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Chen Dongmei, Jean-Michel Delmas, Dominique Hurtaud Pessel, Eric Verdon. Development of a multi-class method to determine nitroimidazoles, nitrofurans, pharmacologically active dyes and chloramphenicol in aquaculture products by liquid chromatography-tandem mass spectrometry. Food Chemistry, Elsevier, 2020, 311, pp.125924. 10.1016/j.foodchem.2019.125924 . anses-02400704

HAL Id: anses-02400704

<https://hal-anses.archives-ouvertes.fr/anses-02400704>

Submitted on 21 Dec 2021

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1 **Development of a multi-class method to determine nitroimidazoles,**
2 **nitrofurans, pharmacologically active dyes and chloramphenicol in**
3 **aquaculture products by liquid chromatography-tandem mass**
4 **spectrometry**

5
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16 Non-Authorised Antimicrobial Veterinary Medicinal Product Residues and for Banned
17 Pharmacologically Active Dyes, Fougères, France

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19 **ABSTRACT**

20 LC-MS/MS method was developed for the efficient identification and quantification of 21
21 banned substances including various nitroimidazoles, nitrofurans, pharmacologically-active
22 dyes and chloramphenicol, respectively in aquaculture products. The sample preparation was
23 started by acid-treatment with 2-nitrobenzaldehyde (NBA) to liberate matrix-bound residues
24 of nitrofurans. A modified QuEChERS method was optimized for the extraction and clean-up
25 of the target analytes. The metabolites of the four conventional nitrofurans (nitrofurantoin,
26 furazolidone, nitrofurazone and furaltadone) and of three other nitrofurans (nifursol,
27 nifuroxazide, and nitrovin), and an underivatizable nitrofuran (nifurpirinol) were
28 simultaneously detected. Furthermore, 21 banned substances were quantified by LC-MS/MS
29 with ESI using one single injection. To evaluate and validate the performance of the method
30 the criteria of the Decision (EC) no 2002/657 were applied. Decision limit ($CC\alpha$) of target
31 analytes ranged 0.067-1.655 $\mu\text{g}/\text{kg}$ in aquaculture products. The recovery ranged
32 77.2%-125.6%, and the relative standard deviations of inter-day analyses (RSD) were less
33 than 25%.

34 **KEY WORDS:** Nitroimidazoles; Nitrofurans; Dyes; Chloramphenicol; Residues; LC-MS/MS;
35 Aquaculture products

36

37 **1. Introduction**

38 Veterinary medicinal products (VMPs), such as nitrofurans (NFs), chloramphenicol (CAP),
39 nitroimidazoles (NIIMs), and also the non-veterinary product pharmacologically-active
40 were used extensively in aquaculture because of their low costs and high effectiveness.
41 According to European Union legislation, the use of these compounds has been banned for
42 food producing animals. They have been classed in table 2 of Commission regulation, except
43 for dye compounds which do not enter this legislation because they never have been recorded as
44 medicinal products (European Commission, 2009). Nevertheless no residues of these
45 substances shall be found in food products. NFs are active broad-spectrum antibacterial drugs
46 and in the past have been widely used in veterinary medicine. The most often used NFs
47 compounds are furazolidone, furaltadone, nitrofurazone and nitrofurantoin. As mentioned by
48 Hoogenboom et al. (Hoogenboom, Berghmans, Polman, Parker, & Shaw, 1992), McCracken &
49 Kennedy (1997), and Zuidema et al. (2004), they can be rapidly metabolized into semicarbazide
50 (SEM), 3-amino-2-oxazolidinone (AOZ), 3-amino-5-morpholinomethyl-2-oxazolidinone
51 (AMOZ), and 1-aminohydantoin (AHD), respectively. Recently, some other NFs including
52 nifursol, nitrovin, nifuroxazide, and nifurpirinol (NPIR) have been notified due to having the
53 similar structure. They are metabolized into 3,5-dinitrosalicylic acid hydrazine (DNSH),
54 aminoguanidin hydrochloride (AMG), and salicylic acid hydrazine (PSH), respectively, and
55 excluding NPIR undertaking no metabolisation. The chemical structures of parent compounds
56 of NFs and their metabolites are shown in Fig. 1. The most frequently used NIIMs including
57 metronidazole (MNZ), dimetridazole (DMZ), ronidazole (RNZ), and ipronidazole (IPZ) help
58 combatting anaerobic bacterial and parasitic infections. The analytical method should cover as

59 well the three metabolites of these NIIMs including hydroxy-metronidazole (MNZOH),
60 2-hydroxymethyl-1-methyl-5-nitroimidazole (HMMNI), and hydroxy-iproimidazole (IPZOH),
61 respectively. Triphenylmethane dyes have been used as antiparasites and antifungus infection
62 agents in aquaculture, including malachite green (MG), crystal violet (CV) and brilliant green
63 (BG). MG and CV can be easily metabolized into leucomalachite green (LMG) and
64 leucocrystal violet (LCV), respectively. CAP is a broad-spectrum antibacterial drug which was
65 also widely used in veterinary treatments.

66 However, these VMPs and Dyes are prohibited globally due to their carcinogenic and
67 mutagenic potency or to the additional risk of causing aplastic anemia in the case of CAP.
68 These compounds are currently regulated at the target level of 1 µg/kg for NFs, 3 µg/kg for
69 NIIMs, 2 µg/kg for Dyes, and 0.3 µg/kg for CAP in the EU in accordance with the respective
70 Minimum Required Performance Limits (MRPL) set for CAP, NFs and MG (Dyes) (European
71 Commission, 2003; European Commission, 2004) and later Reference Points for Action
72 (RPA) set for the same CAP, NFs and MG (Dyes) (EFSA, 2014; EFSA, 2015; EFSA, 2016)
73 and in accordance with the EU-RLs Recommended Limits set for NIIMs (CRL Guidance,
74 2007). In order to reduce the number of implemented analytical methods for the control of
75 banned antimicrobial and dye residues in food from animal origin and especially in
76 aquaculture products, it is of interest to combine them all when possible but with keeping high
77 standard level of reliability of the official control.

78 The recent mostly used methods for the determination of NFs, NIIMs, Dyes and CAP
79 were LC-MS/MS due to the unambiguous identification and accurate quantification
80 possibilities. For NFs, most currently published analytical methods mainly focused on AOZ,

81 AMOZ, SEM, and AHD in animal derived food (Zhang et al., 2017; Kaufmann, Butcher,
82 Maden, Walker, & Widmer, 2015; El-Demerdash, Song, Reel, Hillegas, & Smith, 2015;
83 Verdon, Couedor, & Sanders, 2007; Shendy, Al-Ghobashy, Alla, & Lotfy, 2016). There are
84 very few methods for determination of the four other above-mentioned NFs. Kaufmann et al.
85 (2015) have developed a method to determine PSH, DNSH and AMG. Verdon et al. (2007)
86 reported a method to analyze DNSH in poultry muscle. There were many methods applied to
87 analyze NIIMs in animal-derived food (Zhang et al., 2017; Granja et al., 2013; Tölgyesi,
88 Sharma, Fekete, Fekete, Simon, & Farkas, 2012; Cronly et al., 2009; Hurtaud-Pessel,
89 Delepine, & Laurentie, 2000). There also have been many methods to determine Dyes in
90 aquaculture products (Hurtaud-Pessel, Couëdor, Verdon, & Dowell, 2013; Schneider, &
91 Andersen, 2015; Kaplan, Olgun, & Karaoglu, 2014; Ascari, Dracz, Santos, Lima, Diniz, &
92 Vargas, 2012). For CAP, many analytical methods have also been reported over the 20 past
93 years.

94 However, very few multi-class methods were applied for the simultaneous analysis of
95 these four groups of target analytes in animal-derived food due to their different
96 physicochemical characteristics. Zhang et al. (2017) have reported to analyze four NFs (AOZ,
97 AMOZ, SEM, and AHD), 7 NIIMs and CAP in chicken muscle and eggs. Shendy et al. (2016)
98 described a method to simultaneously determine NFs and NIIMs including AOZ, AHD,
99 AMOZ, SEM, RNZ, and DMZ in honey. There are a few methods to determine NFs and CAP
100 in animal food product (Kaufmann et al., 2015; El-Demerdash et al., 2015; An et al. 2015;
101 Veach, Baker, Kibbey, Fong, Broadaway, & Drake, 2015). Up to now, there are no methods
102 for analysis of all the four groups of target analytes simultaneously in animal-derived food.

103 The sample preparation was a crucial factor for analysis of target analytes. NFs
104 metabolites are strongly bound to proteins, which need to be released through mild acid
105 hydrolysis before analysis. The derivatization of NFs metabolites has been recognized to be
106 essential, since the strong polar (poor retention on RP columns) of underivatized metabolites
107 show poor ionization properties in the electrospray interface of a mass spectrometer. Most
108 methods for NFs analysis applied HCl for hydrolysis and 2-NBA for metabolite derivatization
109 at 37 °C for 14 h followed by either liquid-liquid extraction or solid-phase extraction, which is
110 time-consuming. For NIIM, Dyes and CAP, the mostly used methods extracted these
111 substances either with ethyl acetate or acetonitrile or with a buffer solution, and followed by a
112 solid-phase extraction (SPE) or a liquid-liquid extraction clean-up. For all the four groups of
113 target analytes, the sample preparation became a subtle procedure to elaborate for achieving
114 acceptable extraction and/or efficient clean-up at the same time for all groups.

115 The aim of our project is to develop a multi-residue method based on LC-MS/MS
116 capable of reaching low Reference Point for Action (RPA) levels for the control for
117 chloramphenicol, nitrofurans and their metabolites, and possible dyes or nitroimidazoles in
118 the aquaculture products. In this article, LC-MS/MS in both positive and negative ESI modes
119 is used to detect multi-banned substances. The sample preparation procedure was initiated
120 with hydrolysis and derivatization followed by modified QuEChERS, which simplified the
121 extraction method and reduced the extraction time. Internal standards are used to reach more
122 accurate quantification. The performance of the method was evaluated and validated
123 according to the criteria of the Decision (EC) no 2002/657.

124

125 **2. Materials and methods**

126

127 **2.1 Chemicals and reagents**

128 **Analytes:** 1-amino-hydantoin (AHD), 3-amino-2-oxazolidinone (AOZ),
129 3-amino-5-morpholinomethyl-2-oxazolidinone (AMOZ), semicarbazide (SEM),
130 aminoguanidin hydrochloride (AMG), 3,5-dinitrosalicylic acid hydrazine (DNSH), nifurpirinol
131 (NPIR), salicylic acid hydrazine (PSH), dimetridazole (DMZ), metronidazole (MNZ),
132 ronidazole (RNZ), ipronidazole (IPZ), hydroxy-metronidazole (MNZOH),
133 2-hydroxymethyl-1-methyl-5-nitroimidazole (HMMNI), hydroxy-ipronidazole (IPZOH),
134 malachite green (MG), leuco-malachite green (LMG), crystal violet (CV), leuco-crystal violet
135 (LCV), brilliant green (BG), and Chloramphenicol (CAP).

136 **Internal standards:** NP-AHD-¹³C₃, NP-AOZ-D₄, NP-SEM-¹³C, ¹⁵N₂, NP-AMOZ-D₅,
137 DMZ-D₃, HMMNI-D₃, IPZ-D₃, MNZ-¹³C₂, ¹⁵N₂, IPZOH-D₃, MNZOH-D₂, RNZ-D₃, MG-D₅,
138 LMG-D₅, CV-D₆, LCV-D₆, and CAP-D₅.

139 These analytes and internal standards were obtained from Sigma-Aldrich (St-Quentin
140 Fallavier, France), Witega (Berlin, Germany), LGC standards (Molsheim, France),
141 Ehrenstorfer GmbH (Augsburg, Germany), and CIL Cluzeau (Ste Foy-La-Grande, France).

142 HPLC-MS grade methanol, acetonitrile, ethyl acetate, and iso-hexane were purchased
143 from Fisher Scientific (Illkirch, France). HPLC grade formic acid and ammonium formate were
144 from Fisher Scientific (Illkirch, France). Nitrobenzaldehyde (2-NBA), hydrochloric acid, and
145 sodium chloride were obtained from Sigma-Aldrich (St-Quentin Fallavier, France).
146 Di-potassium hydrogen phosphate was bought from VWR (Fontenay sous Bois, France).

147 **2.2 LC-MS/MS analysis**

148 HPLC system was composed of HPLC pump Ultimate 3000, Autosampler Ultimate 3000,
149 and Column oven 3000 (Dionex, Villebon sur Yvette, France). Symmetry C₁₈ analytical column
150 (5 μm, 100 x 2.1 mm) from Waters Co. (Guyancourt, France) was used to separation the
151 analytes. The mobile phase was 20 mM ammonium formate+0.02% formic acid (A) and
152 methanol (B) with a gradient elution at 1.0 ml/min flow rate as follows (t in min): t₀, A = 90%;
153 t₆, A = 10%; t₉, A = 10%; t_{9.1}, A =90%; t₁₂, A = 90%.

154 Mass spectrometry was TSQ Vantage (Thermo scientific, Villebon sur Yvette, France)
155 with electrospray interface (ESI). The source parameters were as follows: Spray voltage: 3000
156 V; Vaporizer temperature: 300 °C; Sheath gas pressure: 30 psi; Aux gas pressure: 30 psi;
157 Capillary temperature: 300 °C; Cycle time: 0.5 s. The mass spectrometer was operated in a
158 selective reaction monitoring (SRM) mode selecting one precursor ion and two product ions per
159 each target compound, except for internal standard for which only one SRM transition was
160 monitored for each The SRM acquisition parameters, ie. m/z precursor ion, m/z product ion,
161 and collision energy are shown in Table S1.

162 **2.3 Samples**

163 Five hundred grams of aquaculture products, including salmon, trout, and shrimp, were
164 purchased from a local supermarket. After being homogenized in a high-speed food blender,
165 the samples were stored below -20 °C prior to using for the method developed . The samples
166 taken for the validation and the way the sampling was operated. One gram portion of samples
167 was taken to enter the extraction-purification process. The goal was to demonstrate the
168 capacity of the method to deliver accurate data on different fish/aquaproduct species.

169 **2.4 Sample preparation**

170 One gram portion of sampled flesh was added 5 mL of hydrochloric acid (0.1 mol/L) and
171 0.2 mL of 50 mmol/L of nitrobenzaldehyde solution freshly prepared in methanol. After
172 derivatization for 2 h at 60 °C with moderate shaking, 3 mL of 0.5 mol/L di-potassium
173 hydrogen phosphate, 5 mL of iso-hexane, and 2 g NaCl were added and mixed for 5 min.
174 After centrifugation for 5 min at 6000 g at 4 °C, the (upper) iso-hexane phase was discarded.
175 Five milliliters of ethyl acetate were added to the remaining extraction solution, and vortexed
176 for 30 s, and then mixed for 10 min. After centrifugation for 10 min at 6000 g at 4 °C, 5 mL of
177 the (upper) organic phase was transferred to a 10 mL polypropylene tube. Five milliliters of
178 acetonitrile were added to the samples again, and vortexed 30 s and mixed for 10 min. After
179 centrifugation for 10 min at 6000 g at 4 °C, 5 mL of the (upper) organic phase was
180 transferred to a 10 mL polypropylene tube. Then, all the extraction solution was collected and
181 then evaporated under gentle nitrogen flow at 50 °C after adding of 2 g MgSO₄ in order to
182 obtain an oily residual phase. The residues were then dissolved with 0.5 mL of acetonitrile
183 and filtered through a 0.22 µm PVDF filter.

184 ***2.5 Method validation according to Decision No. (EC) 2002/657***

185 2.5.1 Identification parameters

186 The performance of the method was assessed through its qualitative parameters: analyte
187 specificity, molecular identification in terms of retention time (RT), and of transition ion
188 ratios. The specificity of the assay was demonstrated by analyzing 20 representative blank
189 tissue samples and checking interfering peaks at the retention time of target analytes.
190 According to European Union Commission Decision No. 2002/657 (European Commission,
191 2002) with a minimum total score of 3 for Group B (authorized substances) or 4 for Group A

192 (non-authorized substances) identification points, one precursor ion and two product ions
193 were sufficiently monitored to fulfil this requirement. The analytes were additionally
194 identified by matching retention times of peaks with the values of the corresponding standard
195 analyzed under the same experimental conditions. The analyte in the sample should be eluted
196 at the retention time corresponding to the analyte in spiked samples (within a range of relative
197 retention time of $\pm 2.5\%$). Each analyte ion ratio was effectively measured on each
198 chromatogram, corresponding to the less intense SRM transition signal against the most
199 intense SRM transition ion one.

200 2.5.2 Quantitative parameters

201 The performance of the method was assessed through its 3 main quantitative parameters:
202 – 1) accuracy demonstrated in terms of trueness – 2) precision, the precision being expressed
203 as the intra- and inter-day/series repeatabilities, and – 3) confirmatory analytical limits (limit
204 of decision $CC\alpha$ and capacity of detection $CC\beta$). The validation was performed with a set of
205 three series of analyses including salmon, trout, and shrimp. For each of the series, the
206 experiment comprised 24 samples: 6 calibration standard (CS) samples over the range of
207 calibration claimed for each group of substances, and 18 validating standard (VS) samples set
208 at three concentrations and for each concentration being repeated 6 times. One of the series
209 was analysed each day. The CS and VS samples are matrix-spiked samples, ie they are
210 prepared by addition of standards to blank matrix prior to extraction. $CC\alpha$ and $CC\beta$ were
211 recommended in Commission Decision No. (EC) 2002/657. In this article, $CC\alpha$ and $CC\beta$
212 were calculated as described by Verdon et al. (2007).

213 This calibration curve was built from the CS samples using linear regression model
214 $AX+B$ using specific internal standard for each substance. This calibration curve was then
215 used to measure the back-calculated concentration of each compound in the validating

216 samples (VS). The calibration range of the CS samples for each substance was made of 6
217 levels including the negative control: for Nitrofurans, at 0.0, 0.5, 1.0, 1.5, 2.0, and 3.0 $\mu\text{g}/\text{kg}$;
218 for Dyes, at 0.0, 1.0, 2.0, 3.0, 4.0, and 5.0 $\mu\text{g}/\text{kg}$; for Nitroimidazoles, at 0.0, 1.5, 3.0, 4.5, 6.0,
219 and 7.5 $\mu\text{g}/\text{kg}$; and for CAP, at 0.0, 3.0, 4.5, 6.0, 7.5, and 9.0 $\mu\text{g}/\text{kg}$. Internal standard
220 concentration used for Nitrofurans, Dyes, Nitroimidazoles, and CAP were 2.0, 2.0, 5.0, and
221 5.0 $\mu\text{g}/\text{kg}$, respectively.

222 The percentage of trueness for the estimated concentration of the analytes
223 back-calculated from the validating standards (VS) was expressed as the bias measured from
224 the actually spiked concentration and estimated for each analyte at three levels (0.5, 1.0, and
225 2.0 $\mu\text{g}/\text{kg}$ for Nitrofurans; 1.0, 2.0, and 4.0 $\mu\text{g}/\text{kg}$ for Dyes; 1.5, 3.0, and 6.0 $\mu\text{g}/\text{kg}$ for
226 Nitroimidazoles, and 3.0, 4.5, and 6.0 $\mu\text{g}/\text{kg}$ for CAP). Each concentration was repeated 6
227 times within a day for the intra-day precision test. The precision in terms of repeatability and
228 reproducibility was evaluated by calculating the relative standard deviation (RSD) for each
229 analyte at each level of concentration.

230

231 **3. Results and discussion**

232 ***3.1 Optimization of LC-MS/MS***

233 In order to obtain nice resolution and high sensitivity, mobile phase and analytical
234 column should be chosen based on the evaluation of ionization efficiency. Almost all
235 references covering the topic of our project claimed LC separations performed on
236 reversed-phase (RP) materials by applying either methanol/water or acetonitrile/water
237 mixtures as mobile phase operated in the gradient as well as isocratic mode. In some cases,
238 acetic acid, formic acid, and ammonium salts are added to support analyte ionization and
239 improve the chromatographic separation efficiency. So, methanol, acetonitrile, formic acid,
240 and different ratio of ammonium acetate (aqueous ammonia) were tested. Better optimized

241 sensitivity was observed when using a mixture of methanol and of 20 mmol/l ammonium
242 formate with 0.02% formic acid.

243 Four different columns were tested to separate the target compounds, including Acquity
244 BEH C₁₈ (50 X 2.1 mm, 1.7 μm), Accucore Phenyl Hexyl (100 X 2.1 mm, 4.6 μm), Symmetry
245 C₁₈ (150 X 3.9 mm, 5 μm), Colone Hypersil Gold (50 X 2.1 mm, 1.9 μm). The results showed
246 that Symmetry C₁₈ (3.9 X 150 mm, 5 μm) gave both good peak shapes and higher ion
247 response.

248

249 **3.2 Optimization of sample preparation**

250 3.2.1 Optimization of acid hydrolyzed and derivatized conditions

251 Since its easy combination with proteins *in vivo*, Nitrofuran metabolites need to be
252 released from tissues by means of acid hydrolysis, and simultaneously derivatized with
253 2-NBA after its release from the tissue proteins. The commonly used derivatization procedure
254 was 14 h at 37 °C (Kaufmann et al., 2015) or 4 h at 55 °C (Verdon et al., 2007). The
255 derivatization temperature and time have been investigated to improve the extraction efficient
256 and shorten the derivatization time. First, we compared the derivatization temperature and
257 time according to published articles (14 h at 37 °C, and 4 h at 55 °C). We also tested another
258 third condition (2 h at 60 °C) to try shortening the derivatization time. The results showed that
259 derivatization for 2 h at 60 °C delivered slightly higher recovery for most of the nitrofuran
260 analytes (Fig.1).

261 Moreover, different derivatization time (2, 3, and 4 h) were performed at 60 °C to
262 evaluate the extraction efficiency. When increasing the derivatization time, the recovery of
263 nitrofuran metabolites did not significantly improve. So, 2 h was finally selected as extraction
264 time (Fig. 1).

265 In this study, we also confirmed that the acid hydrolysis and derivatization steps both

266 may affect the recovery of NIIMs, Dyes and CAP. When the derivatization time was
267 increased up to 4 h, the recovery showed a start of decrease for Dyes and did not significantly
268 change for NIIMs and CAP.

269 Moreover, the pH value of derivatization was also tested. When adding the hydrochloric
270 acid and the nitrobenzaldehyde solution, the pH value was adjusted to 2.0, 2.5, and 3.0,
271 separately. The results showed that the derivatization efficiency for most of the analytes was
272 decreased with increasing pH values from 2.0 to 3.0.

273 3.2.2 Optimization of the extraction procedure

274 To extract the analytes from animal food matrix with as little interference and as much
275 high recovery as possible, is the most difficult and critical process. Originally, the QuEChERS
276 method involved a single extraction step, i.e. a sample clean up via dispersive solid phase
277 extraction using primary secondary amines (Anastassiades, Lehotay, Stajnbaher, & Schenck,
278 2003). In this study, a modified QuEChERS extraction protocol without sample clean up
279 followed by evaporation was optimized and employed.

280 The most important impacting factors were the extraction solvent and the extraction
281 volume. Some articles have reported to extract nitrofurans using ethyl acetate at pH 7
282 condition (Kaufmann et al., 2015; Verdon et al., 2007; Kim, Kim, Seok-Won, Lee, & Kim,
283 2015), some using acetonitrile (Shendy et al., 2016; An et al., 2015). For nitroimidazoles,
284 most published papers used ethyl acetate (Granja et al., 2013; Boison, Asea, & Matus, 2012)
285 and acetonitrile (Tölgyesi et al., 2012; Cronly et al., 2009) as the extraction solvent. Most
286 articles developed a method to determine Dyes using acetonitrile as the extraction solvent
287 (Hurtaud-Pessel et al., 2013; Schneider et al., 2015; Kaplan et al., 2014; Ascari et al., 2012;
288 Qin et al., 2016; Dowling, Mulder, Duffy, Regan, & Smyth, 2007). For chloramphenicol, ethyl

289 acetate was used most of the time as the extraction solvent.

290 According to the properties of the four-family analytes, ethyl acetate, acetonitrile, and
291 their mixture, at different pH values, were chosen as extraction solvents and compared. The
292 tested matrix was trout. From the results (Fig. 2), we observed the mixture of ethyl acetate
293 and acetonitrile exhibited the best extraction efficiency especially for MG, CV, and BG,
294 which was about more than 50% higher than those exhibited by other solvents. Therefore, the
295 mixture of ethyl acetate and acetonitrile was chosen as the extraction solvent. When
296 comparing extraction volume of ethyl acetate (5 mL and 10 mL), the extraction efficiency was
297 not significantly different. So, 5 mL ethyl acetate was considered sufficient to extract the
298 target VMP residues from fish matrix.

299 After extraction with acetonitrile, several salts were investigated as salting-out agents in
300 order to increase the ratio of the analytes into the acetonitrile layer and decrease water mixing
301 into the acetonitrile layer. After optimization of the parameters, salting out and complete
302 phase separation was achieved via addition of 2 g MgSO₄ and 2 g NaCl for having a higher
303 efficiency of extraction, especially for Dyes.

304 When developing the method using trout as tested matrix, there was no fat-soluble
305 interference. However, when using salmon as matrix, fat-soluble interference was observed.
306 So, before extracting, iso-hexane was added to decrease/remove the fat-soluble interference.
307 In order to determine the targeted compounds in all matrices using the same sample
308 preparation method, iso-hexane was used to remove fat-soluble interference before extraction
309 in the three species: i.e. trout, salmon, and shrimp.

310 After derivatization, the main step described by published articles was to neutralize the

311 solution by adding 5 mL of 0.1M di-potassium hydrogenophosphate followed by adding 1M
312 NaOH to adjust the pH to 7.0 ± 0.5 (controlled by pH strips). In order to simplify the
313 procedure, we tested different volumes and different concentrations of di-potassium
314 hydrogenophosphate to adjust pH to 7.0 ± 0.5 . After optimization, 3 mL of 0.5 mol/L
315 di-potassium hydrogen phosphate was finally kept to adjust pH to 7.0 ± 0.5 .

316 3.2.3 Optimization of reconstituting solvent recovery after evaporation step

317 The composition of the sample solvent medium before injection directly affects both the
318 separation behavior of analytes in the HPLC column and their sensitivity during LC - MS/MS
319 detection. For nitroimidazoles, nitrofurans, and chloramphenicol, most published articles
320 report using the mobile phase to dissolve/reconstitute the residues after the evaporation step.
321 However, for dyes, the residue was usually reconstituted with ammonium acetate-acetonitrile
322 (1:1) (Ascari et al., 2012), acetonitrile-water (60/40) (Huang, Zhao, Dai, Hou, Zhao, & Liang,
323 2016), or acetonitrile (Schneider et al., 2015). To select the optimal solvent medium, different
324 ratios of methanol-water, methanol-formic acid, methanol-ammonium formate solution, and
325 acetonitrile-water, acetonitrile-formic acid, and acetonitrile-ammonium formate solution were
326 compared in this study. The result showed that when using the mixture of methanol-water or
327 acetonitrile-water as reconstituting solvent, the extracting solvent recovery for dyes was very
328 low due to insufficient power of miscibility. When using pure methanol or pure acetonitrile to
329 dissolve the residues, a higher recovery was obtained (Fig. 3). Therefore, acetonitrile was
330 chosen as the dissolving/reconstituting solvent prior to injection. The injection volume for
331 LC-MS/MS mostly used in the published articles was 10 μ L. However, when injecting 10 μ L
332 to LC-MS/MS, the shape of nitroimidazole peaks were very wide. So, different volumes of

333 injection were tested, including 2, 5, 10 μL . The results showed that 2 μL as injection volume
334 led to a satisfactory peak shape for all the analytes.

335

336 **3.3 Method validation**

337 The matrix-spiked calibrations of nitroimidazoles, nitrofurans, dyes, and
338 chloramphenicol for salmon, trout, and shrimp were established, respectively. Matrix-spiked
339 calibration regression' coefficient of correlation R were in the range from 0.9916 to 0.9997
340 for nitrofurans over the concentration range of 0-3.0 $\mu\text{g}/\text{kg}$, from 0.9980 to 1.000 for
341 nitroimidazoles over the concentration range of 0-7.5 $\mu\text{g}/\text{kg}$, from 0.9909 to 0.9999 for dyes
342 over the concentration range of 0-5.0 $\mu\text{g}/\text{kg}$, and from 0.9962 to 0.9976 for dyes over the
343 concentration range of 0-9.0 $\mu\text{g}/\text{kg}$ in aquaculture products.

344 The CCAs of confirmation for nitroimidazoles, nitrofurans, dyes and chloramphenicol in
345 aquaculture products varied from 0.067 to 0.512 $\mu\text{g}/\text{kg}$ for salmon, 0.077 to 0.848 $\mu\text{g}/\text{kg}$ for
346 trout, 0.083 to 1.655 $\mu\text{g}/\text{kg}$ for shrimp, respectively.

347 The recoveries of nitrofurans, nitroimidazoles, dyes and chloramphenicol estimated at
348 three concentrations in aquaculture products ranged 89.8-112.0%, 77.2-104.4%, 83.3-107.6%,
349 and 93.0-125.6%, respectively, except for DNSH at 2.0 $\mu\text{g}/\text{kg}$ and 3.0 $\mu\text{g}/\text{kg}$ in shrimp slightly
350 exceeding the criteria of performance recommended into the Decision 2002/657 to be ranging
351 [-50% - +20%]. The RSD were less than 24.7%, 13.7%, 12.8%, and 16.1% in aquaculture
352 products. The trueness of nitrofurans, nitroimidazoles, dyes and chloramphenicol, expressed
353 as bias, were from -8.8% to +8.9% except for DNSH at 1.0 $\mu\text{g}/\text{kg}$ and 2.0 $\mu\text{g}/\text{kg}$ with a
354 trueness in the range of 16%-18%. All the data of accuracy and precision are showed in Table

355 1.

356 The specificity of the assay was demonstrated by checking interfering peaks at the
357 retention time of target analytes. The results showed that there were no interference peaks
358 co-eluting with target analytes (Fig. 4). All the chromatograms obtained throughout the
359 validation study showed a very good stability of the retention times for all the analytes with
360 relative deviations always remaining below $\pm 2.5\%$. According to European Union
361 Commission Decision 657/2002 (European Commission 2002) with a minimum total score,
362 one precursor ion and two product ions were monitored. This requirement is fulfilled for the
363 method (Table S1). Each analyte ion ratio was effectively measured on each chromatogram,
364 corresponding to the less intense signal against the most intense one. During the validation,
365 the ion ratios measured on the spiked samples were compared with those obtained from the
366 matrix matched calibration curve standards (Table 2). The calculated ion ratio results from
367 this work were in compliance with the ion ratio tolerance laid down in EU Commission
368 Decision No. (EC) 2002/657.

369 ***3.4 Further method improvement***

370 In this study, nitroimidazoles, nitrofurans, dyes and chloramphenicol in aquaculture
371 products were simultaneously determined by LC-MS/MS. However, some further method
372 development should still be carried out to improve the method's performance. In fact, the
373 analytical limits expressed as $CC\alpha$ were satisfactory according to the current RPAs but
374 excluding for CAP, for which the RPA level of $0.3 \mu\text{g/kg}$ is not reached. This method should
375 be further optimized to increase the sensitivity of the detection for CAP. Furthermore, the
376 matrices used to validate the method only included trout, salmon, and shrimp tissues. This

377 method can be more extensively developed to determine residues in other aquaculture
378 products. Finally the pre-treatment of samples is still a crucial step before LC-MS/MS
379 detection. We believe that fast and friendly-environmental techniques will be highly
380 concerned in the future.

381

382 **4. Conclusions**

383 A new LC-MS/MS confirmatory method was tentatively developed to match with the
384 relevant challenge of identifying and quantifying simultaneously the major prohibited
385 veterinary substances in aquaculture products, ie. nitroimidazoles, nitrofurans, dyes, and
386 chloramphenicol. The sample preparation procedure included acid hydrolysis and
387 derivatization steps as to favor the detection of the nitrofuran metabolites. After optimization,
388 the derivatization time was decreased compared to those of published articles dedicated to
389 NFs. The extraction procedure was also simplified by using di-potassium hydrogen phosphate
390 to adjust the pH to 7.0 ± 0.5 . It was the first time that a modified QuEChERS method was used
391 to extract these four family compounds in aquaculture products before analysis. The results
392 indicated that the sample preparation method was able to extract and clean up the residues of
393 nitroimidazoles, nitrofurans, dyes, and chloramphenicol in aquaculture products in a quick
394 and cheap way.

395 The simultaneous detection and quantification of 21 target compounds, ie. 8
396 nitrofurans, 7 nitroimidazoles, 5 dyes, and chloramphenicol were performed by a
397 reversed-phase liquid chromatography coupled with LC-MS/MS. To the authors' knowledge,
398 it is the first time to separate these four family compounds using one single liquid phase

399 separative system. The method demonstrates satisfactory validation characteristics with
400 respect to specificity, trueness, precision, and sensitivity for all of the target compounds
401 excepting a lightly reduced performance in trueness for DNSH and the fact that CAP cannot
402 be detected down to the EU reference point for action “RPA” level currently set at 0.3 µg/kg.
403 It will potentially be a useful tool for accurately monitoring the residues of nitroimidazoles,
404 nitrofurans, dyes, and chloramphenicol in aquaculture products and protecting consumer
405 health.

406

407 **Conflict of interest**

408 The authors declare that they have no known competing financial interests or personal
409 relationships that could have appeared to influence the work reported in this paper.

410

411 **Acknowledgement**

412 The authors thank the research projects National Natural Science Foundation of
413 China (NSFC, 31572570) for the financial support, and the European Commission
414 Directorate-General for Health and Food Safety (European contribution to the European
415 Union Reference Laboratory SI2.726842 & SI2.777451), which enabled this work to be
416 carried out.

417 The authors would like to thank Professor Zonghui Yuan, Dr. Pascal Sanders, Sophie
418 Mompelat for their effort and contribution to this project.

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530

531 **Figure legends**

532 Fig. 1 Comparative results for different derivatized temperatures (A) and time (B).

533 Fig. 2 The effect of extraction solvent on recoveries of analytes fortified in trout samples.

534 Fig. 3 The effect of extraction solvent recovery on injection in LC-MS/MS

535 Fig. 4 LC-MS/MS chromatograms of multi-banned substances in salmon samples.

536 (0.5 μ g/kg for Nitrofurans; 1.0 μ g/kg for Dyes; 1.5 μ g/kg for Nitroimidazoles, and 3.0 μ g/kg
537 for CAP).

538

539

540 **Tables**

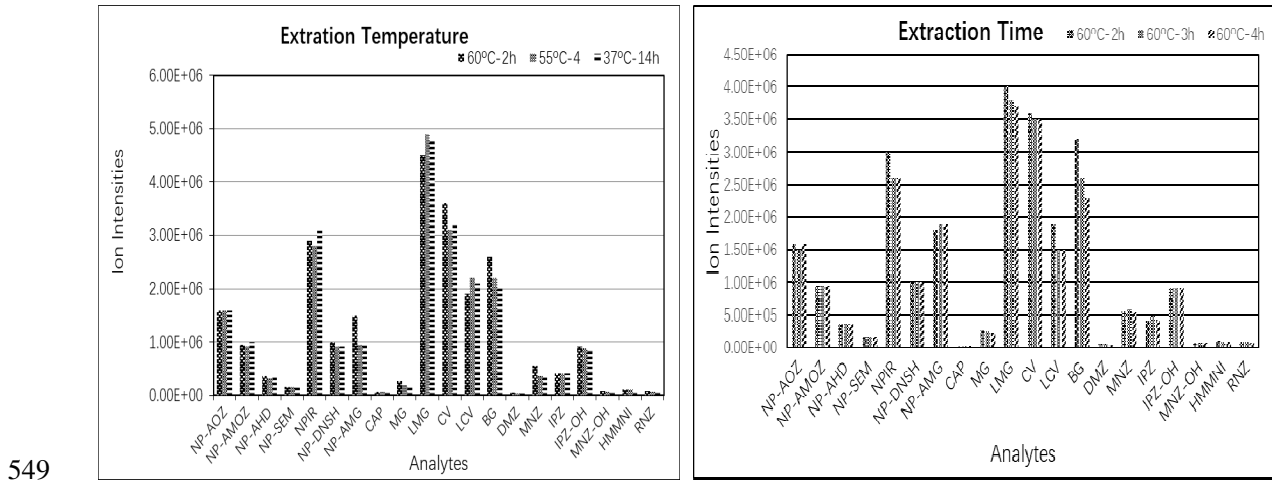
541 Table 1 The accuracy, expressed as recovery for each salmon, trout and shrimp matrix and
542 expressed globally in terms of trueness (bias %) and intra-lab inter-series inter-day
543 reproducibility

544 Table 2 Relative ion abundancy of analytes in matrix-matched standard solutions and in
545 matrix-spiked salmon.

546

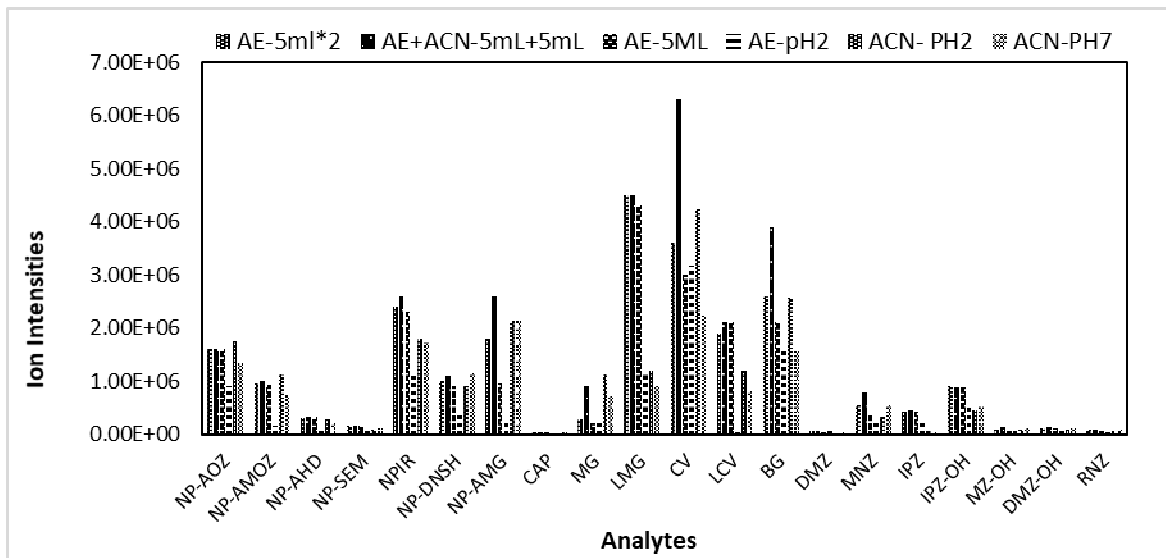
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548 **Figures**



550 Fig. 1 Comparative results for different derivatized temperatures (A) and time (B).

551

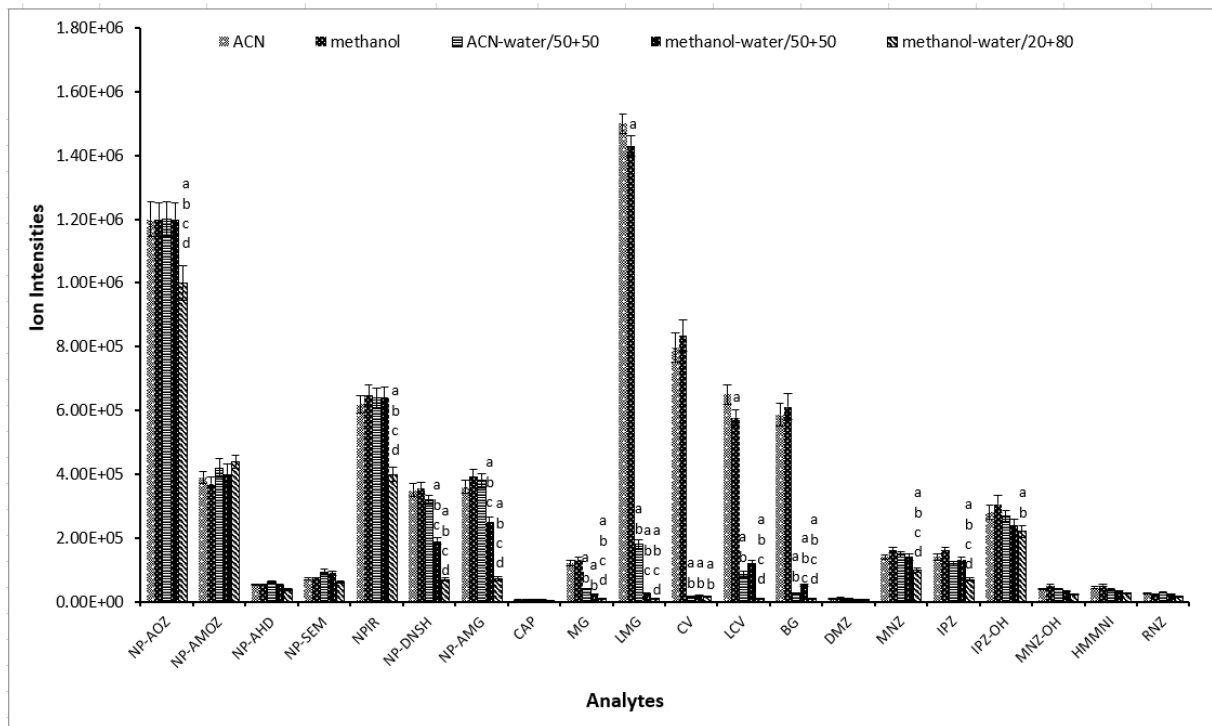


552

553 Fig. 2 The effect of extraction solvent on recoveries of analytes fortified in trout samples.

554

555



556

557 a Statistical significances compared with ACN are $p < 0.05$.

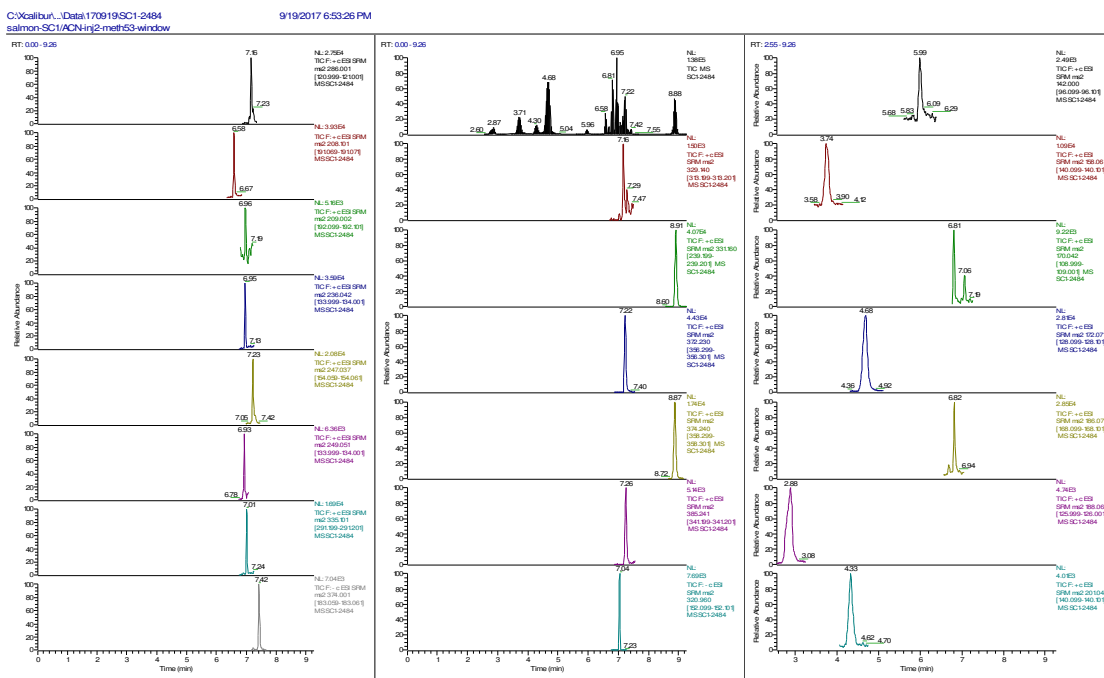
558 b Statistical significances compared with methanol are $p < 0.05$.

559 c Statistical significances compared with ACN-water/50+50 are $p < 0.05$.

560 d Statistical significances compared with methanol-water/50+50 are $p < 0.05$.

561 Fig. 3 The effect of extraction solvent recovery on injection in LC-MS/MS

562



563

564 Fig. 4 LC-MS/MS chromatograms of multi-banded substances in salmon samples.

565 (0.5µg/kg for Nitrofurans; 1.0µg/kg for Dyes; 1.5µg/kg for Nitroimidazoles, and 3.0 µg/kg
566 for CAP).

567 **Tables**568 Table 1 The accuracy, expressed as recovery for each salmon, trout and shrimp matrix and expressed globally in terms of trueness (bias %) and
569 intra-lab inter-series inter-day reproducibility

| Compound | Spiked Concentration (µg/kg) | Salmon | | Trout | | Shrimp | | Trueness (%) (n=18) | Reproducibility inter-series (%) (n=18) |
|----------|------------------------------------|---|-----------------|---|-----------------|---|-----------------|---------------------------|---|
| | | Estimated Concentration (µg/kg) (n=6) | Recovery (%) | Estimated Concentration (µg/kg) (n=6) | Recovery (%) | Estimated Concentration (µg/kg) (n=6) | Recovery (%) | | |
| NP-AHD | 0.5 | 0.562±0.023 | 112.4 | 0.485±0.060 | 97.0 | 0.494±0.038 | 98.8 | 2.8 | 10.5 |
| | 1.0 | 1.041±0.099 | 104.1 | 1.109±0.088 | 110.9 | 1.014±0.133 | 101.4 | 5.5 | 10.4 |
| | 2.0 | 1.920±0.115 | 96.0 | 1.903±0.088 | 95.2 | 1.929±0.154 | 96.5 | 4.1 | 6.0 |
| NP-AOZ | 0.5 | 0.484±0.008 | 96.8 | 0.487±0.013 | 97.4 | 0.474±0.030 | 94.8 | 0.8 | 6.5 |
| | 1.0 | 0.985±0.036 | 98.5 | 1.003±0.061 | 100.3 | 0.861±0.0690 | 86.1 | -2.9 | 5.2 |
| | 2.0 | 1.905±0.097 | 95.3 | 1.951±0.054 | 97.6 | 2.050±0.069 | 102.5 | 2.4 | 5.8 |
| NP-SEM | 0.5 | 0.467±0.051 | 93.4 | 0.466±0.074 | 93.2 | 0.498±0.058 | 99.6 | -4.5 | 12.6 |
| | 1.0 | 1.043±0.088 | 104.3 | 0.898±0.084 | 89.8 | 0.935±0.145 | 93.5 | -4.1 | 9.9 |
| | 2.0 | 1.977±0.109 | 98.9 | 1.966±0.181 | 98.3 | 2.014±0.134 | 100.7 | 0.7 | 6.9 |
| NP-AMTZ | 0.5 | 0.511±0.157 | 102.2 | 0.501±0.574 | 100.2 | 0.500±0.006 | 100.0 | 0.8 | 6.5 |
| | 1.0 | 0.976±0.038 | 97.6 | 0.985±0.058 | 98.5 | 0.950±0.054 | 95.0 | -2.9 | 5.2 |
| | 2.0 | 1.931±0.089 | 96.6 | 2.051±0.113 | 102.6 | 1.877±0.066 | 93.9 | -2.4 | 5.8 |
| NP-AMG | 0.5 | 0.516±0.008 | 103.2 | 0.484±0.037 | 96.8 | 0.496±0.022 | 99.2 | -0.3 | 5.3 |
| | 1.0 | 1.015±0.064 | 101.5 | 0.946±0.072 | 94.6 | 1.061±0.087 | 106.1 | 2.2 | 8.5 |
| | 2.0 | 2.020±0.094 | 101.0 | 1.859±0.128 | 93.0 | 1.991±0.146 | 99.6 | 0.7 | 7.0 |
| NP-DNSH | 0.5 | 0.465±0.036 | 93.0 | 0.532±0.043 | 106.4 | 0.560±0.115 | 112.0 | 3.8 | 15.6 |
| | 1.0 | 1.094±0.043 | 109.4 | 1.000±0.081 | 100.0 | 1.409±0.280 | 140.9 | 16.7 | 20.7 |
| | 2.0 | 2.002±0.24 | 100.1 | 1.920±0.067 | 96.0 | 2.865±0.482 | 143.3 | 18.1 | 24.7 |
| NP-PSH | 0.5 | 0.471±0.041 | 94.2 | 0.571±0.015 | 114.2 | 0.454±0.216 | 90.8 | -0.3 | 11.9 |
| | 1.0 | 1.081±0.092 | 108.1 | 0.904±0.072 | 90.4 | 1.032±0.089 | 103.2 | 0.5 | 11.0 |

| | | | | | | | | | |
|-------|-----|-------------|-------|-------------|-------|-------------|-------|------|------|
| | 2.0 | 1.958±0.167 | 97.9 | 1.885±0.122 | 94.3 | 2.095±0.109 | 104.8 | 1.0 | 7.9 |
| NPIR | 0.5 | 0.500±0.038 | 100.0 | 0.527±0.023 | 105.4 | 0.452±0.029 | 90.4 | -1.4 | 8.8 |
| | 1.0 | 1.036±0.042 | 103.6 | 1.050±0.079 | 105.0 | 0.946±0.102 | 94.6 | 1.1 | 8.7 |
| | 2.0 | 1.933±0.038 | 96.7 | 2.103±0.023 | 105.2 | 2.112±0.029 | 105.6 | 2.5 | 7.6 |
| DMZ | 1.5 | 1.457±0.044 | 97.1 | 1.490±0.054 | 99.3 | 1.158±0.185 | 77.2 | -8.8 | 13.7 |
| | 3.0 | 2.948±0.109 | 98.3 | 2.906±0.158 | 96.9 | 2.843±0.225 | 94.8 | -3.4 | 5.7 |
| | 6.0 | 5.71±0.243 | 95.2 | 6.026±0.213 | 100.4 | 5.929±0.247 | 98.8 | 1.9 | 4.4 |
| HMMNI | 1.5 | 1.473±0.049 | 98.2 | 1.533±0.072 | 102.2 | 1.502±0.072 | 100.1 | 0.2 | 4.4 |
| | 3.0 | 2.910±0.049 | 97.0 | 2.895±0.072 | 96.5 | 2.889±0.098 | 96.3 | -3.4 | 2.5 |
| | 6.0 | 5.503±0.136 | 91.7 | 5.848±0.169 | 97.5 | 6.053±0.118 | 100.9 | -3.3 | 4.6 |
| IPZ | 1.5 | 1.191±0.098 | 79.4 | 1.518±0.083 | 101.2 | 1.421±0.180 | 94.7 | -1.5 | 8.6 |
| | 3.0 | 3.035±0.123 | 101.2 | 2.909±0.147 | 97.0 | 3.033±0.219 | 101.1 | -0.3 | 5.7 |
| | 6.0 | 5.980±0.116 | 99.7 | 5.736±0.359 | 95.6 | 6.126±0.204 | 102.1 | -0.9 | 4.8 |
| MNZ | 1.5 | 1.491±0.024 | 99.4 | 1.507±0.029 | 100.5 | 1.499±0.028 | 99.9 | -0.1 | 1.8 |
| | 3.0 | 2.904±0.072 | 96.8 | 2.911±0.056 | 97.0 | 2.879±0.039 | 96.0 | -3.4 | 1.9 |
| | 6.0 | 5.693±0.078 | 94.9 | 5.855±0.081 | 97.6 | 5.854±0.068 | 97.6 | -3.3 | 1.8 |
| IPZOH | 1.5 | 1.547±0.083 | 103.1 | 1.314±0.036 | 87.6 | 1.475±0.122 | 98.3 | -3.6 | 9.0 |
| | 3.0 | 2.955±0.079 | 98.5 | 2.937±0.109 | 97.9 | 3.042±0.192 | 101.4 | -0.7 | 4.6 |
| | 6.0 | 5.824±0.334 | 97.1 | 5.871±0.109 | 97.9 | 6.021±0.198 | 100.4 | -1.6 | 4.0 |
| MNZOH | 1.5 | 1.515±0.034 | 101.0 | 1.551±0.065 | 103.4 | 1.479±0.024 | 98.6 | 1.0 | 3.4 |
| | 3.0 | 2.874±0.088 | 95.8 | 2.987±0.105 | 99.6 | 2.911±0.064 | 97.0 | -2.5 | 3.3 |
| | 6.0 | 5.832±0.073 | 97.2 | 5.787±0.146 | 96.5 | 5.793±0.082 | 96.6 | -2.3 | 1.8 |
| RNZ | 1.5 | 1.508±0.088 | 100.5 | 1.566±0.047 | 104.4 | 1.555±0.037 | 103.7 | 2.9 | 4.1 |
| | 3.0 | 2.891±0.078 | 96.4 | 3.041±0.114 | 101.4 | 2.984±0.167 | 99.5 | -0.9 | 4.5 |
| | 6.0 | 5.702±0.121 | 95.0 | 6.092±0.129 | 101.5 | 5.995±0.376 | 99.9 | -1.2 | 4.8 |
| MG | 1.0 | 1.004±0.099 | 100.4 | 0.921±0.091 | 92.1 | 1.023±0.172 | 102.3 | -1.7 | 13.0 |
| | 2.0 | 1.897±0.083 | 94.9 | 1.934±0.168 | 96.7 | 2.067±0.205 | 103.4 | -1.7 | 8.6 |
| | 4.0 | 3.977±0.229 | 99.4 | 3.961±0.158 | 99.0 | 4.169±0.235 | 104.2 | -0.9 | 5.5 |
| LMG | 1.0 | 1.029±0.040 | 102.9 | 1.021±0.031 | 102.1 | 0.993±0.029 | 99.3 | 1.4 | 3.5 |

| | | | | | | | | | |
|-----|-----|-------------|-------|-------------|-------|-------------|-------|------|------|
| | 2.0 | 2.042±0.044 | 102.1 | 1.935±0.073 | 96.8 | 2.023±0.018 | 101.2 | 0 | 3.4 |
| | 4.0 | 3.692±0.049 | 92.3 | 3.746±0.075 | 93.7 | 4.024±0.086 | 100.6 | 4.5 | 4.3 |
| CV | 1.0 | 1.011±0.034 | 101.1 | 0.873±0.030 | 87.3 | 0.967±0.029 | 96.7 | 1.3 | 6.8 |
| | 2.0 | 1.928±0.035 | 96.4 | 1.872±0.065 | 93.6 | 1.881±0.043 | 94.1 | -5.3 | 2.8 |
| | 4.0 | 3.815±0.086 | 95.4 | 3.896±0.087 | 97.4 | 3.840±0.031 | 96.0 | -3.7 | 2.0 |
| LCV | 1.0 | 1.012±0.038 | 101.2 | 1.049±0.040 | 104.9 | 0.958±0.057 | 95.8 | 0.6 | 5.8 |
| | 2.0 | 2.046±0.049 | 102.3 | 1.895±0.074 | 94.8 | 2.032±0.041 | 101.6 | -0.4 | 4.4 |
| | 4.0 | 3.984±0.109 | 99.6 | 3.804±0.104 | 95.1 | 4.034±0.120 | 100.9 | 1.5 | 3.7 |
| BG | 1.0 | 0.993±0.082 | 99.3 | 1.048±0.080 | 104.8 | 1.070±0.114 | 107.0 | 3.7 | 9.1 |
| | 2.0 | 1.939±0.218 | 97.0 | 2.011±0.035 | 100.6 | 2.152±0.104 | 107.6 | 1.7 | 7.1 |
| | 4.0 | 3.332±0.455 | 83.3 | 4.113±0.215 | 102.8 | 4.078±0.298 | 102.0 | -3.8 | 12.8 |
| CAP | 3.0 | 2.790±0.368 | 93.0 | 3.243±0.221 | 108.1 | 3.769±0.421 | 125.6 | 8.9 | 16.1 |
| | 4.5 | 4.652±0.503 | 103.4 | 4.511±0.225 | 100.2 | 4.895±0.292 | 108.8 | 4.1 | 8.0 |
| | 6.0 | 6.027±0.284 | 100.5 | 5.981±0.312 | 99.7 | 6.456±0.626 | 107.6 | 2.6 | 7.6 |

570 Table 2 Relative ion abundancy of analytes in matrix-matched standard solutions and in
 571 matrix-spiked salmon.

| Toxin | Average of ion ratios of standard solution (%) | Maximum permitted tolerances | Average of ion ratios of matrix-spiked salmon (%) |
|---------|--|------------------------------|---|
| NP-AHD | 45.7 | 34.2%-57.1% ($\pm 25\%$) | 45.4 |
| NP-AOZ | 60.5 | 48.4%-72.6% ($\pm 20\%$) | 60.7 |
| NP-SEM | 70.6 | 56.5%-84.7% ($\pm 20\%$) | 68.3 |
| NP-AMOZ | 34.7 | 26.0%-43.3% ($\pm 25\%$) | 34.8 |
| NP-AMG | 30.3 | 22.7%-37.9% ($\pm 25\%$) | 29.7 |
| NP-DNSH | 64.3 | 51.4%-71.1% ($\pm 20\%$) | 63.8 |
| NP-PSH | 25.6 | 19.2%-32.0% ($\pm 25\%$) | 23.1 |
| NPIR | 91.3 | 73.0%-109.6% ($\pm 20\%$) | 97.2 |
| DMZ | 77.5 | 62.0%-93.0% ($\pm 20\%$) | 78.7 |
| HMMNI | 22.4 | 16.8%-28.0% ($\pm 25\%$) | 22.6 |
| IPZ | 49.8 | 37.3%-62.2% ($\pm 25\%$) | 52.1 |
| MNZ | 43.7 | 32.8%-54.6% ($\pm 25\%$) | 43.0 |
| IPZOH | 53.0 | 42.4%-63.6% ($\pm 20\%$) | 53.8 |
| MNZOH | 110.5 | 88.4%-132.6% ($\pm 20\%$) | 110.2 |
| RNZ | 15.5 | 10.8%-21.1% ($\pm 30\%$) | 16.0 |
| MG | 25.8 | 19.3%-32.2% ($\pm 25\%$) | 27.1 |
| LMG | 37.6 | 28.2%-47.0% ($\pm 25\%$) | 37.7 |
| CV | 40.8 | 30.6%-51.0% ($\pm 25\%$) | 41.1 |
| LCV | 82.3 | 65.8%-98.8% ($\pm 20\%$) | 83.4 |
| BG | 39.8 | 29.8%-49.8% ($\pm 25\%$) | 39.2 |
| CAP | 91.2 | 72.9%-109.4% ($\pm 20\%$) | 96.1 |

572