

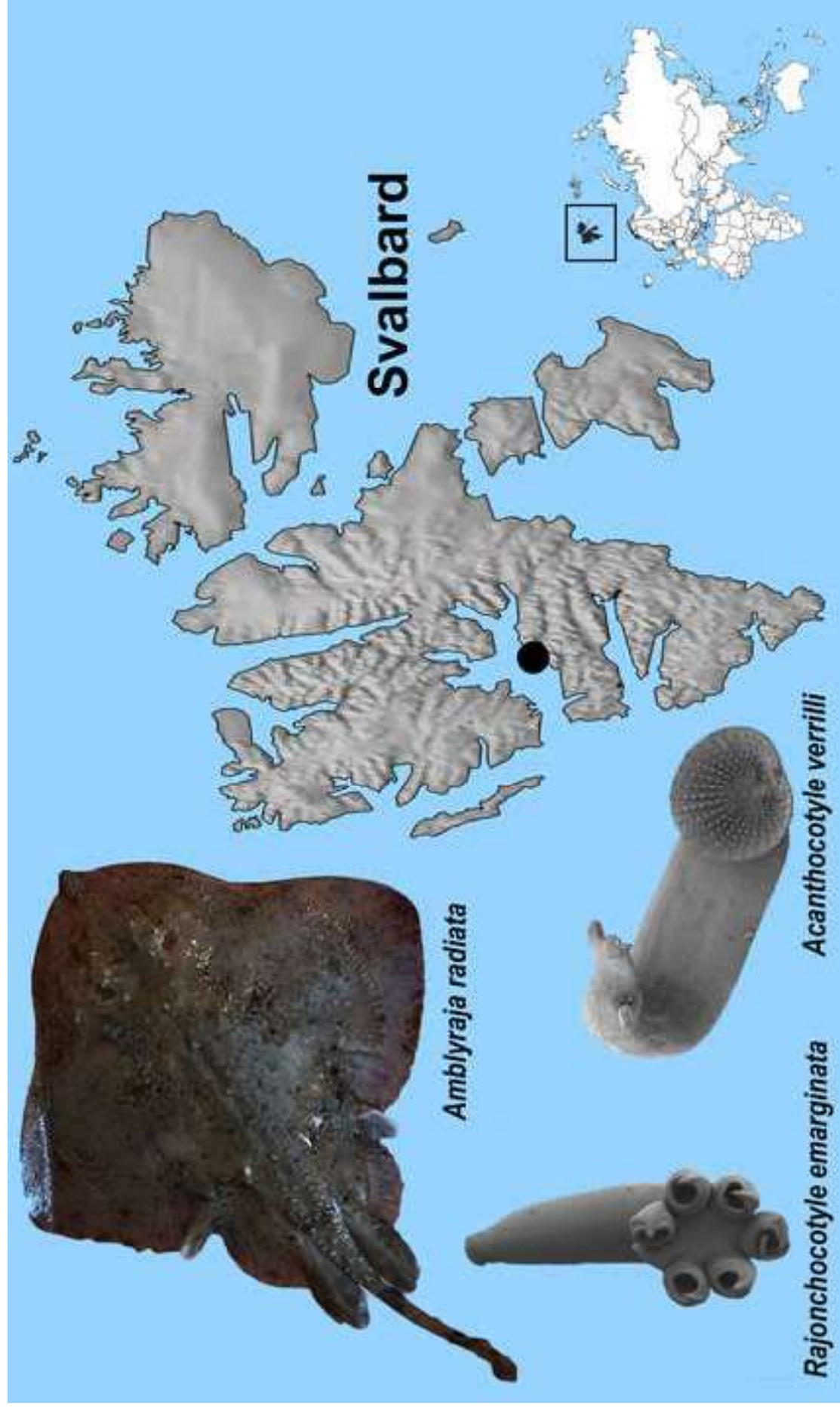
Parasitic flatworms infecting thorny skate, *Amblyraja radiata*: Infection by the monogeneans *Acanthocotyle verrilli* and *Rajonchocotyle emarginata* in Svalbard

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1 **Parasitic flatworms infecting thorny skate, *Amblyraja radiata*:**  
2 **infection by the monogeneans *Acanthocotyle verrilli* and**  
3 ***Rajonchocotyle emarginata* in Svalbard**

4  
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18 **Abstract**

19 Parasite diversity above the Arctic circle remains understudied even for commercially  
20 valuable host taxa. Thorny skate, *Amblyraja radiata*, is a common bycatch species with a  
21 growing commercial value. Its natural range covers both sides of the North Atlantic including  
22 the Arctic zone. Svalbard is a Norwegian archipelago located on the northwest corner of the  
23 Barents Shelf which sustains a spectacular species diversity. So far, several monogenean  
24 species have been reported infecting thorny skate across the Atlantic Ocean. In the present  
25 study, we intend to fill in the knowledge gap on monogenean parasites infecting thorny  
26 skate in the northern part of its range and thus indirectly assess the connectivity between  
27 the thorny skate populations off the Svalbard coast and from previously studied locations.  
28 46 monogenean individuals were recovered from 11 specimens of thorny skate. Following  
29 morphological and molecular assessment, two species of monogeneans, *Acanthocotyle*  
30 *verrilli* and *Rajonchocotyle emarginata*, were identified. The results serve as the  
31 northernmost record for both parasite genera and the first record of monogenean species  
32 off Svalbard. Detailed morphometric evaluation revealed a relatively high level of  
33 morphological variation in *A. verrilli* compared to its congeners. Phylogenetic reconstruction  
34 placed *A. verrilli* in a well-supported clade with *A. imo*. Our study also suggests high  
35 diagnostic significance of sclerotised structures in the identification of *Rajonchocotyle*. Even  
36 though the occurrence of two directly transmitted parasite species supports the previously  
37 suggested long-distance migration of *A. radiata*, future studies employing highly variable  
38 genetic markers are needed to assess the ongoing and historical migration patterns.

39

40 **Highlights**

- 41 • First record of monogenean species in Svalbard
- 42 • Northernmost record for representatives of Acanthocotyliidae and Hexabothriidae
- 43 • Transatlantic occurrence of parasites supports connectivity of thorny skate  
44 populations

45 **Keywords:** Rajidae, Acanthocotyliidae, Hexabothriidae, first monogenean in Svalbard

46 **1. Introduction**

47 Thorny skate (Chondrichthyes, Rajidae) is a common bycatch species with growing  
48 commercial value. It prefers shallow coastal waters with muddy or sandy substrate [1] and  
49 temperatures from -1.4°C to 16°C [2]. Increased fishing effort during the last decades

50 severely impacted the overall biomass of this skate species characterized by low fecundity,  
51 slow growth rate, and late maturity [3,4]. Its known geographic distribution ranges from  
52 South Carolina in the Western part of the Atlantic Ocean to Greenland and the North Sea in  
53 the East and Svalbard in the Arctic zone [5–11].

54

55 Svalbard is an archipelago in the Arctic Ocean located on the northwest corner of the  
56 Barents Shelf. The Arctic Ocean is the smallest of all oceans with a mean depth of 1361 m  
57 and a total area of approximately 10 million km<sup>2</sup> [12,13]. It consists of four abyssal plains  
58 surrounded by continental shelves comprising c. 50% of the total area [13]. The northern  
59 and western margins of the Barents Shelf end in the continental slope down to the Polar  
60 Ocean Basin and the oceanic Norwegian Greenland Sea, respectively [14]. The wide range of  
61 habitats on continental shelves sustains a spectacular biodiversity in this marine ecosystem  
62 [15] and harbours species of Atlantic and Pacific affinities due to ancient connections.  
63 However, continental shelves were an important migration barrier especially to shallow-  
64 water organisms [16] between the Arctic and adjacent oceans [17] during the last  
65 Pleistocene glacial/interglacial cycles. Overall, the inventory of biodiversity on Svalbard is far  
66 from complete because of the focus of most studies on its west coast in view of the better  
67 accessibility of this region. Similar to terrestrial habitats, the inventory of marine biodiversity  
68 off Svalbard's coast has been limited and biased by sampling techniques (e.g., pelagic trawls)  
69 or towards certain taxonomic groups such as Crustacea and Mollusca [18,19].

70

71 ~~In general, parasite biodiversity in the Arctic is mostly understudied and many species~~  
72 ~~remain unknown including fish parasites [20–23]. Although the known distribution of~~  
73 ~~*Amblyraja radiata* covers large parts of the North Atlantic and Arctic region under eternal ice,~~  
74 ~~data about its parasite fauna from those areas are missing. Data about parasite fauna of~~  
75 ~~*Amblyraja radiata* from the North Atlantic and Arctic regions of its distribution are missing.~~  
76 ~~In general, parasite biodiversity in the Arctic is mostly understudied and many species~~  
77 ~~remain unknown including fish parasites [20–23].~~ Scientific exploration of the marine  
78 parasite fauna in this part of the world has been mainly concentrated on Franz Josef Land  
79 and heteroxenous parasite taxa [22]. Recently, [Murzina et al. \[24\]](#) ~~Murzina et al. (2019)~~  
80 reported on the parasite fauna of *Leptoclinus maculatus* (Perciformes, Stichaeidae) (Fries  
81 1838), being the first record of parasitic flatworms ([Trematoda](#)) at the Svalbard coast. [↗](#)

82 ~~total~~[Globally](#), there are almost 1500 parasite species described from 900 elasmobranch  
83 species to date [25]. Helminth infections of the thorny skate were reported worldwide (see  
84 Table 1). So far, representatives of two monogenean families and three species have been  
85 reported. Monogenea is a group of parasitic flatworms (Neodermata, Platyhelminthes) with  
86 worldwide occurrence and a mostly ectoparasitic life-style. They are primarily parasites of  
87 fish characterised by a direct life-cycle and predominantly narrow host-specificity [26,27].  
88 The basic division is between members of blood feeding Polyopisthocotylea and epithelial  
89 feeding Monopisthocotylea, representatives of both of which have been reported to infect  
90 the thorny skate [28,29]. The northernmost distribution of a parasitic flatworm infecting *A.*  
91 *radiata* was *Acanthocotyle verilli* Gotto, 1899 recorded off Tromsø, Norway [30]  
92 (~~Poddubnaya et al., 2016~~, see Table 1). Hence, flatworm infections on *A. radiata* have never  
93 been recorded above the Arctic Circle.

94

95 In the present study, we intend to fill in this gap of knowledge in the distribution of  
96 monogenean parasites infecting thorny skate, *A. radiata*, at the northern part of its known  
97 range, in Spitsbergen, Svalbard.

98

## 99 **2. Material and Methods**

100

### 101 2.1. [Data-Host](#) collection

102 In total, eleven specimens of *A. radiata* were examined for the presence of monogenean  
103 individuals during a field expedition in Spitsbergen, Svalbard, July 2016 organised by the  
104 Centre for Polar Ecology (University of South Bohemia, Czech Republic). Host specimens  
105 were caught in the Adventfjorden near Hotellneset, Spitsbergen, Svalbard (78°15'18"N,  
106 15°30'58"E) using benthic gill nets at a depth of 30–40 m and immediately transported to  
107 the laboratory in seawater containers. Prior to dissection and subsequent examination,  
108 skates were euthanised by overdosing with tricaine methane sulphonate (Sigma-Aldrich,  
109 Darmstadt, Germany).

110

### 111 2.2. Parasite collection and morphological examination

112 The fins, gills and nasal cavity were examined for the presence of monogeneans.  
113 Monogenean individuals were transferred with a needle and mounted on slides using a

114 solution of glycerine ammonium picrate (GAP). Selected specimens were kept in 99%  
115 ethanol and subsequently stained using acetocarmine combined with Gomori trichrome,  
116 cleared with clove oil and mounted in Canada balsam. Two species were found in this study,  
117 *Acanthocotyle verilli* and *Rajonchocotyle emarginata*. Infection parameters per parasite  
118 species namely prevalence (percentage of infected hosts), infection intensity (mean number  
119 of monogenean individuals per infected host) and abundance (mean number of  
120 monogenean individuals per all-examined host) were calculated following Ergens and Lom  
121 [31]. In total, 13 and 21 morphological characters including hard and soft parts following  
122 Kearn et al. [32] ~~Kearn et al. (2016)~~ and Bullard and Dippenaar (2003), respectively, were  
123 measured and photographed using a Leica DM 2500 LED microscope (Leica Microsystems,  
124 Wetzlar, Germany) and the software LasX v3.6.0. Voucher specimens are deposited in the  
125 collection of the Research Group Zoology: Biodiversity and Toxicology at Hasselt University  
126 in Diepenbeek, Belgium (HU) under the following accession numbers: XIII.4.38-50, XIV.1.01-  
127 16xx-xx. Type specimens from the Helminthological Collection of the South Australian  
128 Museum, South Australia, Australia (AHC) and the National Museum of Natural History of  
129 the Smithsonian Institution, Washington, USA (USNM), were examined for comparative  
130 purposes: *Acanthocotyle atacamensis* Ñacari, Sepulveda, Escribano & Oliva, 2019 – 1  
131 paratype (1 slide) USNM 1480281; *A. gurgesiella* Ñacari, Sepulveda, Escribano & Oliva, 2018  
132 – 1 paratype (1 slide) USNM 1422089; *A. imo* Ñacari, Sepulveda, Escribano & Oliva, 2019 – 1  
133 paratype (1 slide) USNM 1480278; *A. lobianchi* Monticelli, 1888 - 2 vouchers (2 slides) AHC  
134 36231, 36232; *A. pacifica* Bonham & Guberlet, 1938 – 2 paratypes (2 slides) USNM 1321942;  
135 *A. pugetensis* Bonham & Guberlet, 1938 – 2 paratypes (2 slides) USNM 1321940; *A. verrilli* –  
136 1 paratype (1 slide) USNM 135051; *A. urolophi* Kearn, Whittington, Chisholm & Evans-  
137 Gowing, 2016 - 4 paratypes (4 slides) AHC 36222, 36223, 36224, 36225; *Branchotentes*  
138 *octohamatus* Glennon, Chisholm & Whittington, 2005 – 2 paratypes (2 slides) AHC28769,  
139 28770; *B. robinoverstreeti* Bullard & Dippenaar, 2003 – 1 holotype (1 slide) USNM 1387687;  
140 *Callorhynchocotyle callorhynchi* (Manter, 1955) – 2 paratype (2 slides) AHC29747; *C.*  
141 *amatoi* Boeger, Kritsky & Pereira, 1989 – 1 paratype (2 slides) AHC29749; *Eropcotyle*  
142 *antarctica* (Hughes, 1928) – 2 paratypes (2 slides) AHC29725; *E. somniosi* – 4 paratype (4  
143 slides) USNM 1349221; *Heteronchocotyle gymnurae* Neifar, Euzet & Ben Hassine, 2001 – 1  
144 paratype (1 slide) USNM 1385030; *Paraheteronchocotyle amazonensis* Mayes, Brooks &  
145 Thorson, 1981 – 2 paratypes (2 slides) USNM 1372658; *Rajonchocotyle emarginata* Olsson,

146 1876 – 1 paratype (1 slide) USNM 1337399; *R. laevis* Price, 1942 – 1 syntype (1 slide)  
147 USNM1337422; *R. wehri* Price, 1942 – 4 paratypes (4 slides) USNM 1337421;  
148 *Squalonchocotyle borealis* (Van Beneden, 1853) Cerfontaine, 1899 – 4 paratypes (4 slides)  
149 USNM 1349221; *S. callorhynchi* – 1 paratype, 1 holotype (2 slides) USNM 1338129; *S. impristi*  
150 – 1 holotype (1 slide) USNM 1338749. Selected specimens of both collected species were  
151 drawn using a drawing tube and afterwards edited using the software GIMP v2.10.20.  
152 Interspecific morphological differences and the level of intraspecific phenotypic variability  
153 were evaluated using measurements relative to the total length of the parasite's body  
154 because of the correlation between morphological variables and the total length [34].  
155 Morphometric parameters were subsequently analysed by principal component analysis  
156 (PCA) in the R software package stats [35] and visualised using ggplot2 [36]. Only specimens  
157 mounted on slides with GAP were part of the analyses to avoid effects of the staining  
158 method on the results [37]. The following variables were used in PCA: (1) the ratios body  
159 width/total length (TL), (2) body length/TL, (3) pharynx length/TL, (4) pharynx width/TL, (5)  
160 diameter of the pseudohaptor/TL, (6) number of sclerite rows/TL, (7) testes maximum  
161 width/TL, (8) germarium length/TL and (9) germarium width/TL. Raw measurements are  
162 provided as Supplementary material (see Table S1&S2).

163

### 164 2.3. Scanning electron microscopy (SEM)

165 Following host examination, live monogenean individuals were fixed with hot 4% neutral  
166 buffered formaldehyde solution and transported to the Parasitology Laboratory of [the](#)  
167 Centre for Polar Ecology in České Budějovice, Czech Republic. Subsequently, samples were  
168 washed in 0.1 M phosphate buffer solution (three times for 15 minutes), post-fixed in 2%  
169 osmium tetroxide solution (for 60 minutes), washed again and dehydrated with an  
170 ascending acetone series (30%, 50%, 70%, 80%, 90%, 95%, 100%), each step for 15 minutes.  
171 Following dehydration, monogenean specimens were dried in liquid CO<sub>2</sub> using a critical point  
172 method, placed on metal targets using double-stick tape, gold coated in a BAL-TEC SCD 050  
173 sputter coater (Bal-Tec, Los Angeles, USA) and observed using a SEM JEOL JSM-7401F  
174 scanning electron microscope (JEOL, Tokio, Japan) at the Laboratory of Electron Microscopy,  
175 Institute of Parasitology, Biology Centre CAS in České Budějovice.

176

### 177 2.4. Molecular data generation



178 The posterior part of the body or complete specimens were used for genomic DNA isolation.  
179 The total genomic DNA was extracted using the Qiagen Blood and Tissue Isolation Kit  
180 (Qiagen, Hilden, Germany) following the manufacturer's instructions. A portion of the large  
181 ribosomal subunit (28S rRNA) gene was amplified using the primer combination C1 (5'-  
182 ACCCGCTGAATTTAAGCAT-3') and D2 (5'-TGGTCCGTGTTTCAAGAC-3') [38]. Each PCR reaction  
183 contained 1.5 unit of *Taq* Polymerase (ThermoFisher Scientific, Waltham, USA), 1X buffer  
184 containing 0.1 mg/ml bovine serum albumin, 1.5 mM MgCl<sub>2</sub>, 200 mM dNTPs, 0.5 mM of each  
185 primer and 50 ng of genomic DNA in a total reaction volume of 30 µl under the following  
186 conditions: initial denaturation at 95 °C for 2 min followed by 39 cycles of 94 °C for 20 s,  
187 annealing at 58 °C for 30 s, extension at 72 °C for 1 min and 30 s, and a final extension step  
188 at 72 °C for 10 min. PCR products were purified using ExoSAP-IT (ThermoFisher Scientific,  
189 Waltham, USA) under the following conditions: 15 min at 37 °C and 15 min at 80 °C.  
190 Targeted DNA fragments were sequenced using the same primers as in the amplification  
191 reactions together with a Big Dye Chemistry Cycle Sequencing Kit v3.1 (ThermoFisher  
192 Scientific, Waltham, USA). Following clean up using the BigDye XTerminator Purification Kit  
193 (ThermoFisher Scientific, Waltham, USA), fragments were visualised on an ABI 3130 capillary  
194 sequencer (ThermoFisher Scientific, Waltham, USA). Electropherograms were visually  
195 inspected and assembled in MEGA7 [39]. The obtained sequences were deposited in NCBI  
196 GenBank under the accession numbers [MW260310-12](#)~~xx-xx~~.

197

## 198 2.5. Phylogeny

199 Phylogenetic placement of collected monogenean species was inferred based on 28S rDNA  
200 at family level. Sequences generated during this study were aligned using MUSCLE [40]  
201 under default distance measures as implemented in MEGA v7 [39], together with 28S rRNA  
202 gene sequences of other *Acanthocotyle* spp. Selected representatives of related  
203 monogenean families (Bothrirematidae, Anoplodiscidae and Udonellidae) were used as  
204 outgroup [41]. Genetic distances among the species of *Acanthocotyle* were calculated as the  
205 pairwise difference (uncorrected p-distance) in MEGA v7 [39]. Poorly aligned and overly  
206 divergent regions were trimmed using Gblocks v0.91b [42] under the less strict flanking  
207 position option and allowing gap positions within the blocks. The final alignment consisted of  
208 825 bp. The most appropriate evolutionary model, the HKY +  $\Gamma$  model [43], was selected  
209 based on the Bayesian information criterion in jModelTest v2 [44]~~and was used in all~~

210 ~~subsequent analyses~~. Phylogenetic relationships were inferred under Bayesian inference (BI  
211 in MrBayes v3.2.0 [45]) based on two independent runs (20<sup>5</sup> generations, sampled every  
212 1,000th generation and with a burn-in of 10%). Parameter convergence and run stationarity  
213 were assessed in Tracer v1.6 [46]. Moreover, a maximum likelihood (ML) search was  
214 performed in RAxML v8.2.12 with tree search conducted using RAxML's standard tree search  
215 algorithm and bootstrap support calculated using the option with an automated number of  
216 replicates to obtain stable support values under the frequency stopping criterion [47].  
217 Phylogenetic trees were edited in FigTree v1.4.2 ([http://tree.bio.ed.ac.uk/](http://tree.bio.ed.ac.uk/software/figtree) software/figtree).

### 218 **3. Results**

219 In total, 37 and 9 monogenean individuals were found on fins and gills, respectively.  
220 Based on a detailed morphological examination, two species of monogeneans were  
221 identified.

222  
223 To comply with the regulations set out in article 8.5 of the amended 2012 version of the  
224 International Code of Zoological Nomenclature [48], details of ~~the species *Acanthocotyle*~~  
225 ~~*verrilli* and *Rajonchocotyle ermaginata*~~ have been submitted to ZooBank based on their  
226 respective original descriptions. ~~The Life Science Identifier (LSID of the article is~~  
227 ~~urn:lsid:zooBank.org:pub:xxx).~~ For each taxon, the [Life Science Identifier \(LSID\)](#) is reported in  
228 the taxonomic summary.

#### 230 **3.1. *Acanthocotyle verrilli* Goto, 1899**

231 Monogenea van Beneden, 1858

232 **Family** Acanthocotylidae Monticelli, 1903

233 **Genus** *Acanthocotyle* Monticelli, 1888

234 **Type-host:** *Amblyraja radiata* (Donovan, 1808) (Chondrichthyes, Rajidae)

235 **Other hosts:** *Bathyraja spinicauda* (Jensen, 1914) (Chondrichthyes, Arhynchobatidae);

236 *Leucoraja erinacea* (Mitchill, 1825) (Chondrichthyes, Rajidae)

237 **Site on host:** Fins.

238 **Prevalence:** 72,7% (8 out of 11 infected)

239 **Intensity of infection:** 4,6 (1-12)

240 **Abundance:** 3, [437](#) (0-12).

241 **Type-locality:** Cape Cod, USA

242 **Other localities:** Adventfjorden, Spitsbergen; Chaleur Bay, Canada; Coast of Maine, USA; off  
243 Bergen, Norway; off Newfoundland, Canada; North-eastern Norwegian Sea

244 **ZooBank registration:** The LSID for ~~the new record of~~ *Acanthocotyle verrilli* is  
245 [urn:lsid:zoobank.org:act:FF7F506B-FB65-446C-9F91-](urn:lsid:zoobank.org:act:FF7F506B-FB65-446C-9F91-255C45BF2E86)  
246 [255C45BF2E86urn:lsid:zoobank.org:act:xxx](urn:lsid:zoobank.org:act:xxx).

247

#### 248 3.1.1. ~~Amended diagnosis~~ Remarks

249 In total, 13 morphological characters including soft body parts and sclerotised structures  
250 were measured (see Table 2, Figs. 1&2). Body elongate, circular pseudohaptor with radial  
251 rows of sclerites (26 – 34) covered by tegument (Fig. 2G). Pseudohaptor with 28 – 34 rows of  
252 sclerites, each row consists of 4 – 10 sclerites. Marginal valve of pseudohaptor with distinct  
253 fringe. The central part of the pseudohaptor, between the radial rows of sclerites, shows a  
254 depression indicating this part to be responsible for the attachment to the body (Fig 2E).  
255 True haptor with 16 marginal hooks located subterminally at the posterior margin of  
256 pseudohaptor (Fig. 2E). Hooks in the true haptor organised into one central pair and 14  
257 hooks in a peripheral row, the latter pointing centrally (Fig. 2G) with shafts free of  
258 tegument (Fig. 2H). Pharynx globular. Three anterior adhesive lobes on each side of head  
259 with sense organs located at the side of each internal lobe (Fig. 2B). Accessory glands at the  
260 level of pharynx. Excretory bladders on each side, anterior to vitellarium field. Eyes absent.  
261 Testes (26 – 47) mainly rounded usually arranged in several (2 – 3) rows. Seminal vesicle  
262 unlobed, anterior part communicates with male genital opening via curved ejaculatory duct  
263 immediately posterior to pharynx. Male accessory gland reservoirs 2, adjacent to ejaculatory  
264 duct. Common opening at the level of intestinal bifurcation. Penis sclerite absent.  
265 Germarium immediately anterior to testes. Small uterine receptaculum seminis adjacent to  
266 germarium. Vagina absent. Germinal appendix not observed. Uterine pore opening on right  
267 side of the body, at the level of posterior part of pharynx. Eggs attached externally by an  
268 abopercular appendage (Fig. 2C). The egg operculum shows a pointed end (Fig. 2D). Vitelline  
269 follicles not discrete with lobed and overlapping margins, extending from level of germarium  
270 to near the posterior end of body proper.

#### 271 3.1.2. Differential diagnosis

272 Based on the recent revision of Acanthocotylidae published by Kearn et al. [32] ~~-(2016)~~  
273 combined with new species descriptions of Ñacari et al. [49,50] ~~-(2018, 2019)~~, there are 12

274 species of *Acanthocotyle* currently considered valid: *Acanthocotyle atacamensis*;  
275 *Acanthocotyle elegans* Monticelli, 1890; *Acanthocotyle greeni* Macdonald & Llewellyn, 1980;  
276 *Acanthocotyle gurgesiella*; *Acanthocotyle imo*; *Acanthocotyle lobianchi*; *Acanthocotyle*  
277 *pacifica*; *Acanthocotyle patagonica* Kuznetsova, 1971; *Acanthocotyle pugetensis*;  
278 *Acanthocotyle urolophi*; *Acanthocotyle verrilli*; *Acanthocotyle williamsi* Price, 1938.  
279 *Acanthocotyle verrilli* most closely resembles *A. atacamensis*, *A. gurgesiella*, *A. imo* and *A.*  
280 *urolophi*. All these species have more than 20 testes, a haptor armed with 21-39 radial rows  
281 of sclerites, and a dextral opening of the uterine pore. *Acanthocotyle urolophi* is  
282 distinguished from the other species by the form of the vitelline follicles. Unlike in *A.*  
283 *atacamensis*, *A. gurgesiella*, *A. imo* and *A. verrilli*, the vitelline follicles of *A. urolophi* are  
284 discrete and easy to count. The number of testes ranges from 26 to 47 (mode 36) in *A.*  
285 *verrilli*, 40 to 58 (mode 50) in *A. atacamensis*, 28 to 43 (mode 30) in *A. gurgesiella*, 32 to 47  
286 (mode 41) in *A. imo* and 40 to 70 (mode 55) in *A. urolophi*. Testes of *A. verrilli* are organised  
287 in numerous rows compared to two rows in the case of *A. imo* and *A. gurgesiella*. Unlike in  
288 *A. atacamensis*, *A. imo* and *A. gurgesiella*, testes of *A. verrilli* have overlapping margins. The  
289 number of radial rows of sclerites range from 28 to 34 (mode 32) in *A. verrilli*, 28 (no  
290 variation) in *A. atacamensis*, 36 to 40 (mode 40) in *A. gurgesiella*, 30 to 35 (mode 32) in *A.*  
291 *imo* and 32 to 37 (mode 35) in *A. urolophi*. In specimens of *A. verrilli* possessing 28 radial  
292 rows of sclerites in the pseudohaptor, there are only 4–5 sclerites in the first row (counting  
293 from the position of the true haptor) compared to 6 in *A. atacamensis*. The difference  
294 between *A. verrilli* and *A. imo* is then visible in a smooth marginal valve of the pseudohaptor  
295 in *A. imo* compared to a distinct fringe in *A. verrilli*. *Acanthocotyle verrilli* can be  
296 distinguished from *A. gurgesiella* by the absence of an armed male genital aperture.

297

### 298 3.1.3. Interspecific differentiation based on multivariate statistics

299 Principal component analysis combining ~~qualitative and quantitative data~~ metric and meristic  
300 data (see Material & Methods) was performed to examine and visualise differences between  
301 three morphologically similar species of *Acanthocotyle* for which raw data are available (Fig.  
302 3A-C). Figure 3A shows the comparison of all three species. The first PC explained 64.5% and  
303 the second 12.3% of the variation in the dataset. In the resulting biplot, specimens of *A.*  
304 *verrilli* collected in this study are clearly distinguished from the other two species along the  
305 first axis and display more intraspecific variability. Figure 3B presents a PC biplot of *A. verrilli*

306 and *A. imo* and shows clear differentiation along the first axis (PC1 explained 54.5% and PC2  
307 17.3% of the variation in the dataset). Figure 3C presents a PCA biplot of *A. verrilli* and *A.*  
308 *atacamensis* and shows clear differentiation along the first axis (PC1 explained 67.1% and  
309 PC2 12.6% of the variation in the dataset). The diameter of the pseudohaptor and the total  
310 body length display the highest contribution to the separation in all datasets of all  
311 parameters-.

#### 313 3.1.4. Phylogenetic reconstruction

314 In total, three identical sequences of the 28S rDNA region from *A. verrilli* were generated in  
315 this study (Genbank accession numbers xx-xx). Phylogenetic reconstruction placed *A. verrilli*  
316 in a well-supported clade with *Acanthocotyle imo* (Fig. 4). Genetic interspecific differences  
317 between the species of *Acanthocotyle* with available 28S rDNA region sequences are  
318 presented in Table 3.

### 319 **3.2. *Rajonchocotyle emarginata* (Olsson, 1876)**

320 Monogenea van Beneden, 1858

321 **Family** Hexabothriidae Price, 1942

322 **Genus** *Rajonchocotyle* Cerfontaine, 1899

323 **Type-host:** *Amblyraja radiata* (Donovan, 1808) (Chondrichthyes, Rajidae)

324 **Other hosts:** *Bathyraja brachyurops* (Fowler, 1910); *Bathyraja magellanica* (Philippi, 1902)  
325 (Chondrichthyes, Arhynchobatidae); *Leucoraja naevus* (Müller & Henle, 1841); *Raja brachyura*  
326 Lafont, 1871; *Raja clavata* Linnaeus, 1758; *Raja microocellata* Montagu, 1818; *Raja*  
327 *montagui* Fowler, 1910; *Raja undulata* Lacepède, 1802 (Chondrichthyes, Rajidae) and  
328 *Psammobatis scobina* (Philippi, 1857) (Chondrichthyes, Arhynchobatidae)

329 **Site on host:** Gills.

330 **Prevalence:** 27,3% (3 out of 11 specimens infected)

331 **Intensity of infection:** 3 (1-6)

332 **Abundance:** 0,8 (0-6).

333 **Type-locality:** Bohuslän Coast, Sweden

334 **Other localities:** Adventfjorden, Spitsbergen; Cardigan Bay, Wales; Chaleur Bay, Canada;  
335 Galway Bay, Ireland; Marine Bahusiae, Scandinavia; Mediterranean Sea, Italy; off Plymouth,  
336 UK; off Roscoff, France; Northwest coast of Spain; Patagonian Shelf; off Tromsø, Norway

337 **ZooBank registration:** The LSID for *Rajonchocotyle emarginata* is  
338 [urn:lsid:zoobank.org:act:865F76DA-FADB-49F1-9B43-](https://zoobank.org/urn:lsid:zoobank.org:act:865F76DA-FADB-49F1-9B43-24F4AAC88256)  
339 [24F4AAC88256urn:lsid:zoobank.org:act:xxx](https://zoobank.org/urn:lsid:zoobank.org:act:xxx).

340

### 341 3.2.1 ~~Remarks~~ Amended diagnosis

342 In total, 21 morphological characters including soft body parts as well as sclerotised  
343 structures were measured (see Table 4, Figs. 5&6). Body elongate with tegument covered by  
344 numerous transverse ridges organised in radial rows (Fig. 6A). Haptor symmetrical with six  
345 suckers, armed with three pairs of C-shaped haptor sucker sclerites of similar shape and  
346 size (Fig. 5) and with a sharp hook (Fig. 6F). Peduncles of suckers of similar size. Each of the  
347 suckers contains a large sclerite ending in a hook pointing to the deep lumen (Fig. 6B&F).  
348 Sclerites form a bulge structure visible at the terminal region of each sucker (Fig. 6G). The  
349 sucker margin surmounted by a rim supporting the sclerite (Fig. 6F&H). Marginal haptor  
350 appendix with a pair of terminal suckers with three valves (Fig. 6D) and of V-shaped hamuli  
351 possessing a sharply pointed and curved tip (Fig. 6E) situated near the distal end of  
352 appendix. Mouth subterminal, situated on the ventral side of the body and formed by the  
353 oral sucker (Fig. 6C). Pharynx spherical, reaching the posterior end of oral sucker. Intestinal  
354 tract bifurcation at the level of pharynx. Testes occupy area in the central part of the body,  
355 irregular in size and shape, number of testes not ascertainable. Other parts of male  
356 reproductive system not distinguishable. Slightly lobed ovary (paratype USNM 1337399) and  
357 Y-shaped structure of vaginal ducts. Vitellaria extending from the level of intestinal  
358 bifurcation (paratype USNM 1337399) to the posterior end of the body into the haptor. Eggs  
359 fusiform with two incipient polar filaments (Fig. 5), located at level of anterior part of  
360 vitellarium.

### 361 3.2.2 Differential diagnosis

362 Up to now, there have been 17 genera of hexabothriid monogeneans described:  
363 *Branchotenthes* Bullard et Dippenaar, 2003; *Callorhynchocotyle* Suriano & Incorvaia, 1982;  
364 *Dasyonchocotyle* Hargis, 1955; *Epicotyle* Euzet & Maillard, 1974; *Erpocotyle* Van Beneden &  
365 Hesse, 1863; *Heteronchocotyle* Brooks, 1934; *Hexabothrium* von Nordmann, 1840;  
366 *Hypanocotyle* Chero, Cruces, Sáez, Camargo, Santos & Luque, 2018; *Mobulicola* Patella &  
367 Bullard, 2013; *Neonchocotyle* Ktari & Maillard, 1972; *Paraheteronchocotyle* Mayes, Brooks &  
368 Thorson, 1981; *Pristonchocotyle* Watson & Thorson, 1976; *Protocotyle* Euzet & Maillard,

369 1974; *Pseudohexabothrium* Brinkmann, 1952; *Rajonchocotyle* Cerfontaine, 1899;  
370 *Rhinobatonchocotyle* Doran, 1953 and *Squalonchocotyle* Cerfontaine, 1899. Species of  
371 *Rajonchocotyle* can be distinguished by the presence of a symmetrical haptor in comparison  
372 to *Callorhynchocotyle*, *Epicotyle*, *Heteronchocotyle*, *Neonchocotyle*, *Paraheteronchocotyle*,  
373 *Pristonchocotyle*, *Pseudohexabothrium* and *Rhinobatonchocotyle*. Species of *Rajonchocotyle*  
374 also differ from those within *Dasyonchocotyle* and *Hexabothrium* by having an unarmed  
375 male copulatory organ. Unlike representatives of *Branchotenthes*, *Erpocotyle*, *Hypanocotyle*,  
376 *Mobulicola* and *Squalonchocotyle* where the vagina is differentiated into muscular and  
377 glandular portions and possesses parallel vaginal ducts, species of *Rajonchocotyle* have an  
378 undifferentiated vagina and Y-shaped vaginal ducts. Species of *Rajonchocotyle* closely  
379 resemble representatives of *Protocotyle* but they differ by having undifferentiated vaginal  
380 ducts that are Y-shaped while the vaginal ducts of *Protocotyle* are also undifferentiated but  
381 parallel [51]. According to Boeger & Kritsky (1989), four species of *Rajonchocotyle* are  
382 currently considered valid: *Rajonchocotyle batis* Cerfontaine, 1899, *R. emarginata* (Olsson,  
383 1876), *Rajonchocotyle laevis* Price, 1942 and *Rajonchocotyle wehri* Price, 1942.  
384 *Rajonchocotyle emarginata* can be distinguished from its congeners by the total and  
385 proportional size of the haptoral sclerites. Unlike in *R. emarginata*, the size of sclerites of *R.*  
386 *batis* and *R. laevis* is not equal. While the anterior pair of *R. batis* is bigger than the posterior  
387 pair, sometimes even twice of the size, the posterior pair of *R. laevis* is just a bit smaller  
388 compared to the anterior pair. The difference between *R. emarginata* and *R. batis* is also  
389 visible in the shape of the anchor roots. The different pairs of haptoral sclerites of *R. wehri*  
390 and *R. emarginata* are all of equal size but the overall size of sclerites of *R. emarginata* is  
391 smaller than in *R. wehri* (median sclerites of 404–561  $\mu\text{m}$  compared to 924–956  $\mu\text{m}$  in *R.*  
392 *wehri*). *Rajonchocotyle emarginata* can also be distinguished from all its other congeners by  
393 having eggs with 2 long filaments (total egg length 353–446  $\mu\text{m}$ ) in comparison to the lack of  
394 polar filaments in *R. batis*, a small knob at each pole in *R. laevis* and two very short, fusiform  
395 egg filaments in *R. wehri* (total egg length 285–300  $\mu\text{m}$ ) [52–54].

396

#### 397 **4. Discussion**

398 Invertebrate diversity is understudied in polar regions, with available information biased  
399 towards a few taxa [18,19]. In general, knowledge about the parasite fauna in cold areas  
400 remains poor, and zoonotic parasitosis received most attention so far [21]. Despite an

401 intensified effort, reflected by the numerous parasitological surveys conducted recently [55–  
402 58], fish flatworms are rarely studied in the Svalbard archipelago. Rokicka (2009) does not  
403 mention a single monogenean infection for over 94 examined fish specimens belonging to 4  
404 species [59]. Our study is the first record of monogenean species off Spitsbergen Island and  
405 the Svalbard archipelago.

406

#### 407 4.1. Species richness and geographic distribution of *Acanthocotyle* and *Rajonchocotyle*

408 In total, 12 currently valid species of Acanthocotylidae have been reported out of 15 species  
409 accounting for 5.6% of the species diversity of Rajiformes. Two teleost fish species,  
410 *Reinhardtius hippoglossoides* (Walbaum, 1792) and *Sebastes alutus* (Gilbert, 1890), were  
411 also recorded as hosts for *Acanthocotyle williamsi* [60,61]. The recorded infection of  
412 *Acanthocotyle* sp. on *Narcine maculata* (Torpediniformes) is rather considered to result from  
413 transfer during fish capture [62]. With an estimation of over 800 species, the number of  
414 potential elasmobranch and holocephalan hosts of species of Acanthocotylidae is high. From  
415 less than 10% of them [63,64], only 59 species of Hexabothriidae have been described [51].  
416 Considering the overall high species richness and rather stricter host-specificity of  
417 monogeneans, the known diversity of both examined monogenean groups infecting  
418 cartilaginous fishes can be assumed to be proportionally minimal.

419 The overall worldwide occurrence of *Acanthocotyle* spp. seems to follow the geographical  
420 distribution of their skate hosts as summarised in [Ñacari et al. \[49\]](#)–~~Ñacari et al. (2019)~~. Host-  
421 specificity ranges from one (9 species of *Acanthocotyle*) to five host species in *A. lobianchi*.  
422 However, the known distribution of *A. lobianchi* is currently mostly restricted to Plymouth,  
423 UK [65] with one record from Naples, Italy [66]. A similar host ~~range~~ (four rajid species) and  
424 ~~occurrence-geographical~~ range was reported for *A. pacifica* [67–69]. The distribution of *A.*  
425 *verrilli* overlaps with the cross-Atlantic occurrence of *A. radiata*. This monogenean species  
426 was further reported from two other rajid species so far (see Table 5). However, reported  
427 differences in host-specificity and distribution patterns of *Acanthocotyle* spp. are suggested  
428 to result from biased sampling toward a few host species.

429 Unlike in *Acanthocotyle*, species of *Rajonchocotyle* seem to be less host specific as several  
430 representatives were recorded from different rajid hosts with a maximum number of 10 in  
431 the case of *R. emarginata*. This difference in host-specificity possibly can be driven by the  
432 site of infection (skin in *Acanthocotyle* spp. versus gills in *Rajonchocotyle* spp.) or mode of



433 reproduction (eggs being attached to the parasite body by stalks in *Acanthocotyle* spp.  
434 [70,71] and floating eggs and free-swimming larvae in *Rajonchocotyle* spp. [72,73]. However,  
435 discovery of cryptic species that are more host specific than the species they were originally  
436 assigned to, has already changed views on parasite species richness and host-specificity [74].  
437 Given the lack of genetic data on species of *Rajonchocotyle* and hexabothriids in general,  
438 and their close interspecific morphological similarity, the presence of cryptic species cannot  
439 be excluded. Further research is needed to verify the level of host-specificity and overall  
440 distribution patterns of both monogenean families as large parts of the host distribution  
441 remains devoid of parasitological investigation.

442

#### 443 4.2. Morphological and genetic diversity

444 The specimens of *A. verrilli* analysed in this study did not differ in most morphological  
445 characteristics from earlier reports but some differences were observed. Goto (1899) and  
446 Sproston (1946) counted 30 or 32 radial rows of sclerites in the haptor while in our  
447 specimens the rows ranged from 28 to 34 (32 most frequently, in 14 of the 34 specimens).  
448 Thus, intraspecific variation in the number of radial rows of sclerites was reported in *A.*  
449 *verrilli*, as in other congeners [32,49,50]. Interestingly, [a relationship between age and](#)  
450 [therelated](#) number of radial rows of polyopisthocotylean monogenean species was  
451 suggested [75]. However, such pattern does not emerge for our data (see Supplementary  
452 Table S1). Moreover, because the number of rows in the pseudohaptor and the number of  
453 rows of testes can overlap, the difference between *A. verrilli* and *A. atacamensis* was  
454 revised. We propose the number of sclerites in the first row (counting from the position of  
455 the true haptor) as an additional diagnostic character. This study confirms that the  
456 morphology and size of sclerotised structures are of a high diagnostic significance in  
457 *Acanthocotyle*. Close morphological similarities of *A. verrilli* with *A. imo* and *A. atacamensis*  
458 are reflected in the genetic distance matrix (Table 3). Interestingly, the two species of  
459 *Acanthocotyle* collected from representatives of *Amblyraja* formed a [monophyletic](#) clade in  
460 the phylogenetic reconstruction (Fig. 6). However, molecular data on the remaining  
461 *Acanthocotyle* spp. as well as haplotypes of *A. verrilli* from other rajid genera are needed to  
462 shed light on the evolutionary history of this parasite-host system.

463 As pointed out in previous studies, boundaries between hexabothriid monogenean species  
464 are mostly defined by variable characters, as they are unstable across different fixation and

465 staining methods [51,76]. Vaughan and Christison (2012), using multivariate statistics,  
466 combined measurements of the hamulus and sucker sclerites to distinguish species of  
467 *Callorhynchocotyle*. Our study confirmed that the morphology of the hamulus and the size of  
468 sucker sclerites is of a high diagnostic significance. As a result, the combination of the  
469 proportional size of sclerotised structures and egg filaments is proposed for species  
470 identification of *Rajonchotyle* spp. Moreover, multiple staining methods should be used for  
471 correct assignment of hexabothriid monogeneans in general and species of *Rajonchocotyle*  
472 in particular.

473

#### 474 4.3. Parasites of *A. radiata* as a tag for host population structure

475 To date, ~~288~~ helminth parasite species were reported from *A. radiata* worldwide. Even  
476 though our study was restricted to monogeneans, ongoing investigations suggest the  
477 presence of at least 9 endoparasitic helminth species infecting *A. radiata* in Svalbard  
478 (unpublished). *Acanthocotyle verrilli* was further reported from *A. radiata* by Sproston, 1946  
479 off Bergen, Norway, North-eastern Norwegian Sea (Rokicki and Berland, 2009), ~~and~~ on the  
480 opposite side of the Atlantic Ocean off Newfoundland and Chaleur Bay in Canada [29,78]  
481 and the northern East coast of the USA [79]. Unlike in *A. verrilli*, the known occurrence of *R.*  
482 *emarginata* on *A. radiata* also spans the southern hemisphere, on the Patagonian Shelf [80].  
483 Moreover, *R. batis* and two monogenean species from Monocotylidae were reported  
484 parasitizing on *A. radiata* in previous studies (see Table 1).

485 Although our results match with the previous records of monogeneans collected from *A.*  
486 *radiata*, the new locality off Svalbard represents the highest known latitude in the northern  
487 hemisphere those two monogenean genera have ever been recorded from (see Tables 5&6).  
488 As monogeneans display a direct life cycle and short-lived larval stage [73,81], their  
489 distribution is primarily affected by the distribution and migration patterns of their host.  
490 Given the previous reports of both monogenean species from *A. radiata* on both sides of  
491 Atlantic Ocean, historical and ongoing overseas connectivity of skate species at the Svalbard  
492 coast is proposed as suggested in Chevolot et al. [82] ~~Chevolot et al., (2007)~~. Differences in  
493 life history traits such as total length and density between the populations of thorny skate  
494 were observed [5,83,84]. In contrast to the previously recorded high level of fidelity and  
495 relatively small home range (mostly fewer than 100 km, ~~[85–88]~~ ~~Daan et al., 2005; Dulvy et~~  
496 ~~al., 2000; Templeman, 1984; Walker et al., 1997~~), long-term connectivity between thorny

497 skate populations from the eastern and western part of the Atlantic facilitated by historical  
498 population expansion was recently documented [82]. These recently suggested large  
499 migratory capacities of thorny skates concur with the occasional records at depths down to  
500 1000 m [89] with continental shelves considered as important migration barrier [16].  
501 However, there are low levels of migration between the North Sea population and other  
502 European areas [82]. Given their expected higher mutation rate and reproduction coefficient  
503 compared to their hosts, parasitic flatworms including monogeneans were proposed as tags  
504 for historical and ongoing host migration [90–92]. However, both reported species of  
505 monogeneans infecting thorny skate are not strictly host specific (see Table 5&6). Other  
506 skate species can therefore contribute to the worldwide occurrence of *R. emarginata* and  
507 the occurrence throughout the northern hemisphere of *A. verrilli*. In general, monogeneans  
508 tend to be less host specific in pelagic and deepwater areas in comparison to littoral habitats  
509 due to the lower host availability connected also with fish population size [93–96]. On the  
510 other hand, an influence of light intensity on the larval hatching of *R. emarginata* as a result  
511 of adaptation to the behavioural differences between the hosts was proposed [72]. Such an  
512 adaptation might therefore have resulted in depth-dependent host specificity of this  
513 parasite species. Keeping in mind the rather plastic nature of currently used morphological  
514 characters in both examined monogenean families, phenotypic evaluation of any  
515 differentiation would need to be employed over a large number of specimens per  
516 population.

517 More variable genetic markers would need to be applied in order to investigate the  
518 historical connectivity of the parasite populations in Svalbard and other areas and evaluate  
519 the level of geographically/host species driven differentiation.

520

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533

#### 534 **Figure captions**

535 Fig. 1: *Acanthocotyle verrilli* ex *Amblyraja radiata*. A) Ventral view of the whole body. B)  
536 Reproductive system. Abbreviations: aa, anterior adhesive lobes; ag, accessory glands; bl,  
537 excretory bladders; ed, ejaculatory duct; ge, germarium; gr, male accessory gland reservoir;  
538 h, haptor; in, intestine; ph, pharynx; sr, seminal receptacle; sv, bipartite seminal vesicle; te,  
539 testes; th, true haptor; up, uterine atrium; ut, uterus; vd, vas deferens; vt, vitelline duct; vi,  
540 vitelline.

541 Fig. 2: SEM observations of *Acanthocotyle verrilli*. A) Ventral view of the whole body, B) View  
542 of the anterior part of the body with adhesive lobes on each side, visible sense organ and  
543 pharynx opening. C) Uterus opening with an externally attached egg. D) Detailed apical view  
544 of an egg with abopercular appendage. E) Posterior part of the body formed by haptor and  
545 convex pseudohaptor, pseudohaptor with marginal valve. F) Detailed view of  
546 pseudophaptoral sclerites organised in rows. G) Detailed view of haptor located at the  
547 posterior margin of pseudohaptor armed with 14 peripheral and 2 central marginal hooks.  
548 H) aa – adhesive lobes, eg – egg operculum, h – haptor, m – marginal valve, mh – marginal  
549 hook, po – pharynx opening, mhs – marginal hooks sclerite, ps – pseudohaptor, so – sense  
550 organ.

551 Fig. 3: Biplots showing the interspecific differences of *Acanthocotyle* spp. based on  
552 proportional morphometric measurements standardized by the total body length. Only the  
553 first two axes are shown. A) Principal component analysis (PCA) of *A. verrilli* (this study), *A.*  
554 *atacamensis* [49] (~~Ñacari et al. 2019~~) and *A. imo* [49] (~~Ñacari et al. 2019~~). B) PCA of *A. verrilli*  
555 (this study) and *A. atacamensis* [49] (~~Ñacari et al. 2019~~). C) PCA of *A. verrilli* (this study) and  
556 *A. imo* [49] (~~Ñacari et al. 2019~~).

557 Fig. 4: Bayesian inference phylogram based on available 28S rDNA sequences of  
558 Acanthocotylidae with specification of the host species. Representatives of [three other](#)  
559 [families of Gyrocotylidea Gyrodactylidea](#) were used as outgroup. Bootstrap percentages for  
560 maximum likelihood (before slashes) and posterior probabilities for Bayesian inference

561 (behind slashes) are shown. The scale bar indicates the expected number of substitutions  
562 per site.

563 Fig. 5: *Rajonchocotyle emarginata* ex *Amblyraja radiata*. A) Ventral view of the whole body.  
564 B) Anterior haptoral sucker sclerites. C) Middle haptoral sucker sclerites. D) Posterior  
565 haptoral sucker sclerites. E) Hamuli. F) Egg. Abbreviations: ap, appendix; eg, egg; h, haptor;  
566 ha, hamuli; it, intestine; os, oral sucker; ph, pharynx; sc, sclerite; su, sucker; te, testes; vd,  
567 vaginal duct; vi, vitellarium.

568 Fig. 6: SEM observations of *Rajonchocotyle emarginata*. A) Ventral view of the whole body,  
569 B) Haptoral structures formed by six suckers each armed with a sclerite, and appendix, C)  
570 View of the anterior part of the body with the oral sucker, D) Lateral view of the posterior  
571 part of the appendix with two terminal suckers. E) Bottom-up view on the posterior part of  
572 the appendix with two terminal suckers. F) Haptoral sucker rounded by the rim with a  
573 sclerite bulge ending in a hook. G) Side view of haptoral suckers with peduncle structure on  
574 the left side. H) Detailed view of the sucker rim and hook. ap – appendix, as – appendicular  
575 sucker, h – hook, hp – haptor, rm – sucker rim, sb – sucker bulge, sp – sucker peduncle os –  
576 oral sucker.

#### 577 **Supplementary material**

578 Table S1: Raw morphometric data for *Acanthocotyle verrilli* ex *Amblyraja radiata*.  
579 Measurement are given in micrometers.

580 Table S2: Raw morphometric data for *Rajonchocotyle emarginata* ex *Amblyraja radiata*.  
581 Measurement are given in micrometers.

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**Table 1:** List of helminth species reported from *Amblyraja radiata* with taxonomic designation and locality of the report.

Class	Species	Family	Locality	Reference
Cestoda	<i>Anthobothrium cornucopia</i> Van Beneden, 1850	Tetraphylidea	off Newfoundland, Canada	[78]
	<i>Calyptrbothrium riggii</i> Monticelli, 1893	Phyllobothriidae	off Kattegat, Denmark	[97]
	<i>Echeneibothrium canadensis</i> Keeling & Burt, 1996	Echeneibothriidae	Barent Sea Bay of Fundy, Canada	Review in [23] [98]
	<i>Echeneibothrium dubium</i> Van Beneden, 1858		Northwestern Atlantic Ocean Chaleur Bay, Canada	[99] [29]
	<i>Echeneibothrium variabile</i> Van Beneden, 1850		Barent Sea	Review in [23]
	<i>Echinobothrium raji</i> Heller, 1949	Echinobothriidae	Barent Sea	Review in [23]
	<i>Grillotia (Grillotia) erinaceus</i> (Van Beneden, 1858)	Lacistorhynchidae	Atlantic Coast of North America Barent Sea	[100] Review in [23]
	<i>Grillotia (Grillotia) brayi</i> Beveridge & Campbell, 2007		Kolkugrunn, Iceland Chaleur Bay, Canada	[101] [29]
	<i>Grillotia</i> sp. Guiart, 1927		North-eastern Norwegian Sea	[102]
			Northwestern Atlantic Ocean	[99]

	<i>Phormobothrium affine</i> (Olsson, 1867)	Phyllobothriidae	Chaleur Bay, Canada	[29]
	<i>Phyllobothrium dagnalium</i> Southwell, 1927		Northwestern Atlantic Ocean	[103]
	<i>Phyllobothrium</i> sp. Van Beneden, 1850		off Newfoundland, Canada	[78]
	<i>Phyllobothrium thridax</i> Van Beneden, 1849		Barent Sea	Review in [23]
	<i>Pseudanthobothrium hansenii</i> Baer, 1956	Echeneibothriidae	North-eastern Norwegian Sea	[102]
	<i>Scyphophyllidium giganteum</i> (Van Beneden, 1858)	Phyllobothriidae	Chaleur Bay, Canada Barent Sea	[29,78] Review in [23]
	<i>Trilocularia acanthiaevulgaris</i> Olsson, 1867	Tetraphyllidea	off Newfoundland, Canada	[78]
	<i>Tritaphros retzii</i> Lönnberg, 1889	Echeneibothriidae	Barent Sea	Review in [23]
<b>Trematoda</b>	<i>Aporocotylidae</i> gen. sp.	Aporocotylidae	Barent Sea	Review in [23]
	<i>Gonocerca phycidis</i> Manter, 1925	Gonocercidae	North Sea south of Fair Isle, UK	[104]
	<i>Hemiurus levinseni</i> Odhner, 1905	Hemiuridae	Barent Sea	[23]
	<i>Otodistomum cestoides</i> (Van Beneden, 1870)	Azygiidae	Chaleur Bay, Canada off Newfoundland, Canada	[29] [78]
			North-eastern Norwegian Sea	[102]
			Barent Sea	Review in [23]
<b>Monogenea</b>	<i>Acanthocotyle verrillii</i> Goto, 1899	Acanthocotylidae	Cape Cod, USA off Bergen, Norway Chaleur Bay, Canada off Newfoundland, Canada	[105] [106] [29] [78]*
			North-eastern Norwegian Sea	[102]*
			Barent Sea	Review in [23]
			Adventfjorden, Spitsbergen	This study
	<i>Dictyocotyle coeliaca</i> Nybelin, 1941		off Trondheim, Norway North Sea, UK	[107] [104,108,109]
			North-western Atlantic Ocean, Canadian coast	[110]
	<i>Rajonchocotyle batis</i> Cerfontaine, 1899	Hexabothriidae	North-eastern Norwegian Sea	[102]

	<i>Rajonchocotyle emarginata</i> (Olsson, 1876)		Chaleur Bay, Canada Barent Sea Patagonian Shelf off Tromsø, Norway Adventfjorden estuary, Spitsbergen	[29] Review in [23] [111] [112] This study
<b>Acanthocephala</b>	<i>Echinorhynchus gadi</i> Zoega in Müller, 1776	Echinorhynchidae	North-eastern Norwegian Sea	[102]
<b>Nematoda</b>	<i>Anisakis</i> sp. Dujardin, 1845	Anisakidae	off Newfoundland, Canada	[78]
	<i>Anisakis simplex</i> (Rudolphi, 1809)		Barent Sea	[23]
	<i>Contracaecum plagiosomorum</i> (Linstow, 1905)		North-eastern Norwegian Sea	[102]
	<i>Contracaecum</i> sp. Railliet & Henry, 1912		West coast of Norway Barent Sea	[113] Review in [23]
	<i>Hysterothylacium aduncum</i> (Rudolphi, 1802)	Raphidascarididae	off Newfoundland, Canada	[78]
	<i>Phocascaris</i> sp. Höst, 1932	Anisakidae	North-eastern Norwegian Sea	[102]
	<i>Porrocaecum</i> sp. Railliet & Henry, 1912	Ascarididae	off Newfoundland, Canada	[78]
	<i>Pseudanisakis tricupola</i> Gibson, 1973	Acanthocheilidae	Bay of Fundy, Canada Barent Sea	[98] Review in [23]

948 **Table 2:** Meristic and morphometric data for *Acanthocotyle verrilli* ex *Amblyraja radiata*  
 949 from Adventfjorden, Spitsbergen. Measurements are given in micrometers.

<i>Acanthocotyle verrilli</i> (n=30)	
Total body length	1069–5855 (3541) <sup>a</sup>
Body proper length	715–4688 (2757) <sup>b</sup>
Body width at the level of ovary	330–925 (632) <sup>b</sup>
Pharynx	
Length	124–340 (233) <sup>b</sup>
Width	77–413 (252) <sup>b</sup>
Pseudohaptor	
Number radial rows of sclerites	28–34 (mode=32) <sup>b</sup>
Length	354–1167 (767) <sup>b</sup>
Width	312–1112 (765) <sup>b</sup>
Testes	
Testicular field length	393–2583 (1522) <sup>c</sup>
Number	18–47 (mode=34) <sup>d</sup>
Maximum width	80–267 (173) <sup>c</sup>
Germarium	
Length	60–419 (215) <sup>c</sup>
Width	98–347 (203) <sup>c</sup>

950 Notes: <sup>a</sup>(n=28); <sup>b</sup>(n=29); <sup>c</sup>(n=26); <sup>d</sup>(n=22)



951 **Table 3:** Uncorrected pairwise genetic distances (%) between *Acanthocotyle* spp. based on  
 952 844bp of 28S rDNA gene portion. GenBank accession numbers are given in brackets.  
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	<i>A. verrilli</i>	<i>A. atacamensis</i>	<i>A. gurgesiella</i>	<i>A. imo</i>
<i>A. verrilli</i> ( <del>MW260310xx-xx</del> )				
<i>A. atacamensis</i> (MH511079)	0.75			
<i>A. gurgesiella</i> (KY379329)	2.01	2.26		
<i>A. imo</i> (MH511085)	0.25	0.50	1.76	
<i>A. urolophi</i> (FJ971962)	3.01	3.01	3.27	2.76

954 **Table 4:** Meristic and morphometric data for *Rajonchocotyle emarginata* ex *Amblyraja*  
 955 *radiata* from Adventfjorden, Spitsbergen. Measurements are given in micrometers.

<i>Rajonchocotyle emarginata</i> (n=5)	
Total body length	8497–10556 (9449) <sup>a</sup>
Body proper length	6465–7943 (7093) <sup>b</sup>
Body width at level of ovary	1604–2601 (2073) <sup>b</sup>
Anterior sucker	
Length	101–530 (280) <sup>a</sup>
Width	120–858 (506) <sup>a</sup>
Pharynx	
Length	113–116 (115) <sup>c</sup>
Width	121–121 (121) <sup>c</sup>
Haptor	
Length	1547–2642 (2199) <sup>a</sup>
Width	2113–3337 (2556) <sup>a</sup>
Haptoral sclerites	
Anterior sclerite length	417–495 (455) <sup>a</sup>
Median sclerite length	402–503 (447) <sup>a</sup>
Posterior sclerite length	316–523 (420) <sup>a</sup>
Appendix	
Length	1913–4078 (3247) <sup>a</sup>
Width	605–939 (749) <sup>a</sup>
Hamulus	
Outer root length	42–51 (46) <sup>c</sup>
Inner root length	37–43 (40) <sup>c</sup>
Testes	
Testicular field length	1822–4075 (3088) <sup>a</sup>
Number	10–36 (mode=24) <sup>a</sup>
Egg	
Proper length	248–303 (280) <sup>a</sup>
Number of filaments	2 <sup>d</sup>
Length of filament	105–143 (123) <sup>c</sup>

956 Notes: <sup>a</sup>(n=4); <sup>b</sup>(n=5); <sup>c</sup>(n=2); <sup>d</sup>(n=3)

957 **Table 5:** List of *Acanthocotyle* spp. with host species designation and locality of the report.

Parasite species	Host species	Family	Locality	Reference
<i>Acanthocotyle atacamensis</i> Ñacari, Sepúlveda, Escribano & Oliva, 2019	<i>Bathyraja peruana</i> McEachran & Miyake, 1984	Arhynchobatidae	off Tocopilla, Chile	[49]
<i>Acanthocotyle elegans</i> Monticelli, 1890	<i>Raja clavata</i> Linnaeus, 1758	Rajidae	off Plymouth, UK	[65]
<i>Acanthocotyle greeni</i> Macdonald & Llewellyn, 1980	<i>Raja clavata</i>	Rajidae	off Naples, Italy off Plymouth, UK	[66] <sup>1</sup> [70]
<i>Acanthocotyle gurgesiella</i> Ñacari, Sepulveda, Escribano & Oliva, 2017	<i>Gurgesiella furvescens</i> de Buen, 1959	Gurgesiellidae	off Valparaiso, Chile	[50]
<i>Acanthocotyle imo</i> Ñacari, Sepúlveda, Escribano & Oliva, 2019	<i>Amblyraja frerichsi</i> Krefft, 1968	Rajidae	off Tocopilla, Chile	[49]
<i>Acanthocotyle lobianchi</i> Monticelli, 1888	<i>Bathyraja brachyurops</i> (Fowler, 1910) <sup>a</sup>	Arhynchobatidae	off Plymouth, UK	[65]
	<i>Leucoraja naevus</i> (Müller & Henle, 1841) <sup>b</sup>	Rajidae		
	<i>Raja clavata</i>	Rajidae	off Naples, Italy off Plymouth, UK	[66] [65]
	<i>Raja microocellata</i> Montagu, 1818	Rajidae		
	<i>Raja montagui</i> Fowler, 1910	Rajidae		
<i>Acanthocotyle pacifica</i> Bonham & Guberlet, 1938	<i>Beringraja binoculata</i> (Girard, 1855) <sup>c</sup>	Rajidae	Puget Sound, USA	[67]
			Friday Harbour, USA	[69]
	<i>Raja rhina</i> Jordan & Gilbert, 1881	Rajidae	Puget Sound, USA Friday Harbour, USA	[68] <sup>2</sup> [69]
<i>Acanthocotyle patagonica</i> Kuznetsova, 1975	<i>Bathyraja brachyurops</i> <sup>a</sup>	Arhynchobatidae	Patagonian Shelf	[80]
<i>Acanthocotyle pugetensis</i> Bonham & Guberlet, 1938	<i>Beringraja binoculata</i> <sup>c</sup>	Rajidae	Friday Harbour, USA off San Francisco, USA	[69] [114]
<i>Acanthocotyle urolophi</i> Kearn, Whittington, Chisholm & Evans-Gowing, 2016	<i>Urolophus cruciatus</i> (Lacepède, 1804)	Urolophidae	off Tasmania, Australia	[32]

<i>Acanthocotyle verrilli</i> Goto, 1899	<i>Amblyraja radiata</i> (Donovan, 1808) <sup>d</sup>	Rajidae	Cape Cod, USA	[105]
	<i>Amblyraja radiata</i> <sup>d</sup>		off Bergen, Norway	[106]
	<i>Amblyraja radiata</i> <sup>e</sup>		Chaleur Bay, Canada	[29]
	<i>Amblyraja radiata</i> <sup>d</sup>		off Newfoundland, Canada	[78] <sup>2</sup>
	<i>Amblyraja radiata</i> <sup>d</sup>		Continental slope between Norway and Spitsbergen	[77] <sup>2</sup>
	<i>Amblyraja radiata</i>		Barent Sea	Review in [23] <sup>2</sup>
			Adventfjorden estuary, Spitsbergen	This study
	<i>Bathyrāja spinicauda</i> (Jensen, 1914)	Arhynchobatidae	Continental slope between Norway and Spitsbergen	[77] <sup>2</sup>
	<i>Leucoraja erinacea</i> (Mitchill, 1825) <sup>f</sup>	Rajidae	Cape Cod, USA	[105]
			Coast of Maine, USA	[79]
<i>Acanthocotyle williamsi</i> Price, 1938	Unidentified skate	-	Bering Sea, Aleutian Island (Salt Island)	[115]
<i>Acanthocotyle</i> sp.	<i>Bathyrāja brachyurops</i> <sup>a</sup>	Rajidae	Northwest coast of Spain	[116]
	<i>Narcine maculata</i> (Shaw, 1804) <sup>g</sup>	Narcinidae	off Plymouth, UK	[62]
	<i>Raja clavata</i>	Rajidae	off Plymouth, UK Belgian Coast	[106]
	<i>Raja microocellata</i>	Rajidae	Northwest coast of Spain	[116]
	<i>Sympterygia bonapartii</i> Müller & Henle 1841	Arhynchobatidae	off Puerto Deseado, Argentina	[117]

958 <sup>a</sup> reported as *Raja brachyurops* Fowler, 1910  
959 <sup>b</sup> reported as *Raja naevus* Müller & Henle, 1841  
960 <sup>c</sup> reported as *Raja binocularata* Müller & Henle, 1841  
961 <sup>d</sup> reported as *Raja radiata* Donovan, 1808  
962 <sup>e</sup> reported as *Raja scabrata* Garman, 1913  
963 <sup>f</sup> reported as *Raja erinacea* Mitchill, 1825  
964 <sup>g</sup> reported as *Raja maculate* Shaw, 1804  
965 <sup>1</sup> reported as *Acanthocotyle oligoterus* Monticelli, 1899  
966 <sup>2</sup> reported as *Pseudoacanthocotyle*

**Table 6:** List of *Rajonchocotyle* spp. with host species designation and locality of the report.

Parasite species	Host species	Family	Locality	Reference
<i>Rajonchocotyle batis</i> Cerfontaine, 1899	<i>Amblyraja hyperborea</i> (Collett, 1879) <sup>a</sup>	Rajidae	Continental slope between Norway and Spitsbergen	[77]
	<i>Amblyraja radiata</i> (Donovan, 1808) <sup>b</sup>	Rajidae	Continental slope between Norway and Spitsbergen	
	<i>Beringraja binoculata</i> (Girard, 1855) <sup>c</sup>	Rajidae	Friday Harbor, Salish Sea, USA	[69]
	<i>Bathyraja spinicauda</i> (Jensen, 1914)	Arhynchobatidae	Continental slope between Norway and Spitsbergen	[77]
	<i>Dipturus batis</i> (Linnaeus, 1758) <sup>d</sup>	Rajidae	off Skagerrak, Denmark	[54]
			off Liège, Belgium	[52]
			off Ostend, Belgium	
			Porcupine Bank, Ireland	[118]
			off Plymouth, UK	[65]
		<i>Raja clavata</i> Linnaeus, 1758	Rajidae	off Roscoff, France
<i>Rajonchocotyle emarginata</i> (Olsson, 1876)	<i>Amblyraja radiata</i> <sup>e</sup>	Rajidae	Chaleur Bay, Canada	[29]
	<i>Amblyraja radiata</i> <sup>b</sup>		Patagonian Shelf	[111]
	<i>Amblyraja radiata</i>		off Tromsø, Norway	[30]
			Barent Sea	Review in [23]
			Adventfjorden estuary, Spitsbergen	This study
	<i>Bathyraja brachyurops</i> (Fowler, 1910) <sup>f</sup>	Arhynchobatidae	Patagonian Shelf	[111]
	<i>Bathyraja magellanica</i> (Philippi, 1902) <sup>g</sup>			
	<i>Leucoraja naevus</i> (Müller & Henle, 1841) <sup>h</sup>	Rajidae	off Plymouth, UK	[65]
	<i>Psammobatis scobina</i> (Philippi, 1857) <sup>i</sup>	Arhynchobatidae	Patagonian Shelf	[111]
	<i>Raja brachyura</i> Lafont, 1871	Rajidae	off Plymouth, UK Northwest coast of Spain	[65] [116]

	<i>Raja clavata</i>	Rajidae	off Plymouth, UK Mari Bahusiae, Scandinavia Mediterranean Sea, Italy Cardigan Bay, Wales off Roscoff, France Patagonian Shelf Galway Bay, Ireland	[119] [65] [120] [121,122] [119] [53] [111] [123]
	<i>Raja microocellata</i> Montagu, 1818	Rajidae	off Plymouth, UK Northwest coast of Spain	[65] [116]
	<i>Raja montagui</i> Fowler, 1910	Rajidae	off Plymouth, UK	[65]
	<i>Raja</i> sp.	Rajidae	Northwest coast of Spain	[116]
	<i>Raja undulata</i> Lacepède, 1802	Rajidae	Northwest coast of Spain	
<i>Rajonchocotyle laevis</i> Price, 1942	<i>Dipturus laevis</i> (Mitchill, 1818) <sup>j</sup>	Rajidae	Woods Hole, USA	[53]
<i>Rajonchocotyle wehri</i> Price, 1942	<i>Raja asterias</i> (Delaroche, 1809) <sup>k</sup>	Rajidae	Friday Harbor, USA	[53]

<sup>a</sup> reported as *Raja hyperborea* Collett, 1879

<sup>b</sup> reported as *Raja radiata* Donovan, 1808

<sup>c</sup> reported as *Raja binocolata* Girard, 1855

<sup>d</sup> reported as *Raja batis* Linnaeus, 1758

<sup>e</sup> reported as *Raja scabrata* Garman, 1913

<sup>f</sup> reported as *Raja brachyurops* Fowler, 1910

<sup>g</sup> reported as *Raja magellanica* Philippi, 1902

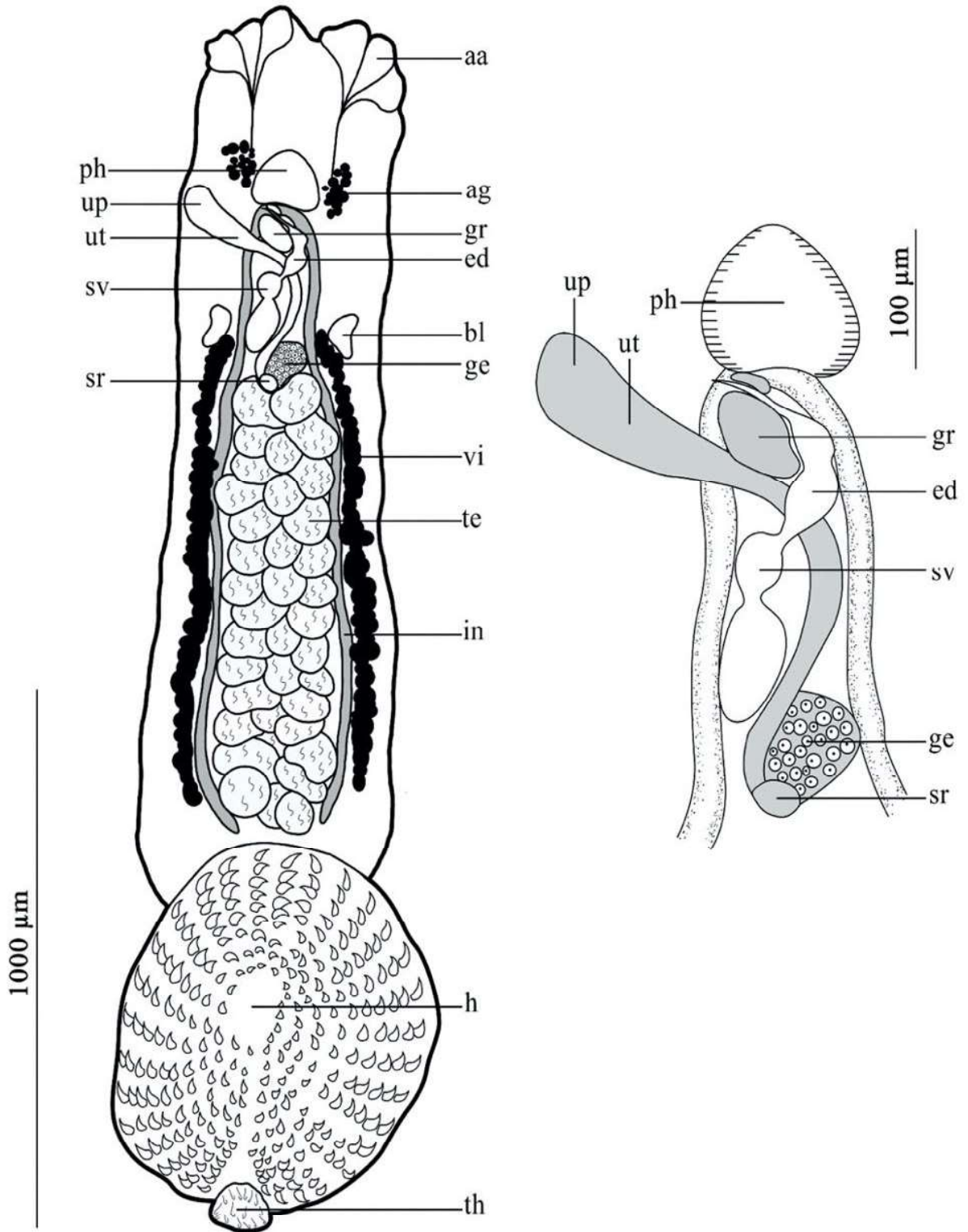
<sup>h</sup> reported as *Raja naevus* Müller & Henle

<sup>i</sup> reported as *Raja scobina* Philippi, 1857

<sup>j</sup> reported as *Raja laevis* Mitchill, 1818

<sup>k</sup> reported as *Raja stellata* Delaroche, 1809

Figure 1



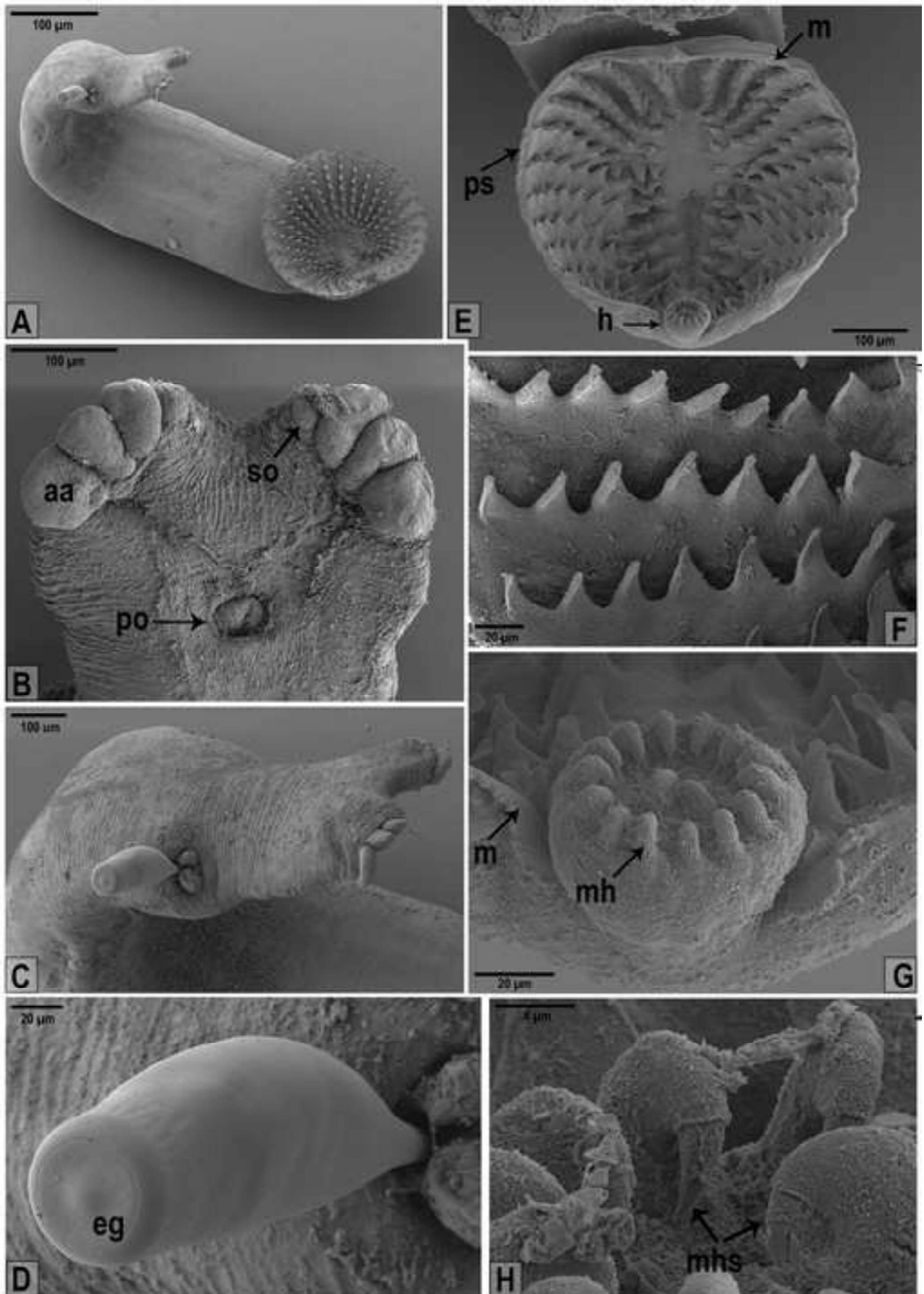
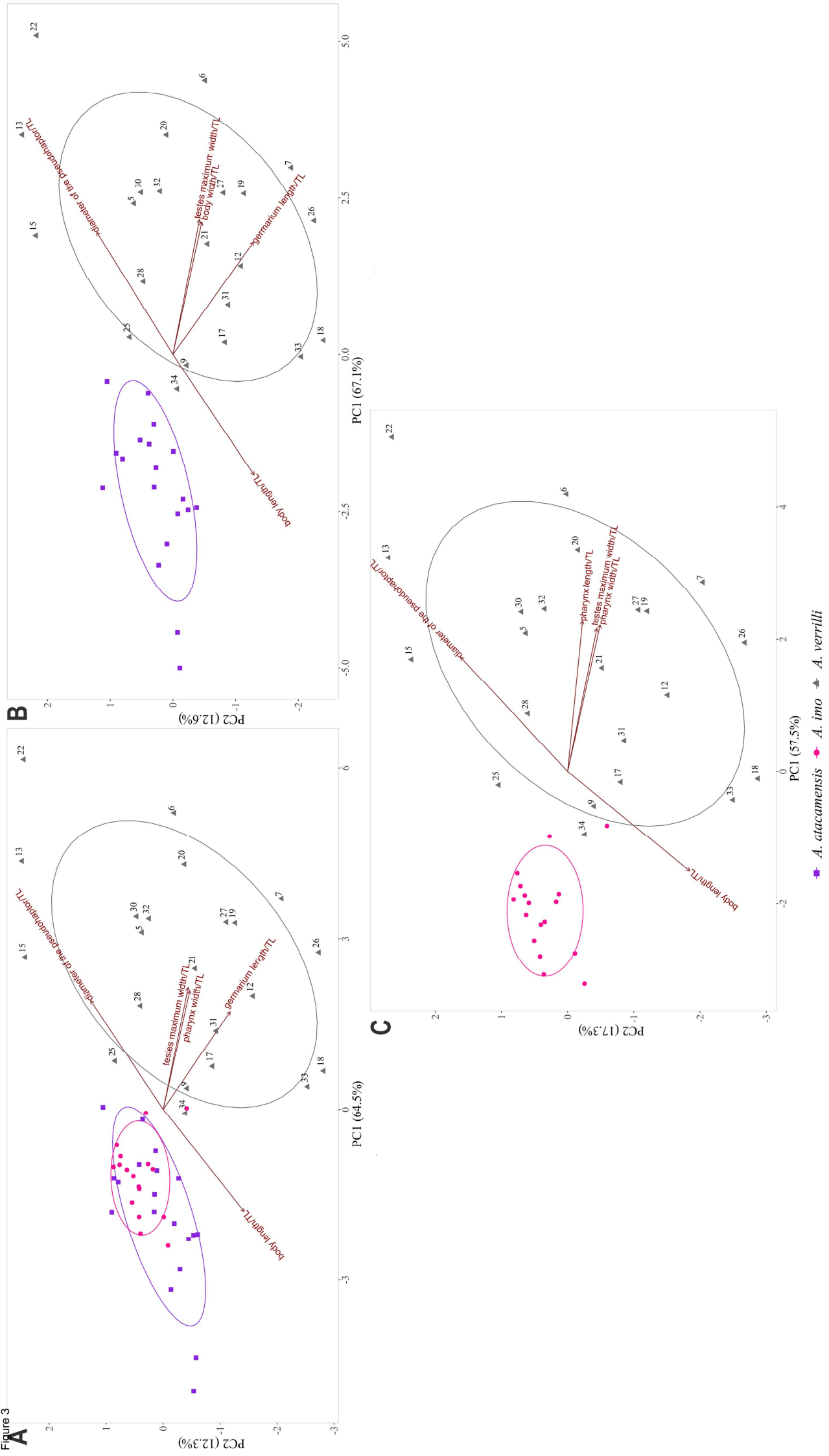
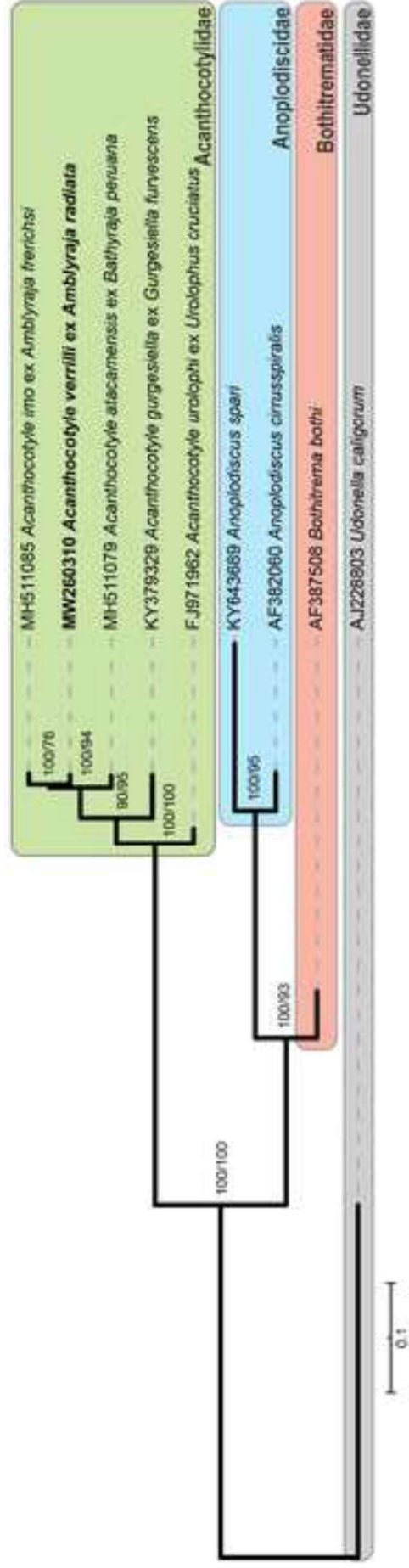




Figure 3





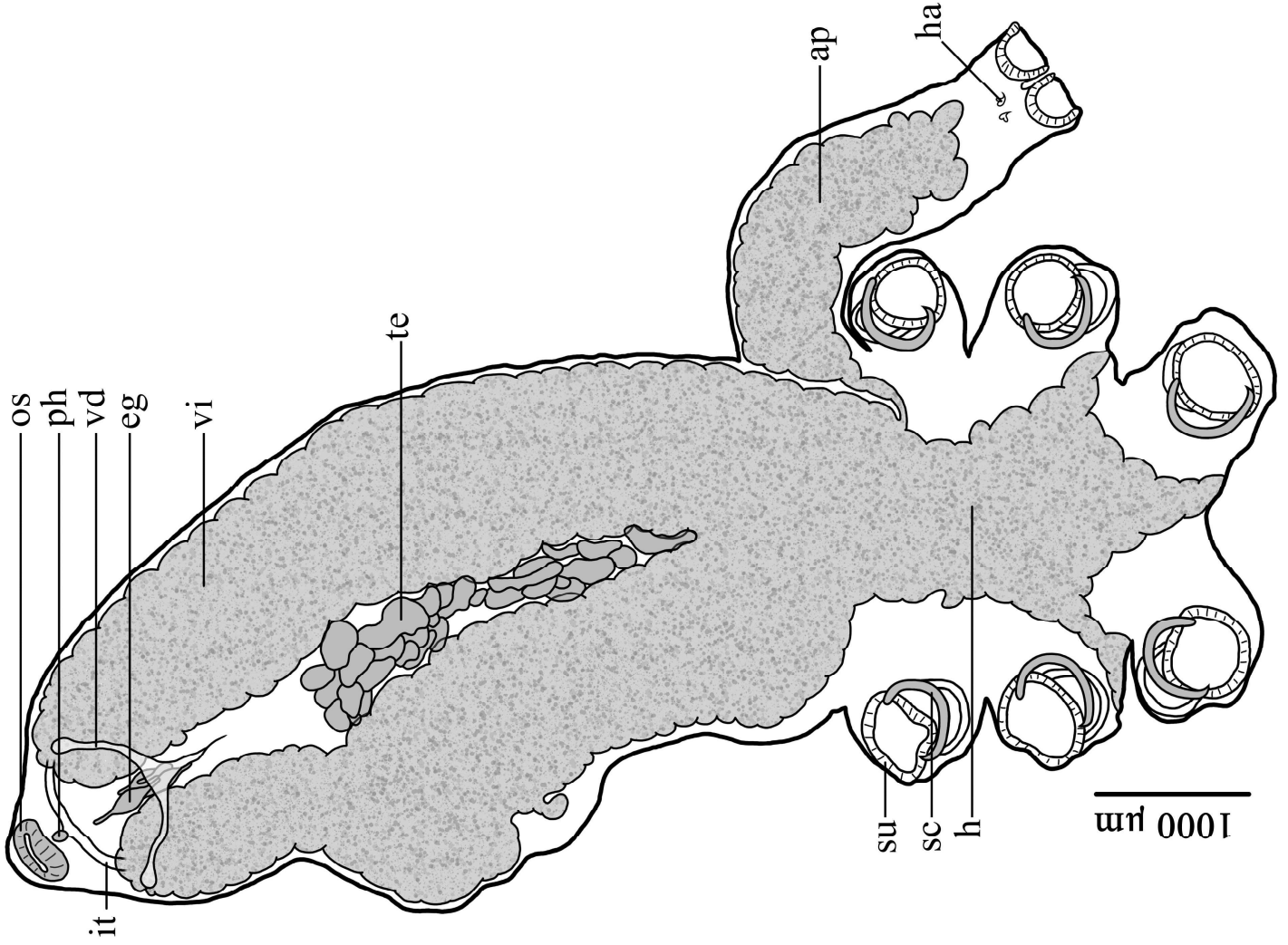
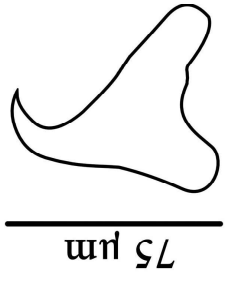
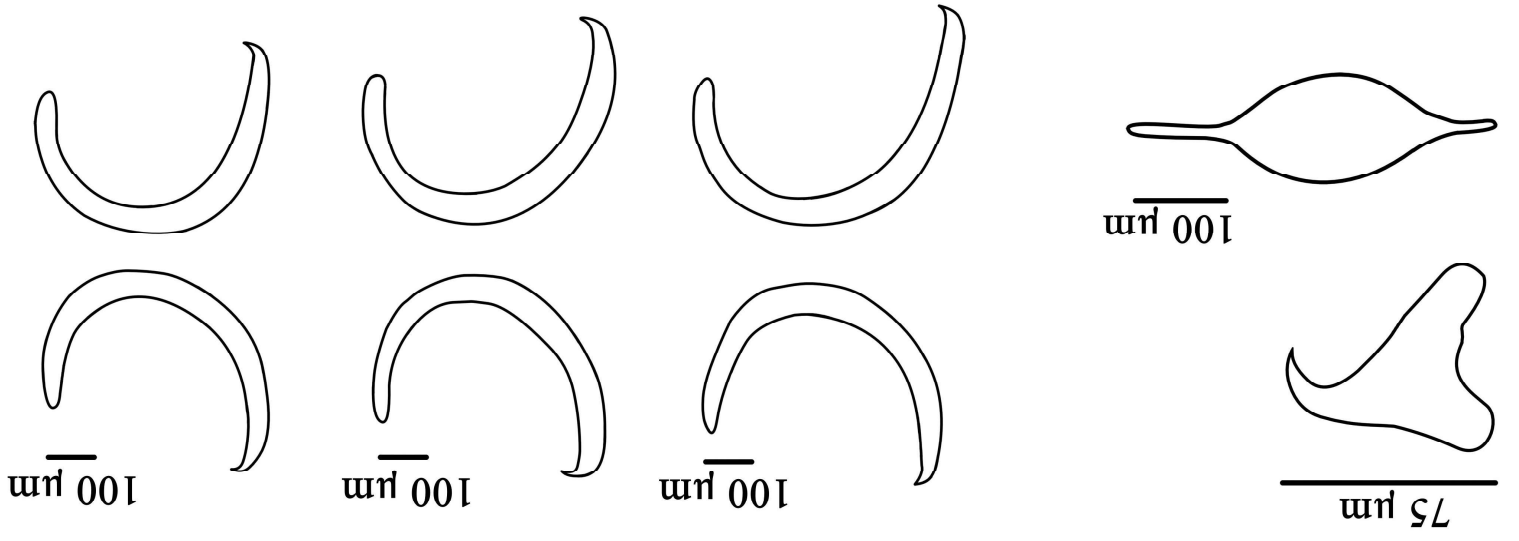


Figure 5

