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Evaluation of combined sewer overflow impacts on short-term pharmaceutical and illicit drug occurrence in a heavily urbanised tidal river catchment (London, UK)

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4 **EVALUATION OF COMBINED SEWER OVERFLOW IMPACTS**
5 **ON SHORT-TERM PHARMACEUTICAL AND ILLICIT DRUG**
6 **OCCURRENCE IN A HEAVILY URBANISED TIDAL RIVER**
7 **CATCHMENT (LONDON, UK)**

8

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29 **Abstract**

30 The occurrence of pharmaceutical and illicit drug residues potentially arising
31 from combined sewer overflows (CSOs) in the Central London portion of the
32 Thames Estuary is presented. Approximately 39 million tonnes of untreated
33 sewage enter the River Thames at 57 CSO points annually. Differential analysis
34 of influents and effluents in a major wastewater treatment plant identified seven
35 potential drug-related CSO markers based on removal rates. Three were
36 present in influent at concentrations $>1 \mu\text{g L}^{-1}$ (caffeine, cocaine and
37 benzoylecgonine). During dry weather, analysis of hourly samples of river water
38 revealed relatively consistent concentrations for most drugs, including CSO
39 markers, over a tidal cycle. River water was monitored over a week in January
40 and July and then daily across six consecutive weeks in November/December
41 2014. Out of 31 compounds monitored, 27 drug residues were determined in
42 the River Thames and, combined, ranged between $\sim 1,000\text{-}3,500 \text{ ng L}^{-1}$. Total
43 drug concentration generally declined during extended periods of drier weather.
44 For CSO markers, short-term increases in caffeine, cocaine and
45 benzoylecgonine concentration were observed $\sim 24 \text{ h}$ after CSO events
46 (especially those occurring at low tide) and generally within one order of
47 magnitude. Timings of elevated occurrence also correlated well with
48 ammonium ion and dissolved oxygen data following CSOs. This work also
49 represents an important study of pharmaceutical occurrence before a major
50 'Super Sewer' infrastructure upgrade in London aiming to reduce CSOs by
51 95 %.

52 **Keywords:** river water monitoring, emerging contaminants, high resolution
53 mass spectrometry, CSOs

54 **1. Introduction**

55 Pharmaceuticals as environmental contaminants have been the focus of
56 much research in the past 20 years. Concentrations, generally in the ng- μ g L⁻¹
57 range, have now been reported in most environmental compartments including
58 wastewater [1-3], surface/ground water [4-6], marine water [7-9], solids [10],
59 biota [11] and even in air [12]. However, the primary source of pharmaceutical
60 and illicit drug contamination in the receiving environment has been identified
61 as outputs from wastewater treatment plants (WWTPs), as either treated
62 effluent or via sludge. In the EU, some pharmaceutical compounds have been
63 placed on a 'watch-list' until sufficient evidence on the full extent of their impacts
64 is known [13]. Environmental contamination and effects of illicit drugs have also
65 been reported, albeit on a smaller scale to pharmaceuticals, and the focus for
66 these has been largely on their measurement in untreated wastewater to
67 estimate community consumption patterns [14-16].

68 As part of the wastewater infrastructure of many developed towns and
69 cities, combined sewers are often used to simultaneously carry storm water and
70 municipal sewage to urban WWTPs. Such sewers are often designed to carry
71 several fold the average dry-weather load, but in extreme cases of runoff,
72 rainfall or snowmelt, capacity can be breached. In these cases, combined
73 sewer overflow (CSO) events occur to avoid back-flooding of streets and
74 homes. Storm flow is normally mixed with treated or untreated wastewater and
75 released directly into a nearby river or water body. Many reports have detailed
76 the resultant changes in water quality [17] and ecosystem impacts [18] arising
77 from faecal matter [19], microbial pathogens [20, 21], priority pollutants [22] and
78 other storm water-related contents [23] .

79 In London, ~39 million tonnes of untreated sewage is discharged into the
80 river Thames every year on average, but following exceptional wet weather and
81 flooding in 2014, that total rose to 62 million tonnes [24]. London is mostly
82 served by a Victorian combined sewer system built by Sir Joseph Bazalgette
83 following the ‘Great Stink’ of 1858. From 1831 until its completion in 1865, an
84 estimated 40,000 Londoners died from cholera. The expansion of London and
85 an increasing population (>8.3 m) has meant that the system is currently
86 running at approximately 80 % of its capacity, resulting in more frequent
87 breaches with CSOs occurring at least once a week, even at times of light
88 rainfall. London’s sewer system contains 57 CSO vents, 36 of which were
89 assessed as having adverse environmental effects [25, 26]. CSO discharges
90 were found to reduce the dissolved oxygen (DO) levels in river, introduce
91 pathogenic organisms and to cause negative aesthetic changes in the river
92 through the release of sewage, sewage litter, grease and scum directly into the
93 river. A potential solution has been the Thames Tideway Tunnel, or ‘Super
94 Sewer’, currently being built ~66 m under the river over 25 km. This major
95 upgrade will intercept 34 CSOs and reroute sewage to a relief WWTP at
96 Beckton in east London. It is due to be completed by 2023 and aims for an
97 average 95 % reduction in sewage discharged to the river [27].

98 In comparison to prioritised pollutants, the impact of CSOs containing
99 multiple pharmaceutical residues on receiving waters has received relatively
100 little attention. A recent study by Kay et al. [28], showed that concentrations of
101 five compounds monitored over 18 months in non-tidal rivers did not decrease
102 even 5 km from the nearest WWTP in Northern England, which may potentially
103 influence risk assessments based on models using first-order decay kinetics in

104 rivers [29]. Repeated sampling was also performed to identify fluctuations
105 across a day, which showed significant variance in measured concentrations
106 and, in some cases, across two-three orders of magnitude. A second study by
107 Benotti and Brownawell near New York City reported concentrations of 12 high-
108 volume pharmaceutical residues in mixed freshwater-saline regions across
109 Jamaica Bay during dry and wet weather conditions [30]. Of these, two
110 compounds had similar or higher concentrations in comparison to dry weather
111 conditions (acetaminophen and nicotine). Despite being a comprehensive
112 spatial study, repeated sampling was not performed to monitor temporal
113 changes at each site. However, this study demonstrated the effect of salinity on
114 pharmaceutical concentrations. Weyrauch et al. showed that compounds with
115 removal efficiencies >95 % during wastewater treatment could result in
116 elevated concentration in river water after CSOs [31]. For example, and though
117 not a pharmaceutical, concentrations of nitrilotriacetic acid in the River Spree
118 increased by 10-fold following a CSO and was well removed by a WWTP in
119 Berlin. Compounds with intermediate removal above ~56 % also showed an
120 increase in some cases, despite dilution with rainwater. Madoux-Humery et al.,
121 performed high resolution temporal sampling of sewage outfalls over a year in
122 Canada [32]. Several CSO markers were monitored and *E. coli* was considered
123 the best overall. However, of four pharmaceuticals monitored, carbamazepine
124 was determined to be the best marker of CSOs due to its persistence, specificity
125 for human use, stability and correlation with *E. coli*. Previous work by the same
126 group showed that caffeine was correlated with faecal coliforms [33] and its use
127 as an indicator of wastewater contamination was also shown by other groups
128 in different parts of the world [34-38]. Acetaminophen was also identified as a

129 suitable CSO marker by other groups [38, 39]. In an alternative approach, Fono
130 et al. showed that chirality could be exploited to identify raw sewage discharges
131 and/or CSOs using the ratio of one of the isomers of propranolol to its total
132 concentration [40]. Aside from CSOs, use of drug markers has also recently
133 been proposed to differentiate sewage from manure contamination [41]. Save
134 for a few studies [42-44], the number of pharmaceuticals and especially illicit
135 drugs included is generally small. More comprehensive analytical methods are
136 required to fully identify the scale of CSO impacts more broadly regarding such
137 compounds. Ideally, these should be more tailored to the catchment at the
138 method development stage. The advent of liquid chromatography-high
139 resolution mass spectrometry (LC-HRMS) has enabled a more flexible
140 approach to multi-residue analysis, by allowing targeted, untargeted and
141 suspect screening to be performed on large numbers of compounds, often
142 simultaneously [45-48]. However, reports using such approaches for CSO
143 impact assessment on receiving waters for pharmaceuticals and illicit drugs are
144 few.

145 The aim of this work was to identify fluctuations in drug concentrations
146 in the Central London catchment of the River Thames potentially arising from
147 CSO events. The objectives were (a) to perform a differential quantitative
148 analysis of influent and effluent wastewater to identify CSO-related drug
149 markers, and (b) to monitor fluctuations in general drug occurrence, as well as
150 ammonium and DO in receiving river water during dry and wet weather. In
151 particular, sampling sites were chosen for their location ~25 km away from any
152 main WWTP effluent discharge points. This project focused on quantitative
153 monitoring of a larger number of pharmaceutical and illicit drug compounds than

154 studied previously (n=31), and measured at high frequency, with an analytical
155 method based on LC-HRMS that was flexibly adapted for the catchment. Also,
156 this work serves as a potential snapshot of drug contamination before a major
157 sewer infrastructure upgrade such as the Thames 'Super Sewer' project.

158

159 2. **Experimental**

160 2.1 *Materials and Reagents*

161 All reagents were of analytical grade or higher. Methanol (MeOH), acetonitrile
162 (MeCN), dichloromethane (DCM) and dimethyldichlorosilane (DMDCS) were
163 purchased from Fisher Scientific (Loughborough, UK). Ammonium acetate and
164 37 % (w/v) hydrochloric acid solution were sourced from Sigma-Aldrich
165 (Gillingham, Dorset, UK). Ultra-pure water was obtained from a Millipore Milli-
166 Q water purification system with a specific resistance of 18.2 MΩ.cm (Millipore,
167 Bedford, USA). All glassware including stock solution vials and evaporation
168 tubes were silanised to reduce loss of analyte through adsorption to the glass
169 surfaces. Each component was rinsed with a 50:50 (v/v) MeOH/H₂O solution
170 before triplicate rinses with DCM. A 10:90 (v/v) DMDCS/DCM solution was then
171 used to rinse the container followed by triplicate rinses with each of DCM, 50:50
172 MeOH:H₂O solution and water. A total of 51 pharmaceuticals, illicit drugs and
173 metabolite reference materials were purchased from Sigma Aldrich
174 (Gillingham, UK) for analytical method development and assessment (See
175 Table S1. Stock solutions (1,000 mg L⁻¹) were prepared in MeOH and working
176 standard solutions prepared weekly in ultrapure water or LC mobile phase A.
177 All solutions were stored in silanised amber glass vials at 4 °C in dark
178 conditions.

179

180 2.2 *Sampling sites and procedures*

181 Wastewater influent (immediately after the fine screen) and treated effluent
182 were taken as seven 24-hour composite samples from a major sewage
183 treatment works in London (population equivalent = 3.5 million) from 11-17th
184 March 2014 to identify pharmaceuticals and illicit drug residues potentially
185 indicative of CSO events. A 12-hour diurnal occurrence study was conducted
186 using 13 hourly grab samples (500 mL) taken on Tuesday 12th August 2014, at
187 Gabriel's Pier, London (51°30'31.0" N; 0°06'35.1" W) covering a period from
188 07:00 to 19:00 and collected at ~0.5 m depths. A moderate temperature (16-23
189 °C), mainly dry day (<1 mm rainfall) was chosen to reflect a normal daily river
190 cycle and free from storm runoff or triggered CSOs. For inter-season
191 occurrence of pharmaceutical and illicit drug CSO marker candidates, samples
192 were taken from two sites, again at ~0.5 m depths each time: Site 1 was at
193 Lambeth Bridge (51°29'42.4"N 0°07'27.8"W) and Site 2 was at Gabriel's Pier
194 (as above). Of 57 vents in total in London, six CSO vents lay in close proximity
195 to Site 2 in both directions, spanning from Westminster Bridge to Blackfriars
196 Bridge. For Site 1, a CSO vent lay within 50 m of the sampling site on the same
197 bank. Following this, a high frequency sampling campaign was conducted by
198 taking grab samples over a 6-week period at 09:00 on weekdays from Site 2
199 from 3rd November-13th December 2014. All samples of wastewater and river
200 water were collected in 500 mL Nalgene bottles, transported immediately to the
201 laboratory (~30-60 min transit time), acidified to < pH 2 with HCl and frozen (-
202 20 °C) until analysis. Tide heights were also recorded at the river sampling site
203 at each timepoint using the local tidal gauge pole. Daily rainfall data for the

204 sampling site was gathered from the published CEH-GEAR dataset by Tanguy
205 et al. [49].

206

207 *2.3 Sample pre-treatment and solid phase extraction*

208 Before extraction, samples were thawed and filtered under vacuum using
209 Whatman GF/F 0.7 μm glass microfiber filters. For matrix-matched standards,
210 acidified 100 mL sample aliquots were spiked volumetrically before solid phase
211 extraction (SPE). HyperSep Retain Polar Enhanced Polymer (PEP) cartridges
212 (200 mg x 6 mL) were selected for SPE of river water and wastewater (Thermo
213 Fisher Scientific, Runcorn, UK). Cartridges were conditioned with 4 mL MeOH
214 and 4 mL ultrapure water. Acidified samples (100 mL) were loaded under
215 vacuum at $\sim 5 \text{ mL min}^{-1}$ and washed thereafter with 4 mL 5:95 (v/v) MeOH:H₂O.
216 The sorbent was dried under vacuum prior to elution for ~ 10 min before elution
217 with 4 mL MeOH. Eluted extracts were evaporated to dryness under N₂ at 35 °C
218 and reconstituted in 100 μL of 10 mM ammonium acetate 90:10
219 water:acetonitrile (mobile phase A) using a positive displacement pipette. The
220 reconstituted samples were then sonicated for ~ 10 min before being transferred
221 to an amber HPLC vial fitted with a silanised insert for analysis.

222

223 *2.4 Instrumentation*

224 For LC-HRMS analysis, an Accela ultra-high performance LC system, an HTS-
225 A5 autosampler (at 10 °C) and an ExactiveTM (Orbitrap) HRMS detector were
226 used throughout. All separations were performed on a Thermo 150 x 2.1 mm,
227 2.6 μm Accucore C₁₈ analytical column fitted with a matching 10 x 2.1 mm, 2.6
228 μm Accucore C₁₈ guard column. The LC flow rate was 0.4 mL min⁻¹, the

229 temperature was maintained at 24 °C and the injection volume was 20 µL. A
230 binary gradient elution profile of 90:10 to 20:80 10 mM ammonium acetate in
231 water:acetonitrile (mobile phase A and B, respectively) was used as follows:
232 0% B for 2.5 min; 0-30% B from 2.5 to 7.5 min; 30% B from 7.5 to 12.5 min; 30-
233 40% B from 12.5 to 15 min; 40-100% B from 15.0 to 20.0 min; 100% B from
234 20.0 to 27.5 min. Re-equilibration time was 7.5 min. The Exactive™ HRMS was
235 fitted with a heated electrospray ionisation source (HESI-II). All samples and
236 model solutions were run separately in either positive or negative ionisation
237 mode at 50,000 FWHM with a scan range of m/z 100–1000. Each acquisition
238 cycle comprised of a full-scan without higher energy collisional dissociation
239 (HCD) followed by a full scan with HCD enabled (collision energy: 20 eV; cycle
240 time: ~2 s). Sheath, auxiliary and sweep gas settings were 50, 10 and 0
241 arbitrary units, respectively. The capillary temperature was 350 °C; the heater
242 temperature was 300 °C; and the positive/negative spray voltages were +4.50
243 kV and –3.00 kV. All acquisition data was processed using Xcalibur v2.0
244 software. The entire analytical method was validated to ICH guidelines in
245 wastewater and river water (see Tables S2-S4) [50]. Method development
246 details are also presented in the Supplementary Information. For wastewater
247 influent and effluent, the method was found to be quantitative for n=33 and n=38
248 compounds in untreated influent and treated effluent, respectively. For river
249 water, the method could reliably quantify n=31 compounds at environmentally
250 relevant concentrations.

251

252 *2.5 Targeted analysis, quantitation and statistical procedures*

253 Confirmation of target analyte occurrence in all samples was based on the
254 accurate mass of the protonated/deprotonated precursor ion and its associated
255 major HCD product ion to within 5 ppm mass accuracy, the ratio between these
256 two ions (<30 % to a matrix-matched standard) and a matching
257 chromatographic retention time (t_R) to within 15 s. For 24-h composite
258 influent/effluent wastewater samples, duplicate aliquots were extracted for each
259 day and determined using matrix-matched calibration using a pooled matrix of
260 all samples across the week-long sampling period. Background correction was
261 performed, as needed. Calibration lines were prepared for $N \geq 5$ points,
262 alongside triplicate background-corrected quality control samples (50 ng/L) to
263 allow the accuracy of the method to be monitored. Given that the river was tidal
264 and brackish, significant variance in analyte matrix effects across days was
265 observed for a number of compounds (data not shown), so all drugs were
266 determined in duplicate using 3-point standard addition in each sample
267 separately for added accuracy. Drug occurrence in all samples is reported as
268 the average of duplicates with error bars representing the larger of the two
269 measurements. For temporal occurrence experiments, measured values over
270 each timeframe were averaged and the associated variance expressed as the
271 standard deviation, unless otherwise specified.

272 All statistical treatment of data was performed in Microsoft Excel. For
273 quantitation/calibration, lines-of-best-fit were applied and coefficients of
274 determination (R^2) calculated. For correlations between tide height/rainfall and
275 drug concentration (Figure S4), the Pearson correlation (R) was calculated and
276 significance tested by considering a p -value threshold of 0.05 to reject the null
277 hypothesis. For statistical comparisons of drug removal efficiency from

278 wastewater, data was first checked for normality and the p -value quoted
279 following application of the specified test.

280

281 *2.6 Suspect screening of wastewater and river water*

282 Suspect screening was performed on wastewater samples only to differentially
283 identify unique drugs/metabolites or those with potentially higher
284 concentrations in influent. Post-acquisition automated peak selection was
285 performed using Thermo TraceFinder™ version 3.1 software which contained
286 a library of HRMS spectra for $n=1,492$ pesticides, herbicides, fungicides,
287 pharmaceuticals, metabolites and illicit drugs. Following this, predicted t_R for
288 potentially new compounds was performed using a previously developed neural
289 network algorithm (Trajan v6.0, Trajan Software Ltd., Lincolnshire, UK) using
290 reference t_R data for 166 pharmaceuticals, illicit drugs and metabolites
291 measured in influent and effluent wastewater extracts [51]. Compounds were
292 tentatively identified using a t_R window of ± 1.3 min and an accurate m/z within
293 5 ppm of its calculated m/z . Lastly, an 80% fit threshold to theoretical isotope
294 profile was set, with an acceptable intensity threshold deviation for each isotope
295 ion set at 25% of the theoretical value.

296

297 *2.7 DO, ammonium and conductivity monitoring*

298 Percentage DO, pH, conductivity (as a measure of salinity), and ammonium
299 concentration were taken at 15-minute intervals by the Environment Agency
300 (EA), UK and analysed at three sites (Putney, Brentford and Hammersmith)
301 using YSI6600 systems (Environmental Monitoring Systems, Herts, UK). DO
302 was measured as % saturation using the YSI optical DO Sensor. The Sonde

303 software automatically compensated for the effect of temperature. River pH was
304 measured using a combination electrode with an Ag/AgCl reference electrode.
305 Ammonium was measured using an YSI ion selective electrode and the
306 reference being provided by the pH combination electrode. Conductivity (μS
307 cm^{-1}) was reported as specific to 25 °C and was calibrated using a solution of
308 KCl. The YSI6600 sensors were calibrated every 4 weeks following standard
309 EA operating procedures.

310

311 **3. Results and Discussion**

312 *3.1 Differential analysis of influent and effluent wastewaters and* 313 *identification of candidate CSO markers*

314 To shortlist a selection of CSO-related pharmaceutical and illicit drug markers,
315 differential analysis of influent and effluent wastewaters was performed. Direct
316 analysis of in-sewer CSO samples was not performed due to limited access.
317 Two important criteria were considered. Candidate CSO drug markers were
318 shortlisted where they were: (a) ideally only present in untreated influent
319 wastewater (i.e. high removal efficiency in the WWTP); and (b) remained at
320 measurable and relatively consistent concentrations every day (i.e., minimal
321 seasonal variation or recreational usage patterns should be evident).

322 All determined drug concentrations are presented in Tables S5 and S6
323 and summarised in Figure 1. A total of 14 compounds were quantifiable almost
324 every day in untreated influent wastewaters and two of these were unique to it,
325 i.e. diazepam and sulfapyridine, present at 76 ± 14 and 184 ± 96 ng L^{-1} ,
326 respectively, which were both selected as candidates. Prescription drug
327 concentrations were generally consistent across the week in both influent and

328 effluent (except for sulfapyridine, which was not detected on one day). Both
329 bezafibrate and furosemide were quantifiable in influent at similar
330 concentrations ($\sim 400 \text{ ng L}^{-1}$), but less than the lower limit of quantification
331 (LLOQ) in effluent. This corresponded to an >10 -fold lower concentration, so
332 both were considered as potential CSO markers. Tramadol exhibited the
333 opposite trend, with significantly higher levels detected in effluent at $1,138 \pm 106$
334 ng L^{-1} ($p = 3 \times 10^{-7}$, Student's two-tailed t -test), with over a two-fold concentration
335 increase observed between both matrices. Nine other compounds were present
336 at quantifiable levels on a regular basis in effluent. Extensive wastewater
337 monitoring over the past five years as part of the £130 m UK Water Industry
338 Research (UKWIR) Chemical Investigation Programme (CIP) Phase 2 (CIP2)
339 has played a key role in the selection of substances and sites for future controls
340 and remedial measures [52, 53]. It included up to 73 individual determinands
341 across 44 WWTPs from 2015-2017 including data for six pharmaceutically-
342 related compounds for which removal rates could be calculated: diclofenac (42
343 $\pm 29 \%$), ibuprofen ($98 \pm 4 \%$), propranolol ($28 \pm 24 \%$), carbamazepine (-8
344 $\pm 35 \%$), carbamazepine epoxide ($30 \pm 28 \%$) and fluoxetine ($43 \pm 22 \%$) [54].
345 The London-based WWTP studied here was not included within the 44 CIP2
346 sites. Comparative removal rates for this WWTP could be calculated reliably
347 here for carbamazepine (-61% , i.e., more concentrated in the effluent) and
348 propranolol (34%), and an estimation made for fluoxetine (65% ; occurrence
349 was $< \text{LLOQ}$, but $> \text{LOD}$ in influent).

350 For the selected illicit drugs, most were quantifiable during the week
351 except for methylenedioxymethamphetamine (MDMA) and generally increased
352 over the weekend. This was consistent with recreational consumption trends

353 seen previously [15]. Ketamine was eliminated as a candidate CSO marker, as
354 it was present at slightly higher concentrations in effluents than influents (58 ± 5
355 and 42 ± 9 ng L⁻¹, respectively) and measurements also lay close to the LLOQ.
356 Ketamine has been shown to display partial transformation in sewer transit
357 (<25 %) [55], as well as variable and even negative removal rates following
358 wastewater treatment [56, 57]. Possible reasons for higher concentrations in
359 effluent include residence times below 24 h, as well cleavage of conjugated
360 metabolites and desorption from particulate matter during treatment [58-60].
361 Mephedrone was detected at low levels in all samples and quantifiable at 83
362 ± 45 ng L⁻¹ in six out of seven influent samples (<LLOQ in effluent). Interestingly,
363 concentrations of cocaine and its metabolite benzoylecgonine remained high in
364 influent wastewater across the week with only a relatively minor increase in
365 occurrence over the weekend (%RSD <10 % for benzoylecgonine and <25 %
366 for cocaine), which is not consistent with many other cities. London is known as
367 one of the highest consumers of cocaine and this result suggested everyday
368 usage [16]. Cocaine was detected at significantly higher levels in influent (p
369 $= 3 \times 10^{-5}$; Student's two-tailed t -test) as well as analyte concentrations in effluent
370 at ~30-fold lower levels, which represented >99 % removal efficiency at this
371 WWTP. While WWTP removal performances can differ between sites, similar
372 removal of cocaine and benzoylecgonine from influent has been reported in
373 other parts of UK and globally, even up to 100 % [57, 61]. Given their metabolic
374 linkage, both were given further consideration as CSO markers. In addition to
375 these compounds, caffeine was also detected only in influent. However, its
376 concentration was so high that it lay outside of the quantifiable range when
377 using background corrected matrix-matched standard addition. However,

378 previous work using stable isotope internal standards showed that caffeine
379 concentration in untreated wastewater from London was quite stable at 23 ± 2
380 $\mu\text{g L}^{-1}$ across a full week [15]. Caffeine has also been shown to be removed
381 almost completely by wastewater treatment processes by both aerobic and
382 anaerobic degradation [57, 62]. Caffeine was therefore retained as a candidate
383 CSO marker and more reliable measurements in river water matrix were
384 possible when present at a diluted concentration. Another compound, salicylic
385 acid, was present at excessively high concentrations to quantify it in influent
386 and was not detected in effluent. However, the poor method performance for
387 this compound, observed in all three matrices assessed, meant it was not
388 suitable for quantitative monitoring and was eliminated for use.

389 Application of HRMS database searching (TraceFinder) and reference
390 to matching predicted chromatographic retention times resulted in tentative
391 identification of $n=32$ more drug residues in influent and $n=28$ more in effluent
392 across the week (Tables S7 and S8). For influent only, two detectable
393 chromatographic peaks were present for four compounds in extracted ion
394 chromatograms within their 1.3 min retention window even at 5 ppm mass
395 accuracy/isotope profile matching (i.e., matching hydrocortisone, salbutamol,
396 testolactone and acetylsalicylic acid, but not confirmed with reference
397 standards). A total of 14 compounds were detected in influent at higher signal
398 intensities than effluent at least once across the week (Figure 2 and Table S9).
399 Eleven compounds were tentatively identified in effluent every day, including
400 nine also present in influent every day. However, two unresolved isomers
401 (quinine and quinidine) were present at markedly higher signal intensities in
402 influent and were used together as a combined signal as potential CSO

403 markers. It was expected that of the two, quinine was likely to be the dominant
404 compound given its widespread use in tonic waters.

405 A total of seven target analytes (bezafibrate, benzoylecgonine, caffeine,
406 diazepam, sulfapyridine, cocaine and furosemide) were shortlisted as
407 candidate CSO markers quantitatively. Quinine and quinidine were used
408 together as qualitative CSO markers. For the six-week monitoring study, all
409 other compounds were still included for river water monitoring, even if not
410 considered as potential CSO markers to assess the potential contribution of
411 CSOs in general.

412

413 3.2 *Diurnal variation in drug concentrations in the River Thames*

414 The river sampling sites in Central London lay within the Thames Estuary,
415 where river levels often change by up to seven metres, twice a day. River flow
416 is relatively small compared with the volume of the tide and therefore, is well
417 mixed. Generally, the entire water mass travels in and out of the estuary with
418 tidal cycles. When CSOs discharge to the river, it takes approximately one
419 month for litter and sewage to exit the estuary to the sea in Winter and up to
420 three months in Summer [24]. River water is also brackish to the top of the
421 estuary at Teddington Lock, which lies west of the city. Previous research has
422 shown that varying salinity, dissolved organic carbon (DOC) and/or suspended
423 particulate matter (SPM) can influence drug concentrations in tidal waters [42,
424 43]. Therefore, fluctuations in drug concentration were monitored over a tidal
425 cycle on a day free from storm water runoff or CSOs to understand the impact
426 of fresh/saline water changes. From a qualitative perspective, n=24/31
427 compounds included in the validated method were detected at least once

428 across the day at Site 2 (Table S10) showing that the selection of compounds
429 was highly relevant to this catchment and benefited greatly from the use of
430 flexible full-scan LC-HRMS-based methods. Of these, n=18 drug residues were
431 quantifiable and n=13 of those determined at all sampled time points. Figure
432 3(a) shows that four potential CSO marker drugs were quantifiable and
433 remained relatively low in concentration. As perhaps expected, caffeine was
434 present at the highest concentration across the day at $112 \pm 48 \text{ ng L}^{-1}$, and it
435 presented a minor correlation with tide. No obviously apparent correlation with
436 tide was observed for the other three CSO markers and all remained below ~ 20
437 ng L^{-1} . Figure 3 (b)-(d) show the other determined pharmaceutical residues,
438 again most of which showed low and relatively consistent concentration
439 profiles. Tramadol and carbamazepine concentrations were the highest
440 between $\sim 100\text{-}300 \text{ ng L}^{-1}$ over the 12-hour period. Tramadol occurrence has
441 been linked to hospital effluent contribution to CSOs, but was present at lower
442 concentrations in untreated wastewaters here [63]. Trimethoprim,
443 sulfamethazine, carbamazepine and ketamine were the only obvious cases
444 showing any correlation with tide or water conductivity. These almost doubled
445 in concentration at high tide which was in contrast to observations for
446 pharmaceuticals by some other researchers [42, 43]. Three of London's five
447 WWTPs (Beckton, Riverside and Crossness) discharge treated wastewater into
448 the Thames $\sim 25\text{-}30 \text{ km}$ to the east of the Central London location (Site 2) and
449 serve a combined population equivalent of ~ 5.9 million ($\sim 71 \%$ of Greater
450 London). The remainder of the population is served mainly by Mogden WWTP,
451 which discharges effluent $\sim 25 \text{ km}$ west of Site 2 (~ 2 million population
452 equivalent). Therefore, concentration rises with high tide are likely due to drug

453 residues from more treated effluent entering downstream being swept inland
454 towards Site 2. Therefore, and in general, drug residues were not removed from
455 the sampling site by a tidal cycle and concentrations largely remained relatively
456 consistent. This was particularly useful for CSO markers considering that river
457 water conductivity changed from ~650-1,000 $\mu\text{S cm}^{-1}$ across the tidal cycle on
458 this date showing the salt water influx/efflux.

459

460 3.3 *Inter-season occurrence of pharmaceutical and illicit drug CSO marker* 461 *candidates*

462 CSOs were categorised into two main types. CSO Type 1 comprised of storm
463 water combined with untreated sewage, which was discharged directly into the
464 river. CSO Type 2 represented heavily diluted storm water that was screened,
465 settled in tanks and mixed with fully treated wastewater at a major WWTP
466 before release to the river. Public notifications of either CSO type corresponded
467 to two monitored sites in London: (a) Hammersmith pumping station (CSO Type
468 1) and (b) Mogden WWTP (CSO Type 2). Weather in January 2014 was one of
469 the wettest on record since 1910 with ~135 mm rainfall and available data from
470 Hammersmith Pumping Station alone revealed ~1,637,456 m^3 of CSO Type 1
471 discharge and 2,505,000 m^3 of Type 2 from Mogden WWTP [64]. However, the
472 total volume of either CSO type was likely much higher given that several more
473 pumping stations and CSO vents exist across the Central London catchment.
474 Across 2014, 16 million tonnes of untreated sewage were discharged into the
475 River Thames from just the central London CSO vents covering the two
476 sampling points selected. Three of these (the Hammersmith, Lots Road, and
477 Western Pumping Stations) contributed 11 million tonnes to that total. One Type

478 1 CSO event occurred during the week sampled in winter on 16th January, 2014
479 at 21:50 hours, but after a grab sample was taken. However, concentrations of
480 caffeine and benzoylecgonine increased at both Sites 1 and 2 on the following
481 day (Figure 4). Furthermore, at Site 1 increases in concentration were also
482 observed for bezafibrate and cocaine, most likely as it lay so close to a CSO
483 vent, but this trend was not observed at Site 2. Caffeine had the highest
484 concentration overall and reached a maximum of 1,520 ng L⁻¹ at Site 1 and
485 ~13 h after this Type 1 CSO. Its high concentration was prolonged in this
486 instance and took roughly two days to return to baseline concentrations. No
487 CSOs occurred during the week of sampling in July, 2014. Only ~44 mm rainfall
488 was recorded for the month with 24,000 m³ of Type 1 CSO discharge from
489 Hammersmith Pumping Station and no Type 2 CSO discharge from Mogden
490 WWTP. By comparison, caffeine concentrations were much lower in Summer
491 and rarely reached >200 ng L⁻¹. Detection of all other substances was
492 intermittent. Interestingly, baseline concentrations of bezafibrate and
493 benzoylecgonine remained relatively consistent with the January samples,
494 despite recorded rainfall and tidal height differences of >3.5 m across all
495 sampling timepoints. At this time of year, salinity of the river was also much
496 higher and more affected by tide as its freshwater composition was much lower
497 (conductivity of ~600-700 µS in the Winter dates studied versus 900-3,000 µS
498 in Summer)

499

500 *3.4 Longitudinal daily monitoring of pharmaceutical and illicit drug*
501 *occurrence in the River Thames over six weeks*

502 Site 2 was selected for a longitudinal occurrence study of all 31
503 pharmaceuticals given its convenience, reliability and safety of access during
504 bad weather across six weeks in Autumn and Winter, 2014. Furthermore, it
505 represented an equidistant point in the river between the major west and east
506 WWTP discharge points (~25 km in either direction). A total of 27 drug residues
507 were determined in the River Thames (Figure 5). The total (summed)
508 concentration of all compounds monitored varied from ~1-3.5 $\mu\text{g L}^{-1}$.

509 Over the course of the study, 13 CSOs were triggered due to heavy
510 rainfall (Table S11). In all, six Type 1 CSOs were recorded over the six-week
511 period, which were most relevant to this study. Of these, four samples were
512 taken within 24 hours following a CSO event. Available Type 1 CSO-related
513 records from the Hammersmith, Lotts Road and Western pumping stations
514 showed that a combined total of 1,883,485 and 204,150 m^3 of untreated
515 sewage mixed with storm water was discharged into the Central London region
516 of the River Thames in November and December months, respectively [64].
517 Measured total concentrations of illicit drugs and pharmaceuticals decreased in
518 general throughout November and December (Figure 5 and Table S12).
519 Approximately 75 % (~80-90 mm) of the total rainfall fell in the first three weeks.
520 Dilution with freshwater arising from the upper Thames may have been a
521 contributor to this decline, amongst other factors such as changing temporal
522 consumption patterns, varying WWTP performance, changing river water
523 chemistry (e.g., salinity, etc.), molecular stability and biological activity. On the
524 other hand, prolonged elevated concentrations following CSOs could have
525 arisen here where several events occurred in rapid succession, especially in
526 the first three weeks, and which were slowly removed by the tide. The top five

527 most concentrated compounds on average across the six weeks were caffeine
528 ($477 \pm 313 \text{ ng L}^{-1}$), diazepam ($305 \pm 558 \text{ ng L}^{-1}$), tramadol ($220 \pm 75 \text{ ng L}^{-1}$),
529 carbamazepine ($154 \pm 99 \text{ ng L}^{-1}$) and amitriptyline ($102 \pm 57 \text{ ng L}^{-1}$). Temporal
530 variance in measured concentrations across the 30 sampled days was, as
531 perhaps expected, high and not likely to only include any impact of CSOs, but
532 also changes in community consumption behaviour, illness/disease treatments
533 or seasonal consumption patterns influencing the concentrations in treated
534 wastewater effluents [65]. Where Type 1 CSOs occurred, no readily identifiable
535 spikes in total concentration of all drugs determined were observed within a 24
536 to 48hour period, nor any correlations with tide height, daily rainfall, or a ratio
537 of both ($R^2 < 0.1$ in all cases). Principal component analysis did not yield any
538 further classification between daily concentrations determined for all 27
539 compounds (Figure S2). In addition, five out of six Type 1 CSOs were also
540 accompanied by Type 2 CSOs, which may have served to dilute untreated
541 wastewater entering the Thames Tideway further. Some additional interesting
542 observations were made. The illicit drugs ketamine and mephedrone were
543 detected almost every day at $12 \pm 4 \text{ ng L}^{-1}$ and $9 \pm 2 \text{ ng L}^{-1}$, respectively. The
544 latter was banned in the UK in 2010, but was still determined in wastewater
545 influent, effluent and river water here in 2014. However, despite being present
546 at higher concentrations in influent, its concentration flux did not align with
547 CSOs, likely in part due to recreational use increasing over the weekend.

548 When focussing on the seven shortlisted candidate CSO markers, some
549 trends became more evident, but were very complex to interpret. Firstly,
550 concentrations of caffeine, cocaine and its metabolite benzoylecgonine in river
551 water showed a correlation with some CSOs. As their concentrations in

552 untreated wastewater was regularly $>1 \mu\text{g L}^{-1}$, this was perhaps expected over
553 the other four compounds. Elevated concentrations were mainly detected in
554 samples taken on the following day (Figure 6) especially following the two
555 heaviest rainfall events and CSOs on 23rd November and 11th December, 2014,
556 both during the lower portion of incoming flood tidal phases. For the latter date,
557 two CSOs were triggered on the following day at 06:25 (Type 1) and 08:58
558 (Type 2) just before the sample was taken and which enabled subsequent
559 determination of all compounds at higher concentrations, even within 3 hours
560 following a Type 1 discharge. However, neither cocaine nor benzoylecgonine
561 were detected at obviously elevated levels following Type 1 CSOs on the 4th or
562 14th November. On both occasions, the river was at the top of its tidal phase
563 and dilution may have occurred. As before, elevated caffeine concentration
564 following CSOs seemed prolonged over several days in comparison to cocaine,
565 especially after the heaviest rain event on the 22nd/23rd November.
566 Concentrations of diazepam were high across the first two weeks of the
567 campaign and then decreased markedly thereafter and did not correlate with
568 any one CSO event directly. Short-term elevated concentrations may be more
569 prolonged for this compound given its potential for sorption to sediment [66].
570 Following the CSO event on the 4th November, elevated concentrations of
571 sulfamethazine and sulfamethoxazole occurred, and a mild rise in
572 concentration of sulfapyridine over the following 48 h. However, sulfapyridine
573 was not useful to indicate other Type 1 CSO events across the remainder of
574 the campaign. Lastly, furosemide and bezafibrate yielded no apparent trends
575 and were removed from further interpretations.

576 The majority of compounds tentatively identified during suspect
577 screening as being indicative of influent wastewater were not present in river
578 water. However, the combined signal for the stereoisomers quinine/quinidine
579 was detected every day ($[M+H]^+$ m/z 325.1910), but revealed no obvious co-
580 incidence with CSO events (Figure S3). However, achieving chromatographic
581 resolution of both compounds and quantification is still required to fully evaluate
582 their individual value as CSO markers. Furthermore, the use of signal intensities
583 from LC-HRMS analysis was likely subject to variable matrix interference due
584 to the influence of seawater with tide, especially over the first week of the
585 sampling campaign (Figure 7(a)). However, for the majority of the six weeks,
586 conductivity measurements indicated that the river was predominantly
587 composed of freshwater (600-800 μS), mainly arising from influx of upstream
588 sources to Teddington Lock experiencing heavy rainfall and run-off.

589

590 *3.5 Ammonium, pH and %DO*

591 Comparison of drug concentrations with ammonium and %DO data
592 gathered simultaneously from Putney, Hammersmith and Brentford (each
593 ~5-7 km apart) in the west of the city revealed correlations with most Type 1
594 CSOs (Figure 7 (b)-(d)). Interestingly, and despite their distances apart, the
595 changes in ammonium/%DO concentrations at each site aligned well with each
596 other, indicating that CSOs may be triggered across the length of the network
597 simultaneously. However, and in agreement with some of the drug
598 measurements here, poorly discernable changes in ammonium concentration
599 or %DO were observed for Type 1 CSOs on the 4th, 8th or 9th November (only
600 observed clearly at the Brentford site). The pH of the river remained relatively

601 constant over the six weeks ($\text{pH} = 7.77 \pm 0.09$), and very minor reductions of
602 <0.25 pH units were observed during periods of elevated ammonium
603 concentration.

604 The duration of CSO impacts could be interpreted from ammonium
605 and %DO data (unfortunately, data for CSO duration and discharge volumes
606 were not available for specific dates). Generally, and like CSO drug markers,
607 changes occurred within 24 h after a CSO and returned to normal levels ~ 24 h
608 later. A mild positive, but statistically significant correlation ($R = 0.6023$;
609 $p = 0.0049$) existed between total concentrations of the three main CSO drug
610 markers determined on the following day with tide height:daily rainfall ratio at
611 the time of sampling (Figure S4). Therefore, it was concluded that there exists
612 a fine balance between tide height/direction, rainfall and time (<24 h here)
613 before an influent wastewater-specific drug can be measured in the river to
614 potentially indicate CSO influx. The Type 1 CSO event on the 23rd of November
615 2014 was the most prominent and prolonged from these data which explains
616 why concentrations of some CSO drug markers increased so markedly. The
617 Putney site is closest by distance to Site 2 chosen for drug monitoring (~ 11 km).
618 Despite being more central, smaller changes in ammonium and %DO were
619 observed across the six-week period. Therefore, proximity to a local CSO vent
620 will likely affect measurements overall. Ideally, more sites should be monitored
621 across this catchment to more fully understand spatial impacts of
622 pharmaceuticals and illicit drugs from CSOs on receiving waters. However,
623 despite short-lived peaks in concentration, longer term concentrations of
624 pharmaceuticals and illicit drugs in CSO material may decline overall upon

625 completion of the Thames Tunnel, which aims to reduce annual sewage
626 discharge via CSOs by 95 % [27].

627

628 **Conclusions**

629 Of 31 compounds monitored quantitatively, 27 pharmaceuticals and illicit drug
630 residues were determined in river water in the Thames Tideway in daily
631 measurements over six weeks. However, occurrence and total concentrations
632 of pharmaceuticals and illicit drugs as a whole showed no short-term correlation
633 with specific CSO events (total concentration lay between $\sim 1.0\text{-}3.5 \mu\text{g L}^{-1}$).
634 Following differential analysis of influent and effluent wastewater, seven
635 compounds were shortlisted as potentially being influent wastewater specific
636 and three of these were present at concentrations $>1,000 \text{ ng L}^{-1}$ in influent (i.e.
637 caffeine, cocaine and benzoylecgonine). In river water, these three compounds
638 showed noticeably elevated concentrations $\sim 24\text{-}48 \text{ h}$ after CSO events
639 following major rainfall events and aligned with ammonium and %DO data. It
640 was found that there existed a fine balance between tide height, direction and
641 rainfall, before any elevated concentrations of these CSO markers were
642 recorded. Therefore, CSO releases should be ideally aligned with the onset of
643 the ebb tidal phase to enable sufficient dilution to occur. However, even with
644 dilution, more research is required to understand the longer-term impacts of
645 CSOs on drug occurrence in receiving waters and particularly any potential
646 improvements following a major infrastructure upgrade such as that planned in
647 London to mitigate them.

648

649

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662

663 **References**

- 664 [1] Y. Luo, W. Guo, H.H. Ngo, L.D. Nghiem, F.I. Hai, J. Zhang, S. Liang,
665 X.C. Wang, A review on the occurrence of micropollutants in the aquatic
666 environment and their fate and removal during wastewater treatment, *Science*
667 *of the Total Environment*, 473-474 (2014) 619-641.
- 668 [2] T.A. Ternes, A. Joss, H. Siegrist, Scrutinizing pharmaceuticals and
669 personal care products in wastewater treatment, *Environmental Science and*
670 *Technology*, 38 (2004) 392A-399A.
- 671 [3] P. Verlicchi, M. Al Aukidy, E. Zambello, Occurrence of pharmaceutical
672 compounds in urban wastewater: Removal, mass load and environmental risk
673 after a secondary treatment-A review, *Science of the Total Environment*, 429
674 (2012) 123-155.
- 675 [4] K.O. K'Oreje, L. Vergeynst, D. Ombaka, P. De Wispelaere, M. Okoth,
676 H. Van Langenhove, K. Demeestere, Occurrence patterns of pharmaceutical
677 residues in wastewater, surface water and groundwater of Nairobi and
678 Kisumu city, Kenya, *Chemosphere*, 149 (2016) 238-244.
- 679 [5] D.W. Kolpin, E.T. Furlong, M.T. Meyer, E.M. Thurman, S.D. Zaugg,
680 L.B. Barber, H.T. Buxton, Pharmaceuticals, hormones, and other organic
681 wastewater contaminants in U.S. streams, 1999-2000: A national
682 reconnaissance, *Environmental Science and Technology*, 36 (2002) 1202-
683 1211.
- 684 [6] J.P. Bound, N. Voulvoulis, Predicted and measured concentrations for
685 selected pharmaceuticals in UK rivers: Implications for risk assessment,
686 *Water Research*, 40 (2006) 2885-2892.

687 [7] N.A. Alygizakis, P. Gago-Ferrero, V.L. Borova, A. Pavlidou, I.
688 Hatzianestis, N.S. Thomaidis, Occurrence and spatial distribution of 158
689 pharmaceuticals, drugs of abuse and related metabolites in offshore
690 seawater, *Science of the Total Environment*, 541 (2016) 1097-1105.

691 [8] S. Gaw, K.V. Thomas, T.H. Hutchinson, Sources, impacts and trends
692 of pharmaceuticals in the marine and coastal environment, *Philosophical
693 Transactions of the Royal Society B: Biological Sciences*, 369 (2014).

694 [9] R. Moreno-González, S. Rodriguez-Mozaz, M. Gros, D. Barceló, V.M.
695 León, Seasonal distribution of pharmaceuticals in marine water and sediment
696 from a mediterranean coastal lagoon (SE Spain), *Environmental Research*,
697 138 (2015) 326-344.

698 [10] J. Beausse, Selected drugs in solid matrices: A review of
699 environmental determination, occurrence and properties of principal
700 substances, *TrAC - Trends in Analytical Chemistry*, 23 (2004) 753-761.

701 [11] T.H. Miller, N.R. Bury, S.F. Owen, J.I. MacRae, L.P. Barron, A review
702 of the pharmaceutical exposome in aquatic fauna, *Environmental Pollution*,
703 239 (2018) 129-146.

704 [12] A. Cecinato, C. Balducci, M. Perilli, Illicit psychotropic substances in
705 the air: The state-of-art, *Science of The Total Environment*, 539 (2016) 1-6.

706 [13] Carvalho R.N. , Ceriani L. , Ippolito A. , L. T., Development of the First
707 Watch List under the Environmental Quality Standards Directive, European
708 Commission Joint Research Center, Ispra, Italy 2015,

709 [14] E. Archer, B. Petrie, B. Kasprzyk-Hordern, G.M. Wolfaardt, The fate of
710 pharmaceuticals and personal care products (PPCPs), endocrine disrupting
711 contaminants (EDCs), metabolites and illicit drugs in a WWTW and
712 environmental waters, *Chemosphere*, 174 (2017) 437-446.

713 [15] D.R. Baker, L. Barron, B. Kasprzyk-Hordern, Illicit and pharmaceutical
714 drug consumption estimated via wastewater analysis. Part A: Chemical
715 analysis and drug use estimates, *Science of the Total Environment*, 487
716 (2014) 629-641.

717 [16] C. Ort, A.L.N. van Nuijs, J.D. Berset, L. Bijlsma, S. Castiglioni, A.
718 Covaci, P. de Voogt, E. Emke, D. Fatta-Kassinos, P. Griffiths, F. Hernández,
719 I. González-Mariño, R. Grabic, B. Kasprzyk-Hordern, N. Mastroianni, A.
720 Meierjohann, T. Nefau, M. Östman, Y. Pico, I. Racamonde, M. Reid, J.
721 Slobodnik, S. Terzic, N. Thomaidis, K.V. Thomas, Spatial differences and
722 temporal changes in illicit drug use in Europe quantified by wastewater
723 analysis, *Addiction*, 109 (2014) 1338-1352.

724 [17] M. Riechel, A. Matzinger, E. Pawlowsky-Reusing, H. Sonnenberg, M.
725 Uldack, B. Heinzmann, N. Caradot, D. von Seggern, P. Rouault, Impacts of
726 combined sewer overflows on a large urban river – Understanding the effect
727 of different management strategies, *Water Research*, 105 (2016) 264-273.

728 [18] J.W. Armstrong, R.M. Thom, K.K. Chew, Impact of a combined sewer
729 overflow on the abundance, distribution and community structure of subtidal
730 benthos, *Marine Environmental Research*, 4 (1980) 3-23.

731 [19] R.P. Eganhouse, P.M. Sherblom, Anthropogenic organic contaminants
732 in the effluent of a combined sewer overflow: Impact on Boston Harbor,
733 *Marine Environmental Research*, 51 (2001) 51-74.

734 [20] T.T. Fong, M.S. Phanikumar, I. Xagorarakis, J.B. Rose, Quantitative
735 detection of human adenoviruses in wastewater and combined sewer

736 overflows influencing a Michigan river, Applied and Environmental
737 Microbiology, 76 (2010) 715-723.

738 [21] K.N. Irvine, G.W. Pettibone, Dynamics of indicator bacteria populations
739 in sediment and river water near a combined sewer outfall, Environmental
740 Technology (United Kingdom), 14 (1993) 531-542.

741 [22] J. Gasperi, S. Zgheib, M. Cladière, V. Rocher, R. Moillon, G.
742 Chebbo, Priority pollutants in urban stormwater: Part 2 - Case of combined
743 sewers, Water Research, 46 (2012) 6693-6703.

744 [23] J.H. Lee, K.W. Bang, Characterization of urban stormwater runoff,
745 Water Research, 34 (2000) 1773-1780.

746 [24] Creating a River Thames fit for our future: An updated strategic and
747 economic case for the Thames Tideway Tunnel, 2015, Department for
748 Environment, Food and Rural Affairs (DEFRA),
749 [https://assets.publishing.service.gov.uk/government/uploads/system/uploads/
750 attachment_data/file/471847/thames-tideway-tunnel-strategic-economic-
751 case.pdf](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/471847/thames-tideway-tunnel-strategic-economic-case.pdf)

752 [25] N. Lloyd, Thames Tunnel Evidence Assessment Final Report,
753 Department of Environment, Food and Rural Affairs (DEFRA), 2012,
754 [https://assets.publishing.service.gov.uk/government/uploads/system/uploads/
755 attachment_data/file/69532/pb13748-thamestunnel-evidence-assessment.pdf](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/69532/pb13748-thamestunnel-evidence-assessment.pdf)

756 [26] Thames Tideway Strategic Study: Executive Summary Thames Water
757 Utilities Ltd. Doc Ref: 8.1.1 2013,
758 [https://infrastructure.planninginspectorate.gov.uk/wp-
759 content/ipc/uploads/projects/WW010001/WW010001-001255-
760 8.1.1_Thames_Tideway_Strategic_Study_Executive_Summary.pdf](https://infrastructure.planninginspectorate.gov.uk/wp-content/ipc/uploads/projects/WW010001/WW010001-001255-8.1.1_Thames_Tideway_Strategic_Study_Executive_Summary.pdf)

761 [27] The Thames Tideway Scheme, [https://www.tideway.london/the-
762 tunnel/our-solution/](https://www.tideway.london/the-tunnel/our-solution/), Date Accessed: 03/10/2018

763 [28] P. Kay, S.R. Hughes, J.R. Ault, A.E. Ashcroft, L.E. Brown, Widespread,
764 routine occurrence of pharmaceuticals in sewage effluent, combined sewer
765 overflows and receiving waters, Environmental Pollution, 220 (2017) 1447-
766 1455.

767 [29] D. Schowanek, K. Fox, M. Holt, F.R. Schroeder, V. Koch, G. Cassani,
768 M. Matthies, G. Boeije, P. Vanrolleghem, A. Young, G. Morris, C. Gandolfi,
769 T.C.J. Feijtel, GREAT-ER: A new tool for management and risk assessment
770 of chemicals in river basins contribution to GREAT-ER #10, Water Science
771 and Technology, 2001, pp. 179-185.

772 [30] M.J. Benotti, B.J. Brownawell, Distributions of pharmaceuticals in an
773 urban estuary during both dry- and wet-weather conditions, Environmental
774 Science and Technology, 41 (2007) 5795-5802.

775 [31] P. Weyrauch, A. Matzinger, E. Pawlowsky-Reusing, S. Plume, D. von
776 Seggern, B. Heinzmann, K. Schroeder, P. Rouault, Contribution of combined
777 sewer overflows to trace contaminant loads in urban streams, Water
778 Research, 44 (2010) 4451-4462.

779 [32] A.-S. Madoux-Humery, S. Dorner, S. Sauvé, K. Aboufadi, M.
780 Galarneau, P. Servais, M. Prévost, Temporal variability of combined sewer
781 overflow contaminants: Evaluation of wastewater micropollutants as tracers of
782 fecal contamination, Water Research, 47 (2013) 4370-4382.

783 [33] S. Sauvé, K. Aboufadi, S. Dorner, P. Payment, G. Deschamps, M.
784 Prévost, Fecal coliforms, caffeine and carbamazepine in stormwater collection
785 systems in a large urban area, Chemosphere, 86 (2012) 118-123.

786 [34] I.J. Buerge, T. Poiger, M.D. Müller, H.R. Buser, Caffeine, an
787 anthropogenic marker for wastewater contamination of surface waters,
788 *Environmental Science and Technology*, 37 (2003) 691-700.

789 [35] N. Sankararamakrishnan, Q. Guo, Chemical tracers as indicator of
790 human fecal coliforms at storm water outfalls, *Environment International*, 31
791 (2005) 1133-1140.

792 [36] I.J. Buerge, T. Poiger, M.D. Müller, H.R. Buser, Combined sewer
793 overflows to surface waters detected by the anthropogenic marker caffeine,
794 *Environmental Science and Technology*, 40 (2006) 4096-4102.

795 [37] K.A. Peeler, S.P. Opsahl, J.P. Chanton, Tracking anthropogenic inputs
796 using caffeine, indicator bacteria, and nutrients in rural freshwater and urban
797 marine systems, *Environmental Science and Technology*, 40 (2006) 7616-
798 7622.

799 [38] M.A. Launay, U. Dittmer, H. Steinmetz, Organic micropollutants
800 discharged by combined sewer overflows – Characterisation of pollutant
801 sources and stormwater-related processes, *Water Research*, 104 (2016) 82-
802 92.

803 [39] B. Kasprzyk-Hordern, R.M. Dinsdale, A.J. Guwy, Illicit drugs and
804 pharmaceuticals in the environment - Forensic applications of environmental
805 data, Part 2: Pharmaceuticals as chemical markers of faecal water
806 contamination, *Environmental Pollution*, 157 (2009) 1778-1786.

807 [40] L.J. Fono, D.L. Sedlak, Use of the chiral pharmaceutical propranolol to
808 identify sewage discharges into surface waters, *Environmental Science and*
809 *Technology*, 39 (2005) 9244-9252.

810 [41] C. Fenech, K. Nolan, L. Rock, A. Morrissey, An SPE LC–MS/MS
811 method for the analysis of human and veterinary chemical markers within
812 surface waters: An environmental forensics application, *Environmental*
813 *Pollution*, 181 (2013) 250-256.

814 [42] P.A. Lara-Martín, E. González-Mazo, M. Petrovic, D. Barceló, B.J.
815 Brownawell, Occurrence, distribution and partitioning of nonionic surfactants
816 and pharmaceuticals in the urbanized Long Island Sound Estuary (NY),
817 *Marine Pollution Bulletin*, 85 (2014) 710-719.

818 [43] H. Zhao, J.L. Zhou, J. Zhang, Tidal impact on the dynamic behavior of
819 dissolved pharmaceuticals in the Yangtze Estuary, China, *Science of The*
820 *Total Environment*, 536 (2015) 946-954.

821 [44] Q. Sun, Y. Li, M. Li, M. Ashfaq, M. Lv, H. Wang, A. Hu, C.-P. Yu,
822 PPCPs in Jiulong River estuary (China): Spatiotemporal distributions, fate,
823 and their use as chemical markers of wastewater, *Chemosphere*, 150 (2016)
824 596-604.

825 [45] R. Bade, L. Bijlsma, T.H. Miller, L.P. Barron, J.V. Sancho, F.
826 Hernández, Suspect screening of large numbers of emerging contaminants in
827 environmental waters using artificial neural networks for chromatographic
828 retention time prediction and high resolution mass spectrometry data analysis,
829 *Science of the Total Environment*, 538 (2015) 934-941.

830 [46] F. Hernández, M. Ibáñez, R. Bade, L. Bijlsma, J.V. Sancho,
831 Investigation of pharmaceuticals and illicit drugs in waters by liquid
832 chromatography-high-resolution mass spectrometry, *TrAC - Trends in*
833 *Analytical Chemistry*, 63 (2014) 140-157.

834 [47] C.B. Mollerup, M. Mardal, P.W. Dalsgaard, K. Linnet, L.P. Barron,
835 Prediction of collision cross section and retention time for broad scope

836 screening in gradient reversed-phase liquid chromatography-ion mobility-high
837 resolution accurate mass spectrometry, *Journal of Chromatography A*, 1542
838 (2018) 82-88.

839 [48] L.P. Barron, G.L. McEneff, Gradient liquid chromatographic retention
840 time prediction for suspect screening applications: A critical assessment of a
841 generalised artificial neural network-based approach across 10 multi-residue
842 reversed-phase analytical methods, *Talanta*, 147 (2016) 261-270.

843 [49] M. Tanguy, H. Dixon, I. Prosdocimi, D.G. Morris, V.D.J. Keller, Gridded
844 estimates of daily and monthly areal rainfall for the United Kingdom (1890-
845 2012) [CEH-GEAR], NERC Environmental Information Data Centre, 2014,
846 <https://doi.org/10.5285/5dc179dc-f692-49ba-9326-a6893a503f6e>

847 [50] Validation of analytical procedures: text and methodology Q2(R1),
848 International Conference on Harmonisation tripartite guidelines, 2005.

849 [51] K. Munro, T.H. Miller, C.P.B. Martins, A.M. Edge, D.A. Cowan, L.P.
850 Barron, Artificial neural network modelling of pharmaceutical residue retention
851 times in wastewater extracts using gradient liquid chromatography-high
852 resolution mass spectrometry data, *Journal of Chromatography A*, 1396
853 (2015) 34-44.

854 [52] M. Gardner, S. Comber, M. Scrimshaw, E. Cartmell, J. Lester, B. Ellor,
855 The significance of hazardous chemicals in wastewater treatment works
856 effluents, *Science of the Total Environment*, 437 (2012) 363-372.

857 [53] S. Comber, M. Gardner, V. Jones, B. Ellor, Source Apportionment of
858 Trace Contaminants in Urban Sewer Catchments, *Environmental Technology*
859 36 (2014) 573-587.

860 [54] S. Comber, M. Gardner, P. Sorme, D. Leverett, B. Ellor, Active
861 Pharmaceutical Ingredients Entering the Aquatic Environment From
862 Wastewater Treatment Works: A Cause for Concern?, *Science of the Total*
863 *Environment*, 613-614 (2018) 538-547.

864 [55] J. Li, J. Gao, P.K. Thai, X. Sun, J.F. Mueller, Z. Yuan, G. Jiang,
865 Stability of Illicit Drugs as Biomarkers in Sewers: From Lab to Reality,
866 *Environmental Science and Technology*, 52 (2018) 1561-1570.

867 [56] M.J. Andrés-Costa, N. Rubio-López, M. Morales Suárez-Varela, Y.
868 Pico, Occurrence and removal of drugs of abuse in Wastewater Treatment
869 Plants of Valencia (Spain), *Environmental Pollution*, 194 (2014) 152-162.

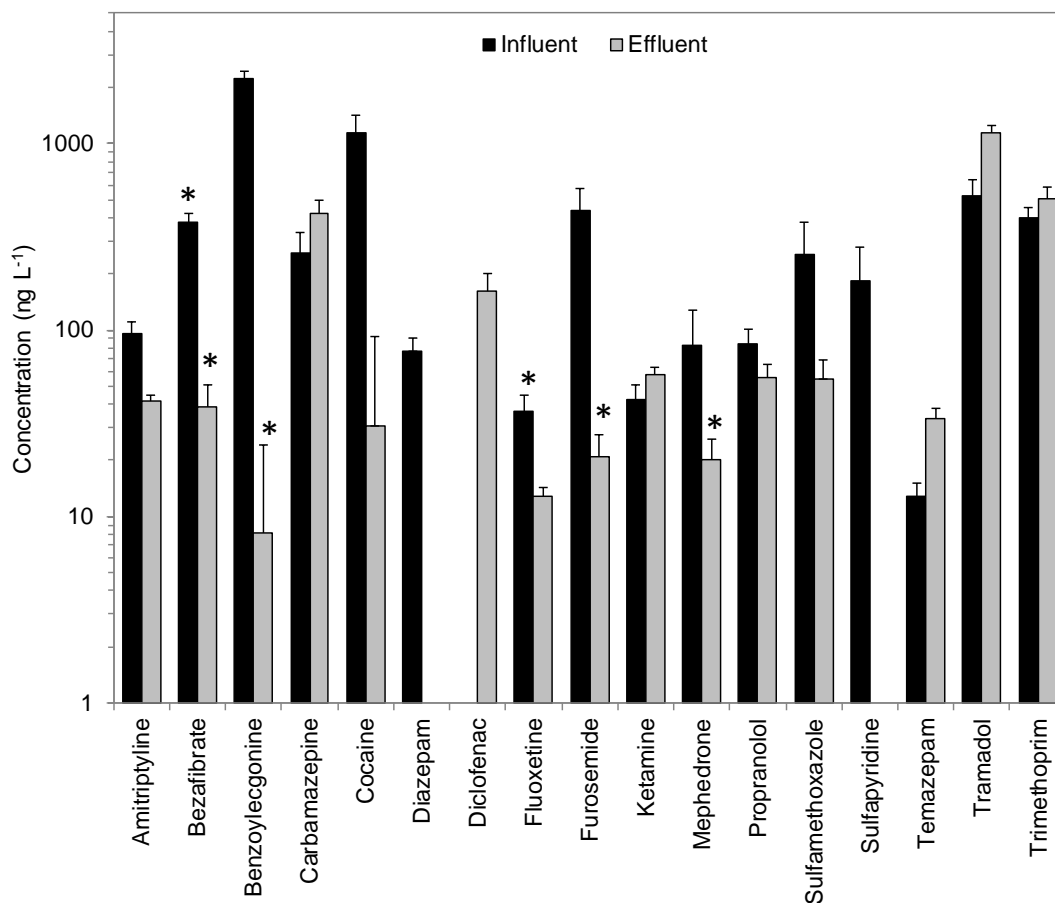
870 [57] D.R. Baker, B. Kasprzyk-Hordern, Spatial and temporal occurrence of
871 pharmaceuticals and illicit drugs in the aqueous environment and during
872 wastewater treatment: New developments, *Science of the Total Environment*,
873 454-455 (2013) 442-456.

874 [58] D.R. Baker, V. Očenášková, M. Kvicálová, B. Kasprzyk-Hordern, Drugs
875 of abuse in wastewater and suspended particulate matter — Further
876 developments in sewage epidemiology, *Environment International*, 48 (2012)
877 28-38.

878 [59] C. Postigo, M.J. López de Alda, D. Barceló, Drugs of abuse and their
879 metabolites in the Ebro River basin: Occurrence in sewage and surface water,
880 sewage treatment plants removal efficiency, and collective drug usage
881 estimation, *Environment International*, 36 (2010) 75-84.

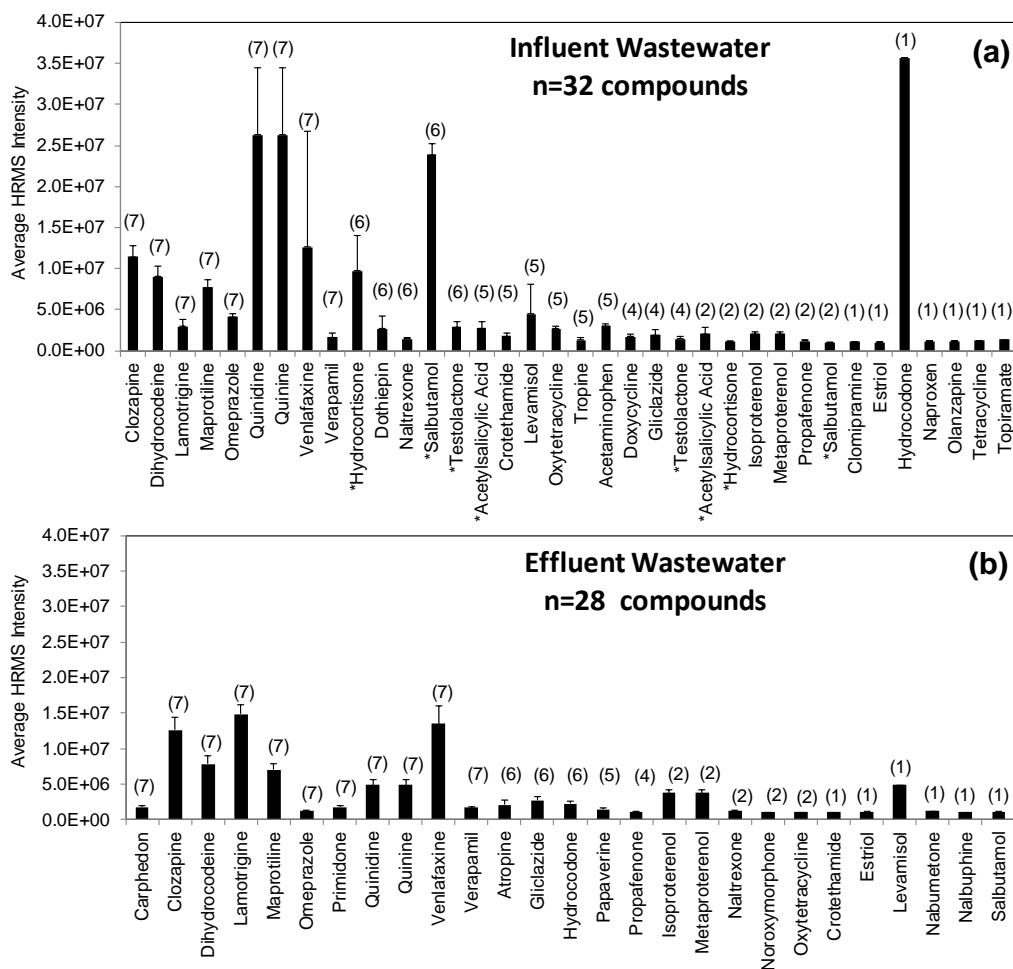
882 [60] J. Wieber, R. Gugler, J.H. Hengstmann, H.J. Dengler,
883 Pharmacokinetics of ketamine in man, *Anaesthetist*, 24 (1975) 260-263.

884 [61] P. Ramin, F. Polesel, A.L. Brock, B.G. Plósz, The impact of
885 temperature on the transformation of illicit drug biomarkers in wastewater,
886 Science of the Total Environment, 644 (2018) 1612-1616.
887 [62] M. Chtourou, M. Mallek, M. Dalmau, J. Mamo, E. Santos-Clotas, A.B.
888 Salah, K. Walha, V. Salvadó, H. Monclús, Triclosan, carbamazepine and
889 caffeine removal by activated sludge system focusing on membrane
890 bioreactor, Process Safety and Environmental Protection, 118 (2018) 1-9.
891 [63] B. Reoyo-Prats, D. Aubert, A. Sellier, B. Roig, C. Palacios, Dynamics
892 and sources of pharmaceutically active compounds in a coastal
893 Mediterranean river during heavy rains, Environmental Science and Pollution
894 Research, 25 (2018) 6107-6121.
895 [64] Thames Sewage Events, Thames Anglers Conservancy,
896 <http://www.rivertac.org/thames-sewage-events/>, Date Accessed: 01/10/2018
897 [65] W.C. Scott, C.S. Breed, S.P. Haddad, S.R. Burket, G.N. Saari, P.J.
898 Pearce, C.K. Chambliss, B.W. Brooks, Spatial and temporal influence of
899 onsite wastewater treatment systems, centralized effluent discharge, and
900 tides on aquatic hazards of nutrients, indicator bacteria, and pharmaceuticals
901 in a coastal bayou, Science of the Total Environment, 650 (2019) 354-364.
902 [66] K. Stein, M. Ramil, G. Fink, M. Sander, T.A. Ternes, Analysis and
903 Sorption of Psychoactive Drugs onto Sediment, Environmental Science &
904 Technology, 42 (2008) 6415-6423.
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Figure 1. Differential analysis of drug occurrence in untreated influent and treated effluent wastewaters from a major treatment works in London in n=7 consecutive 24-h composite samples in March, 2014. Bars marked with * represent semi-quantitative measurements as values were <LLOQ, but >LOD. Error bars represent the standard deviation of the means of all measurements for each compound across the 7-day period.

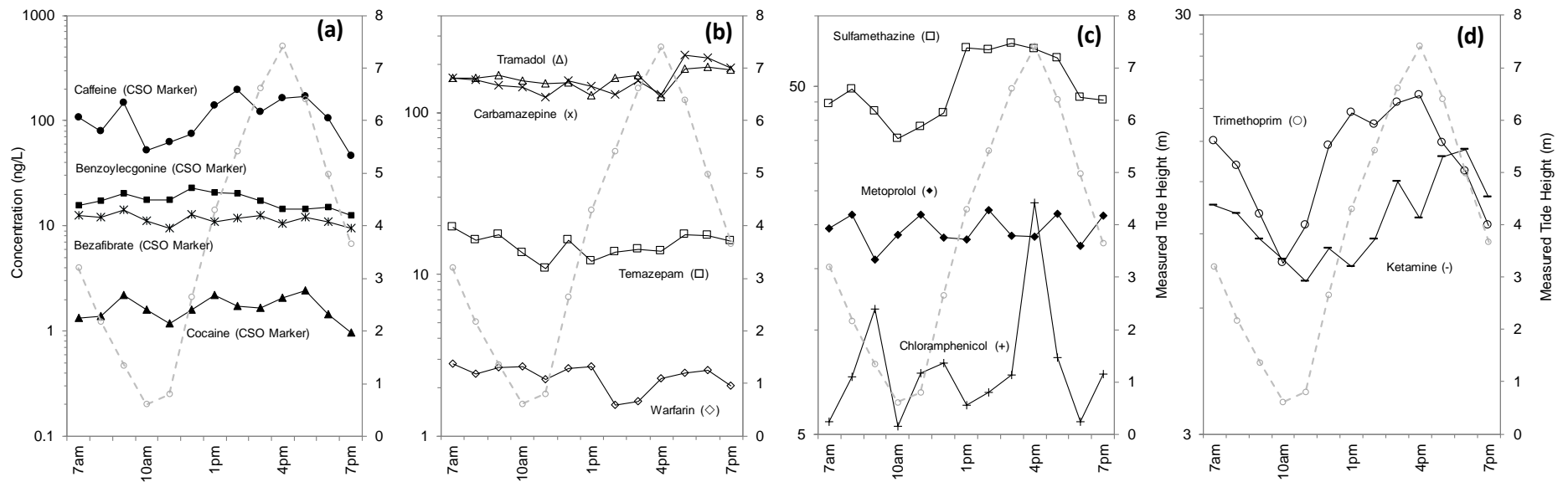


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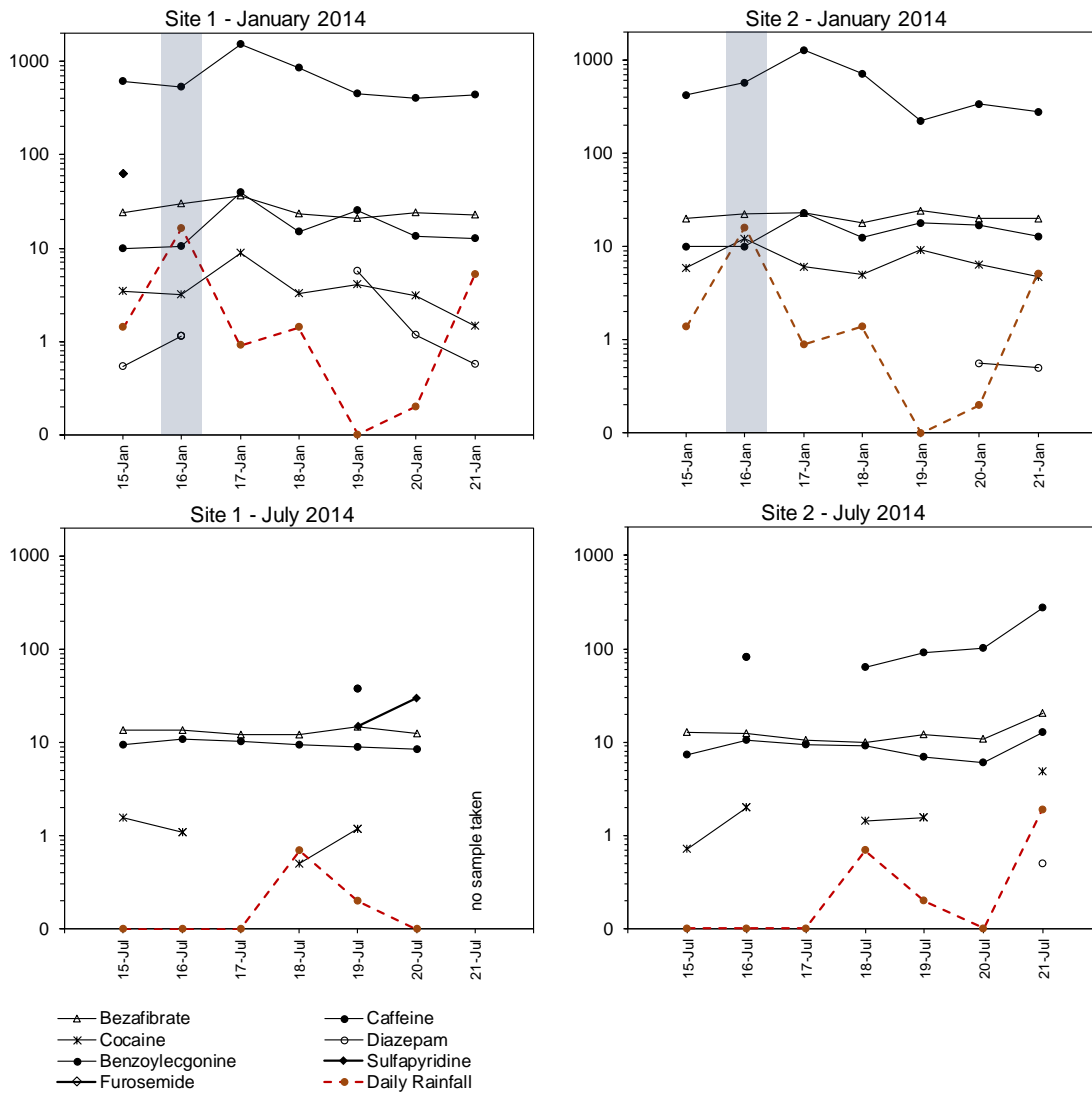
Figure 2. Average signal intensity for each compound tentatively identified by retrospective *in silico* suspect screening in (a) untreated influent and (b) treated effluent wastewaters. Their corresponding occurrence frequency out of 7 days is shown in parenthesis. Bars represent the mean and whiskers represent the standard deviation of that number of daily measurements in (c) and (d). Compounds marked with * represent those where two matching predicted t_R values (± 1.30 min threshold) and HRMS signals ($\delta < 5$ ppm for $[M+H]^+$ or $[M-H]^-$) were obtained.

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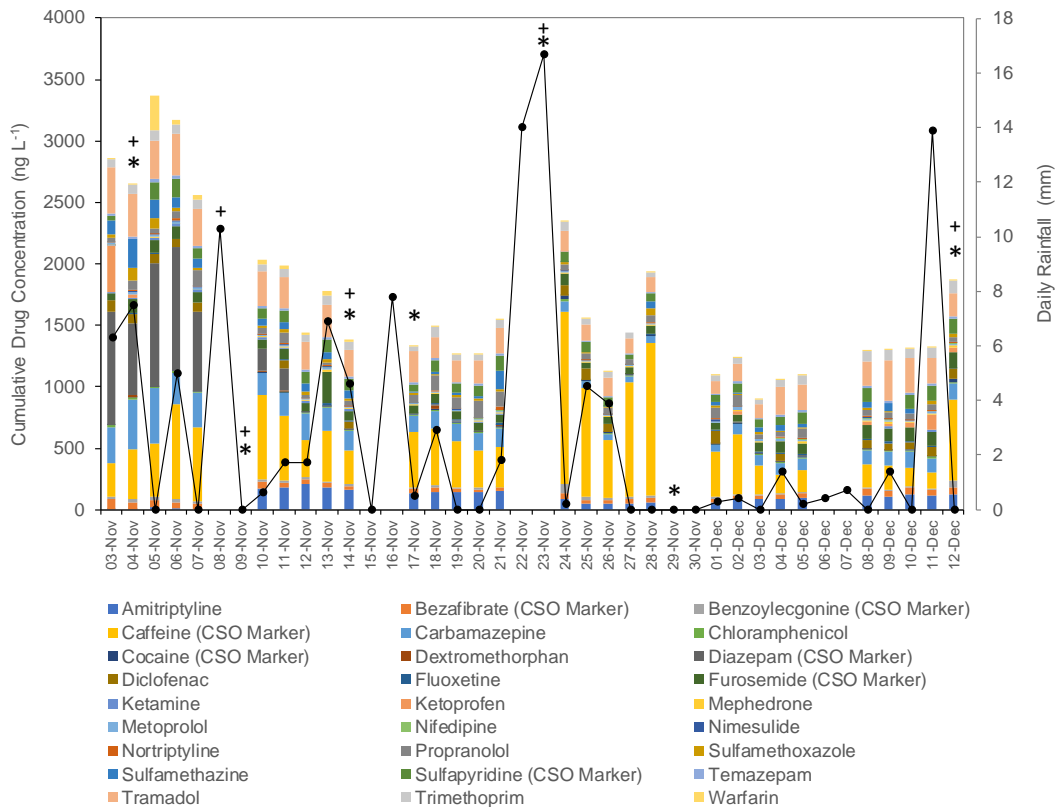
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Figure 3. Diurnal variation in (a) CSO marker drug compounds and (b)-(d) all other drug compounds determined above the LLOQ in the River Thames on the 14th August, 2014. Black datapoints represent the mean of n=2 replicate grab sample analyses. Grey dashed lines represent the measured tide height at the time of sampling. No CSOs occurred on this day (<1 mm rainfall).



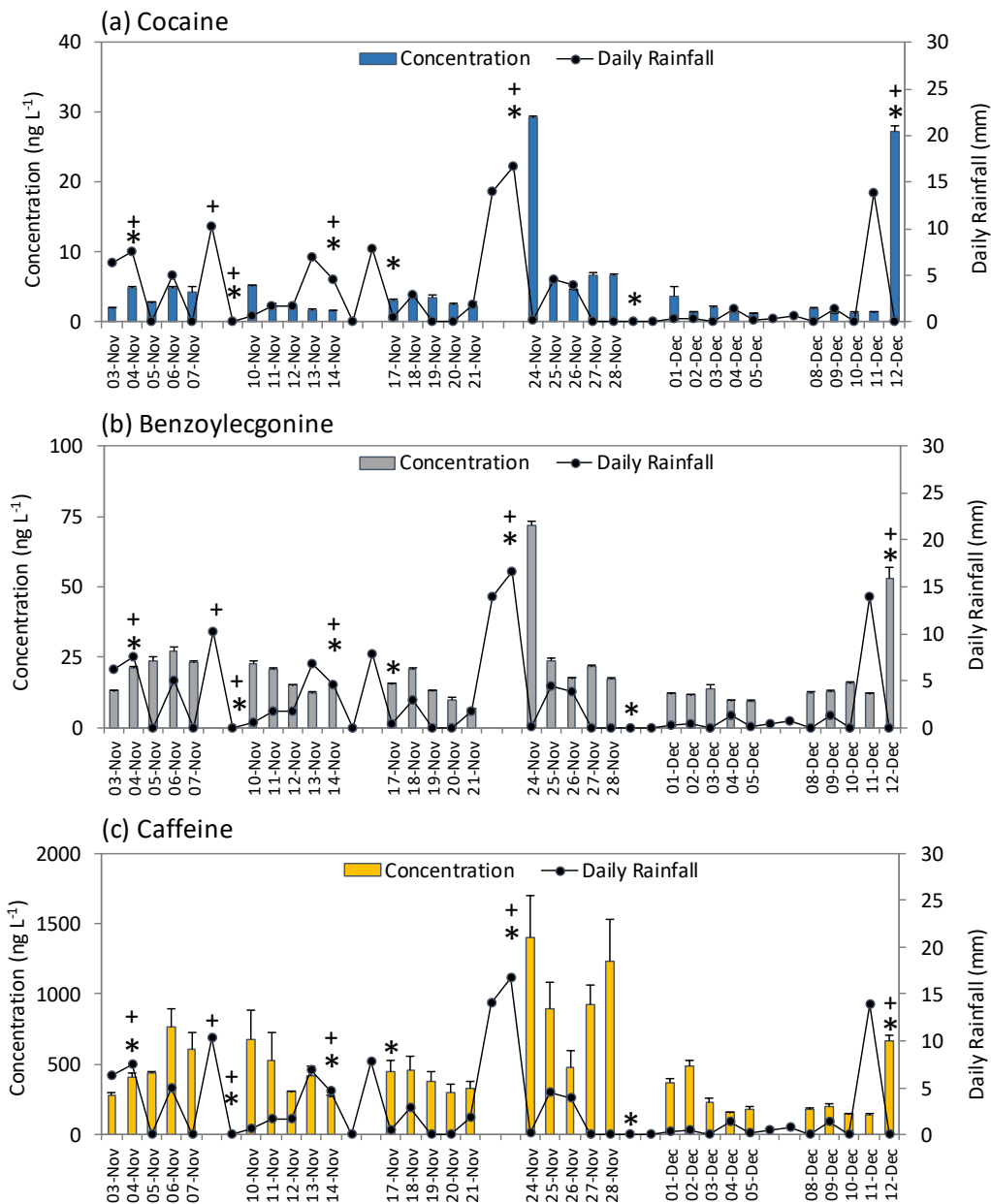
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Figure 4. Measured concentrations of seven shortlisted candidate drug CSO markers in samples of Thames River water from two sites in January and July 2014 and overlaid with daily rainfall. A Type 1 CSO occurred on on 17th January, 2014 at 21:50 hours (shaded in grey). Note: No sample was taken from Site 1 on 21st July, 2014. All measurements represent the mean of n=2 replicates.



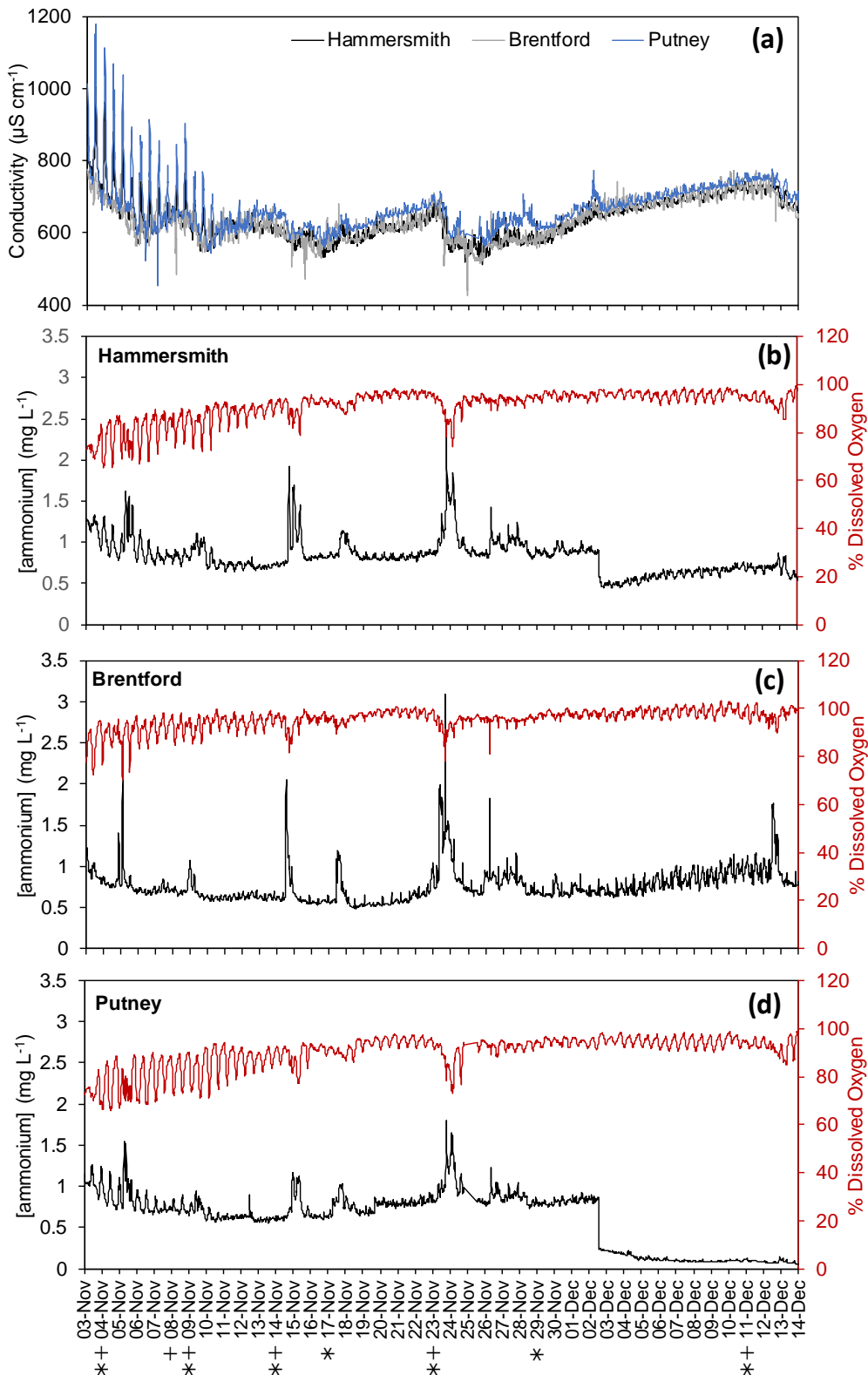
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Figure 5. Cumulative concentration of all drug residues determined on weekdays in the River Thames across Nov-Dec, 2014. Dates marked with + are Type 1 CSOs where storm water and untreated sewage were combined and released directly into the river. Dates marked with * represent Type 2 CSO events where storm water was mixed with treated wastewater effluent at a WWTP and then released into the river (where both + and * exist, two such CSOs occurred on the same date, also see Table S11).



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Figure 6. Occurrence of three drug CSO markers in river water from the Thames over six weeks in Nov-Dec, 2014 (overlaid with daily rainfall). Dates marked with + or * are as in Figure 5. Bars represent the mean of two replicates and whiskers represent the maximum value measured.



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957 **Figure 7.** Continuous monitoring data at three sites on the River Thames in
 958 Nov-Dec, 2014 for (a) conductivity and (b)-(d) % DO (red)/ammonium
 959 concentration (black) at Hammersmith, Brentford and Putney sites,
 960 respectively. Data-acquisition frequency =15 min. Dates marked with +/*
 961 represent CSO Types 1 and/or 2, respectively.