

2018-02

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<http://hdl.handle.net/10026.1/10024>

10.1016/j.scitotenv.2017.09.101

Science of The Total Environment

Elsevier BV

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Accepted journal article in Science of the Total Environment – please refer to website

<https://doi.org/10.1016/j.scitotenv.2017.09.101>

Received 30 June 2017, Revised 8 September 2017, Accepted 11 September 2017, Available online 26 September 2017.

Active Pharmaceutical Ingredients Entering the Aquatic Environment From Wastewater Treatment Works: A Cause for Concern?

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Abstract

This work reports on the variation in wastewater treatment works (WwTW) influent concentrations of a wide variety of active pharmaceutical ingredients (APIs), their removal efficiency, effluent concentrations and potential risks to the aquatic environment. The research is based on data generated from two large UK-wide WwTW monitoring programmes. Taking account of removal of parent compound from the aqueous phase during treatment in combination with estimates of dilution available it is possible to prioritise the APIs of greatest risk of exceeding estimates of predicted no effect concentrations (PNEC) in receiving waters for all WwTW in the UK. The majority of substances studied were removed to a high degree, although with significant variation, both within and between WwTW. Poorer removal (between influent and effluent) was observed for ethinyloestradiol, diclofenac, propranolol, the macrolide antibiotics, fluoxetine, tamoxifen and carbamazepine. All except the last two of these substances were present in effluents at concentrations higher than their respective estimated PNEC (based on measurement of effluents from 45 WwTW on 20 occasions). Based on available dilution data as many as 890 WwTW in the UK (approximately 13% of all WwTW) may cause exceedances of estimated riverine PNECs after mixing of their effluents with receiving waters. The overall degree of risk is driven by the toxicity value selected, which in itself is controlled by the availability of reliable and relevant ecotoxicological data and

37 consequently the safety factors applied. The dataset and discussion, provides information to assist in
38 the future management of these types of chemicals.

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40 **Key words: pharmaceuticals, API, wastewater, effluent, fate, risk assessment**

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45 **1. INTRODUCTION**

46

47 The use and environmental prevalence of pharmaceuticals increases on an annual basis due to a
48 variety of reasons including the widening array of medical treatments available, greater availability of
49 medicines across the world, affordability, population growth, population ageing (in some countries)
50 and changing perspectives towards, for example, pain (Jelic et al., 2011). Active Pharmaceutical
51 Ingredients (API) are detected throughout the environment in water, soil, sediment, sludge as well as
52 in drinking waters in some countries (Kasprzyk-Hordern et al., 2008; Zorita et al., 2009; Wahlberg et al.,
53 2011; Jones et al., 2014; Lees et al., 2016). Although the mere presence of pharmaceutical is not
54 always associated with harm to the environment or human health, concerns are rising associated with
55 antimicrobial resistance and chronic impacts on biodiversity including endocrine disrupting effects on
56 fish (Levado et al., 2004; Jobling et al., 2005; Tyler et al., 2008). The main source of occurrence of
57 APIs in the river environment is from human use of pharmaceuticals, via the continuous discharge of
58 effluent from the Wastewater Treatment Works (WwTW) (Gardner et al., 2012; Melvin et al., 2016).
59 Hence, investigating the occurrence, fate and risk of APIs is currently of great interest to regulators
60 and the water industry alike, with a focus to better understand the loadings entering WwTW and the
61 observed within and between works variation in removal efficiencies and concentrations often
62 observed for APIs (Gardner, 2013).

63

64 The range of concentrations found for pharmaceuticals studied in the UK is similar to that observed in
65 continental Europe as well as in the USA (Kolpin et al., 2002; Ashton et al., 2004; Hope et al., 2012;
66 Bradley et al., 2016; Burns et al., 2017). Table 1 provides examples of other reported data for APIs
67 determined as part of this research, rather than a complete list of all APIs detected in effluent and
68 receiving waters. Other studies have also shown that there is a clear association between the number
69 of pharmaceuticals used in a society and the levels of API found in receiving water bodies ranging
70 from API concentration of typically less than 100 ng/l in the surface and groundwater and below 50
71 ng/l in treated drinking water (WHO, 2011; Furlong et al., 2017) to higher levels reported adjacent to
72 production facilities (Phillips et al., 2010). Predicted no effect concentrations (PNECs) have been
73 reported for some APIs below 1 ng/l and APIs such as diclofenac (CAS 15307-79-6), 17-beta-

74 estradiol (E2) (CAS 50-28-2) and 17-alpha-ethinylestradiol (EE2) (CAS 57-63-6) are on the European
75 Water Framework Directive (WFD) 'watch list' (EU, 2013). This requires member states to gather
76 monitoring data in order to assess risk to the environment, leading to significant sources of APIs
77 needing to be quantified and factors controlling the discharge of APIs carefully considered along with
78 impacts on receiving water ecology, including effects of mixtures (Bound and Voulvoulis, 2006).
79

80 **Table 1. Average aquatic concentrations for APIs of interest to this research found in river**
 81 **water, as well as usage, excretion and removal in WwTW.**

82

API	Therapeutic Class	Upstream (µg/l)	Influent (µg/l)	Effluent (µg/l)	WwTW removal (%)	Down stream (µg/l)	UK consumption (ton/year), 2009 and 2011	Excreted unchanged compound (%)
Aspirin (acetylsalicylic acid)	Anti-inflammatory/analgesics	NA	NA	NA	NA	<0.0005 ^b	130 ^d	<1 ^b
Atenolol	Beta blocker	NA	NA	NA	NA	0-0.56 ^b	28 ^e	90 ^f
Azithromycin	Antibiotic	NA	0.163 ^l	0.030 ^l	90 ^l	NA	NA	NA
Carbamazepine	Antiepileptic	NA	2.593 ^b	3.117 ^b	ND ^b	0.0005-0.356 ^b	48 ^e	3 ^b
Ciprofloxacin	Antibiotic	NA	1.090 ^l	0.052 ^l	97 ^l	NA	NA	NA
Clarithromycin	Antibiotic	NA	0.524 ^l	0.092 ^l	91 ^l	NA	NA	NA
Diclofenac	Anti-inflammatory	<0.020 ^a	0.107-0.981 ^c	0.599 ^a	70-92 ^c	0.154 ^a	28 ^e	15 ^f
Erythromycin	Antibiotic	<0.010 ^a	2.0 ^k	0.109 ^a	25-91 ^l	0.159 ^a	3 ^d	25 ^f
Oestrogen (E1)	Natural hormone	NA	0.042 ^g	0.011-0.025 ^g	58-96 ^g	NA	NA	NA
Oestradiol (E2)	Contraceptive	NA	0.016 ^g	0.0013-0.0039 ^g	89-96 ^g	NA	NA	NA
Ethinylestradiol (EE2)	Contraceptive	NA	0.0017 ^g	0.00033-0.00078 ^g	53-71 ^g	NA	NA	NA
Fluoxetine	Psychiatric drugs	NA	0.070 ^k	0.023 ^j	33-100 ^h	NA	6.4 ^m	NA
Ibuprofen	Analgesic	0.432 ^a	14.0 ^k	4.201 ^a	90-100 ^l	1.105 ^a	258 ^e	10 ^f
Oxytetracycline	Antibiotic	NA	1.09 ^l	0.029 ^l	99 ^l	NA	NA	NA
Ofloxacin	Antibiotic	NA	0.081 ^l	0.023 ^l	89 ^l	NA	NA	NA
Propranolol	Antihypertensive	0.010 ^a	0.542 ^b	0.093 ^a 0.388 ^b	28 ^b	0.041 ^a	15 ^e	<0.5 ^b
Tamoxifen	Anti-cancer	<0.010 ^a	0.0002-0.015 ^c	<0.010 ^a	32-45 ^c	<0.010 ^a	NA	NA

83

84 ND = not detected; NA = not available. ^aAshton et al., 2006; ^b[Kasprzyk-Hordern et al., 2008](#); ^cZhou et al., 2009;
 85 ^d2006 sales data for Wales; [Kasprzyk-Hordern et al., 2008](#); ^eIMS figure on active ingredient sales; ^fWHO, 2011;
 86 ^gHeffley et al., 2014; ^hClara et al., 2005; ⁱLi et al., 2014; ^jGardner et al., 2012; ^kGardner et al., 2013; ^lSinger et
 87 al., 2014; ^mBoxall et al., 2014

88

89 Many countries have therefore started monitoring programs to investigate the exposure of APIs in
 90 order to gain a better understanding of their sources, fate and risk (Falås et al., 2012). The Chemical
 91 Investigation Program (CIP) in the UK is a large ongoing investment being undertaken by the water
 92 industry to assist the UK in meeting its obligations under the WFD to monitor concentrations of
 93 priority chemicals including APIs in WwTW influent, intermediate processes and effluent as well as
 94 assessing their risk to receiving waters (Gardner et al., 2013). The first phase of the CIP (named CIP1
 95 here) was a project that ran from 2012-2015 with one of its aims to investigate the fate of trace
 96 substances (including 11 APIs) in influent, effluent and intermediate WwTW processes of 25 WwTW.
 97 Some of results from this program have been reported previously (Gardner et al., 2012, Gardner et al.;
 98 2013, Jones et al.; 2013 and Comber et al., 2014). The £140 million investment in the second phase of
 99 the CIP (labelled CIP2 in this work) program builds on the outputs from CIP1 but extends the range
 100 of WwTW monitored and the number of determinands in order to in some cases measure (for WFD
 101 priority substances and priority hazardous substances) and in some cases predict (for emerging

102 chemicals such as APIs) the impact on receiving waters. The CIP2 determinands include 19 APIs and
103 4 metabolites at currently 45 WwTW on 20 occasions. In total, over 60 000 samples are to be taken,
104 with over 2 million determinations. This study reports on the findings for APIs from the CIP1 and
105 CIP2 programmes.

106

107 WwTWs are primarily designed to serve the purpose of removing pathogens, suspended solids and
108 gross organic and inorganic matter, rather than the removal of the increasing numbers of modern
109 chemicals generally present in the $\mu\text{g/l}$ range or less (Melvin, 2016). It has also been observed that
110 there is a wide variation in removal rates for different substances, both within and between WwTWs.
111 This difference in removal rate creates large uncertainty factors for the prediction and modeling of
112 effluent concentrations and therefore creates a challenge in conducting meaningful risk assessments.
113 There are currently no statutory consents applied to APIs in WwTW effluent, however, there is an
114 urgent need to better understand the risk posed by APIs in effluents to receiving waters in order to
115 inform future investment and to design and implement better risk assessment (Gardner, 2013). The
116 presence of APIs is not measured on a routine basis for most WwTWs owing to cost and lack of
117 legislative drivers. Consequently, there are a number of previous studies modelling the impact of APIs
118 based on consumption, WwTW removal and dilution but the cost of analysis generally prevents the
119 actual measurement of APIs in effluent (Johnson et al., 2013a,b; 2015).

120

121 This study utilizes CIP 1 (11 APIs, from 25 WwTW sampled on up to 15 occasions) and the more
122 recent CIP2 program (19 parent APIs and 4 metabolites, from WwTW sampled on 20 occasions).
123 Although the APIs studied represent only a fraction of the total APIs in use, financial and practical
124 constraints associated with sampling, preservation, analysis and replication meant the number of
125 determinands needed to be controlled. However, APIs were prioritised on potential risk to the aquatic
126 environment and all of the main classes of API have been represented (Table A1). Concentrations in
127 the WwTW effluent have been compared with derived PNECs in receiving waters in order to generate
128 a priority list of APIs of potential concern.

129

130 **2. MATERIALS AND METHODS**

131 **2.1 Selection of Pharmaceuticals**

132 The selection of chemicals for CIP1 is discussed elsewhere (Gardner et al., 2012). The list of
133 candidate APIs for inclusion in CIP2 was based primarily on a prioritization study undertaken by
134 UKWIR in 2014 (UKWIR, 2014). Unlike many previous prioritisations, which focused on usage and
135 concentrations detected in surface waters/effluents, problem sites or substances, this study adopted a
136 risk assessment approach by comparing the estimated environmental concentrations of nearly 150
137 pharmaceuticals (screened on usage and perceived hazard from a list of thousands of candidate

138 substances) with data for their respective effect concentrations on a variety of receptor organisms in
139 the aquatic environment.

140

141 For the purposes of CIP2, this list was further refined by selection of substances that were considered
142 to have the greatest potential as candidate WFD priority substances. The criteria for this selection
143 were a) that the risk characterisation ratio (predicted concentration divided by the highest probable no
144 effect concentration (PEC/PNEC) ranked higher than 1 in the overall 2014 UKWIR prioritisation and
145 b) that the data supporting the derivation of a PNEC were relatively reliable and complied with the
146 WFD approach to PNEC derivation (EU 2011). In effect, this meant that PNECs were derived using
147 experimental rather than modelled effects, long-term effects in organisms from different trophic levels
148 were available (though short term exposure was also considered) and assessment factors were applied
149 according to WFD guidance (EU 2011).

150

151 The APIs prioritised were then further reviewed for their relevance to wastewater treatment, and the
152 likelihood that the substance might be present in sewage effluents and hence discharged to surface
153 waters (rather than being partitioned to sewage sludge). This resulted in the list of substances
154 tabulated in Table A1 of the Electronic Supplementary Information (ESI). For the purposes of
155 estimating risks, the PNEC values derived in the UKWIR prioritization (UKWIR 2014) were then re-
156 examined and (where available) they were substituted with the latest estimates derived by the EU
157 Joint Research Centre (JRC, 2015), by the pharmaceutical industry (Astra Zeneca, 2016; NSF, 2016)
158 or published in the open literature (Murray-Smith et al., 2012). Where no PNEC was available from
159 these sources, the ecotoxicology data applied in deriving the PNECs reported by UKWIR (UKWIR
160 2014) were used to deterministically estimate PNECs, according to WFD guidance (EU 2011) (Table
161 2 and ESI Table A1). It is recognized that as new ecotoxicity data becomes available, substance
162 PNECs are subject to update, and the estimates of PNECs applied in the present study may not, in
163 every case, reflect the most up to date applied or proposed PNEC for regulatory purposes (e.g. under
164 the WFD or European Medicines Agency (EMA) Environmental Risk Assessments. However, the
165 estimated PNECs reported here were applied in the CIP for the purposes of selection for monitoring,
166 preliminary risk assessment and prioritization, and so remain relevant in this context, and it is beyond
167 the objectives of the present study to derive new PNECs for each of the APIs monitored.

168

169 **2.2 Sampling programme**

170

171 WwTWs were selected for the CIP program on the basis of broadly representing the distribution of
172 UK WwTWs (A1, ESI), predominantly activated sludge plants (ASP) and trickling biofilters (TF) but
173 also Membrane bioreactors (MBR) and oxidation ditches (OD) (Table A2 of ESI).

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Data used for this research were (Table A2 of ESI):

- **CIP1 program:** 25 WwTW data for primary, secondary and tertiary process for 11 APIs. Sampling for this element of the programme was conducted over a two-year period between 2011 and 2013. In this part of the programme two samples (spaced more than 4h apart to provide a degree of replication) were taken on between 10 and 15 occasions.
- **CIP2 program:** 19 APIs and 4 metabolites were sampled on 20 occasions at 45 WwTWs in the influent and effluent (not intermediate process stages, unlike CIP1) over a two-year period between 2015 and 2017.

Samples were collected on a stratified/random spot sampling basis (i.e. grab samples taken at discrete times rather than multiple integrated sampling), with sampling occasion spaced at approximately monthly intervals. A minimum of 15% of samples was taken at non-working hours (evenings and weekends) to ensure a wide a range possible of sampling intervals.

2.3 Sampling and analysis

The samples were collected in stainless steel samplers, stored in glass container and transported at 4° C to the analysis laboratories. The samples were stored a maximum of 5 days prior to analysis. This period was shown to be appropriate as not leading to more than a 20% change in determinand concentration; as confirmed before the start of the CIP sampling programme by undertaking tests of sample stability. Samples for the determination of steroid oestrogens were preserved by adding 30% hydrochloric acid and copper nitrate (Gardner, 2012). All analysis was by laboratories with ISO17025 accreditation. Prior to the programme candidate laboratories were required to undertake tests of analytical performance to demonstrate that they met the stated programme requirements for limit of detection, precision and recovery in relevant sample matrices at relevant concentrations – that is, proof of performance was required, rather than methods being stipulated. Methods used for the determination of pharmaceuticals were all based on variants of High Performance Liquid Chromatograph–Mass Spectrometry (HPLC-MS) or Gas Chromatography-Mass Spectrometry (GC-MS). Quality assurance/quality control (QA/QC) procedures, including the use of field blanks, were observed and reported for sample collection. Within laboratory QC sample pre-treatment and analysis for both laboratory tests and field sampling. Laboratories also took part in a bespoke proficiency testing scheme for pharmaceuticals. Details of the proficiency testing scheme used to ensure quality assurance is provided in A2 of the ESI. Where reported concentrations were below LOD (for the majority of substances apart from ibuprofen and tamoxifen this applied to fewer than 10% of the approximately 1000 results reported), the result was substituted at half face value - as stipulated in the

210 relevant daughter Directive (EC, 2009) of the WFD. There were significant instances of inter-
 211 laboratory bias or inter-regional variation, which would otherwise indicate if there was a bias in the
 212 procedure of sample handling and analysis methodology.

213

214

215 **Table 2. Determinand abbreviations and required limits of detection and total error**

Code	Determinand	Concentration (µg/l)			P% ²
		PNEC ¹	Required LOD effluent	Required LOD river	
ATNL	Atenolol	148	0.01	0.01	50
ATOV	Atorvastatin	1.7	0.01	0.01	50
ATOV _o	Ortho-hydroxy-atorvastatin	1.7	0.01	0.01	50
ATOV _p	Para-hydroxy-atorvastatin	1.7	0.01	0.01	50
AZMY	Azithromycin	0.09	0.005	0.005	50
CBAZ	carbamazepine	2.5	0.1	0.1	50
CBAZe	10,11- epoxy-carbamazepine	2.5	0.1	0.1	50
CIPR	Ciprofloxacin	0.089	0.01	0.01	50
CLMY	Clarithromycin	0.13	0.01	0.01	50
DCF	Diclofenac	0.05	0.01	0.01	50
ERMY	Erythromycin	0.2	0.1	0.1	50
ERMY _n	Norerythromycin	0.2	0.1	0.1	50
E1	Oestrone	0.003	0.001	0.001	50
E2	17β oestradiol	0.001	0.0003	0.0003	50
EE2	17α ethinyloestradiol	0.0001	0.00003	0.00003	50
FLXT	Fluoxetine	0.047	0.01	0.01	50
IBPF	Ibuprofen	0.01	0.01	0.01	50
METF	Metformin	13.45	0.1	0.1	50
PRPL	Propranolol	0.1	0.01	0.01	50
RNTD	Ranitidine	0.31	0.1	0.1	50
SERT	Sertraline	0.121	0.01	0.01	50
SERT _n	Norsertaline	0.121	0.01	0.01	50
TMXF	Tamoxifen	0.49	0.005	0.005	50

216 ¹ Estimated PNEC (ESI Table 1). ²The target maximum tolerable error is equal to:

$$217 \left[(targetLOD)^2 + \left(\frac{A \times P\%}{100} \right)^2 \right]^{\frac{1}{2}}$$

218

219 Where the target maximum LOD and P% are given in the table and A is the determinand concentration in the
 220 sample. Performance testing should seek to demonstrate that the tolerable total error limit is achieved by
 221 showing that precision (2 x standard deviation) and bias are respectively no larger than half the target maximum
 222 total error. Thus, for example, for a total error limit of 100 units, standard deviation should be shown not to be
 223 larger than 25 and bias should not exceed 50. LOD was defined as 3.3x the standard deviation of blank-
 224 corrected results of determinations made on a sample containing essentially no determinand (where possible in a
 225 relevant sample matrix) (Thompson and Ellison, 2013) In many cases, it was not possible to find effluent
 226 samples free from determinands in which case a synthetic sample was used.)

227

228 2.4 Data handling and analysis

229 The data handling and the statistical analysis were conducted with either Microsoft Excel (2016) or

230 IBM SPSS Statistics software (version 20).

231

232 In the data handling, the replicates were averaged and this value was then used for further statistical
233 calculations. Mean, median, maximum, minimum and percentiles were calculated from the daily
234 average. Fraction remain was calculated from the influent concentration as a fraction of the various
235 stages of the process. The removal was calculated as percentage from the concentration (C):

236

$$237 \text{ Removal (\%)} = (C_{\text{influent}} - C_{\text{effluent}}) / C_{\text{influent}}$$

238

239 For the purpose of this research the term 'removal' relates to the loss of specified compounds from
240 the aqueous phase between influent and effluent (and intervening process steps where quoted). It
241 should be noted that the term removal does not necessarily mean degradation of the API; the loss of
242 the parent compound may be a result of a combination of partitioning to particulates and/or
243 degradation to metabolites.

244

245 **2.5 Risk assessment approach**

246 **2.5.1 Face value risk ranking**

247 A "face value" exceedance is one in which the mean effluent concentration is greater than the relevant
248 estimated PNEC; a "high confidence" exceedance is one for which the lower part of the 90%
249 confidence interval about the mean effluent concentration is greater than the estimated PNEC i.e.
250 there is 95% confidence that the mean is larger than the estimated PNEC.

251

252 **2.5.2 Refined risk assessment based on estimated available dilution**

253 Previous research has used a combination of modelled average river flows (Comber et al., 2013) and
254 average WwTW discharge volumes to estimate dilution of effluent with receiving water. Effluent
255 flow data was derived from measured values for larger WwTW, but estimated for works serving less
256 than 2000 population equivalent based on water company estimates of connected population and
257 per capita wastewater discharge to sewer (200 l/head/day- including an allowance for runoff)
258 (Comber et al., 2007). A matrix (Table 3) of available dilution was then generated.

259

260

261 **Table 3. Estimated dilutions available for UK WwTW**

262

	Dilution ratio band									Total no. works
	0-1	1-2	2-5	5-10	10-15	15-20	20-50	50-100	100+	
Midpoint dilution	1	1.5	3.5	7.5	12.5	17.5	35	75	100	
Combined dist' dil' ¹	0 ²	4.4	10.6	23.6	36.5	35	70	150	200	
Population served										
<250	86	75	54	54	11	21	86	86	2656	3127
251-500	0	6	0	6	25	0	50	81	605	774
501-2000	0	5	20	20	51	46	351	285	544	1322
2001-10000	17	25	151	160	130	84	202	93	130	993
10001-50000	82	67	160	103	24	30	48	15	18	548
50001-200000	54	29	27	12	5	12	20	5	0	164
200001-1m	74	0	11	0	0	0	0	0	0	85
>1m	4	4	0	0	0	0	0	0	0	7
Total	316	211	423	355	246	194	756	564	3954	7020
%	5	3	6	5	4	3	11	8	56	

263 ¹ Values used to calculate PECs in river using a Combined Distribution simulation (see A3 and Table
 264 A3). ² A worst case scenario of zero dilution.

265

266 The next step was to generate a cumulative percentile distribution of effluent concentration data (in
 267 10%ile intervals between 10 and 100%). This was achieved by averaging the effluent concentrations
 268 for each of the 45 WwTW sampled as part of the CIP2 survey. Step three was to divide each
 269 percentile concentration by the dilution available (using the value from the combined distribution
 270 estimate – See A3 of the ESI) to generate a PEC. The PEC can then be compared with estimated API
 271 PNECs to determine the number of WwTW at risk of exceeding the PNEC for any given each dilution
 272 band and percentile effluent concentration. An example of the risk assessment is provided in Table
 273 A4 of the ESI.

274

275 **3. RESULTS and DISCUSSION**

276

277 **3.1 Removal efficiency for APIs**

278

279 The CIP1 study generated removal data for APIs across all stages of treatment, influent, after primary
 280 settlement, secondary biological treatment and where applied, post tertiary treatment. To gain a better
 281 understand of the fate of the 11 pharmaceuticals through the treatment train, the fraction of API
 282 remaining in the effluent after treatment was calculated across all 25 WwTW in the CIP1 program
 283 (Table 4).

284

285 Each cycle of sampling was treated as an isolated entity (averaging the samples within the same day),
 286 thus simplify the ability to compare APIs removal across the diverse range of works. As seen from the

287 data in Table 4, most APIs are removed in the secondary biological treatment process and very little
 288 through further tertiary treatment. This corresponds well with previously published data (Stockholm
 289 Vatten, 2010). The absolute effluent concentrations (Table 4) also correspond well with those reported
 290 elsewhere for predominantly UK effluents (Table 1).

291

292 **Table 4. CIP1 data for API fraction remaining throughout the process stages in the WwTW, as**
 293 **well as the absolute effluent concentration**
 294

Fraction of API remaining in effluent after treatment												
API	Primary Process			Secondary Process			Tertiary Process			Effluent Concentration (µg/l)		
	Median	5-%ile	95-%ile	Median	5-%ile	95-%ile	Median	5-%ile	95-%ile	Median	5-%ile	95-%ile
Diclofenac (DCF)	0.76	0.40	1.6	0.52	0.18	1.2	0.44	0.16	1.0	0.20	0.084	0.51
Erythromycin (ERMY)	0.79	0.26	1.7	0.52	0.11	1.2	0.44	0.08	1.1	0.43	0.052	2.0
Ethinylestradiol (EE2)	0.96	0.36	2.4	0.54	0.13	1.9	0.49	0.10	3.5	0.0003	0.0001	0.0020
Oestrone (E1)	1.0	0.59	2.1	0.28	0.02	2.4	0.10	0.01	1.2	0.0048	0.0007	0.058
Oestradiol (E2)	0.97	0.44	1.6	0.11	0.01	0.8	0.05	0.01	0.80	0.0009	0.0001	0.012
Fluoxetine (FLXT)	0.79	0.38	1.5	0.48	0.08	1.2	0.46	0.09	1.1	0.032	0.0050	0.066
Ibuprofen (IBPF)	0.83	0.39	1.3	0.04	0.00	0.2	0.01	0.00	0.21	0.19	0.0050	2.9
Ofloxacin (OFLX)	0.88	0.12	2.2	0.45	0.08	1.4	0.34	0.05	1.0	0.016	0.0050	0.14
Oxytetracycline (OXTCY)	0.66	0.13	1.6	0.16	0.00	0.6	0.13	0.01	0.54	0.21	0.019	1.1
Propranolol (PRPL)	0.91	0.52	1.4	0.68	0.14	1.2	0.65	0.16	1.2	0.14	0.042	0.32
Salicylic acid (SLCYA)	0.85	0.28	1.6	0.01	0.00	1.1	0.01	0.00	0.33	0.18	0.017	3.8

295

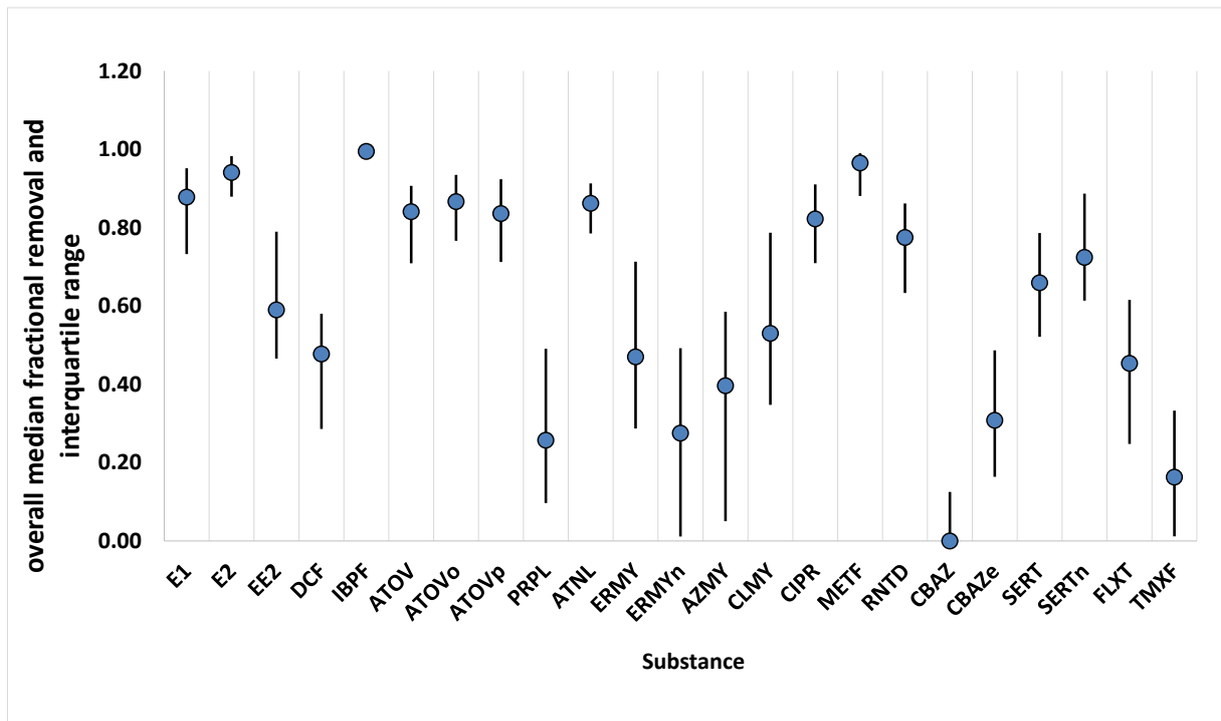
296

297 ERMY, DCF, FLXT and OXTCY were all shown to have similar removal efficiencies throughout the
 298 primary and secondary treatment processes based on the CIP1 dataset. The primary process relies
 299 mostly on removal of APIs through adsorption onto sludge (Stockholm Vatten, 2010) as retention
 300 times are relatively low and so this fits well to the data found for OXTCY, as it is previously known
 301 to adsorb strongly onto solids (Verlicchi, 2012) and found at higher concentration (4 mg/kg) in sludge
 302 compared with other APIs such as DCF, ERMY and FLXT (0.07, 0.05 and 0.12 mg/kg, respectively)
 303 (Jones et al., 2014). PRPL had overall poor removal of 35% and 26% (0.65 and 0.74 fraction
 304 remaining) between influent and effluent for CIP1 and CIP2 respectively (Table 4 and 5), which also
 305 corresponded well with previously published data of 28% removal efficacy (0.72 fraction remaining)
 306 in WwTW (Kasprzyk-Hordern et al., 2008).

307

308 In the CIP2 data set (Table 5) there was high total removal (based on comparison of influent and
 309 effluent API concentrations) of IBPF, METF, E2, ATNL, ATOV_p, E1, ATOV, CIPR, ATOV_o, which
 310 all had fraction remaining ratios of 0.2 or lower (i.e. better than 80% removal efficiency) (Figure 1,

311 and Table A5). This suggested either rapid biodegradation of the parent compound and/or adsorption
312 to sludge. None of the substances are considered sufficiently volatile to suggest any significant loss to
313 the atmosphere. The intermediate set of APIs consisting of SERTn, RNTD, CLMY, EE2, FLXT, DCF
314 and ERMV, which all had fraction remaining below 0.6 (i.e. greater than 40% removal efficiency).
315 PRPL, CBAZe, ERMVn, AZMY and CBAZ all showed poor removal through the WwTW process
316 (Figure 1 and Table A5).



317
318 **Figure 1. Fractional removal for APIs in CIP2**

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320

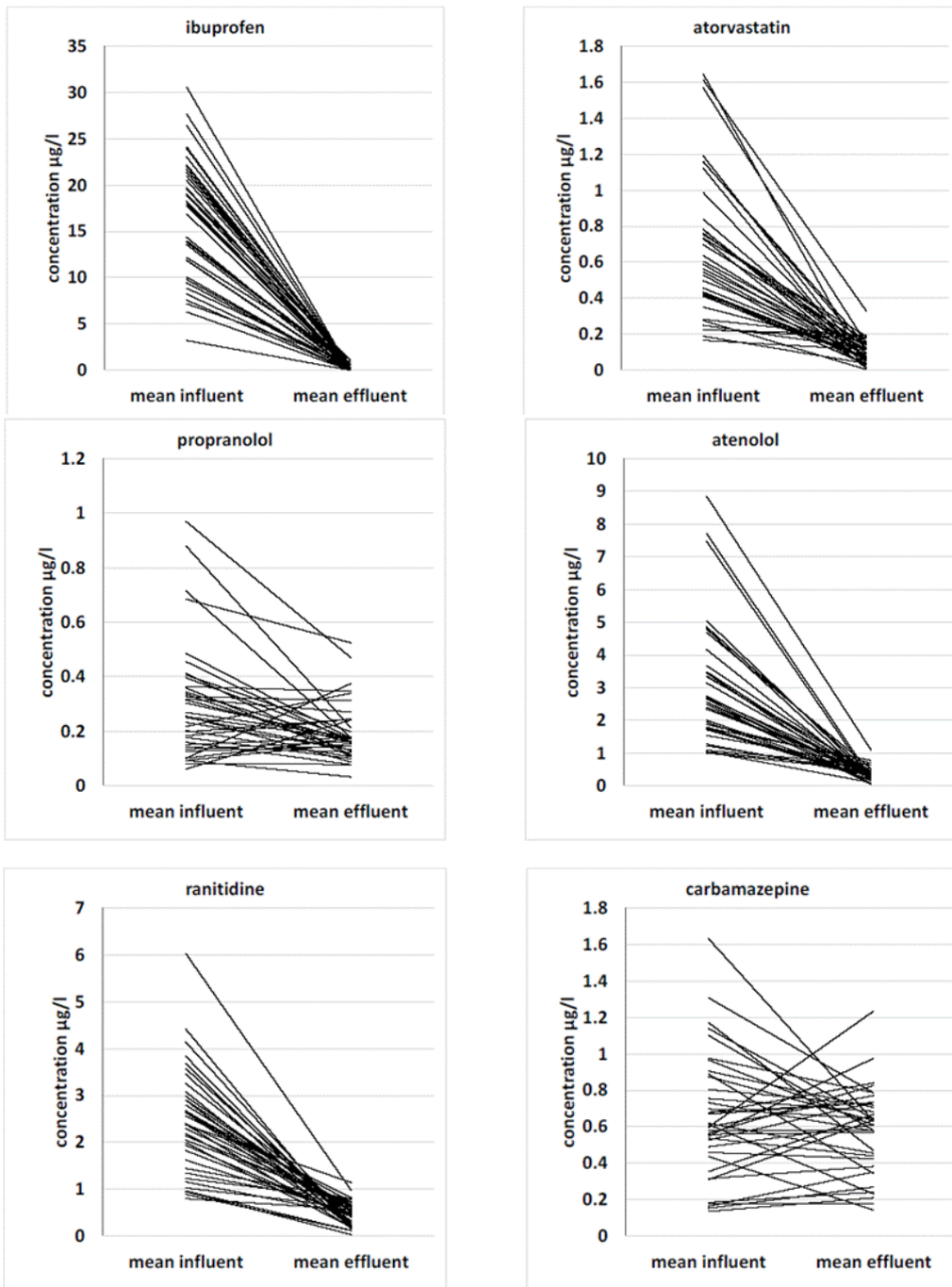
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Table 5. Summary concentration values for CIP2 APIs (45 WwTW sampled on 20 occasions)

Substance	Influents (µg/l)			Effluents (µg/l)		
	Median of WwTW average values	25%ile	75%ile	Median of WwTW average values	25%ile	75%ile
E1	0.038	0.030	0.049	0.004	0.002	0.014
E2	0.014	0.011	0.019	0.001	0.0003	0.002
EE2	0.00051	0.00041	0.00097	0.00020	0.00014	0.00040
DCF	0.54	0.40	0.76	0.29	0.20	0.41
IBPF	18.13	12.08	21.32	0.11	0.02	0.56
ATOV	0.61	0.41	1.01	0.10	0.06	0.16
ATOV _o	1.33	0.81	1.76	0.17	0.08	0.29
ATOV _p	1.33	0.82	2.03	0.21	0.12	0.35
PRPL	0.260	0.171	0.354	0.174	0.119	0.245
ATNL	2.600	1.872	3.297	0.323	0.210	0.463
ERMY	0.733	0.551	1.161	0.350	0.190	0.558
ERMY _n	0.060	0.050	0.091	0.050	0.027	0.050
AZMY	0.351	0.171	0.748	0.202	0.095	0.425
CLMY	0.953	0.684	1.564	0.400	0.265	0.711
CIPR	0.861	0.385	1.510	0.147	0.067	0.276
METF	129	104	208	4.8	1.7	15
RNTD	2.35	1.68	3.06	0.529	0.286	0.730
CBAZ	0.60	0.43	0.84	0.641	0.477	0.756
CBAZ _e	0.18	0.11	0.42	0.117	0.072	0.292
SERT	0.18	0.12	0.27	0.063	0.037	0.081
SERT _n	0.12	0.10	0.21	0.033	0.016	0.045
FLXT	0.10	0.07	0.15	0.051	0.036	0.079
TMXF	0.0034	0.0026	0.0047	0.0025	0.0025	0.0028
TXP	0.0050	0.0028	0.0052	0.0050	0.0026	0.0050
BZT	2.16	1.60	3.97	1.38	1.08	2.62
TZT	1.59	1.19	2.60	1.27	0.88	1.96

323
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Figure 2 below represents mean concentrations in the influent and effluent for selected APIs with the others shown in Figure A2 and demonstrates the degree of variability for APIs between WwTW.



329

330

331

332 **Figure 2. Graphic representation of mean concentrations of APIs in influent and effluent of**
 333 **individual CIP2 WwTWs**

334

335 In some cases there appears to be an increase in API concentrations in the effluent compared with the
 336 influent (Figure 2 and A2 of ESI). There are three main reasons for this:

- 337 1) The hydraulic retention time (HRT) within a WwTW means that samples of influent and
 338 effluent collected at the same time (a practical requirement of the work) may not reflect actual
 339 removal efficiency owing to within works management practices, e.g. batch flow, sludge

340 return pumping, taking place at the time of sampling. Given HRTs vary vastly between works
341 and types of works it was not practical to calculate nor practically sample WwTW based on
342 their HRTs.

343 2) The APIs were detected at ng/l levels in a highly complex matrix (particularly the influent)
344 therefore analytical errors may lead to apparent increase in concentrations during treatment
345 (Jelic et al., 2011).

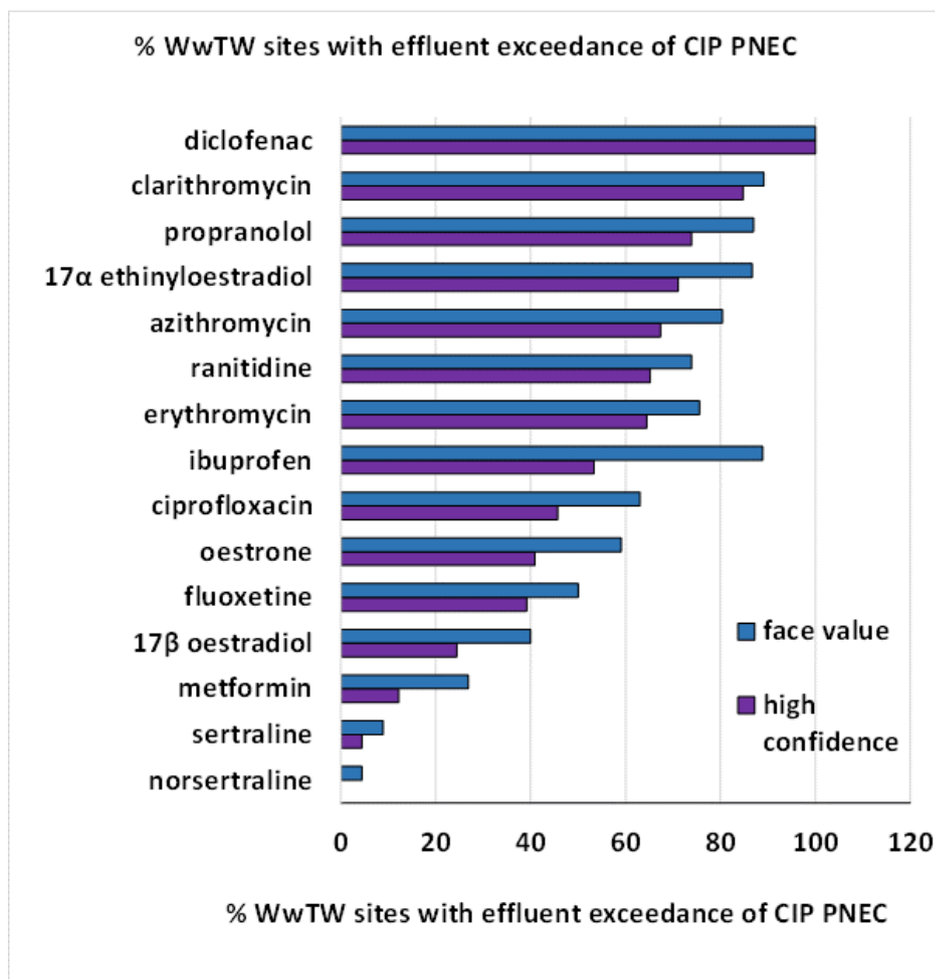
346 3) In some cases this is a real effect, for example E1 is a degradation product of E2 (Heffley et
347 al., 2014) and so if the rate of loss of E1 during treatment is less than that of E2, then an
348 apparent increase in E1 will occur.

349

350 **3.2 What is the environmental risk of the APIs being discharged in WwTW effluent?**

351 The median and interquartile concentration values of pharmaceuticals in influents and effluents are
352 summarised in Table 5. Figure 3 shows a summary risk ranking of the CIP pharmaceutical group of
353 substances in relation to the estimated predicted no-effect concentrations applied in CIP (CIP
354 PNECs). A “face value” exceedance is one in which the mean effluent concentration is greater than
355 the relevant estimated PNEC; a “high confidence” exceedance is one for which the lower part of the
356 90% confidence interval about the mean effluent concentration is greater than the estimated PNEC i.e.
357 there is 95% confidence that the mean is larger than the estimated PNEC. Substances not shown do
358 not figure as noteworthy exceedances.

359



360

361 **Figure 3. Risk ranking of CIP2 APIs**

362

363 Figure 3 above illustrates the severity of potential non-compliances for pharmaceuticals as the ratio of
 364 the observed concentrations in effluents to the relevant estimated PNEC. This ratio represents the
 365 dilution that would be required to achieve compliance, assuming zero upstream concentrations. An
 366 important proportion of UK wastewater treatment discharges are not subject to very much greater than
 367 a twofold dilution so the potential for downstream non-compliance with PNEC values does exist on
 368 the basis of a single effluent discharge alone. Table 3 shows that over 500 WwTW has estimated
 369 dilutions of less than 2, 8% of all the WwTW in the UK. Added to this concern must be a
 370 consideration of the pharmaceutical concentrations already present in a receiving watercourse
 371 upstream of the discharge. Whilst the CIP2 programme did not include the determination of
 372 pharmaceuticals in upstream river samples such analysis was undertaken for a range of Priority
 373 Substances, including trace organic compounds that like pharmaceuticals, are primarily discharged as
 374 a result of domestic inputs to wastewater. The evidence obtained from these investigations is that the
 375 burden of upstream contamination is far from irrelevant and that discharges in the higher parts of a
 376 river catchment, for example from septic tanks and small WwTW, can raise concentrations to values

377 that subsequent discharges lower in the catchment only serve to maintain (Phillips et al., 2015). This
378 is an aspect that deserves careful future examination in the context of pharmaceuticals.

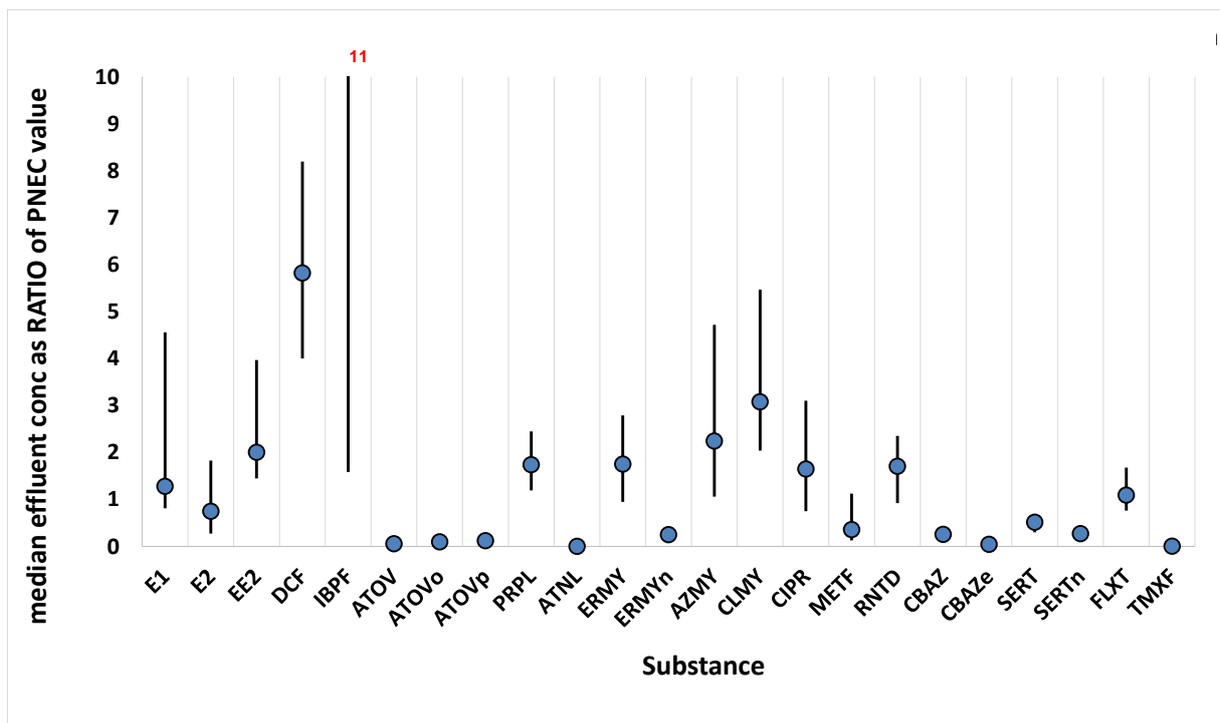
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380 Figure 4 shows that several pharmaceuticals have been shown to be present in effluents at
381 concentrations close to, or in many cases in excess of, values that might form the basis of future
382 regulatory limit values.

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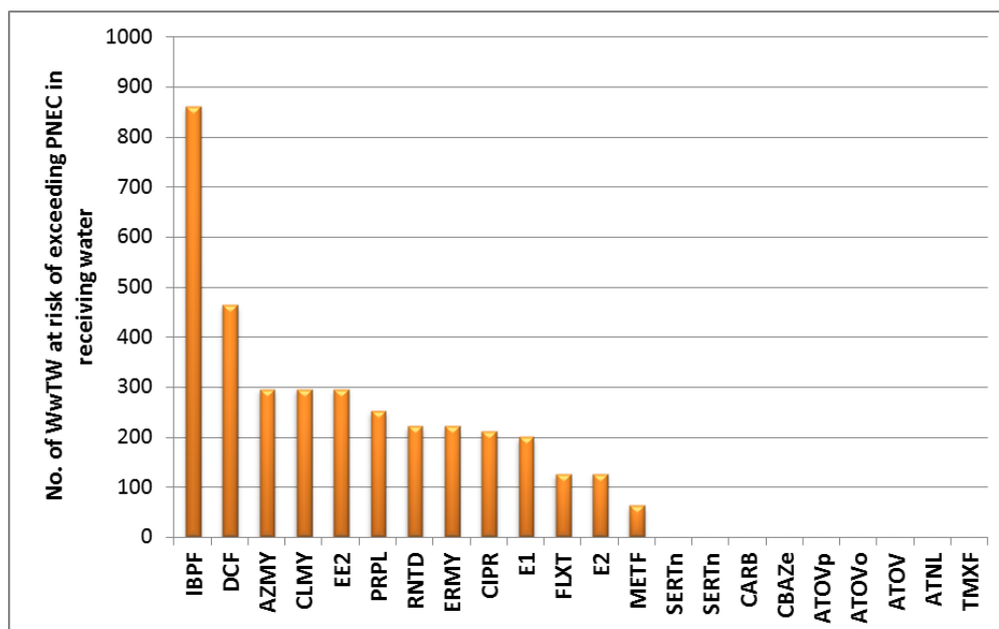
387 **Figure 4. Required within river dilution of WwTW effluent for API concentrations to be**
388 **less than their estimated PNEC.**

389 **Note median effluent concentration for ibuprofen (IBPF) as a ratio of estimated PNEC is 11.**

390

391 Applying a more realistic risk assessment using estimates of available dilution for UK WwTW
392 effluents discharged to receiving waters, combined with the measured API concentrations from the
393 CIP2 dataset generates a similar priority ranking list in terms of the number of WwTW potentially
394 exceeding downstream estimated PNECs after the effluent has mixed with receiving water (Figure 5).
395 For IBPF this equates to 890 WwTW or 13% of all WwTW in the UK. This estimate is also based on
396 the assumption that there are no significant inputs of API upstream of the WwTW in question.

397



398

399 **Figure 5. Number of WwTW at risk of exceeding estimated PNEC downstream of receiving**
 400 **water**

401

402 DCF, AZMY, CLMY, EE2, PRPL, CIPR, RNTD and E1 are all predicted to exceed downstream
 403 PNECs in over 200 WwTW. Required mean removal efficiency for any tertiary treatment would
 404 range from 35% to 61% depending on the API (Table A6 of ESI). Whether the same tertiary treatment
 405 technology could be applied to all of the APIs largely depends on their physico-chemical
 406 characteristics. The use of granulated activated carbon (GAC) would require an API to have a
 407 reasonable affinity for carbon (i.e. a relatively high octanol:water coefficient - logKow) which may
 408 not always be the case for APIs with a high degree of polarity, particularly those that are charged at
 409 typical effluent pH (pH 7.5) which would include DCF, IBPF, ATOV and to a degree CIPR (pKa =
 410 6.09). Furthermore, as can be seen from Figures 1 and 2, there are considerable variations in the
 411 removal rates between WwTW and so it may be expected to observe a similarly wide variation in
 412 removal rates and/or final effluent API concentration, if additional tertiary treatment were to be
 413 applied. This would obviously lead to a degree of uncertainty regarding possible compliance with any
 414 given in river PNEC or water quality standard.

415

416 Figures 1 and 2 show that WwTW in general, have a high (but variable) removal rate for most
 417 substances with only E2, EE2 propranolol, the macrolide antibiotics, carbamazepine fluoxetine and
 418 tamoxifen exhibiting poor removal. It is clear (and unsurprising) that more complex factors, such as
 419 the contaminant load on the WwTW, residence time in the works, overall strength of the influent,
 420 questions of operation and maintenance as well as the presence of absence of tertiary treatment “add-
 421 ons”, combine at each location to result in the observed treatment performance (Zorita et al., 2009;
 422 Le-Minh et al., 2010; Deegan et al., 2011). In the wider context, the persistence of pharmaceuticals in
 423 surface waters will be determined by the degree of upstream contamination from other (in this case,

424 presumably WwTW) inputs higher in the river catchment. As has been seen in elements of the CIP2
425 programme dealing with Priority Substances, upstream contamination and lack of headroom for
426 downstream discharges can often be more important than the local impact of a given WwTW. The
427 likely importance of upstream inputs for pharmaceuticals is unclear. Whilst upstream inputs are
428 inevitable in all sites except those at the top of catchments (where there may still be influences from
429 septic tanks) the effect of such inputs is not known, but if smaller WwTW are less efficient than the
430 predominantly larger works selected for the CIP programmes, then the risk to surface waters of
431 exceeding estimated PNECs for APIs may be significant. Persistence and the degree and rate of
432 breakdown in the environment are critical in this context. To fulfil their purpose pharmaceuticals need
433 to be absorbed by the patient, to remain for sufficient time to have the desired effect and then be
434 excreted. This means that in terms of their structure and hence fate and behaviour, pharmaceuticals
435 tend to occupy a middle ground between substances on the one hand that are non-polar, hydrophobic,
436 insoluble, and persistent and those that are highly polar, soluble, mobile and relatively readily
437 biodegradable. This suggests that some degree of degradation in-river might mean that input of
438 pharmaceuticals upstream may not be as great a risk as it is for other persistent, highly mobile priority
439 substances such as some metals, persistent pesticides and industrial compounds.

440

441 It should, however, be noted that this assessment is based on the mixing of single APIs in effluent and
442 receiving water under average flow conditions for a fraction of the APIs currently available and used.
443 During summer months river flows are significantly lower than average values, yet effluent flows will
444 remain relatively stable (accepting rain events contributing to flow in combined sewerage systems)
445 leading to generally lower dilution available and therefore higher concentrations of effluent derived
446 contaminants in receiving waters. Seasonal pattern of use for some APIs, antihistamines in summer,
447 flu vaccines in winter etc, would also lead to a variable distribution of APIs in WwTW effluent and
448 hence variable risk to receiving waters. The potential risk of mixtures is complex and requires detailed
449 knowledge of ecotoxicology for the APIs of interest. Such assessments along with determining
450 temporal variations in risk, require more a significantly detailed dataset (not necessarily currently
451 available) and as such is beyond the scope of this broader risk assessment.

452

453 Whilst the objective of this research has not been to estimate costs for compliance, drawing on
454 previous estimates of costs for API treatment based on fitting sand filters and granulated carbon
455 sorption technology, the whole life cost (based on 2007 data) for achieving downstream compliance
456 with the estimated IBPF PNEC would approximately £9bn (Comber et al., 2007). So for illustrative
457 purposes it is evident that achieving compliance for all API estimated PNECs would be a substantial
458 investment by the water industry. These estimates are only based on mixing downstream of receiving
459 water and effluent and do not take account of any biodegradation or sorption to particulates leading to

460 reduced exposure which would need to be considered as part of a more detailed risk assessment prior
461 to considering any remedial action regarding removal of APIs from WwTW effluent.

462

463 Much in relation to future compliance (and therefore cost to the water companies) will depend on the
464 derivation method and data used to set water quality standards. The outputs of the CIPs in this case
465 constitute a valuable risk assessment of the likely impact of whatever regulations might be introduced
466 in the future. Of the pharmaceuticals / likely future Priority Substances, the so-called WFD watch list
467 substance diclofenac, the steroids as well as possibly ibuprofen appear to be at risk of causing
468 widespread exceedances of estimated PNECs in UK rivers. With respect to these substances, options
469 of regulated use and control of patient behaviour relating to disposal of unused medicines might be
470 enough to make a substantial difference. However, wastewater treatment solutions might turn out to
471 be essential for the steroids, at least in the case of EE2.

472

473 **4. CONCLUSIONS**

474 As has been observed for the CIP1 program there are a high variability in the removal of APIs
475 observed between and within the individual plants. This variation may be due to many factors such as
476 process technology as well as regional variation. Rates of removal in wastewater treatment have also
477 been determined. The majority of substances studied are removed to a high degree, but with a wide
478 variation in performance. Those that are less substantially reduced in concentration are
479 ethinyloestradiol, diclofenac, propranolol, the macrolide antibiotics, fluoxetine, tamoxifen and
480 carbamazepine. All except the last two of these substances are present in effluents at concentration
481 higher than their estimated respective PNECs.

482

483 If the PNECs applied in the present study were all implemented as regulatory quality standards under
484 the WFD, the risk assessment undertaken suggests that over a 10 times dilution would be required, to
485 ensure that some APIs (ibuprofen in this case) meet their downstream quality standards, assuming no
486 upstream contribution to background concentrations. This could entail treatment at up to 890 WwTW
487 to meet current PNECs.

488

489 Much in relation to the need for future action by dischargers depends on whether or not these
490 substances are regulated and the water quality standard chosen, but if the CIP estimated PNECs are a
491 guide to regulatory limits, then there is potential for localised non-compliance in surface waters; at
492 least in the case of ethinyloestradiol, diclofenac, ibuprofen, propranolol and the macrolide antibiotics.
493 Further monitoring of pharmaceuticals in surface waters to determine the temporal variations in river
494 concentrations associated with changing river flows (and hence dilution), the persistence, and the
495 bioavailability of APIs needs to be considered.

496

497 **Acknowledgements**

498 The authors wish to thank the co-ordinator of the CIP programme – UK Water Industry Research
499 (UKWIR) for authorising the use of the information reported here, and the UK Water Utility
500 companies Anglian, Dwr Cymru, Northumbrian, Scottish, Severn Trent, Southern, South West,
501 Thames, United Utilities, Wessex and Yorkshire Water for their considerable efforts in generating it.

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503

504 **REFERENCES**

505 Ashton, D., Hilton M. And Thomas K. V. Investigating the environmental transport of human
506 pharmaceuticals to streams in the United Kingdom. *Sci Total Environ.* 2004;333:167-184.

507
508 Astra Zeneca (2016) Environmental Risk Data Relating to our medicine, accessed, 21/3/17.
509 [https://www.astrazeneca.com/content/dam/az/our-](https://www.astrazeneca.com/content/dam/az/our-company/Sustainability/Environmental_risk_data_relating_to_our_medicines.pdf)
510 [company/Sustainability/Environmental risk data relating to our medicines.pdf](https://www.astrazeneca.com/content/dam/az/our-company/Sustainability/Environmental_risk_data_relating_to_our_medicines.pdf)

511
512 Bound, J.P., and Voulvoulis, N. Predicted and measured concentrations for selected pharmaceuticals
513 in UK rivers: implications for risk assessment. *Water Res* 2006; 40: 2885-2892.

514
515 Boxall, A. B. A., Keller V.D.J. Straub J.O., Monteiro S.C., Fussell R. and Williams R.J. Exploiting
516 monitoring data in environmental exposure modelling and risk assessment of pharmaceuticals.
517 *Environ. Internat.* 2014;73:176-185.

518
519 Bradley, P.M., Journey, C.A., Button, D.T., Carlisle, D.M., Clark, J.M., Mahler, B.J., Nakagaki, N.,
520 Qi, S.L., Waite, I.R., VanMetre, P.C. Metformin and other pharmaceuticals widespread in Wadeable
521 streams of the southeastern United States. *Environ. Sci. Technol. Lett.* 2016;3:243-249.

522
523 Burns, E.E., Thomas-Oates, J., Kolpin, D.W., Furlong, E.T., Boxall, A.B.A. Are exposure predictions,
524 used for prioritization of pharmaceuticals in the environment, fit for purpose? *Environ. Toxicol.*
525 *Chem.* 2017;in press.

526
527 Clara, M., Kreuzinger N., Strenn B., Gans O. And Kroiss H. The solids retention time—a suitable
528 design parameter to evaluate the capacity of wastewater treatment plants to remove micropollutants.
529 *Water Res.* 2005;39:97-106.

530
531 Comber, S., Gardner, M., Georges, K., Thornton A. (2007) Dangerous Substances and Priority
532 Hazardous Substances/Priority Substances under the Water Framework Directive (07/WW/17/7). UK
533 Water Industry Research (UKWIR), 1 Queen Anne's Gate, London, UK. ISBN: 1 84057 464 X.

534
535 Comber S., Smith R., Daldorph P., Gardner M., Constantino C. and Ellor B. Development of a
536 chemical source apportionment decision support framework for catchment management. *Environ Sci*
537 *Technol.* 2013;47:9824-9832.

538
539 Comber S., Gardner M., Jones, V. and Ellor B. Source Apportionment of Trace Contaminants in
540 Urban Sewer Catchments. *Environ Technol.* 2014;36(5):573-587.

541
542 Deegan, A.M., Shaik B., Nolan K., Urell K., Oelgemöller M., Tobin J., Morrissey A. Treatment
543 options for wastewater effluents from pharmaceutical companies. *Int J Environ Sci Technol.* 2011;
544 8:649–666.

545
546 EC, 2009: Technical Specifications for Chemical Analysis and Monitoring of Water Status. Directive
547 2009/90/EC.

548
549 EU (2011) European Union Technical Report - 2011 – 055. Common Implementation Strategy
550 for the Water Framework Directive (2000/60/EC). Guidance Document No. 27 Technical Guidance
551 For Deriving Environmental Quality Standards. ISBN : 978-92-79-16228-2. DOI : 10.2779/43816.

552
553 EU (2013) European Union Directive 2013/39/EU of the European Parliament and of the Council of
554 12 August 2013 amending Directives 2000/60/EC and 2008/105/EC as regards priority substances in
555 the field of water policy.

556

557 Falås, P., Andersen H.R., Ledin A., and La Cour Jansen J. Occurrence and reduction of
558 pharmaceuticals in the water phase at Swedish wastewater treatment plants. *Water Sci Technol.*
559 2012;66(4):783-791.
560

561 Furlong, E.T., Batt, A.L., Glassmeyer, S.T., Noriega, M.C., Kolpin, D.W., Mash, H., Schenck, K.M.
562 Nationwide reconnaissance of contaminants of emerging concern in source and treated drinking
563 waters of the United States: Pharmaceuticals. *Sci. Total Environ.* 2017;579:1629-1642.
564

565 Gardner M., Comber S., Scrimshaw M., Cartmell E., Lester J. and Ellor B. The significance of
566 hazardous chemicals in wastewater treatment works effluents. *Sci Total Environ.* 2012;437:363-372.
567

568 Gardner M., Jones, V., Comber S., Scrimshaw M., Coello-Garcia, T., Cartmell E., Lester J. and Ellor
569 B. Performance of UK wastewater treatment works with respect to trace contaminants. *Sci Total*
570 *Environ.* 2013;456-457:359-369.
571

572 Heffley J., Comber S., Wheeler B. and Redshaw C. Developing a modelling approach to predict
573 pharmaceutical discharges from UK sewage treatment works using steroid estrogens as a case study.
574 *Environ Sci: Processes and Impacts* 2014;16:2571-2580.
575

576 Henze, M., *Biological wastewater treatment: principles, modelling and design.* IWA publishing,
577 2008.
578

579 Hope, B.K., Pillsbury L., and Boling B A state-wide survey in Oregon (USA) of trace metals and
580 organic chemicals in municipal effluent. *Sci Total Environ* 2012;417:263-272.
581

582 IMS (2016) Quantiles IMS (Intercontinental Marketing Services) Data for active ingredient sales.
583 <http://www.imshealth.com/>; accessed 24th May 2016.
584

585 Jelic, A, Gros M., Ginebreda A., Cespedes-Sanchez R., Ventura F., Petrovic M. and Barcelo D.
586 Occurrence, partition and removal of pharmaceuticals in sewage water and sludge during wastewater
587 treatment. *Water Res.* 2011;45(3):1165-1176.
588

589 Johnson A., Dumont E., Williams R., Oldenkamp R., Cisowska I. and Sumpter J. Do concentrations
590 of ethinylestradiol, estradiol, and diclofenac in European rivers exceed proposed EU Environmental
591 Quality Standards? *Environ. Sci. Tech.* 2013a;47(21):12297-12304.
592

593 Johnson A., Oldenkamp R., Dumont E. and Sumpter J. Predicting concentrations of the cytostatic
594 drugs cyclophosphamide, carboplatin, 5-fluorouracil, and capecitabine throughout the sewage
595 effluents and surface waters of Europe. *Environ. Tox. And Chem.* 2013b;32(9):1954-1961.
596

597 Johnson A., Keller V., Dumont E. and Sumpter J. Assessing the concentrations and risks of toxicity
598 from the antibiotics ciprofloxacin, sulfamethoxazole, trimethoprim and erythromycin in European
599 rivers. *Sci of the Tot. Environ.*, 2015, 747-755.
600

601 Jones, V., Gardner M. and Ellor B. Concentrations of trace substances in sewage sludge from 28
602 wastewater treatment works in the UK. *Chemosphere* 2014;111:478-484.
603

604 Jobling, S., Williams R., Johnson A., Taylor A., Gross-Sorokin M., Nolan M., Tyler C., van Aerle R.,
605 Santos E. and Brighty G. Predicted exposures to steroid estrogens in UK rivers correlate with
606 widespread sexual disruption in wild fish populations. *Environ Health Persp.* 2005;114:32-39.
607

608 JRC (2015) Ibuprofen Dossier, Draft Version 3.
609

610 JRC (2015) Summary Dossier Review, Draft.
611

612 JRC (2016) Development of the first Watch List under the Environmental Quality Standards
613 Directive. Joint Research Centre, Report EUR 27142 EN, by Raquel N. Carvalho, Lidia Ceriani,
614 Alessio Ippolito and Teresa Lettieri, 2015, Accessed 21/3/17
615 <http://publications.jrc.ec.europa.eu/repository/bitstream/JRC95018/lbna27142enn.pdf>
616
617 Kasprzyk-Hordern, B., Dinsdale R.M. and Guwy A.J. The occurrence of pharmaceuticals, personal
618 care products, endocrine disruptors and illicit drugs in surface water in South Wales, UK. *Water Res*
619 2008;42(13): 3498-3518.
620
621 Kolpin, D.W., Furlong, E.T., Meyer, M.T., Thurman, E.M., Zaugg, S.D., Barber, L.B., and Buxton,
622 H.T. Pharmaceuticals, hormones, and other organic wastewater contaminants in U.S. streams, 1999-
623 2000: A national reconnaissance. *Environ. Sci. Technol.* 2002;36:1202-1211.
624
625 Le-Minh N., Khan S.J., Drewes J.E., Stuetz R.M. Fate of antibiotics during municipal water recycling
626 treatment processes. *Water Res.* 2010; 44(15):4295–4323.
627
628 Lees K., Fitzsimons M., Tappin A., Snape J. and Comber S. Pharmaceuticals in soils of lower income
629 countries: physico-chemical fate and risks from wastewater irrigation. *Environ. Int.* 2016; .94:712-
630 723.
631
632 Levado R., Thibaut R., Raldua D, Martin R., and Porte C. First evidence of endocrine disruption in
633 feral carp from the Ebro River. *Tox. And Appl. Pharma.* 2004;196(2):247-257.
634
635 Li, C., Cabassud C. and Guigui C. Evaluation of membrane bioreactor on removal of pharmaceutical
636 micropollutants: a review. *Desal and Water Treat* 2015; 55(4):845-858.
637
638 Melvin, S.D. and Leusch F.D.L. Removal of trace organic contaminants from domestic wastewater: A
639 meta-analysis comparison of sewage treatment technologies. *Environ International* 2016; 92: 183-188.
640
641 Murray-Smith, R.J. Coombe, V.T., Grönlund, M.H., Waern F., Baird J.A. Managing emissions of
642 active pharmaceutical ingredients from manufacturing facilities: An environmental quality standard
643 approach. *Int. Env. Ass. and Man.* 2012;8(2):320–330.
644
645 NSF (2016) National Science Foundation, Water and Environmental Technology Centre,
646 Pharmaceutical PNEC list. Accessed 21/3/17. [http://www.nsfwetcenter.org/wp-](http://www.nsfwetcenter.org/wp-content/uploads/2016/10/WET-Center-Phamaceutical-PNEC-list-3.pdf)
647 [content/uploads/2016/10/WET-Center-Phamaceutical-PNEC-list-3.pdf](http://www.nsfwetcenter.org/wp-content/uploads/2016/10/WET-Center-Phamaceutical-PNEC-list-3.pdf)
648
649 Phillips, P.J., Smith, S.G., Kolpin, D.W., Zaugg, S.D., Buxton, H.T., Furlong, E.T., Esposito, K.,
650 Stinson, B. Pharmaceutical formulation facilities as sources of opioids and other pharmaceuticals to
651 wastewater-treatment-plant effluents. *Environ. Sci. Technol.* 2010;44:4910-4916.
652
653 Phillips, P.J., Schubert, C., Argue, D., Fisher, E., Furlong, E.T., Foreman, W., Gray, J., Chalmers, A.
654 Concentrations of hormones, pharmaceuticals and other micropollutants in groundwater affected by
655 septic systems in New England and New York. *Sci. Total Environ.* 2015;512-513:43-54.
656
657 Rowett C., Comber S. and Hutchinson T. The impact of natural and anthropogenic Dissolved Organic
658 Carbon (DOC), and pH on the toxicity of triclosan to the crustacean *Gammarus pulex* (L.). *Sci Total*
659 *Environ.* 2016;565:222-231.
660
661 Singer, Andrew C., Järhult J.D., Grabic R., Khan G.A., Lindberg R.H., Fedorova G., Fick J., Bowes,
662 M.J., Olsen B. and Söderström H. Intra-and inter-pandemic variations of antiviral, antibiotics and
663 decongestants in wastewater treatment plants and receiving rivers. *PLoS One* 9.9 2014;9(9):108621.
664

665 Stockholm Vatten, 2010. Läkemedelsrester i Stockholms vattenmiljö Förekomst, förebyggande
666 åtgärder och rening av avloppsvatten
667 www.stockholmvatten.se/globalassets/pdf1/rapporter/avlopp/avloppsrening/lakemedelsrapport_slutra
668 [pport.pdf](http://www.stockholmvatten.se/globalassets/pdf1/rapporter/avlopp/avloppsrening/lakemedelsrapport_slutra) (accessed sept, 2016)
669
670 Thompson, M. and Ellison, S.L.R. Towards an uncertainty paradigm of detection capability,
671 *Analytical Methods* 2013;5:5857-61.
672
673 Tyler C., Jobling S. and Sumpter J. Endocrine disruption in wildlife: a critical review of the evidence.
674 *Crit. Rev. in Toxicol.* 2008;28:319-361.
675
676 UKWIR (2014) Risk based Prioritisation of Pharmaceuticals UKWIR Report 14/WW/17/16 (August
677 2014) ISBN: 1840577355.
678
679 Verlicchi P, Al Aukidy M, Zambello E. Occurrence of pharmaceutical compounds in urban
680 wastewater: removal, mass load and environmental risk after a secondary treatment-a review. *Sci*
681 *Total Environ.* 2012;429:123-55.
682
683 Wahlberg, C., Björlenius B. and Paxéus N. Fluxes of 13 selected pharmaceuticals in the water cycle
684 of Stockholm, Sweden. *Water Sci and Technol.* 2011;63(8):1772-1780.
685
686 WHO, 2011: Pharmaceuticals in Drinking-water, Public Health and Environment Water, Sanitation,
687 Hygiene and Health WHO/HSE/WSH/11.05
688 www.who.int/water_sanitation_health/publications/2011/pharmaceuticals_20110601.pdf (accessed
689 Sept, 2016)
690
691 Zhou, J. L., Zhang Z.L., Banks E., Grover D. and Jiang J.Q. Pharmaceutical residues in wastewater
692 treatment works effluents and their impact on receiving river water." *J Hazard Mat.* 2009;166(2):655-
693 661.
694
695 Zorita S., Mårtensson L. and Mathiasson L., Occurrence and removal of pharmaceuticals in a
696 municipal sewage treatment system in the south of Sweden. *Sci Total Environ.* 2009;407(8): 2760–
697 2770.
698