Modelling the photochemical fate of ibuprofen in surface waters

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MODELLING THE PHOTOCHEMICAL FATE OF IBUPROFEN IN SURFACE WATERS

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Abstract

We show that the main photochemical processes involved in the phototransformation of anionic ibuprofen (IBP) in surface waters are the reaction with \(^{•}\)OH, the direct photolysis and possibly the reaction with the triplet states of chromophoric dissolved organic matter (\(^{3}\)CDOM\(^*\)). These conclusions were derived by use of a model of surface-water photochemistry, which adopted measured parameters of photochemical reactivity as input data. The relevant parameters are the polychromatic UVB photolysis quantum yield (\(\Phi_{IBP} = 0.33\pm0.05\), \(\mu\pm\sigma\)), the reaction rate constant with \(^{•}\)OH (\(k_{IBP,^{•}\text{OH}} = (1.0\pm0.3)\times10^{10}\) M\(^{-1}\) s\(^{-1}\)), the \(^{1}\)O\(_2\) rate constant (\(k_{IBP,^{1}\text{O}_2} = (6.0\pm0.6)\times10^{4}\) M\(^{-1}\) s\(^{-1}\)), while the reaction with CO\(_3\)\(^*\) can be neglected. We adopted anthraquinone-2-sulphonate (AQ2S) and riboflavin (Ri) as CDOM proxies and the reaction rate constants with the respective triplet states were \(k_{IBP,^{3}\text{AQ2S}*} = (9.7\pm0.2)\times10^{9}\) M\(^{-1}\) s\(^{-1}\) and \(k_{IBP,^{3}\text{Ri}*} = 4.5\times10^{7}\) M\(^{-1}\) s\(^{-1}\). The reaction with \(^{3}\)CDOM\(^*\) can be an important IBP sink if its rate constant is comparable to that of \(^{3}\)AQ2S\(^*\), while it is unimportant if the rate constant is similar to the \(^{3}\)Ri\(^*\) one. The photochemical pathways mainly lead to the transformation (oxidation and/or shortening) of the propanoic lateral chain of IBP, which
appears to be significantly more reactive than the isobutyl one. Interestingly, none of the detected intermediates was produced by substitution on the aromatic ring.

**Keywords**: pharmaceuticals; PCCPs; non-steroidal anti-inflammatory drug; surface-water photochemistry; sensitised phototransformation; hydroxyl radical; singlet oxygen.

**Introduction**

Ibuprofen (2-(4-(2-methylpropyl)phenyl)propanoic acid, hereafter IBP) is a non-steroidal anti-inflammatory drug that is widely used nowadays as the active principle of many “over the counter” pharmaceutical products. Excretion by humans after partial metabolism and incorrect drug disposal are two important pathways of IBP to sewage waters. For instance, it has been the sixth most abundant pharmaceutical detected in the influent flow of wastewater treatment plants (WWTPs) in Italy (Castiglioni et al., 2006). Although it is susceptible to biodegradation (Tiehm et al., 2011; de Graaf et al., 2011), IBP is only partially removed by WWTPs (Oulton et al., 2010; Morasch et al., 2010) and the treatment efficiency undergoes a considerable decrease during the winter months (Castiglioni et al., 2006; Santos et al., 2009), when the IBP environmental inputs tend to increase considerably (Danshvar et al., 2010). The combination of widespread use and incomplete WWTP elimination results into an extensive occurrence of IBP in surface waters. Indeed, this compound has been detected in water systems all over the world (Zuccato et al., 2005; Zhao et al., 2009; Yu et al., 2009; Pailler et al., 2009; Moldovan et al., 2009; Fernandez et al., 2010; Camacho-Munoz et al., 2010; Loos et al., 2010; Tejon et al., 2010; Zhao et al., 2010; Helenkar et al., 2010; Waiser et al., 2011; Wang et al., 2011; Lewandowski et al., 2011).

IBP has been found to accumulate in fatty fish tissues and muscles (Zhang et al., 2010) and to cause disturbance to both amphipoda (De Lange et al., 2009) and cnidaria (Quinn et al., 2008). IBP effects can be increased when it is present in mixture with other pharmaceuticals. Negative effects at environmentally significant levels have been observed for both cnidaria (Quinn et al., 2009) and human embryonic cells (Pomati et al., 2006). The latter finding is potentially of very high concern when considering the occurrence of IBP in drinking water, at very variable levels depending on the location (Kleywegt et al., 2011; Valcarcel et al., 2011).

Biodegradation and phototransformation are potentially important transformation processes for IBP in surface waters (Lin and Reinhard, 2005). Phototransformation might involve direct
photolysis and indirect reaction with transient species (\( ^{\cdot}\)OH, CO\(_3\)^{\cdot}\), \( ^{1}\)O\(_2\), \( ^{3}\)CDOM\(^{*}\)) generated by CDOM, nitrite and nitrate under irradiation (Hoigné, 1990; Peuravuori and Pihlaja, 2009). As far as the direct photolysis is concerned, IBP is a poor absorber of sunlight (Lin and Reinhard, 2005; Peuravuori and Pihlaja, 2009) but its highest-wavelength absorption band (maximum around 265 nm) is characterised by a quite efficient photolysis (Yuan et al., 2009). Poor sunlight absorption and elevated photolysis quantum yield are contrasting issues as far as the environmental importance of direct photolysis is concerned. A similar case holds for the herbicide 2-methyl-4-chlorophenoxy acetic acid (MCPA), and we have recently found that MCPA direct photolysis can be a significant process but it is highly influenced by the environmental variables (Vione et al., 2010). The available literature data do not allow a clear assessment of the importance of the direct photolysis as a potential transformation pathway for IBP in surface waters (Packer et al., 2003; Lin and Reinhard, 2005; Matamoros et al., 2009; Peuravuori and Pihlaja, 2009).

We have recently developed a model approach to describe the transformation kinetics of dissolved compounds as a function of water chemical composition, column depth and photochemical reactivity (i.e. direct photolysis quantum yield and reaction rate constants with \( ^{\cdot}\)OH, CO\(_3\)^{\cdot}\), \( ^{1}\)O\(_2\) and \( ^{3}\)CDOM\(^{*}\)) (Maddigapu et al., 2011). Such an approach was able to effectively predict field data in the case of 2,4-dichloro-6-nitrophenol in the Rhône delta lagoons (Chiron et al., 2007). The goal of the present paper is to assess the role of direct and indirect photochemistry in the degradation of IBP in surface waters. To this purpose, we combine a kinetic study of IBP transformation by direct photolysis and reaction with \( ^{\cdot}\)OH, CO\(_3\)^{\cdot}\), \( ^{1}\)O\(_2\) and \( ^{3}\)CDOM\(^{*}\), with a photochemistry model. The latter enables the prediction of IBP transformation kinetics as a function of photoreactivity data and of key environmental variables.

IBP phototransformation intermediates, formed in the first step of the environmentally most significant processes were also identified. IBP has pK\(_a\) = 4.4 (Martell et al., 1997) and its anionic form would prevail under most conditions that are relevant to surface waters. In this work, irradiation was carried out at both pH 2 and 8 where the neutral and the anionic IBP forms prevail, respectively.

**Experimental section**

**Reagents and materials**

Ibuprofen (IBP, purity grade 98%), anthraquinone-2-sulphonic acid, sodium salt (AQ2S, 97%), furfuryl alcohol (98%), NaNO\(_3\) (>99%), NaHCO\(_3\) (98%), anhydrous Na\(_2\)SO\(_4\) (99%), NaCl (99.5%),
HClO₄ (70%) and H₃PO₄ (85%) were purchased from Aldrich, NaOH (99%), methanol and 2-propanol (both LiChrosolv gradient grade) and dichloromethane (GC Suprasolv) from VWR Int., Rose Bengal (RB) from Alfa Aesar, riboflavin (>98%) from Sigma.

**Irradiation experiments**

Solutions to be irradiated (5 mL) were placed inside Pyrex glass cells (4.0 cm diameter, 2.3 cm height, 295 nm cut-off wavelength) and magnetically stirred during irradiation. Irradiation of IBP + nitrate to study reactions with •OH and CO₃•, and irradiation of IBP alone to study the direct photolysis were carried out under a Philips TL 01 UV-Vis lamp, with emission maximum at 313 nm and 3.0±0.2 W m⁻² UV irradiance in the 300-400 nm range, measured with a power meter by CO.FO.ME.GRA. (Milan, Italy) equipped with a UV-sensitive probe. The incident photon flux in solution was actinometrically determined using the ferrioxalate method (Kuhn et al., 2004). By knowing, as a function of the wavelength, the fraction of radiation absorbed by Fe(C₂O₄)₃³⁻, the quantum yield of Fe²⁺ photoproduction and the shape of the lamp spectrum (*vide infra*), it is possible to use the measured formation rate of Fe²⁺ to fix the value of the incident spectral photon flux density \( p^o(\lambda) \). The photon flux of the UV-Vis lamp between 300 and 500 nm was

\[
P_o = \int p^o(\lambda) d\lambda = 2.0 \times 10^{-5} \text{ Einstein L}^{-1} \text{ s}^{-1}.
\]

The transformation of IBP photosensitised by AQ2S was studied under a Philips TL K05 UVA lamp, with emission maximum at 365 nm, 28 W m⁻² UV irradiance (300-400 nm), and 2.1\( \times 10^{-5} \) Einstein L⁻¹ s⁻¹ incident photon flux in solution between 300 and 500 nm. The photodegradation of IBP sensitised by Rose Bengal (RB) via \( ^1\text{O}_2 \) was studied under a Philips TL D 18W/16 yellow lamp, with emission maximum at 545 nm and 11 W m⁻² irradiance in the visible, measured with the CO.FO.ME.GRA. power meter equipped with a probe sensitive to visible radiations.

The mixtures IBP-riboflavin were irradiated at 365 nm in a cuvette in parallel beam, using a Xenon lamp (1600 W) equipped with a Bausch and Lomb monochromator. The incident photon flux in solution, measured by ferrioxalate actinometry, was 1.2\( \times 10^{-6} \) Einstein L⁻¹ s⁻¹.

The choice of the lamps had the purpose of exciting each photosensitiser as selectively as possible. The direct photolysis of IBP was studied under a lamp having maximum emission in the UVB region, upon consideration of the IBP absorption spectrum. The emission spectra of the lamps were taken with an Ocean Optics SD 2000 CCD spectrophotometer and normalised to the actinometry results, also taking into account the absorbance of the Pyrex glass walls of the irradiation cells. Note that, irrespective of the presence of the Pyrex glass, no lamp emitted radiation.
below 300 nm. The absorption spectra of the relevant compounds were taken with a Varian Cary 100 Scan UV-Vis spectrophotometer. The various emission and absorption spectra are reported in Figure 1.

**Monitoring of IBP transformation**

After the scheduled irradiation time, the cells were withdrawn from the lamp and the irradiated solutions were analysed by high-performance liquid chromatography (HPLC-UV) to monitor the time evolution of IBP. The adopted Merck-Hitachi instrument was equipped with an autosampler AS2000A (100 µL sample volume), pumps L-6200 and L-6000 for high-pressure gradients, a reverse-phase column Merck LiChrocart RP-C18 packed with LiChrospher 100 RP-18 (125 mm × 4.6 mm × 5 µm), and a UV-Vis detector L-4200 (detection wavelength 279 nm). It was adopted an isocratic elution with a 80:20 mixture of CH$_3$OH:aqueous H$_3$PO$_4$ (pH 2.8) at a flow rate of 1.0 mL min$^{-1}$. The retention time of IBP was 3.3 minutes, the column dead time 0.90 min.

The time evolution of furfuryl alcohol to quantify the formation rate of $^{1}$O$_2$ under the yellow lamp was also monitored by HPLC-UV, as reported in Minella et al. (2011).

**Identification of IBP transformation intermediates**

Intermediate identification was carried out with gas chromatography coupled with mass spectrometry. To this purpose, aqueous solutions after irradiation were extracted with 3 mL dichloromethane, dehumidified with anhydrous Na$_2$SO$_4$ and evaporated to dryness. Each sample was reconstructed with 100 µL dichloromethane. The solution was transferred into a vial and injected into a capillary gas chromatograph (Agilent 6890) coupled with a mass spectrometer (Agilent 5973 inert). The injection system used was a Gerstel CIS4 PTV. Initial injection temperature was 40 °C, programmed at 5 °C/s; final temperature was 320 °C, held for 9 min. The injection volume was 2 µl in the splitless mode. The capillary column used was a HP-5MS, 30 m × 0.25 mm × 0.25 µm film thickness. Initial column temperature was 40 °C and was increased by 15 °C/min to 300 °C. The carrier gas was ultrapure He (1.0 mL/min; SIAD, Bergamo, Italy). The ionization source worked in the electronic impact (EI) mode and the mass spectrometer worked in the Scan mode from 44 to 450 Th. Identification of spectra was performed by using the Wiley 7n library (Agilent Part No. G1035B).
Kinetic data treatment

Reaction rates were determined by fitting the time evolution data of IBP with pseudo-first order equations of the form \( C_t = C_o \exp(-k t) \), where \( C_t \) is the concentration of IBP at the irradiation time \( t \), \( C_o \) its initial concentration and \( k \) the pseudo-first order degradation rate constant. The initial degradation rate is \( R_{DCNP} = k C_o \). The reported errors on the rates (\( \pm \sigma \)) were derived by curve fitting and depend on the scattering of the experimental data around the fitting curve. The reproducibility of repeated runs was around 10-15%. The data plots, the fits and the numerical integrations to determine the absorbed photon fluxes were all carried out with the Fig.P software package (BIOSOFT, Cambridge, UK).

Results and discussion

Kinetics of IBP photochemical transformation

Direct photolysis. IBP (initial concentration 20 µM) was irradiated under the TL 01 lamp (emission maximum at 313 nm, see Figure 1) at pH 2 and 8. The time evolution of IBP under the adopted experimental set-up is reported in Figure S1-SM (SM = Supplementary Material). Note that significant IBP degradation was obtained at both pH values and that the adopted lamp did not emit radiation below 300 nm. Such results are in agreement with Matamoros et al. (2009). The transformation rate was \( \text{Rate}_{IBP} = (8.84 \pm 1.07) \times 10^{-11} \text{ M s}^{-1} \) at pH 2 and \( (4.28 \pm 0.65) \times 10^{-11} \text{ M s}^{-1} \) at pH 8. The photon flux absorbed by IBP can be expressed as

\[
P^a_{IBP} = \int_{\lambda} p^\gamma(\lambda) \left[ 1 - 10^{-\epsilon_{IBP}(\lambda) b[IBP]} \right] d\lambda,
\]

where \( p^\gamma(\lambda) \) is the incident spectral photon flux density of the lamp, \( \epsilon_{IBP}(\lambda) \) the molar absorption coefficient of IBP at pH 2 (neutral form) or pH 8 (anionic one, see Figure 1), \( b \) the optical path length in solution (0.4 cm), and \([IBP]\) the initial IBP concentration (20 µM). One gets \( P^a_{IBP} = 8.73 \times 10^{-11} \text{ einstein L}^{-1} \text{ s}^{-1} \) at pH 2 and \( 1.30 \times 10^{-10} \text{ einstein L}^{-1} \text{ s}^{-1} \) at pH 8. From these data it is possible to obtain the polychromatic photolysis quantum yield of IBP, \( \Phi_{IBP} \), in the UVB region where the spectra of the lamp and IBP overlap. It is \( \Phi_{IBP} = \text{Rate}_{IBP} (P^a_{IBP})^{-1} = 1.01 \pm 0.12 \) at pH 2 and \( 0.33 \pm 0.05 \) at pH 8 (\( \mu \pm \sigma \)). The latter datum is not too far from the quantum yield value of 0.2 obtained by Yuan et al. (2009) upon UVC irradiation of ibuprofen at pH 7. This high quantum yield of photolysis is consistent with the observed photodecarboxylation reactions (\textit{vide infra}), which are reported to be very efficient photolysis pathways (Budac and Wan, 1992).
**Reaction with •OH.** The reaction rate constant between IBP and •OH was determined upon competition kinetics with 2-propanol, using nitrate photolysis as the •OH source. The main reactions that are expected to take place in the system under consideration are the following (Mack and Bolton, 1999; Buxton et al., 1988):

\[
\begin{align*}
\text{NO}_3^- + \text{hv} + \text{H}^+ & \rightarrow \cdot\text{OH} + \cdot\text{NO}_2 & \left[\Phi_1 = 0.01\right] & (1) \\
2\text{-Propanol} + \cdot\text{OH} & \rightarrow \text{Products} & \left[k_2 = 1.9 \times 10^9 \text{ M}^{-1} \text{s}^{-1}\right] & (2) \\
\text{IBP} + \cdot\text{OH} & \rightarrow \text{Products} & \left[k_3\right] & (3)
\end{align*}
\]

Upon application of the steady-state approximation to [•OH], one gets the following expression for the initial transformation rate of IBP in the presence of 2-propanol:

\[
\text{Rate}_{\text{IBP}} = \frac{R_{\cdot\text{OH}} \cdot k_3 \cdot [\text{IBP}]}{k_3 \cdot [\text{IBP}] + k_2 \cdot [2\text{-Propanol}]} \quad (4)
\]

Figure 2 reports \(\text{Rate}_{\text{IBP}}\) as a function of the concentration of 2-propanol, upon irradiation of 10 mM NaNO\(_3\) + 20 µM IBP at pH 2 (open squares) and at pH 8 (solid stars). For each pH value it is also reported the fit curve of equation (4) to the experimental data and the two 95% confidence bands (dotted for pH 2, dashed for pH 8).

It is evident that in both cases the alcohol inhibited the transformation of IBP, which is consistent with the scavenging of •OH in reaction (2). Under the adopted conditions (irradiation time up to 4 h) the rate of IBP direct photolysis could be neglected. The experimental rate data were fitted with equation (4), fixing the values of \(k_2\) and [IBP] and letting \(R_{\cdot\text{OH}}\) and \(k_3\) vary. From the fit we obtained \(k_3 = (1.5 \pm 0.1) \times 10^{10} \text{ M}^{-1} \text{s}^{-1}\) at pH 2 and \((1.0 \pm 0.3) \times 10^{10} \text{ M}^{-1} \text{s}^{-1}\) at pH 8. These values are very near previous results by Parij et al. (1995). Packer et al. (2003) have obtained a somewhat different value, \(k_3 = (6.5 \pm 0.2) \times 10^9 \text{ M}^{-1} \text{s}^{-1}\) at pH 3.5 upon adoption of the Fenton reaction as •OH source. Considering that the Fenton process yields other oxidising species (e.g. ferryl) in addition to •OH (Bossmann et al., 2004; Prousek, 2007), which are usually less reactive than •OH itself, it cannot be excluded that use of the Fenton reaction could lead to a slight underestimation of the •OH rate constant.
Reaction with CO$_3$$^\cdot$. A semi-quantitative assessment of the reactivity of a substrate with CO$_3$$^\cdot$ can be carried out by studying the effect of bicarbonate on the transformation photoinduced by nitrate (Vione et al., 2009a). Figure 3 reports the initial transformation rate of 20 µM IBP upon irradiation of 10 mM NaNO$_3$, with the addition of variable NaHCO$_3$ concentrations. In the presence of bicarbonate up to 10 mM, the solution pH gradually increased from 6.5 to 8.5. The trend of the IBP rate with irradiated nitrate is also shown upon addition of a phosphate buffer (NaH$_2$PO$_4$ + Na$_2$HPO$_4$). The total concentration of phosphate and the ratio NaH$_2$PO$_4$ / Na$_2$HPO$_4$ were chosen to have the same concentration values as for NaHCO$_3$, and the same pH within ±0.1 units. Finally, the transformation rate of IBP in the presence of NaHCO$_3$ without nitrate is also reported.

From Figure 3 it can be derived that: (i) the direct photolysis of IBP (irradiation without nitrate) was negligible under the adopted experimental conditions; (ii) bicarbonate inhibited the transformation of IBP upon nitrate irradiation, while phosphate had practically no effect. Nitrate photolysis yields •OH (reaction 1), and the addition of NaHCO$_3$ induces the production of CO$_3$$^\cdot$ by reaction between the hydroxyl radical and HCO$_3^\cdot$ / CO$_3^{2\cdot}$ (Buxton et al., 1988). There would be competition between IBP and HCO$_3^\cdot$ / CO$_3^{2\cdot}$ for reaction with •OH, but CO$_3$$^\cdot$ (although it is less reactive than •OH) could also contribute to the transformation of IBP. Vione et al. (2009a) have found that the formation rate of •OH + CO$_3$$^\cdot$ by irradiation of nitrate and bicarbonate is higher than the rate of •OH formation with nitrate alone. A likely explanation is that •OH and •NO$_2$, photogenerated upon nitrate photolysis, undergo recombination to nitrate and H$^+$ when still in the solvent cage and before diffusing to the solution bulk. An excess bicarbonate could react with cage •OH and inhibit recombination, thereby enhancing the production of reactive species upon nitrate photolysis (reactions (5-8), Bouillon and Miller, 2005; Nissenson et al., 2010).

\[
\begin{align*}
\text{NO}_3^- + h\nu + H^+ & \rightarrow [\cdot\text{OH} + \cdot\text{NO}_2]_{\text{cage}} \quad (5) \\
[\cdot\text{OH} + \cdot\text{NO}_2]_{\text{cage}} & \rightarrow \cdot\text{OH} + \cdot\text{NO}_2 \quad (6) \\
[\cdot\text{OH} + \cdot\text{NO}_2]_{\text{cage}} & \rightarrow \text{NO}_3^- + H^+ \quad (7) \\
[\cdot\text{OH} + \cdot\text{NO}_2]_{\text{cage}} + \text{HCO}_3^- & \rightarrow \text{CO}_3^\cdot + \text{H}_2\text{O} + \cdot\text{NO}_2 \quad (8)
\end{align*}
\]

Because it induces the formation of a higher amount of a less reactive species, the addition of bicarbonate to irradiated nitrate enhances the transformation of the compounds that show a ratio of the rate constants with CO$_3$$^\cdot$ and •OH higher than 0.01 (Vione et al., 2009a). The same compounds could also undergo significant transformation by CO$_3$$^\cdot$ in the environment (Vione et al., 2009b). In contrast, inhibition of transformation by bicarbonate is observed for molecules that would also
undergo insignificant reaction with $\text{CO}_3^{-\bullet}$ in surface waters, such as 4-nitrophenol. However, considering that the addition of bicarbonate would also alter the solution pH, with implications for nitrate photolysis and possibly for substrate reactivity, enhancement or inhibition by bicarbonate are defined by comparison with phosphate buffers at the same pH values (Vione et al., 2009a).

In the case of IBP, the inhibition of transformation by bicarbonate compared to phosphate suggests that the reaction between IBP and $\text{CO}_3^{-\bullet}$ in surface waters would not be important (the relevant second-order rate constant is expected to be significantly below $10^8 \text{M}^{-1} \text{s}^{-1}$).

**Reaction with $^1\text{O}_2$.** Figure 4 reports the initial transformation rate of IBP, as a function of its initial concentration, upon irradiation of 10 µM Rose Bengal (RB) adopted as a source of $^1\text{O}_2$ (reaction (9); Miller, 2005). Irradiation was carried out only at pH 8, because at pH 2 RB is protonated and unable to absorb visible radiation to a significant extent. The reaction (10) between IBP and $^1\text{O}_2$ would be in competition with the thermal deactivation of singlet oxygen (reaction (11); Rodgers and Snowden, 1982):

$$
\text{RB} + \text{hv} + \text{O}_2 \rightarrow \text{RB} + ^1\text{O}_2 \quad (9)
$$

$$
\text{IBP} + ^1\text{O}_2 \rightarrow \text{Products} \quad [k_{10}] \quad (10)
$$

$$
^1\text{O}_2 \rightarrow \text{O}_2 + \text{heat} \quad [k_{11} = 2.5 \cdot 10^5 \text{s}^{-1}] \quad (11)
$$

Upon application of the steady-state approximation to $^1\text{O}_2$ one gets the following expression for the initial transformation rate of IBP ($\text{Rate}_{\text{IBP}}$):

$$
\text{Rate}_{\text{IBP}} = \frac{R_{1\text{O}_2} \cdot k_{10} \cdot [\text{IBP}]}{k_{11} + k_{10} \cdot [\text{IBP}]} \quad (12)
$$

where $R_{1\text{O}_2}$ is the formation rate of $^1\text{O}_2$ by 10 µM RB under the adopted irradiation device. The measurement of $R_{1\text{O}_2}$ was carried out upon irradiation at pH 8 of 10 µM RB + 0.1 mM furfuryl alcohol (FFA), which reacts with $^1\text{O}_2$ with a rate constant $k_{\text{FFA}} = 1.2 \cdot 10^8 \text{M}^{-1} \text{s}^{-1}$ (Haag et al., 1984). The initial transformation rate of FFA under the adopted conditions was $\text{Rate}_{\text{FFA}} = (1.26 \pm 0.06) \cdot 10^{-7} \text{M s}^{-1}$. Photogenerated $^1\text{O}_2$ could undergo deactivation or reaction with FFA, and upon application of the steady-state approximation to $[^1\text{O}_2]$ one gets:

$$
R_{1\text{O}_2} = \text{Rate}_{\text{FFA}} \frac{k_{11} + k_{\text{FFA}} \cdot [\text{FFA}]}{k_{\text{FFA}} \cdot [\text{FFA}]} \quad (13)
$$
From equation (13) one gets $R_{1O2} = (2.75 \pm 0.13) \cdot 10^{-6}$ M s$^{-1}$. The linear trend of $Rate_{IBP}$ vs. $[IBP]$ in Figure 4 suggests that, in equation (12), $k_{10} \ [IBP] \ll k_{11}$. If this approximation holds, one gets $Rate_{IBP} = R_{1O2} \ (k_{11})^{-1} \ [IBP]$. From Figure 4 one obtains $Rate_{IBP} = (6.00 \pm 0.33) \cdot 10^{-7}$ [IBP], and from the known values of $R_{1O2}$ and $k_{11}$ one gets $k_{10} = (6.00 \pm 0.58) \cdot 10^{-4}$ M$^{-1}$ s$^{-1}$ as the reaction rate constant between IBP and $^{1}O_2$. This result confirms that the hypothesis $k_{10} \ [IBP] \ll k_{11}$ was justified.

**Reaction with irradiated AQ2S.** The choice of AQ2S is motivated by the fact that this compound has a reactive triplet state but it does not yield either $^{•}$OH or $^{1}O_2$ under irradiation, thus avoiding formation of potentially interfering transients (Loeff et al., 1983; Maurino et al., 2008, Maddigapu et al., 2010c). Figure 5 reports the initial transformation rate of IBP as a function of its initial concentration, upon UVA irradiation of 0.1 mM AQ2S at pH 2 and pH 8. From the linear trend of the plot one gets $Rate_{IBP} = (4.66 \pm 0.17) \cdot 10^{-4}$ [IBP] at pH 2 and $(3.19 \pm 0.09) \cdot 10^{-4}$ [IBP] at pH 8. AQ2S would be by far the main radiation absorber at both pH values. From the absorption spectra (Figure 1), one gets that at 330 nm the absorbance of 0.1 mM AQ2S would be $10^4$ times or more higher than that of IBP at the highest adopted concentration value (10 µM). Therefore, it is possible to simply determine the photon flux absorbed by AQ2S as $P_a^{AQ2S} = \int \sigma(\lambda) [1 - 10^{-\varepsilon_{AQ2S}(\lambda)ib[AQ2S]}] d\lambda = 2.01 \cdot 10^{-6}$ einstein L$^{-1}$ s$^{-1}$ ($\sigma(\lambda)$ is the spectral incident photon flux density of the UVA lamp and $b = 0.4$ cm). The polychromatic quantum yield of IBP phototransformation by AQ2S would thus be $\Phi_{IBP,AQ2S} = Rate_{IBP} (P_a^{AQ2S})^{-1} = (2.32 \pm 0.08) \cdot 10^2$ [IBP] at pH 2 and $(1.59 \pm 0.04) \cdot 10^2$ [IBP] at pH 8.

The reactive triplet state $^{3}AQ2S^*$ accounts for the degradation processes that take place with AQ2S under irradiation. It has a formation quantum yield $\Phi_{3AQ2S^*} = 0.18$ and a deactivation rate constant $k_{3AQ2S^*} = 1.1 \cdot 10^7$ s$^{-1}$ (Loeff et al., 1983). The formation rate of $^{3}AQ2S^*$ would be $R_{3AQ2S^*} = \Phi_{3AQ2S^*} \cdot P_a^{AQ2S}$, and its deactivation would be in competition with the reaction with IBP (rate constant $k_{3AQ2S^*,IBP}$). Therefore, the transformation rate of IBP by irradiated AQ2S could be expressed as follows:

$$Rate_{IBP} = \Phi_{3AQ2S^*} \cdot P_a^{AQ2S} \cdot \frac{k_{3AQ2S^*,IBP} \cdot [IBP]}{k_{3AQ2S^*} + k_{3AQ2S^*,IBP} \cdot [IBP]}$$

(14)

Under the hypothesis that $k_{3AQ2S^*,IBP} \ [IBP] \ll k_{3AQ2S^*}$, one gets $Rate_{IBP} = \Phi_{3AQ2S^*} \cdot P_a^{AQ2S} \cdot k_{3AQ2S^*,IBP} (k_{3AQ2S^*})^{-1} \ [IBP]$, which is compatible with the linear trend reported in Figure 5. It is also $\Phi_{IBP,AQ2S}$
Rate_{IBP} (P_a^{AQ2S})^{-1} = \Phi_{3AQ2S^{*}} k_{3AQ2S^{*},IBP} (k_{3AQ2S^{*}})^{-1} [IBP]. By comparison with the experimental data, \(\Phi_{IBP,AQ2S} = (2.32\pm0.08)\times10^{2} \) [IBP] at pH 2 and \((1.59\pm0.04)\times10^{2}\) [IBP] at pH 8, one gets \(k_{3AQ2S^{*},IBP} = (1.42\pm0.05)\times10^{10}\) and \((9.70\pm0.24)\times10^{9}\) M\(^{-1}\) s\(^{-1}\) at pH 2 and 8, respectively. This finding confirms that the hypothesis \(k_{3AQ2S^{*},IBP} \ll k_{3AQ2S^{*}}\) was reasonable.

**Reaction with irradiated riboflavin.** Riboflavin is another well-known sensitiser. It produces \(^1\text{O}_2\) under irradiation and its excited triplet state (\(^3\text{Ri}^{*}\)) also reacts through electron transfer with electron-donor molecules (Barbieri et al., 2008). The solutions containing riboflavin (5.4 µM) and IBP (10 µM) were buffered at pH 8 and irradiated in aerated solution for 1 h, to limit riboflavin loss to 10%. The rate of IBP consumption was quite small: \(5.5\times10^{-11}\) M s\(^{-1}\). Considering that the absorbance of riboflavin at 365 nm is 0.053, the incident photon flux is \(1.2\times10^{-6}\) Einstein L\(^{-1}\) s\(^{-1}\) and the quantum yield of \(^3\text{Ri}^{*}\) formation is 0.375 in neutral water (Islam et al., 2003), the rate of \(^3\text{Ri}^{*}\) production in our system would be \(5.1\times10^{-8}\) M s\(^{-1}\). In another set of experiments, we observed that \(^3\text{Ri}^{*}\) is efficiently quenched by oxygen with a rate constant of \(1.06\times10^{9}\) M\(^{-1}\) s\(^{-1}\). The pseudo first-order rate constant of \(^3\text{Ri}^{*}\) deactivation (\(k_{3\text{Ri}^{*}}\)) in neutral deoxygenated aqueous solution is \(1.5\times10^{5}\) s\(^{-1}\) (Kouras et al., unpublished results), which raises to \(4.2\times10^{5}\) s\(^{-1}\) in aerated systems because of the reaction between \(^3\text{Ri}^{*}\) and \(^1\text{O}_2\). When considering both the reaction between \(^3\text{Ri}^{*}\) and IBP and the production of \(^1\text{O}_2\) by \(^3\text{Ri}^{*}\), followed by reaction of \(^1\text{O}_2\) with IBP, it is possible to write the rate of IBP transformation as follows:

\[
Rate_{IBP} = 5.1\times10^{-8}\cdot\left(\frac{k_{\text{Ri}^{*},IBP} \cdot [IBP]}{4.2\times10^{5} + k_{\text{Ri}^{*},IBP} \cdot [IBP]} + \frac{27\times10^{5}}{4.2\times10^{7} k_{10} + k_{10} \cdot [IBP]}\right)
\]  

The first term in the bracket corresponds to the oxidation of IBP by \(^3\text{Ri}^{*}\) and the second to the oxidation of IBP by \(^1\text{O}_2\). Using \(k_{10} = (6.00\pm0.58)\times10^{4}\) M\(^{-1}\) s\(^{-1}\) and \(k_{11} = 2.5\times10^{5}\) s\(^{-1}\), the rate of IBP consumption through \(^1\text{O}_2\) is equal to \((7.9\pm0.8)\times10^{-14}\) M s\(^{-1}\) and is thus negligible. This means that in our experimental conditions IBP mainly disappears by direct reaction with \(^3\text{Ri}^{*}\). The rate constant of the reaction \(k_{3\text{Ri}^{*},IBP}\) is estimated to be \((4.5\pm0.4)\times10^{7}\) M\(^{-1}\) s\(^{-1}\). This value is two orders of magnitude lower than that measured in the case of AQ2S.
The availability of the direct photolysis quantum yield and of the reaction rate constants of the anionic IBP form with •OH, 1O2 and 3AQ2S*/3Ri* (proxies for 3CDOM*) allow the calculation of the first-order rate constant of IBP transformation (kIBP) as a function of water chemical composition and column depth. The model approach to link substrate reactivity and environmental features is described in detail elsewhere (Albinet et al., 2010a; Maddigapu et al., 2010a; Vione et al., 2010; Hatipoglu et al., 2010; Maddigapu et al., 2011). Figure S2-SM reports the sunlight spectrum adopted for the calculations, which corresponds to a 22 W m\(^{-2}\) irradiance in the UV (Frank and Klöpffer, 1988). A major issue for the assessment of outdoor reactivity is that sunlight irradiance is not constant. The time unit adopted in the cited model is the summer sunny day (SSD), which corresponds to a fair-weather 15 July at 45°N latitude. The incident UV energy in a SSD is equivalent to 10 hours continuous irradiation at 22 W m\(^{-2}\) UV irradiance (Maddigapu et al., 2010b). When applying the model, the first-order rate constant kIBP of IBP phototransformation is given in units of SSD\(^{-1}\), while the half-life times \(\tau_{IBP} = \ln 2 \times (k_{IBP})^{-1}\) have units of SSD.

Inclusion in the model of the IBP kinetic data obtained here suggests that direct photolysis and reactions with •OH and possibly 3CDOM* would be the main IBP transformation pathways in surface waters. Under the hypothesis that \(k_{3CDOM*,IBP} \sim k_{3AQ2S*,IBP} = 9.7 \times 10^9 \text{ M}^{-1} \text{s}^{-1}\), with low dissolved organic carbon (DOC, expressed as NPOC, Non-Purgeable Organic Carbon) the order of importance of the three relevant pathways would be •OH > photolysis > 3CDOM*. At high NPOC it would be 3CDOM* > photolysis > •OH. Such a scenario could be compatible with previous results of IBP photodegradation in surface water samples under irradiation, which suggest that reaction with •OH would not be the only relevant process (Packer et al., 2003). More recently, significant reactivity has been found between IBP and fulvic acids under irradiation (Jacobs et al., 2011). On the other hand, taking \(k_{3CDOM*,IBP} \sim k_{3Ri*,IBP} = 4.5 \times 10^7 \text{ M}^{-1} \text{s}^{-1}\), the reaction with 3CDOM* would be minor even at high NPOC.

Figure 6 reports the model results for kIBP as a function of nitrite and NPOC, with fixed nitrate (51 µM), bicarbonate (2.1 mM) and carbonate (26 µM) and with a water column depth of 1 m. In Figure 6a it is hypothesised that IBP reacts with 3CDOM* with \(k_{3CDOM*,IBP} = 9.7 \times 10^9 \text{ M}^{-1} \text{s}^{-1}\), following the results obtained with AQ2S. In Figure 6b it is hypothesised \(k_{3CDOM*,IBP} = 4.5 \times 10^7 \text{ M}^{-1} \text{s}^{-1}\), in analogy with the riboflavin data. As far as Figure 6a is concerned, at low nitrite kIBP has a minimum for 1 mg C L\(^{-1}\) NPOC. In the system described by Figure 6a nitrite would be the main •OH source (with the exclusion of the highest NPOC values, where CDOM would prevail as
source), DOM the main $^\text{•OH}$ sink, and $^3\text{CDOM}^*$ would obviously be produced by organic matter alone. At low NPOC, the $k_{\text{IBP}}$ decrease with NPOC is due to $^\text{•OH}$ scavenging by DOM. The increase of $k_{\text{IBP}}$ after the minimum would be accounted for by the role of $^3\text{CDOM}^*$ in the transformation of IBP. If the reaction between IBP and $^3\text{CDOM}^*$ is negligible, a continuous decrease of $k_{\text{IBP}}$ with increasing NPOC is expected as shown in Figure 6b. In the latter case, DOM would scavenge $^\text{•OH}$ and its chromophoric moieties would compete with IBP for radiation absorption, thereby inhibiting the IBP direct photolysis. The data of Figure 6a foresee an IBP half-life time $\tau_{\text{IBP}} = \ln 2 (k_{\text{IBP}})^{-1}$ of less than one week, while those of Figure 6b suggest that the half-life time could vary from a few days to a couple of months. Also note that the adopted column depth of 1 m is quite favourable to photochemistry.

An assessment of the importance of the reaction between IBP and $^3\text{CDOM}^*$ can be obtained by application of our model to the IBP degradation kinetics reported by Packer et al. (2003) upon irradiation of Mississippi river water. Water data were 63 $\mu$M nitrate, 8.9 mg C L$^{-1}$ NPOC, bicarbonate and carbonate not reported (but they are minor $^\text{•OH}$ sinks and the reaction between IBP and $\text{CO}_3^{\text{−•}}$ is negligible) (Packer et al., 2003). The solutions have been placed in quartz bottles with diameter $d$ = 4.0 cm, which would yield a geometrical slab-equivalent optical path length $b = 2 d / \pi = 2.5$ cm (Albinet et al., 2010b). Note however that, for such small $b$ values where the inner-filter effect of the solution is expected to be small, the IBP phototransformation kinetics would depend very little on the path length. The adopted lamp was approximately five times more intense than sunlight. With this apparatus, Packer et al. (2003) obtained a first-order rate constant of 0.45 day$^{-1}$, which would become 0.09 day$^{-1}$ under normal sunlight. With the input data reported above, our model yielded $k_{\text{IBP}} = 0.24$ SSD$^{-1}$ with $k_{^3\text{CDOM}^*,\text{IBP}} = 4.5 \cdot 10^7$ M$^{-1}$ s$^{-1}$ and 0.73 SSD$^{-1}$ with $k_{^3\text{CDOM}^*,\text{IBP}} = 9.7 \cdot 10^9$ M$^{-1}$ s$^{-1}$. Under the hypothesis that 1 SSD = 1 day of Packer et al., it is quite likely that CDOM in Mississippi river water is poorly reactive toward IBP. Moreover, our model foresees that direct photolysis would be the main transformation reaction in the system, with $^\text{•OH}$ only contributing to 0.02 SSD$^{-1}$. Note that the model adopts the sunlight spectrum reported in Figure S2-SM, which might be significantly different from that of the lamp used by Packer et al. (medium-pressure Hg-vapour lamp filtered by borosilicate glass), in particular at the wavelengths around 300 nm that are the most significant for the direct photolysis of IBP.

It is also possible to compare the IBP elimination rate constant in late summer months, derived from the field data of Tixier et al. (2003) in the epilimnion of the Swiss Lake Greifensee (5 m depth) with the model results, to get insight into the potential importance of photochemistry in the fate of IBP in surface waters. Tixier et al. (2003) obtained a field rate constant of 0.022 day$^{-1}$,
which would include all possible processes such as outflow, sedimentation, •OH-induced photodegradation and possibly biodegradation. Note that the known processes accounted for approximately one half of the observed transformation. With the Lake Greinfensee data (0.1 mM nitrate, 3.5 mg C L\(^{-1}\) NPOC, 2 mM bicarbonate, 10 µM carbonate) (Canonica et al., 2005) and \(d = 5\) m, one obtains from our model \(k_{\text{IBP}} \sim 0.030\) SSD\(^{-1}\) for the sum of •OH reaction and direct photolysis, of which 0.024 SSD\(^{-1}\) would be accounted for by direct photolysis alone. Tixier et al. (2003) report that unfavourable meteorology would approximately halve the importance of photochemistry in the period under study, while our model hypothesizes constant good weather. By halving \(k_{\text{IBP}}\) and its two components (•OH and photolysis) in our model, one gets a direct photolysis contribution of 0.012 SSD\(^{-1}\). Under the hypothesis that 1 SSD = 1 day, one gets a reasonable match between model results concerning direct photolysis (0.012 SSD\(^{-1}\)) and the IBP degradation kinetics in Lake Greinfensee that could not be accounted for by the known processes (0.006 – 0.011 day\(^{-1}\)). Such a comparison suggests that direct photolysis might be a significant pathway for IBP transformation in the lake epilimnion during the summer months.

**Identification of IBP phototransformation intermediates**

To identify the intermediates of the potentially most important photochemical transformation pathways of IBP in surface waters (direct photolysis, •OH, \(^3\)AQ2S*), a significantly higher initial IBP concentration (1 mM) was used than in the kinetic studies. It was adopted pH 8, adjusted with NaOH. Other conditions were as follows: 1 M NaNO\(_3\) for •OH, 1 mM AQ2S for triplet state reactivity. The adopted irradiation times were up to 6 h for •OH, up to 3 h for \(^3\)AQ2S*, and up to 3 days for the direct photolysis. Table 1 reports the IBP transformation intermediates identified by GC-MS following the different pathways. Experimental mass spectra and the comparisons with the spectra libraries are reported as SM. The direct photolysis of IBP yielded III and IV as detected phototransformation intermediates. Interestingly, the same compounds have been detected upon 254-nm irradiation of IBP (Szabó et al., 2011), which is reasonable because both UVC irradiation and our conditions (UVB) would excite the highest-wavelength absorption band of IBP. Compound III was also detected in the presence of •OH, and IV was detected in all the three pathways under study (•OH, direct photolysis and \(^3\)AQ2S*). Considering that all the three systems were irradiated, direct photolysis could be operational at some extent also in the presence of nitrate and AQ2S. However, III has been identified upon IBP transformation in photo-Fenton systems (Méndez-Arriaga et al., 2010), where it probably originated by •OH substitution at the carbon in alpha to the
carboxylic group (probably via H abstraction followed by \(^\cdot\)OH addition) followed by decarboxylation/oxidation. Moreover, we detected IV in the system IBP + AQ2S that was irradiated under the UVA lamp, where the direct photolysis of IBP should be low to negligible. The same compound has been detected by Jacobs et al. (2011) in the presence of irradiated fulvic acids, under conditions where the direct photolysis of IBP was negligible.

Compound I was identified with IBP + \(^3\)AQ2S\(^*\), upon transformation of the lateral chain containing the carboxylic group. Transformation of the COOH-containing (propanoic) chain was also observed for III and IV (already discussed). Compound II, only detected in the presence of \(^3\)AQ2S\(^*\), would be produced upon shortening of both lateral chains of IBP. Interestingly, this is the only detected intermediate that underwent transformation of the isobutyl chain, which is thus expected to be much less reactive compared to the propanoic one.

The identification of V (only formed with \(^3\)AQ2S\(^*\)) was only tentative, because of the lack of the relevant mass spectrum in the adopted library. Identification was mainly based on the mass of the molecular ion that yielded the likely chemical formula C\(_{13}\)H\(_{18}\)O, namely an oxygen atom less than IBP. If this is the case, V would arise from IBP + \(^3\)AQ2S\(^*\) following a reductive pathway, which might possibly involve the reduced radical species AQ2S\(^{-}\) and/or AQ2S-H\(^*\), or superoxide/hydroperoxide. The latter may be formed via the following reactions (Maurino et al., 2008; Maddigapu et al., 2010c; Maurino et al., 2011) (ISC = Inter-System Crossing, S-H = substrate):

\[
\begin{align*}
AQ2S + h\nu & \xrightarrow{ISC} \ ^3AQ2S^* \quad (16) \\
\ ^3AQ2S^* \rightarrow AQ2S \text{ or products} & \quad (17) \\
\ ^3AQ2S^* + AQ2S \rightarrow AQ2S^{-}\ + AQ2S^{**} & \quad (18) \\
\ ^3AQ2S^* + S-H \rightarrow AQ2S-H^* + S^* & \quad (19) \\
\ ^3AQ2S^* + S-H \rightarrow AQ2S^{-}\ + S-H^{**} & \quad (20) \\
AQ2S^{-}\ + O_2 \rightarrow AQ2S + O_2^{-}\ & \quad (21) \\
AQ2S-H^* + O_2 \rightarrow AQ2S + HO_2^* & \quad (22)
\end{align*}
\]

Scheme 1 summarises the transformation pathways leading to the detected IBP intermediates. The dashed arrows connecting IBP with III and IV via \(^3\)AQ2S\(^*\) and \(^\cdot\)OH account for the theoretical possibility for them to be produced by IBP direct photolysis when nitrate and AQ2S are irradiated. Interestingly, the transformation intermediates detected in the present study do not match those identified upon IBP degradation with irradiated TiO\(_2\) (Méndez-Arriaga et al., 2008), which suggest
that IBP in surface waters and upon TiO$_2$ photocatalysis would follow quite different reaction pathways.

**Conclusions**

- Photochemistry can be an important transformation process of IBP in surface waters. The prevailing pathways involved are the reaction with $^\cdot$OH, the direct photolysis and possibly, depending on IBP reactivity, transformation induced by $^3$CDOM$^\ast$.
- Anionic IBP, which prevails at the pH values of surface waters has a polychromatic UVB photolysis quantum yield $\Phi_{IBP} = 0.33\pm0.05$, a reaction rate constant with $^\cdot$OH $k_{IBP,\cdot OH} = (1.0\pm0.3)\times10^{10}$ M$^{-1}$ s$^{-1}$, a $^1$O$_2$ rate constant $k_{IBP,^1O2} = (6.00\pm0.58)\times10^4$ M$^{-1}$ s$^{-1}$, a $^3$AQ2S$^\ast$ rate constant $k_{IBP,^3AQ2S^\ast} = (9.70\pm0.24)\times10^9$ M$^{-1}$ s$^{-1}$, and a $^3$Ri$^\ast$ rate constant $k_{IBP,^3Ri^\ast} = (4.5\pm0.4)\times10^7$ M$^{-1}$ s$^{-1}$. Moreover, IBP does not react with CO$_3^{\cdot-}$ to a significant extent. The results obtained with AQ2S and riboflavin as model systems for CDOM are quite contrasting, showing that the reactivity of IBP with oxidant triplets is greatly influenced by the triplet nature.
- We compared our model predictions with literature data of irradiation of Mississippi water samples and with a field study carried out in Lake Greinfensee, Switzerland. In both cases it is suggested a poor reactivity of $^3$CDOM$^\ast$ toward IBP and an important role of the direct photolysis.
- IBP phototransformation mainly takes place via lateral chain shortening, and the propanoic chain is much more reactive than the isobutyl one. The IBP aldehyde, which would arise upon IBP reduction, was also tentatively identified in the presence of AQ2S under irradiation.

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Table 1. IBP transformation intermediates detected under the different studied conditions. Near each formula it is reported the GC retention time ($t_R$) and the reaction pathway that generated the relevant intermediate. DP = direct photolysis, n/a = not applicable, (*) = tentative identification. The match between the experimental mass spectra and the library ones is reported in the Supplementary Material. Identification implied an overall mass spectrometric match $> 85\%$ and the match of the most abundant ions and of the molecular one.

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<th>Formation pathway</th>
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<td>14.72</td>
<td>$^3\text{AQ2S}^*$</td>
</tr>
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</table>
Figure 1. (a) Absorption spectrum of ibuprofen (IBP) at pH 2 and 8. Incident spectral photon flux density $p^0(\lambda)$ of the adopted TL 01 lamp (emission maximum at 313 nm).
(b) Absorption spectrum of Rose Bengal (RB). Incident spectral photon flux density of the yellow lamp (TL D 18W/16 Yellow).
(c) Absorption spectrum of antraquinone-2-sulphonate (AQ2S). Incident spectral photon flux density of the UVA lamp (TL K 05).
Figure 2. Initial transformation rates of 20 µM IBP upon irradiation of 10 mM NaNO₃, as a function of the concentration of added 2-propanol, at pH 2 (open squares, dotted curves) and 8 (solid stars, dashed curves). The pH values were adjusted with HClO₄ and NaOH, respectively. Note that there are three curves for each data set. The central curve is the fit one, the lateral curves are the 95% confidence bands. nM = 10⁻⁹ M. Irradiation was carried out under the TL 01 lamp.
Figure 3. Initial transformation rates upon irradiation of (□) 20 µM IBP and 10 mM NaNO₃, as a function of the concentration of NaHCO₃; (Δ) 20 µM IBP and 10 mM NaNO₃, as a function of the concentration of added phosphate buffer (same concentration as NaHCO₃ and same pH, within 0.1 units); (◊) 20 µM IBP without nitrate, as a function of NaHCO₃ concentration. nM = 10⁻⁹ M. Irradiation was carried out under the TL 01 lamp.
Figure 4. Initial transformation rates of IBP upon irradiation of 10 µM Rose Bengal (RB) under the yellow lamp (Philips TL D 18W/16), as a function of the IBP concentration. The solution pH was 8, adjusted with NaOH. The lamp and RB spectra are reported in Figure 1b. pM = 10^{-12} M. The regression line is dashed, the 95% confidence bands are dotted.
Figure 5. Initial transformation rates of IBP upon UVA irradiation of 1 mM AQ2S, as a function of the concentration of IBP at pH 2 and 8. The pH values were adjusted with HClO₄ and NaOH, respectively. The regression lines are dashed, the 95% confidence bands are dotted.
Figure 6. (a) First-order transformation rate constant of IBP, as a function of nitrite concentration and the NPOC values. Other conditions: 51 µM nitrate, 2.1 mM bicarbonate, 26 µM carbonate. Here it is hypothesised $k_{IBP,3CDOM^*} = 9.7 \times 10^9 \text{ M}^{-1} \text{ s}^{-1}$.

(b) First-order transformation rate constant of IBP, under the hypothesis that $k_{IBP,3CDOM^*} = 4.5 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$. Other conditions are same as before.
Scheme 1. A summary of the processes leading to the detected intermediates under the studied irradiation conditions. Note that identification of V is only tentative, as is the assignment of III and IV as IBP transformation intermediates upon reaction with •OH and/or $^3$AQ2S* (see dashed arrows). DP = direct photolysis; (*) = tentative identification.