



Molecular circumscription of new species of *Gyrocotyle* Diesing, 1850 (Cestoda) from deep-sea chimaeriform holocephalans in the North Atlantic

Rodney A. Bray · Andrea Waeschenbach · D. Timothy J. Littlewood ·
Odd Halvorsen · Peter D. Olson

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Abstract Chimaeras, or ratfishes, are the only extant group of holocephalan fishes and are the sole host group of gyrocotylidean cestodes, which represent a sister group of the true tapeworms (Eucestoda). These unique, non-segmented cestodes have been known since the 1850s and multiple species and genera have been erected despite a general agreement that the delineation of species on the basis of morphology is effectively impossible. Thus, in the absence of molecular studies, the validity of gyrocotylid taxa

and their specific host associations has remained highly speculative. Here we report the presence of *Gyrocotyle* spp. from rarely-caught deep-sea chimaeras collected in the North-East Atlantic, and describe two new species: *G. haffii* n. sp. from the bent-nose chimaera, *Harriota raleighana* Goode & Bean, and *G. discoveryi* n. sp. from the large-eyed rabbit fish, *Hydrolagus mirabilis* (Collett). Nuclear ribosomal sequence data were generated for individual parasites taken from different host species collected on different dates and from different localities and were combined with previously published sequences. Phylogenetic analyses supported the recognition of independent lineages and clusters, indicative of species, but were indecisive in recovering the root of the tree in analyses that included non-gyrocotylid outgroup taxa. The molecular data reveal variation not reflected in morphology and point to a complex picture of genetic divergence shaped by both isolation and migration in the deep-sea environment.

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R. A. Bray · A. Waeschenbach · D. T. J. Littlewood ·
P. D. Olson (✉)
Division of Parasites and Vectors, Department of Life
Sciences, Natural History Museum, London SW7 5BD,
UK
e-mail: P.Olson@nhm.ac.uk

O. Halvorsen
Natural History Museum, University of Oslo,
P.O. Box 1172, Blindern, 0318 Oslo, Norway

Introduction

Holocephalans are deep-sea, cartilaginous fish of the chondrichthyan subclass Holocephali and its only order, the Chimaeriformes, commonly known as rat fishes or ghost sharks. Although highly successful in the Palaeozoic, resulting in a rich palaeontological fauna, the group is now represented by only 39 species in five genera (Inoue et al., 2010). Molecular studies confirm that the group is the sister taxon of the Elasmobranchii and three families are recognised, the Callorhynchidae, Chimaeridae and Rhinochimaeridae, where the former is the sister group to the latter two (Inoue et al., 2010). According to a relaxed molecular clock method employed by Inoue et al. (2010), the Holocephali arose in the Silurian Period (c.410–447 Ma), the Callorhynchidae diverged from its sister group in the Jurassic Period (c.161–190 Ma) and the other families diverged in the mid-Cretaceous Period (c.98–146 Ma). Licht et al. (2012) expanded the representation of holocephalans and reported similar results, with the group diverging between the late Silurian and the early Devonian. This ancient and distinctive host group harbours similarly distinctive parasites, and in this study, we report on those of the cestode order Gyrocotylidea.

Gyrocotylideans are non-segmented tapeworms known together with the Amphilinidea as cestodarians and are the putative sister group of the true tapeworms, or eucestodes (Waeschenbach et al., 2012). Within the order, species of the only accepted genus *Gyrocotyle* Diesing, 1850 are common, well-reported parasites of holocephalans. Their systematics have been reviewed previously (e.g. Colin et al., 1986; Bandoni & Brooks, 1987; Williams et al., 1987; Gibson, 1994) and the morphological characters for distinguishing species have been thoroughly discussed and, according to Williams et al. (1987), been found wanting. Moreover, in addition to the lack of reliable morphological characters for species identification, the method of preservation has been shown to have a significant effect on their morphology, making it difficult to provide reliable species identifications post-preservation (Colin et al., 1986). In the absence of such characters, it appears that many specimens have been identified historically on the basis of their host species. Meanwhile, sequence data are available for only three putative species.

In this paper we use partial large nuclear ribosomal subunit (*lsrDNA*; domains D1–D3) and complete small nuclear ribosomal subunit (*ssrDNA*) sequences to reconstruct a phylogenetic network of *Gyrocotyle* spp. from five chimaeras: *Chimaera monstrosa* L., *Hydrolagus mirabilis* (Collett), *Hydrolagus colliei* (Lay & Bennett) (Chimaeridae), *Harriotta raleighana* Goode & Bean (Rhinochimaeridae) and *Callorhynchus milii* Bory de Saint-Vincent (Callorhynchidae). Based on these data, we recognise and name two new species of *Gyrocotyle* from the bent-nose chimaera, *Ha. raleighana*, and the large-eyed rabbit fish, *Hy. mirabilis*. These deep-sea host species are not commonly seen and, although probably not rare at depths greater than 1,000 metres, are difficult to capture, requiring specialised equipment and considerable effort.

Materials and methods

Specimen collection, preservation and morphological study

Chimaeras were collected by RAB using a semi-balloon otter trawl during three research cruises in the North-East Atlantic aboard the National Environmental Research Council research vessel RRS *Discovery* (April 2001 and September/October 2002). The fishes were immediately dissected and worms extracted and fixed briefly in Berland's fluid and preserved in 80% ethanol for morphological examination, and in 100% ethanol for molecular analysis. In some cases, worms were cut into separate parts before fixation. Whole-mounts were stained with Mayer's paracarmine, cleared in beechwood creosote and mounted in Canada balsam. Measurements were made through a drawing tube on an Olympus BH-2 microscope using a Digicad Plus digitising tablet and Carl Zeiss KS100 software adapted by Imaging Associates, and are quoted in micrometres. Where two-dimensions are given length precedes width. Type- and voucher material has been submitted to the Natural History Museum, London, UK (NHMUK). Additional specimens for molecular analysis were obtained from waters off Norway (Tromsø, Finnmark and Bergen) and Tasmania, Australia, and published sequences of three *Gyrocotyle* species were included in the analyses. A list of taxa including collection information and sequence accession numbers is given in Table 1.

Table 1 *Gyrocotyle* species collection information and GenBank sequence accessions

Species	Label	Host	Locality	Coordinates	Depth (m)	Collection date	<i>IsrDNA</i>	<i>ssrDNA</i>
<i>Gyrocotyle haffii</i> n. sp.	Gcot6	<i>Hariotta raleighana</i> Goode & Bean	North Atlantic, Goban Spur ^a	49.7666°N, – 2.3500°W	1,631–1,653	15.iv.2001	MN657006	MN655880
<i>Gyrocotyle discoveryi</i> n. sp.	Gcot	<i>Hydrolagus mirabilis</i> (Collett)	North Atlantic, Goban Spur ^a	49.8166°N, – 11.7333°W	1,175–1,250	15.iv.2001	MN657011	–
	Gcot2		North Atlantic, Goban Spur ^a	49.8166°N, – 11.7333°W	1,175–1,250	15.iv.2001	MN657003	MN655879
	Gcot3		North Atlantic, Goban Spur ^a	49.8166°N, – 11.7333°W	1,175–1,250	15.iv.2001	MN657004	–
	Gcot4		North Atlantic, Goban Spur ^a	49.8166°N, – 11.7333°W	1,175–1,250	15.iv.2001	MN657005	–
	Gcot7		North Atlantic, Porcupine Bight ^a	51.2500°N, – 11.9166°W	1,200	30.ix.2002	MN657007	MN655881
	Gcot8		North Atlantic, Porcupine Bight ^a	51.2500°N, – 11.9166°W	1,200	30.ix.2002	MN657008	–
	Gcot9		North Atlantic, Goban Spur ^a	49.8333°N, – 12.0833°W	1,360–1,240	19.x.2002	MN657009	–
<i>Gyrocotyle nybelini</i> (Furhmann, 1931)	Gyrol	<i>Chimaera monstrosa</i> L.	Coast of Finnmark, Norway ^b	na	na	2.xi.2001	MN657016	MN655885
<i>Gyrocotyle confusa</i> van der Land & Dienske, 1968	Gycon	<i>Chimaera monstrosa</i> L.	Coast of Finnmark, Norway ^b	na	na	2.xi.2002	MN657014	–
<i>Gyrocotyle urna</i> (Grube & Wagener in Wagener, 1852)	Gcot11	<i>Chimaera monstrosa</i> L.	North Atlantic, Goban Spur ^a	49.8333°N, – 12.0833°W	1,360–1,240	19.x.2002	MN657010	MN655882
	Guma		Coast of Troms, Norway ^b	na	na	11.xi.2002	MN657013	MN655884
	Guma2		Coast of Finnmark, Norway ^b	na	na	2.xi.2001	MN657012	–
	Gyro2		Unknown fjord, Norway ^c	na	na	10.xi.2003	MN657015	MN655883
	Gyro		Unknown fjord, Bergen, Norway ^a	na	na	6.vi.1996	AF286924.2	AJ228782
<i>Gyrocotyle rugosa</i> Diesing, 1850	Grg	<i>Hydrolagus colliei</i> (Lay & Bennett)	Gulf of Alaska, Torch Bay, Alaska ^d	58.3142°N, – 136.8036°W	na	1997	AF286925.2	AF124455
<i>Gyrocotyle</i> sp.	Gyc	<i>Callorhynchus militi</i> Bory de Saint-Vincent	Tasman Sea, off Hobart, Australia ^e	na	na	1.x.1997	EU343735	EU343741

^aCollector: R. A. Bray; ^bCollector: C. Vøllekvist; ^cCollector: K. MacKenzie; ^dCollector: G. Tyler; ^eCollector: K. Rohde

Molecular analysis

Ethanol was removed from tissue samples by soaking in tris-EDTA buffer overnight or by evaporation at room temperature. Total genomic DNA was extracted using the DNeasy blood and tissue kit (Qiagen). Partial *lsrDNA* (domains D1-D3; c.1,400 bp) was amplified using LSU5 or ZX-1 + 1200R or 1500R primers; in the case of *Gyrocotyle confusa* van der Land & Dieneske, 1968, only a short fragment of 512 bp could be sequenced from a fragment amplified using primers 900F + 1500R. Complete *ssrDNA* (c.2,000 bp) was amplified using WormA and WormB primers for a subset of the taxa (see Table 1). PCRs were carried out in 25 µl reaction volumes using puRe Taq Ready-to-go PCR beads (Amersham Biosciences, Little Chalfont, UK) and 1 µl of a 10 µM solution of each primer. Cycling conditions included an initial denaturation for 5 min at 95 °C, followed by 40 cycles of 30 s at 95 °C denaturation, 30 s at 55 °C (*lsrDNA*) or 54 °C (*ssrDNA*) and 2 min at 72 °C, followed by a final hold of 7 min at 72 °C. Amplicons were purified using a QIAquick Gel Extraction Kit or a QIAquick PCR Purification Kit (Qiagen, Hilden, Germany). Sequencing of both strands was carried out on an Applied Biosystems 3730 DNA Analyser, using Big Dye version 1.1. PCR and internal sequencing primers for *lsrDNA* are given in Littlewood et al. (2000), except for ZX-1 which was modified from van der Auwera et al. (1994) as shown in bold: ACC CGC TGA **ATT** TAA GCA TAT. Primers for *ssrDNA* are given in Littlewood & Olson (2001). Contigs were assembled using Sequencher 4.5 (GeneCodes Corporation, Ann Arbor, USA) and manually checked for ambiguous and incorrect base calls. Sequence identity was verified using the Basic Local Alignment Search Tool (BLAST) (www.ncbi.nlm.nih.gov/BLAST/).

Phylogenetic analysis

Gene-specific alignments were made for all available gyrocotylidean sequences along with multiple representatives of either caryophyllidean or spathebothriidean species (see Olson et al., 2008) used as outgroups. In addition, a gyrocotylidean-only sequence alignment was made. Sequences were aligned with MAFFT version 7.149b (Kato, 2008) using 1,000 cycles of iterative refinement and the *genafpair* algorithm. Alignment masks for ambiguously aligned positions were generated using GBLOCKS (Castresana, 2000; Talavera &

Castresana, 2007) using less stringent settings, and were further refined by eye in Mesquite version 3.5 (Maddison & Maddison, 2018). Alignments with indicated exclusion sets are available from the NHM Data Portal at <https://doi.org/10.5519/0003327>. MrModeltest2 (Nylander, 2004) was used to select a model of nucleotide substitution using Akaike's information criterion. Data were partitioned into three character sets: (i) partial *lsrDNA*; (ii) complete *ssrDNA*; and (iii) partial *lsrDNA* + complete *ssrDNA*. Phylogenetic trees were constructed using Bayesian inference with MrBayes, version 3.2 (Ronquist & Huelsenbeck, 2003). Likelihood settings were set to nst = 6, rates = invgamma, ngammacat = 4 (equivalent to the GTR+I+G model of evolution). In the combined analysis, parameters were estimated separately for each gene. Four chains (temp = 0.2) were run for 15,000,000 generations and sampled every 1,000th generation; 10,000,000 generations were discarded as 'burn-in'. The 'burn-in' period was determined as the point when the average standard deviation of split frequency values were < 0.01.

To comply with the regulations set out in article 8.5 of the amended 2012 version of the *International Code of Zoological Nomenclature* (ICZN, 2012), details of all new taxa have been submitted to ZooBank. For each new taxon, the Life Science Identifier (LSID) is reported in the taxonomic summary.

Results

Molecular analyses

Bayesian inference analysis of the combined *ssr/lsrDNA* data is shown in Fig. 1, and the results of analyses of the individual gene partitions are given in Supplementary Figures S1 and S2. Nodes supported by < 0.95 posterior probabilities were collapsed. Table 2 gives the corrected (GTR+I+G) pairwise distances estimated for each gene. Analyses including either caryophyllidean or spathebothriidean outgroup taxa failed to robustly resolve relationships among the gyrocotylidean samples, as a consequence of the need to exclude large numbers of sites that lacked clear positional homology between ingroup and outgroup sequences. For example, the *lsrDNA* alignment including caryophyllidean outgroup taxa required

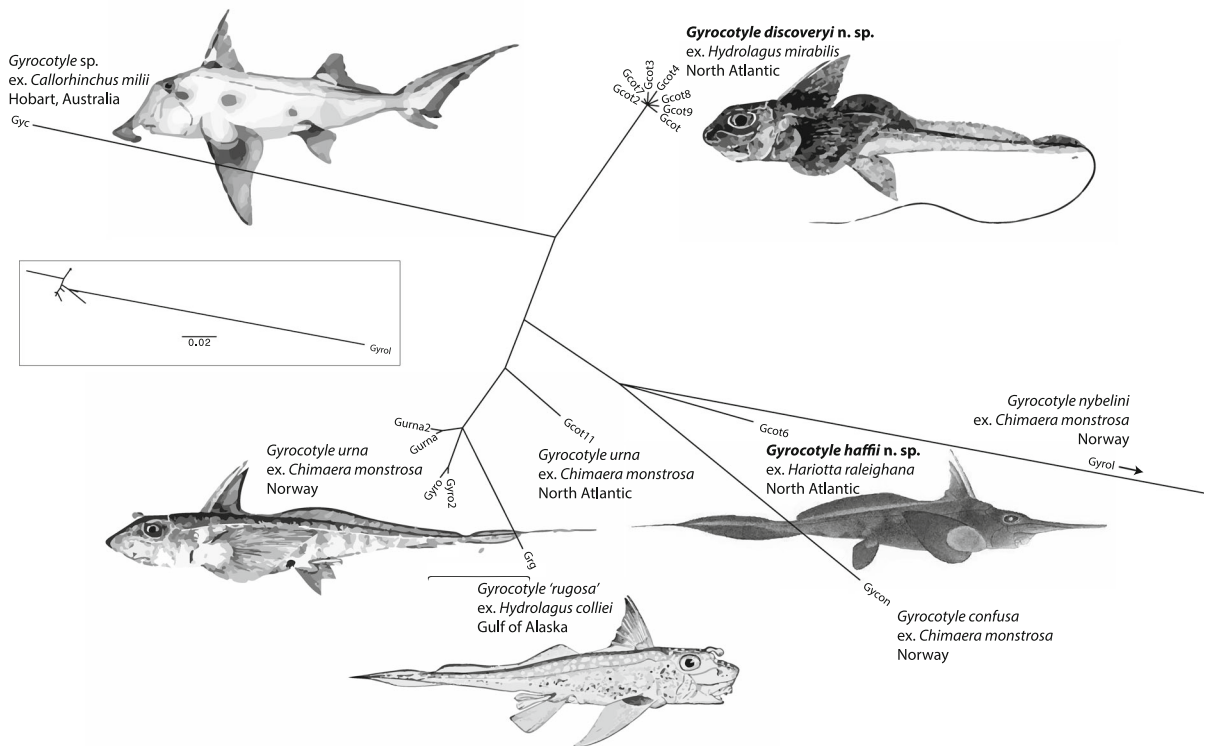


Fig. 1 Unrooted, consensus network of *Gyrocotyle* species based on combined complete *ssr* + partial *lsrDNA*. Nodes supported by < 0.95 posterior probabilities have been collapsed. Boxed inset shows the topology including the full branch subtending *G. nybelini*. Sample labels are given in Table 1

60% of the sites to be excluded whereas an alignment including only gyrocotylidean sequences required only 25% and thus included a greater number of informative characters among the ingroup sequences. We therefore chose to maximise the number of informative sites by aligning the gyrocotylidean sequences to themselves and consequently present our results as un-rooted networks.

All data partitions showed that the most divergent taxon by an order of magnitude was *G. nybelini* (Fuhrmann, 1931) Bandoni & Brooks, 1987 from *C. monstrosa* collected off Norway (Table 2); inset boxes in Fig. 1 and Supplementary Figures S1 and S2 were required to depict its full branch length relative to the other taxa. This was followed by *Gyrocotyle* sp. from *C. milii* off Australia. Among the samples collected from the North-East Atlantic, those from *Hy. mirabilis* formed a tight cluster with good separation from the other branches of the network and are described below as *Gyrocotyle discoveryi* n. sp. A specimen from *Ha. raleighana* similarly formed a distinct lineage in the

network and has been described below as *Gyrocotyle haffii* n. sp. This lineage was connected in an unresolved node with *G. confusa* and *G. nybelini*, also from *C. monstrosa* off Norway. Samples identified as *G. urna* (Grube & Wagener in Wagener, 1852) Wagener, 1858 from *C. monstrosa* showed considerable divergence, both among Norwegian fjords and between these and the North Atlantic, whereas the sample identified as *G. rugosa* Diesing, 1850 from *Hy. colliciei* from the Gulf of Alaska was closer to the *G. urna* samples from Norway than they are to the *G. urna* sample from the North-East Atlantic. The possibility that *G. urna*/*G. rugosa* represents a single, variable species is discussed below.

Class Cestoda

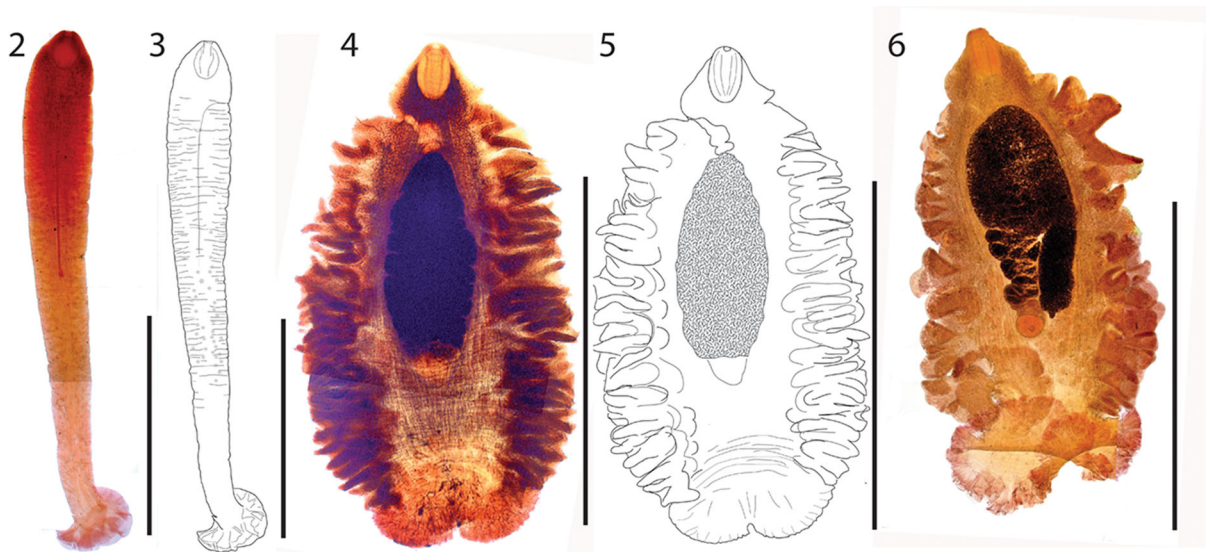
Order Gyrocotylidea Poche, 1926

Family Gyrocotylidae Benham, 1901

Genus *Gyrocotyle* Diesing, 1850

Table 2 Corrected (GTR+I+G) pairwise distances between samples (above the diagonal: *ssrDNA*; below the diagonal: *IsrDNA*)

	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17
	Gyrol	Gcot2	Gcot3	Gcot4	Gcot7	Gcot8	Gcot9	Gcot11	Gcot	Gurna2	Gurna	Gycon	Gyro2	Gyro	Gcot6	Grg	Gyc
1	Gyrol	0.099			0.099			0.104		0.104			0.101	0.101	0.099	0.097	0.12
2	Gcot2	0.253			0			0.01		0.01			0.011	0.011	0.015	0.014	0.027
3	Gcot3	0.252	0.001														
4	Gcot4	0.177	0	0.001													
5	Gcot7	0.185	0	0.028	0			0.01		0.01			0.011	0.011	0.015	0.014	0.027
6	Gcot8	0.185	0	0.028	0	0											
7	Gcot9	0.185	0	0.028	0	0											
8	Gcot11	0.157	0.029	0.066	0.022	0.024	0.024			0.005			0.005	0.006	0.013	0.008	0.029
9	Gcot	0.256	0	0.002	0	0.001	0.001	0.03									
10	Gurna2	0.127	0.03	0.076	0.019	0.025	0.025	0.007	0.031				0	0	0.014	0.003	0.029
11	Gurna	0.12	0.03	0.075	0.019	0.025	0.025	0.007	0.031	0							
12	Gycon	0.273	0.115	0.111	0.08	0.071	0.071	0.033	0.116	0.043	0.043						
13	Gyro2	0.165	0.03	0.067	0.02	0.025	0.025	0.01	0.031	0.005	0.005	0.043		0	0.014	0.004	0.03
14	Gyro	0.215	0.03	0.03	0.021	0.03	0.03	0.012	0.031	0.006	0.006	0.063	0.001		0.014	0.004	0.034
15	Gcot6	0.194	0.045	0.045	0.032	0.045	0.045	0.024	0.045	0.023	0.024	0.045	0.025	0.026		0.016	0.032
16	Grg	0.228	0.038	0.04	0.022	0.039	0.039	0.017	0.038	0.013	0.013	0.071	0.014	0.014	0.027		0.03
17	Gyc	0.189	0.039	0.047	0.036	0.041	0.041	0.039	0.04	0.04	0.04	0.092	0.037	0.036	0.05	0.04	



Figs. 2–6 Images and drawings of the new *Gyrocotyle* species. 2, Photomicrograph of *Gyrocotyle haffii* n. sp. holotype (NB: the specimen is immature) ex *Harriotta raleighana*, Goban Spur (13951/14); 3, Line-drawing of *Gyrocotyle haffii* n. sp., holotype; 4, Photomicrograph of *Gyrocotyle discoveryi* n. sp. holotype ex *Hydrolagus mirabilis* (Goban Spur; 15063/103a); 5, Line-drawing of *Gyrocotyle discoveryi* n. sp., holotype; 6, *Gyrocotyle discoveryi* n. sp. paratype ex *Hydrolagus mirabilis* (Goban Spur; 15066/124a). Scale-bars: 10 mm

Gyrocotyle haffii n. sp.

Type-host: *Harriotta raleighana* Goode & Bean (Chimaeriformes: Rhinochimaeridae), bent-nosed chimaera.

Type-locality: Goban Spur (49°46'N, 12°21'W, depth 1,631–1,653 m, 22–23.iv.2001; RRS *Discovery* Cruise 252, No. 13951/14), North-East Atlantic.

Type-material: Holotype (NHMUK.2019.11.21.1), paratype (NHMUK.2019.11.21.2).

Site in host: Spiral intestine.

Representative DNA sequences: MN655880 (*ssrDNA*); MN657006 (*lsrDNA*, domains D1–D3).

ZooBank registration: The Life Science Identifier (LSID) for *Gyrocotyle haffii* n. sp. is urn:lsid:zoobank.org:act:7717F9D6-4C9D-4C59-8D0A-4C5E306B4671.

Etymology: The species is named in honour of our late colleague and friend Professor Harford 'Haffi' Williams in recognition of his contribution to the understanding of the Gyrocotylidea.

Description

[Based on a single intact, immature whole worm and second immature worm from which the central portion

had been excised for molecular analysis; Figs. 2, 3]. With characters of the order. Body elongate with minute annular ridges; no large lateral flap. Length 23,504; greatest width near anterior extremity, 3,131. Rosette relatively small with few crenulations, 1,943 long. Anterior sucker large, oval, 1,750 × 1,223. Reproductive system immature; anlagen commences 2,852 from anterior extremity, 8,191 long; consisting of a long, narrow patch of stained tissue reaching, and a branched section passing, towards lateral margin of worm; apparently opening at *c.*268 from anterior extremity. Only other evidence of reproductive organs is putative vitelline glands scattered around posterior extremity of anlagen.

Diagnosis

Gyrocotyle haffii n. sp. can be diagnosed from other congeners on the basis of unique nucleotide characters in our rDNA alignments (listed as alignment position-nucleotide): *ssrDNA*: 218-T, 723-A, 746-C, 747-A, 748-G, 1,158-T, 1,654-G, 1,673-C, 2,115-C; *lsrDNA*: 612-T, 837-A, 875-T, 1,306-A, 1,395-C, 1,402-A, 1,501-G.

Remarks

As far as we are aware there is only one previous report of a gyrocotylidean from *Ha. raleighana*, the bent-nose chimaera. Parukhin (1966) reported “*Gyrocotylodes nybelini* Fuhrmann, 1931” in this host from the South Atlantic Ocean. Parukhin (1968) repeated this report saying (in translation) “Found in *Callorhynchus capensis*. Two adult parasites were found in two fish. In addition, six larvae were found in one of them. In addition to *C. capensis*, specimens were found in two *Hariota* [sic] *raleighana*. In both cases there were two specimens. Previously, this species was observed in the Atlantic in *Chimaera monstrosa*”. In addition, it seems likely that the records of ‘cestode adults’ from *Ha. raleighana*, *Hy. mirabilis* and *C. monstrosa* from the Rockall Trough off NW Scotland by Mauchline & Gordon (1984) refer to *Gyrocotyle* spp.

There is no reliable morphological character to differentiate this species or indeed any of the gyrocotylidean species circumscribed by molecular means. Therefore, the species is diagnosed by its relatively marked sequence divergence from those of recognised species.

***Gyrocotyle discoveryi* n. sp.**

Type-host: *Hydrolagus mirabilis* (Collett) (Chimaeriformes: Chimaeridae), large-eyed rabbitfish.

Type-locality: Goban Spur, North-East Atlantic.

Other localities: Porcupine Seabight (51°09′N, 11°55′W, depth 1,200 m, 30.xi.2002, RRS *Discovery* Cruise, No. 15048-14, 15); Goban Spur (49°49′N, 11°44′W, depth 1,175–1,250 m, 27.iv.2001, RRS *Discovery* Cruise 252, No. 13963/17, 20, 24, 72; 49°41′N, 11°53′W, depth 1,053–1,077 m, 23.iv.2001, RRS *Discovery* Cruise 252, No. 13962/4; 49°47′N, 11°58′W, depth 1,240–1,360 m, 19.x.2002, RRS *Discovery* Cruise D266, No. 15066-124, 125; 51°09′N, 11°55′W, depth 1200 m, 30.ix.2002, RRS *Discovery* Cruise D266, No. 15063-103), North-East Atlantic.

Type-material: Holotype (NHMUK 2019.11.21.3), paratypes (NHMUK.2019.11.21.4–13 from Goban Spur; NHMUK.2019.11.21.14–19 from Porcupine Sea Bight).

Site in host: Spiral intestine.

Representative DNA sequences: MN655879 and MN655881 (*ssrDNA*); MN657003–MN657005,

MN657007– MN657009, MN657011 (*lsrDNA*, domains D1–D3).

ZooBank registration: The Life Science Identifier (LSID) for *Gyrocotyle discoveryi* n. sp. is urn:lsid:zoobank.org:act:7B028A0B-B8EB-495E-A9F1-F29DDB60B89A.

Etymology: The species is named after the RRS *Discovery*, the NERC research vessel on which the specimens were collected.

Description

[Based on 17 specimens; Figs. 4–6.] With characters of the order. Body relatively squat, with deeply crenulated margins, 8,634–17,586 × 4,996–8,439 (12,071 × 6,676), width 36–98 (60)% of length. Anterior sucker distinct, 1,149–1,621 (1,347) long, 596–1,022 (771) wide. Uterus large, in central part of body, 1,546–2,996 (2,232) from anterior extremity, 2,453–6,226 (4,533) long, 35–43 (38)% of body length. Rosette distinct, fairly complex, 1,797–3,092 (2,575) long, junction with soma not clear. Eggs tanned, operculate, 85–97 × 39–56 (89 × 49).

Diagnosis

Gyrocotyle discoveryi n. sp. can be diagnosed from other congeners on the basis of unique nucleotide characters in our rDNA alignments (listed as alignment position-nucleotide): *ssrDNA*: 176-G, 782-G, 862-G, 973-C; *lsrDNA*: 573-A, 800-T, 1,245-C, 1,246-C, 1,247-G, 1,360-T, 1,369-T, 1,375-C, 1,379-G, 1,382-G, 1,391-A, 1,449-T, 1,468-T, 1,477-T.

Remarks

Mauchline & Gordon (1984) reported a “cestode” in *Hy. mirabilis* from the Rockall Trough off NW Scotland, which is, as far as we are aware, the only possible record of a *Gyrocotyle* from this host. Two species of *Gyrocotyle*, *G. major* van der Land & Templeman, 1968 and *G. abyssicola* van der Land & Templeman, 1968, have been reported from its congener, the small-eyed rabbit fish *Hydrolagus affinis* (de Brito Capello) on the edges of the continental shelf off the eastern coast of Newfoundland (van der Land & Templeman, 1968). These two species are illustrated as much more elongate than our specimens, with less complex lateral wrinkling and

small rosettes. The worms were recovered from frozen hosts, so the gross morphology may well not be of significance in differentiating these species. Subsequently, these two taxa have been reported from the same host species off south-western Greenland by Karlsbakk et al. (2002) and, puzzlingly, from a rhinochimaerine species, the straight-nosed rabbit fish *Rhinochimaera atlantica* Holt & Byrne, off the Scotian Shelf by Hogans & Hurlbut (1984). In the North-East Atlantic, *R. atlantica* has not been found harbouring *Gyrocotyle*, but it does harbour the strobilate tapeworm *Chimaerocestos prudhoei* Williams & Bray, 1984 and a congeneric host, the Pacific spookfish *R. pacifica* (Mitsukuri, 1895) also harbours a species of *Chimaerocestos* Williams & Bray, 1984 (see Caira et al., 1999, 2014). Other records of *Gyrocotyle* spp. from *Hydrolagus* spp. are from the Pacific Ocean (see Bandoni & Brooks, 1987).

There are no reliable morphological characters to differentiate this species or indeed any of the gyrocotylidean species circumscribed by molecular means. Therefore, the species is diagnosed by its relatively marked sequence divergence from those of recognised species.

Discussion

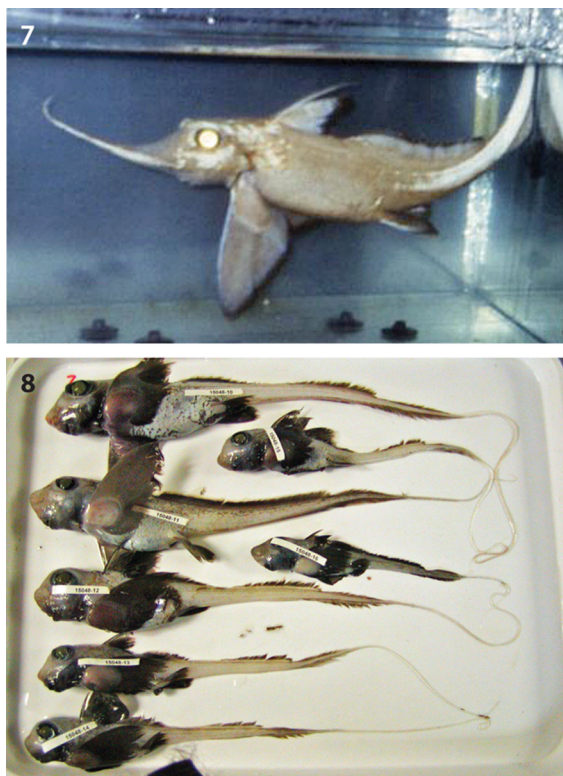
Colin et al. (1986) made a careful study, based on 1,361 specimens, of the morphological characters used for distinguishing species of *Gyrocotyle* and concluded that, due to the great contractibility of the worms, their reactions to different fixation techniques and the state of the worms at fixation (e.g. alive, dead, from frozen hosts), some characters were of limited or no value, i.e. total length and breadth, the degree of lateral crenulation, the complexity of the rosette, the distribution of body spines and the morphology of the eggs. In effect, they came to the conclusion that *Gyrocotyle* spp. could not be reliably identified using morphological characters. Indeed, these authors considered *Gyrocotyle confusa* and *Gyrocotyloides nybelini* as synonyms of *G. urna*, and the genus *Gyrocotyloides* Fuhrmann, 1930 as synonymous with *Gyrocotyle*. When an unidentified “chimaera cestodarian” was reported in the Caribbean chimaera *Chimaera cubana* Howell Rivero by Bunkley-Williams & Williams (2004), they reckoned that “most authors agree that only one morphologically highly

variable species of cestodarian is found in chimaeras, but some confusion exists about calling it *Gyrocotyle rugosa* Diesing, 1850 or *G. urna* (Grube & Wagener, 1852)”.

Despite the difficulties in identifying *Gyrocotyle* spp. on the basis of morphology and the controversies in the literature relative to the specific and generic status of various morphological forms (e.g. Colin et al., 1986; Bandoni & Brooks, 1987; Williams et al., 1987), there have been few investigations utilising molecular data. Simmons et al. (1972) utilised DNA hybridisation to confirm the distinctness of four species of *Gyrocotyle* from the Pacific Ocean. Bristow & Berland (1988), Berland et al. (1990) and Bristow (1992), using electrophoresis, fatty acid chemistry and biological characteristics, retained three species as distinct, but did not recognise the genus *Gyrocotyloides*. Olson & Caira (1999) generated an *ssrDNA* sequence of *Gyrocotyle rugosa* (Grg, Table 1), Olson et al. (2001) added partial *lsrDNA* data for this species and generated *ssrDNA* and *lsrDNA* data for *G. urna* (Gyro, Table 1), and Olson et al. (2008) generated *ssr/lsrDNA* sequences for *Gyrocotyle* sp. (Gyc, Table 1). In 2007, Waeschenbach et al. completed the *lsrDNA* sequence of *G. urna* (i.e. Gyro) and in 2012 complemented this with large fragments of mitochondrial genome data (Waeschenbach et al., 2012).

Each of the latter studies were aimed at resolving higher-level interrelationships of eucestodes and did not attempt to address the interrelationships or validity of named species and genera within the order. In this paper we make a first attempt at this, using ribosomal sequences from a variety of gyrocotylids, including some identified by other workers. The inability to root the resulting trees negated the ability to define clades, but the results still provide a picture of the relative genetic distances between samples and how they are interconnected within the network. The species *G. nybelini*, *G. confusa* and *G. haffii* n. sp. are part of an unresolved trichotomy, but are separated by long branches from the other samples and from each other. This indicates that *G. haffii* n. sp. is not conspecific with *G. nybelini*, suggesting in turn that Parukhin (1966, 1968) may have been incorrect in reporting *G. nybelini* from the host *Ha. raleighana*.

Gyrocotyle sp. from *Callorhynchus milii* off Hobart, Australia, forms another long branch in the network and, on this basis, is likely to represent an undescribed species. The hosts of this lineage of



Figs. 7, 8 Host images. 7, *Harriota raleighana* (longnose chimaera) suspended in tank for photography. 8, Some of the *Hydrolagus mirabilis* (large-eyed rabbitfish) specimens investigated

Gyrocotyle are unusual chimaeras known commonly as ghost sharks or elephant fish and are restricted to the temperate coasts of Australia and New Zealand. Not deep-sea dwelling, they constitute part of the fisheries in both countries and are commonly taken, suggesting that the collection and study of their gyrocotylid parasites should make the circumscription of this putatively novel species easier than for species whose hosts are rarely obtained.

Gyrocotyle urna, with six samples from *C. monstrosa* clustering with ‘*G. rugosa*’ from *Hydrolagus colliei* off Alaska, may well represent a complex of similar species, or is a single widespread species in northern waters with contacts *via* the deep Arctic Ocean. There is also distinct divergence between *G. urna* specimens from different Norwegian fjords. These may be as deep as 1,300 m and, like most fjords, are deeper than the adjacent sea and generally have a sill at their mouth formed by the glacier’s terminal moraine. This topology may explain the

apparent isolation of *Gyrocotyle* populations from different fjords as indicated by their genetics.

Gyrocotyle discoveryi n. sp., represented by seven samples from *Hy. mirabilis* in the North-East Atlantic, is almost genetically homogeneous. The regions of the two sites of collection are adjacent, with the Goban Spur forming the relatively shallow bank at the southern margin of the Porcupine Seabight. The samples formed a tight cluster that most likely represents a clade specific to the large-eyed rabbitfish.

As far as we are aware, the only gyrocotylids previously reported from the North-East Atlantic are the three species known from *C. monstrosa*, the commonly found holocephalan in the region. These are *G. urna*, the most commonly reported, and two rarer forms, *G. confusa* and *G. nybelini*. As stated previously, the latter species has been housed by various authors in the genus *Gyrocotylodes*, but this has been more commonly accepted as a synonym of *Gyrocotyle* (see Gibson, 1994). In contrast, our data lend some support to the recognition of *Gyrocotylodes* as a distinct genus, given its far greater genetic divergence in comparison to the other samples, including those obtained from far reaching parts of the globe.

Conclusions

The Gyrocotylidea is a small, but common group of cestodes of holocephalans with a widespread distribution characteristic of a relictual parasite group restricted to a relictual host group. The mostly deep-sea habitat of their hosts represents an unusually stable environment in which this host-parasite system evolved and likely explains their long-term persistence. Other features of the deep-sea, including fjords, are likely to have structured these systems in ways that are not immediately obvious until topography and mechanisms of isolation are considered, and may account for why genetic divergences do not strongly correlate with the degree of geographical separation among samples. Their systematics has been hitherto reliant on morphology and host-associations which in most cases have failed to satisfactorily distinguish species. It is therefore imperative that molecular investigations be employed to guide the circumscription of natural groups. Our results indicate that *Gyrocotyle* comprises not one cosmopolitan, non-

specific species, but a group of distinct, mostly host-specific, species that cannot be distinguished by morphology. Although the recognition of individual lineages and sequence clusters as species is problematical and probably provisional, establishing these conceptions now is justified by the fact that the hosts of the new species are rarely seen; the specimen of the long-nosed chimaera *Ha. raleighana* (Fig. 7) is the only one RAB has examined in over 30 years of marine trawling, whereas *Hy. mirabilis* (Fig. 8) is found in numbers, but only at particular depths and localities.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval All applicable institutional, national and international guidelines for the care and use of animals were followed.

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