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Insights from new cestodes of the crocodile shark, *Pseudocarcharias kamoharai* (Lamniformes: Pseudocarchariidae), prompt expansion of *Scyphyophyllidum* and formal synonymization of seven phyllobothriidean genera – at last!

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Abstract

Three new cestode species are described from the crocodile shark (Pseudocarcharias kamoharai) in Ecuador. All three were examined with light and scanning electron microscopy. The unique combination of morphological features in one of the new species prompted formal investigation of the non-monophyly of Paraorygmatobothrium relative to the morphologically similar genera Doliobothrium, Guidus, Marsupiobothrium, Nandocestus, Orectolobicestus, Ruhnkecestus and Scyphophyllidium. Sequence data generated for part of the 28S rDNA gene were subjected to maximum likelihood (ML) analysis. The resulting tree led to the synonymization of six of these seven genera with Scyphophyllidium, and transfer of their species to the latter genus. With the new species, the number of described members of Scyphophyllidium is now 45. The diagnosis of Scyphophyllidium is revised to accommodate these species. In addition, to expedite future descriptions, eight categories of Scyphophyllidium species are circumscribed, based largely on bothridial features. Scyphophyllidium timvickiorum n. sp. is a category 1 species. Beyond being the smallest category 1 species, it bears, rather than lacks, apical suckers and lacks, rather than bears, strobilar scutes. The two other new species are members of Clistobothrium. Clistobothrium amyae n. sp. differs from its congeners in bothridial shape, elongate cephalic peduncle and tiny size. Clistobothrium gabywalterorum n. sp. differs from the two of its congeners that also possess foliose bothridia in overall size and testis number. Despite their substantial morphological differences, the ML tree indicates they are sister taxa. Both are unique among their congeners in possessing cephalic peduncle spinitriches. The diagnosis of *Clistobothrium* is revised accordingly.

Introduction

The past decade or so has seen the erection of a series of seven novel genera found parasitizing the spiral intestines of members of three orders of sharks as well as one species of stingray (Caira & Durkin, 2006; Ruhnke et al., 2006a; Reyda, 2008; Caira et al., 2011; Cutmore et al., 2011; Ruhnke & Workman, 2013), and one species of skate (Ivanov, 2006), which, despite sharing a series of unique ultrastructual features and similarities in overall proglottid anatomy (see Ruhnke, 2011; Cutmore et al., 2017), differ considerably in bothridial morphology. For example, the bothridia of Doliobothrium Caira, Malek & Ruhnke, 2011 each lack an apical sucker and possess a proximal aperture; those of Orectolobicestus Ruhnke, Caira & Carpenter, 2006 each possess an apical sucker and marginal loculi; those of *Ruhnkecestus* Caira & Durkin, 2006 lack an apical sucker but bear facial loculi; those of Hemipristicola Cutmore, Theiss, Bennett & Cribb, 2011 bear an apical sucker and a deep central cavity; and those of Alexandercestus Ruhnke & Workman, 2013 bear an apical sucker and are highly foliose. The bothridia of Guidus Ivanov, 2006 are highly globose (Ivanov, 2006). The bothridia of Nandocestus Reyda, 2008 resemble those of Orectolobicestus in bearing marginal loculi, but this genus is unique among these genera in that it parasitizes a freshwater stingray rather than sharks. It also bears circumcortical, rather than lateral, vitelline follicles (Reyda, 2008). Bothridial features also serve to distinguish the above genera from three allied genera of shark tapeworms erected 25 or more years ago (see Woodland, 1927; Yamaguti, 1952; Ruhnke, 1994). For example, the bothridia of Scyphopyllidium Woodland, 1927 and Marsupiobothrium Yamaguti, 1952 bear apical suckers and are globose in form. The bothridia

As molecular phylogenetic analyses have expanded to include greater representation of these genera, the close affinities among these genera have been confirmed. However, these works have also served to call the monophyly of the speciose, yet morphologically uniform, Paraorygmatobothrium into question relative to at least a subset of the above genera (Cutmore et al., 2011, 2017; Caira et al., 2014a; Ruhnke et al., 2020). This suggests that a critical re-evaluation of these genera is in order. The discovery of a new species parasitizing the crocodile shark, Pseudocarcharias kamoharai (Matsubara), prompted us to formally tackle that issue here. Not only does this species exhibit a blend of the diagnostic morphological features of several of the above genera, but also phylogenetic analysis of a portion of the 28S rDNA gene suggests that it is most closely related to a subset of species of Paraorygmatobothrium. In identifying the most effective strategy for aligning the classification of these taxa with their phylogenetic relationships and morphologies, we also examined new material of the poorly known genera Marsupiobothrium and Scyphophyllidium to determine the conditions of several key morphological features.

As the only member of the family Pseudocarchariidae Compagno, *P. kamoharai* was also interesting from a comparative standpoint because it represents the only family of lamniform sharks that has not yet been examined for cestodes. In addition to the problematic new phyllobothriidean species referred to above, this shark species was found to host two new, relatively morphologically divergent, species of the phyllobothriidean genus *Clistobothrium* Dailey & Vogelbein, 1990, both of which are also described below.

Materials and methods

Specimen collection

Eight specimens of P. kamoharai, consisting of five females (97-106 cm in total length [TL]) and three males (84.5-108 cm in TL), were examined. All eight specimens were collected between May 22 and June 2 of 2014 from a fish market in Santa Elena (2°12'24.4"S, 80°56'58.1"W), Ecuador. Additional information on each host can be obtained from the Global Cestode Database (www.elasmobranchs.tapewormdb. uconn.edu) by entering the specimen numbers (i.e. EC-4, EC-5, EC-8, EC-9, EC-35, EC-36, EC-54 and EC-55). A small sample of liver tissue was taken from each animal and preserved in 95% ethanol for molecular verification of host identity. In each case, the spiral intestine was removed and opened with a mid-ventral longitudinal incision and then washed with seawater. Washings were either fixed in 10% seawater-buffered formalin (9:1) for morphological work or in 95% ethanol for molecular work. In some cases, the resulting washings were examined for cestodes under a stereomicroscope in the field prior to fixation and a subset of specimens found was fixed in 10% seawater-buffered formalin and a subset was fixed in 95% ethanol. Spiral intestines of five animals were then fixed in 10% seawater-buffered formalin and two were fixed in 95% ethanol. After one or two weeks, all formalin-fixed material was transferred to 70% ethanol for storage. Material preserved in 95% ethanol was stored in a -20°C freezer.

Morphological methods

Whole mounts of worms from P. kamoharai were prepared as follows for examination with light microscopy: specimens were hydrated in a graded ethanol series, stained in Delafield's haematoxylin, differentiated in tap water, destained in acidic 70% ethanol, neutralized in basic 70% ethanol, dehydrated in a graded ethanol series, cleared in methyl salicylate and mounted on glass slides under coverslips in Canada balsam diluted with methyl salicylate. They were then placed in a drying oven at 55°C for one week. Measurements were taken with a Zeiss Axioskop 2 Plus compound microscope (Thornwood, New York, USA) using SPOT Diagnostic Instrument Digital Camera Systems and SPOT software, version 4.6 (SPOT Imaging Solutions, Sterling Heights, Michigan, USA). Unless otherwise stated, measurements are presented in micrometres as ranges, followed in parentheses by the mean, standard deviation, total number of specimens measured and total number of measurements taken when more than one measurement was made per worm. With the exception of testes number, all proglottid measurements come from the terminal-most mature proglottid. Line drawings were made with a camera lucida attached to the Zeiss Axioskop 2 Plus compound microscope.

Temporary whole mounts of eggs were prepared as follows: gravid proglottids were transferred to a 1:10 mixture of glycerine and 70% ethanol, teased apart with a fine needle to release the eggs and placed in an open container in a fume hood overnight. They were then mounted in the same mixture on glass slides under coverslips, the edges of which were sealed with two coats of clear nail polish. Images were taken using the camera system described above.

Museum abbreviations used are as follows: LRP, Lawrence R Penner Parasitology Collection, Department of Ecology and Evolutionary Biology, University of Connecticut, Storrs, Connecticut, USA; MEPN, Museo de Colecciones Biológicas Gustavo Orcés, Escuela Politécnica Nacional, Ladrón de Guevara E11-253, Quito, Ecuador; MPM, Meguro Parasitological Museum, Tokyo, Japan; USNM, National Museum of Natural History, Smithsonian Institution, Washington, DC, USA. Microthrix terminology follows Chervy (2009). Ruhnke *et al.* (2017) was used as the source of valid phyllobothriidean species, except that *Paraorygmatobothrium musteli* (van Beneden, 1850) Ruhnke, 2011 was also included.

Specimens from *P. kamoharai* were prepared for scanning electron microscopy (SEM) as follows: they were hydrated in a filtered graded series of ethanols, transferred to a solution of 1% osmium tetroxide and placed in a refrigerator overnight; they were then dehydrated in a filtered graded series of ethanols, placed in hexamethyl-disilazane (Ted Pella Inc., Redding, California, USA) and allowed to air-dry in a fume hood for approximately 1 h. They were then mounted on aluminium stubs using double-sided PELCO carbon tabs (Ted Pella Inc.), sputter coated with 35 nm of gold/palladium and examined with a FEI Nova NanoSEM 450 field emission scanning electron microscope (FEI, Hillsboro, Oregon, USA).

In addition, two specimens of *Marsupiobothrium gobelinus* Caira & Runkle, 1993, taken from the same specimen of the goblin shark (*Mitsukurina owstoni* Jordan) from which the type material of this species was collected, were prepared for and examined with SEM as described above. The whole mounts of the hologenophores of *Marsupiobothrium* sp. 1, for which 28S rDNA and 18S rDNA data (LRP nos 8336 and 8337, respectively) were generated by Caira *et al.* (2014a), were examined. In Downloaded from https://www.cambridge.org/core. IP address: 32.218.8.192, on 30 Mar 2020 at 22:17:34, subject to the Cambridge Core terms of use, available at https://www.cambridge.org/core/terms. https://doi.org/10.1017/S0022149X20000036

Current name	Previous name	Category	Host order	Type host species	Apical sucker	Proximal aperture	Marginal loculi	Facial loculi	Bothridial shape	Scutes	Spinitriches on proximal bothridial surfaces	Spinitriches on distal bothridial surfaces	Egg shape	Source of data
Scyphophyllidium alopias (Yamaguti, 1952) Caira, Jensen & Ruhnke n. comb.	Marsupiobothrium alopias Yamaguti, 1952 (type)	1	Lamniformes	Alopias vulpinus	Yes	Yes	No	No	Globose	Yes	?	?	?	Yamaguti, 1952; Ivanov, 2006; Ruhnke, 2011
Scyphophyllidium angustum (Linton, 1889) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium angustum (Linton, 1889) Ruhnke, 2011	5	Carcharhiniformes	Carcharhinus obscurus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Linton, 1889; Ruhnke, 2011
Scyphophyllidium arnoldi (Ruhnke & Thompson, 2006) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium arnoldi Ruhnke & Thompson, 2006	5	Carcharhiniformes	Negaprion acutidens	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke & Thompson, 2006
Scyphophyllidium bai (Ruhnke & Carpenter, 2008) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium bai Ruhnke & Carpenter, 2008	6	Carcharhiniformes	Mustelus mustelus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	Spindle-shaped	Ruhnke & Carpenter, 2008
Scyphophyllidium barberi (Ruhnke, 1994) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium barberi Ruhnke, 1994	3	Carcharhiniformes	Triakis semifasciata	Yes	No	No	With two facial muscle bands	Flat	Yes	Serrate gladiate	Gongylate gladiate	Spherical	Ruhnke, 1994; Ruhnke <i>et al.</i> , 2020
Scyphophyllidium bullardi (Ruhnke, Daniel & Jensen, 2020) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium bullardi Ruhnke, Daniel & Jensen, 2020	5	Carcharhiniformes	Carcharhinus brevipinna	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium campbelli (Ruhnke, Daniel & Jensen, 2020) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium campbelli Ruhnke, Daniel & Jensen, 2020	5	Carcharhiniformes	Rhizoprionodon terraenovae	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke <i>et al.</i> , 2020
<i>Scyphophyllidium</i> cf. <i>giganteum</i> (of Caira <i>et al.</i> , 2014a)		4	Carcharhiniformes	Galeorhinus galeus	Yes	No	No	No	Globose	No	Gladiate (unmodified)	?	?	Caira et al., 2001
Scyphophyllidium chiloscyllii (Subhapradha, 1955) Caira, Jensen & Ruhnke n. comb.	Orectolobicestus chiloscyllii (Subhapradha, 1955) Ruhnke, Caira & Carpenter, 2006	2	Orectolobiformes	Chiloscyllium griseum	Yes	No	Yes	No	Flat	?	?	?	?	Subhapradha, 1955
Scyphophyllidium christopheri (Cutmore, Bennett, Miller & Cribb, 2017) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium christopheri Cutmore, Bennett, Miller & Cribb, 2017	6	Carcharhiniformes	Carcharhinus sorrah	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Cutmore <i>et al.</i> , 2017

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Current name	Previous name	Category	Host order	Type host species	Apical sucker	Proximal aperture	Marginal loculi	Facial loculi	Bothridial shape	Scutes	Spinitriches on proximal bothridial surfaces	Spinitriches on distal bothridial surfaces	Egg shape	Source of data
Scyphophyllidium deburonae (Ruhnke, Daniel & Jensen, 2020) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium deburonae Ruhnke, Daniel & Jensen, 2020	7	Carcharhiniformes	Carcharhinus isodon	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate gladiate	?	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium exiguum (Yamaguti, 1935) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium exiguum (Yamaguti, 1935) Ruhnke, 1994	6	Lamniformes	Alopias vulpinus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	Spindle-shaped	Yamaguti, 1935; Ruhnke, 1994
Scyphophyllidium filiforme (Yamaguti, 1952) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium filiforme (Yamaguti, 1952) Ruhnke, 1996	8	Lamniformes	Alopias vulpinus	Yes	No	No	No	Flat	Yes	?	?	?	Yamaguti, 1952; Ruhnke, 2011
Scyphophyllidium floraformis (Southwell, 1912) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium floraformis (Southwell, 1912) Ruhnke, 2011	6	Carcharhiniformes	Carcharhinus sorrah	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Southwell, 1912 Ruhnke, 2011
Scyphophyllidium giganteum (van Beneden, 1858) Woodland, 1927 (type)		4	Carcharhiniformes	Galeorhinus galeus	Yes	No	No	No	Globose	Yes	?	?	Spindle-shaped	van Beneden, 1858; Woodland 1927; Euzet, 1959; Ruhnke, 2011
Scyphophyllidium gobelinus (Caira & Runkle, 1993) Caira, Jensen & Ruhnke n. comb.	Marsupiobothrium gobelinus Caira & Runkle, 1993	4	Lamniformes	Mitsukurina owstoni	Yes	No	No	No	Globose	No	Gladiate (unmodified)	Trifid	?	Caira & Runkle, 1993; this study
Scyphophyllidium guariticus (Marques, Brooks & Lasso, 2001) Caira, Jensen & Ruhnke n. comb.	Nandocestus guariticus (Marques, Brooks & Lasso, 2001) Reyda, 2008 (type)	2	Myliobatiformes	Paratrygon aiereba	Yes	No	Yes	No	Flat	Yes	Cyrillionate	Serrate gladiate	Spherical	Marques <i>et al.</i> , 2001; Reyda, 2008
Scyphophyllidium harti (Cutmore, Bennett, Miller & Cribb, 2017) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium harti Cutmore, Bennett, Miller & Cribb, 2017	5	Carcharhiniformes	Carcharhinus Ieucas	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium haselii (Caira, Malek & Ruhnke, 2011) Caira, Jensen & Ruhnke n. comb.	Doliobothrium haselii Caira, Malek & Ruhnke, 2011 (type)	1	Carcharhiniformes	Carcharhinus dussumieri ^a	No	Yes	No	No	Globose	Yes	Serrate gladiate	Serrate gladiate	?	Caira et al., 201
Scyphophyllidium janineae (Ruhnke, Healy & Shapero, 2006) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium janineae Ruhnke, Healy & Shapero, 2006	2	Carcharhiniformes	Hemipristis elongata	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke et al., 2006b

Scyphophyllidium kelleyae (Ruhnke, Caira & Carpenter, 2006) Caira, Jensen & Ruhnke n. comb.	Orectolobicestus kelleyae Ruhnke, Caira & Carpenter, 2006	2	Orectolobiformes	Chiloscyllium indicum	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Ruhnke <i>et al.</i> , 2006a
Scyphophyllidium kirstenae (Ruhnke, Healy & Shapero, 2006) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium kirstenae Ruhnke, Healy & Shapero, 2006	5	Carcharhiniformes	Hemigaleus microstoma	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke et al., 2006b
Scyphophyllidium latipi (Caira & Durkin, 2006) Caira, Jensen & Ruhnke n. comb.	Ruhnkecestus latipi Caira & Durkin, 2006 (type)	3	Carcharhiniformes	Scoliodon macrorhynchus ^b	No	No	No	Yes	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Caira & Durkin 2006
Scyphophyllidium leuci (Watson & Thorson, 1976) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium leuci (Watson & Thorson, 1976) Ruhnke, 2011	8	Carcharhiniformes	Carcharhinus Ieucas	Yes	No	No	No	Flat	?	?	?	With or without small knobs	Watson & Thorson, 1976
Scyphophyllidium lorettae (Ruhnke, Caira & Carpenter, 2006) Caira, Jensen & Ruhnke n. comb.	Orectolobicestus lorettae Ruhnke, Caira & Carpenter, 2006	2	Orectolobiformes	Chiloscyllium cf. punctatum	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Ruhnke <i>et al.</i> , 2006a
Scyphophyllidium mattisi (Ruhnke, Daniel & Jensen, 2020) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium mattisi Ruhnke, Daniel & Jensen, 2020	7	Carcharhiniformes	Rhizoprionodon terraenovae	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate gladiate	?	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium mobedii (Malek, Caira & Haseli, 2010) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium mobedii Malek, Caira & Haseli, 2010	6	Carcharhiniformes	Carcharhinus dussumieri [®]	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Malek <i>et al.</i> , 2010
Scyphophyllidium mukahensis (Ruhnke, Caira & Carpenter, 2006) Caira, Jensen & Ruhnke n. comb.	Orectolobicestus mukahensis Ruhnke, Caira & Carpenter, 2006	2	Orectolobiformes	Chiloscyllium indicum	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Ruhnke <i>et al.</i> , 2006a
Scyphophyllidium musculosum (Subhapradha, 1955) Caira, Jensen & Ruhnke n. comb.	Doliobothrium musculosum (Subhapradha, 1955) Caira, Malek & Ruhnnke, 2011	1	Carcharhiniformes	Rhizoprionodon acutus	No	Yes	No	No	Globose	Yes	Serrate gladiate	Serrate gladiate	?	Subhapradha, 1955; Caira et al., 2011
Scyphophyllidium musteli (van Beneden, 1850) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium musteli (van Beneden, 1850) Ruhnke, 2011	8	Carcharhiniformes	Mustelus mustelus ^c	Yes	No	No	No	Flat	Yes	?	?	?	van Beneden, 1850; Ruhnke, 2011
Scyphophyllidium nicaraguensis (Watson & Thorson, 1976) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium nicaraguensis (Watson & Thorson, 1976) Ruhnke, 2011	8	Carcharhiniformes	Carcharhinus leucas	Yes	No	No	No	Flat	?	?	?	Spindle-shaped	Watson & Thorson, 1976

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Table 1. (Continued.)

Table 1. (continued.)											Spinitriches	Spinitriches		
Current name	Previous name	Category	Host order	Type host species	Apical sucker	Proximal aperture	Marginal loculi	Facial loculi	Bothridial shape	Scutes	on proximal bothridial surfaces	on distal bothridial surfaces	Egg shape	Source of data
Scyphophyllidium orectolobi (Butler, 1987) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium orectolobi (Butler, 1987) Ruhnke, 2011	2	Orectolobiformes	Orectolobus maculatus	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Serrate gladiate	Spindle-shaped	Butler, 1987; Ruhnke, 2011; Cutmore <i>et al.</i> , 2017
Scyphophyllidium paulum (Linton, 1897) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium paulum (Linton, 1897) Ruhnke, 2011	5	Carcharhiniformes	Galeocerdo cuvier	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	Spindle-shaped	Linton, 1897; Ruhnke, 2011
Scyphophyllidium prionacis (Yamaguti, 1934) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium prionacis (Yamaguti, 1934) Ruhnke, 1994 (type)	5	Carcharhiniformes	Prionace glauca	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	Spindle-shaped	Yamaguti, 1934; Ruhnke, 1994
Scyphophyllidium randyi (Ruhnke, Caira & Carpenter, 2006) Caira, Jensen & Ruhnke n. comb.	Orectolobicestus randyi Ruhnke, Caira & Carpenter, 2006	2	Orectolobiformes	Chiloscyllium hasselti	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Ruhnke <i>et al.</i> , 2006a
Scyphophyllidium roberti (Ruhnke & Thompson, 2006) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium roberti Ruhnke & Thompson, 2006	5	Carcharhiniformes	Negaprion brevirostris	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Ruhnke & Thompson, 2006
Scyphophyllidium rodmani (Ruhnke & Carpenter, 2008) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium rodmani Ruhnke & Carpenter, 2008	6	Carcharhiniformes	Mustelus antarcticus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	Spindle-shaped	Ruhnke & Carpenter, 2008
Scyphophyllidium sinclairtaylori (Cutmore, Bennett, Miller & Cribb, 2017) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium sinclairtaylori Cutmore, Bennett, Miller & Cribb, 2017	6	Carcharhiniformes	Carcharhinus sorrah	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sinuspersicense (Malek, Caira & Haseli, 2010) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium sinuspersicense Malek, Caira & Haseli, 2010	6	Carcharhiniformes	Carcharhinus dussumeri ^a	Yes	No	No	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Malek <i>et al.,</i> 2010
Scyphophyllidium taylori (Cutmore, Bennett & Cribb, 2009) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium taylori Cutmore, Bennett & Cribb, 2009	3	Carcharhiniformes	Hemigaleus australiensis	Yes	No	No	With two facial muscle bands	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Cutmore <i>et al.</i> , 2009
Scyphophyllidium timvickiorum Caira, Hayes & Jensen n. sp.		1	Lamniformes	Pseudocarcharias kamoharai	Yes	Yes	No	No	Globose	No	Gladiate (unmodified)	Gongylyate columnar	Spindle-shaped	This study

Scyphophyllidium triacis (Yamaguti, 1952) Caira, Jensen & Ruhnke n. comb.	Paraonygmatobothrium triacis (Yamaguti, 1952) Ruhnke, 1996	8	Carcharhiniformes	Triakis scyllium	Yes	No	No	No	Flat	Yes	?	?	?	Yamaguti, 1952; Ruhnke, 1996
Scyphophyllidium tyleri (Ruhnke, Caira & Carpenter, 2006) Caira, Jensen & Ruhnke n. comb.	<i>Orectolobicestus tyleri</i> Ruhnke, Caira & Carpenter, 2006 (type)	2	Orectolobiformes	Chiloscyllium punctatum	Yes	No	Yes	No	Flat	Yes	Serrate gladiate	Gongylate columnar	?	Ruhnke <i>et al.,</i> 2006a
Scyphophyllidium typicum (Subhapradha, 1955) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium typicum (Subhapradha, 1955) Ruhnke, 2011	5	Carcharhiniformes	Rhizoprionodon acutus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Subhapradha, 1955; Ruhnke, 2011
Scyphophyllidium ullmanni (Cutmore, Bennett, Miller & Cribb, 2017) Caira, Jensen & Ruhnke n. comb.	Paraorygmatobothrium ullmanni Cutmore, Bennett, Miller & Cribb, 2017	5	Carcharhiniformes	Carcharhinus cautus	Yes	No	No	No	Flat	Yes	Serrate gladiate	Serrate gladiate	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium uraguayense Brooks, Marques, Perroni & Sidagis, 1999		4	Carcharhiniformes	Mustelus mento	Yes	No	No	No	Globose	Yes	?	?	?	Brooks <i>et al.</i> , 1999
Scyphophyllidium sp. 1	Paraorygmatobothrium sp. 1 (of Cutmore et al., 2017)	8	Carcharhiniformes	Carcharhinus cautus ^d	Yes	No	No	No	Flat	?	?	?	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sp. 2	Paraorygmatobothrium sp. 2 (of Cutmore et al., 2017)	8	Carcharhiniformes	Sphyrna lewini ^d	Yes	No	No	No	Flat	?	?	?	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sp. 3	Paraorygmatobothrium sp. 3 (of Cutmore et al., 2017)	8	Carcharhiniformes	Carcharhinus amboinensis ^d	Yes	No	No	No	Flat	Yes	?	?	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sp. 4	Paraorygmatobothrium sp. 4 (of Cutmore et al., 2017)	8	Carcharhiniformes	Carcharhinus limbatus ^d	Yes	No	No	No	Flat	?	?	?	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sp. 5	<i>Doliobothrium</i> sp. (of Cutmore <i>et al.</i> , 2017)	1	Carcharhiniformes	Rhizoprionodon taylori ^d	No	Yes	No	No	Globose	?	?	?	?	Cutmore <i>et al.</i> , 2017
Scyphophyllidium sp. 6	Marsupiobothrium sp. 1 (of Caira et al., 2014)	1	Lamniformes	Alopias pelagicus ^d	Yes	Yes	No	No	Globose	No	Gladiate (unmodified)	Gongylate columnar	Spindle-shaped	This study
Alexandercestus gibsoni Ruhnke & Workman, 2013 (type)		N/A	Carcharhiniformes	Negaprion acutidens	Yes	No	No	No	Foliose	Yes	Gladiate (unmodified)	Serrate gladiate (tiny)	?	Ruhnke & Workman, 2013
Alexandercestus manteri Ruhnke & Workman, 2013		N/A	Carcharhiniformes	Negaprion brevirostris	Yes	No	No	No	Foliose	Yes	?	?	?	Ruhnke & Workman, 2013
<i>Guidus antarcticus</i> (Wojciechowska, 1991) Ivanov, 2006		N/A	Rajiformes	Bathyraja maccaini	Yes	No	No	No	Globose	?	?	?	?	Wojciechowska, 1991; Ivanov, 2006

Table 1. (Continued.)														
Current name	Previous name	Category	Host order	Type host species	Apical sucker	Proximal aperture	Marginal loculi	Facial loculi	Bothridial shape	Scutes	Spinitriches on proximal bothridial surfaces	Spinitriches on distal bothridial surfaces	Egg shape	Source of data
Guidus argentinense Ivanov, 2006 (type)		N/A	Rajiformes	Bathyraja brachyurops	Yes	Q	o Z	N	Globose	Yes	د	6	Spindle-shaped	Ivanov, 2006
<i>Guidus awii</i> (Rocka & Zdzitowiecki, 1998) Ivanov, 2006		N/A	Rajiformes	Bathyraja maccaini	Yes	N	°N N	No	Globose	°N N	~	2	~	Rocka & Zdzitowiecki, 1998; Ivanov, 2006
Hemipristicola gunterae Cutmore, Theiss, Bennett & Cribb, 2011 (type)		N/A	Carcharhiniformes	Hemipristis elongata	Yes	No	°N N	With central cavity	Flat	Yes	Serrate gladiate	Serrate gladiate	~	Cutmore <i>et al.</i> , 2011
Thysanocephalum (Linton, 1889) Braun, 1900 (type)		N/A	Carcharhiniformes	Galeocerdo cuvier	No	°N N	°N N	No	Foliose	Modified	Serrate gladiate	Serrate gladiate	ć	Linton, 1889; Caira <i>et al.</i> , 1999, 2001
^a As Carcharhinus cf. dussumieri. ^b As Scoliodon laticaudus. ^c As Mustelus vulgaris.	dussumieri. udus. s.													

addition, five specimens collected from the pelagic thresher shark (Alopias pelagicus Nakamura) in Taiwan in 2013 and 2017 that we believe are conspecific with *Marsupiobothrium* sp. 1 of Caira et al. (2014a) were prepared for and examined with SEM and as whole mounts for characterization of scolex features and egg morphology as described above. Although we were not able to borrow the type material of Marsupiobothrium alopias Yamaguti, 1952 (MPM no. SY7149) from the MPM, Iwaki Takashi kindly provided us with a series of images taken at intervals throughout the depth of the bothridia of the type specimen to help us evaluate the nature of the feature located in the centre of the bothridia that was interpreted as a sucker by Ivanov (2006) and Ruhnke (2011). Also examined were the whole mounts of the hologenophore (LRP no. 8346) of Scyphophyllidium cf. giganteum of Caira et al. (2014a) collected from Galeorhinus galeus (L) in the Chatham Rise off New Zealand, three specimens identified as Scyphophyllidium giganteum (van Beneden, 1858) Woodland, 1927 collected from G. galeus off the coast of California by Nathan Riser (LRP nos. 2742-2744), and the specimen prepared for SEM collected by Riser from G. galeus off California included in Caira et al. (2001).

Molecular methods

The D1–D3 region of the 28S rDNA gene were sequenced for one specimen of each of the three new species we collected from *P. kamoharai*. The centre portion of each worm was removed for DNA extraction; the remainder of each worm was prepared as a whole mount to serve as a hologenophore (*sensu* Pleijel *et al.*, 2008) for the specimens sequenced.

Total genomic DNA was extracted using a MasterPure[™] DNA Purification Kit (EpiCentre Technologies, Madison, Wisconsin, USA) following manufacturer's instructions. Specimens were then incubated at 65° C for 1 h and left at room temperature overnight with gentle shaking to dissolve DNA into solution. DNA quantity and quality were assessed using a NanoDrop 2000 microvolume spectrophotometer (Thermo Fisher Scientific, Waltham, Massachusetts, USA). Amplification of the D1-D3 region of the 28S rDNA gene was done in a 10 µl volume with 1 µl of DNA template, 0.1 µl of 10 M of each primer, 5 µl of GoTaq® Green Master Mix (Promega, Fitchburg, Wisconsin, USA) and 3.8 µl of water. The following primer pair was used for amplification: LSU-5 (5'-TAGGTCGACCCGCTGAAYTTA-3') (Littlewood et al., 2000) (5'-GCTATCCTGGAGGGAAACTTCG-3') LSU-1500R and (Tkach et al., 2003). Polymerase chain reaction product was purified using ExoSAP-IT.7 (Affymetrix, Inc., Santa Clara, California, USA). Sequencing was done using the primer pair LSU-55F (5'-AACCAGGATTCCCCTAGTAACGGC-3') (Bueno & Caira, 2017) and LSU-1200R (5'-GCATAGTTCACCATCTTTCGG-3') (Littlewood et al., 2000). Both strands were sequenced on an ABI PRISM® 3100 Genetic Analyzer (Applied Biosystems, Foster City, California, USA) using ABI Big Dye^{1M} dideoxy terminators (version 3.1). Contigs for the three specimens were assembled Geneious, v. 10.1.3 (Biomatters, Newark, New Jersey, USA).

Molecular phylogenetic analysis

For comparative purposes, sequence data for a portion of the 28S rDNA gene were obtained from GenBank for a total of 33 species of Alexandercestus, Doliobothrium, Guidus, Hemipristicola, Marsupiobothrium, Nandocestus, Orectolobicestus, Paraorygmatobothrium, Ruhnkecestus, Scyphophyllidium and

Host of specimen sequenced.

Thysanocephalum Linton, 1890. Also included were 14 species belonging to eight other genera of phyllobothriideans (see table 2), including *Clistobothrium*. Based on the phylogenetic relationships indicated in the tree resulting from the analyses of Caira *et al.* (2014a), *Disculiceps* sp. 1 of Caira *et al.* (2014a) and *Cathetocephalus thatcheri* Dailey & Overstreet, 1973 were included as outgroups, again using data obtained from GenBank. These sequences ranged in length from 726 bp to 1214 bp. Details of the specimens included in the molecular phylogenetic analysis are given in table 2.

Sequences were originally aligned and trimmed in Geneious, version 10.1.3. They were then aligned using PRANK (Löytynoja & Goldman, 2010) on the webPRANK Server using the default settings, but with the '+F flag' removed. The bestfitting model of evolution was determined using jModelTest, v. 2.1.10 (Guindon & Gascuel, 2003; Darriba et al., 2012) based on the evaluation of 88 models on the CIPRES Science Gateway (Miller et al., 2010). Sample-size corrected Akaike Information Criterion values were used to evaluate goodness of fit. A maximum likelihood (ML) analysis was conducted using Garli, v. 2.01 (Zwickl, 2006), also on the CIPRES Science Gateway (Miller et al., 2010). Default Garli configuration settings were used with the following exceptions: the starting tree topology was set to 'random', the number of attachment branches evaluated per terminal was set to 84 (i.e. twice the number of terminals in the matrix) and the number of independent search replicates was set to 100. Based on the results of the jModelTest analysis, TVM + I + G was employed as the model of evolution. Bootstrap (BS) values resulting from 1000 BS replicates were also generated with Garli v. 2.01 using the configuration settings specified above. BS values were displayed on the best tree using SumTrees v. 4.0.0 in DendroPy v. 4.0.3 (Sukumaran & Holder, 2010).

Results

Morphology and ultrastructure of poorly known genera

Marsupiobothrium alopias Yamaguti, 1952 (fig. 1a)

When he erected *Marsupiobothrium* in 1952, Yamaguti established *M. alopias*, from a host identified as the common thresher shark (*Alopias vulpinus* Bonnaterre), in Japan, as the type species. The bothridia were described as pear-shaped sacs with sphincterlike muscles and a submarginal apical sucker. We are unaware of any additional material of this species having been collected since that time. Despite the global distribution of *A. vulpinus* (see Compagno, 1984), we have not encountered this tapeworm in any of the over 50 specimens of common thresher sharks we have examined for cestodes at shark tournaments off New England, USA, or in fish markets in Taiwan. As a consequence, the type material remains the only available material of this species and this species has yet to be included in a molecular phylogenetic analysis or examined using SEM.

Marsupiobothrium alopias was re-described by both Ivanov (2006) and Ruhnke (2011) based on their examination of the type material. Their work raises an interesting question regarding the nature of a feature found on the centre of the proximal surface of the globose bothridia of this species. No mention of such a feature was made by Yamaguti (1952). However, both Ivanov (2006) and Ruhnke (2011) reported the presence of a sucker on the centre of each bothridium. Our examination of the images of the bothridia of the type specimen provided to us by the MPM

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(fig. 1a) indicates that this feature is actually a proximal aperture, rather than a sucker. Unfortunately, beyond scutes on the strobila, the microtriches on the scolex of M. *alopias* have not yet been characterized.

Marsupiobothrium gobelinus *Caira & Runkle*, 1993 (fig. 2a-c) This species has also not been reported since the time of its original description from the goblin shark (M. owstoni Jordan) by Caira & Runkle (1993). Its bothridia were characterized as globose, each with an apical sucker; no mention was made of a proximal aperture. Details of the surfaces of the scolex given in the original description were limited to mention of the fact that all surfaces of the bothridia and stalks (referred to as peduncles) were covered with 'slender, blade-like microtriches' (Caira & Runkle, 1993: 85); no SEM images were provided. Our examination of two additional specimens with SEM here confirmed the absence of apertures on the proximal surfaces of the bothridia (fig. 2a), and also that the capilliform filitriches on the anterior regions of the strobila are not arranged in scutes. The distal bothridial surfaces were found to bear a relatively unique form of spinithrix (fig. 2b). These spinitriches most closely resemble the trifurcate form of Chervy (2009). However, only their tips are trifid and, rather than bearing three extensions of similar length, these spinitriches bear one long central extension flanked on each side by a much shorter extension. In addition, the distal tips of all three extensions are rounded, rather than pointed (inset fig. 2b). Filitriches were not seen on this surface. The proximal bothridial surfaces were found to be covered with densely arranged narrow gladiate spinitriches and capilliform filitriches (fig. 2c). Inclusion of this species in molecular phylogenetic analyses, and, thus, confirmation of its phylogenetic position, awaits the collection of material preserved in ethanol for molecular work.

Marsupiobothrium sp. 1 (figs 1b and 2d-h)

Previously characterized, based solely on molecular data, the new material allowed us to characterize some of the basic morphological features of this species for the first time. This species was found to conspicuously differ from M. gobelinus in that it bears apertures on its proximal bothridial surfaces (fig. 2d). The distal bothridial surfaces were found to bear gongylate columnar spinitriches (fig. 2e). The proximal bothridial surfaces near the bothridial rims were found to bear a band of densely arranged simple gladiate spinitriches interspersed with capilliform filitriches (fig. 2f); the proximal surfaces away from the rim were seen to bear only capilliform filitriches (fig. 2g). In terms of its utrastructural features, the capilliform filitriches on the anterior regions of the strobila were not arranged as scutes (fig. 2h); in this respect, this species also differs conspicuously from M. alopias. The availability of gravid proglottids allowed us to characterize the eggs of this species as being spindle-shaped with bipolar filaments that are uneven in length (fig. 1b).

Scyphophyllidium giganteum (van Beneden, 1858) Woodland, 1927 (fig. 2i) and S. cf. giganteum

When Woodland (1927) erected *Scyphyophyllidium*, he did so in a somewhat cursory fashion. His knowledge of the species was based on a single specimen, 95 mm in length, collected from the spiral intestine of a triakid shark identified as *Galeus vulgaris* Fleming (synonym of *Galeorhinus galeus*) collected off Plymouth, UK. He considered this specimen to be conspecific with the species identified by van Beneden (1858) as *Anthobothrium giganteum* van Beneden, 1858 collected off Belgium from a shark he

Table 2. Taxa used in the phylogenetic analysis, with their revised names, host species, GenBank numbers and source of data.

Current name	Previous name	Host species	GenBank no. (D1–D3) 28S rDNA	Source of data
Alexandercestus gibsoni		Negaprion acutidens	KC505623	Ruhnke & Workman, 2013
Calyptrobothrium sp. 1		Torpedo nobiliana	KF685754	Caira et al., 2014a
Cathetocephalus thatcheri		Carcharhinus leucas	KF685884	Caira et al., 2014a
Chimaerocestus n. sp. 1		Rhinochimaera pacifica	KF685882	Caira et al., 2014a
Chimaerocestus n. sp. 2		Rhinochimaera pacifica	KF685758	Caira et al., 2014a
Clistobothrium amyae n. sp.		Pseudocarcharias kamoharai	MN706184	This study
Clistobothrium carcharodoni		Carcharodon carcharias	HM856633 ^a	Randhawa, 2011
Clistobothrium cf. montaukensis		Lamna nasus	JF436971 ^a	Randhawa & Brickle 2011
Clistobothrium gabywalterorum n. sp.		Pseudocarcharias kamoharai	MN706183	This study
Clistobothrium montaukensis		Isurus oxyrinchus	EF095259	Caira et al., 2014a
Crossobothrium cf. dohrnii		Heptranchus perlo	KF685759	Caira et al., 2014a
Crossobothrium laciniatum		Hexanchus griseus	KF685883	Caira et al., 2014a
Disculiceps sp. 1		Carcharhinus limbatus	KF685761	Caira et al., 2014a
Guidus sp.		Bathyraja multispinis	MH688710	Beer <i>et al.</i> , 2019
Hemipristicola gunterae		Hemipristis elongata	HQ680624	Cutmore et al., 2011
New genus 10 n. sp. 1		Sphyrna lewini 1	KF685889	Caira et al., 2014a
Orygmatobothrium cf. musteli 1		Mustelus mustelus	KF685768	Caira et al., 2014a
Orygmatobothrium cf. musteli 2		Mustelus mustelus	KF685891	Caira et al., 2014a
Phyllobothrium cf. lactuca		Mustelus mento	KF685770	Caira <i>et al.</i> , 2014a
Phyllobothrium squali		Squalus acanthias	KF685897	Caira et al., 2014a
Scyphophyllidium arnoldi	Paraorygmatobothrium arnoldi	Negaprion acutidens	MN686528	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium bai	Paraorygmatobothrium bai	Mustelus mustelus	KC505625	Ruhnke & Workman 2013
Scyphophyllidium bullardi	Paraorygmatobothrium bullardi	Carcharhinus brevipinna	GQ470001	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium campbelli	Paraorygmatobothrium campbelli	Rhizoprionodon terraenovae	GQ470009	Ruhnke et al., 2020
Scyphophyllidium cf. giganteum		Galeorhinus galeus	KF685901	Caira et al., 2014a
Scyphophyllidium christopheri	Paraorygmatobothrium christopheri	Carcharhinus sorrah	MG008931	Cutmore <i>et al.</i> , 2017
Scyphophyllidium deburonae	Paraorygmatobothrium deburonae	Carcharhinus isodon	GQ470041	Ruhnke <i>et al.</i> , 2020
Scyphophyllidium exiguum	Paraorygmatobothrium exiguum	Alopias vulpinus	KF685769	Caira et al., 2014a
Scyphophyllidium guariticus	Nandocestus guariticus	Paratrygon aiereba	KF685888	Caira et al., 2014a
Scyphophyllidium harti	Paraorygmatobothrium harti	Carcharhinus leucas	MG008939	Cutmore et al., 2017
Scyphophyllidium janineae	Paraorygmatobothrium janineae	Hemipristis elongata	HQ680625	Cutmore et al., 2011
Scyphophyllidium kirstenae	Paraorygmatobothrium kirstenae	Hemigaleus microstoma	KC505626	Ruhnke & Workman 2013
Scyphophyllidium latipi	Ruhnkecestus latipi	Scoliodon macrorhynchus	KF685900	Caira <i>et al.</i> , 2014a
Scyphophyllidium mattisi	Paraorygmatobothrium mattisi	Rhizoprionodon terraenovae	GQ470009	Ruhnke et al., 2020
Scyphophyllidium orectolobi	Paraorygmatobothrium orectolobi	Orectolobus maculatus	MG008940	Cutmore et al., 2017
Scyphophyllidium paulum	Paraorygmatobothrium paulum	Galeocerdo cuvier	HQ680630	Cutmore et al., 2011

⁽Continued)

Table 2. (Continued.)

Current name	Previous name	Host species	GenBank no. (D1–D3) 28S rDNA	Source of data
Scyphophyllidium prionacis	Paraorygmatobothrium prionacis	Prionace glauca	KF685892	Caira <i>et al.</i> , 2014a
Scyphophyllidium randyi	Orectolobicestus randyi	Chiloscyllium hasselti	KF685767	Caira <i>et al.</i> , 2014a
Scyphophyllidium sinclairtaylori	Paraorygmatobothrium sinclairtaylori	Carcharhinus sorrah	MG008933	Cutmore et al., 2017
Scyphophyllidium taylori	Paraorygmatobothrium taylori	Hemigaleus australiensis	HQ680631	Cutmore et al., 2011
Scyphophyllidium timvickiorum n. sp.		Pseudocarcharias kamoharai	MN706182	This study
Scyphophyllidium tyleri	Orectolobicestus tyleri	Chiloscyllium punctatum	KF685890	Caira et al., 2014a
Scyphophyllidium typicum	Paraorygmatobothrium typicum	Rhizoprionodon cf. acutus 3	MN686558	Ruhnke et al., 2020
Scyphophyllidium ullmanni	Paraorygmatobothrium ullmanni	Carcharhinus cautus	MG008942	Cutmore et al., 2017
Scyphophyllidium sp. 1	Paraorygmatobothrium sp. 1	Carcharhinus cautus	MG008938	Cutmore et al., 2017
Scyphophyllidium sp. 2	Paraorygmatobothrium sp. 2	Sphryna lewini	MG008932	Cutmore et al., 2017
Scyphophyllidium sp. 3	Paraorygmatobothrium sp. 3	Carcharhinus amboinensis	MG008926	Cutmore et al., 2017
Scyphophyllidium sp. 4	Paraorygmatobothrium sp. 4	Carcharhinus limbatus	MG008944	Cutmore et al., 2017
Scyphophyllidium sp. 5	Doliobothrium sp.	Rhizoprionodon taylori	MG008928	Cutmore et al., 2017
Scyphophyllidium sp. 6	Marsupiobothrium sp. 1	Alopias pelagicus	KF685771	Caira <i>et al.</i> , 2014a
Thysanocephalum thysanocephalum		Galeocerdo cuvier	KF685902	Caira et al., 2014a
Trilocularia gracile		Squalus acanthias	KF685776	Caira et al., 2014a

^aTaxa for which only D2 data are available.

identified as 'milandre' (also considered to be G. galeus). Woodland provided no formal generic diagnosis. Instead, he described the details of his specimen and included the designation 'gen. n.' after the name of the species in the heading of that treatment. Following examination of specimens from sharks identified as Eugaleus galeus (L) (also a synonym of G. galeus) from Sète, France, Euzet (1959: 59) provided the following brief diagnosis of the genus: 'Scolex à 4 bothridies ovoides globuleuses, à ouverture antérieure, ne pouvant s'étaler. Pas de ventouse accessoire. Ovaire tétralobé. Vagin débouchant antérieurement à la poche du cirre. Vitellogènes latéraux. Pores génitaux alternant irrégulièrement', along with illustrations of a scolex and proglottid. ("Scolex with 4 ovoid globose bothridia, an anterior aperture, unable to spread. Accessory sucker lacking. Ovary tetralobed. Vagina opening anterior to the cirrus pouch. Vitellaria lateral.") Euzet (1994) subsequently added several additional details to the diagnosis, including the fact that the strobila was acraspedote and apolytic, the testes were numerous and post-vaginal testes were present on the poral side.

Based on the examination of three specimens from Nathan Riser, we were able to confirm the interpretation by Caira *et al.* (2001) of the bothridia as highly globose (fig. 2i); there is no evidence of the presence of proximal bothridial apertures. This work also confirmed the observation of Ruhnke (2011) that the bothridia each bear a small apical sucker. Furthermore, these specimens clearly bear capilliform filitriches on the strobila that are arranged as scutes. The proglottids of these specimens are consistent with the illustrations of van Beneden (1858) and Euzet (1959) in being longer than wide.

The only representative of *Scyphophyllidium* included in molecular phylogenetic work to date is the species identified as

S. cf. *giganteum* by Caira *et al.* (2014a) from *G. galeus* off the Chatham Rise in New Zealand. The morphology of the hologenophore is consistent with *S. giganteum* in most respects. Its bothridia are globose, bear apical suckers and lack proximal apertures. It also clearly bears scutes on its strobila. However, unlike the mature proglottids of *S. giganteum*, which are longer than wide, those of this specimen are substantially wider than long. Thus, we concur with Caira *et al.* (2014a) that this specimen should continue to be provisionally referred to as *S. cf. giganteum*.

Molecular phylogenetic analysis

The tree resulting from our ML analysis is shown in fig. 3. Two of the three new species discovered in P. kamoharai grouped as members of a clade that also included the three other species of Clistobothrium for which data were available in GenBank. The third new species grouped most closely with the species identified as Marsupiobothrium sp. 1 by Caira et al. (2014a) (referred to as Scyphophyllidium sp. 6 in fig. 3); this clade was sister to two of the 22 species included in the analysis that were originally assigned to Paraorygmatobothrium. The clade consisting of these four species grouped as sister taxon (with a BS value of 98%) to a larger clade that included the 20 remaining species originally assigned to Paraorygmatobothrium included in our analysis, as well as the species included in our analysis that were originally assigned to Doliobothrium, Nandocestus, Orectolobicestus, Ruhnkecestus and Scyphyophyllidium. The specimen of Hemipristicola gunterae grouped as the sister taxon to this larger clade, with Alexandercestus gibsoni as the sister taxon to this group. In contrast, the specimen of Guidus sp. grouped well outside of the above clade,

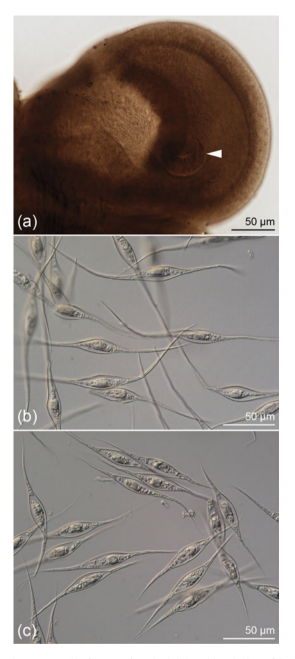


Fig. 1. Photomicrographs of species of *Scyphophyllidium*. (a) Bothridium of the holotype of *Marsupiobothrium alopias* (now *Scyphophyllidium alopias*) (MPM no. SY7149) from *Alopias vulpinus* in Japan; proximal aperture indicated by arrowhead. (b) Eggs of *Marsupiobothrium* sp. 1 of Caira *et al.* (2014a) (now *Scyphophyllidium* sp. 6) from *Alopias pelagicus* in Taiwan. (c) Eggs of *Scyphophyllidium timvickiorum* Caira, Hayes & Jensen n. sp. from *Pseudocarcharias kamoharai* in Ecuador.

as the sister taxon of a clade consisting of *Phyllobothrium squali* Yamaguti, 1952 and *Calyptrobothrium* sp. 1.

Revision of classification of seven genera in question

Our examination of material of *Marsupiobothrium* and *Scyphyophyllidium* expanded our understanding of the key features of both genera, and also served to confirm the latter as a valid genus. To help inform the development of an optimal solution for resolving the non-monophyly of *Paraorygmatobothrium* relative to six of the above genera (i.e. *Doliobothrium*,

Marsupiobothrium, Nandocestus, Orectolobicestus, Ruhnkecestus and Scyphophyllidium), we also examined the key morphological features exhibited by their 44 described species. As can be seen from table 1, the majority of these features are present in a variety of non-unique combinations across the members of these seven genera. In light of these data and the results of our molecular phylogenetic analysis, synonymization of six of these genera is in order. As the oldest valid name, Scyphophyllidium is hereby designated as the valid name for the larger, more inclusive genus; Doliobothrium, Marsupiobothrium, Nandocestus, Orectolobicestus, Paraorygmatobothrium and Ruhnkecestus are designated as its junior synonyms. The 42 described species and six undescribed species currently assigned to the latter six genera are hereby transferred to Scyphophyllidium. The new combinations and their appropriate authority citations are given in table 1. A revised, expanded diagnosis of the genus that accommodates all of these species is provided below.

Scyphophyllidium Woodland, 1927 revised

Synonyms. Doliobothrium Caira, Malek & Ruhnke 2011, Marsupiobothrium Yamaguti, 1952, Nandocestus Reyda, 2008, Orectolobicestus Ruhnke, Caira & Carpenter, 2006, Paraorygmatobothrium Runke, 1994 and Ruhnkecestus Caira & Durkin, 2006.

Diagnosis. Phyllobothriidea. Worms euapolytic or apolytic. Scolex with four bothridia, with or without cephalic peduncle. Bothridia globose or flat, often with band of muscle fibres around perimeter, without or occasionally with two facial semi-circular muscle bands, with or without marginal loculi, without or occasionally with facial loculi, without or occasionally with proximal aperture. Proximal bothridial surfaces with serrate gladiate or occasionally gladiate spinitriches. Distal bothridial surfaces with serrate gladiate, gongylate columnar or gongylate gladiate spinitriches. With or without neck. Capilliform filitriches on strobila usually arranged in scutes. Immature proglottids wider than long; usually becoming longer than wide with maturity. Genital pores usually alternating irregularly, located in anterior half of proglottid. Testes numerous, one to two layers deep in cross section; post-vaginal testes present. Cirrus sac containing coiled, armed cirrus. Ovary near posterior end of proglottid, H-shaped in dorsoventral view, tetralobed in cross section. Uterus ventral to vagina, median, extending anteriorly from anterior margin of ovary to well past cirrus sac. Vitellarium follicular; follicles usually arranged in two lateral bands, rarely circumcortical; bands extending almost entire length of proglottid, usually reduced or interrupted by ovary and cirrus sac. Excretory ducts in two lateral pairs. Eggs generally spindle-shaped, occasionally spherical.

Type species. Scyphophyllidium giganteum (van Beneden, 1858) Woodland, 1927.

Additional species. Forty-three listed in table 1. Parasites of the spiral intestines of sharks of the orders Carcharhiniformes, Lamniformes and Orectolobiformes, and occasionally batoids of the order Myliobatiformes. Cosmopolitan.

Remarks

As a result of these generic synonymizations, 44 species are now recognized as valid members of *Scyphophyllidium*. In order to expedite the future description of new species in this genus, the implementation of a strategy to help simplify comparisons, following Ghoshroy & Caira (2001) for the speciose onchoproteoce-phalidean genus *Acanthobothrium* Blanchard, 1848, is in order. To this end, we have circumscribed eight categories of

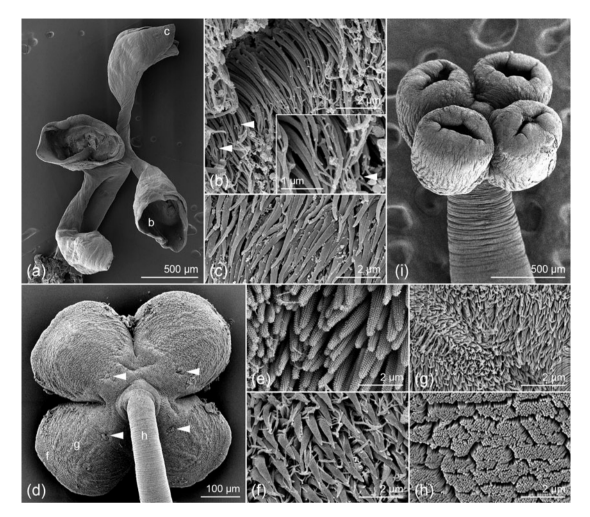


Fig. 2. Scanning electron micrographs of *Marsupiobothrium gobelinus* (now *Scyphophyllidium gobelinus*) from *Mitsukurina owstoni* in Australia. (a) Scolex; small letters indicate locations of details in (b) and (c). (b) Distal bothridial surface (with close-up inset); arrowheads indicate trifid tips of gladiate spinitriches. (c) Proximal bothridial surface. Scanning electron micrographs of *Marsupiobothrium* sp. 1 of Caira *et al.* (2014a) (now *Scyphophyllidium* sp. 6) from *Alopias pelagicus* in Taiwan. (d) Scolex, proximal view; small letters indicate locations of details in (f–h); arrowheads indicate proximal apertures on each bothridium. (e) Distal bothridial surface. (f) Proximal bothridial surface adjacent to rim. (g) Proximal bothridial surface away from rim. (h) Surface of strobila. (i) Scanning electron micrograph of *Scyphophyllidium* cf. *giganteum* from *Galeorhinus galeus* in New Zealand (modified from Caira *et al.*, 2001).

Scyphophyllidium based on the most conspicuous and informative scolex features. Each of the 44 described species and seven undescribed species has been assigned to one of these categories (see table 1). This strategy facilitates future descriptions by limiting the comparisons required to establish novelty to only those species belonging to the same category as each new species. The inclusion of ultrastructural features in this categorization strategy is appropriate because the characterization of microtriches has become essentially routine in the description of new species in these phyllobothriideans. We note that these categories do *not* reflect the phylogenetic relationships of their members; they were designated solely to help expedite and shorten future descriptions.

The eight categories are as follows. Category 1: species with globose bothridia, each with a proximal aperture; the species given this category designation are the three species formerly assigned to *Doliobothrium* (now *S. haselii*, *S. musculosum* and *Scyphophyllium* sp. 5) and the species formerly known as *M. alopias* (now *S. alopias*) and *Marsupiobothrium* sp. 1 (now *Scyphophyllidium* sp. 6). Category 2: species with bothridia bearing marginal loculi; species given this category designation are the six formerly assigned to *Orectolobicestus* (now *S. chiloscyllii*, *S. kelleyae*, *S. lorettae*, *S. mukahensis*, *S. randyi* and *S. tyleri*), the single species

formerly assigned to Nandocestus (i.e. S. guariticus) and, based on re-interpretation of SEMs, also S. orectolobi (of Cutmore et al., 2017) and S. janineae (of Ruhnke et al., 2006b). Category 3: species with facial loculi or facial semi-circular muscle bands; taxa given this category designation are the two species with facial semicircular muscle bands formerly assigned to Paraorygmatobothrium (now S. barberi and S. taylori) as well as the single species with facial loculi formally assigned to Ruhnkecestus (now S. latipi). Category 4: species with globose bothridia that lack proximal apertures; species given this category designation are the two described and one undescribed original members of Scyphyophyllidium (i.e. S. giganteum, S. cf. giganteum and S. uruguayense) and the species formerly referred to as M. gobelinus (now S. gobelinus). Category 5: species with bothridia that are essentially flat, lack proximal apertures, marginal loculi, facial loci and semi-circular muscle bands, and bear serrate gladiate spinitriches on their distal bothridial surfaces; species currently given this category designation are the following 11 species formerly assigned to Paraorygmatobothrium: S. angustum, S. arnoldi, S. bullardi, S. campbelli, S. harti, S. kirstenae, S. paulum, S. prionacis, S. roberti, S. typicum and S. ullmanni. Category 6: species with bothridia that are essentially flat, lack proximal apertures, marginal loculi, facial loculi and semi-circular

muscle bands, and bear gongylate columnar spinitriches on their distal bothridial surfaces; species currently given this category designation are the following eight species formerly assigned to Paraorygmatobothrium: S. bai, S. christopheri, S. exiguum, S. floraformis, S. mobedii, S. rodmani, S. sinclairtaylori and S. sinuspersicense. Category 7: species with bothridia that are essentially flat, lack proximal apertures, marginal loculi, facial loculi and semicircular muscle bands, and bear gongylate gladiate spinitriches on their distal bothridial surfaces; species currently given this category designation are two species formerly assigned to Paraorygmatobothrium, now S. deburonae and S. mattisi. Category 8: this is a temporary category designation that currently includes species with flat, unmodified bothridia (all previously assigned to Paraorygmatobothrium), the surfaces of which have yet to be characterized using SEM. SEM characterization of species in this category will allow them to be transferred to category 5, 6 or 7, depending on whether their distal bothridial surfaces are found to bear serrate gladiate, gongylate columnar or gongylate gladiate spinitriches, respectively. Species given this category designation are the following five species, all formerly assigned to Paraorygmatobothrium: S. filiforme, S. leuci, S. musteli, S. nicaraguensis and S. triacis, as well as four of the six undescribed species formerly assigned to Paraorygmatobothrium (now Scyphophyllidium sp. 1 through 4, respectively).

Scyphophyllidium timvickiorum Caira, Hayes & Jensen n. sp. (figs 1c, 4 and 5)

ZooBank number for species: urn:lsid:zoobank.org:act:9A4F7760-BF71-490A-A6E2-66133921276D.

Based on four gravid, eight mature and one immature worm, two scoleces examined with SEM and whole mounts of its strobila, and four preparations of eggs from detached gravid proglottids. Worms euapolytic, slightly craspedote, 4.2–9.8 (6.4 ± 1.7 ; 13) mm long; 7–21 (13 ± 4 ; 13) total proglottids per worm; maximum width 646–872 (743 ± 69 ; 13) near middle of scolex. Scolex 594–855 (723 ± 71 ; 11) long, with two dorsal and two ventral bothridia; cephalic peduncle lacking. Bothridia globose, highly muscular, with proximal apertures (fig. 5b) and single loculus, 267–451 (352 ± 34 ; 13; 39) long by 308–453 (376 ± 29 ; 12; 41) wide, bearing apical sucker; apical sucker 46–87 (67 ± 10 ; 12; 38) long by 57–99 (78 ± 10 ; 12; 40) wide; rims of bothridia with conspicuous band of circular muscle fibres.

Distal bothridial surfaces covered with gongylate columnar spinitriches and acicular filitriches (fig. 5d). Proximal bothridial surfaces with gladiate spinitriches and capilliform filitriches on rim (fig. 5e), with band of densely arranged gladiate spinitriches adjacent to rim (fig. 5f), with capilliform filitriches only on remainder of surfaces (fig. 5g). Capilliform filitriches on strobila not arranged in scutes.

Immature proglottids 6–19 (11 ± 4; 12) in number, approximately rectangular, becoming conspicuously longer than wide with maturity. Mature proglottids 1–4 (1 ± 1; 12) in number; terminal mature proglottid 688–1438 (1036 ± 198; 12) long by 192–267 (229 ± 23; 12) wide; length:width ratio 2.8–6.5:1 (4.6 ± 1.1; 12). Gravid proglottids 0–1 (n = 4). Testes 35–51 (42 ± 5; 13; 31) in total number, 13–19 (16 ± 2; 13; 31) in postporal field, 22–49 (36 ± 6; 10, 30) long by 23–70 (41 ± 9; 10, 30) wide, length: width ratio 0.49–1.63:1 (0.9 ± 0.25; 8; 30), arranged in 2–4 irregular columns anterior to cirrus sac, 1–2 columns in poral or aporal fields. Cirrus sac pyriform, 90–175 (143 ± 26; 12) long by 79–108 (99 ± 9; 12) wide, containing coiled cirrus; cirrus covered with

spinitriches (fig. 4c, d). Vas deferens minimal, coiled at anteriomedial margin of cirrus sac. Genital pores unilateral (n = 2) or irregularly alternating (n = 11), 74–86% (79 ± 3; 12) of proglottid length from posterior end. Vagina weakly sinuous, extending from ootype, along midline of proglottid to anterior margin of cirrus sac then laterally along anterior margin of cirrus sac to open into common genital atrium. Ovary near posterior end of proglottid, H-shaped in frontal view, 94-235 (166 ± 36 ; 12) long by 108–178 (129 \pm 20; 11) wide, tetralobed in cross section, weakly lobulated. Mehlis' gland posterior to ovary. Vitellarium follicular; follicles irregular in shape, arranged in two lateral bands; each band consisting of 2-3 columns of follicles, extending throughout length of proglottid interrupted by cirrus sac and ovary. Uterus ventral, extending from ovarian bridge to anterior to cirrus sac. Four excretory ducts, in one dorsal and one ventral pair. Eggs spindle-shaped with bipolar filaments; filaments unequal in length (fig. 1c).

Taxonomic summary

Type and only known host. Pseudocarcharias kamoharai (Matsubara), crocodile shark (Lamniformes: Pseudocarchariidae).

Site of infection. Spiral intestine.

Type locality. Santa Elena (2°12′24.4″S, 80°56′58.1″W), Santa Rosa de Salinas, eastern Pacific Ocean, Ecuador.

Additional localities. None.

Etymology. This species honours CH's parents, Tim and Vicki Hayes, for their unwavering support of her academic pursuits.

Specimens deposited. Holotype (MEPN no. 49443); two paratypes consisting of one immature and one gravid worm (MEPN nos 49444–49445); five paratypes consisting of four mature and one gravid worm (LRP nos 10138–10142) and five slides with glycerine mounts of eggs (LRP nos 10144–10148); SEM voucher (LRP no. 10143); five paratypes consisting of one gravid and four mature worms (USNM nos 1608084–1608088). Specimen examined with SEM retained in JNC's personal collection.

Molecular sequence data. 28S rDNA hologenophore (LRP no. 9311 [EC-5-P1V]) for GenBank no. MN706182.

Remarks

Scyphophyllidium timvickiorum n. sp. is a category 1 species in that its bothridia bear proximal apertures. It differs from the three other species assigned to this category in its possession of fewer testes (i.e. 35–51 vs 69–74, 74–92 and 155–187, in *S. musculosum*, *S. alopias* and *S. haselii*, respectively). It further differs from *S. haselii* and *S. musculosum* in that its bothridia bear, rather than lack, apical suckers. In addition, it is a much smaller worm than *S. alopias* (4.2–9.8 vs 25.4–26.2 mm TL). Unlike *S. alopias* and *S. haselii*, it also lacks scutes on its strobila.

Clistobothrium amyae Caira, Hayes & Jensen n. sp. (figs 6 and 7a-e)

ZooBank number for species: urn:lsid:zoobank.org:act:42A65147-09A3-4950-8DC8-EFD58A495651.

Based on whole mounts of seven mature worms, three scoleces examined with SEM and whole mounts of their strobila. Worms euapolytic, acraspedote, 7.7–16.5 (11.4 ± 3.6 ; 7) mm long; 25–40 (32 ± 6 ; 7) total proglottids per worm; maximum width at level of scolex. Strobila without distinct dorsomedian muscle band.

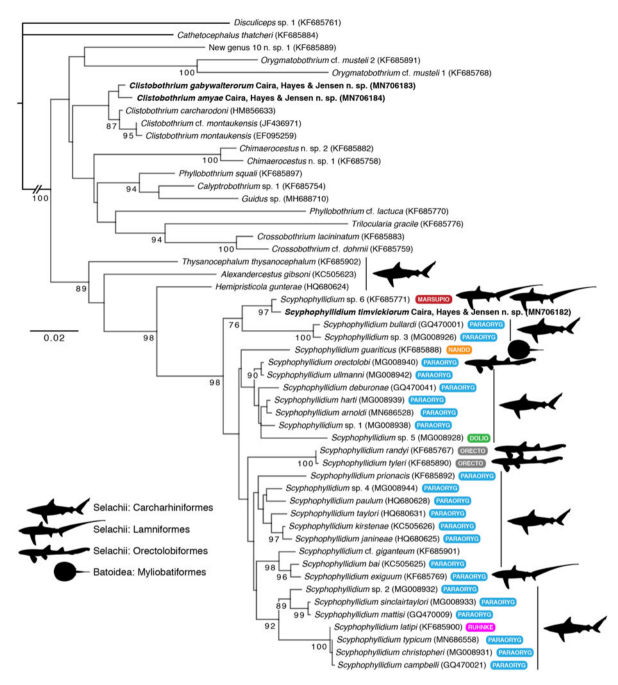


Fig. 3. Phylogenetic tree resulting from maximum likelihood (ML) analysis of a portion of the 28S rDNA gene for select phyllobothriideans, with focus on species previously assigned to *Paraorygmatobothrium* and allied genera, showing placement of three new species (in bold); *Cathetocephalus thatcheri* and *Disculiceps* sp. 1 were used as outgroups. Taxon labels include new generic assignments, GenBank numbers and abbreviations of previous generic assignments. Nodal support values presented as ML bootstrap values; only values greater than or equal to 70 are shown). Scale bar indicates nucleotide substitutions per site. Abbreviations: DOLIO, *Doliobothrium*; MARSUPIO, *Marsupiobothrium*; NANDO, *Nandocestus*; ORECTO, *Orectolobicestus*; PARAORYG, *Paraorygmatobothrium*; RUHNKE, *Ruhnkecestus*.

Scolex with cruciform apical region, two dorsal and two ventral round, stalked bothridia, and cephalic peduncle. Scolex proper 281–759 (547 ± 158 ; 6) long by 499–970 (821 ± 154 ; 7) wide; bothridia 203–428 (374 ± 64 ; 6; 11) long by 200–464 (367 ± 79 ; 6; 14) wide; each bothridium with one circular, apical sucker 68–125 (89 ± 18 ; 4; 18) long by 70–123 (95 ± 18 ; 6; 16) wide; stalks 141–469 (351 ± 182 ; 2; 3) long by 80–224 (146 ± 55 ; 4; 6) wide. Cephalic peduncle conspicuous, 1020–3840 (2226 ± 1015 ; 7) long.

Distal bothridial surfaces covered with slender, aristate gladiate spinitriches and capilliform filitriches (fig. 7b). Proximal bothridial

surfaces densely covered with slender gladiate spinitriches and capilliform filitriches (fig. 7c). Apex of cruciform region covered with sparsely arranged capilliform filitriches (fig. 7d). Cephalic peduncle densely covered with moderately sized, slender gladiate spinitriches and sparsely arranged capilliform filitriches (fig. 7e).

Immature proglottids 24-39 (31 ± 6 ; 7) in number, wider than long, becoming longer than wide with maturity. Mature proglottids one in number, longer than wide; terminal proglottid 423– 2283 (1434 ± 589 ; 7) long by 207–453 (378 ± 84 ; 7) wide; length-to-width ratio 1.1-3.7:1 (2.3 ± 1.1 ; 7). Testes 73–106 (87

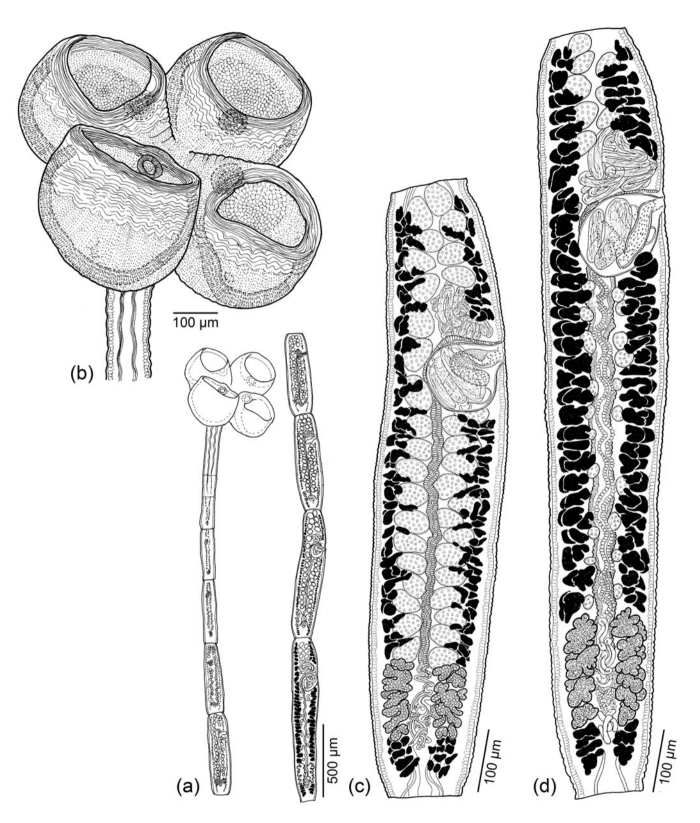


Fig. 4. Line drawings of *Scyphophyllidium timvickiorum* Caira, Hayes & Jensen n. sp. (a) Whole worm (holotype: MEPN no. 49443). (b) Scolex (holotype: MEPN no. 49443). (c) Subterminal mature proglottid showing testes (holotype: MEPN no. 49443). (d) Terminal mature proglottid (holotype: MEPN no. 49443).

 \pm 11; 7; 13) in total number, 7–15 (11 \pm 3; 7; 11) in postporal field, distributed in 4–6 (4.7 \pm 0.9; 10) columns anterior to cirrus sac, round to oblong, 24–54 (35 \pm 8; 6; 24) long by 24–58 (45 \pm 7; 6; 24) wide. Cirrus sac J-shaped, 181–343 (227 \pm 59; 6) long by

56–128 (87 ± 24; 7) wide, containing coiled cirrus; cirrus covered with minute spinitriches. Vas deferens minimal, coiled at anteriomedial and medial margins of cirrus sac. Genital pores lateral, irregularly alternating, 44-61% (51 ± 6; 7) of proglottid length

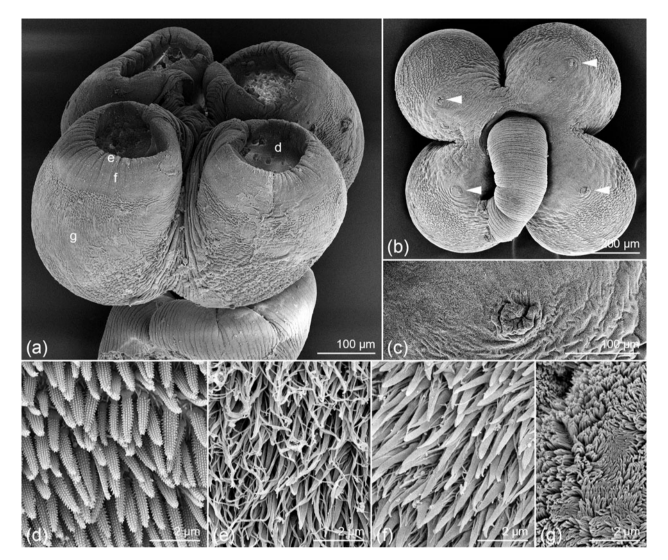


Fig. 5. Scanning electron micrographs of *Scyphophyllidium timvickiorum* Caira, Hayes & Jensen n. sp. (a) Scolex, lateral view; small letters indicate locations of details in (d–g). (b) Scolex, proximal view; arrowheads indicate proximal apertures on each bothridium. (c) Close-up of proximal aperture of one bothridium. (d) Distal bothridial surface. (e) Proximal bothridial surface near rim. (f) Proximal bothridial surface of gladiate spinithrix band adjacent to rim. (g) Proximal bothridial surface away from rim.

from posterior end. Vagina sinuous, extending from ootype, along midline of proglottid mid-level of cirrus sac, crossing cirrus sac ventrally then extending along anterior margin of cirrus sac to enter genital atrium anterior to cirrus. Ovary near posterior end of proglottid, H-shaped in frontal view, 162-341 (234 ± 72 ; 6) long by 78–109 (95 ± 13 ; 6) wide, bilobed in cross section, weakly lobulated. Mehlis' gland posterior to ovarian bridge. Vitellarium follicular; follicles irregular in shape, arranged in two lateral bands; each band consisting of 3–5 columns of follicles, extending throughout length of proglottid, interrupted by cirrus sac. Uterus ventral, extending from ovarian bridge to posterior margin of cirrus sac. Four excretory ducts, in one dorsal and one ventral pair.

Taxonomic summary

Type and only known host. Pseudocarcharias kamoharai (Matsubara), crocodile shark (Lamniformes: Pseudocarchariidae).

Site of infection. Spiral intestine.

Type locality. Santa Elena (2°12′24.4″S, 80°56′58.1″W), Santa Rosa de Salinas, eastern Pacific Ocean, Ecuador.

Additional localities. None.

Etymology. This species honours Dr Amy Donahue for her enthusiastic and innovative support of outreach science activities in her role as Vice Provost for Academic Operations at the University of Connecticut.

Specimens deposited. Holotype (MEPN no. 49441); one paratype (MEPN no. 49442); three paratypes (LRP nos 10132–10134); three SEM vouchers (LRP nos 10135–10137); two paratypes (USNM nos 1608082–1608083). Specimens examined with SEM retained in the JNC's personal collection.

Molecular sequence data. 28S rDNA hologenophore (LRP no. 10109 [EC-54-100 V]) for GenBank no. MN706184.

Remarks

Clistobothrium amyae n. sp. differs conspicuously from all three of its described congeners in its possession of an extremely elongate cephalic peduncle with gladiate spinitriches, rather than a cephalic peduncle that is extremely short as in *C. carcharodoni*

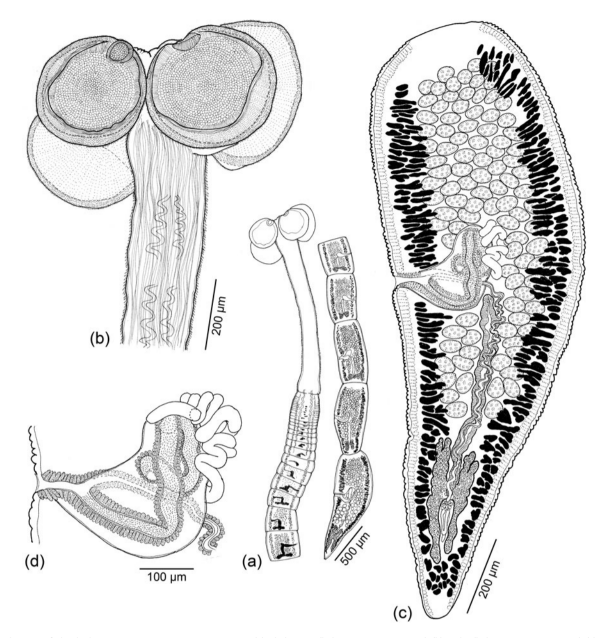


Fig. 6. Line drawings of *Clistobothrium amyae* Caira, Hayes & Jensen n. sp. (a) Whole worm (holotype: MEPN no. 49441). (b) Scolex (holotype: MEPN no. 49441). (c) Terminal mature proglottid (holotype: MEPN no. 49441). (d) Detail of terminal genitalia (holotype: MEPN no. 49441).

Dailey & Vogelbein, 1990 and lacks spinitriches or is essentially lacking as in both *Clistobothrium montaukensis* Ruhnke, 1993 and *Clistobothrium tumidum* (Linton, 1922) Ruhnke, 1993. The apical suckers of the bothridia of this new species are also substantially smaller than those of its three congeners (50–115 vs 280–360 and 310–500 in diameter, respectively in *C. tumidum* and *C. montaukensis*, and 417–461 long by 333–398 wide in *C. carcharodoni*). This new species further differs from *C. tumidum* and *C. montaukensis* in that its bothridia are flat rather than foliose. In addition, *C. amyae* n. sp. is a much shorter worm than both *C. carcharodoni* and *C. montaukensis* (5.6–15.8 vs 24–40 and 38.5–119.5 mm TL, respectively). It also bears many fewer proglottids than *C. montaukensis* and *C. tumidum* (30–66 vs more than 100 in both of the latter species).

Across the D2 region of the 28S rDNA alignment, this new species differs from the undescribed species reported from the porbeagle shark by Randhawa & Brickle (2011) by 22 bp, and, thus, likely represents a distinct species.

Clistobothrium gabywalterorum Caira, Hayes & Jensen n. sp. (figs 7f-j and 8)

ZooBank number for species: urn:lsid:zoobank.org:act: F6B8EDF1-D078-45A0-B185-091BA120FA5E.

Based on one mature worm, two immature worms, one scolex observed with SEM and the whole mount of its strobila. Worms euapolytic, acraspedote, 23.8 (n = 1) mm long; maximum width 1850–2163 (2007 ± 221; 3) at level of scolex; 127 (n = 1) total proglottids per worm. Strobila without distinct dorsomedian muscle band. Scolex consisting of four bothridia and cephalic peduncle; cruciform apical region lacking. Scolex proper 1456–1581 (1512 ± 64; 3) long by 1850–2163 (2003 ± 157; 3) wide. Bothridia

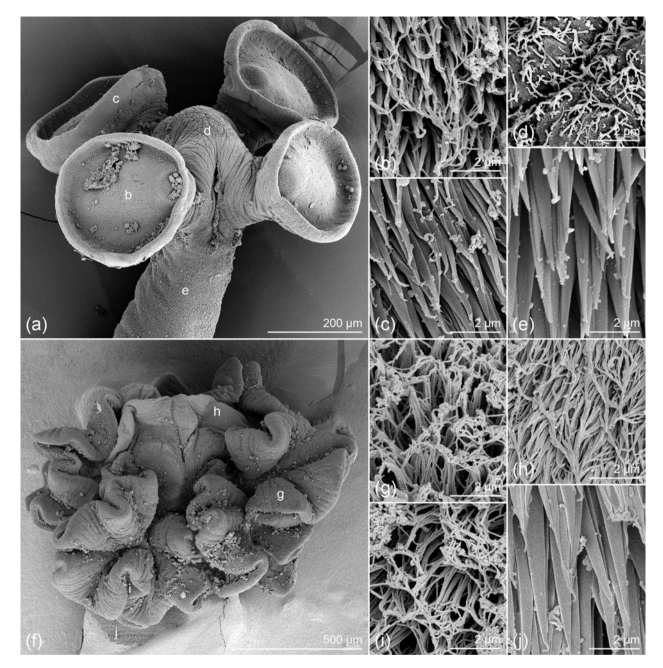


Fig. 7. Scanning electron micrographs of *Clistobothrium amyae* Caira, Hayes & Jensen n. sp. (a) Scolex; small letters indicate locations of details in (b–e). (b) Distal bothridial surface. (c) Proximal bothridial surface. (d) Surface of cruciform apical region. (e) Surface of cephalic peduncle. Scanning electron micrographs of *Clistobothrium gabywalterorum* Caira, Hayes & Jensen n. sp. (f) Scolex; small letters indicate locations of details in (g–j). (g) Distal bothridial surface. (h) Distal surface of apex of bothridia anterior to apical sucker. (i) Proximal bothridial surface. (j) Cephalic peduncle surface.

foliose, 697–1227 (915 ± 222; 3; 8) long by 907–1518 (1277 ± 247; 3; 7) wide; each bothridium with single apical sucker; apical suckers 203–356 (292 ± 48; 3; 10) long by 284–373 (325 ± 35; 3; 11) wide. Cephalic peduncle 2579–3016 (2798 ± 309; 2) long.

Distal surfaces of apex of bothridia anterior to apical sucker with slender gladiate spinitriches and capilliform filitriches (fig. 7h). Distal surfaces of bothridia and apical suckers densely covered with capilliform filitriches (fig. 7g). Proximal bothridial surfaces covered with slender, aristate gladiate spinitriches and capilliform filitriches (fig. 7i). Cephalic peduncle densely covered with gladiate spinitriches (fig. 7j); filitriches not seen in this region. Immature proglottids 123 (n = 1) in number, wider than long; mature proglottids four (n = 1) in number; terminal mature proglottid 1401 (n = 1) long by 591 (n = 1) wide; length:width ratio 2.4:1 (n = 1). Testes 164–185 $(175 \pm 15; 2)$ in total number, 44–51 $(48 \pm 5; 2)$ in postporal field, spherical, 36–54 $(47 \pm 5.8; 1; 6)$ in diameter, arranged in 7–10 irregular columns anterior to cirrus sac, 3–4 irregular columns in poral and aporal fields. Vas deferens minimal, coiled, medial, extending slightly anterior to and posterior to cirrus sac. Cirrus sac J-shaped, 138 (n = 1) long by 203 (n = 1)wide, containing coiled, armed cirrus. Genital pores lateral, irregularly alternating, 68% (n = 1) of proglottid length from posterior end of terminal proglottid. Vagina weakly sinuous, extending

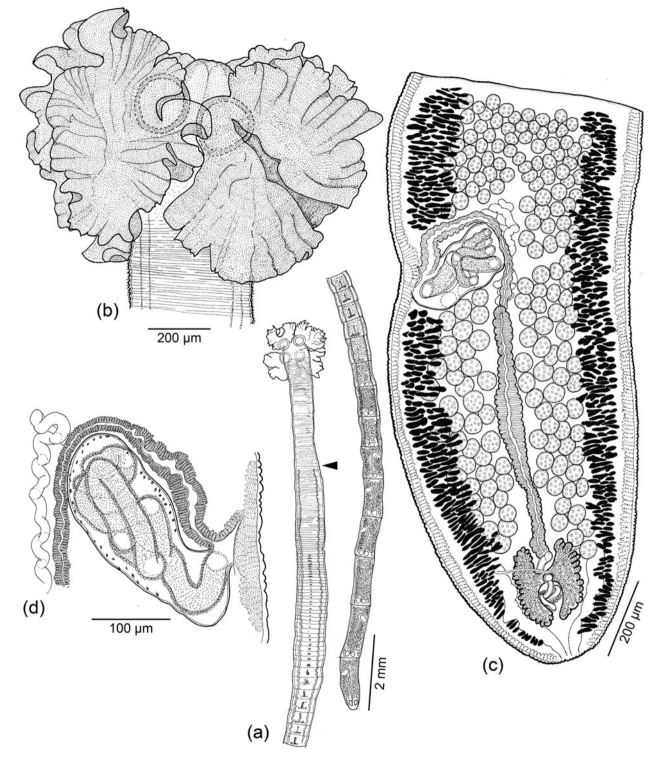


Fig. 8. Line drawings of *Clistobothrium gabywalterorum* Caira, Hayes & Jensen n. sp. (a) Whole worm; arrowhead indicates posterior extent of cephalic peduncle (paratype: USNM no. 1608081). (b) Scolex (holotype: MEPN no. 49440). (c) Terminal mature proglottid (paratype: USNM no. 1608081). (d) Detail of terminal genitalia of terminal mature proglottid (paratype: USNM no. 1608081). (d) Detail of terminal genitalia

from ootype along midline of proglottid to anterior margin of cirrus sac, then laterally along anterior margin of cirrus sac to open into common genital atrium anterior to cirrus. Ovary posterior, H-shaped in frontal view, 210 (n = 1) long by 68 (n = 1) wide, weakly lobate, bilobed in cross section. Mehlis' gland posterior to ovarian bridge. Vitellarium follicular; follicles irregular in shape, arranged in two lateral bands; each band consisting of six to eight columns of follicles, interrupted by the cirrus sac. Uterus ventral, extending from level of ovary to posterior margin of cirrus sac. Four excretory ducts, in one dorsal and one ventral pair.

Taxonomic summary

Type and only known host. Pseudocarcharias kamoharai (Matsubara), crocodile shark (Lamniformes: Pseudocarchariidae).

Site of infection. Spiral intestine.

Type locality, Santa Elena (2°12′24.4″S, 80°56′58.1″W), Santa Rosa de Salinas, eastern Pacific Ocean, Ecuador.

Additional localities. None.

Etymology. This species honours Gabriela del Pilar Flores Rivera and Walter Gilberto Tigrero González of the Ministerio de Producción, Comercio Exterior, Inversiones y Pesca, Ecuador, for sharing their extensive knowledge of local elasmobranch catches and assistance with all aspects of the fieldwork and permitting process that made our collections in Ecuador possible.

Specimens deposited. Holotype (MEPN no. 49440); one immature paratype (LRP no. 10130); SEM voucher (LRP no. 10131); one paratype (USNM no. 1608081). Scolex examined with SEM retained in JNC's personal collection.

Molecular sequence data. 28S rDNA hologenophore (LRP no. 8673 [EC-54-1V]) for GenBank no. MN706183.

Remarks

The description of a new species based on the limited amount of material available here is typically not advisable. However, this new species exhibits clear morphological and molecular differences from its four described congeners. *Clistobothrium gabywalterorum* n. sp. differs conspicuously from *C. carcharodoni* and *C. amyae* in that its bothridia are foliose, rather than flat. In addition, it is a much smaller worm than *C. montaukensis* (23.8 vs 38.5–119.5 mm) and a much larger worm than *C. amyae* (23.8 vs 5.6–15.8 mm). It exhibits a greater number of testes than *C. amyae* and *C. carcharodoni* (164–185 vs 77–127 and 91–123, respectively) and fewer testes than *C. montaukensis* and *C. tumidum* (165–185 vs 198–263 and 234–307, respectively). Unlike all species except *C. amyae*, this new species also possesses a long cephalic peduncle that bears gladiate spinitriches.

Across the 728 bp in the D2 region of the 28S rDNA alignment, which includes data for all five species of *Clistobothrium* for which sequence data are now available, this species differs from *C. carcharodoni* by 11 bp, from *C. montaukensis* by 24 bp, from *C. amyae* by 12 bp and from the undescribed species reported from the porbeagle shark by Randhawa & Brickle (2011) identified as *Clistobothrium* cf. montaukensis by 16 bp.

The most recent diagnosis of *Clistobothrium*, which is that of Ruhnke (2011), is revised below to accommodate both of the above new species.

Clistobothrium Dailey & Vogelbein, 1990 revised

Diagnosis. Phyllobothriidea. Worms apolytic or euapolytic. Strobila with or without distinct longitudinal dorsomedian band of muscles. Scolex with two dorsal and two ventral bothridia, usually with dome-shaped or cruciform apical region. Each bothridium with apical sucker and one flat or foliose loculus, with or without conspicuous stalk. Cephalic peduncle short or elongate. Immature proglottids wider than long; mature proglottids at least twice as long as wide. Testes numerous; postporal testes present. Cirrus sac containing coiled cirrus; cirrus armed with small spinitriches. Genital atrium present. Vagina crossing or extending

anterior to cirrus sac, opening anterior to cirrus sac. Ovary posterior, H-shaped in dorsoventral view, bilobed in cross section. Uterus ventral, extending from ovary to posterior margin of cirrus sac in mature proglottids, extending to anterior margin of cirrus sac in gravid proglottids. Eggs spherical; surface mamillated or spinose.

Type species. Clistobothrium carcharodoni Dailey & Vogelbein, 1990.

Additional species: C. amyae Caira, Hayes & Jensen n. sp., C. montaukensis Ruhnke, 1994, C. tumidum (Linton, 1922) Ruhnke, 1994, C. gabywalterorum Caira, Hayes & Jensen n. sp. Parasites of the spiral intestine of sharks of the order Lamniformes. Cosmopolitan.

Discussion

As of this writing, a total of 45 described species are considered to belong to *Scyphophyllidium*. An additional seven undescribed species that have been included in molecular phylogenetic analyses from previously unexplored host species, should also now be considered to belong to the genus. To avoid future confusion, six of these seven species are formally assigned the following unique numerical designations (see table 1): *Paraorygmatobothrium* sp. 1 through 4 of Cutmore *et al.* (2017) are assigned the designations *Scyphophyllidium* sp. 1 through 4, respectively. *Doliobothrium* sp. of Cutmore *et al.* (2017) is assigned the designation *Scyphophyllidium* sp. 5. *Marsupiobothrium* sp. 1 of Caira *et al.* (2014a) is assigned the designation *Scyphophyllidium* sp. 6. *Scyphophyllidium* cf. *giganteum* of Caira *et al.* (2014a) is considered to represent a distinct, seventh species for the reasons described above.

It is interesting to consider the host associations of these 52 species of *Scyphophyllidium*. By far the majority of the described species (i.e. 32 of 45) parasitize carcharhiniform sharks (i.e. ground sharks). In terms of the remaining 13 described species, seven parasitize orectolobiform sharks (i.e. carpet sharks), five parasitize lamniform sharks (i.e. mackerel sharks) and one parasitizes a freshwater stingray. The majority of the known undescribed species (i.e. six of seven) also parasitize carcharhiniform sharks; the remaining one species parasitizes a lamniform shark.

The topology of the tree resulting from our phylogenetic analysis suggests that instances of associations with hosts other than carcharhiniform sharks represent host-switching events from carcharhiniform sharks in either an immediate or slightly more distant ancestor. At a minimum, there appear to have been two switches to lamniform sharks, two switches to orectolobiform sharks and one switch to a stingray. Given the trophic nature of cestode transmission, it is tempting to suggest these host switches may reflect similarities in diet. Jensen & Bullard (2010), who conducted the most extensive work to date identifying the final intermediate hosts of Scyphophyllidum species formerly assigned to Paraorygmatobothrium, determined that bony fish play this role in the life cycles of all of the species they examined, several of which are among the carcharhiniform-hosted species represented in our study. Indeed, the diets of Orectolobus maculatus (Bonnaterre), Chiloscyllium punctatum Müller & Henle, A. vulpinus (Bonnaterre), A. pelagicus Nakamura and even that of Paratrygon aiereba (Müller & Henle) all include bony fish to a greater or lesser extent (Compagno, 1984; Last & Stevens, 2009; de Carvalho et al., 2003, respectively). However, this does not

explain the presence of *S. randyi* in *Chiloscyllium hasselti* (Bleeker), which does not appear to consume bony fish (Compagno & Neim, 1998). Nor does it explain the lack of reports of this genus from the many other species of Orectolobiformes, Lamniformes and stingrays, which are too numerous to list here, the diets of which also include bony fish.

Table 1 summarizes what is known about the distribution of some of the key morphological and ultrastructural features in the 52 (described and undescribed) species of Scyphyophyllidium, subsets of which were used to establish the six genera now considered to be junior synonyms of the latter genus. The topology of our phylogenetic tree indicates that many of these characters are either homoplasious or unique to single species (i.e. autapomorphies). Examples of homoplasious characters include: the presence of marginal loculi (S. orectolobi, S. janineae, S. randyi and S. tyleri, and S. guariticus); globose rather than flat bothridia (S. cf. giganteum and S. timvickiorum and Scyphophyllidium sp. 6); the presence of proximal bothridial apertures (Scyphophyllidium sp. 5 and S. timvickiorum). An example of an autapomorphic feature is the presence of facial loculi in S. latipi. Also intriguing is the fact that the majority (i.e. ten) of the 13 species of Scyphophyllidium, for which gravid proglottids have been observed, exhibit spindle-shaped eggs. The exceptions are S. barberi and S. guariticus, both of which have spherical eggs and S. leuci with eggs that Watson & Thorson (1976) reported were either with or without knobs. It would be interesting to determine the full extent of spindle-shaped eggs across the other species of Scyphophyllidium.

Evidence supporting the close affinities among at least subsets of the genera synonymized here with Scyphophyllidium also comes from SEM. The majority of these species exhibit the somewhat unusual conditions of capilliform filitriches on the strobila that are arranged in scutes, serrate gladiate spinitriches on the proximal bothridial surfaces and serrate gladiate, gongylate columnar or gongylate gladiate spinitriches on their distal bothridial surfaces. To our knowledge, the only other cestode taxa that possess one or more of these ultrastructural features for which sequence data are also available, are species of Alexandercestus, Guidus, Hemipristicola, Orygmatobothrium Diesing, 1863 and, possibly also in modified form (see below), Thysanocephalum Linton, 1890. The topology of our molecular phylogenetic tree suggests that Thysanocephalum is the sister taxon of the clade consisting of Scyphophyllidium + Hemipristicola + Alexandercestus, in which case, all three ultrastructural features may have originated in the shared common ancestor of these four genera. In contrast, the presence of these features in the more distantly related Orygmatobothrium appears to be homoplasious.

We have taken a relatively conservative approach here with respect to the genera we have synonymized with *Scyphophyllidium*. However, in the future, serious consideration should be given to whether *Hemipristicola* and possibly also *Alexandercestus* should also be synonymized with *Scyphophyllidium*. Beyond sharing subsets of the above unique ultrastructural features with *Scyphophyllidium*, Cutmore *et al.* (2017) found the monotypic *Hemipristicola* to nest deeply among species now assigned to *Scyphyophyllidium* in the trees resulting from both their Bayesian and ML phylogenetic analyses of NADH1 amino acid data. Morphologically, *H. gunterae* differs from species of *Scyphophyllidium* in its possession of a deep central cavity in each of its bothridia. But, it is possible this feature will ultimately also be found to represent an autapomorphy for this species. Both species of *Alexandercestus* can be distinguished from existing

members of *Scyphophyllidium* in their possession of foliose bothridia, but the bothridia of *Alexandercestus manteri* Ruhnke & Workman, 2013 are only weakly foliose. It will be interesting to see the results of future phylogenetic work that includes *A. manteri*. Fortunately, even if both genera are ultimately determined to be synonyms of *Scyphophyllidium*, the latter remains the oldest, and thus valid, name for the genus. Although *Guidus* shares highly muscular, globose bothridia, and filitriches arranged in scutes with subsets of species of *Scyphophyllidium*, its placement well outside of all of these taxa in the tree resulting from our phylogenetic analysis indicates that these features are homoplasious in this skate-hosted taxon.

In contrast, the bothridia of *Thysanocephalum* are distinctive in consisting of 'a small specialized anterior loculus followed by an extensive posterior loculus that is narrow at its connection to the anterior loculus, but expands almost immediately into a large, extensively folded, bifid structure' (Caira *et al.*, 1999: 103). Furthermore, rather than scutes, the surfaces of the strobila of *T. thysanocephalum* bear 'leaf-like' structures (Caira *et al.*, 1999: 125). In combination, these distinctive morphological features and the topology of our molecular phylogenetic tree justify retaining this as a valid genus.

The placement of six phyllobothriidean genera into synonymy with Scyphophyllidium was a major action that necessitated substantial revision of the classification of the order. While the molecular phylogenetic analyses (Cutmore et al., 2011, 2017; Caira, et al., 2014a; Ruhnke et al., 2020; our analyses here) supporting this action have all been based on data from a limited selection of genes (i.e. D1-D3 of 28S rDNA, 18S rDNA and/or NADH1), the taxon sampling has been relatively robust. In terms of described species, our analysis included the only species originally assigned to Hemipristicola, one of two species originally assigned to Marsupiobothrium, the only species originally assigned to Nandocestus, two of five species originally assigned to Orectolobicestus, the only species originally assigned to Ruhnkecestus, one of two species originally assigned to Scyphophyllidium. In addition, our analysis included 22 of the 34 species previously assigned to Paraorygmatobothrium, with representation from all three orders of sharks known to host species previously assigned to this genus. With respect to undescribed species, our taxon sampling included one species originally assigned to Doliobothrium, one originally assigned to Marsupiobothrium, one originally assigned to Scyphophyllidium and four species originally assigned to Paraorygmatobothrium. Although we believe this work has paved the way for the development of a more thorough understanding of the evolution and host associations of these cestodes, a larger sampling of molecular markers is necessary for the elucidation of robust clades within the genus.

The shark order Lamniformes is unusual among elasmobranchs in its extremely high ratio of families to genera – the 15 extant species of lamniforms are currently assigned to nine genera in seven families, four of which are monotypic. Work to date on the cestodes of lamniforms indicates that the cestode faunas of each family are highly divergent relative to one another (Linton, 1889, 1922; Yamaguti, 1935, 1952; Dailey, 1969, 1971; Kurochkin & Slankis, 1973; Beveridge & Campbell, 1988; Dailey & Vogelbein, 1990; Caira & Runkle, 1993; Ruhnke, 1993, 2011; Palm, 2004; Caira *et al.*, 1997, 2014b; Olson & Caira, 2001). Our interest in examining the cestodes of the monotypic Pseudocarchariidae was motivated largely by the fact that this family had not been examined for cestodes. This host species has eluded examination previously in large part because, unlike many of the other lamnid species, its flesh generally has little appeal for human consumption (Compagno, 1984) and, thus, this shark is infrequently landed in fish markets around the world. Our arrival in Ecuador during what is locally considered to be 'crocodile shark season' (i.e. May and early June) when this species is landed, at least in the region of Santa Elena, was thus, extremely fortuitous.

The two new species of *Clistobothrium* reported here bring the total number of described species to five; Randhawa & Brickle's (2011) report of the undescribed species *C. cf. montaukensis* expands the total to six. While the two species described here parasitize the monotypic Pseudocarchariidae, the remaining four species parasitize members of the Lamnidae – specifically, *Carcharodon carcharias* L., *Isurus oxyrinchus* Rafinesque and *Lamna nasus* Bonnaterre. Thus, it would be extremely interesting to examine the two remaining species of lamnids (i.e. *Isurus paucus* Guitart and *Lamna ditropis* Hubbs & Follett), neither of which has been examined for *Clistobothrium*. We believe both are highly likely to host additional members of the genus.

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Conflicts of interest. None.

Ethical standards. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and international guides on the care and use of animals.

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