P13 lost –COCH₃ generated fragment ion of m/z 97, indicated the molecular structure of P13 included –NH₂COCH₃.

P14 and P15 had same molecular ions m/z 212 under ESI+ mode, indicating the MWs of P14 and P15 were 211. For P14, the fragment ion of m/z 184 was produced by the loss of –(CH₃)₂, the fragment ion of m/z 170 was produced by the loss of –CH=CH₂ and the fragment ion of m/z 142 was produced by the loss of –(CH₃)₂ and –CH=CH₂. The MS/MS spectra of P15 suggested the fragment ion of m/z 170 was produced by the loss of –CH₃, –CH₂ and –CH₂, the fragment ion of m/z 103 and 86 were produced by the cleavage of s-triazine ring.

P16, P17 and P18 were detected under ESI+ mode with the same m/z 214, indicating the MWs of P16, P17 and P18 were 211. For P16, the fragment ion of m/z 173 was produced by the loss of –NHCH=CH₂. The MS/MS spectra of P17 suggested the fragment ion of m/z 196 was produced by the loss of –OH, the fragment ion of m/z 170 was produced by the loss of –CH₃, –CH₂ and –OH, the fragment ion of m/z 143 was produced by the loss of –CH₃ and –NHCH(CH₃)₂ and the fragment ion of m/z 129 was produced by the cleavage of s-triazine ring.

Figure 4. The extracted ion chromatograms (EICs) of ATZ solution after 90 min irradiation in UV/H₂O₂ process: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
P19 was detected under ESI+ modes with MW of 195 Da. P19 generated fragment ion of \( m/z \) 154 by loss of –CH₃ and –CH₂CH₃, and lost –NHC₃H₅ and –CH₃ generated fragment ion of \( m/z \) 127. For P18, P20 and P21, no fragment ions were detected in MS/MS spectra. The molecular structure of P18, P20 and P21 were deduced by MWs and previous researches²¹,³¹.

Based on these identified photooxidation intermediates, the oxidation pathway of ATZ in UV/H₂O₂ process was proposed, as shown in Fig. 5. Apart from these intermediates detected in direct UV irradiation treatment, several new oxidation products (P12–P21) were identified in UV/H₂O₂ oxidation system. These intermediates were probably generated by the attack of ·OH³²–³⁴. Two feasible oxidation modes of ·OH onto ATZ molecular existed in UV/H₂O₂ system, (1) abstraction of hydrogen atom (P12, P14, P15, P16 and P21), (2) the hydroxylation attack (P14, P15, P17, P18 and P20)³⁵,³⁶.

**Figure 5.** The proposed degradation pathway of ATZ in UV/H₂O₂ process.

Photooxidation intermediates and degradation pathway in UV/TiO₂ process. The EICs of ATZ solutions after UV/TiO₂ oxidation are exhibited in Fig. 6. The effect of solution pH on the oxidation pathway of ATZ in UV/TiO₂ process was remarkable. At pH 4.0, thirteen oxidation intermediates were identified, including P1–P8, P11, P18, P22, P25 and P26. However, in case of pH 7.0, ATZ and seventeen oxidation intermediates were detected, including P1–P8, P11, P18, P22, P25, P26 and 4 new appeared peaks (P17, P19, P23 and P24). As for pH 10.0, the number of detected intermediates was changed again, thirteen peaks were extracted. Compared to pH 4.0, peak P19 replaced P22 at pH 10.0. The MS fragment information for the new intermediates of ATZ in UV/TiO₂ process are shown in Supplementary Table S12.

According to previous studies and the MS spectra, the molecular structures of P22–P26 were deduced³⁸,³¹,³⁵, as shown in Supplementary Table S13. Compared to UV and UV/H₂O₂ process, five new photooxidation products of ATZ were identified during UV/TiO₂ oxidation. UV/TiO₂ treatment is an indirect heterogeneous photooxidation process, due to the addition of TiO₂ nanoparticles³⁹. Five steps occurred in heterogeneous system, as stated below: (1) ATZ molecular transferred from the bulk to TiO₂ particle; (2) adsorption of ATZ onto the photon activated TiO₂ surface; (3) photooxidation reaction; (4) desorption of oxidation products from TiO₂ particles surface; (5) transfer of oxidation products from TiO₂ surface to the bulk³⁷,³⁸. Photogenerated holes (\( h^+_{\text{VB}} \)) and ·OH were produced when TiO₂ particles were irradiated by UV light. Furthermore, other free radicals (superoxide radical
and hydroperoxide radical) were produced in UV/TiO₂ system39, increased the types of oxidation reactions of ATZ. On the basis of identified photooxidation intermediates, the oxidation pathway of ATZ in UV/TiO₂ process was proposed, as shown in Fig. 7.

Chlorine demand in post-chlorination. Previous studies focus on DBPs have proved that chlorine demand is an important element related with the formation of DBPs during chlorination process40,41. The chlorine demand of ATZ solution after UV, UV/H₂O₂ and UV/TiO₂ treatment under different pH values are shown in Fig. 8.

Generally, the amount of chlorine consumed in chlorination treatment process increased along with the increase of photooxidation time. The increase of chlorine demand indicated the production of various phototization intermediates aggrandized the reactivity of oxidized solution toward chlorine. In order to confirm the chlorine demand of original ATZ solution, controlled trials were conducted, obtaining chlorine demand of 0.776 ± 0.08 with solution pH value as 7.0. In UV process, the chlorine demand increased with UV fluence, as shown in Fig. 8a. After 300 min irradiation, the chlorine demand reached 4.997 ± 0.16, 4.491 ± 0.11 and 4.121 ± 0.14 mg/L with photooxidation pH value at 4.0, 7.0 and 10.0, respectively. Further increase of UV irradiation time to 780 min, the increasing tendency of chlorine demand became slow, except the photooxidation pH of 4.0. Although chlorination process consumed the most chlorine under photooxidation pH of 4.0, the amounts of chlorine consumed under different photooxidation pH values were close. In UV/H₂O₂ process, the amount of chlorine consumed in chlorination process firstly increased sharply with the increase of irradiation time from 0 to 540 min, as shown in Fig. 8b. After 540 min irradiation, the chlorine demand reached 4.636 ± 0.21, 7.325 ± 0.18

Figure 6. The extracted ion chromatograms (EICs) of ATZ solution after 90 min irradiation in UV/TiO₂ process: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
and 10.003 ± 0.15 mg/L with photooxidation pH value at 4.0, 7.0 and 10.0, respectively. Further increase of photooxidation time to 780 min, the consumed chlorine exhibited slight downward trend. The influence of solution pH during photooxidation on chlorine demand was remarkable, the amount of chlorine consumed was the highest under photooxidation pH of 10.0. In UV/TiO₂ process, the chlorine demand increased rapidly with the increase of irradiation time from 0 to 300 min, as shown in Fig. 8c. After 300 min irradiation, the chlorine demand reached 4.491 ± 0.11, 5.645 ± 0.13 and 6.402 ± 0.14 mg/L with photooxidation pH value at 4.0, 7.0 and 10.0, respectively. Further increase of UV irradiation made unconspicuous impact on chlorine demand. The key parameter that had an important effect on the chlorine demand was the solution pH during photooxidation, the amount of chlorine consumed was the highest as photooxidation pH of 10.0 under different photooxidation time.

In comparison, the amount of chlorine consumed by ATZ solution after UV/H₂O₂ treatment was the highest than direct UV photolysis and UV/TiO₂ oxidation, especially under photooxidation pH of 10.0 after prolonged irradiation. The increase of chlorine demand depended on the production of specific oxidation intermediates. As for UV/H₂O₂ oxidation under solution pH 10.0, seventeen oxidation intermediates were identified including P20 and P21 only detected in this condition.

**Formation of DBPs.** Five kinds of DBPs, including DCAA, TCAA, TCP, TCM and CHP, were detected after 24 h chlorination (under solution pH of 7.0) of the UV, UV/H₂O₂ and UV/TiO₂ oxidized ATZ solution. In order to evaluate the effects of irradiation time and photooxidation pH on the formation of DBPs, raw ATZ solutions were chlorinated under solution pH of 7.0. The concentrations of DCAA, TCAA, TCP and TCM reached 0.425 ± 0.015, 0.533 ± 0.01, 0.114 ± 0.01 and 0.678 ± 0.105 μg/L, respectively, after 24 h chlorination. Meanwhile, CHP was undetected in the samples.

**Formation of DBPs after UV treatment.** As shown in Fig. 9, irradiation time and photooxidation pH made distinct influences on the formation of DBPs. Generally, TCM dominated the DBPs generation and direct photolysis significantly changed DBPs formation and speciation. The effect of irradiation time in UV photolysis system under solution pH of 4.0 on the formation of DBPs is depicted in Fig. 9a. The formation of TCM and TCP during chlorination process increased sharply with the increase of irradiation time. The concentration of DCAA and CHP increased slowly with prolonged UV oxidation. However, the concentration of TCAA increased slightly at the early stage of reaction time, then decreased as the irradiation time exceeded 210 min. The effects of irradiation time in UV photolysis system under solution pH of 7.0 and 10.0 on the formation of DBPs is depicted in Fig. 9b, c, respectively. The concentrations of TCM and TCP under photooxidation pH of 7.0 and 10.0 declined significantly compared to the output under pH 4.0. Under photooxidation pH of 7.0, the concentration of DBPs basically remains stable after 210 min irradiation, except TCP. However, under photooxidation pH of 10.0, the concentrations of CHP, TCP, DCAA and TCAA exhibited downward trend as the irradiation time increased.

![Figure 7. The proposed degradation pathway of ATZ in UV/TiO₂ process.](https://www.nature.com/scientificreports/)
Figure 8. The chlorine demand of oxidized ATZ solutions after 24 h chlorination, (a) UV process; (b) UV/H$_2$O$_2$ process; (c) UV/TiO$_2$ process.
Figure 9. DBPs formations of UV oxidized ATZ solutions: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
Figure 10. DBPs formations of UV/H$_2$O$_2$ oxidized ATZ solutions: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
Figure 11. DBPs formations of UV/TiO$_2$ oxidized ATZ solutions: (a) pH = 4.0; (b) pH = 7.0; (c) pH = 10.0.
It is worth noting that CHP was undetected during chlorination of raw ATZ solution. This phenomenon indicated that ATZ did not react with chlorine to produced CHP in the bulk. The variation tendency of DBPs observed might have been related to the oxidation of ATZ by direct UV photolysis and the production oxidation intermediates. It has been reported that direct photolysis of organics could break down the hydrophobic parts, forming hydrophilic and polar intermediates, and increased DBPs formation potentials.

**Formation of DBPs after UV/H2O2 treatment.** The effects of irradiation time and UV/H2O2 oxidation pH on the formation of DBPs are depicted in Fig. 10. Compared to other DBPs, much more TCM and TCP were formed in UV/H2O2 treated ATZ solutions. Under photooxidation pH of 4.0, the TCM and TCP formations were greatly increased with increase of irradiation time at initial reaction stage, as shown in Fig. 10a. After 210 min oxidation in UV/H2O2 system, the concentration of TCAA began to decrease. Furthermore, the formation of TCP was suppressed after longer oxidation. Figure 10b shows the formations of DBPs in UV/H2O2 treated ATZ solutions under photooxidation pH of 7.0. The influence of the irradiation time on the formation of DBPs was also remarkable, for the destruction of ATZ during photooxidation process improved the reactivity of solution toward chloride. TCM concentration increased rapidly, reaching its maximum level after 540 min photooxidation. The formation of TCP increased with increase of irradiation time, on the contrary, the formations of TCAA and CHP were suppressed after longer oxidation. The formations of DBPs in UV/H2O2 treated ATZ solutions under photooxidation pH of 10.0 is shown in Fig. 10c. The formations of TCM and TCP were vastly promoted by UV/H2O2 oxidation at the entire stage of reaction time. However, the concentrations of TCAA, DCAA and CHP in photo-oxidized solutions only increased within early stage of reaction time.

At pH 4.0, fourteen oxidation intermediates were identified, furthermore, fifteen and seventeen intermediates were detected at pH 7.0 and 10.0, respectively. The variation tendency of DBPs in the UV/H2O2 oxidized ATZ solutions under different pH values during chlorination process might due to the diverse species and concentrations of ATZ intermediates.

**Formation of DBPs after UV/TiO2 treatment.** The concentrations of DBPs formed under present experimental conditions are shown in Fig. 11. Without UV/TiO2 oxidation, only small amounts of DBPs were formed. However, the formations of DBPs were vastly promoted in the UV/TiO2 treated ATZ solutions during 24 h chlorination, and TCM and TCP exhibited the highest concentrations compared to DCAA, TCAA and CHP. Figure 11a shows the formations of DBPs in UV/TiO2 treated ATZ solutions under photooxidation pH of 4.0. The formations of TCM and TCAA were greatly promoted at the early stage of reaction time, then decreased as the oxidation reaction continues. Figure 11b, c show the formations of DBPs in UV/TiO2 treated ATZ solutions under photooxidation pH of 7.0 and 10.0, respectively, and the formation trend of DBPs were similar to the case of pH 4.0. The similar tendency of DBPs in UV/TiO2 oxidized ATZ solutions under different pH values during chlorination process might due to the similar main intermediates of ATZ (P1–P8, P11, P18).

**Conclusion**

This study investigated the effect of solution pH on the oxidation pathway of ATZ in UV, UV/H2O2, and UV/TiO2 oxidation system and the impact of photooxidation on the DBPs formation of ATZ solution during post-chlorination was followed. The structures of the main photooxidation intermediates were deduced on the basis of MS and MS/MS spectra, which showed that the photooxidation of ATZ in UV/H2O2 and UV/TiO2 system was significantly pH-dependent processes. The plausible degradation pathways of ATZ in photooxidation systems were proposed. The photooxidation pH and irradiation time had distinctly impact on the DBPs formation in oxidized ATZ solutions during post-chlorination. The formations of DBPs were enormously promoted in the early stage of photooxidation reaction time, TCM and TCP were the main DBPs formed. The increased DBPs concentrations in oxidized ATZ solutions might have been related to the combustion of ATZ and the production of oxidation intermediates. The observed phenomena in this study indicated that the DBPs formation in micropollutants containing water after pre-oxidation might be promoted. This tendency should be seriously evaluated and appropriately resolved when photooxidation processes are applied to water treatment.

Received: 29 May 2020; Accepted: 22 October 2020
Published online: 23 November 2020

**References**

1. Singh, A. et al. Advances in controlled release pesticide formulations: Prospects to safer integrated pest management and sustainable agriculture. *J. Hazard. Mater.* 385, 121525.1-121525.20 (2020).
2. Glinski, D. A., Purucker, S. T., Van Meter, R. J., Black, M. C. & Henderson, W. M. Henderson, Analysis of pesticides in surface water, stemflow, and throughfall in an agricultural area in South Georgia, USA. *Chemosphere* 209, 496–507 (2018).
3. Sjerps, R. M. A., Kooij, P. J. F., van Loon, A. & Van Wezel, A. P. Occurrence of pesticides in Dutch drinking water sources. *Chemosphere* 235, 510–518 (2019).
4. Reentsma, T., Alder, L. & Banasiak, U. Emerging pesticide metabolites in groundwater and surface water as determined by the application of a multimethod for 150 pesticide metabolites. *Water Res.* 47(15), 5535–5545 (2013).
5. Chen, N. et al. Transfer and degradation of the common pesticide atrazine through the unsaturated zone of the Chalk aquifer (Northern France). *Environ. Pollut.* 255, 113125.1-113125.10 (2019).
6. Pérez-Iglesias, J. M., Franco-Belussi, L., Natale, G. S. & de Oliveira, C. Biomarkers at different levels of organisation after atrazine formulation (SIPTRAN 500SC) exposure in *Rhinella schneideri* (Anura: Bufonidae) Neotropical tadpoles. *Environ. Pollut.* 244, 733–746 (2019).
7. Wang, D. et al. Strong promoted catalytic ozonation of atrazine at low temperature using tourmaline as catalyst: Influencing factors, reaction mechanisms and pathways. *Chem. Eng. J.* 354, 113–125 (2018).
8. Li, K. et al. Effect of pre-oxidation on low pressure membrane (LPM) for water and wastewater treatment: A review. Chemosphere 231, 287–300 (2019).
9. Zhang, N., Hu, K. & Shan, B. Ballast water treatment using UV/TiO₂ advanced oxidation processes: An approach to invasive species prevention. Chem. Eng. J. 243, 7–13 (2014).
10. Zhang, Y., Xiao, Y., Zhong, Y. & Lim, T. Comparison of amoxicillin photodegradation in the UV/H₂O₂ and UV/permanganate systems: Reactivity, kinetics, degradation pathways, and antibacterial activity. Chem. Eng. J. 372, 420–428 (2019).
11. Liu, Z. et al. Pilot-scale UV/H₂O₂ process for drinking water treatment—analysis and comparison of different activated carbon columns. Chem. Eng. J. 382, 123044.1-123044.9 (2020).
12. Wang, Y., Li, Y., Yi, P. & Zhang, H. Degradation of clofibric acid by UV, O₃ and UV/O₃ processes: Performance comparison and degradation pathways. J. Hazard. Mater. 379, 120771.1-120771.11 (2019).
13. Riga, A., Soutsas, K., Ntampegiotis, K., Karayannis, V. & Papapolymerou, G. Effect of system parameters and of inorganic salts on the decolorization and degradation of Procion H-ex dyes. Comparison of H₂O₂/UV/Penton, Fenton, Peroxone, TiO₂/UV and TiO₂/UV/H₂O₂ processes. Desalination 211, 72–86 (2007).
14. Datcu, A. et al. UV–vis light induced photocatalytic activity of TiO₂/graphene oxide nanocomposite coatings. Catal. Today 321–322, 81–86 (2019).
15. Brillas, E. & Sirés, I. & Oturan, M. A.,. Electro–Fenton process and related electrochemical technologies based on Fenton's reaction chemistry. Chem. Res. 109(12), 6570–6631 (2009).
16. Chen, H., Chen, C. & Wang, G. Performance evaluation of the UV/H₂O₂ process on selected nitrogenous organic compounds: Reductions of organic contents vs. corresponding C-, N-DBPs formations. Chemosphere 85(4), 591–597 (2011).
17. Zhang, M., Xu, B., Wang, Z., Zhang, T. & Gao, N. Formation of iodinated trihalomethanes after ferrate pre-oxidation during chlorination and chloramination of iodine-containing water. J. Taiwan Inst. Chem. E. 60, 453–459 (2016).
18. Ao, X. et al. The impact of UV treatment on microbial control and DBPs formation in full-scale drinking water systems in northern China. J. Environ. Sci. China 87, 398–410 (2020).
19. Hua, Z. et al. DBP alteration from NOM and model compounds after UV/UVs treatment with post chlorination. Water Res. 158, 237–245 (2019).
20. Sun, J. et al. The influence of the UV/chlorine advanced oxidation of natural organic matter for micropollutant degradation on the formation of DBPs and toxicity during post-chlorination. Chemosphere 137, 870–879 (2019).
21. Liu, Y. et al. Influence of solution pH on degradation of atrazine during UV and UV/H₂O₂ oxidation: Kinetics, mechanism, and degradation pathways. RSC Adv. 9(61), 35847–35861 (2019).
22. Liu, Y., Duan, J., Li, W., Beecham, S. & Mulcahy, D. Effects of organic matter removal from wastewater secondary effluent by aluminum sulfate coagulation on haloacetic acids formation. Environ. Sci. Technol. 33(7), 484–493 (2016).
23. Liu, Y. et al. Determination of volatile disinfection byproducts in water by gas chromatography–triple quadrupole mass spectrometry. Anal. Lett. 48(1), 188–203 (2014).
24. Ku, Y., Chang, J. & Cheng, S. Effect of solution pH on the hydrolysis and photolysis of diazinon in aqueous solution. Water Air Soil Pollut. 108(3–4), 445–456 (1998).
25. Li, C., Zhang, D., Peng, J. & Li, X. The effect of pH, nitrate, iron (III) and bicarbonate on photodegradation of oxytetracycline in aqueous solution. J. Photoch. Photobiol. A 356, 239–247 (2018).
26. Buxton, G. V., Greenstock, C. L., Helman, W. P. & Ross, A. B. Critical review of rate constants for reactions of hydrated electrons, hydrogen atoms and hydroxyl radicals (·OH/·O⁻) in aqueous solution. J. Phys. Chem. Ref. Data 17(2), 513–886 (1988).
27. Uyguner, C. S., Suphankad, S. A., Kerc, A. & Bekbolet, M. Evaluation of adsorption and coagulation characteristics of humic acids preceded by alternative advanced oxidation techniques. Desalination 210(1–3), 183–193 (2007).
28. Shankar, M. V., Cheralathan, K. K., Arabindo, B., Palanichamy, M. & Marugesan, V. Enhanced photocatalytic activity for the destruction of monocrotophos pesticide by TiO₂/H₂O₂. J. Mol. Catal. A Chem. 233(1–2), 195–200 (2004).
29. Chen, C. Photolytic destruction of endocrine disruptor atrazine in aqueous solution under UV irradiation: Products and pathways. J. Hazard. Mater. 172(1–2), 675–684 (2009).
30. Imoberdorf, G. & Mohseni, M. Kinetic study and modeling of the vacuum–UV photoinduced degradation of 2,4-T. Chem. Eng. J. 187, 114–122 (2012).
31. Khan, J. A. et al. Kinetic and mechanism investigation on the photochemical degradation of atrazine with activated H₂O₂, S₂O₈²⁻ and HSO₅⁻. Chem. Eng. J. 252, 393–403 (2014).
32. Acosta-Rangel, A., Sánchez-Polo, M., Polo, A. M. S., Rivera-Utrilla, J. & Barber-Mendoza, M. S. Sulfonamides degradation assisted by UV, UV/H₂O₂ and UV/K₂S₂O₈: Efficiency, mechanism and byproducts cytotoxicity. J. Environ. Manag. 225, 224–231 (2018).
33. Liu, T. et al. The role of reactive oxygen species and carbonate radical in oxcarbazepine degradation via UV, UV/H₂O₂: Kinetics, mechanisms and toxicity evaluation. Water Res. 147, 204–213 (2018).
34. Yang, Y. et al. Degradation of sulfamethoxazole by UV, UV/H₂O₂ and UV/peracetic acid (PDS): Formation of oxidation products and effect of bicarbonate. Water Res. 118, 196–207 (2017).
35. Shi, H. et al. Mechanism investigation on the enhanced and selective photoelectrochemical oxidation of atrazine on molecular imprinted mesoporous TiO₂. Appl. Catal. B Environ. 246, 50–60 (2019).
36. Al-Mamun, M. R., Kader, S., Islam, M. S. & Khan, M. Z. H. Photocatalytic activity improvement and application of UV-TiO₂ photocatalysis in textile wastewater treatment: A review. J. Environ. Chem. Eng. 7(5), 103248.1-103248.17 (2019).
37. Fujishima, A., Zhang, X. & Tryk, D. A. Heterogeneous photocatalysis: From water photolysis to applications in environmental cleanup. Int. J. Hydrog. Energy 32(14), 2664–2672 (2007).
38. Konstantinou, I. K. & Albanis, T. A. TiO₂-assisted photocatalytic degradation of azo dyes in aqueous solution: Kinetic and mechanistic investigations. A review. Appl. Catal. B Environ. 49(1), 1–14 (2004).
39. Bagwasi, S., Tian, B., Zhang, J. & Nasir, M. Synthesis, characterization and application of bismuth and boron Co-doped TiO₂: A visible light active photocatalyst. Chem. Eng. J. 217, 108–118 (2013).
40. Hu, C., Li, A., Lin, Y., Ling, X. & Cheng, M. Degradation kinetics and DBP formation during chlorination of metribuzin. J. Taiwan Inst. Chem. E. 80, 255–261 (2017).
41. Wang, H. et al. Formation of DBPs during chlorination of antibiotics and control with permanganate/bisulfite pretreatment. Chem. Eng. J. 392, 123701.1-123701.11 (2019).
42. Chu, W., Gao, N., Yin, D., Krasner, S. W. & Mitch, W. A. Impact of UV/H₂O₂ pre-oxidation on the formation of haloacetamides and other nitroguanine disinfection byproducts during chlorination. Environ. Sci. Technol. 48(20), 12190–12198 (2014).
43. Mao, Y. et al. Effects of conventional ozonation and electro–peroxozone pretreatment of surface water on disinfection by-product formation during subsequent chlorination. Water Res. 130, 322–332 (2017).
44. Dotson, A. D., Keen, V. O. S., Metz, D. & Linden, K. G. UV/H₂O₂ treatment of drinking water increases post-chlorination DBP formation. Water Res. 44(12), 3703–3713 (2010).
45. Sarathy, S. & Mohseni, M. Effects of UV/H₂O₂ advanced oxidation on chemical characteristics and chlorine reactivity of surface water natural organic matter. Water Res. 44(14), 4087–4096 (2010).
Acknowledgements
This work was financially supported by the Natural Science Foundation of Shandong Province (No. ZR2017BEE016), Science Fund of Yantai University (No. TM17B19) and Project of Introducing and Cultivating Young Talent in the Universities of Shandong Province. The authors gratefully acknowledge the support of the three foundations.

Author contributions
Y.L., K.Z., H.Z. and Q.L. participated in the study design. Y.L., L.H. and K.Z. performed the experiments. Y.L. and K.Z. completed the analysis. Y.L. and K.Z. drafted the manuscript. K.Z. and Q.L. supervised the analysis and critically revised the manuscript. Y.L. and Q.L. provided research supervision and funding. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-77006-0.

Correspondence and requests for materials should be addressed to K.Z. or Q.L.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020