Large scale patterns of trematode parasite communities infecting *Cerastoderma edule* along the Atlantic coast from Portugal to Morocco

Simão Correia, Luísa Magalhães, Rosa Freitas, Hocein Bazairi, Meriam Gam, Xavier de Montaudouin

PII: S0272-7714(19)30679-1

DOI: https://doi.org/10.1016/j.ecss.2019.106546

Reference: YECSS 106546

To appear in: Estuarine, Coastal and Shelf Science

Received Date: 10 July 2019

Revised Date: 14 November 2019

Accepted Date: 13 December 2019

Please cite this article as: Correia, Simã., Magalhães, Luí., Freitas, R., Bazairi, H., Gam, M., de Montaudouin, X., Large scale patterns of trematode parasite communities infecting *Cerastoderma edule* along the Atlantic coast from Portugal to Morocco, *Estuarine, Coastal and Shelf Science* (2020), doi: https://doi.org/10.1016/j.ecss.2019.106546.

This is a PDF file of an article that has undergone enhancements after acceptance, such as the addition of a cover page and metadata, and formatting for readability, but it is not yet the definitive version of record. This version will undergo additional copyediting, typesetting and review before it is published in its final form, but we are providing this version to give early visibility of the article. Please note that, during the production process, errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

© 2019 Published by Elsevier Ltd.



Credit author statement

Simão Correia: Formal analysis, Investigation, Writing – Original draft.

Luísa Magalhães: Conceptualization, Methodology, Formal analysis, Investigation, Writing – Review & Editing.

Rosa Freitas: Conceptualization, Methodology, Resources, Supervision.

Hocein Bazairi: Conceptualization, Methodology, Investigation, Resources. Meriam Gam: Investigation.

Xavier de Montaudouin: Conceptualization, Methodology, Formal analysis, Investigation, Resources, Writing – Review & Editing, Supervision.

Journal Pression

1	Large	scale	patterns	of	trematode	parasite	communities	infecting
2	Cerasto	oderma	edule alonç	g the	Atlantic coas	st from Po	rtugal to Moroco	0
3								
4	Correia,	Simãoª,	Magalhães	, Luí	sa ^{a*} , Freitas, I	Rosa ^ª , Baza	iiri, Hocein⁰, Gan	n, Meriam ^c ,
5	de Mont	audouin	, Xavier ^ь					

- ^a Departamento de Biologia & CESAM, Universidade de Aveiro, 3810-193 Aveiro, Portugal
- ^b Université de Bordeaux, EPOC, UMR 5805 CNRS, 2, rue du Pr Jolyet, F-33120 Arcachon,

8 France

- 9 ^c University Mohammed V in Rabat, Laboratory of Biodiversity, Ecology and Genomics,
- 10 BioBio Research Center, Faculty of Sciences, 4 Av. Ibn Battota, Rabat, Morocco
- ^{*}Corresponding author: Tel.: +351 234 370 350; fax. +351 234 372 587
- 12 E-mail address: luisa.magalhaes@ua.pt
- 13

14 Abstract

15 In this study, spatial sampling was performed from North Portugal to South Morocco to 16 analyse trematode communities of the widespread bivalve Cerastoderma edule, host of several trematode species. From the twelve trematode species found in this study, nine 17 were present in multiple aquatic systems demonstrating high trematode dispersal ability, 18 19 driven by the presence of all the hosts. Multivariate analysis related to trematode communities in cockles clustered: 1) Portuguese aquatic systems influenced by cold waters, 20 21 leading to low trematode abundance; 2) coastal systems characterized by dominance of 22 trematode Parvatrema minutum and muddy sediments; 3) lagoons (or bays) with high 23 oceanic influence and high trematode diversity. These findings suggested that, besides host 24 species presence, temperature is an important trigger for parasite infection, with coastal 25 upwelling operating as a shield against trematode infection in Portugal and masking

- 26 latitudinal gradients. Results highlighted the possible consequences of thermal modification
- 27 mediated by oceanographic global circulation change on cockle populations.
- 28
- 29 Keywords
- 30 Cockle; Spatial survey; Host-parasite system; Abiotic drivers; Coastal waters;
- 31 Community composition

32 **1. Introduction**

33 Coastal aquatic systems are known for their high diversity and importance to 34 populations of migratory and resident species (Levin et al., 2001). These aquatic systems have high economic value providing many natural resources and settlement sites for human 35 population (Basset et al., 2013). In these ecosystems, a considerable part of the animal 36 37 biological diversity are parasitic species, corresponding approximately, to 2/5 of eukaryotic 38 total species richness (Dobson et al., 2008; Hudson et al., 2006). These parasites are also ubiquitous in every free-living community. However, they remain neglected in most global 39 40 community analyses. In terms of functional diversity, parasites play a key structuring role in 41 communities, providing information about the ecosystem functioning (Hudson et al., 2006; Marcogliese, 2004; Dairain et al., 2019). They impose adverse effects on their hosts 42 phenotype and health (Marcogliese, 2004) which may result into high mortality at population 43 44 scale (Curtis, 1995; Fredensborg et al., 2005; O'Connell-Milne et al., 2016). At higher scale, 45 they induce significant impact on environmental functions (Thomas et al., 1999), 46 exacerbated in a climate change framework (Mouritsen et al. 2005, Marcogliese 2008). 47 Thus, parasites are considered ecosystem engineers by interfering with the host local 48 population and consequently affecting the value and impact of these species in the 49 ecosystem (Thomas et al., 1999). This role is magnified when the host species itself is also 50 an ecosystem engineer (Dairain et al., 2019).

51 In coastal waters, including bays, estuaries and coastal lagoons, trematodes are the 52 most abundant and common macroparasites (Lauckner, 1983; Roberts et al., 2009). They 53 have a complex and heteroxenous life cycle, infecting more than one host species to 54 complete their life cycle (Bartoli and Gibson, 2007) with alternation between asexual 55 multiplication and sexual reproduction phases (Whitfield, 1993). The typical trematode life 56 cycle includes a free-living larva (miracidium) that hatches from the egg and infects first 57 intermediate hosts, most of the time a mollusc. At this stage, the parasite transforms into a 58 mother sporocyst that will develop, depending on the trematode species, into a mature 59 sporocyst or into redia. Within the sporocysts or rediae, cercariae free-living stages develop

by asexual multiplication. Cercariae will emerge from the first host and reach the second intermediate host, an invertebrate or a vertebrate species depending on the trematode species. The cercariae penetrate the second intermediate host and metamorphose into metacercariae. The cycle is complete when the parasitized second intermediate host is predated by the final host, a vertebrate species. Then, each metacercaria transforms into an adult form that will reproduce sexually and produce eggs (Bartoli and Gibson, 2007; Esch, 2002; Roberts et al., 2009).

67 The complex life cycle described here, shows not only how important host diversity is in the distribution of trematode parasites, but also highlights that environmental parameters 68 must have a key role in the modulation of parasite population dynamics, especially by their 69 70 impact on infective free-living larvae transmission and infection success (Anderson and 71 Sukhdeo, 2010; de Montaudouin et al., 2016a, 2016b; Koprivnikar and Poulin, 2009; Studer and Poulin, 2013). Indeed, a greater host diversity is usually reflected in a greater diversity 72 73 of parasites, particularly for those with complex life cycles (Sukhdeo and Sukhdeo, 2004) 74 and in the same sense, a higher abundance of suitable hosts increases the abundance and 75 prevalence of parasites in the ecosystem (Combes, 1991). As an example, Hechinger and 76 Lafferty (2005) found a positive correlation between bird communities composition (the final 77 hosts of several trematode species) and trematode communities composition in a snail host. 78 Similarly, Thieltges and Reise (2007) demonstrated higher metacercariae abundance (the 79 trematode parasitic stage occurring in the second intermediate host) positively correlated to 80 an increase in abundance and diversity of the higher trophic level host communities. On the 81 other hand, higher host density can also promote a dilution effect and therefore decrease the 82 parasite burden in a specific host (Buck et al., 2017; Magalhães et al., 2016; Mouritsen et al., 2003). Concerning the abiotic environmental factors, different conditions of temperature 83 (Achiorno and Martolrelli, 2016; de Montaudouin et al., 2016a), salinity and pH (Koprivnikar 84 et al., 2010, 2014; Mouritsen, 2002; Studer and Poulin, 2013), among others, can also have 85 an influence on trematode dynamics (Wilson et al., 2002). As an example, a higher cercariae 86 87 emission from the first intermediate host is related to an increase on the water temperature

(de Montaudouin et al., 2016a) and consequently a higher trematode infectivity (Thieltges
and Rick, 2006). Favourable salinity conditions have been reported to benefit, as well,
trematode emergence from the first intermediate host (Koprivnikar et al., 2014; Lei and
Poulin, 2011).

92 Bivalves (along with several other molluscs) are suitable and frequent first and/or second intermediate hosts for trematode parasites (Lauckner, 1983), especially because 93 94 they are easily invaded by trematodes free-living stages through their suspension-feeding 95 activity. Cerastoderma edule, the edible cockle, is among the most common and widely distributed bivalve species of the northeast Atlantic coast, from Norway (Dabouineau and 96 Ponsero, 2011) to Mauritania (Honkoop et al., 2008). Cockles are extensively commercially 97 98 exploited presenting therefore high socio-economic value. Besides, cockles display a crucial 99 ecological role, linking primary producers to higher trophic levels (key species) and acting as ecosystem engineer (Ciutat et al., 2006; Morgan et al., 2013; Rakotomalala et al., 2015). 100 101 This bivalve species acts as first and/ or second intermediate host of several trematode 102 species (de Montaudouin et al., 2009; Longshaw and Malham, 2013) and, when compared 103 to other bivalves, parasitic communities of cockles are particularly diverse and abundant. For 104 these reasons, cockles and their associated trematode fauna are a good model to study 105 host/parasite interactions (e.g. de Montaudouin et al., 2009; Lauckner, 1983; Thieltges et al., 106 2006).

107 The often so called Latitudinal Diversity Gradient, describing that species richness 108 increases from the poles to the tropics, is a pattern widely recognized and applicable to 109 many terrestrial and marine species (Hillebrand, 2004). However, the causes that determine 110 this gradient are not yet fully understood, with hypotheses that go from higher migration 111 rates to the tropics (Jablonski et al., 2006) and/ or a lower climate variation that allows the accumulation of species (Guo and Ricklefs, 2000), to the greater opportunity of species to 112 specialize due to the high productivity and environmental stability, characteristic of tropical 113 habitats (Harrison and Cornell, 2008; Mittelbach et al., 2007). Nonetheless, when referring to 114 115 parasitic species, the knowledge is more limited and the observation of latitudinal patterns,

especially in the case of parasites with complex life cycles, it is not so evident (Poulin and 116 117 Leung, 2011; Stephens et al., 2016). Despite the recent study efforts on latitudinal patterns 118 of parasites (Poulin and Morand, 2004; Studer et al., 2013; Thieltges et al., 2009, 2011; 119 Torchin et al., 2015), results are usually conflicting. For instance, Rohde and Heap (1998) 120 observed an increase in the diversity and abundance of monogenean parasites towards the 121 tropics, however, in the same study, no latitudinal gradient was observed for digenean 122 parasites. Studer et al. (2013) have not found evidences of latitudinal patterns when working with the cockle Austrovenus stutchburyi and trematodes as host-parasite model, while 123 Poulin and Mouritsen (2003) have demonstrated an increase of trematode diversity in a snail 124 125 intermediate host at higher latitudes. An opposite pattern was observed by Thieltges et al. (2009) for trematode communities infecting a crustacean host. 126

127 The present study aimed to provide, for the first time, a large spatial survey of 128 trematode communities infecting *Cerastoderma edule* in the southern range of its distribution 129 area, i.e. from the North of Portugal to the South of Morocco. The tested hypotheses were: 130 1) trematode communities follow a latitudinal gradient driven by abiotic latitudinal-related 131 factors and 2) trematode communities abundance and/ or diversity is dependent on the type 132 of studied system (lagoon vs. estuary vs. bay).

134

2. Material and Methods

135 *2.1 Study Area*

136 The present study was conducted in a total of seventeen semi-diurnal tidal marine systems distributed along Portugal and Morocco coastline (Figure 1). Six aquatic systems 137 138 were sampled in Portugal from July to October 2016: the Ria de Aveiro coastal lagoon, Óbidos coastal lagoon, Albufeira coastal lagoon, Sado estuary, Mira estuary and Ria 139 140 Formosa coastal lagoon. In Morocco, eleven aquatic systems were sampled from November 2007 to January 2008: the Tahaddart estuary, Loukkos estuary, Merja Zerga coastal lagoon, 141 142 Sebou estuary, Oum Er Rbia estuary, Sidi Moussa coastal lagoon, Oualidia coastal lagoon, 143 Souss estuary, Chbika estuary, Khnifiss coastal lagoon and Dakhla bay.

144 Through literature review, information on annual variation of water temperature and 145 salinity for each sampled aquatic system at each respective sampled year (whenever 146 possible) was obtained and gathered in Table 1.

Throughout this manuscript, the term 'coastal lagoon' is used when referring to 147 coastal water bodies, connected to the ocean through one or more inlets and separated by a 148 barrier. The lagoons vary from oligonaline to hypersaline conditions (Gooch et al., 2015). 149 'Estuaries' were considered as semi-enclosed coastal water bodies, that have a connection 150 151 with the open sea and within which seawater is measurably diluted with fresh water derived from land drainage (Pritchard, 1967). When in the presence of large bodies of water that 152 153 enters through the coast and are intimately connected to an ocean by a wide entrance, the 154 term 'bay' was used (UN, 1982).

- 155
- 156

2.2 Field sampling and parasite identification

157 At each sampling area, sediment samples were collected to perform grain size 158 analysis following the method described by Quintino et al. (1989). Silt and clay fraction (fine 159 particles with diameter below 63 μ m) were assessed by wet sieving and the remaining

fractions (sand and gravel) were determined by sieving through a column of five sieves with
decreasing mesh sizes (2.00, 1.00, 0.50, 0.250 and 0.125 mm).

162 Cockles were collected in the intertidal zone (with exception of Óbidos lagoon) using 163 six quadrats (0.25 m² each) randomly placed along a 100 m parallel to the water transect 164 and by sieving the sediment through a 1-mm mesh. The number of cockles per square 165 meter (density) was then estimated. In the Óbidos lagoon, a subtidal area where the quadrat 166 method was impossible to perform, samples were collected with a hand dredge and cockle 167 density (d) was calculated following the equation:

$$d(ind.m^{-2}) = \frac{n}{a \times t \times l}$$

where "n" corresponds to the total number of cockles collected; "a" to the hand dredge area (m²); "t" the mean number of trawls (mean number of times the dredge was dragged in the sediment per launch); and "I" the total number of launches.

171 Shell length (SL) of each cockle was measured to the lowest mm with a calliper. 172 From each aquatic system, according to availability, a variable number (between 11 and 65) 173 of adult cockles (23 – 30 mm) were dissected. In Albufeira lagoon and Mira estuary, due to 174 low abundance of cockles from this length class, twenty cockles representing the SL of each 175 area (16 – 31 mm and 8 – 18 mm, respectively) were dissected. Cockle flesh was squeezed 176 between two glass slides and observed under a stereomicroscope. All trematodes were 177 identified to the species level following de Montaudouin et al. (2009) identification key.

Parasite abundance (mean number of trematode metacercariae per cockle), prevalence (percentage of infected cockles by trematode species) and trematode species richness (number of trematode species present) were calculated according to Bush et al. (1997).

182

183 2.3 Data analysis

184 To test the influence of latitude (independent variable) on the variation of trematode 185 species richness (TSR), trematode total prevalence (TTP) and trematode mean abundance 186 (TMA) (dependent variables), individual regression analyses were performed using the
187 SPSS v.25 software.

188 The Chbika estuary, due to the lack of abiotic information, was excluded from all the 189 following multivariate analyses and used only on trematodes descriptive information.

190 The data matrix with the abundance of trematode per site [abundance per trematode species metacercariae x sampling site] was square root transformed and the Bray-Curtis 191 192 (Legendre and Legendre, 1998) similarity calculated between sites. To identify the biological affinity groups, the resemblance matrix was then analysed using a hierarchical clustering 193 194 analysis tool. The affinity groups were characterized according to environmental and 195 biological features by calculating the mean value of annual maximum and minimum water 196 temperature, annual maximum and minimum water salinity, median grain-size, cockle density at sampling time and by calculating the rarefaction index, i.e. an estimation of the 197 trematode species diversity through standardization of the number of samples (Gotelli and 198 Colwell, 2001). Differences in terms of trematode metacercariae abundance were then 199 tested among affinity groups, type of aquatic system (lagoons vs estuaries) and 200 201 geographical position (north vs. south of strait of Gibraltar). Differences were tested using 202 permutational multivariate analysis of variance (Anderson et al., 2008) following unrestricted 203 permutation of the raw data (9999 permutations) and the calculation of type III sums of 204 squares. Similarity Percentages (SIMPER) were used to characterize the type of aquatic 205 system and the geographical position by the species that most contributed to the 206 dissimilarity between groups. Affinity groups differences were visualized through Principal 207 Coordinates Ordination analysis (PCO) after a distance among centroids resemblance 208 (Clarke and Warwick, 2001). The abiotic variables that were highly correlated (Spearman ρ > [0.7]) to samples ordination were represented as superimposed vectors in the PCO graph. 209

To model the relationship and provide quantitative measures of abiotic and biotic data (annual maximum and minimum water temperature, annual maximum and minimum water salinity and median grain-size and cockle density) on metacercariae community of each aquatic system, a Distance-based linear model (DistLM) was performed (Anderson et

al., 2008). The results were obtained using appropriate permutation (9999 permutations) and
following the step-wise method and Akaike information criterion (AICc). This criterion
balances between data fitness and the most parsimonious model (the model with lower
power loss). Thus, it is considered the best model, among the possible ones, the one
showing the lowest AIC (Symonds and Moussalli, 2011). Distances among aquatic systems
were visualized through a dbRDA plot.

220

All multivariate analyses were performed using the PRIMER v.6 software.

221

222 **3. Results**

223 3.1 Trematode species richness, prevalence and abundance

During this study, 398 cockles were dissected, 280 were infected by a total of twelve trematode species. *Bucephalus minimus* and *Monorchis parvus* infecting cockles as first intermediate host, *Gymnophalus choledochus* using cockles as first and second intermediate host and nine species of trematodes at metacercariae stage, i.e. infecting cockles as second intermediate host, *Curtuteria arguinae, Diphterostomum brusinae, Himasthla continua, H. elongata, H. interrupta, H. quissetensis, Parvatrema minutum, Psilostomum brevicolle* and *Renicola roscovitus* (Table 2).

231 P. minutum was the most prevalent and abundant trematode species, representing approximately 90 % of total metacercariae abundance. This species was found in twelve out 232 233 of the seventeen sampled systems and therefore throughout the whole sampled latitudinal 234 gradient. H. elongata was exclusively found in three aquatic systems located north of 38 °N (northern Portuguese aquatic systems) whereas, H. continua was only present in five 235 236 aquatic systems located south of 35 °N. R. roscovitus was the rarest and the least abundant species, represented by 1 metacercariae identified in the Óbidos lagoon (Figure 2; Table 4). 237 238 C. arguinae, H. quissetensis (representing 5 % of total trematode abundance) and P. 239 brevicolle were common in several aquatic systems south of 37 °N, found in the two aquatic systems nearest to the north Atlantic Ocean – Mediterranean transition (strait of Gibraltar, 36 240

°N) and in several aquatic systems south of this biogeographical barrier (Figure 2). *G. choledochus, D. brusinae* and *H.* interrupta, all the other species infecting cockles as second
intermediate host, showed low abundance, representing each less than 1.5 % of total
abundance (Figure 3).

Overall, trematode species richness (TSR, considering all identified trematode species) increased linearly from north to south but accounting for only 22% of the total variation ($R^2 = 0.22$). When excluding Chbika and Sebou, two small estuaries that displayed the lowest species richness in this study (1 trematode species each), the linear model fitted better with latitude, explaining 61% of total variation (Table 3).

Trematode total prevalence (TTP, considering all identified trematode species) decreased with the latitude as demonstrated by the quadratic equation that explained 82% of the TTP variety ($R^2 = 0.82$; Table 3). The Óbidos lagoon (39 °N, Portugal) showed the lowest TTP value (6%) whereas, almost every aquatic system south of 36 °N presented 100% of TTP (except Loukkos estuary, 35 °N) (Table 4).

Trematode mean abundance (TMA, considering trematode species infecting cockles as second intermediate host) did not follow significantly any model. Nonetheless, a higher TMA was observed in the central zone of the total extent of the sampled area in comparison to the aquatic systems located at the northern and southern edges of the total area sampled in this study. The Óbidos lagoon (39 °N, Portugal) presented the lowest TMA (0.03 ± 0.17 metacercariae.cockle⁻¹) whereas, Oum Er Rbia (33 °N, Morocco) presented the highest TMA (820 ± 595 metacercariae.cockle⁻¹).

262

263

3.2 Multivariate analysis

Cluster analysis, at a similarity distance of 70%, allowed to assemble the different areas in three affinity groups (Figure 4A): group A, subdivided into A1 composed by Tahaddart, Loukkos, Sebou and Souss estuaries and A2 composed by Merja Zerga lagoon, Oum Er Rbia estuary and Khnifiss lagoon; group B, divided in B1 composed by Sidi Moussa

268 and Oualidia lagoons and group B2, splitted into B2.1 composed by Dakhla bay and B2.2 269 composed by all Portuguese systems (Ria de Aveiro lagoon, Óbidos lagoon, Albufeira 270 lagoon, Sado estuary, Mira estuary and Ria Formosa lagoon). Cockle density showed to be 271 significantly higher in group B1 and lower in group B2.1. Sediment median grain-size was 272 significantly higher in group B2.2 and lower in group B2.1 (Table 5). Concerning water physico-chemical characterization, all descriptors showed no significant differences among 273 274 the affinity groups. Nevertheless, group A1 displayed the highest values of maximum and minimum water temperature while, group B2.2 registered the lowest values for the same 275 276 variables. Maximum and minimum water salinity presented the highest values for group B2.1 and the lowest for groups B1 and B2.2, respectively (Table 5). 277

The obtained affinity groups were represented on the PCO ordination graph (Figure 278 279 4B). The axis 1 of the PCO explained 59% of the total variation, separating the group A 280 which is composed by the majority of the aquatic systems south of 36 °N and classified as 281 estuaries (positive side of the axis) from the group B2 (negative side of the axis), that 282 comprises all aquatic systems north of 36 °N, considered as costal lagoons, and the southernmost sampled aquatic system (Dakhla bay, 23 °N). PERMANOVA results confirmed 283 284 a significant difference between sites located North and South from the Gibraltar strait 285 (PERMANOVA: 240.37, p < 0.001) mostly driven by the presence of *P. minutum* and *H.* 286 quissetensis (Table 6) and a significant difference between estuaries and coastal lagoons (PERMANOVA: 38.94, p < 0.001) particularly dependent also on P. minutum and H. 287 288 quissetensis (Table 6). Minimum water temperature and cockle density presented a positive 289 correlation to this axis and sediment median grain-size a negative correlation. In turn, axis 2 290 described 27% of total variation dividing group B2.2 and group A, both in the positive side of 291 the axis, from group B1 in the negative side of the axis. The geographical position (north vs. 292 south) in relation to the strait of Gibraltar (36 °N) showed again an important effect on these groups separation that was also driven by cockle density (negative correlation). 293

294 The best model obtained through the DistLM analysis included the variables 295 sediment median grain-size and cockle density as predictors of trematode community 296 composition, explaining 42 % of the total variation ($R^2 = 0.42$). When transposed to the 297 dbRDA plot, axis 1 (representing 29 % of total variation), led by the differences in terms of 298 median grain size, separated the aquatic systems north of the strait of Gibraltar (Portuguese 299 aquatic systems) from the southern (Moroccan) aquatic systems with the exception of Sidi 300 Moussa coastal lagoon and Merja Zerga estuary. Axis 2 (explaining 13 % of total variation) 301 was positively correlated to cockle density separating two coastal lagoons of Morocco (Sidi 302 Moussa and Oualidia coastal lagoons) and Oum Er Rbia estuary from the other aquatic 303 systems (Figure 5).

Sonution

304 **4. Discussion**

305 Trematode parasites can have a significant impact on population dynamics of their 306 hosts, implying modification on host growth (Wegeberg and Jensen, 2003) or inducing 307 higher mortalities rates (Desclaux et al., 2004), which highlights the importance to study 308 host-parasite interactions. This study is among the few reports on large-scale trematode 309 communities infecting cockles (e.g. de Montaudouin et al., 2009; Magalhães et al., 2015) 310 and represents the first large-scale assessment of latitudinal gradient (and abiotic related 311 factors) as a driver of trematode communities composition in Cerastoderma edule as a host 312 model. In addition, this is the first exhaustive study on trematode species composition in 313 Moroccan cockles.

There are sixteen trematode species infecting Cerastoderma edule in its 314 315 distributional range that use this bivalve as first and/ or second intermediate host (de 316 Montaudouin et al., 2009). In the present study, a total of 12 species were identified along 317 the whole sampled latitudinal gradient. Nine (Bucephalus minimus, Curtuteria arguinae, 318 Diphterostomum brusinae, Gymnophallus choledochus, Himasthla continua, H. interrupta, H. 319 quissetensis, Psilostomum brevicolle and Parvatrema minutum) showed a great dispersal 320 ability, since they were present in a wide range, i.e. along the sampled latitudes, possibly 321 related to migration of birds and fish which are trematode final hosts (Feis et al., 2015). For a 322 trematode species, to complete its life cycle, the three intermediate/final host species should be present in the same aquatic system (Bustnes and Galaktionov, 1999). Nonetheless, 323 biogeographical barriers, such as oceans in the case of continental species, are among the 324 325 most important factors contributing to isolation and prevention of species exchange among 326 regions (Cox and Moore, 1980; Ricklefs and Schluter, 1993).

In the present study, the strait of Gibraltar (36 °N), a known geographical barrier, showed some influence on trematode communities, dividing the observed trematode species in three different categories. The first category gathers seven trematode species (*Monorchis parvus, Bucephalus minimus, Psilostomum brevicolle, Gymnophallus choledochus,*

331 Parvatrema minutum, Curtuteria arguinae and Himasthla interrupta) for which there is no 332 influence of the biogeographical barrier (*i.e.* Gibraltar) and which occurrence is mediated by 333 the predominant presence of the intermediate hosts in the sampled area (Peringia ulvae, 334 Scrobicularia plana, Gobius spp., among others (cf. Table 2)). In this study H. continua was 335 only found south of Gibraltar (36 °N), however, its published distribution (de Montaudouin et 336 al., 2009) places this species in this ubiquitous group of parasites. The life cycle of C. arguinae, is unknown (Desclaux et al., 2006) preventing any hypothesis concerning the link 337 with first intermediate and final hosts distribution. *M. parvus* was only observed at the 338 339 Portuguese coast, still this isolated occurrence is more likely related to the fact that 340 trematodes infecting first intermediate host, usually display very low prevalence (Granovitch 341 and Johannesson, 2000; Islam et al., 2009; Tigga et al., 2014). Indeed, *M. parvus* has been previously recorded along the whole studied latitudinal gradient (de Montaudouin et al., 342 343 2009). The second category, contrastingly, gathers two trematode species with a Southern 344 distribution, Diphterostomum brusinae and H. quissetensis, possibly mediated by the 345 biogeographical barrier and first intermediate host, Tritia reticulata, an abundant gastropod 346 in southern areas, from France to Morocco (but its actual northern latitude corresponds to Baltic Sea (Pizzolla, 2005). Thirdly, H. elongata and Renicola roscovitus displayed a 347 348 Northern distribution likewise their first intermediate host Littorina littorea, a marine 349 gastropod that inhabits the intertidal zone from the White Sea to Gibraltar, on the Atlantic 350 east coast (Johannesson, 1988). Thus, this study confirms that at this latitudinal scale, the 351 distribution of the trematode species is mainly driven by the occurrence of the first 352 intermediate host (de Montaudouin and Lanceleur, 2011; Thieltges, 2007; Thieltges et al., 2009), considering that final hosts have generally a wider distribution area (Magalhães et al., 353 354 2015).

Trematode species richness (TSR) and trematode total prevalence (TTP) increased towards south (from 40 °N to 23 °N). This latitudinal dependent pattern of TSR follows the same trend described for benthic macrofaunal diversity which show higher values with

decreasing latitudes (Macpherson, 2002; Martins et al., 2013; Roy et al., 2004). However,
this pattern was not observed in the case of Chbika and Sebou estuaries, located at 28 °N
and 34 °N, respectively, that presented the lowest TSR.

361 Chbika is not a permanent estuary ('Oued'), which might induce lower macrofaunal and parasite diversity. Conversely, the Sebou river is one of the biggest North African rivers, 362 363 crossing several riverside populations and then impacted by several anthropogenic 364 activities, including agricultural activities (Perrin et al., 2014). The constant use of pesticides 365 and fertilizers by the agricultural activities, plus the connection with untreated sewage from 366 peripheral cities, result in very pollutant loads that contaminate the river to its estuary (Perrin 367 et al., 2014). This presumable poor health and ecological status of this ecosystem might be 368 inducing a low trematode parasites diversity, opposing to the latitudinal trend. Indeed, 369 trematodes, especially free-living stages, are sensitive to water disturbance, reducing their 370 survival, preventing transmission between hosts (Pietrock and Marcogliese, 2003; 371 Koprivnikar et al., 2007) and can be used as indicators of retrograde condition (MacKenzie, 372 1999). Therefore, our results showed that such as free-living species, parasitic fauna seems 373 to follow a latitudinal gradient of decreasing species richness from tropical to extra-tropical areas, one of the oldest recognized ecological patterns (Wright et al., 1993; Willig et al., 374 375 2003). This latitudinal pattern is usually not clear for communities of parasites with complex life cycles (Poulin and Leung, 2011; Studer et al., 2013) due to the complexity of host/ 376 377 parasite interactions, but also because these cycles (and in particular infection success stage) depend on several abiotic factors, including anthropogenic stressors (Rohr et al., 378 2008; Altman and Byers, 2014). 379

Among abiotic factors, temperature is one of the strongest drivers of trematode activity (propagule dispersal, survival, infection (Thieltges and Rick, 2006)) as well as of host infection levels (higher values observed during the warmer seasons (Goater, 1993; Desclaux et al., 2004)). As an example, in Arcachon bay, a synchrony was observed between parasites emergence from the first intermediate host and the infection in the second

385 intermediate host when water temperature was above 15 °C (de Montaudouin et al., 2016a). 386 In the case of the present study, the minimum water temperature of the aquatic systems 387 south of 36 °N was around 15°C, 3 °C above north of 36 °N. This temperature difference can 388 explain why Morocco displays higher TTP and TMA than Portugal. In fact, this study showed that abundance and prevalence of trematode communities presented higher values in 389 390 ecosystems characterized by higher maximum water temperature (affinity group A). In 391 contrast, all the northern sampled areas (Portuguese aquatic systems located north of the 392 strait of Gibraltar) were all gathered in the same affinity group (group B2.2) characterized by 393 the lowest trematode abundance and prevalence. The Portuguese coast proximity to an 394 upwelling front and consequent occurrence of cold vertical currents (Queiroz et al., 2012) 395 may operate as a shield against trematode infection.

396 In the present study, the discrimination of the trematode assemblages was also 397 explained by the cockle density, the sediment median grain-size and the type of aquatic 398 system, some of these factors being possible confounding. Cockles density was higher in 399 the studied areas characterized by higher maximum temperature. This can result either from 400 increased recruitment success (Gam et al., 2010; Magalhães et al., 2016) or from better 401 conditions in terms of food quality and quantity. Nevertheless, the system presenting the 402 highest cockle density (Sidi Moussa lagoon) was also the system with one of the lowest trematode abundance suggesting that when a certain density threshold is passed, cercariae 403 404 are diluted among second intermediate host and consequent lower mean metacercariae infection is observed, as previously highlighted by Magalhães et al. (2016). 405

The type of aquatic system (estuaries, lagoons or bays) showed also to be determinant for the trematode community composition. There was a clear separation between three particular areas South of 36 °N, that presented higher trematode biodiversity: Oualidia and Sidi Moussa lagoon, and Dakhla bay corresponding to group B1 and B2.1, respectively. These areas are classified as coastal lagoons (group B1) or bays (group B2.1) and therefore characterized by higher oceanic influence and hydrodynamics (Kjerfve and

Magill, 1989) and lower water temperature variations, especially Dakhla bay which is located 412 413 in the interaction between the Canary stream and the subtropical ridge currents (Orbi et al., 414 1999). Leung et al. (2009) and Mouritsen and Poulin (2005) referred that these combinations of characteristics are usually related to higher trematode diversity and abundance, which 415 416 was confirmed by our results (the highest rarefaction indices were found in affinity groups B1 and B2.1). Additionally, coastal lagoons tend to be shallower compared to estuaries (Kierfve, 417 1986). This enables light to penetrate up to the bottom, allowing benthic plants to thrive 418 (higher productivity) (Kennish and Paerl, 2010) which could lead to an increase of benthic 419 communities (first hosts of trematode species) diversity and abundance. As previously 420 discussed, higher hosts diversity and abundance is usually correlated to higher trematode 421 422 diversity (Hechinger and Lafferty, 2005).

423 Cockles from Óbidos lagoon, located 40 °N, displayed extreme values of infection, 424 with the lowest TTP and TMA registered. Most of the hosts of the trematode species found 425 in this study have been previously reported in this lagoon (e.g. Carvalho et al., 2011; 426 Lourenço, 2006), however this aquatic system presented the lowest values of water 427 temperature, showing once again the great importance that water temperature has on 428 trematode infection. Besides, this system also has the peculiarity that cockles were collected 429 in a subtidal position. Despite this assumption is based on only one sampling point, these 430 results could suggest that cockles tidal position can also be a driver of trematode infection. In fact, similar results were found by Gam et al. (2008) in Merja Zerga coastal lagoon 431 (Morocco) where cockles living in subtidal zones presented lower trematode infection that 432 433 intertidal sympatric specimens. This was interpreted as a differential distance to upstream 434 intertidal first intermediate host.

The temporal gap and the seasonal differences among some of the samples could of course bias our spatial analysis. However, at these scales, we assume that spatial effects (17° in latitude corresponding to approx. 3,000 km) are stronger than temporal effects (8 years), once trematode composition at a given site is often stable at multiannual scale. As an illustration, de Montaudouin et al. (2012) showed that trematode communities of

440 Arcachon bay, France, were the same after 8 years, the same temporal gap as in this study, 441 due to environmental stability along time. In the same direction, in the Ria de Aveiro coastal 442 lagoon, one of the sampled systems, diversity of trematode was equivalent after 6 years (Freitas et al., 2014; Russell-Pinto et al., 2006). On the other hand, concerning seasonal 443 444 differences, actually, trematode abundance (infecting cockles as second host) follows a seasonal pattern, increasing their infections during the warmer seasons, contrary to what is 445 observed in the colder seasons (Desclaux et al., 2004; Goater, 1993). This could mean a 446 positive influence of the results (in terms of infection intensity) in the Portuguese aquatic 447 systems (systems sampled during summer). However, due to the low trematode abundance 448 on the Portuguese systems, seasonal effects are not so evident (Magalhães et al., 2018). 449 450 Moreover, the present study findings demonstrated a higher trematode abundance on 451 Moroccan aquatic systems (winter samples), which means that seasonal effects were not 452 exacerbated. Finally, we believe that the present findings are of high impact because of the 453 interest in trematode communities knowledge improvement in each sampled area and each 454 country.

455

456 5. CONCLUSION

The present study showed that at the studied scale, trematode abundance and prevalence in cockles seemed to follow a latitudinal pattern. This latitudinal gradient of trematode, however, was more related to temperature than to latitude. In fact, the type of aquatic system, namely coastal lagoons or bays, also demonstrated to have an impact, correlated with the oceanic influence. Nevertheless, the occurrence of trematode species is only possible when all the hosts of that species were present in the ecosystem.

Furthermore, the obtained results highlighted the ubiquity of trematode parasites in the different aquatic systems and alert to a possible change on the trematode fauna composition and abundance in cockle populations driven by thermal modification mediated by oceanographic global circulation. Besides global temperature monitoring, it is then imperative to incorporate trematodes communities assessment in ecological studies, due to

their highly integrative significance, to better predict potential negative impact on host populations and communities sustainability. Nonetheless, due to the seasonal and temporal differences between sampling efforts it is important to perform more studies to fully comprehend what drives trematode communities in a latitudinal gradient, along with studies that support temporal consistency in trematode communities.

473

474 Acknowledgments

This work was supported by the research project COCKLES (EAPA 458/2016 COCKLES 475 Co-Operation for Restoring CocKle SheLlfisheries & its Ecosystem-Services in the Atlantic 476 Area) and by the financial support from the project 18571 CNRST (Morocco) - CNRS 477 478 (France). Simão Correia and Luísa Magalhães benefited from research fellowships funded 479 INTERREG-ATLANTIC program through the research project bv COCKLES (EAPA_458/2016 COCKLES Co-Operation for Restoring CocKle SheLlfisheries & its 480 481 Ecosystem-Services in the Atlantic Area). Rosa Freitas was funded by national funds (OE), 482 through FCT – Fundação para a Ciência e a Tecnologia, I.P., in the scope of the framework 483 contract foreseen in the numbers 4, 5 and 6 of the article 23, of the Decree-Law 57/2016, of 484 August 29, changed by Law 57/2017, of July 19. Thanks are also due to FCT/MCTES for the financial support to CESAM (UID/AMB/50017/2019), through national funds. 485

486 Authors are thankful to all the institutions and fishermen that contributed for the 487 accomplishment of this study.

488 **References**

Achab, M. (2011). Les plages et les vasières des environs des embouchures des oueds
Tahaddart et Gharifa (NW du Maroc) : dynamique morphosédimentaire et impact des
aménagements sur leur évolution récente. In: Sandy beaches and coastal zone
management. Proceedings of the Fifth International Symposium on Sandy Beaches.
Travaux de l'Institut Scientifique, Rabat, 6, 1-12.

Achiorno, C.L. and Martorelli, S.R. (2016). Effect of temperature changes on the cercarialshedding rate of two trematodes. Iheringia, Série Zoologia, 106, e2016020.
doi:<u>10.1590/1678-4766e2016020</u>.

Altman, I. and Byers, J.E. (2014). Large-scale spatial variation in parasite communities
influenced by anthropogenic factors. Ecology, 95, 1876-1887. doi:<u>10.1890/13-0509.1</u>.

Anajjar, E.M., Chiffoleau, J.F., Bergayou, H., Moukrim, A., Burgeot, T., Cheggour, M. (2008).
Monitoring of trace metal contamination in the souss estuary (South Morocco) using the
clams *Cerastoderma edule* and *Scrobicularia plana*. Bulletin of Environmental
Contamination and Toxicology, 80(3), 283–288. doi:10.1007/s00128-008-9364-2.

503 Anderson, T. and Sukhdeo, M. (2010). Abiotic versus biotic hierarchies in the assembly of 504 parasite populations. Parasitology, 135(4), 743-754. doi:10.1017/S0031182009991430.

505 Anderson, M.J., Gorley, R.N., Clarke K.R. (2008). PERMANOVA + for PRIMER: guide to 506 software and statistical methods. University of Auckland and PRIMER-E, Plymouth.

Bao, R., Freitas, M.D.C., Andrade, C. (1999). Separating eustatic from local environmental
effects: A late-Holocene record of coastal change in Albufeira Lagoon, Portugal.
Holocene, 9(3), 341–352. doi:10.1191/095968399675815073.

Bartoli, P. and Gibson, D.I. (2007). Synopsis of the life cycles of Digenea (Platyhelminthes)
from lagoons of the northern coast of the western Mediterranean. Journal of Natural
History, 41(25–28), 1553–1570. doi:10.1080/00222930701500142.

513 Basset, A., Elliott, M., West, R.J., Wilson, J.G. (2013). Estuarine and lagoon biodiversity and

- 514 their natural goods and services. Estuarine, Coastal and Shelf Science, 132(June), 1–4.
- 515 doi:10.1016/j.ecss.2013.05.018.
- 516 Buck, J.C., Hechinger, R.F., Wood, A.C., Stewart, T.E., Kuris, A.M., Lafferty, K.D. (2017).
- 517 Host density increases parasite recruitment but decreases host risk in a snail– 518 trematode system. Ecology, 98(8), 2029-2038.
- Bush, A.O., Lafferty, K.D., Lotz, J.M., Shostak, A.W. (1997). Parasitology Meets Ecology on
 Its Own Terms: Margolis et al. Revisited. The Journal of Parasitology, 83(4), 575.
 doi:10.2307/3284227.
- Bustnes, J.O. and Galaktionov, K. (1999). Anthropogenic influences on the infestation of
 intertidal gastropods by seabird trematode larvae on the southern Barents Sea coast.
 Marine Biology, 133(3), 449–453. doi:10.1007/s002270050484.
- 525 Carvalho, S., Pereira, P., Pereira, F., de Pablo, H., Vale, C., Gaspar, M.B. (2011). Factors
 526 structuring temporal and spatial dynamics of macrobenthic communities in a eutrophic
 527 coastal lagoon (Óbidos lagoon, Portugal). Marine Environmental Research, 71, 97-110.
 528 https://doi.org/10.1016/j.marenvres.2010.11.005
- 529 Ciutat, A., Widdows, J., Readman, J. W. (2006). Influence of cockle *Cerastoderma edule*530 bioturbation and tidal-current cycles on resuspension of sediment and polycyclic
 531 aromatic hydrocarbons. Marine Ecology Progress Series 328: 51-64.
 532 doi:10.3354/meps328051.
- 533 Clarke, K.R. and Gorley, R.N. (2006). PRIMER v.6: User Manual/Tutorial (Plymouth
 534 Routines in Multivariate Ecological Research). PRIMER-E, Plymouth.
- 535 Clarke, K.R. and Warwick, R. (2001). Change in Marine Communities: An approach to 536 statistical analysis and interpretation. PRIMER-E, Plymouth
- 537 Combes, C. (2001). Parasitism: the Ecology and Evolution of Intimate Interactions Chicago:
 538 The University of Chicago Press.

539 Cox, C.B. and Moore, P.D. (1980). Biogeography: An Ecological and Evolutionary Approach.

540 Halsted Press, NY, U.S.A.

- 541 Curtis, L.A. (1995). Growth, trematode parasitism, and longevity of a long-lived marine 542 gastropod (*Ilyanassa obsoleta*). Journal of the Marine Biological Association of the 543 United Kingdom, 75, 913-925. <u>doi:10.1017/S0025315400038248</u>.
- 544 Curtis, L.A., Kinley, L., Tanner, N.L. (2000). Longevity of oversized individuals: growth,
 545 parasitism, and history in an estuarine snail population. Journal of the Marine Biological
 546 Association of the United Kingdom, 80(2000), 811–820.
 547 doi:10.1017/S0025315400002782.
- 548 Dabouineau, L. and Ponsero, A. (2011). Synthesis on biology of Common European Cockle
 549 *Cerastoderma edule*.
- Dairain, A., Legeay, A., de Montaudouin, X. (2019). Influence of parasitism on bioturbation:
 from host to ecosystem functioning. Marine Ecology Progress Series, 619, 201-214.
 <u>doi:10.3354/meps12967.</u>
- 553 de Montaudouin, X. and Lanceleur, L. (2011). Distribution of parasites in their second 554 intermediate host, the cockle *Cerastoderma edule*: Community heterogeneity and 555 spatial scale. Marine Ecology Progress Series 428, 187–199. doi:<u>10.3354/meps09072</u>.
- de Montaudouin, X., Binias, C., Lassalle, G. (2012). Assessing parasite community structure
 in cockles *Cerastoderma edule* at various spatio-temporal scales. Estuarine, Coastal
 and Shelf Science 110, 54-60. doi:10.1016/j.ecss.2012.02.005.
- 559 de Montaudouin, X., Blanchet, H., Desclaux-Marchand, C., Lavesque, N., Bachelet, G.
- 560 (2016a). Cockle infection by *Himasthla quissetensis* I. From cercariae emergence to
 561 metacercariae infection. Journal of Sea Research, 113, 99-107.
- de Montaudouin, X., Blanchet, H., Desclaux-Marchand, C., Bazairi, H., Alfeddy, N.,
 Bachelet, G. (2016b). Cockle infection by *Himasthla quissetensis* II. The theoretical
 effects of climate change. Journal of Sea Research, 113, 108-114.

de Montaudouin, X., Thieltges, D. W., Gam, M., Krakau, M., Pina, S., Bazairi, H.,
Dabouineau, L., Russell-Pinto, F., Jensen, K. T. (2009). Digenean trematode species in
the cockle *Cerastoderma edule*: Identification key and distribution along the NorthEastern Atlantic Shoreline. Journal of the Marine Biological Association of the United
Kingdom, 89(3), 543–556. doi:10.1017/S0025315409003130.

- 570 Desclaux, C., de Montaudouin, X., Bachelet, G. (2004). Cockle *Cerastoderma edule*571 population mortality: role of the digenean parasite *Himasthla quissetensis*. Marine
 572 Ecology Progress Series, 279, 141–150. doi:10.3354/meps279141.
- 573 Desclaux, C., Russell-Pinto, F., de Montaudouin, X., Bachelet, G., (2006). First record and
 574 description of metacercariae of *Curtuteria arguinae* n. sp. (Digenea: Echinostomatidae),
 575 parasite of cockles *Cerastoderma edule* (Mollusca: Bivalvia) in Arcachon bay, France.
- 576 Journal of Parasitology 92(3), 578-587. <u>doi:10.1645/GE-3512.1.</u>
- Dias, J. M., Lopes, J. F., Dekeyser, I. (2000). Tidal propagation in Ria de Aveiro lagoon,
 Portugal. Physics and Chemistry of the Earth, Part B: Hydrology, Oceans and
 Atmosphere, 25(4), 369–374. doi:10.1016/S1464-1909(00)00028-9.
- Dobson, A., Lafferty, K.D., Kuris, A.M., Hechinger, R.F., Jetz, W. (2008). Homage to
 Linnaeus: How many parasites? How many hosts?. Proceedings of the National
 Academy of Sciences of the United States of America, 105, 11482-11489.
 doi:10.1073/pnas.0803232105.
- 584 Dzikowski, R., Diamant, A., Paperna, I. (2003). Trematode metacercariae of fishes as
 585 sentinels for a changing limnological environment. Diseases of Aquatic Organisms, 55,
 586 145-150.
- 587 Esch, G.W. (2002). The Transmission of Digenetic Trematodes: Style, Elegance,
 588 Complexity. Integrative and Comparative Biology, 42(2), 304–312.
 589 doi:10.1093/icb/42.2.304.
- 590 Feis, M.E., Thieltges, D.W., Olsen, J.L., de Montaudouin, X., Jensen, K.T., Bazairi, H.,

- 591 Culloty, S.C., Luttikhuizen, P.C. (2015). The most vagile host as the main determinant
 592 of population connectivity in marine macroparasites. Marine Ecology Progress Series,
 593 520, 85–99. doi:10.3354/meps11096.
- Fortunato, A. B., Nahon, A., Dodet, G., Rita Pires, A., Conceição Freitas, M., Bruneau, N.,
 Azevedo, A., Bertin, X., Benevides, P., Andrade, C., Oliveira, A. (2014). Morphological
 evolution of an ephemeral tidal inlet from opening to closure: The Albufeira inlet,
 Portugal. Continental Shelf Research, 73, 49–63. doi:10.1016/j.csr.2013.11.005.
- Fredensborg, B.L., Mouritsen, K.N., Poulin, R. (2006). Relating bird host distribution and
 spatial heterogeneity in trematode infections in an intertidal snail-from small to large
 scale. Marine Biology, 149, 275-283. doi:10.1007/s00227-005-0184-1.
- Freitas, R., Martins, R., Campino, B., Figueira, E., Soares, A.M.V.M., Montaudouin, X.
 (2014). Trematode communities in cockles (*Cerastoderma edule*) of the Ria de Aveiro
 (Portugal): Influence of inorganic contamination. Marine Pollution Bulletin, 82, 117–126.
 doi:10.1016/j.marpolbul.2014.03.012.
- Gam, M., Bazaïri, H., Jensen, K. T., de Montaudouin, X. (2008). Metazoan parasites in an
 intermediate host population near its southern border: The common cockle
 (*Cerastoderma edule*) and its trematodes in a Moroccan coastal lagoon (Merja Zerga).
 Journal of the Marine Biological Association of the United Kingdom, 88(2), 357–364.
 doi:10.1017/S0025315408000611.
- Gam, M., de Montaudouin, X., Bazairi, H. (2010). Population dynamics and secondary
 production of the cockle *Cerastoderma edule*: A comparison between Merja Zerga
 (Moroccan Atlantic Coast) and Arcachon Bay (French Atlantic Coast). Journal of Sea
 Research, 63(3–4), 191–201. doi:10.1016/j.seares.2010.01.003.
- Gamito, S. and Erzini, K. (2005). Trophic food web and ecosystem attributes of a water
 reservoir of the Ria Formosa (south Portugal). Ecological Modelling, 181(4), 509–520.
 doi:10.1016/j.ecolmodel.2004.02.024.

- Geawhari, M.A., Huff, L., Mhammdi, N., Trakadas, A., Ammar, A. (2014). Spatial-temporal
 distribution of salinity and temperature in the Oued Loukkos estuary, Morocco: using
 vertical salinity gradient for estuary classification. Computational Statistics, 3(1), 1–9.
 doi:10.1186/2193-1801-3-643.
- Goater, C.P. (1993). Population biology of *Meiogymnophallus minutus* (Trematoda:
 Gymnophallidae) in cockles from the Exe Estuary. Journal of the Marine Biological
 Association of the United Kingdom, 73, 163–177. doi:10.1017/S0025315400032707.
- Gooch, D.G., Lillebø, A., Stålnacke, P., Alves, F.L., Bielecka, M., Krysanova, V. (2015).
 Challenges in the Policy–Environment–Modelling management context (Chapter 01).
 In: Coastal Lagoons in Europe: Integrated Water Resource Strategies" (Eds.: Lillebø A.,
 Stålnacke P., Gooch G.D.). International Water Association (IWA), UK. 1-9. ISBN:
 9781780406282.
- Gotelli, N.J. and Colwell, R.K. (2001). Quantifying biodiversity: procedures and pitfalls in the
 measurement and comparison of species richness. Ecological Letters, 4, 379–391.
 doi:10.1046/j.1461-0248.2001.00230.x.
- Granovitch, A. and Johannesson, K. (2000). Digenetic trematodes in four species of Littorina
 from the West Coast of Sweden. Ophelia, 53(1), 55–65.
 doi:10.1080/00785326.2000.10409435.
- Guo, Q.F. and Ricklefs, R.E. (2000). Species richness in plant genera disjunct between
 temperate eastern Asia and North America. Botanical Journal of the Linnean Society,
- 637 134, 401–423. doi:<u>10.1111/j.1095-8339.2000.tb00538.x</u>.
- Haddout, S., Maslouhi, A., Magrane, B., Igouzal, M. (2016). Study of salinity variation in the
 Sebou River Estuary (Morocco). Desalination and Water Treatment, 57(36), 17075–
 17086. doi:10.1080/19443994.2015.1091993.
- Harrison, S. and Cornell, H. (2008). Toward a better understanding of the regional causes of
 local community richness. Ecology Letters, 11(9), 969–979. doi:<u>10.1111/j.1461-</u>

643 0248.2008.01210.x.

- Hechinger, R.F. and Lafferty, K.D. (2005). Host diversity begets parasite diversity: bird final
 hosts and trematodes in snail intermediate hosts. Proceedings of the Royal Society B,
 272, 1059–1066. doi:10.1098/rspb.2005.3070.
- Hillebrand, H. (2004). On the Generality of the Latitudinal Diversity Gradient. The American
 Naturalist, 163(2), 192-211. doi:10.1086/381004.
- Hilmi, K., Koutitonsky, V.G., Orbi, A., Lakhdar, J.I., Chagdali, M. (2005). Oualidia lagoon,
 Morocco: An estuary without a river. African Journal of Aquatic Science, 30(1), 1–10.
 doi:10.2989/16085910509503828.
- Honkoop, P.J.C, Berghuis, E.M., Holthuijsen, S., Lavaleye, M.S.S., Piersma, T. (2008).
 Molluscan assemblages of seagrass-covered and bare intertidal flats on the Banc
 d'Arguin, Mauritania, in relation to characteristics of sediment and organic matter.
 Journal of Sea Research, 60, 235-243. doi:10.1016/j.seares.2008.07.005.
- Hudson, P.J., Dobson, A.P., Lafferty, K.D. (2006). Is a healthy ecosystem one that is rich in
 parasites?. Trends in Ecology & Evolution, 21, 381-385. doi:10.1016/j.tree.2006.04.007.
- Islam, Z., Alam, M.Z., Akter, S., Roy, B.C., Mondal, M.M.H. (2012). Distribution patterns of
 vector snails and Trematode cercaria in their vectors in some selected areas of
 Mymensingh. Journal of Environmental Science and Natural Resources, 5(2), 37-46.
 doi:10.3329/jesnr.v5i2.14599.
- Jablonski, D., Roy, K., Valentine, J.W. (2006). Out of the tropics: evolutionary dynamics of
 the latitudinal diversity gradient. Science, 314, 102–106. doi:10.1126/science.1130880.
- Johannesson, K. (1988). The paradox of Rockall why is a brooding gastropod (*Littorina* saxatilis) more widespread than one having a planktonic larval dispersal stage (*L. littorea*)?. Marine Biology, 99, 507–513.
- 667 Kennish, M.J. and Paerl, H.W. (2010). Coastal Lagoons Critical Habitats of Environmental

- 668 Change. In: Coastal lagoons: critical habitats of environmental change. M. J. Kennish 669 and H. W. Paerl (eds.), 1-16. USA: Marine science series, CRC Press.
- Khalki, A. E. and Moncef, M. (2007). Etude du peuplement de copepodes de l'estuaire de
 l'Oum Er Rbia (cote Atlantique du Maroc): Effets des marees et des lachers de
 barrages. Lebanese Science Journal, 8(1), 3–18.
- Kjerfve, B. (1986). Comparative oceanography of coastal lagoons. In: Estuarine Variability.
 D. A. Wolfe (ed.), 63-81. USA: Academic Press.
- Kjerfve, B. and Magill, K.E. (1989). Geographic and Hydrodynamic Characteristics of
 Shallow Coastal Lagoons. Marine Geology, 88, 187–199.
 doi:10.1038/bonekey.2013.128.
- Koprivnikar, J. and Poulin, R. (2009). Effects of temperature, salinity and water level on the
 emergence of marine cercariae. Parasitology Research, 10, 957-965.
- 680 Koprivnikar, J., Ellis, D., Shim, K.C., Forbes, M.R. (2014) Effects of temperature and salinity
- on emergence of *Gynaecotyla adunca* cercariae from the intertidal gastropod *Ilyanassa obsoleta*. Journal of Parasitology, 100, 242-245. doi:<u>10.1645/13-331.1</u>.
- Koprivnikar, J., Forbes, M.R., Baker, R.L. (2007). Contaminant effects on host-parasite
 interactions: atrazine, frogs, and trematodes. Environmental Toxicology and Chemistry,
 26, 2166-2170. doi:10.1897/07-220.1.
- Koprivnikar, J., Lim, D., Fu, C., Brack, S.H.M. (2010). Effects of temperature, salinity, and
 pH on the survival and activity of marine cercariae. Parasitology Research, 106, 1167-
- 688 1177. doi:<u>10.1007/s00436-010-1779-0</u>.
- Lauckner, G. (1983). Diseases of mollusca: bivalvia. In: Kinne, O. (Ed.), Diseases of Marine
 Animals. Biologische Helgoland, Hamburg, Germany, 477–879.
- Lauckner, G. (1987). Ecological effects of larval trematode infestation on littoral marine
 invertebrate populations. International Journal for Parasitology, 17(2), 391–398.

693 doi:10.1016/0020-7519(87)90114-7.

- Legendre, P. and Legendre, L. (1998). Numerical ecology. Developments in Environmental
 Modelling. Elsevier Science, Amsterdam.
- Lei, F. and Poulin, R. (2011) Effects of salinity on multiplication and transmission of an
 intertidal trematode parasite. Marine Biology, 158, 995-1003.
- Leung, T.L.F., Donald, K.M., Keeney, D.B., Koehler, A.V, Peoples, R.C., Poulin, R. (2009).
 Trematode parasites of Otago Harbour (New Zealand) soft-sediment intertidal
 ecosystems: Life cycles, ecological roles and DNA barcodes. New Zealand Journal of
 Marine and Freshwater Research, 43(4), 857–865. doi:10.1080/00288330909510044.
- Levin, L.A., Boesch, D.F., Covich, A., Dahm, C., Erseus, C., Ewel, K.C., Kneib, R.T.,
 Moldenke, A., Palmer, M.A., Snelgrove, P., Strayer, D., Weslawski, J.M. (2001). The
 function of marine critical transition zones and the importance of sediment biodiversity.
 Ecosystems, 4(5), 430-451. <u>doi:10.1007/s10021-001-0021-4.</u>
- Lillebø, A.I., Ameixa, O.M.C.C., Sousa, L.P., Sousa, A.I., Soares, J.A., Dolbeth, M., Alves,
 F.L. (2015). The physio-geographical background and ecology of Ria de Aveiro. In:
 Lillebø, A.I., Stålnacke, P., Gooch, G.D. (Eds.), Coastal Lagoons in Europe: Integrated
 Water Resource Strategies. International Water Association (IWA), London, UK, 21–28.
- Longshaw, M. and Malham, S.K. (2013). A review of the infectious agents, parasites,
 pathogens and commensals of European cockles (*Cerastoderma edule* and *C. glaucum*). Journal of the Marine Biological Association of the United Kingdom, 93(1),
 227–247. doi:10.1017/S0025315412000537.
- Lourenço, P.M. (2006). Seasonal abundance of aquatic birds at Óbidos lagoon. Airo, 16, 2329.
- Maanan, M., Zourarah, B., Carruesco, C., Aajjane, A., Naud, J. (2004). The distribution of
 heavy metals in the Sidi Moussa lagoon sediments (Atlantic Moroccan Coast). Journal
 of African Earth Sciences, 39(3–5), 473–483. doi:10.1016/j.jafrearsci.2004.07.017.

719	Machado,	M.M.	and	Costa,	A.M.	(1994).	Enzymatic	and	morphological	criteria	for
720	disting	guishin	g betv	ween <i>Ca</i>	rdium	edule and	d C. galucun	n of th	ne Portuguese c	oast. <i>M</i> a	irine
721	Biolog	y, 120	, 535-	-544.							

MacKenzie, K. (1999). Parasites as pollution indicators in marine ecosystems: a proposed
early warning system. Marine Pollution Bulletin, 38, 955–959. doi:10.1016/s0025326x(99)00100-9.

Macpherson, E. (2002). Large-scale species-richness gradients in the Atlantic Ocean.
Proceedings of the Royal Society: Biological Sciences, 269(1501), 1715–1720.
doi:10.1098/rspb.2002.2091.

Magalhães, L., Correia, S., de Montaudouin, X., Freitas, R. (2018). Spatio-temporal variation
of trematode parasites community in *Cerastoderma edule* cockles from Ria de Aveiro

730 (Portugal). Environmental Research, 164, 114-123. doi:10.1016/j.envres.2018.02.018.

- Magalhães, L., Freitas, R., de Montaudouin, X. (2015). Review: *Bucephalus minimus*, a
 deleterious trematode parasite of cockles *Cerastoderma* spp. Parasitology Research,
 114, 1263-1278. doi:10.1007/s00436-015-4374-6.
- Magalhães, L., Freitas, R., de Montaudouin, X. (2016). Cockle population dynamics:
 recruitment predicts adult biomass, not the inverse. Marine Biology, 163(1), 1–10.
 doi:10.1007/s00227-015-2809-3.
- Malhadas, M. S., Leitão, P. C., Silva, A., Neves, R. (2009). Effect of coastal waves on sea
 level in Óbidos Lagoon, Portugal. Continental Shelf Research, 29(9), 1240–1250.
 doi:10.1016/j.csr.2009.02.007.
- Marcogliese, D. J. (2004). Parasites: Small Players with Crucial Roles in the Ecological
 Theater. EcoHealth, 1, 151-164. doi:10.1007/s10393-004-0028-3.
- Marcogliese, D.J. (2008). The impact of climate change on parasites and infectious disease
 of aquatic animals. Revue Scientifique et Technique de l'Office International des
 Epizooties, 27, 467-484.

745	Martins, F., Leitão, P.C., Silva, A., Neves, R. (2001). 3D modelling in the Sado estuary using
746	a new generic vertical discretization approach. Oceanologica Acta, 24, 51-62.
747	doi:10.1016/S0399-1784(01)00092-5.

- Martins, R., Magalhães, L., Peter, A., San Martín, G., Rodrigues, A. M., Quintino, V. (2013).
 Diversity, distribution and ecology of the family Syllidae (Annelida) in the Portuguese
 coast (Western Iberian Peninsula). Helgoland Marine Research, 67(4), 775–788.
 doi:10.1007/s10152-013-0362-3.
- Mittelbach, G.G., Schemske, D.W., Cornell, H.V., Allen, A.P., Brown, J.M., Bush, M.B.,
 Harrison, S.P., Hurlbert, A.H., Knowlton, N., Lessios, H.A., McCain, C.M., McCune,
 A.R., McDade, L.A., McPeek, M.A., Near, T.J., Price, T.D., Ricklefs, R.E., Roy, K., Sax,
 D.F., Schluter, D., Sobel, J.M., Turelli, M. (2007). Evolution and the latitudinal diversity
 gradient: speciation, extinction and biogeography. Ecology Letters, 10, 315-331.
 doi:10.1111/j.1461-0248.2007.01020.x.
- Morgan, E., O'Riordan, R.M., Culloty, S.C. (2013). Climate change impacts on potential
 recruitment in an ecosystem engineer. Ecology and Evolution, 3, 581-594.
 doi:10.1002/ece3.419.
- Mouritsen, K.N. (2002). The *Hydrobia ulvae-Maritrema subdolum* association: influence of
 temperature, salinity, light, water-pressure and secondary host exudates on cercarial
 emergence and longevity. Journal of Helminthology, 76, 341-347.
 doi:<u>10.1079/JOH2002136</u>.
- Mouritsen, K.N. and Poulin, R. (2002). Parasitism, community structure and biodiversity in
 intertidal ecosystems. Parasitology, 124, S101–S117.
 doi:10.1017/S0031182002001476.
- Mouritsen, K.N. and Poulin, R. (2005). Parasitism can influence the intertidal zonation of non-host organisms. Marine Biology, 148(1), 1–11. doi:10.1007/s00227-005-0060-z.
- 770 Mouritsen, K.N., McKechnie, S., Meenken, E., Toynbee, J. L., Poulin, R. (2003). Spatial

heterogeneity in parasite loads in the New Zealand cockle: The importance of host
condition and density. Journal of the Marine Biological Association of the United
Kingdom, 83(2), 307–310. doi:10.1017/S0025315403007124h.

- Mouritsen, K.N., Tompkins, D.M., Poulin, R. (2005). Climate warning may cause a parasiteinduced collapse in coastal amphipod populations. Oecologia, 146, 476-484.
- O'Connell-Milne, S.A., Poulin, R., Savage, C., Rayment, W. (2016). Reduced growth, body
 condition and foot length of the bivalve *Austrovenus stutchburyi* in response to parasite
 infection. Journal of Experimental Marine Biology and Ecology, 474, 23-28.
 doi:10.1016/j.jembe.2015.09.012.
- Oliveira, A., Fortunato, A.B., Rego, J.R.L. (2006). Effect of morphological changes on the
 hydrodynamics and flushing properties of the Óbidos lagoon (Portugal). Continental
 Shelf Research, 26(8), 917–942. doi:10.1016/j.csr.2006.02.011.
- Orbi, A., Dafir, J.E., Berraho, A., Sarf, F. (1999). Etude Pluridisciplinaire de la Baie de
 Dakhla. Note d'information (interne) sur la planification de l'aquaculture dans la baie de
 Dakhla, Maroc.
- Perrin, J.L., Raïs, N., Chahinian, N., Moulin, P., Ijjaali, M. (2014). Water quality assessment
 of highly polluted rivers in a semi-arid Mediterranean zone Oued Fez and Sebou River
 (Morocco). Journal of Hydrology, 510, 26–34. doi:10.1016/j.jhydrol.2013.12.002.
- Pietrock, M. and Marcogliese, D. J. (2003). Free-living endohelminth stages: at the mercy of
 environmental conditions. Trends in Parasitology, 19, 293-299. doi:10.1016/S14714922(03)00117-X.
- Pizzolla, P.F. (2005). *Tritia reticulata* Netted dog whelk. In Tyler-Walters H. and Hiscock K.
 (eds) Marine Life Information Network: Biology and Sensitivity Key Information
 Reviews. Plymouth: Marine Biological Association of the United Kingdom.
- Poulin, R. and Leung, T.L.F. (2011). Latitudinal gradient in the taxonomic composition of
 parasite communities. Journal of Helminthology, 85, 228-233.

797 doi:10.1017/s0022149x10000696.

- Poulin, R. and Morand, S. (2004). Parasite biodiversity. Washington DC, Smithsonian
 Institution Press.
- 800 Pritchard, D.W. (1967). What Is an Estuary: Physical Viewpoint. In Estuaries. AAAS,
 801 Washington DC.
- Queiroz, N., Humphries, N.E., Noble, L.R., Santos, A.M., Sims, D.W. (2012). Spatial
 dynamics and expanded vertical niche of blue sharks in oceanographic fronts reveal
 habitat targets for conservation. PLoS ONE, 7(2). doi:10.1371/journal.pone.0032374.
- Quintino, V., Rodrigues, A.M., Gentil, F., (1989). Assessment of macrozoobenthic
 communities in the lagoon of Óbidos, western coast of Portugal. Scientia Marina, 53,
 645–654.
- Rakotomalala, C., Grangeré, K., Ubertini, M., Forêt, M., Orvain, F. (2015). Modelling the
 effect of *Cerastoderma edule* bioturbation on microphytobenthos resuspension towards
 the planktonic food web of estuarine ecosystem. Ecological Modelling, 316, 155–167.
 doi:10.1016/j.ecolmodel.2015.08.010.
- Ricklefs, R.E. and Schluter, D. (1993). Species diversity: regional and historical influences.
 In: Ecological Communities: Historical and Geographical Perspectives (eds Ricklefs,
 R.E. & Schluter, D.). University of Chicago Press, Chicago, IL, 350-363.
- Roberts, L. S., Janovy, J., Schmidt, G. D. (2009). Gerald D. Schmidt & Larry S. Roberts'
 foundations of parasitology. Boston: McGraw-Hill.
- Rohde, K. and Heap, M. (1998). Latitudinal differences in species and community richness
 and in community structure of metazoan endo-and ectoparasites of marine teleost fish.
 International Journal for Parasitology, 28(3), 461-474.
- 820 Rohr, J.R., Raffel, T.R., Sessions, S.K., Hudson, P.J. (2008). Understanding the net effects
- of pesticides on amphibian trematode infections. Ecological Applications, 18(7), 1743-

822 1753. doi:10.1890/07-1429.1.

- Roy, K., Jablonski, D., Valentine, J.W. (2004). Beyond Species Richness: Biogeographic
 Patterns and Biodiversity Dynamics Using Other Metrics of Diversity. In: Lomolino,
 M.V., Heaney, L.R. (Eds.), Frontiers of Biogeography: New Directions in the Geography
 of Nature. Sinauer, Sunderland, USA, 151–170.
- Russell-Pinto, F., Goncalves, J.F., Bowers, E. (2006). Digenean larvae parasitizing
 Cerastoderma edule (Bivalvia) and Nassarius reticulatus (Gastropoda) from Ria de
 Aveiro, Portugal. Journal of Parasitology, 92, 319–332. doi:10.1645/ge-3510.1.
- Schulte-Oehlmann, U., Oehlmann, J., Fioroni, P., Bauer, B. (1997). Imposex and
 reproductive failure in *Hydrobia ulvae* (Gastropoda: Prosobranchia). Marine Biology,
 128(2), 257–266. doi:10.1007/s002270050090.
- Semlali, A., Chafik, A., Talbi, M., Budzinski, H. (2012). Origin and Distribution of Polycyclic
 Aromatic Hydrocarbons in Lagoon Ecosystems of Morocco. The Open Environmental
 Pollution & Toxicology Journal, 3, 37–46. doi:<u>10.2174/1876397901203010037</u>.
- Silva, G., Costa, J.L., de Almeida, P.R., Costa, M.J. (2006). Structure and dynamics of a
 benthic invertebrate community in an intertidal area of the Tagus estuary, western
 Portugal: A six year data series. Hydrobiologia. 555(1), 115–128. doi:10.1007/s10750005-1110-8.
- Sousa, W.P. (1983). Host life history and the effect of parasitic castration on growth: a field
 study of *Cerithidea californica*. Journal of Experimental Marine Biology and Ecology, 73,
- 842 273–296. <u>doi:10.1016/0022-0981(83)90051-5</u>.
- Stephens, P.R., Altizer, S., Smith, K.F., Alonso Aguirre, A., Brown, J.H., Budischak, S.A.,
 Byers, J.E., Dallas, T.A., Jonathan Davies, T., Drake, J.M., Ezenwa, V.O., Farrell, M.J.,
 Gittleman, J.L., Han, B.A., Huang, S., Hutchinson, R.A., Johnson, P., Nunn, C.L.,
 Onstad, D., Park, A., Vazquez-Prokopec, G.M., Schmidt, J.P., Poulin, R. (2016). The
 macroecology of infectious diseases: a new perspective on global-scale drivers of

848 pathogen distributions and impacts. Ecological Letters, 19, 1159-1171.

849 doi:<u>10.1111/ele.12644.</u>

Studer, A. and Poulin, R. (2013) Cercarial survival in an intertidal trematode: a multifactorial
experiment with temperature, salinity and ultraviolet radiation. Parasitology Research,
112, 243-249. doi:10.1007/s00436-012-3131-3.

Studer, A., Widmann, M., Poulin, R., Krkošek, M. (2013). Large scale patterns of trematode
parasitism in a bivalve host: No evidence for a latitudinal gradient in infection levels.
Marine Ecology Progress Series, 491, 125–135. doi:org/10.3354/meps10483.

Sukhdeo, M.V. and Sukhdeo, S.C. (2004). Trematode behaviours and the perceptual worlds
of parasites. Canadian Journal of Zoology, 82(2), 292–315. doi:org/10.1139/z03-212.

Symonds, M.R.E. and Moussalli, A. (2011). A brief guide to model selection, multimodel
inference and model averaging in behavioural ecology using Akaike's information
criterion. Behavioral Ecology and Sociobiology, 65(1), 13-21.

Thieltges, D. (2007). Habitat and transmission – effect of tidal level and upstream host
density on metacercarial load in an intertidal bivalve. Parasitology, 134(4), 599-605.
doi:10.1017/S003118200600165X.

Thieltges, D.W. and Reise, K. (2007). Spatial heterogeneity in parasite infections at different
spatial scales in an intertidal bivalve. Oecologia, 150(4), 569–581. doi:10.1007/s00442006-0557-2.

Thieltges, D.W. and Rick, J. (2006). Effect of temperature on emergence, survival and infectivity of cercariae of the marine trematode *Renicola roscovita* (Digenea: Renicolidae). Diseases of Aquatic Organisms, 73, 63–68. doi:10.3354/dao073063.

Thieltges, D.W., Ferguson, M.A., Jones, C.S., Noble, L.R., Poulin, R. (2009).
Biogeographical patterns of marine larval trematode parasites in two intermediate snail
hosts in Europe. Journal of Biogeography, 36(8), 1493-1501. doi:10.1111/j.13652699.2008.02066.x.

35

- Thieltges, D.W., Hof, C., Dehling, D.M., Brändle, M., Brandl, R., Poulin, R. (2011). Host diversity and latitude drive trematode diversity patterns in the European freshwater fauna. Global Ecology and Biogeography, 20, 675-682.
- Thieltges, D.W., Krakau, M., Andresen, H., Fottner, S., Reise, K. (2006) Macroparasite
 community in molluscs of a tidal basin in the Wadden Sea. Helgoland Marine Research,
 60, 307–316.
- Thomas, F., Poulin, R., De Meeüs, T., Guégan, J., Renaud, F. (1999). Parasites and
 Ecosystem Engineering: What Roles Could They Play?. Oikos, 84(1), 167-171.
 doi:10.2307/3546879.
- Tigga, M.N., Bauri, R.K., Deb, A.R., Kullu, S.S. (2014). Prevalence of snail's intermediate
 host infected with different trematodes cercariae in and around Ranchi. Veterinary
 World, 7(8), 630-634. doi:10.14202/vetworld.2014.630-634.
- Torchin, M.E., Miura, O., Hechinger, R.F. (2015). Parasite species richness and intensity of
 interspecific interactions increase with latitude in two wide-ranging hosts. Ecology, 96,
 3033-3042.
- UN General Assembly, Convention of the Law of the Sea, 10 December, 1982, available at:
 https://www.refworld.org/docid/3dd8fd1b4.html [accessed 9 April 2019].
- Wegeberg, A.M. and Jensen, K.T. (1999). Reduced survivorship of *Himasthla* (Trematoda,
 Digenea)-infected cockles (*Cerastoderma edule*) exposed to oxygen depletion. Journal
- 893 of Sea Research, 42, 325–331. <u>doi:10.1016/S1385-1101(99)00035-0</u>.
- Whitfield, P.J. (1993). Parasitic helminths. In: F. E. G. Cox (Ed.), Modern parasitology, 2d
 ed. Oxford: Blackwell Scientific Publications, 24-52.
- Willig, M.R., Kaufman, D.M., Stevens, R.D. (2003). Latitudinal gradients of biodiversity:
 Pattern, process, scale, and synthesis. Annual Review of Ecology Evolution and
 Systematics, 34, 273-309. doi:10.1146/annurev.ecolsys.34.012103.144032.

899	Wilson, K., Bjørnstad, O.N., Dobson, A.P., Merler, S., Poglayen, G., Randolph, S.E., Read,
900	A.F., Skorping, A. (2002). Heterogeneities in macroparasite infections: patterns and
901	processes. In the Ecology of Wildlife Diseases (ed. Hudson, P. J., Rizzoli, A., Grenfell,
902	B. T., Heesterbeek, H. and Dobson, A. P.), 6–44. Oxford University Press, Oxford.
903	Wright, D. H., Currie, D. J., Maurer, B. A. (1993). Energy supply and patterns of species
904	richness on local and regional scales. in R. E. Ricklefs and D. Schluter, editors. Species
905	diversity in ecological communities: historical and geographical perspectives, 66-74.
906	University of Chicago Press, Chicago, Illinois, USA.

- 207 Zidane, H., Mannan, M., Mouradi, A., Maanan, M., El Barjy, M., Zourarah, B., Blais J.-F.
- 908 (2017). Environmental Science and Pollution Research, 24, 7970. doi:10.1007/s11356-

JournalPre

909 <u>017-8367-0.</u>

Fig. 1 Study area. Geographical location of the 17 aquatic systems distributed along the Portuguese and Moroccan coastlines. Sampling sites: 1 - Ria de Aveiro; 2 - Óbidos lagoon; 3 - Albufeira lagoon; 4 - Sado estuary; 5 - Mira estuary; 6 - Ria Formosa; 7 - Tahaddart estuary; 8 - Loukkos estuary; 9 - Merja Zerga lagoon; 10 - Sebou estuary; 11 - Oum Er Rbia estuary; 12 - Sidi Moussa lagoon; 13 - Oualidia lagoon; 14 - Souss estuary; 15 - Chbika estuary; 16 - Khnifiss lagoon and 17 - Dakhla bay.

Fig. 2 Latitudinal distribution of the 12 trematode species found in *Cerastoderma edule*. Dashed line: Strait of Gibraltar. Grey bar: Presumable distributional range.

Fig. 3 Trematode mean abundance per aquatic system for the two most representative species, *Parvatrema minutum* (black bar) and *Himasthla quissetensis* (dark grey bar), and for the other cercariae (light grey bar).

Fig. 4 Cluster analysis based on trematode parasites communities in seventeen aquatic systems from Portugal and Morocco (**A**) and Principal coordinates ordination (PCO) showing the variables that better explained samples distribution (**B**). MGS: Sediment median grain-size; M Temp: Maximum annual water temperature; density: cockle density. Sampling sites: 1 - Ria de Aveiro; 2 - Óbidos lagoon; 3 - Albufeira lagoon; 4 - Sado estuary; 5 - Mira estuary; 6 - Ria Formosa; 7 - Tahaddart estuary; 8 - Loukkos estuary; 9 - Merja Zerga lagoon; 10 - Sebou estuary; 11 - Oum Er Rbia estuary; 12 - Sidi Moussa lagoon; 13 - Oualidia lagoon; 14 - Souss estuary; 15 - Chbika estuary; 16 - Khnifiss lagoon and 17 - Dakhla bay.

Fig. 5 Distance-based redundancy analysis plot and the correlated variables that explained aquatic systems distribution based on trematode abundance. Sampling sites: 1 - Ria de Aveiro; 2 - Óbidos lagoon; 3 - Albufeira lagoon; 4 - Sado estuary; 5 - Mira estuary; 6 - Ria Formosa; 7 - Tahaddart estuary; 8 - Loukkos estuary; 9 - Merja Zerga lagoon; 10 -

Sebou estuary; 11 - Oum Er Rbia estuary; 12 - Sidi Moussa lagoon; 13 - Oualidia lagoon; 14

- Souss estuary; 15 - Chbika estuary; 16 - Khnifiss lagoon and 17 - Dakhla bay.

ournal Pre-proof

Table 1. Characterization of each sampled area in terms of type of aquatic system (1 = lagoon; 2 = estuary; 3 = bay), latitude (LAT), longitude
(LON), surface area (km ²), maximum annual water temperature (M Temp, °C), minimum annual water temperature	e (m Temp, °C), maximum
annual water salinity (M Sal), minimum annual water salinity (m Sal) and sediment median grain-size (MGS, mm). ND	– no data

System	Туре	LAT (N)	LON (W)	Surface Area (km ²)	M Temp	m Temp	M Sal	m Sal	MGS	References
Ria Aveiro	1	40°38'	8°44'	83.0	22.0	15.0	28.0	12.0	0.277	Dias et al., 2000 Lillebø et al., 2015
Óbidos	1	39°24'	9°12'	7.0	22.0	10.0	37.0	26.0	0.392	Malhadas et al., 2009 Oliveira et al., 2006
Albufeira	1	38°30'	9°10'	1.3	22.5	11.0	36.0	31.0	0.404	Fortunato et al., 2014
Sado	2	38°28'	8°50'	240.0	21.9	13.8	36.0	10.9	0.304	Bao et al., 1999 Martins et al., 2001
Mira	2	37°43'	8°46'	16.0	22.5	12.0	35.0	27.0	0.353	Silva et al., 2006
Ria Formosa	1	36°58'	7°52'	170.0	26.0	12.0	36.5	13.0	0.326	Gamito and Erzini, 2005
Tahaddart	2	35°46'	5°42'	10.0	26.0	13.0	41.0	21.0	0.194	Achab, 2011
Loukkos	2	35°07'	06°00'	72.0	27.0	15.0	34.0	22.0	0.203	Geawhari et al., 2014
Merja Zerga	1	34°51'	06°16'	27.0	28.0	11.0	35.0	27.0	0.272	Gam et al., 2010
Sebou	2	34°16'	06°39'	17.5	30.0	16.0	35.0	12.0	0.219	Haddout et al., 2015
Oum Er Rbia	2	33°28'	08°34'	1.5	25.0	15.0	35.0	30.0	0.146	Khalki and Moncef, 2007
Sidi Moussa	1	32°54'	08°49'	4.2	27.0	15.0	33.0	22.0	0.273	Maanan et al., 2004
Oualidia	1	32°45'	08°30'	3.0	21.0	16.0	36.0	28.0	0.181	Hilmi et al., 2005
Souss	2	30°21'	09°35'	16.0	25.0	19.0	39.0	35.0	0.202	Anaijar et al., 2008

Chbika	2	28°14'	11°42'	0.2	ND	ND	ND	ND	0.184	
Khnifiss	1	28°03'	12°15'	65.0	22.0	16.0	38.0	34.0	0.150	Semlali et al., 2012
Dakhla	3	23°45'	15°50'	400.0	26.0	14.0	39.5	37.0	0.120	Zidane et al., 2018

Journal Pre-proof

Table 2. Digenean trematode species found in *Cerastoderma edule* from the Atlantic coasts of Portugal and Morocco with indication of the hosts involved in their life cycle. Adapted from de Montaudouin et al. (2009). *: Probable final host of *Curtuteria arguinae*.

Trematode species	1 st intermediate host	2 nd intermediate host	Final host
Bucephalus minimus	Cerastoderma edule	Pomatoschistus spp.	Dicentrarchus labrax
Monorchis parvus	Cerastoderma edule	Cerastoderma edule	Diplodus spp.
Gymnophalus choledochus	Cerastoderma edule	Cerastoderma edule	Water birds
Curtuteria arguinae	Unidentified species	Cerastoderma edule	Water birds (*)
Diphterostomum brusinae	Tritia reticulata	Cerastoderma edule	Blennius, Sargus. Symphodus, Oblata
Himasthla continua	<i>Peringia</i> spp.	Cerastoderma edule	Water birds
Himasthla elongata	Littorina littorea	Cerastoderma edule	Water birds
Himasthla interrupta	Peringia spp.	Cerastoderma edule	Water birds
Himasthla quissetensis	Tritia reticulata	Cerastoderma edule	Water birds
Parvatrema minutum	Scrobicularia plana	Cerastoderma edule	Haemotopus ostralegus
Psilostomum brevicolle	Peringia spp.	Cerastoderma edule	Water birds
Renicola roscovitus	Littorina littorea	Cerastoderma edule	Water birds

1011

Table 3. Results of the regression analyses performed to test the influence of latitude (independent variable) on trematode species richness (TSR) and total trematode prevalence (TTP) in *Cerastoderma edule*. SE: standard error. w/o: without. Sampling sites: Se – Sebou estuary; Ch – Chbika estuary.

			p - value	R ²
Constant	11.265	3.272	0.004	
Latitude	-0.225	0.096	0.034	
Model		1.164	0.034	0.217
		~	3	
Constant	14.501	2.344	<0.001	
Latitude	-0.307	0.068	0.001	
Model		1.164	0.001	0.608
	. ? `	•		
Constant	-4.158	1.454	0.013	
Latitude	0.371	0.091	0.001	
Latitude ²	-0.007	0.001	<0.001	
Model		0.136	<0.001	0.821
-	Latitude Model Constant Latitude Model Constant Latitude Latitude ²	Latitude -0.225 Model Constant 14.501 Latitude -0.307 Model Constant -4.158 Latitude 0.371 Latitude ² -0.007	Latitude -0.225 0.096 Model 1.164 Constant 14.501 2.344 Latitude -0.307 0.068 Model 1.164 Constant -4.158 1.454 Latitude 0.371 0.091 Latitude ² -0.007 0.001	Latitude -0.225 0.096 0.034 Model 1.164 0.034 Constant 14.501 2.344 <0.001

Table 4. Prevalence of trematode parasites infecting cockle populations in the aquatic systems of Portugal and Morocco. Sampling sites: RA - Ria de Aveiro; OB - Óbidos lagoon; AL - Albufeira lagoon; SA - Sado estuary; MI - Mira estuary; RF - Ria Formosa; TA - Tahaddart estuary; LO - Loukkos estuary; MZ - Merja Zerga lagoon; SE - Sebou estuary; OR - Oum Er Rbia estuary; SM - Sidi Moussa lagoon; OA - Oualidia lagoon; SO - Souss estuary; CH - Chbika estuary; KH - Khnifiss lagoon and DA - Dakhla bay. N: number of dissected cockles.

		B. minimus	M. parvus	G. choledochus	D. brusinae	H. continua	H. elongata	H. interrupta	H. quissetensis	C. arguinae	P. brevicolle	P. minutum	R. roscovitus	Total parasite Prevalence
	RA						11					26		35
	(n=65)					\sum		_			_			
	OB	3				7							3	6
	(n=35)	Ū	_	_			_	_	_	_	_	_	Ū	Ū
	AL		40		20		10							55
gal	(n=20)	_	40		20	_	10	_	_	_	_	_	-	55
Portugal	SA		5				48	5						52
а.	(n=21)	_		_	-	_	40	5	_	_	_	_	-	52
	MI			50							5			55
	(n=20)	-	_	50	-	_	-	_	-	_	0	_	-	55
	RF				4				36	20	4	8		48
	(n=25)	-	_	_	7	_	-	_	50	20	4	0	-	40
	TA	8			15	54						100		100
	(n=13)	U	_	_	10	04	_	_	_	_	_	100	_	100
000	LO	10			10							90		90
Morocco	(n=20)	10	-	_	10	-	_	_	_	_	_	30	_	30
2	MZ				30				70	100		100		100
	(n=20)	-	_	-	50	-	_	_	10	100	_	100	-	100

					ournal	Pre-	proof						
 SE (n=17)	_	_	_	_	_	_	_	_	_	_	100	_	100
OR (n=20)	_	_	_	_	80	_	65	_	_	45	95	_	100
SM (n=15)	7	_	-	47	_	_	100	100	100	13	67	_	100
OA (n=20)	5	_	_	30	_	_	75	100	100	20	_	_	100
SO (n=20)	_	_	5	_	10	_	60	_	-	£	95	_	100
CH (n=11)	_	_	_	_	_	_	_	-	0	0	100	_	100
KH (n=20)	10	_	5	85	100	_	60	R	_	45	100	_	100
DA (n=36)	_	_	_	92	47	2	39	_	75	25	11	_	100

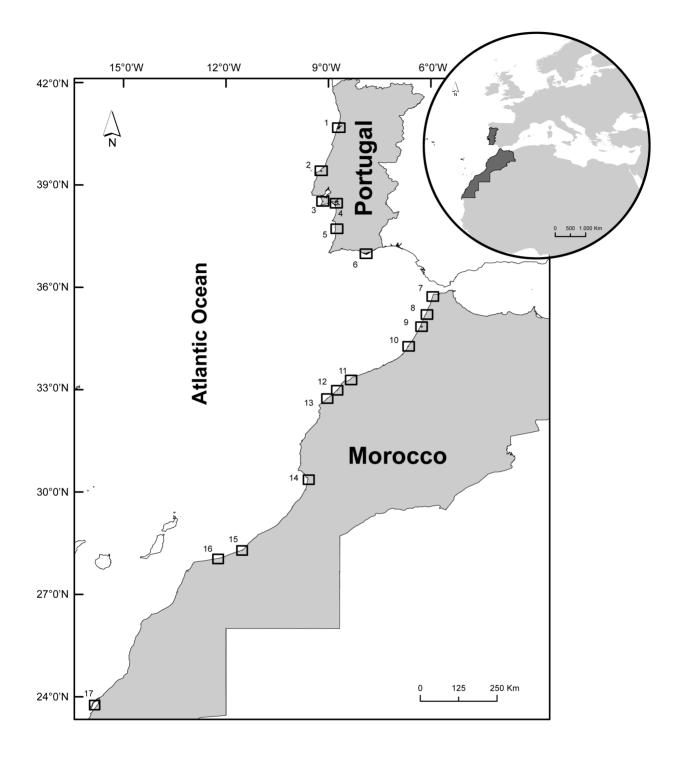
Jonula

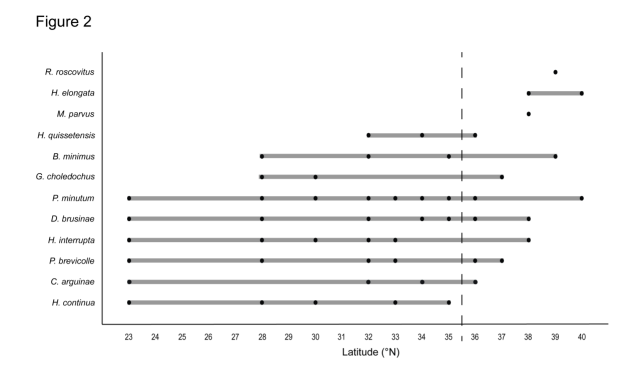
Table 5. Characterization of each affinity group in terms of maximum annual water temperature (M Temp, °C), minimum annual water temperature (m Temp, °C), maximum annual water salinity (M Sal), minimum annual water salinity (m Sal), sediment median grain size (MGS, mm), cockles density (ind.m⁻²) and Rarefaction index for a standardized number of 10 individuals (ES(10)). Significant differences among affinity groups are represented with different letters (p < 0.05). Sampling sites: RA - Ria de Aveiro; OB - Óbidos lagoon; AL - Albufeira lagoon; SA - Sado estuary; MI - Mira estuary; RF - Ria Formosa; TA - Tahaddart estuary; LO - Loukkos estuary; MZ - Merja Zerga lagoon; SE - Sebou estuary; OR - Oum Er Rbia estuary; SM - Sidi Moussa lagoon; OA - Oualidia lagoon; SO - Souss estuary; CH - Chbika estuary; KH - Khnifiss lagoon and DA - Dakhla bay.

Affinity groups	Sampling areas	M Temp	m Temp	M Sal	m Sal	MGS	Cockle density	ES (10)
A1	TA, LO, SE, SO	27.0 ± 2.2	15.8 ± 2.5	37.3 ± 3.3	22.5 ± 9.5	0.2 ± 0.0^{a}	404 ± 457^{a}	1.0
A2	MZ, OR, KH	25.0 ± 3.0	14.0 ± 2.6	36.0 ± 1.7	30.3 ± 3.5	$0.2 \pm 0.1^{a,b}$	1643 ± 2267 ^{a,b}	1.3
B1	SM, OA	24.0 ± 4.2	15.5 ± 0.7	34.5 ± 2.1	25.0 ± 4.2	$0.2 \pm 0.1^{a,b}$	6092 ± 1114 ^b	2.4
B2.1	DA	26.0	14.0	39.5	37.0	0.1 ^b	15 [°]	2.4
B2.2	RA, OB, AL, SA, MI, RF	22.8 ± 1.6	12.3 ± 1.8	34.8 ± 3.4	20.0 ± 9.0	$0.3 \pm 0.0^{\circ}$	135 ± 304 ^a	1.0

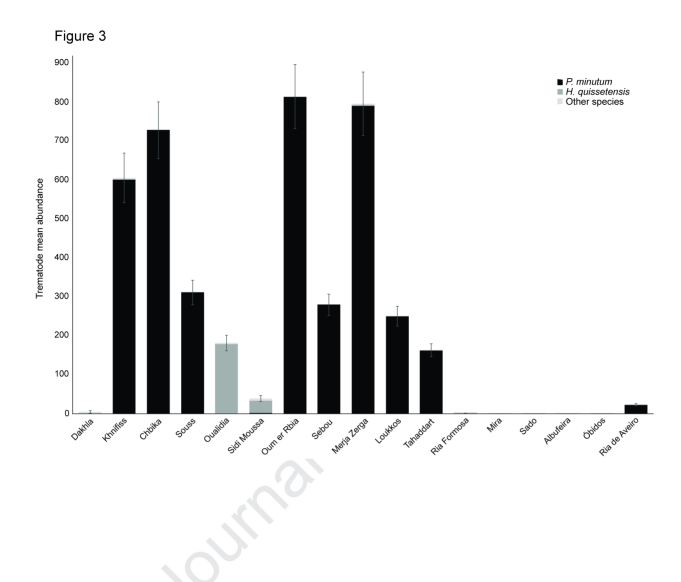
Table 6. Results of the Similarity Percentages (SIMPER) used to characterize the type of aquatic systems (coastal lagoons vs. estuaries) and the geographical position (northern systems vs. southern systems) by the trematode species (of *Cerastoderma edule*) that most contributed to dissimilarity between groups.

	Species	Average abundance group 1	Average abundance group 2	Dissimilarity Contribution (%)
	Parvatrema minutum	5.42	12.22	69.28
0	Himasthla quissetensis	1.70	0.00	8.76
Coastal lagoons (1) vs	Himasthla interrupta	0.66	0.29	4.64
Estuaries (2)	Curtuteria arguinae	0.91	0.00	3.83
	Himathla elongata	0.05	0.13	3.74
	Species	Average abundance group 1	Average abundance group 2	Dissimilarity Contribution (%)
	Parvatrema minutum	0.84	13.15	58.62
	Himasthla quissetensis	0.06	1.81	10.26
Northern systems (1) vs	Dipheterostomum brusinae	0.04	1.04	9.12
Southern systems (2)	Curtuteria arguinae	0.00	1.30	8.48
	Himasthla interrupta	0.01	0.99	6.02



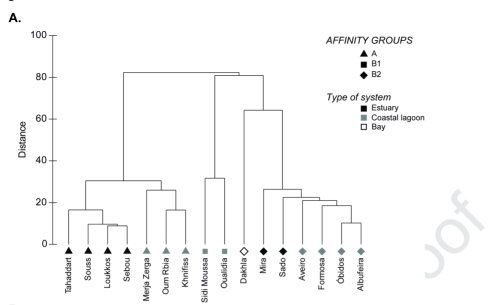


Jonuly

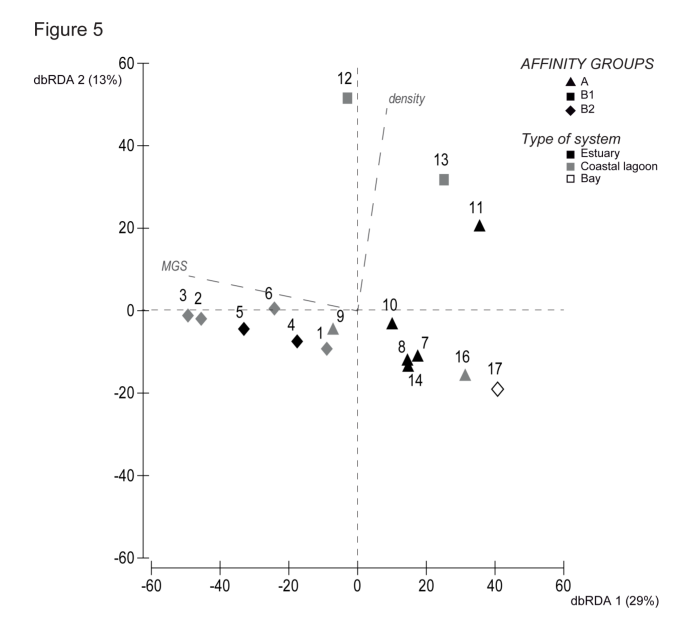




В.



60 T PCO2 (27%) 40-MGS 8 7 10 ▲7 ▲ 14▲11 ▲16 20 2 5 0 9 ▲ MTemp 17 \$ -20density -40-13¹² -60--40 -20 20 40 -60 ò



1 Highlights

- 2 Trematode diversity in the southern distribution area of *Cerastoderma edule*.
- 3 Trematode species occurrence driven by the presence of the first intermediate host.
- 4 Trematode prevalence and abundance was positively correlated to temperature.
- 5 At large scale, cockle density positively influences trematode abundance.
- 6 Systems under buffered environmental variations presented higher trematode
- 7 abundance.
- 8 Cockle density influences positively trematode abundance within certain values

Declaration of interests

 \boxtimes 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.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Journal Prerk