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Spatial and temporal occurrence of pharmaceuticals in UK estuaries

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Abstract

There is a lack of data on the occurrence of pharmaceuticals in estuaries worldwide, with little understanding of their temporal and spatial variations globally. Ibuprofen, paracetamol, diclofenac, trimethoprim and citalopram were measured in twelve estuaries in the UK. Initially, these compounds were monitored in the Humber Estuary, where samples were taken every two months over a twelve month period in order to assess their spatial and temporal variations. Ibuprofen was found at some of the highest concentrations ever measured in an estuary globally (18 – 6297ng l⁻¹), with paracetamol also measured at relatively high concentrations (4 – 917 ng l⁻¹) in comparison to the other compounds. In terms of spatial distribution, a pattern was observed where the highest concentrations were found at a site at which wastewater is discharged, whilst compound concentrations were often lower upstream and downstream of this site. The downstream profile of pharmaceuticals differed temporally with concentrations highest downstream when input from wastewater effluent was highest. Eleven further estuaries were sampled around the UK in order to put the occurrence of pharmaceuticals in the Humber Estuary into a wider context. Pharmaceutical concentrations in the other estuaries sampled were less than 210 ng l⁻¹ but, again, ibuprofen and paracetamol were found at concentrations higher than other compounds, whereas diclofenac and citalopram were absent in many estuaries. The

29 Humber, which is the receiving environment for the sewage effluent of approximately 20%
30 (13.6 million people) of the population of England, was observed to have the highest overall
31 concentration of pharmaceuticals in contrast to the other estuaries sampled, thereby
32 representing a worst case scenario for pharmaceutical pollution.

33 Keywords: Pharmaceuticals; Emerging Contaminants; Estuary; Occurrence; Temporal
34 Distribution; Spatial Distribution

35 **1. Introduction**

36 Despite the extensive and long-term use of pharmaceuticals, it has only been in the past few
37 decades that interest in pharmaceutical pollution has gained popularity and now hundreds of
38 pharmaceuticals have been detected in the aquatic environment (Hughes et al. 2013; Gaw
39 et al. 2014). Their presence is sustained through continuous input from wastewater
40 treatment plants (WWTPs), as well as from improper disposal, agriculture and aquaculture
41 (Godoy et al. 2015). Pharmaceuticals are designed to be biologically active, often at low
42 levels, and their presence in surface water has led to concern over their potential biological
43 effect (Santos et al. 2010). Many pharmaceuticals (e.g. diclofenac and fluoxetine) have been
44 found to illicit a negative response on biota in laboratory exposures at concentrations similar
45 to those found in the aquatic environment (Eades and Waring 2010; Franzellitti et al. 2013;
46 Miguez et al. 2016).

47 The fate of pharmaceuticals is best understood in the freshwater environment, with input,
48 environmental conditions, biological degradation and sediment-related processes playing a
49 prominent role in their spatial and temporal distribution (Li 2014). Pharmaceuticals often
50 show a decline in concentration downstream from input sources as the result of dilution,
51 degradation and partitioning to sediment (Kunkel and Radke 2012). However, due to the
52 prevalence of WWTPs, this leads to the continuous input of pharmaceuticals into the
53 environment. As a result, these processes are not enough to sufficiently remove compounds
54 leading to their high detection in the aquatic environment and, potentially, transportation into
55 estuaries and coastal waters (Ebele et al. 2017).

56 Estuaries are receiving waters, often for many rivers, acting as a confluence for
57 contaminants and therefore increasing the potential risk of pharmaceutical pollution in these
58 environments (Ridgway and Shimmiel 2002). Estuaries are ecologically important to
59 ecosystem services, providing habitat for many species and acting as an area for recreation
60 and transport (Ridgway and Shimmiel 2002). Despite this, few studies have measured the
61 occurrence of pharmaceuticals in estuaries, and those that do exist typically lack the
62 resolution to determine spatial and temporal patterns (Table 1). Studies which have
63 investigated the spatial and temporal patterns of pharmaceuticals are often locally focused,
64 monitoring only one estuary (for example Tamtam et al., 2012; Hedgespeth et al. 2012;
65 Cantwell et al. 2017), and it is important to determine if any patterns seen are relevant at a
66 wider scale and represent a risk to the environment.

67 This study aimed to further contribute to the overall picture of pharmaceutical contamination
68 in estuaries. Five target compounds, ibuprofen, paracetamol, diclofenac, trimethoprim and
69 citalopram, were chosen for the present study, based on their prevalent usage and predicted
70 risk to the aquatic environment (National Health Service 2017; Roos et al. 2012). To the
71 authors' knowledge, citalopram has not previously been monitored in the estuarine
72 environment (Table 1). Moreover, monitoring of the aforementioned compounds is limited,
73 with some of these measurements dating back almost fifteen years. The target compounds
74 were measured every other month over a twelve month period at various sites in the Humber
75 Estuary to determine their spatial and temporal occurrence. In addition, eleven further
76 estuaries, located in other parts of the UK, were selected in order to determine whether
77 concentrations observed in the Humber were representative of other estuaries.

78 **Table 1:** Maximum concentrations of ibuprofen, paracetamol, diclofenac and trimethoprim detected in
 79 estuaries globally (ng l⁻¹). Citalopram has not previously been monitored in any estuaries.

Region	Estuary	Ibuprofen	Paracetamol	Diclofenac	Trimethoprim	Reference
Asia	Jiulong, China	21	13	11		Sun et al. (2016)
	Hailing Bay, China				37	Chen et al. (2015)
	Qinzhou Bay, China			7		Cui et al. (2019)
	Yangtze, China			<MDL		Yang et al. (2011)
	Yangtze, China				330	Zhang et al. (2012)
	Yangtze, China		<MDL			Zhao et al. (2015)
Europe	Seine, France				45	Tamtam et al. (2008)
	Elbe, Germany	1		1		Weigel et al. (2002)
	Arade, Portugal	28	88	31		Gonzalez-Rey et al. (2015)
	Douro, Portugal				16	Madureira et al. (2010)
	Tejo, Portugal	<MDL	11	52	8	Reis-Santos et al. (2016)
	Bilbao, Spain		440	650	2046	Mijangos et al. (2018)
	Plentzia, Spain		49	22	6	Mijangos et al. (2018)
	Urdaibai, Spain		321	35	3	Mijangos et al. (2018)
	Belfast Lough, UK	376	<MDL	<MDL	32	Thomas and Hilton (2004)
	Mersey, UK	386	<MDL	195	569	Thomas and Hilton (2004)
	Tees, UK	88	<MDL	191	17	Thomas and Hilton (2004)
	Thames, UK	928	<MDL	125	<MDL	Thomas and Hilton (2004)
Thames, UK				19	Munro et al. (2019)	
Tyne, UK	755		90	46	Thomas and Hilton (2004)	
North America	Charleston Harbour, USA	8	28			Hedgespeth et al. (2012)
America	Jamaica Bay, USA	38	156		125	Benotti and Brownawell (2007)
	Narragansett Bay, USA		60		18	Cantwell et al. (2017)

	New York Bay, USA	162	14	Cantwell et al. (2018)
	San Francisco, USA		4	Klosterhaus et al. (2013)
Oceania	Sydney, Australia	31		Birch et al. (2015)

80

81 **2. Methods**

82 **2.1 Study Area**

83 The Humber Estuary is a macrotidal estuary located in Yorkshire, on the East Coast of
84 England, UK (Figure 1). It is 303 km², has an average depth of 6.5 m and is the confluence
85 of the Rivers Ouse, Trent and Hull which pass through some of the largest urban areas in
86 the UK, thus it is the receiving water for approximately 20% of UK effluent (European
87 Environment Agency, 2019; Table 2). Samples were collected from nine sites along a 65 km
88 stretch on the North side of the estuary (Figure 1). Two of these were located in the River
89 Ouse; A1 was the furthest upstream and A2 was located less than 1km upstream from the
90 confluence with the Humber Estuary. The furthest site upstream in the Humber Estuary (R1)
91 was the receiving site for effluent from Melton WWTP, which serves a population equivalent
92 (PE) of 12,255 (European Environment Agency, 2017). Three sites (R2-R4) were positioned
93 every 2 km downstream from R1. Three final sites (A3-A5) were located 20km from R1 in the
94 lower estuary and 15 km from the mouth. Further information on site location can be found in
95 supplementary material S1. The Humber Estuary is an important site for conservation and
96 has been designated as a Special Protection Area (SPA), also containing a Special Area of
97 Conservation (SAC). It is also a vital habitat for many species of international importance,
98 providing habitat for 4.1% of the red knot (*Calidris canutus*) and 5.7% of the common
99 redshank (*Tringa tetanus*) international populations, and as a result has also been
100 designated as a RAMSAR site (Buck et al. 1997)

101 Samples were also collected from eleven further estuaries which encompassed a range of
102 estuary types, tidal ranges and sizes (Table 2). The total PE was calculated for the WWTPs

103 in the catchment area of each estuary (Table 2); further information on the proximity of
 104 WWTPs to the sampling sites in each estuary can be found in supplementary material S2.
 105 Many of these estuaries have been designated as SACs, SPAs and RAMSAR sites as the
 106 result of the sensitive and important species resident to them.

107 **Table 2:** Information on the type and size of estuaries sampled (Davidson et al.1991).
 108 Information on the number of WWTPs and the population equivalent served in 2014 was
 109 calculated from an interactive wastewater treatment map (European Environment Agency
 110 2019).

Estuary	Type	Estuary Area (km²)	Tidal Type	Number of WWTPs in Catchment	Total PE (000s)
Cromarty	Complex	92.3	Mesotidal	3	15.6
Forth	Complex	84.0	Macrotidal	33	1 613.3
Humber	Coastal Plain	303.6	Macrotidal	304	13 674.7
Mersey	Coastal Plain	89.1	Macrotidal	30	3 689.7
Portsmouth	Ria	15.9	Macrotidal	2	383
Severn	Coastal Plain	556.8	Macrotidal	171	6 724.4
Solway	Complex	420.6	Macrotidal	20	314.9
Tay	Complex	121.3	Mesotidal	12	167.6
Tees	Coastal Plain	13.5	Macrotidal	9	844.9
Thames	Coastal Plain	46.5	Macrotidal	198	16 510.5
Tyne	Complex	7.9	Macrotidal	6	1 092.8
Ythan	Barbuilt	2.8	Mesotidal	1	11.2

111

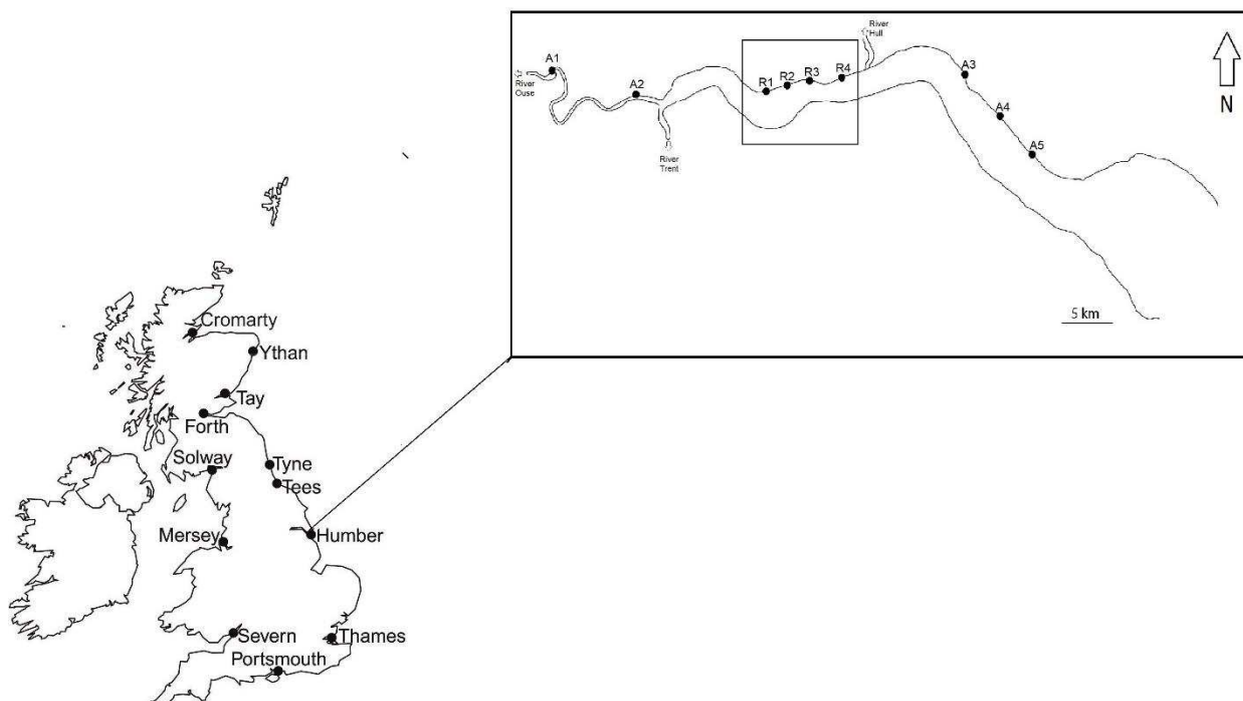
112 **2.2 Sampling**

113 2.2.1 Seasonal Monitoring

114 Sampling was carried out in the Humber Estuary, UK, every two months from October 2016
115 to August 2017 at sites R1-R4 (Figure 1). Samples were also collected from four additional
116 sites (A1-A2 and A4-A5) in October, February and June, and a further site (A3) in February
117 and June (Figure 1). Sampling was carried out during a high neap tide (\pm 3 hours) to
118 minimise differences in diurnal concentrations as the result of tides (Lara-Martin et al. 2014).
119 At each site, 3 x 1 L of surface seawater were collected in amber glass bottles and
120 temperature, pH and dissolved oxygen determined using a HACH meter and salinity (0 – 27
121 ppt) measured with a refractometer (supplementary material S1). Water samples were kept
122 on ice or in the fridge at 4 °C and extracted within 48 hours for analysis of pharmaceuticals.

123 2.2.2 UK Wide Monitoring

124 Sampling was carried out in August and September 2017 during high tides (\pm 3 hours) in
125 eleven additional UK estuaries in order to provide a wider context for the concentrations of
126 pharmaceuticals seen in the Humber Estuary (Figure 1). Within each estuary, sites were
127 chosen in the upper, middle and lower parts of the estuary and 1 L of water was collected at
128 each of these in amber glass bottles. Temperature, pH, dissolved oxygen and salinity (0-34
129 ppt) were determined as above and samples stored and extracted in the same manner
130 (supplementary material S2).



131

132 **Figure 1** Map of field sites for seasonal and UK wide monitoring of selected
 133 pharmaceuticals. The sites in the box (R1-R4) indicate those which were sampled every two
 134 months whilst A1-A2 and A3-A5 were sampled every four months.

135

136 2.3 Chemical Analysis

137 2.3.1 Study Compounds

138 Five study compounds, ibuprofen, paracetamol, diclofenac, trimethoprim and citalopram,
 139 were chosen for monitoring (Table 3) and standards of diclofenac sodium (≥ 98.5),
 140 acetaminophen ($\geq 99\%$), citalopram (≥ 98), ibuprofen ($\geq 98\%$), and trimethoprim ($\geq 98\%$) were
 141 supplied by Sigma-Aldrich Ltd. (Dorset, UK).

142 **Table 3:** Physico-chemical characteristics of the study compounds. Physico-chemical data obtained
 143 from Alygizakis et al. (2016), Bayen et al. (2013) and Kasprzyk-Hordern et al. (2007). Prescription
 144 data obtained from National Health Service (2019); supplementary material S3.

Compound	Therapeutic Use	Prescriptions (kg year ⁻¹)	Water Solubility (mg l ⁻¹)	Log _{KOW}	Molecular Weight	pKa
Ibuprofen	Nonsteroidal anti-inflammatory drug (NSAID)	82,756	41.05	3.97	206.29	9
Paracetamol	Painkiller	2,169,244	22.7	0.9	151.16	9.9
Diclofenac	NSAID	5459	4.52	4.51	296.15	4.2
Trimethoprim	Antibiotic	8444	171.1	1.4	290.32	7.1
Citalopram	Antidepressant	9204	4.02	3.74	324.39	9.4

145

146 2.3.2 Solid Phase Extraction

147 A composite sample was made by combining the 3 x 1L surface water samples collected
 148 from each site during seasonal monitoring, or from each of the estuaries during the UK-wide
 149 survey, by adding them together in a 5 L beaker and stirring vigorously for two minutes. A
 150 500 mL sub-sample was filtered through a 0.45 µm cellulose filter (Scientific Laboratory
 151 Supplies, Hessle, UK) under vacuum. Solid phase extraction was performed on the filtered
 152 water samples using Oasis HLB cartridges (Waters Corporation, Massachusetts, USA),
 153 which were conditioned with 5 mL 100% methanol followed by 5 mL deionised water at a
 154 rate of 1 mL min⁻¹. The sample was loaded on to the cartridge at a rate of 10 mL min⁻¹,
 155 during which care was taken not to let the sorbent material dry out. The cartridges were then

156 rinsed with 5 mL deionised water and the sorbent was dried under vacuum for 15 minutes to
157 remove excess water prior to elution. Elution was performed with 2 x 5 mL 0.1% trifluoroacetic
158 acid in methanol. The eluent was evaporated to dryness using a rotary evaporator (40°C,
159 speed 7) and reconstituted with methanol: water (10:90).

160 SPE recovery was evaluated by spiking known concentrations (100, 200, and 1000 ng l⁻¹) of
161 all study compounds into three replicates each of artificial seawater made up to 20 ppt in
162 deionised water (supplementary material S4). The mean recovery across all concentrations
163 was used to correct the measured environmental concentrations (Table 4).

164 **Table 4:** Mean method detection limits (\pm standard deviation), mean method quantification levels (\pm
165 standard deviation) and mean recovery (\pm standard deviation) of target compounds.

Compound	MDL (ng l ⁻¹)	MQL (ng l ⁻¹)	Recovery (%)
Citalopram	0.34 (0.25)	1.18 (0.85)	43 (5.5)
Diclofenac	1.77 (1.35)	5.91 (4.49)	20 (11.0)
Ibuprofen	1.45 (0.41)	4.83 (1.38)	73 (34.0)
Paracetamol	3.28 (1.82)	10.93 (6.07)	86 (34.1)
Trimethoprim	0.07 (0.04)	0.24 (0.12)	63 (10.6)

166

167 2.3.3 UltraperformanceTM-ESI-(QqLIT) MS/MS analysis

168 Analysis was carried out according to Gros et al. (2012). Briefly, chromatographic
169 separations were performed with a Waters Acquity Ultra-Performance liquid chromatograph
170 system equipped with two binary pump systems (Milford, Massachusetts, USA) and coupled
171 to a 5500 QTRAP hybrid quadrupole-linear ion trap mass spectrometer with a turbo ion
172 spray source (Applied Biosystems, Foster Systems, Foster City, CA, USA). Citalopram and
173 trimethoprim were analysed under positive electrospray ionisation (PI) using an Acquity HSS
174 T₃ column (50 mm x 2.1 mm, 1.8 μ m particle size) and ibuprofen, paracetamol and diclofenac
175 were analysed under negative ion (NI) electrospray using an Acquity BEH C₁₈ column (5 mm
176 x 2.1 mm, 1.7 μ m particle size), both from Waters Corporation.

177 All data acquisition was performed in Analyst 2.1 software. Quantification of analytes was
178 performed by selective reaction monitoring (SRM), monitoring two transitions for each
179 compound as described in Gros et al. (2012). Method detection limits (MDL) and
180 quantification levels (MQL) were determined for each of the compounds based on a signal-
181 to-noise ratio of 3 and 10 respectively (Table 4).

182

183 **2.4 Statistical Analysis**

184 Statistical analysis was performed in R 3.3.1. In order to determine if there was a difference
185 in the occurrence of pharmaceuticals between sampling months, concentrations from R1-4
186 were grouped together, as these sites were sampled during all of the sampling periods. A
187 Friedman's Test followed by a Nemenyi post-hoc test were conducted using the PMCMR
188 package (Pohlert 2014). All data is presented in graphs created by the ggplot2 package
189 (Wickham 2016).

190

191 **3. Results**

192 **3.1 Humber Estuary**

193 Pharmaceuticals were frequently detected (58 - 97% of samples for individual study
194 compounds) in the Humber Estuary (Table 5) and concentrations followed the order of
195 ibuprofen>paracetamol>diclofenac>trimethoprim>citalopram. Whilst mean concentrations
196 were in the order of 100 ng l⁻¹ or below, maximum concentrations were approximately five to
197 ten times higher (Table 5; supplementary material S5). Maximum levels of ibuprofen and
198 paracetamol detected in the Humber are the highest concentrations reported in estuaries to
199 date (Table 1).

200

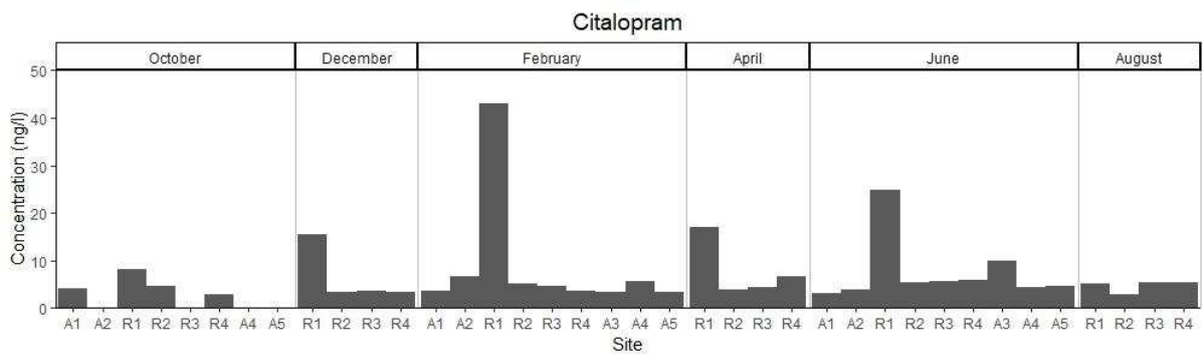
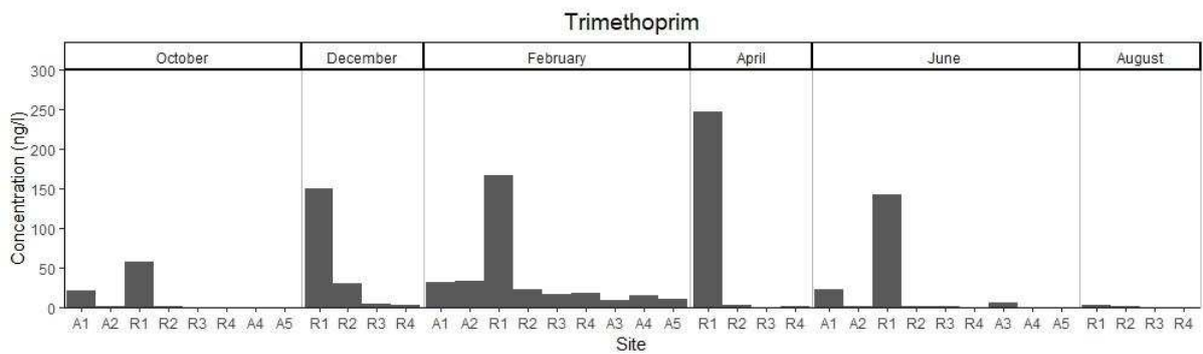
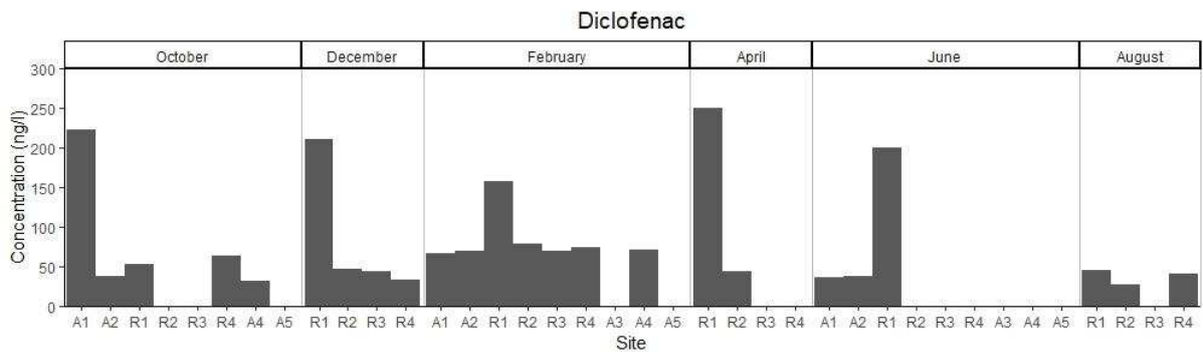
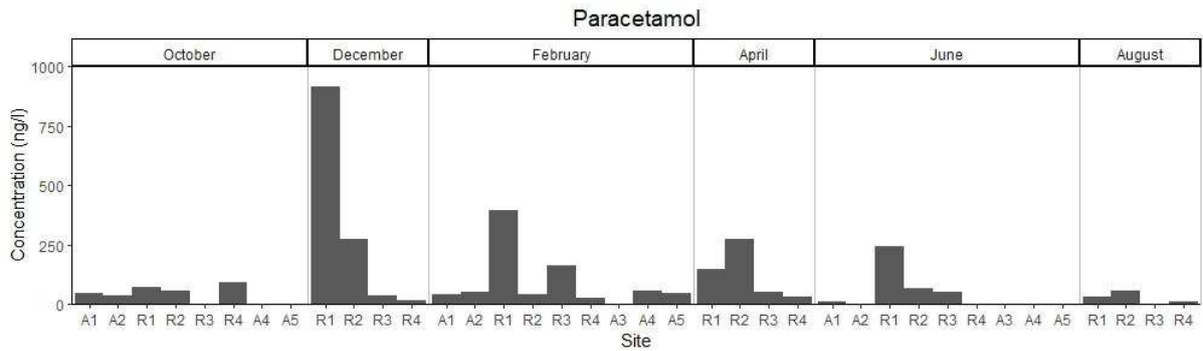
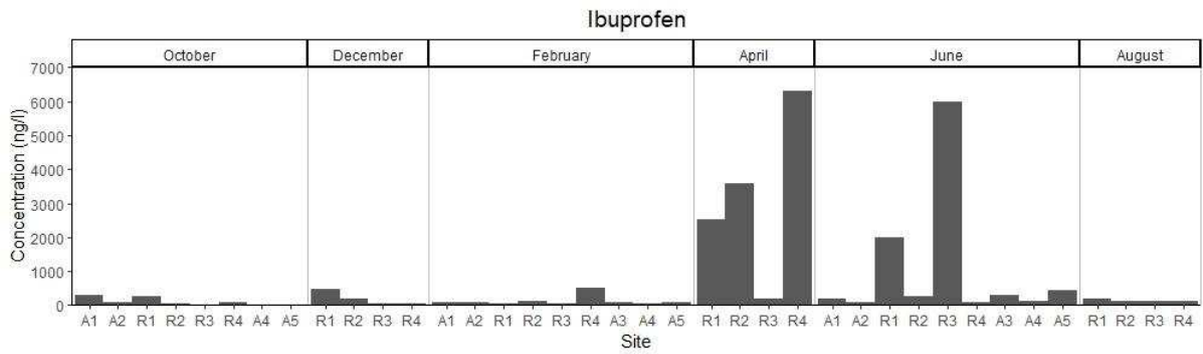
201 **Table 5:** Pharmaceutical concentrations (ng l⁻¹) in the Humber Estuary (n=38) during a 12 month
 202 sampling campaign. Values were corrected based on mean recovery values (Table 3). Max =
 203 maximum concentration, SD = standard deviation. Detection rate is the amount of samples above the
 204 method quantification limit (MQL).

Compound	Detection Rate	Max	Mean	SD
	(%)	(ng l⁻¹)	(ng l⁻¹)	
Ibuprofen	97.37	6297.14	665.58	1481.49
Paracetamol	73.68	916.88	88.65	163.66
Diclofenac	57.89	250.8	51.44	68.29
Trimethoprim	92.11	247.02	27.43	54.56
Citalopram	89.47	42.93	6.39	7.66

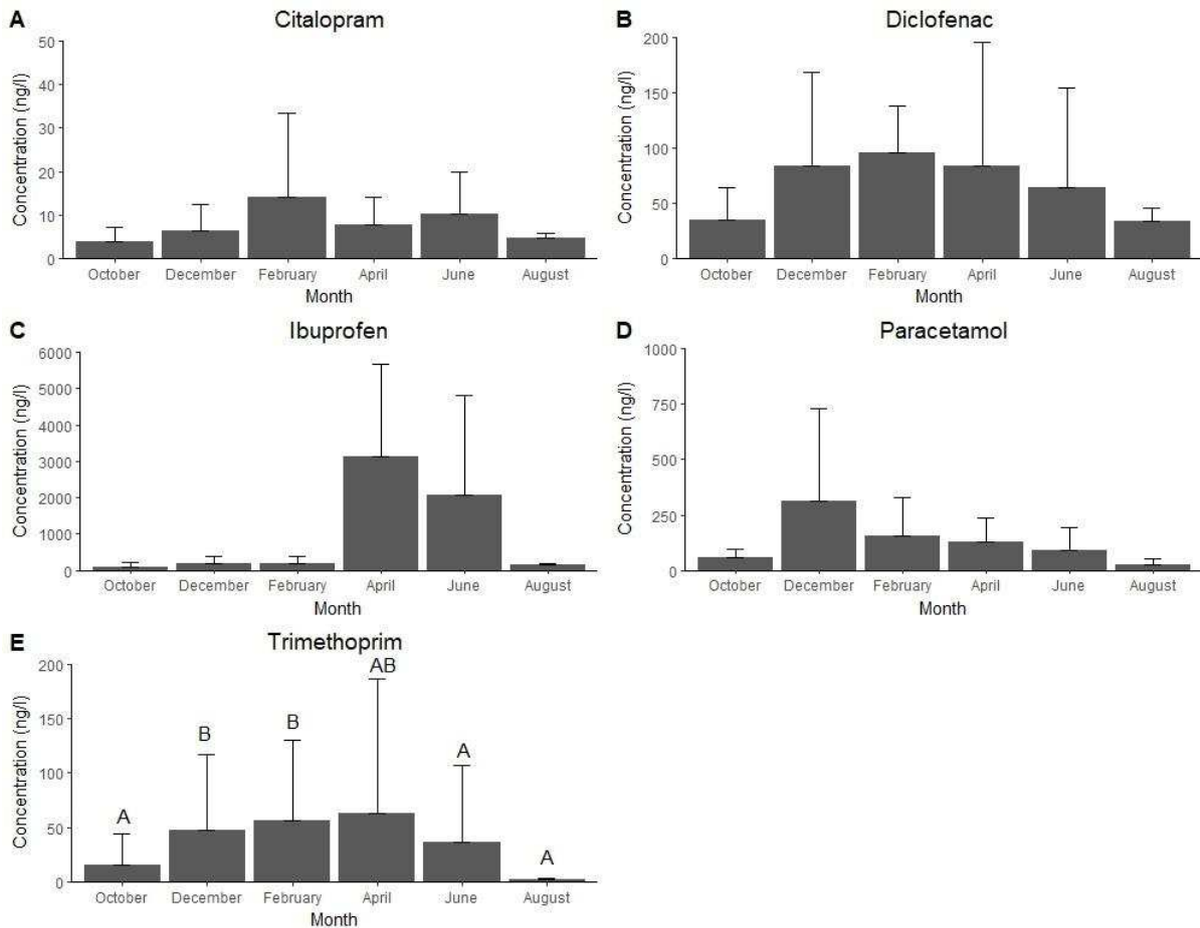
205
 206 A general pattern was observed in the occurrence of pharmaceuticals in the Humber with
 207 concentrations peaking at sampling site R1 (Figures 2) and those up (samplings sites A1-
 208 A2) and downstream (sampling sites R2-A5) of this site similar to each other. However, this
 209 pattern was not entirely consistent and sometimes concentrations declined downstream (A3-
 210 A5). Maximum concentrations were generally seen at sampling site R1 although during
 211 some of the sampling periods they occurred at sites R2-R4.

212 Of the three months where all sites were sampled February had the highest detection rates
 213 and concentrations of pharmaceuticals at downstream sites (A3-A5), whilst many of the
 214 compounds were absent at these sites in October and June (Figure 2). In contrast, ibuprofen
 215 was an exception to this with concentrations found at these sites during all of the sampling
 216 periods. Citalopram also showed little decline in downstream concentrations in June and
 217 was present at A3-A5 at concentrations similar to or higher than many of the sites further
 218 upstream (Figure 2). There appeared to be a relationship between the concentration of
 219 pharmaceuticals at R1 and those seen at the other sites; typically, a higher concentration at
 220 R1 resulted in a higher presence at sites further downstream (Figure 2).

221 Trimethoprim was the only compound to show a statistically significant difference between
222 sampling months (Friedman's Test, chi-squared = 14.71, $p < 0.05$) with concentrations
223 significantly higher in winter (December and February; 3.29 – 166.54 ng l⁻¹) compared to
224 October and the summer months (June and August; 0 – 142 ng l⁻¹; Figure 3). Nevertheless,
225 the difference was almost significant for ibuprofen ($p = 0.054$) and citalopram ($p = 0.051$).
226 For citalopram, February had the highest concentrations (3.74 – 42.93 ng l⁻¹), whereas
227 ibuprofen concentrations were higher in April and June (186.37 – 6297.14 ng l⁻¹; Figures 3)
228 in comparison to the other sampling periods. All compounds were found at their lowest mean
229 concentrations in August (Figures 3), with no peaks seen at sampling site R1 (Figure 2).



231 **Figure 2** Concentrations of target analytes at nine sites in the Humber Estuary. Values were
 232 corrected based on mean recovery values (Table 3). Sites are listed from furthest upstream (A1) to
 233 furthest downstream (A5). R1-R4 were sampled every sampling event whilst the other sites were only
 234 sampled in October, February and June, except for A1 which was not sampled in October.



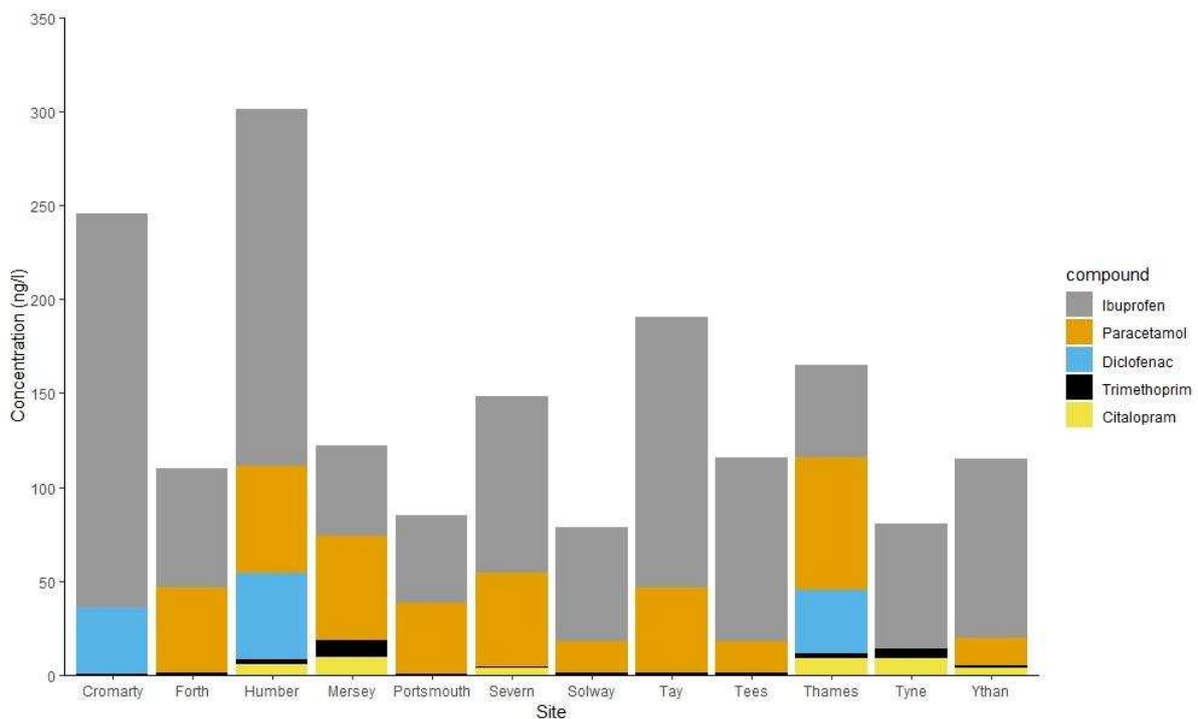
235
 236 **Figure 3** Mean bi-monthly concentrations (\pm one standard deviation) of (A) Ibuprofen (B) Paracetamol
 237 (C) Diclofenac (D) Trimethoprim and (E) Citalopram at the four sites monitored most frequently (R1-
 238 R4). Values were corrected based on mean recovery values (Table 3). Letters denote statistically
 239 significant difference (Friedman's Test).

240
 241 **3.2 UK-wide sampling**

242 Pharmaceuticals were detected in all of the estuaries sampled around the UK but only at
 243 concentrations in the low ng l⁻¹ range, generally lower than those detected in the Humber
 244 Estuary (Figure 4). Relative concentrations were similar to that found in the Humber;

245 ibuprofen>paracetamol>diclofenac>citalopram>trimethoprim (supplementary material S6).
 246 Ibuprofen and trimethoprim were present in all of the estuaries sampled whereas diclofenac
 247 was only detected in two of the other estuaries, the Cromarty and Thames (Figure 4). The
 248 Thames and Humber were the only estuaries to contain all of the compounds. The Humber
 249 had the overall highest concentration of pharmaceuticals and only the Cromarty and Tay had
 250 a total concentration of pharmaceuticals over 200 ng l⁻¹ (Figure 4).

251



252

253 **Figure 4** Concentrations of citalopram, diclofenac, ibuprofen, paracetamol and trimethoprim across
 254 eleven estuaries in the UK. Concentrations have been corrected for recovery (Table 3).
 255 Concentrations reported for the Humber are maximum concentrations measured in August, when the
 256 wider UK survey was undertaken.

257

258 4. Discussion

259 Most monitoring studies to date have been carried out in freshwater systems as it was
 260 originally thought that estuaries and coastal waters would dilute compounds so that they
 261 would be undetectable (Fabbri and Franzellitti 2016). Despite this hypothesis,

262 pharmaceutical contamination was found to be widespread and all of the estuaries
263 monitored in the current study contained at least three of the target analytes at levels of a
264 similar magnitude to those found in the freshwater environment, and higher than those
265 measured in many other estuaries (Hughes et al. 2013; Table 1). The levels of
266 pharmaceuticals detected in this study contribute to the overall picture on pharmaceutical
267 pollution and add to the growing evidence of this global issue (aus der Beek et al. 2016). Our
268 work indicates that the limited monitoring carried out to date may not have captured peak
269 concentrations that occur in these environments and clearly highlights that further work is
270 needed.

271 Ibuprofen was detected at the highest concentrations and in all of the estuaries sampled,
272 with its occurrence not only exceeding levels detected in other estuaries (Table 1) but also
273 those seen in river water both in the UK (Barbara Kasprzyk-Hordern et al. 2008; Kay et al.
274 2017; Burns et al. 2017, 2018) and globally (Hughes et al. 2013). Ibuprofen has only been
275 measured in seven estuaries previously, with maximum concentrations all under 1000 ng l⁻¹
276 (Table 1). Further monitoring studies should include ibuprofen as a priority to determine if
277 high concentrations seen in the UK are similar to those elsewhere.

278 Concentrations of paracetamol, diclofenac and trimethoprim were similar to those seen in
279 other global estuaries, with mean concentrations less than 100 ng l⁻¹ (Table 1). Whilst
280 maximum concentrations of paracetamol were similar to those detected in rivers (Barbara
281 Kasprzyk-Hordern et al. 2008; Burns et al. 2017), concentrations of diclofenac and
282 trimethoprim were considerably lower (Hughes et al. 2013; Nakada et al. 2017). In the
283 present study, water samples were collected at high tide, when concentrations would be
284 expected to be lowest, so it is possible that these levels could be higher at other points in the
285 tidal cycle (Yang et al. 2016). This is the first study to measure the occurrence of citalopram
286 although concentrations were low and did not exceed 50 ng l⁻¹, in agreement with previous
287 studies which have monitored citalopram in rivers (Hughes et al. 2013). Nevertheless,
288 PNECs for citalopram are below this level (Minguez et al. 2016).

289 Whilst widespread occurrence of pharmaceuticals was seen in the UK patterns in their
290 spatial and temporal distributions within and between estuaries were observed.

291

292 4.1 Humber Estuary

293 4.1.1 Spatial Variation

294 It is generally expected that pharmaceutical concentrations will decrease downstream due to
295 physical processes in an estuary leading to their breakdown and removal (Daughton 2016).
296 The spatial pattern of pharmaceutical occurrence in the Humber Estuary followed this
297 pattern to a degree; peak concentrations were found in the middle of the estuary, particularly
298 at R1, where samples were collected next to an outlet from a wastewater treatment plant
299 (WWTP), indicating that WWTPs could be a significant source of pharmaceuticals in the
300 Humber Estuary. Input from WWTPs has been attributed as the largest source of
301 pharmaceutical pollution in the aquatic environment (Caldwell 2016). In some cases
302 maximum concentrations were detected outside of this site; in April and June maximum
303 concentrations for paracetamol and ibuprofen occurred at sites R2-4. It is difficult to
304 determine what caused these peaks although these sites are within 6km from R1, so it is
305 possible that the large increases seen at these sites are still due to input at R1, and
306 fluctuations between these sites are the result of sampling timing (Ort et al. 2010). The site
307 (R4) which showed the highest levels ($6.2 \mu\text{g l}^{-1}$) of ibuprofen was also 7km upstream from
308 the confluence of the River Hull. Transport of pharmaceuticals from this tributary upstream
309 during high tide could also account for the increases seen. The River Trent, located near the
310 confluence with the Ouse (Figure 1), could also account for the addition of further
311 pharmaceuticals. Inputs of pharmaceuticals in other studies have also been attributed to
312 other sources such as improper disposal, leaching from landfills or through veterinary usage
313 and subsequent runoff of these compounds into the aquatic environment, which could also
314 account for these differences (Bound and Voulvoulis 2005; Ebele et al. 2017).

315 Dilution plays a key role in the fate of pharmaceuticals in the aquatic environment and the
316 decrease in concentrations after R1 is presumably caused by dilution away from the input
317 source (Baker and Kasprzyk-Hordern 2013). Decline of pharmaceutical concentrations
318 downstream in the estuary was observed to a greater extent for some compounds and,
319 therefore, is unlikely to be entirely due to dilution. Degradation has been found to be a
320 significant factor affecting the fate of pharmaceuticals and could account for these
321 differences (Caracciolo et al. 2015). Citalopram experienced the lowest decrease in
322 concentration downstream and was typically found at the same concentration or higher at A5
323 compared to A1, which could be explained by the low degradation which has been observed
324 in other studies (Metcalf et al. 2010; Styrihave et al. 2011). Ibuprofen, paracetamol and
325 trimethoprim also showed little decline in concentration beyond initial dilution after R1, which
326 is consistent with what has been seen at other sites. These compounds have been found up
327 to 10 km downstream from a WWTP (Bendz et al. 2005, Kay et al. 2017, Burns et al. 2018),
328 and trimethoprim has even been found 200 km downstream from a WWTP (Tamtam et al.
329 2008). Further WWTPs are located within the estuary (European Environment Agency,
330 2019) which could also account for this lack in decline. Diclofenac on the other hand, was
331 not detected at A3 or A5 during any of the sampling periods, but was found at A4. The
332 downstream decline of diclofenac has been found to be variable, with some studies finding it
333 to be more persistent than others (Bendz et al. 2005; Wilkinson et al. 2017). Removal of
334 compounds through degradation and sorption to sediment has been found to be highly
335 dependent on environmental conditions and sediment type.

336 4.2.1 Temporal Variation

337 Seasonal differences of pharmaceuticals have been observed in a number of studies and
338 these are often attributed to changes in usage and local environmental conditions (Golovko
339 et al. 2014b; Moreno-González et al. 2014). Trimethoprim was the only compound to show
340 significant temporal differences in concentrations (at sites R1-R4), with average winter
341 concentrations more than double that of those found during the summer months. Previous

342 studies have explained the occurrence of antibiotics in winter due to their higher usage in
343 those months to treat seasonal infections (Verlicchi and Zambello 2016). The temporal
344 differences seen in the occurrence of trimethoprim in the Humber Estuary appeared to follow
345 this pattern as prescriptions were highest in October 2016 to March 2017 and lowest in
346 August 2017 (supplementary material S4). Trimethoprim has been observed to have higher
347 winter concentrations in some studies (Golovko et al. 2014b) but not in others (Burns et al.
348 2018). Burns et al. (2018) found higher levels of trimethoprim during spring in the Ouse
349 (upstream from A1), which was attributed to hydrological differences seen between the
350 seasons sampled. As a result, it is likely that the temporal differences in trimethoprim are the
351 result of different site specific conditions or daily variations. Temporal variations in other
352 studies have also been explained by lower temperatures leading to lower degradation
353 (Golovko et al. 2014a), however, input at R1 was highest in April. The other target
354 compounds have exhibited seasonal patterns in other studies but did not in the Humber.
355 Paracetamol, for instance, has been detected at high concentrations in spring in some rivers
356 but winter in others, whilst other studies found no temporal variations (Paíga et al. 2016; Ma
357 et al. 2017; Burns et al. 2018).

358 Temporal variations in the downstream pattern of pharmaceuticals were also observed, with
359 the greatest variation seen at the sites furthest downstream (A3-A5). Pharmaceuticals were
360 mostly absent from these sites in October, with the exception of ibuprofen, where
361 concentrations were reduced. Sampling at high tide could account for the absence of these
362 pharmaceuticals downstream as the result of increased dilution or transport of contaminants
363 upstream (Munro et al. 2019). Pharmaceutical concentrations often fluctuate diurnally as the
364 result of the timing of effluent discharges from WWTPs and combined sewer overflows
365 (CSOs), as well as variations in wastewater quantities as the result of consumption patterns
366 (Xu et al. 2007). To an extent, there was a pattern in the presence of compounds at R1
367 consistent with those seen downstream in the estuary, so it is possible that the temporal
368 variations could be the result of these daily variations, instead of conditions seen seasonally.

369 The concentration of pharmaceuticals at R1 were lowest in October and the low input could,
370 in part, account for the absence of compounds seen at sites furthest downstream (A3-A5).
371 Likewise, concentrations for the majority of compounds were highest at R1 during February
372 where concentrations were highest at sites furthest downstream (A3-A5). This is further
373 evidence that there is a difference in input from WWTPs. R1 is not the only site at which
374 wastewater is discharged but if these other sites exhibit the same temporal variations then it
375 could explain the differences observed in concentrations at A3-A5. WWTP removal has been
376 found to be less efficient during the winter time due to lower temperatures and decreased
377 biodegradation, leading to higher concentrations in effluent (Vieno et al. 2005). At R1,
378 concentrations for all compounds were lowest in August when temperatures were warmest
379 (supplementary material S1).

380

381 4.3 UK Estuaries

382 The Humber Estuary was shown to represent a worst case scenario in terms of
383 pharmaceutical pollution, with all five pharmaceuticals present at relatively high
384 concentrations. Of the estuaries sampled, it was the second highest impacted by WWTPs,
385 with a PE of approximately 13.7 million people. The Thames, which was the most impacted,
386 was the only other estuary to contain all five compounds. A higher presence of
387 pharmaceuticals is frequently seen in large urban areas due to their increased usage (Hong
388 et al. 2018). With the exception of both the Humber and the Thames estuaries, there was no
389 apparent relationship between the number of WWTP and concentrations (Table 2). The
390 Cromarty Firth, which was the receiving water of only 3 WWTPs (15,600 PE), exhibited
391 similar levels of pharmaceuticals to the Humber. This could be explained by differences in
392 WWTP efficiency, as technology used in WWTPs can greatly affect the removal of
393 pharmaceuticals. For example, ibuprofen removal has been reported to be between 7% and
394 99% at different WWTPs (Radjenovic et al. 2007; Jelic et al. 2015). It is possible that the
395 removal efficiency of WWTPs could differ between areas, with rural areas being less efficient

396 as they are serving smaller populations. Rural areas are more likely to have a higher
397 occurrence of septic tanks, which could contribute to the elevated levels seen in the
398 Cromarty (Hanamoto et al. 2018). Whilst the Humber experienced the lowest concentration
399 in August, it is possible that seasonal variations in population in areas like the Scottish
400 Highlands (a tourist destination), where the Cromarty is located, could be responsible for
401 these higher concentrations, increasing pressure on WWTPs. Pharmaceuticals in a
402 Portuguese river have previously shown higher concentrations which was thought to be the
403 result of increased summer populations (Rocha et al. 2014).

404 The presence of pharmaceuticals is greatly influenced by environmental conditions and
405 proximity of the sampling site to input sources, possibly accounting for some of the apparent
406 differences in concentrations observed between estuaries. Water samples from different
407 locations in the estuary were mixed together and a subsample was taken to obtain a
408 snapshot of the presence of pharmaceuticals, and it is likely that these concentrations will
409 vary depending on these factors. This could possibly explain the absence of diclofenac,
410 which in the Humber study was frequently undetected in sites downstream in the estuary.
411 Citalopram also had a low detection (50%) in estuaries, however, it was detected in
412 estuaries which have the highest PE.

413 There are also likely to be more complex interactions in play which further affect the
414 occurrence of pharmaceuticals in estuaries and which can help to explain the spatial
415 differences seen. Differences in site specific conditions such as salinity profiles and
416 hydrology can affect sorption processes, degradation and dilution. Undoubtedly, these
417 processes, in conjunction with daily variations in rainfall and temperature, are likely to be
418 responsible for differences in concentrations in estuaries between sampling periods, yet it is
419 still clear that pharmaceutical pollution is a ubiquitous problem in estuaries (Tamtam et al.
420 2008).

421 Ibuprofen, paracetamol, diclofenac and trimethoprim were previously monitored in the
422 Mersey, Thames, Tees and Tyne estuaries (as well as Belfast Lough) in 2002 (Thomas and

423 Hilton, 2004). It was also found that ibuprofen was present at the highest concentrations.
424 Paracetamol, however, was not detected in any of the estuaries sampled in 2002, which
425 indicates that the occurrence of this compound could be rising. A rise in pharmaceuticals
426 would be consistent with what has been found in other areas. For example, analysis of
427 sediment cores in Jamaica showed an overall rise in pharmaceutical concentrations over
428 time, with these concentrations doubling over the last decade (Lara-Martin et al. 2015). This
429 highlights the importance of establishing baseline measurements of pharmaceuticals, in
430 order to determine areas most at risk and therefore requiring continued monitoring. The
431 Humber Estuary likely poses the greatest risk, particularly due to the high level
432 concentrations of ibuprofen. Other large urban estuaries (such as the Thames and Severn)
433 may also warrant a further detailed study. However, as seen with the Cromarty, focus on
434 monitoring should be extended to rural areas as well.

435

436 **5. Conclusion**

437 All five target analytes, ibuprofen, paracetamol, diclofenac, trimethoprim and citalopram,
438 were detected in twelve estuaries in the UK. Diclofenac is a compound that has been
439 highlighted by EU legislation as a potential concern yet paracetamol and ibuprofen were
440 consistently detected at higher concentrations and at levels which could be toxic to aquatic
441 organisms (Vestel et al. 2016). In particular, the concentrations of ibuprofen measured
442 indicate that the limited monitoring of pharmaceuticals in estuaries around the globe to date
443 has not accurately quantified peak concentrations. Whilst trimethoprim was detected in every
444 sample it was only present at concentrations in the low ng l⁻¹ range and although citalopram
445 was present at lowest concentrations it showed the least change in concentration
446 downstream. A more intensive monitoring regime of the Humber Estuary showed that
447 pharmaceutical input from WWTPs is a significant source and could explain the overall
448 higher concentrations of pharmaceuticals in large urban estuaries. Despite this, a rural
449 estuary had the highest concentration of ibuprofen which may be due to lower removal at

450 smaller rural sewage works. More detailed studies need to be undertaken in order to
451 understand the complex interactions taking place in estuaries which could affect the fate of
452 pharmaceuticals.

453 Whilst there was little significant variation of pharmaceutical concentrations between
454 sampling periods in the Humber Estuary, August typically had the lowest input from WWTPs
455 and overall lowest concentrations, which is when samples were taken from estuaries
456 throughout the UK. Consequently, it could be expected that pharmaceutical concentrations
457 may exceed those measured. Additionally, samples were taken on a high tide when it would
458 be expected that concentrations are lowest due to dilution. This study provides an important
459 baseline of pharmaceutical measurements in the UK and highlights ibuprofen as a
460 compound which may warrant further assessment. This work provides further evidence to
461 the growing problem of pharmaceutical pollution, highlighting that it is not only an urban and
462 localised issue.

463

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