

1 **REMOVAL AND ENVIRONMENTAL RISK ASSESSMENT OF**
2 **CONTAMINANTS OF EMERGING CONCERN FROM**
3 **IRRIGATION WATERS DURING MICROALGAE TREATMENT**

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55 **ABSTRACT**

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57 The present study evaluated the efficiency of a semi-closed, tubular, horizontal photobioreactor
58 (PBR) to treat a mixture of irrigation and rural drainage water, focusing in the removal of different
59 contaminants of emerging concern (CECs), and evaluating the environmental impact of the
60 resulting effluent. Target CECs included pharmaceuticals, personal care products and flame
61 retardants. Of the 13 compounds evaluated, 11 were detected in the feed water entering the PBR,
62 and diclofenac (DCF) (1,107 ng L⁻¹) and N,N-diethyl-toluamide (DEET) (699 ng L⁻¹) were those
63 present at the greatest concentrations. The best removal efficiencies were achieved for the
64 pharmaceuticals diazepam (94%), LZP (83%) and OXA (71%), and also for ibuprofen (70%). For
65 the rest of the CECs evaluated, attenuation was similar to that obtained after conventional
66 wastewater treatment, ranging from basically no elimination (CBZ and tris-(2-chloroethyl)
67 phosphate (TCEP)) to medium efficiencies (DCF and tributyl phosphate (TBP) (50%)).
68 Environmental risk assessment based on hazard quotients (HQs) resulted in HQ values <0.1 (no
69 risk associated) for most of the compounds and most of the trophic levels considered. Values
70 between 1 and 10 (moderate risk) were obtained for tonalide (fish) and CBZ (invertebrates). The
71 most sensitive trophic levels were green algae, whereas fish and aquatic plants were the most
72 resilient. Our results suggest that microalgae-based treatments could become a green, cost-
73 effective alternative to conventional wastewater treatment regarding the efficient elimination of
74 these contaminants.

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Keywords: personal care products, pharmaceuticals, microalgae, ecotoxicity, environmental hazard, green treatments

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93 **1. INTRODUCTION**
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95 Agricultural activities and animal feeding operations (without regulated slurry or manure
96 pits) are becoming more intensive in order to satisfy the also increasing food demand, leading to
97 a constant increase in the use of veterinary pharmaceuticals in cattle farming activities, and
98 inorganic fertilizers and/or synthetic pesticides in agriculture (Oerke, 2006; Popp et al., 2013).
99 This is causing relevant amounts of diffuse pollution affecting both surface and groundwater
100 systems (Dolliver and Gupta, 2008; García-Galán et al., 2010; Sabourin et al., 2009).
101 Furthermore, crops irrigation with reclaimed wastewater has become a common practice in
102 countries under a significant water scarcity (such as those in the Mediterranean area). Wastewater
103 effluents are considered as one of the main entrance pathways for a broad variety of organic
104 micropollutants into the aquatic environment, as these are not fully removed even after tertiary
105 and/or advanced treatments such as UV radiation, MBR, reverse osmosis (RO) or nanofiltration
106 (NF) (Biel-Maeso et al., 2018; Mamo et al., 2018; Racar et al., 2020). In consequence, this practice
107 can only contribute to increase the environmental occurrence of the so-called contaminants of
108 emerging concern (CECs), which include compounds such as pharmaceuticals and personal care
109 products (PPCPs), fundamental in our daily routine, but also high production volume chemicals
110 such as plasticizers, preservatives or flame retardants, which are frequently used in industrial
111 processes (Krzeminski et al., 2017; Loos et al., 2009; van Wezel et al., 2018). Currently, there is
112 no European legislation regarding reclaimed water quality. Spain is the European country with
113 the highest volume of wastewater reuse, and this practice is regulated by the RD1620/2007,
114 describing the water quality required depending on its final use. However, CECs are never
115 mentioned. Last of all, the application of cattle manure or biosolids from urban wastewater
116 treatment plants (WWTPs) as organic amendment should not be neglected, as these may still
117 contain residues of non-polar CECs (Langdon et al., 2010; Sabourin et al., 2009). Overall,
118 depending on the polarity of these pollutants, irrigation or storm events can lead to the
119 translocation of these CECs from the crop fields (Ccanccapa et al., 2016; Langdon et al., 2010;
120 Postigo et al., 2016). Drainage channels (and also open irrigation channels) can receive a large

121 amount of this rural run-off; these channels usually discharge into rivers, as their diversion into
122 main collectors towards WWTPs is usually unfeasible. Thus, these pollutants eventually spread
123 in aquatic ecosystems and may indirectly affect a huge variety of non-target species, endangering
124 the natural equilibrium of river and streams (García-Galan et al., 2017; Proia et al., 2013). For
125 instance, bioaccumulation of anti-inflammatories such as diclofenac (DCF) and ibuprofen (IBU)
126 has been observed in larvae of caddisflies and leeches at concentrations up to 183 ng g⁻¹ (Huerta
127 et al., 2015), and the anxiolytic oxazepam (OXA) was found in the freshwater shrimp *Gammarus*
128 *fossarum* at levels up to 200 ng g⁻¹ and 30 ng g⁻¹ respectively (Maria Jesus García-Galan et al.,
129 2017). The corroborated spread of antibiotic resistance genes and the endocrine disruption caused
130 by the plasticizer bisphenol A and other synthetic hormones in certain fish species are amongst
131 the most critical environmental issues to tackle nowadays (Cacace et al., 2019; Huerta et al.,
132 2016).

133 Nature-based, low-cost treatment systems, such as constructed wetlands (CWs) or
134 microalgae-based treatments, are gradually becoming a feasible and more appropriate alternative
135 to conventional WWTPs for small populations in rural areas. These alternative technologies are
136 being intensively investigated and, so far, promising results regarding CECs removal have been
137 observed, performing both as secondary and tertiary treatments (Ávila et al., 2014; García-Galán
138 et al., 2018; Matamoros et al., 2015; Vassalle et al., 2020a). Specifically, microalgae-based
139 treatments have received a renewed consideration due to their high efficiency removing nutrients
140 and organic matter within a more sustainable frame than conventional wastewater treatments.
141 Microalgae biomass grows fixating CO₂ and assimilating the nutrients (mostly nitrogen (N) and
142 phosphorus (P)) present in the influent wastewater. Oxygen is generated through photosynthesis
143 and used up by heterotrophic aerobic bacteria to degrade the organic matter present in the water
144 (including CECs). Microalgae systems have the dual capacity of treating wastewater efficiently
145 and producing microalgae biomass which, after an appropriate harvesting/separation technique
146 from the aqueous phase, can be further profited to produce bioenergy (biogas) or other added-
147 value products such as pigments, biofertilizers or even bioplastics (Arashiro et al., 2020; Khan et
148 al., 2019; Rueda et al., 2020). In consequence, if this biomass is managed properly, the waste

149 generated during microalgae treatment is considerably reduced, as well as the operation and
150 maintenance (O&M) costs when compared to conventional systems, as external aeration is no
151 required due to photosynthesis.

152 There are two basic types of microalgae treatment systems, open and closed reactors. Open
153 systems or high rate algal ponds (HRAPs) are the most frequently used systems, mainly due to
154 their lower O&M costs, but cultures are more exposed to external contamination, and the different
155 growth and environmental parameters (temperature, sunlight) can hardly be regulated (Park and
156 Craggs, 2010). Closed systems are presented as horizontal tubular photobioreactors (PBRs), as
157 vertical cylinders (column PBRs) or flat plate PBRs (consisting of flat, thin panels). They are
158 mostly used for commercial production of microalgae biomass (growing single, pure cultures), as
159 the biomass yields are typically higher, microalgae cultures are more protected against external
160 contamination and control of the operation parameters is better. Yet, the costs of O&M are higher
161 (higher energy requirements for mixing), dissolved oxygen (DO) may accumulate within the tubes
162 to toxic levels and biofouling may also appear in their inner walls. Recently, an innovative design
163 of a hybrid or semi-closed PBR (combining the advantages and avoiding the limitations of both
164 open and closed systems) have been tested, evaluating their efficiencies in wastewater
165 bioremediation and biomass yield (Díez-Montero et al., 2020) and also in the removal of different
166 antibiotics, sunscreens, plasticizers and pesticides (García-Galán et al., 2020b, 2018; Vassalle et
167 al., 2020b), with favorable outcomes. To the author's knowledge, the use of closed or semi-closed
168 PBRs is not frequent, as HRAPs are predominant in wastewater treatment systems.

169 The present study aims to investigate the capacity of a semi-closed, horizontal tubular PBR,
170 acting as a tertiary treatment, to remove 13 different CECs from irrigation water, including 6
171 pharmaceuticals, 4 personal care products, 2 flame retardants and one surfactant. The different
172 removal pathways within the PBR have been discussed, and the potential ecotoxicity of the PBR
173 effluent has been evaluated, estimating the risk quotients associated to the CECs.

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178 **2. MATERIALS AND METHODS**

179 **2.1. Description and operation of the semi-closed tubular horizontal PBR**

180 An innovative, new prototype of a semi-closed tubular horizontal PBR was conceived,
181 deployed and validated within the framework of the H2020 EU project INCOVER “Innovative
182 Eco-technologies for Resource Recovery from Wastewater” (<http://incover-project.eu/> GA
183 689242). Three PBRs were the core of a more complex pilot plant at demonstrative scale, which
184 main objective was to use agricultural drainage water and domestic wastewater as a valuable
185 resource to produce different added-value products. The plant was located in the Agròpolis
186 experimental campus of the Universitat Politècnica de Catalunya-BarcelonaTech (UPC), next to
187 the agricultural area of the Llobregat Delta that belongs to the Baix Llobregat Agrarian Park. The
188 park comprises 2900 Ha of fruit and vegetable crops located in the alluvial plains of the Llobregat
189 Delta and the lower valley of the Llobregat River (Montasell i Dorda and Callau i Berenguer,
190 2008).

191 A detailed description of the PBRs, the start-up of the plant and the main outcomes
192 regarding wastewater treatment can be found elsewhere (García et al., 2018; Uggetti et al., 2018).
193 Briefly, each PBR consisted of two open tanks of polypropylene connected by 16 horizontal tubes
194 (Figure 1). The useful volume of each PBR was 11.7 m³. Paddlewheels were installed in the
195 middle of each open tank to promote and favor the homogeneous distribution and mixing of the
196 liquor and also the release of the excess DO accumulated along the closed tubes. They also
197 contributed to create a water level difference (0.2 m) between both tanks, which made the mixed
198 liquor flow by gravity from one tank to the opposite one (Figure 1). The PBRs operated under a
199 hydraulic residence time (HRT) regime of 5 d (feeding of 2.3 m³ d⁻¹ approximately). Online
200 sensors of pH (Hach Lange Spain S.L.), DO (Neurtek, Spain) and temperature (Campbell
201 Scientific Inc., USA) were installed in one of the two open tanks of the PBR.

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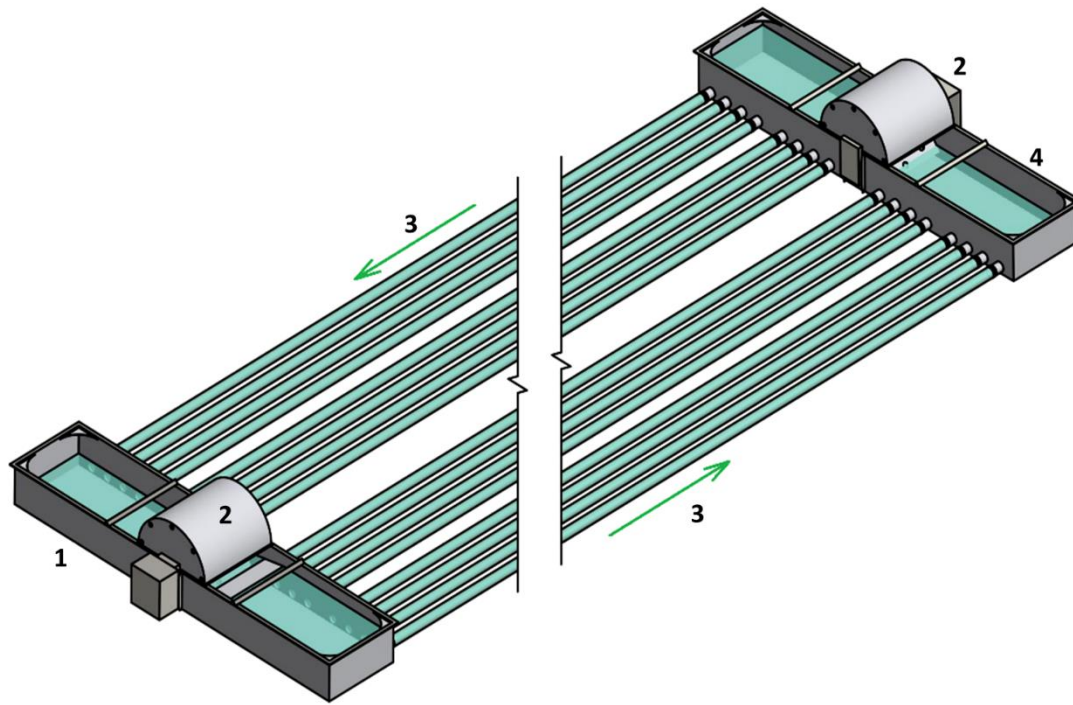


Figure 1. Scheme of the semi-closed tubular closed photobioreactor used in this study. 1:inflow from the homogenization tank; 2: paddle wheel; 3: direction of the flow within the tubes; 4: outflow to the storage tanks. Samples were taken in 1 and 4.

2.3. Sampling strategy

The three PBRs were fed daily with water from an open channel near the facilities, which carried both reclaimed wastewater from an urban WWTP nearby and agricultural run-off from the surrounding agricultural land (from now on, irrigation water). The WWTP serves 375,000 PE and has been designed to treat 64,000 m³ d⁻¹. Wastewater treatment consisted of a primary physicochemical treatment, followed by MBR and disinfection by means of UV and chlorination. Biosolids are not applied in the crop fields of this area. The water collected from the channel was mixed with domestic wastewater from a septic tank (7:1, v:v), in order to provide more nutrients for biomass growth. This feed water was mixed in a homogenization tank with constant stirring, right before the feeding operation (it was filled up anew every day). Sampling was carried out during two consecutive weeks during summer (July), three days per week and always at the same time, 10 am. Feed water of the PBR (PBR influent) and effluent mixed liquor were taken in one of the PBRs (n=12 samples). For physicochemical characterization of the samples, they were taken in PVC bottles and directly analyzed in the laboratory. For CECs analyses, samples were

223 collected in amber glass bottles and immediately filtered through 0.45 μm PVDF membrane filters
224 (Millipore, USA) and frozen upon arrival to the laboratory (amber glass bottles).

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226 **2.4. Analytical methodologies**

227 **2.4.1. Samples characterization**

228 Both influent and effluent samples were analyzed on the following wastewater quality
229 parameters: DO and temperature (EcoScan DO 6, ThermoFisher Scientific, USA) and pH (Crison
230 506, Spain) which were also measured on-site; turbidity (Hanna HI 93703, USA); total suspended
231 solids (TSS), volatile suspended solids (VSS), alkalinity, chemical oxygen demand (COD)
232 following Standard Methods (APHA-AWWA-WEF, 2012); $\text{NH}_4^+\text{-N}$ according to Solórzano
233 method (Solórzano, 1969). The ions $\text{NO}_2^-\text{-N}$, $\text{NO}_3^-\text{-N}$ and $\text{PO}_4^{3-}\text{-P}$ were measured by ion
234 chromatography (ICS-1000, Dionex Corporation, USA). Total carbon (TC), total phosphorus
235 (TP) and total nitrogen (TN) were measured by a TOC analyzer (multi N/C 2100S, Analytik Jena,
236 Germany). All the analyses were done in triplicate and results are given as average values. Mixed
237 liquor samples were examined under an optic microscope (Motic, China) for qualitative
238 evaluation of microalgae populations, employing taxonomic books and databases for their
239 identification.

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241 **2.4.2. CECs analysis**

242 Thirteen target compounds were selected based on their occurrence in WWTP effluents
243 and surface water bodies (Couto et al., 2019; Loos et al., 2013; Margenat et al., 2017; Serra-Roig
244 et al., 2016). These included 6 pharmaceuticals (diazepam (DZP), carbamazepine (CBZ), DCF,
245 IBU, lorazepam (LZP) and OXA), 2 organophosphate flame retardants (tributyl phosphate
246 (TBPh) and tri-(2-chloroethyl) phosphate (TCEP)), 3 fragrances (galaxolide (HHCB), tonalide
247 (AHTN) and methyl dihydrojasmonate (MDHJ), 1 insect repellent (N,N-diethyl-toluamide
248 (DEET)) and 1 surfactant (2,4,7,9-tetramethyl-5-decyne-4,7-diol, also known as Surfynol-104
249 (TMDD)). Further information on their physico-chemical characteristic are given in Table S1 of
250 Supplementary Material. Analytical standards for all the compounds were purchased from

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251 Sigma–Aldrich (Steinheim, Germany), including the deuterated compounds atrazine-d₅,
252 mecoprop-d₃, tonalide- d₃ and dihydroCBZ. Trimethylsulfonium hydroxide (TMSH) was
253 obtained from Fluka (Buchs, Switzerland). Strata-X polymeric cartridges (200 mg) were
254 purchased from Phenomenex (Torrance, CA, USA). The 1-2 µm glass fiber filters (Ø 47 mm) and
255 0.45 µm PVDF membrane filters were obtained from Whatman (Maidstone, UK) and (Millipore,
256 USA), respectively.

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258 *GC-MS-MS analysis*

259 For the determination of the different target analytes, samples were analyzed by gas
260 chromatography coupled to mass spectrometry (GC-MS/MS), adapting the methodology by
261 Matamoros and Bayona (2006). For both influent and effluent water samples, 100 mL were
262 preconcentrated using a previously activated polymeric solid-phase extraction cartridge (200 mg
263 Strata X, Phenomenex, US). Further information on pretreatment and GC-MS/MS methodology
264 validation and application is given elsewhere (Margenat et al., 2017; Matamoros and Bayona,
265 2006).

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267 **2.5. Environmental risk assessment**

268 In order to evaluate the potential ecotoxicological risk of those CECs still present in the PBR
269 effluent, hazard quotients (HQ) have been estimated as indicated in equation 1,

270 [1]:

$$271 \quad HQ = \frac{MEC}{PNEC}$$

272 where MEC is the measured environmental concentration, and PNEC is the predicted-no effect
273 concentration. When PNEC data are not available, alternative PNECs can be derived by dividing
274 the toxicity endpoint values found in the literature (EC₅₀ or LC₅₀) by an uncertainty factor of up
275 to 1000 (Sanderson et al., 2004). HQ values < 0.1 mean that no adverse effects are expected.
276 When 0.1<HQ<1, the risk is low but it should not be neglected; when 1<HQ<10, a moderate risk
277 is implied, and HQ > 10 means a relevant ecological hazard (EMEA, 2006).

278 Eventually, for the purpose of evaluating the overall ecotoxicity risk of the PBR effluent,
279 cumulative HQs were calculated for each trophic level considered, adding all HQs calculated for
280 each individual CEC detected in the effluent.

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282 **3. RESULTS**

283 **3.1. Water quality parameters**

284 On-line measurements of temperature, DO and pH are given in the Supplementary
285 Material (Figure S1). The photosynthetic activity of microalgae caused daily variations of DO
286 and pH, characteristic of these systems, with DO ranging from 8 to 14 mg L⁻¹ and pH from 8 to
287 10.5. The mixed liquor temperature increased during daylight, reaching values up to 41°C, due to
288 the high solar radiation and ambient temperature. At night, the mixed liquor was cooled,
289 decreasing to approximately 24°C.

290 Data on the performance of the PBR were already published elsewhere (Vassalle et al., 2020b),
291 and are included in the Supplementary Material (Table S2). Briefly, the average biomass
292 productivity in the PBR was low (6.9 ± 0.7 g VSS m⁻² d⁻¹) due probably to the low concentration
293 of total inorganic nitrogen (TIN), N-NH₄⁺ and phosphate (P-PO₄³⁻) in the PBR feed water.
294 Average VSS concentration in the PBR effluent (mixed liquor) was 215 mg L⁻¹, corresponding to
295 a 74% of the TSS, which is in accordance with the values generally observed in microalgae-based
296 systems (García-Galán et al., 2018; Gutiérrez et al., 2016). The registered pH values > 8 promoted
297 precipitation of inorganic salts of different nature, leading to an increase of the VSS/TSS ratio.
298 COD concentration increased a 16% during PBR treatment, which is generally linked to the
299 release of a fraction of the carbon fixed during photosynthesis as dissolved organic carbon (DOC)
300 by microalgae (Arbib et al., 2013; García-Galán et al., 2018; García et al., 2006).

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302 **3.2. Occurrence of contaminants of emerging concern in the irrigation water**

303 *3.2.1. Pharmaceuticals*

304 The six targeted pharmaceuticals, 4 psychiatric drugs and 2 non-steroidal anti-
305 inflammatory drugs (NSAIDs), were detected in the PBR feed water all the sampling days (Figure

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2A). CBZ was the most abundant psychiatric drug (660-830 ng L⁻¹), followed by LZP, OXA and DZP. The concentration of CBZ was in agreement with that found in previous studies on the same site (García-Galán et al., 2018) and in the Baix Llobregat area (Margenat et al., 2017). The ubiquity of this anticonvulsant in the aquatic environment has been frequently demonstrated, being currently considered as one of the most reliable anthropogenic pollution tracers given its resilience to biodegradation during conventional wastewater treatment (WWT), and also to photodegradation (Guo and Krasner, 2009; Hai et al., 2018). Its presence in agricultural run-off waters has also been reported by Pedersen et al., (2005), who detected CBZ in agricultural run-off from crop fields irrigated with effluent wastewater in California, at levels between 320 and 440 ng L⁻¹. Lower concentrations were reported in rural run-off in Mexico (1-35 ng L⁻¹) (Moeder et al., 2017) and also by Tran et al. (2019) in both urban and agricultural run-off. LZP was present at average concentration of 511 ng L⁻¹, slightly higher than levels previously detected in irrigation water in the same area by Margenat et al. (2017) and in the Llobregat river by Proia et al. (2013), probably due to the mixing of the irrigation water with the septic tank wastewater. DZP was detected at much lower levels (5.3-8.1 ng L⁻¹), similar to those reported by Proia et al. (2013). OXA was detected at concentrations between 216 ng L⁻¹ and 371 ng L⁻¹. This psychiatric drug is also the final degradation product of DZP and LZP aforementioned, which are amongst the most highly consumed anti-depressants worldwide (Kosjek et al., 2012). It has been detected at similar levels other irrigation channels near our study site, also fed with reclaimed wastewater (178 ng L⁻¹), but also in irrigation channels fed with surface water (25-36 ng L⁻¹) and groundwater (<2 ng L⁻¹) (Margenat et al., 2017). It was also present in surface water influenced by agricultural run-off in the UK (White et al., 2019), and frequently detected in WWTP effluents all over Europe (81 out of the 90 effluent samples analyzed in 18 countries) at average concentration of 162 ng L⁻¹ (Loos et al., 2013). Regarding the NSAIDs evaluated, DCF was present at levels in the range 860-1,306.8 ng L⁻¹, higher than concentrations reported in a previous campaign on the same site (García-Galán et al., 2018) (similarly to LZP, it is probably due to the mix with the water from the septic tank, which could have had residual DCF). and also higher than those found in rural run-off in Mexico or Singapore (Moeder et al., 2017; Tran et al., 2019). IBU was also detected at

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334 concentrations in the range 321-512 ng L⁻¹, data which is in agreement with that detected in the
335 aforementioned work by Moeder et al. (2017). Similar levels were detected by White et al. (2019)
336 in surface waters receiving rural run-off in the UK.

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338 3.2.2. *Personal care products*

339 Two of the three fragrances investigated, HHCB and AHTN, were detected at average
340 concentrations of 191 ng L⁻¹ and 127 ng L⁻¹, similar levels to those detected in a previous sampling
341 campaign in the same location (García-Galán et al., 2018). Their presence in surface waters is
342 frequent and usually attributed to wastewater effluent discharges and not to agricultural run-off
343 (Blum et al., 2018; Celeiro et al., 2019; Corada-Fernández et al., 2017; Gómez et al., 2012). The
344 insect repellent DEET was present at concentrations ranging from 502-698 ng L⁻¹, higher than
345 those found by Margenat et al. (2017) in other irrigation channels nearby. This high levels were
346 due to the marked seasonal variability associated to this compound, as its usage is much higher
347 during summer and mosquitoes proliferation (Merel et al., 2015). Despite it is clearly a compound
348 of domestic application and, therefore, from wastewater origin (Gago-Ferrero et al., 2017; Launay
349 et al., 2016), its environmental ubiquity has been demonstrated in several studies, including
350 stormwater run-off, surface waters and groundwaters (Brausch and Rand, 2011; Burant et al.,
351 2018; Rehr et al., 2020), and also in agricultural run-off and at similar levels than those observed
352 in this study (Tran et al., 2019).

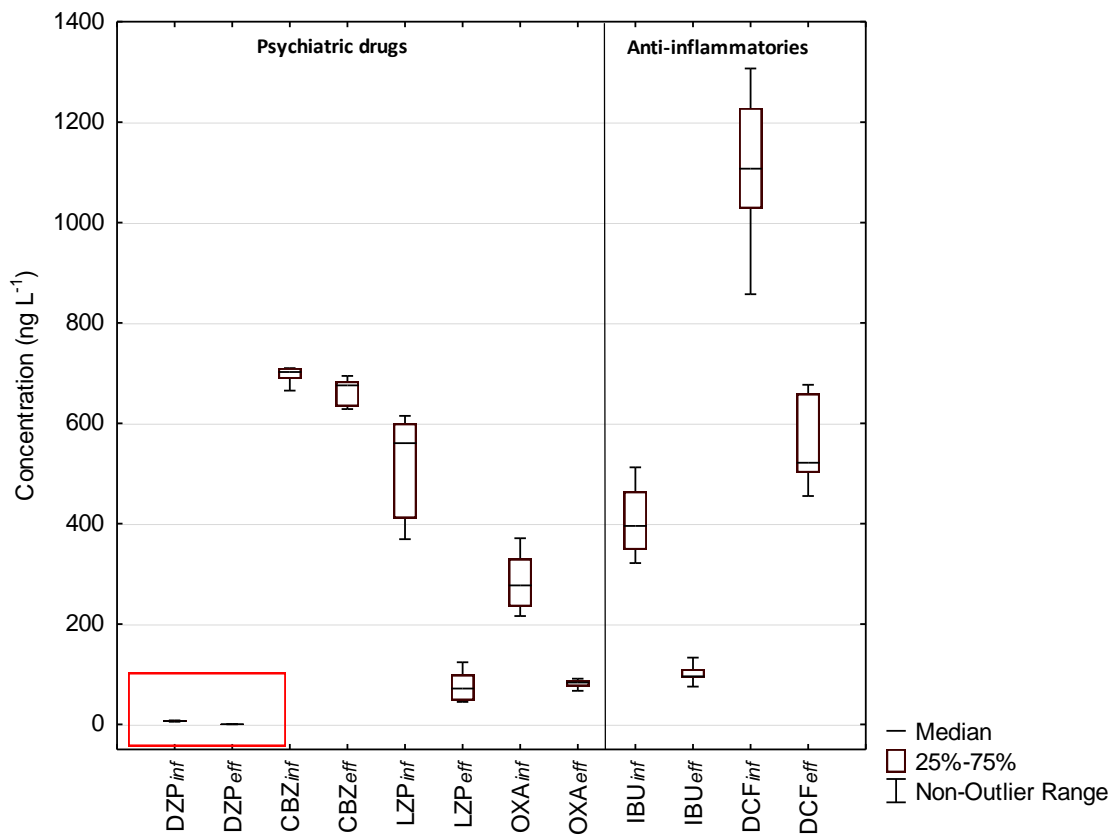
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354 3.2.3. *Organophosphate flame retardants and surfactants*

355 The organophosphate flame retardants TBP and TCEP were present at average values of
356 34.8 ng L⁻¹ and 284 ng L⁻¹ respectively (Figure 2B), levels slightly higher than those detected in
357 other irrigation channels (Margenat et al., 2017). These compounds have been detected in
358 basically all the environmental compartments due to their broad range of applications (pesticides
359 solvents, detergents antifoaming, additives, etc.) and their extensive use in industrial activities, as
360 well as the progressive disuse of polybrominated flame retardants (Yang et al., 2017). Different
361 authors have also found both TBP and TCEP in stormwater run-off at similar or higher

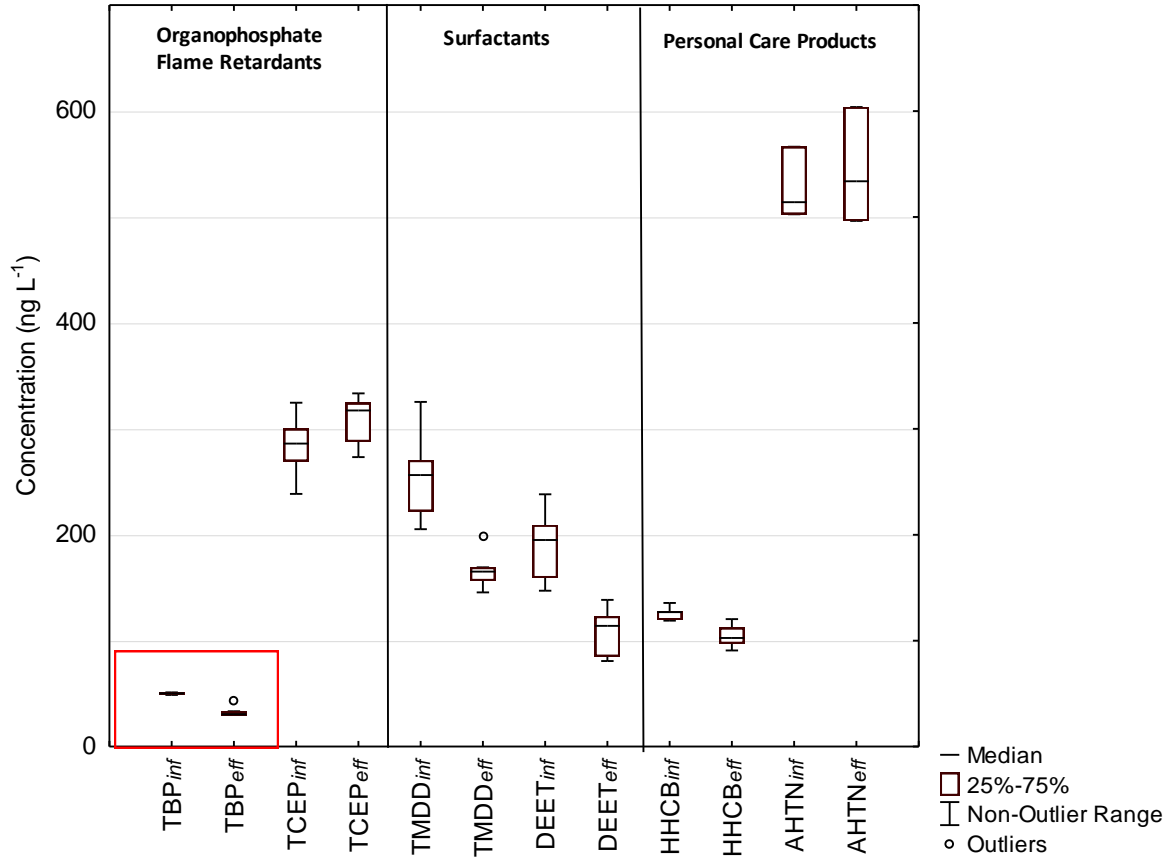
362 concentration ranges, and also in precipitation water (rain and snow) (Burant et al., 2018; Regnery
 363 and Püttmann, 2010). These authors stated that, when used as additives, these compounds do not
 364 bind to the matrix and so they can be released to the environment by volatilization and dissolution.
 365 Precipitation wash-off and dry deposition, together with WWTPs effluents discharges, are their
 366 main entry pathways. The surfactant and anti-foaming TMDD, known with the commercial name
 367 of Surfynol 104®, was present at concentrations ranging from 205 ng L⁻¹ to 325 ng L⁻¹. TMDD is
 368 used in the industry to reduce the surface tension of coatings, adhesives, paints and printing inks,
 369 but it is also used in pesticide formulations and in toilet and kitchen paper in the domestic context
 370 (Guedez and Püttmann, 2014). It has been detected in surface waters impacted by WWTPs
 371 effluents, in concentrations ranging from 16 to 240 ng L⁻¹ (Blum et al., 2018), and up to the µg L⁻¹
 372 level in rivers impacted by industrial activities (Guedez and Püttmann, 2014).

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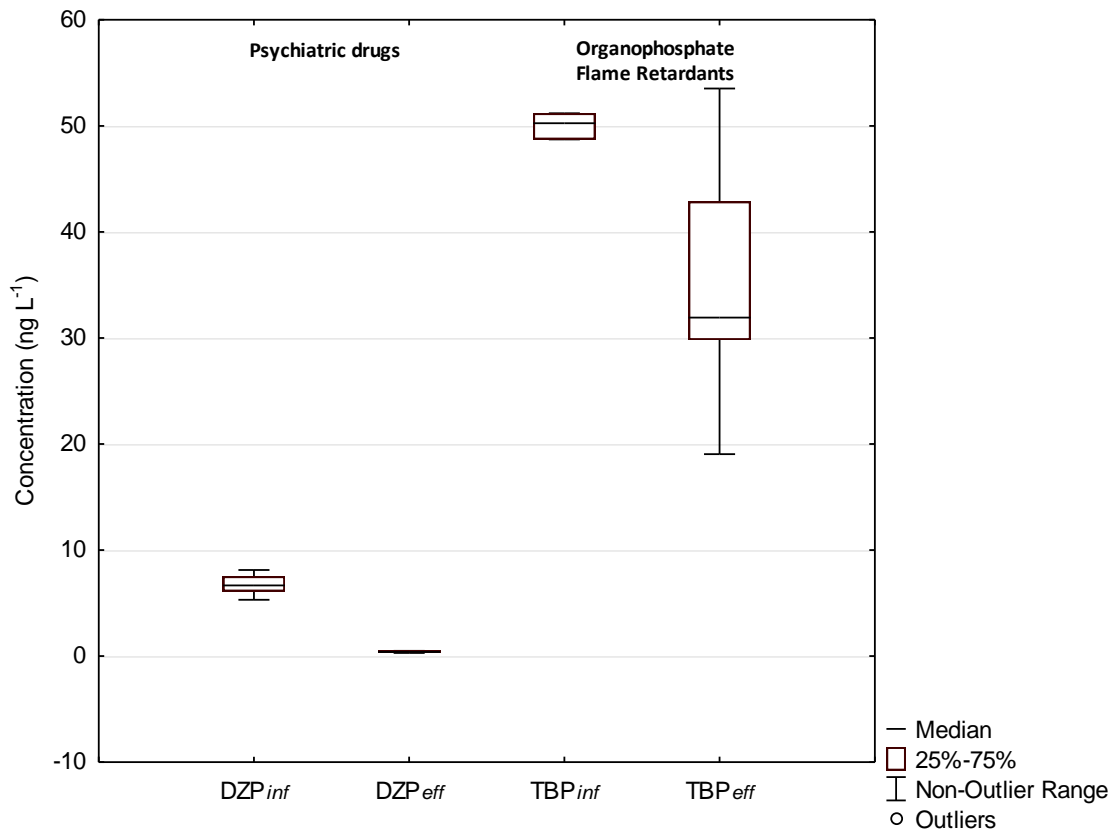
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395 **Figure 2.** Box plots of the concentrations of pharmaceuticals and fragrances (A) and other contaminants of
 396 emerging concern (B) in rural run-off (influent) and effluent samples of the PBR. Marked compounds have
 397 been zoomed in graph C'.
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400 3.3. Removal of CECs during PBR treatment

401 3.3.1. Pharmaceuticals

402 The pharmaceuticals entering the PBR have been classified according to their removal
 403 efficiency (RE%): efficiently removed (>70%) namely IBU, DZP, LZP and OXA; moderately
 404 removed (35-50%), namely DCF, and poorly removed (<25%), namely CBZ (Table 1).

405 The good removal of DZP (94%±5) is significant, as it is usually inefficiently removed during
 406 conventional WWTs. Indeed, many studies have reported RE% ranging from negative
 407 eliminations to barely a 18% (García-Galán et al., 2016; Mamo et al., 2018; Rodriguez-Mozaz et
 408 al., 2015), although also better removals have been observed (30-60%) (Gros et al., 2012; Mira
 409 et al., 2019). West and Rowland (2012) studied direct and indirect photodegradation of DZP (also

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410 OXA) and concluded that the presence of humic substances increased its photodegradation rate;
411 in the case of our PBR, both the humic acids present in the open channel and the carbon exudates
412 from the microalgae within the reactor could have enhanced the photodegradation of this drug.
413 On the contrary, the aforementioned authors also observed that humic substances seemed to slow
414 down the photodegradation of OXA. This drug is highly resilient to both aerobic and anaerobic
415 biodegradation and also to photodegradation (Kosjek et al., 2012; Calisto et al., 2011; Loos et al.,
416 2013), and some authors have indicated that it is likely to persist in water for decades (Klaminder
417 et al. (2015). Considering its high log K_{ow} (3.3), adsorption onto the microalgae biomass seems to
418 be the main removal pathway within the PBR, despite its recalcitrance during conventional
419 WWTs. In a previous study by Gojkovic et al. (2019), removals in the range 2-27% were obtained
420 in a laboratory-scale flat panel PBR, using different microalgae species. In that study, OXA was
421 indeed detected in the biomass (37% maximum). To the author's knowledge, there are no previous
422 studies on the elimination of OXA in full scale microalgae systems. Last of all, it should be
423 regarded that both DZP and OXA (and other benzodiazepines) can persist in soils long enough
424 after irrigation to be uptaken by different crops, as demonstrated by Carter et al., (2018) with
425 radish and silverbeet. The excellent removals obtained in the present study highlight the feasibility
426 of microalgae-based treatment to remove these drugs before water reclamation.

427 Regarding LZP (83%±5 removal), given its low solubility and high log K_{ow} , it seems that
428 microalgae uptake is the most likely removal pathway, although photodegradation cannot be
429 neglected either (Calisto et al., 2011). Lower removals (30-57%) were obtained by Hom-Diaz et
430 al. (2017) in a smallerscale closed PBR operating as secondary treatment, with a similar TSS
431 concentration than the PBR in this work, but with higher HRT (8-12 h). LZP is also incompletely
432 removed during conventional WWTs (<50%) (Dolar et al., 2012; Mira et al., 2019).

433 For IBU (RE% of 70%±12), the results obtained in the present study agree with those obtained in
434 removals in HRAPs operating as secondary treatments (García-Galán et al., 2020a; Villar-
435 Navarro et al., 2018). These authors attributed its removal mostly to aerobic biodegradation, as
436 adsorption onto biomass was very low. Indeed, despite its high log K_{ow} (3.97), IBU is charged
437 negatively at the pH of the PBR ($pK_a = 5.3$), being repelled by the negative charge of the

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438 microalgae cell walls (Matamoros et al., 2016). Ding et al., (2017) obtained lower removals for
439 IBU (20%-60%) in laboratory batch experiments with the fresh-water diatom *Navicula* sp. The
440 higher initial concentrations (1-50 mg L⁻¹) could be responsible of these lower eliminations due
441 to toxicity events against the diatom. In a different study, IBU removal in the presence of
442 microalgae was attributed to indirect photodegradation rather than to sorption, due to the presence
443 of dissolved organic matter acting as a photocatalyst of the reaction (de Wilt et al., 2016).
444 CBZ was poorly removed (11% ± 8). Different studies have also reported low removals in HRAPs
445 operating as secondary treatments, ranging from no removal (García-Galán et al., 2020a) to
446 eliminations in the range of 9-23% with HRT of 6 d (Villar-Navarro et al., 2018). Matamoros et
447 al. (2015) obtained removals of 46% (4 d of HRT) and 62% (8 d HRT) also during the warm
448 season, highlighting that even under the best conditions for microalgae-based treatment efficiency
449 (summer campaigns), CBZ is highly stable towards photodegradation and aerobic biodegradation.
450 Díaz-Garduño et al. (2017) obtained similar results in laboratory scale experiments (RE% in the
451 range 0-23%); removals in the range of 10-30% have been obtained with different species of green
452 algae (*Chlorella* sp., *Scenedesmus* sp., *Coelastrum astroideum* and *Chlamydomonas mexicana*
453 (de Wilt et al., 2016; Gojkovic et al., 2019; Matamoros et al., 2016; Xiong et al., 2016). These
454 authors reached the conclusion that bioadsorption and/or bioaccumulation were negligible, being
455 biodegradation the main elimination mechanism. On the contrary, García-Galán et al. (2020a)
456 observed concentrations of CBZ in the biomass equivalent to the 39% of the initial concentration
457 in the influent, but yet it was not eliminated in the system, but still present in the effluent and at
458 higher than those in the influent. These results indicated a clear bioaccumulation in the biomass
459 of this drug. Some authors also point out that glucuronide moieties of CBZ have never been
460 included in monitoring studies (due to the lack of commercial standards), and demonstrated the
461 presence and cleavage of these metabolites during conventional wastewater treatments (Vieno et
462 al., 2007). Bahlmann et al. (2014) even suggested a concentration increase of CBZ of nearly
463 100% during wastewater treatment due to this cleavage.

464 DCF was removed by a 52% ± 6 on average. These results agree with those obtained in
465 previous studies in HRAPs acting as secondary treatment, with removals of 55% (Vassalle et al.,

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466 2020a), 39-74% (Villar-Navarro et al., 2018) and 51-55% (García-Galán et al., 2020a). The latter
467 pointed out that bioadsorption/bioaccumulation played a relevant role in its removal from the
468 aqueous phase ($\log K_{ow}=4.5$), given the high concentrations detected in the biomass (267.9 ng g^{-1}),
469 whereas biodegradation was low. On the other hand, de Wilt et al. (2016) attributed the removal
470 of DCF in different types of wastewater (40-60%) to phototransformation, as they observed its
471 elimination in laboratory batches without microalgae inoculum. Photodegradation of DCF in
472 surface waters has been previously reported (Kunkel and Radke, 2012; Zhang et al., 2008). In
473 HRAPs, Matamoros et al. (2015) observed that the removal of this drug was considerably higher
474 during the warm/summer season (82-92%) compared to the cold season (21-29%). Other factors
475 such as the transparency of the plastic material of the tubes in the PBR (Harris et al., 2013) may
476 also reduce the light penetration and the photodegradation rates of photosensitive compounds,
477 compared to those observed in open systems.

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479 3.3.2. *Personal Care Products*

480 The fragrances HHCB and AHTN were only partially removed, with average RE% of
481 45% and 20%, respectively. These results are lower than those obtained in a previous campaign
482 in the same location (García-Galán et al., 2018), and also to those obtained in open systems
483 operating as secondary treatments ($51\% \pm 12$ for HHCB and $46\% \pm 7$ for AHTN) (Matamoros et
484 al., 2015). Laboratory scale assays also yielded higher removals (near 100%) after 7-10 d (Díaz-
485 Garduño et al., 2017; Matamoros et al., 2016). Both compounds have high $\log K_{ow}$ (>5) and \log
486 K_{oc} (>3.7) and a very low biodegradability, being biomass adsorption the most probable removal
487 pathway. Regarding DEET, an average negative removal was obtained for DEET. Díaz-Garduño
488 et al. (2017) obtained removal efficiencies for DEET ranging from negative values ($n=2$) to 55%
489 ($n=1$) in laboratory scale batch reactors.

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491 3.3.3. *Organophosphate Flame Retardants and Surfactants*

492 TBP was removed by a $43\% \pm 7$, whereas TCEP concentrations in the effluent of the PBR were
493 similar than those of the influent. This is probably due to the plastic components of the PBR

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494 system, which may release TCEP to the aqueous phase during treatment, as already suggested by
495 (Rodil et al., 2012, 2009) in conventional WWTPs facilities. In a previous sampling campaign in
496 the same location, TCEP was removed only in a 20% (García-Galán et al., 2018). In open systems
497 acting as secondary treatments, Matamoros et al. (2015) obtained RE% in the range 15-39% for
498 TCEP and 24-82% for TBP under HRT of 4 d, reaching better results with longer HRTs (8 d).
499 High Henry constant for TBP (K_H of $0.3 \text{ atm m}^3 \text{ mol}^{-1}$) may be indicative of volatilization events
500 and its partial removal in HRAPs, and also in the open tanks of the semi-closed PBR. Indeed,
501 aerated batch reactors at laboratory scale confirmed the recalcitrance of TCEP, which was
502 removed $< 20\%$ after 10 d, whereas that TBP was more efficiently removed (Matamoros et al.,
503 2016). TCEP is a highly hydrophilic compound ($\log k_{ow}=1.44$) so it is not likely to be adsorbed
504 onto the biomass either, contrary to TBP ($\log k_{ow}= 4$). Furthermore, TCEP is a highly stable
505 molecule, not prone to biodegradation, which, together with its high solubility, makes it a highly
506 mobile and persistent pollutant once discharged into environmental waters (Blum et al., 2018;
507 Reemtsma et al., 2008; Rodil et al., 2012). Last of all, the surfactant TMDD was removed by a
508 33% in the PBR, and considering its low solubility and high $\log K_{ow}$, adsorption onto biomass
509 seems a feasible removal pathway within the system. Conventional WWT is generally quite
510 inefficient in removing this surfactant, with barely no elimination (Blum et al., 2018, 2017;
511 Guedez and Püttmann, 2014).

521 **Table 1.** Maximum, median and average concentrations (\pm SDV) detected for the different CECs evaluated,
 522 and removal efficiency (R%) after PBR treatment.
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FAMILY	NAME	Maximum (ng L ⁻¹)	Median (ng L ⁻¹)	Average (ng L ⁻¹)	Removal (R%)
PHARMACEUTICALS	Carbamazepine (CBZ)	833	702	717 \pm 59	11 \pm 8
	Diclofenac (DCF)	1307	1107	1106 \pm 111	52 \pm 6
	Ibuprofen (IBU)	512	395	406 \pm 138	70 \pm 12
	Lorazepam (LZP)	615	560	511 \pm 113	83 \pm 6
	Oxazepam (OXA)	371	277	284 \pm 60	71 \pm 7
	Diazepam (DZP)	8	7	7 \pm 1	94 \pm 5
PERSONAL CARE PRODUCTS	Galaxolide (HHCB)	238	195	191 \pm 27	45 \pm 14
	Tonalide (AHTN)	136	127	126 \pm 6	20 \pm 5
	N,N-diethyl-m-toluamide (DEET)	1328	574	699 \pm 90	- 4 \pm 12
ORGANOPHOSPHATE FLAME RETARDANTS	Tributyl phosphate (TBP)	81	50	54 \pm 4	43 \pm 7
	Tris(2-chloroethyl) phosphate (TCEP)	325	286	284 \pm 29	- 4 \pm 5
SURFACTANTS	Surfinol 104 (TMDD)	325	256	256 \pm 42	33 \pm 7

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527 3.3. Environmental risk assessment

528 As indicated in section 2.5, hazard quotients (HQs) were calculated for those CECs not
 529 fully removed during PBR treatment, following equation [1]. To estimate the PNEC, toxicity data
 530 for different standard test species covering different trophic levels were obtained from the
 531 ECOTOX database of the Environmental Protection Agency (EPA). Chronic exposure indicators
 532 (NOEC) would be preferable in the case of CECs, as non-target species are exposed to low
 533 concentrations of these contaminants during long periods of time, so unexpected long-term effects
 534 could eventually appear. However, as chronic toxicity data are frequently unavailable, PNECs
 535 were calculated using EC₅₀ and LC₅₀ as indicators of acute toxicity (regarding immobilization and
 536 mortality, respectively). These values were divided by an uncertainty factor (1000) to become
 537 more representative values of the real situation under environmental conditions (longer periods
 538 of exposure) (Sanderson et al., 2004; Valcárcel et al., 2011). HQs were estimated for green algae,
 539 invertebrates, crustaceans and fish (standard test species, see Table 2). In order to establish a worst
 540 case scenario, when different toxicity endpoints were available for a given compound, the lowest

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541 toxicity value was used (Table S3 in Supplementary Material). Given their homogeneity, the
542 average measured concentrations in the PBR effluent for each CEC were used. HQ values are
543 shown in Table 2. Calculations were subjected to the availability of the toxicity data; in
544 consequence, risk evaluation for LZP and OXA could not be done. However, both drugs have a
545 similar low solubility and high log K_{ow} - K_{oc} values that indicate a high tendency to adsorb onto
546 biomass and bioaccumulate, as demonstrated in previous studies (García-Galan et al., 2017;
547 Lagesson et al., 2016). Amongst the different CECs still present in the PBR effluent, HQ values
548 between 1 and 10 were obtained only for AHTN (fish) and CBZ (invertebrates), implying a
549 moderate hazard in the receiving water body. HQ values between 0.1 and 1 (low risk) were
550 obtained for the fragrances AHTN and HHCB in most cases, for IBU against fish and for DEET
551 against crustaceans. Nevertheless, the majority of the compounds yielded HQs < 0.1, meaning
552 that no environmental risk would be derived from their discharge on the PBR effluent. Given the
553 results obtained, and considering the cumulative HQs in the effluent, the sensitivity of the
554 different trophic levels would be as follows: invertebrates > fish > crustaceans > green algae
555 (Figure 3). Despite the overall good removal efficiency of the PBR for the different CECs studied,
556 the decrease of the derived ecotoxicity risk was only moderate (Figure 3), with a 38% reduction
557 for fish, 15% for invertebrates, 16% for crustaceans and only a 3% for green algae. The low
558 removals of CBZ or AHTN would lead to a higher impact against different species, which are
559 mostly unaffected by the presence of the other CECs. Indeed, different authors have reported a
560 moderate to high environmental risk derived from the CBZ presence in European surface waters
561 (Palma et al., 2020; Zhou et al., 2019), and Díaz-Garduño et al. (2017) obtained HQ>1 for AHTN
562 and green algae after microalgae treatment. In the prioritization study for pharmaceuticals
563 performed by Zhou et al., (2019) in different European countries, DCF, IBU and CBZ posed the
564 highest risk to aquatic ecosystems. However, the levels obtained after microalgae treatment in the
565 present study yielded HQ<0.1 for all of them (except for CBZ against invertebrates). It should be
566 considered that PBR effluent concentrations will be subjected to further dilution once discharged
567 in the receiving water bodies. Therefore, the estimated risk derived from exposure would be
568 considerably lower. On the contrary, the number of CECs potentially present in the water

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569 analyzed is much higher than the 15 compounds considered in the present study. Furthermore, it
570 should be taken into account that conventional risk assessment of CECs is usually based on this
571 concentration addition for estimating the mixture toxicity (European Commission, 2009), ignoring
572 toxicity derived from synergies, additive effects or antagonistic effects (Baek et al., 2019).
573 Likewise, addition of HQs of pharmaceuticals with similar modes of action (i.e. psychiatric drugs
574 or anti-inflammatories) could result in the overestimation of adverse effects. Mixture toxicity is
575 out of the scope of the present study, but it is actually a hot topic within the scientific community,
576 which is currently devoting a huge effort to discern and evaluate more realistic toxicity scenarios.

Table 2. Average PBR effluent concentrations (ng L⁻¹) (used as measured environmental concentrations, MEC), ecotoxicity endpoints used for the different trophic levels considered (mg L⁻¹) and hazard quotients (HQ) estimated.

	Average MEC (ng L ⁻¹)	TOXICITY ENDPOINTS (mg L ⁻¹)				HQ (effluent)				
		Green algae	Invertebrate	Crustaceans (<i>Daphnia magna</i>)	Fish	Green algae	Invertebrate	Crustaceans	Fish	
PHARMACEUTICALS	Diazepam (DZP)	0.38 ± 0.07	-	47.3 ^{5*}	4.3 ⁶	0.3 ^{9*}	-	8.2E-06	9.2E-05	1.4E-03
	Carbamazepine (CBZ)	665.1 ± 27.5	74 ²	0.4 ⁵	13.8 ⁶	54.2 ⁹	8.9E-03	1.8	4.8E-02	1.2E-02
	Ibuprofen (IBU)	101.4 ± 68.6	315 ²	-	>45 ⁶	0.7 ^{7*}	3.2E-04	-	2.2E-03	1.5E-01
	Diclofenac (DCF)	555.9 ± 90.4	72 ²	-	28.1 ⁶	71 ^{10*}	7.7E-03	-	1.9E-02	7.8E-03
PERSONAL CARE PRODUCTS	Galaxolide (HHCB)	109.2 ± 22.8	0.7 ¹	0.3 ^{4*}	2.7 ⁶	3.6 ^{9*}	1.5E-01	3.8E-01	4.1E-02	7.8E-01
	Tonalide (AHTN)	101.1 ± 8.5	0.5 ¹	0.5 ^{4*}	0.2 ⁶	0.1 ^{7*}	2.2E-01	2.2E-01	4.1E-01	1.01
	N,N-diethyl-toluamide (DEET)	544.6 ± 54.6	-	-	1 ⁶	71.2 ^{8*}	-	7.8E-02	5.4E-01	7.6E-03
ORGANOPHOSPHATE FLAME RETARDANTS	Tributyl phosphate (TBP)	28.2 ± 6.3	1.1 ²	12.5 ⁵	35 ⁶	1.3 ^{7*}	2.6E-02	2.2E-03	8.1E-04	2.8E-02
	Tris-(2-chloroethyl) phosphate (TCEP)	308.9 ± 23.5	51 ²	-	330 ⁶	3.7 ^{9*}	6.1E-03	-	9.4E-04	8.2E-02
SURFACTANTS	2,4,7,9-Tetramethyl-5-decyne- 4,7-diol (TMDD)	167.2 ± 20	-	-	91 ⁶	36	-	-	2.2E-03	5.5E-03

1- *Pseudokirchneriella subcapitata*; 2-*Desmodesmus subspicatus*; 3- *Tetrahymena pyriformis*; 4-*Chironomus riparius*; 5- *Brachionus calyciflorus*; 6- *Daphnia magna* 7- *Pimephales promelas*.

8- *Oncorhynchus mykiss*. 9- *Danio rerio*. 10- *Cyprinus carpio*

*: LC₅₀ values (the other toxicity endpoints are EC₅₀ values)

Values for TMDD are taken from Guedez and Puttman, 2014.

Green algae and crustaceans endpoint values for TCEP are taken from Cristale et al. 2013.

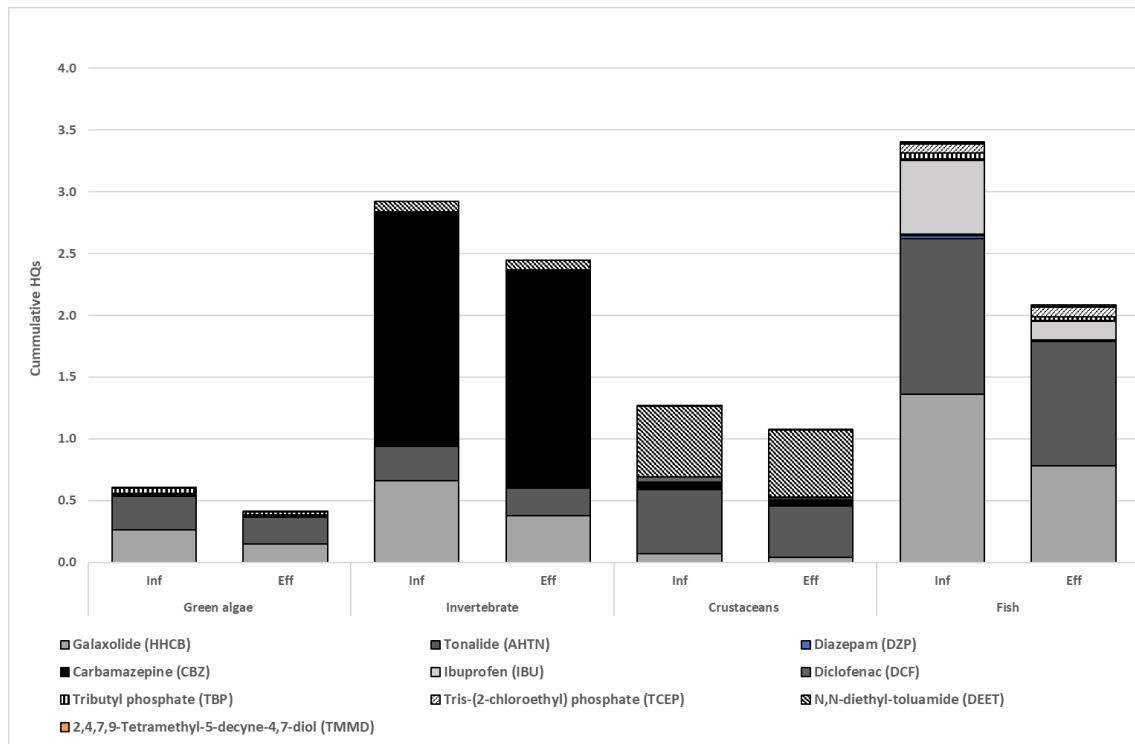


Figure 3. Cumulative HQs for each of the trophic levels considered in the feed water of the PBR (first column) and effluent (second column). For all the compounds, HQs were available for at least 3 of the 4 trophic levels considered, with the exception of TMDD (only crustaceans and fish).

4. CONCLUSIONS

The capacity of a semi-closed, tubular horizontal PBR to remove 13 contaminants of emerging concern (CECs) detected in water from an agricultural irrigation channel was evaluated. Removal efficiencies ranged from efficiently removed (>70%) namely IBU, DZP, LZP and OXA; moderately removed (35-70%), namely DCF, HHCB, TBP and TMDD; and poorly removed (<35%), AHTN, CBZ, TCEP and DEET. Nevertheless, for most of the compounds their removal were comparable to those obtained in conventional WWTPs. On the other hand, very good elimination efficiencies were obtained for the benzodiazepines OXA (highly recalcitrant) and DZP, which are generally barely removed in conventional treatment systems. An environmental toxicity evaluation has been performed to fathom out the impact of the PBR effluent in the receiving water body. Despite most of the compounds have an HQ <0.1, implying no risks associated, the cumulative assessment has highlighted a low to moderate risk for the different trophic levels except for green algae. The PBR treatment reduced the environmental risk between a 3% (green algae) and 38% (fish). Overall, the good treatment efficiency of the PBR, together

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2 with the related low O&M costs and sustainability, makes this treatment approach a feasible
3 alternative to conventional treatment.
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7
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