

Accepted Manuscript

Title: Development of a simple analytical method for the simultaneous determination of paracetamol, paracetamol-glucuronide and *p*-aminophenol in river water

Author: Lúcia H.M.L.M. Santos Paula Paíga Alberto N. Araújo Angelina Pena Cristina Delerue-Matos M. Conceição B.S.M. Montenegro



PII: S1570-0232(13)00243-2
DOI: <http://dx.doi.org/doi:10.1016/j.jchromb.2013.04.032>
Reference: CHROMB 18384

To appear in: *Journal of Chromatography B*

Received date: 2-11-2012
Revised date: 16-4-2013
Accepted date: 23-4-2013

Please cite this article as: L.H.M.L.M. Santos, P. Paíga, A.N. Araújo, A. Pena, C. Delerue-Matos, M.C.B.S.M. Montenegro, Development of a simple analytical method for the simultaneous determination of paracetamol, paracetamol-glucuronide and *p*-aminophenol in river water, *Journal of Chromatography B* (2013), <http://dx.doi.org/10.1016/j.jchromb.2013.04.032>

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2 **Development of a simple analytical method for the simultaneous determination**
3 **of paracetamol, paracetamol-glucuronide and *p*-aminophenol in river water**

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11 Lúcia H.M.L.M. Santos^a, Paula Paíga^b, Alberto N. Araújo^a, Angelina Pena^c, Cristina
12 Delerue-Matos^b, M. Conceição B.S.M. Montenegro^{a*}

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20
21
22 ^aREQUIMTE, Faculty of Pharmacy, University of Porto, Rua Jorge Viterbo
23 Ferreira 228, 4050-313 Porto, Portugal

24
25
26
27 ^bREQUIMTE, Instituto Superior de Engenharia do Porto, Instituto Politécnico do
28 Porto, Rua Dr. António Bernardino de Almeida, 431, 4200-072 Porto, Portugal

29
30
31
32 ^cGroup of Health Surveillance, Center of Pharmaceutical Studies, University of
33 Coimbra, Health Sciences Campus, Azinhaga de Santa Comba, 3000-548 Coimbra,
34 Portugal

35
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37
38 *Corresponding author: Phone: +351 220428677; fax: +351 226093390; e-mail:
39 mcbranco@ff.up.pt

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23 **Abstract**

24 Paracetamol is among the most worldwide consumed pharmaceuticals. Although its
25 occurrence in the environment is well documented, data about the presence of its
26 metabolites and transformation products is very scarce. The present work describes the
27 development of an analytical method for the simultaneous determination of
28 paracetamol, its principal metabolite (paracetamol-glucuronide) and its main
29 transformation product (*p*-aminophenol) based on solid phase extraction (SPE) and high
30 performance liquid chromatography coupled to diode array detection (HPLC-DAD).
31 The method was applied to analysis of river waters, showing to be suitable to be used in
32 routine analysis. Different SPE sorbents were compared and the use of two Oasis WAX
33 cartridges in tandem proved to be the most adequate approach for sample clean up and
34 pre-concentration. Under optimized conditions, limits of detection in the range of 40 to
35 67 ng/L were obtained, as well as mean recoveries between 60 and 110% with relative
36 standard deviations (RSD) below 6%. Finally, the developed SPE-HPLC/DAD method
37 was successfully applied to the analysis of the selected compounds in samples from
38 seven rivers located in the north of Portugal. Nevertheless all the compounds were
39 detected, it was the first time that paracetamol-glucuronide was found in river water at
40 concentrations up to 3.57 µg/L.

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44 **Keywords:** Paracetamol, paracetamol-glucuronide, *p*-aminophenol, solid phase
45 extraction, HPLC, river water

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1. Introduction

Paracetamol (acetaminophen or *N*-acetyl-4-aminophenol) is one of the most popular and widely used medicines for the treatment of pain and fever, both as an over-the-counter (OTC) and as a prescribed medicine. It can be used in a wide range of patients, including children, pregnant women or the elderly. Following oral administration, approximately 90% of paracetamol is metabolized, being conjugated with glucuronide (40–67%) and, in a less extent, with sulphate (20–46%), to form inactive metabolites, which are eliminated in urine together with a small fraction of unchanged paracetamol (<5%) [1]. Although paracetamol presents a high removal efficiency (approximately 99%) in WWTPs [2, 3], it has been detected in their effluents at concentrations up to low microgram per litre [4-6], contributing to its entrance into surface waters [7-9]. Once in the environment, paracetamol is mainly degraded by microorganisms, which are capable of using it as carbon and energy sources [10].

Although paracetamol is not highly persistent in the environment, continuous input overrules its high transformation rate [11], thus, it can adversely affect aquatic organisms. Acute toxicity effects in the invertebrate *Daphnia magna* (EC₅₀ ranging from 26.6 to 50 mg/L) [12-14], the marine bacterium *Vibrio fischeri* (EC₅₀ = 549.7 mg/L) and the fish *Oryzias latipes* (EC₅₀ = >160 mg/L) [12] have been reported. Effects on cell cultures with EC₅₀ values of 19 mg/L have also been described [13].

Nowadays analytical methodologies described in literature are mainly focused in multi-residues methods that allow the simultaneous determination of paracetamol together with a large number of pharmaceuticals from several therapeutic groups [15-17]. Most of them are principally focused in parent compounds and rarely analyze metabolites and/or transformation products. At present, methods reported for the determination of paracetamol metabolites are focused in biological matrices [18-20]

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72 rather than in environmental ones [21]. On the other hand, paracetamol may also be
73 degraded, both during wastewater treatment and in the environment, giving different
74 transformation products [22, 23]. *p*-Aminophenol was identified as its main
75 transformation product, and its presence in wastewater samples was reported [23].
76 However the origin of *p*-aminophenol cannot only be attributed to the degradation of
77 paracetamol, since it is also widely used in industrial applications and is known as a
78 transformation product from pesticides. Furthermore, *p*-aminophenol was also described
79 as the primary degradation product of paracetamol during the storage of its medicinal
80 formulations [24].

81 High-performance liquid chromatography (HPLC) coupled to tandem mass
82 spectrometry (MS/MS) has been designated as technique of choice for the determination
83 and quantification of pharmaceuticals in environmental samples [25]. However, these
84 equipments are still very expensive and they are not available in many laboratories for
85 routine analysis. On the other hand, almost all laboratories have HPLC systems with
86 diode array UV absorbance and/or fluorescence detection that may effectively be used
87 for the analysis of pharmaceuticals in environmental samples [26-29].

88 Due to the complexity of environmental samples, analysis of pharmaceuticals has to
89 be preceded by a pre-concentration step, which allows the detection of low
90 concentrations and simultaneously removes the interferences. This is often performed
91 by solid phase extraction (SPE). Generally, polymeric sorbents, like Oasis HLB, are the
92 most used for pre-concentration of pharmaceuticals from aqueous matrices [16, 17, 30-
93 32], though mixed-mode ion-exchange sorbents have also been described [33, 34].

94 Thus, the present work describes the development and validation of an analytical
95 method based on off-line SPE, using a mixed mode reversed phase/ anionic exchange
96 sorbent, followed by LC-DAD for the determination of paracetamol, its main metabolite

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97 (paracetamol-glucuronide) and its principal transformation product (*p*-aminophenol) in
98 river waters. The performance and application of this method is important, since allows
99 the simultaneous monitoring of parent compound, metabolite and transformation
100 product as well as the evaluation of their environmental interdependence, using one of
101 the most worldwide consumed pharmaceuticals (paracetamol) as example.

102 Finally, the developed methodology was successfully applied to the analysis of the
103 selected compounds in seven rivers from north of Portugal. To our knowledge this is the
104 first time that paracetamol-glucuronide was found in surface waters.

107 2. Materials and methods

108 2.1. Chemicals and reagents

109 Paracetamol (PCT) (acetaminophen), *p*-aminophenol (PAP) (4-aminophenol) and
110 paracetamol-glucuronide (PCT-G) (ρ -acetamidophenyl β -D-glucuronide) sodium salt
111 were purchased from Sigma-Aldrich (Steinheim, Germany). All standards were of high
112 purity grade (>93%). HPLC-grade methanol, HPLC-grade acetonitrile and HPLC-grade
113 acetone, *n*-hexane and formic acid (purity \geq 98%) were obtained from Merck
114 (Darmstadt, Germany), hydrochloric acid 37% and glacial acetic acid (purity \geq 99.7%)
115 were purchased from Carlo Erba (Rodano, Italy), ammonia 25% was obtained from
116 Panreac (Barcelona, Spain), ammonium hydroxide solution, ammonium acetate (purity
117 \geq 98%), ethyl acetate and dichloromethane were purchased from Sigma-Aldrich
118 (Steinheim, Germany). HPLC-grade water (18.2 M Ω cm) was obtained by purifying
119 deionised water in a Milli-Q Simplicity 185 system (Millipore, Molsheim, France).

120 Individual stock standard solutions were prepared for each compound by
121 dissolving 10 mg of powder in 10 mL of methanol, obtaining a final concentration of

122 1000 mg/L, and stored at -20 °C. Stock standard solutions were renewed every week.

123 An intermediate standard solution was daily prepared by mixing the three individual
124 stock solutions and diluting with a mixture methanol-water (10:90, v/v) to give a final
125 concentration of 10 mg/L and kept at 4 °C. Working standard solutions were also
126 prepared in a mixture methanol-water (10:90, v/v) by dilution of appropriate amounts of
127 the intermediate solution. Amber glassware was used to prevent light degradation.
128 These working standard solutions were used for preparation of the calibration curve and
129 for spiking samples in the validation study.

130 All standard solutions and sample extracts were filtered through a 0.20 µm
131 PTFE syringe filter (Teknokroma, Barcelona, Spain) and homogenised using a vortex
132 mixer (VWR, Radnor, Delaware, USA). All chromatographic solvents were filtered
133 through a 0.20 µm nylon membrane filter (Supelco, Bellefonte, PA, USA) using a
134 vacuum pump (Dinko D-95, Barcelona, Spain) and degassed for 15 min in an ultrasonic
135 bath (Raypa[®] Trade, Terrassa, Spain).

136 SPE cartridges used were Oasis[®] MAX (60 mg, 3 mL), Oasis[®] WAX (150 mg, 6
137 mL), Oasis[®] MCX (150 mg, 6 mL) and Oasis[®] HLB (200 mg, 6 mL) from Waters
138 (Mildford, MA, USA), LiChrolut[®] EN/RP-18 (EN 40-120 µm, 100 mg (bottom) and
139 RP-18 40-63 µm, 200 mg (top), 6 ml) from Merck (Poland), Strata[™]-SDB-L (500 mg,
140 6 mL) and Strata[™]-X (200 mg, 3 mL) from Phenomenex (USA), and Enviro-clean[®] (C₈
141 and quaternary amine, 1000 mg, 6 mL) from Unit Chemical Technologies (UCT), Inc.
142 (Bristol, PA, USA).

143

144 2.2. Sample collection

145 River water (2.5 L) was collected from seven rivers located in the north of Portugal,
146 which is one of the most densely populated areas of the country. Sample collection,

147 preservation and storage were done according to the US EPA Method Guideline [35].
148 River samples were collected on the river side in amber glass bottles and kept
149 refrigerated ($\pm 4^{\circ}\text{C}$) during the transport to the laboratory. Samples were collected along
150 one week in September 2011.

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152 **2.3. Sample pre-treatment and extraction**

153 River water samples were vacuum filtered through 1.2 μm glass microfiber filters
154 (GF/C, Whatman, UK), followed by 0.20 μm nylon membrane filters (Supelco,
155 Bellefonte, PA, USA) and stored at -20°C , until extraction.

156 For the SPE procedure a vacuum manifold system (Phenomenex, USA) was
157 used. Two Oasis WAX cartridges were initially conditioned, in separate, with 2 mL of
158 methanol, 2 mL of HPLC-grade water, and 2 mL of HPLC-grade water pH 7 (pH
159 adjusted with ammonia) at a flow rate of 1 mL/min. After that, the SPE cartridges were
160 connected in tandem and 50 mL of river water (pH adjusted to 7 with ammonia) were
161 loaded onto the cartridges at a flow rate of 1 mL/min. Finally, analytes were eluted with
162 5 mL of methanol and 5 mL of 5% ammonium hydroxide in methanol at a flow rate of 1
163 mL/min, and the eluates were pooled in one single collection vial. Extracts were
164 evaporated to dryness under a gentle stream of nitrogen and reconstituted in 250 μL of a
165 mixture methanol-water (10:90, v/v), allowing a pre-concentration factor of 200.

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167 **2.4. Liquid chromatography**

168 Chromatographic analysis was performed on a Nexera Ultra-High Performance
169 Liquid Chromatography system (Shimadzu Corporation, Kyoto, Japan) equipped with
170 two solvent delivery modules LC-30 AD, a column oven CTO-20 AC, an autosampler
171 SIL-30 AC and an UV/Vis photodiode array detector SPD-M20A. The system was

172 controlled by a system controller CBM-20A. Two chromatographic systems were used
173 in this work in order to evaluate the performance of the method for HPLC and fast-
174 HPLC analysis. On the former, separation was carried out using a Luna C18(2) column
175 (150 x 4.6 mm i.d., 5 μ m particle size) (Phenomenex, USA) that was kept at 25 $^{\circ}$ C,
176 while sample vials were kept at 4 $^{\circ}$ C. An injection volume of 40 μ L was used. The
177 optimized mobile phase consisted of 10 mM ammonium acetate/acetic acid (pH 6) as
178 solvent A and acetonitrile as solvent B, using a flow rate of 1 mL/min. The gradient
179 elution was performed as follow: 0–9.0 min, 97–74.8% A; 9.0–10.0 min, return to initial
180 conditions; 10.0–15.0 min equilibration of the column. For fast-HPLC analysis, a Shim-
181 pack XR-ODS (100 x 4.6 mm i.d., 2.2 μ m particle size) (Shimadzu, Kyoto, Japan)
182 column was used and it was kept at 25 $^{\circ}$ C. The separation was performed using the
183 same mobile phase composition (10 mM ammonium acetate/acetic acid (pH 6) as
184 solvent A and acetonitrile as solvent B) at a flow rate of 2 mL/min. The gradient elution
185 was: 0–1.0 min, 96–91.6% A; 1.0–1.5 min, 91.6–80% A; 1.5–3.0 min, 80% A; 3.0–3.5
186 min, returned to initial conditions; 3.5–4.0 min, equilibration of the column. An
187 injection volume of 10 μ L was used and sample vials were kept at 4 $^{\circ}$ C. The analytes
188 were monitored at their maximum UV wavelengths, namely 243, 232 and 245 nm for
189 paracetamol-glucuronide, *p*-aminophenol and paracetamol, respectively. Lab Solutions
190 software (Shimadzu Corporation, Kyoto, Japan) was used for control and data
191 processing.

193 2.5. Method validation

194 Method validation was performed for both HPLC and fast-HPLC analysis, in
195 order to evaluate which chromatographic technique might be more suitable for
196 application to real samples.

197 Target analytes were identified in the chromatograms by comparison of the
198 retention time of the peaks obtained with these ones of a standard solution.

199 Simultaneously the identification of the analytes was also confirmed comparing the
200 corresponding UV spectra of the peaks of the sample and of standard solution
201 chromatograms. For the quantification of the analytes in real samples the standard
202 addition method was used.

203 The linearity of the method was established by setting calibration curves using linear
204 regression analysis over the concentration range of 50 to 1500 $\mu\text{g/L}$; however final
205 concentrations tested depend of the sensitivity reached for each analyte. Method
206 detection limit (MDL) and method quantification limit (MQL) were determined as the
207 minimum amount detectable of analyte with a signal-to-noise ratio of 3 and 10,
208 respectively.

209 Method accuracy (expressed as recovery percentage) and precision (expressed in
210 terms of relative standard deviation (RSD)) were evaluated by recovery studies of the
211 analytes in river water, spiked at different final concentration levels (0.75, 2.50, 3.75
212 and 5.00 $\mu\text{g/L}$). Experiments were performed in triplicate ($n = 3$). Recoveries were
213 determined comparing the concentrations obtained with the initial spiking levels. As
214 river water may contain the target analytes, blanks (samples without standard solution
215 addition) were analyzed in order to determine their concentrations, which were then
216 subtracted to the spiked river water.

217 Method precision was determined by repeated intra- and inter-day analysis, through
218 five successive injections of a river water sample spiked with a standard mixture
219 containing all the analytes at a final concentration of 5.00 $\mu\text{g/L}$ in one day and its
220 injection in five consecutive days, respectively.

221 The influence of the matrix in the UV signal was evaluated by preparing a
222 calibration curve in river extract, and comparing it with one achieved for the standards
223 prepared in a mixture of methanol-water (10:90, v/v). A blank (sample with no addition
224 of the standards) was simultaneously assayed in order to subtract the levels of the target
225 analytes present in the sample.

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228 3. Results and discussion

229 3.1. Solid phase extraction optimization

230 The SPE method was optimized using ultra-pure and river water.

231 The selection of the most adequate SPE sorbent is a critical point in the
232 development of an extraction procedure; therefore the performance of different SPE
233 cartridges was evaluated and the recoveries in ultra-pure water, pH adjusted to 7, are
234 shown in Figure 1. As can be seen, paracetamol could be efficiently recovered by
235 almost all SPE cartridges (with the exception of Enviro-clean[®]), while for its metabolite
236 higher recoveries (72–103%) were obtained with the mixed mode reversed
237 phase/anionic exchange sorbents Oasis MAX and WAX, since at pH 7, paracetamol-
238 glucuronide is negatively charged (Figure S1, Supplementary data), establishing ionic
239 interactions with those sorbents. Therefore, polymeric sorbents such as Oasis HLB,
240 Strata-X and Strata SDB-L, which extract analytes by reversed phase mechanisms, were
241 not able to recover this compound. Furthermore, *p*-aminophenol showed the lowest
242 recovery, which did not exceed 43%, using the Oasis WAX cartridges, and it was not
243 recovered at all with Enviro-clean[®], given that at pH 7, *p*-aminophenol is in a neutral
244 form (Figure S1, Supplementary data), and this mixed mode sorbent (cationic exchange
245 and C₈) is not able to interact with *p*-aminophenol by reversed phase mechanisms. As it

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246 is depicted in Figure 1, the hydrophilic polymeric sorbents Oasis HLB and Strata X
247 yielded similar recoveries for all analytes, which is in agreement with their similar
248 physico-chemical properties [36]. Comparing the different SPE sorbents tested, it can be
249 seen that highest average recoveries were achieved for all the analytes with Oasis WAX
250 (Figure 1).

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Insert Figure 1

254 Mixed-mode ion-exchange sorbents allow two types of interaction mechanisms,
255 namely reversed phase and ionic-exchange. Thus, taking special attention to pH and
256 solvents employed in each step of SPE protocol, analytes can be selectively eluted [37].
257 In this context, the effect of sample's pH was studied for the mixed-mode ion-exchange
258 sorbents (Oasis MAX, WAX and MCX) in order to enhance the *p*-aminophenol
259 recovery. Figure 2 shows the recoveries obtained in ultra-pure water. For paracetamol-
260 glucuronide, better recoveries were achieved using reversed phase/anionic exchange
261 sorbents (Oasis MAX and WAX) at a pH range between 3 and 7, which is in agreement
262 with its pKa (Table 1). At this pH range paracetamol-glucuronide is negatively charged
263 (Figure S1, Supplementary data), being able to bind to the sorbent by ionic interactions,
264 since Oasis MAX and WAX are a strong and a weak anionic-exchange polymeric
265 sorbent, respectively, that are based on Oasis HLB (poly(*N*-vinylpyrrolidone-
266 divinylbenzene) copolymer) chemically modified with quaternary amine groups
267 (dimethylbutylamine) and piperazine groups, respectively [37]. On the other hand,
268 Oasis MCX is a strong cation-exchange polymeric sorbent that has an Oasis HLB
269 skeleton chemically modified with sulfonic groups [37], therefore is not able to interact

270 with a negatively charged compound, and for that reason the recoveries for Oasis MCX
271 are very low (<20%).

272

273 Insert Table 1 and Figure 2

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275 Like paracetamol-glucuronide, the highest recoveries of *p*-aminophenol were
276 also achieved with a reversed-phase/anionic exchange sorbent, namely Oasis WAX, at a
277 pH range between 6 and 8 (Figure 2b). According to *p*-aminophenol pKa, at these pH
278 values the molecule is in a neutral form (Figure S1, Supplementary data), being
279 extracted through reversed phase mechanisms. However, recoveries of *p*-aminophenol
280 using Oasis MCX were similar to that one obtained with Oasis WAX for a pH up to 6.
281 At this pH range *p*-aminophenol presents the amine group positively charged (Figure
282 S1, Supplementary data) and, therefore, is able to establish ionic interactions with the
283 sulfonic groups of the sorbent.

284 Relatively to paracetamol, all the mixed-mode ion-exchange sorbents tested
285 provided good recoveries (higher than 70%) in all the studied pH range (Figure 2c),
286 since it is in a neutral form (Figure S1, Supplementary data), being extracted through
287 reversed phase mechanisms. Based on the results obtained, a sample's pH of 7 was
288 chosen for further studies.

289 In contrast to the excellent recoveries of paracetamol and its metabolite obtained
290 with Oasis WAX, the recovery of *p*-aminophenol may be improved using two SPE
291 cartridges in tandem. Different possible combinations were tested using Oasis WAX,
292 MCX and HLB. Oasis HLB was used in this study due to have been proved that, at pH
293 7, *p*-aminophenol is extracted by reversed phase mechanisms, so an increase in the mass
294 sorbent of the lipophilic/hydrophilic balance Oasis HLB sorbent might improve the

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295 recovery of *p*-aminophenol. Results showed that two Oasis WAX columns in tandem
296 allowed increasing the recovery of *p*-aminophenol (64%), maintaining good recoveries
297 of the others analytes (Table S1, Supplementary data).

298 Moreover, different organic solvents were tested namely methanol, acetonitrile,
299 acetone, ethyl acetate, *n*-hexane and dichloromethane, as well as different proportions of
300 ammonium hydroxide in methanol. No improvement on *p*-aminophenol recovery was
301 observed comparatively to the elution protocol suggested by the manufacturer that is
302 methanol, for the extraction of neutral compounds extracted by reversed phase
303 mechanisms, and 5% ammonium hydroxide in methanol, to revert the ionic interactions
304 and release the compounds extracted through anionic exchange mechanisms.

305 After the SPE protocol have been settled, different sample volumes were studied
306 using two Oasis WAX cartridges in tandem. For paracetamol and paracetamol-
307 glucuronide, recoveries around 100% were achieved for sample volumes up to 100 mL,
308 although the recovery of *p*-aminophenol decreased with increasing sample volume
309 (Figure S2, Supplementary data). Thus, attending that the main goal of this work was to
310 develop an extraction procedure for the determination of paracetamol, its main
311 metabolite (paracetamol-glucuronide) and its main transformation product (*p*-
312 aminophenol) in river waters, a sample volume of 50 mL was chosen as a commitment
313 between the recovery of the analytes and the intended application of the developed
314 method.

315

316 **3.2. Optimization of chromatographic conditions**

317 In order to optimize the chromatographic separation, different mobile phases were
318 tested, using methanol, acetonitrile or a mixture of methanol and acetonitrile as organic
319 solvent and water with different additives, such as ammonium acetate and acetic acid at

320 different concentrations as aqueous phase. The best separation was achieved using 10
321 mM ammonium acetate/acetic acid (pH 6) and acetonitrile. This mobile phase was used
322 for both developed chromatographic procedures.

323

324 **3.2.1. HPLC analysis**

325 After the mobile phase composition had been established, the elution gradient
326 and flow rate were adjusted in order to improve the chromatographic resolution as well
327 as the peaks shape and to get the shorter analysis time. The optimum flow rate was set
328 at 1.0 mL/min and the elution was performed on a 15 minutes gradient. Different
329 column temperatures were also studied (23 °C, 25 °C and 30 °C) and for all the analytes
330 the best peak shape and resolution were obtained using a temperature of 25 °C. Finally,
331 the injection volume was tested (20, 30 and 40 µL) and the chromatographic response
332 was improved, without loss of resolution or peak shape, when an injection volume of 40
333 µL was used. A representative chromatogram of a 10 µg/mL standard mixture of the
334 compounds analyzed is presented in Figure 3a.

335

336 Insert Figure 3

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338 **3.2.2. Fast-HPLC analysis**

339 Upon reducing the particle size from chromatographic column from 5 µm to 2.2 µm,
340 it was possible to use higher flow rates, obtaining a faster separation. Parameters such
341 as elution gradient and flow rate were optimized in order to find the best
342 chromatographic resolution and to obtain narrower peaks in the shortest analysis time.
343 This was achieved using a gradient elution performed on 4 minutes with an optimal
344 flow rate of 2.0 mL/min. The influence of different temperatures in the range of 25 °C to

345 35 °C was also studied and the best peak shapes was obtained for a temperature of 25
346 °C. Lastly, the injection volume was varied from 5 to 10 µL in order to improve the
347 chromatographic response without loss of the peak shapes. This was achieved for an
348 injection volume of 10 µL. An example of a chromatogram obtained with the optimized
349 chromatographic conditions is depicted in Figure 3b.

350

351 3.3. Method validation

352 The performance of the developed methods was validated in terms of sensitivity,
353 linearity, recoveries, precision (intra- and inter-day) and interference of the matrix,
354 using river water. Detailed analytical quality assurance data is shown in Table 2.

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Insert Table 2

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358 As it is depicted in Table 2a, HPLC and fast-HPLC methods showed differences in
359 what concern to sensitivity, being the lowest MDL and MQL obtained for the former.

360 Linearity was studied in the range of 50 to 1500 µg/L, setting calibration curves
361 using linear regression analysis. Depending on the sensitivity reached by the
362 chromatographic system employed, each analyte presents different linear responses.
363 Both methods gave correlation coefficients (r^2) higher than 0.999 for all analytes (Table
364 2a). A six point calibration curve for each compound was daily performed and the
365 possible fluctuation in signal intensity was checked by injecting a standard solution at
366 two concentration levels after each eight injections.

367 Accuracy of the method was estimated from recovery experiments of the target
368 analytes at different concentration levels. Four fortification levels in river water were
369 tested for both methods; however fast-HPLC method could not be validated to the 0.75

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370 $\mu\text{g/L}$ level due to its lower sensitivity comparatively to HPLC method, but it was
371 validated to the other fortification levels (2.50, 3.75 and 5.00 $\mu\text{g/L}$) as is shown in Table
372 2b. The recoveries obtained were satisfactory for all the compounds (recovery from 60
373 to 120%, with RSD values lower than 6%), except for *p*-aminophenol. In this case, a
374 value lower than 60% for 0.75 $\mu\text{g/L}$ spiking level was obtained, which could be related
375 with losses during the pre-concentration step due to the physico-chemical properties of
376 the compound.

377 The precision of the method was evaluated in terms of repeatability (intra-day) and
378 reproducibility (inter-day), exhibiting RSD values below to 3% and 6%, respectively,
379 for both methods (Table 2a).

380 The influence of the matrix in the UV signal was evaluated by comparing a
381 calibration curve in river extract with one achieved for the standards prepared in a
382 mixture of methanol-water (10:90, *v/v*). The calibration curves obtained for all the
383 compounds exhibited similar slopes and higher Y-intercept values for the river extract,
384 indicating that there is a slightly matrix effect, however this is similar within the linear
385 range. In order to correct the matrix effect, the standard addition method was used.

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387 **3.4. Application to real samples**

388 In order to demonstrate the applicability of the developed method, samples from
389 seven rivers located in the north of Portugal were analyzed. As SPE-HPLC/DAD
390 showed to be more sensitive, with lower MDL and MQL for all the compounds,
391 samples were only analyzed using this method. Quantification of the river samples was
392 done using the standard addition method and the confirmation of the positive findings
393 was carried out by comparing the UV spectra of the peaks present in the samples with
394 that one obtained in the standard solution chromatogram. An example of a

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2 395 chromatogram of a river sample obtained with the developed methodology is shown in
3 396 Figure 4.

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11 400 All the compounds were detected in Portuguese rivers (Table 3). As expected,
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13 paracetamol reported the lowest concentrations (up to 0.25 µg/L), since it is efficiently
14 401 removed (approximately 99%) in WWTPs [2]. The results are in agreement with
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16 402 concentrations of paracetamol reported in literature in rivers from Spain [17], Serbia
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18 403 [16] and United Kingdom [9].
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30 408 On the other hand, the principal metabolite of paracetamol (paracetamol-
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32 glucuronide) and its main transformation product (*p*-aminophenol) could be detected in
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34 409 Portuguese rivers at levels up to 3.57 µg/L and 1.63 µg/L, respectively, usually having
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36 410 higher concentrations than paracetamol (Table 3). Given that paracetamol-glucuronide
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38 411 entry in the environment through human excretion and it was previously detected in a
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40 412 WWTP effluent at levels up to 462 µg/L [21], WWTP effluents may be pointed out as
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42 413 the principal source of entrance of this metabolite into surface waters.
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48 415 *p*-Aminophenol was identified as a transformation product of paracetamol and its
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50 416 presence reported in wastewaters [23]. Our results showed that *p*-aminophenol could
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52 417 also be found in surface waters, being detected in five out of seven studied rivers, with
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54 418 concentrations from below MQL (<0.23 µg/L) to 1.63 µg/L (Table 4). Although these
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56 419 concentrations are lower than those one reported in two Romanian rivers [40].
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4. Conclusions

A rapid, simple and sensitive method for the simultaneous determination of paracetamol, paracetamol-glucuronide and *p*-aminophenol in river water was developed.

The extraction procedure allowed the analysis of three related compounds with different physico-chemical properties in a single step using two Oasis WAX cartridges in tandem and a sample pH of 7. Recoveries higher than 60% were obtained for all the compounds. The SPE procedure allows the elimination of interferences and, at the same time, a pre-concentration of the analytes, depicted in MDLs from 40 to 67 ng/L.

Comparatively to HPLC-MS/MS, HPLC-DAD has the advantageous of being an inexpensive analytical technique that can be seen as an affordable, useful and cost-effective alternative for routine analysis of pharmaceuticals, their metabolites and transformation products in environmental waters.

The developed method was applied to the determination of the selected analytes in samples from seven Portuguese rivers, showing their occurrence in surface waters with levels up to few micrograms per litre. The results obtained shows that besides pharmaceuticals, scientific community should also focus its attention in the evaluation of the presence of metabolites and transformation products in the environment, seeing that, sometimes, they may be at higher concentrations than the parent compounds. The proposed methodology may be applied to monitoring the behaviour of the selected analytes during wastewater treatment as well as their ability to adsorb to soil and sediments.

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445 **Acknowledgments**

1
2 446 This work has been supported by Fundação para a Ciência e a Tecnologia (FCT)
3
4 447 through the grant no. Pest-C/EQB/LA0006/2011, project PTDC/ECM/103141/2008 and
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7 448 project PTDC/AAC-AMB/120889/2010. Lúcia H.M.L.M. Santos thanks to FCT and
8
9 449 FSE/POPH for her PhD grant (SFRH/BD/48168/2008).
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515 **Figure captions:**

516

517 **Figure 1** – Recoveries obtained for the target analytes in ultra-pure water with
518 different SPE cartridges. PCT-G – Paracetamol-glucuronide; PAP – *p*-Aminophenol;
519 PCT – Paracetamol

520

521 **Figure 2** – Recoveries obtained in ultra-pure water at different sample's pH. a)
522 Paracetamol-glucuronide; b) *p*-Aminophenol; c) Paracetamol

523

524 **Figure 3** – Example of a chromatogram of a standard mixture 10 µg/mL for the
525 selected analytes analyzed by a) HPLC and b) Fast-HPLC. 1 – Paracetamol-
526 glucuronide; 2 – *p*-Aminophenol; 3 – Paracetamol

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528 **Figure 4** – Chromatogram of Douro river sample. 1 – Paracetamol-glucuronide; 3 –
529 Paracetamol

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Highlights

- Determination of paracetamol, paracetamol-glucuronide and *p*-aminophenol in rivers
- Development of a SPE procedure for their simultaneous determination
- All compounds were detected in rivers with concentrations up to few $\mu\text{g/L}$
- First time that paracetamol-glucuronide was found in river water

Accepted Manuscript

Figure 1

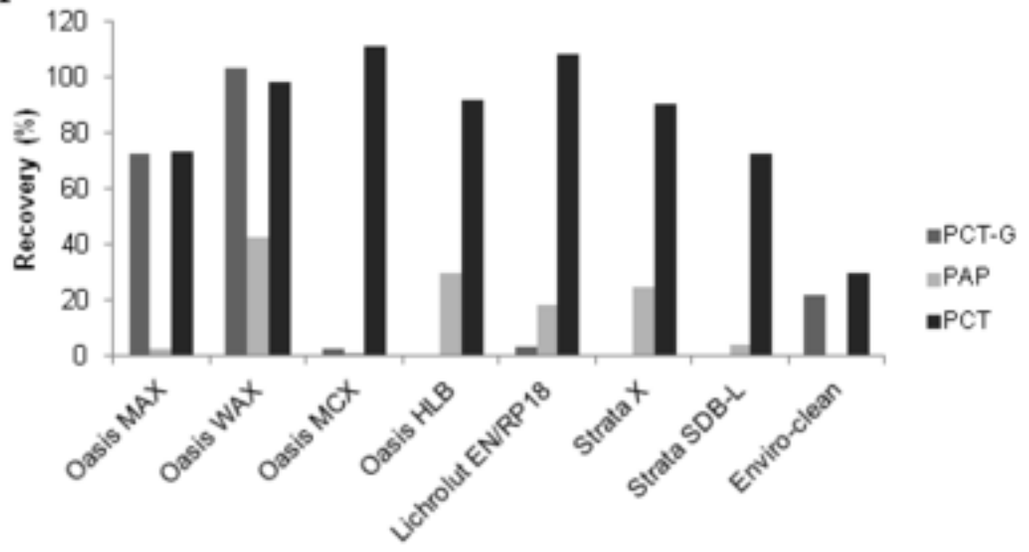
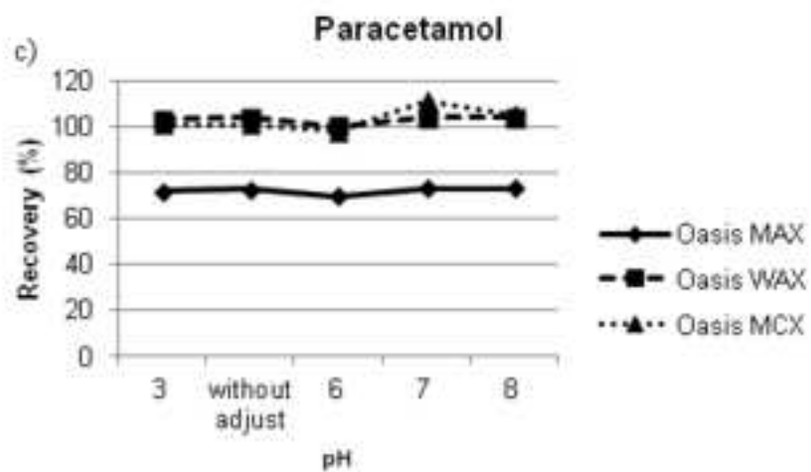
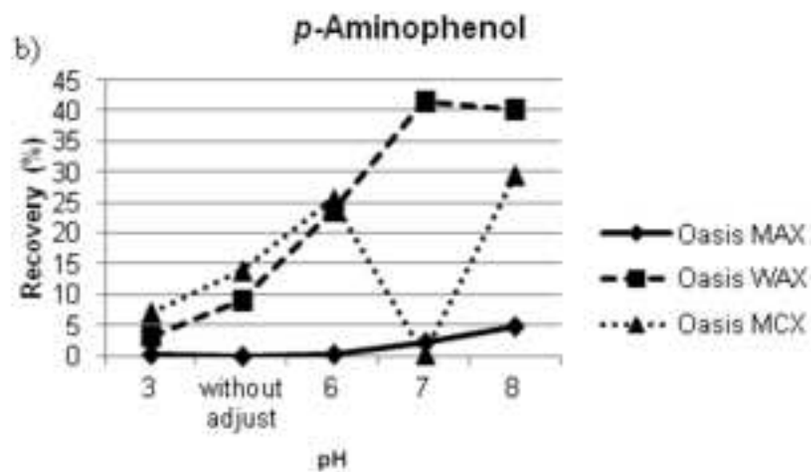
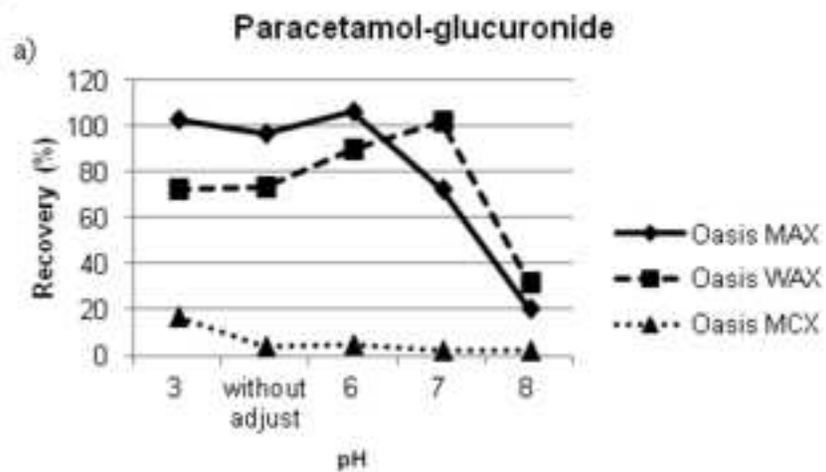
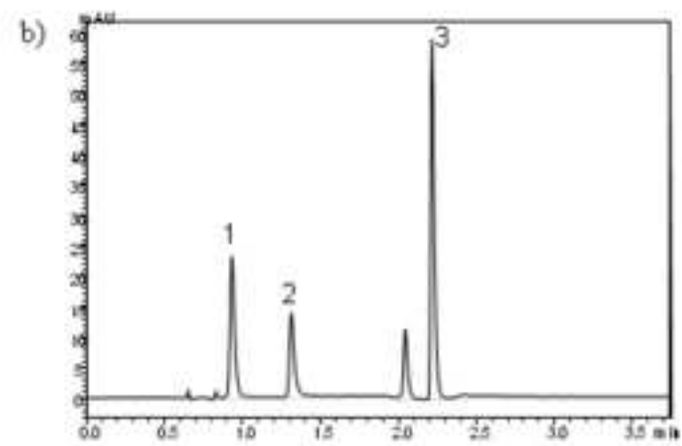
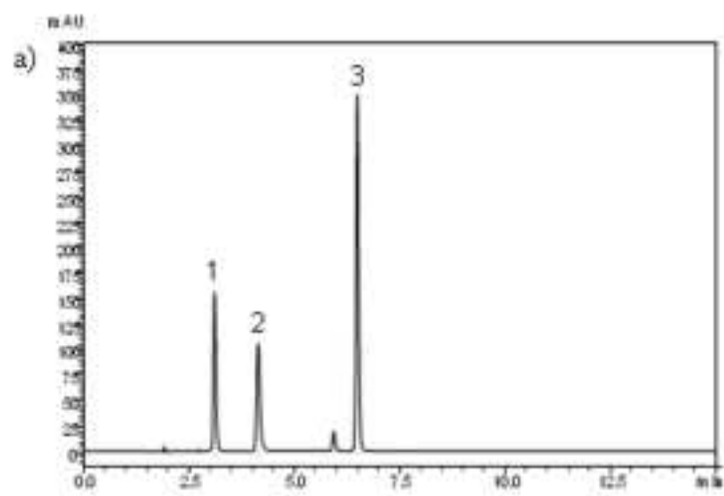


Figure 2



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Figure 3



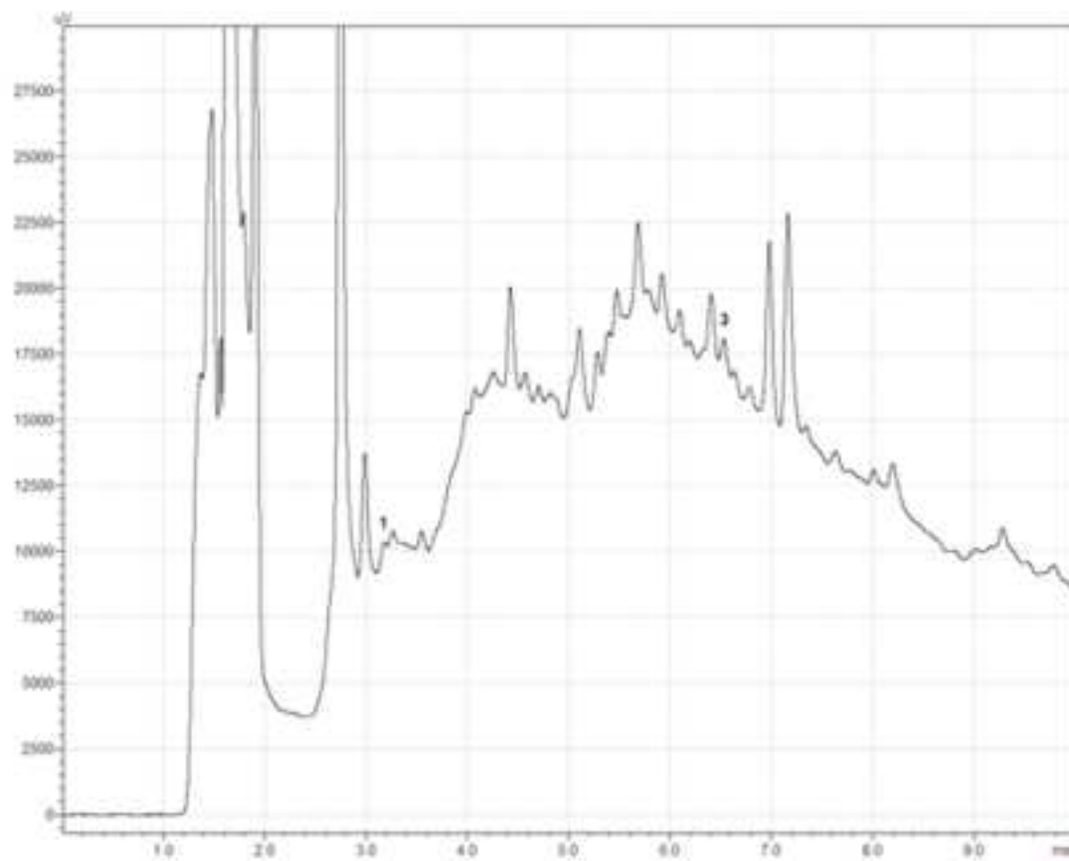
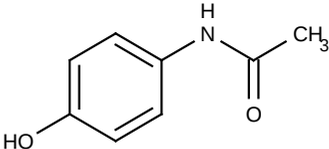
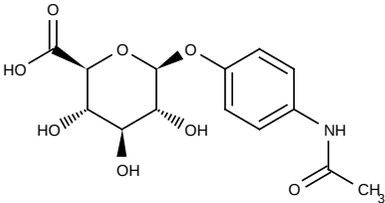
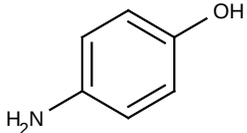


Table 1 – Physico-chemical properties of the selected compounds

	Chemical structure	Formula	CAS no.	M _w	Solubility (g L ⁻¹)	pK _a ^a	log K _{OW} ^b
Paracetamol (PCT)		C ₈ H ₉ NO ₂	130-90-2	151.16	12.8	9.46	0.46
Paracetamol-glucuronide (PCT-G)		C ₁₄ H ₁₇ NO ₈	120595-80-4	327.29	27.7	3.17; 12.22	-1.23
<i>p</i> -Aminophenol (PAP)		C ₆ H ₇ NO	123-30-8	109.13	15.0	5.43; 10.40	0.24

^aAdapted from reference [38]^bAdapted from reference [39]

Table 2 – a) Linearity, detection and quantification limits of the method (MDL, MQL), precision intra- and inter-day of the developed SPE-HPLC/DAD and SPE-fast-HPLC/DAD methods. **b)** Recoveries obtained, expressed in percentage (%), for the selected analytes in river water for the developed methods.

a)

Compound	SPE-HPLC/DAD						SPE-fast-HPLC/DAD					
	Linear range (µg/L)	Correlation coefficient (r^2)	MDL (µg/L)	MQL (µg/L)	Precision intra-day (% RSD)	Precision inter-day (% RSD)	Linear range (µg/L)	Correlation coefficient (r^2)	MDL (µg/L)	MQL (µg/L)	Precision intra-day (% RSD)	Precision inter-day (% RSD)
PCT-G	50-1500	0.9999	0.040	0.134	0.547	0.836	50-1500	0.9997	0.199	0.664	2.75	5.06
PAP	50-1500	0.9999	0.067	0.225	0.368	0.632	500-1250	0.9990	0.275	0.905	2.14	4.39
PCT	50-1500	0.9999	0.042	0.141	0.097	0.958	50-1500	0.9996	0.210	0.701	0.913	3.56

b)

Compound	SPE-HPLC/DAD				SPE-fast-HPLC/DAD		
	Recoveries (RSD %)						
	Spiking level (µg/L)				Spiking level (µg/L)		
	0.75	2.50	3.75	5.00	2.50	3.75	5.00
PCT-G	100 (2.4)	117 (2.1)	110 (0.7)	112 (3.0)	108 (2.5)	99 (0.7)	112 (1.2)
PAP	53 (5.5)	62 (4.3)	64 (0.5)	60 (0.2)	62 (3.7)	65 (4.0)	59 (2.7)
PCT	99 (1.5)	103 (2.3)	107 (3.9)	109 (0.02)	106 (3.5)	105 (4.0)	108 (0.9)

Table 3 – Concentration of the analyzed compounds, expressed in $\mu\text{g/L}$, in river waters from the north of Portugal. Standard deviation (SD) values are indicated in brackets.

Sample	Paracetamol-glucuronide	<i>p</i> -Aminophenol	Paracetamol
Cabrum river	<MQL	n.d.	<MDL
Douro river	<MQL	n.d.	<MQL
Ave river	0.36 (± 0.02)	1.63 (± 0.05)	0.17 (± 0.002)
Leça river	3.57 (± 0.06)	1.25 (± 0.02)	0.25 (± 0.01)
Tâmega river	<MQL	0.40 (± 0.01)	<MDL
Lima river	<MQL	<MQL	<MDL
Minho river	0.18 (± 0.01)	0.52 (± 0.05)	<MDL

n.d. – not detected; <MDL – below method detection limit; <MQL – below method quantification limit