1	Toxicity of the Non-Steroidal Anti-inflammatory Drugs (NSAIDs)
2	acetylsalicylic acid, paracetamol, diclofenac, ibuprofen and naproxen
3	towards freshwater invertebrates: a review
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16 Abstract

Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) represent one of the main therapeutic class of 17 molecules contaminating aquatic ecosystems worldwide. NSAIDs are commonly and extensively 18 used for their analgesic, antipyretic and anti-inflammatory properties to cure pain and inflammation 19 in human and veterinary therapy. After use, NSAIDs are excreted in their native form or as 20 metabolites, entering the aquatic ecosystems. A number of monitoring surveys has detected the 21 presence of different NSAIDs in freshwater ecosystems in the ng/L - µg/L concentration range. 22 Although the concentrations of NSAIDs in surface waters are low, the high biological activity of 23 these molecules may confer them a potential toxicity towards non-target aquatic organisms. The 24 present review aims at summarizing toxicity, in terms of both acute and chronic toxicity, induced by 25 the main NSAIDs detected in surface waters worldwide, namely acetylsalicylic acid (ASA), 26 27 paracetamol (PCM), diclofenac (DCF), ibuprofen (IBU) and naproxen (NPX), both singularly and in mixture, towards freshwater invertebrates. Invertebrates play a crucial role in ecosystem 28 29 functioning so that NSAIDs-induced effects may result in hazardous consequences to the whole freshwater trophic chain. Acute toxicity of NSAIDs occur only at high, unrealistic concentrations, 30 while sub-lethal effects arise also at low, environmentally relevant concentrations of all these drugs. 31 32 Thus, further studies represent a priority in order to improve the knowledge on NSAID toxicity and mechanism(s) of action in freshwater organisms and to shed light on their real ecological hazard 33 34 towards freshwater communities.

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36 Keywords: Non-Steroidal Anti-inflammatory Drugs (NSAIDs); freshwater ecosystems;
37 invertebrates; toxicity

1. Pharmaceuticals as emerging contaminants of freshwaters

In the last two decades, pharmaceutical compounds have been identified as emerging contaminants 40 for aquatic ecosystems. Pharmaceutical compounds are extensively and increasingly used both in 41 human and veterinary therapy, including agriculture and aquaculture (Boxall et al., 2015). 42 Pharmaceuticals have been designed to have a specific mode of action, targeting specific organs, 43 44 metabolic pathways or receptors to modulate physiological functions, to treat a disease and to restore the health of the organism. For these reasons, pharmaceuticals play a pivotal role in our 45 society, which commonly uses, and often abuses, a number of these molecules. For instance, in the 46 47 European Union (EU) alone, it has been estimated that about 3,000 different substances are used in human therapy, including anti-inflammatory drugs, contraceptives, antibiotics, β-blockers, lipid 48 regulators, neuroactive drugs and many others (Fent et al., 2006). After their use, pharmaceuticals 49 are excreted unchanged or as metabolites entering the sewage. As wastewater treatment plants 50 (WWTPs) own a limited removal efficiency for several drugs, they are discharged in WWTP 51 52 effluents contributing to the contamination of surface waters and, rarely, of groundwater and drinking water (Santos et al., 2010). According to the trend of production and use, as well as their 53 pharmacokinetic and chemico-physical properties, different pharmaceuticals are detected in aquatic 54 55 ecosystems in the ng/L to mg/L concentration range worldwide (Santos et al., 2010; Al Aukidy et al., 2014; Bagnis et al., 2018; Fekadu et al., 2019). In fact, a recent comprehensive review of 56 measured environmental concentrations (MECs) for both human and veterinary pharmaceuticals on 57 a global scale showed that 631 different substances were found water samples worldwide (aus der 58 Beek et al., 2016). Thus, the presence of pharmaceuticals in aquatic ecosystems represents one of 59 the main concerns that ecotoxicology has to face (Fent et al., 2006; Santos et al., 2010; Boxall et al., 60 2015). 61

63 2. Non-Steroidal Anti-inflammatory Drugs (NSAIDs) in freshwater ecosystems

Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) represent one of the most relevant therapeutic 64 class found in aquatic ecosystems worldwide (aus der Beek et al., 2016). NSAIDs are administered 65 for their analgesic, antipyretic and anti-inflammatory properties to cure pain and inflammation in 66 67 both human and veterinary therapy. NSAIDs inhibit the synthesis and the release of prostaglandins from arachidonic acid, acting as non-selective inhibitors of cyclooxygenase (COX) enzymes, 68 namely cyclooxygenase-1 (COX-1) and cyclooxygenase-2 (COX-2) isoforms (Gierseet al., 1995). 69 70 Different NSAIDs have been prescribed extensively or are commercialized over-the-counter worldwide. For instance, more than 70 million prescriptions are written each year in the United 71 States, while considering the over-the-counter sale, more than 30 billion NSAID doses are 72 consumed annually in the United States alone (Wiegand and Vernetti, 2017). Because of their huge 73 and increasing use, coupled with their specific pharmacokinetic properties, NSAIDs reach 74 75 detectable concentrations in both sewage and surface water, accounting for 15% of pharmaceuticals measured in aquatic ecosystems worldwide (Santos et al., 2010). NSAIDs and analgesics are the 76 most frequently detected class of pharmaceuticals in the environment (Fekadu et al., 2019), as a 77 number of monitoring surveys have reported levels of NSAIDs exceeding 1 µg/L in influent and 78 79 effluents of WWTPs, while lower concentrations (in the ng/L range) have been found in surface waters (Santos et al., 2010; aus der Beek et al., 2016; Bagnis et al., 2018; Fekadu et al., 2019). 80 Diclofenac is the most frequently detected pharmaceutical in environmental samples, while also 81 ibuprofen and naproxen were detected nearly as often as diclofenac globally (aus der Beek et al., 82 2016). Moreover, five NSAIDs, including acetylsalicylic acid (ASA), paracetamol (PCM), 83 diclofenac (DCF), ibuprofen (IBU) and naproxen (NPX) are included in the list of sixteen 84 substances that were detected in surface, drinking, and groundwater of all the five United Nations 85 (UN) regional groups (i.e., Africa Group, Asia-Pacific Group, Eastern Europe Group, Latin 86

American and Caribbean States Group, and Western Europe and Others Group, which also includes 87 North America, Australia and New Zealand), with global average concentrations ranging between 88 0.032 and 0.922 µg/L (aus der Beek et al., 2016). Although the concentrations of NSAIDs in 89 90 freshwaters can be considered as relatively low, their high biological activity may pose a serious risk towards non-target species at different levels of the ecological hierarchy, leading to dissimilar 91 toxic effects. To date, some previous reviews or meta-analyses have summarized the occurrence, 92 toxicity and/or environmental risk of diverse pharmaceuticals (e.g., aus der Beek et al., 2016; 93 Bagnis et al., 2018; Fekadu et al., 2019; Ohoro et al., 2019) or a specific compound (e.g., 94 diclofenac; Acuña et al., 2015; Lonappan et al., 2016) in aquatic ecosystems, but none has 95 specifically focused on the toxicity of drugs belonging to a specific class of pharmaceuticals 96 towards freshwater invertebrates. Thus, the present review aimed at summarizing the toxicity 97 induced by the exposure to ASA, PCM, DCF, IBU and NPX towards freshwater invertebrates. 98 99 Freshwater invertebrate species globally account for approximately 2% (150,000 estimated species grouped in 17 phyla; Strayer, 2006) of an estimated 6.7 million invertebrate species (Collen et al., 100 101 2012). Although individually small and inconspicuous, aquatic invertebrates play a pivotal role in 102 ecosystem functioning, including the transfer of energy from autotrophs to higher levels of the food web and the recycling of nutrients (e.g., Pingram et al., 2014; Macadam and Stockan, 2015). 103 Moreover, many invertebrates species are easy to be cultured and maintained under laboratory 104 conditions, and are very sensitive to exogenous stresses, including the exposure to environmental 105 contaminants, making them excellent model organisms in ecotoxicological surveys. A systematic 106 literature research was performed in Google Scholar, Scopus and Web of Science databases. 107 Literature research was focused on papers published in the 2000-2019 period of time, using for each 108 single pharmaceutical compound different combinations of keywords dealing with their effects on 109 freshwater invertebrates, including pharmaceutical drugs, non-steroidal anti-inflammatory drugs, 110 freshwater, invertebrates, effects, toxicity. 111

3. Features of focal Non-Steroidal Anti-inflammatory Drugs (NSAIDs)

Acetylsalicylic acid, diclofenac, naproxen, ibuprofen and paracetamol are the most common 113 NSAIDs detected in aquatic environments (Fekadu et al., 2019). The main physico-chemical 114 properties of the investigated NSAIDs are reported in Table 1. Acetylsalicylic acid (ASA; 2-115 116 (acetyloxy)benzoic acid) has remained for over 90 years as one of the most prescribed analgesics in human medical care worldwide (Katzung, 2015). ASA is commonly used to reduce pain, fever, or 117 inflammation and after oral administration it overwhelms hepatic metabolic reactions that transform 118 119 it into conjugates (e.g., glucoronides) ease to be excreted. Although as much as 80% of a single therapeutic doses of ASA is metabolized in the liver, the remaining part is excreted as unchanged 120 parent compound, entering the sewage. ASA was detected in sewage effluent and surface water at 121 maximum levels of 1.5 and 3.1 µg/L (e.g., Ternes, 1998; Schulman et al., 2002) respectively, even 122 if concentrations up to 13 µg/L (Santos et al., 2010 and references therein) and even 59.6 µg/L were 123 detected in wastewater treatment plants from Spain (Metcalfe et al., 2003). ASA was detected in 124 surface waters, groundwater and/or tap or drinking waters from 15 Countries worldwide, with 125 global average and maximum measured environmental concentrations of 0.922 µg/L and 20.96 126 µg/L, respectively (aus der Beek, 2016). 127

Paracetamol (PCM; N-(4-hydroxyphenyl)acetamide), also known as acetaminophen, is an analgesic and antipyretic drug. Although PCM does not own a proper anti-inflammatory action, it is usually included in the NSAID therapeutic group by a toxicological point of view because its mechanism of action is similar to that of NSAIDs (Misra et al., 1990). PCM can be purchased as an over-thecounter sale drug in most Countries worldwide and represents one of the most frequently detected pharmaceuticals in surface waters, wastewaters and drinking water. A recent review showed that PCM was detected in surface waters, groundwater and/or tap or drinking waters from 29 Countries worldwide, with global average and maximum measured environmental concentrations of 0.161 μ g/L and 230 μ g/L, respectively (aus der Beek, 2016).

Diclofenac (DCF; 2-[(2,6-dichlorophenyl)amino] phenylacetic acid) is a phenylacetic acid used to 137 138 reduce inflammation and pain associated with arthritis, osteoarthritis, rheumatoid arthritis, and ankylosing spondylitis (Todd and Sorkin, 1988). As DCF can be sold both as an over-the-counter 139 sale drug and under medical prescriptions, it is one of the main drugs used worldwide and, 140 consequently, one of the main pharmaceuticals contaminating the aquatic ecosystems. As WWTPs 141 have a limited efficiency of removal, DCF is commonly detected at low µg/L range in WWTP 142 effluents of Europe and North and South America (Roberts and Thomas, 2006; Gómez et al., 2007). 143 Accordingly, DCF was commonly detected also in surface waters, in concentrations ranging 144 145 between low ng/L up to low µg/L (Metcalfe et al., 2003; Bound and Voulvoulis, 2006; Gros et al., 2006). Because of its occurrence in aquatic ecosystems and potential toxicity, the European Union 146 has included DCF to the list of the Water Framework Directive (2013/39/EU) as priority molecules 147 to be monitored in aquatic ecosystems. However, as sufficient high-quality monitoring data were 148 obtained for DCF, the EU commission decided that this substance should be removed from the 149 watch list (commission implementing decision 2018/840/EU). DCF was detected in surface waters, 150 groundwater and/or tap or drinking waters from 50 Countries worldwide, with global average and 151 maximum measured environmental concentrations of 0.032 µg/L and 18.74 µg/L, respectively (aus 152 der Beek, 2016). 153

Ibuprofen (IBU; ((+/-)-2-(p-isobutylphenyl) propionic acid with R and S isomers) is used to relieve the symptoms of arthritis, rheumatic disorders, pain and fever (Hayashi et al., 2008). IBU represents one of the core pharmaceuticals included in the "Essential Drug List" of the World Health Organization (WHO), and it is produced in large amounts worldwide (Heckmann et al., 2007). Because of its over-the-counter sale, large prescription volume and high excretion rate (~70-80% of

the therapeutic dose), IBU has been identified as one of the main pharmaceuticals in aquatic 159 ecosystems. Moreover, IBU has relatively high mobility into aquatic environments, but a lower 160 persistence in comparison with other pharmaceuticals (Buser et al., 1999). IBU was detected in 161 moderate to high concentrations both in the effluents of WWTPs and in surface waters during 162 surveys carried out in both Europe and North America (Metcalfe et al., 2004; Santos et al., 2010). 163 IBU was detected in surface waters, groundwater and/or tap or drinking waters from 47 Countries 164 165 worldwide, with global average and maximum measured environmental concentrations of 0.108 μ g/L and 303 μ g/L, respectively (aus der Beek, 2016). 166

Naproxen (NPX; (S) 6-methoxy- α -methyl-2-naphthalene acetic acid) is a prototypical member of 167 NSAIDs, commonly used in the treatment of migraine, rheumatoid arthritis, and osteoarthritis. 168 169 Naproxen is metabolized in the liver and eliminated in its unchanged form (10% of the dose) or as metabolites (60% of the dose) through the urine and feces. NPX can undergo diverse 170 biotransformation pathways, the most important being conjugation with glucuronic acid to form 171 naproxen-β-1-O-acyl glucuronide, as well as O-dealkylation made by CYP2C9 and CYP1A2 172 enzymes, leading the production of 6-Odesmethylnaproxen (Hug, 2006). NXP was commonly 173 detected in surface waters at concentrations up to 32 µg/L in Pakistan, 4.5 mg/L in Canada, 0.328 174 mg/L in China, and 0.24 mg/L in Japan (Brun et al., 2006; Komori et al., 2013; Zhao et al., 2010). 175 A recent review showed that NPX was detected in surface waters, groundwater and/or tap or 176 drinking waters from 45 Countries worldwide, with global average and maximum measured 177 environmental concentrations of 0.050 µg/L and 32 µg/L, respectively (aus der Beek, 2016). 178

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4. Accumulation and biotransformation of NSAIDs in freshwater invertebrates

The studies on accumulation of NSAIDs in freshwater invertebrates were performed on different 183 species belonging to different taxa. NSAID residues in invertebrates were relatively higher 184 compared with other classes of pharmaceuticals and the most frequent NSAIDs accumulated and 185 measured in invertebrates were DCF and IBU (Miller et al., 2018). Focusing on freshwater 186 invertebrates, some studies investigated the accumulation of NSAIDs in invertebrates from different 187 ecosystems worldwide. Measureable concentrations of DCF (12.4 ng/g dry weight) and IBU (183 188 ng/g dry weight) were found in Hydropsyche spp. individuals from the River Segre (Spain; Huerta 189 190 et al., 2015). An in-situ study aimed at developing an analytical method to determine emerging 191 contaminants in benthic invertebrates, namely Gammarus fossarum, Potamopyrgus antipodarum and Chironomus riparius, organisms were exposed upstream and downstream of a WWTP located 192 on a French river that drains dense urban and industrial areas in the Southeast of France (Berlioz-193 Barbier et al., 2014). IBU was measured in G. fossarum specimens (concentration range 60.6 -194 105.4 ng/g wet weight) but not in P. antipodarum specimens (< limit of quantification), while DCF 195 was measured in C. riparius specimens (concentration range 26 - 51.5 ng/g wet weight; Berlioz-196 Barbier et al., 2014). Diclofenac was detected in measurable concentrations in Planorbis spp. (13 197 198 ng/g wet weight), Hyalella azteca (20 ng/g wet weight), Utterbackia imbecillis (15 ng/g wet weight) and Corbicula fluminea (23 ng/g wet weight) from the North Bosque River (Texas, USA; Du et al., 199 2015). Diclofenac was accumulated in diverse mussel species from the Taihu Lake (China), 200 including Anodonta spp. (2.45 ng/g dry weight), Bellamya spp. (3.14 ng/g dry weight), 201 Corbiculidae (2.59 ng/g dry weight), and in Siberian prawn Exopalaemon modestus (6.7 ng/g dry 202 weight), while IBU was measured at higher mean concentrations than DCF in tissues of Bellamya 203 spp. (71.23 ng/g dry weight), Corbiculidae (41.6 ng/g dry weight), and in Siberian prawn (24.96 204 ng/g dry weight) (Xie et al., 2015). Gabricova and co-authors (2015) measured DCF in Erpobdella 205

octoculata specimens (19.66 ng/g wet weight) from the Zivny Stream (Czech Republic), while Ruhi 206 et al. (2016) found DCF in Hydropsyche (9 ng/g dry weight) and IBU in Hydropsyche (182.7 dry 207 weight) and Phagocata vitta (30.9 dry weight) from the Segre River (Spain). A study by Ikkere and 208 209 coauthors (2018) investigated the presence of different NSAIDs (i.e., tolfenamic acid, meloxicam, carprofen, flunixin, diclofenac, ibuprofen, phenylbutazone, ketoprofen and mefenamic acid) in soft 210 tissues of four freshwater mussels, namely Unio tumidus, Anodonta anatina, Anodonta cygnea and 211 Dreissena polymorpha, from Latvian ecosystems. Only IBU was detected in half of analyzed 212 samples in concentrations ranging between 0.52 and 109 ng/g wet weight. A recent study by Yang 213 and co-authors (2020) investigated the levels, bioaccumulation, and trophic transfer of 45 214 pharmaceuticals and personal care products, including some DCF and IBU, in highly urbanized 215 rivers, namely the New Qinhuai River, the Qinhuai River and a section of the Yangtze River 216 (China). DCF and IBU were detected in measureable concentrations in phytoplankton (DCF 217 218 concentration range = 1.3 - 8.4 ng/g wet weight; IBU concentration range = 14.5 - 35.8 ng/g wet weight), zooplankton (concentration range = 2.1 - 12.4 wet weight; IBU concentration range = 20.9219 220 -48.9 ng/g wet weight) and three invertebrate species (i.e., freshwater shrimps, mussels and snails; 221 concentration range = 1.1 - 5.9 wet weight; IBU concentration range = 4.8 - 11.6 ng/g wet weight) from the three rivers (Yang et al., 2020). 222

After uptake, NSAIDs can undergo biotransformation processes. Cytochrome P450 mixed function oxidase (MFO) systems play a crucial role in oxidation of drugs and xenobiotics in humans and in a number of species, including bacteria, plants, fish and aquatic invertebrates (Snyder, 2000; Rewitz et al., 2006; Gottardi et al., 2016). Different P450 gene families (CYP) have been characterized in fish and invertebrates (Stegeman and Livingstone, 1998). The CYP2 family, particularly the subfamily CYP2C9, has been identified as the responsible for NSAID biotransformation (Blanco et al., 2005; Zanger et al., 2008). Another pathway of biotransformation of carboxylate NSAIDs, such

as ASA, DCF, NPX and IBU) involves the glucuronic acid conjugation catalyzed by the uridine 230 diphosphoglucuronosyl transferase superfamily of enzymes, resulting in acyl glucuronides 231 (Pritchard, 1993). These compounds are reactive intermediates that can undergo acyl migration and 232 hydrolysis and can also form adducts with nucleophilic amino acid residues (Pritchard, 1993). 233 Many NSAID-derived acyl glucuronides, including those obtained from DCF and IBU, have been 234 shown to form covalent bonds with intra and extracellular proteins, with toxicological consequences 235 236 (Boelsterli, 2007). Information on biotransformation products is currently limited to surface waters and biota. In detail, they have been scarcely determined across invertebrates, with only 12 reported 237 concentrations (Miller et al., 2018). However, to date none of such studies included NSAIDs. Thus, 238 239 developing new methods to measure biotransformation products in invertebrates represents a priority in NSAID ecotoxicology. The measurement of accumulated levels of NSAIDs and their 240 biotransformation products in organisms should allow to perform a more reliable risk assessment 241 242 for these compounds in the environment and to address prioritization of hazardous compounds, as well as to study potential pharmacological or toxicological effects for the understanding of the risk. 243 244 In fact, the quantification of NSAIDs associated with effect-based studies should allow to shed light on the cause-effect relationship and threshold associated with the onset of the effect, avoiding 245 extrapolation of exposure concentrations to observed effects (Miller et al., 2018). 246

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5. Toxicity of NSAIDs towards freshwater invertebrates

Toxic effects induced by the exposure to the main NSAIDs measured in freshwater ecosystems, namely acetylsalicylic acid, paracetamol, diclofenac, ibuprofen and naproxen, in freshwater invertebrates were performed on different model species belonging to different *taxa* (Figure 1). The number of studies investigating NSAID chronic toxicity were higher than those focused of acute

toxicity. Crustaceans, mainly the Cladoceran Daphnia magna, were the main model organisms used 253 to explore both acute and chronic toxicity of NSAIDs, followed by Mollusca. In Table 2-7 are 254 summarized the studies investigating the acute or chronic toxicity of selected NSAIDs and mixtures 255 256 towards freshwater invertebrates. Acute toxicity describes the effects induced by either a single exposure or multiple exposures in a short time period and appears as lethal endpoints (e.g., 257 mortality or immobilization). Chronic toxicity describes the onset of adverse effects resulting from 258 259 prolonged and repeated exposure to stressors, which appears as sub-lethal endpoints (e.g., growth inhibition, molecular or biochemical alterations, behavioral changes). 260

261 5.1. Toxic effects induced by acetylsalicylic acid (ASA)

262 The toxicity of acetylsalicylic acid (ASA) was investigated on two crustacean species (Daphnia magna and Daphnia longispina) and a planarian species (Dugesia japonica) (Table 1). Acute and 263 chronic toxicity of ASA was investigated on Daphnia magna and Daphnia longispina through the 264 265 assessment of survival (i.e., immobilization or mortality) or reproduction and growth (Marques et al., 2004a). After 48 hrs of exposure, the 50% Effect Concentration (EC₅₀) of ASA for D. 266 longispina was 647.31 mg/L and it was about half compared to that calculated for D. magna 267 (1,293.05 mg/L). These results were different from those found by other studies of D. magna, 268 showing that the 50% Lethal Concentration (LC₅₀) at 48 hrs of ASA was 88.33 mg/L (Gómez-269 Oliván et al., 2014) and the EC₅₀ at 48 hrs was 88.1 mg/L (Cleuvers, 2004). Increasing 270 concentrations of ASA significantly affected the fecundity of D. magna and D. longispina. Similar 271 No Observed Effect Concentration (NOEC) and Lowest Observed Effect Concentration (LOEC) for 272 273 both the Cladoceran species (i.e., NOEC = 1.00 mg/L and LOEC = 1.80 mg/L) were found (Marques et al., 2004a). Considering the population intrinsic growth rate (r), ASA boosted the 274 275 population growth in D. longispina, whereas an opposite trend was observed for D. magna 276 (Marques et al., 2004a). Sub-lethal effects were induced by the exposure to 17.9 mg/L of ASA in terms of defects in eye regeneration, unstimulated behavior, and scrunching in regenerating tails,
but not in full worms, in the freshwater planarian *D. japonica*, suggesting developmental injuries
(Zhang et al., 2019).

Chronic toxicity of ASA was also investigated at biochemical level by the application of a battery 280 of different biomarkers of oxidative stress and genotoxicity, namely lipid peroxidation, protein 281 carbonyl content, activity of the antioxidant enzymes superoxide dismutase, catalase and 282 glutathione peroxidase and DNA fragmentation (Gómez-Oliván et al., 2014). The exposure to a 283 single concentration (8.8 mg/L) of ASA, corresponding to equal to the Lowest Observed Adverse 284 Effect Level (LOAEL) obtained from a previous acute assay, induced the modulation of the activity 285 of antioxidant enzymes, as well as the increase of lipid peroxidation and DNA fragmentation in 286 287 treated specimens compared to controls (Gómez-Oliván et al., 2014).

288 5.2 Toxic effects induced by paracetamol (PCM)

The toxicity of paracetamol (PCM) was assessed on diverse freshwater invertebrate species, including a cnidarian (*Hydra vulgaris*) and a rotifer (*Plationus patulus*) species, different crustacean (*Daphnia magna, Daphnia longispina* and *Moina macrocopa*) and bivalve (*Dreissena polymorpha* and *Corbicula fluminea*) species, as well as a planarian species (*Dugesia japonica*) (Table 2).

PCM median lethal concentrations (LC₅₀) for *D. magna* was 224, 40.0, 8.06 and 5.32 mg/L at 24, 293 48, 96 hrs and 21 days, respectively (Du et al., 2016). In the same species, the PCM median effect 294 concentrations (EC_{50}) for body length, number of carapaces per adult, number of broods per female 295 and egg production per female was 4.78, 4.21, 2.38 and 1.12 mg/L, respectively (Du et al., 2016). A 296 study performed by Nunes and coauthors (2014) investigated the toxicity of PCM towards different 297 298 freshwater species, including D. magna and D. longispina. PCM toxicity was widely variable among species, even among species that were phylogenetically related. Considering acute toxicity 299 in terms of EC₅₀ for invertebrates, D. magna (4.7 mg/L) was more sensitive than D. longispina 300

(67.9 mg/L). Moreover, PCM caused mortality during a chronic toxicity reproduction test with D. 301 302 magna at the highest tested concentrations (between 1.2 and 1.7 mg/L). Although treated specimens generated offspring, they did not survive over the whole duration of the experiment. A different 303 304 response was observed for *D. longispina*, which showed a significant delay in the first reproductive event and a reduction in the fecundity, but no mortality. A study by Sarma and coauthors (2014) 305 exposed the rotifer *Plationus patulus* and the cladoceran *Moina macrocopa* to increasing 306 concentrations of PCM (concentration range 2 - 32 mg/L) in order to assess changes in population 307 growth. Population growth curves of both the species were affected by PCM concentrations, 308 showing a decrease in organism density with increasing levels of the drug. Moreover, the daily rate 309 of population growth was negatively affected by PCM exposure in both the zooplanktonic species. 310 A 7-days exposure to increasing PCM concentrations (concentration range 0.01 - 10 mg/L) did not 311 affect the survival of *Hydra vulgaris* specimens at concentrations up to 1.0 mg/L, while no adverse 312 313 effects on feeding and bud formation were induced after 17 days of exposure to the same concentration range. Moreover, the ability of dissected polyps to regenerate hypostome, tentacles 314 315 and foot was not altered (Pascoe et al., 2003). All the studies mentioned above showed that acute 316 effects of PCM occur only at mg/L concentrations, while sub-lethal effects arise also at lower, often environmentally relevant, concentrations. A decreased unstimulated speed in day 12 regenerating 317 tails was observed in *D. japonica* specimens exposed to 15.5 mg/L of PCM (Zhang et al., 2019). 318 Chronic toxicity of PCM was also investigated at low levels of the biological organization. 319 Biochemical effects of PCM exposure were investigated in the freshwater clam Corbicula fluminea 320 following short- (96-hrs) and long-term (28-days) exposures to increasing PCM concentrations 321 322 (Brandão et al., 2014). No mortality was observed in clams over short- or long-term exposures.

PCM did not modulate catalase activity but induced a significant decrease of glutathione Stransferase (GST) and glutathione reductase (GR) activity over both short- and long-term exposures. A significant increase of lipid peroxidation was noted at the end of short- and long-term

exposure to the highest PCM tested concentrations. These results indicated that the exposure to 326 increasing PCM concentration caused notable changes in the cellular redox status of C. fluminea 327 (Brandão et al., 2014). The in-vitro cytogenotoxicity of PCM was investigated through the 328 application of a battery of four biomarkers (i.e., the comet test to investigate DNA fragmentation 329 and frequency of apoptotic and necrotic cells, and the neutral red retention assay - NRRA) on 330 hemocytes collected from the zebra mussel Dreissena polymorpha exposed for 1 hour to 30, 150 331 and 450 µg/L (Parolini et al., 2009). Dose-dependent decrease in the stability of lysosome 332 membranes (NRRA), coupled with a significant increase in both primary (DNA fragmentation) and 333 fixed (frequency of apoptotic and necrotic cells) genetic damage was induced by PCM. In-vivo 96-334 hrs exposures of *D. polymorpha* specimens to three, environmentally relevant PCM concentrations 335 $(0.154; 0.75 \text{ and } 1.51 \text{ }\mu\text{g/L})$ showed that this drug can alter the oxidative status of this bivalve 336 species (Parolini et al., 2010). Low PCM concentrations did not cause neither mortality of zebra 337 338 mussel over the duration of the experiment nor changes in hemocyte viability. Although PCM did not induce primary genetic injuries in zebra mussel hemocytes at all the tested concentration, a 339 340 significant increase of fixed genetic damage, in terms of both micronuclei and apoptotic frequency, was noted at the end of the exposure to the highest tested concentrations. Moreover, a significant 341 destabilization of lysosomal membranes and significant modulation of catalase, glutathione 342 peroxidase (GPx) and GST activity was induced by the exposure to 0.75 and 1.51 µg/L of PCM. 343

344 5.3. Toxic effects induced by Diclofenac (DCF)

Acute and chronic toxicity of diclofenac (DCF) towards freshwater invertebrates was explored on a rotifer (*Plationus patulus*), two crustaceans (*Daphnia magna* and *Moina macrocopa*), a chironomid (*Chironomus riparius*), a bivalve (*Dreissena polymorpha*) and a gastropod (*Lymnea stagnalis*) species (Table 3).

Mortality of *D. magna* specimens arose after only 24 hrs of exposure to a high DCF concentration 349 (486 mg/L). DFC exposure caused 50% mortality (EC₅₀) in *D. magna* after 21 days of exposure to 350 2.00 mg/L and a significant reduction of egg production at the lowest exposure concentrations of 351 352 0.50 mg/L (Du et al., 2016). A study by de Oliveira and coauthors (2016) calculated that the EC₅₀ for *D. magna* was 123.3 mg/L, but no effects on population growth was noted after the exposure to 353 a range of increasing DCF concentrations (concentration range 29.5 - 75 mg/L). A 21-days 354 exposure to four increasing DCF concentrations (concentration range 5 - 5,000 µg/L) did not cause 355 significant changes in molting frequency, number of eggs produced in the first brood, total number 356 of eggs per individual, total number of broods per individual, body length and growth rate in D. 357 358 magna specimens (Liu et al., 2017). In the same study, 96-hrs exposure to 50 µg/L of DCF induced significant changes in the expression of some genes related to detoxification, growth, development 359 and reproduction, which were inhibited after 24 hrs and overexpressed after 48 hrs of exposure (Liu 360 361 et al., 2017). In contrast, the exposure to increasing concentrations of DCF (concentration range 2 -32 mg/L) affected the population growth curves of the rotifer *Plationus patulus* and the cladoceran 362 Moina macrocopa, leading to a decrease in organism density with increasing levels of drug and 363 negative effects on the daily rate of population increase (Sarma et al., 2014). 364

Chronic toxicity of DCF was investigated also at molecular and biochemical level in different 365 invertebrate species. A research by Haap and co-authors (2008) investigated the toxicity of DCF at 366 biochemical level in *D. magna* by assessing the modulation of heat shock protein 70 (hsp70) level 367 as a biomarker for proteotoxicity, showing that the modulation of such protein occurred only at 368 concentrations of DCF higher than 40 mg/L. The cyto-genotoxicity of DCF was investigated 369 through an *in-vitro* approach by exposing for 1 hour hemocytes collected from the zebra mussel D. 370 polymorpha to 60, 126 and 250 µg/L (Parolini et al., 2009). A significant cytotoxic effect, in terms 371 of destabilization of the lysosomal membranes, was noted only after the exposure to 250 µg/L of 372 DCF, while both primary genetic lesions (i.e., DNA fragmentation) and fixed damage to DNA (i.e., 373

frequency of apoptotic and necrotic cells) occurred after the exposures to all the tested 374 concentrations. A further *in-vitro* experiment (Parolini et al., 2011a) investigated the toxicity of 375 increasing DCF concentrations (0.001, 0.01, 0.1, 1 and 10 mg/L) on three different cell typologies 376 of the zebra mussel, namely hemocytes, gill and digestive gland cells. After 96 hrs of exposure, the 377 viability of DCF treated gill cells was significantly reduced already at the lowest tested 378 concentration. Moreover, the viability of DCF-treated digestive gland cells was significantly 379 reduced already after 48 hours exposure to 0.01 mg/L, while hemocyte viability was reduced 380 already at the lowest concentration (0.001 mg/L). An *in-vivo* 96-hrs exposure of the zebra mussels 381 to three increasing concentrations (95, 318 and 637 ng/L) of DCF showed a negligible cyto-382 genotoxicity. In fact, only a slight decrease of lysosomal membrane stability was observed at the 383 end of exposure to the highest tested concentration (637 ng/L), while no other effects arose (Parolini 384 et al., 2011b). DCF sub-lethal toxicity in terms of immunotoxicity was assessed on the gastropod 385 386 Lymnaea stagnalis exposing specimens for 3 days to environmentally relevant (concentration range $1 - 10 \mu g/L$) and therapeutic concentrations (concentration range $100 - 1,000 \mu g/L$) of DCF 387 388 (Boisseaux et al., 2017). Diclofenac induced immune responses, while no immunosuppression was 389 observed. DCF significantly affected the immunocapacity and the immunoefficiency of the snails' hemocytes. This effect is typical of an inflammatory response, confirmed by the increase of the 390 NADPH-oxidase activity, mainly at 1,000 µg/L. A 10-days chronic toxicity test with C. riparius 391 was performed to assess effects on survival, growth and developmental stage, in terms of biomass, 392 as well as emergence rates and sex ratio after 21 days of exposure to DCF-spiked sediments. No 393 effects on survival and no change in the sex-ratio was induced by DCF exposure. In contrast, DCF 394 decreased the emergence ratio in organisms exposed at concentrations of 34.0 µg/g of DCF (Nieto 395 et al., 2017). 396

397 5.4. Toxic effects induced by Ibuprofen (IBU)

Acute and chronic toxicity of ibuprofen (IBU) towards freshwater invertebrates was investigated on a crustacean (*Daphnia magna*), a cnidarian (*Hydra vulgaris*), two bivalve (*Dreissena polymorpha* and *Corbicula fluminea*) and a gastropod (*Planorbis carinatus*) species (Table 4).

Mortality of all D. magna specimens was caused after 24 hrs of exposure to high levels of IBU (200 401 mg/L), while EC₅₀ was calculated as 3.97 (Du et al., 2016). The 48 hrs EC₅₀ for immobilization was 402 estimated as 108 mg/L, while the reproduction was reduced above 10 mg/L and that survival was 403 unaffected at concentrations up to 40 mg/L, with no survival above 80 mg/L (Heckmann et al., 404 2005). A further study revealed that a 14-days exposure of D. magna to increasing IBU 405 concentrations affected life-history traits and population performance of the species (Heckmann et 406 407 al., 2007). Population growth rate significantly decreased at all the IBU tested concentrations, while survival was affected only at the end of the exposure to 80 mg/L of IBU. In contrast, reproductive 408 effort was affected also by the exposure to the lowest IBU tested concentrations, whereby the 14-409 410 days EC₅₀ was calculated as 13.4 mg/L, while it was completely inhibited at 80 mg/L. Similar results were obtained by Hayashi and coauthors (2008), who exposed 5-days old D. magna 411 412 specimens to the same range of IBU concentrations tested by Heckmann and coauthors (2007) for 413 10 days. Specimens exposed to 40 mg/L of IBU generated significantly fewer offspring than controls, while no reproduction occurred at 80 mg/L. Moreover, a significant delay of the first 414 reproductive event occurred at all the tested IBU concentrations. Daphnia magna survival was 415 affected after the exposure to 80 mg/L during the 10-day exposure, while the population grew after 416 the exposure to control, 20 and 40 mg/L of IBU. In contrast, a significant decrease of population 417 growth was noted at the end of the exposure to 80 mg/L of IBU (Hayashi et al., 2008). A 7-days 418 419 exposure to increasing IBU concentration did not affect the survival of the cnidarian H. vulgaris at concentrations up to 1 mg/L, while after 17 days of IBU administration neither feeding nor bud 420 421 formation nor ability of dissected polyps to regenerate a hypostome, tentacles and foot were affected (Pascoe et al., 2003). However, another study showed that the exposure to 5 mg/L of IBU 422

inhibited the regeneration of the cnidarian, with a 96-hrs IC₅₀ (i.e., the concentration that inhibits 423 50% of the embryos to develop) was calculated as 3.84 mg/L (Quinn et al., 2008). Acute and sub-424 lethal effects of IBU were also investigated on the freshwater Keeled rams horn snails (Planorbis 425 426 *carinatus*) exposed for 72-hrs or 21-days to different IBU concentrations. The 48- and 72-hrs LC_{50} were 17.1 mg/L at both the time points, while the 21-days LOEC and NOEC based on the survival 427 of specimens were calculated as 45.36 and 5.36 mg/L, respectively. In addition, the 21-days LOEC 428 and NOEC calculated for snail reproduction (i.e., hatching success) were 5.36 and 2.43 mg/L, 429 respectively, while for growth were 2.43 and 1.02 mg/L, respectively (Pounds et al., 2008). 430

Chronic toxicity of IBU was investigated also at molecular and biochemical level through the 431 application of different techniques. A study by Wang and coauthors (2016) investigated the 432 modulation of the expression of CYP360A, CYP314, and GST genes involved in the detoxification 433 434 process and the responses of their associated enzymes activity, as well as in some physiological parameters (e.g., growth and reproduction) in D. magna specimens exposed to three 435 436 environmentally relevant concentrations of IBU. The exposure to IBU did not affect the reproduction of cladocerans, in terms of total amount of eggs produced and total number of clutches 437 per female, as well as of body length of treated specimens. The same experiment showed that the 438 439 treatment with 0.5 µg/L of IBU inhibited the expression of CYP360A gene, while at 50 µg/L an overexpression of such gene occurred. A similar trend was also induced by the exposure to at 0.5 440 µg/L of IBU towards the GST gene, while CYP314 expression was inhibited after a short time 441 exposure (6 hrs), while was overexpressed after prolonged exposure time (48 hrs). Similarly, 442 erythromycin N-demethylase and aminopyrine N-demethylase were both inhibited after 6-hrs 443 exposure but overexpressed after 48-hrs exposure to 0.5 µg/L. Moreover, an induction of 444 445 glutathione S-transferase (GST), superoxide dismutase (SOD), and catalase (CAT) activity was observed in short-term exposure to IBU, while a dose-dependent increase of EROD activity and 446 447 methane dicarboxylic aldehyde (MDA) content occurred (Wang et al., 2016). The cyto-genotoxicity

of IBU was investigated through an *in-vitro* approach by 1-h exposure of zebra mussel hemocytes 448 to different IBU concentrations (Parolini et al., 2009). The stability of lysosomal membranes was 449 reduced at the endo of the exposure to 450 and 909 µg/L of IBU, while genotoxicity, in terms of 450 DNA fragmentation and frequency of apoptotic cells, occurred in response to all the tested 451 concentrations. A further 96-hrs in-vivo exposure of the zebra mussel showed that IBU induced a 452 slight cyto-genotoxicity on hemocytes at 0.2 µg/L of IBU, while higher concentrations (2 and 8 453 µg/L) significantly affected lysosomal membrane stability and arose both primary and fixed genetic 454 damage. In addition, IBU modulated the activity of antioxidant and detoxifying enzymes at all the 455 tested concentrations, suggesting that this drug can imbalance the oxidative status of mussels and 456 provoke the onset of oxidative stress also at low, environmental concentrations (Parolini et al., 457 2011c). Similar results were obtained on the clam Corbicula fluminea (Aguirre-Martínez et al., 458 2015). IBU induced a destabilization of lysosomal membrane at all the tested concentrations, as 459 460 well as the increase of the activity of phase I and II enzymes, including the activation of GR and GPx at the highest tested concentration (50 mg/L). Moreover, an increase of lipid peroxidation, but 461 not of DNA damage was observed at the end of the exposure to 50 µg/L (Aguirre-Martínez et al., 462 2015). Another study performed on the zebra mussel investigated the effects of 7-days exposure to 463 IBU at molecular level, exploring the changes in the expression of mRNA of enzymes and other 464 proteins involved in the prevention of protein damage (hsp70) and oxidative stress (SOD, CAT and 465 metallothionein), in the biotransformation (GST, aryl hydrocarbon receptor) and elimination (P-466 glycoprotein) of xenobiotics, as well as in reversible protein post-translational modification (protein 467 phosphatase 2A). Zebra mussel specimens exposed to the lowest tested concentration of IBU 468 experienced an oxidative stress situation as pointed out by the induction of mRNA levels observed 469 for SOD, CAT and metallothionein in the digestive gland after 1 or 4 days of treatment. At the 470 471 higher concentrations, an increase in levels of transcripts for GST occurred, suggesting the

472 activation of biotransformation processes of IBU or by-products deriving from oxidative stress473 (Contardo-Jara et al., 2011).

474 5.5 Toxic effects induced by Naproxen (NPX)

Acute and chronic toxicity induced by naproxen (NPX) towards freshwater invertebrates was 475 investigated on a rotifer (Brachionus calyciflorus), four crustacean (Thamnocephalus platyurus, 476 Ceriodaphnia dubia, Daphnia magna, Hyalella azteca), a cnidarian (Hydra magnipapillata) and a 477 bivalve (Elliptio complanata) species (Table 5). Acute and chronic toxicity of NPX was assessed 478 through standard bioassays using rotifers (Brachionus calyciflorus) and 479 crustaceans (Thamnocephalus platyurus, Ceriodaphnia dubia). Naproxen acute toxicity for T. platyurus and B. 480 481 *calvciflorus* (LC₅₀; mortality) was calculated as 62.48 mg/L and 84.09 mg/L, respectively, while for C. dubia (EC₅₀; immobilization) was 66.37 mg/L (Isidori et al., 2005). Other studies showed that 482 acute toxicity of NPX, in terms of immobilization EC₅₀ to crustaceans, were in the same order of 483 484 magnitude (Cleuvers, 2004; Gheorghe et al., 2016; Kwak et al., 2018). The study by Isidori and coauthors (2005) showed that chronic tests were responsive at lower concentrations compared to 485 acute ones. Chronic toxicity in terms of inhibition or population growth for rotifers and crustaceans 486 (EC₅₀) was 0.56 (0.40–0.62) mg/L for T. platyurus and 0.33 (0.11–0.63) mg/L for C. dubia, 487 respectively. Similar results were obtained by Kwak and coauthors (2018) on D. magna and M. 488 489 macrocopa.

Sub-lethal effects of NPX were also focused on effects at molecular and biochemical levels. The exposure to NPX and its photoproducts din not induce genotoxicity neither in *E. coli* nor in *S. typhimurium* (Isidori et al., 2005). Immunotoxic effects of NPX, in terms of phagocytosis, intracellular esterase activity, adherence to microplate wells and lipid peroxidation, was tested on hemolymph collected from the freshwater mussel *Elliptio complanata* exposed *in vitro* to 2.5-100

 μ M (concentration range 0.57 – 23 mg/L) of NPX. Threshold effect level for phagocytosis, 495 intracellular esterase activity and adherence to microplate wells was 35, 152 and 4 µM of NPX, 496 respectively, while a significant decrease in lipid peroxidation occurred (Gagné et al., 2006). A 497 study of the amphipod Hyalella azteca investigating the onset of oxidative stress and the subsequent 498 oxidative damage to genetic material induced by NPX sodium enriched sediments showed that 48 499 hrs of exposure to sediments enriched with 76.6 and 339.2 mg/kg of NPX induced a dose-dependent 500 increase of genotoxicity, lipid peroxidation and protein carbonylation, as well as an increase of 501 502 SOD and CAT activity coupled with a decrease of GPx activity (García-Medina et al., 2015). Similar effects were found in D. magna exposed to 0.017 mg/L of NAP that showed a significant 503 increase of SOD and CAT activity and lipid peroxidation, while a decrease was noted for GPx. 504 Moreover, a significant increase of genetic damage (i.e., DNA fragmentation) was noted after 48 505 and 96 hrs of exposure to NAP (Gómez-Oliván et al., 2014). Lastly, a recent work by Yamindago 506 507 and coauthors (2019) showed that NPX median lethal concentrations (LC₅₀) in *H. magnipapillata* was 51.99 mg/L, 44.93 mg/L, and 42.50 mg/L after 24, 48, and 72 hrs of exposure, respectively. 508 509 Morphological observation of the exposed cnidarian showed that 40 mg/L of NPX stimulated the 510 contraction of body column and tentacles after 24 hrs. A KEGG pathway analysis of the genes differentially expressed in *H. magnipapillata* after NPX exposure for 6, 24, or 48 hrs pointed out 511 various cellular and metabolic effects, including protein processing in the endoplasmic reticulum, 512 Wnt signaling, and tryptophan metabolism (Yamindago et al., 2019). 513

514 5.6 Toxic effects induced by mixtures of NSAIDs

Acute and chronic toxicity induced by mixtures of NSAIDs towards freshwater invertebrates was investigated on two crustaceans (*Daphnia magna* and *Hyalella azteca*) and a bivalve species (*Dreissena polymorpha*) only (Table 6). A study of Cleuvers (2004) investigated the ecotoxicity of four NSAIDs, namely DCF, IBU, NPX and ASA, on *D. magna*. Toxicities of single NSAIDs were

relatively low, with half-maximal EC_{50} values in *Daphnia magna* ranging from 68 to 166 mg/L. 519 The toxicity of the mixture composed by the four NSAIDs at different concentrations ($EC_5/4$, 520 $EC_{10}/4$, $EC_{20}/4$, $EC_{50}/4$, and $EC_{80}/4$) was considerable, even at concentrations that caused no or 521 slight effects when the organism were exposed to the single substances. Parolini and Binelli (2012) 522 investigated the sub-lethal effects induced by a mixture of three common NSAIDs, namely 523 diclofenac, ibuprofen and paracetamol, on the freshwater bivalve zebra mussel (Dreissena 524 polymorpha). Zebra mussels were exposed to three mixtures in which each single drug was 525 included at the median value of surface waters (Low), wastewater effluents (Mid) and predicted 526 environmental concentration (PEC; High), similar to those previously tested in experiments 527 assessing the toxicity of each single molecule independently (Parolini et al., 2010; 2011b,c; 2013). 528 The exposure to the three mixtures induced a significant cellular stress in bivalves, in terms of 529 destabilization of the lysosome membranes (i.e., NRRA) probably caused by the alteration of the 530 531 oxidative status of the bivalves, as indicated by the modulation of the antioxidant enzyme activity. Moreover, the mixtures induced significant increase of both primary (i.e., DNA fragmentation) and 532 fixed (i.e., apoptosis and micronuclei) genetic damage. The effects induced by the mixture resulted 533 higher than those observed in previous studies when the single drugs were tested individually at 534 concentrations similar to those included in the mixtures (Parolini et al., 2010; 2011b,c). Another 535 study investigating the toxicity of binary mixtures of DCF with PCM, IBU, NPX, and ASA 536 compared to that of the same NSAIDs in their isolated form, on the amphipod Hyalella azteca 537 showed that NSAIDs induced oxidative stress both in isolated form and in binary mixtures (Gómez-538 Oliván et al., 2014). In detail, modulation of SOD, CAT, and GPx activity, as well as increase of 539 lipid peroxidation and protein carbonylation occurred with NSAIDs in isolated form and at higher 540 extend when NSAIDs where in binary mixtures. 541

543 **6.** Conclusions and future needs

The exposure to the most common NSAIDs found in the aquatic ecosystems, namely acetylsalicylic 544 acid, diclofenac, ibuprofen, naproxen, paracetamol, as well as to their mixtures, might represent a 545 risk for non-target, freshwater invertebrates. Although data from standardized tests, in terms of 546 EC_{50} or LC_{50} from acute or chronic toxicity tests, are comparable, it is not simple to draw a scale of 547 548 toxicity of NSAIDs because of some contrasting results among studies testing the toxicity of the same drug towards the same model species or the lack of information for specific drugs. However, 549 according to EC₅₀ values obtained on *D. magna*, which was the model species on which all the 550 551 NSAIDs were tested, DCF and IBU can be identified as the more toxic drugs, followed by PCM, NPX and ASA. Simultaneously, although different responses occurred within phylogenetic-related 552 species, cladocerans can be indicated as the more sensitive organisms to NSAID exposure. 553 Although acute toxicity of NSAIDs occurs only at high, unrealistic concentrations, much higher 554 than those currently measured in freshwaters worldwide, sub-lethal effects due to long-term 555 exposures cannot be neglected. A growing number of studies performed on diverse invertebrate 556 species belonging to different levels of the ecological hierarchy showed that the exposure to low, 557 environmentally relevant concentrations of NSAIDs, both independently and in mixture, can cause 558 559 a variety of adverse effects at molecular, biochemical and cellular level, while effects at individual level (e.g., growth, survival, reproduction) seem to be less probable, although they cannot be 560 underestimated considering that are used for risk assessment. For instance, concentrations of PCM 561 562 similar to those measured in environments worldwide induced ecologically relevant effects on two species belonging to Cnidaria and Mollusca taxa (i.e., Hydra vulgaris and Corbicula fluminea, 563 respectively). Despite these alarming findings, it is important to bearing in mind that chronic 564 toxicity data, in term of sub-lethal effects at individual or sub-individual levels, come from two 565 different approaches, the first one performed on whole animals exposed in vivo and the second one 566

on single cells isolated or collected by animals and then exposed in vitro to NSAIDs. These 567 approaches result in different outcomes, which can lead to different considerations concerning the 568 toxicity of NSAIDs towards aquatic organisms. For instance, EC₅₀ or IC₅₀ from whole animal 569 experiments support the conclusion that NSAIDs in the environment are at much lower 570 concentrations than those causing toxicity, while biochemical or molecular investigations from both 571 in vitro and in vivo studies pointed out a potential hazard of these drugs also at environmentally 572 relevant concentrations. However, some caveats need to be considered, mainly related to the 573 574 uncertainty of the real amount of xenobiotic reaching the cells, the realism of exposures and the lack of cells mechanisms owned by cells to mitigate the effects of xenobiotics, as instead the whole 575 576 organism have, that might result in a potential overestimation of the effect. In addition, although NSAIDs-induced molecular or biochemical effects were induced in whole body or tissues/organs 577 isolated after *in vivo* exposures, they are early, and often specific, responses that the organism 578 579 activate as a consequence of an exposure to these drugs. To date, there is a dearth of information concerning the effects at cellular and tissue levels induced by NSAIDs in freshwater species, 580 581 including invertebrates, as well as on the propagation of the effects from the lowest levels of the biological organization to the highest ones. Moreover, sub-lethal effects highlighted by short- and 582 mid-term exposures might be also more worrisome considering that in natural ecosystems, 583 584 invertebrates are exposed to measurable NSAID concentrations for their whole lifespan. For all the reasons mentioned above, the assessment of the risk of NSAIDs towards aquatic species comparing 585 acute or sub-lethal effects only could not be accurate. Thus, an approach using measured 586 environmental concentrations (MECs) combined with predicted no effect concentrations (PNECs) 587 was proposed by European commission (2003) to screen compounds with potential environmental 588 risks and it was recently refined to properly identify the priority pollutants that should be regularly 589 590 monitored in surface waters (Zhou et al., 2019). According to this approach, DCF and IBU were prioritized as high risk pharmaceuticals, PCM as moderate risk one (Zhou et al., 2019; Palma et al., 591

2020), while ASA as a negligible risk compound (Gómez-Canela et al., 2019). To date, the 592 environmental risk of NPX and NSAID mixture was not assessed. This approach is crucial 593 considering that the increasing production and use of NSAIDs worldwide might result in a notable 594 increase in their environmental levels occurring in aquatic ecosystems, with a consequent 595 enhancement of the risk related to the exposure to these pharmaceuticals towards non-target, 596 freshwater invertebrates. For these reasons, further studies should be needed to enlarge the 597 knowledge on NSAID toxicity towards aquatic organisms, not only invertebrates but also 598 599 vertebrates, considering long-term exposures and the use of alternative and innovative assays to shed light on the mechanism(s) of action of these pharmaceutical compounds. Moreover, as the 600 most of the studies on NSAID toxicity were performed on native, parental compounds, exploring 601 the toxicity of NSAID metabolites, which might results also more toxic than the parental ones (e.g., 602 Marques et al., 2004b; Isidori et al., 2005), should contribute to understand the hazard of these 603 604 drugs. Lastly, considering that NSAIDs occur in aquatic ecosystems in complex 'cocktails' and few previous studies demonstrated that NSAID mixtures induced higher effects than the single 605 606 compounds, further studies aimed at investigating the toxicity of binary or complex NSAID mixtures should be a priority to shed light on adverse effects, mechanism(s) of action and ecological 607 risk of these therapeutics towards freshwater communities. 608

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614 **7. References**

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Figure and Table captions

Figure 1: Studies investigating acute (panel a) or chronic (panel b) toxicity of acetylsalicylic acid
(ASA), paracetamol (PCM), diclofenac (DCF), ibuprofen (IBU), naproxen (NPX) and NSAID mixture
(Mixture) towards different freshwater invertebrates.

Table 1: main physico-chemical properties of the investigated Non-steroidal anti-inflammatorydrugs (NSAIDs).

Table 2: List of studies investigating the adverse effects induced by acetylsalicylic acid (ASA) exposure
towards freshwater invertebrates. Test species, range of ASA concentrations used, duration of the exposure
and investigated endpoints are reported.

Table 3: List of studies investigating the adverse effects induced by paracetamol (PCM) exposure towards
freshwater invertebrates. Test species, range of PCM concentrations used, duration of the exposure and
investigated endpoints are reported.

Table 4: List of studies investigating the adverse effects induced by diclofenac (DCF) exposure towards
freshwater invertebrates. Test species, range of DCF concentrations used, duration of the exposure and
investigated endpoints are reported.

881 Table 5: List of studies investigating the adverse effects induced by ibuprofen (IBU) exposure towards 882 freshwater invertebrates. Test species, range of IBU concentrations used, duration of the exposure and 883 investigated endpoints are reported.

884 Table 6: List of studies investigating the adverse effects induced by Naproxen (NPX) exposure towards 885 freshwater invertebrates. Test species, range of NAP concentrations used, duration of the exposure and 886 investigated endpoints are reported.

Table 7: List of studies investigating the adverse effects induced by the exposure to complex mixtures of
NSAIDs towards freshwater invertebrates. Test species, range of concentrations of NSAIDs included in the
mixture, duration of the exposure and investigated endpoints are reported.