

Heterogeneous microgeographic genetic structure of the common cockle (*Cerastoderma edule*) in the Northeast Atlantic Ocean: biogeographic barriers and environmental factors

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9 Heterogeneous microgeographic genetic structure of the common cockle (*Cerastoderma*
10 *edule*) in the Northeast Atlantic Ocean: biogeographic barriers and environmental
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39 **Abstract**

40 Knowledge of genetic structure at the finest level is essential for conservation of
41 genetic resources. Despite no visible barriers limiting gene flow, significant genetic
42 structure has been shown in marine species. The common cockle (*Cerastoderma edule*)
43 is a bivalve of great commercial and ecological value inhabiting the Northeast Atlantic
44 Ocean. Previous population genomics studies demonstrated significant structure both
45 across the Northeast Atlantic, but also within small geographic areas, highlighting the
46 need to investigate fine-scale structuring. Here, we analysed two geographic areas that
47 could represent opposite models of structure for the species: 1) the SW British Isles
48 region, highly fragmented due to biogeographic barriers, and 2) Galicia (NW Spain), a
49 putative homogeneous region. 9,250 SNPs genotyped by 2b-RAD on 599 individuals
50 from 22 natural beds were used for the analysis. The entire SNP dataset mostly
51 confirmed previous observations related to genetic diversity and differentiation,
52 however, neutral and divergent SNP outlier datasets enabled disentangling physical
53 barriers from abiotic environmental factors structuring both regions. While Galicia
54 showed a homogeneous structure, the SW British Isles region was split into four
55 reliable genetic regions related to oceanographic features and abiotic factors, such as
56 sea surface salinity and temperature. The information gathered supports specific
57 management policies of cockle resources in SW British and Galician regions also
58 considering their particular socio-economic characteristics; further, these new data will
59 be added to those recently reported in the Northeast Atlantic to define sustainable
60 management actions across the whole distribution range of the species.

61 **Introduction**

62 Knowledge of genetic diversity distribution is crucial for the sustainable management
63 and conservation of natural resources (Leary et al. 2009; Sa-Pinto et al. 2012). This
64 distribution is affected by larval connectivity, demographic parameters and selective
65 processes operating on species populations. Scarcity of physical barriers in marine
66 environments is expected to promote higher connectivity among populations in
67 comparison to terrestrial species (Waples 1998). Moreover, marine species usually
68 show large population sizes, which along with pelagic larval stages, often lasting
69 several weeks, facilitate population genetic homogenization across wide regions (Sa-
70 Pinto et al. 2012; do Prado et al. 2018). Despite these general features, genetic studies
71 on marine organisms have frequently detected genetic differentiation, even at local
72 scales (i.e., below the geographic scale of effective dispersal of the species studied,
73 known as chaotic genetic patchiness (CGP); see Eldon et al. 2016), which can be
74 explained by historical and reproductive/demographic factors (e.g. high fecundity and
75 high mortality in early life stages, sweepstakes reproductive success; see Parrondo et al.
76 2022), natural selection associated with environmental conditions (Vilas et al. 2015; do
77 Prado et al. 2018; Vera et al. 2019) and oceanic features such as residual currents,
78 bathymetry, coastline shape, upwelling, fronts, gyres and eddies (Vera et al. 2016;
79 Coscia et al. 2020; Handal et al. 2020; Fisher et al. 2022; Vera et al. 2022).

80 Different types of ocean fronts have been described across the Northeast Atlantic
81 region, encompassing tidal mixing fronts, shelf break fronts, and freshwater fronts
82 separating estuarine freshwater and higher salinity coastal waters (Sharples and
83 Simpson 2019). Examples of these frontal systems on the NW European Shelf include
84 the Celtic Sea Front (NE Celtic Sea), the Irish Sea Front (NW Irish Sea), the Alderney
85 Race (with one of the strongest current in Europe) and the Ushant Front (W English

86 Channel) (Suberg et al. 2019). These fronts may influence genetic structure acting as
87 barriers to cross-front planktonic dispersal and as conduits through along-front dispersal
88 by frontal jets, with important influences on the pelagic distribution of larvae of marine
89 species (Galarza et al. 2009). Biogeographical barriers can also limit dispersal in marine
90 environments. In the Northeast Atlantic region, Cape Finisterre, the Cornwall
91 Peninsula, the tip of Brittany, the Llyn Peninsula, and the Alderney race along Cotentin
92 Peninsula have been identified as potential barriers to the connectivity of marine
93 organisms due to their oceanographic features, including fish (Abaunza et al. 2008;
94 Larmuseau et al. 2009) and molluscs (Dupont et al. 2007; Piñeira et al. 2008; Martinez
95 et al. 2015; Handal et al. 2020; Vera et al. 2022).

96 The common cockle, *Cerastoderma edule*, is a bivalve mollusc naturally distributed
97 throughout the Northeast Atlantic coast, from Senegal, West Africa, to Norway,
98 northern Europe, where it inhabits on intertidal and shallow subtidal soft sediments
99 (Hayward and Ryland 1995). The species is commercially exploited and provides a
100 wealth of services to coastal communities mainly in Ireland, United Kingdom, France,
101 Spain and Portugal, where it is harvested (Flach and de Bruin 1994; Carss et al. 2020;
102 Jackson-Bue et al. 2022). Cockle harvest has been reduced since the 1980s (> 100,000
103 tonnes) to nowadays (~ 25,000 tonnes in 2019) due to changes in fisheries policies,
104 overfishing, variable recruitment and mass mortalities produced by pollution, climate
105 events and parasites (Villalba et al. 2014; Mahony et al. 2020; Pampin et al. 2023).

106 Furthermore, cockles are considered keystone for ecosystem due to their role as reef
107 engineers, agents of carbon sequestration and their linking between primary producers
108 and higher trophic levels (Norris et al. 1998; Carss et al. 2020). The species is dioecious
109 and can live up to 10 years displaying fast sexual maturation (reached in the first year of
110 life) and high fecundity (Honkoop and van der Meer 1998). The reproductive period

111 occurs from April to August (Malham et al. 2012), but it can be extended to September
112 in more southern European countries such as Portugal (Mahony et al. 2021), and
113 planktonic larvae can remain in the water column for 30 days facilitating widespread
114 dispersal (de Montaudouin et al. 2003; Dare et al. 2004).

115 Genetic studies throughout the natural cockle's distribution have identified three main
116 population genetic units: i) a southern group encompassing the Atlantic coast from
117 Morocco to the Bay of Biscay; ii) a central group comprising of the Celtic and Irish
118 Seas, the English Channel and the southern North Sea; and iii) a northern group
119 consisting of the northern North Sea (Beaumont et al. 1980; Hummel et al. 1994;
120 Martínez et al. 2013; 2015). These results have been recently confirmed by Vera et al.
121 (2022) through a wide genome scan (~10,000 single nucleotide polymorphisms, SNPs),
122 but additionally enabled identifying substructure within the main genetic groups using
123 outlier loci under divergent selection, mostly in accordance with residual current
124 patterns and environmental variables.

125 However, due to the limited number of markers and/or the scale of sample collection, a
126 comprehensive picture of population connectivity in the common cockle is still
127 incomplete. Information at the microgeographic level, always considering the dispersal
128 capacity of the species (Eldon et al. 2016; Vera et al. 2022), is relevant for the
129 management of fisheries (Bernatchez et al. 2017). Using a wide SNP genomic
130 screening, Coscia et al. (2020) identified three genetic clusters (global $F_{ST} = 0.021$) of
131 cockles in the Celtic and Irish seas and that could be associated with residual ocean
132 currents, salinity and geographical proximity using information on larval dispersal.

133 This study aimed to analyse the genetic structure of the common cockle at a
134 microgeographic scale using 2b Restriction Associated DNA sequencing (2b-RADseq).
135 Two regions were investigated: (1) the SW British Isles and the English Channel,

136 characterised by putative habitat fragmentation due to tidal mixing fronts and
137 biogeographical barriers; and (2) the Northwest coast of Spain (Galicia), representing a
138 quite homogeneous region according to previous information on other mollusc species
139 (Diz and Presa 2009; Vera et al. 2016). The results confirmed the significant
140 differentiation of cockles' populations at microgeographic scale, but also the power of
141 larval dispersal to homogenize rather wide coastal areas, thus providing essential
142 information for proper management of this valuable resource.

143

144 **Material and methods**

145 *Sample area and oceanography*

146 Two geographic areas along the Northeast Atlantic coast were investigated (Fig. 1).
147 The first was focused on the British Isles and English Channel (hereafter called the SW
148 British Isles region), where previous, though incomplete information, supported
149 significant genetic sub-structuring (e.g. Coscia et al. 2020; Vera et al. 2022). The
150 second area was Galicia (Northwest Spain), which may be genetically homogeneous
151 according to information in other mollusc species (Diz and Presa 2009; Vera et al.
152 2016).

153 Over the cockle reproductive season (May to September; Mahony et al. 2020), the
154 coastline of Galicia is characterised by wind-driven upwelling of cold waters resulting
155 in sea surface temperatures (SSTs) that are several degrees colder than off-shore SSTs
156 (Supplementary Fig. 1b). Also driven by the predominantly northerly winds in the
157 summer months, the Portugal coastal current transports waters southwards along the
158 coastline of Iberia (Teles-Machado et al. 2016) with residual current strengths along the
159 Galician coastline exceeding 0.15 m/s (Supplementary Fig. 1d). The SW British Isles
160 region is divided into distinct oceanographic regions (the English Channel, the Celtic

161 Deep, the Celtic Sea and the Irish Sea) by diverging current or seasonal frontal systems
162 (Galparsoro et al. 2014). Several tidal mixing fronts separate seasonally stratified and
163 mixed waters (Supplementary Fig. 1a): the Ushant Front (Group “Grepma”, 1988), the
164 Celtic Sea Front, and the Irish Sea Front (Simpson and Pingree, 1978). The Celtic Sea
165 is characterised by northward flow along the western coast of Cornwall which merges
166 into the Celtic Sea Front jet and links into the Irish Coastal Current which transports
167 water clockwise along the south and west coast of Ireland (Supplementary Fig. 1c;
168 Brown et al. 2003; Fernand et al. 2006). Northward currents along the Ushant Front
169 link the American Shelf with the Celtic Sea. The southern English Channel coast is
170 dominated by northeastward flow, with the strongest currents occurring around the
171 Cotentin Peninsula.

172

173 *Sample collection*

174 A total of 374 cockles from 14 wild natural beds were collected across the
175 aforementioned two regions in the period 2017-2020 and stored in 100% ethanol for
176 analyses (Table 1). Additionally, 231 cockles from eight beds previously analysed
177 (Vera et al. 2022: identified as IDA_18, IDC_18, WDE_17, WBY_17, FBS_17,
178 FAR_17, SNO_17 and SLO_17, where 17 and 18 in the codes represent 2017 and
179 2018, respectively) were included in the analysis to achieve a comprehensive picture of
180 the areas studied, thus providing an overall total of 605 cockles. To avoid generation
181 overlapping, all samples belonged to the 0+ year age class of their sampling year. No
182 temporal replicates were included considering the temporal genetic stability previously
183 reported by Vera et al. (2022).

184

185 *Single Nucleotide Polymorphism (SNP) genotyping*

186 Total DNA was extracted from gills using the E.Z.N.A. E-96 mollusc DNA kit
187 (OMEGA Bio-tek), following manufacturer recommendations. 2b-RAD libraries (~ 90
188 cockles per run) were constructed using the Alfl Iib restriction enzyme and sequenced
189 in an Illumina NextSeq 500 platform following Maroso et al. (2018; 2019). Bowtie
190 1.1.2 (Langmead et al. 2009) was used to align reads to the cockle's genome (Bruzos et
191 al. 2022) allowing a maximum of three mismatches and a unique valid alignment (-v 3 -
192 m 1). The reference-based mode with default parameters in the gstacks module of
193 STACKS 2.0 (Catchen et al. 2013) was used for SNP calling. For genotyping, SNPs
194 were filtered following Vera et al. (2022): *i*) SNPs genotyped in > 60% individuals; *ii*)
195 MAC (minimum allele count) ≥ 3 ; *iii*) conformance to Hardy-Weinberg expectations
196 (i.e. SNPs with significant F_{IS} values ($P < 0.05$) in at least 25% of the populations were
197 removed); and *iv*) the most polymorphic SNP within each RAD-tag was retained.
198 Individuals with less than 250,000 reads were discarded.

199

200 *Genetic diversity and population structure*

201 Estimates of genetic diversity (i.e. mean number of alleles per locus (N_a), observed
202 (H_o) and expected (H_e) heterozygosity, proportion of polymorphic loci), departure
203 from Hardy-Weinberg equilibrium (HWE) and inbreeding coefficients (F_{IS}) were
204 estimated using GENEPOP v4.0 (Rousset 2008) and ARLEQUIN v3.5 (Excoffier and
205 Lischer 2010). Because a minimum allele frequency (MAF) filtering was not applied,
206 ARLEQUIN was also used to estimate H_o , H_e and F_{IS} exclusively with polymorphic
207 loci (Minimum Allele Frequency (MAF) > 0.017 according to sample size) for
208 comparison with previous studies.

209 Global and pairwise coefficients of population differentiation (F_{ST}) between cockle
210 beds were calculated with ARLEQUIN v3.5 using 10,000 permutations to test for

211 significance. The variational Bayesian clustering method implemented in the package
212 fastSTRUCTURE v2.3.4 (Raj et al. 2014) was used to estimate the number of genetic
213 population units (K) in the whole studied area and in each region testing from K = 1 to
214 K = number of beds + 1, with an admixture ancestry model, convergence criterion of 1
215 $\times 10^{-8}$, five cross-validated sets and the simple prior (flat-beta prior). The most likely
216 number of K was estimated using the “chooseK.py” program included in the
217 fastSTRUCTURE which gives the best K value and the K corresponding with weak
218 population structure in the data using heuristic scores. Summarised outputs were carried
219 out using the software POPHELPER (Francis 2017). Discriminant analyses of principal
220 components (DAPC) were run in ADEGENET package (Jombart et al. 2010; Jombart
221 and Ahmed 2011) for the R platform (R Development Core Team, 2014; [http://www.r-](http://www.r-project.org)
222 [project.org](http://www.r-project.org)) with the whole dataset and for each region. Data were transformed using
223 PCA (Principal Component Analysis) and the optimal number of principal components
224 (PC) was calculated using the `optim.a.score()` command (see Miller et al. 2020).
225 Isolation by distance (IBD) was checked by the correlation between geographical
226 (measured as the shortest oceanic distance between two beds in Km) and genetic
227 distance (measured as $F_{ST}/1-F_{ST}$; Rousset 1997) matrices using a Mantel test with
228 10,000 permutations using NTSYS v.2.1 (Rohlf 1993).

229

230 *Outlier tests*

231 The Bayesian F_{ST} -based method implemented in BAYESCAN v2.1 (Foll and
232 Gaggiotti 2008) was used to identify outlier loci subjected to selection. BAYESCAN
233 was run using default parameters (i.e. 20 pilot runs; prior odds value of 10; 100,000
234 iterations; burn-in of 50,000 iterations and a sample size of 5,000, hereafter “BY10”),
235 but we also explored increasing prior odds value to 1000 (hereafter “BY1000”). Despite

236 high prior odds tend to remove false positives, they also reduce the power for detection
237 loci under selection (Foll 2012). Loci with a False Discovery Rate (FDR, q-value) <
238 0.05 were considered as outliers. Moreover, the principal components-based method
239 implemented in R package PCADAPT v4.0 (Luu et al. 2017; Prive et al. 2020) was also
240 applied. This method renders low false-positive rates and uses individual information,
241 not requiring *a priori* population assignment. For the analysis, the number of principal
242 components (PC) retained was performed with the “chooseK” option. The outlier
243 identification was carried out with an FDR < 0.05. We considered as outliers those loci
244 identified by any of the two approaches, but additionally those shared between all
245 approaches as the most confident ones.

246

247 *Seascape analyses*

248 Effects of spatial (latitude and longitude) and relevant abiotic factors in coastal and
249 marine environments (sea surface temperature (SST, °C); sea bottom temperature (SBT,
250 °C); sea surface salinity (SSS, psu); sea bottom salinity (SBS, psu); bottom shear stress
251 (BSS, N·m⁻²); net primary productivity (NPP, mg·m⁻³·day⁻¹); see Coscia et al. 2020 and
252 Vera et al. 2022) shaping genetic differentiation across beds in the studied areas were
253 assessed using a canonical redundancy analysis (RDA) implemented in the VEGAN
254 software (Oksanen 2015) in R. This abiotic information was retrieved as monthly
255 averages from the IBI_REANALYSIS_PHYS_005_002 ocean reanalysis model
256 (https://resources.marine.copernicus.eu/?option=com_csw&task=results?option=com_csw&view=details&product_id=IBI_REANALYSIS_PHYS_005_002) and
257 IBI_REANALYSIS_BIO_005_003 model
258 (https://resources.marine.copernicus.eu/?option=com_csw&task=results?option=com_csw&view=details&product_id=IBI_REANALYSIS_BIO_005_003) for the period
260

261 2014-2018 (Supplementary Table 1), respectively. The nearest model cell classified as
262 ocean was selected to extract the data (average distance between the sampling location
263 and centre of the nearest model grid cell edge = 11.6 km). Then, averages for the
264 spawning season (i.e., from April to September, see Malham et al. 2012; Mahony et al.
265 2020), winter (i.e., from January to March) and summer (i.e., from July to September),
266 were calculated for each bed. Allele frequencies were calculated for each bed with
267 ADEGENET package using the “makefreq” option. Loci with missing values were
268 removed from the analysis. The significance of the variance associated to the different
269 variables was tested with 1,000 random permutations. Variance inflation factor (VIF)
270 was estimated to explore collinearity (correlation) between seascape variables in the
271 dataset, with VIF values > 10 suggesting important collinearity problems (Marquardt,
272 1970). The selection model was performed using automatic stepwise model building
273 algorithm based on permutation p-values tests. This procedure was performed with the
274 *ordistep* function included in VEGAN. The reduced panel of explanatory variables was
275 used to recalculate the total proportion of genetic variation in the variance partitioning.
276 The weight of the different loci on the significant environmental vectors was calculated
277 using VEGAN. All these analyses were performed separately for the whole, neutral and
278 divergent outlier SNP datasets in the regions studied.

279 Potential correlations between allele frequencies and seascape variables were
280 investigated with BAYENV2 (Coop et al. 2010; Gunther and Coop 2013) and results
281 were compared with the mentioned RDA analyses. The method implemented in this
282 software allows controlling the neutral genetic structure, because the fit improvement
283 for a given genetic variant between a model including the environmental factor and a
284 model including only neutral genetic structure is tested (Rellstab et al. 2015).

285 BAYENV2 was carried out using the whole SNP datasets from SW British Isles and

286 Galicia, respectively. First, analyses were performed with 100,000 iterations across five
287 independent runs to obtain the average covariance matrix for each subset. Secondly, the
288 correlation between each SNP and the different variables was calculated using 100,000
289 iterations to obtain Bayes factors (BF). As in the previous step, five independent runs
290 were used. Only SNPs with a BF > 10 and Spearman's coefficient (ρ) thresholds >
291 1% for any variable in all runs were considered as a well-supported environment-
292 associated SNPs. Finally, significantly correlated SNPs were compared with the outliers
293 identified in the BAYESCAN and PCADAPT analyses.

294

295 *Gene mining and functional enrichment*

296 RAD-tags including divergent outlier SNPs were mapped in the *C. edule* genome (Bruzos
297 et al. 2022) and their position compared with the consistent genomic windows under
298 divergent selection previously reported by Vera et al. (2022) in the Northeast Atlantic
299 Ocean. The very low genetic differentiation with neutral markers in the studied areas
300 precluded the detection of consistent genomic regions under stabilizing selection. Thus,
301 we could verify in more restricted geographical scenarios (SW British Isles and Galicia)
302 the consistency of the genomic regions under divergent selection previously detected.
303 Additionally, we looked for new regions under selection considering the singularity of
304 the new sample collections of this study following a similar methodology to that proposed
305 by Vera et al. (2022). Briefly, we defined a consistent window when ≥ 2 consecutive
306 outliers were detected; then, we expanded the region ± 250 kb from the external outliers
307 of the seed to define a genomic window for mining. Genes included in those genomic
308 windows were identified using the cockle's transcriptome assembled and annotated by
309 Pardo et al. (2022), which was used as reference to detect Gene Ontology (GO) functional
310 enrichment of the genomic regions under selection (FDR 5%) using GOfuncR (Grote

311 2022). Furthermore, we also analysed genomic windows around the SNPs correlated with
312 environmental variables for mining; since we could not identify consecutive SNPs as with
313 outliers, we were more conservative and defined smaller windows around each SNP (\pm
314 100 kb).

315

316 **Results**

317 *Genetic diversity and differentiation: whole sample and SNP dataset*

318 A total of 599 cockles were analysed, since six specimens that exhibited a low number
319 of reads (< 250.000 reads) from WDE_17 (two individuals), SAN_17 (one individual),
320 SVI_17 (two individuals) and SMO_17 (one individual), were removed. After quality
321 filtering, the number of SNPs retained in the whole dataset was 9,250. This number was
322 slightly lower than the number used in the macrogeographical study carried out by Vera
323 et al. (2022) (9,309 markers), because 59 of these markers were monomorphic in the
324 studied regions. All the 9,250 markers were included in the “9,309 markers” dataset
325 and their genomic information is available at
326 <https://onlinelibrary.wiley.com/doi/10.1111/eva.13340>, where the SNP code from Vera
327 et al. (2022) has been maintained for comparison between studies.

328 Observed (H_o) and expected (H_e) heterozygosities ranged respectively from 0.070
329 (SMO_17, Spain) to 0.080 (IWC_20, IGC_20 and IKF_20, Ireland; mean \pm SD = 0.075
330 \pm 0.003) and from 0.076 (WDE_17, Wales) to 0.087 (SNO_17, Spain and IKF_20,
331 Ireland; mean \pm SD = 0.082 \pm 0.003) (Table 1). All F_{IS} values per locus and bed were
332 positive, suggesting heterozygote deficit, but low (always < 0.115) and not
333 significant; and all beds met to HW expectations ($P < 0.0022$; 0.05/22 populations), an
334 expected outcome considering the HW filtering applied to retain SNPs. The percentage
335 of polymorphic loci ranged from 25.5 % in SMO_17 (Spain) to 52.8 % in SNO_17

336 (Spain) (mean \pm SD = $41.3 \pm 7.2\%$). When only polymorphic loci within each bed were
337 considered, H_o ranged from 0.137 in SLO_17 (Spain) to 0.181 in SMO_17 (Spain)
338 (mean \pm SD = 0.155 ± 0.014), showing these two beds also the lowest (0.156) and
339 highest (0.200) H_e (mean \pm SD = 0.171 ± 0.012). No differences in genetic diversity
340 were found between the SW British Isles and Galician regions (Mann-Whitney U tests
341 $P > 0.250$ for H_o , H_e with all loci and with polymorphic loci). Genetic diversity was in
342 the range of previous values reported by Vera et al. (2022) for the whole Atlantic area
343 using the same methodology.

344 Global F_{ST} for all beds was 0.02118 ($P < 0.001$), pairwise F_{ST} ranged from 0 (non-
345 significant $\neq 0$) for many bed pairs up to a maximum of 0.05040 ($P < 0.001$) between
346 IDC_18 and SVI_17 (Supplementary Table 2). Most pairwise comparisons were
347 significant excluding those from Galicia. Average pairwise F_{ST} between the SW British
348 Isles and Galicia was 0.03400 ($P < 0.001$), while 0.01374 ($P < 0.001$) within the SW
349 British Isles and -0.00529 ($P = 1.000$) within Galicia. The two beds from the Cotentin
350 Peninsula (FBV_19 and FGO_19, SW British Isles region), separated by 190
351 kilometres, showed significant genetic differentiation ($F_{ST} = 0.01207$, $P < 0.001$). The
352 most likely K values inferred by fastSTRUCTURE were 1 and 3. When K = 3 was
353 plotted, two main groups were identified differentiating the SW British Isles (IGC_20,
354 IKF_20, IWC_20, IDA_18, IDC_18, WDE_17, WBY_17, ECE_20, FBS_17, FBV_19
355 and FGO_19) from Galicia (plus Arcachon) (FAR_17, SBA_17, SMI_17, SAN_17,
356 SNO_17, SLO_17, SSA_17, SVI_17, SCA_17, SMO_17, SBI_18) (Fig. 2A). FGO_19
357 (France, Cotentin Peninsula) showed a high component of the southern group, also
358 detectable in all samples from the English Channel (ECE_20, FBS_17 and FBV_18),
359 suggesting some introgression between the two groups. The DAPC representation on
360 the SW British Isles also suggested differentiation of the English Channel samples from

361 the northernmost populations across the second component, while the first one,
362 indicated a remarkable divergence of the IKF_20 sample from the remaining ones (Fig.
363 3A). The DAPC from Galicia showed most of the samples grouped excluding SNO_17,
364 in the middle of the distribution, below Cape Finisterre, and SBI_18, the southernmost
365 one (Fig. 4A).

366

367 *Genetic structure within regions: demographic and selective factors*

368 To understand the factors underlying genetic differentiation within the SW British Isles
369 and Galician regions, we first identified those loci under selection using three different
370 statistical approaches. BY10, BY1000 and PCADAPT detected 159, 47 and 84 outliers
371 in the SW British Isles, respectively, all of them under divergent selection and
372 representing a total of 186 different outliers (Supplementary Table 3). Thirty-five
373 markers were shared between the three methods. The number of outliers in Galicia was
374 much lower (BY10 = 15, BY1000 = 2, PCADAPT = 39), two of them shared between
375 the three methods and representing a total of 51 outliers, all of them putatively under
376 divergent selection. Among the whole outlier dataset, 15 were shared between SW
377 British Isles and Galicia. Then, by discounting the total number of outliers to the whole
378 dataset in each region, a total of 9,064 neutral markers were identified in the SW British
379 Isles and 9,199 in Galicia, representing the neutral datasets for each region.

380 Small but significant genetic differentiation was detected among the SW British Isles
381 beds using neutral markers ($F_{ST} = 0.00778$, $P < 0.001$), suggesting limitations to larval
382 dispersion in this area by biogeographical barriers. As expected, the 186 total outliers
383 rendered a much higher global F_{ST} (0.10959, $P < 0.001$), being more accentuated when
384 using the shared set of outliers between methods ($F_{ST} = 0.17411$, $P < 0.001$), which
385 suggests selective factors increasing structuring. Pairwise F_{ST} ranged from -0.03095

386 (IKF_20 – WDE_17) to 0.02366 (IGC_20 – FGO_19 pair) for neutral markers; from
387 0.00061 (IDA_18 – IDC_18 pair) to 0.17142 (IDA_18 – FGO_19 pair) for the 186 total
388 outliers; and from -0.00345 (IDA_18 – IDC_18 pair) to 0.27313 (IKF_20 – FBS_17
389 pair) for the 35 shared outliers (Supplementary Table 4). IBD was significant with the
390 shared and total outlier datasets ($r = 0.63169$ and 0.55136 , respectively; $P < 0.001$), but
391 not with the neutral dataset ($r = 0.08578$, $P = 0.330$). These results suggest that
392 correlations could be a by-product of the unequal spatial distribution of the
393 environmental factors responsible of selective forces shaping the cockle's genome,
394 since IBD patterns should be reflected by the balance between drift and migration on
395 neutral markers. The fastSTRUCTURE analyses identified $K = 1$, $K = 2$ and $K = 3$ as
396 the most likely values for the neutral, 35 shared outlier and 186 total outlier datasets,
397 respectively (Fig. 2B), which consistently differentiated the Celtic Sea and the North-
398 west Irish cluster (IGC_20, IKF_20 and IWC_20), not studied to date, and the English
399 Channel cluster (ECE_20, FBS_17, FBV_19, FGO_19). In contrast, the Irish Sea
400 appeared as a rather differentiated group with the 186 outliers, which was split into two
401 clusters, the Irish side (IDA_18 and IDC_18) most closely associated with the Celtic
402 and Atlantic Ocean cluster, and the Welsh side (WDE_17 and WBY_17), most closely
403 linked with the English Channel cluster, when using the 35 outlier loci. The
404 differentiation of the Irish Sea from the other samples, and the contrast between the
405 Welsh and Irish (east and west, respectively) samples of the Irish Sea, was shown when
406 exploring a scenario with a larger K value, with both datasets displaying a very similar
407 structure with $K = 4$ (Supplementary Figures 2 and 3). The DAPC analysis with neutral
408 markers showed a very similar picture to that described with the whole SNP dataset
409 (Fig. 3B), however, the 186 and 35 outlier datasets displayed a very distinct picture,
410 both separating the English Channel (ECE_20, FGO_19, FBV_19 and FBS_17) from

411 the Welsh populations, but also from the Irish populations, which were further divided
412 into two groups, the westernmost Northeast Atlantic Ocean group (IWC_20, ICG_20
413 and IKF_20) and the Irish/Celtic Seas group (IDC_18 and IDA_18) (Figs. 3C and 3D).
414 In contrast to the SW British Isles, no population differentiation was found in Galicia
415 with the neutral dataset ($F_{ST} = 0.00552$, $P = 1.000$), also supported by the
416 fastSTRUCTURE ($K = 1$) and DAPC, as previously outlined with whole dataset (Figs.
417 4A and B). However, low but significant differentiation was detected with the 51
418 outliers ($F_{ST} = 0.00870$, $P < 0.001$), the pairwise F_{ST} supporting a significant
419 differentiation of the two northernmost samples (SMI_17 and especially SBA_17;
420 Supplementary Table 5) from the rest. This differentiation was not disclosed with
421 fastSTRUCTURE ($K = 1$; see Supplementary Fig. 4) and only suggested with DAPC
422 (Fig. 4C).

423

424 *Seascape analysis*

425 RDA analyses in SW British Isles region suggested longitude as the main driver for the
426 observed differentiation with all datasets and seasons (Table 2). Latitude was also
427 supported as driver for many models, especially for those related to the 186 total
428 outliers. Sea bottom salinity (SBS) was suggested for all seasons with the 186 outlier
429 dataset, while bottom shear stress (BSS) was for reproductive and summer seasons
430 using the whole and neutral datasets (Table 2). When longitude and latitude were
431 removed, sea surface temperature (SST) was suggested for all the datasets in the
432 summer season, and in the reproductive and winter seasons only with the whole and
433 186 outlier datasets, respectively. Sea bottom temperature (SBT) was suggested for the
434 reproductive and summer season with the 186 outlier dataset. SBS and sea surface
435 salinity (SSS) were suggested with the 186 outlier dataset for the summer and winter

436 seasons, respectively. Net primary production (NPP) was suggested for all datasets in
437 the winter season and for the 186 outlier dataset for the reproductive season. Finally,
438 BSS was suggested in all seasons for the neutral dataset and in the reproductive and
439 summer seasons for the complete dataset. In Galician region, no associations were
440 found, except for latitude in all periods analysed using the 51 outliers, and for BSS
441 during winter when latitude and longitude were removed (Table 2). However, VIF
442 values were usually high (> 10), suggesting that results should be taken with caution
443 due to the high collinearity among the variables in many cases.

444 While no correlations were identified in Galicia with BAYENV2, a total of 54 markers
445 were correlated with different environmental variables in the SW British Isles
446 (Supplementary Table 6). Thirty of these markers (55.6%) were previously identified as
447 outliers by the different methodologies applied. Markers were mainly correlated with
448 latitude, longitude, temperature and salinity. The main variable correlated with genetic
449 markers in the reproductive season and summer scenarios was SBT, while SSS and
450 NPP were in the winter scenario.

451

452 *Gene mining around outliers and environmental correlated markers*

453 Genetic markers associated with divergent selection or correlated with environmental
454 variables were mapped in the common cockle genome to look for functional
455 interpretation (Supplementary Tables 3 and 6). Outliers identified in the SW British
456 Isles area were scattered across all chromosomes, between one in C18 and 22 in C3,
457 while five chromosomes (C8, C11, C13, C14, C17) did not bear any outlier in Galicia,
458 the maximum being detected in C1 (11 outliers) (Table 3). The 51 outliers detected in
459 Galicia only identified a single consistent genomic region (window) under selection
460 according to our criteria and other five outliers were distributed across four confident

461 genomic windows previously reported by Vera et al. (2022) (Supplementary Tables 7
462 and 8). However, among the 186 outliers detected in the SW British Isles, 14 defined
463 five new consistent genomic windows under divergent selection and other 45 mapped
464 on genomic windows previously reported by Vera et al. (2022) (Supplementary Tables
465 7 and 8). Most outliers detected in Galicia were specific to this region, while an
466 important number of outliers from the SW British Isles were shared with the Northern
467 region previously analysed by Vera et al. (2022) (Supplementary Fig. 5). Still, a notable
468 proportion of outliers in the North were specific of each study (North-Vera et al.
469 (2022): 137 vs SW British Isles: 101) suggesting specific evolutionary factors related to
470 each scenario. Among the genes annotated in the five new windows, several related to
471 oxidative stress, hypoxia and immunity were identified in a 200 kb region in C2 and in
472 a 340 kb region in C3 (Supplementary Table 9) (Gerdol and Venier 2015; Grandi et al.
473 2016; Sokolov et al. 2019). Also, in a 480 kb region in C5, some genes involved in
474 signalling and detoxification (Wang et al. 2018; Kron 2022; Thoma et al. 2022) were
475 identified. Finally, a gene associated with ocean acidification (Lim et al. 2021) was
476 identified in C19. Despite the low number of genes handled, a significantly enriched
477 GO Molecular Function was detected (protein serine/threonine phosphatase activity;
478 GO:0004722) taking as background the common cockle transcriptome reported by
479 Pardo et al. (2022).

480 Markers correlated with environmental variables were scattered across most
481 chromosomes, excluding C7, C14 and C18, and the higher number (seven markers)
482 were detected in two big chromosomes (C2 and C4) (Table 3). An important number of
483 correlated markers were also identified as outliers for divergent selection (55.6 %),
484 some of them associated with consistent genomic windows (Supplementary Table 8).
485 Of note, the three markers detected in one of the most consistent genomic windows in

486 C4. We also mined the cockle genome around the correlated marker dataset
487 (Supplementary Table 10) and detected several genes related to nervous system
488 development and physiology. These genes were mostly clustered at C1 around
489 142462_31 (correlated with SBT) and C2 around 210318_7 (correlated with SST),
490 respectively. Furthermore, some of these genes were previously associated with
491 temperature stress and oxygen depletion stress or differentially expressed under specific
492 experimental conditions in other mollusc species (Woo et al. 2011; Chen et al. 2022).
493 Another important group of genes scattered around different markers in the cockle
494 genome were related to immunity and defence and had been previously reported in
495 other mollusc species in response to viruses and bacteria (Barbosa et al. 2022; Saco et
496 al. 2023) (Supplementary Table 9).

497 **Discussion**

498 Assessment of the distribution of genetic variability across populations, incorporating
499 historical processes and local adaptation framed within the dispersal range of the focal
500 organism (Richardson et al. 2014), is essential to develop management actions to
501 preserve exploited species (Bernatchez et al. 2017). In the present study, two different
502 patterns of genetic structure at microgeographic scale were identified in two regions
503 within the natural distribution of *C. edule*, highlighting the need to perform analyses at
504 multiple spatial scales (Hoffman et al. 2012), to provide information supporting the
505 management of this valuable resource.

506 *Heterogeneous pattern of microgeographic structure in the common cockle*

507 The two geographic areas studied, the SW British Isles region and Galicia, were
508 selected by their different habitat fragmentation patterns. Both areas were slightly
509 differentiated ($F_{ST} = 0.03400$), in accordance with their location in the major northern
510 and southern regions of the species' range separated around French Brittany (Vera et al.

511 2022), but did not show differences in genetic diversity, unlike Vera et al. (2022), who
512 reported a slight, but significant higher diversity in the southern region.

513 The extensive analysis performed in Galicia (10 natural beds) suggested the presence of
514 a single panmictic unit in this area, as previously reported for other molluscs, with
515 similar pelagic larval periods (*Donax trunculus*, Nantón et al. 2017; *Ensis siliqua*,
516 Arias-Pérez et al. 2012; *Mitilus galloprovincialis*, Diz and Presa 2009; *Ostrea. edulis*,
517 Vera et al. 2016; *Polititapes rhomboides*, Chacón et al. 2021), and for other marine
518 species (*Hippocampus guttulatus*; Lopez et al. 2015; *Pollicipes pollicipes*, Parrondo et
519 al. 2022). Our data does not support Cape Finisterre as a biogeographical barrier for the
520 species as previously suggested (Lopez-Jamar et al. 1992; Piñeira et al. 2008; Martinez
521 et al. 2013; Cruz et al. 2020), since no differentiation was detected between beds at both
522 sides of the Cape with the whole and neutral datasets. However, when using outlier
523 loci, the two northernmost Galician beds showed significant differentiation with the
524 remaining ones, especially the bed closest to the Cantabrian Sea (SBA_17) (average F_{ST}
525 = 0.03145), which could be related to the higher temperature regime in the Cantabrian
526 Sea (Marquina et al. 2015), but a more detailed study in the Cantabrian Sea would be
527 necessary to confirm this observation. Oceanographic dynamics in the Galician coast
528 indicate that the cold-upwelled water usually penetrates estuaries on the west, while it
529 only occurs during very intense events on the north (Alvarez et al. 2010). Thus, water
530 temperature decreases from north to west, with an SST average value of 19.5 °C in the
531 Cantabrian coast compared with the 18.5 °C measured in the west coast for the 1985-
532 2005 period (Gomez-Gesteira et al. 2008). Larval dispersal modelling carried out in
533 Vera et al. (2022) (see their Fig. 7) confirmed that cockle beds are well connected with
534 each other by larval transport in Galicia, but the connection between the Rias and the
535 sites to the northeast of Cape Finisterre was weaker, though present. Furthermore,

536 whilst the beds along the northwest coast of the Iberian Peninsula are affected by very
537 similar oceanographic conditions, during the late spring and late summer, temperatures
538 at the most north-easterly site can differ markedly from those at the other beds due to
539 its location at the edge of the upwelling system and at the inception point of the
540 Portugal Coastal Current (STT two degrees higher in the northern beds (mean = 18.51
541 °C) than in the southern ones (mean = 16.47 °C) during the summer; see Supplementary
542 Table 1). Despite the genotype-environment associations methods did not identify sea
543 temperature as driver, latitude, which is highly correlated with temperature, was
544 suggested by the RDA analysis as potential driver in the region.

545 Previous data from the SW British Isles suggested significant structure in *C. edule*
546 related both to current dynamics as well as to abiotic factors, such as salinity and
547 temperature (Coscia et al. 2020; Vera et al. 2022), as reported in other shellfish species
548 such as the horse mussel *Modiolus modiolus* (Gormley et al. 2015) and the great scallop
549 *Pecten maximus* (Vendrami et al. 2019; Hold et al. 2021). However, some regions in
550 this area are still poorly sampled in the common cockle (English Channel) or without
551 information (West Irish coast, North-east Atlantic). Outlier markers showed a moderate
552 pairwise genetic differentiation between beds ($F_{ST} = 0.10959$ and 0.17411 with the 186
553 and 35 outlier datasets, respectively), higher than that observed with neutral markers
554 ($F_{ST} = 0.00778$), as expected, suggesting selective factors shaping specific genomic
555 regions in a small geographic area. An important proportion of divergent outliers (68
556 markers) were shared with those reported by Vera et al. (2022) for the northern group
557 (210 outliers), which gives robustness to our observations; however, data also suggests
558 specific selective factors shaping the cockle's genome associated with the new
559 sampling in the SW British Isles (117 new outlier loci; 31 within consistent genomic
560 windows). In fact, five new confident genomic windows were identified including

561 relevant genes related to oxidative stress and immunity that would deserve further
562 studies as candidates to explain the association observed with environmental factors.
563 Despite biotic factors, such as pathogens, could not be contemplated in our study, their
564 diversity and distribution (influenced by abiotic factors) are important drivers shaping
565 the genome and distribution of species (Theodosopoulos et al. 2019) and specifically in
566 cockles (Vera et al. 2022; Pampin et al. 2023). Furthermore, we also deepened into the
567 correlation of specific SNPs with environmental factors and could identify, by mining
568 in the cockle genome several genes related to nervous transmission and immunity,
569 arranged in clusters or scattered in different chromosomes, that had been previously
570 reported in other mollusc associated with temperature or oxidative stress (Woo et al.
571 2011; Barbosa et al. 2022; Chen et al. 2022).

572 The population structure observed in the SW British Isles region may be in part
573 explained by the residual ocean currents and ocean fronts that characterise this area, but
574 also by selective factors such as salinity gradients, variable bottom shear stress (due to
575 large tidal variability) and sea temperature gradients (driven by ocean currents,
576 stratification and mixing, and latitudinal gradients); however, spatial seascape results
577 should be taken with caution due to the collinearity detected among variables. Both
578 outlier datasets could identify four genetic clusters following two main west-east and
579 north-south axes, which could explain the correlation observed between genetic and
580 geographic distances for outlier loci, but also the identification of longitude and latitude
581 as two main drivers in the seascape analysis. According to the outlier information, the
582 new sampled beds from Western Ireland (IGC_20, IKF_20 and IWC_20) (Northeast
583 Atlantic) would constitute a new cluster. These sites are connected by the Irish coastal
584 current (Brown et al. 2003; Fernand et al. 2006) and larval dispersal modelling (see Fig.
585 7 Vera et al. 2022) showed that the beds along the southwest coast of Ireland are well

586 interconnected. The Irish Sea can be split into two different clusters associated with the
587 Irish and Welsh sides, as previously suggested by Coscia et al. (2020). Sites along the
588 southeast coast of the Irish Sea are generally connected by northward currents and sites
589 along the north coast of Wales by eastward currents. In contrast, the two sites on the
590 west coast of the Irish Sea (IDC_18 and IDA_18), appear genetically separated from
591 the remainder of the Irish Sea; This may be driven by the Irish Sea Front acting as a
592 barrier which also drives warmer temperatures in the northwest Irish Sea than in the
593 well-mixed northeast Irish Sea. Finally, the English Channel forms a fourth cluster
594 including the ECE_20 bed from Cornwall with the southern beds limited by the Ushant
595 front. Interestingly, the Cotentin Peninsula, previously identified as a physical barrier to
596 dispersal in other molluscs, such as the slipper limpet *Crepidula fomicata* (Dupont et al.
597 2007) and *P. maximus* (Nicolle et al. 2016; Handal et al. 2020), showed a significant
598 differentiation between samples on its west and east sides (FGO_19 and FBV_19) with
599 neutral markers ($F_{ST} = 0.01045$, $P < 0.001$) and higher with outlier loci (F_{ST} 35 outliers
600 = 0.06654, $P < 0.001$; F_{ST} 186 outliers = 0.05876, $P < 0.001$), suggesting additional
601 selective factors differentiating both sides. Oceanic distance between the two Cotentin
602 beds (~ 190 km) is shorter than the longest distance between Galician beds (~ 300 km),
603 where no genetic differentiation was detected with neutral markers. Of note, FGO_19
604 showed an important genomic component of the South group, suggesting introgression
605 from the south especially in the west coast of the Cotentin Peninsula.

606

607 *Management implications*

608 The present study represents a refined analysis of the population structure of *C. edule* in
609 two geographic areas of small-medium size representing differentiated models that could
610 aid to obtain a more comprehensive picture for improving the management and

611 conservation of this valuable commercial and ecological resource. Galician beds were
612 suggested to constitute a panmictic population and this region could be managed as a
613 single genetic unit. The fisheries in this region are exclusively commercial and their
614 exploitation management can be through territorial concessions leased by shellfisher
615 guilds or directly by Galician regional government (i.e. free access shellfish areas). This
616 genetic information should be included in the Galician legislation, thus allowing
617 translocations from high production areas (Ría de Noia) to depleted ones by different
618 factors, such as the parasite *M. cochillia* (Ría de Arousa; Villalba et al. 2014). However,
619 caution should be taken considering biotic factors not evaluated in our study, such as
620 emergent pathologies (e.g. marteiliosis), which will require specific recommendations
621 within the general framework depicted in our study. A sharp fragmentation was displayed
622 by the SW British Isles region, especially with divergent outliers, mostly representing
623 adaptive management units (AMU, Bernatchez et al. 2017). Thus, Western (Northeast
624 Atlantic) Irish beds would represent a differentiated group from those previously
625 described, while subtle genetic sub-structuring was identified along the English Channel,
626 with a significant effect at the Cotentin Peninsula representing as a biogeographic barrier.
627 Furthermore, the Irish Sea, a narrow water body mass between Wales and Ireland, appears
628 to represent differentiated units at both sides of the Irish Sea according to our information.
629 All these population units should be individually managed, avoiding translocations
630 between them. Finally, our results could help to improve cockles' production by founding
631 appropriate broodstock to enhance depleted populations and by tracing samples to check
632 undesirable transferences among regions.

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638

639 **Author Contributions**

640 MV, AV and PM designed and supervised the study. DI, AC, KM, FO, SKM, SL
641 performed field collections. PM, SCC, SL, PER, SKM and FO provided funding. FM,
642 MH and AB analysed bioinformatically genomic sequences and created genotyping
643 files. SBW and PER provided information about oceanography, environmental
644 variables and developed geographic maps included in the figures. MV, CB, ACC, AB
645 and PM performed the genetic analyses. MV and PM wrote the manuscript with
646 contributions from all authors. All of them read the manuscript and gave their approval.

647

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652

653 **Conflict of interest**

654 The authors declare no conflict of interest.

655

656 **Data archiving**

657 Data for this study are available at Dryad Digital Repository (<http://to be completed after>
658 *manuscript is accepted for publication*) and Supplementary material.

659 **References**

660 Abaunza P, Murta AG, Campbell N, Cimmaruta R, Comesana AS, Dahle G, Santamaria
661 MTG, Gordo LS, Iversen SA, MacKenzie K, Magoulas A, Mattiucci S, Molloy
662 J, Nascetti G, Pinto AL, Quinta R, Rarnos P, Sanjuan A, Santos AT, Stransky C,
663 Zimmermann C (2008) Stock identity of horse mackerel (*Trachurus trachurus*)
664 in the Northeast Atlantic and Mediterranean Sea: Integrating the results from
665 different stock identification approaches. Fish. Res. 89:196-209

666 Alvarez I, Gomez-Gesteira M, DeCastro M, Gomez-Gesteira JL, Dias JM (2010)
667 Summer upwelling frequency along the western Cantabrian coast from 1967 to
668 2007. J. Mar. Syst. 79:218-226

669 Alvarez I, Gomez-Gesteira M, deCastro M, Lorenzo MN, Crespo AJC, Dias JM (2011)
670 Comparative analysis of upwelling influence between the western and northern
671 coast of the Iberian Peninsula. Cont. Shelf Res. 31:388-399

672 Arias-Perez A, Fernandez-Tajes J, Gaspar MB, Mendez J (2012) Isolation of
673 microsatellite markers and analysis of genetic diversity among East Atlantic
674 populations of the sword razor shell *Ensis siliqua*: a tool for population
675 management. Biochem. Genet. 50: 397-415

676 Barbosa M, Schwaner C, Pales Espinosa E, Allam B (2022) A Transcriptomic analysis
677 of phenotypic plasticity in *Crassostrea virginica* larvae under experimental
678 acidification. Genes 13: 1529

679 Beaumont AR, Day TR, Gade G (1980) Genetic variation at the octopine
680 dehydrogenase locus in the adductor muscle of *Cerastoderma edule* (L) and 6
681 other bivalve species. Mar.Biol. Letters 1: 137-148

682 Bernatchez L, Wellenreuther M, Araneda C, Ashton DT, Barth JMI, Beacham TD,
683 Maes GE, Martinsohn JT, Miller KM, Naish KA, Ovenden JR, Primmer CR,

684 Suk HY, Therkildsen NO, Withler RE (2017) Harnessing the Power of
685 Genomics to Secure the Future of Seafood. *Trends Ecol. Evol.* 32:665-680

686 Brown J, Carrillo L, Fernand L, Horsburgh KJ, Hill AE, Young EF, Medler KJ (2003)
687 Observations of the physical structure and seasonal jet-like circulation of the
688 Celtic Sea and St. George's Channel of the Irish Sea. *Cont. Shelf Res.* 23:533–
689 561

690 Bruzos AL, Santamarina M, Garcia-Souto D, Diaz S, Rocha S, Zamora J, Lee Y, Viña-
691 Feas A, Quail MA, Otero I, Pequeño-Valtierra A, Temes J, Rodriguez-Castro J,
692 Villanueva A, Costas D, Rodriguez R, Prieto T, Tomas L, Alvariño P, Alonso J,
693 Cao A, Iglesias D, Carballal MJ, Amaral AM, Balseiro P, Calado R, El Khalfi B,
694 Izagirre U, de Montaudouin X, Pade NG, Probert I, Ricardo F, Ruiz P, Skazina
695 M, Smolarz K, Pasantes JJ, Villalba A, Ning Z, Ju YS, Posada D,
696 Demeulemeester J, Baez-Ortega A, Tubio JMC (2022) The evolution of two
697 transmissible leukaemias colonizing the coasts of Europe. *Biorxiv*
698 doi10.1101/2022.08.06.503021

699 Carss DN, Brito AC, Chainho P, Ciutat A, de Montaudouin X, Fernandez Otero RM,
700 Incera Filgueira M, Garbutt A, Goedknecht MA, Lynch SA, Mahony KE, Maire
701 O, Malham SK, Orvain F, Olivier AvdS, Jones L (2020) Ecosystem services
702 provided by a non-cultured shellfish species: The common cockle *Cerastoderma*
703 *edule*. *Mar. Environ. Res.* 158:104931

704 Catchen J, Hohenlohe PA, Bassham S, Amores A, Cresko WA (2013) Stacks: an
705 analysis tool set for population genomics. *Mol. Ecol.* 22:3124-3140

706 Chacon GM, Arias-Perez A, Freire R, Martinez L, Ojea J, Insua A (2021) Genetic
707 characterization of wild, broodstock and seed samples of *Polittapes rhomboides*

708 (Bivalvia: Veneridae): Implications for hatchery seed production. Aquacult.
709 Rep. 20:100658

710 Chen J, Leng T, Jiang YM, Chen XB, Liu ZM (2022) RNA-seq analysis of the
711 differential response to low-temperature stress in two morphs of mud crabs
712 (*Scylla paramamosain*). Comp. Biochem. Physiol. Part D Genomics Proteomics
713 43:101010

714 Coop G, Witonsky D, Di Rienzo A, Pritchard JK (2010) Using environmental
715 correlations to identify loci underlying local adaptation. Genetics 185:1411-1423

716 Coscia I, Wilmes SB, Ironside JE, Goward-Brown A, O'Dea E, Malham SK, McDevitt
717 AD, Robins PE (2020) Fine-scale seascape genomics of an exploited marine
718 species, the common cockle *Cerastoderma edule*, using a multimodelling
719 approach. Evol. Appl. 13:1854-1867

720 Cruz A, da Costa F, Fernandez-Perez J, Nanton A, Fernandez-Boo S, Insua A, Mendez
721 J (2020) Genetic variability in *Ruditapes decussatus* clam combined with
722 *Perkinsus* infection level to support founder population selection for a breeding
723 program. PeerJ 8:e9728

724 Dare PJ, Bell MC, Walker P, Bannister RCA (2004) Historical and current status of
725 cockle and mussel stocks in The Wash. CEFAS.

726 de Montaudouin X, Bachelet G, Sauriau PG (2003) Secondary settlement of cockles
727 *Cerastoderma edule* as a function of current velocity and substratum: a flume
728 study with benthic juveniles. Hydrobiologia 503:103-116

729 Diz AP, Presa P (2009) The genetic diversity pattern of *Mytilus galloprovincialis* in
730 Galician Rias (NW Iberian estuaries). Aquaculture 287:278-285

731 do Prado FD, Vera M, Hermida M, Bouza C, Pardo BG, Vilas R, Blanco A, Fernandez
732 C, Maroso F, Maes GE, Turan C, Volckaert FAM, Taggart JB, Carr A, Ogden
733 R, Nielsen EE, Martinez P, Aquatrace C (2018) Parallel evolution and
734 adaptation to environmental factors in a marine flatfish: Implications for
735 fisheries and aquaculture management of the turbot (*Scophthalmus maximus*).
736 *Evol. Appl.* 11:1322-1341

737 Dupont L, Ellien C, Viard F (2007) Limits to gene flow in the slipper limpet *Crepidula*
738 *fornicata* as revealed by microsatellite data and a larval dispersal model. *Mar.*
739 *Ecol. Prog. Ser.* 349:125-138.

740 Eldon B, Riquet F, Yearsley J, Jollivet D, Broquet T (2016) Current hypotheses to
741 explain genetic chaos under the sea. *Curr. Zool.* 62:551-566

742 Excoffier L, Lischer HEL (2010) Arlequin suite ver 3.5: a new series of programs to
743 perform population genetics analyses under Linux and Windows. *Mol. Ecol.*
744 *Resour.* 10:564-567

745 Fernand L, Nolan GD, Raine R, Chambers CE, Dye SR, White M, Brown J (2006) The
746 Irish coastal current: A seasonal jet-like circulation. *Cont. Shelf Res.* 26:1775–
747 1793

748 Fisher MC, Helser TE, Kang S, Gwak W, Canino MF, Hauser L (2022) Genetic
749 structure and dispersal in peripheral populations of a marine fish (Pacific cod,
750 *Gadus macrocephalus*) and their importance for adaptation to climate change.
751 *Ecol. Evol.* 12:e8474

752 Flach EC, de Bruin W (1994) Does the activity of cockles, *Cerastoderma edule* (L.) and
753 lugworms, *Arenicola marina* L., make corophium-volutator pallas more

754 vulnerable to epibenthic predators: a case of interaction modification. *J. Exp.*
755 *Mar. Biol. Ecol.* 182:265-285

756 Foll M, Gaggiotti O (2008) A Genome-Scan Method to Identify Selected Loci
757 Appropriate for Both Dominant and Codominant Markers: A Bayesian
758 Perspective. *Genetics* 180:977-993

759 Foll M (2012) BayeScan v2.1 User Manual. Available at:
760 http://cmpg.unibe.ch/software/BayeScan/files/BayeScan2.1_manual.pdf

761 Francis RM (2017) POPHELPER: an R package and web app to analyse and visualize
762 population structure. *Mol. Ecol. Resour.* 17:27-32

763 Galarza JA, Carreras-Carbonell J, Macpherson E, Pascual M, Roques S, Turner GF,
764 Rico C (2009) The influence of oceanographic fronts and early-life-history traits
765 on connectivity among littoral fish species. *Proc. Natl. Acad. Sci. U.S.A.*
766 106:1473-1478

767 Galparsoro I, Borja A, Uyarra MC (2014) Mapping ecosystem services provided by
768 benthic habitats in the European North Atlantic Ocean. *Front. Mar. Sci.* 1:23

769 Gerdol M, Venier P (2015) An updated molecular basis for mussel immunity. *Fish*
770 *Shellfish Immunol.* 46:17-38

771 Gomez-Gesteira M, de Castro M, Alvaez I, Gomez-Gesteira JL (2008) Coastal sea
772 surface temperature warming trend along the continental part of the Atlantic Arc
773 (1985-2005). *J. Geophys. Res. Oceans* 113:C04010

774 Gormley K, Mackenzie C, Robins P, Coscia I, Cassidy A, James J, Hull A, Piertney S,
775 Sanderson W., Porter J (2015) Connectivity and Dispersal Patterns of Protected
776 Biogenic Reefs: Implications for the Conservation of *Modiolus modiolus* (L.) in
777 the Irish Sea. *PLOS One* 10:e0143337

778 Grandi A, Santi A, Campagnoli S, Parri M, De Camilli E, Song C, Jin B, Lacombe A,
779 Castori-Eppenberger S, Sarmientos P, Grandi G, Viale G, Terracciano L,
780 Chiarugi P, Pileri P, Grifantini R (2016) ERMP1, a novel potential oncogene
781 involved in UPR and oxidative stress defense, is highly expressed in human
782 cancer. *Oncotarget* 7:63596-63610

783 Grote S (2022) GOfuncR: Gene ontology enrichment using FUNC. R package version
784 1.18.0

785 Group “Grepma” (1988) A physical, chemical and biological characterization of the
786 Ushant tidal front. *Int. Rev. ges. Hydrobiol. Hydrogr.* 73:511–536

787 Gunther T, Coop G (2013) Robust Identification of Local Adaptation from Allele
788 Frequencies. *Genetics* 195:205-220

789 Handal W, Szostek C, Hold N, Andrello M, Thiebaut E, Harney E, Lefebvre G, Borcier
790 E, Jolivet A, Nicolle A, Boye A, Foucher E, Boudry P, Charrier G (2020) New
791 insights on the population genetic structure of the great scallop (*Pecten*
792 *maximus*) in the English Channel, coupling microsatellite data and demogenetic
793 simulations. *Aquat. Conserv.* 30:1841-1853

794 Hayward PJ, Ryland JS (1995) Handbook of the marine fauna of north-west Europe.
795 Oxford University Press.

796 Hoffman JI, Clarke A, Clark MS, Fretwell P, Peck LS (2012) Unexpected fine-scale
797 population structure in a broadcast-spawning Antarctic marine mollusc. *PLOS*
798 *One* 7:e32415

799 Hold N, Robins P, Szostek CL, Lambert G, Lincoln H, Le Vay L et al (2021) Using
800 biophysical modelling and population genetics for conservation and

801 management of an exploited species, *Pecten maximus* L. Fish. Oceanogr.
802 30:740-756.

803 Honkoop PJC, van der Meer J (1998) Experimentally induced effects of water
804 temperature and immersion time on reproductive output of bivalves in the
805 Wadden Sea. Journal of Experimental Mar. Biol. Ecol. 220:227-246

806 Hummel H, Wolowicz M, Bogaards RH (1994) Genetic variability and relationships for
807 populations of *Cerastoderma edule* and of the *C. glaucum* complex. Netherlands
808 J. Sea Res. 33:81-89

809 Jackson-Bue M, Brito AC, Cabral S, Carss DN, Carvalho F, Chainho P, Ciutat A,
810 Sanchez EC, de Montaudouin X, Otero RMF, Filgueira MI, Fitch A, Garbutt A,
811 Goedknecht MA, Lynch SA, Mahony KE, Maire O, Malham SK, Orvain F,
812 Rocroy M, Olivier AvdS, Jones L (2022) Inter-country differences in the
813 cultural ecosystem services provided by cockles. People Nat. 4:71-87

814 Jombart T, Ahmed I (2011) adegenet 1.3-1: new tools for the analysis of genome-wide
815 SNP data. Bioinformatics 27:3070-3071

816 Jombart T, Devillard S, Balloux F (2010) Discriminant analysis of principal
817 components: a new method for the analysis of genetically structured
818 populations. BMC Genetics 11:94

819 Kron NS (2022) In search of the *Aplysia* immunome: an in silico study. BMC Genomics
820 23:543

821 Langmead B, Trapnell C, Pop M, Salzberg SL (2009) Ultrafast and memory-efficient
822 alignment of short DNA sequences to the human genome. Genome Biol. 10:R25

823 Larmuseau MHD, Van Houdt JKJ, Guelinckx J, Hellemans B, Volckaert FAM (2009)
824 Distributional and demographic consequences of Pleistocene climate

825 fluctuations for a marine demersal fish in the north-eastern Atlantic. *J. Biogeogr.*
826 36:1138-1151

827 Leary D, Vierros M, Hamon G, Arico S, Monagle C (2009) Marine genetic resources: A
828 review of scientific and commercial interest. *Mar. Policy* 33:183-194

829 Lim Y-K, Cheung K, Dang X, Roberts SB, Wang X, Thiagarajan V (2021) DNA
830 methylation changes in response to ocean acidification at the time of larval
831 metamorphosis in the edible oyster, *Crassostrea hongkongensis*. *Mar. Environ.*
832 Res. 163:105214

833 Lopez A, Vera M, Planas M, Bouza C (2015) Conservation genetics of threatened
834 *Hippocampus guttulatus* in vulnerable habitats in NW Spain: Temporal and
835 spatial stability of wild populations with flexible polygamous mating system in
836 captivity. *PLOS One* 10:0117538

837 Lopez-jamar E, Cal RM, Gonzalez G, Hanson RB, Rey J, Santiago G, Tenore KR
838 (1992) Upwelling and outwelling effects on the benthic regime of the
839 continental-shelf off Galicia, NW Spain. *J. Mar. Res.* 50:465-488

840 Luu K, Bazin E, Blum MGB (2017) Pcadapt: an R package to perform genome scans
841 for selection based on principal component analysis. *Mol. Ecol. Resour.* 17:67-
842 77

843 Mahony KE, Lynch SA, Egerton S, Cabral S, de Montaudouin X, Fitch A, Magalhaes
844 L, Rocroy M, Culloty SC (2020) Mobilisation of data to stakeholder
845 communities. Bridging the research-practice gap using a commercial shellfish
846 species model. *PLOS One* 15:0238446

847 Mahony KE, Lynch SA, Egerton S, Laffan RE, Correia S, de Montaudouin X, Mesmer-
848 Dudons N, Freitas R, Culloty SC (2021) Latitudinal influence on gametogenesis
849 and host-parasite ecology in a marine bivalve model. *Ecol. Evol.* 11:7029-7041

850 Malham SK, Hutchinson TH, Longshaw M (2012) A review of the biology of European
851 cockles (*Cerastoderma* spp.). *J. Mar. Biol. Assoc. U. K.* 92:1563-1577

852 Maroso F, Casanova A, do Prado FD, Bouza C, Pardo BG, Blanco A, Hermida M,
853 Fernandez C, Vera M, Martinez P (2018) Species identification of two closely
854 exploited flatfish, turbot (*Scophthalmus maximus*) and brill (*Scophthalmus*
855 *rhombus*), using a ddRADseq genomic approach. *Aquat. Conserv.* 28:1253-1260

856 Maroso F, Perez de Gracia C, Iglesias D, Cao A, Diaz S, Villalba A, Vera M, Martinez
857 P (2019) A useful SNP panel to distinguish two cockle species, *Cerastoderma*
858 *edule* and *C. glaucum*, co-occurring in some European beds, and their putative
859 hybrids. *Genes* 10:760

860 Marquardt DW (1970). Generalized inverses, ridge regression, biased linear estimation
861 and nonlinear estimation. *Technometrics* 12:59.

862 Marquina D, Angel Fernandez-Alvarez F, Norena C (2015) Five new records and one
863 new species of Polycladida (Platyhelminthes) for the Cantabrian coast (North
864 Atlantic) of the Iberian Peninsula. *J. Mar. Biol. Assoc. U. K.* 95:311-322

865 Martinez L, Mendez J, Insua A, Arias-Perez A, Freire R (2013) Genetic diversity and
866 population differentiation in the cockle *Cerastoderma edule* estimated by
867 microsatellite markers. *Helgol. Mar. Res.* 67: 179-189

868 Martinez L, Freire R, Arias-Perez A, Mendez J, Insua A (2015) Patterns of genetic
869 variation across the distribution range of the cockle *Cerastoderma edule* inferred
870 from microsatellites and mitochondrial DNA. *Mar. Biol.* 162:1393-1406

871 Miller JM, Cullingham CI, Peery RM (2020) The influence of a priori grouping on
872 inference of genetic clusters: simulation study and literature review of the DAPC
873 method. *Heredity* 125:269-280

874 Nanton A, Arias-Perez A, Freire R, Fernandez-Perez J, Novoa S, Mendez J (2017)
875 Microsatellite variation in *Donax trunculus* from the Iberian Peninsula, with
876 particular attention to Galician estuaries (NW Spain). *Estuar. Coast. Shelf Sci.*
877 197:27-34

878 Nicolle A, Moitie R, Ogor J, Dumas F, Foveau A, Foucher E, Thiebaut E (2016)
879 Modelling larval dispersal of *Pecten maximus* in the English Channel: A tool for
880 the spatial management of the stocks. *ICES J. Mar. Sci.* 74:1812–1825.

881 Norris K, Bannister RCA, Walker PW (1998) Changes in the number of oystercatchers
882 *Haematopus ostralegus* wintering in the Burry Inlet in relation to the biomass of
883 cockles *Cerastoderma edule* and its commercial exploitation. *J. Appl. Ecol.*
884 35:75-85

885 Oksanen J. (2015) Multivariate analysis of ecological communities in R: vegan tutorial.
886 <http://cc.oulu.fi/~jarioksa/opetus/metodi/vegantutor.pdf>

887 Pampin M, Casanova A, Fernandez C, Blanco A, Hermida M, Vera M, Pardo BG,
888 Coimbra RM, Cao A, Iglesias D, Carballal MJ, Villalba A, Martinez P (2023)
889 Genetic markers associated with divergent selection against the parasite
890 *Marteilia cochillia* in common cockle (*Cerastoderma edule*) using
891 transcriptomics and population genomics data. *Front. Mar. Sci.* 10:1057206

892 Pardo BG, Fernandez C, Pampin M, Blanco A, Iglesias D, Cao A, Carballal MJ,
893 Villalba A, Martinez P (2022) Transcriptome characterization of the common

894 cockle (*Cerastoderma edule*) after exposure to a *Marteilia cochillia* outbreak.
895 Biorxiv doi10.1101/2022.10.18.512677

896 Parrondo M, Moran P, Ballenghien M, Acuna JL, Aguion A, Arrontes J, Chiss J, Cruz
897 T, Fernandes JN, Garcia-Florez L, Garcia-Vazquez E, Geiger KJ, Macho G,
898 Thiebaut E, Weidberg N, Jollivet D, Borrell YJ (2022) Chaotic genetic
899 patchiness in the highly valued atlantic stalked barnacle *Pollicipes pollicipes*
900 from the iberian peninsula: implications for fisheries management. Front. Mar.
901 Sci. 9:801780

902 Piñeira J, Quesada H, Rolan-Alvarez E, Caballero A (2008) Genetic discontinuity
903 associated with an environmentally induced barrier to gene exchange in the
904 marine snail *Littorina saxatilis*. Mar. Ecol. Prog. Ser. 357:175-184

905 Prive F, Luu K, Vilhjalmsson BJ, Blum MGB (2020) Performing highly efficient
906 genome scans for local adaptation with R Package pcadapt Version 4. Mol. Biol.
907 Evol. 37:2153-2154

908 Raj A, Stephens M, Pritchard JK (2014) fastSTRUCTURE: Variational inference of
909 population structure in large SNP data sets. Genetics 197:573-U207

910 Rellstab C, Gugerli F, Eckert AJ, Hancock AM, Holderegger R (2015) A practical guide
911 to environmental association analysis in landscape genomics. Mol. Ecol.
912 24:4348-4370

913 Richardson JL, Urban MC, Bolnick DI, Skelly DK (2014) Microgeographic adaptation
914 and the spatial scale of evolution. Trends Ecol. Evol. 29:165-176

915 Rohlf F (1993) NTSYS-pc. Numerical taxonomy and multivariate analysis system,
916 Version 2.1. Setauket, New York.

917 Rousset F (1997) Genetic differentiation and estimation of gene flow from F-statistics
918 under isolation by distance. *Genetics* 145:1219-1228

919 Rousset F (2008) GENEPOP '007: a complete re-implementation of the GENEPOP
920 software for Windows and Linux. *Mol. Ecol. Resour.* 8:103-106

921 Saco A, Suárez H, Novoa B, Figueras A (2023) A genomic and transcriptomic analysis
922 of the C-type lectin gene family reveals highly expanded and diversified
923 repertoires in bivalves. *Mar. Drugs* 21:254

924 Sa-Pinto A, Branco MS, Alexandrino PB, Fontaine MC, Baird SJE (2012) Barriers to
925 gene flow in the marine environment: insights from two common intertidal
926 limpet species of the Atlantic and Mediterranean. *PLOS One* 7:0050330

927 Sharples J, Simpson JH (2019) Shelf sea and shelf slope fronts. In: Reference Module in
928 Earth Systems and Environmental Sciences. Vol. 1, pp 24-34. Elsevier.

929 Simpson JH, Pingree RD (1978). Shallow Sea Fronts Produced by Tidal Stirring. In:
930 Bowman, M.J., Esaias, W.E. (eds) *Oceanic Fronts in Coastal Processes*.
931 Springer, Berlin, Heidelberg.

932 Sokolov EP, Markert S, Hinzke T, Hirschfeld C, Becher D, Ponsuksili S, Sokolova IM
933 (2019) Effects of hypoxia-reoxygenation stress on mitochondrial proteome and
934 bioenergetics of the hypoxia-tolerant marine bivalve *Crassostrea gigas*. *J.*
935 *Proteomics* 194:99-111

936 Suberg LA, Miller PI, Wynn RB (2019) On the use of satellite-derived frontal metrics
937 in time series analyses of shelf-sea fronts, a study of the Celtic Sea. *Deep Sea*
938 *Res. Part I Oceanogr. Res. Pap.* 149:103033

939 Theodosopoulos AN, Hund AK, Taylor SA (2019) Parasites and host species barriers in
940 animal hybrid zones. *Trends Ecol. Evol.* 34:19-30

941 Thoma J, Stenitzer D, Grabherr R, Staudacher E (2022) Identification, characterization,
942 and expression of a beta-Galactosidase from *Arion* species (Mollusca).
943 *Biomolecules* 12:1578

944 Vendrami DLJ, De Noia M, Telesca L, Handal W, Charrier G, Boudry P, Eberhart-
945 Phillips L, Hoffman JI (2019) RAD sequencing sheds new light on the genetic
946 structure and local adaptation of European scallops and resolves their
947 demographic histories. *Sci. Rep.* 9:7455

948 Vera M, Carlsson J, El Carlsson J, Cross T, Lynch S, Kamermans P, Villalba A, Culloty
949 S, Martinez P (2016) Current genetic status, temporal stability and structure of
950 the remnant wild European flat oyster populations: conservation and restoring
951 implications. *Mar. Biol.* 163:239

952 Vera M, Maroso F, Wilmes SB, Hermida M, Blanco A, Fernandez C, Groves E,
953 Malham SK, Bouza C, Robins PE, Martinez P, Cockles C (2022) Genomic
954 survey of edible cockle (*Cerastoderma edule*) in the Northeast Atlantic: A
955 baseline for sustainable management of its wild resources. *Evol. Appl.* 15:262-
956 285

957 Vera M, Pardo BG, Cao A, Vilas R, Fernandez C, Blanco A, Gutierrez AP, Bean TP,
958 Houston RD, Villalba A, Martinez P (2019) Signatures of selection for
959 bonamiosis resistance in European flat oyster (*Ostrea edulis*): New genomic
960 tools for breeding programs and management of natural resources. *Evol. Appl.*
961 12:1781-1796

962 Vilas R, Vandamme SG, Vera M, Souza C, Maes GE, Volckaert FAM, Martinez P
963 (2015) A genome scan for candidate genes involved in the adaptation of turbot
964 (*Scophthalmus maximus*). *Mar. Genom.* 23: 77-86

965 Villalba A, Iglesias D, Ramilo A, Darriba S, Parada JM, No E, Abollo E, Molaes J,
966 Carballal MJ (2014) Cockle *Cerastoderma edule* fishery collapse in the Ria de
967 Arousa (Galicia, NW Spain) associated with the protistan parasite *Marteilia*
968 *cochillia*. Dis. Aquat. Org. 109:55-80.

969 Wang M, Wang L, Ni D, Yi Q, Wang X, Jia Z, Song L (2018) The mRNA expression
970 profiles demonstrating versatile roles of glutathione S-transferase genes in the
971 mollusk *Chlamys farreri*. Invertebr. Surviv. J. 15:302-315

972 Waples RS (1998) Separating the wheat from the chaff: Patterns of genetic
973 differentiation in high gene flow species. J. Hered. 89:438-450

974 Woo S, Jeon HY, Kim SR, Yum S (2011) Differentially displayed genes with oxygen
975 depletion stress and transcriptional responses in the marine mussel, *Mytilus*
976 *galloprovincialis*. Comp. Biochem. Physiol. Part D Genomics Proteomics 6:
977 348-356

978

979 **Figure legends:**

980 Figure 1. Geographical distribution of the *Cerastoderma edule* beds analysed in the
981 present study. Ocean bathymetry is shaded in blue. Summer sea surface ocean currents
982 are schematically depicted with magenta coloured arrows. Tidal mixing fronts are
983 indicated with purple dashed lines. 'UF' stands for Ushant Front, 'CSF' for Celtic Sea
984 Front, and 'ISF' for Irish Sea Front. Location codes are shown in Table 1. Beds
985 previously analysed by Vera et al. (2022) are marked with asterisks.

986 Figure 2. Population structure of *Cerastoderma edule* at different geographical scales
987 using fastSTRUCTURE. Each vertical bar represents one individual, and the colour
988 proportion of each bar represents the posterior probability of assignment of each
989 individual to the different clusters (K) inferred by the program. The most likely K = 3
990 using the whole dataset (A), and K = 2 using the 57 shared divergent outliers between
991 methodologies and K = 3 using the total 186 divergent outliers (B) for the SW British
992 Isles are represented. Codes are shown on Table 1. Plots for all the K values tested for
993 the different datasets are shown in Supplementary Figures.

994 Figure 3. Discriminant Analysis of Principal Components (DAPC) plots of
995 *Cerastoderma edule* beds belonging to the SW British Isles. The weight of retained
996 discriminant analysis (DA) and principal components selected are shown on left bottom
997 box and right bottom box, respectively. Results using the complete whole dataset (A),
998 the neutral dataset (B), the 35 shared divergent outliers between methodologies (C) and
999 the 186 divergent outliers (D) are represented. Codes are shown on Table 1.

1000 Figure 4. Discriminant Analysis of Principal Components (DAPC) plots of
1001 *Cerastoderma edule* beds belonging to Galicia. The weight of retained discriminant
1002 analysis (DA) and principal components selected are shown on left bottom box and

1003 right bottom box, respectively. Results using the complete dataset (A), neutral dataset
1004 (B) and 51 divergent outliers (C) are represented. Codes are shown on Table 1.

1005 Table 1. *Cerastoderma edule* beds analysed in the present study. Location, sampling year, geographical coordinates, code, country, number of
 1006 individuals collected (N initial) and analysed after quality filtering (N), observed heterozygosity (Ho), expected heterozygosity (He), inbreeding
 1007 coefficient (F_{IS}) for all dataset and for polymorphic loci are shown. Locations in italics were previously analysed by Vera et al. (2022).

Location	Year	Lat (deg N)	Lon (deg E)	Code	Country	N initial	All dataset				Polymorphic loci (MAF > 0.017)				
							N	Ho	He	F _{IS}	Polymorphic loci	% Polymorphic loci	Ho	He	F _{IS}
Galway Connemara	2020	53.306	-9.846	IGC_20	Ireland	27	27	0.080	0.085	0.067	4205	45.5	0.166	0.179	0.074
Kerry- Feale	2020	52.488	-9.652	IKF_20	Ireland	30	30	0.080	0.087	0.080	4362	47.2	0.161	0.176	0.088
West Cork-Cockle Beach	2020	51.463	-9.744	IWC_20	Ireland	30	30	0.080	0.085	0.064	4294	46.4	0.163	0.175	0.070
<i>Dundalk Bay-Annagassan</i>	<i>2018</i>	<i>53.884</i>	<i>-6.341</i>	<i>IDA_18</i>	<i>Ireland</i>	<i>29</i>	<i>29</i>	<i>0.074</i>	<i>0.080</i>	<i>0.082</i>	<i>2993</i>	<i>32.4</i>	<i>0.161</i>	<i>0.177</i>	<i>0.090</i>
<i>Dundalk Bay-Cooley</i>	<i>2018</i>	<i>53.996</i>	<i>-6.287</i>	<i>IDC_18</i>	<i>Ireland</i>	<i>22</i>	<i>22</i>	<i>0.077</i>	<i>0.083</i>	<i>0.081</i>	<i>3493</i>	<i>37.8</i>	<i>0.173</i>	<i>0.190</i>	<i>0.087</i>
<i>Dee Estuary</i>	<i>2017</i>	<i>53.343</i>	<i>-3.174</i>	<i>WDE_17</i>	<i>Wales</i>	<i>30</i>	<i>28</i>	<i>0.071</i>	<i>0.076</i>	<i>0.074</i>	<i>2561</i>	<i>27.7</i>	<i>0.167</i>	<i>0.182</i>	<i>0.084</i>
<i>Burry</i>	<i>2017</i>	<i>51.643</i>	<i>-4.166</i>	<i>WBY_17</i>	<i>Wales</i>	<i>30</i>	<i>30</i>	<i>0.073</i>	<i>0.080</i>	<i>0.091</i>	<i>3529</i>	<i>38.2</i>	<i>0.148</i>	<i>0.165</i>	<i>0.101</i>
Camel Estuary (Cornwall)	2020	50.531	-4.930	ECE_20	England	24	24	0.073	0.081	0.105	3667	39.6	0.159	0.180	0.114
<i>Somme Bay</i>	<i>2017</i>	<i>50.201</i>	<i>1.627</i>	<i>FBS_17</i>	<i>France</i>	<i>30</i>	<i>30</i>	<i>0.071</i>	<i>0.080</i>	<i>0.111</i>	<i>3438</i>	<i>37.2</i>	<i>0.147</i>	<i>0.167</i>	<i>0.119</i>
Baie des Veys (Brévands)	2019	49.365	-1.150	FBV_19	France	26	26	0.079	0.081	0.022	3579	38.7	0.169	0.174	0.030
Gouville sur mer	2019	49.105	-1.612	FGO_19	France	23	23	0.077	0.082	0.059	4055	43.8	0.163	0.174	0.063
<i>Arcachon Bay</i>	<i>2017</i>	<i>44.580</i>	<i>-1.238</i>	<i>FAR_17</i>	<i>France</i>	<i>30</i>	<i>30</i>	<i>0.074</i>	<i>0.083</i>	<i>0.111</i>	<i>4335</i>	<i>46.9</i>	<i>0.140</i>	<i>0.159</i>	<i>0.120</i>
O Barqueiro	2017	43.722	-7.701	SBA_17	Spain	30	30	0.076	0.085	0.107	4595	49.7	0.140	0.158	0.115
Miño	2017	43.361	-8.206	SMI_17	Spain	30	30	0.073	0.081	0.102	4159	45.0	0.140	0.157	0.110
Anllóns	2017	43.220	-8.943	SAN_17	Spain	30	29	0.073	0.081	0.101	3526	38.1	0.143	0.161	0.110
<i>Ría de Noia</i>	<i>2017</i>	<i>42.790</i>	<i>-8.923</i>	<i>SNO_17</i>	<i>Spain</i>	<i>30</i>	<i>30</i>	<i>0.078</i>	<i>0.087</i>	<i>0.099</i>	<i>4885</i>	<i>52.8</i>	<i>0.139</i>	<i>0.156</i>	<i>0.107</i>
<i>Lombos do Ulla</i>	<i>2017</i>	<i>42.629</i>	<i>-8.775</i>	<i>SLO_17</i>	<i>Spain</i>	<i>30</i>	<i>30</i>	<i>0.075</i>	<i>0.085</i>	<i>0.113</i>	<i>4602</i>	<i>49.8</i>	<i>0.137</i>	<i>0.156</i>	<i>0.120</i>
Sarrido	2017	42.507	-8.826	SSA_17	Spain	30	30	0.074	0.083	0.103	4317	46.7	0.140	0.158	0.112
Vilanova	2017	42.561	-8.831	SVI_17	Spain	27	25	0.072	0.081	0.112	3175	34.3	0.153	0.173	0.118
Campelo	2017	42.421	-8.685	SCA_17	Spain	30	30	0.073	0.082	0.115	4287	46.3	0.140	0.159	0.121
Moaña	2017	42.286	-8.730	SMO_17	Spain	20	19	0.070	0.077	0.088	2361	25.5	0.181	0.200	0.093
Baiona	2018	42.117	-8.822	SBI_18	Spain	17	17	0.075	0.082	0.092	3609	39.0	0.174	0.192	0.097

1008

1009 Table 2. Results of the redundancy analysis (RDA) on the SW British Isles region of *Cerastoderma edule*. Only variables included by the
 1010 forward selection model are shown.

1011

SW British Isles			Complete dataset		Neutral dataset		186 total outlier dataset	
Model	Season	Variable	P-value	Adjusted R ²	P-value	Adjusted R ²	P-value	Adjusted R ²
All seascape variables	Reproductive period	Latitude	-		-		0.005	
		Longitude	0.001	0.102	0.001	0.098	0.001	0.414
		SBS	-		-		0.051	
		BSS	0.004		0.001		-	
	Winter	Latitude	0.001		0.001		0.009	
		Longitude	0.016	0.087	0.015	0.080	0.001	0.423
		SBS	-		-		0.030	
	Summer	Latitude	-		-		0.012	
		Longitude	0.002	0.102	0.003	0.098	0.001	0.410
		SBS	-		-		0.055	
		BSS	0.001		0.001		-	
	Only abiotic variables	Reproductive period	SST	-		-		0.013
SBT			-	0.053	-	0.053	0.001	0.319
BSS			0.002		0.002		-	
NPP			-		-		0.027	
Winter		SST	0.003		-		-	
		SSS	-	0.067	-	0.069	0.041	0.202
		BSS	-		0.010		-	

1013

	NPP	0.017		0.008		0.019	1012
Summer	SST	0.011		0.014		0.001	
	SBT	-	0.086	-	0.083	0.009	0.355
	SBS	-		-		0.083	
	BSS	0.001		0.001		-	

Galicia			Complete dataset		Neutral dataset		51 total outlier dataset	
Model	Season	Variable	P-value	Adjusted R²	P-value	Adjusted R²	P-value	Adjusted R²
All seascape variables	Reproductive period	Latitude	-	-	-	-	0.012	0.124
	Winter	Latitude	-	-	-	-	0.011	0.124
	Summer	Latitude	-	-	-	-	0.008	0.124
Only abiotic variables	Reproductive period	-	-	-	-	-	-	-
	Winter	BSS	-	-	-	-	0.045	0.083
	Summer	-	-	-	-	-	-	-

1014

Adjusted R² and P-value associated to each variable of its selection stage. SST: Sea Surface Temperature; SBT: Sea Bottom Temperature; SSS: Sea Surface Salinity; SBS:

1015

Sea Bottom Salinity; BSS: Bottom Shear Stress; NPP: Net Primary Production.

1016

1017 Table 3. Distribution of divergent outliers and markers correlated with environmental
 1018 variables in the SW British Isles and Galicia across the *Cerastoderma edule* genome
 1019 (version 4.0).

Mega-scaffold (chromosome)	Chromosome length (bp)	Outlier loci (divergent selection)			Markers correlated environmental variables*
		British Isles	Galicia	Shared	
C1	64,609,245	21	11	3	5 (3)
C2	56,319,168	14	2		7 (3)
C3	55,987,847	22	3	1	6 (5)
C4	52,087,795	18	5	2	7 (6)
C5	50,828,891	11	1	1	3 (2)
C6	40,237,005	13	3	2	4 (1)
C7	39,934,596	2	1		
C8	39,684,391	9			2 (1)
C9	39,070,162	11	3		2 (1)
C10	38,264,924	14	8	2	1
C11	38,197,540	2			1
C12	36,327,582	6	1		1 (1)
C13	35,955,507	10			5 (2)
C14	33,816,358	5			
C15	31,726,440	3	1	1	3 (1)
C16	31,510,408	10	8	1	2 (2)
C17	26,587,828	4			2 (1)
C18	22,603,465	1	1	1	
C19	21,711,631	4	1	1	1
Other scaffolds		6	2		2 (1)
Total		186	51	15	54 (30)

1020

1021 * Only detected in SW British Isles; in parentheses those markers also identified as
 1022 outliers for divergent selection

1023

1024

1025

Fig.1

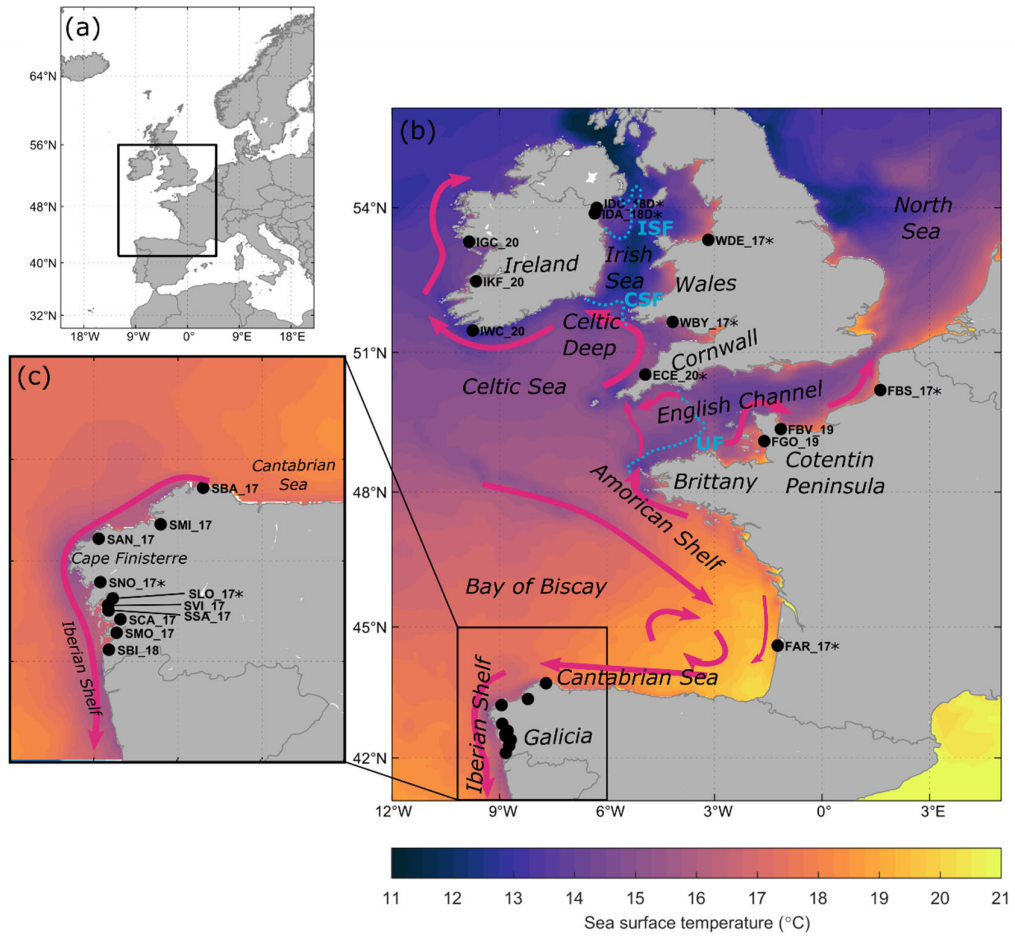


Fig.2

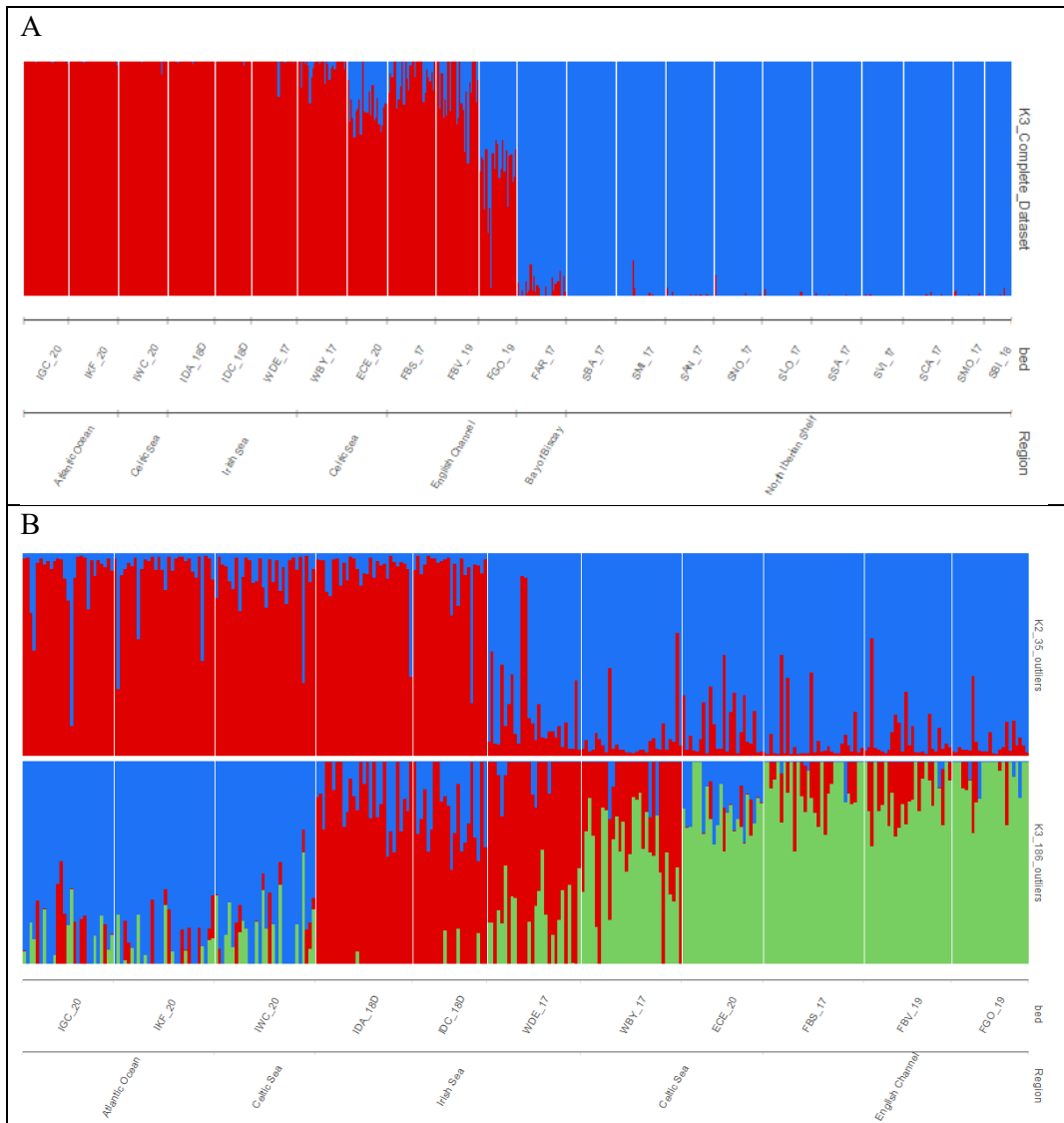
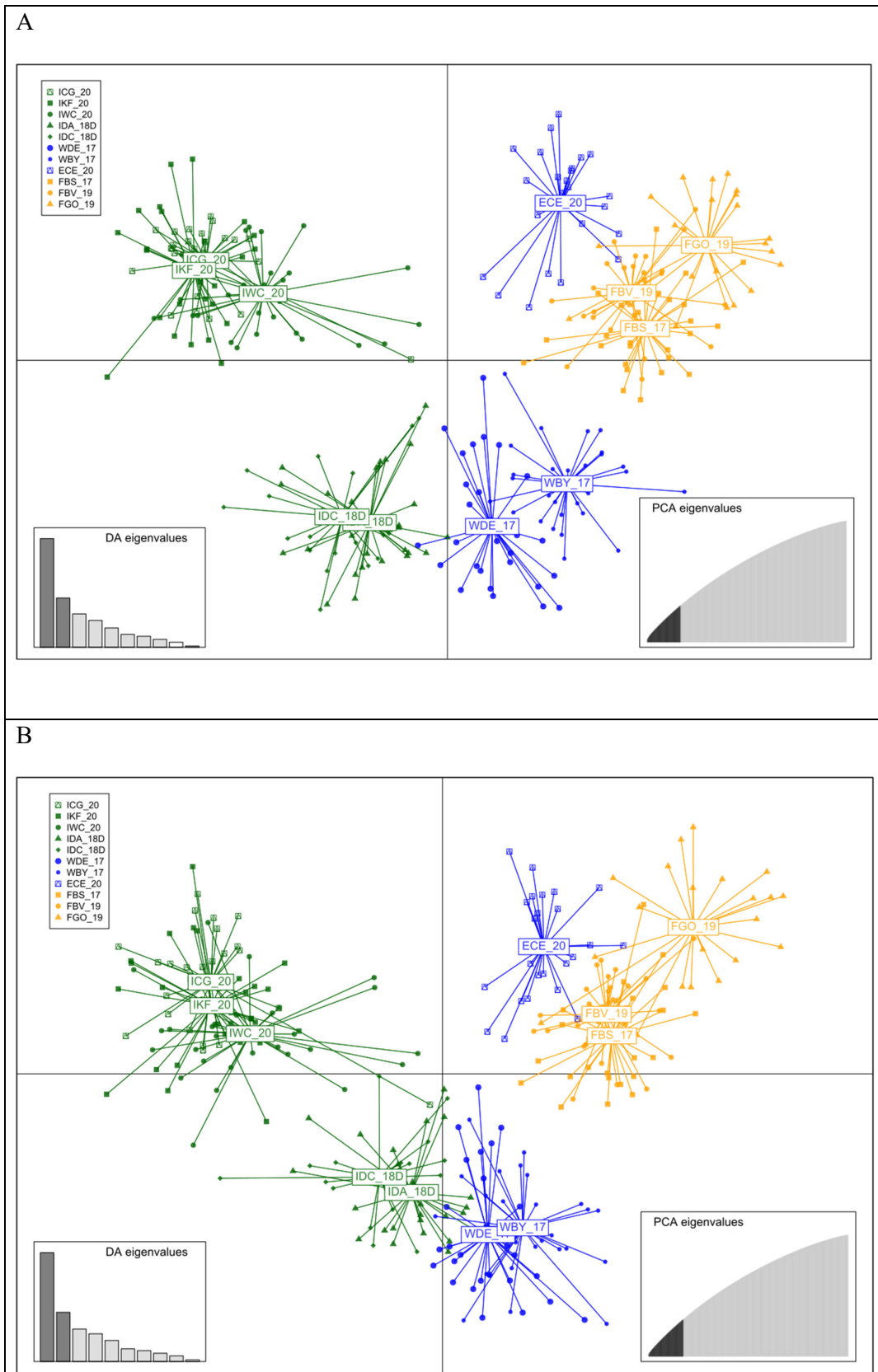
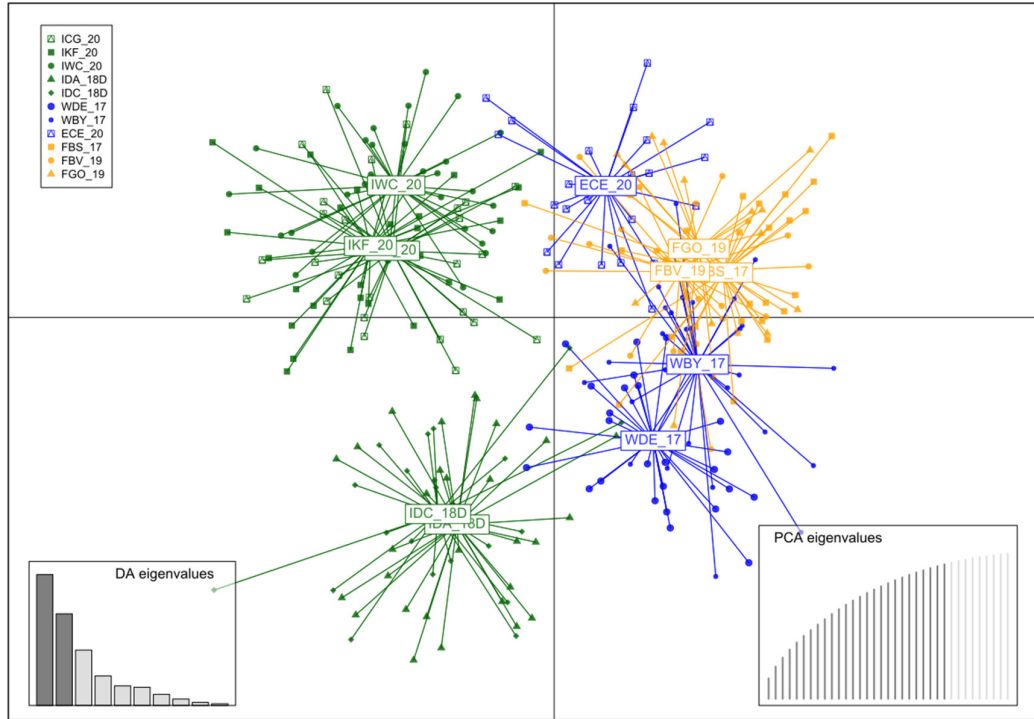


Fig.3



C



D

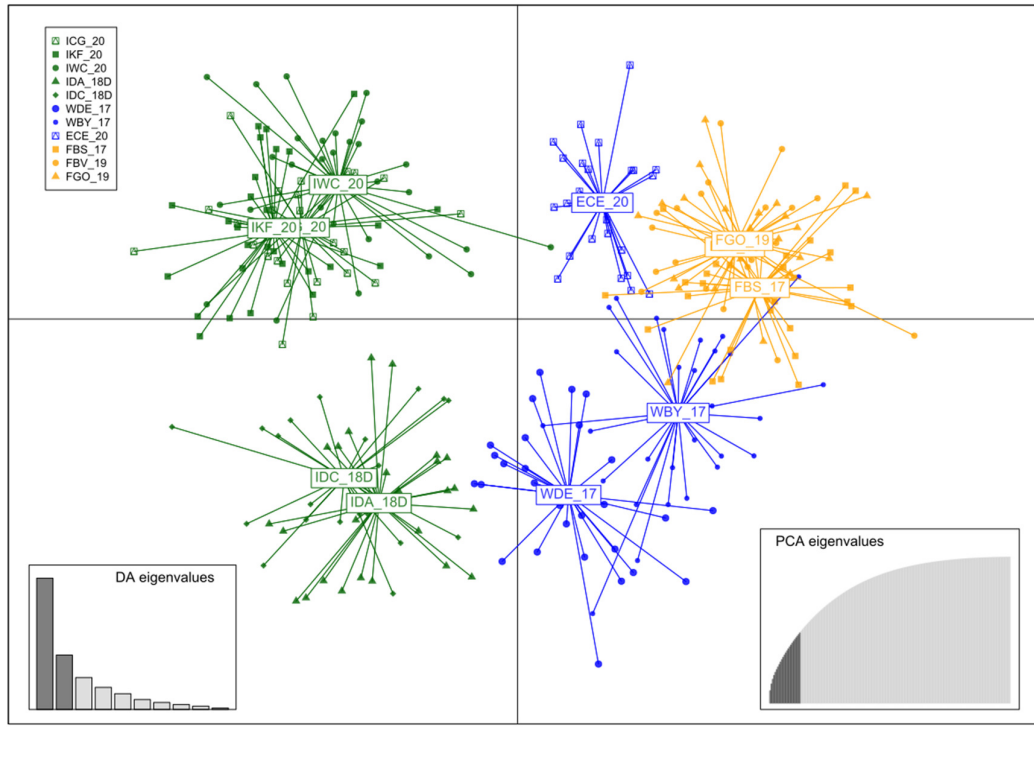


Fig. 4

