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Baseline pathological data of the wedge clam *Donax trunculus* from the Tyrrhenian Sea (Mediterranean Basin)

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8 ABSTRACT: In recent years, a collapse in *Donax trunculus* fishing yields has occurred in the 9 Tyrrhenian Sea (Mediterranean Basin). There is little information available on the impact disease 10 may have had on D. trunculus populations. For the first time, a pathological survey was performed on the natural beds of the bivalve on the Campania and Lazio coasts, western Italy. Detected 11 12 pathogens and related diseases were analysed, and their prevalence and mean intensity values were calculated. Viral particles, Chlamydia-like organisms, ciliates, coccidians, microcells and 13 14 trematodes were observed. An unknown ciliate was linked to severe inflammatory and necrotic 15 lesions in the digestive gland. Metacercariae of the trematode Postmonorchis sp. were also 16 strongly represented in almost all samples, reaching high levels of infection; however, none of the pathogens described required the World Organisation for Animal Health to be notified. I nitial 17 results indicated that further surveys related to environmental data are necessary in order to assess 18 the relevance of these early observations in managing the declining D. trunculus population in 19 20 the Tyrrhenian Sea.

- 21 KEY WORDS: Wedge clam Protozoa Bivalve disease Trematodes
- 22

23 1. INTRODUCTION

24 Donax trunculus (Bivalvia: Donacidae) is an Atlantic-Mediterranean warm-temperate bivalve 25 species found in the Mediterranean Sea, the Black Sea and in the Atlantic Ocean in coastal regions 26 from Senegal to France. In these environments, D. trunculus populations are distributed in sandy beds between depths of 0-2 m in the Mediterranean, and from 0-6 m on the Atlantic coasts, with 27 the highest densities at 0-3 m (Deval 2009). This bivalve represents an important fishery resource 28 29 due to its high economic value. Landings in Europe over the last 12 yr (since 2006) were 11202 30 t, with a maximum yield of 1355 t in 2005, followed by a steady decline, with only 516 t in 2011 31 (FAO 2013). A recent collapse in yields has occurred in the Tyrrhenian Sea of the Mediterranean 32 Basin (de la Huz et al. 2002, Rambaldi et al. 2010, Marie et al. 2016). The causes of this collapse 33 are unclear. Abiotic factors resulting from anthropogenic activity—such as beach nourishment and chemical pollutants—along with extreme meteorological events related to global warming, 34 35 have been advocated as causing significant persistent changes to the environment of the sandy 36 beds of the Tyrrhenian coast, with serious consequences for the bivalve resources (Rambaldi et 37 al. 2010). However, other environmental stressors associated with climate change may also have 38 facilitated the spread of diseases into previously unaffected shellfish populations, exacerbated 39 disease where it already exists or facilitated the emergence of novel pathogens, which also may 40 have caused severe losses. There is very little information available on the prevalence and 41 distribution of disease in D. trunculus populations in the Mediterranean Sea. Although pathogens 42 such as Rickettsia/ Chlamydia-like organisms (RLOs/CLOs), nematodes and trematodes have 43 been described as parasites of *Donax* spp. (Sindermann & Rosenfield 1967, Comps & Raimbault 1978, Ansell 1983, de Montaudouin et al. 2014), few studies have reported specifically on D. 44 45 trunculus, and little is known of the pathogens that may adversely affect the health of these 46 populations. In general, trematodes are the most often described parasites of the family 47 Donacidae, including Fellosto- midae and Monorchidae (Palombi 1933a,b, Carella et al. 2013, de 48 Montaudouin et al. 2014).

In the present study, we sampled *D. trunculus* from the Tyrrhenian Sea over several years and describe the pathogens and diseases found using histopathol- ogy and molecular techniques to provide baseline data on the health status of this species.

52 2. MATERIALS AND METHODS

53 2.1 Sampling

From 2008 to 2015, 330 wedge clams D. trunculus were collected during the summer or autumn 54 from different geographical areas. The sampling sites in the Campania region were the Volturno 55 56 River mouth (41° 0' 57" N, 13° 56' 50" E) from July 2008 to 2010 (90 clams) and Litorale Domitio 57 (41°4'8"N, 13° 51'4"E) from June and July 2012 (60 clams) and June 2014 and 2015 (60 clams). In September 2012, samples were also collected at Formia (41° 14'21"N, 13° 36' 43" E) (30 clams) 58 in the Lazio region. In October 2013, clams were collected in 3 areas of Salerno Province 59 including Foce Sele (30 clams) (40° 29' 02" N, 14° 56' 29" E), Foce Tusciano (30 clams) (40° 34' 60 61 50" N, 14° 52' 58" E) and Solofrone (30 clams) (40° 22' 10" N, 15°00'0"E) (Fig. 1). The clams were sampled from the shore using a hand-held semi-circular dredge as used by the local 62 63 fishermen.

64 2.2 Animal tissue processing

The animals were opened in the laboratory and examined for external signs of pathology under a 65 dissecting microscope (Nikon SMZ-10). The bivalves were then fixed in Davidson's fixative for 66 24 to 48 h and processed for routine histopathology. Additionally, pieces of digestive gland, gills 67 68 and gonads were also preserved from each animal for further molecular and transmission electron 69 microscopy (TEM) analysis. For histopathology, 2 transverse sections, approx. 5 mm thick, 70 including the mantle, gonad, digestive gland, gills, kidney and foot, were excised from each clam and placed into histological cassettes. The tissue samples were embedded in paraffin wax, sliced 71 in 5 pm sections and subsequently stained with haematoxylin and eosin (H&E). For better 72 73 characterization of the tissue, additional stains, such as PAS-BA and Masson's Trichrome (Mazzi 74 1977), were also used on selected slides from every sampled area. Furthermore, to improve the 75 characterization of the prokaryote-like inclusions, additional staining techniques were used: 76 Gram's method for Gram-positive and Gram-negative bacteria, Macchiavello stain (Mazzi 1977) 77 and Giemsa (Howard & Smith 1983). All micrographs were captured using a Nikon DS-Fi1 video 78 camera mounted on a Nikon 50i microscope connected to a computer. Histological sections were 79 examined for the presence of parasites and pathological alterations. Pathogen prevalence (P) and 80 mean intensity (I) of the different parasites were calculated. For evaluation of the infection 81 intensity of trematodes such as Postmonorchis sp. present in gill tissue, data were obtained by 82 histological examination (histological infection density, HID) as described by Carella et al. 83 (2013).

The percentage of clams affected by each parasite and their pathological condition were
determined for each sampled bed included in the survey. The overall prevalence (and 95% CI) of
each parasite and pathological condition were calculated for the whole study.

87 2.3 TEM

After histological examination, small pieces of digestive gland parasitized by protozoa and
individuals with digestive glands that had a foamy aspect at the level of A cells under light
microscopy were selected from the stored tissues for ultrastructural analysis by TEM.

91 Tissues were fixed in 2% glutaraldehyde in filtered seawater for 24 h at 4°C, post-fixed in 2%

- tetroxide osmium (2 h at 4°C) and embedded in EPON. Ultrathin sections (70–90 nm) were cut,
- the tissues were stained with uranyl acetate and lead citrate and were observed using a JEOL 100
- 94 CX2 transmission electron microscope at the Centro Interdipartimentale di Servizio per la
- 95 Microscopia Elettronica (CISME), University of Naples.
- 96

97 2.4 Molecular identification of RLOs/CLOs

For RLO/CLO identification, DNA isolation was performed from pieces of digestive gland
previously preserved in 96% EtOH. DNA extractions were performed employing the DNeasy
Blood and Tissue Kit (Qiagen) following the manufacturer's instructions. The DNA concentration
was measured using a Nanodrop 2000c spectrophotometer (ThermoScientific), and DNA quality
was verified by electrophoresis on a 1 % agarose gel stained with ethidium bromide.

103 The 16S primers RCF/RCR described by Costa et al. (2012) for Rickettsia/Chlamydia-like bacteria were used. The PCR mixture consisted of 0.4 pM of each primer RCF/RCR (Sigma), 104 105 PCR reactions were carried out using PCR buffer at 1x concentration, 1.5 mM MgCl₂, 200 pM of each deoxynucleotide (dNTP mixture; Takara Bio) and 1 unit of Taq DNA Polymerase 106 (Invitrogen), 2 pl of template DNA was added to the mixture to make a final volume of 25 pl. 107 After a denaturation step at 94°C for 5 min, 30 cycles were run with an Eppendorf thermal cycle 108 109 as follows: denaturation at 94°C for 1 min, annealing at 51°C for 1 min and elongation at 72°C 110 for 1 min. A final elongation step at 72°C for 10 min was performed.

The PCR products were run on 1.2% agarose gel. The amplified fragments were gel-eluted and
directly sequenced. The sequencing reactions were run on a 310 Automated Sequencer (Applied
Biosystems). The best BLAST hits for the obtained nucleotide sequences were downloaded and

aligned with their corresponding sequences using ClustalW. The alignment files were employed to construct neighbour icining trees using the software MECA y = 7 (Kumer et al. 2016)

to construct neighbour-joining trees using the software MEGA v.7 (Kumar et al. 2016).

116 3. RESULTS

117 A total of 330 wedge clams from the Lazio and Campania regions were examined. Histopathologi-

cal analysis revealed different symbionts and pathological conditions affecting *Donax trunculus*.

119 Observed pathogens and their tropism are shown in Fig. 2; see also Table 1 & Figs. 3-9.

3.1 Viral infection

121 Under light microscopy, in specimens from Litorale Domitio from 2014 and 2015, some A cells 122 of the digestive epithelium were vacuolated and foamy. TEM examination of the digestive tubules 123 from 3 cases from 2014 and 2 from 2015 showed the presence of viral particles 60 nm in 124 dimension in A cells. The virions exhibited an icosahedral symmetry (Fig. 3). Moreover, 125 cytoplasmic viral factories (where viral replication and assembly likely take place) constituted by 126 both electron-dense cytoplasmic inclusions and membrane vesicles were visible (Fig. 3 insert).

127 3.2 Prokaryote infection by CLOs

128 During histology, light microscopy observations of H&E-stained tissues revealed rounded 129 intracellular inclusions of prokaryote-like colonies. The inclusions were (mean \pm SD) 7.65 \pm 1.79 pm in length and were found in the epithelium of the digestive gland in clams from all localities. 130 These colonies occupied the cytoplasm of the A cells of the digestive tubules, in some cases 131 causing hypertrophy or destruction (Fig. 4A,B). In some cases, an infiltrative mild/strong 132 133 haemocytic reaction was detected. The colonies were positive for Rickettsia/Chlamydia by Macchiavello and Giemsa staining (Fig. 4C,D). Most infected clams had 8 to 20 inclusions 134 135 section⁻¹, but a few clams had 20 to 30 inclusions section⁻¹. The maximum P and mean I were recorded in Litorale Domitio in 2015 (83%), with 30 colonies per histological section. No 136 prokaryote-like colonies were observed at Foce Sele and Solofrone in October 2013 (Table 1). 137

PCR amplification was run on 3 samples from Volturno (2008 and 2009), 3 samples from Litorale
Domitio (2012) and 2 samples from Formia (2012) on a 350 bp region produced by the partial
16SrRNA. BLAST analysis revealed 100% identity among all the sequences, and the best
nucleotide identity score (92%) was obtained for various strains of *Clamydia*- like bacteria, such
as *Chlamdya pecorum, C. ibidis* and *Estrella lausannensis* (Fig. 5). The sequences were deposited

in Genebank (accession no. MH492627).

3.3 Protists

145 Two different types of ciliates were observed: one located in the gills and another in the digestive gland. Moreover, oocytes of the gregarine Nematop- sis sp. were also detected in the connective 146 147 tissue of the digestive gland and gills (Fig. 6). These oocysts were oval in shape, had a maximum dimension of (mean \pm D) 9.49 \pm 1.24 pm (n = 20) and usually contained 1 sporozoite oocyst⁻¹. 148 149 Ciliated protozoa wereobserved adjacent to the gills (Fig. 6A). They were oval-shaped, had dense ciliature and measured 10 to 12 pm in length. They may have been Hypocomella sp., which have 150 151 been described in other bivalve species, but no reports have occurred in the wedge clam. The ciliates had 2 nuclei (macro- and micronucleus). They did not appear to cause any specific host 152 153 response. The maximum P were 36% in Foce Sele and 40% in Tusciano in 2013.

In 2 samples from 2014 and 1 sample from 2015, the presence of small protozoans similar to
haplosporidian microcell parasites were detected in different tissues of *D. trunculus* from Litorale
Domitio (June 2015), particularly in muscular and connective tissues and the intracellular space
within haemocytes (Fig. 6C,D). The infection was focal and low for any given organ. They mostly
resembled the *Mikrocytos*-like parasite previously described by Garcia et al. (2018), while for the
immune cell tropism they were similar to other microcells belonging to *Bonamia* species.

160

161 An unidentified ciliate endoparasite was detected in the interstitial spaces of digestive tubules in the digestive gland (Fig. 7). The parasite was present in every sampled area, showing a high 162 prevalence and infection intensity, with an overall P of 219 (Fig. 7A). The parasite was always 163 164 accompanied by an intense inflammatory reaction, and in some cases it was asso ciated with focal 165 necrosis of the digestive tubules and intestinal epithelia (Fig. 7B). Generally, the parasites were grouped with others and were often present at the haemo-lymphatic sinus level, eating host 166 haemo- cytes (Fig. 7C,D). Morphologically, the ciliates were ovoid with a tapered anterior end 167 168 and round in crosssection. They had an estimated maximum body width

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170 3.4 Metazoa

171 Metacercarie of the trematode *Postmonorchis* sp., as previously described by Carella et al. (2013), 172 were also detected (Fig. 8A-C). They were located at the gill level and were distributed across the 173 upper, middle and lower part of filaments, in many cases reaching high infection intensity and 174 macroscopic visibility (Fig. 8A). The mean diameter of the cysts was (mean \pm SD) 180-220 \pm 175 23.82 pm (n = 50). In a few cases, the parasite was detected in the foot, the stomach and the 176 kidney-pericardium system. The inflammatory response included encapsulation and ranged from 177 mild to intense in gill tissues, eliciting hypertrophy and hyperplasia of the gill epithelia.

Different phases of development of the metacer- caria were observed. In fact, some of them had
a smaller dimension (~160 pm) and a thinner cyst wall compared to other metacercariae (Fig. 8B).
In these cases, the pathogen appeared to be more susceptible to the host inflammatory response.
Other histopatho- logical findings included mild hypertrophy or hyperplasia of mucus cells and
the connective tissue of the gills, as demonstrated by PAS-BA and Trichrome staining. A higher
prevalence was detected in the northern part of the Campania region, at the Volturno River mouth
in 2008 and 2009.

185 Sporocysts of the Fellodistomidae trematode *Bac- ciger bacciger* (Rudolphi 1819) were observed 186 in D. trunculus in all localities except for the areas of Salerno Province. Sporocysts containing 187 developing cercariae were found mostly in the gonad, but also in the foot, mantle and/or the whole visceral mass (Fig. 8D). The fully developed cercaria showed a seti- ferous tail, eosinophilic 188 189 staining and an ovoid body of 310 ± 52 pm (n = 10) (Fig. 8E). When present, the parasite generally showed heavy infection intensity. The tissues of the gonad and digestive gland of these clams 190 191 were replaced by a sporocysts and cercariae that caused intense damage and host castration. 192 Animals did not show a haemocyte reaction to the pathogen. Within the sporocyst, the cercariae 193 were typically in the same range of deveolopmental stages.

194 195

4. DISCUSSION AND CONCLUSIONS 196

197 This study revealed, for the first time, the occurrence of different pathogens and related 198 pathological conditions affecting the wedge clam *Donax trunculus* along the Campania and Lazio 199 coasts.

200 The presence of viral particles in A cells of the digestive gland were observed in our study. 201 Previous reports described 2 types of birnavirus in the bivalve Tellina tenuis, tellina virusl (TV-1) and tellina virus2 (TV-2) (Chung & Paetzel 2013, Nobiron et al. 2008), in digestive tubule 202 203 cells. TV-1 was described in connection with a population of diseased *T. tenuis* with watery flesh. 204 Infected animals had atrophic digestive tissues and were markedly thinner than healthier 205 specimens (Renault & Novoa 2004, Nobiron et al. 2008). The virus showed morphological and physio- chemical characteristics similar to infectious pancre atic necrosis virus (IPNV), an 206 aquabirnavirus that infects salmonid fishes (Hill 1976, Underwood et al. 1977). On the basis of 207 208 their location and dimension, the virions observed in our case were in concordance with the 209 above-mentioned reports; however, further studies are needed to definitely assign them to this 210 group.

211 RLOs and CLOs are obligate, intracellular bacterial parasites associated with a variety of 212 vertebrate and invertebrate hosts (Gollas-Galvan et al. 2014). Several authors have documented the presence of RLOs/CLOs in many aquatic animals, including fish, molluscs and crustaceans 213 (Gulka & Chang 1984, Wang & Gu 2002, Sun & Wu 2004), with most of the literature focused 214 215 on teleost fishes and bivalve molluscs. In some cases, these pathogens are causative agents of severe mortality outbreaks in farmed aquatic species such as the Rickettsia Candidatus 216 Xenohaliotis californiensis that causes mass mortalities of Haliotis spp. in many countries 217 218 (Crosson et al. 2014). However, little is known about the pathogen's life cycle or host range 219 (Ferrantini et al. 2009). In this study, the observed intracellular bacteria were recorded within the 220 cytoplasmic vacuoles of the digestive gland epithelium of D. trunculus. In many cases, the 221 immune response of the host was mild. Only some specimens had digestive epithelia destruction 222 as reported by Comps & Raimbault (1978). Our molecular study revealed that the prokaryotic 223 inclusion in the digestive gland belonged to *Chlamydi- ales*, as also reported by Costa et al. (2012) 224 in *Rudi- tapes decussatus*. Interestingly, in contrast to RLOs, which have been described in 225 different tissues in bivalves, CLOs have been reported only in digestive tubules. In particular, a 226 study by Renault & Cochen- nec (1995) reported the presence of *Chlamydia* sp. in the Japanese 227 oyster Crassostrea gigas, where indirect fluorescent and peroxidase conjugated antibody tests suggested that this agent might share common antigens with the prokaryotic agent Chlamydia 228 229 psittaci, strain ovis. Infections by members of the genera Chlamydia have been described in a variety of adult bivalve molluscs, with reports in bivalves in Chesapeake Bay by Harshbarger et 230 al. (1977), in Mercenaria mercenaria from the Great South Bay (Meyers 1979) in Argopecten 231 232 irradians in Canada (Morrison & Shum 1982) and in mussels from Basque Country (Cajaraville 233 & Angulo 1991).

234 Regarding protists, the gregarine *Nematopsis* sp. was not highly represented, with no possible species identification. This parasite has not been previously reported in D. trunculus, but 235 236 Nematopsis sp. has been reported in different bivalve species such as Mytilus galloprovincialis (Carella 2010), R. decussa- tus and Cerastoderma glaucum (Longshaw & Mal- ham 2013). 237

238 Ciliate parasites are very common organisms in bivalves (Bower et al. 1994). In this study, 2 types 239 of ciliates were observed. One was present adjacent to the gills and caused evident damage. An 240 unidentified ciliate was observed in the digestive tissues of specimens in all the sampled areas and had high prevalence and infection intensity. The protozoa were characterized by the presence 241 242 of cilia, a rounded nucleus and an elongated, tube-shaped body structure. The cytoplasm of the ciliates was typically full of food vacuoles, which appeared to contain debris. Our study showed 243 that ciliate infections can cause significant morbidity related to intense tissue damage. Other 244 245 studies reported the detrimental effects of ciliated protozoa on the digestive tissue of bivalves. 246 Elston et al. (1999) described mortality episodes linked to an invasive opportunistic ciliate 247 belonging to Scuticociliatida. The same group of protozoa has been described in other bivalve 248 species and in crustaceans where infection is fatal (Morado & Small 1994, 1995, Cawthorn et al. 249 1996). We still do not have a specific taxonomic affiliation for the observed protozoa in D. 250 trunculus from the Tyrrhenian area. Further molecular analyses are needed to identify the 251 observed pathogen.

252 Microcell parasites are small intracellular protozoans mostly detected in molluscs, and they can

253 be associated with high mortality rates. They are essentially mollusc parasites represented by 2

254 genera: Bonamia and Mikrocytos (Carnegie & Cochennec- Laureau 2004). Microcell-like

255 parasites were detected at low prevalence in our samples in the muscles and connective tissues, 256 and the infection was focal and low for any given organ. This was the first time that a microcell

parasite has been detected in Italy. Further molecular analyses are required in order to classify 257 258 the organism.

259 Metazoan parasites detected in this survey included different phases of development of digenean

260 trematodes. B. baccigerwas reported by Ramon et al. (1999), with adult phases originally discovered by Rudolphi (1819) in marine fishes. In bivalves, the literature reports that heavy 261

262 sporocyst infiltration of *B. bacciger* can cause complete castration and depletion of body

energy, with host soft tissues becoming empty and watery (Lauckner 1983). On the French 263

- 264 Atlantic and Italian Mediterranean coasts, the parasite caused large-scale fluctuations in the
- 265 abundance of *Ruditapes* spp. and *D. vittatus* populations. It was also previously reported in the 266 Campania region by Palombi (1933a,b) at Fusaro Lagoon, with higher prevalence during the spring from March to May. In our study, the samples were all collected during the summer 267 season, with a higher infection prevalence in Litorale Domitio in 2014 and 2015. 268

269 Metacercariae of *Postmonorchis* sp. have already been described by Carella et al. (2013). The

- 270 prevalence of the pathogen ranged from 76 to 100%, while the infection intensity fluctuated in
- 271 the study areas. An inflammatory response was always recorded in parasitized animals and was
- 272 characterized by inflammatory capsules that adhered to and enclosed the foreign body; such

273 responses were, in some cases, ineffective against completely formed metacercariae with a 274 thicker covering, as already reported (Carella et al. 2013).

275 Young (1953) described Postmonorchis donacis from the Pacific coast of the United States. The 276 adult phases were found in teleosts belonging to the families Sciaenidae and Embiotocidae, and

277 cercariae and metacercariae were detected in the bean clam D. gouldii. The reported

- 278 metacercariae of Postmonorchis sp. in D. trunculus showed a combination of features of the 3
- 279 described species of *Postmonorchis* present in the literature: *P. variabilis*, *P. orthopristis* and *P.* 280 donacis.

281 In conclusion, the present study was the first attempt to identify pathogens of *D. trunculus* col-

282 lected from a natural shellfish bed on the Tyrrhenian coast, combining histological and

283 molecular analyses that should be useful as a reference in future health surveys. Gill ciliates,

284 Nematopsis sp., and Chlamydialike colonies seemed to have limited pathogenicity and light infection intensity. On the other hand, based on their pathogenic potential, the observed viral 285

286 particles, ciliated protozoa, bucephalid sporo- cysts and monorchidae metacercariae caused 287 heavy infections.

288 Among the protozoa, the unidentified ciliate protozoan in the digestive gland was significant for

- 289 its high prevalence and was possibly detrimental to the wedge clam populations. None of the 290 observed parasites are on the list of obligatory notification of the World Organization for
- 291 Animal Health (OIE).

292 In this study, infections with microcell parasites were observed in a few specimens of D.

293 trunculus. In France, these parasites caused animal mortality (Garcia et al. 2018). Based on

294 morphological characteristics and its detection in the wedge clam, we consider that the pathogen 295 probably belongs to the novel Mikrocytos species detected in France. This underlines the

296 possible importance of including microcells on the list of potential disease agents.

- 297 Our data is limited to one season with corresponding environmental parameters. However, these
- 298 initial results show that further surveys related to environmental data and seasons are necessary

299 in order to assess the relevance of the pathogens and diseases observed in this declining D.

300 trunculus population in the Tyrrhenian Sea.

301

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