

1 Disruption of the prostaglandin metabolome and  
2 characterization of the pharmaceutical exposome in fish  
3 exposed to wastewater treatment works effluent as revealed  
4 by nanoflow-nanospray mass spectrometry-based  
5 metabolomics

6 *Arthur David,<sup>†</sup> Anke Lange,<sup>‡</sup> Alaa Abdul-Sada,<sup>†</sup> Charles R. Tyler,<sup>‡</sup> and Elizabeth M. Hill<sup>†\*</sup>*

7 <sup>†</sup>School of Life Sciences. University of Sussex. Brighton. U.K. BN1 9QG.

8 <sup>‡</sup>Biosciences, College of Life & Environmental Sciences. University of Exeter, Exeter. U.K.

9 EX4 4QD.

10 \*To whom correspondence should be addressed:

11 Tel: +44 1273 678382

12 Fax: 44 1273 877586

13 email: [E.M.Hill@sussex.ac.uk](mailto:E.M.Hill@sussex.ac.uk)

14

15

16

17

18 **Abstract.**

19 Fish can be exposed to a complex mixture of chemical contaminants, including  
20 pharmaceuticals, present in discharges of wastewater treatment works (WwTWs) effluents.  
21 There is little information on the effects of effluent exposure on fish metabolism, especially  
22 the small molecule signaling compounds which are the biological target of many  
23 pharmaceuticals. We applied a newly developed sensitive nanoflow-nanospray mass  
24 spectrometry non-targeted profiling technique to identify changes in the exposome and  
25 metabolome of roach (*Rutilus rutilus*) exposed to a final WwTWs effluent for 15 days. Effluent  
26 exposure resulted in widespread reduction (between 50% and 90%) in prostaglandin (PG)  
27 profiles in fish tissues and plasma with disruptions also in tryptophan/ serotonin, bile acid and  
28 lipid metabolism. Metabolite disruptions were not explained by altered expression of genes  
29 associated with the PG or tryptophan metabolism. Of the 31 pharmaceutical metabolites that  
30 were detected in the effluent exposome of fish, 6 were non-steroidal anti-inflammatory drugs  
31 but with plasma concentrations too low to disrupt PG biosynthesis. PGs, bile acids and  
32 tryptophan metabolites are important mediators regulating a diverse array of physiological  
33 systems in fish and the identity of wastewater contaminants disrupting their metabolism  
34 warrants further investigation on their exposure effects on fish health.

35

## 36 1. Introduction

37 Fish inhabiting lowland rivers are often exposed to reductions in water quality caused  
38 by discharges of effluents from wastewater treatment works (WWTWs). Treated wastewaters  
39 contain a wide range of biologically active man-made chemicals including pharmaceuticals,  
40 personal care products and pesticides, which maybe incompletely removed during influent  
41 treatment resulting in their temporally varying concentrations in the receiving waters.<sup>1</sup> Many  
42 studies have reported the presence of numerous pharmaceuticals from different therapeutic  
43 classes in final effluents and surface waters including non-steroidal anti-inflammatory drugs  
44 (NSAIDs), selective serotonin reuptake inhibitors (SSRIs), antipsychotics, statins and fibrates,  
45 beta blockers, and anticoagulants.<sup>1-3</sup> Pharmaceuticals are designed to interact with high  
46 affinity with specific biological targets (e.g. receptors, enzymes) in humans and livestock (for  
47 veterinary pharmaceuticals) and these targets can be evolutionary conserved and functional  
48 in fish and other vertebrates,<sup>4</sup> which may lead to unwanted pharmacological effects in non-  
49 target species. A challenge for the environmental community is to prioritize chemicals for risk  
50 assessment and monitoring in order to improve the quality of surface and ground waters.

51 The read-across hypothesis assumes that internal drug concentrations in blood plasma  
52 of wildlife which are similar to human therapeutic plasma concentrations (HtPC) may result  
53 in expected 'therapeutic' effects in non-target organisms.<sup>5</sup> For example, exposure to some  
54 anti-depressant SSRIs affect behavioural responses in fish at plasma concentrations similar to  
55 the HtPC.<sup>6,7</sup> However, because fish populations can be exposed to complex mixtures of  
56 biologically active chemicals,<sup>8-10</sup> identifying associations between exposure to their mixtures  
57 and health outcomes is extremely difficult.<sup>11</sup> To date, most analytical methods monitoring  
58 exposure of fish to effluents have used a targeted approach based on selection of key

59 contaminants and metabolite pathways and are thus limited in their assessments of the risk  
60 of exposure to chemical mixtures.<sup>e.g 8-10</sup> An untargeted analytical approach based on liquid  
61 chromatography high resolution mass spectrometry (LC-MS) combined with chemometric  
62 pattern recognition can identify chemical mixtures accumulating in fish (i.e., the exposome)  
63 and simultaneously investigate for any associated disruption of metabolite profiles (the  
64 metabolome). Using this approach, previous studies have demonstrated that exposure to  
65 contaminants in wastewater effluents results in disruptions to lipid, steroid and bile acid  
66 metabolism and uptake of a variety of household and personal care products in fish blood  
67 plasma.<sup>e.g 12-14</sup> However, until now metabolite profiling methods have not been able to detect  
68 trace levels of contaminants, endocrine or other signaling compounds present in tissues or  
69 plasma extracts. Recently, we have demonstrated that the detection of less abundant  
70 components in samples can be improved by using highly sensitive nanoflow/nanospray LC-  
71 MS combined with appropriate sample preparation methods.<sup>15-17</sup> An additional advantage of  
72 the method is that the chemical exposome, including pharmaceuticals, can also be profiled  
73 alongside the metabolome in the same sample. Hence, an untargeted analytical approach,  
74 based on nanoflow ultraperformance liquid chromatography-nanoelectrospray ionization-  
75 time-of-flight mass spectrometry (nUPLC-nESI-TOFMS) offers a very promising tool for  
76 advancing ecotoxicological investigations in fish exposed to the complex mixtures of  
77 contaminants present in WwTWs effluents.

78         The aim of this study was to investigate the effects of exposure to a final treated  
79 wastewater effluent and the resultant uptake of a complex chemical exposome on the tissue  
80 and plasma metabolome of fish. Extracts of blood plasma and tissues from effluent-exposed  
81 and control fish were profiled using our newly developed nUPLC-nESI-TOFMS method in order

82 to reach unprecedented levels of sensitivity and enable the detection of signaling compounds  
83 and contaminants present at very low concentrations. Our work revealed, for the first time,  
84 widespread disruption of signaling metabolites including prostaglandins (PGs), in tissues of  
85 effluent-exposed fish which could not be explained solely by the components of the  
86 pharmaceutical exposome known to target PG biosynthesis. As PGs are important signaling  
87 agents mediating a number of fish physiological processes, this study highlights the  
88 importance of understanding the mixture effects of contaminants present in wastewaters on  
89 fish health.

## 90 **2. Materials and Methods**

91 Sources of chemicals and standards are given in the Supporting Information.

### 92 2.1 Effluent characteristics

93 The WwTW chosen for this study received 95% influent from a domestic source  
94 (population equivalent of 117,574) and 5% input from industrial wastewaters. The influent  
95 was treated by fine screens, chemically assisted settlement, biological aerated flooded filter  
96 processes and UV disinfection. During the exposure period, the pH of the effluent was 7.3,  
97 concentrations of suspended solids 21 mg/l, biochemical oxygen demand 11 mg/l, chemical  
98 oxygen demand 67 mg/l and total ammonia 29 mg/L (South West Water data).

### 99 2.2 Fish exposure

100 A mixed-sex population of sexually mature roach [*Rutilus rutilus*, age 2+, length of 14.5  
101  $\pm$  1.3 cm and weight 45.4  $\pm$  12.1 g (mean  $\pm$  SEM)] was exposed to either a final treated WwTWs  
102 effluent or clean water (control). Roach were exposed in triplicate tanks (20 fish per 200 L

103 tank) containing either dechlorinated water or 100% effluent for 15 days. The flow rate was  
104 10 L/min and the tanks were continually aerated with an average temperature of 12 °C ± 1  
105 and a light dark cycle of 15h:9h. During the first 7 days of exposure, mortality occurred  
106 amongst 8 out of 60 of effluent-exposed fish, while no mortalities occurred in the control  
107 group. At the end of the experimental period, fish were terminated using an overdose of  
108 phenoxyethanol according to UK Home Office regulations. Blood was collected from the  
109 caudal vein using heparinized needles and samples were centrifuged (5 min, 10,000 RCF) and  
110 the plasma supernatant collected. Gonads, kidney (posterior), gill and liver tissues were  
111 dissected out and immediately snap frozen in liquid nitrogen. All samples were stored at -70  
112 °C until analysis.

113 Wastewater effluent samples (2 x 2.5 litres) were collected in solvent washed, acid  
114 rinsed amber glass bottles on days 0, 7, 15 of the exposure period. Samples were stored at 4  
115 °C with the addition of acetic acid (1%) and methanol (5%) and were extracted within 12 h  
116 after collection.

### 117 2.3 Sample preparation for nUHPLC-nESI-TOFMS profiling

118 Plasma samples were analyzed from up to 5 fish per replicate tank and in total 30  
119 females and 25 males were analyzed (males were present in lower proportion than females  
120 in the mixed sex population). Plasma samples (85-120 µL) were prepared as described in David  
121 et al.<sup>15</sup> Previous studies have shown that this method, using phospholipid removal followed by  
122 mixed-mode cation exchange plates, extracts non polar, cationic and anionic metabolites  
123 (e.g., amino acids, neurotransmitters, bile acids, lipids, eicosanoids and steroids) as well as  
124 contaminants such as pharmaceuticals from plasma samples. Briefly, methanol with 1%

125 formic acid (FA) was added to each sample to a final proportion of 80/20 methanol/plasma  
126 (v/v). An internal standard (IS) mixture containing a mix of seven compounds (venlafaxine-d6,  
127 propranolol-d7, carbamazepine-d10 and fluoxetine-d5 for the +ESI mode and prostaglandin  
128 E2-d4, diclofenac-d4 and triclosan-C13 for the –ESI mode) was added to the plasma extracts.  
129 The amount of IS added were adjusted to give a final concentration of 5 ng IS/120 µL of  
130 plasma. Full details of plasma extraction using Phree and SPE plates are given in the  
131 Supporting Information.

132 Testes (0.28-0.47 g), kidney (0.09-0.17 g), gill (0.28-0.47 g) and liver (0.09-0.20 g),  
133 tissues were extracted in glass tubes (5 mL) using three volumes of methanol and a Microson  
134 XL2000 ultrasonic probe (Misonix Farmingdale) (15W x 30s).<sup>18</sup> Before extraction, the mix of  
135 seven IS was added to each tissue extract to give a final concentration of 5 ng IS to 0.4 g of  
136 testes, to 0.15 g of kidney or gill and to 0.35 g of liver. After extraction, samples were,  
137 centrifuged (13,000 RCF for 20 min) and supernatants collected. The samples were diluted to  
138 20% water (HPLC grade) and extracted using the Phree and SPE plates (see Supporting  
139 Information).

#### 140 2.4 Extraction of wastewater samples for nUHPLC-nESI-TOFMS analysis of NSAIDs.

141 Final wastewater effluent samples (1 L) were spiked with the IS mix, and filtered  
142 through glass wool and filter paper Whatman No1 (Whatman, Maidstone, UK). Samples were  
143 extracted through an Oasis HLB (20 mL, 1 g) cartridge, which was pre-conditioned with 10 mL  
144 of methanol and 10 mL HPLC water. The cartridge was washed with 20 mL of distilled water  
145 and dried under vacuum. Analytes were eluted with 10 mL of methanol and 10 mL of ethyl

146 acetate. Extracts were dried and reconstituted in 1 mL of methanol and 300 mL of the extract  
147 evaporated and purified using the Strata-X-C SPE protocol as described for tissue samples.

## 148 2.5 nUHPLC-nESI-TOFMS

149 The methods used for the nUHPLC-nESI-TOFMS profiling of small molecules have been  
150 established in our laboratory.<sup>15, 17</sup> The nUHPLC-nESI-TOFMS injections were performed on a  
151 Waters nanoAcquity UPLC linked to a Waters Xevo G2 TOFMS, equipped with a nESI source  
152 (Waters, Manchester, U.K) (see Supporting Information for analytical details).

## 153 2.6. Quality control

154 For the metabolomics analysis, two work-up samples (i.e. extraction without sample)  
155 per analytical batch were prepared to monitor for background contaminants. Quality control  
156 (QC) samples comprising a composite sample of the control extracts (diluted two fold) were  
157 prepared in order to monitor for nUHPLC-nESI-TOFMS repeatability and sensitivity during  
158 analysis of a sample run. Each run commenced with the injection of the blank work-up  
159 samples followed by injection of 3 QC samples to equilibrate the source. The samples were  
160 injected randomly with QC samples analysed after every 6 samples. In total, 151 samples  
161 (excluding QCs) were injected in both +ESI and -ESI MS modes.

## 162 2.7 Chemometric analyses

163 Mass spectra were collected in full scan mode (50 to 1000 m/z) and spectral peaks  
164 were deconvoluted and aligned using TransOmics algorithm (Waters, Manchester, UK). Peak  
165 picking was performed creating an aggregate dataset from all sample files in which the MS  
166 features were binned into retention time (Rt) x m/z values. Datasets were exported to SIMCA



167 software (Umetrics, Crewe, UK) for multivariate analysis (principal component analysis, PCA)  
168 in order to examine class separating differences between control and exposed groups. Data  
169 sets were log transformed and pareto scaled, and models were developed using the database  
170 of  $n$  (from 10 to 15) different fish per class. PCAs were performed in order to observe  
171 clustering between treatments, and models were evaluated through an examination of  
172 explained variation (R2X) and predictive ability (Q2X) parameters. In order to determine  
173 which contaminants and metabolites influence discrimination between two classes, the  
174 datasets were modelled using orthogonal partial least squares to latent structures (OPLS)  
175 method which were assessed by the explained variation (R2Y) and predictive ability (Q2Y). Rt  
176 x m/z signals pertaining to discriminatory contaminants or metabolites were detected using  
177 an 'S' plot of the OPLS model. e.g. 13, 18, 19 Signals associated with the presence of chemical  
178 contaminants arising from effluent exposure were identified as only present in effluent-  
179 exposed fish and not detected in any replicates from control fish or from work-up blank  
180 samples. Signals corresponding to metabolite markers were either decreased in control fish  
181 or increased in effluent-exposed fish but also present in control fish but not in work-up blanks.  
182 The signal intensity of suspected metabolite markers of effluent exposure were normalized  
183 to the area of the closest IS. Analysis, using the Kruskal-Wallis test, of replicate fish from each  
184 of the triplicate tanks from the two exposure treatments revealed no significant intertank  
185 variability for any metabolite marker. Hence, metabolites from fish from replicate tanks were  
186 analyzed as one treatment for each exposure by non-parametric Mann-Whitney U-test (SPSS,  
187 IBM) with a correction for a false discovery rate of 5%.<sup>20</sup>

188           The putative identities of contaminants and metabolites were determined from  
189 screening the m/z of molecular ion using Metabosearch

190 (<http://omics.georgetown.edu/metabosearch.html>) which searched databases of the Human  
191 Metabolome Database (<http://www.hmdb.ca/>), PubChem  
192 (<http://pubchem.ncbi.nlm.nih.gov/>), METLIN (<http://metlin.scripps.edu/>) and LipidMaps  
193 (<http://www.lipidmaps.org/>). Chemical identity was confirmed from accurate mass, isotopic  
194 fit and fragmentation data obtained from high energy collisional-induced dissociation and  
195 from comparison in-house with standard compounds or with open source databases.

## 196 2.8 Targeted analysis of NSAIDs by nUHPLC-nESI-TOFMS

197 NSAIDs were quantified in the profiling datasets of tissues and plasma and also after  
198 nUHPLC-nESI-TOFMS analysis of wastewater extracts (see Supporting Information for details)  
199 Method detection and method quantification limits (MDLs, MQLs) are given in Tables S1 and  
200 S5.

## 201 2.9 Gene expression analysis

202 Quantitative real-time RT-PCR (RT-QPCR) was performed on male and female gill, liver  
203 and gonad tissue samples to quantify the expression profiles of transcripts encoding enzymes  
204 for which related metabolites were found to change in the prostaglandin and serotonin  
205 pathways. The targets were prostaglandin-endoperoxide synthase 1 (*ptgs1*), prostaglandin-  
206 endoperoxide synthase 2b (*ptgs2b*), prostaglandin E synthase (*ptges*) and cytosolic  
207 phospholipase A2 (*pla2g4*) in the prostaglandin pathway and monoamine oxidase (*mao*),  
208 dopa decarboxylase (aromatic amino acid decarboxylase, *ddc*), aldehyde oxidase 1 (*aox1*),  
209 tryptophan 2,3-dioxygenase a (*tdo2a*) and tryptophan 2,3-dioxygenase b (*tdo2b*) in the  
210 serotonin pathway. Details on RNA analyses, primer sequences, sizes of PCR products and  
211 PCR assay conditions are provided in Supporting Information and Table S2. Samples from 3

212 fish per tank were analysed and the Chauvenet's criterion<sup>21</sup> was applied to detect any outliers  
213 in the transcript expression data. Identified outliers were removed from the data set prior to  
214 statistical analyses. Data were analyzed using a t-test comparing transcript expression  
215 between all control and exposed fish and results of all tests were accepted as significant at  $p$   
216  $< 0.05$  (IBM SPSS Statistics for Windows, Version 21.0.0).

217

### 218 **3. Results and Discussion**

#### 219 **3.1 nUHPLC-nESI-TOFMS profiling of roach plasma and tissues**

220 Extracts of blood plasma from male and female roach exposed to either a treated  
221 WwTWs effluent or to clean water were profiled by nUPLC-nESI-TOFMS in both +ESI and -ESI  
222 modes. PCA modeling of the datasets revealed clear separation between control and effluent-  
223 exposed roach of both sexes on the first component of the scores plots (-ESI mode Figure 1,  
224 +ESI mode, Figure S1). In these plots, the QC samples clustered together indicating that the  
225 profiling methods were highly repeatable with little analytical variability. The datasets were  
226 further analyzed by OPLS modeling which revealed highly predictable models for control and  
227 exposed datasets from either male or female roach ( $R^2Y$  of  $>0.99$  and  $Q^2Y > 0.69$ ). No  
228 separation according to sex was observed in effluent-exposed roach in either the PCA or OPLS  
229 models indicating that the response (in terms of plasma chemical profiles) to effluent  
230 exposure was similar in female and male fish. Therefore, female tissue samples (gills, liver  
231 and the posterior kidney which contains both renal cells and macrophages) were analyzed as  
232 they were available in greater numbers than males. Extracts of testes tissue from males were  
233 also profiled because it is known that PGs play important roles in sexual differentiation<sup>22</sup> and

234 the association between exposure to WwTWs effluents and feminization of male fish has  
235 been well documented.<sup>e.g.23</sup> The PCA of the datasets of all tissue extracts revealed a clear  
236 separation between samples from control and exposed fish (Figure S2).

### 237 3.2 Metabolite markers of effluent exposure

238 From analysis of the 'S' loading plots of the OPLS models of the plasma datasets, it was  
239 apparent that most of the chemicals driving the separation between control and effluent-  
240 exposed fish were contaminants which were only detected in plasma of exposed fish (see  
241 section 3.3 below). However, ten metabolite structures were identified as significant markers  
242 of effluent exposure (Table 1). In agreement with a previous study,<sup>14</sup> exposure to WwTWs  
243 effluent resulted in disrupted metabolism of plasma bile acids and lipids. After effluent  
244 exposure, the plasma concentrations of two conjugated secondary bile acids were increased  
245 by 5-12 fold (female and male fish) for taurodeoxycholic acid, and 39-56 fold for a  
246 tauroolithocholic acid-like structure. No changes in the levels of primary bile acids, such as  
247 taurocholic acid and cyprinol sulfate, were detected. Little is known about the effect of  
248 chemical exposures on disruption of bile acid profiles in fish or mammals, however secondary  
249 bile acids are formed by the metabolism of gut microflora.<sup>24,25</sup> It is possible that the observed  
250 increase in metabolism of secondary bile acids was a result of disruption of the community or  
251 gene expression of resident gut bacteria as a result of exposure to chemical components in  
252 the effluent. Increases in the lipid metabolite sphinganine (9- fold) and its potential precursor,  
253 a palmitoyl serine type metabolite (177-227 fold), were also observed indicating disruption in  
254 sphingolipid signaling.

255 In our study, the use of more sensitive nanoflow/nanospray LC-MS profiling methods  
256 resulted in detection of new metabolites in effluent-exposed fish that were associated with

257 other signaling pathways. These included a 5-11 fold increase in the plasma concentration of  
258 tryptophan and a 2-fold decrease in its metabolite acetylserotonin, indicating effects on the  
259 serotonin pathway (Table 1). 'S' plot analysis of OPLS models of metabolite profiles in tissues  
260 revealed that concentrations of indolepyruvate, a metabolite of tryptophan, were increased  
261 by 13 and 32-fold in female gills and male testes, respectively (Table S3), indicating that  
262 tryptophan metabolism was also altered in these tissues as well as in blood plasma of effluent-  
263 exposed roach. In order to explore the mechanisms responsible for these metabolite changes,  
264 selected gene transcripts associated with tryptophan/serotonin metabolism were analyzed  
265 by qRT-PCR in gills, liver and gonads of male and female fish (Figure S3). Tryptophan can be  
266 metabolized by dioxygenases and a decrease in their expression would result in an increased  
267 substrate for metabolism to acetyl serotonin or indolepyruvate.<sup>26</sup> However, the dioxygenase  
268 (*tdo2a*) transcript was only significantly decreased in male and female liver from exposed fish  
269 and was increased in male gills. Significant increases in the dopa decarboxylase transcript  
270 levels (*ddc*) which metabolizes tryptophan to serotonin, and monamine oxidase expression  
271 (*mao*) which further converts serotonin to hydroxyindole acetic acid,<sup>27</sup> were observed only in  
272 female gills and for *ddc* also in female liver. Taken together, effluent exposure did not result  
273 in a widespread alteration in the transcription of key genes that could explain the changes  
274 involved in tryptophan metabolism in roach tissues.

275 Concentrations of a number of eicosanoid type metabolites were also altered in  
276 plasma samples from effluent-exposed male and female fish. These included a 10-15-fold  
277 reduction in prostaglandin F2 $\alpha$  (PGF2 $\alpha$ ) levels, a 5-fold decrease in a tetranor PGE1-like  
278 structure, and a 13-16- fold increase in an oxygenated C20 metabolite (dicarboxy leukotriene  
279 B4-like) (Table 1). Examination of the 'S' plots from OPLS models of female gills, liver, kidney

280 and male testes revealed numerous signals, mainly corresponding to prostaglandin  
281 structures, which were prevalent (depressed) markers of effluent exposure (Table S3). PGE<sub>2</sub>  
282 and PGF<sub>2</sub>α levels were reduced by 52-90% in all the tissues analyzed, together with a mean  
283 93% reduction observed for PGF<sub>2</sub>α in plasma (Figure 2). Reductions in levels of a further 7  
284 different prostaglandin metabolites were detected in various body tissues (Table S3) and  
285 further work is needed to determine their structural identity.

286         The widespread reduction of prostaglandin structures in tissues and in blood plasma  
287 indicates a general inhibition of the prostaglandin metabolism in effluent-exposed fish.  
288 Prostaglandins are formed from the action of prostaglandin endoperoxide synthases (PGTS),  
289 also termed cyclooxygenase enzymes, on the C<sub>20</sub> polyunsaturated fatty acid, arachidonic  
290 acid, which in turn is released from membrane phospholipids by the action of cytosolic  
291 phospholipase A<sub>2</sub> (PLA<sub>2</sub>). At least two PTGS genes (*pgts* 1 and 2) are present in fish and qRT-  
292 PCR analysis of gills, liver and gonads of male and female fish revealed that their expression  
293 was not altered as a result of effluent exposure (Figure S3). Expression of *pla2g4*, associated  
294 with substrate release by PLA<sub>2</sub>, was reduced in liver and gills of males but not in females or  
295 in gonads of either sex. Likewise, *pgtes*, which encodes PGE synthase and conversion of the  
296 PGH<sub>2</sub> intermediate to PGE<sub>2</sub>, was reduced in liver and gills of males but not females, nor in  
297 gonads of either sex. These data indicate that the widespread reduction of PG levels in the  
298 various body tissues were not due to changes in expression of genes associated with their  
299 biosynthesis, but rather may arise from direct inhibition of enzyme activity instead.

300         The functional role and the identities of the full complement of biologically active  
301 prostaglandin structures has not been not fully determined in fish systems. However, PGE<sub>2</sub>  
302 regulates ion transport in fish gills and kidney<sup>28</sup>, macrophage differentiation in the posterior

303 kidney, and is an important inflammatory and blood clotting mediator produced by leucocytes  
304 and thrombocytes.<sup>29</sup> PGF2 $\alpha$  is can regulate reproductive behavior, synchronous spawning,  
305 and ovulation in females.<sup>30</sup> PGs are also important signaling agents during cell development  
306 and differentiation.<sup>31</sup> This indicates that widespread reduction of PG levels may have  
307 deleterious effects affecting numerous endpoints in fish including development,  
308 reproduction, the immune response and osmoregulation.

309         The role of other metabolites identified in our study has yet to be fully understood in  
310 fish physiology. However, there remains the possibility that the roles of serotonin, bile acid  
311 and PG metabolites maybe linked, or act in concert, to affect health in fish. For instance, a  
312 follow-up study of brain tissue sampled from fish used in this study revealed significant  
313 increases of serotonin concentrations in selected brain regions of the effluent-exposed fish  
314 (A David, pers comm). Serotonin regulates aggression<sup>32</sup> and PGF2 $\alpha$  mediates courtship in  
315 male fish,<sup>33</sup> which indicates that further investigations on the effects of wastewater exposures  
316 on disruption of fish behaviour are warranted. Similarly, serotonin and PG metabolites are  
317 both signalling agents in the hypothalamus-pituitary-gonad axis of fish and regulate gonadal  
318 development.<sup>22,34</sup> They are also both important modulators of the immune and inflammatory  
319 response of vertebrates,<sup>29, 35</sup> and both regulate epithelial cell function in gills.<sup>28,36</sup> In  
320 mammals, some secondary bile acids, including taurodeoxycholic acid, can act as anti-  
321 secretory agents in the colon,<sup>37</sup> whereas both serotonin and prostaglandin metabolites  
322 promote epithelial mucosal secretion and gastrointestinal protection<sup>38,39</sup> raising the question  
323 as to whether interactions occur between these different metabolites affecting  
324 gastrointestinal function in fish.

325

### 326 3.3 The link between the exposome and metabolome of effluent-exposed fish

327 Interrogation of the 'S' loading plots of signals unique to effluent-exposed fish  
328 revealed the accumulation of a complex mixture of contaminants in blood plasma and tissues.  
329 These contained 31 pharmaceuticals from 11 different classes including NSAIDs (naproxen,  
330 mefenamic acid, diclofenac, ibuprofen, felbinac, celecoxib) and SSRIs (paroxetine, citalopram,  
331 sertraline, fluoxetine, norfluoxetine, nortriptyline and nortriptyline) (Figure 3 and Table S4).  
332 In addition, a wide range of endocrine disruptors, personal care products, pesticides,  
333 antibacterial and human dietary products were also detected in the plasma and tissues of  
334 effluent-exposed but not control fish (Table S4).

335 Due to matrix interferences, the concentrations of SSRIs in the different sample types  
336 could not be determined in this study and their quantitative analysis required further method  
337 development. However, the concentrations of NSAIDs, which are potent inhibitors of PGTS  
338 enzymes in vertebrate systems, were determined in effluent and fish samples. During the  
339 exposure period, NSAIDs levels in the effluent were all below 0.5 µg/l, except for naproxen  
340 which was present at a mean ± SD of 3.7 ± 2.3 µg/l (Tables 2 and S5). Studies to date indicate  
341 that these concentrations of NSAIDs were too low to affect PG or PGTS gene expression levels  
342 in fish.<sup>40-42</sup> In addition, NSAID concentrations in plasma and tissues of effluent-exposed fish  
343 were determined and compared with reported human therapeutic plasma concentrations,  
344 HtPC<sup>43</sup> (Table 2). NSAIDs were detected in plasma and most of the tissues analyzed, but with  
345 the exception of naproxen and ibuprofen, their mean concentrations were < 1ng/ml plasma  
346 or 1 ng/g tissue. Mean concentrations of naproxen and ibuprofen in the plasma were higher  
347 than in the effluent indicating bioconcentration of these contaminants in the fish. The effluent  
348 concentrations of naproxen were above the MQL, which allowed the determination of a  
349 plasma bioconcentration factor of 4, similar to that already reported for this compound.<sup>44</sup>



350 However, the plasma concentrations of all NSAIDs were between 650 and 14,000-fold below  
351 the HtPC. As PG levels were significantly reduced in blood plasma and body tissues, this would  
352 indicate that either NSAIDs and their mixtures were more potent in fish compared with  
353 humans, or that other contaminants were present that inhibited PG biosynthesis in the fish.  
354 Previous reports suggest that many other compounds can inhibit PGTS enzyme activity,  
355 including pesticides and some of the endocrine disrupting chemicals detected in fish in this  
356 study (Table S4).<sup>45,46</sup> Further work is needed to identify other PGTS inhibitors amongst the  
357 diverse range of contaminants which bioconcentrate in fish exposed to WwTW effluents.

### 358 3.4 Implications of the work and future studies.

359 This study was conducted with undiluted effluent, which reflects the exposures of  
360 aquatic animals, including fish, in some countries where there is little or no dilution of the  
361 discharges into the receiving waters.<sup>47</sup> However, there was a mortality of 13% during effluent  
362 exposure indicating that fish were subjected to high physiological stress. Further studies are  
363 required to determine whether similar metabolite disruptions are observed with diluted  
364 effluents that typify ambient waters.

365 Through the application of newly developed sensitive non-targeted chemical profiling  
366 techniques, a novel finding emerging is the widespread reduction in PG metabolism in the  
367 roach exposed to a final treated WwTWs effluent. This finding could not have been predicted  
368 from an analysis of pharmaceuticals in the effluent which are designed to target PG  
369 metabolism, and there is a need to identify the key causative contaminants disrupting PG  
370 biosynthesis in WwTW effluents. Moreover, we highlight the need for further understanding  
371 on the identity and function of the numerous PG metabolites that were detected in fish  
372 tissues. Furthermore, PGs, serotonin and bile acid metabolites are components of key

373 signaling pathways which mediate gastrointestinal function, ion transport, immune and  
374 reproductive processes in vertebrates, and disruption in their levels may potentially impact  
375 on a range of health endpoints in contaminated fish.

376 Our study further emphasizes the importance of understanding mixture effects of  
377 contaminants (including their metabolites) on the health of wildlife. Improvements in  
378 discovery-based chemical profiling technologies are likely to increase coverage of the  
379 exposome and metabolome, and advance our understanding of the link between exposure to  
380 complex mixtures of contaminants and their impact on multiple metabolite pathways.  
381 However, it will require a step change in knowledge to understand the link between  
382 disruption of multiple interacting pathways and their consequent effects on wildlife or human  
383 health. Without this knowledge, environmental policymakers may have to consider whether  
384 regulation of chemical contaminants should be proactive and informed by changes in  
385 molecular profiles with the *potential* for physiological disruption, rather than a reliance on  
386 demonstrated health endpoints which may take many decades to investigate, particularly in  
387 respect to exposure of chemical mixtures.

### 388 **Supporting Information**

389 Supporting information comprises further analytical details, PCA score plots of metabolomics  
390 datasets and tables of identification of markers of effluent exposure. This material is available  
391 free of charge via the Internet at <http://pubs.acs.org>.

### 392 **Acknowledgments**

393 This research was supported by a Marie Curie Intra European Fellowship within the European  
394 Community Seventh Framework Programme (FP7/2007-2013, grant agreement number

395 302097). We gratefully acknowledge Jan Shears, Victoria Jennings, the fish team at Exeter  
396 University and Andrew Chetwynd (Sussex) for help with the effluent exposure.

397

## 398 References

399

400 1. Li, W. C., Occurrence, sources, and fate of pharmaceuticals in aquatic environment and soil.  
401 *Environ. Pollut.* 2014, 187, 193-201.

402 2. Kostich, M. S.; Batt, A. L.; Lazorchak, J. M., Concentrations of prioritized pharmaceuticals in  
403 effluents from 50 large wastewater treatment plants in the US and implications for risk  
404 estimation. *Environ. Pollut.* **2014**, 184, 354-359.

405 3. der Beek, T. A.; Weber, F. A.; Bergmann, A.; Hickmann, S.; Ebert, I.; Hein, A.; Kuster, A.,  
406 Pharmaceuticals in the environment-global occurrences and perspectives. *Environ. Toxicol.*  
407 *Chem.* **2016**, 35, (4), 823-835.

408 4. Gunnarsson, L.; Jauhiainen, A.; Kristiansson, E.; Nerman, O.; Larsson, D. G. J., Evolutionary  
409 conservation of human drug targets in organisms used for environmental risk assessments.  
410 *Environ Sci Technol.* **2008**, 42, (15), 5807-5813.

411 5. Rand-Weaver, M.; Margiotta-Casaluci, L.; Patel, A.; Panter, G. H.; Owen, S. F.; Sumpter, J.  
412 P., The Read-Across Hypothesis and Environmental Risk Assessment of Pharmaceuticals.  
413 *Environ Sci Technol.* **2013**, 47, (20), 11384-11395.

414 6. Valenti, T. W.; Gould, G. G.; Berninger, J. P.; Connors, K. A.; Keele, N. B.; Prosser, K. N.;  
415 Brooks, B. W., Human Therapeutic Plasma Levels of the Selective Serotonin Reuptake  
416 Inhibitor (SSRI) Sertraline Decrease Serotonin Reuptake Transporter Binding and Shelter-  
417 Seeking Behavior in Adult Male Fathead Minnows. *Environ. Sci. Technol.* **2012**, 46, (4), 2427-  
418 2435.

419 7. Margiotta-Casaluci, L.; Owen, S. F.; Cumming, R. I.; de Polo, A.; Winter, M. J.; Panter, G. H.;  
420 Rand-Weaver, M.; Sumpter, J. P., Quantitative Cross-Species Extrapolation between Humans  
421 and Fish: The Case of the Anti-Depressant Fluoxetine. *PLoS One.* **2014**, 9, (10).

422 8. Ramirez, A. J.; Brain, R. A.; Usenko, S.; Mottaleb, M. A.; O'Donnell, J. G.; Stahl, L. L.; Wathen,  
423 J. B.; Snyder, B. D.; Pitt, J. L.; Perez-Hurtado, P.; Dobbins, L. L.; Brooks, B. W.; Chambliss, C. K.,  
424 Occurrence of pharmaceuticals and personal care products in fish: results of a national pilot  
425 study in the United States. *Environ Toxicol Chem.* **2009**, 28, (12), 2587-2597.

426 9. Meador, J. P.; Yeh, A.; Young, G.; Gallagher, E. P., Contaminants of emerging concern in a  
427 large temperate estuary. *Environ Pollut.* **2016**, 213, 254-267.

428 10. Tanoue, R.; Nomiya, K.; Nakamura, H.; Kim, J. W.; Isobe, T.; Shinohara, R.; Kunisue, T.;  
429 Tanabe, S., Uptake and Tissue Distribution of Pharmaceuticals and Personal Care Products in  
430 Wild Fish from Treated-Wastewater-Impacted Streams. *Environ Sci Technol.* **2015**, 49, (19),  
431 11649-11658.

432 11. Hamilton, P. B.; Cowx, I. G.; Oleksiak, M. F.; Griffiths, A. M.; Grahn, M.; Stevens, J. R.;  
433 Carvalho, G. R.; Nicol, E.; Tyler, C. R., Population-level consequences for wild fish exposed to  
434 sublethal concentrations of chemicals – a critical review. *Fish Fish.* **2015**, 17, (3), 545-566

435 12. Skelton, D. M.; Ekman, D. R.; Martinovic-Weigelt, D.; Ankley, G. T.; Villeneuve, D. L.; Teng,  
436 Q.; Collette, T. W., Metabolomics for in situ environmental monitoring of surface waters  
437 impacted by contaminants from both point and nonpoint sources. *Environ. Sci. Technol.* **2014**,  
438 48, (4), 2395-2403.

- 439 13. Alvarez-Munoz, D.; Al-Salhi, R.; Abdul-Sada, A.; Gonzalez-Mazo, E.; Hill, E. M., Global  
440 metabolite profiling reveals transformation pathways and novel metabolomic responses in  
441 Solea senegalensis after exposure to a non-ionic surfactant. *Environ. Sci. Technol.* **2014**, *48*,  
442 (9), 5203-5210.
- 443 14. Al-Salhi, R.; Abdul-Sada, A.; Lange, A.; Tyler, C. R.; Hill, E. M., The xenometabolome and  
444 novel contaminant markers in fish exposed to a wastewater treatment works effluent.  
445 *Environ Sci Technol.* **2012**, *46*, (16), 9080-9088.
- 446 15. David, A.; Abdul-Sada, A.; Lange, A.; Tyler, C. R.; Hill, E. M., A new approach for plasma  
447 (xeno)metabolomics based on solid-phase extraction and nanoflow liquid chromatography-  
448 nano-electrospray ionisation mass spectrometry. *J Chromatog.r A.* **2014**, *1365*, 72-85.
- 449 16. Chetwynd, A. J.; Abdul-Sada, A.; Hill, E. M., Solid-phase extraction and nanoflow liquid  
450 chromatography-nano-electrospray ionization mass spectrometry for improved global urine  
451 metabolomics. *Anal. Chem.* **2015**, *87*, (2), 1158-1165.
- 452 17. Chetwynd, A. J.; David, A.; Hill, E. M.; Abdul-Sada, A., Evaluation of analytical performance  
453 and reliability of direct nanoLC-nanoESI-high resolution mass spectrometry for profiling the  
454 (xeno)metabolome. *J Mass Spectrom.* **2014**, *49*, (10), 1063-1069.
- 455 18. Flores-Valverde, A. M.; Hill, E. M., Methodology for profiling the steroid metabolome in  
456 animal tissues using ultraperformance liquid chromatography-electrospray-time-of-flight  
457 mass spectrometry. *Anal. Chem.* **2008**, *80*, (22), 8771-8779.
- 458 19. Wiklund, S.; Johansson, E.; Sjostrom, L.; Mellerowicz, E. J.; Edlund, U.; Shockcor, J. P.;  
459 Gottfries, J.; Moritz, T.; Trygg, J., Visualization of GC/TOF-MS-based metabolomics data for  
460 identification of biochemically interesting compounds using OPLS class models. *Anal. Chem.*  
461 **2008**, *80*, (1), 115-122.
- 462 20. Benjamini, Y.; Hochberg, Y., Controlling the False Discovery Rate - a Practical and Powerful  
463 Approach to Multiple Testing. *J Roy Stat Soc B Met.* **1995**, *57*, (1), 289-300.
- 464 21. Chauvenet, W., Methods of least squares. **1863**, *Reprinted (1960) 5th edition.* Dover,  
465 NY., 469-566.
- 466 22. Pradhan, A.; Olsson, P. E., Juvenile Ovary to Testis Transition in Zebrafish Involves  
467 Inhibition of Ptges. *Biol. Reprod.* **2014**, *91*, (2), 15.
- 468 23. Lange, A.; Paull, G. C.; Hamilton, P. B.; Iguchi, T.; Tyler, C. R., Implications of persistent  
469 exposure to treated wastewater effluent for breeding in wild roach (*Rutilus rutilus*)  
470 populations. *Environ. Sci. Technol.* **2011**, *45*, (4), 1673-1679.
- 471 24. Reyes, H., Phylogenetic diversity of bile acids. *Revista Medica De Chile* **1994**, *122*, (8), 944-  
472 950.
- 473 25. Ridlon, J. M.; Kang, D. J.; Hylemon, P. B., Bile salt biotransformations by human intestinal  
474 bacteria. *Journal of Lipid Research* **2006**, *47*, (2), 241-259.
- 475 26. Cortes, J.; Alvarez, C.; Santana, P.; Torres, E.; Mercado, L., Indoleamine 2,3-dioxygenase:  
476 First evidence of expression in rainbow trout (*Oncorhynchus mykiss*). *Dev. Comp. Immunol.*  
477 **2016**, *65*, 73-78.
- 478 27. Sainio, E. L.; Pulkki, K.; Young, S. N., L-tryptophan: Biochemical, nutritional and  
479 pharmacological aspects. *Amino Acids* **1996**, *10*, (1), 21-47.
- 480 28. Avella, M.; Part, P.; Ehrenfeld, J., Regulation of Cl<sup>-</sup> secretion in seawater fish  
481 (*Dicentrarchus labrax*) gill respiratory cells in primary culture. *J. Physiol.-London.* **1999**, *516*,  
482 (2), 353-363.
- 483 29. Gomez-Abellan, V.; Sepulcre, M. P., The role of prostaglandins in the regulation of fish  
484 immunity. *Mol. Immunol.* **2016**, *69*, 139-145.

- 485 30. Goncalves, D.; Costa, S. S.; Teles, M. C.; Silva, H.; Ingles, M.; Oliveira, R. F., Oestradiol and  
486 prostaglandin F2 alpha regulate sexual displays in females of a sex-role reversed fish. *Proc. R.*  
487 *Soc. B-Biol. Sci.* **2014**, *281*, (1778), 1-9.
- 488 31. Cha, Y. I.; Solnica-Krezel, L.; DuBois, R. N., Fishing for prostanoids: Deciphering the  
489 developmental functions of cyclooxygenase-derived prostaglandins. *Dev. Biol.* **2006**, *289*, (2),  
490 263-272.
- 491 32. Winberg, S.; Thornqvist, P. O., Role of brain serotonin in modulating fish behavior. *Curr.*  
492 *Zool.* **2016**, *62*, (3), 317-323.
- 493 33. Yabuki, Y.; Koide, T.; Miyasaka, N.; Wakisaka, N.; Masuda, M.; Ohkura, M.; Nakai, J.; Tsuge,  
494 K.; Tsuchiya, S.; Sugimoto, Y.; Yoshihara, Y., Olfactory receptor for prostaglandin F-2 alpha  
495 mediates male fish courtship behavior. *Nat. Neurosci.* **2016**, *19*, (7), 897-904.
- 496 34. Prasad, P.; Ogawa, S.; Parhar, I. S., Role of serotonin in fish reproduction. *Front. Neurosci.*  
497 **2015**, *9*, 1-9.
- 498 35. Shajib, M. S.; Khan, W. I., The role of serotonin and its receptors in activation of immune  
499 responses and inflammation. *Acta Physiol.* **2015**, *213*, (3), 561-574.
- 500 36. Shakarchi, K.; Zachar, P. C.; Jonz, M. G., Serotonergic and cholinergic elements of the  
501 hypoxic ventilatory response in developing zebrafish. *J. Exp. Biol.* **2013**, *216*, (5), 869-880.
- 502 37. Keating, N.; Mroz, M. S.; Scharl, M. M.; Marsh, C.; Ferguson, G.; Hofmann, A. F.; Keely, S.  
503 J., Physiological concentrations of bile acids down-regulate agonist induced secretion in  
504 colonic epithelial cells. *J. Cell. Mol. Med.* **2009**, *13*, (8B), 2293-2303.
- 505 38. Fidalgo, S.; Ivanov, D. K.; Wood, S. H., Serotonin: from top to bottom. *Biogerontology* **2013**,  
506 *14*, (1), 21-45.
- 507 39. Dey, I.; Lejeune, M.; Chadee, K., Prostaglandin E-2 receptor distribution and function in  
508 the gastrointestinal tract. *Br. J. Pharmacol.* **2006**, *149*, (6), 611-623.
- 509 40. Bhandari, K.; Venables, B., Ibuprofen bioconcentration and prostaglandin E2 levels in  
510 the bluntnose minnow *Pimephales notatus*. *Comp. Biochem. Physiol. C-Toxicol. Pharmacol*  
511 **2011**, *153*, (2), 251-257.
- 512 41. Morthorst, J. E.; Lister, A.; Bjerregaard, P.; Van der Kraak, G., Ibuprofen reduces zebrafish  
513 PGE(2) levels but steroid hormone levels and reproductive parameters are not affected.  
514 *Comp. Biochem. Physiol. C-Toxicol. Pharmacol.* **2013**, *157*, (2), 251-257.
- 515 42. Mehinto, A. C.; Hill, E. M.; Tyler, C. R., Uptake and Biological Effects of Environmentally  
516 Relevant Concentrations of the Nonsteroidal Anti-inflammatory Pharmaceutical Diclofenac in  
517 Rainbow Trout (*Oncorhynchus mykiss*). *Environ. Sci. Technol.* **2010**, *44*, (6), 2176-2182.
- 518 43. Schulz, M.; Iwersen-Bergmann, S.; Andresen, H.; Schmoldt, A., Therapeutic and toxic blood  
519 concentrations of nearly 1,000 drugs and other xenobiotics. *Crit Care.* **2012**, *16*, (4), R136.
- 520 44. Brown, J. N.; Paxeus, N.; Forlin, L.; Larsson, D. G. J., Variations in bioconcentration of  
521 human pharmaceuticals from sewage effluents into fish blood plasma. *Environ. Toxicol.*  
522 *Pharmacol.* **2007**, *24*, (3), 267-274.
- 523 45. Kugathas, S.; Audouze, K.; Ermler, S.; Orton, F.; Rosivatz, E.; Scholze, M.; Kortenkamp, A.,  
524 Effects of Common Pesticides on Prostaglandin D2 (PGD2) Inhibition in SC5 Mouse Sertoli  
525 Cells, Evidence of Binding at the COX-2 Active Site, and Implications for Endocrine Disruption.  
526 *Environ. Health Perspect.* **2016**, *124*, (4), 452-459.
- 527 46. Kristensen, D. M.; Skalkam, M. L.; Audouze, K.; Lesne, L.; Desdoits-Lethimonier, C.;  
528 Frederiksen, H.; Brunak, S.; Skakkebaek, N. E.; Jegou, B.; Hansen, J. B.; Junker, S.; Leffers, H.,  
529 Many Putative Endocrine Disruptors Inhibit Prostaglandin Synthesis. *Environ. Health Perspect.*  
530 **2011**, *119*, (4), 534-541.

531 47. Keller, V. D. J.; Williams, R. J.; Lofthouse, C.; Johnson, A. C., Worldwide estimation of river  
532 concentrations of any chemical originating from sewage treatment plants using dilution  
533 factors. *Environ. Toxicol. Chem.* **2014**, *33*, (2), 447-452.

534

535

**Table 1. The effect of exposure to wastewater effluent on blood plasma metabolite profiles of roach.**

Retention time	Observed molecular ion	Molecular ion species	$\Delta$ ppm	Formula of ion	Fragment ions	Kegg no	Identity	Fold change	p value	Fold change	P value
<b><i>Metabolites decreased in plasma</i></b>								<b>Female</b>		<b>Male</b>	
6.19	219.1136	M+H	0.9	C12H15N2O2	160.0766	C00978	acetylserotonin <sup>a</sup>	1.6	0.014	2.7	2.8x10 <sup>-4</sup>
13.61	353.2326	M-H	-0.6	C20H33O5	309.2058	C00639	PGF2 $\alpha$ <sup>a</sup>	10.6	3.7x10 <sup>-5</sup>	14.7	2.6x10 <sup>-4</sup>
14.20	297.1702	M-H	0	C16H25O5	nd	-	tetranor PGE1 like	5.2	4.0x10 <sup>-6</sup>	4.8	8.5x10 <sup>-5</sup>
<b><i>Metabolites increased in plasma</i></b>											
15.80	383.2068	M-H	-0.5	C20H31O7	nd	-	dicarboxy LTB4 like	16.6	6.1 x 10 <sup>-6</sup>	13.3	8.1 x 10 <sup>-5</sup>
14.74	498.2889	M-H	0	C26H44NO6S	nd		taurodeoxycholic acid <sup>a</sup>	5.1	1.3x10 <sup>-4</sup>	12.1	1.2x10 <sup>-4</sup>
19.29	482.2939	M-H	-0.2	C26H44NO5S	nd	C05122 C02592	tauroolithocholic acid like	39.3	2.0x10 <sup>-6</sup>	56.0	2.6x10 <sup>-5</sup>
6.01	205.0979	M+H	0.1	C11H13N2O2	188.0713, 146.0606	C00078	tryptophan <sup>a</sup>	5.0	4x10 <sup>-3</sup>	10.7	4.0x10 <sup>-4</sup>
4.27	204.0662	M+H	0.5	C11H10NO3	158.0605, 130.0658	C00331	Indolepyruvate <sup>a</sup>	22.9	2.1x10 <sup>-5</sup>	77.6	5.6x10 <sup>-5</sup>
15.59	302.3060	M+H	0.3	C18H40NO2	284.2952	C00836	sphinganine	9.1	6.4 x 10 <sup>-5</sup>	9.4	5.2 x 10 <sup>-5</sup>
25.19	352.2828	M+Na	0	C19H39NO3Na	312.2898		palmitoyl serine like	227	4.5x10 <sup>-5</sup>	177	3.0x10 <sup>-4</sup>

<sup>a</sup> metabolite identity confirmed by a pure standard. Other metabolite structures determined from comparison of CID fragments with MS databases.  $\Delta$  ppm difference between observed and calculated molecular ion. nd= not detected. Metabolites disrupted by effluent exposure were determined from P values calculated using Mann Whitney test and after correction for a false discovery rate of 5%. n=10 and n=13 for control and exposed males; n=13 and n=16 for control and exposed females.

**Table 2. Concentrations of nonsteroidal anti-inflammatory drugs in effluent, blood plasma and tissues of effluent-exposed roach and human therapeutic plasma concentrations (HtPC).**

		Effluent <sup>a</sup> ng/ml (n=6)	plasma ♀ ng/ml (n=16)	plasma ♂ ng/ml (n=13)	gills ng/g (n=10)	kidney ng/g (n=12)	liver ng/g (n=12)	testes ng/g (n=12)	HtPC <sup>b</sup> ng/ml
<b>Naproxen</b>	Range	2.0 - 7.7	5.2 - 27	6.7 - 22	3.3 - 6.9	3.5 - 16	12 - 351	<0.11 <sup>d</sup> - 1.9	20000 - 50000
	Mean±SD	3.7 ± 2.3	13 ± 6.2	14 ± 4.6	5.1 ± 1.2	8.2 ± 3.8	75 ± 91	0.93 ± 0.48	
<b>Felbinac</b>	Range	<0.25 <sup>c</sup>	0.36 - 1.1	<0.19 <sup>d</sup> - 0.84	0.19-0.23	0.58 - 0.97	3.5 - 8.7	0.056 - 0.11	400 - 1000
	Mean±SD	<0.25 <sup>c</sup>	0.65 ± 0.15	0.57 ± 0.27	0.21 ± 0.01	0.77 ± 0.1	5.7 ± 1.6	0.091 ± 0.013	
<b>Diclofenac</b>	Range	0.22 - 0.38	<0.30 <sup>d</sup> - 1.9	<0.1 - 0.82	0.35 -1.0	<0.24 <sup>d</sup> - 0.35	<0.21 <sup>d</sup> - 0.31	<0.03 <sup>c</sup> - 0.32	500 - 3000
	Mean±SD	0.35 ± 0.10	0.43 ± 0.45	0.39 ± 0.27	0.58 ± 0.21	0.13 ± 0.10	0.21 ± 0.09	0.09 ± 0.09	
<b>Ibuprofen</b>	Range	<0.01 <sup>c</sup>	<7.5 <sup>d</sup> - 13	<7.5 <sup>d</sup> - 14	<2.0 <sup>c</sup> - <6.0 <sup>d</sup>	<2.0 - <6.0 <sup>d</sup>	<1.7 <sup>c</sup> - <5.1 <sup>d</sup>	<0.75 <sup>c</sup> - < 2.3 <sup>d</sup>	15000 - 30000
	Mean±SD	<0.01 <sup>c</sup>	5.2 ± 3.7	5.8 ± 4.3	<2.0 <sup>c</sup>	<2.0 <sup>c</sup>	<1.7 <sup>c</sup>	<0.75 <sup>c</sup>	
<b>Celecoxib</b>	Range	0.013 – 0.018	0.30 - 0.32	0.29 - 0.31	0.23 - 0.25	0.35 - 0.37	0.36 - 0.41	<0.019 <sup>c</sup> - 0.094	360 - 800
	Mean±SD	0.015 ± 0.002	0.31 ± 0.01	0.30 ± 0.01	0.24 ± 0.01	0.36 ± 0.003	0.38 ± 0.01	0.091 ± 0.001	
<b>Mefenamic acid</b>	Range	0.08 – 0.15	<0.94 <sup>d</sup> - 1.2	<0.94 <sup>d</sup> - 1.3	<0.25 <sup>c</sup> - 0.75 <sup>d</sup>	0.94 - 1.0	1.0 - 1.7	<0.094 <sup>c</sup> - <0.28 <sup>d</sup>	2000 - 10000
	Mean±SD	0.10 ± 0.03	0.70 ± 0.40	0.79 ± 0.42	<0.25 <sup>c</sup>	0.98 ± 0.03	1.2 ± 0.18	<0.09 <sup>c</sup>	

<sup>a</sup>Effluent samples measured at 0, 7, and 15 days of exposure. Plasma and tissue samples analysed at the end of the 15 day exposure period. <sup>b</sup>HtPC from Shultz et al.<sup>43</sup> <sup>c</sup> = MDL value, <sup>d</sup> = MQL value. SD= standard deviation. For the mean calculations, all concentrations that were over the limits of detection (≥MDL) but below the limits of quantification (<MQL) were assigned the value considered as the MDL in each case. Concentrations below the MDL were considered to be zero. Gills, kidney and liver samples were from females only. NSAID levels in control fish were below the MDL values given in Table S1.



## Figure captions.

### **Figure 1. Principal component analysis scores plot of the chemical profiles in plasma of roach (male and female) exposed either to a final treated WwTWs effluent or clean water.**

Extracts were profiled in -ESI mode by nUPLC-nESI-TOFMS. Quality control (QC) samples were used to monitor the analytical performance of the MS platform. % R2X values are given on the respective components. MC= male control, FC=female, control, FE=female effluent, ME= male effluent.

### **Figure 2. Effect of effluent exposure on PGE2 and PGF2 $\alpha$ concentrations in fish tissues and blood plasma**

PGE2 and PGF2 $\alpha$  concentrations were measured from the mean ( $\pm$  standard deviation) relative abundance compared to the internal standard. The % reduction in PG concentrations in the effluent-exposed group were determined from the mean values in the control fish which had been normalised to 100% for each tissue type. \*\*p<0.01;\*\*\*p<0.001;\*\*\*\*p<0.0001. n=10 and n=13 for control and exposed male plasma; n=13 and n=16 for control and exposed female plasma; n=12 and n=12 for control and exposed testes; n=10 and n=12 for control and exposed kidney; n=14 and n=12 for control and exposed liver; n=11 and n=10 for control and exposed gills, respectively. PGE2 was not detected in any plasma sample, MDL; method detection limit.

### **Figure 3. Number and classes of pharmaceuticals detected in the blood plasma and tissues of roach exposed to a WwTW effluent.**

NSAIDs, non-steroidal anti-inflammatory drugs. SSRIs, selective serotonin reuptake inhibitors. In total, 31 pharmaceuticals were detected across the different tissues and the identities of 26 were confirmed by pure standards (see Table S4). No pharmaceuticals were detected in samples from control fish.

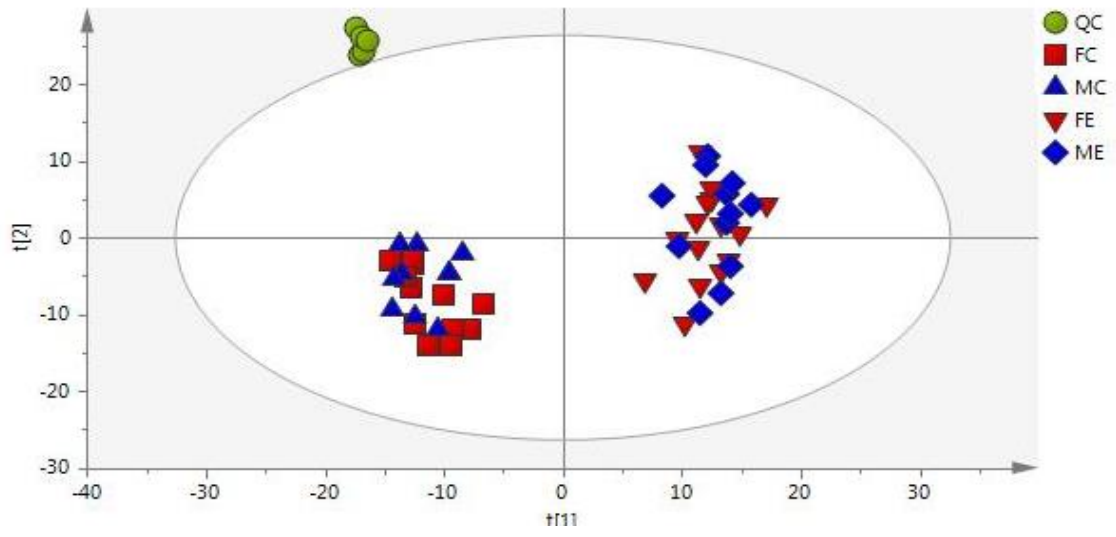


Figure 1.

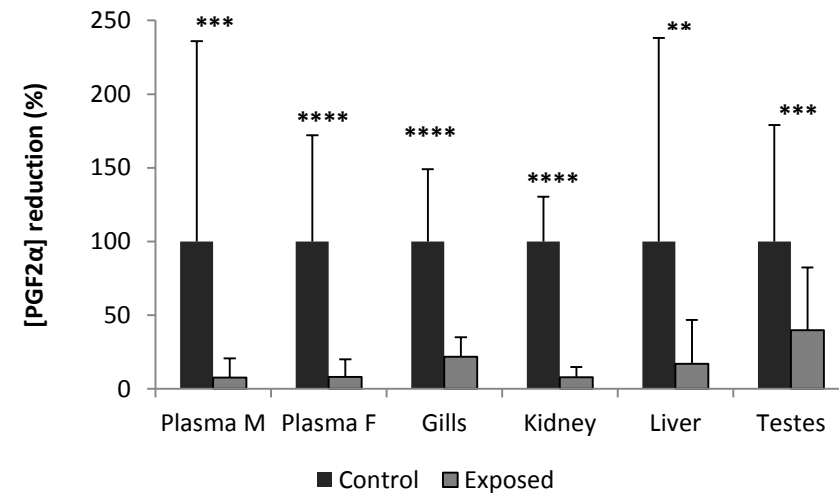
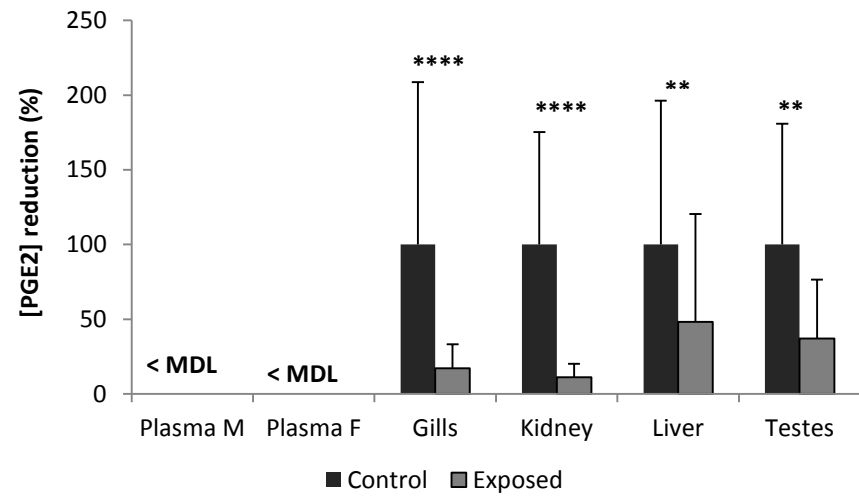


Figure 2.

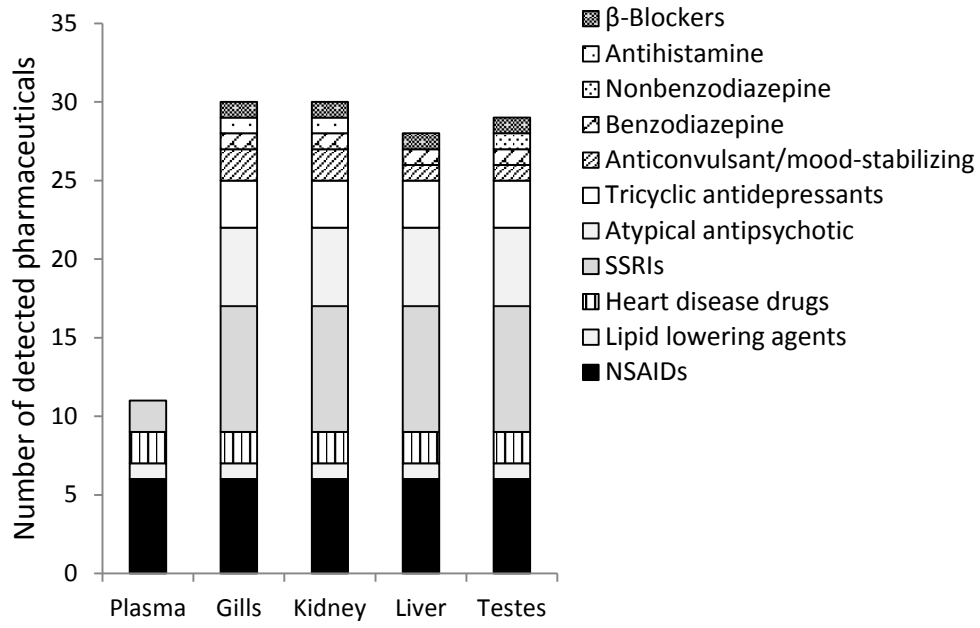


Figure 3.