

1 **Limitation of flow effect on passive sampling accuracy**
2 **using POCIS with the PRC approach or o-DGT: a pilot-scale**
3 **evaluation for pharmaceutical compounds.**

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9 **Abstract**

10 Flow velocity is known to alter passive sampling accuracy. We investigated the POCIS (Polar
11 Organic Chemical Integrative Sampler) with PRC (Performance Reference Compounds) approach and
12 Diffusive Gradients in Thin Films samplers (o-DGT) to limit the effect of flow on the quantification
13 accuracy of ten model pharmaceuticals compounds ($0.16 \leq \log K_{ow} \leq 4.51$). POCIS and o-DGT
14 samplers were exposed for seven days in controlled pilot-scale (hundreds of liters) experiments
15 under quiescent or flowing ($2 < V < 18 \text{ cm s}^{-1}$) conditions. Under flowing conditions, both POCIS-PRC
16 and o-DGT efficiently limited the flow effect ~~for the other compounds~~ and led, in most cases, to
17 biases within analytical uncertainty (20%). Under quiescent conditions, o-DGT performed accurately
18 (bias < 30% for most compounds) whereas the PRC approach was unsuitable to improve upon the
19 accuracy of POCIS (PRC was unable to desorb). Therefore, both approaches are helpful in limiting the
20 effects of flow on accuracy, but only o-DGT is efficient in quiescent conditions. However, o-DGT
21 currently suffers from poorer sensitivity compared to POCIS, but the future development of o-DGT
22 devices with wider windows could overcome this limitation.

23 **Keywords**

24 Polar Organic Chemical Integrative Sampler (POCIS)

25 Diffusive Gradients in Thin Films (DGT)

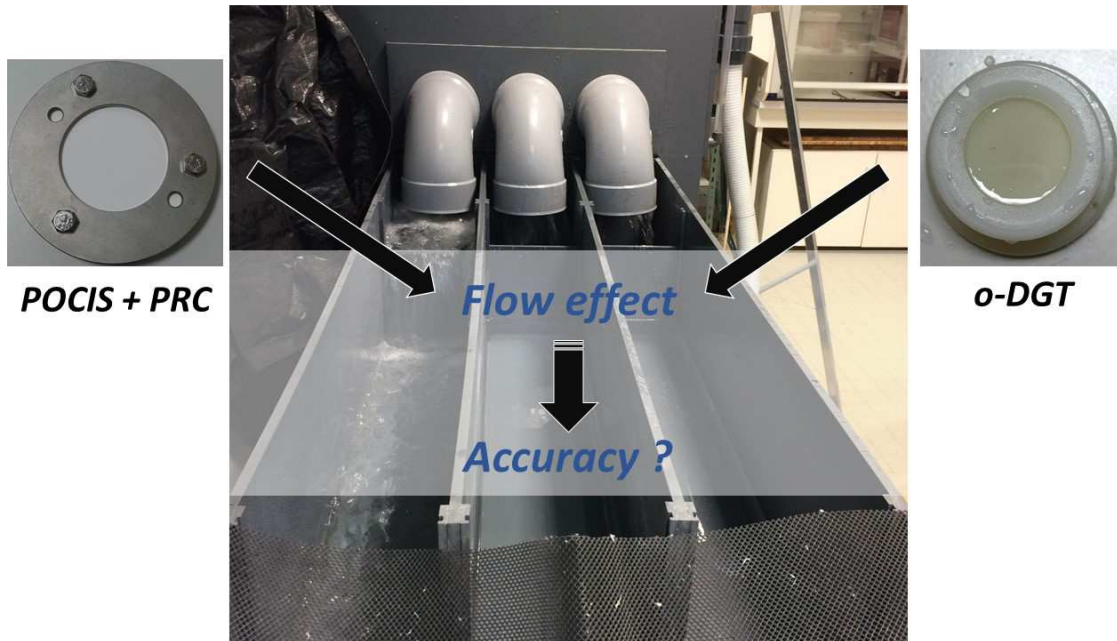
26 Time-weighted average concentration

27 Flow velocity

28 Accuracy

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30 **Graphical abstract.**



31

32 **1. Introduction**

33 Passive sampling has been used since 1999 for polar and semi-polar organic micropollutant
34 monitoring in aquatic systems (Stuer-Lauridsen, 2005). Compared to conventional sampling (*i.e.*, grab
35 sampling), passive sampling allows easier access to time-weighted average concentrations and
36 increased sensitivity, thanks to its *in situ* concentration ability. However, passive sampling accuracy is
37 known to be altered by several environmental factors (Gong et al., 2018; Harman et al., 2012; Li et

38 al., 2011). Among them, flow velocity is of particular concern, and conventional calibrations (*i.e.*
39 determination of sampling rates) under laboratory controlled conditions can lead to more than 100%
40 inaccuracy for field deployments (Poulier et al., 2014). Several strategies have been identified to
41 correct flow effects (Fauvelle et al., 2017). The first one is to establish empirical relationships
42 between sampling rate and flow velocity. This has been done for several polar organic compounds (Li
43 et al., 2010), but these relationships do not allow sampling rate estimations at zero flow. Moreover,
44 correcting for significant flow variations during the samplers' exposure could be tricky with such a
45 strategy. In this context, *in situ* correction is preferable. Performance Reference Compounds (PRC)
46 have been proposed as an *in situ* correction method for the effects of environmental factors,
47 including flow velocity. Initially developed for hydrophobic compounds in Semipermeable Membrane
48 Devices (Huckins et al., 2002), the PRC approach corrects the targeted compound sampling rate
49 relative to the *in situ* desorption rate of a reference compound, assuming isotropic exchange. The
50 PRC approach has been successfully developed for several polar pesticides (Mazzella et al., 2010), but
51 it might not work for all compounds and all exposures (Booij and Chen, 2018; Harman et al., 2011).
52 The last strategy identified is to increase the sampler's membrane resistance to limit the influence of
53 its surrounding conditions. This strategy is implemented in the Diffusive Gradients in Thin Films (DGT)
54 technique, which was initially developed for metals (Davison and Zhang, 1994), was recently adapted
55 for organic compounds (Challis et al., 2016; Chen et al., 2012; Guibal et al., 2017) (o-DGT). Compared
56 to other passive samplers, DGT devices include a diffusive gel (typically 0.8 mm thick) that constrains
57 mass transfer mostly to the diffusion rate within the gel (Davison and Zhang, 2012). However, this
58 tool is ~~however~~ not totally independent of flow velocity, as previously shown for metals (Gimpel et
59 al., 2001). In fact, under low flow conditions, a significant diffusive boundary layer (DBL) is created in
60 front of the sampler (Challis et al., 2016), altering the diffusion path to be considered. Although
61 promising, all of these strategies have limitations. Their implementation is limited in the literature,
62 and data on their efficiency are too scarce to provide guidance for choosing a relevant strategy to
63 correct flow effects.

64 This work focuses on the implementation of PRC with POCIS (Polar Organic Chemical
65 Integrative Sampler) and o-DGT for correcting the effects of flow on the passive sampling of ten
66 model pharmaceuticals compounds ($0.16 \leq \log K_{OW} \leq 4.51$). POCIS and o-DGT samplers were
67 submitted to different flow velocities in controlled pilot-scale (hundreds of liters) experiments. The
68 accuracy of the different strategies (POCIS without correction, POCIS with PRC and o-DGT) was
69 evaluated for quiescent and flowing conditions.

70 2. Material and methods

71 2.1. Chemicals

72 A Gradient A10 Milli-Q system from Millipore produced ultra-pure water (UPW). Unless stated
73 otherwise, all solvents were of LC-MS grade and reagents of analytical grade. The following
74 pharmaceuticals were used and had a purity greater than 97%: atenolol (ATE), carbamazepine (CAR),
75 diclofenac (DIC), fluoxetine (FLU), ketoprofen (KET), metoprolol (MET), paroxetine (PAR), propranolol
76 (PRO), sulfamethoxazole (SUL) and trimethoprim (TRI). Stock pharmaceutical solutions (100 mg L^{-1}),
77 working solutions (containing each pharmaceuticals at 1 mg L^{-1}) and internal standard solutions (10
78 mg L^{-1} , presented in Table 1) were prepared in methanol and stored at -18 C . Deuterated atrazine
79 desisopropyl (DIA-d5) was used as PRC for POCIS samplers.

80 *Table 1: Pharmaceuticals characteristics after HPLC separation and mass detection and internal standards used.*

Pharmaceutical	Log K_{OW}	Retention time (min)	Mass (g mol ⁻¹)	Internal standard
Atenolol	0.16	2.32	266.1630	Salbutamol-d3
Carbamazepine	2.45	7.70	236.0950	Carbamazepine-d10
Diclofenac	4.51	9.07	295.0167	Diclofenac-d4
Fluoxetine	4.05	7.88	309.1341	Carbamazepine-d10
Ketoprofen	3.12	8.08	254.0943	Diclofenac-d4

Metoprolol	1.88	6.52	267.1834	Carbamazepine-d10
Paroxetine	1.23	6.69	329.1427	Carbamazepine-d10
Propranolol	3.48	7.30	259.1572	Propranolol-d3
Sulfamethoxazole	0.89	5.86	253.0521	Sulfamethoxazole-d4
Trimethoprim	0.91	4.75	290.1379	Trimethoprim-d3

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82 2.2. Description of pilots

83 Control of experimental conditions is required to isolate the effects of flow. Pilot scale
84 experiments offer the best method to mimic field conditions while maintaining experimental control.
85 Two pilots were used to mimic the flow conditions a passive sampler is exposed to: an artificial river
86 and a tank (Figure S1). The tank (200 L) was used to simulate a quiescent system ($V_0 = 0 \text{ cm s}^{-1}$). The
87 artificial river (500 L) was fed with a pump (flow rate = $13 \text{ m}^3 \text{ h}^{-1}$) and divided into 3 channels (20.3
88 cm width and 152 cm length) with different flow velocities using a gate system. The mean velocities
89 measured during exposures were $V_1 = 2.5 \pm 1.2 \text{ cm s}^{-1}$; $V_2 = 6.3 \pm 1.0 \text{ cm s}^{-1}$ and $V_3 = 17.5 \pm 2.1 \text{ cm s}^{-1}$
90 ($n=8$). Temperature was recorded every 10 min using a Tinytag temperature logger (TG-4100) and
91 were $17 \pm 2^\circ\text{C}$ and $15 \pm 1^\circ\text{C}$ during POCIS and o-DGT exposure, respectively.

92 All pilots were fed with tap water (composition shown in Table S1) spiked with the 10
93 pharmaceutical compounds at an initial concentration of $0.5 \mu\text{g L}^{-1}$ each. Continuous renewal of the
94 water was performed at a rate of 15% volume per day. Water samples were taken four times during
95 each experiment (at the beginning and after 3, 5 and 7 days) for pharmaceutical analysis (average
96 concentrations are displayed in Table S2). A total of 100 mL of sample was filtered, adjusted to pH 7
97 and spiked with 10 μL of surrogates (simazine-d5, monuron-d6 and prometryn-d6) to identify any
98 loss during the extraction step. Water samples were then transferred to SPE cartridges
99 (Chromabond® HR-X; 60 mg, 3 mL 85 μm , preconditioned with 5 mL of methanol then 5 mL of UPW)

100 using a GX-241 automated system from Gilson. Water sample cartridges were finally dried and stored
101 at -18°C before extraction.

102 2.3. Passive sampler preparation and exposure

103 POCIS were prepared by enclosing 200 mg of Oasis® HLB receiving phase within two
104 polyethersulfone (PES) membranes, held by two stainless steel rings (20.5 cm² window). The Oasis®
105 HLB was previously spiked with 4 µg g⁻¹ of DIA-d5 (PRC) (Mazzella et al., 2010), and the PES
106 membranes (90 mm diameter and 0.1 µm pore size from Pall Corporation) were previously washed
107 according to Guibal et al. (2015). Duplicate POCIS were exposed parallel to the flow for 7 days at each
108 flow velocity. Potential re-adsorption of PRC was checked using duplicate of POCIS blank (without
109 PRC) exposed alongside the POCIS with PRC and was found to be insignificant (<1%). After exposure,
110 POCIS were dismantled and the receiving phases were transferred to SPE cartridges for extraction.
111 The exact mass of the receiving phase recovered for extraction was determined for estimation of
112 compound concentration in water.

113 o-DGT samplers were prepared by enclosing a disc of binding gel and of diffusive gel in a piston
114 type holder (3.14 cm² window, purchased from DGT Research). Gels were prepared according to
115 Challis et al. (2016) and were composed of Oasis® HLB receiving phase embedded in 1.5% agarose gel
116 (25 mg Oasis® HLB per disc, nominal) and 1.5% agarose gel only for the binding and diffusive gels,
117 respectively. Diffusive gels were prepared with four different thicknesses: 0.16, 0.41, 0.61 and 0.84
118 mm (RSD ≤ 1.5%, n=4). Triplicate o-DGT samplers of each thickness were exposed parallel to the flow
119 for 7 days at each flow velocity. After exposure, o-DGT samplers were dismantled and binding gel
120 discs were recovered for extraction.

121 2.4. Pharmaceutical extraction and analysis

122 POCIS and water sample cartridges were eluted with 3 mL of methanol followed by 3 mL of a
123 mixture of 75:25 v:v methanol:ethyl acetate. After elution, ten µL of a solution with internal
124 standards were added to the water extracts. Extracts were then evaporated to dryness under

125 nitrogen flow and then reconstituted with 1 mL of methanol for POCIS extracts or 90:10
126 UPW:methanol for water extracts. POCIS extracts were diluted 10 times, and 10 µL of internal
127 standards solution was added.

128 o-DGT binding gel discs (containing Oasis® HLB receiving phase) were extracted with 3x3 mL of
129 methanol under sonication (210 W for 2 minutes). Extracts were spiked with 10 µL of internal
130 standards solution and evaporated to dryness under nitrogen flow. Extracts were finally
131 reconstituted with 1 mL of 90:10 UPW:methanol.

132 All samples were analyzed with an HPLC Infinity 1290 coupled with a Q-ToF 6540 equipped with
133 a Jet Stream electrospray ionization source (Agilent). The procedure is fully detailed in Guibal et al.
134 (2018). Briefly, chromatographic separation was performed with a RP18+ Nucleoshell column
135 (Macherey-Nagel), and UPW and methanol (with 5 mM ammonium formate and 0.1% acid formic for
136 both) were used as eluent during a 16 min analytical gradient. Autosampler and column
137 temperatures were kept at 4°C and 30°C, respectively. Mass acquisition was operated in the “all-
138 ions” positive mode (collision energies: 0, 10, 20 and 40 V). QA/QC was used to control any
139 deviations during analysis. Pharmaceutical characteristics after HPLC separation and mass detection
140 are displayed in Table 1 and analytical performances in Table S3.

141 2.5. Exposure concentration estimation

142 Time-weighted average concentrations for the pilots (C_w) were estimated using both POCIS
143 and o-DGT. Two kinds of estimations were done: a “standard” and an “advanced” estimation (see
144 details below). The “standard” estimation aims to represent a routine estimation and was performed
145 with simple and widespread strategies (devices and data treatment) from literature. The “advanced”
146 determination was performed using more sophisticated strategies from the literature that are
147 believed to correct for flow effect.

148 2.5.1. Estimation based on POCIS

149 The “standard” estimation of C_w based on POCIS (n=2) was performed using Eq. 1 (Alvarez et
150 al., 2004):

151
$$C_w = m/R_S t \quad (1)$$

152 where m is the accumulated mass of compound in the sampler, R_S is the sampling rate, and t
153 is the deployment time. For routine purposes, dedicated calibrations are not conceivable, and R_S
154 values must be taken from the literature. For a given compound, the most relevant value (*i.e.*,
155 adequacy between calibration conditions and our study’s conditions) was chosen from literature,
156 among references using a similar POCIS configuration (200 mg Oasis® HLB) (Table S4). We considered
157 R_S values only if linear velocity or quiescence was specified. Then, we kept the R_S determined with
158 the closest flow velocity from our pilot study. When several values matched this criterion, we
159 considered the relevance of water matrix or doping level.

160 The PRC approach was used as an “advanced” procedure to correct R_S values for flow effects.
161 DIA-d5 was selected as PRC (Carpinteiro et al., 2016; Li et al., 2018a; Mazzella et al., 2010), and
162 corrections were made using the procedure detailed in Mazzella et al. (2010). Briefly, the elimination
163 rate constant of DIA-d5 (k_{ePRC}) was determined for each flow velocity, and a corrected sampling rate
164 (R_S^{cor}) was calculated using Eq. (2):

165
$$R_S^{cor} = R_S^{ref} \times \left(\frac{k_{ePRC}}{k_{ePRC}^{ref}} \right) \quad (2)$$

166 where R_S^{ref} and k_{ePRC}^{ref} are the sampling rate and the PRC elimination rate constant,
167 respectively, for the reference condition. The highest flow velocity ($V_3 = 17.5 \pm 2.1 \text{ cm s}^{-1}$) was taken
168 as the reference condition and sampling rates determined in this condition (R_S^{ref}) are presented in
169 Table S5. C_w was finally calculated using R_S^{cor} in Eq. 1.

170 2.5.2. Estimation based on o-DGT

171 The “Standard” estimation of C_w was based on o-DGT samplers equipped with 0.84 mm
172 diffusive gels (n=3) and performed using Eq.3 (Zhang and Davison, 1995):

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$$C_w = \frac{m\Delta g}{A_g t D} \quad (3)$$

174 where m is the accumulated mass of compound in the sampler, Δg is the diffusive gel thickness, D is
 175 the diffusion coefficient in the diffusive gel, A_g is the geometric exposure area (3.14 cm²), and t is the
 176 deployment time. D values were taken from Challis et al. (Challis et al., 2016) and corrected for
 177 temperature (T) using the Stokes-Einstein equation (Eq.4) where η is the water viscosity (taken from
 178 Lemmon et al., NIST chemistry WebBook):

179
$$\frac{D_1 T_1}{\eta_1} = \frac{D_2 T_2}{\eta_2} \quad (4)$$

180 The “advanced” estimation of C_w was made using a more sophisticated model (Eq.5, Santner
 181 et al., 2015) that considers the thickness of the DBL (δ) and lateral diffusion within the sampler:

182
$$C_w = \frac{m}{k_{ld} A_g t} \left(\frac{\Delta g}{D} + \frac{\delta}{D_w} \right) \quad (5)$$

183 where k_{ld} is the lateral diffusion flux increase coefficient (calculated according to Santner et al.,
 184 2015), and D_w is the diffusion coefficient in water (modeled using the Hayduk-Laudie equation
 185 described in Schwarzenbach et al., 1993). C_w was estimated alongside δ through direct adjustment of
 186 Eq.5 with Statistica software (version 6.1, Statsoft) on the full set of exposed o-DGT (*i.e.*, triplicate
 187 samplers equipped with 0.16; 0.41; 0.61 or 0.84 mm diffusive gels). δ values are displayed in Table
 188 S6.

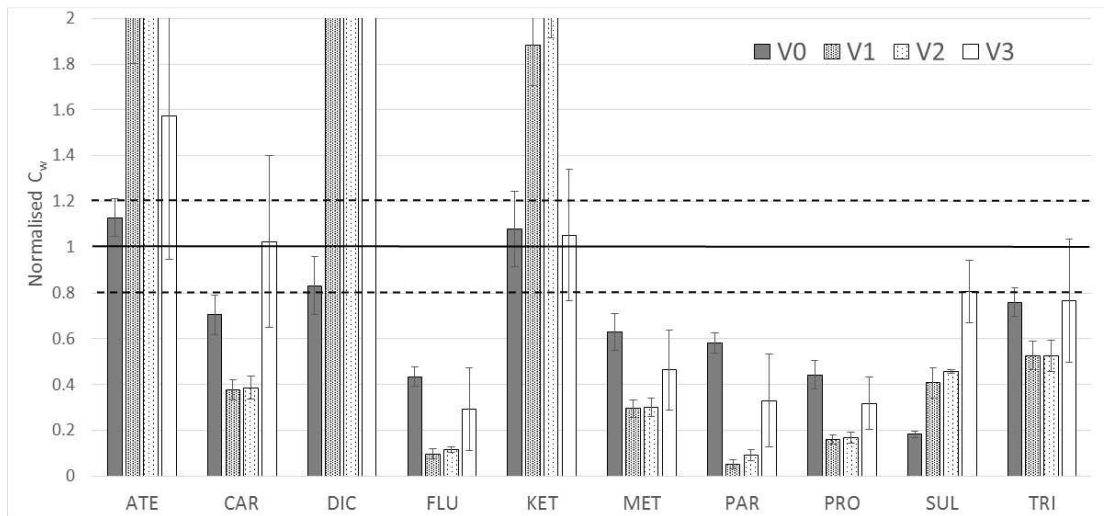
189 **2.5.3. Performance of passive samplers**

190 To discuss the accuracy on time-weighted average concentrations in pilots estimated using
 191 passive samplers, the average measured concentration determined on grab samples was taken as the
 192 reference concentration. The performance of passive samplers is consequently discuss through the
 193 comparison of C_w values determined with a given passive sampling procedure (C_w^{PS} ; *i.e.* POCIS or o-
 194 DGT and “standard” or “advanced”) and the corresponding concentration determined with grab
 195 sampling (C_w^{GS}). Given that C_w^{GS} were not identical for POCIS and o-DGT exposure and for V_0 and V_{1-3}
 196 flow velocities, the comparison between passive sampling procedures and flow velocities is made on
 197 the normalized value of C_w^{PS} to C_w^{GS} for clarity.

198 3. Results and discussion

199 3.1. POCIS C_w estimation without PRC

200 Normalized water concentrations determined using POCIS sampling with R_s from literature
201 (“standard” procedure) are presented in Figure 1. Except for diclofenac, concentration differences
202 with grab sampling were lower than 140%, which agrees with the 138% uncertainty reported by
203 Poulhier et al. (2014) during field deployment for pesticide monitoring. The important concentration
204 differences reported for diclofenac indicate that the R_s values chosen were not relevant for our
205 system. Considering that these values (Di Carro et al., 2014) were determined in conditions very close
206 to our study (flow velocities, doping level, water matrix and temperature), the R_s robustness for this
207 compound appears low. For the other compounds, concentration difference with grab sampling was
208 lower than 100% in non-quiescent conditions (V_{1-3}) with few exceptions (atenolol V_1 and V_2 ,
209 ketoprofen V_2). Moreover, most of concentration differences were between 50 and 75% (normalized
210 values between 0.25 and 0.5). These differences are higher than analytical uncertainty (typically 20%)
211 and probably arise from discrepancies between calibration conditions and our studied system,
212 although the R_s values were selected with the aim of limiting such discrepancies. Among them, flow
213 velocity could be of concern but other parameters should not be excluded. Positioning of the POCIS
214 (*i.e.*, parallel or perpendicular to flow) will affect hydrodynamics but is rarely indicated in the
215 literature. When considering quiescent conditions only (V_0), estimations were satisfactory
216 (concentration differences $\leq 57\%$, except sulfamethoxazole). Good performance for this condition
217 probably arose from the ease of choice for R_s , since calibrations in quiescent conditions were
218 available for each studied compound. This highlight the necessity, for the accurate use of POCIS, to
219 utilize calibrations performed in conditions very similar to the studied system.



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Figure 1: Mean C_w concentration (\pm SD) determined with POCIS and the “standard” procedure normalized to concentration determined with grab sampling. Flow velocities V_0 , V_1 , V_2 and V_3 were 0, 2.5, 6.3 and 17.5 cm s^{-1} , respectively. Dashed lines indicate typical analytical uncertainty (i.e. 20%). Values reach 2.2 and 2.4 for ATE at V_1 and V_2 , respectively; 2.5, 3.4 and 4.5 for DIC at V_1 , V_2 and V_3 , respectively; 2.1 for KET at V_2 .

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3.2. PRC approach for POCIS

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DIA-d5, proposed as a PRC for POCIS samplers (Mazzella et al., 2010), was investigated to correct C_w estimations for flow variations. For quiescent conditions (V_0), no PRC desorption from POCIS could be quantified ($k_e=0$). This approach is therefore not suitable for quiescent systems. For non-quiescent conditions, k_e values were 0.024, 0.031 and 0.046 d^{-1} for V_1 , V_2 and V_3 , respectively. These values are in the same range with values found in the literature: 0.034 d^{-1} (Carpinteiro et al., 2016), 0.044 d^{-1} (Belles et al., 2014), 0.046 d^{-1} (Li et al., 2018a) and 0.057 d^{-1} (Mazzella et al., 2010). However, the value of Mazzella et al. (2010) is twice the magnitude as the value we determined at the same flow velocity (V_1 ; 2-3 cm s^{-1}). This discrepancy possibly arose from the different exposure duration used to determine each value (i.e. 7 days in this study versus 28 days in Mazzella et al. study).

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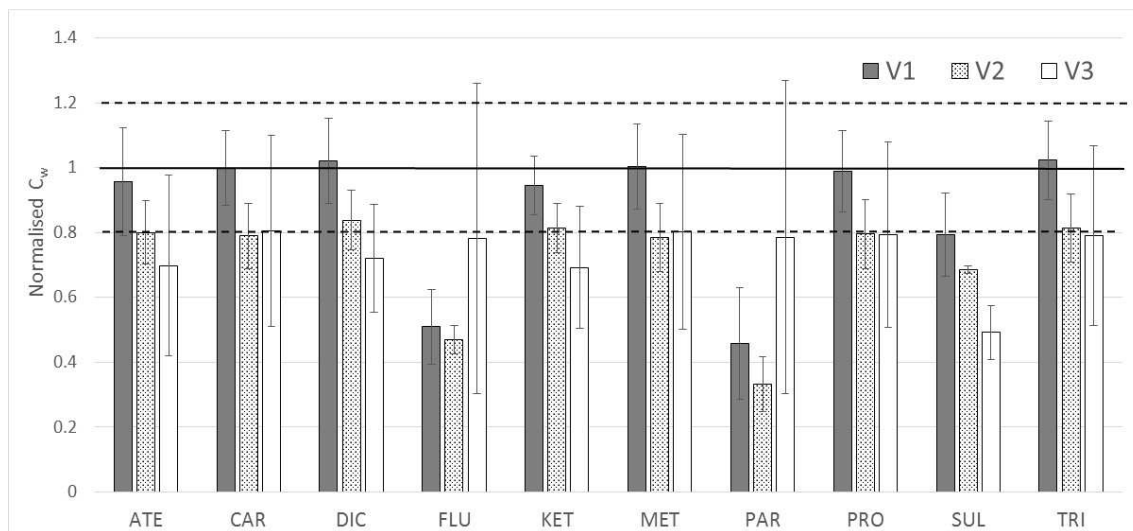
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Normalized water concentrations determined using POCIS sampling with the PRC approach are displayed in Figure 2. For the reference condition (V_3) concentration differences with grab sampling were $\leq 31\%$ except for sulfamethoxazole (51%). Values are consistent with target analytical accuracy (typically 20%) and highlight the adequacy of the reference sampling rate (R_S^{ref}) for the studied

240 system. This not surprising given that R_S^{ref} were determined using the same pilot and identical
241 conditions as this study.



242

243 *Figure 2: Mean C_w concentration (\pm SD) determined using POCIS with PRC normalized to concentration determined with grab*
244 *sampling. Dashed lines indicate typical analytical uncertainty (i.e. 20%). No value could be calculated for quiescent condition*
245 *(V_0) because there was no PRC desorption.*

246 Compared to C_w estimation without correction of R_S^{ref} , C_w estimations with PRC correction was
247 closer from estimation with grab sampling of approximately 25-30% at V_2 and of approximately 40-
248 50% for V_1 (Figure S2). The use of the PRC approach to correct sampling rate for flow discrepancy
249 allows therefore improving water concentration estimation. Compared to the reference condition
250 (V_3), the PRC approach allowed similar accuracy for V_2 and improved accuracy for V_1 except for
251 fluoxetine and paroxetine (Figure 2). The PRC approach appears to be a useful technique to handle
252 flow discrepancies between POCIS calibration and field application for several compounds, consistent
253 with the findings of the Mazzella et al. (2010) study. However, PRC correction was less efficient for
254 fluoxetine and paroxetine. This in agreement with Booij and Chen (2018) or Harman et al. (2011) who
255 stressed that the PRC approach has some limitations and might not work for all compounds.

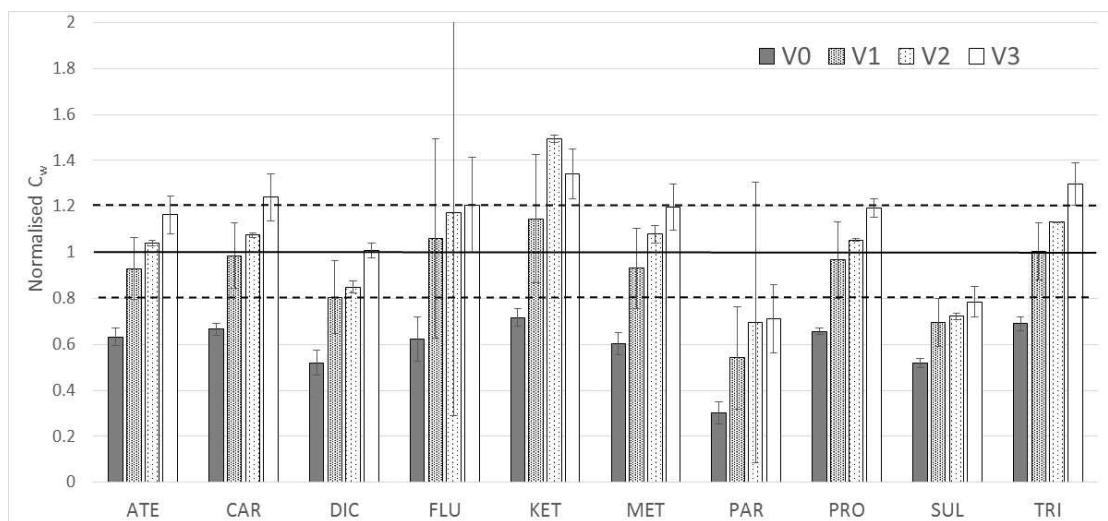
256 It has been suggested that the use of DIA-d5 as a PRC might be limited to deployment times
257 shorter than 15 days (Booij and Chen, 2018). Given that our results were obtained following 7 days
258 deployment times, the efficiency of the PRC approach was further investigated for 15 and 21 days
259 deployment times (Figure S3). Considering non-quiescent conditions only (V_{1-3}), no differences were

260 found for carbamazepine, diclofenac, ketoprofen, metoprolol, propranolol and trimethoprim between
261 C_w estimated with 7, 15 and 21 days of deployment. For atenolol, fluoxetine, paroxetine and
262 sulfamethoxazole, some differences were observed between C_w estimated with the different
263 exposure times. However, given that part of these differences were almost the same between the
264 reference condition (V_3 , no PRC correction possibility) and the other conditions (V_{1-2} , PRC correction
265 applied), these differences can be only partly a consequence a potential time-related limitation of
266 the PRC correction. PRC correction efficiency appears therefore only little affected by the
267 deployment time and the above conclusion for 7 days deployments might hold for more typical
268 deployments (*i.e.* 2-3 weeks). Considering quiescent conditions, significant desorption of DIA-d5 was
269 observed for 15 and 21 days deployments, enabling PRC correction. Although efficiency of the PRC
270 correction increased with deployment time, C_w estimation with POCIS was never found to match C_w
271 estimation with grab sampling. Discrepancies between POCIS and grab sampling estimations were
272 about 50-80% and 40-70% for 15 and 21 days deployments, respectively. Therefore, in quiescent
273 conclusion, efficiency of the PRC correction will depend on the deployment time but will never allow
274 full correction of flow effect for typical field deployments (*i.e.* 2-3 weeks).

275 3.3. C_w estimation with o-DGT

276 Figure 3 presents normalized water concentrations determined using o-DGT sampling with the
277 “standard” procedure, ~~except for clarithromycin, erythromycin, gemfibrozil and roxithromycin~~
278 ~~(discussed at the end of the section)~~. When the system was not quiescent (*i.e.*, $2 \leq V_{1-3} \leq 18 \text{ cm s}^{-1}$),
279 concentration difference with grab sampling was <50% for any compound. Given that most
280 differences were in the same range of the analytical accuracy (typically 20%), o-DGT accuracy
281 appears little affected by flow conditions between 2-18 cm s^{-1} . These results are in accord with
282 previous studies on metals (Davison and Zhang, 2012; Gimpel et al., 2001). When the system was
283 quiescent ($V_0 = 0 \text{ cm s}^{-1}$), the concentration difference with grab sampling was increased for all
284 compounds and reached between 28% (ketoprofen) and 70% (paroxetine). This altered accuracy has

285 previously been described for metals (Davison and Zhang, 2012) and is attributed to the formation of
 286 a significant diffusive boundary layer in front of the sampler that increases the diffusion path of
 287 compounds. Inaccuracies consequently arise from the simplicity of Eq. 3, which does not account for
 288 this diffusion path increase.

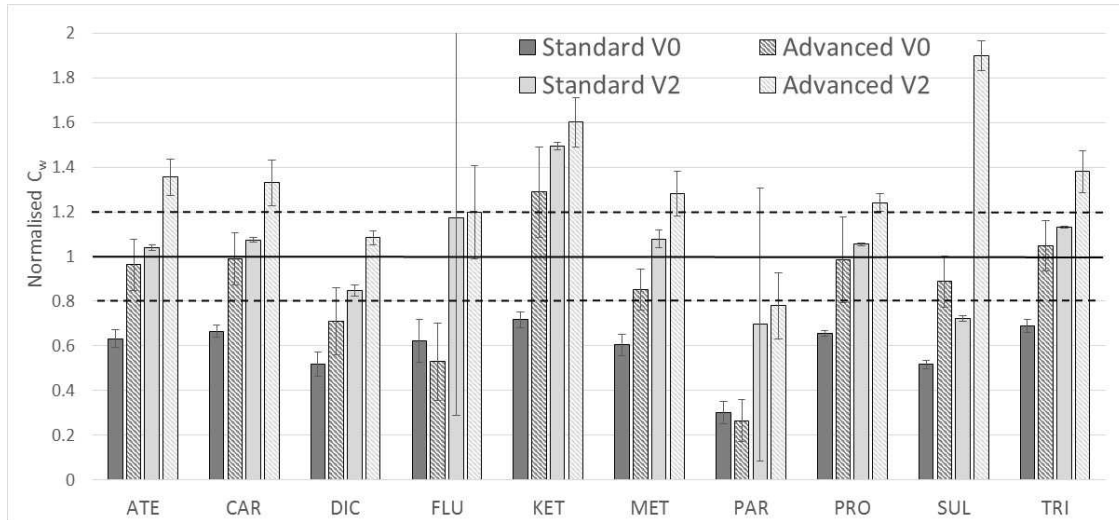


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290 Figure 3: Mean C_w concentration (\pm SD) determined with o-DGT and the “standard” procedure normalized to concentration
 291 determined with grab sampling. Dashed lines indicate typical analytical uncertainty (i.e. 20%).

292 Similar to what is done for metals (Garmo et al., 2006), such inaccuracy could be limited by use
 293 of the “advanced” procedure based on the deployment of samplers with various diffusive gel
 294 thicknesses and data treatment with a model that considers DBL formation (Eq. 5). Normalized water
 295 concentrations determined with this procedure are displayed in Figure S4. Surprisingly, when the
 296 system was not quiescent (V_{1-3}), in most cases the “advanced” procedure produced higher
 297 concentration differences between C_w estimations based on grab and o-DGT sampling compared to
 298 the “standard” procedure (the example of V_2 is displayed in Figure 4). This indicates a limitation of
 299 the model that might arise from the inaccuracy of its input parameters. Indeed, the lateral diffusion
 300 flux increase coefficient (k_{ld}) was initially established for phosphates (Santner et al., 2015) that have a
 301 greater ability to diffuse (i.e., higher diffusion coefficients) compared to the studied pharmaceuticals.
 302 D_w might also be questioned since values were estimated with an empirical model (Schwarzenbach et
 303 al., 1993). Conversely, for a quiescent system (V_0), the “advanced” procedure led to better C_w
 304 estimations except for fluoxetine, ketoprofen and paroxetine, which have similar levels of accuracy

305 when calculated with the “standard” procedure (Figure 4). The “advanced” procedure allows
 306 estimations within analytical uncertainty (<20%) except for diclofenac (29%), fluoxetine (47%),
 307 ketoprofen (29%) and paroxetine (73%). Therefore, the “advanced” procedure appears to be
 308 promising for producing accurate estimations in quiescent systems.



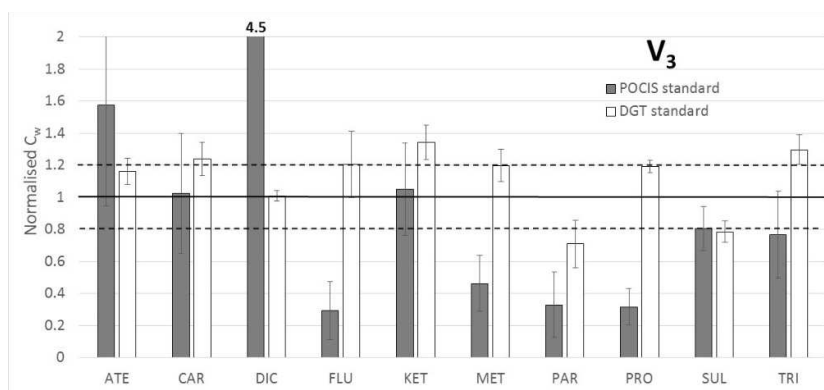
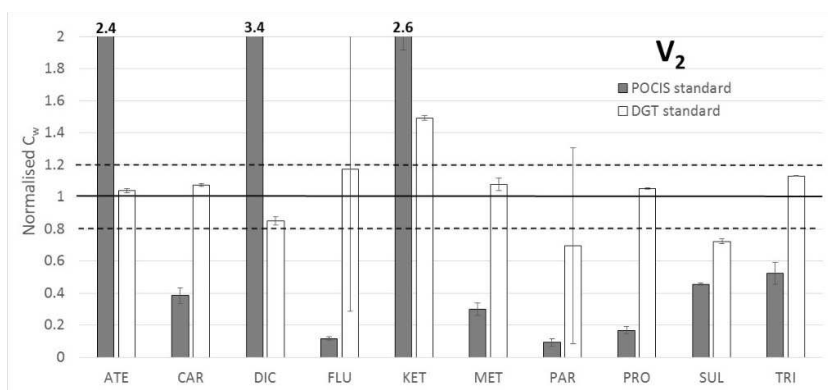
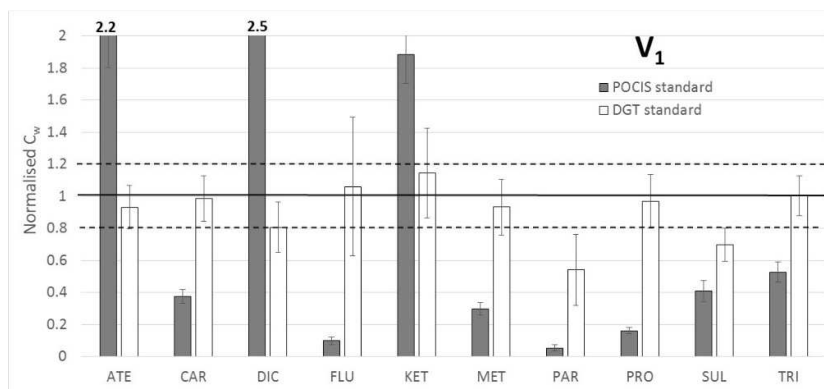
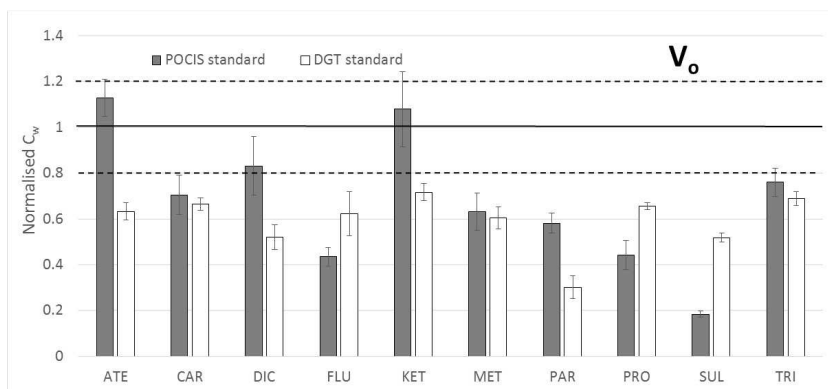
309

310 *Figure 4 : Mean C_w concentration (± SD) determined for V₀ and V₂ flow velocities using o-DGT with “standard” or “advanced”*
 311 *procedure normalized to concentration determined with grab sampling. Dashed lines indicate typical analytical uncertainty*
 312 *(i.e. 20%).*

313

314 **3.4. Comparison of POCIS and o-DGT**

315



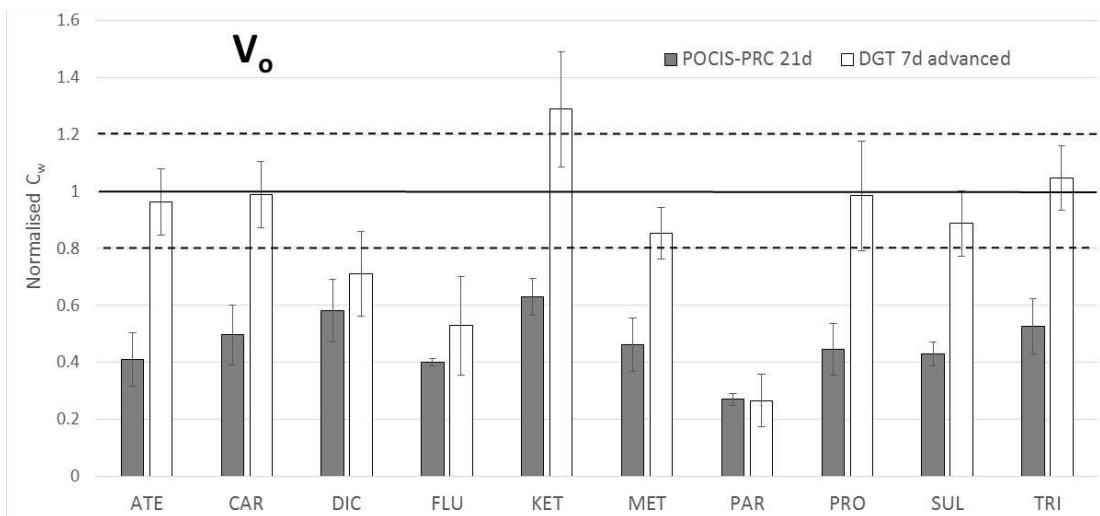
316 Figure 5: Mean C_w concentration (\pm SD) determined using POCIS or o-DGT (both with "standard" procedure) normalized to concentration determined with grab sampling. Dashed lines indicate
 317 typical analytical uncertainty (i.e. 20%).

318

319 For routine water surveys, simple procedures are required, and “standard” procedures are likely
320 to be used. In this context, o-DGT shows better accuracy (Figure 5) in non-quiescent conditions (V_{1-3})
321 with some exceptions for V_3 where accuracy is similar (carbamazepine, ketoprofen, sulfamethoxazole
322 and trimethoprim). Quiescent conditions (V_0) led to more varied results (Figure 5). POCIS was more
323 accurate for atenolol, diclofenac, ketoprofen and paroxetine whereas o-DGT was more accurate for
324 fluoxetine, propranolol, and sulfamethoxazole. Both tools performed similarly for carbamazepine,
325 metoprolol and trimethoprim. For routine purposes, this study currently supports the use of o-DGT
326 to limit the effect of flow in non-quiescent conditions, whereas in quiescent conditions, the choice
327 depends on the target compounds.

328 In cases where the use of more sophisticated procedure is conceivable, estimations can be
329 improved with the “advanced” procedure using PRCs. For non-quiescent conditions, o-DGT
330 performance was already close to the analytical accuracy (typically 20%) and no improvement over
331 the “standard” procedure was required. For POCIS, the use of DIA-d5 as a PRC was required to obtain
332 similar accuracies for most of the studied compounds. However, this approach was less efficient for
333 fluoxetine and paroxetine. In quiescent conditions, o-DGT estimations were improved with the
334 “advanced” procedure for most compounds (Figure 4). Given that in quiescent conditions PRC was
335 unable to desorb for 7 days deployment time, o-DGT estimations with the “advanced” procedure
336 were compared to POCIS with the “advanced” procedure (PRC approach) using 21 days deployment
337 time (Figure 6). For atenolol, carbamazepine, metoprolol, propranolol, sulfamethoxazole and
338 trimethoprim, C_w estimations based on o-DGT were closer from estimation based on grab sampling,
339 compared to estimation based on POCIS. O-DGT and grab sampling estimation were even identical
340 when considering analytical uncertainty. For diclofenac, fluoxetine, ketoprofen and paroxetine, o-
341 DGT and POCIS showed similar performances with the advanced procedure in C_w estimation.

342



343

344 *Figure 6: Mean C_w concentration (\pm SD) determined for V_o using POCIS (21 days deployment with PRC) and o-DGT (7 days*
 345 *deployment with “advanced” procedure) normalized to concentration determined with grab sampling. Dashed lines indicate*
 346 *typical analytical uncertainty (i.e. 20%).*

347 Overall, if implementation of the PRC approach is conceivable for POCIS, flow effects will be
 348 limited for both POCIS and o-DGT and each will perform with accuracies within acceptable analytical
 349 error (<20%) for most of the studied compounds. However, only o-DGT currently achieves such
 350 accuracy in quiescent conditions. Nevertheless, an important advantage of POCIS compared to o-DGT
 351 is its higher sampling rates, especially at high flow velocities (Table S7), resulting in a greater POCIS
 352 sensitivity. Although a recent comparison of POCIS and o-DGT showed similar detection frequencies
 353 in four freshwater systems (Challis et al., 2018), lower sensitivity of o-DGT (up to 67 times for
 354 fluoxetine) may be an issue for some compounds depending on the targeted system contamination.
 355 POCIS use may be therefore preferred in such context. The choice of strategy to limit flow effects will
 356 be driven by the study constrains in terms of sensitivity, accuracy and flow conditions. If sensitivity is
 357 favored (*e.g.*, low-contaminant systems), then POCIS with PRC should be chosen. Conversely, if
 358 sensitivity is not a limiting constraint (*e.g.*, contaminated systems), both POCIS with PRC or o-DGT
 359 can be chosen. However, if quiescent (or nearly quiescent) conditions are expected, then o-DGT
 360 should be chosen to achieve better accuracy. Anyway, it must be stressed, that flow velocity is not
 361 the only condition that affects passive sampling in aquatic systems (*e.g.*, pH, temperature,
 362 biofouling), and the above recommendations may be reconsidered in the future.

363 3.5. Future improvements

364 For a given sampler, different behaviors were observed among the studied compounds and
365 some failures were identified whatever the sampler used. For POCIS, correction with DIA-d5 as PRC
366 was not fully satisfactory for fluoxetine and paroxetine. Therefore, more suitable PRCs still have to be
367 identified for these two compounds. Indeed, Li et al. (2018b) identified very recently Antipyrine-d3 as
368 a promising PRC for pharmaceuticals compounds. Furthermore, when many compounds are
369 targeted, simultaneous implementation of several PRCs will probably be required for extensive flow
370 effect correction.

371 For any study, it may be of interest to limit the number of tools required and the complexity of
372 procedures used. However, given that each sampler has limitations, improvements are required to
373 generalize the use of one sampler. A current limitation of POCIS is accuracy in quiescent conditions
374 because of limitation of the PRC approach. Improving POCIS accuracy in quiescent systems will
375 require the development of a PRC able to desorb in this condition or the development of a new
376 strategy. Therefore, further improvements on POCIS are currently speculative. Conversely, several
377 improvements can be proposed for o-DGT. First, decreasing biases in quiescent systems to allow
378 accurate use of the “standard” procedure could be achieved with an increase in diffusive gel
379 thickness. Using thicker gel will result in limiting the impact of DBL formation on the diffusion path
380 and therefore limit the inaccuracy of Eq. 3. It was estimated (Table S8) that using 2.5-mm-thick
381 diffusive gels and the “standard” procedure would allow achieving accuracies better than 25% in
382 quiescent systems. However, such gel thickness increase will also alter o-DGT method sensitivity by a
383 2.8 factor because of the length increase of the diffusion path. Second, the lack of sensitivity of o-
384 DGT compared to POCIS could be overcome. Sensitivity differences are a consequence of sampling
385 rate differences (Table S7). This difference was previously demonstrated for several antibiotics (Chen
386 et al., 2013) and anionic pesticides (Guibal et al., 2017) to be largely due to sampling area difference.
387 Therefore, increasing the sampling area of o-DGT should result in increased sampling rates (Eq. S1
388 and S2) and higher sensitivity. It can be calculated (Table S9) that a 160 cm² sampling area (~7 cm

389 radius) should allow sampling rates not lower than conventional POCIS ones (except for fluoxetine
390 and ketoprofen), even if gel thickness is increased for quiescent systems. This design would result in
391 similar sensitivity compared with POCIS. Such a theoretical o-DGT configuration must be tested in the
392 field, since physical constraints on these larger gels could be significant.

393 4. Conclusions

394 Both POCIS with PRC approach and o-DGT were shown to limit flow effect on passive sampling
395 accuracy. However, none of these approaches is currently universal and each one has some
396 drawbacks. Therefore, before choosing an approach, preliminary validation on the targeted
397 compounds and deployment conditions is advisable. Limitation of flow effect using both approaches
398 was not fully satisfactory for all compounds. To overcome these limitations, it will require
399 improvement of calibrations for both samplers and development of new PRCs for POCIS. Therefore,
400 future improvements are probably more speculative for POCIS than for o-DGT samplers. The o-DGT
401 approach has probably a better potential to allow in the future limitation of flow effect on a wide
402 range of compounds. Conversely, POCIS samplers already propose better sensitivity compared to o-
403 DGT samplers. Therefore, unless new designs allowing better sensitivity of o-DGT samplers are
404 proposed and validated, POCIS samplers could still be preferred for micropollutants monitoring in
405 natural systems.

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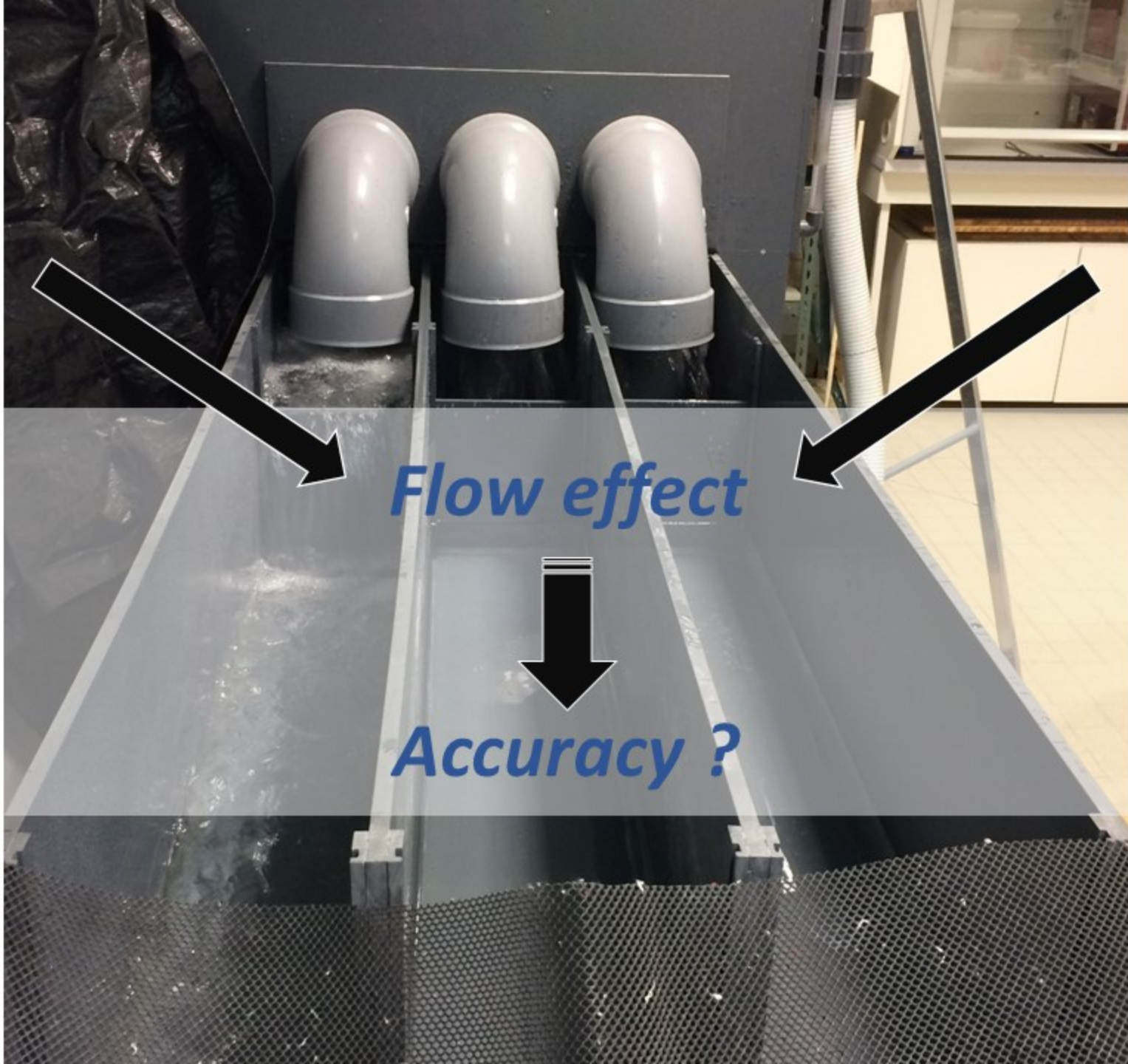
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505



POCIS + PRC



Flow effect

Accuracy ?



o-DGT