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1 Multibiomarker biomonitoring approach using three bivalve species in the Ebre Delta

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16 **Abstract**

17 Bivalves have proved to be useful bioindicators for environmental pollution. In the
18 present study, mussels (*Mytilus galloprovincialis*), cockles (*Cerastoderma edule*) and
19 razor shells (*Solen marginatus*) were collected in the Ebre Delta, an extensive area
20 devoted to rice farming and affected by pesticide pollution, from April to July, the
21 heaviest rice-field treatment period. Possible effects of pollution were assessed through
22 biochemical markers (carboxylesterase (CE), antioxidant and neurotoxicity-related
23 enzymes and lipid peroxidation levels). Data on environmental variables, bivalve
24 reproductive condition and presence of organic pollutants, marine phycotoxins,
25 pathogens or histopathological conditions in bivalve's tissues were also evaluated.
26 Although the bioaccumulated pesticides did not explain the patterns observed for
27 biochemical responses, the obtained results point to an effect of environmental pesticide
28 pollution on enzymatic markers, with a prominent contribution of CE to such changes.
29 Mussels and razor shells provided a more sensitive biochemical response to pollution
30 than cockles. Environmental variables, bivalves reproductive condition and marine
31 phycotoxins did not seem to have a relevant effect on the biomarkers assessed.

32
33 **Keywords:** mussel, cockle, razor shell, biomarkers, contaminants, pesticides, histology,
34 phycotoxins

38 **1. Introduction**

39 Estuaries are semi-enclosed coastal areas characterized by high biomass, biodiversity
40 and primary production, which favour the proliferation of aquatic organisms but are also
41 highly exposed to anthropogenic impacts. Since these regions are often devoted to
42 agriculture, pesticides derived from this activity are an important source of pollution
43 that can threaten water quality (Mañosa et al. 2001; Ochoa et al. 2013; Rodrigues et al.
44 2018).

45 Since concentrations of toxic agents in environmental samples do not fully inform on
46 their biological effects in organisms, environmental chemical analyses must be
47 completed with the use of biomarkers defined as sub-individual responses such as
48 molecular, biochemical and physiological responses that occur in exposed organisms
49 and that allow identifying the effects of toxic agents in natural populations. In this
50 sense, bivalves play a prominent role as bioindicators due to their filter-feeding
51 behavior, widespread distribution and easy sampling and have been widely used for
52 ecotoxicological purposes (Farcy et al. 2013; Rodrigues et al. 2018).

53 Alterations in the activity levels of enzymes involved xenobiotic metabolism,
54 neurotoxicity and oxidative stress are biochemical markers known to respond to
55 environmental stress related to chemical exposure, and their combined use is strongly
56 recommended (Capó et al. 2015; Mejdoub et al. 2017; Solé and Sanchez-Hernandez
57 2018). Carboxylesterases (CEs; EC 3.1.1.1) are hydrolases of wide specificity that
58 hydrolyze esterified xenobiotics to the corresponding alcohol and carboxylic acid
59 (Wheelock and Nakagawa 2010) and are inhibited by several different compounds, such
60 as the oxon-forms of organophosphate (OP) insecticides, carbamates or sulfonamides
61 (Wheelock et al. 2005). The enzyme acetylcholinesterase (AChE; EC 3.1.1.7) catalyzes
62 the hydrolysis of the neurotransmitter acetylcholine at cholinergic nerve terminals and is

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63 also inhibited by OP and carbamate pesticides (Kristoff et al. 2010). Exposure to
64 organic toxicants enhances the production of reactive oxygen species (ROS) that can
65 react with important macromolecules, such as DNA, proteins and lipids. Antioxidant
66 enzymes, such as glutathione peroxidase (GPX; EC 1.11.1.9), glutathione reductase
67 (GR; EC 1.8.1.10) and glutathione-S-transferases (GST; EC 4.4.1.20) carry out specific
68 reactions preventing the adverse effects of ROS (Regoli and Giuliani 2014). These
69 enzymes interact in a complex network, and, alongside with lipid peroxidation (LPO),
70 are frequently used as indicators of oxidative stress induced by chemical pollution in
71 bivalves (Matozzo et al. 2018a). A wide range of other effect markers, such as
72 histological alterations, microbiological measurements or pathogenic conditions have
73 been commonly used to reveal signs of altered health status in bivalves in response to
74 pollutants (Farcy et al. 2013; Izagirre et al. 2014; Matozzo et al. 2018b and references
75 therein). Multidisciplinary studies that combine different sets of biomarkers that
76 respond to both natural and anthropogenic stressors (e.g. contaminants) provide a
77 broader perspective and a better understanding of the observed dynamics than more
78 restricted approaches, and are highly recommended in ecotoxicological studies
79 (Cajaraville et al. 2000; Galloway et al. 2004; Matozzo et al. 2018b).
80 Many biomarkers are also known to vary according to environmental (e.g. temperature)
81 and/or biological factors (e.g. nutritive status or reproductive conditions) in bivalves
82 (Moore et al. 2007; Farcy et al. 2013; González-Fernández et al. 2015a, b). The
83 physiologic alterations derived from these factors can cause misinterpretation of
84 biomarker responses, and, consequently, potentially confounding factors should be
85 included in bivalve biomonitoring studies.
86 Mussels have been extensively used in ecotoxicological field studies as bioindicators
87 (Farcy et al. 2013; Mundhe et al. 2016; Mejdoub et al. 2017; Matozzo et al. 2018b; Solé

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88 and Sanchez-Hernandez 2018). In contrast, knowledge on the potential use as
89 bioindicator species of cockles and razor shells in the field is more limited, although a
90 few studies have addressed the use of their enzymatic responses as biomarkers (Jebali et
91 al. 2011; Nilin et al. 2012; Velez et al. 2016; Ferrante et al. 2014; Nunes and Resende
92 2017; Pearce and Mann 2006).

93 From this perspective, the present study aims to provide a multi-biomarker approach
94 with the use of the Mediterranean mussel, the common cockle and the grooved razor
95 shell in an estuarine area devoted to mariculture, under the impact of pesticides and
96 other anthropogenic chemicals, in order to find the most suitable bioindicator. Changes
97 in the levels of different biochemical markers (activity of CE, antioxidant and
98 neurotoxicity-related enzymes, and LPO levels) were assessed under the hypothesis of a
99 response mainly to pesticides derived from agricultural activity, but also to other
100 organic contaminants potentially present in the study area. Additional factors that may
101 influence biomarker responses, such as environmental variables, bivalve reproductive
102 condition and phycotoxins occurrence, or act as markers themselves, such as the
103 presence of pathogens and/or histopathological conditions were also taken into account.

104 **2. Materials and methods**

106 *2.1. Sampling area and specimen collection*

107 The Ebre Delta is an extensive estuarine area (320 km²) located at the mouth of the Ebre
108 River (Catalonia, NE Spain). It constitutes one of the most important aquatic
109 environments in the western Mediterranean and is the most important bivalve and rice
110 producer in the region (Guallar et al. 2016; Mañosa et al., 2001).
111 Mussels (*Mytilus galloprovincialis*), cockles (*Cerastoderma edule*) and razor shells
112 (*Solen marginatus*) (approximate shell size range (cm): 4.5–7.0, 1.9–3.3 and 8.3–11.5,

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113 respectively) were collected in April, May, June and July of 2017 from the Alfacs Bay
114 of the Ebre Delta. Mussels were collected at 0.5 m depth from suspended cords located
115 at the central part of the northern margin of the bay (40° 37.24'N, 0° 39.22'E), and
116 cockles and razor shells were sampled by traditional techniques from the northern (40°
117 37.58'N, 0° 39.66'E) and southern (40°36.17'N, 0°40.18'E) margins of the bay at
118 subtidal areas located at 0.5 and 1 m depth, respectively (Fig. 1). The number of
119 specimens collected of each species and for each analysis is specified in the
120 corresponding following sections.

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122 *2.2. Environmental variables*

123 Records for temperature (in °C), salinity (in S_p), dissolved O₂ concentration (in mg/L)
124 and chlorophyll-a concentration (in µg/L) were taken at the northern margin of the
125 Alfacs bay for mussels and cockles (40° 37.33'N, 0° 39.49'E) and at the southern
126 margin for razor shells (40° 36.45'N, 0° 39.37'E) using an YSI multiparameter sounder
127 (Fig. 1). Each parameter was measured in triplicate at a weekly frequency: on the
128 sampling day and the two previous weeks. Data provided in Table 1 are the result of the
129 mean of these three measurements.

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131 *2.3. Analysis of organic contaminants*

132 A multi-residue analytical method (Álvarez-Muñoz et al. en prep) was used for the
133 extraction and quantification of a mixture of contaminants including pesticides,
134 endocrine disruptors (EDCs) and pharmaceuticals (PhACs) in bivalves' soft tissues. A
135 pool containing between 5 and 15 specimens was prepared for each species (i.e. *M.*
136 *galloprovincialis*, *C. edule* and *S. marginatus*) and sampling period (i.e. April, May,
137 June and July). Each pool was homogenized using a grinder and freeze-dried prior to

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138 the analysis. Approximately 1 g of sample was extracted per duplicate for each pool
139 using QuEChERS (Bekolut® Citrat-Kit-01) with acetonitrile in aqueous condition
140 followed by the addition of 4 g MgSO₄, 1 g NaCl, 1 g NaCitrate and 0.5 g disodium
141 citrate sesquihydrate. Purification was carried out by means of dispersive Solid Phase
142 Extraction (dSPE) using a sorbent mixture of 400 mg PSA and 400 mg C18 (Bekolut®
143 PSA-Kit-04A). Samples were passed through a phospholipid removal plate (Ostro™
144 Pass-through sample preparation, from Waters) prior injection in Ultra-High
145 Performance Liquid Chromatography-High Resolution Mass Spectrometry (UHPLC-
146 HRMS) Orbitrap Q Exactive™ (Thermo Fischer Scientific, San Jose, CA, USA) for the
147 identification and quantification of the target compounds.

150 *2.4. Biochemical determinations*

151 A total of 40 specimens of *M. galloprovincialis*, 80 of *C. edule* and 40 of *S. marginatus*
152 (mean shell length 5.54 ± 0.54 , 2.72 ± 0.36 and 10.27 ± 0.9 cm, respectively) were used
153 for biochemical determinations. Ten individual samples were considered for each
154 sampled month in the case of mussels and razor shells. In the case of cockles, due to
155 their small size, two specimens were included in each sample.
156 In the case of mussels, haemolymph was extracted from the adduct muscle immediately
157 upon sampling, using a 1 mL syringe with a 0.21 gauge needle. Haemolymph was
158 frozen at -80 °C and centrifuged ($5,000$ g \times 5 min at 4 °C) just before analyses to obtain
159 a cell-free supernatant. Mussel gills and digestive glands of the three species were
160 carefully dissected avoiding contamination by other tissues and immediately frozen in
161 liquid nitrogen and stored at -80 °C until further analyses.

162 As demonstrated by Solé and Sanchez-Hernandez (2018), the digestive gland is the
163 most suitable tissue for measuring changes in CE activity in mussels. Therefore, and
164 considering the special focus on CEs in the present study, this tissue was chosen as
165 target for comparing biochemical results among the three species.

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167 *2.4.1. Tissue preparation*

168 Tissues were homogenised (1:5, w/v) in ice-cold homogenisation buffer using a
169 Polytron® blender. In the case of gills, homogenization was carried out in a phosphate
170 buffer (50 mM, pH 7.4) containing 1 mM ethylenediaminetetraacetic acid (EDTA),
171 while for digestive glands a phosphate buffer (100 mM, pH 7.4) containing 150 mM
172 KCl, 1 mM EDTA, and 1mM dithiothreitol (DTT) was used. The resulting homogenates
173 were centrifuged at 10,000 g for 30 min at 4 °C, and the post-mitochondrial supernatants
174 were used for the enzymatic determinations (S10).

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176 *2.4.2. Enzymatic assays*

177 Enzymatic activities quantified in digestive glands of the three species and in mussel
178 gills were: carboxylesterase (CE) (using the commercial colorimetric substrates p-
179 nitrophenyl acetate (pNPA), p-nitrophenyl butyrate (pNPB), 1-naphthyl acetate (1-NA)
180 and 1-naphthyl butyrate (1-NB)), glutathione reductase (GR), glutathione peroxidase
181 (GPX) and glutathione S-transferase (GST). The use of an array of substrates for
182 assessing CE activity has been recommended, since a variety of CE isozymes, with
183 dissimilar ability to hydrolyse different substrates can be found in the same tissue
184 (Wheelock et al. 2005). CE activity was also assayed in mussel haemolymph using the
185 substrate 1-NA, the only one for which CE shows high activity in this tissue (Solé and
186 Sanchez-Hernandez 2018). Acetylcholinesterase (AChE) activity was also determined

187 in mussel haemolymph and gills, where it shows higher activity than in the digestive
188 gland. Lipid peroxidation (LPO) levels, as indicator of oxidative stress damage, were
189 also quantified in mussel digestive gland and gills.
190 CE and AChE activity determinations were carried out as described by Solé et al.
191 (2018a). Sample volumes were: 25 µl for CE (further S10 dilutions for substrates pNPA
192 and pNPB were 1:2 for mussel gills, 1:10 for mussel and cockle digestive gland and
193 1:20 for razor shell digestive gland; for substrates 1-NA and 1-NB, 1:5 for mussel gills,
194 1:20 for mussel and cockle digestive gland, 1:40 for razor shell digestive gland and
195 undiluted haemolymph for mussel and 25 µl of undiluted sample for AChE
196 determinations. Regarding oxidative-stress-related determinations, GR, GPX and GST
197 activities were determined as described in Koenig and Solé (2012). Sample volumes
198 were: 20 µl for GR (except for 10 µl in razor shell digestive gland), 10 µl for GPX
199 (undiluted) and 25 µl for GST (1:10 in all cases). LPO assay in mussel gills and
200 digestive gland was performed as described in Dallarés et al. (2016).
201 Assay conditions were kept similar and only the sample volume adjusted in order to
202 maintain linearity in the enzymatic measurements. All assays were carried out in
203 triplicate at 25°C in 96-wellplates using a TECAN Infinite M200 microplate reader and
204 blanks (sample free) accompanied the sample batches to correct for non-enzymatic
205 reactivity of the substrates. Enzymatic activities are expressed as nmol/min/mg protein.
206 LPO levels are expressed in nmol MDA (malondialdehyde)/g wet weight. Total protein
207 content was determined by the Bradford method (Bradford 1976) adapted to microplate
208 and using the Bradford Bio-Rad Protein Assay reagent and bovine serum albumin
209 (BSA) as standard (0.05–1 mg/mL). Absorbance was read at 595 nm.

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211 *2.5. Histological assessment*

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212 A total of 40 specimens of *M. galloprovincialis*, 40 *C. edule* and 40 *S. marginatus* (ten
213 specimens for each sampled month, mean shell length 5.18 ± 0.62 , 2.23 ± 0.21 and 9.51
214 ± 2.27 cm, respectively) were used for histological purposes. After dissection, a 5 mm
215 section of each individual containing all main organs was fixed in Davidson's fixative
216 (composition: 10% glycerine, 20% formalin, 30% ethanol (95%), 30% seawater and
217 10% glacial acetic acid) during 24–48h for further histological processing. The rest of
218 the body was conserved in 96% ethanol for further potential molecular assays. After
219 fixation in Davidson's solution, tissues were embedded in paraffin, sectioned at 3 μm ,
220 mounted on slides, stained with Haematoxylin and Eosin and examined under an
221 Optech Biostar B5ICS light microscope.

222 The presence of pathogens, the condition of the different tissues and gonadal
223 development were also evaluated.

224 225 *2.6. Microbiological and marine phycotoxins analysis*

226 A minimum of 10 specimens of *M. galloprovincialis*, *C. edule* and *S. marginatus* were
227 used for these analyses in order to obtain 100 g of homogenate tissue per species.

228 The presence of *Escherichia coli* was assessed in bivalves' tissues following the EU
229 reference method BMS is ISO 16449-3. The procedure was based on the most probable
230 number (MPN) method divided in two stages. The first stage consists of a five-tube
231 three dilution with mineral modified glutamate broth (MMGB) inoculated with dilutions
232 of shellfish homogenates (incubation $37^{\circ}\text{C} \pm 1^{\circ}\text{C}$ for 24h \pm 2h). The presence of *E.*
233 *coli* was confirmed by subculturing acid producing and color change tubes in tryptone
234 bile x-glucuronide medium (TBX) agar. The presence of blue-green colonies is positive
235 for *E. coli* positive β -glucuronidase (incubation $44^{\circ}\text{C} \pm 1^{\circ}\text{C}$ for 21h \pm 3h).

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236 Lipophilic marine toxins were analysed by LC-MS/MS analysis according to the EU-
237 Harmonised Standard Operating Procedure (SOP) procedure (ver. 5, 2015). Samples
238 were analyzed under alkaline elution conditions (Garcia-Altarets et al. 2013). Briefly, an
239 Agilent 1200 LC (Agilent Inc. Palo Alto, CA) coupled to a 3200 QTRAP mass
240 spectrometer (AB Sciex, Concord, ON, Canada) was used. Analytical separation was
241 performed on a X-Bridge C8 column (2.1 × 50 mm, 3.5 µm) protected with a pre-
242 column (2.1 × 10 mm, 3.5 µm) from Waters (Milford, MA, USA). A binary gradient
243 was programmed with water (mobile phase A) and acetonitrile/water (mobile phase B)
244 both containing 6.7 mM of ammonium hydroxide.

245 Amnesic shellfish poisoning toxins (ASP) were analysed by LC-UV analysis according
246 to the EU-Harmonised SOP procedure for determination of domoic acid in shellfish by
247 RP-HPLC using UV detection (ver. 1, 2008). For LC-UV analyses, an Alliance LC
248 (Waters) was used. Analytical separation was performed on a X-Bridge C18 column
249 (4.6 × 250 mm, 4.6 µm) protected with a pre-column (2.1 × 10 mm, 3.5 µm) from
250 Waters (Milford, MA, USA). A mobile phase of acetonitrile/water (15:85) containing
251 0.1% formic acid was used. All runs were carried out at 40 °C using a flow rate of 1.2
252 mL/min. The injection volume was 20 µL and the autosampler was set at 4 °C.
253 Detection was performed at 242 nm.

254 Paralytic shellfish poisoning toxins (PSP) were determined by the Mouse Bioassay
255 (MBA) method according to the EURLMB SOP ver.1 (2004). Briefly, acetone
256 extraction of the whole flesh or the hepatopancreas of molluscs was followed by
257 evaporation and resuspension of the residue in a 1% solution of Tween 60 surfactant.
258 One milliliter aliquots of the extract were ip injected into three male mice and observed
259 for 24h. The death of two of the three mice within 24h was interpreted as a positive

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260 result. On the contrary, if none or only one of the mice died within this time, the test
261 was considered to be negative.

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263 *2.7. Data analyses*

264 General Liner Models (GLM) were applied to test the null hypothesis of no differences
265 among the four sampling months for each enzymatic activity quantified, setting the
266 variable month as factor, and post-hoc S-N-K analyses. Prior to these analyses, data of
267 some enzymes were logarithmically or square-root-transformed to meet both normality
268 and homoscedasticity.

269 Permutation multivariate analyses (PERMANOVA) were also performed considering
270 individual samples as replicates to test the null hypothesis of no differences in the
271 enzymatic pool composition among the four sampling months for the three bivalve
272 species addressed. The analyses were carried out using PERMANOVA+ for PRIMER
273 v6 (Anderson et al. 2008) on a Bray-Curtis similarity matrix derived from
274 untransformed data. Permutation p-values were obtained under unrestricted permutation
275 of raw data (9,999 permutations). A similarity percentages analysis (SIMPER) was
276 carried out using individual samples as replicates to identify the enzymatic activities
277 that contributed most to the similarity/dissimilarity of individual samples within/among
278 the samples of the four months sampled. Moreover, with the aim of visualizing patterns
279 of dissimilarity in the enzymatic pool of the three bivalve species across the four
280 months sampled, factorial correspondence analyses (FCA) were performed using
281 STATISTICA v7 (StatSoft, Inc. 2004) on data matrices containing enzymatic data of
282 each species. Hierarchical cluster analyses (Bray-Curtis similarity, average grouping
283 method) were simultaneously performed based on the coordinates of the first two axes
284 obtained in the corresponding FCA to identify month-related groups clearly. The

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285 previous multivariate analyses were not applied to enzymatic data of mussel
286 haemolymph, due to the low number of biochemical markers assessed. In order to make
287 the enzymatic pool activity patterns of the three bivalve species comparable, data of
288 LPO levels in mussels were omitted. Finally, Spearman rank correlations were used to
289 test the null hypothesis of no association among CE substrates and antioxidant enzymes
290 within the tissues of the three bivalve species.

291

292 **3. Results**

293 *3.1 Environmental variables*

294 Most environmental variables measured presented temporal variation throughout the
295 sampling period (Table 1). Water temperature markedly increased from April to July in
296 the northern and southern margins, this trend being more marked in the latter locality.
297 While salinity increased by nearly two points in the northern margin, it decreased over
298 one point on the southern locality. Oxygen concentration did not show a clear trend in
299 either locality. Finally, while chlorophyll-a concentration showed a marked decrease in
300 the northern margin, it remained fairly stable in the southern one.

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302 *3.2. Contaminants concentration*

303 Mean concentration, expressed in ng/g dry weight (dw) \pm relative standard deviation
304 (n=2 replicates), of contaminants quantified in bivalves' soft tissues across the four
305 sampled months are shown in Table 2. The following target chemicals were below the
306 method detection limit in all cases and are thus not shown in the table: the organitrogen
307 pesticides metolachlor, simazine and deethylatrazine, the organophosphorus pesticide
308 malathion, the herbicides bentazone, MCPA and propanil, the insecticides acetamiprid
309 and imidacloprid, the endocrine disruptors bisphenol A, triclosan and triclocarban and

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310 the pharmaceuticals sulfamethoxazole, venlafaxine and cabamazepine. Three of the
311 quantified chemicals were pesticides, namely atrazine, thiabendazole and diazinon, with
312 levels ranging from below method quantification limit (MQL) to 14 ng/g dw of atrazine
313 in razor shells from June (supplementary material, Table 1). The other five
314 contaminants detected were compounds considered endocrine disruptors such as
315 caffeine, methylparaben, ethylparaben, propylparaben and 1-H-benzotriazole. The
316 levels found ranged from below MQL up to 51 ng/g dw of caffeine measured in razor
317 shells in July (supplementary material, Table 1). Actually, caffeine was the contaminant
318 presenting the highest concentrations in the three bivalves species analyzed. The mean
319 levels of the majority of the contaminants measured were quite stable across months,
320 only atrazine measured in razor shell showed an increasing trend from April to July
321 (Table 2).

322 3.3. Biochemical determinations

324 Activity levels of the different enzymes assayed in the three bivalve species, as well as
325 LPO levels, are illustrated in Figs. 2–5.
326 In the case of mussels digestive gland, activity of CE progressively decreased with time,
327 showing significantly lower activity values in July samples than in the other three
328 months for the four substrates analysed (GLM, $F_{(3, 36)}=14.858$, $p<0.001$ for 1-NA; $F_{(3, 36)}=6.587$, $p=0.001$ for 1-NB; $F_{(3, 36)}=12.424$, $p<0.001$ for pNPA and $F_{(3, 36)}=3.939$,
329 $p=0.016$ for pNPB) (Figs. 2A–D). The same trend was observed in cockles (GLM, $F_{(3, 36)}=7.710$, $p<0.001$ for 1-NA; $F_{(3, 36)}=5.763$, $p=0.003$) for pNPA and $F_{(3, 36)}=12.923$,
330 $p<0.001$ for pNPB), with the exception of 1-NB, which displayed the opposite pattern,
331 although without showing significant differences among months (Figs. 2A–D). A
332 similar decreasing trend was found in razor shells, although significantly lower

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335 activities were observed in April for the substrates 1-NA and pNPA (GLM, $F_{(3, 36)}$,
336 $F_{(3, 36)}=4.190$, $p=0.012$ and $F_{(3, 36)}=4.709$, $p=0.007$, respectively) (Figs 2A–D).
337 A higher variability was observed among the responses of antioxidant enzymes: GR
338 activity values significantly decreased from April (mussels) and May (razor shells) to
339 July samplings (GLM, $F_{(3, 36)}=11.944$, $p<0.001$ and $F_{(3, 35)}=6.577$, $p=0.001$,
340 respectively), while a progressive increase in the activity of GPX and GST from April
341 to July was observed in razor shells (GLM, $F_{(3, 36)}=10.305$, $p<0.001$ and $F_{(3, 35)}=6.139$,
342 $p=0.002$, respectively) (Figs. 3A–C). No clear trends were detected for these enzymes in
343 the case of cockles (GLM, $p>0.05$).
344 Regarding enzymatic determinations in mussel gills, significant decreasing trends
345 through time were found for CE activity with the substrates 1-NA, pNPA and pNPB
346 (GLM, $F_{(3, 36)}=10.122$, $p<0.001$; $F_{(3, 36)}=11.128$, $p<0.001$ and $F_{(3, 36)}=4.359$, $p=0.01$,
347 respectively) (Figs. 4A–C), as also for GPX and GST activities (GLM, $F_{(3, 36)}=3.005$,
348 $p=0.043$ and $F_{(3, 36)}=11.663$, $p<0.001$, respectively) (Figs. 5B, C). In the case of the
349 enzymes tested in mussel haemolymph (i.e. CE with substrate 1-NA and AChE), no
350 significant differences among months were detected in any case (GLM, $p>0.05$).
351 The permutational multivariate analyses (PERMANOVA) applied to individual samples
352 showed a significant effect of the factor month in the enzymatic pool of the digestive
353 gland of the three bivalve species (Pseudo- $F_{(3, 36)} = 6.3068$, $p_{(perm)} = 0.0003$ for mussels,
354 Pseudo- $F_{(3, 36)} = 5.0565$, $p_{(perm)} = 0.0003$ for cockles and Pseudo- $F_{(3, 35)} = 4.2551$, $p_{(perm)}$
355 $= 0.0004$ for razor shells), and of the gills of mussels (Pseudo- $F_{(3, 36)} = 9.2979$, $p_{(perm)} =$
356 0.0001). Figure 6 shows the resulting dendrograms of the cluster analyses
357 simultaneously performed to the FCAs, which show the patterns of similarity among the
358 enzymatic pools of the four sampled months in the three bivalve species. In digestive
359 gland of mussels and razor shells (Fig. 6A, D), April, May and June clustered together

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360 while July remained as a separate clade. For mussel gills (Fig. 6B), the two earliest
361 sampling were separated from the two latter ones. In the case of cockles (Fig. 6C),
362 enzymatic pools of April and June were most related while May and July formed
363 independent clades. The similarity percentages analysis (SIMPER) allowed the
364 identification, for each bivalve species, of the enzymatic activities that contributed most
365 to the similarity/dissimilarity of the enzymatic pools within/between sampled months.
366 These were: CE with all substrates assayed (i.e. pNPA, pNPB, 1-NA and 1-NB) and
367 GST in the case of mussels, cockles and razor shells digestive gland and CE with
368 substrates 1-NB and pNPB, GST and GR for mussel gills (Supplementary material,
369 Table 2).
370 Significant positive correlations were found among all CE substrates in mussel digestive
371 gland ($r_s=0.700-0.915$, $p<0.01$) and gills ($r_s=0.663-0.865$, $p<0.01$) and in razor shells
372 digestive gland ($r_s=0.460-0.832$, $p<0.01$). In the case of cockles digestive gland,
373 significant positive associations were detected among the CE substrates pNPA, pNPB
374 and 1-NA ($r_s=0.755-0.818$, $p<0.01$). In the case of antioxidant enzymes, significant
375 positive correlations among GR, GPX and GST were detected in mussels digestive
376 gland ($r_s=0.377-0.503$, $p<0.05$), between GR and GPX and between GST and GPX in
377 mussel gills ($r_s=0.339-0.439$, $p<0.05$), between GST and GR in cockles digestive gland
378 ($r_s=0.534$, $p<0.05$) and between GST and GPX in razor shells digestive gland ($r_s=0.733$,
379 $p<0.05$).

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381 *3.4. Histology*

382 The histological study revealed that the three bivalve species were in maturation and
383 spawning reproductive phases during the sampling period. In the case of mussels, late

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384 spawning and gonadal reabsorption phases could be also observed in May and June,
385 while non observable gonads were found in July.
386 The most relevant histopathological observations in mussels were unspecific lesions of
387 hypertrophic nuclei with peripheral chromatin in the digestive gland in May, June and
388 July. Moreover, one sample was infected with moderate levels of the protozoan
389 *Marteilia* sp. In the case of cockles, *Rickettsia*-like organisms (RLOs) were observed in
390 gills in all samplings (Fig. 7), as well as ciliates of the genus *Trichodina* in the stomach
391 and intestinal lumens in June. In regard to razor shells, RLOs were also detected in all
392 samplings, in gills and the digestive system.

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394 3.5. Microbiological and phycotoxins analysis

395 Low levels of *E. coli* were found in tissues of mussels and razor shells (20–45 and 18–
396 78 MPN/100 g, respectively) through the different samplings. In contrast, much higher
397 numbers of these bacteria were detected in cockles, increasing in number from 790
398 MPN/100 g in April, to 9200 MPN/100 g in July.
399 Levels of hydrophilic toxins (comprising PSP, paralytic shellfish poisoning and ASP,
400 amnesic shellfish poisoning toxins) were low and did not reach the maximum permitted
401 levels (MPLs) in edible shellfish tissues for human consumption according to the
402 European Union (Regulation EC 853/2004: 20 mg domoic acid/kg (ASP) and 800 µg eq
403 STX/kg (PSP)). Regarding lipophilic toxins, only two phycotoxins were detected:
404 traces of yessotoxin in mussel samples in May, and pinnatoxin-G at low concentrations
405 in mussel samples of the four months (range 3.2–6 µg/kg) and in razor shell samples in
406 late June (2.5 µg/kg). Pinnatoxin-G levels in mussel samples were higher in late June
407 than in the other three sampling dates. Levels detected for lipophilic toxins did not reach
408 the MPLs according to the EU (Regulation EC 853/2004 and Regulation EC 786/2013:

1 409 160 µg/kg okadaic acid (OA), equivalents for OA, dinophysistoxins (DTXs) and
2 410 pectenotoxins (PTXs) together; 3.75 mg/kg for yessotoxins (YTXs) and 160 µg/kg for
3
4 411 azaspiracids (AZAs)). Other lipophilic marine toxins are not yet regulated in the
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7 412 European Union, like cyclic imines mainly comprising spirolides (SPXs),
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9 413 gymnodimines (GYMs) and pinnatoxins (PnTXs).
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11 414 12 13 14 415 **4. Discussion**

15 16 416 *4.1. Biochemical responses and relationship to local contaminants*

17 417 A general and coordinated change of enzymatic markers throughout the
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19 418 sampling period was revealed after multivariate analysis performed on enzymatic data
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21 419 by the PERMANOVA tests and cluster analyses, and after GLMs for independent
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23 420 biochemical markers. The results of the SIMPER analyses highlighted the particular
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25 421 contribution of CE and, to a lesser extent, also GST to this response in the digestive
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27 422 gland of the three studied bivalves and also in mussel gills. However, the
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29 423 bioaccumulated pesticides apparently did not explain the patterns observed for the
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31 424 biochemical responses. Indeed, most of the herbicides and pesticides tested were not
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33 425 detected in any biological sample, and those that did, seemed to be below the threshold
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35 426 limit for altering the activity of the enzymes addressed in the present study, as deduced
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37 427 from the lack of correspondence between their concentrations and the biochemical
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39 428 responses observed (Table 2). Previous studies on the same bay indicated that after the
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41 429 period of pesticide application in the rice fields of the Ebre Delta, which takes place
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43 430 from April to June (Terrado et al. 2007; Köck et al. 2010; Suárez-Serrano et al. 2010),
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45 431 the maximum levels of pesticides in its drainage channels and bays are attained during
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47 432 the spring and summer period (when this study takes place), soon after the field waters
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49 433 are discharged into the bays (Escartín and Porte 1997; Santos et al. 2000). It has already
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434 been reported that the measurement of contaminants concentration in bivalves' tissues
435 could lead to underestimation of the real pollution load in the surrounding waters
436 (Lehotay et al. 1998; Köck et al. 2010). Indeed, these authors detected inexistent or very
437 low levels of contaminants in oyster and mussel tissues despite that the concentrations
438 of these same chemicals were high in water, which agrees with the low levels of
439 atrazine, thiabendazole and diazinon in the present research. The possible effect of
440 environmental contaminants in the studied organisms, even though the former are non-
441 detectable in their tissues, makes biomarkers highly recommended tools usually
442 incorporated in ecotoxicological studies (Farcy et al. 2013; Solé and Sanchez-
443 Hernandez 2018).

444 The presence of other anthropogenic chemicals accumulated in biota was low
445 (few ng/g dw) except for caffeine, with levels much higher than previously reported in
446 the study area and other locations (Dodder et al. 2014; Álvarez-Muñoz et al. 2015). No
447 clear temporal trends were observed for the organic contaminants measured except for
448 caffeine in mussel and razor shells, which increased in the summer months, and for
449 atrazine in razor shells, which increased from April to June. Caffeine has been reported
450 to produce alterations on metabolic activity and oxidative stress biomarkes in bivalves
451 (Cruz et al. 2016). Therefore, this compound might explain part of the patterns observed
452 for biochemical markers (see below), although as far as we are concerned, no reference
453 concentrations have been recorded in bivalves' tissues; all studies addressing effects of
454 caffeine on invertebrates are based on water concentrations. In line with this influx of
455 urban residues during the summer period, it stands out a higher presence of the fecal
456 bacteria *E. coli* detected in cockle samples collected in July, which are in all likelihood
457 explained by the raw sewage discharges of a nearby town during the touristic summer
458 season.

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459 Regarding CE activities, a former study conducted with mussels in the same area
460 and over a longer period of time reported an inhibition of CE activity (1-NA as
461 substrate) in gills and digestive gland after the arrival of pollutants from rice fields to
462 the Alfacs Bay in early summer (Escartín and Porte 1997), similarly to the pattern
463 observed in the present study. Moreover, Solé et al. (2018a) characterized baseline
464 enzymatic activities for CEs in the digestive glands of the same three bivalve species
465 addressed herein and collected from the same sites, and after *in vitro* exposure to the OP
466 metabolite chlorpyrifos oxon, also concluded that CEs are potentially good indicators of
467 pesticide pollution in bivalves. With regard to the tissues addressed, Solé and Sanchez-
468 Hernandez (2018) and Escartín and Porte (1997) reported higher inhibition of CE in
469 mussel gills than in digestive gland when exposed to pollutants in *in vitro* conditions,
470 which waited to be confirmed under real field situations. However, a similar inhibitory
471 trend on CE activity was observed in the two selected tissues in mussels from April-
472 May to July for the substrates 1-NA and pNPA, and higher inhibition in digestive gland
473 than in gills was detected for the longer-chain carbon esters pNPB and 1-NB (Fig. 4A-
474 D), non-confirming these previous expectations, and rather suggesting higher
475 detoxification activity in the former tissue under field conditions. It could be
476 hypothesized that the complexity of field conditions, where mixtures of chemicals
477 occur, could yield a different pattern of CE activity in the selected tissues, with the
478 digestive gland showing a greater participation in detoxification processes than when
479 faced with *in vitro* conditions. Solé and Sanchez-Hernandez (2018) suggested that the
480 substrates pNPB and 1-NB were potentially more suitable for detecting inhibition of CE
481 activity in the field in mussels than the substrates pNPA and 1-NA based on their lower
482 IC50 when exposed *in vitro* to the OP pesticide dichlorvos, but also in response to other
483 pharmaceuticals and personal care products (PPCPs). This is not corroborated by

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484 present field results, since CE seemed more sensitive to inhibition from May to July
485 when using the substrates 1-NA and pNPA both in gills and digestive gland. Although
486 these and other authors (e.g. Otero and Kristoff 2016) have noted that the selection of
487 appropriate substrates is species and tissue-specific, present results indicate that there
488 might be additional variables influencing CE activity even within the same tissue and
489 species. The inhibitory CE pattern observed in mussels was also apparent in *C. edule*
490 and *S. marginatus*, although cockles showed an unexpected increase in activity with
491 time when using the substrate 1-NB, which can explain the lack of correlation between
492 this and the other three substrates. This result contrasts with the outcomes of the study
493 by Solé et al. (2018a), in which all four substrates were significantly correlated in the
494 three bivalves selected, including cockle. This result could point to a different
495 contribution of CE isozymes in cockles compared to mussels and razor shells under
496 conditions of pollution exposure. Razor shells demonstrated less sensitivity regarding
497 the longer-chain carbon esters (i.e. pNPB and 1-NB), apparently due to a higher
498 variability in the data (Fig. 2A–D). The observed substrate-specific variability in CE for
499 the three bivalves addressed highlights the importance of using a battery of substrates
500 for assessing the inhibition of this enzyme by pollutants.

501 The lack of effect on AChE activity in mussel gills and hemolymph supports former
502 observations in bivalves that suggested that CE is a more adequate indicator of OP
503 exposure (Galloway et al. 2002; Wheelock and Nakagawa 2010; Otero and Kristoff
504 2016; Sole et al 2018b).

505 The responses of the antioxidant enzymes assessed were not consistent across
506 the three species addressed. For GPX and GST, clear activation patterns were observed
507 in razor shells (Fig. 3B, C), which point to chemical stress by pollutants and the
508 involvement of these enzymes in the associated detoxification processes. Conversely,

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509 inhibitory patterns were observed for GR in mussel and razor shells digestive glands
510 (Fig. 3A), as already reported by Mundhe et al. (2016) in the presence of the
511 organophosphate pesticide monocrotophos. As regards tissue responsiveness in mussels,
512 GR was inhibited in the summer months in gills and digestive gland (Fig. 5A), although
513 in the former organ this inhibition was not significant ($p>0.05$) due to high variability.
514 By contrast, GPX and GST were more responsive in gills than in digestive gland,
515 although an inhibition in the activity was revealed in July for GPX and in June and July
516 for GST (Figs. 5B, C). The lack of a clear antioxidant response in mussels was
517 concordant with no clear increase of oxidative stress damage measured as enhanced
518 LPO levels, and both suggest that the chemical threat, rather than acting over ROS
519 production, could be more selective towards CE inhibition in this case. Antioxidant
520 responses in bivalves are complex and controversial because their activation can take
521 place at low oxidative-stress conditions, but depletion of antioxidant activities can occur
522 in situations of severe oxidative stress, since antioxidant enzymes can be a target of
523 ROS themselves (Regoli and Giuliani 2014).

524

525 *4.2. Influence of temperature on biochemical markers*

526 Temperature is considered one of the most important confounding factors in
527 bivalve monitoring studies (Farcy et al. 2013), provided that it is known to increase
528 metabolic rate and the production of ROS, to modify catalytic efficiency and influence
529 phytoplankton abundance (Somero 1995; Pörtner 2002), among others. Very few
530 studies have addressed CEs response to temperature variations in animals, and have
531 yielded different conclusions: at higher temperatures Owusu et al. (1994) reported an
532 increase in CE activity in aphids while Escartín and Porte (1997) found no significant
533 change in mussels due to this factor. However, higher susceptibility of CE to pollutants

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534 under increasing temperatures has been validated in aquatic species (Laetz et al. 2014;
535 Freitas et al. 2017) and would suggest that the generalized inhibition on CE activity
536 with time due to pesticide pollution might have been enhanced in the warmer months.
537 Activity levels of AChE are known to increase with higher temperatures (Pfeifer et al.
538 2005), and an increase of its activity across the four samplings would thus be expected
539 in the absence of a pollution effect. The opposite trends observed in the present study,
540 similar to those by Escartín and Porte (1997) in the same area, support the hypothesis of
541 an effect of pesticide pollution on bivalves. Higher temperatures have also been
542 reported to significantly increase the activity of antioxidant enzymes, yielding an
543 oxidative stress-like response in mussels (Hu et al. 2015), which could be explained
544 either by an increased production of ROS or by alterations of enzymatic catalytic
545 efficiency (Somero 1995; Almeida and Mascio 2011). In this respect, a generalized
546 increase in the activity of antioxidant enzymes was not observed, for we believe that the
547 influence of temperature in this case might have either been weak or masked by other
548 processes of greater influence.

549

550 *4.3. Influence of reproductive condition on biochemical markers*

551 Another important confounding factor in the assessment of pollution effects in
552 bivalves is the reproductive condition, which interferes with biomarker responses and is
553 closely related to temperature and nutritive status (Farcy et al. 2013; González-
554 Fernández et al. 2015a, b). Although no specific measurements of nutritive condition
555 were performed in the present study, the low variability in chlorophyll a concentration
556 across samplings suggests a fairly constant supply of food to the bivalves throughout the
557 study. Reproduction represents a critical period with a major influence in gene
558 expression, metabolism and immune function, among others (Farcy et al. 2013;

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559 González-Fernández et al. 2017). All these energetically demanding processes can result
560 in poorer physiological condition, and make coping with stressful events difficult
561 (Berthelin et al. 2000). This is why it is considered a major confounding factor for the
562 interpretation of biomonitoring data, and assessing the effect of xenobiotics during the
563 reproductive phase has been recommended (Farcy et al. 2013). Accordingly, the
564 histological analysis revealed that bivalve populations were at the active reproductive
565 stage during the sampling period. In the case of cockles and razor shells, the uniform
566 reproductive condition across samplings suggests that observed changes in biomarkers
567 should not be driven by reproduction events. In the case of mussels, in which final
568 spawning and resting stages could be observed in June and July, alterations in
569 biomarkers due to the reproductive condition could have occurred during these two
570 months. However, decreasing trends in mussel CE and GR are concordant to those
571 observed for cockles and/or razor shells, and no clear trends were observed for GPX and
572 GST. We believe that reproduction effects on enzymatic activities in this case might
573 have been weak and not clearly identified.

574

575 *4.4. Histopathological analysis*

576 Histological analysis of tissue damage was screened in all sampling groups
577 (species and times) and no significant histological alterations could be associated to
578 pesticide pollution. The frequent detection of RLOs in gills of cockles and razor shells
579 could potentially be associated to some type of contamination. For example, *Rickettsia*
580 infections of gut/digestive tubules of oysters have been found to have a significant
581 correlation with nickel contamination on the American coast (Kim et al. 1998; Morley
582 2010), and have also been found to be significantly higher in deep-sea mussels exposed
583 to petroleum seeps (Powell et al., 1999), although further studies are required to

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584 demonstrate a direct effect to such exposure. The high prevalence and intensity reported
585 for the ciliate *Trichodina* sp. in cockle gills in June is indicative of a great abundance of
586 this ciliate in the environment, in some cases linked to environmental eutrophication
587 (Palm and Dobberstein 1999). Bivalve gill ciliates have been found to be most common
588 at heavily polluted sites on the northeast coast of America (Morley 2010). A unique
589 sample of mussel was infected with *Marteilia* sp. (probably *M. refringens*), a pathogen
590 of obligatory declaration according to the World Organisation for Animal Health (OIE).
591 This parasite has been historically detected in mussels and flat oysters in the Ebro Delta
592 production areas. However, in this case, prevalence seems to be low in an optimal
593 period for the parasite (Carrasco et al. 2008).

594 595 4.5. *Phycotoxins analysis*

596 Recently, marine phycotoxins have been proposed as an additional confounding
597 factor in studies assessing effects of pollution (Farcy et al. 2013). However, they were
598 found at very low concentrations in the present study, for it is unlikely that they have
599 interfered with biomarkers response. Harmful hydrophilic toxins ASP and PSP did not
600 seem to pose a threat to bivalve consumption during the period of study as they were all
601 below the legislation EU threshold.

602 603 **Conclusions**

604 The consistent responses of CE across species and tissues (in mussels) with
605 respect to AChE and antioxidant enzymes suggest that CE activity can be a more
606 sensitive and robust biomarker when evaluating pesticide pollution in bivalves. Among
607 the three bivalve species, mussels provided the most sensitive response regarding CE
608 activity. With respect to oxidative stress, it was better reflected by razor shells

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609 enzymatic responses. In turn, cockles seemed to provide the less sensitive response,
610 both considering CE and antioxidant enzymes. This inter-species dependence of the
611 responses of different enzymatic functional groups points to the use of more than one
612 bioindicator as the best approach to ecotoxicological studies using bivalves.
613 Furthermore, biochemical markers seemed to provide a much more robust and sensitive
614 response than histological ones. None of the confounding factors potentially influencing
615 biomarker responses seemed to be relevant enough to modulate the assessed enzymatic
616 activities in the study area.

617

618 **Conflict of interest**

619 The authors declare that they have no conflict of interest.

620

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Table 1. Physico-chemical water parameters quantified in the two areas sampled throughout the length of the study. T: temperature; S: salinity; O₂: oxygen concentration; Chla: Chlorophyll-a.

Area	Date	T (°C)	S (psu)	O₂ (mg/L)	Chla (µg/L)
Northern margin (mussels and cockles)	April	15.77	36.92	7.36	4.51
	May	17.80	37.57	6.41	4.70
	June	21.60	38.22	5.88	2.98
	July	23.63	38.62	6.41	2.34
Southern margin (razor shells)	April	16.57	36.57	7.21	2.48
	May	18.77	36.60	5.73	2.27
	June	22.90	35.47	6.00	2.29
	July	27.47	35.53	5.84	2.75

Table 2. Mean concentration (ng/g dry weight) \pm relative standard deviation (RSD) (n=2 replicates) of the contaminants quantified in soft tissues of mussels (*Mytilus galloprovincialis*), cockles (*Cerastoderma edule*) and grooved razor shells (*Solen marginatus*) in the four sampling performed in the Alfacs Bay, Ebre Delta. MDL: method detection limit; MQL: method quantification limit.

Family	Compounds	<i>Mytilus galloprovincialis</i>				<i>Cerastoderma edule</i>				<i>Solen marginatus</i>			
		April	May	June	July	April	May	June	July	April	May	June	July
Organitrogen pesticides	Atrazine	1.56 \pm 0.32	1.29 \pm 0.10	<MQL	<MQL	0.82 \pm 0.08	<MDL	<MDL	<MDL	3.04 \pm 0.84	5.15 \pm 1.02	13.63 \pm 2.75	8.44 \pm 0.23
Organophosphorus pesticides	Thiabendazole	<MDL	<MDL	<MDL	<MDL	<MDL	0.93 \pm 0.18	0.43 \pm 0.003	0.60 \pm 0.00	<MDL	<MDL	<MDL	<MDL
	Diazinon	<MDL	<MDL	0.46 \pm 0.00	0.51 \pm 0.00	0.57 \pm 0.00	1.53 \pm 0.02	<MDL	<MDL	<MDL	<MDL	0.56 \pm 0.01	0.55 \pm 0.01
Endocrine Disruptors (EDCs)	Caffeine	<MDL	<MDL	<MQL	11.82 \pm 1.93	22.60 \pm 5.48	33.62 \pm 5.51	<MDL	<MDL	<MQL	<MQL	46.95 \pm 6.29	50.96 \pm 15.12
	Methylparaben	2.69 \pm 0.02	1.01 \pm 0.24	0.54 \pm 0.08	0.83 \pm 0.02	2.00 \pm 0.06	3.54 \pm 0.05	1.01 \pm 0.11	1.13 \pm 0.19	1.91 \pm 0.59	1.71 \pm 0.05	1.19 \pm 0.08	1.03 \pm 0.12
	Ethylparaben	1.85 \pm 0.12	<MDL	<MDL	0.54 \pm 0.06	0.55 \pm 0.02	0.36 \pm 0.03	0.73 \pm 0.04	0.35 \pm 0.02	<MDL	<MDL	<MQL	<MQL
	Propylparaben	<MDL	<MDL	<MDL	<MQL	1.38 \pm 0.04	0.93 \pm 0.06	0.94 \pm 0.02	1.18 \pm 0.06	<MDL	<MDL	<MQL	<MQL
	1H-benzotriazole	0.88 \pm 0.76	1.38 \pm 0.18	0.83 \pm 0.51	0.66 \pm 0.01	1.14 \pm 0.53	2.24 \pm 1.28	<MQL	1.44 \pm 0.29	1.32 \pm 0.16	1.26 \pm 0.11	2.01 \pm 0.24	1.73 \pm 0.23

Figure captions

Figure 1. Study area showing bivalves sampling sites (●) and localities where environmental variables were measured (◇) in the Alfacs Bay (Ebre Delta).

Figure 2. Levels of carboxylesterase activity using four different colorimetric substrates (p-nitrophenyl acetate, pNPA; p-nitrophenyl butyrate, pNPB; 1-naphthyl acetate, 1-NA and 1-naphthyl butyrate, 1-NB) in the digestive gland of mussels, cockles and grooved razor shells collected in the Alfac's Bay of the Ebre Delta in April, May, June and July of 2017. Different letters indicate statistical differences among months during the sampling period ($p < 0.05$).

Figure 3. Levels of antioxidant enzymatic activities (glutathione reductase, GR; glutathione peroxidase, GPX and glutathione S-transferases, GST) determined in the digestive gland of mussels, cockles and grooved razor shells collected in the Alfac's Bay of the Ebre Delta in April, May, June and July of 2017. Different letters indicate statistical differences among months during the sampling period ($p < 0.05$).

Figure 4. Levels of carboxylesterase activity using four different colorimetric substrates (p-nitrophenyl acetate, pNPA; p-nitrophenyl butyrate, pNPB; 1-naphthyl acetate, 1-NA and 1-naphthyl butyrate, 1-NB) and acetylcholinesterase activity (AChE) in the digestive gland, gills and haemolymph of mussels collected in the Alfac's Bay of the Ebre Delta in April, May, June and July of 2017. Different letters indicate statistical differences among months during the sampling period ($p < 0.05$).

Figure 5. Levels of antioxidant enzymatic activities (glutathione reductase, GR; glutathione peroxidase, GPX and glutathione S-transferases, GST) and lipid peroxidation (LPO) levels determined in the digestive gland and gills of mussels collected in the Alfac's Bay of the Ebre Delta in April, May, June and July of 2017. Different letters indicate statistical differences among months during the sampling period ($p < 0.05$).

Figure 6. Dendrograms resulting from the hierarchical cluster analyses based on enzymatic data in mussel digestive gland (A), mussel gills (B), cockles digestive gland (C) and razor shells digestive gland (D) collected in the Alfac's Bay of the Ebre Delta in April, May, June and July of 2017.

Figure 7. *Rickettsia*-like organisms (RLOs) (arrowheads) in gills of cockles collected in the Alfac's Bay of the Ebre Delta in June of 2017.













