

Comparative study of domoic acid accumulation, isomer content and associated digestive subcellular processes in five marine invertebrate species

José Luis García-Corona, Hélène Hegaret, Malwenn Lassudrie, Amélie Derrien, Aouregan Terre-Terrillon, Tomé Delaire, Caroline Fabioux

▶ To cite this version:

José Luis García-Corona, Hélène Hegaret, Malwenn Lassudrie, Amélie Derrien, Aouregan Terre-Terrillon, et al.. Comparative study of domoic acid accumulation, isomer content and associated digestive subcellular processes in five marine invertebrate species. Aquatic Toxicology, 2024, 266, pp.106793. 10.1016/j.aquatox.2023.106793 . hal-04347002

HAL Id: hal-04347002 https://hal.univ-brest.fr/hal-04347002v1

Submitted on 15 Dec 2023

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1	Comparative study of domoic acid accumulation, isomer content and associated
2	digestive subcellular processes in five marine invertebrate species
3	
4	José Luis García-Corona ¹ , Hélène Hegaret ¹ , Malwenn Lassudrie ² , Amélie Derrien ² , Aouregan
5	Terre-Terrillon ² , Tomé Delaire ¹ , Caroline Fabioux ¹ *
6	
7	¹ Laboratoire des Sciences de l'Environnement Marin, UMR 6539 LEMAR
8	(UBO/CNRS/IRD/Ifremer). Institut Universitaire Européen de la Mer, rue Dumont d'Urville,
9	Technopôle Brest-Iroise, 29280 Plouzané, France.
10	
11	² Ifremer, LITTORAL LER BO, Station de Biologie Marine, Place de la Croix, BP 40537,
12	29900 Concarneau Cedex, France.
13	
14	*Corresponding author: Caroline Fabioux
15	
16	Laboratoire des Sciences de l'Environnement Marin, UMR 6539
17	(CNRS/UBO/IFREMER/IRD). Institut Universitaire Européen de la Mer, Technopôle Brest-
18	Iroise 29280, Plouzané, France.
19	
20	e-mail: <u>caroline.fabioux@univ-brest.fr</u>

22 Abstract

Despite the deleterious effects of the phycotoxin domoic acid (DA) on human health, and the 23 permanent threat of blooms of the toxic *Pseudo-nitzschia* sp. over commercially important 24 fishery-resources, knowledge regarding the physiological mechanisms behind the profound 25 differences in accumulation and depuration of this toxin in contaminated invertebrates remain 26 27 very scarce. In this work, a comparative analysis of accumulation, isomer content, and subcellular localization of DA in different invertebrate species was performed. Samples of 28 scallops *Pecten maximus* and *Aequipecten opercularis*, clams *Donax trunculus*, slippersnails 29 Crepidula fornicata, and seasquirts Asterocarpa sp. were collected after blooms of the same 30 concentration of toxic Pseudo-nitzschia australis. Differences (P < 0.05) in DA accumulation 31 were found, wherein *P. maximus* showed up to 20-fold more DA in the digestive gland than 32 the other species. Similar profiles of DA isomers were found between P. maximus and A. 33 opercularis, whereas C. fornicata was the species with the highest biotransformation rate 34 35 (~10%) and D. trunculus the lowest (~4%). DA localization by immunohistochemical analysis revealed differences (P < 0.05) between species: in *P. maximus*, DA was detected 36 37 mainly within autophagosome-like vesicles in the cytoplasm of digestive cells, while in A. opercularis and C. fornicata significant DA immunoreactivity was found in post-autophagy 38 residual bodies. A slight DA staining was found free within the cytoplasm of the digestive 39 cells of D. trunculus and Asterocarpa sp. The Principal Component Analysis revealed 40 similarities between pectinids, and a clear distinction of the rest of the species based on their 41 42 capacities to accumulate, biotransform, and distribute the toxin within their tissues. These findings contribute to improve the understanding of the inter-specific differences concerning 43 the contamination-decontamination kinetics and the fate of DA in invertebrate species. 44

45 Keywords: domoic acid, shellfish, DA isomers, autophagy, interspecific differences.

46 **1. Introduction**

47 Domoic acid (DA) is an extremely dangerous phycotoxin responsible of the illness referred as

- 48 amnesic shellfish poisoning (ASP) syndrome in humans (Perl *et al.*, 1990, Pulido, 2008; La
- 49 Barre *et al.*, 2014). This highly potent neuroexcitatory amino acid is naturally produced by
- some diatoms of the genus *Pseudo-nitzschia* (Bates *et al.*, 1998, 2018), wherein the species
- 51 *Pseudo-nitzschia australis* is one of the most toxigenic (Lelong *et al.*, 2012; La Barre *et al.*,
- 52 2014). The recurrent presence of toxic blooms of *Pseudo-nitzschia* sp., and the subsequent
- 53 production of DA, frequently affect fishery resources on the North Atlantic coasts of France.
- 54 Indeed, suspension-feeding invertebrates are capable of ingesting toxic *Pseudo-nitzschia* cells
- leading to high amounts of DA accumulated in their tissues (Basti *et al.*, 2018; Dusek
- Jennings *et al.*, 2020) seriously threatening human health through contaminated seafood
- 57 consumption (Pulido, 2008; La Barre *et al.*, 2014). Over the last two decades, these blooms
- have caused numerous and persistent harvest closures for some economically important
- 59 species (Amzil *et al.*, 2001; Husson *et al.*, 2016).
- 60 Notwithstanding, profound inter-specific variability in the toxicokinetics of accumulation and
- 61 depuration rates of DA burdens have been reported between several invertebrate species in the
- 62 same affected area (Costa *et al.*, 2004, 2005a,b; Bogan *et al.*, 2007a,b,c; Lage et al., 2012;
- Ben haddouch et al., 2016; Dusek Jennings et al., 2020; Blanco et al., 2021; Kvrgić et al.,
- 64 2022). Thus, invertebrates have been broadly classified as "fast" or "slow" DA-depurators
- 65 (Blanco *et al.*, 2002a,b; Basti *et al.*, 2018). Larger scallops, such as King scallops *Pecten*
- 66 maximus (Blanco et al., 2002a; García-Corona et al., 2022) and giant scallops Placopecten
- 67 *magellanicus* (Gilgan, 1990; Haya *et al.*, 1991), some big-clams, such as razor clams *Siliqua*
- 68 *patula* (Horner *et al.*, 1993; Dusek Jennings *et al.*, 2020), and some cephalopod mollusk such
- as Octopus vulgaris (Costa et al., 2004) and Eledone moschata (Costa et al., 2005b) as well as
- the common cuttlefish *Sepia officinalis* (Costa *et al.*, 2005a; Ben haddouch *et al.*, 2015) are
- capable of accumulating high amounts of DA, principally in the digestive gland, and require
- from many months to a couple of years to depurate the toxin from their tissues. Therefore,
- these species have been considered as slow DA-depurators. Notwithstanding, during *Pseudo-*
- *nitzschia* outbreaks, the king scallop *P. maximus* is usually amongst the most contaminated
- species (James *et al.*, 2005; Blanco *et al.*, 2002a, 2021). Levels of DA exceeding up to 5-fold
- the European regulatory limit of 20 mg kg⁻¹ are not unusual in *P. maximus* (Blanco *et*
- *al.*,2006; Bogan *et al.*, 2007a,b; García-Corona *et al.*, 2022). Conversely, mussels (Novaczek
- *et al.*, 1992 ; Blanco *et al.*, 2002b; Mafra *et al.*, 2010), and even smaller scallops, such as

Argopecten purpuratus (Álvarez et al., 2020) are known as fast DA-depurators since they can
depurate up to 90 % of total DA burdens over hours to days. These species-specific
differences in DA accumulation-depuration represent a real issue for fishery economy and
management after ASP-blooms. Thus, understanding the physiological mechanisms behind
this phenomenon is of high interest.

84 Mauriz and Blanco (2010), as well as Lage et al. (2012) found that nearly 90% of total DA accumulated in P. maximus and O. vulgaris, respectively, was free in a soluble form in the 85 cytoplasm of the digestive cells. García-Corona et al. (2022) observed, using an 86 immunohistochemical subcellular localization of DA in P. maximus, that DA is trapped into 87 small-spherical membrane-bound vesicles localized in the cytoplasm of digestive cells, 88 89 suggesting that autophagy could be one of the potential physiological mechanisms behind the long retention of a part of DA in this species. Nevertheless, to date, the immunohistochemical 90 (IHC) localization of DA has not been applied to any other invertebrate species contaminated 91 92 with DA, which greatly hinders the comparison of the subcellular mechanisms involved in the accumulation and retention of this toxin between affected species. Autophagy is a highly 93 94 regulated and dynamic "self-eating" catabolic system related to the intracellular ingestion and digestion (Cuervo, 2004; Wang et al., 2019; Zhao et al., 2021). Through autophagy the 95 lysosomes receive autophagosomic vesicles (autophagosomes) containing cytoplasmic 96 cellular components, such as macromolecules, damaged or misfolded proteins, and entire 97 organelles, as well as extracellular-derived molecular cargo from endocytosis and 98 99 phagocytosis for degradation, digestion, recycling, or excretion (Klionsky et al., 2014; McMillan, 2018; Wang et al., 2019). These distinctive capabilities establish an essential role 100 101 of autophagy in maintaining metabolic homeostasis and cellular health in bivalves (Balbi et al., 2018; Picot et al., 2019; Rodríguez-Jaramillo et al., 2022). 102

- 103 Not only untransformed DA, but also some structural isomers of the toxin (*i.e.* isoA, isoD,
- 104 isoE, and epi-DA) are frequently detected in seafood during ASP-monitoring. The
- 105 concentrations of DA-isomers commonly range from 0.5 to ~20% of total DA burdens
- 106 (Wright et al., 1990a; Costa et al., 2005; Takata et al., 2009; Zheng et al., 2022). Despite
- 107 some studies pointing out some degree of species-specific biotransformation of DA in
- 108 bivalves (Wright *et al.*, 1990b; Blanco *et al.*, 2010), fish and shellfish (Vale and Sampayo,
- 109 2001), and cephalopods (Costa *et al.*, 2005), no work has ever compared the
- 110 biotransformation profiles of DA against the subcellular localization of this toxin in
- 111 contaminated invertebrates. This information could be useful to elucidate differences in DA

- 112 uptake and allocation, as well as the potential implication of subcellular mechanisms on
- 113 depuration of this toxin between species.
- 114 This study compared biotransformation and subcellular localization of DA in five invertebrate
- species simultaneously exposed to natural toxic *P. australis* blooms to answer the question:
- 116 How do invertebrate species differ in their ability to accumulate, process, and allocate DA in
- 117 their tissues?
- 118

119 **2. Materials and methods**

2.1. Sample collection and *Pseudo-nitzschia australis* bloom-associated environmental data

A total of 38 invertebrate samples were collected in 2021 in the northwest coast of Brittany, 122 France. The samples consisted in clams *Donax trunculus* (n = 11) collected on the 30th of 123 March in the Bay of Douarnenez, and scallops *P. maximus* (n = 5), *A. opercularis* (n = 10), 124 slippersnail Crepidula fornicata (n =7), and sea squirt Asterocarpa sp. (n = 5) collected on the 125 8th of April in Camaret-sur-Mer (Fig. 1). Animals were collected eight days after blooms of 126 similar intensity of the DA-producing P. australis according to the French national 127 phytopankton monitoring network (French Observation and Monitoring program for 128 Phytoplankton and Hydrology in coastal waters, REPHY) in both sampling sites ($[2.6 \times 10^5]$ 129 cell.L⁻¹] on March 23, 2021 in the Bay of Douarnenez), and $[1.1 \times 10^5 \text{ cell.L}^{-1}]$ on March 30, 130 2021 (in Camaret-sur-Mer), respectively, https://bulletinrephytox.fr/accueil) (Fig 1). Once at 131 the laboratory, the digestive gland (DG) of the scallops (P. maximus and A. opercularis) was 132 carefully dissected from the rest of the tissues, and subsequently sectioned in two halves. For 133 the rest of the species with diffuse visceral mass (C. fornicata, D. trunculus, and Asterocarpa 134 135 sp.) the soft body (*i.e.* total flesh) was divided into two equal portions at the mid visceral level, including a section of the DG on each. For each individual, one of these DG/visceral 136 sections was fixed in Davidson's solution (Kim et al., 2006) for histology, and the second 137 DG/visceral sections section was stored at -20 °C for toxin analysis. 138

139 2.2. Toxin quantification and DA-isomer analysis by liquid chromatography-tandem 140 mass spectrometry (LC-MS/MS)

Since the DG accumulates most of DA (Mauriz and Blanco, 2010), only this tissue wasconsidered for toxin analysis in this work. For the non-pectinid species, the DG was separated

- 143 from the rest of the visceral mass once the tissues were frozen. DA was extracted from the
- 144 DG following the procedure described by Quilliam *et al.*, (1989). Samples were homogenized
- from 200 ± 10 mg of frozen DG in 1 mL of 50% MeOH/H₂O using a Fastprep-24 5G system
- 146 (MP Biomedicals, Sta. Ana, CA, USA). The extract was clarified by centrifugation at 19,000
- 147 \times g at 4 °C for 10 min and the supernatant was isolated, filtered through a 0.2 µm nylon
- 148 centrifugal filter (VWR International, Radnor, PA, USA), and stored at -20 °C until analysis.
- 149 The quantification of total DA (tDA = ensemble of all DA isomers) and each isomer of the
- toxin in the DG was carried out by LC-MS/MS according to Ayache et al. (2019) with
- 151 modifications, using a Shimadzu UFLCxr system coupled to a quadruple hybrid mass
- spectrometer API400Q-Trap (Sciex, Concord, ON, Canada) equipped with a heated
- electrospray ionization (ESI) source. Chromatographic separation was carried out on a
- reversed-phase column Phenomenex Luna Omega C18 (150×2.1 mm, 3 µm, Phenomenex,
- 155 Torrance, CA, USA). The separation was carried out using a mobile phase consisting of
- aqueous eluent A (100% $H_2O + 0.1\%$ H-COOH) and organic eluent B (95% $CH_3CN/5\%$ H_2O
- + 0.1% H-COOH). The run started following a gradient from A to B as follows: 5% at min 0,
- 158 18.6% at 17 min, 95% at 17.5 min, 95% at 19.5 min, 5% at 20 min, and 5% at 25 min. The
- flow rate was 200 μ L.min⁻¹ and the injection volume was 5 μ L. The column temperature was maintained at 30 °C.
- The ESI interface was operated with a curtain gas of 20 psi, temperature of 550 °C, gas1 55 161 162 psi, gas2 60psi, and an ion spray voltage of 5500 V. The detection of DA was achieved by multiple reaction monitoring (MRM) in positive ion mode. The transition 312.1 > 266.1163 164 (collision energy = 22 V) was used for quantification and 312.1 > 161.1 (collision energy = 33 V) for confirmation. The quantification was performed relative to the DA standard 165 166 (National Research Council Canada, NRCC) with a 6-point calibration curve. The Limit of 167 Quantification (LOQ) (S/N = 10) and the Limit of Detection (LOD) (S/N = 3) of the method were 0.25 and 0.08 ng DA mL⁻¹, respectively, which corresponded to 1.25 and 0.4 ng DA g⁻¹ 168 in tissue. 169

170 2.3. Immunodetection of DA and quantitative histology

171 Tissue samples fixed in Davidson's solution were dehydrated in ethanol series of progressive

- 172 concentrations (70%, 80%, 95%, and 100%), cleared in xylene, and embedded in paraffin
- 173 (Paraplast Plus, Leica Bio-systems, Richmond, IL, USA). Paraffin blocks were cut in 4-µm-
- thick sections using a rotary microtome (Leica RM 2155, Leica Microsystems) and sections

- mounted on polylysine-coated glass slides (Sigma-Aldrich, St. Louis, MO, USA). A series of
- three consecutive sections was performed for each sample, which were used for (i)
- immunohistochemical detection of DA, (ii) multichromic staining, and (iii) hematoxylin/eosin
- staining, as described below. Sections were deparaffinized in xylene and rehydrated in ethanol
- 179 series of regressive concentrations before staining.
- 180 In order to detect the presence of DA at the subcellular level in the tissue sections, an
- 181 immunohistochemical DA labeling technique was applied following the procedure described
- in García-Corona *et al.* (2022) on the first slide of each sample. Briefly, tissue sections were
- incubated overnight with a Goat polyclonal anti-DA primary antibody $(0.01 \text{ mg.mL}^{-1},$
- 184 Eurofins Abraxis[®], Warminster, PA, USA) at 4°C, and the next day the slides were incubated
- at 37 °C for 2h with an HRP sharped IgG Rabbit anti-Goat secondary antibody (0.001 mg.mL⁻
- ¹, abcam[®], Cambridge, UK). Then, samples were washed and revealed with diaminobenzidine
- 187 (DAB+ Chromogen Substrate Kit, abcam[®], Cambridge, UK) for 1 h in darkness at room
- temperature and counterstained with Harry's hematoxylin.
- 189 The second slide from each sample was stained with a multichromic procedure (Costa and
- 190 Costa, 2012). This technique consists of a combination of Alcian Blue and Periodic Acid-
- 191 Schiff's for the demonstration of acid mucopolysaccharides and neutral glycoconjugates, in
- blue and magenta tones, respectively, Hematoxylin blueing for nuclear materials, and Picric
- 193 Acid to identify proteins in yellow hues.
- The last set of tissue sections was stained with Hematoxylin–Eosin as reference (Kim *et al.*,
 2006). The slides were examined under a Zeiss Axio Observer Z1 light microscope.
- 196 For quantitative histological analysis, five randomly selected regions $(63\times; -1.3 \text{ mm}^2)$ from
- each DG section treated for immunohistochemical DA detection, multichromic, and
- hematoxylin-eosin staining were digitized at high resolution (600 dpi). A total of 570 images
- 199 (*i.e.* 114 micrographs by species) were used to obtain the following data: (a) DA chromogenic
- signal (DAcs) corresponds to the coverage area, in pixels, occupied by the positive anti-DA
- staining. This was manually calculated using an operator-driven digital image analysis system
- 202 (Image Pro Plus software v. 4.5, Media Cybernetics, Silver Spring, MD, USA) (Gómez-
- Robles *et al.*, 2005). The area reported as the DA chromogenic signal was calculated as DAcs
- 204 = (DA chromogenic signal area/ total area occupied by the DG on the analyzed region of the
- slide) \times 100. Since almost all the DA chromogenic signal detected in DG is trapped in
- 206 membrane-bound vesicles present in the cytoplasm of digestive cells (García-Corona et al.,

2022), the (b) Total autophagy (Ta) and total DA autophagy (DAa) were calculated by 207 counting the total number of autophagosome-like vesicles, and the number of 208 autophagosome-like vesicles with DA chromogenic signal, respectively, on each digitized 209 210 image. A fraction of the DA chromogenic signal is also observed in post-autophagic residual bodies within the digestive cells (García-Corona et al., 2022), thus the frequencies of (c) Total 211 residual bodies (Trb) and DA residual bodies (DArb) were assessed as the total number of 212 residual bodies and the total number of residual bodies with DA chromogenic signal, 213 respectively, on each digitized image. Finally, (d) Cell vacuolization (Vac), measured as an 214 215 indicator of potential histopathologies related to DA accumulation in the DG, represents the total number of vacuoles within the digestive cells of each invertebrate species on each 216

- 217 digitized image.
- 218

219 **2.4. Statistical analysis**

All statistical analyses were performed in the R computing environment (R v. 4.2.2, R Core 220 221 Team, 2022). A priori Lilliefors (Kolmogorov-Smirnov) and Bartlett tests were applied to 222 confirm the normality of frequencies and homogeneity of variances of the residuals of the data, respectively (Hector, 2015). All data were transformed (log, $1/\chi$, or $\sqrt{\chi}$) prior to analysis 223 224 to meet a priori assumptions. The percentage-expressed values were also arcsine (arcsine \sqrt{P}) transformed (Zar, 2010), but all data are reported untransformed as the means \pm standard 225 226 errors (SE). Separate one-way analyses of variance (ANOVA, type II Sum of Squares) were applied to assess statistically significant differences of toxin accumulation in the DG, 227 228 proportion of DA isomers, and quantitative histological features between species. As needed, 229 post hoc comparisons of means with Tukey's honest significance test (HSD) were performed 230 to identify differences between means (Hector, 2015; Zar, 2010). Principal component 231 analysis (PCA) was performed using the FactoMineR package with the factoextra package for data visualization into smaller factorial clusters within a 95% confidence interval. All data 232 matrices were auto-scaled before PCA analysis. The corrplot package was run to calculate the 233 correlation coefficients and their significance between variables within their given PCs. All 234 graphics were generated using the package ggplot2. The level of statistical significance was 235 set at $\alpha = 0.05$ for all analyses (Zar, 2010). 236

237

238 **3. Results**

239 **3.1. Toxin accumulation and biotransformation**

- 240 Significant differences in the amounts of total DA (tDA) accumulated in the digestive glands
- 241 (DG) were found between the different invertebrate species sampled after blooms of the toxic
- 242 *P. australis* (Fig. 2). The significantly higher burdens of tDA were observed in the scallop *P*.
- 243 *maximus*, with $638.6 \pm 35.5 \text{ mg.kg}^{-1}$, followed by those of the snail *C. fornicata*, with $48.5 \pm$
- 14.2 mg.kg⁻¹, the scallop A. opercularis (22.7 \pm 2.6 mg kg⁻¹), and the clam D. trunculus (12 \pm
- 1.7 mg kg⁻¹)., The lowest values (P < 0.05) of tDA were found in the ascidian Asterocarpa sp.
- 246 $(4.2 \pm 1.5 \text{ mg kg}^{-1})$. Moreover, as shown in Fig. 2, an important intraspecific variability in
- tDA accumulation was also observed in *P. maximus* and *C. fornicata*, with values ranging
- from 530 to 731 mg kg⁻¹, and from 0.2 to 93.8 mg kg⁻¹, respectively.
- 249 The toxin analysis carried out by LC-MS/MS revealed differences in biotransformation of DA
- in the digestive glands among the different invertebrate species (Table I). For all species,
- relative concentration levels of DA isomers were <10 % of the tDA burdens. Nonetheless, *C*.
- *fornicata* was the species with the highest proportions (P < 0.001) of DA isomers (9.3 ± 1.1)
- 253 %), while *D. trunculus* showed significantly low DA isomer amounts $(4.2 \pm 0.3 \%)$.
- 254 Concerning the analysis of DA isomers proportion, *P. maximus* and *A. opercularis* showed
- similar biotransformation profiles of the toxin since similar amounts of each DA isomer were
- reported in both species. Furthermore, as shown in Table I, among the five species, the lowest
- ratio of isoE (P < 0.05) was measured in *Asterocarpa* sp., and a significantly higher proportion
- of isoD was recorded in *C. fornicata*, while the smallest amounts (P < 0.05) of isoA and epi-
- 259 DA were quantified in *D. trunculus*.

260 **3.2. DA subcellular localization and histological measurements**

The microanatomical observations of histological sections evidenced qualitative differences in the localization of DA and the subcellular features linked to the accumulation of the toxin

among the invertebrate species analyzed in this study (Fig. 3, and supplementary materials

- 264 S1-5). DA detected by immunohistochemistry (IHC) appeared as a brown chromogenic signal
- 265 (cs) on slides (Fig 3A, 3D, 3G, 3J, 3M, and S1A-B, S2A-B, S3A-B, S4A-B, S5A-B).
- In the digestive gland of *P. maximus* DA was detected mainly trapped within small (~1-2.5
- 267 µm diameter) autophagosome-like vesicles (a) distributed throughout the cytoplasm of the
- 268 digestive cells (dc). A narrow fraction of DA-immunoreactivity was also observed in residual
- bodies (rb) distributed in the acinar region (ar) of the digestive diverticula (dd) (Fig. 3A, S1A-
- B). The presence of membrane-bounded vesicles (a) with positive DA-signal (cs) in the

- tubular region (tr) of the digestive diverticula (dd) was confirmed by means of the
- 272 multichromic staining (MC), which produces a dark violet/blue hueing in membrane-bounded
- structures (Fig. 3B, S1C-D). Hematoxylin-Eosin (H&E) staining (Fig. 3C, S1E-F) highlighted
- a moderate vacuolization (v) within the cytoplasm of the digestive cells of *P. maximus*.
- 275 Neither the autophagosomes (a) nor the residual bodies (rb) acquired any coloration with the
- 276 H&E staining but residual bodies appeared yellow-green.
- 277 In the queen scallop *A. opercularis*, a strong DA-chromogenic signal (cs) was found in the
- residual bodies (rb) of the digestive diverticula (dd) (Fig. 3D, S2A-B). No DA chromogenic
- signal was observed in the autophagosome-like vesicles (a) present in the cytoplasm of the
- digestive cells of the digestive diverticula (dd) (Fig. 3E-F, S2A-B). An intense process of
- vacuolization (v) of the digestive cells of *A. opercularis* was found (Fig. 3E-F, S2C-D), while
- H&E staining (Fig. 3F, S2E-F) showed that the autophagosomes seem to gather giving rise to
- the residual bodies (rb) in the cytoplasm of the adipocyte-like digestive cells (al) of the
- 284 digestive diverticula (dd).
- A similar result was found for *C. fornicata*, since most of the brown DA-chromogenic
 staining (cs) was found in small residual bodies (rb) present in the basal cytoplasmic region
 (bl) of the digestive cells (dc) (Fig. 3G, S3A-B), while autophagosome-like vesicles (a) that
 are distributed in the apical region of the digestive cells (dc) (Fig. 3H-I, S3A-B) did not show
 any DA-immunoreactivity.
- 290 A slight-blurred DA-chromogenic signal (cs) was also observed only free in the cytoplasm of
- the digestive cells of *D. trunculus* (Fig 3J, S4A-B). The presence of autophagosome-like
- vesicles (a, small blue colored vesicles distributed in the cytoplasm, Fig 3K, S4C-D) and
- residual bodies (rb, larger round non-colored structures present within adipocyte-like cells,
- Fig 3L, S4C-D) was confirmed in the digestive cells (dc) of clams (Fig. 3K-L, S4C-F).
- 295 Meanwhile, in sea squirts (Asterocarpa sp.) DA-chromogenic signal (cs) was rarely identified
- and was located as small brown points (Fig. 3M, S5A-B) distributed through the digestive
- epithelium (pse) of the blind ampulla (ba) (Fig. 3N-O, S5C-F).
- 298 The results of the quantitative analysis of histological parameters are shown in Fig. 4. The
- 299 coverage area of the DA chromogenic signal (%DAcs, Fig. 4A) was significantly higher in
- the most contaminated invertebrate species (*P. maximus* = 4.8 ± 0.4 %, and *C. fornicata* = 5.3
- ± 0.4 %). In addition, differences (*P* < 0.05) were found in the amount of DA chromogenic

- signal in *A. opercularis* $(3.2 \pm 0.2 \%)$ compared to the species contaminated with the lowest DA burdens (*D. trunculus* = 0.2 %, and *Asterocarpa* sp. = 0%).
- 304 On the other hand, as seen in Fig. 4B, total autophagy (Ta) reached its highest values (P <0.05) in the bivalve species, with frequencies of 185.4 ± 18 autophagosomes, area⁻¹ in P. 305 maximus, 123.2 ± 12.6 autophagosomes, area⁻¹ in *D. trunculus*, and 102.9 ± 9.7 306 autophagosomes. area⁻¹ in A. opercularis. The proportion of total autophagy (Ta) was 307 significantly lower in *C. fornicata* (60.9 ± 5.8 autophagosomes. area⁻¹) and *Asterocarpa* sp. 308 $(18.3 \pm 2.9 \text{ autophagosomes. area}^{-1})$. Nevertheless, the frequency of autophagosomes with 309 positive DA-chromogenic signal (DAa) significantly peaked in P. maximus (99.7 \pm 9.7 310 autophagosomes. area⁻¹, corresponding to 53.8% of the Ta), followed by C. fornicata (39.8 \pm 311 4.6 autophagosomes. area⁻¹, corresponding to 65.3% of the Ta). The lowest proportions (P312 <0.05) of autophagosomes with positive DA-chromogenic signal (DAa) were observed in A. 313 opercularis, D. trunculus, and Asterocarpa sp, with ≤ 7 autophagosomes. area⁻¹, 314 which corresponded to 8.4, 1.2, and 0% of the total autophagy (Ta), respectively (Fig. 4B). In 315 contrast, the frequencies of total residual bodies (Trb) and residual bodies with DA 316 chromogenic signal (DArb) significantly peaked in C. fornicata (92.4 \pm 5.2 rb. area⁻¹, and 317 51.9 ± 4.1 rb. area⁻¹, respectively), while the frequencies of both subcellular parameters 318 319 showed their lowest values (P < 0.05) in the rest of the species (Fig. 4C). It is important to highlight that the percentage of residual bodies with DA chromogenic signal (%DArb) 320 321 compared to total residual bodies (Trb) was significantly higher in A. opercularis, with a 67.1 \pm 3%, followed by *C. fornicata* and *P. maximus*, with rates of 58 \pm 3.8% and 35.4 \pm 3.3%, 322 323 respectively. The lowest % DArb (P < 0.05) was reported for D. trunculus ($2.2 \pm 1.3\%$) and Asterocarpa sp. (0%). Finally, the highest frequency of cell vacuolization (Vac) of the 324 digestive cells was measured in A. opercularis (67.4 \pm 6.7 vacuoles. area⁻¹, P <0.05), followed 325 by *P. maximus* $(31.6 \pm 2.4 \text{ vacuoles. area}^{-1})$. Significantly lower vacuolization (Vac) rates 326 were reported for the rest of the species (<8 vacuoles. area⁻¹, Fig. 4D). 327

328 3.3. Integrative analysis compiling DA accumulation/biotransformation and subcellular 329 features

- A principal component analysis (PCA) was computed to summarize all variables measured in
 this study on the five invertebrate species studied: DA accumulation, biotransformation, and
- subcellular parameters (Fig. 5). The PCA described two-thirds (66.6 %) of the total variance
- 333 of the data along the first two principal dimensions. For the whole data set, the clustering-

PCA provided a clear distinction between species, except for the two pectinid species, which 334 slightly overlapped (Fig. 5A). In the scatter plot, P. maximus and A. opercularis showed 335 similar scores on the principal components and were different from the rest of the species. 336 Meanwhile, D. trunculus, C. fornicata, and Asterocarpa sp., were grouped separately from 337 each other (Fig. 5A). As shown in Fig. 5B, the dimension/principal component 1 (PC1, 42.3 338 % of the total variance) mainly explained the accumulated untransformed DA, isoD and isoA, 339 as well as the histological parameters such as domoic acid chromogenic signal (%DAcs). 340 domoic acid autophagy (DAa), total residual bodies (Trb), and residual bodies with DA signal 341 342 (DArb). In this PC1, the fraction of isoA was strongly and positively correlated to the %DAcs and DArb (r = 0.5 and 0.6, P < 0.05, respectively). Likewise, a strong and significant 343 344 correlation was found between the untransformed DA and DAa (r = 0.8), and between DArb and %DAcs (r = 0.8) in this dimension. The amounts of isoE and epi-DA, as well as total 345 346 autophagy (Ta) and vacuolization (Vac), were the strongest correlated variables to dimension/principal component 2 (24.3 % of the explained variance). A positive correlation (r 347 348 = 0.5, P < 0.05) between total DA (tDA) and isoE was found with Ta within the PC2. As observed in Fig. 5, P. maximus and A. opercularis were associated with higher tDA and isoE, 349 350 as well as the maximum frequencies of Ta and Vac. Meanwhile, C. fornicata was related to 351 higher amounts of isoD, epi-DA, Trb, and D. trunculus with the highest fraction of

352 untransformed DA.

353 **4. Discussion**

In this study, we compared domoic acid (DA) accumulation and isomer profiles with the subcellular localization of this toxin among naturally contaminated invertebrates to progress in the understanding of interspecific differences in DA fate in marine invertebrates.

The DA contents measured in invertebrate tissues are the result of the accumulated and the subsequently depurated toxin. Moreover, differences in DA accumulation in the organisms are strongly dependent on the toxicity of the *Pseudo-nitzschia* cells, the duration of the ASP blooms, the time through the animals were exposed to toxic microalgae, and the moment at which the organisms were sampled during the bloom. In this work, DA contaminated animals were collected 8 days after maximum cell densities of *P. australis* bloom of similar intensity, duration and origin.

Since DA is a highly water-soluble molecule, it is expected to be easily accumulated in the 364 majority of forager species (Trainer et al., 2012; La Barre et al., 2014). Nonetheless, the 365 scallops, but notably P. maximus, as well as C. fornicata, remained significantly more 366 contaminated than the rest of the species in this study. These important differences in DA 367 accumulation in the digestive gland at the interspecific level are in accordance with 368 considerably high variability in DA amounts frequently detected in these species (Bogan et 369 al., 2007a,b,c; Basti et al., 2018, Blanco et al., 2021) resulting from differences in the 370 accumulation but also in the depuration rates of DA reported mostly for bivalve species (Vale 371 372 and Sampayo, 2001; Blanco et al., 2010; Dusek Jennings et al., 2020). Notably, within the pectinidae family, some large scallops like P. maximus can accumulate up to 3,200 mg 373 DA.kg⁻¹ in their DG (James et al., 2005; Blanco et al., 2006), which is 5-fold more than the 374 DA accumulated in the DG of the same species in this work. In contrast, smaller scallops, 375 such as A. opercularis (Ventoso et al., 2019), A. purpuratus (Álvarez et al., 2020) and A. 376 irradians (O'Dea et al., 2012) accumulate lower DA burdens (~7-30 mg DA.kg⁻¹) similar to 377 378 those recorded in A. opercularis in this work, in the same organs. Depuration kinetics of the toxin differ also between these species. Whereas P. maximus exhibits depuration rates as slow 379 as 0.007 day⁻¹ in the DG, remaining highly contaminated for months or even a few years 380 (Blanco et al., 2002a, 2006), other scallops such as A. purpuratus show decontamination 381 debits near to 10 day^{-1} in the DG, allowing to depurate ~90% of total DA burdens within 382 hours or a couple of days (Álvarez et al., 2020). Thus, after all the differences in 383 accumulation and depuration rates of DA between invertebrate species discussed above, a 384 possible event of rapid depuration of DA in A. opercularis, D. trunculus, and Asterocarpa sp. 385 before sampling can be part of the interspecific differences of DA concentrations measured in 386 this study. Several factors could explain variability in DA decontamination: the transfer of 387 DA in other tissues than DG, its biotransformation and its depuration. 388

Differential tissue distribution of DA may not explain more than 20% of the interspecific
variability observed in this study since the digestive gland accumulates more than 80% of

total DA burdens in most invertebrates (Blanco *et al.*, 2002a; Costa *et al.*, 2005a,b). For all

the five species of this study, three bivalve molluscs (*P. maximus*, *A. opercularis* and *D.*

393 trunculus), one gasteropod mollusc (C. fornicata) and one ascidian (Asterocarpa sp.) DA

isomers were observed in digestive gland with significant interspecific differences between

the proportions of isomers E, D, A and epi-DA; iso-E being more represented in molluscs

396 compared to ascidian. Although it is known that DA isomerization can occur within toxic

Pseudo-nitzschia cells (Amzil et al., 2001; Bates et al., 2018; Quilliam et al., 1989; Wright et 397 398 al., 1990a), in the present study all invertebrate species were exposed to the same Pseudonitzschia toxic bloom. These two sets of information demonstrate that metabolic conversion 399 400 of DA occurs in marine invertebrates as hypothesized first by Vale and Sampayo (2001) and is species-specific. The integrative analysis revealed a close and significant relationship 401 402 between some subcellular features (vacuolization, autophagy, presence of residual bodies) and the isomer profile of the toxin. Understanding DA compositional changes is important not 403 only as a means of predicting toxicity, but also because biotransformation could participate in 404 405 the prolonged retention of this toxin in invertebrate species by means of some of the 406 subcellular mechanisms analyzed here. Notwithstanding, biotransformation does not appear to 407 be the main route of DA elimination in these species since it represents less than 10% of total DA of the digestive gland measured in these five species, as well as in previous studies (Costa 408 409 et al., 2005a; Blanco et al., 2010; Zheng et al., 2022). There is only one study showing some insights of DA biotransformation linked to apparent augmentation of the overall DA 410 411 detoxification rate in the cuttlefish Sepia officinalis, wherein DA isomers comprise a relevant percentage of the toxin profile in the branchial hearts, suggesting that this organ has an 412 important function in system detoxification of DA (Costa et al., 2005a). 413

Furthermore, it is worth to mention that king scallops were slightly contaminated (~5 mg DA kg⁻¹, data from the REPHY French monitoring program) before the bloom of *P. australis* occurred in late March 2021, after which they became highly contaminated (~ 650 mg DA kg⁻¹). Therefore, it is inferred that the concentrations of DA isomers found in the digestive glands of *P. maximus*, and consequently, in all the invertebrate species analyzed in this work, were the result of the bloom of *P. australis* occurred in late March 2021.

420 Despite the enormous differences in DA concentrations between the marine invertebrates 421 analyzed in this work, the physiological mechanisms behind this phenomenon remain poorly understood. To date, only a few hypotheses about the biological processes potentially 422 423 involved in the large accumulation and long retention of DA in some bivalve species have been proposed. On the one hand, Trainer and Bill (2004) characterized tissue-specific 424 425 expression of high and low affinity glutamate receptors in S. patula, inferring that this species might selectively express low affinity glutamate receptors in all tissues, and high affinity sites 426 in specific tissues that retained DA for long periods of time. On another hand, Mauriz and 427 Blanco (2010) hypothesized that one of the causes of the long retention of DA in the DG of P. 428 429 maximus was not the binding of the toxin to some cellular component as previously discussed,

but the lack of efficient membrane transporters in the scallops to excrete the toxin. Recently, 430 431 using immunostaining of DA, García-Corona et al. (2022) revealed that in P. maximus, once entered the cells, a part of DA was localized in the cytoplasm of digestive cells of the 432 digestive diverticula, trapped within autophagosome-like vesicles. Moreover, transcriptomic 433 analyses revealed the upregulation of genes related to autophagy and vesicle-mediated 434 transport in the DG of P. maximus injected with DA in the adductor muscle (Ventoso et al., 435 2021), as well as in the DG of A. opercularis after exposure to DA-producing Pseudo-436 nitzschia (Ventoso et al., 2019). Taken together, these data suggest that the formation of 437 438 autophagosomal structures could be part of the explanation for the long retention of DA in P. maximus. The results obtained in this work cope with these findings, since most of the DA-439 440 labeling was found within a large number of autophagosomes distributed throughout the cytoplasm of the digestive cells in *P. maximus*. Additionally, a strong DA-chromogenic signal 441 442 was found within the post-autophagic residual bodies present in the adipocyte-like cells in A. opercularis, and in the basal region of the digestive diverticula in C. fornicata. During 443 444 autophagy the lysosomes in the digestive cells of these species receive DA trapped within autophagosomic-vesicles. Nonetheless, the evidence of this work indicates that a fraction of 445 DA remains accumulated within autophagosomic structures instead being excreted or used by 446 447 the cells, leading to its accumulation within the autophagosomes, and consequently blocking its excretion outside the cell by exocytosis (Cuervo, 2004; Zhao et al., 2021). This eventually 448 triggers the aggregation of autophagosomes with sequestered DA to form residual bodies that 449 can remain in the cytoplasm of the digestive cells indefinitely. There is evidence of the long 450 451 retention of exogenous compounds through specialized cellular mechanisms in animals. A concrete example is the dynamics of phagocytosis displayed by dermal macrophages, 452 explaining both persistence and strenuous removal of tattoo ink in mammalian skin. Baranska 453 454 et al. (2018) demonstrated that upon tattooing, pigment particles are captured by dermal macrophages. Eventually, macrophages laden with tattoo ink die and release the pigment 455 particles, which remain in an extracellular form at the site of tattooing where they are 456 457 recaptured by neighboring or incoming macrophages. Through adult life, several cycles of ink capture-release-recapture can occur, accounting for long-term tattoo persistence (Baranska et 458 al., 2018). Macrophagy and autophagy are analogous processes. 459 During macrophagy specialized cells called macrophages use their 460 461 cytoplasmic membranes to engulf large extracellular particles (\geq 0.5 µm, *i.e.* bacteria and debris) via endocytosis, giving rise to internal vesicular 462

compartments called phagosomes. Phagosomes with cargo materials fuse with lysosomes, 463 forming phagolysosomes, leading to enzymatic degradation (Flannagan et al., 2012; Gordon, 464 2016). Like autophagy, macrophagy is a major mechanism used to remove pathogens and 465 466 cellular debris for detoxification or nutrient recycling purposes, in which macrophages can have lifespans of months to a few years (Baranska et al., 2018). The discussion above raises a 467 new hypothesis suggesting that a part of DA that is not excreted from the cells due to the lack 468 of efficient membrane transporter (Mauriz and Blanco, 2010), may undergo successive cycles 469 of capture-release-recapture by autophagosomes through the regenerative cycle of digestive 470 471 cells in some invertebrates, without any or very few toxin vanishing from months to years. 472 Therefore, long-term DA persistence could rely on autophagosome renewal or on potential 473 longevity of residual bodies. A close relationship between early autophagy and DA 474 sequestration can be established in *P. maximus*, whereas in *A. opercularis* and *C. fornicata* 475 toxin accumulation seems to be closely linked to late autophagy and the formation of residual bodies in the DG. This evidence strengthens the hypothesis stated by García-Corona et al. 476 477 (2022), where autophagy was proposed as one of the possible causes of the prolonged retention of part of DA initially accumulated, now not only in *P. maximus*, but also in other 478 479 marine invertebrates. The next step is to decipher the fate and life-spent of autophagosomes 480 and residual bodies with anti-DA immunolabelling within a scenario of contamination and decontamination. 481

Although the IHC method for the *in situ* detection of DA in contaminated invertebrates used 482 in this work has a high-sensitivity (~1 mg DA.kg⁻¹, García-Corona et al., 2022) only a slight-483 blurred DA chromogenic signal was found in the cytoplasm of the digestive cells of D. 484 *trunculus*, and *Asterocarpa* sp. This would suggest that in these species, intracellular DA is 485 not bound to any subcellular structure or component. Consequently, the feeble amounts of 486 toxin free in the cytoplasm of the digestive cells could be quickly depurated after DA 487 contamination but a part of DA, could also be lost by washing during histological process. 488 489 Furthermore, when all species are compared, the proportion of DA chromogenic signal seems not correspond to the total amount of toxin accumulated in the DG of the animals. Despite the 490 large difference in DA concentration between P. maximus and A. opercularis (638.6 mg DA 491 kg^{-1} vs 22.7 mg DA kg^{-1} , respectively), the difference in DA signal was small (~2 % between 492 493 both species). Therefore, it is possible that a fraction of the DA accumulated in the DG of both species is free and dissolved in the cytoplasm of the digestive cells as reported for P. 494 495 maximus (Mauriz and Blanco, 2010) and for O. vulgaris (Lage et al., 2012), and that P.

maximus effectively lacks efficient membrane transporters to excrete the toxin out of the cell
(Mauriz and Blanco, 2010), thus the chromogenic signal observed in the DG of both pectinids
could correspond to the fraction of DA trapped by the autophagic system, and not to the total
DA burdens in the DG. Further analyzes will be necessary to corroborate all the ideas
discussed above.

501 Scallops, P. maximus but even more so A. opercularis contaminated by DA in this study have significantly higher digestive cell vacuolization rates in their digestive gland compared to 502 other species. Cell vacuolization is a common histopathological lesion in bivalves under 503 stressful environmental conditions (Rodríguez-Jaramillo et al., 2022). According to Shubin et 504 al. (2016) this is a well-known subcellular phenomenon observed in animal cells which often 505 506 accompanies cell death after exposure to artificial or natural low-molecular-weight 507 compounds, such as DA. The scarce literature related to the effects of *Pseudo-nitzschia spp*. 508 or DA on invertebrates indicates that DA could potentially disturb behavioral, metabolic, 509 molecular, and physiological processes in some bivalves such as P. maximus (Ventoso et al., 2021; Liu et al., 2007a,b), A. opercularis (Ventoso et al., 2019), A. irradians (Chi et al., 510 511 2019), and some mussels, like M. edulis (Dizer et al., 2001) and M. galloprovinciallis (Pazos et al., 2017). Nevertheless, no lethal effects resulting from exposure to DA have been reported 512 in any of these species, suggesting either a low sensitivity to the toxin or yet unnoticed 513 514 negative effects. Further research is needed in order to decipher how DA exposure and its biotransformation modulate cell vacuolization, as well as its potential detrimental effects on 515 516 the digestive cells of pectinids, and possibly, over other invertebrates, as reported for other phycotoxins in other bivalve species (Hegaret et al., 2010; Lassudrie et al., 2014). 517

518 Furthermore, as discussed above, the highest proportions of total autophagy, and production 519 of residual bodies reported in P. maximus, A. opercularis, and C. fornicata, seems to directly 520 correspond to the sequestration of DA within these subcellular structures, which indicates that autophagy could be also considered as a sign of homeostatic impairment, as reported in other 521 marine bivalve species when activated as an auxiliary mechanism for recycling internal 522 energy to cope with detrimental environmental conditions (Moore, 2008; Rodríguez-Jaramillo 523 et al., 2022), or to depurate toxicological agents (Moore, 2004; Picot et al., 2019). The 524 525 particularly highest proportions of DA-autophagy in *P. maximus* analyzed here stress out the 526 need to carry out the measurement of the frequency of these subcellular features in a DA contamination and decontamination scenario. This basic knowledge is necessary to confirm 527

these physiological processes are the actual reasons for the long retention of a part of thistoxin in this species.

530 The findings presented in this work put in evidence DA biotransformation in invertebrate 531 species, and strongly suggest the role of subcellular mechanisms such as early and late 532 autophagy, in the accumulation, localization and long retention of DA in some marine 533 invertebrates.

534 **5.** Conclusions

The evidence presented in this work corroborates the profound interspecific differences in the 535 536 accumulation of DA between different species of marine invertebrates, as well as species-537 specific profiles of toxin biotransformation among the analyzed species. Similar profiles of 538 DA isomers were found between P. maximus and A. opercularis, whereas C. fornicata was the species with the highest biotransformation rate, and D. trunculus the lowest. In P. 539 540 maximus, A. opercularis and C. fornicata the DA chromogenic signal was detected mainly within autophagosomic-structures in the cytoplasm of digestive cells, while in D. trunculus 541 and Asterocarpa sp. DA signal was found free in the cytoplasm of the digestive cells. This 542 543 evidence indicates that localization of DA and its effects at the subcellular level appear to be species-specific, and the integrative analysis revealed that these parameters could be 544 potentially influenced by the biotransformation profiles of the toxin. All this new information 545 is highly valuable to strengthen ASP-monitoring systems since most of the invertebrate 546 species analyzed in this work could be used as sentinels of DA contamination in affected 547 areas. Furthermore, this study provides a set of innovative histological parameters developed 548 to assess quantitatively some subcellular mechanisms potentially involved in the 549 accumulation and long-retention of DA among contaminated invertebrates. This quantitative 550 551 information may be integrated into numerical models that allow estimating and predicting toxicokinetics of contamination and depuration in fishery-stocks frequently affected during 552 553 blooms of toxic Pseudo-nitzschia sp.

554 Acknowledgments

555 The authors are grateful to Sylvain Enguehard (Novakits, Nantes) for providing the non-

commercial primary antibodies necessary to carry out this study, as well as Nicolas Chomerat

557 (from Ifremer, Concarneau) for sample transporting, and Adeline Bidault and Morgan

558 Perennou (from LEMAR, Brest) for their support during sampling and dissections. We also

thank Marie Calvez and Nelly Le Goïc (LEMAR, Brest) for their assistance with tissue

sectioning, and Carmen Rodríguez-Jaramillo (CIBNOR, La Paz) for her advices to optimize
non-commercial antibodies for the IHC analysis. Thanks to Alejandra L. Peña for English
edition.

563 **Declaration of competing interest**

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

566 **Funding**

- 567 This work received financial support from the research project "MaSCoET" (Maintien du
- 568 Stock de Coquillages en lien avec la problématique des Efflorescences Toxiques) financed by
- 569 France Filière Pêche and Brest Métropole. JLGC is recipient of a doctorate fellowship from
- 570 CONACyT, Mexico (REF: 2019- 000025-01EXTF-00067).

571 Data availability statement

- 572 The evidence and data that support the findings of this study are available from the
- 573 corresponding author upon reasonable request.

574 Ethics statements

- 575 The organisms used in this work were transported and handled according to the International
- 576 Standards for the Care and Use of Laboratory Animals. The number of sampled organisms
- 577 contemplated "the rule of maximizing information published and minimizing unnecessary
- 578 studies". In this sense, 38 individuals were considered the minimum number of organisms
- 579 needed for this work.

580 Author contributions

- 581 Conceived the study: CF, HH, JLGC. Sampling: JLGC, HH, CF, ML, TD, AT. Processed the
- samples: JLGC, TD, AD, AT. Analyzed the data: JLGC, AD. Interpretation of data: JLGC,
- 583 CF, HH, AD. Contributed reagents/materials/analysis tools: CF, HH, AD, AT, ML. Wrote the
- 584 first draft of the manuscript: JLGC. Writing review & editing: CF, HH, JLGC, ML, AD,
- 585 AT.

586 Literature cited

- Álvarez, G., Rengel, J., Araya, M., Álvarez, F., Pino, R., Uribe, E., Díaz, P.A., Rossignoli,
 A.E., López-Rivera, A., Blanco, J., 2020. Rapid domoic acid depuration in the scallop *Argopecten purpuratus* and its transfer from the digestive gland to other organs. *Toxins*,
 12, 698. <u>https://doi.org/10.3390/toxins12110698</u>.
- Amzil, Z., Fresnel, J., Le Gal, D., Billard, C., 2001. Domoic acid accumulation in French
 shellfish in relation to toxic species of *Pseudo-nitzschia multiseries* and *P. pseudodelicatissima. Toxicon*, 39(8), 1245–1251. <u>https://doi.org/10.1016/s0041-</u>
 0101(01)00096-4
- Ayache, N., Hervé, F., Martin-Jézéquel, V., Amzil, Z., Caruana, A. M. N., 2018. Influence of
 sudden salinity variation on the physiology and domoic acid production by two strains of *Pseudo-nitzschia australis*. In T. Mock (Ed.), *Journal of Phycology*, 55(1), 186–195.
 https://doi.org/10.1111/jpy.12801.
- Balbi, T., Cortese, K., Ciacci, C., Bellese, G., Vezzulli, L., Pruzzo, C., Canesi, L., 2018.
 Autophagic processes in *Mytilus galloprovincialis* hemocytes: effects of *Vibrio tapetis*. *Fish & Shellfish Immunology*. 73, 66–74. https://doi.org/10.1016/j.fsi.2017.12.003.
- 602 Baranska, A., Shawket, A., Jouve, M., Baratin, M., Malosse, C., Voluzan, O., Vu Manh, T.-603 P., Fiore, F., Bajénoff, M., Benaroch, P., Dalod, M., Malissen, M., Henri, S., Malissen, B., 2018. Unveiling skin macrophage dynamics explains both tattoo persistence and strenuous 604 removal. *Experimental* Medicine, 215(4), 1115-1133. 605 Journal of https://doi.org/10.1084/jem.20171608 606
- Basti, L., Hégaret, H., Shumway, S.E., 2018. Harmful Algal Blooms and Shellfish. In:
 Harmful Algal Blooms: A Compendium Desk Reference, First Edition. Shumway, S.E.,
 Burkholder, J.M., Morton, S.L. (eds). John Wiley & Sons Ltd.
- Bates S.S., Garrison D.L., Horner R.A., 1998. Bloom dynamics and physiology of domoicacid-producing *Pseudo-nitzschia* species. In: Physiological ecology of harmful algal
 multiseries.In: Harmful algal blooms 2000 (Ed. by G.M. Hallegraeff, S.I. Blackburn, C.J.
 Bolch & R.J. Lewis), pp. 320–323. Intergovernmental Oceanographic Commission of
 UNESCO, Paris.
- Bates, S.S., Hubbard, K.A., Lundholm, N., Montresor, M., Leaw, C.P., 2018. *Pseudo- nitzschia*, *Nitzschia*, and domoic acid: new research since 2011. *Harmful Algae*, 79, 3-43.
 https://doi.org/10.1016/j.hal.2018.06.001.

- Ben Haddouch, A., Taleb, H., Elmortaji, H., Ben Brahim, S., Ennafah, B., Menchih, K.,
 Boumaz, A., Mzaki, F., Radi, A., Loutfi, M., 2016. Accumulation and tissue distribution of
 domoic acid in the common cuttlefish, *Sepia officinalis* from the south Moroccan coast. *American Academic Scientific Research Journal for Engineering, Technology, and*
- 622 *Sciences*, 15(1), 252–264.
- Blanco, J., Acosta, C., Bermúdez de la Puente, M., Salgado, C., 2002a. Depuration and 623 anatomical distribution of the amnesic shellfish poisoning (ASP) toxin domoic acid in the 624 king scallop Pecten maximus. Aquatic Toxicology, 60 (1-2),111-121. 625 https://doi.org/10.1016/s0166-445x(01)00274-0. 626
- 627 Blanco, J., Acosta, C.P., Mariño, C., Muñiz, S., Martín, H., Moroño, A., Correa, J., Arévalo,
- F., Salgado, C., 2006. Depuration of domoic acid from different body compartments of the
- 629 king scallop *Pecten maximus* grown in raft culture and natural bed. *Aquatic Living*
- 630 *Resources*, 19 (3), 257–265. <u>https://doi.org/10.1051/alr:2006026</u>.
- Blanco, J., Bermúdez, M., Arévalo, F., Salgado, C., Moroño, A., 2002b. Depuration of
 mussels (*Mytilus galloprovincialis*) contaminated with domoic acid. *Aquatic Living Resources*, 15, 53–60. <u>https://doi.org/10.1016/S0990-7440(01)01139-1</u>.
- Blanco, J., Livramento, F., Rangel, I. M., 2010. Amnesic shellfish poisoning (ASP) toxins in
 plankton and molluscs from Luanda Bay, Angola. *Toxicon*, 55(2–3), 541–546.
 https://doi.org/10.1016/j.toxicon.2009.10.008.
- Blanco, J., Moroño, A., Arévalo, F., Correa, J., Salgado, C., Rossignoli, A., Lamas, P., 2021.
 Twenty-Five Years of Domoic Acid Monitoring in Galicia (NW Spain): Spatial, Temporal
 and Interspecific Variations. *Toxins*, 13(11):756. https://doi.org/10.3390/toxins13110756.
- Bogan, Y. M., Kennedy, D. J., Harkin, A. L., Gillespie, J., Vause, B. J., Beukers-Stewart, B.
 D., Hess, P., Slater, J.W., 2007a. Variation in domoic acid concentration in king scallop
 (*Pecten maximus*) from fishing grounds around the Isle of Man. *Harmful Algae*, 6, 81–92.
- 643 <u>https://doi.org/10.1016/j.hal.2006.07.002</u>.
- Bogan, Y., Bender, K., Hervas, A., Kennedy, D., Slater, J., Hess, P., 2007c. Spatial variability
 of domoic acid concentration in king scallops *Pecten maximus* off the southeast coast of
 Ireland. *Harmful Algae*. 6(1): 1-14. <u>https://doi.org/10.1016/j.hal.2006.05.004</u>

- Bogan, Y.M., Harkin, A.L., Gillespie, J., Kennedy, D.J., Hess, P., Slater, J.W., 2007b. The
 influence of size on domoic acid concentration in king scallop, *Pecten maximus* (L.). *Harmful Algae*, 6, 15–28. https://doi.org/10.1016/j.hal.2006.05.005.
- Chi, C., Zhang, C., Liu, J., Zheng, X., 2019. Effects of marine toxin domoic acid on innate
 immune responses in bay scallop. Journal of Marine Science and Engineering, 7(11), 407.
 https://doi.org/10.3390/jmse7110407.
- Costa, P. R., Rosa, R., Duarte-Silva, A., Brotas, V., Sampayo, M. A. M., 2005a.
 Accumulation, transformation and tissue distribution of domoic acid, the amnesic shellfish
 poisoning toxin, in the common cuttlefish, *Sepia officinalis. Aquatic Toxicology*, 74(1),
 82–91. https://doi.org/10.1016/j.aquatox.2005.01.011
- Costa, P. R., Rosa, R., Pereira, J., M. Sampayo., 2005b. Detection of domoic acid, the
 amnesic shellfish toxin, in the digestive gland of *Eledone cirrhosa* and *E. moschata*(Cephalopoda, Octopoda) from the Portuguese coast. *Aquatic Living Resources*, 18(4),
 395–400). https://doi.org/10.1051/alr:2005041.
- Costa, P. R., Rosa, R., Sampayo, M. A. M., 2004. Tissue distribution of the amnesic shellfish
 toxin, domoic acid, in *Octopus vulgaris* from the Portuguese coast. *Marine Biology*,
 144(5), 971–976. https://doi.org/10.1007/s00227-003-1258-6.
- Costa, P., Costa, M.H., 2012. Development and application of a novel histological
 multichrome technique for clam histopathology. *Journal of Invertebrate Pathology*, 110,
 411–414. https://doi.org/10.1016/j.jip.2012.04.013.
- Cuervo, A.M., 2004. Autophagy: many paths to the same end. *Molecular and Cellular Biochemistry*, 263 (1/2), 55–72. <u>https://doi.org/10.1023/b:mcbi.0000041848.57020.57</u>.
- Dizer, H., Fischer, B., Harabawy, A.S.A., Hennion, M.C., Hansen, P.D., 2001. Toxicity of
 domoic acid in the marine mussel *Mytilus edulis*. *Aquatic Toxicology*, 55: 149-156.
 https://doi.org/10.1016/s0166-445x(01)00178-3.
- Dusek Jennings, E., Parker, M. S., Simenstad, C. A., 2020. Domoic acid depuration by
 intertidal bivalves fed on toxin-producing *Pseudo-nitzschia multiseries*. *Toxicon*, 6,
 100027. https://doi.org/10.1016/j.toxcx.2020.100027.

- Flannagan, R. S., Jaumouillé, V., Grinstein, S., 2012. The Cell Biology of Phagocytosis. *Annual Review of Pathology: Mechanisms of Disease*, 7(1), 61–98.
 https://doi.org/10.1146/annurey-pathol-011811-132445.
- García-Corona, J. L., Hégaret, H., Deléglise, M., Marzari, A., Rodríguez-Jaramillo, C., 678 679 Foulon, V., Fabioux, C., 2022. First subcellular localization of the amnesic shellfish toxin, domoic acid, in bivalve tissues: Deciphering the physiological mechanisms involved in its 680 681 long-retention in the king scallop *Pecten* maximus. Harmful Algae. 116. https://doi.org/10.1016/j.hal.2022.102251. 682
- Gilgan, M.W., Burns B.G., Landry, G.J., 1990. Distribution and magnitude of domoic acid
 contamination of shellfish in Atlantic Canada. In: E. Graneli, B. Sundstrom, L. Edler, D.M.
 Anderson (Eds.), Toxic Marine Phytoplankton. Elsevier, N.Y. pp. 469- 474.
- 686 Gómez-Robles, E., Rodríguez-Jaramillo, C., Saucedo, P.E., 2005. Digital image analysis of
- 687 lipid and protein histochemical markers for measuring oocyte development and quality in
- pearl oyster *Pinctada mazatlanica* (Hanley, 1856). Journal of Shellfish Research, 24(4),
- 689 1197-1202. http://dx.doi.org/10.2983/0730-8000(2005)24[1197:DIAOLA]2.0.CO;2.
- Gordon, S., 2016. Phagocytosis: An Immunobiologic Process. *Immunity*, 44(3), 463–475.
 https://doi.org/10.1016/j.immuni.2016.02.026.
- Haya, K., Martin, J.L., Burridge, L.E., Waiwood, B.A., Wildish, J., 1991. Domoic acid in
 shellfish and plankton from the bay of Fundy, New Brunswick, Canada. *Journal of Shellfish Research*, 10, 113–118.
- Hector, A., 2015. The new statistics with R: an introduction for biologists, 1st ed. OxfordUniversity Press, New York.
- Hégaret, H., Smolowitz, R. M., Sunila, I., Shumway, S. E., Alix, J., Dixon, M., Wikfors, G.
 H., 2010. Combined effects of a parasite, QPX, and the harmful-alga, *Prorocentrum minimum* on northern quahogs, *Mercenaria mercenaria*. *Marine Environmental Research*,
- 700 69(5), 337–344. <u>https://doi.org/10.1016/j.marenvres.2009.12.008</u>.
- Horner, R.A., Kusske, M.B., Moynihan, B.P., Skinner, R.N., Wekell, J.C., 1993. Retention of
 Domoic Acid by Pacific Razor Clams, *Siliqua patula* (Dixon, 1789): Preliminary Study. *Journal of Shellfish Research*, 12, 451–456.

- Husson, B., Hernández-Fariñas, T., Le Gendre, R., Schapira, M., Chapelle, A., 2016. Two
 decades of *Pseudo-nitzschia* spp. blooms and king scallop (*Pecten maximus*)
 contamination by domoic acid along the French Atlantic and English Channel coasts:
 Seasonal dynamics, spatial heterogeneity and interannual variability. *Harmful Algae*, 51,
 26–39. https://doi.org/10.1016/j.hal.2015.10.017
- James, K.D., Gillman, M., Fernández-Amandi, M., López- Rivera, A., Fernández Puente, P.,
 Lehane, M., Mitrovic, S., Furey, A., 2005. Amnesic shellfish poisoning toxins in bivalve
 molluscs in Ireland. *Toxicon*, 46, 852–858. https://doi.org/10.1016/j.toxicon.2005.02.009.
- Jones, T.O., Whyte, J.N.C., Ginther, N.G., Townsend, L.D., Iwama, G.K., 1995. Haemocyte
 changes in the pacific oyster, *Crassostrea gigas*, caused by exposure to domoic acid in the
 diatom *Pseudo-nitzschia pungens* f. *multiseries. Toxicon*, 33 (3), 347–353.
- 715 <u>https://doi.org/10.1016/0041-0101(94)00170-</u>.
- Kim, Y., Ashton-Alcox, K.A., Powell, E.N., 2006. Histological Techniques for Marine
 Bivalve Molluscs: update. NOAA Technical Memorandum NOS NCCOS 27, Maryland.
- Klionsky, D. J., Eskelinen, E.-L., Deretic, V., 2014. Autophagosomes, phagosomes,
 autolysosomes, phagolysosomes, autophagolysosomes... Wait, I'm confused. *Autophagy*,
 10 (4), 549–551. <u>https://doi.org/10.4161/auto.28448</u>.
- Kvrgić, K., Lešić, T., Džafić, N., Pleadin, P., 2022. Occurrence and seasonal monitoring of
 domoic acid in three shellfish species from the Northern Adriatic Sea. *Toxins*, 14(1), 33;
 https://doi.org/10.3390/toxins14010033.
- La Barre, S., Bates, S.S. Quilliam, M.A., 2014. Domoic acid. In. Outstanding marine
 molecules: chemistry, biology, analysis. Edited by S. La Barre and J.-M. Kornprobst.
 Wiley-VCH Verlag GmbH & Co. KgaA, Weinheim, Germany, pp. 189–216.
- Lassudrie, M., Soudant, P., Richard, G., Henry, N., Medhioub, W., da Silva, P. M., Donval,
- A., Bunel, M., Le Goïc, N., Lambert, C., de Montaudouin, X., Fabioux, C., Hégaret, H.,
- 729 2014. Physiological responses of Manila clams Venerupis (=Ruditapes) philippinarum
- with varying parasite Perkinsus olseni burden to toxic algal Alexandrium ostenfeldii
- r31 exposure. *Aquatic Toxicology*, 154, 27–38. <u>https://doi.org/10.1016/j.aquatox.2014.05.002</u>.
- Lelong, A., Hégaret, H., Soudant, P., Bates, S.S., 2012. *Pseudo-nitzschia* (Bacillariophyceae)
 species, domoic acid and amnesic shellfish poisoning: revisiting previous paradigms. *Phycologia*, 51 (2), 168–216. https://doi.org/10.2216/11-37.1.

- Liu, H., Kelly, M.S., Campbell, D.A., Dong, S.L., Zhu, J.X., Fang, J.G., Wang, S.F., 2007a.
 Exposure to domoic acid affects larval development of king scallop *Pecten maximus*(Linnaeus, 1758). *Aquatic Toxicology*, 81, 152–158.
 https://doi.org/10.1016/j.aquatox.2006.11.012.
- Liu, H., Kelly, M.S., Campbell, D.A., Dong, S.L., Zhu, J.X., Wang, S.F., 2007b. Ingestion of
 domoic acid and its impact on king scallop *Pecten maximus* (Linnaeus, 1758). *Journal of Ocean University of China*, 6, 175–181. https://doi.org/10.1007/s11802-007-0175-6.
- Mafra, L.L., Bricelj, V.M., Fennel, K., 2010. Domoic acid uptake and elimination kinetics in
 oysters and mussels in relation to body size and anatomical distribution of toxin. *Aquatic Toxicology*, 100 (1), 17–29. https://doi.org/10.1016/j.aquatox.2010.07.0.
- Mauríz, A., Blanco, J., 2010. Distribution and linkage of domoic acid (amnesic shellfish poisoning toxins) in subcellular fractions of the digestive gland of the scallop *Pecten maximus*. *Toxicon*, 55 (2-3), 606–611. https://doi.org/10.1016/j.toxicon.2009.10.
- McMillan, D.B., Harris, R.J., 2018. The Animal Cell. In An Atlas of Comparative Vertebrate
 Histology (pp. 3–25). Elsevier. <u>https://doi.org/10.1016/b978-0-12-410424-2.00001-9</u>.
- Moore, M.N., 2004. Diet restriction induced autophagy: a lysosomal protective system against
 oxidative- and pollutant-stress and cell injury. *Marine Environmental Research*, 58 (2–5),
- 752 603–607. <u>https://doi.org/10.1016/j.marenvres.2004.03</u>.
- Moore, M.N., 2008. Autophagy as a second level protective process in conferring resistance
 to environmentally-induced oxidative stress. *Autophagy*, 4 (2), 254–256.
 <u>https://doi.org/10.4161/auto.5528</u>.
- Novaczek, I., Madhyastha, M.S., Ablett, R.F., Donald, A., Johnson, G., Nijjar, M.S., Sims,
 D.E., 1992. Depuration of domoic acid from live blue mussels (*Mytilus edulis*). *Canadian Journal of Fisheries and Aquatic Sciences*, 49 (2), 312–318. <u>https://doi.org/10.1139/f92-</u>
 035.
- O'Dea, S.N., 2012. Occurrence, Toxicity, and Diversity of *Pseudo-nitzschia* in Florida
 Coastal Waters. University of South Florida ProQuest Dissertations Publishing. 1515837.
- Pazos, A.J., Ventoso, P., Martínez-Escauriaza, R., Pérez-Parallé, M.L., Blanco, J., Triviño,
 J.C., Sánchez, J.L., 2017. Transcriptional response after exposure to domoic acid-

- producing *Pseudo-nitzschia* in the digestive gland of the mussel *Mytilus galloprovincialis*.
 Toxicon, 140, 60–71. https://doi.org/10.1016/j.toxicon.2017.10.002.
- Perl, T.M., Bédard, L., Kosatsky, T., Hockin, J.C., Todd, E.C.D., Remis, R.S., 1990. An
 Outbreak of Toxic Encephalopathy Caused by Eating Mussels Contaminated with Domoic
 Acid. *N. Engl. J. Med.* 322(25): 1775–1780. https://doi:10.1056/NEJM199006213222504.
- 769 Picot, S., Morga, B., Faury, N., Chollet, B., Dégremont, L., Travers, M.A., Renault, T., Arzul,
- I., 2019. A study of autophagy in hemocytes of the Pacific oyster Crassostrea gigas.
- 771 *Autophagy*, 1–9. <u>https://doi.org/10.1080/15548627.2019.1596</u>.
- Pulido, O.M., 2008. Domoic Acid Toxicologic Pathology: A Review. *Marine Drugs*, 6, 180219. https://doi.org/10.3390/md20080010.
- Quilliam, M.A., Sim, P.G., McCulloch, A.W., McInnes, A.G., 1989. High-performance liquid
 chromatography of domoic acid, a marine neurotoxin, with application to shellfish and
 plankton. *International Journal of Environmental Analytical Chemistry*, 36 (3), 139–154.
 https://doi.org/10.1080/03067318908026867.
- R Core Team (2020). R: a language and environment for statistical computing. R Foundation
 for Statistical Computing, Vienna, Austria. URL <u>https://www.R-project.org/</u>.
- Rodríguez-Jaramillo, C., García-Corona, J.L., Zenteno-Savín, T., Palacios, E., 2022. The
 effects of experimental temperature increase on gametogenesis and heat stress parameters
 in oysters: Comparison of a temperate-introduced species (*Crassostrea gigas*) and a native
- tropical species (*Crassostrea corteziensis*). Aquaculture, 561, 738683.
 <u>https://doi.org/10.1016/j.aquaculture.2022.738683</u>.
- Shubin, A. V., Demidyuk, I. V., Komissarov, A. A., Rafieva, L. M., & Kostrov, S. V., 2016.
 Cytoplasmic vacuolization in cell death and survival. *Oncotarget*, 7(34), 55863–55889.
 https://doi.org/10.18632/oncotarget.10150.
- 788 Silvert, W. and Subba R.D.V., 1992. Dynamic model of the flux of domoic acid, a neurotoxin,
- through a *Mytilus edulis* population. *Canadian Journal of Fisheries and Aquatic Sciences*,
 49, 400- 405. <u>https://doi.org/10.1139/f92-045</u>.
- 791 Takata, Y., Sato, S., Ha, D. V., Montojo, U. M., Lirdwitayaprasit, T., Kamolsiripichaiporn, S.,
- 792 Kotaki, Y., Fukuyo, Y., Kodama, M., 2009. Occurrence of domoic acid in tropical
- 793 bivalves. *Fisheries Science*, 75(2), 473–480. <u>https://doi.org/10.1007/s12562-009-0073-5</u>.

- Trainer, V.L., Bates, S.S., Lundholm, N., Thessen, A.E., Cochlan, W.P., Adams, N.G., 2012. *Pseudo-nitzschia* physiological ecology, phylogeny, toxicity, monitoring and impacts on
 ecosystem health. *Harmful Algae*, 14, 271–300. <u>https://doi.org/10.1016/j.hal.2011.10.025</u>.
- Trainer, V.L., Bill, B.D., 2004. Characterization of a domoic acid binding site from Pacific
 razor clam. *Aquatic Toxicology*, 69, 125–132.
 https://doi.org/10.1016/j.aquatox.2004.04.012.
- Vale, P., Sampayo, M.A.M., 2001. Domoic acid in Portuguese shellfish and fish. *Toxicon*,
 39:893–904. https://doi.org/10.1016/s0041-0101(00)00229-4.
- 802 Ventoso, P., Pazos, A.J., Blanco, J., Pérez-Parallé, M.L., Triviño, J.C., Sánchez, J.L., 2021. Transcriptional Response in the Digestive Gland of the King Scallop (Pecten maximus) 803 804 After the Injection of Domoic Acid. Toxins, 13. 339. 805 https://doi.org/10.3390/toxins13050339.
- Ventoso, P., Pazos, A.J., Pérez-Parallé, M.L., Blanco, J., Triviño, J.C., Sánchez, J.L., 2019.
 RNA-Seq Transcriptome Profiling of the Queen Scallop (*Aequipecten Opercularis*)
 Digestive Gland after Exposure to Domoic Acid-Producing *Pseudo-nitzschia. Toxins*, 11,
 97. https://doi.org/10.3390/toxins11020097.
- Wang, L., Ye, X., Zhao, T., 2019. The physiological roles of autophagy in the mammalian life
 cycle. *Biological Reviews*, 94, 503–516. <u>https://doi.org/10.1111/brv.12464</u>.
- 812 Wright, J.L., Bird, C.J., de Freitas, A.S., Hampson, D., McDonald, J., Quilliam, M.A., 1990a.
- Chemistry, biology, and toxicology of domoic acid and its isomers. Canada Diseases
 Weekly Report = Rapport Hebdomadaire des Maladies au Canada. Pp. 21-26.
- 815 Wright, J.L.C., Falk, M., McInnes, A.G., Walter, J.A., 1990b. Identification of isodomoic acid
- D and two new geometrical isomers of domoic acid in toxic mussels. *Canadian Journal of Chemistry*, 68(1), 22–25. https://doi.org/10.1139/v90-005.
- 818 Zar, J. H., 2010. Biostatistical Analysis. 5th Ed. Pearson, Westlake Village, CA, 251 pp.
- Zhao, Y.G., Codogno, P., Zhang, H., 2021. Machinery, regulation and pathophysiological
 implications of autophagosome maturation. *Nature Reviews Molecular Cell Biology*.
 https://doi.org/10.1038/s41580-021-00392-4.
- Zheng, G., Wu, H., Guo, M., Peng, J., Zhai, Y., Tan, Z., 2022. First observation of domoic
- acid and its isomers in shellfish samples from Shandong Province, China. Journal of

824 *Oceanology and Limnology*, 40(6), 2231–2241. <u>https://doi.org/10.1007/s00343-022-2104-</u>

825 <u>3</u>.

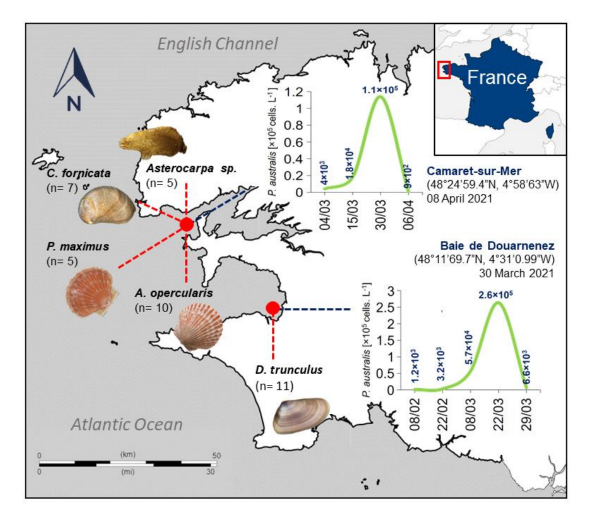
Table 1. Relative abundance of DA and its isomers in the digestive glands of the scallops *P. maximus* (n =5) and *A. opercularis* (n =10), the clam

827 *D. trunculus* (n =11), the slippersnail *C. fornicata* (n =7), and the sea squirt *Asterocarpa* sp. (n =5) contaminated during *P. australis* blooms in 828 the northwest coast of Brittany, France between March-April 2021.

	-	Statistical analysis					
	P. maximus	A. opercularis	D. trunculus	C. fornicata	Asterocarpa sp.	Statistical analysis	
DA (%)	93.3 ± 0.6^{b}	93.6 ± 0.3^{b}	$95.8\pm0.3^{\rm a}$	90.7 ± 1.1^{c}	94.5 ± 0.1^{ab}	$F_{(df=4,33)} = 11.8, P < 0.0001$	
isoE (%)	$4.3\pm0.3^{\rm a}$	4.3 ± 0.3^{a}	$3.5\pm0.3^{\mathrm{a}}$	$3.2\pm0.4^{\mathrm{a}}$	1.6 ± 0.1^{b}	$F_{(df=4,33)} = 10.9, P < 0.0001$	
isoD (%)	$1.5 \pm 0.3^{\rm bc}$	$1 \pm 0.1^{\mathrm{bc}}$	$0.5\pm0.1^{ m c}$	$4\pm0.8^{\mathrm{a}}$	$2.1\pm0.0^{ m b}$	$F_{(df=4,33)} = 17.3, P < 0.0001$	
isoA (%)	$0.4\pm0.0^{ m ab}$	$0.7\pm0.0^{\mathrm{a}}$	$0.2\pm0.0^{ m b}$	0.6 ± 0.1^{a}	$0.5\pm0.0^{\mathrm{a}}$	$F_{(df=4,33)} = 10.4, P < 0.0001$	
epi-DA (%)	$0.4\pm0.1^{ m b}$	$0.4 \pm 0^{\circ}0^{\mathrm{b}}$	$0\pm0.0^{ m c}$	1.5 ± 0.1^{a}	1.3 ± 0.0^{a}	$F_{(df=4,33)} = 156.4, P < 0.0001$	

Results are expressed as mean \pm SE. Data were analyzed using species (five levels) as factor in separate one-way ANOVAs (P < 0.05). The F-test statistic and degrees of freedom (df) are reported. Different superscript letters indicate significant differences between species. The level of

831	statistical	significance	was	set	at	α	=0.05.
-----	-------------	--------------	-----	-----	----	---	--------



832

Figure 1. Sampling sites of the scallops *P. maximus* (n =5) and *A. opercularis* (n = 10), the clam *D. trunculus* (n =11), the slippersnail *C. fornicata* (n =7), and the sea squirt *Asterocarpa sp.* (n =5) and cell densities (cells. L^{-1}) of *P. australis* during toxic blooms in the northwest coast of Brittany, France between February and-April 2021.

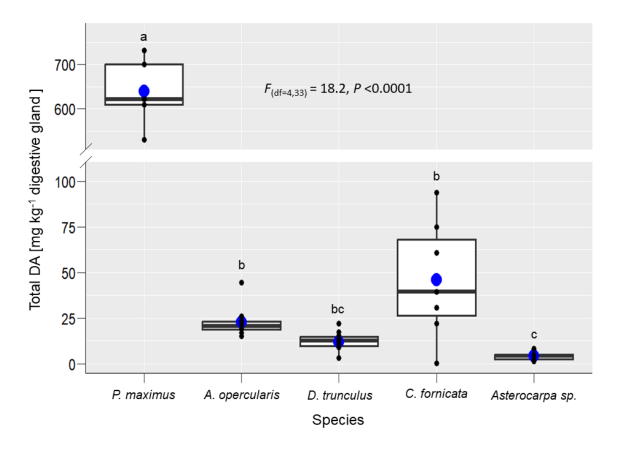


Figure 2. Total domoic acid (tDA) concentration in the digestive glands of the scallops P. 838 *maximus* (n = 5) and *A. opercularis* (n = 10), the clam *D. trunculus* (n = 11), the slippersnail *C*. 839 fornicata (n =7), and the sea squirt Asterocarpa sp. (n =5) contaminated during P. australis 840 blooms in the northwest coast of Brittany, France between on the 30th of March (for the 841 scallops P. maximus, A. opercularis, the slippersnail C. fornicata, and the sea squirt 842 Asterocarpa sp.) and on the 8th of April, 2021 (for the clam D. trunculus). The upper and 843 lower limits of the boxes are the quartiles, the middle horizontal line is the median, the 844 845 extremes of the vertical lines are the upper and lower limits of the observations, and black dots are the individual observations. The blue dots are the means for each species. Data were 846 847 analyzed using species (five levels) as factor using a one-way ANOVA (P < 0.05). The F-test statistic and degrees of freedom (df) are reported. Different superscript letters indicate 848 significant differences between species. The level of statistical significance was set at $\alpha = 0.05$. 849

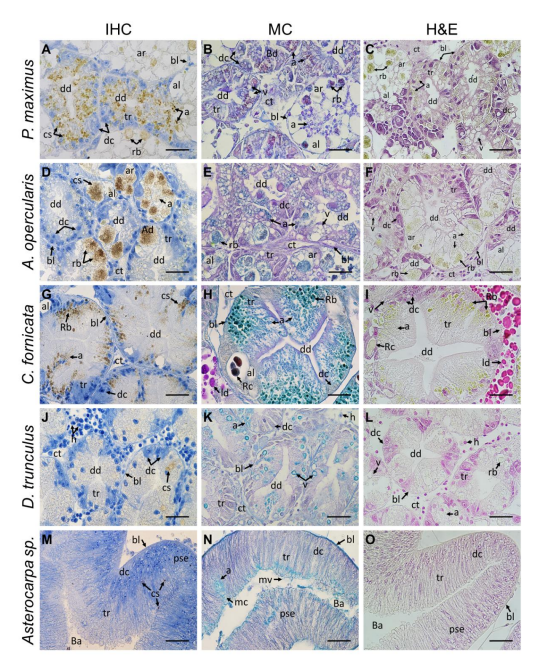


Figure 3. Microphotographs of digestive glands of the scallops P. maximus (A, B, C), A. 852 opercularis (D, E, F), the slippersnail C. fornicata (G, H, I), the clam D. trunculus (J, K, L), 853 and the sea squirt Asterocarpa sp. (M, N, O) contaminated with domoic acid (DA) during P. 854 australis blooms in the northwest coast of Brittany, France in March-April, 2021. IHC (A, D, 855 856 G, J, M) = Immunohistochemical detection of DA using specific anti-DA antibody (0.08 mg. mL^{-1}); MC (B, E, H, K, N) = multichromic histochemical staining of neutral carbohydrates 857 (violet-magenta dyes), acid glycoconjugates (blue hues), and proteins (yellowish tones); H&E 858 (C, F, I, L, O) = conventional histological Hematoxylin-Eosin staining. a = autophagosomic-859 like vesicles, al = adipocyte-like cell, ar = acinar region, Ba = blind ampulla, bl = basal860 lamina, cs = DA chromogenic signal, ct = connective tissue, dc = digestive cells, dd =861 digestive diverticulum, hc = hemocytes, ld = lipid droplets, mc = mucus, mv = microvilli, pse 862 = pseudostratified epithelium, rb = residual bodies, rc = residual concretions, tr = tubular 863 region, v = vacuoles. Scale bar: $63 \times = 30 \ \mu m$. 864

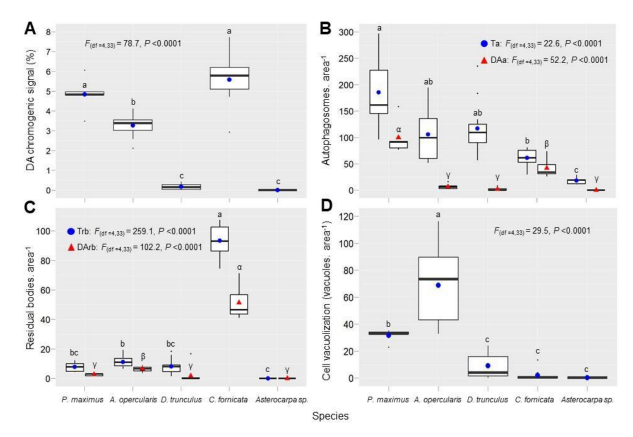


Figure 4. Quantitative analysis of DA localization and subcellular features in the digestive 866 glands of the scallops P. maximus (n = 5) and A. opercularis (n = 10), the clam D. trunculus (n = 10)867 =11), the slippersnail C. fornicata (n = 7) and the sea squirt Asterocarpa sp. (n = 5)868 contaminated with DA during P. australis blooms in the northwest coast of Brittany, France, 869 in March-April, 2021. (A) DA chromogenic signal (%); (B) Autophagy (autophagosomes. 1.3 870 mm^2 , Ta = total autophagy, DAa = DA autophagy); (C) Residual bodies (residual bodies. 1.3) 871 mm^2 , Trb = total residual bodies, DArb = DA in the residual bodies); (D) Cell vacuolization 872 (vacuoles. 1.3 mm²). The upper and lower limits of the boxes are the quartiles, the middle 873 874 horizontal line is the median, the extremes of the vertical lines are the upper and lower limits 875 of the observations, and black dots are the outliers (values that deviate from the median more than 1.5 times the interquartile range). The blue dots and red triangles are the means of each 876 variable. Data were analyzed using species (five levels) as factor in separate one-way 877 ANOVA's (P < 0.05). The F-test statistic and degrees of freedom (df) are reported. Different 878 superscript letters indicate significant differences between species. The level of statistical 879 significance was set at $\alpha = 0.05$. 880

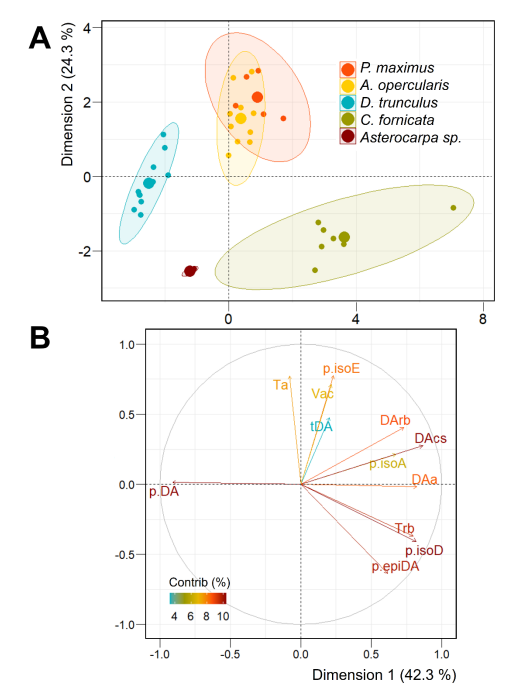


Figure 5. Principal component analysis (PCA) summarizing data from the scallops P. 882 *maximus* (n = 5) and *A. opercularis* (n = 10), the clam *D. trunculus* (n = 11), the slippersnail *C*. 883 fornicata (n =7), and the sea squirt Asterocarpa sp. (n =5) contaminated with domoic acid 884 (DA) during P. australis blooms in the northwest coast of Brittany, France, between March-885 April 2021. Dimension 1 and dimension 2 together describe 66.6 % of the total variance. (A) 886 887 Scatter plot of individuals from each species. Larger symbols are the barycenter of each group, confidence ellipses level was fixed at $\alpha = 0.05$. (B) Variable contribution plot. The 888 direction of the arrows shows the correlations of variables (tDA = total DA, DAcs = DA889 890 chromogenic signal, Ta = total autophagy, DAa = DA autophagy (%), Trb = total residual bodies, DArb = DA in the residual bodies (%), Vac = cell vacuolization, and the percentages 891 (p) of DA isomers, p.DA = untransformed DA, p.isoE = isoE, p.isoD = isoD, p.isoA = isoA, 892

- 893 p.epiDA = epiDA) with given PCs, and its color intensity shows their contribution (Contrib
- 894 %) to the explained variance.