- 1 Robust phylogenetic position of the enigmatic hydrozoan Margelopsis
- 2 haeckelii revealed within the family Corymorphidae
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- 20 Keywords: Cnidaria, Hydrozoa, Corymorphidae, Margelopsidae, *Margelopsis haeckelii*, 21 Molecular phylogeny
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23 Abstract

24 The life-cycle and polyp morphology of representatives of Margelopsidae are very different from all other species in the hydrozoan clade Aplanulata. Their evolutionary origin and phylogenetic 25 26 position has been the subject of significant speculation. A recent molecular study based only on 27 COI data placed Margelopsidae as a sister group to all Aplanulata, an unexpected result because margelopsid morphology suggests affiliation with Tubulariidae or Corymorphidae. Here we used 28 multigene analyses, including nuclear (18S rRNA and 28S rRNA) and mitochondrial (16S rRNA 29 and COI) markers of the hydroid stage of the margelopsid species Margelopsis haeckelii Hartlaub, 30 31 1897 and the medusa stage of Margelopsis hartlaubii Browne, 1903 to resolve their phylogenetic position with respect to other hydrozoans. Our data provide strong evidence that *M. haeckelii*, the 32 type species of Margelopsis, is a member of the family Corymorphydae. In contrast, M. hartlaubii 33 34 Browne, 1903 is sister to *Plotocnide borealis* Wagner, 1885, a member of Boreohydridae. These results invalidate the family Margelopsidae. The phylogenetic signal of polyp and medusa stages 35 is discussed in light of concept of inconsistent evolution and molecular phylogenetic analysis. 36

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38 Introduction

Species in the family Margelopsidae Mayer, 1910 (Aplanulata, Hydrozoa, Cnidaria) have intriguing life histories. The family is exclusively represented by hydrozoans with holopelagic life-

cycles, where medusae and solitary vasiform polyps float freely throughout the water column. 41 42 Interestingly, siphonophore specialists used margelopsid species as a model to explain the origin of siphonophoran colonies (Totton and Bargmann, 1965). Margelopsidae is comprised of three 43 genera; Margelopsis Hartlaub, 1897; Pelagohydra Dendy, 1902; and Climacocodon Uchida, 44 45 1924, none of which have been sampled for comprehensive molecular analyses. Phylogenetic analysis using only COI sequences (Ortman et al, 2010) of Margelopsis hartlaubii Browne, 1903 46 47 suggested that Margelopsidae might be the sister group to the rest of Aplanulata. However, authors have not recovered strong support for this placement (Nawrocki et al., 2013). The 48 49 systematics and phylogenetic position of Margelopsidae is solely based on insufficient 50 morphological data. Given their polyp morphology, species of Margelopsidae show affinities with Tubulariidae or Corymorphydae, but their unique medusa morphology was used to justify their 51 52 original erection as a separate family. Thus, sampling with more DNA markers and specimens especially including the type species Margelopsis haeckelii - has been needed to determine the 53 54 scope and phylogenetic position of the family Margelopsidae.

55 Despite difficulties of sampling margelopsid hydroids, we were finally able to collect representatives of Margelopsis haeckelii Hartlaub, 1897 and Margelopsis hartlaubii Browne, 1903 56 for molecular studies. Margelopsis haeckelii is the most studied species of its family, yet, 57 58 documented collection records and morphological examinations have been very few (Hartlaub, 1897; Hartlaub, 1899; Lelloup, 1929; Werner; 1955, Schuchert, 2006). Polyps of M. haeckelii 59 closely resemble tubulariid hydranths, having two whorls of tentacles but lacking both a 60 hydrocaulus and stolonal system (Fig. 1, A, B). Free-swimming medusae develop from medusa 61 62 buds located between whorls of polyp tentacles (Fig. 1, B, C, D). Eggs of *M. haeckelii* develop on the manubrium of the medusa (Fig. 1, C, D) and transform directly or through an encysted stage 63 64 into a hydranth that never fixes to a substrate, exhibiting a continuous planktonic lifestyle (Werner; 1955). It is thought that eggs of this species are parthenogenetic, as no male gonads have ever 65 66 been reliably documented. There is less information about *M. hartlaubii*, which is only known from the medusa stage. The medusa of *M. hartlaubii* can readily be distinguished from the medusa of 67 M. haeckelii by its thick apical mesoglea of the bell without apical canal and two tentacles per bulb 68 69 (Fig. 1, C, D, E) (Schuchert, 2006).

In our study we obtained full-length sequences of 18S rRNA and 28S rRNA and partial sequences of the mitochondrial ribosomal 16S rRNA and cytochrome oxidase subunit I (COI) in order to phylogenetically place *M. haeckelii* and *M. hartlaubii* within as comprehensive sampling of hydrozoan taxa as possible. Using this approach, we provide the first molecular evidence that *M. haeckelii* should be placed within the family Corymorphydae. Our findings further showed that the previously sequenced *M. hartlaubii* is a relative of the family Boreohydridae, and is only distantly related to *Margelopsis haeckelii*, the type species of the genus.

77 Methods and materials

78 Animal sampling.

Some *M. haeckelii* polyps were collected in the North Sea (loc. Belgium, Ostend, 51.218028°, 2.879417°) (Fig. 1, F, J). Polyps were collected with a plankton net in the coastal area. Collected animals were used to set up a lab culture. The obtained culture was maintained throughout the year in aquaria using artificial sea water (salt Red Sea Coral Pro, salinity 30–32‰) at the Department of Embryology, Lomonosov Moscow State University, Russia, Moscow. For both polyp and medusa stages, *Artemia salina* nauplii, at least 3 days after hatching, were used for feeding. Animals were fed once a day.

Also, *M. haeckelii* medusae were collected in the Atlantic Ocean, Atlantic Coast of North America
(loc. USA, New York, 40.560556°, -73.882333°). Medusae were collected with a plankton net in
the coastal area, about 10 meters out from the shore. Collected animals were fixed and stored in
96% ethanol (Fig. 1, F, J).

M. hartlaubii DNA was a gift from Dr. Peter Schuchert (Schuchert, 2022). The medusa was collected in Norway, Raunefiord (60.2575°, 05.1393°) with a plankton net from 200 to 0 m depth on 14-JUN-2016.

93 Meiobenthic polyps of *Plotocnide borealis* (formerly known as *Boreohydra simplex*; Pyataeva et 94 al., 2016) were collected in the White Sea near the N.A. Pertsov White Sea Biological Station of 95 the Moscow State University, Kandalaksha Bay, Russia (66.528056°, 33.185556°). Fine mud with 96 polyps was collected with a light hyperbenthic dredge from depth 20-40 m. Collected individuals 97 were fixed and stored in 96% ethanol.

98 Identification of COI, 16S rRNA, 18S rRNA and 28S rRNA sequences.

COI, 16S rRNA, 18S rRNA and 28S rRNA sequence fragments were amplified from genomic
DNA using PCR methods. Genomic DNA was extracted using standard phenol/chloroform
protocols. This method involved tissue digestion with proteinase K (20 mg/mL) in a lysis buffer
(20 mM Tris-CL pH 8.0, 5 mM EDTA pH 8.0, 400 mM NaCl, 2%SDS), extraction with
phenol/chloroform (1:1), precipitation with 0.1 vol 3M Sodium acetate and 1 vol. 100% Isopropanol
and elution in mQ water.

105 For amplification, we used the following primers pairs:

106 16SAR (TCGACTGTTTACCAAAAACATAGC) and 16SBR (ACGGAATGAACTCAAATCATGTAAG) for 16S rRNA (Cunningham and Buss, 1993); and 107 108 iGLCO1490 (TITCIACIAAYCAYAARGAYATTGG) and iGHCO2198 (TAIACYTCIGGRTGICCRAARAAYCA) for COI (Geller et al., 2013). Amplification programs used 109 for 16S rRNA and COI are as previously described in Prudkovsky et al., 2019. 110

111 18S-EukF (WAYCTGGTTGATCCTGCCAGT) and 18S-EukR (TGATCCTTCYGCAGGTTCACCTAC) for rRNA (Medlin et 112 18S al., 1988). F97 113 (CCYYAGTAACGGCGAGT), R2084 (AGAGCCAATCCTTTTCC), F1383 (GGACGGTGGCCATGGAAGT) and R3238 (SWACAGATGGTAGCTTCG) for 28S rRNA (Evans 114 115 et al., 2008). Amplification programs used for 18S rRNA and 18S rRNA are as previously described in Evans et al., 2008. 116

Full-length 18S rRNA and 28S rRNA sequences of *M. haeckelii* from the North Sea were obtained 117 118 from the reference transcriptome available in our laboratory. For transcriptome sequencing, total RNA was extracted from a mixture of various *Margelopsis* life and developmental stages. Total 119 RNA extraction was conducted using the Zymo Research Quick-RNA MiniPrep Plus Kit according 120 121 to the manufacturer's instructions. Poly-A RNA enrichment, cDNA library construction and sequencing were carried out at Evrogen (Russia). The cDNA library was sequenced using the 122 123 Illumina NovaSeq 6000 SP flow cell to produce with 150-bp paired-end reads. The high-quality 124 reads were employed for the *M. haeckelii* transcriptome assembly with the SPAdes assembler 125 (v.3.13.1) (Bankevich et al., 2012).

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127 Phylogenetic analyses

Nucleotide sequences were aligned using the MUSCLE algorithm in MUSCLE software (v3.8.31)
 (Edgar et al., 2004) and trimmed with the TrimAL tool (v.1.2rev59) (Capella-Gutiérrez et al., 2009).
 A heuristic approach "automated1" was used to select the best automatic method for trimming our alignments.

Phylogenetic analyses were performed using Maximum Likelihood methods in IQTree v.2.0-rc2 software (Minh, et al., 2020) according to the optimal models for each gene. Individual marker analyses and a concatenated gene analysis were performed. The best models of nucleotide substitution were chosen using ModelFinder (Kalyaanamoorthy et al., 2017). The GTR+F+I+G4 was found to be optimal for the COI dataset; GTR+F+I+G4 for 16S rRNA; TIM3+F+R3 for 18S
 rRNA; and TIM3+F+R5 for 28S rRNA. One thousand bootstrap replicates were generated for
 each individual analysis, as well as for the combined analysis.

The concatenated COI+16S+18S+28S alignment was constructed using Sequence Matrix (https://github.com/gaurav/taxondna). The concatenated dataset was analyzed using IQTree (v.2.0-rc2) with partitioned analysis for multi-gene alignments (Chernomor, et al., 2016). The set of selected species for concatenated analysis was chosen mainly according to Nawrocki et al. (2013) and considering the availability of individual gene sequences in GenBank for COI, 16S rRNA, 18S rRNA and 28S rRNA.

145 Trees were visualized in FigTree v1.4.4 and processed with Adobe Illustrator CC. No alterations 146 were made to the tree topology or the branch lengths.

An approximately unbiased (AU) test (Hidetoshi, 2002) was performed using IQTree software for
 testing alternative phylogenetic hypotheses.

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150 Data availability

Sequences obtained in this study have been deposited in GenBank under the following accession
numbers: *Margelopsis haeckelii* (OK129327, OK139084, OK142735, OK127861, ON391039,
ON391070), *Margelopsis hartlaubii* (ON237369, ON237671, ON237710), *Plotocnide borealis*(OK110252).

155

156 **Results**

Our phylogenetic investigation of phylogenetic affinities of species of Margelopsidae was 157 conducted employing Maximum likelihood analysis for all single gene datasets as well as our final 158 159 concatenated four-gene dataset (COI, 16S rRNA, 18S rRNA, 28S rRNA). All taxa used in our 160 analysis are arranged taxonomically in Table 1. All *M. haeckelii* and *M. hartlaubii* sequences (COI, 16S rRNA, 18S rRNA, 28S rRNA) were newly generated for this study. M. hartlaubii had 161 previously only had COI available on GenBank (Ortman et al., 2010). Maximum Likelihood 162 163 bootstrapping (MLB) analysis of the concatenated dataset recovered a relatively well resolved tree and recovered Margelopsidae paraphyly. M. hartlaubii was recovered sister to Plotocnide 164 165 borealis Wagner, 1885 (MLB=100), forming a clade that affiliate with the family Boreohydridae, a sister taxon to all other Aplanulata genera (MLB = 100) (Fig. 2). Each individual COI, 16S rRNA, 166 167 18S rRNA or 28S rRNA analysis also recovered a strong supported affiliation of M. hartlaubii 168 within Boreohydridae (MLB = 100) (Fig. 2). At the same time, both *M. haeckelii* from different locations nested within the clade of the Corymorphidae (MLB=89). This clade comprised two 169 subclades, each well supported, one for genus Euphysa, including the type species Euphysa 170 171 aurata Forbes, 1848, and the other for Corymorpha + M. haeckelii, including the type species, 172 Corymorpha nutans M. Sars (Fig 2). M. haeckelii is nested inside the clade Corymorpha bigelowi Maas, 1905, Corymorpha nutans M. Sars, 1835, Corymorpha sarsii Steenstrup, 1855 and 173 Corymorpha pendula L. Agassiz (MLB=89). Clade Euphysa+Corymorpha+M. haeckelii was 174 recovered to be the sister to Tubulariidae (MLB=85), which together with Branchiocerianthus 175 176 imperator Allman, 1885 constitute the superfamily Tubularioidea. Tubularioidea is recovered as sister to Hydridae (MLB=91). General topology of our phylogenetic tree obtained in combined 177 178 analysis coincides with the Aplanulata tree published by Nawrocki et al., 2013.

Separate COI and 16S rRNA analysis recovered, that individuals of *Margelopsis haeckelii* from
the opposite sides of the Atlantic Ocean are representatives of the same species (Fig. 1S, 2S).
No nucleotide substitutions were identified in analyzed sequences of *Margelopsis haeckelii* from
the waters of Belgium (51.218028°, 2.879417°) and the USA (40.560556°, -73.882333°).

At the same time, *M. hartlaubii* COI sequences analysis revealed five mismatches between sequences obtained in this study (ON237369) and sequence published in Ortman et al., 2010 (GQ120058.1) (Fig. 1S). However, COI sequences of *M. hartlaubii* published in Ortman et al., 2010 (GQ120058.1 and GQ120059.1) also are not identical and have three mismatches.

Phylogenetic hypothesis testing (AU test) was performed to test the statistical significance of tree
topologies in our Maximum Likelihood analysis. The AU test rejected the phylogenetic hypothesis
of the monophyly of *M. haeckelii* and *M. harlaubii*, providing strong evidence for the polyphyly of *Margelopsis*. Also, as our two individual marker analyses (16S and 28S) (Supp. 2, 3) placed *M. haeckelii* as a sister to *Corymorpha*, two hypotheses of alternative placements of *M. haeckelii*were evaluated: *M. haeckelii* is inside or outside *Corymorpha*. Results of the testing significantly
support (p < 0.05) the hypothesis that *M. hackelii* is within *Corymorpha*. (Fig. 5S).

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195 Discussion

Our concatenated dataset (COI+16S+18S+28S), which included a comprehensive taxonomic 196 sampling of hydrozoans, recovered Margelopsis haeckelii within Corymorphidae, nested within a 197 198 clade consisting of several Corymopha species. This result is consistent with previous findings 199 based solely on polyp morphology, where Margelopsidae was grouped with Tubulariidae and Corymorphidae in the superfamily Tubularoidea (Rees, 1957). Being quite small (1-2 mm), 200 201 hydrocaulus-lacking pelagic polyps of the Margelopsidae are similar to those sessile polyps of 202 corymophids and tubulariids despite the latter having a well-developed hydrocaulus and reaching 203 up to ten centimeters in height. For all three families, hydranth tentacles are arranged into two, 204 oral and aboral whorls and blastostyles are situated in the inter-tentacular region (Fig. 3, A, C). 205 Our phylogenetic data support assertions that polyp tentacle patterns may be an important 206 morphological character for identifying lineages in Aplanulata (Rees, 1957, Nawrocki et al. 2013).

207 Interestingly, *M. haeckelii* jellyfish are atypical in having radial symmetry, which more usually is 208 bilateral in Aplanulata. The *M. haeckelii* jellyfish has 3-4 tentacles per bulb instead of one long 209 tentacle per medusa, something typically seen among Corymorpha medusae. Even in Euphysa, 210 the sister group to Corymorpha, radially symmetric adult medusae develop asymmetrically in 211 contrast to medusae of *M. haeckelii*. The medusae of *Euphysa flammea* Hartlaub, 1902 only have 212 a single tentacle in their youngest stage, with a second, third and fourth being added successively 213 over time (Schuchert, 2010). Radially symmetric medusae in the species *P. borealis*, which is 214 deeply nested in our phylogenetic analyses of Aplanulata (Pyataeva et al., 2016; this study) 215 suggests that radial symmetry has re-evolved in *M. haeckelii*, a manifestation of the original body 216 plan symmetry for medusae of Aplanulata. The presence of an apical canal in the umbrella may be a phylogenetically significant character warranting further investigation, as this character is 217 218 shared both by *M. haeckelii* and all *Corymorpha* medusae (Fig. 3, A, C, marked orange). 219 Reproductive characters appear to also reflect phylogenetic relationships in Aplanulata. Among 220 all of Tubularoidea, only Corymorpha embryos undergo encystment similar to that of M. haeckelii 221 (Petersen, 1990).

222 Surprisingly, our concatenated gene dataset, as well as our single gene COI dataset, recovered the medusa known as *M. hartlaubii* to be a close relative of *Plotocnide borealis*, and not closely 223 224 related to *M. haeckelii* nor group within Corymorphidae. This result is further supported by 225 independent morphological data showing several similarities between medusae of M. hartlaubii and P. borealis, including thick apical mesoglea of the bell (Fig. 3, marked blue), lack of an 226 227 umbrella apical canal and nematocyst batteries being located at the distal parts of tentacles (Fig. 228 3, marked violet) (Schuchert, 2006). Based on our findings, medusae described by Browne (1903) 229 have been wrongly attributed to the genus Margelopsis. Nawrocki et al. (2013) suggested that the 230 hypothesis of *M. hartlaubii* as the sister to the rest of Aplanulata was uncertain due to low bootstrap support and that more genetic markers were needed to understand the phylogenetic 231

placement of the species. Based on our multi-marker phylogenetic analysis and morphological
data (Browne, 1903; Schuchert, 2006) we hypothesize that *M. hartlaubii* has a mud-dwelling,
meiobenthic polyp like *P. borealis* (Fig. 3), and that the two species combined represent the sister
group to the rest of Aplanulata.

In addition to *M. haeckelii* and *M. hartlaubii*, there are several other suspected species in the 236 237 genus Margelopsis, including Margelopsis gibbesii (McCrady, 1859) and Margelopsis australis 238 (Browne, 1910). Following Schuchert (2007), the World Register of Marine Species (https://www.marinespecies.org/) lists Margelopsis gibbesii as invalid. This stems from the fact 239 240 that the original material used to describe this species, as Nemopsis gibbesii, consisted of a 241 margelopsid polyp and a bougainvilliid medusa, the latter subsequently recognized as a medusa 242 of Nemopsis bachei (L. Agassiz, 1862). This situation has generated subsequent nomenclatural confusion. More recently, Calder and Johnson (2015) stabilized the situation by designating the 243 244 hydroid specimen illustrated by McCrady (1859) in Plate 10, Figure 7 as a lectotype for the 245 margelopsid species. Calder and Johnson (2015) went on to provide evidence casting doubt on the distinction between M. gibbesii and M. haeckelii but maintained the two species given the 246 247 geographic locations on either side of the north Atlantic and pending further study. In this study, 248 however, using molecular phylogenetics, we have shown that Margelopsis from the western North Atlantic, and M. haeckelii from the eastern North Atlantic is the same species as M. haeckelii, 249 250 Margelopsis gibbesii invalid. The lack of any nucleotide substitution in COI and 16S sequences of Margelopsis representatives from both sides of Atlantic Ocean makes it possible to suggest 251 252 that these two populations are not isolated.

Margelopsis australis is only known from its original collection and is based on a single medusa specimen, lacking reliable characters for distinguishing it from *M. hartlaubii* (Browne 1910). Moreover, the single specimen was described as being "somewhat contracted and in a crumbled condition" (Browne 1910). Based on the available morphological data, we cannot state with any degree of certainty that *M. australis* is a valid species, or that it is a member of *Margelopsis*.

258 Medusae are a useful means of identifying species, genera and even family ranks (Rees, 1957; 259 Bouillon, et al., 2006). A change in morphology of the typical jellyfish form within a family is usually due to the reduction of the medusa stage, something that is widespread throughout 260 Anthoathecata and Leptothecata (Cornelius, 1992; Leclere et al., 2009; Cartwright, Nawrocki, 261 262 2010). However, M. haeckelii is a normally developed medusa, distinctly different from those typical of *Corymorpha*, despite their close relationship recovered by our phylogenetic analysis. 263 264 Recent studies using molecular phylogenetic methods have revealed several such cases in which 265 related taxa have very different jellyfishes or species with similar jellyfishes are only distantly related. The morphologically aberrant jellyfish Obelia is so different from other Companulariidae 266 267 that a hypothesis was proposed for the re-expression of this jellyfish after its evolutionary reduction (Boero, Sara, 1987). However, this hypothesis was not supported by molecular 268 phylogenetic analysis and Obelia may have originated from a Clytia-like ancestor (Cunha et al. 269 270 2017; Govindarajan et al., 2006; Leclere et al., 2019). Larsonia pterophylla (Haeckel, 1879) was previously assigned to the genus Stomotoca due to similarity of their jellyfishes (Larson, 1982). 271 272 Interestingly, the structure of the polyps in the genera *Larsonia* and *Stomotoca* are so dissimilar 273 that they could be attributed to different families (Boero, Bouillon, 1989). And indeed, according 274 to molecular data, L. pterophylla and Stomotoca atra L. Agassiz, 1862 are not closely related. Rather, L. pterophylla is closely related to Hydrichthys boycei from the Pandeidae family, and S. 275 276 atra is ungrouped with most species (Schuchert, 2018; Woodstock et al., 2019). Inclusion of the 277 genus Cytaeis in Bougainvilliidae or the genera Polyorchis and Scrippsia in Corynidae is 278 surprising due to the discrepancy between the jellyfishes of these genera and those typical of the 279 respective families (Nawrocki, Cartwright, 2010; Prudkovsky et al., 2017). Finally, we conclude 280 that appearance of atypical jellyfishes in hydrozoan families can indicate a great evolutionary 281 plasticity of the medusa stage morphology. In contrast, the morphology of the hydroids appear to be more phylogenetically constant. For example, the morphology of Cytaeis hydroids is similar to 282

the structure of Bougainvillidae hydroids with stolonal colonies, and Obelia-like polyps are typical for the family Campanulariidae (Prudkovsky et al., 2017; Leclere et al., 2019).

285 Concepts of 'mosaic' or 'inconsistent evolution' were proposed for these cases in which closely 286 related hydroids can produce very different medusae or vice versa (Naumov 1956, 1960; Rees, 287 1957). Inconsistent evolution was explained by differences in the rate and direction of evolution 288 in the two life cycle stages. Some incongruences between hydroid and medusa systems seem to 289 result from weaknesses in a classification system (Petersen, 1990), but our work provides new 290 reason to return to the discussion of this concept.

291 **Taxonomic recommendations**

Based on our results, as well as a number of previous studies, we formally recommend the following changes to the taxonomy of Margelopsidae and its component species:

- a) As multigene phylogenetic analyses nested *Margelopsis haeckelii*, the type species of
 Margelopsis, within genus *Corymorpha*, we recommend to redesignate it into *Corymorpha haeckelii*.
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298 Corymorpha M. Sars, 1835

299 Type species: *Corymorpha nutans* M. Sars, 1835 by monotypy.

- Diagnosis: Solitary hydroids with more or less vasiform hydranth, with long caulus or with 300 301 short, squat polyp with broad head. Rarely a hydrant without a caulus. Hydranth with 302 one or several closely set whorls of 16 or more moniliform or filiform tentacles and one 303 or more aboral whorls of 16 or more long, non-contractile moniliform or filiform tentacles. 304 Gastrodermal diaphragm parenchymatic or without parenchymatic specializations of the gastrodermis. Hydrocaulus, if present, stout, covered by thin perisarc, filled with 305 306 parenchymatic gastrodermis, with long peripheral canals; aboral end of caulus with papillae turning more aborally into rooting filaments, rooting filaments scattered or 307 gathered in a whorl, rooting filaments composed of epidermis and solid gastrodermis, 308 sometimes tips with non-ciliated statocysts. Otherwise, hydroid planktonic and 309 310 hydrocaulus reduced, with a central depression. With or without asexual reproduction through constriction of tissue from aboral end of hydrocaulus. Gonophores develop on 311 312 blastostyles arranged in a whorl over aboral tentacles. Gonophores remain either fixed as sporosacs, medusoids, or are released as free medusae. Medusa bell apex dome-shaped 313 or pointed, with apical canal. Four marginal bulbs present, lacking long exumbrellar 314 315 spurs. With a single tentacle or three short tentacles and one long tentacle that differs not merely in size, but also in structure. Rarely with 1-6 tentacle per bulb. Manubrium thin-316 317 walled, sausage-shaped with flared mouth rim, reaching to umbrella margin. Cnidome 318 comprises stenoteles, desmonemes, and haplonemes, with or without euryteles. 319 Remarks: This diagnosis for the most part corresponds to Schuchert, 2010 (Schuchert,
- 2010), Petersen, 1990 (Petersen, 1990) and Nawrocki et al., 2013 (Nawrocki et al., 2013),
 but with modifications (indicated in bold) to polyp and medusa body shape, and cnidome
 description to include *Margelopsis (Corymorpha) haeckelii.*
 - b) We suggest moving *Margelopsis hartlaubii* into family Boreohydridae and recommend to redesignate it into Plotocnide *hartlaubii*.
- 327 *Plotocnide* Wagner, 1885

328 Type species: Plotocnide borealis Wagner, 1885 by monotypy.

Diagnosis: Medusa umbrella evenly rounded with thick apical jelly and scattered groups of exumbrellar nematocysts; manubrium half as long as bell cavity, with or without broad, dome-shaped apical chamber; **without apical canal**; mouth simple, with ring of nematocysts; gonad forming thick ring around manubrium; four narrow radial canals and

narrow ring canal; four marginal bulbs each with 1-3 solid tentacles per bulb; tentacles 333 334 terminate in ovoid knob studded with nematocysts. No ocelli. Cnidome comprises desmonemes and stenoteles, with or without mastigophores. Hydroids, if known, 335 solitary, small, with one whorl of reduced tentacles, capitate or not, located in the oral or 336 337 median part of body; perisarc covering of base filmy or absent; gametes in body wall. Remarks: This diagnosis for the most part corresponds to Schuchert, 2006, 2010 338 (Schuchert, 2006; Schuchert, 2010), but with modifications (indicated in bold) to medusa 339 body shape, and cnidome description to include Margelopsis (Plotocnide) hartlaubii. 340

- 341 342
- c) We suggest that Margelopsidae should no longer be used, and both *Pelagohydra* and
 Climacocodon should be moved to Aplanulata *incertae sedis* until additional molecular
 phylogenetic analyses can clarify their phylogenetic placement.

345 **Conclusion**

Our results clarify the phylogenetic picture of Aplanulata, by revealing the phylogenetic position 346 of M. haeckelii, type species of the genus Margelopsis as falling within Corymorpha and M. 347 hartlaubii as being a close relative of *Plotocnide* in the family Boreohydridae. On the case of the 348 349 latter species, this phylogenetic result conflicts with the century old hypothesis that Margelopsis 350 belongs to Tubulariidae or Corymorphidae (Nawrocki et al., 2013). However, by showing that M. 351 haeckelii falls within the genus Corymorpha, our investigation presents strong evidence in support of this traditional hypothesis. Because *M. haeckelii* is a hydrozoan belonging to Corymorphidae, 352 353 we can infer that this lineage evolutionarily lost their hydrocaulus and stolon, likely as an adaptation to a holopelagic life-cycle. It was previously suggested that the foundation for this type 354 355 of changes in body plan, and accompanying life-style, might lead to speciation and could be 356 reflected by changes in the expression of Wnt signaling components (Duffy, 2011). Based on our 357 results, *M. haeckelii* might be a prime candidate for testing this hypothesis.

Unfortunately, due to the few and extremely irregular documented collection records of hydroids from the supposedly sister genera of *Margelopsis*, *Pelagohydra* and *Climacocodon*, the phylogenetic relationships within this group are still obscured. It remains unclear if Pelagohydra and Climacodon form a clade with either M. hartlaubii or M. haeckelii, or neither. Thus, the number of origin of a secondarily specialized pelagic polyp stage is still not known. The possible relationships between these three genera, as well as their phylogenetic placement, still need to be verified by additional studies when molecular data become available.

365 Acknowledgements

We thank Dr. Peter Schuchert for the gift of *Margelopsis hartlaubii* DNA. We are grateful to Dr. Allen G. Collins for sequencing of the COI and 16S of *Margelopsis haeckelii* from NY (USA). We also thank Dr. Brett Gozales for the help with text and grammar editing. This study was supported by federal project 0088-2021-0009 of the Koltzov Institute of Developmental Biology of the Russian Academy of Sciences. This work was also supported by RSF, grant number 22-14-00116.

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506 Figure legends

Fig. 1. Morphology of collected Margelopsidae representatives and the locations of its samplings.
(A-D) *Margelopsis haeckelii* Hartlaub, 1897. (A) Newly hatched polyp, (B) Mature polyp with
medusa buds, (C, D) Mature medusa. (E) Mature medusa of *Margelopsis hartlaubii* Browne, 1903.
Photo Credit: Dr. Peter Schuchert (Schuchert, 2022). (F, J) Geographic locations of sampling
sites. Abbreviations, ac – apical canal, at – aboral tentacles, e – embryos, h – hypostome, md –
medusoid ot – oral tentacles, tb – tentacle bulb, yp – young polyp.

Fig.2. Analysis of phylogenetic position of *Margelopsis haeckelii* and *Margelopsis hartlaubii* in Aplanulata. Phylogenetic hypothesis of *Margelopsis haeckelii* relationships based on the combined mitochondrial and nuclear dataset (CO1+16S+18S+28S). Node values indicate bootstrap support from 1000 replicates. *Margelopsis haeckelii* and *Margelopsis hartlaubii* are in red. *WGS84 51.218028°, 2.879417°, ** WGS84 40.560556°, -73.882333°

Fig. 3. Comparison of morphological characters of (A) *Margelopsis hartlaubii*, (B) *Margelopsis haeckelii*, (C) *Corymorpha nutans* and (D) *Plotocnide borealis*. Scalebar – 0.4 mm. Color coding:
yellow – oral and aboral whorls of polyp tentacles, pink– region of medusa budding, green – the
region of gametes formation, orange – apical canal, blue – medusa umbrella with clusters of
exumbrellar nematoblasts, violet – clusters of nematocysts located at the distal parts of tentacles. *Margelopsis hartlaubii, Margelopsis haeckelii, Corymorpha nutans* and *Plotocnide borealis*modified from Schuchert (2006; 2010)

Fig. 1S. Phylogenetic hypothesis of *Margelopsis haeckelii* and *Margelopsis hartlaubii* relationships based on nuclear cytochrome oxidase subunit I (COI). Node values indicate bootstrap support from 1000 replicates. *Margelopsis haeckelii* and *Margelopsis hartlaubii* are in red.

Fig. 2S. Phylogenetic hypothesis of *Margelopsis haeckelii* and *Margelopsis hartlaubii* relationships based on the mitochondrial 16S rRNA. Node values indicate bootstrap support from
 1000 replicates. *Margelopsis haeckelii* and *Margelopsis hartlaubii* are in red.

Fig. 3S. Phylogenetic hypothesis of *Margelopsis haeckelii* and *Margelopsis hartlaubii* relationships based on the 28S rRNA large ribosomal subunit. Node values indicate bootstrap support from 1000 replicates. *Margelopsis haeckelii* and *Margelopsis hartlaubii* are in red.

Fig. 4S. Phylogenetic hypothesis of *Margelopsis haeckelii* and *Margelopsis hartlaubii* relationships based on the 18S rRNA small ribosomal subunit. Node values indicate bootstrap support from 1000 replicates. *Margelopsis haeckelii* and *Margelopsis hartlaubii* are in red.

538 Fig. 5S. Testing of the phylogenetic hypotheses with AU test.

Table 1. List of the species included in the study and corresponding GenBank accession numbersof all analyzed sequences.

541

suborder	family	species	16S rRNA	18S rRNA	28S rRNA	COI	vouchers
Aplanulata	Boreohydridae	Plotocnide borealis	KU721822. 1	KU721833 .1	OK11025 2	KU721812.1	RU087.2
	Candelabridae	Candelabrum cocksii	EU876535. 1	AY920758 .1	AY920796 .1	GU812438.1	MHNGINVE29591
	Corymorphyda e	Branchioceriant hus imperator		JN594046. 2	JN594035. 2	JX121580.1	MHNG:INVE 74105
		Corymorpha bigelowi	EU448099	EU876564 .1	EU272563 .1	JX121581.1	KUNHM 2829

	Corymorpha nutans	EU876532.	EU876558 .1	EU879931 .1	JX121586.1	MHNG:INVE 48745
	Corymorpha	EU876538.	EU876565	EU305510	JX121583.1	KUNHM DIZ2962
	pendula	1	.1	.1		
	Corymorpha sarsii	KP776787. 1	JN594049. 2	JN594038. 2	JX121585.1	MHNG:INVE 68950
	Euphysa aurata	EU876536. 1	EU876562 .1	EU879934 .1	JX121587.1	MHNG:INVE 48753
	Euphysa intermedia	EU876531. 1	AY920759 .1	EU879930 .1	JX121582.1	
	Euphysa japonica	KP776802. 1	EU301605 .1	JX122505. 1	MF000498.1	
	Euphysa tentaculata	EU876537. 1	EU876563 .1	EU879935 .1	JX121588.1	
	Hataia parva	JN594033. 1	JN594045. 2	JN594034. 2	JX121608.1	UF:5407
Hydridae	Hydra hymanae	GU722762. 1	JN594051. 2	JN594040. 2	GU722849.1	
	Hydra oligactis		JN594052. 2	JN594041. 2	GU722871.1	
	Hydra utahensis		JN594053. 2	JN594042. 2	GU722861.1	
	Hydra vulgaris	EU876543. 1	JN594054. 2	JN594043. 2	GU722914.1	
	Hydra viridissima		EU876569 .1	EU879940 .1	GU722845.1	
Margelopsidae	Margelopsis	OK129327	OK139084	OK142735	OK127861	
	haeckelii	ON391070			ON391039	
	Margelopsis	ON287278	ON237671	ON237710	ON237369	
	hartlaubii				GQ120058.1	
Protohydridae	Protohydra leuckarti	KU721828. 1	KU721835 .1		KU721813.1	Protohydra2010072 7.6
Tubuldariidae	Ectopleura crocea	EU876533. 1	KF699111. 1	EU879932 .1	JX121589.1	MHNG:INVE 34010
	Ectopleura dumortierii	FN687542. 1	EU876561 .1	EU879933 .1	JX121590.1	
	Ectopleura larynx		EU876572 .1	EU879943 .1	JX121591.1	MHNG-INVE- 54563
	Ectopleura marina	EU883542. 1	EU883547 .1	EU883553 .1	JX121592.1	
	Ectopleura wrighti	FN687541. 1	JN594055. 2	JN594044. 2	JX121593.1	MHNG:INVE 27331

		Hybocodon chilensis	EU876539. 1	EU876566 .1	EU879937 .1	JX121594.1	MHNG:INVE 36023
		Hybocodon prolifer	FN687544. 1	EU876567 .1	EU879938 .1	JX121595.1	
		Hydractinia sp	EU305477. 1	EU305495 .1	EU305518 .1		KUNHM2876
		Ralpharia gorgoniae	EU305482. 1	EU272633 .1	EU272590 .1	GU812437.1	KUNHM2778
		Tubularia indivisa	FN687530. 1	EU876571 .1	EU879942 .1	JX121596.1	
		Zyzzyzus warreni	EU305489. 1	EU272640 .1	EU272599 .1	JX121597.1	KUNHM 2777
Capitata	Asyncorynidae	Asyncoryne ryniensis	EU876552. 1	EU876578 .1	GQ424289 .1		KUNHM 2639
	Cladocorynida e	Cladocoryne floccosa	AY512535. 1	EU272608 .1	EU272551 .1		personal:A. Lindner:AL1407
	Cladonematida e	Staurocladia vallentini	GQ395332. 1	GQ424322 .1	GQ424293 .1	MF000500.1	Sch522
		Staurocladia wellingtoni	AY787882. 1	GQ424323 .1	EU879948 .1	MF000486.1	
	Corynidae	Coryne uchidai	GQ395319. 1	GQ424332 .1	GQ424305 .1	KT981912.1	
		Sarsia tubulosa	EU876548. 1	EU876574 .1	EU879946 .1		MHNGINV35763
		Stauridiosarsia ophiogaster	EU305473. 1	EU272615 .1	EU272560 .1		KUNHM 2803
	Moerisiidae	Odessia maeotica	GQ395324. 1	GQ424341 .1	GQ424314 .1		MHNG INVE53642
	Pennariidae	Pennaria disticha	AM088481 .1	GQ424342 .1	GQ424316 .1		MHNG INVE29809
	Porpitidae	Porpita porpita	AY935322. 1	GQ424319 .1	EU883551 .1	LT795124.1	RM3_747
	Solanderiidae	Solanderia secunda	EU305484. 1	AJ133506. 1	EU305533 .1	JX121599.1	KUNHM 2611
	Zancleidae	Zanclea costata	EU876553. 1	EU876579 .1	EU879951 .1		MHNGINV26507
		Zanclea prolifera	EU305488. 1	EU272639 .1	EU272598 .1		KUNHM 2793
Fillifera	Eudendriidae	Eudendrium capillare	AY787884. 1		EU305514 .1	JX121602.1	KUNHM2625
	Proboscidactyli dae	Proboscidactyla flavicirrata	EU305480. 1	EU305500 .1	EU305527 .1	JX121600.1	USNM:1074994

Ptilocodiidae	Hydrichthella epigorgia	EU305478. 1	EU272622 .1	EU272569 .1	JX121601.1	KUNHM 2665
Stylasteridae	Lepidopora microstylus	EU645329. 1	EU272644 .1	EU272572 .1	JX121603.1	USNM:1027724

Figure 1

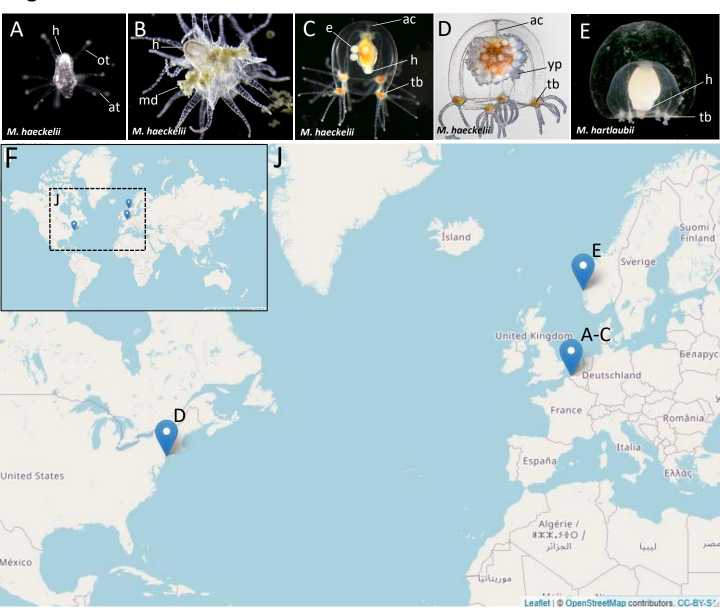
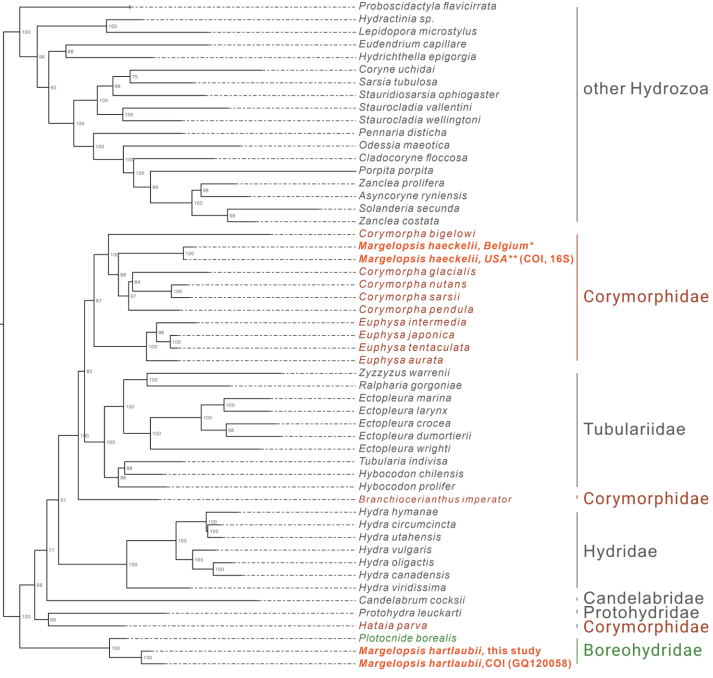


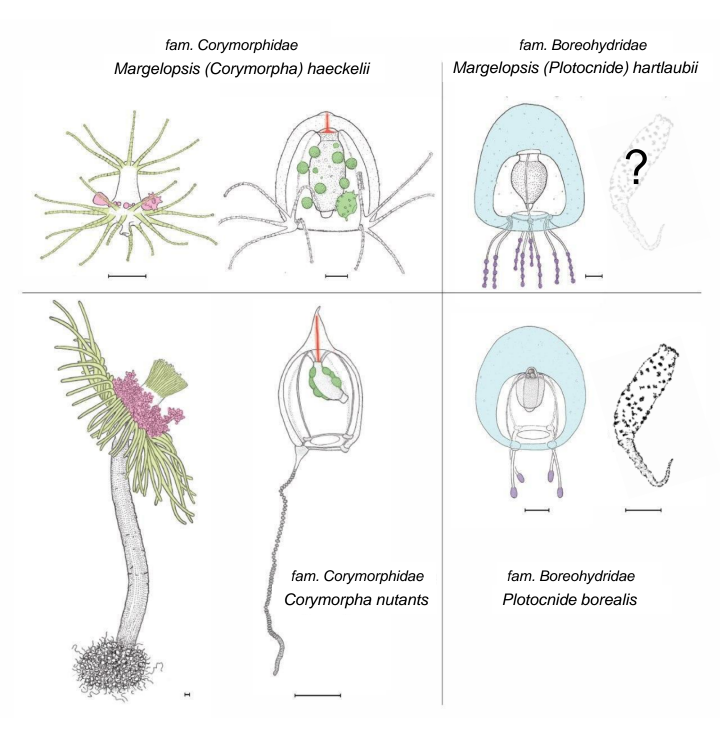
Figure 2

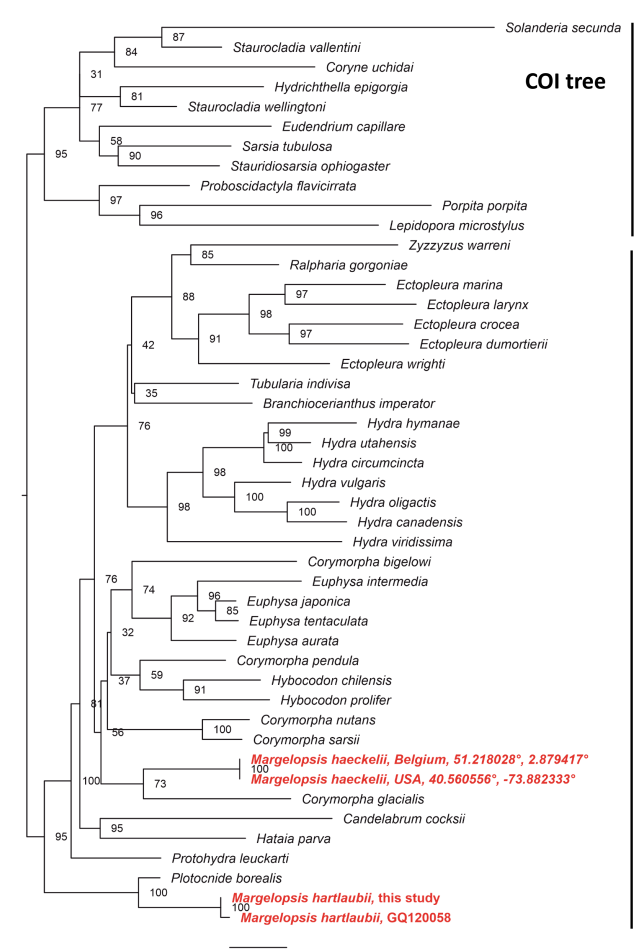


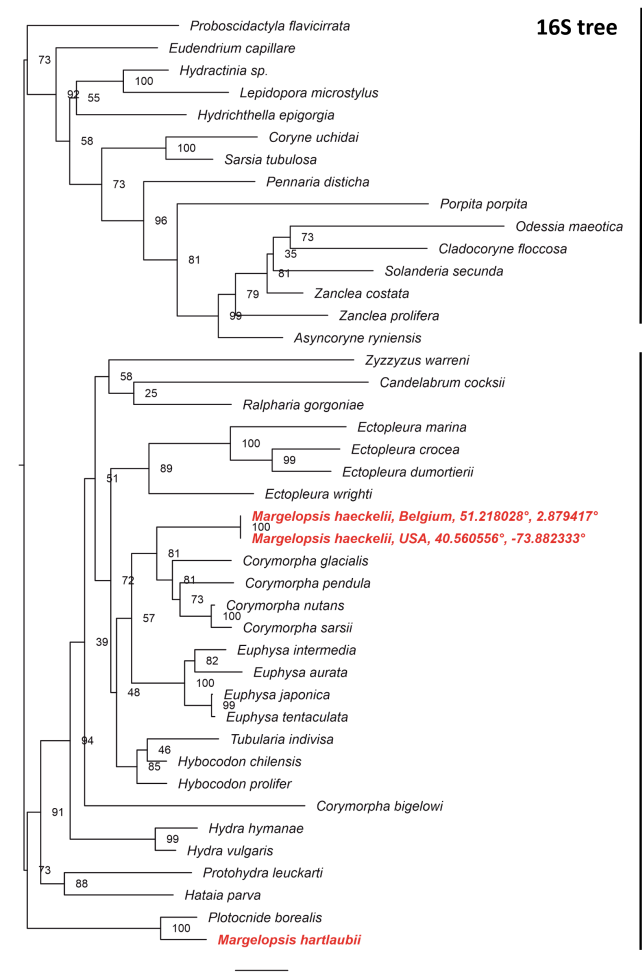
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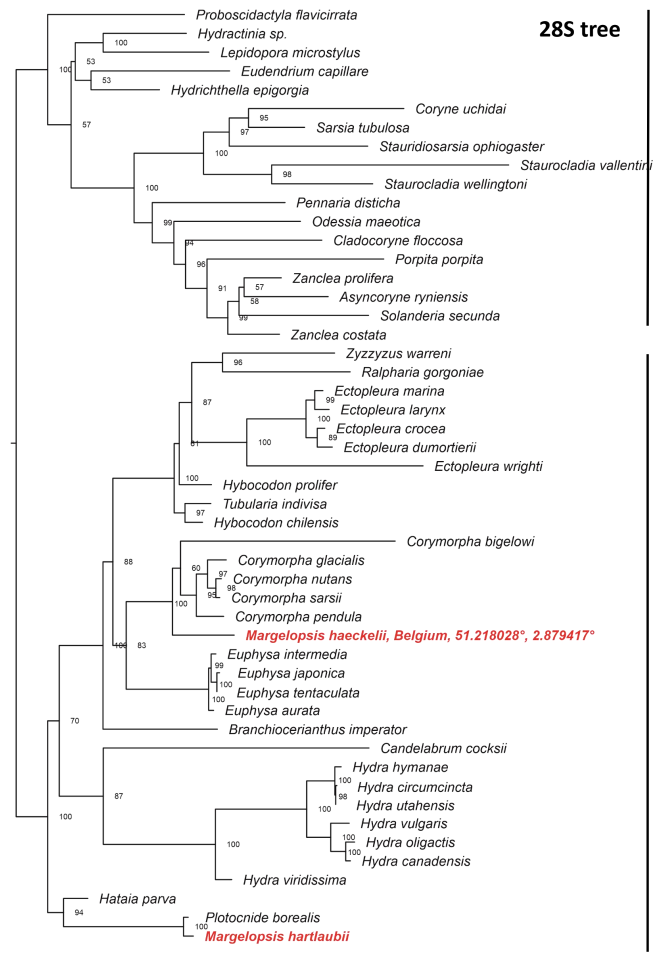
Figure 3



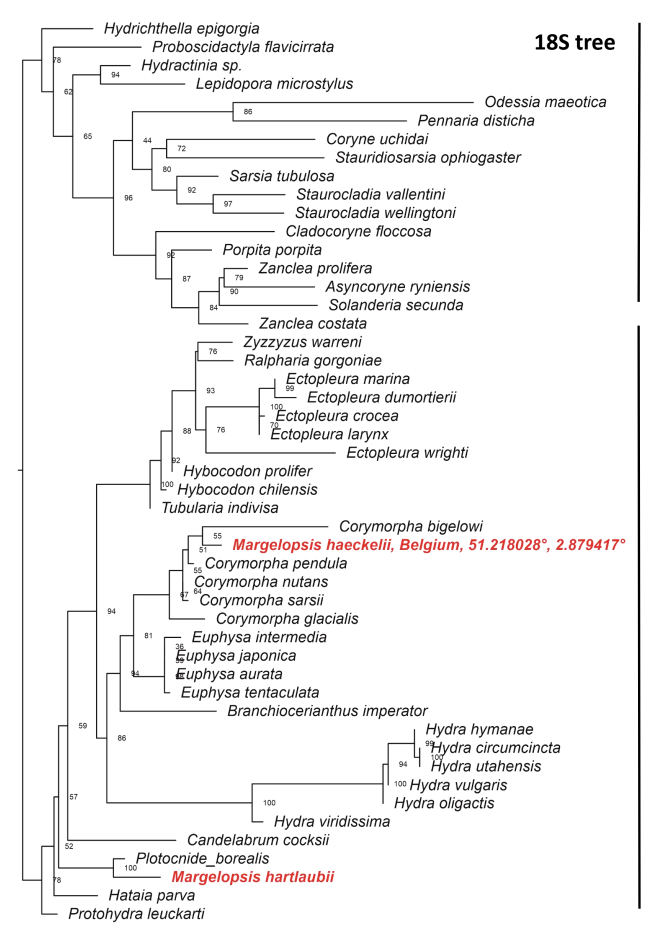




Aplanulata



Aplanulata

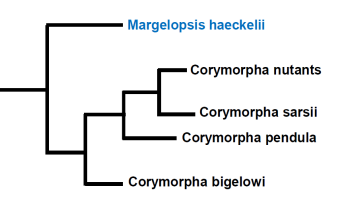


other Hydrozoa

Aplanulata

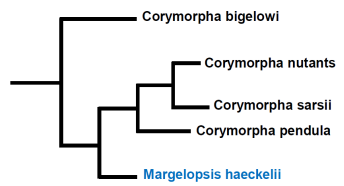
Phylogenetic hypothesis #1

Margelopsis haeckelii is a sister to the genus Corymorpha, AU test



Phylogenetic hypothesis #2

Margelopsis haeckelii nested within the genus Corymorpha, AU test



Rejected, **p** > 0.05

Supported, p < 0.05