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Parasitological and pathological findings in Fin Whales (*Balaenoptera physalus*) stranded along Italian coastlines

Parasitological findings in Mediterranean Fin Whales

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Abstract

Mediterranean fin whale population faces many threats to its conservation, including both anthropic and natural issues. Few records on the parasitofauna of this species are present for this geographical area. The aim of this survey was to investigate presence and impact of parasitic diseases in Mediterranean fin whales (*Balaenoptera physalus*). Seven animals, stranded along Italian coastlines in the period 2006-2015, were submitted to necropsy and parasitological examination. One protozoan parasite, *Toxoplasma gondii* was detected in one fin whale and, for the first time in mysticetes, was successfully genotyped as a type II strain with 15 microsatellite markers.

One crustacean (*Pennella* spp.) and four helminth taxa (*Crassicauda boopis*, *Ogmogaster antarcticus*, *Tetrabothrium ruudi* and *Bolbosoma* sp.) were overall detected and morphologically identified. Cases of infestation by adult *P. balaenopterae* included variable parasitic burdens; impairment of immune system was suspected to be present in most severe cases, as already described for other cetacean species. Immature stages of *Pennella* sp. were also detected in two animals and are here described for the first time in cetaceans. Infection by *C. boopis* was observed or suspected in five cases. Parasitic thrombi, involving renal vessels and vena cava, fibrosis of renal parenchyma and renal impairment were observed in association to this parasite. Larval nematodes associated to arteriosclerosis of the mesenteric arteries were detected. To our knowledge, this is the first report of *C. boopis* inside the Mediterranean Sea and these findings should prompt further investigation to evaluate the prevalence of this severe infection in Mediterranean fin whales.

Keywords: *Balaenoptera physalus*, helminth parasites, crustacean parasites, *Toxoplasma gondii*, Genotyping, Mediterranean Sea

1. Introduction

Fin whales *Balaenoptera physalus* (Linnaeus, 1758) are the most abundant mysticetes in the Mediterranean Sea. Genetic studies confirmed the existence of a resident Mediterranean population (Bérubé et al. 1998) listed as vulnerable by the International Union for Conservation of Nature (IUCN) Red List, since it is progressively decreasing.

Major concerns for their conservation are attributable to anthropic activities, with ship strikes considered as the most relevant threats (Panigada & Notarbartolo Di Sciara 2012). In addition, natural diseases are considered a problem for their conservation: two outbreaks of Dolphin Morbillivirus have been described in the Pelagos Sanctuary involving fin whale specimens during the period 2011-2013 (Mazzariol et al. 2016). Epidemics of this viral disease should be also regarded as one of the major threats for this species. In some cases, in cetaceans, this viral infection is associated to opportunistic infection, such as those related to *Toxoplasma gondii*. Despite the fact that most of the toxoplasmosis infections are referred to toothed whales, with the protozoan being recognized as responsible for abortion, encephalitis and systemic lethal disease (Migaki et al. 1990, Di Guardo et al. 2010), 2 reports of *T. gondii* exist in baleen whales, in particular a young fin whale stranded in Italy (Mazzariol et al. 2012), and a seropositive humpback whale (*Megaptera novaeangliae*) stranded in England (Forman et al. 2009).

As for *T. gondii*, only few data on other parasites in Mediterranean fin whales are reported (Tamino 1953, Malatesta et al. 1998, Cicek et al. 2007, Giorda et al. 2017). The large copepod *Pennella balaenopterae* Koren and Danielssen, 1877 (Copepoda, Pennelliidae) has been described here since long time and its presence on whales' skin in the Pélagos Sanctuary is considered almost regular (Tamino 1953, Notarbartolo Di Sciara et al. 2003). Its presence is easily detectable even in live animals, since the posterior part of their body emerges from the host's skin and trails free in the water, while the head penetrates the skin and blubber until reaching the muscular fascia. In small odontocetes, the intensity of infection has been positively correlated to deficiency of immune system

26 in striped dolphins (Vecchione & Aznar 2014). Among endoparasites, the trematode *Ogmogaster*
27 *antarcticus* Johnston, 1931 (Digenea, Notocotylidae) was described in a fin whale stranded on the
28 north-eastern coast of Tyrrhenian Sea. As for nematodes, infection by *Crassicauda* sp. was recently
29 reported in a specimen of *B. physalus*, associated with a severe nephropathy and severe mesenteric
30 lesions (Giorda et al. 2017). Infection by the species *Crassicauda boopis* Baylis 1920 is known to
31 cause serious illness in baleen whales, since the localization of adult worms in the vascular and renal
32 district can lead to lethal outcomes. This species has been described in fin whales from the Atlantic
33 and Pacific Ocean, but no reports in the Mediterranean are present in literature (Lambertsen 1986,
34 1992, Lempereur et al. 2017).

35 In order to give a throughout picture on the effect of the parasites in the conservation of this species
36 in the Mediterranean Sea, the parasitic findings and associated pathological changes in fin whales
37 stranded along the Italian coastline in a ten years' period (2006-2015) have been described in this
38 paper.

39

40 **2. Material and Methods**

41 Between October 2006 and January 2015 seven carcasses of fin whales, stranded along the Italian
42 coastlines (Tyrrhenian and Ligurian Sea), were necropsied on the stranding sites following standard
43 protocols (McLellan et al. 2004) by the Cetacean stranding Emergency Response Team (C.E.R.T.),
44 established by the Ministry for Environment, Territory and Sea at Padova University, in collaboration
45 with the local Units of Health Institutions.

46 Biometric data, carcass condition code, body condition score, age estimation and sex of the stranded
47 animals are reported in Table 1.

48

49 *Parasitological and pathological analyses for metazoan parasites*

50 Following post-mortem examination, whenever it was possible, parasitological analyses were carried
51 out on skin, blubber and respiratory, gastrointestinal, cardiovascular, and urogenital systems. In

particular, portions of skin and blubber with embedded crustacean parasites were collected; the bronchial tree was longitudinally opened and pulmonary parenchyma was dissected and inspected for parasites. Vessels and liver ducts were dissected and examined for lesions and parasites; portion of the liver was sliced, washed and the sediment was examined under a stereomicroscope. Stomach chambers and portions of intestine (small and large intestine at least 10m each) were separately sampled and the contents were filtered with 1.0 and 0.5 mm mesh sieves; the material was then observed by stereomicroscope. As previously described by Lambertsen (1992), intestinal mucosa and mesenteric arteries were examined to detect lesions due to the migration of parasitic larvae of the genus *Crassicauda*. Renal vessels and vena cava were also examined to search for adult specimens of *Crassicauda boopis* or related lesions. Ureteral ductworks were opened and accurately examined to isolate the tails of this nematode. When possible, urine sediment, obtained by centrifugation at 2000 rpm for 5 minutes, was analyzed for the detection of eggs or larvae of *Crassicauda*. All parasites recovered during post mortem examination were washed in physiological saline, counted and fixed in 70% ethanol.

Morphometric characteristics of crustacean species of the genus *Pennella* were studied by stereomicroscopy. The nematodes and immature stages of crustaceans were examined as wet-mounts in clearing agents (glycerin or Amman's lactophenol), whereas flatworms were stained with Semichon's Acid Carmine and mounted in Canada balsam. All parasites were measured under light microscope (Nis Elements D software, Nikon).

Crustacean parasites were compared with descriptions by Thompson (1905), Hogans (1987) and Abaunza et al. (2001). Dichotomous keys and literature data were used to identify the helminths (Delyamure 1955, Margolis & Pike 1955, Lambertsen 1985, Raga et al. 1986, Bray et al. 2008).

Specimens of *C. boopis* were deposited at the Natural History Museum of London (NHMUK) (accession number: *Crassicauda boopis* 2015.10.11.1-4).

Tissue samples showing lesions were fixed in 10% buffered neutral formalin, paraffin-embedded, cut (4 μ m thickness) and routinely stained with hematoxylin and eosin for microscopic examination.

78 Further histochemical techniques were used on selected sections in order to gain better information
79 on pathogens and tissue changes (PAS and Masson's Trichromic).

80

81 *Analyses for protozoan parasites*

82 During necropsy, tissue samples (brain, spinal cord, heart, lymph nodes, skeletal muscle, lung, spleen,
83 liver and kidney) were collected; a portion of each tissue was processed for routine histological
84 examination and an aliquot was stored at -20°C and subsequently analyzed by molecular methods to
85 detect parasites of the family Apicomplexa.

86 DNA extraction and PCR assay for detection of *Toxoplasma gondii*, *Neospora*, and *Sarcocystis*

87 DNA extraction was performed on all the aforementioned tissues using NucleoSpin® Tissue kit
88 (Macherey-Nagel, Germany). The PCR reaction was carried out in 30µl volume containing 1X PCR
89 buffer, 2 mM MgCl₂, 200 µM each of the dNTPs, 2 U Platinum® Taq DNA Polymerase (Invitrogen,
90 UK), 1µM of each primer, as described previously (Ho et al. 1996), and 1-3 µl of DNA extract. This
91 PCR assay permits to amplify a conserved region of the *nss-rRNA* gene (300 bp in size) of *Neospora*
92 spp. and other Apicomplexa coccidian as *Sarcocystis* and *Toxoplasma*. The reaction mixture was first
93 treated at 95°C for 5 min, followed by 12 cycles at 94°C for 30 sec and 58°C for 30 sec; 23 cycles at
94 94°C for 30 sec, 52°C for 30 sec and 72°C for 30 sec and a final extension step at 72°C for 7 min.
95 DNA extracted from *T. gondii* oocysts, isolated from a domestic cat, were used as positive control.
96 PCR products were analysed by electrophoresis in SYBR Safe stained (Invitrogen, UK) 2% agarose
97 gel, visualised with Geldoc XR (Bio-Rad Laboratories, USA) under UV light, subsequently purified
98 and sequenced at BMR-Genomics (Padova, Italy). The sequences were analysed using ChromasPro
99 (version 1.42, Technelysium Pty Ltd., Australia) and compared in GenBank™ database using
100 BLASTn program (<http://www.ncbi.nlm.nih.gov/>).

101 Extracted DNA was also tested by a commercial Real Time PCR assay (*Toxoplasma* Q- PCR Alert
102 Kit, Nanogen Advanced Diagnostics S.p.a., Buttigliera Alta, Italy) on an ABIPRISM 7300 (Applied

103 Biosystem Carlsbad, USA) following the manufacturer's instructions. The real-time PCR assay
104 targeted the 529 bp repeat region (*REP529*, GenBank accession no. AF146527) of *T. gondii* DNA
105 (Homan et al. 2000). All DNA samples were tested in triplicate and each assay was considered
106 positive if at least one test of the triplicate was positive. Each PCR run included a negative control
107 without DNA and, to check the absence of PCR inhibitors, each sample was coamplified with an
108 internal control consisting of beta-globin gene.

109 Genotyping of *T. gondii* strains

110 DNA samples extracted from tissues that tested positive for *T. gondii* DNA (C_t value < 32) were
111 submitted to a genotyping analysis using 15 microsatellite markers distributed on 10 of 14
112 chromosomes, as described previously (Ajzenberg et al. 2010). Briefly, for each primer pair, the
113 forward one was 5'-end labeled with fluorescein to allow sizing of PCR products electrophoresed in
114 an automatic sequencer. PCR was carried out in a 25- μ L reaction mixture consisting of 12.5 μ L of
115 2X QIAGEN Multiplex PCR Master Mix (Qiagen, France), 5 pmol of each primer and 5 μ L of DNA.
116 Cycling conditions were 15 min at 95°C; 30 s at 94°C, 3 min at 61°C, and 30 s at 72°C (35 cycles);
117 and 30 min at 60°C. One microliter of the PCR product was mixed with 0.5 μ L of a dye-labeled size
118 standard (ROX 500, Applied Biosystems) and 23.5 μ L of deionized formamide (Applied
119 Biosystems). This mixture was denatured at 95°C for 5 minutes and then electrophoresed using an
120 automatic sequencer (ABI PRISM 3130xl, Applied Biosystems). The sizes of the alleles in bp were
121 estimated using GeneMapper analysis software (version 4.0, Applied Biosystems).

122

123 **3. Results**

124 *Metazoan parasites and histopathological findings*

125 Six out of seven fin whales were positive for one or more parasitic species. Overall, 1,164 parasites
126 were collected, belonging to one crustacean and four helminth taxa (Table 2).

127 Adult females of the mesoparasitic copepod *Pennella balaenopterae* (Copepoda, Pennellidae) were
128 detected in five whales. Mild infection was observed in four animals, in which the parasites were

129 anchored mainly on the back and on the abdominal region. In one case, a severe infestation (>200
130 parasites), involving the entire body surface, was observed (whale #3).

131 Immature specimens of parasitic copepods of the family Pennellidae Burmeister, 1835 (Fig. 1) were
132 collected from two animals. The head of the parasites was strongly embedded into the skin and
133 blubber without reaching the muscular fascia, while the trunk and the abdomen emerged from the
134 body of the host. These specimens were small in size (n=6; mean length: 33.9 mm), showed a distinct
135 cephalothorax dorsoventrally flattened and flecked with black pigment. Beyond the border of the
136 cephalothorax the hamate second antennae projected, while the first pair of antennae was delicate and
137 setose. In the ventral side of the cephalothorax, a prominent rostrum was present and the sturdy
138 maxillipeds were situated posteriorly; ventrally in the anterior third of the cephalothorax the cuticle
139 folds forming an hourglass design with a pair of spines in the central part. At the end of the
140 cephalothorax a pair of cuticular pointed structures were laterally present. The filiform thorax
141 extended to a length ten times that of the cephalothorax; the four pairs of thoracic limbs appeared
142 well developed with bristles in the last two segments; the first and second limbs bifurcated in two
143 extremities. The abdomen was filiform; eighteen pairs of bulges or short unbranched lateral
144 appendages (with different degrees of development in the specimens) were visible along the
145 abdomen, which terminated with a deep notch. On either side of this was a small bisetose appendage.
146 The morphometric data of the immature specimens of *Pennella* sp. Oken, 1815 are reported in Table
147 3.

148 Specimens of cestodes were collected from the small intestine of one adult whale. Morphometric
149 characteristics of the parasite and the morphology of the scolex (presence of well visible ear-like
150 appendages and well-developed median lobes on the apical organ) permitted to ascribe the specimens
151 to the species *Tetrabothrius ruudi* Nybelin, 1928 (Eucestoda, Tetrabothriidae). One specimen of
152 acanthocephalan *Bolbosoma* sp. Porta, 1908 (Acanthocephala, Polymorphidae) was detected from
153 the small intestine of one fin whale. The digenean *Ogmogaster antarcticus* Johnston, 1931
154 (Trematoda, Notocotylidae) was found in the rectum of three fin whales.

Crassicaudosis was suspected in five out of seven examined whales. Adult worms of the nematode *Crassicauda boopis* were found in four cases (Table 2). A total of ninety-five tails of *C. boopis* (range: 7-40 parasites/host) were found inside the ureters of the positive animals. In three cases both kidneys were parasitized, while in one case a unilateral infection was observed. The anterior part of the tails was found penetrating the wall of the urinary ductworks and extending into the renal parenchyma (Fig. 2F). Histopathological examinations of kidney revealed multifocal chronic nephritis with massive fibrosis, glomerular sclerosis and tubular atrophy in two animals (#4 and #5). In fin whale #4, larvated ova were histologically observed within the kidney pelvis among epithelial cells, in lumen of arterial renal vessels and in adrenal glands. Additionally, larvated eggs of *Crassicauda* sp. were also found in urinary sediment in fin whale #5.

Cephalic ends of *C. boopis* protruded free and flowing in the vena cava lumen of two animals (#4 and #5). Large pendulous masses (Fig 2B) and fragments of mineralized parasites were grossly observed within the lumen of arterial renal vessels of other two whales (#2 and #3 respectively).

Adult nematodes, peduncolate, knobby and mineralized proliferations obstructed partially or totally blood passage into the vessels. Microscopic exams revealed the presence of parasitic elements also inside these proliferations, containing large numbers of ova, completely embedded by connective tissue layers, multifocally infiltrated by mild chronic inflammatory population (Fig 2D). Nematodes observed lying within the lumen of the renal and adrenal arteries (Fig. 2E) were also partially embedded by a mixed chronic inflammatory response with a massive eosinophils prominence and a severe fibroblastic invasion. These changes multifocally infiltrated also the arteries' walls along with severe edema and hyperemia.

In fin whale #5 mesenteric arteries were characterized by a mural thickening with intimal hyperemia and hemorrhages, multifocal necrosis with cavitation and mineralized area (Fig 2C): these findings reduced vascular lumina. Microscopic examination confirmed a chronic mineralizing endo-arteritis with abundant and severe eosinophilic infiltration sometimes associated to parasitic larvae. Several nodules were grossly observed in the mucosal layer of the intestine of the same animal (# 5) (n >90;

181 diameter: 6-7mm), containing amorphous material, occasionally associated to the presence of a single
182 nematode larvae (Fig. 2A). Gross examination revealed a massive and diffuse greenish discoloration
183 of muscular tissues in this fin whale. Severe eosinophilic infiltration was microscopically observed
184 in several lymph nodes.

185 Epicardial and endocardial granulomas were detected in fin whales #1 and #6 with PAS positive
186 traslucid remains consistent with parasitic fragments digested by the inflammatory reactions, but no
187 parasites were isolated.

188 Additionally, nematode larvae were found free inside the lumen of the intestine of the calf whale #6;
189 some of them carried hints of pre and post-cloacal papillae, indicating a preadult stage.

190

191 *Protozoan parasites*

192 One (fin whale #3) of the seven examined animals was positive for *T. gondii* DNA that was detected
193 in heart, skeletal muscle, mesenteric lymph node and kidney samples (Table 4); the sequences showed
194 100% homology with *T. gondii* (GenBankTM accession no. AY 663792). The DNA sample that was
195 genotyped was extracted from the muscle sample, successfully amplified at 3 microsatellite markers,
196 and was identified as *T. gondii* Type II.

197 No lesions related to toxoplasmosis were observed in the histological sections of the examined animal
198 tissues.

199

200 **4. Discussion**

201 Complete parasitological surveys on fin whales living in the Mediterranean Sea are still lacking due
202 to the technical complexities encountered during necropsy, the large size of these animals and,
203 sometimes, to the difficulty of reaching the stranding site. Moreover, the preservation status of the
204 carcasses affects the analyses that can be carried out.

205 The presence on Mediterranean fin whales of ectoparasitic copepods *Pennella* sp. has been known
206 for a long time (Anthony & Calvet 1905). *Pennella* spp. are large mesoparasites infecting both teleost

207 fishes and marine mammals. The high variability of morphological features of species and stages of
208 parasite development have generated much debate amongst taxonomists (Kabata 1979). Although
209 numerous species of *Pennella* have been described, *P. balaenopterae* is the only recorded copepod
210 species that parasitizes marine mammals (Hogans 1987, Abaunza et al. 2001). The life cycle of *P.*
211 *balaenopterae* is poorly understood and only the adult female and the first naupliar stage have been
212 identified (Abaunza et al. 2001, Arroyo et al. 2002). Intensity of infection by *Pennella balaenopterae*
213 is considered an indicator of health status in cetaceans and long-term, cumulative tendencies of *P.*
214 *balaenopterae* infestation can be associated with challenged dolphin's immune system, debilitating
215 viral infection and high levels of polychlorinated biphenyls (Vecchione & Aznar 2014). Two cases
216 of severe infestation by *Pennella* spp. were described in fin whales stranded alive along Italian and
217 Turkish coastlines (Benvenuti et al. 1991, Çiçek et al. 2007). In our study, the degree of infestation
218 was mild in most cases, and comparable to that reported by other authors in fin whales from Antarctic
219 (Nishiwaki & Hayashi 1950, Mizue & Murata 1952) and Atlantic waters (Raga & Sanpera 1986). In
220 one animal, positive to *Dolphin Morbillivirus* and *Toxoplasma gondii* infection (Mazzariol et al.
221 2012), a severe infestation was already reported. An impairment of the immune system can be
222 suspected in the animal included in this study, mainly due to the presence of the viral infection, thus
223 the higher parasitic burden matches with assumptions in literature.

224 Other two whales, which have encountered DMV (Mazzariol et al. 2016), showed the presence of
225 immature stages of crustacean parasites belonging to the family Pennellidae. The morphological
226 features of these parasites appear similar to those of the youngest individual of *Pennella filosa*
227 (Linnaeus, 1758) described by Thompson (1905) in fish. This is the only report in literature for
228 immature stages of *Pennella* genus from definitive host. Molecular analyses could be useful to ascribe
229 these specimens to the species *P. balaenopterae*. The finding of young individuals of *Pennella* sp. in
230 these two sick young whales suggests that the animals had decreased their mobility shortly before
231 death, allowing during this period the colonization by the parasites, as proposed by Aznar et al. (1994)
232 in dolphins affected by a viral epizootic disease.

233 Adult tapeworms of the families Tetrabothriidae and Diphylobothriidae are described in cetaceans.
 234 Life cycles of these species involve a zooplankton crustacean as first intermediate host and marine
 235 mammals as their definitive hosts (Raga et al. 2008). Only the genera *Priapocephalus* Nybelin, 1922
 236 and *Tetrabothrius* (Eucestoda: Tetrabothriidae) have been described in mysticetes and members of
 237 the genus *Tetrabothrius* have been isolated from fin whales worldwide. *Tetrabothrius affinis*
 238 (Lönnerberg 1891) (syn. *Tetrabothrius wilsoni*) is reported by Delyamure (1955) in Norway, South
 239 Africa, New Zealand and Antarctica (South Shetlands islands); the same author reported *T. ruudi* in
 240 West Norway, France and Russian Pacific coast and Antarctica, but this genus had never been
 241 reported before in the Mediterranean.

242 Acantocephalan of the genus *Bolbosoma* are intestinal parasites of cetaceans. Most of the species are
 243 typical of baleen whales, which get infected most probably by ingestion of the cystacanth larvae
 244 contained in an intermediate or paratenic host (Gazzonis & Merella 2012). Euphasiids and copepods
 245 have been demonstrated to carry larvae of the genus *Bolbosoma* (Shimazu 1975, Tsimbalyuk 1980,
 246 Gregori et al. 2012). Five species are described in *Balaenoptera* spp., i.e. *B. brevicolle* (Malm, 1867),
 247 *B. nipponicum* Yamaguti, 1939, *B. turbinella* (Diesing, 1851), *B. balaenae* (Gmelin, 1790) and *B.*
 248 *hamiltoni* Baylis, 1929 (Delyamure 1955). The pathogenicity is linked to the anchorage of the
 249 proboscis to the intestinal wall, which is reported to cause ulceration and even perforation (Gibson et
 250 al. 1998). No gross lesions were observed in the intestine of this animal, probably due to the presence
 251 of a single specimen.

252 The genus *Ogmogaster* Jägerskiöld, 1891 include six species, that are identified by the number of
 253 longitudinal ridges on the ventral surface, the presence or absence of spines on the tegument and body
 254 size (Raga et al. 1986). *Ogmogaster antarticus* was reported in fin whales from the Spanish Atlantic
 255 coasts and from the Mediterranean basin (Raga et al. 1986, Malatesta et al. 1998). The species shows
 256 wide diffusion and low host specificity, being reported in both cetaceans and pinnipeds.

257 *Crassicauda boopis* has been reported in fin whales from Atlantic and Pacific Ocean (Lambertsen
 258 1986) and is considered endemic in the Atlantic population (Lambertsen 1992). Lambertsen widely

described the diffusion and the mechanisms of this parasitic disease in Atlantic fin whales (1986, 1992). Though the number of analyzed animals is limited, our data prove for the first time that this parasitic disease exists among the Mediterranean population. The degree of severity of the infestation appears to be from moderate to severe in our survey, considering the lesions observed. The localization of the parasite inside the hosts confirm the descriptions in literature, with the female specimens getting to the lumen of the vena cava with the cephalic portion and the male's head trapped in the renal venous vessels. Hypothetically, the localization of the head inside the lumen of vessels allows the worms to feed on host's blood (Lambertsen 1986). Depressed packed red cell volume (PCV) in infected whales, potentially due to chronic consumption of blood by the worms, was reported by Lambertsen but due to the post mortem conditions we could not perform a complete blood count to confirm the anemic condition. The massive inflammatory reaction of the host's tissues to the parasite was broadly observed, causing severe lesions that appeared very similar to those described in literature (Lambertsen 1986). Wide thrombotic masses occupying the lumen of vessels and diffuse flogosis and fibrosis of renal parenchyma were observed in all infected animals, with varying degrees of severity. Multidigitate masses pending in renal veins are also reported by Lambertsen, as consequence of the host's attempt to capsule the parasite. Such lesions are made up of fibrocellular tissue and they depart from the tunica media of the vessels. In chronic cases, a mineralized core can be observed right around the worm. Occlusion of the renal veins can occur as a consequence of the tissue reaction around parasite's body. The lesions observed in this survey completely overlap this description, showing different degrees of severity, that appeared greater in the animal with the higher parasitic burden (n=40). Thrombotic processes found in the kidney of this animal probably had reduced renal blood flow leading to organ impairment. Severe impact of the infection on the host was demonstrated also by serum biochemistry. High concentration of creatinine (6.95mg/dL), urea (157.8 mg/dl) and electrolytes (phosphate and potassium: 24.06mg/dL and 29.5mEq/L, respectively) were found in serum of the unique samples animal (Mignone, personal communication), significantly higher than those reported in literature in infected whales (Lambertsen 1992). In this individual,

eosinophils infiltration was also appreciated in several lymph nodes and muscular tissues and not only associated to parasites: these findings supports a possible increase in eosinophils blood count. Hypereosinophilia has been supposed to affect brain function in marine mammals (Di Guardo 2011) and this condition could have played a role in the stranding together with the renal function impairment suggested by blood chemistry. Other important parasitic lesions, that could have affected the health of this individual, were due to the contemporary presence of larval elements within intestinal and mesenteric arteries' walls along with a severe and chronic inflammatory reaction. Reasonably this finding could support the hypothesis of larval migration of *C. boopis* from intestine to renal vessels through the tunica media of mesenteric arteries, as speculated by Lambertsen (1992) and similarly to what occurs for *Crassicauda* sp. in Cuvier's beaked whales (Diaz-Delgado et al. 2016). Nevertheless, since the animal was previously affected by DMV determining a possible impairment of immune system, the association between larval migration and poor health condition cannot be clearly claimed.

In conclusion, crassicaudosis could represent a cause of concern for the Mediterranean population of fin whales. Particular attention should be paid to the probable wide diffusion of the parasite among the host population and to the high severity of the disease, inheriting further sampling effort.

Four types of Apicomplexa coccidians (Sarcocystiidae) are reported in cetaceans: *Cystoisospora delphini* in bottlenose dolphins, *Sarcocystis* spp. in toothed whales and in a striped dolphin, *Neospora caninum* in bottlenose dolphins, and *T. gondii* in four dolphin species, in one harbor porpoise (*Phocoena phocena* [Linnaeus, 1758]) and recently in one sperm whale (*Physeter microcephalus* Linnaeus, 1758) (Domingo et al. 1992, Di Guardo et al. 1995, Cabezon et al. 2004, Raga et al. 2008, Mazzariol et al. 2011). As for Mysticetes, the reports concerning Sarcocystiidae are limited; Akao (1970) described *Sarcocystis balaenopterale* n. sp. in muscle tissues of a Sei Whale (*Balaenoptera borealis* Lesson, 1828) and Forman et al. (2009) reported the presence of *T. gondii*-specific antibodies in a Humpback whale (*Megaptera novaeangliae*, Borowsky, 1781) using the Sabin Feldman Dye

310 Test. The presence of a coinfection by *T. gondii* and Dolphin morbillivirus was reported by Mazzariol
 311 et al. (2012), in one fin whale included in this study.

312 The result of genotyping indicates the presence of Type II. This genotype seems common in marine
 313 mammals since it was isolated in the California sea otter (*Enhydra lutris* [Linnaeus, 1758]), striped
 314 dolphin (*Stenella coeruleoalba* [Meyen, 1833]), bottlenose dolphin (*Tursiops truncatus* [Montagu,
 315 1821]) and Walrus (*Odobenus rosmarus* [Linnaeus, 1758])(Cole et al. 2000, Miller et al. 2004,
 316 Sundar et al. 2008, Di Guardo et al. 2011); Non-type II genotypes were reported in marine mammals
 317 such as those belonging to haplogroup 12 in North America (formerly Type X and A) in the California
 318 sea otter (*E. lutris*) (Khan et al. 2011), harbor seal (*Phoca vitulina* Linnaeus, 1758) and Californian
 319 sea lion (*Zalophus californianus* [Lesson, 1828]), (Cole et al. 2000, Miller et al. 2004, Sundar et al.
 320 2008, Van Bressem et al. 2009).

321 This study is the first genotyping attempt of a *T. gondii* strain from Mysticetes. Genotype II
 322 predominates not only in terrestrial mammals and birds of North America and Europe, but also in the
 323 marine environment of these areas.

324 It seems that the route of infection by *T. gondii* in cetaceans include ingestion of oocysts with
 325 contaminated water, as these animals feed mainly on cold-blooded animals as fish and invertebrates,
 326 which are not expected to support the tachyzoite and bradyzoite life-stages of *T. gondii* (Forman et
 327 al. 2009). The oocysts, under controlled laboratory conditions, can sporulate and remain viable in
 328 seawater for several months (Lindsay & Dubey 2009) and they can remain infectious in filter-feeding
 329 fish's alimentary canals for some hours post-exposure (Massie et al. 2010). DNA of *T. gondii* was
 330 also detected in shellfish in USA, Brazil and in Italy (Putignani et al. 2011).

331 The infected fin whale was observed swimming in shallow water in front of the Tuscany coast some
 332 days before stranding and it could be supposed that the animal might have been infected in this period
 333 from coastal waters. The animal showed also a dolphin morbillivirus infection and high concentration
 334 levels of organochlorine pollutants in the tissues. These data confirm the opportunistic nature of

335 *Toxoplasma*, which can infect immune-depressed animals, already threatened by infectious diseases
336 and environmental contaminants (Mazzariol et al. 2012).

337

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350

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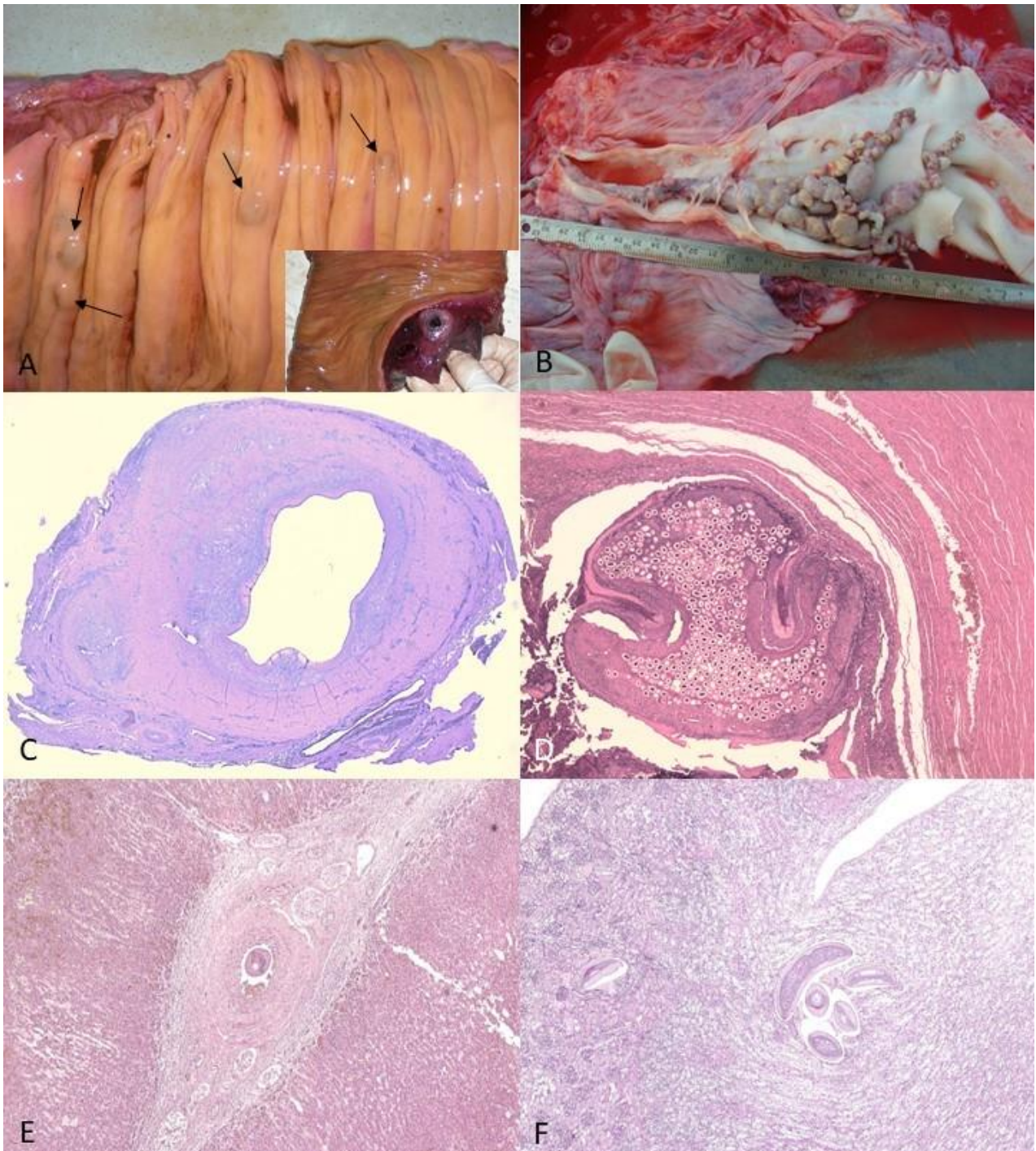
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Fig.1: Immature stages of *Pennella* sp. (A). Abdominal portion of the parasite protruding from the host's skin (bar=0.5 cm); (B) Lateral view of cephalothorax after removing the surrounding tissues (bar=0.5 cm); (C) Ventral view of the head with first (black arrow) and second (black arrowhead) antennae, rostrum (*) and maxillipeds (white arrow) (bar=300µm); (D) Cephalothorax, details of the limbs (ventral view)(bar=500µm); (E) Abdomen of the parasite at two different developmental stages of the appendages.



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520 Fig.2: Nematode infections. (A) #5. Multifocal, well-defined nodules (arrows), containing nematode
 521 larvae (insert) visible on the surface of the intestinal mucosa. (B) #2. Knobby and mineralized
 522 formation in the vascular lumen of a renal vessel, partially obstructing blood passage. (C) #5.
 523 Histological section of a mesenteric artery characterized by a severe mural thickening with intimal
 524 hyperemia and hemorrhages, multifocal necrosis with cavitation and mineralized area. 1X
 525 magnification, Hematoxylin and Eosin (HE). (D) #2. Parasitic thrombus formed by an adult element

526 of *C. boopis*. 4X magnification, HE. (E) #4. Adult element of *C. boopis* in the lumen of an artery in
527 the adrenal parenchyma, characterized by a severe fibroblastic invasion. 4X magnification, HE. (F)
528 #6. Some adult elements of *C. boopis* in the renal parenchyma (medulla). 4X magnification, HE.
529

530 Table 1. Data of fin whales specimens analyzed in this study

Fin whale	ID	CCC	BCS	Age	Sex	Body lenght (m)	Year	Stranding site	DMV*
#1	109	3	Poor	Newborn	M	5.57	2006	Alassio (SV)	-
#2	134	2	Moderate	Juvenile	M	13.40	2008	Giannella (GR)	-
#3	194	3	Poor	Adult	M	16.7	2011	San Rossore (PI)	RT-PCR + (liver, spleen, lung)
#4	208	2	Poor	Juvenile	F	10.78	2011	Capo Testa (OT)	RT-PCR + (liver, spleen, lymph node, muscle)
#5	211	2	Poor	Juvenile	M	10	2011	Savona (SV)	VN +
#6	297	2	Poor	Newborn	F	5	2013	Marciana (LI)	RT-PCR + (brain, spleen, lung, thymus), IHC + (brain, thymus)
#7	342	3	Moderate	Adult	M	17	2015	Camaione (LU)	-

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ID = Identification code; CCC = Carcass Condition Code, according to Geraci and Lounsbury, 2005; BCS = Body Condition Score; F = Female; M = Male; DMV=Dolphin morbillivirus; RT-PCR = reverse transcription PCR; VN = virus neutralization; IHC = immunohistochemical analysis; * = Mazzariol et al. 2016

536 Table 2. Metazoan parasites collected and identified from the examined fin whales.

Organs	Parasite	Fin whale #1	Fin whale #2	Fin whale #3	Fin whale #4	Fin whale #5	Fin whale #6	Fin whale #7
Skin-blubber	<i>Pennella balaenopterae</i> (Copepoda; Pennellidae)	Neg	10	24	3	2	Neg	16
	Immature stage of <i>Pennella</i> sp. (Copepoda; Pennellidae)	Neg	Neg	Neg	15	Neg	3	Neg
Intestine	<i>Ogmogaster antarcticus</i> (Trematoda; Notocotylidae)	Neg	712	254	Neg	Neg	Neg	1
	<i>Tetrabothis ruudi</i> . (Eucestoda; Tetrabothisidae)	Neg	Neg	Neg	Neg	8	Neg	Neg
	<i>Bolbosoma</i> sp. (Acanthocephala; Polymorphidae)	Neg	Neg	Neg	Neg	1	Neg	Neg
	Nematoda larvae (inside nodules)	Neg	Neg	Neg	Neg	7	Neg	Neg
	Nematoda larvae (free into the lumen)	Neg	Neg	Neg	Neg	Neg	9	Neg
Kidney /ureters	<i>Crassicauda boopis</i> tails	Neg	40 (36 M; 4 F)	Neg	38 (24 M; 14 F)	10 (8M; 2 F)	Neg	7
Vena cava	<i>Crassicauda boopis</i> heads	Neg	Neg	Neg	11	Pos	Neg	Neg
Mesenteric arteries	Nematoda larvae (into the vessels wall)	Neg	Neg	Neg	Neg	4	Neg	Neg.

537 Neg = negative; Pos = positive for the presence of *C. boopis*, but not quantitatively determined; F = female; M = male

538

539 Table 3. Mean and individual measurements (in μm) of six immature specimens of *Pennella* sp. Oken, 1815 in different developmental stages.

Total length			33899	21582	26641	27245	32302	43525	52097	540
Cephalothorax	Total length including limbs	2573	2404	2516	2420	2644	2694	27541	27541	541
	Head length	1910	1726	1967	1847	1908	1978	2034	2034	542
	Head width	781	665	747	797	817	824	837	837	543
										544
Thoracic region	Diameter in the middle region	444	392	344	402	472	494	559	545	545
	Total length	25257	14655	18260	18475	23192	34271	42689	42689	546
										547
Abdomen	Total length	6068	4522	5865	6351	6467	6560	6645	6645	548
	Width in the middle region	364	340	365	370	310	324	476	476	549
	Width including appendages	631				630	668	555	550	550
	Appendages		Poorly developed	Poorly developed	Poorly developed	Present	Present	Present	Present	551
										552

553

554

555 Table 4. Results of *T. gondii*, *Neospora*, and *Sarcocystis* detection by PCR and sequencing.

Tissues	Fin whale #1	Fin whale #2	Fin whale #3 °	Fin whale #4	Fin whale #5	Fin whale #6	Fin whale #76
Brain	Negative	Negative	NOT DONE	Negative	Negative	Negative	Negative
Spinal cord	Negative	Negative	NOT DONE	Negative	NOT DONE	Negative	Negative
Heart	NOT DONE	Negative	<i>T. gondii</i> *	Negative	Negative	Negative	NOT DONE
Skeletal muscle	Negative	Negative	<i>T. gondii</i> *	Negative	Negative	Negative	Negative
Lymph nodes	Negative	NOT DONE	<i>T. gondii</i> *	Negative	Negative	Negative	Negative
Spleen	NOT DONE	NOT DONE	Negative	NOT DONE	NOT DONE	NOT DONE	NOT DONE
Liver	NOT DONE	Negative	Negative	Negative	Negative	Negative	Negative
Lung	Negative	Negative	Negative	Negative	Negative	Negative	Negative
Kidney	NOT DONE	Negative	<i>T. gondii</i> *	Negative	Negative	Negative	Negative

567 * *T. gondii*, GenBank accession number AY663792; ° Mazzariol et al. 2012

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