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Photoinduced transformation of pyridinium-based ionic liquids, and implications for their photochemical behavior in surface waters

Paola Calza⁽¹⁾, Giorgio Noè⁽¹⁾, Debora Fabbri⁽¹⁾, Valentina Santoro⁽²⁾, Claudio Minero⁽¹⁾, Davide Vione* ⁽¹⁾, Claudio Medana⁽²⁾

(1) Department of Chemistry, University of Torino, via P. Giuria 5, 10125 Torino, Italy

(2) Department of Molecular Biotechnology and Health Sciences, University of Torino, via P. Giuria 5, 10125

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* Corresponding author. E-mail: davide.vione@unito.it

10 Abstract

The photochemical reactivity of three ionic liquids (1-ethylpyridinium tetrafluoroborate, 1-butyl-4methylpyridinium tetrafluoroborate, and 1-(3-cyanopropyl)pyridinium chloride) was studied by combining laboratory experiments and photochemical modeling, to get insight into the possible behavior in surfacewater environments. Among the studied compounds, phototransformation in sunlit surface waters could be an important attenuation pathway for 1-butyl-4-methylpyridinium tetrafluoroborate (BMPOTFB). In this case the reaction with the carbonate radicals (CO₃^{-•}) would prevail at low values of the dissolved organic carbon (DOC), while the direct photolysis would be important at intermediate to high DOC values. The sensitization by the triplet states of chromophoric dissolved organic matter could play a significant role at high DOC, especially in the presence of a considerable fraction of highly photoreactive pedogenic organic matter derived from soil runoff. The main processes that account for the phototransformation of BMPOTFB and produce the main detected transformation products (TPs) are hydroxylation, detachment/shortening of the butyl chain and double bond formation. Interestingly, there is a considerable overlap between the intermediates formed by direct photolysis and those produced by indirect photochemistry. Some of the TPs formed from BMPOTFB are more toxic than the parent compound towards Vibrio fischeri bacteria, and account for the increase in toxicity of the irradiated mixtures. Differently from BMPOTFB, in the case of the other two studied ionic liquids the photodegradation would play a negligible role in environmental attenuation, with the possible exception of very shallow waters with low DOC.

Keywords: pollutants in surface waters; environmental attenuation of pollution; sunlight-induced reactions; direct and indirect photochemistry.

1. Introduction

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Ionic liquids (ILs) have several interesting properties such as good thermal stability, wide electrochemical potential window, high electric conductivity and miscibility with water or organic solvents, which permit application mainly as solvents for organic synthesis and as electrolytes (Plechkova and Seddon, 2008; Holbrey and Seddon, 1999). Room-temperature ILs have received huge attention as green and high-tech reaction media (Stepnowski and Zaleska, 2005) because, differently from conventional organic solvents, they have negligible vapor pressure and do not emit toxic vapors (Kralisch et al 2005). However, a broad implementation of the use of ILs in industry may have some drawbacks. For instance, due to their high stability, significant amounts of ILs are expected to occur in natural waters and soil in the near future (Bubalo et al. 2014; Czerwicka et al. 2009; Richardson and Ternes, 2014). Surface water is a major environmental compartment in which ILs may be emitted as industrial wastes, and the most hydrophilic of these compounds should preferentially be found in aqueous phases. Therefore, information concerning the fate of ILs in the aquatic environment is crucial. Some studies have shown that ILs are more toxic to cells than conventional solvents (Stepnowski and Zaleska, 2005), and the most common imidazolium-based compounds also showed high resistance against microbial degradation (Czerwicka et al, 2009). Photochemistry is a potentially important attenuation route for ILs in surface waters (Calza et al., 2015). Generally speaking, phototransformation in surface waters can be divided into direct photolysis (a xenobiotic absorbs sunlight, which triggers its degradation) and indirect photochemistry. In the latter, sunlight is absorbed by naturally-occurring photosensitizers (most notably nitrate, nitrite and chromophoric dissolved organic matter, CDOM) to produce reactive transient species (*OH, ¹O₂ and CDOM triplet states, ³CDOM*) (Dong et al., 2015; Rosario-Ortiz and Canonica, 2016). More precisely, nitrate and nitrite yield OH, while irradiated CDOM produces ³CDOM*, which can form ¹O₂ by reaction with oxygen (McNeill and Canonica, 2016).

Irradiated CDOM yields OH as well (Lee et al., 2013; Page et al., 2014), at least partially through a photo-Fenton process (Chiwa et al., 2015; Mostofa and Sakugawa, 2016; Giannakis et al., 2016a/b). An additional transient, the carbonate radical CO₃•-, is produced by oxidation of carbonate and bicarbonate by OH and of carbonate by CDOM* (Canonica et al., 2005; Bahnmuller et al., 2014; Janssen et al., 2014; Trivella et al., 2015; Bintou et al., 2015). These transients are then involved into the phototransformation of xenobiotics (Silva et al., 2015), including the xenobiotics that do not undergo direct photolysis (Janssen et al., 2015). Dissolved organic matter (DOM, not necessarily chromophoric), which is usually quantified as the dissolved organic carbon (DOC), is the main sink of both OH and CO₃-• (Huang and Mabury, 2000; Canonica et al., 2005; Keen et al., 2014). This issue, combined with the fact that CDOM is a major radiation absorber, explains why •OH and CO₃-• are usually more concentrated in low-DOC waters. In contrast, the highest levels of ³CDOM* and ¹O₂ usually occur in high-DOC environments, because organic matter is involved in the production of these transients but not in their scavenging. Actually, the quenching of ³CDOM* occurs by reaction with O₂ or interval conversion, while ¹O₂ is quenched by collision with the water solvent (Wenk et al., 2013; Vione et al., 2014). Within this framework, we studied the photofate of three pyridinium-based ILs under conditions that are relevant to sunlit environmental waters: 1-ethylpyridinium tetrafluoroborate (EPTFB), 1-(3cyanopropyl)pyridinium chloride (CPPC) and 1-butyl-4-methylpyridinium tetrafluoroborate (BMPOTFB). The known physical properties of the studied ILs (Zhao et al., 2003; Bandres et al., 2008; Guerrero et al., 2012) are summarized in Table SM1 in the Supplementary Material (hereafter SM). We paid attention to the identification of the intermediate products formed during ILs degradation, by using the HPLC-MS/MS technique. The identification of transformation products (TPs) is a crucial aspect because, in addition to providing important information on the mechanism of degradation, the formed TPs may have a very different impact on the environment

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compared to the parent molecules. To our knowledge, this is the first study about the photochemical transformation of EPTFB, CPPC and BMPOTFB.

2. Experimental section

2.1. Materials and Reagents

1-Ethylpyridinium tetrafluoroborate (**EPTFB**) (98%), 1-(3-cyanopropyl)pyridinium chloride (**CPPC**) (98.5%), 1-butyl-4-methylpyridinium tetrafluoroborate (**BMPOTFB**) (≥97,0%), sodium nitrate (≥99.9%), sodium bicarbonate (≥99,7%), acetonitrile (≥99%), acetaminophen (APAP), antraquinone-2-sulfonic acid, sodium salt (98%), hydrogen peroxide (35%), formic acid (98%), sodium 1-hexanesulfonate (98%), and phosphoric acid (85%) were purchased from Sigma Aldrich, Italy. Rose Bengal (80%) was purchased from Alfa Aesar. All aqueous solutions were prepared with ultra-pure water (Millipore Milli-QTM).

2.2. Irradiation Experiments

Irradiation experiments were performed in magnetically stirred cylindrical closed cells (Pyrex glass, $40 \text{ mm} \text{ i.d.} \times 25 \text{ mm}$), on 5 mL aqueous solutions containing each studied IL at $20 \text{ }\mu\text{M}$ initial concentration. The goal of these experiments was not to directly reproduce environmental conditions in the laboratory (which is indeed problematic, in particular as far as the water depth is concerned), but rather to obtain reasonably accurate measurements of the main photochemical parameters such as direct photolysis quantum yields and second-order reaction rate constants. These data, used as input values for a photochemical model, allow inferences to be made on the photochemical fate of the studied compounds in environmental waters. Direct photolysis experiments were carried out on each IL taken separately, under UVB irradiation with a Philips TL 20 W/01 RS lamp having emission maximum at 313 nm. The lamp radiation reached the irradiated solutions mainly from the top.

Indirect photolysis experiments followed the competition kinetics approach (Puskas et al., 2005; Dantas et al., 2007; Katsoyiannis et al., 2011; Sun et al., 2016). They were run on systems containing a mixture of all the studied ILs and acetaminophen (APAP), each at 20 µM initial concentration. The rationale for the use of APAP as reference compound is that its second-order reaction rate constants are known for all the transient species relevant to this study (•OH, CO₃-•, ¹O₂ and ³CDOM*; De Laurentiis et al., 2014). An important issue in these competition kinetics experiments is the need to minimize the occurrence of multiple reaction pathways, which are problematic for the assessment of the reaction rate constants. For the competition approach to work, one needs a single prevailing reaction pathway to be operational. Therefore, the radical *OH was produced by irradiating 100 mM NaNO₃ under the TL 01 RS lamp described above, which was also used to produce CO₃-• upon irradiation of 100 mM NaNO₃ + 100 mM NaHCO₃. In both cases, a high concentration of nitrate was used to minimize the IL direct photolysis. Elevated bicarbonate was needed to scavenge the majority of photogenerated •OH, thereby minimizing the interference of •OH on the CO₃-• reactions. Anthraquinone-2-sulfonate (AQ2S) was used as CDOM proxy to study the reactivity of ³CDOM*. The use of AQ2S has pros and cons: on the one side it does not yield •OH or ¹O₂ upon irradiation, which allows for the study of a pure triplet-sensitized process (Maddigapu et al., 2010). In the case of atrazine, a remarkably good agreement has also been found between the reaction rate constants with ³AQ2S* and with natural ³CDOM* (Marchetti et al, 2013; Zeng and Arnold, 2013). However, the triplet state ³AQ2S* may be more reactive than average ³CDOM*, which could lead to an overestimation of the triplet-sensitization rate constants when using AQ2S as CDOM proxy (De Laurentiis et al., 2014; Avetta et al., 2016). The latter issue should be (and was here) taken into account when using kinetic data derived from AQ2S photochemistry. In the AQ2S experiments, solutions containing 0.2 mM AQ2S and 20 µM ILs/APAP mixtures were irradiated under a Philips TLK 05 lamp, with emission maximum at 365 nm. Finally, measures of reactivity with ¹O₂ were performed using a lamp Philips TL D 18W/16

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with emission maximum at 545 nm. The dye Rose Bengal (10 µM initial concentration) was chosen as the ${}^{1}\text{O}_{2}$ source. The irradiation conditions already described were also used for the study of IL transformation products (TPs) *via* the aforementioned photochemical pathways. In this case, however, each IL was irradiated separately and at a relatively high loading (20 mg L⁻¹), to simplify TPs detection and identification.

2.3. Analytical Procedures

- 135 *2.3.1. Liquid Chromatography UV detection (HPLC-UV)*
- After the scheduled irradiation times the cells where withdrawn from the lamp, and their content 136 was analyzed by HPLC-UV to monitor the time evolution of the studied ILs and, when relevant, of 137 APAP as well. The instrument used was a Merck-Hitachi HPLC equipped with AS-2000A 138 autosampler (100 µL injection volume), L-6200 and L-6200A pumps for high-pressure gradients, 139 L-4200 UV-Vis detector and a reverse-phase column Phenomenex Kinetex® 100Å C18 (250 mm × 140 4.6 mm × 5 µm). The eluent consisted of an aqueous solution of the ionic coupler sodium 1-141 hexanesulfonate 0.01 M (eluent A) and acetonitrile (eluent B). After 3 min of isocratic conditions 142 (90/10 v/v A/B), the acetonitrile fraction was increased up to 60/40 v/v in 19 min. For the tests with 143 the dye Rose Bengal these conditions were followed by a second gradient up to 40/60 v/v A/B over 144 145 4 min, which was held for 5 min to allow elution of the photosensitizer. Re-equilibration of the 146 HPLC column was achieved by going back to the starting conditions, which were held for 5 min 147 before the next injection. The mobile phase flow rate was 1 mL/min and the detection wavelength was 210 nm. With these elution conditions the retention times were 5.3 min for APAP, 6.2 min for 148 EPTFB, 6.8 min for CPPC, and 14.6 min for BMPOTFB. 149
- 150 2.3.2. Liquid Chromatography High Resolution Mass Spectrometry (HPLC-HRMS)
- This technique was used to detect and identify the TPs of the studied ILs under irradiation. For HPLC-HRMS analyses of the irradiated aqueous solutions, the chromatographic separations were carried out with a Phenomenex C18 Gemini® NX 110-Å column, 150 mm \times 2.1 mm \times 3 μ m

particle size (Phenomenex, Bologna, Italy), using a Ultimate 3000 HPLC instrument (Dionex, 154 Thermo Scientific, Milan, Italy). The injection volume was 20 µL and the flow rate 200 µL/min. A 155 gradient mobile phase composition was used, going from 5/95 v/v of acetonitrile/ 156 heptafluorobutyric acid (5 mM in water) to 20/80 v/v in 15 min, followed by a second gradient step 157 up to 95/5 v/v in 5 min. 158 A LTQ Orbitrap mass spectrometer (Thermo Scientific, Milan, Italy) was interfaced to the HPLC 159 instrument through an ESI ion source. The HPLC column effluent was delivered into the ion source 160 using nitrogen as both sheath and auxiliary gas. The capillary voltage and tube lens voltage in the 161 ESI source were maintained at 28 V and 70 V, respectively. The source voltage was set to 4.5 kV in 162 the positive ion mode. The capillary temperature was maintained at 270°C. The acquisition method 163 was optimized beforehand in the tuning sections (capillary, magnetic lenses and collimating 164 165 octapole voltages) for the parent compound, to achieve maximum sensitivity. Mass accuracy of the recorded ions (vs calculated) was ± 5 millimass units (mmu, without internal calibration). 166

The MS spectra were acquired using full scan MS (50-1000 m/z range) and MS² acquisition in the positive ion mode, with a resolution of 30000 (500 m/z FWHM) in FTMS (full transmission) mode. The ions submitted to MS² acquisition were chosen on the basis of full MS spectra abundance, without using automatic dependent scan. Collision energy was set to 30% for all of the MS² acquisition methods, and the MS² acquisition range was between the value of ion trap cut-off and the m/z value of the (M+H)⁺ ion. Xcalibur software (Thermo Scientific, Milan, Italy) was used for both acquisition and data analysis.

2.3.3. Toxicity Measurements

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The toxicity of the reaction mixtures collected at different irradiation times was measured with a Microtox® Model 500 Toxicity Analyzer (Milan, Italy). Acute toxicity was assessed with a bioluminescence inhibition assay using the marine bacterium *Vibrio fischeri*, by monitoring changes in the natural emission of the luminescent bacteria when challenged with toxic compounds. Freeze-

dried bacteria, reconstitution solution, diluent (2% NaCl) and an adjustment solution (non-toxic 22% sodium chloride) were purchased from Azur (Milan, Italy). Samples were tested in five dilutions in a medium containing 2% sodium chloride, and luminescence was recorded after 5, 15, and 30 min of incubation at 15°C. Because no substantial changes in luminescence were observed between 5 and 30 minutes, only the percent toxicity recorded at 15 minutes will be discussed. Inhibition of luminescence, compared with a toxic-free control to give the percentage inhibition, was calculated following the established protocol of the Microtox® calculation program.

2.4. Kinetic data treatment

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The time evolution of the studied ILs was fitted with a pseudo-first order equation of the form 187 $C_t^{IL} = C_o^{IL} e^{-k_{IL}t}$, where C_t^{IL} is the IL concentration at the time t, C_o^{IL} the initial concentration, and k_{IL} 188 the pseudo-first order degradation rate constant. The IL initial degradation rate was calculated as 189 $R_o^{IL} = k_{IL} C_o^{IL}$. The initial degradation rate of APAP (R_o^{APAP}) was calculated in a similar way. 190 With a mixture of APAP and of the studied ILs, under conditions where a reactive species X is 191 generated (with $X = {}^{\bullet}OH$, $CO_3^{-\bullet}$, ${}^{1}O_2$ or ${}^{3}AQ2S^*$), if reaction with X only is operational, the initial 192 rate is $R_o^{IL} = k_{X,IL}[X]C_o^{IL}$ for each relevant IL, and $R_o^{APAP} = k_{X,APAP}[X]C_o^{APAP}$ for APAP. Here, $k_{X,IL}$ 193 and $k_{X,APAP}$ are the respective second-order reaction rate constants of X with each studied IL and 194 with APAP, and [X] is the steady-state concentration of the transient species. Because ILs and 195 APAP occur together in the same mixture, [X] is the same in all the relevant expressions of R_o^{IL} and 196 R_o^{APAP} . Moreover, we chose the initial conditions so that $C_o^{IL} = C_o^{APAP}$. On this basis it is possible to 197 derive $R_o^{IL}(R_o^{APAP})^{-1} = k_{X,IL}(k_{X,APAP})^{-1}$, from which the second-order rate constant of the reaction 198 between each IL and X can be obtained as follows: $k_{X,IL} = k_{X,APAP} R_o^{IL} (R_o^{APAP})^{-1}$ (Herrmann et al., 199 2010). Note that the initial rates are derived from the experimental data, while $k_{X,APAP}$ is known 200 from a previous study (De Laurentiis et al., 2014). 201

For the determination of the direct photolysis quantum yields, the photon flux absorbed by each IL (P_a^{IL}) was calculated as follows: $P_a^{IL} = \int_{\lambda} p^{\circ}(\lambda) [1-10^{-\varepsilon_{IL}(\lambda)b[IL]}] d\lambda$, where $p^{\circ}(\lambda)$ is the incident spectral photon flux density of the used UVB lamp, $\varepsilon_{IL}(\lambda)$ is the IL molar absorption coefficient, b=0.4 cm is the optical path length inside the solution, and [IL] = 20μ L is the initial molar concentration of the relevant IL. The values of $p^{\circ}(\lambda)$ were determined by combining spectral measurements with a calibrated Ocean Optics USB2000 CCD spectrophotometer, and chemical actinometry with 2-nitrobenzaldehyde. The detailed procedure has been reported by Marchisio et al. (2015). The wavelength trends of $p^{\circ}(\lambda)$ and $\varepsilon_{IL}(\lambda)$ are shown in **Figure 1A**. The direct photolysis quantum yield was calculated as $\Phi_{IL} = R_o^{IL}(P_a^{IL})^{-1}$.

2.5. Photochemical modeling

The model assessment of ILs photodegradation was carried out with the APEX software (Aqueous Photochemistry of Environmentally-occurring Xenobiotics), available for free as Electronic Supplementary Information of Bodrato and Vione (2014). APEX predicts photochemical reaction kinetics from photoreactivity parameters of xenobiotics (absorption spectra, direct photolysis quantum yields and second-order reaction rate constants with transient species), and from data of water chemistry and depth (Bodrato and Vione, 2014). APEX predictions have been validated by comparison with field data of pollutant phototransformation kinetics in surface freshwaters (Marchetti et al., 2013; Avetta et al., 2016). The standard solar spectrum used in APEX is referred to fair-weather conditions during summertime at mid latitude (Frank and Klöpffer, 1988). Sunlight irradiance is not constant in the natural environment, due to fluctuations in meteorological conditions (not included in APEX) and to diurnal and seasonal cycles. To allow easier comparison between model results and environmental conditions, taking the day-night cycle into account, APEX uses a summer sunny day (SSD) as time unit, equivalent to fair-weather 15 July at 45° N latitude. The absorption of radiation by the photosensitisers (CDOM, nitrate and nitrite) and the

studied substrates is calculated based on competition for sunlight irradiance in a Lambert-Beer approach (Bodrato and Vione, 2014; Braslavsky, 2007). APEX applies to well-mixed waters and gives average values over the water column, with contributions from the well-illuminated surface layer and from darker water in the lower depths, where irradiance is very low (Loiselle et al., 2008).

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3. Results and Discussion

3.1. Photochemical reactivity of the studied ILs

First of all, the direct photolysis of **EPTFB**, **CPPC** and **BMPOTFB** was studied by irradiating the relevant compounds taken separately (no mixtures were used in this case) under a UVB lamp. The ILs time trends are reported in Figure 1B, which shows that the photochemical degradation rates followed the order **BMPOTFB** > **CPPC** > **EPTFB**. The relevant direct photolysis quantum yields are reported in Table 1. The comparison between quantum yield values (Table 1) and photodegradation kinetics (Figure 1B) suggests that EPTFB was the compound featuring both the lowest quantum yield of direct photolysis (about one order of magnitude smaller than the other two ILs) and the slowest direct phototransformation. Compared with the other studied ILs, EPTFB featured an intermediate degree of UVB radiation absorption (Figure 1A). CPPC was the studied IL with the highest quantum yield of direct photolysis but, because of relatively low radiation absorption, its photodegradation was slower compared to BMPOTFB. The reactivity between the studied ILs and the transient species under consideration ($X = {}^{\bullet}OH$, CO₃-•, ¹O₂ or ³AQ2S*) was studied in mixtures of **EPTFB**, **CPPC**, **BMPOTFB** and APAP (hereafter, substrates). All of the four substrates had the same initial concentration in the mixture ([Substrate] $_0 = 20 \mu M$). The reaction with •OH was studied upon UVB irradiation of the mixture in the presence of 0.10 M NaNO₃, and the reactivity order was **BMPOTFB** > **EPTFB** > **CPPC**. The substrates time evolution is shown in Figure 2A, while the second-order reaction rate constants of the studied ILs with *OH are listed in **Table 1**.

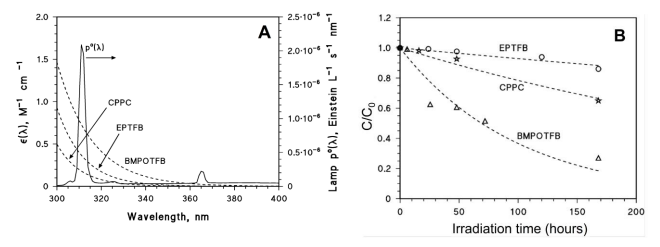


Figure 1. (**A**) Absorption spectra ($\varepsilon(\lambda)$) of the studied ILs, and emission spectrum ($p^{\circ}(\lambda)$) of the used UVB lamp (Philips TL 01). (**B**) Time trends of the studied ILs (each had a 20 μ M initial concentration in a distinct one-component solution) upon UVB irradiation in Milli-Q water (direct photolysis experiments).

Table 1. Photoreactivity parameters (direct photolysis quantum yields and second-order reaction rate constants with photoinduced transients) of the studied ILs and of APAP. For modeling purposes, the range of $k_{IL,^3CDOM^*}$ was varied between 0.05 $k_{IL,^3AQ2S^*}$ and $k_{IL,^3AQ2S^*}$ as explained in the text. Note that Substrate = EPTFB, BMPOTFB, CPPC, or APAP. The APAP data are taken from De Laurentiis et al. (2014). Φ_{APAP} is reported for a sake of completeness, but it was not used to calculate Φ_{IL} . The error bounds, representing the σ-level uncertainty, were taken from the literature in the case of APAP. For the studied ILs, they were derived from duplicate experiments.

	EPTFB	BMPOTFB	CPPC	APAP
$\Phi_{Substrat}$, unitless	$(4.81\pm0.62)\cdot10^{-2}$	$(4.53\pm0.62)\cdot10^{-1}$	$(7.77\pm0.67)\cdot10^{-1}$	$(4.57\pm0.17)\cdot10^{-2}$
$k_{\scriptscriptstyle Substrat\ensuremath{\mathcal{E}}OH},\mathrm{M}^{-1}\;\mathrm{s}^{-1}$	$(3.33\pm2.23)\cdot10^7$	$(2.84\pm1.22)\cdot10^8$	$(1.55\pm0.91)\cdot10^7$	$(1.87\pm0.56)\cdot10^9$
$k_{Substrat \& CO_3^{-\bullet}}, \mathrm{M}^{-1} \; \mathrm{s}^{-1}$	$(7.04\pm5.97)\cdot10^5$	$(4.85\pm2.59)\cdot10^6$	$(1.52\pm0.86)\cdot10^6$	$(3.8\pm1.1)\cdot10^8$
$k_{Substrat \not = O_2}$, ${\rm M}^{-1}~{\rm s}^{-1}$	$(1.40\pm1.30)\cdot10^6$	$(1.50\pm1.10)\cdot10^6$	$(1.30\pm1.10)\cdot10^6$	$(3.68\pm0.73)\cdot10^7$
$k_{Substrat\hat{\ell}AQ2S^*},\mathrm{M}^{-1}\;\mathrm{s}^{-1}$	$(3.71\pm1.17)\cdot10^7$	$(1.99\pm1.18)\cdot10^9$	$(2.64\pm0.99)\cdot10^7$	$(1.08\pm0.16)\cdot10^{10}$

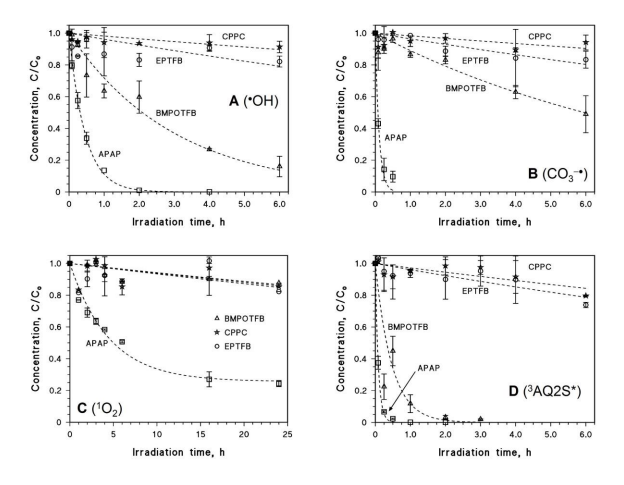


Figure 2. Time trends of APAP and the three studied ILs in the same mixture (20 μM initial concentration for each substrate), in the presence of: (**A**) 0.1 M NaNO₃ under UVB irradiation; (**B**) 0.1 M NaNO₃ + 0.1 M NaHCO₃ under UVB irradiation; (**C**) 10 μM Rose Bengal under yellow-lamp irradiation; (**D**) 0.2 mM AQ2S under UVA irradiation. The error bars represent the standard error of duplicate experiments.

The reaction with CO₃^{-•} was studied upon UVB irradiation of 0.1 M NaNO₃ + 0.1 M NaHCO₃, under which conditions the •OH radicals generated by nitrate photolysis would be largely (>95%) scavenged by the HCO₃⁻ and CO₃²⁻ ions to produce CO₃^{-•}. The •OH scavenging fractions were assessed on the basis of the known •OH reaction rate constants of HCO₃⁻ and CO₃²⁻ (Buxton et al., 1988), of the rate constants of the studied substrates (**Table 1**), and on the initial concentrations in

278 solution. The ILs order of reactivity with CO₃^{-•} was **BMPOTFB** > **EPTFB** > **CPPC**. The relevant time evolutions are shown in Figure 2B, and the second-order reaction rate constants are listed in 279 280 Table 1. The transient ¹O₂ was generated by irradiation of 10 µM Rose Bengal under yellow light. The three 281 ILs showed comparably low reactivity with ¹O₂, much lower than for APAP (see Figure 2C and 282 **Table 1**). Finally, the reactivity by triplet sensitization was studied upon UVA irradiation of 0.2 283 284 mM AQ2S, a concentration value that was chosen to minimize the reaction between excited and ground-state AQ2S (Bedini et al., 2012). Also in this case the reactivity order was BMPOTFB > 285 **EPTFB** > **CPPC** (see **Figure 2D** and **Table 1**). 286 As explained before, ³AQ2S* may be more reactive than natural ³CDOM* and may provide an 287 overestimation of the ³CDOM* rate constants (De Laurentiis et al., 2014; Avetta et al., 2016). For 288 this reason, and based on the results of a recent study (Avetta et al., 2016), the ³AQ2S* rate 289 constants $(k_{IL^3AO2S^*})$ were taken as upper limits for those with $^3\text{CDOM}^*$ $(k_{IL^3CDOM^*})$. The 290 corresponding lower limits were taken as $k_{IL,^3CDOM^*} = 0.05 \ k_{IL,^3AQ2S^*}$. By assuming that $k_{IL,^3CDOM^*}$ 291 varies between 0.05 $k_{IL,^3AQ2S^*}$ and $k_{IL,^3AQ2S^*}$, one gets some insight into the potentially variable 292 reactivity of ³CDOM*. Indeed, triplet sensitisation processes are quite fast when triggered by 293 294 runoff-derived soil organic matter, and they are much slower in the presence of aquagenic CDOM (De Laurentiis et al., 2012). 295 In the above experiments of indirect photochemistry it was generally found that BMPOTFB had a 296 significantly higher reactivity than the other studied ILs, but it was less reactive than APAP. All 297 three ILs have an electron-poor pyridinium ring, which might account for lower reactivity with 298 299 oxidizing transient species when compared to the electron-rich phenolic ring of APAP. However, differently from the other ILs, BMPOTFB also has an electron-donating methyl substituent that 300

could somewhat increase the electron density on the aromatic ring and, therefore, make it more reactive than the other studied ILs.

3.2. Modeling of ILs phototransformation in surface waters

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The photoreactivity parameters reported in **Table 1** were used as input data for the APEX software, using either $k_{IL,^3CDOM^*} = k_{IL,^3AQ2S^*}$ or $k_{IL,^3CDOM^*} = 0.05$ $k_{IL,^3AQ2S^*}$ in different sets of calculations. Because depth and dissolved organic carbon (DOC) are the water body features that most influence photochemical reactions (Vione et al., 2014), the ILs half-life times were computed as a function of these two variables. Reasonable values were assumed for other water parameters of photochemical significance, i.e., 0.1 mM nitrate, 1 µM nitrite, 1 mM bicarbonate, and 10 µM carbonate (Vione et al., 2014). On this basis and for fair-weather summertime irradiation, APEX returned the half-life times reported in Figure 3 (3A,D: BMPOTFB; 3B,E: EPTFB; 3C,F: CPPC; 3A,B,C: $k_{IL,^3CDOM^*} = k_{IL,^3AQ2S^*};$ **3D,E,F**: $k_{IL,^3CDOM^*} = 0.05 k_{IL,^3AQ2S^*}).$ First of all, note that all the half-life times increase with increasing depth and that they also increase with DOC, with the partial exception of a trend with a maximum (**EPTFB** with $k_{EPTFR^3CDOM^*} = k_{EPTFR^3AQ2S^*}$, see **Figure 3B**). The rationale of the depth trend is that the lower depths of deep water bodies are poorly illuminated by sunlight, a phenomenon that offsets the elevated photoreactivity at the water surface. Increasing DOC means increasing DOM and CDOM. The occurrence of CDOM inhibits the direct photolysis (due to competition for irradiance between CDOM and the substrate(s)), while DOM causes an even larger inhibition of the *OH/CO₃-* reactions (due to scavenging of the radicals) (Vione et al., 2014). The inhibition of direct photolysis, of *OH and CO₃-* processes is often not offset by the enhancement of the reactions triggered by ³CDOM* and ¹O₂, of which CDOM is the only source. Actually, CDOM tends to reach absorption saturation in deep waters with high DOC, in which case the production of ³CDOM* is little enhanced by a CDOM increase. In contrast, the scavenging of •OH/CO₃-• and the competition for irradiance do not undergo saturation effects (Avetta et al.,

2016). There are some exceptions, however: if the reactivity of a substrate with ³CDOM*/¹O₂ is remarkably high, these processes can significantly increase the degradation kinetics at elevated DOC. In this case one can observe a maximum in the half-life times as a function of the DOC (Fabbri et al., 2015), which happened here with **EPTFB** if $k_{EPTFB^3CDOM^*} = k_{EPTFB^3AQ2S^*}$ (**Figure 3B**). Among the studied ILs, **BMPOTFB** is predicted to be by far the most photolabile compound in the natural environment. This prediction follows the results of the irradiation experiments, which already suggested that BMPOTFB was more photoreactive than EPTFB or CPPC via both direct photolysis and indirect photochemistry. If $k_{BMPOTFB^3CDOM^*} = k_{BMPOTFB^3AQ2S^*}$, phototransformation would be an important attenuation pathway for BMPOTFB under all the water conditions considered in **Figure 3** (depth = 0-10 m, DOC = 0-12 mg C L⁻¹). If $k_{BMPOTFB^3CDOM^*}$ = $0.05\,k_{{\it BMPOTFB}^3{\it AO2S}^*}$, the photodegradation half-life times would be quite long in deep and high-DOC waters. When considering the half-life time data reported in Figure 3, it should be noted that one cannot have year-long fair-weather summertime conditions in mid-latitude environments. Therefore, the longer lifetimes are certainly underestimated. Differently from BMPOTFB, the predicted photochemical half-life times of EPTFB and CPPC are very high and they can reach years to decades in deep water bodies with high DOC. Under such circumstances, other processes will prevail over phototransformation in the environmental attenuation of these ILs. Photochemistry could still play some role for EPTFB and CPPC, but only in water bodies that are shallow and have low DOC at the same time.

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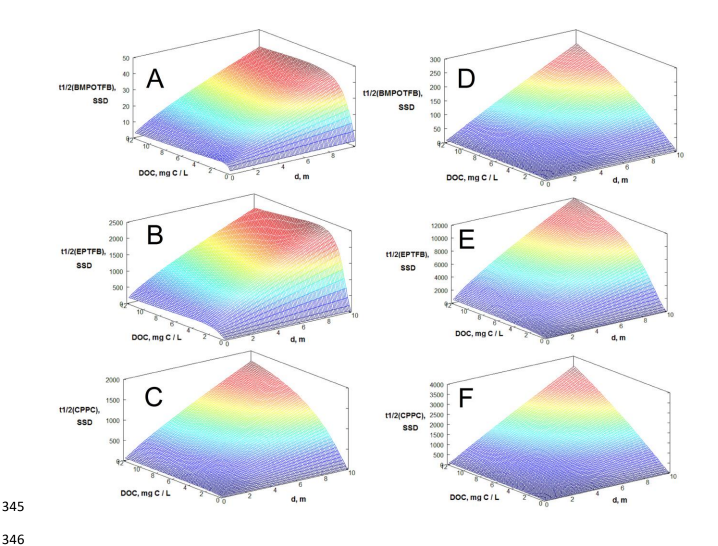


Figure 3. Half-life times predicted by APEX (SSD = summer sunny days equivalent to fair-weather 15 July at 45°N latitude), as a function of the DOC and the water depth d, for: (A) BMPOTFB, with $k_{BMPOTFB^3CDOM^*} = k_{BMPOTFB^3AQ2S^*}$; (B) EPTFB, with $k_{EPTFB^3CDOM^*} = k_{EPTFB^3AQ2S^*}$; (C) CPPC, with $k_{CPPC,^3CDOM^*} = k_{CPPC,^3AQ2S^*}$; (D) BMPOTFB, with $k_{BMPOTFB^3CDOM^*} = 0.05 k_{BMPOTFB^3AQ2S^*}$; (E) EPTFB, with $k_{EPTFB^3CDOM^*} = 0.05 k_{EPTFB^3AQ2S^*}$; (F) CPPC, with $k_{CPPC,^3CDOM^*} = 0.05 k_{CPPC,^3AQ2S^*}$. Other water conditions: 0.1 mM nitrate, 1 μM nitrite, 1 mM bicarbonate, 10 μM carbonate.

Figure 4 reports, as a function of the DOC, the fractions of ILs phototransformation that are accounted for by the different photoreaction pathways, for a water depth of 5 m and other conditions as per the above discussion (4A,D: BMPOTFB; 4B,E: EPTFB; 4C,F: CPPC; 4A,B,C: $k_{IL,^3CDOM^*} = k_{IL,^3AQ2S^*};$ **4D,E,F**: $k_{IL,^3CDOM^*} = 0.05 k_{IL,^3AQ2S^*}).$ The model predicts that the reactions with CO₃^{-•} can be important for all ILs at low DOC (usually below 2 mg C L⁻¹), while •OH and ¹O₂ can be important only in the case of **EPTFB**. The reactions with ³CDOM* can be major pathways at if $k_{IL,^3CDOM^*} = k_{IL,^3AO2S^*}$, while they high DOC become secondary processes if $k_{IL,^3CDOM^*}$ =0.05 $k_{IL,^3AQ2S^*}$. In the latter case, the direct photolysis is expected to strongly dominate the phototransformation of both BMPOTFB and CPPC. By crossing these results with the half-life time data, which show the environmental water conditions where photochemistry can be important in IL attenuation, one gets that: (i) the direct photolysis is very relevant for BMPOTFB, especially in water bodies where the occurrence of CDOM of mainly aquatic origin would limit the triplet-sensitized processes. Other significant **BMPOTFB** transformation pathways are CO₃^{-•} at low DOC and possibly ³CDOM* at high DOC, in the presence of significant amounts of soil-derived organic matter that would enhance the reactions of triplet sensitization (De Laurentiis et al., 2012); (ii) the photodegradation of **EPTFB** may only be significant in low-DOC waters, and in these conditions the prevailing pathways are the direct photolysis and the reactions with ${}^{\bullet}\text{OH}$ and $\text{CO}_3^{-\bullet}$. The exact value of $k_{EPTFR^3CDOM^*}$ is of limited interest because, for the ³CDOM* processes to play a significant role, one needs relatively high-DOC conditions where **EPTFB** photochemistry cannot be important; (iii) a similar issue holds for CPPC, although in this case the *OH process can be neglected and the only important low-DOC pathways are the direct photolysis and the CO₃^{-•} reaction.

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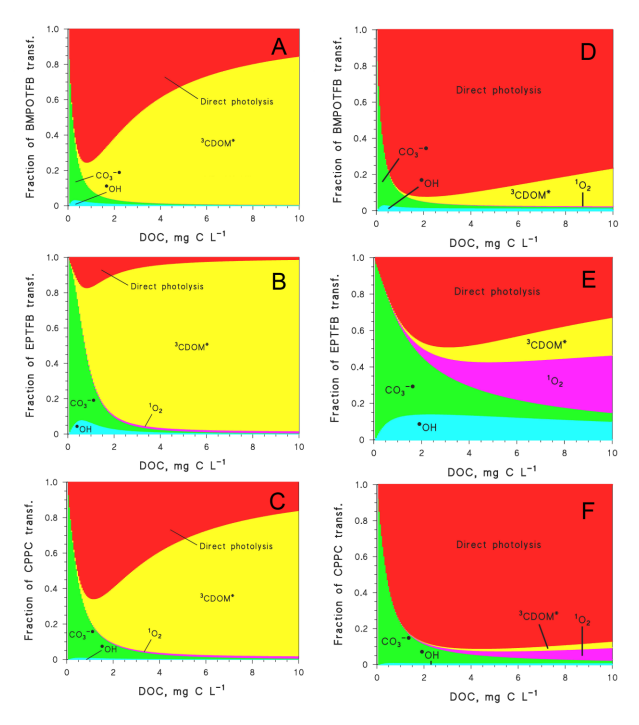


Figure 4. Fractions of ILs phototransformation accounted for by the different photoinduced pathways, as predicted by the APEX software for summertime irradiation conditions. (A) **BMPOTFB**, with $k_{BMPOTFB^3CDOM^*} = k_{BMPOTFB^3AQ2S^*}$; (B) **EPTFB**, with $k_{EPTFR^3CDOM^*} = k_{EPTFR^3AQ2S^*}$; (C) **CPPC**, with $k_{CPPC^3CDOM^*} = k_{CPPC^3AQ2S^*}$; (D) **BMPOTFB**, with $k_{BMPOTFB^3CDOM^*} = 0.05$ $k_{BMPOTFB^3AQ2S^*}$; (E) **EPTFB**, with $k_{EPTFR^3CDOM^*} = 0.05$ $k_{EPTFR^3AQ2S^*}$; (F) **CPPC**, with $k_{CPPC^3CDOM^*} = 0.05$ $k_{CPPC^3AQ2S^*}$. Other water conditions: 5 m depth, 0.1 mM nitrate, 1 μM nitrite, 1 mM bicarbonate, 10 μM carbonate.

3.3. ILs toxicity assessment upon photoinduced degradation

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Acute toxicity was monitored over time upon UVB direct photolysis of the studied ILs. The choice of the direct photolysis pathway for the toxicity study has two reasons: (i) experimental simplicity, because the assessment of toxicity in indirect phototransformation is complicated by the possible production of toxic species/intermediates from the photosensitizers (De Luis et al., 2010). The indirect phototransformation processes might be incompletely accounted for in control experiments carried out without the substrates, because the substrates themselves are major scavengers of transient species (e.g. in the case of *OH and CO₃-*), and the transient steady-state concentrations may vary dramatically when the substrates are absent; (ii) environmental significance. The direct photolysis is an important process in the phototransformation of BMPOTFB, for which photochemistry may be a significant attenuation pathway in surface waters. The direct photolysis is less important for EPTFB (see Figure 4), but the phototransformation kinetics of this compound are predicted to be very slow except for low-DOC waters (Figure 3). In these conditions, the direct phototransformation pathway could play a remarkable role. The phototransformation of **CPPC** can only be significant under low-DOC conditions, where the direct photolysis is an important degradation pathway. The acute toxicity tests were carried out by monitoring changes in the natural emission of the luminescent bacteria Vibrio fischeri, when exposed to potentially toxic compounds. Out of the various available bioassays, this test is sensitive, rapid, cost-effective, reproducible, and it can be used for almost all kinds of toxic compounds (Parvez et al., 2006, Matsushita et al., 2015). The toxicity is here expressed as the percentage of inhibition of the bacteria luminescence. Results obtained on samples subjected to different irradiation times are plotted in Figure 5 (5A: time trend of ILs in the irradiated solutions; **5B**: time trend of the toxicity in the same solutions). The studied ILs are not toxic, as shown by the absence or near absence of a toxic effect before irradiation. Under UVB light the photoinduced transformation of **EPTFB** proceeded through the formation of almost

harmless TPs, and a slight toxicity increase was only observed after 7 days of irradiation. Unfortunately, this compound is the studied IL for which phototransformation (either direct or indirect) is expected to play the least important role in the natural environment (see **Figure 3**). In contrast, the toxicity of irradiated **CPPC** and **BMPOTFB** increased over time, thereby suggesting that their phototransformation proceeded through the formation of moderately toxic compounds.



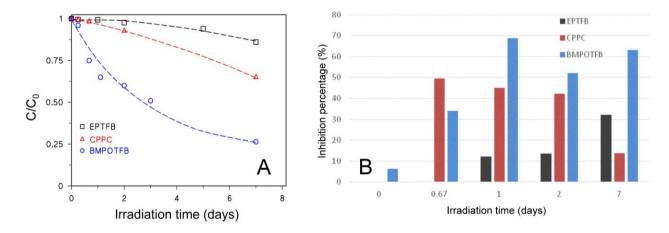


Figure 5. (**A**) Time trends of **BMPOTFB**, **EPTFB** and **CPPC** (20 mg L⁻¹ initial loading in separate solutions) upon direct photolysis under UVB irradiation. (**B**) Toxicity evolution (inhibition of *Vibrio fischeri* emission) in the same irradiated solutions shown in the (A) panel.

3.4. Characterization of the transformation products

The direct photolysis of **EPTFB**, **CPPC** and **BMPOTFB** yielded several transformation products (TPs), which were detected in ESI positive mode via HPLC-HRMS. **Tables SM2-SM4** in the SM report the measured m/z ratios and the most probable empirical formulas for all the detected TPs. The relevant time evolution profiles are provided in **Figure 6.** Among the TPs of **BMPOTFB**, a few of them were formed at early reaction times and disappeared during the course of the irradiation experiments, which lasted for up to eight days. Several other TPs appeared at relatively long

- 429 irradiation times, and their concentration increased during irradiation. The various TPs could be
- formed upon direct photolysis through three different pathways shown in **Scheme 1** and involving:
- (a) hydroxylation; (b) detachment/shortening of the butyl chain; (c) double bond formation.
- The MS^2 spectrum of **BMPOTFB** has a product ion at 94.0652 m/z, presumably produced by the
- loss of the butyl chain (see **Table SM2**). This peculiar loss was helpful in attributing the structures
- of unknown TPs. A TP having 166.1233 m/z (**TP-166**) is consistent with a monohydroxyderivative.
- The formation of a product ion at 71.0447 m/z (hydroxylated butyl moiety) in the MS² spectrum of
- 436 **TP-166**, combined with the loss of C₂H₆, permits to locate the OH group on the butyl chain and to
- confine it on C1 or C2.

- Four species with 164.1077 m/z (**TP-164A** through **D**) were detected and attributed to keto
- derivatives. For **TP-164A** it was not possible to acquire enough structural information, while for the
- other compounds the presence of the product ion at $94.0634 \, m/z$ allows the placement of the
- hydroxyl group on the butyl chain. Additionally, **TP-164B** presents the structural diagnostic loss of
- formaldehyde that allows the keto group to be located on C4 (McLafferty and Turecek, 1993).
- Three species with 182.1184 m/z (TP-182A through C) are consistent with BMPOTFB
- dihydroxyderivatives. All these TPs show the product ion at 94.0652 m/z in their MS² spectrum,
- thereby allowing the two OH groups to be located on the butyl chain. **TP-182C** should have one
- OH group on the C4 atom of the butyl chain, as suggested by the loss of methanol.
- Three species with 180.0995 m/z (**TP-180A** through **C**) should be dihydroxylated/oxidized
- compounds. In the case of **TP-180C**, the loss of formic acid and the formation of the precursor ion
- at $94.0634 \, m/z$ suggest the presence of a carboxylic group on the butyl chain.

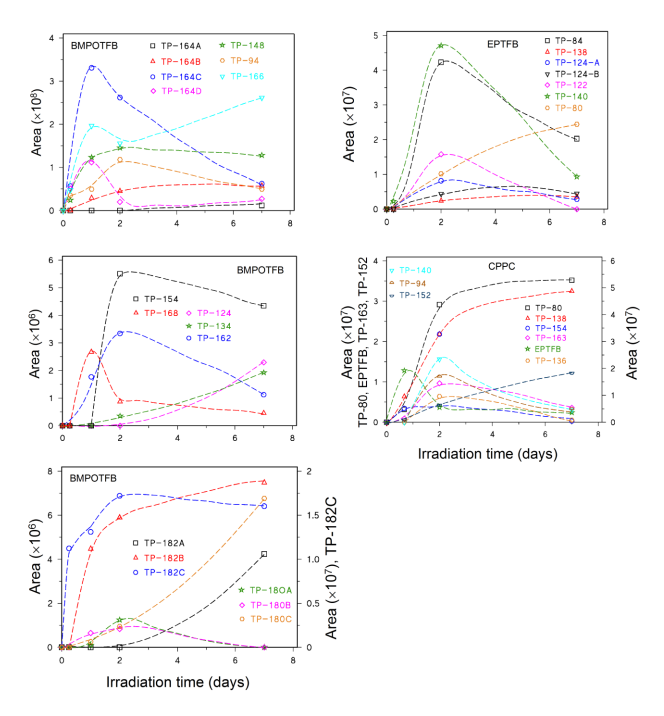


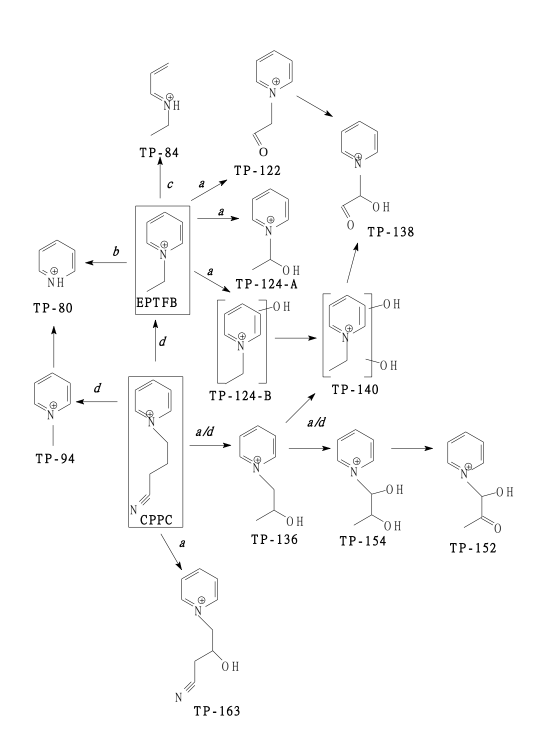
Figure 6. Time evolution of the TPs formed upon UVB irradiation of (left side) **BMPOTFB** and of (right side) **EPTFB** (top) and **CPPC** (bottom). Multiple plots are provided for **BMPOTFB** to show TPs with different peak areas, and for readability issues as well. The dashed curves are just a guide for the eye. In a couple of cases data are plotted against different Y-axes, located on the right hand and on the left hand of each plot. The least numerous group of compounds is listed on the given axis. The remaining compounds, not explicitly mentioned, are plotted against the other axis.

Scheme 1. Proposed transformation pathways followed by **BMPOTFB** under direct photolysis.

Three compounds at 162.0891, 148.1126 and 134.0967 m/z with empirical formulas $C_{10}H_{12}ON$, $C_{10}H_{14}ON$ and $C_{9}H_{12}N$, respectively, are consistent with progressive dealkylation and with formation of a double bond on the alkyl chain. For **TP-162**, the most abundant ion in MS² exhibits a $C_{4}H_{6}O$ loss that suggests the occurrence of a keto group on the butyl chain.

- 465 **TP-168** and **TP-154** may be formed *via* the detachment of a methyl group and/or *via* side-chain
- shortening, but insufficient mass information was available to propose a univocal structure.
- 467 TP-94 and TP-124 are formed upon detachment of the butyl chain. TP-94 is consistent with 4-
- methylpyridine, while **TP-124** would involve methyl oxidation and possibly ring hydroxylation.
- 469 Unfortunately, the information contained in the MS spectra did not allow for the proposal of a
- proper location for the OH group in **TP-124**.
- As far as the indirect photochemistry processes are concerned, TP-148 was also detected upon
- reactions with •OH, CO₃-• and ³AQ2S*, **TP-164 B,D** with •OH, **TP-164 C,D** and **TP-166 C,D** with
- 473 ³AQ2S*, and **TP-94** with CO₃^{-•}. The formation of partially overlapping TPs in different pathways
- 474 is quite common in phototransformation reactions (De Laurentiis et al., 2014).
- In the case of **EPTFB**, three main transformation pathways could be detected under direct photolysis conditions. These pathways involve: (a) hydroxylation; (b) detachment of the ethyl chain, and (c) ring opening (see **Scheme 2**). The MS² spectrum of **EPTFB** has the ion at 80.0492 m/z as base peak, formed through the detachment of the ethyl chain (C₂H₄, see **Table SM3**). Upon
- photolysis, pathway a likely yields two isobaric species with 124.0757 m/z and empirical formula
- 480 $C_7H_{10}ON$, which can be attributed to hydroxyderivatives. In the case of **TP-124A**, the loss of
- 481 C₂H₄O in the MS² spectrum suggests that the OH substituent occurs on the alkyl chain. Moreover,
- the absence of methanol loss tentatively suggests that OH is located on the ethyl carbon atom bound
- 483 to nitrogen (C1). For **TP-124B**, the absence of MS² product ions prevented any hypothesis
- concerning the OH position. The formation of **TP-122**, with empirical formula C₇H₈ON, likely
- involved the oxidation of the alcoholic group to a keto one. The loss of C₂HON in the MS²
- spectrum supports the occurrence of a keto group on the alkyl chain, which should be located on C2
- as suggested by the CO loss (McLafferty and Turecek, 1993). Pathway a also involved the
- formation of TPs with empirical formula C₇H₁₀O₂N (**TP-140**) and C₇H₈O₂N (**TP-138**), which are

well-matched with dihydroxylated and dihydroxylated/oxidized derivatives, respectively. However, the absence of MS^2 product ions prevented any hypothesis about the location of the two OH groups.



Scheme 2. Proposed transformation pathways followed by compounds **EPTFB** and **CPPC** upon direct photolysis.

Further transformation (pathway b) involved the detachment of the alkyl chain with formation of pyridine (TP-80). An alternative route (pathway c) would proceed through ring opening to yield compound TP-84. The most abundant compounds (as far as peak areas are concerned) were TP-140, TP-84 and TP-80, thereby suggesting that the three pathways had similar importance. Interestingly, the compound **TP-84** was also detected upon reactions with •OH, CO₃-• and ³AQ2S*, and TP-140 upon reaction with *OH as well. In a close analogy with **EPTBF**, the MS² spectrum of **CPPC** presents as base peak the pyridinium ion at 80.0492 m/z. This ion is likely formed through the detachment of the cyano-propyl chain (see **Table SM4**). The direct photolysis of **CPPC** involved hydroxylation (see pathway *a* in **Scheme 2**). TP-163 is well-matched with a monohydroxylated derivative, and the detected MS² product ions are consistent with losses of C₂H₃N and C₃H₃ON. Therefore, the OH substituent should not occur on the cyano carbon (C4) or on the carbon attached to the cyano group (C3) (McLafferty and Turecek, 1993). The occurrence of OH on the carbon atom bound to the ring nitrogen (C1) also looks little likely, leaving C2 as the most reasonable position for the OH group. The MS data are also consistent with chain shortening (pathway d) and with a combined chain shortening/hydroxylation (pathway a/d). The detachment of the methylcyano group accounts for the formation of a compound with 108.0809 m/z, recognized as **EPTFB** and confirmed by standard match. A further chain shortening would produce **TP-94**, which was identified as 1-methylpyridine. The latter could yield **TP-80**, matched with pyridine (also detected during **EPTFB** photolysis), through detachment of the methyl group. Pathway a/d yields TP-136, TP-154 and TP-152 through cyano group detachment and hydroxylation. In all cases, the OH groups were added to the propyl chain as assessed by the product ion at 80.0478 m/z (pyridinium ion) in the relevant MS² spectra. The presence of some structurally-diagnostic ions allowed a univocal location of the oxygen atoms

on the alkyl chain. TP-136 formation would imply the detachment of the cyano group and the

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oxidation of the alkyl chain. Actually, the absence of formaldehyde loss combined with C₂H₂O and

 C_3H_4O losses allowed the location of the keto group on C2.

Based on its exact mass, **TP-152** might have an OH group and a keto one. The oxygen-containing functions should be both located on the alkyl chain, as suggested by the occurrence of the pyridinium ion in the MS² spectrum. The absence of any methanol loss is not consistent with the occurrence of OH on C3. Together with the loss of ketene, this suggests that the keto group should be located on C2 and the hydroxyl one on C1. Therefore, **TP-152** could be tentatively identified as the oxidized form of **TP-154**. The formation of **TP-138** and **TP-140** likely involved methylcyanogroup detachment and hydroxylation. The retention times and fragmentation spectra of the TPs derived from **CPPC** matched well with the TPs formed from **EPTFB** photolysis. As far as peak areas are concerned, the main TPs were **TP163**, formed through pathway *a*, **TP108** and **TP80**, involving pathway *d*. While **EPTFB** was formed from **CPPC** at early reaction times and reached its maximum concentration after 1 day of irradiation, all the other TPs were more persistent and either peaked at two days irradiation, or continued accumulating up to at least eight days. No TPs were detected upon indirect phototransformation of **CPPC**, probably because of the scarce reactivity of this substrate with the studied transient species.

The occurrence of definite chemical functions allows some hypotheses to be advanced concerning the potential toxicity of TPs (Mayo-Bean et al., 2012). On this basis and from the above discussion one may assume that among the detected TPs of **BMPOTFB**, those for which a univocal chemical structure could be proposed are unlikely to be more toxic than the parent compound, because no potentially toxic functionalities were added (Mayo-Bean et al., 2012). A toxicity increase could be expected in the (possible but not confirmed) circumstance that dihydroxylation involved the aromatic ring, especially if two OH groups occurred in para position (hydroquinone derivatives) (Mayo-Bean et al., 2012). Similar considerations hold for **CPPC** and **EPTFB**.

4. Conclusions

- Phototransformation in sunlit surface waters is a potentially important attenuation pathway for
 BMPOTFB but not for EPTFB and CPPC, with the possible exception of shallow water
 bodies with very low DOC.
- The phototransformation of **BMPOTFB** is expected to proceed through CO₃^{-•} reactions at low DOC and *via* direct photolysis at higher DOC. In water bodies that are rich in pedogenic organic matter (humic and fulvic substances), also the ³CDOM* reactions could play an important role at elevated DOC.
 - The phototransformation of **EPTFB** and **CPPC** is extremely slow in high-DOC waters, thus the most relevant photochemical attenuation pathways (provided that the water body is shallow) are those prevailing at low DOC: CO₃-•, direct photolysis and, for **EPTFB** alone, •OH.
 - The direct photolysis of **BMPOTFB** and **CPPC** yields TPs that are more toxic than the parent compound. The main identified reaction pathways upon direct photolysis are hydroxylation, lateral-chain shortening and ring opening, and toxic TPs might be formed by multiple hydroxylation if this process yields hydroquinone-like derivatives.

Acknowledgments

Water JPI Pilot Call.

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References

- Avetta, P., Fabbri, D., Minella, M., Brigante, M., Maurino, V., Minero, C., Pazzi, M., Vione, D.,
- 2016. Assessing the phototransformation of diclofenac, clofibric acid and naproxen in surface
- waters: Model predictions and comparison with field data. Water. Res. 105, 383-394.
- Bandres, I., Giner, B., Artigas, H., Royo, F. M., Lafuente, C., 2008. Thermophysic comparative
- study of two isomeric pyridinium-based ionic liquids. J. Phys. Chem. B 112, 3077-3084.
- Bahnmüller, S., von Gunten, U., Canonica, S., 2014. Sunlight-induced transformation of
- sulfadiazine and sulfamethoxazole in surface waters and wastewater effluents. Water Res. 57,
- 575 183-192.
- Bedini, A., De Laurentiis, E., Sur, B., Maurino, V., Minero, C., Brigante, M., Mailhot, G., Vione,
- D., 2012. Phototransformation of anthraquinone-2-sulphonate in aqueous solution.
- 578 Photochem. Photobiol. Sci. 11, 1445-1453.
- Bintou, A. T., Bianco, A., Mailhot, G., Brigante, M., 2015. A new insight into ethoxyquin fate in
- surface waters: Stability, direct and indirect photochemical behaviour and the identification of
- main products. J. Photochem. Photobiol. A: Chem. 311, 118-126.
- Bodrato, M., Vione, D., 2014. APEX (Aqueous Photochemistry of Environmentally occurring
- Xenobiotics): A free software tool to predict the kinetics of photochemical processes in
- surface waters. Environ. Sci.: Processes Impacts 16, 732-740.
- Braslavsky, S.E., 2007. Glossary of terms used in photochemistry. third edition. Pure Appl. Chem.
- *79*, 293-465.
- Bubalo, M.C.; Radošević, K.; Redovniković, I.R.; Halambek, J.; Srček, V.G., 2014. A brief
- overview of the potential environmental hazards of ionic liquids, Ecotoxicol. Environ. Saf.,
- 589 2014,99,1-12.

- 590 Buxton, G. V., Greenstock, C. L., Helman, W. P., Ross, A. B., 1988. 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, 513-886.
- Calza, P., Vione, D., Fabbri, D., Aigotti, R., Medana, C., 2015. Imidazolium-based ionic liquids in
- water: Assessment of photocatalytic and photochemical transformation. Environ. Sci.
- 595 Technol. 49, 10951-10958.
- Canonica, S., Kohn, T., Mac, M., Real, F.J., Wirz, J., von Gunten, U., 2005. Photosensitizer method
- 597 to determine rate constants for the reaction of carbonate radical with organic compounds.
- 598 Environ. Sci. Technol. 39, 9182-9188.
- 599 Chiwa, M., Higashi, N., Otsuki, K., Kodama, H., Miyajima, T., Takeda, K., Sakugawa, H., 2015.
- Sources of hydroxyl radical in headwater streams from nitrogen-saturated forest.
- 601 Chemosphere 119, 1386-1390.
- 602 Czerwicka, M., Stolte, S., Müller, A., Siedlecka, E. M., Gołebiowski, M., Kumirska J., Stepnowski,
- P., 2009. Identification of ionic liquid breakdown products in an advanced oxidation system.
- 604 J. Hazard. Mater. 171, 478-483.
- Dantas, R. F., Canterino, M., Marotta, R., Sansa, C., Esplugas, S., Andreozzi, R., 2007. Bezafibrate
- removal by means of ozonation: Primary intermediates, kinetics, and toxicity assessment.
- 607 Water Res. 41, 2525-2532.
- De Laurentiis, E., Minella, M., Maurino, V., Minero, C., Brigante, M., Mailhot, G., Vione, D.,
- 609 2012. Photochemical production of organic matter triplet states in water samples from
- mountain lakes, located below or above the tree line. Chemosphere 88, 1208-1213.
- De Laurentiis, E., Prasse, C., Ternes, T.A., Minella, M., Maurino, V., Minero, C., Sarakha, M.,
- Brigante, M., Vione, D., 2014. Assessing the photochemical transformation pathways of

- acetaminophen relevant to surface waters: transformation kinetics, intermediates, and
- 614 modelling. Water Res. 53, 235-248.
- De Luis, A. M., Lombrana, J. I., Menendez, A., Sanz, J., 2011. Analysis of the toxicity of phenol
- solutions treated with H₂O₂/UV and H₂O₂/Fe oxidative systems. Ind. Eng. Chem. Res. 50,
- 617 1928-1937.
- Dong, M. M., Trenholm, R., Rosario-Ortiz, F. L., 2015. Photochemical degradation of atenolol,
- carbamazepine, meprobamate, phenytoin and primidone in wastewater effluents. J. Hazard.
- 620 Mater. 2015, 282, 216-223.
- Fabbri, D., Minella, M., Maurino, V., Minero, C., Vione, D., 2015. Photochemical transformation
- of phenylurea herbicides in surface waters: A model assessment of persistence, and
- 623 implications for the possible generation of hazardous intermediates. Chemosphere 119, 601-
- 624 607.
- Frank, R., Klöpffer, W., 1988. Spectral solar photo irradiance in Central Europe and the adjacent
- north Sea, Chemosphere 17, 985-994.
- 627 Giannakis, S., Polo Lopez, M. I., Spuhler, D., Sanchez Perez, J. A. Fernandez Ibanez, P., Pulgarin,
- 628 C., 2016a. Solar disinfection is an augmentable, in situ-generated photo-Fenton reaction-Part
- 1: A review of the mechanisms and the fundamental aspects of the process. Appl. Catal. B:
- Environ. 199, 199-223.
- 631 Giannakis, S., Polo Lopez, M. I., Spuhler, D., Sanchez Perez, J. A. Fernandez Ibanez, P., Pulgarin,
- 632 C., 2016b. Solar disinfection is an augmentable, in situ-generated photo-Fenton reaction Part
- 2: A review of the applications for drinking water and wastewater disinfection. Appl. Catal. B:
- Environ. 198, 431-446.

- 635 Guerrero, H., Garcia-Mardones, M., Cea, P., Lafuente, C., Bandres, I., 2012. Correlation of the
- volumetric behaviour of pyridinium-based ionic liquids with two different equations.
- 637 Thermochim. Acta 531, 21-27.
- 638 Herrmann, H., Hoffmann, D., Schaefer, T., Bräuer, P., Tilgner, A., 2010. Tropospheric aqueous-
- phase free-radical chemistry: Radical sources, spectra, reaction kinetics and prediction tools.
- ChemPhysChem 11, 3796-3822.
- Holbrey J. D., Seddon K. R., 1999. Ionic liquids. Clean Technol. Environ. 1, 223-236.
- Huang, J. P., Mabury, S. A., 2000. Steady-state concentrations of carbonate radicals in field waters.
- Environ. Toxicol. Chem. 19, 2181-2188.
- Katsoyiannis, I. A., Canonica, S., von Gunten, U., 2011. Efficiency and energy requirements for the
- transformation of organic micropollutants by ozone, O₃/H₂O₂ and UV/H₂O₂. Water Res. 45,
- 646 3811-3822.
- Keen, O. S., McKay, G., Mezyk, S. P., Linden, K. G., Rosario-Ortiz, F. L., 2014. Identifying the
- factors that influence the reactivity of effluent organic matter with hydroxyl radicals. Water
- Res. 50, 408-419.
- Kralisch, D., Stark, A., Korsten, S., Kreisel G., Ondruschka, B., 2005. Energetic, environmental and
- economic balances: Spice up your ionic liquid research efficiency. Green Chem. 7, 301-309.
- Janssen, E. M. L., Erickson, P. R., McNeill, K., 2014. Dual roles of dissolved organic matter as
- sensitizer and quencher in the photooxidation of tryptophan. Environ. Sci. Technol. 48, 4916-
- 654 4924.
- Janssen, E. M. L., Marron, E., McNeill, K., 2015. Aquatic photochemical kinetics of benzotriazole
- and structurally related compounds. Environ. Sci.: Processes Impacts 17, 939-946.
- Lee, E., Glover, C. M., Rosario-Ortiz, F. L., 2013. Photochemical formation of hydroxyl radical
- from effluent organic matter: Role of composition. Environ. Sci. Technol. 47, 12073-12080.

- Loiselle, S. A., Azza, N., Cozar, A., Bracchini, L., Tognazzi, A., Dattilo, A., Rossi, C., 2008.
- Variability in factors causing light attenuation in Lake Victoria. Freshwater Biol. 53, 535-545.
- Maddigapu, P. R., Bedini, A., Minero, C., Maurino, V., Vione, D., Brigante, M., Mailhot, G.,
- Sarakha, M., 2010. The pH-dependent photochemistry of anthraquinone-2-sulfonate.
- Photochemical and Photobiological Sciences 9, 323-330.
- Mayo-Bean, K., Moran, K., Meylan, B., Ranslow, P., 2012. Methodology Document for the
- ECOlogical Structure-Activity Relationship Model (ECOSAR) Class Program. US-EPA,
- Washington DC, 46 pp.
- Marchetti, G., Minella, M., Maurino, V., Minero, C., Vione, D., 2013. Photochemical
- transformation of atrazine and formation of photointermediates under conditions relevant to
- sunlit surface waters: Laboratory measures and modelling. Wat. Res. 47, 6211-6222.
- Marchisio, A., Minella, M., Maurino, V., Minero, C., Vione, D., 2015. Photogeneration of reactive
- transient species upon irradiation of natural water samples: Formation quantum yields in
- different spectral intervals, and implications for the photochemistry of surface waters. Water
- 673 Res. 73, 145-156.
- Matsushita, T., Kobayasgu, N., Hashizuka, M., Sakuma, H., Kondo, T., Matsui, Y., Shirasaki, N.,
- 675 2015. Changes in mutagenicity and acute toxicity of solutions of iodinated X-ray contrast
- 676 media during chlorination, Chemosphere 135, 101-107.
- McLafferty, F. W., Turecek, F., 1993. Interpretation of mass spectra. Fourth edition, University
- 678 Science Books, Mill Valley, California.
- McNeill, K., Canonica, S., 2016. Triplet state dissolved organic matter in aquatic photochemistry:
- reaction mechanisms, substrate scope, and photophysical properties. Environ. Sci. Processes
- 681 Impacts 18, 1381-1399.

- Page, S. E., Logan, J. R., Cory, R. M., McNeill, K., 2014. Evidence for dissolved organic matter as
- the primary source and sink of photochemically produced hydroxyl radical in arctic surface
- waters. Environ. Sci. Processes Impacts 16, 807-822.
- Parvez, S., Venkataraman, C., Mukherji, S., 2006. A review on advantages of implementing
- luminescence inhibition test (Vibrio fischeri) for acute toxicity prediction of chemicals.
- Environ. Int. 32, 265-268.
- Plechkova, N. V., Seddon, K. R., 2008. Applications of ionic liquids in the chemical industry
- 689 Chem. Soc. Rev. 37, 123-150.
- 690 Puskas, J. E., Chan, S. W. P., McAuley, K. B., Shaikh, S., Kaszas, G., 2005. Kinetics and
- mechanisms in carbocationic polymerization: The quest for true rate constants. J. Polym. Sci.
- 692 Pol. Chem. 43, 5394-5413.
- Richardson, S. D., Ternes, T. A., 2014. Water analysis: emerging contaminants and current issues.
- 694 Anal. Chem. 86, 2813-2848.
- Rosario-Ortiz, F. L., Canonica, S., 2016. Probe compounds to assess the photochemical activity of
- dissolved organic matter. Environ. Sci. Technol. 50, 12532-12547.
- 697 Silva, M. P., Mostafa, S., McKay, G., Rosario-Ortiz, F. L., Teixeira, A. C. S. C., 2015.
- Photochemical fate of amicarbazone in aqueous media: Laboratory measurement and
- simulations. Environ. Eng. Sci. 32, 730-740.
- Stepnowski, P., Zaleska, A., 2005. Comparison of different advanced oxidation processes for the
- degradation of room temperature ionic liquids J. Photochem. Photobiol. A. 170, 45-50.
- Sun, B., Dong, H. Y., He, D., Rao, D. D., Guan, X. H., 2016. Modeling the kinetics of contaminants
- oxidation and the generation of manganese(III) in the permanganate/bisulfite process.
- 704 Environ. Sci. Technol. 50, 1473-1482.
- 705 Trivella, A., Stawinoga, M., Dayan, F. E., Cantrell, C. L., Mazellier, P., Richard, C., 2015.
- Photolysis of natural beta-triketonic herbicides in water. Water Res. 78, 28-36.

- Vione, D., Minella, M., Maurino, V., Minero, C., 2014. Indirect photochemistry in sunlit surface
- waters: Photoinduced production of reactive transient species. Chemistry Eur. J. 20, 10590-
- 709 10606.
- 710 Wenk, J., Eustis, S. N., McNeill, K., Canonica, S., 2013. Quenching of excited triplet states by
- dissolved natural organic matter. Environ. Sci. Technol. 47, 12802-12810.
- 712 Zeng, T., Arnold, A., 2013. Pesticide photolysis in prairie potholes: Probing photosensitized
- 713 processes. Environ. Sci. Technol. 47, 6735-6745.
- 714 Zhao, H., Malhotra, S. V., Luo, R. G., 2003. Preparation and characterization of three room-
- temperature ionic liquids. Phys. Chem. Liq. 41, 487-492.