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Microscopic ossicle analyses and the complete mitochondrial genome sequence of *Holothuria (Roweothuria) polii* (Echinodermata; Holothuroidea) provide new information to support the phylogenetic positioning of this sea cucumber species

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(Article begins on next page)

1 **Microscopic ossicle analyses and the complete mitochondrial genome sequence of *Holothuria***  
2 **(*Roweothuria*) *polii* (Echinodermata; Holothuroidea) provide new information to support the**  
3 **phylogenetic positioning of this sea cucumber species**

4

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13

14 **Short title:**

15 Ossicles and mtDNA information from *Holothuria polii*

16 **Abstract**

17 Sea cucumbers (Holothuroidea) are ecologically important organisms for their bioturbation and  
18 alkalization activities of the seabed. These species are extensively fished as they are considered  
19 luxury food. Sea cucumbers are also relevant for biomedical studies and the production of bioactive  
20 compounds. A few initiatives are recently evaluating sea cucumbers as novel aquaculture species.  
21 The aim of this study was to provide morphological and genetic information useful for the  
22 identification of *Holothuria polii*, the white spot sea cucumber (a common species of the  
23 Mediterranean Sea). We generated the complete sequence of the mitochondrial DNA (mtDNA)  
24 genome of this species and combined it with a detailed ossicle characterization of the sequenced  
25 specimen by scanning electron microscopic analysis. Ossicles (known also as sclerites) are  
26 anatomical features that can discriminate Holothuroidea species, including the closely related ones of  
27 the genus *Holothuria*. The complete mitochondrial genome was assembled, functionally annotated  
28 and then used to evaluate the phylogenetic relationship of *H. polii* against the other few Holothuroidea  
29 species for which the whole mtDNA was available. The 15,907 bp *H. polii* mtDNA sequence has the  
30 same gene order already reported for *H. scabra*, *H. forskali* and other species of the same class. *CoxI*  
31 and 16S gene sequences were informative for species identification across the genus and could be  
32 used for the authentication of commercialized *Holothuria* spp. The mitochondrial genome sequence  
33 presented here provides the basis to a future analysis of the variability of *H. polii* populations in the  
34 Mediterranean region.

35

36 **Keywords:** mtDNA; phylogeny; sandfish; sclerite; species identification; white spot sea cucumber.

## 37 **1. Introduction**

38           Within the phylum Echinodermata (Leuckart, 1854), which comprises five classes of marine  
39 invertebrates, the class Holothuroidea (known as sandfishes or sea cucumbers) includes about 1400  
40 species (Pawson, 2007). This class represents 90% of the deep-sea floor biomass, therefore its species  
41 are considered amongst the most dominant organisms in the world (e.g.: Pawson and Pawson, 2008;  
42 De Leo et al., 2010).

43           Holothuroid phylogeny is subject to controversies and ambiguities and demands close  
44 inspection and perhaps re-evaluations. Kerr and Kim (2001) used 47 morphological traits and  
45 performed a cladistic analysis to assess the relationship among the orders belonging to the class  
46 Holothuroidea initially established by Bronn (1860). Recent molecular data obtained from both  
47 mitochondrial and nuclear markers identified seven orders (Miller et al., 2017).

48           Within this class, to date the nuclear genome of only three species has been preliminarily  
49 assembled and genome scaffolds are available for *Apostichopus parvimensis*, *A. japonicus* and  
50 *Australostichopus mollis* (Kudtarkar and Cameron, 2017). A total of 10 complete mitochondrial  
51 genomes has been obtained for species of this class (only nine of which are available in GenBank;  
52 Long et al., 2016; Kudtarkar and Cameron, 2017; Wang et al., 2019).

53           In several regions of the world, holothurian fishing and rearing are practiced to supply specific  
54 food markets, mainly driven by Asian countries (Han et al., 2016). Therefore, several efforts have  
55 been made to study and catalogue sea cucumber exploitation hotspots all over the world. Over-  
56 exploitation caused genetic flow loss among populations and in some places their complete extinction  
57 (Friedman et al., 2011; Soliman et al., 2016). One of the main reasons of this excessive harvest is the  
58 rising demand of the Asian luxury food and traditional medicine markets (Purcell et al., 2014).  
59 Moreover, sandfishes consumption, as traditional and valuable food, has raised interests on the  
60 nutritional properties of these animals, highlighting the presence of antioxidant molecules together  
61 with a high protein content and a low fat level, also in the dried form, known as “bêche-de-mer” or  
62 *trepang* (Wen et al., 2010; Roggatz et al., 2016). Other reasons of interests on sea cucumbers derive

63 by their use as models for tissue and organ regeneration, by their peculiar adaptations and by their  
64 relevance for bioactive compounds production, such as holothurins (García-Arrarás and Dolmatov,  
65 2010; Jaeckle and Strathmann, 2013; Zhang et al., 2017).

66 *Holothuria (Roweothuria) polii* (Delle Chiaje, 1823), also known as white spot cucumber, is a  
67 neritic marine organism belonging to the Holothuriidae family (Aspidochirotida). This species lives  
68 in a depth between 0 to 250 m along the coasts of Mediterranean and Black Seas, Suez Gulf and in  
69 some spots of the Atlantic Ocean coasts (Coll et al., 2010). *Holothuria polii*, as other holothurians,  
70 plays an important ecological role as a detritivorous in benthic communities doing both a bioturbation  
71 of the marine sediments and a buffer activity counteracting the effects of water acidification and  
72 showing physiological plasticity (Vergara-Chen et al., 2010; Yuan et al., 2018). *Holothuria polii* is  
73 also one of the most exploited species in terms of commercial trade for food and pharmaceutical  
74 purposes in the Mediterranean area. Turkey has the main sea cucumber commercial fishery where  
75 about 80% of the harvested holothurians belongs to *H. polii* species (González-Wangüemert et al.,  
76 2014). The high market demand, mainly from Asian countries, is increasing the risk to deplete wild  
77 stocks of this species, with a reduction of biodiversity and benthic biomass, which might break the  
78 ecological and chemical marine balances (Purcell et al., 2016; Pawson and Pawson 2008). Therefore,  
79 efforts are necessary to preserve wild sea cucumber populations and to establish new efficient rearing  
80 methods and aquaculture production systems for this species (Bell et al., 2007; Purcell et al., 2013;  
81 Ren et al., 2014; Beltran-Gutierrez et al., 2016).

82 In parallel to conservation actions, it is important to develop new methods and tools to simplify  
83 the identification of *H. polii*. This is needed for species authentication of luxury holothurian food  
84 products to identify frauds and illegal trades of this species, recently studied as a new candidate for  
85 aquaculture (Conand et al., 2018; Rakaj et al., 2019). Morphological identification of species within  
86 the *Holothuria* genus is mainly based on the shape, size and fine details of endodermal ossicles (or  
87 sclerites) which are calcified structures that are part of the echinoderm endoskeleton (Koehler, 1924;  
88 Tortonese, 1965; Aydin and Erkan, 2015). However, the morphological analysis has often led to

89 wrong species assignment of holothurians because ossicles, within species, can change shape,  
90 typologies and location in different body regions (Cutress, 1996; Massin et al., 2000). In addition, a  
91 detailed and complete characterization of ossicles is not available for most species. Hybridization  
92 events between sympatric species and subspecies, that can lead to animals with mixed morphological  
93 features might add confounding factors (Uthicke et al., 2005; Yoshida et al., 2012; Kim et al., 2013).  
94 Thus, molecular information is therefore necessary in order to complete and clarify the inter and  
95 intraspecific diversity of sea cucumbers (Aydin and Erkan, 2015; Dettai et al., 2011; Valente et al.,  
96 2015). At present, molecular phylogeny of the Holothuriidae family, based only on *cox1*, 16S  
97 mitochondrial DNA (mtDNA) sequences and 18S nuclear sequences, is still unresolved for some  
98 subgenera resulting in paraphyletic groups including *Roweothuria*, which in turn includes *H. polii*  
99 (Kerr et al., 2005; Honey-Escandon et al., 2011; Borrero-Pérez et al., 2010).

100 In this study, as a first step for a detailed description of the genetic variability in *H. polii*  
101 populations, we sequenced the complete mitochondrial genome of this species and compared this  
102 mitogenome with the available mtDNA sequences of other holothurian species. In addition,  
103 mitochondrial genome information of *H. polii* was evaluated in a comparative analysis with ossicle  
104 morphology and distribution obtained using detailed microscopy inspection. The produced molecular  
105 and morphological results filled a gap in the phylogenetic analysis of *H. polii* and provided important  
106 classification and identification tools, also useful for the authentication of this species in food and  
107 drug preparations.

108

## 109 **2. Materials and Methods**

### 110 **2.1. Specimen and morphological characterization**

111 A sea cucumber specimen was collected in the western coast of Sardinia (Oristano province).  
112 The identification of the species was conducted following dichotomous keys (Tortonese, 1965;  
113 Koehler, 1924) and using the criteria reported by Aydin and Erkan (2015) who suggested the use of  
114 complementary information on bathymetry and body coloration and shape.

115

## 116 **2.2. Microscopic analyses of the ossicles**

117 Ossicles were analysed by using optical microscopy and scanning electron microscopy (SEM).  
118 Microscopy samples were prepared starting from 1 g of different tissues using sodium hypochlorite  
119 to eliminate non-calcified material. Preparations were from the internal tegument portion of the  
120 *bivium* (dorsal part of the animal) including papillae, tentacles, anal tegument or wall and the  
121 tegument around the calcareous ring. After the digestion of the organic matrix, specimens were  
122 carefully washed with bi-distilled water, taking care to preserve the ossicles. Optical microscopy  
123 analysis of the dorsal body wall was performed to check the cleanliness of the digested organic matrix.  
124 The visualization and the image acquisition were performed using an optical microscope Laborlux  
125 12 (Leitz, Wetzlar, Germany), resuspending purified ossicles in a solution of bi-distilled water. The  
126 ossicles were visualized with a 40X magnification and acquired by Infinity 1-5C camera software  
127 (Teledyne Lumenera, Lumenera Corp. 7 Capella Crt. Ottawa, Ontario, Canada K2E 8A7). For the  
128 SEM analyses, purified ossicle samples were mounted on a glass slide stuck on an aluminum stub  
129 using Silver conductive glue (Silver Print, Provac AG). Specimens were sputtered with 2 nm gold  
130 particles using K500 instrument (Emitec, Lohmar, Germany) at 30 mA for 2 min. Ossicles were  
131 observed using a SEM 515 microscope (Philips, Electronic Instruments, Eindhoven, The  
132 Netherlands) at 10 kV with a spot size of 20 nm. Images acquisition was performed with a K-5 camera  
133 (Pentax, Tokyo, Japan).

134

## 135 **2.3. DNA extraction, library preparation and next generation sequencing**

136 DNA extraction was performed using the Wizard® Genomic DNA Purification Kit (Promega  
137 Corporation, Madison, Wisconsin, USA) following the manufacturer's instructions. Genomic DNA  
138 was quality checked performing an electrophoresis on a 0.8% agarose gel and quantified using the  
139 Qubit 2.0 fluorometer (Invitrogen, Thermo Fisher Scientific, Waltham, Massachusetts, USA).

140 Next generation sequencing included the following procedures for the library preparation.  
141 Genomic DNA was sheared through sonication. Next, after fragment end repair, sequencing adapters  
142 were ligated to both ends and DNA was amplified in an indexing PCR. Then library size distribution  
143 was confirmed using Bioanalyzer instrument (Agilent, Santa Clara, CA, USA) and size selection was  
144 performed using the BluePippin System (Sage Science, Beverly, Massachusetts, USA) with a 2%  
145 agarose gel cassette. Finally, the library was sequenced on an Illumina (San Diego, CA, USA) HiSeq  
146 2500 with  $2 \times 150$  PE rapid run chemistry and 100 bp reads were obtained with an insert size of 450  
147 bp and an inner distance of 250 bp.

148

#### 149 **2.4. Next generation sequencing reads, mtDNA assembly and annotation**

150 Quality of the reads was evaluated using FastQC v.0.1.1.7  
151 (<https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>) that highlighted very high-quality  
152 reads. No other filtering procedures were adopted. Reads were assembled via the iterative approach  
153 implemented in MITObim 1.9.1 (Hahn et al., 2013) using the cytochrome-oxidase subunit 1 (*coxI*)  
154 sequence of *H. polii* (GenBank: KJ493895.1) as reference. To evaluate the quality and the reliability  
155 of the assembled genome, reads were subsequently mapped on it with BWA tool 0.7.17 (Li and  
156 Durbin, 2009) by computing with SAMtool v1.7 (Li et al., 2009) the following parameters: i) breadth  
157 and depth of coverage and ii) the length of the insert size. The annotation of the constructed mtDNA  
158 genome was obtained by using MITOS WebServer (<http://mitos.bioinf.uni-leipzig.de>) and by  
159 manually curating gene boundaries using the NCBI tool ORFfinder  
160 (<https://www.ncbi.nlm.nih.gov/orffinder/>). RNA genes were validated with MFold  
161 (<http://unafold.rna.albany.edu>) and ARWEN (<http://mbioserv2.mbioekol.lu.se/ARWEN/>) software  
162 tools (Zuker, 2003; Bernt et al., 2013; Laslett and Canbäck, 2007). Then, the mitochondrial genome  
163 map was prepared using GenomeVx (<http://wolfe.gen.tcd.ie/GenomeVx/>), setting the *coxI* gene as  
164 the starting point of the mtDNA (Conant and Wolfe, 2008). The complete annotated mitochondrial

165 sequence has been deposited in ENA within the project number PRJEB31737 and the accession  
166 number NC\_045029.

167

## 168 **2.5 Molecular species assignment and phylogenetic analyses**

169 Molecular analyses were performed using the most representative sequences of the species of  
170 holothurians present in NCBI nucleotide database. A portion of 412 bp of all holothurian *cox1* gene  
171 sequences including other sequences from *H. polii* and including the *de novo* assembled *H. polii*  
172 portion of *cox1* (from positions 312 to 723 of the mtDNA genome) was used to confirm the obtained  
173 species assignment using BLASTn analysis (Altschul et al., 1990). Similarly, a portion of 443 bp the  
174 16S rRNA sequence (from position 15,206 to 15,648 of the mtDNA genome) was compared with the  
175 most representative 16S holothurian sequences in order to confirm the *cox1* outputs.

176 The MEGA X software suite (Kumar et al., 2018) was used to compute codon usage, nucleotide  
177 composition statistics and to carry out the phylogenetic analyses. Three phylogenetic trees were  
178 computed. The first one was based on the complete aminoacidic sequence of *cox1* gene of the  
179 holothurians for which the complete mtDNA was available in NCBI (<https://www.ncbi.nlm.nih.gov/>)  
180 while the second one was based on the complete 16S rRNA sequences of the same species of  
181 holothurians. A total of ten species plus the outgroup *Strongylocentrotus purpuratus* (accession  
182 NC\_001453) were considered (Table S2). Initially, BLASTn analysis was used to compare *cox1* and  
183 16S rRNA gene sequences and confirm species identification. *Cox 1* nucleotide sequences were  
184 translated using the Echinoderm mitochondrial genetic code and the multiple sequence alignment  
185 (MSA) was obtained using the CLUSTALW algorithm (Thompson et al., 1994). A *maximum*  
186 *likelihood* (ML) phylogenetic tree was obtained using default settings (molecular evolutionary model:  
187 Jones – Tailor – Thorton) with 1,000 bootstrap replicates (Jones et al., 1992). The second  
188 phylogenetic tree was computed starting from the MSA of 16S rRNA sequences obtained using the  
189 Q-INS-i algorithm implemented in the online version of MAFFT v7.427 (Katoh et al., 2019) in which  
190 secondary structure of RNA were considered. Then, the evolutionary history was inferred by using

191 the *maximum likelihood* (ML) method and General Time Reversible model with 1,000 bootstrap  
192 replicates (Nei and Kumar, 2000).

193 The third phylogenetic tree was built by considering the complete mitochondrial DNA sequence  
194 of nine holothurian species plus the outgroup *Strongylocentrotus purpuratus* (accession  
195 NC\_001453.1) excluding the *Peniagone sp.* mtDNA because of the lack of *trnC* gene in the  
196 annotation. Genomes were aligned using MAFFT v7.427 with default settings (Katoh and Standley,  
197 2013). The MSA was manually curated and a Neighbour Joining (NJ) phylogenetic tree was obtained  
198 by computing evolutionary distances using the Maximum Composite Likelihood method (Tamura et  
199 al., 2004). The rate variation among sites was modelled with a gamma distribution (shape parameter  
200 = 1).

201 Finally, the gene order of the *H. polii* mtDNA genome was compared with those of the other  
202 holothurians mitochondrial genomes in order to detect putative rearrangements within the  
203 Holothuroidea class.

204

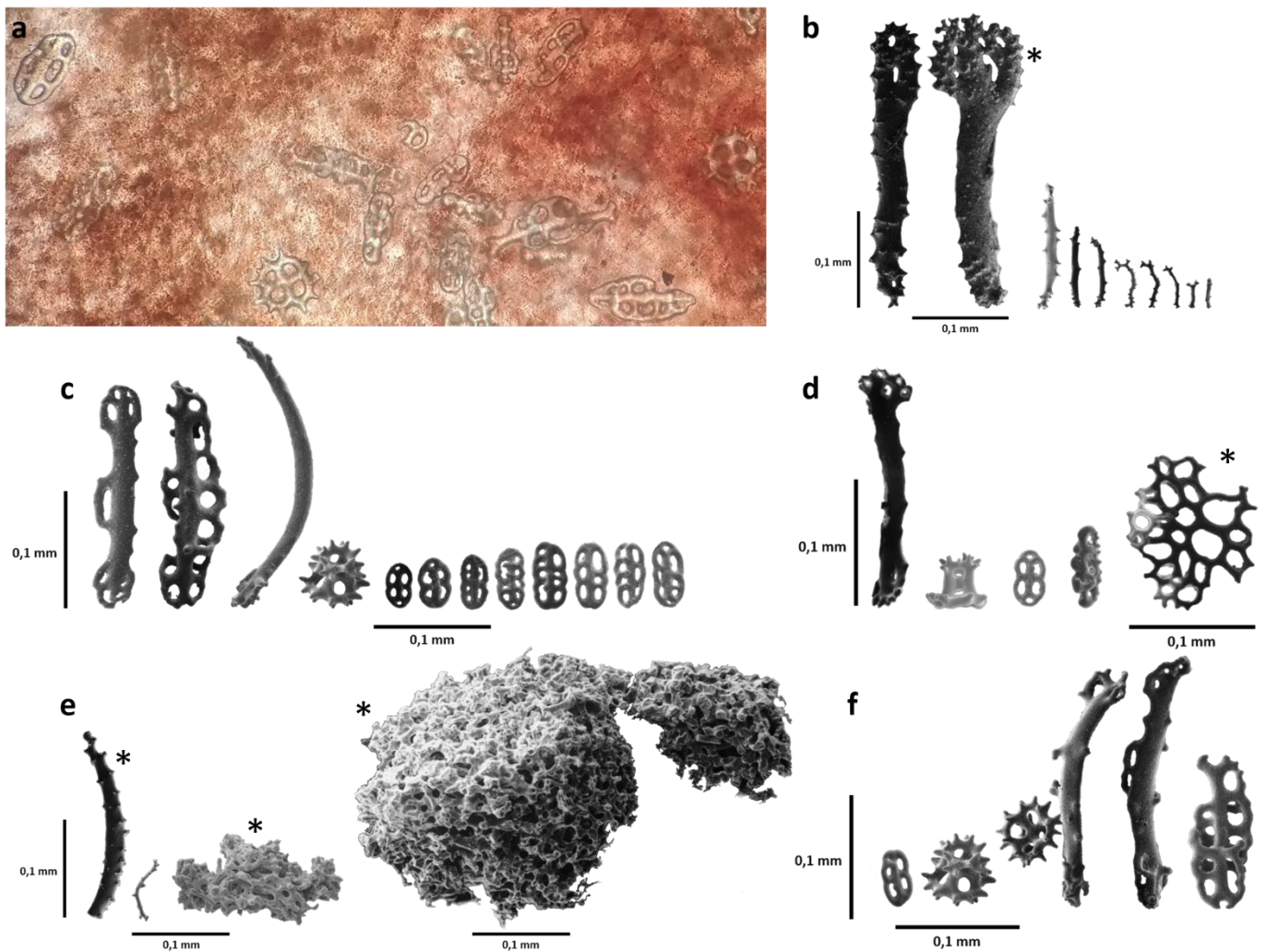
## 205 **3. Results**

### 206 **3.1. Species identification based on ossicle analysis**

207 The application of dichotomous keys (Tortonese, 1965; Koehler, 1924) and the use of  
208 information on the bathymetry and body coloration and shape (Aydin and Erkan, 2015) indicated that  
209 the analysed specimen was from *H. polii*. To confirm this assignment, ossicles were microscopically  
210 analysed.

211 SEM analyses of the ossicles revealed smooth surfaces, perforated buttons and tables as  
212 common features of the ossicle morphology; some examples are shown in Figure 1. The different  
213 analysed parts of the body showed various types of ossicles, some of which having different shapes,  
214 as already reported on the dichotomous keys (Tortonese, 1965; Koehler, 1924). In particular, the  
215 *papillae* tegument contained spiny rods, coral-like rods and some large concretions not described in  
216 the literature yet. The internal tegument covering the calcareous ring showed the presence of both

217 regular and irregular smooth buttons, knobbed buttons and smooth tables. In addition, rods with  
 218 smooth surface and enlarged perforated ends were present. Tentacles showed only rods of different  
 219 size and shape. The external anal tegument contained smooth tables, buttons and rods, the latter with  
 220 irregular shape. A few other ossicles had a rod-like convoluted shape, with large and numerous holes.  
 221 Finally, the dorsal tegument (*bivium*) showed smooth tables, regular and irregular buttons, curved  
 222 rods perforated at the end and rod-like convoluted ossicles (Figure 1). Most of these ossicles were  
 223 already described by Moussa and Wirawati (2018) in *H. polii* and led to the identification of this  
 224 species. However, thus far several shapes described here have not been reported in this species. Figure  
 225 1 describes the novel ossicle types.



226  
 227 **Figure 1.** a. Example of sclerites included in the organic matrix of *Holothuria polii* seen with optical microscope (40X).

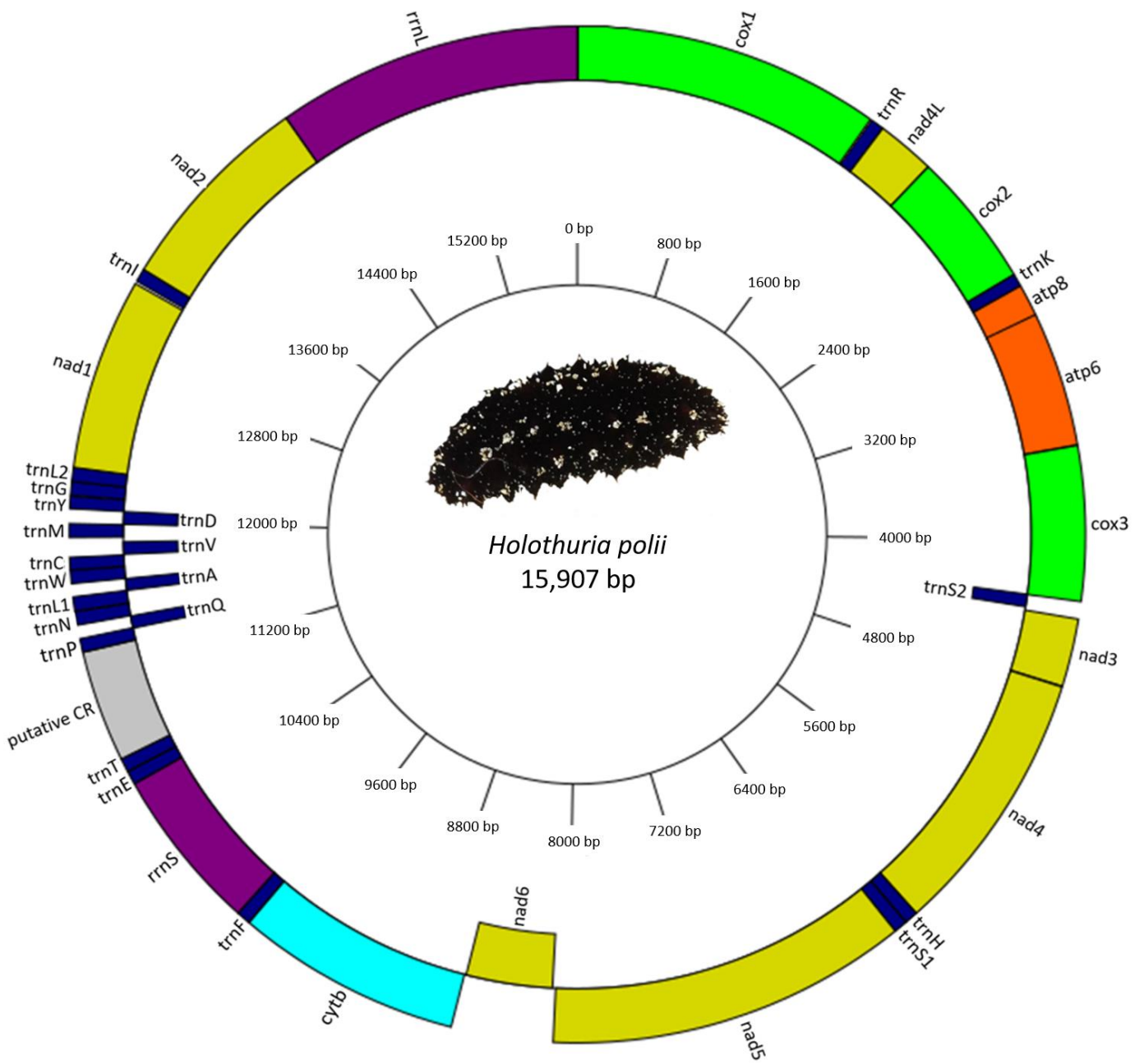
228 b. c. d. e. f. Sclerites seen with SEM with scalebars near and below for each body portion analysed. b: Tentacles. c: *bivium*.  
229 d: Internal tegument of calcareous ring. e: Dorsal *papilla*. f: Anal tegument.  
230 \* indicates novel sclerites not yet described in this species.

### 231 232 **3.2. Description of the complete mtDNA genome of *Holothuria polii***

233 The complete mitochondrial genome of *H. polii* was obtained from whole DNA sequencing  
234 raw data (38,340 read pairs). Breadth and depth of coverage were 100% and 480X, respectively, with  
235 an average inner distance of 250 bp, as expected. No polymorphisms were detected after variant  
236 calling analysis, excluding the presence of heteroplasmy. Table S1 shows the complete annotation  
237 and organization of the *H. polii* genome. This mitogenome consisted of 15,907 bp and included 22  
238 tRNA genes, 13 protein-coding genes, 2 rRNA genes and a putative control region (D-Loop). The  
239 light strand encoded 5 tRNAs and the *nad6* gene, whereas the heavy strand encodes 17 tRNAs, 12  
240 protein-coding genes, the 2 rRNAs and the longest unassigned non-coding region of 566 bp (Figure  
241 2). Given that the second longest unassigned region was 26 bp long, the 566 bp unassigned sequence  
242 was considered the putative control region, which had an A-T content of 63.8% (the highest value  
243 over the whole mtDNA sequence). This mtDNA genome had an overall A-T content equal to 58.3%.  
244 The nucleotide frequencies were: A=30.9%, T=27.4, C=25.6% and G=16.1%. The coding region  
245 consisted of 3,775 amino acids with a total of 3,786 codons. The most frequently used amino acids  
246 were leucin (16.6%) and isoleucin (9.9%), whereas the most frequently used codon was ATA (Ile)  
247 with a frequency of 5.55%, followed by CTA (Leu; 5.49%).

248 All 13 protein-coding genes started with the ATG codon (Met). Nine genes terminated with the  
249 TAA codon, 2 genes terminated with TAG and 2 genes had incomplete stop codons (T--) which are  
250 quite common in animal mitochondrial genes (Ojala et al., 1981). No rearrangements were found in  
251 this species and the gene order was the same as that of the mtDNA genome of *Apostichopus japonicus*,  
252 *Parastichopus californicus*, *P. nigripunctatus*, *P. parvimensis*, *H. scabra*, *H. forskali* and also with

253 that of the animal model *Strongylocentrotus purpuratus* (Echinoidea). The strand position of the  
 254 genes was shared among all these species (Figure S1).



255

256

257 **Figure 2.** Circular visualization and organization of the complete *Holothuria polii* mtDNA. External genes of the circle  
 258 are encoded by the positive strand (5'→3') and internal genes are encoded by the negative strand (3'→5'). The specimen  
 259 picture is reported inside the circle.

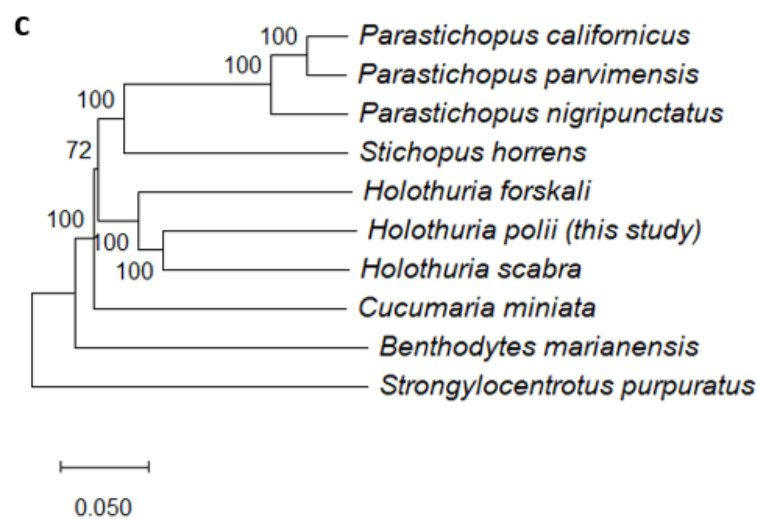
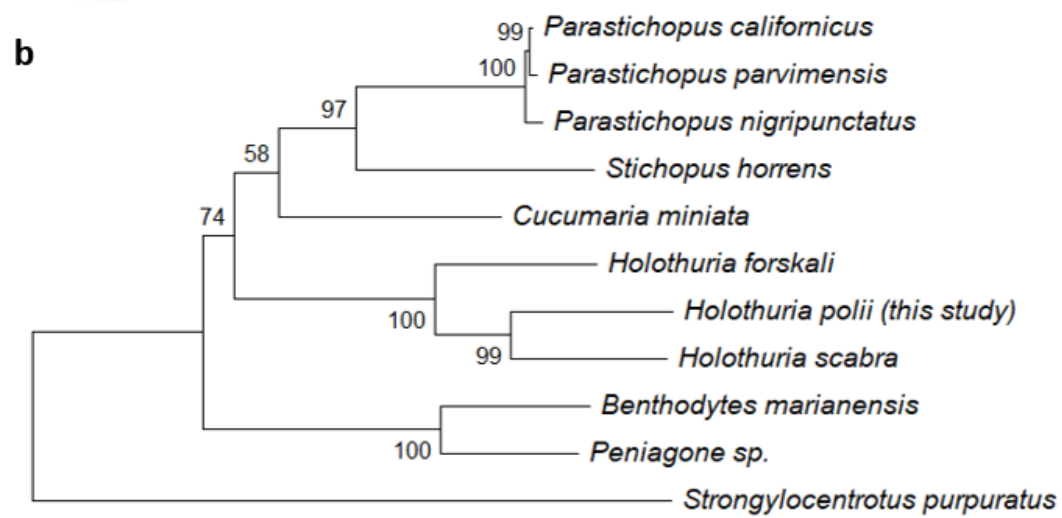
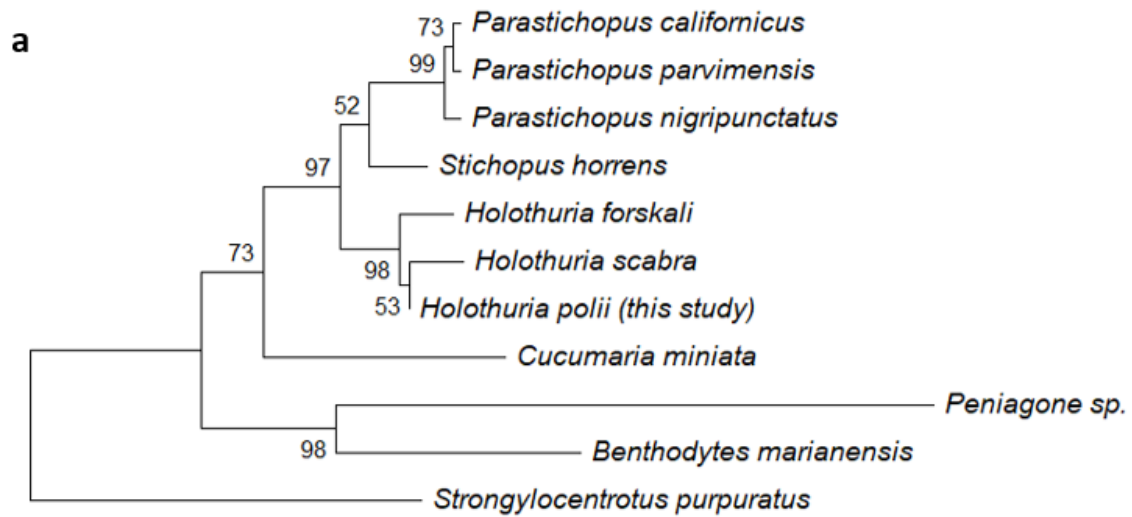
260

261 **3.3. Molecular identification and phylogenetic analyses**

262 The molecular identification of the species based on the selected portions of *cox1* gene using  
263 BLASTn analysis showed a range from 98% to 100% of identity with *cox1* of *H. polii* sequences  
264 already deposited in NCBI, whereas the second higher identity (90%) was obtained for *H. tubulosa*  
265 (GenBank: KJ719549.1). The BLASTn analysis with the portion of the 16S rRNA sequence obtained  
266 similar results (identity ranged from 98% to 100% with *H. polii* sequences and was 94% with the  
267 second closest species). These results confirmed that these two genes (or some portions of them) of  
268 the *H. polii* mtDNA genome are informative in terms of species identification (Uthicke et al., 2010;  
269 Kerr et al., 2005). Table S3 shows a comparison between the main ossicle types and *cox1* and 16S  
270 molecular markers; in particular, the main rod, button and table morphology have been compared  
271 together with the results of BLASTn analysis among our *H. polii* specimen, *H. polii* from other  
272 studies, *H. tubulosa* and *H. scabra* (Moussa and Wirawati 2018; Aydin and Erkan 2015; Massin et  
273 al., 2000).

274 Regarding the phylogenetic analyses, in order to obtain a robust phylogenetic evaluation, a  
275 *maximum-likelihood* phylogenetic tree was constructed based on the entire aminoacidic sequence of  
276 the *cox1* gene of 10 species belonging to 4 orders of the Holothuroidea class for which the whole  
277 mtDNA was available in GenBank (Table S2). Parallely, another ML tree was obtained based on the  
278 sequences of 16S rRNA using the same dataset and in both cases the outgroup *Strongylocentrotus*  
279 *purpuratus* was chosen. The phylogenetic trees are reported in Figure 3.

280 *H. polii cox1* and 16S rRNA sequence obtained in this study grouped with the two extant  
281 mtDNA of the genus with a high bootstrap value showing a strong reliability of the node.



283 **Figure 3.** a. Maximum Likelihood (lnL -2,753.67) phylogenetic tree obtained using the *cox1* aminoacidic sequence of  
284 the available mtDNA of different holothurian species, rooted with the outgroup *Strongylocentrotus purpuratus*. The  
285 bootstrap test values (1,000 replicates) are shown next to the branches.

286 b. Maximum Likelihood (lnL -10,936.53) phylogenetic tree obtained using the 16S rRNA sequence of the available  
287 mtDNA of different holothurian species, rooted with the outgroup *Strongylocentrotus purpuratus*. The bootstrap test  
288 values (1,000 replicates) are shown next to the branches.

289 c. Neighbour-Joining phylogenetic tree obtained with complete mtDNA genome sequences of holothurian species.  
290 The bootstrap test values (1,000 replicates) are shown next to the branches.

291

292 Generally, with the exception of *Cucumaria miniata* position in the 16S tree, all the  
293 phylogenetic trees show similar topologies. The NJ phylogenetic tree based on the complete  
294 sequences of holothurians confirmed the topology obtained with the ML tree based on aminoacidic  
295 *cox1* sequences: *H. polii* clustered with *H. scabra* and together they formed a clade with *H. forskali*.  
296 The three *Parastichopus* species clustered together while *Benthodytes marianensis* was the closest  
297 species to the outgroup, the echinoid *Strongylocentrotus purpuratus* (Figure 3).

298

## 299 **Discussion**

300 The main informative morphological features used to identify holothurian species are the  
301 ossicles, that are calcite structures inserted into the integument, which provide the typical rigidity of  
302 the body of the animal (Cutress et al., 1996). It is important to highlight that the whole intraspecific  
303 diversity of the shape and of the surface of the ossicles, as well as their position on the different body  
304 parts is still largely unknown for most holothurian species, including *H. polii* (Borrero-Pèrez et al.,  
305 2009). *Holothuria polii* is described to have different types of ossicles (buttons, tables, rods and  
306 others) with a smooth surface (Tortonese, 1965; Koehler, 1924). Despite the predominance of smooth  
307 ossicles in the analysed specimen, the presence of some knobbed and rugose buttons could have  
308 potentially led to an uncorrected species assignment. These types of ossicles are characteristics of *H.*  
309 *tubulosa*, which is described to have only rugose ossicles, in addition to tables similar to that of *H.*

310 *polii*. It is still unclear if hybridization events could occur in sympatric species (like *H. polii* and *H.*  
311 *tubulosa*), and if mixed ossicle types (and in which proportion and body positions) could be present.  
312 The ossicle morphological phenotype could be misleading as already observed by other authors  
313 (Moussa and Wirawati, 2018). The broad spectrum of ossicle shapes described in this study using  
314 the SEM images included types that, to our knowledge, were never detected before in *H. polii*. These  
315 observations might suggest that the entire variability of this morphological trait is not completely  
316 known and further investigations are needed also to evaluate the potential confounding hybridization  
317 events.

318 Despite these not completely clarified questions, all other morphological elements, in addition  
319 to molecular evidences based on mtDNA support the assignment of the analysed specimen to *H. polii*.  
320 Molecular analyses of *cox1* and 16S sequences were concordant to unambiguously assign the  
321 specimen to the *H. polii* suggesting that molecular data may be more robust than morphological  
322 descriptors, as already noted by others (Miller et al., 2017). Moreover, in some contexts it is not  
323 possible to analyse ossicles (for instance in highly processed sea cucumber food or in environmental  
324 studies).

325 The complete mitochondrial genome of *H. polii* shows an A-T content in line with the other  
326 holothurian mitogenomes analysed. The comparison of the gene order showed that *H. polii* shares the  
327 gene position on mtDNA with that of all other holothuroids (with the exception of *Stichopus horrens*,  
328 *Cucumaria miniata*, *Benthodytes marianensis*) and also with that of sea urchin *Strongylocentrotus*  
329 *purpuratus*, suggesting that this may represent the ancestral condition. Phylogenetic analyses  
330 confirmed that the represented four orders are clearly supported by robust nodes corroborating the  
331 phylogenetic position of *H. polii* closer to *H. scabra* in respect of *H. forskali*. In particular, these trees  
332 highlighted a distinct clade of abyssal sea cucumbers *Peniagone sp.* and *Benthodytes marianensis*  
333 (Elasipodida) while the Synallactida, Holothuriida and Dendrochirota orders form a cluster together  
334 confirming the Pneumonophora as sister group to Elasipodida (Miller et al., 2017). The *Cucumaria*  
335 *miniata* different position in the 16S rRNA tree in respect of *cox1* and complete mtDNA trees can be

336 explained with the less suitability of mitochondrial rRNA genes for species delimitation in DNA  
337 barcoding and for the low gap between intra and interspecific divergences in these portions in respect  
338 of coding genes (Nijman and Aliabadian, 2010; Vences et al., 2005).

339 *H. polii* plays an important ecological role for the Mediterranean marine environment. This  
340 species is subject to massive illegal overfishing and extensive over utilization (Friedman et al., 2011;  
341 Purcell et al., 2013; 2014). For these reasons, it is important to develop and validate methods and  
342 tools that could be used for species identification and to monitor the distribution and biodiversity of  
343 *H. polii* and related species. Therefore, molecular tools based on mtDNA sequences could become  
344 very relevant for these purposes, in addition to the use for the authentication of food products based  
345 on holothurian species. The mtDNA information we provided for *H. polii* filled an important gap in  
346 this context.

347

348

## 349 **Conclusions**

350 A more extensive description of ossicle variability is still needed to obtain a better description  
351 of holothurian species. It appeared however clear that ossicle data should be coupled with molecular  
352 data to confirm species identification as we did in our study. The mtDNA sequence can be used for  
353 species identification which might be relevant in particular to monitor the trades of holothurians for  
354 the extensive use of these species in the food luxury market. The complete mtDNA sequence of *H.*  
355 *polii* will be also useful as a starting point to monitor genetic diversity of this species and evaluate  
356 the success of conservation programmes of the marine ecological communities that are highly  
357 dependent by sea cucumbers.

358

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362 **Conflict of interest**

363 The authors declare that there is no conflict of interests regarding the publication of this article.

364

365 **References**

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555 **Supplementary materials:**

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557 **Table S1.** Functional annotation of the complete mitochondrial DNA of *Holothuria (Roweothuria) polii*.

Gene	Start	End	Strand	Lenght (bp)	tRNA codon	Intergenic nucleotides	Start codon	Stop codon	Aminoacids
cox1	1	1557	H	1557		0	ATG	TAG	518
trnR	1566	1632	H	67	CGA	8			
nad4L	1633	1929	H	297		0	ATG	TAA	98
cox2	1930	2617	H	688		0	ATG	T—	229
trnK	2619	2682	H	64	AAG	1			
atp8	2683	2850	H	168		0	ATG	TAA	55
atp6	2844	3527	H	684		-7	ATG	TAA	227
cox3	3530	4312	H	783		2	ATG	TAA	260
trnS2	4311	4381	L	71	TCA	-2			
nad3	4400	4744	H	345		18	ATG	TAA	114
nad4	4748	6104	H	1357		3	ATG	T—	452
trnH	6106	6172	H	67	CAC	1			
trnS1	6174	6241	H	68	AGC	1			
nad5	6242	8074	H	1833		0	ATG	TAG	610
nad6	8092	8580	L	489		17	ATG	TAA	162
cytb	8589	9731	H	1143		8	ATG	TAA	380
trnF	9733	9803	H	71	TTC	1			
rrnS	9804	10632	H	829		0			
trnE	10633	10701	H	69	GAA	0			
trnT	10703	10772	H	70	ACA	1			
Putative Control Region	10773	11338	H	566		0			
trnP	11339	11407	H	69	CCA	0			
trnQ	11404	11473	L	70	CAA	-4			
trnN	11476	11545	H	70	AAC	2			
trnL1	11547	11618	H	72	CTA	1			
trnA	11618	11684	L	67	GCA	-1			
trnW	11685	11752	H	68	TGA	0			
trnC	11753	11816	H	64	TGC	0			
trnV	11820	11889	L	70	GTA	3			
trnM	11916	11985	H	70	ATG	26			
trnD	11987	12058	L	72	GAC	1			
trnY	12059	12123	H	65	TAC	0			
trnG	12126	12195	H	69	GGA	2			
trnL2	12195	12265	H	71	TTA	-1			
nad1	12266	13237	H	972		0	ATG	TAA	323
trnI	13251	13318	H	68	ATC	13			
nad2	13319	14362	H	1044		0	ATG	TAA	347
rrnL	14363	15907	H	1545		0			













558 **Table S2.** List of the Holothuroidea used for phylogenetic analyses available in GenBank including *Holothuria polii*  
 559 obtained in this study.

Species	Accession number	mtDNA Length (bp)
<i>Benthodytes marianensis</i>	MH208310	17,567
<i>Holothuria forskali</i>	NC_013884	15,841
<i>Holothuria scabra</i>	NC_027086	15,779
<i>Holothuria polii</i> (this study)	NC_045029	15,907
<i>Cucumaria miniata</i>	NC_005929	17,538
<i>Parastichopus nigripunctatus</i>	NC_013432	16,112
<i>Parastichopus californicus</i>	NC_026727	16,727
<i>Parastichopus parvimensis</i>	NC_029699	16,120
<i>Peniagone sp.</i> *	KF915304	15,507
<i>Stichopus horrens</i>	NC_014454	16,257

560 \* This species has not been used for the complete mtDNA NJ tree.

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562 **Table S3.** Comparison between the main morphological and molecular markers used for *Holothuria polii* species  
 563 identification. The percentages represent the identity and the coverage of the first more similar result and the related  
 564 accession number in BLASTn for each species. Images of the holothurians different from our specimen derive from  
 565 Moussa and Wirawati 2018; Aydin and Erkan 2015; Massin et al. 2000.  
 566

	Buttons	Tables	Rods	<i>cox1</i> (% BLASTn identity; % coverage, accession number)	16S (% BLASTn identity; % coverage, accession number)
<i>Holothuria polii</i> this study				100% LR694133	100% LR694133
<i>Holothuria polii</i> other studies				99.76%; 100% JN207607	99.77%; 99% LC176660
<i>Holothuria tubulosa</i>				89.95%; 96% KJ719549	93.24%; 100% GU797633
<i>Holothuria scabra</i>				84.69%; 98% GQ920784	86.94%; 100% JQ657264

567 Moussa, R., Wirawati, I. 2018. Observations on some biological characteristics of *Holothuria polii* and *Holothuria*  
 568 *sanctori* from Mediterranean Egypt. Int. J. Fish. Aquat. Stud., 6 (3), 351-357.  
 569 Aydin, M., Erkan, S., 2015. Identification and some biological characteristics of commercial sea cucumber in the Turkey  
 570 coast waters. Int. J. Fish. Aquat. Stud., 3 (1), 260-265.  
 571 Massin, C., Mercier, A., Hamel, J. F., 2000. Ossicle change in *Holothuria scabra* with a discussion of ossicle evolution  
 572 within the Holothuriidae (Echinodermata). Acta Zool., 81 (1), 77-91. doi: 10.1046/j.1463-6395.2000.00039.x.

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**Figure S1.** Gene order comparison between the outgroup *Strongylocentrotus purpuratus* (Echinoidea) and the holothurians with the mtDNA available in NCBI. The underlined genes are encoded by the internal strand.

*Strongylocentrotus purpuratus*

nad1	trnI	nad2	rrnL	cox1	trnR	nad4L	cox2	trnK	atp8	atp6	cox3	<u>trnS2</u>	nad3	nad4	trnH	trnS1	nad5	<u>nad6</u>	cob	trnF	rrnS	trnE	trnT	trnP	<u>trnQ</u>	trnN	trnL1	<u>trnA</u>	trnW	trnC	<u>trnV</u>	trnM	<u>trnD</u>	trnY	trnG	trnL2
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*Parastichopus parvimensis*, *Parastichopus nigripunctatus*, *Parastichopus californicus*, *Apostichopus japonicus*, *Holothuria forskali*, *Holothuria scabra*, *Holothuria polii* (this study)

nad1	trnI	nad2	rrnL	cox1	trnR	nad4L	cox2	trnK	atp8	atp6	cox3	<u>trnS2</u>	nad3	nad4	trnH	trnS1	nad5	<u>nad6</u>	cob	trnF	rrnS	trnE	trnT	trnP	<u>trnQ</u>	trnN	trnL1	<u>trnA</u>	trnW	trnC	<u>trnV</u>	trnM	<u>trnD</u>	trnY	trnG	trnL2
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*Stichopus horrens*

nad1	trnI	nad2	rrnL	cox1	trnR	nad4L	cox2	trnK	atp8	atp6	cox3	<u>trnS2</u>	nad3	nad4	trnH	trnS1	nad5	<u>nad6</u>	cob	trnF	rrnS	trnE	trnT	trnP	<u>trnQ</u>	trnN	trnL1	<u>trnA</u>	trnW	trnC	<u>trnV</u>	<u>trnD</u>	trnY	trnG	trnM	trnL2
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*Cucumaria miniata*

nad1	trnI	nad2	rrnL	cox1	trnR	trnE	trnP	trnN	trnL1	trnW	<u>trnV</u>	nad4L	cox2	trnK	atp8	atp6	cox3	<u>trnS2</u>	nad3	nad4	trnH	trnS1	nad5	<u>nad6</u>	cob	trnF	rrnS	trnT	<u>trnQ</u>	<u>trnA</u>	trnC	trnM	<u>trnD</u>	trnY	trnG	trnL2
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*Benthodytes marianensis*

nad1	trnI	nad2	trnT	rrnL	cox1	trnR	nad4L	cox2	trnK	atp8	atp6	cox3	<u>trnS2</u>	nad3	nad4	trnH	trnS1	nad5	<u>nad6</u>	cob	trnF	rrnS	trnE	<u>trnA</u>	trnW	trnM	trnG	trnP	<u>trnQ</u>	trnN	trnL1	trnC	<u>trnV</u>	<u>trnD</u>	trnY	trnL2
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