

Research Article

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
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Author for correspondence:

Kaitlyn M. Dalrymple,
E-mail: kaitlyn.dalrymple@gmail.com

Hexabothriidae and Monocotylidae (Monogenoidea) from the gills of neonate hammerhead sharks (Sphyrnidae) *Sphyrna gilberti*, *Sphyrna lewini* and their hybrids from the western North Atlantic Ocean

Kaitlyn M. Dalrymple^{1,2} , Isaure de Buron², Kristina M. Hill-Spanik², Ashley S. Galloway³, Amanda Barker⁴, David S. Portnoy⁴, Bryan S. Frazier³ and Walter A. Boeger¹

¹Laboratory of Biological Interactions, Universidade Federal do Paraná, Curitiba, PR 81531-980, Brazil; ²Department of Biology, College of Charleston, Charleston, SC 29424, USA; ³South Carolina Department of Natural Resources, Marine Resources Research Institute, 217 Fort Johnson Drive, Charleston, SC 29412, USA and ⁴Marine Genomics Laboratory, Department of Life Sciences, Texas A&M University-Corpus Christi, 6300 Ocean Drive, Corpus Christi, TX 78412, USA

Abstract

Neonates of hammerhead sharks (Sphyrnidae), *Sphyrna lewini* (Griffith and Smith, 1834), the sympatric cryptic species, *Sphyrna gilberti* Quattro *et al.*, 2013, and their hybrids were captured in the western North Atlantic, along the coast of South Carolina, USA, between 2018 and 2019 and examined for gill monogenoids. Parasites were identified and redescribed from the gills of 79 neonates, and DNA sequences from partial fragments of the nuclear 28S ribosomal RNA (rDNA) and cytochrome c oxidase I mitochondrial DNA (COI) genes were generated to confirm species identifications. Three species of monogenoids from Hexabothriidae Price, 1942 and Monocotylidae Taschenberg, 1879 were determined and redescribed. Two species of Hexabothriidae, *Erpocotyle microstoma* (Brooks, 1934) and *Erpocotyle sphyrnae* (MacCallum, 1931), infecting both species of *Sphyrna* and hybrids; and 1 species of Monocotylidae, *Loimosina wilsoni* Manter, 1944, infecting only *S. lewini* and hybrids. *Loimosina wilsoni* 28S rDNA sequences matched those of *Loimosina* sp. from the southern coast of Brazil. Based on limited morphological analysis, *Loimosina parawilsoni* is likely a junior synonym of *L. wilsoni*. This is the first taxonomic study of monogenoids infecting *S. gilberti* and hybrids of *S. gilberti* and *S. lewini*.

Introduction

Two cryptically similar and sister species of hammerhead sharks (Sphyrnidae), *Sphyrna lewini* (Griffith and Smith, 1834) and *Sphyrna gilberti* Quattro *et al.*, 2013 occur along the coast of South Carolina (SC), USA (Quattro *et al.*, 2013). These species are commonly known as the scalloped and Carolina hammerhead, respectively. They can be distinguished only by a difference in the number of precaudal vertebrae (*S. gilberti* has about 10 fewer than *S. lewini*) and genetic data. Their divergence was estimated to have occurred 4.5 million years ago (Pinhal *et al.*, 2012; Quattro *et al.*, 2013). In addition, these species are capable of hybridization (Barker *et al.*, 2019). Because of the morphological similarity, *S. gilberti* was described only recently from the Carolina coasts (South and North Carolina states, USA) in the western North Atlantic Ocean (Quattro *et al.*, 2013). Hence, knowledge about its biology and distribution is limited, and adults' range remains uncertain. However, specimens of *S. gilberti* have been found along the Atlantic coast of Florida (FL), prior to being officially recognized as a distinct species (Abercrombie *et al.*, 2005; Duncan *et al.*, 2006; Quattro *et al.*, 2006). Barker *et al.* (2021) showed that neonates of *S. gilberti* are most common along coastal nurseries in SC, with abundances decreasing latitudinally to be at its lowest in southern FL, and were not found in the Gulf of Mexico. Furthermore, at least 3 adult specimens of *S. gilberti* were captured in southeast Brazil (Pinhal *et al.*, 2012). Knowledge of the parasite fauna of *S. gilberti* is sparse. Presently, the only parasite known for the Carolina hammerhead is a nematode in the spiral valve (Moravec *et al.*, 2020), although it is possible that specimens of *S. gilberti* have been unknowingly included in previous parasitic fauna studies of *S. lewini* in geographic locales where both species occur, and vice versa.

By contrast, the scalloped hammerhead, *S. lewini*, has been formally recognized as a species since the 19th century. It has a global range (Compagno, 1984). Five species of monogenoids were described from this shark. Two of these species are members of Hexabothriidae Price, 1942: *Erpocotyle microstoma* (Brooks, 1934), originally described in *Sphyrna zygaena* (Linnaeus, 1758) (type host) from the coast of Beaufort, North Carolina (NC), USA and reported in *S. lewini* from the western South Atlantic Ocean (Uruguay; Suriano and

Labriola, 1998); and *Erpocotyle sphyrynae* (MacCallum, 1931), also described from *S. zygaena* off Woods Hole, Massachusetts (MA), USA and reported in *S. lewini* from the eastern North Atlantic Ocean (Senegal; Euzet and Maillard, 1967) and the Pacific Ocean (Hawaii, USA; Yamaguti, 1968). The other 3 species are members of Monocotyliidae Taschenberg, 1879: *Cathariotrema selachii* (MacCallum, 1916), originally described from *S. zygaena* (exact locale unknown, but is likely off Woods Hole, MA – see Bullard *et al.*, 2021) and reported in *S. lewini* from the Northern Gulf of Mexico off Mississippi, Alabama and Louisiana in Bullard *et al.* (2021); *Loimosina wilsoni* Manter, 1944, originally described from *S. zygaena* from Montego Bay, Jamaica and reported in *S. lewini* from Alligator Harbor, FL, USA in Hargis (1955); and *Loimosina parawilsoni* Bravo-Hollis, 1970, described in *S. lewini* from the eastern tropical Pacific Ocean (Sinaloa, Mexico; Bravo-Hollis, 1970) (the latter 2 were formerly considered Loimoidae – see Boeger *et al.*, 2014).

In the present study, monogenoids infecting the gills of neonates of *S. gilberti*, *S. lewini* and their hybrids were identified, sequenced, illustrated and redescribed based on specimens collected from the western North Atlantic Ocean (SC, USA) and their types.

Materials and methods

Sampling and collection

A total of 87 neonates, all moribund upon capture, were collected. Because the shark species identification could not be determined at time of capture, the number of sharks sampled was necessary to gain a sufficient sample size for both species of hammerheads, *S. lewini* and *S. gilberti*, and their hybrids.

Sharks were captured using a 231 m long, 3 m deep gillnet with a stretched mesh of 10.3 cm in Bulls Bay, SC (Five Fathom Creek, 33.0095/-79.4853), a nursery area where both species of hammerhead and their hybrids are found in sympatry (Barker *et al.*, 2019). Fresh carcasses were kept on ice, individually labelled and fin clips from each specimen preserved in 20% salt-saturated DMSO and sent to the Marine Genomics Laboratory at Texas A&M University – Corpus Christi, USA, for molecular identification following the methods of Barker *et al.* (2019).

Monogenoids from each hammerhead specimen were identified before the host species identifications were determined. Gills were resected from each host within 10 h post-capture, flooded and shaken rapidly in hot water (68°C) to relax, kill and detach worms from the gill filaments. Some monogenoids were processed immediately – the haptor was fixed in 95% ethanol (EtOH) and the anterior end in 10% neutral buffer formalin (NBF) to generate hologenophores *sensu* Pleijel *et al.* (2008); the remaining specimens were fixed with either 10% NBF or 100% EtOH to obtain final concentrations of 5% NBF and 70% EtOH, respectively. Other hologenophores were generated *via* some EtOH-fixed specimens by removing a small lateral part of their body.

Morphology

Parasites were stained with either acetocarmine or Gomori's trichrome (Humason, 1979) and mounted using Canada balsam or Permunt Mounting Medium (Fisher Chemical, Fairlawn, New Jersey, USA). The haptor of some specimens were mounted separately in Hoyer's medium to examine sclerite and anchor morphology. Drawings were made using an Olympus BX50 differential interference contrast compound microscope mounted with camera lucida. Measurements of the haptoral hook-like sclerites and anchors were obtained using ImageJ software

(www.nih.org) following measurements schemes of MacCallum (1931), Euzet and Maillard (1967) and Bullard and Dippenaar (2003) (Fig. 1A–E). Anchors were measured as indicated in Fig. 2. Measurements are in micrometres; the range is presented followed by the average and the number of measured structures (*n*) in parentheses. Haptoral sucker pairs and their respective sclerites are numbered 1–3, with 1 being closest to the point of attachment for the haptoral appendix.

Vouchers and hologenophores are deposited at the National Museum of Natural History, Smithsonian Institution (USNM) in Washington DC, USA, and the Harold W. Manter Laboratory of Parasitology (HWML) in Nebraska, USA.

For comparison, micrographs of the following specimens from USNM, HWML and Nacional Collection of Helminths, Institute of Biology, National Autonomous University of Mexico (CNHE), Mexico (available at <http://unibio.unam.mx>) were examined: *E. microstoma* (syntypes: USNM 132155, HWML 1437), *E. sphyrynae* (syntypes: USNM 1320885 and USNM 1320884), *L. wilsoni* (syntypes: USNM 1337561, USNM 1337564, HWML 1425) and *L. parawilsoni* (holotype: CNHE 153, syntype CNHE 154).

Molecular analyses

DNA was extracted from parasite tissue using a DNeasyBlood and Tissue kit (Qiagen, Valencia, California, USA) and manufacturer's protocol. Sequences of DNA from portions of the nuclear-encoded 28S ribosomal RNA (rDNA; 28S) and the mitochondrially encoded cytochrome *c* oxidase I (COI) genes were amplified and sequenced for species comparisons and confirmations. Primers LSU5 (5'-TAGGTCGACCCGCTGAAYTTAAGCA-3'; Jensen and Bullard, 2010) and 28S_ECD2 (5'-CTTGGTCCGTGTTTCAAGACGGG-3'; Tkach *et al.*, 2003) were used to amplify a 1100 base pair portion of the 28S gene: a 25 μ L total volume reaction contained 1 \times polymerase chain reaction (PCR) buffer (Promega, Madison, Wisconsin, USA), 3 mM MgCl₂, 0.4 mM deoxynucleotide triphosphates (dNTPs) (Promega), 0.4 \times Rediload gel loading buffer (Invitrogen, Waltham, MA, USA), 0.6 μ M of each primer, 1 U GoTaq[®] Flexi DNA Polymerase (Promega) and 3 μ L template DNA. Cycling was as follows: 5 min initial denaturation at 94°C, followed by 40 cycles of denaturation at 94°C for 45 s, annealing at 62°C for 45 s, extension at 72°C for 45 s, then a final extension at 72°C for 5 min. Primers JB3 (5'-TTTTTTGGGCATCCTGAGGTTTAT-3') and JB4.5 (5'-TAAAGAAAGAACATAATGAAAATG-3'; Bowles *et al.*, 1995) were used to amplify a portion of COI: a 25 μ L total volume reaction contained 1 \times PCR buffer (Promega), 3 mM MgCl₂, 0.2 mM dNTPs (Promega), 0.4 \times Rediload gel loading buffer (Invitrogen), 0.2 μ M of each primer, 1 U GoTaq[®] Flexi DNA Polymerase (Promega) and 1 μ L template DNA. Cycling was as follows: 4 min at 94°C, followed by 40 cycles at 94°C for 30 s, annealing at 48°C for 40 s, extension at 72°C for 50 s and final extension at 72°C for 7 min.

All products were electrophoresed through 1% agarose gels stained with GelRed (Biotium, Fremont, California, USA) and visualized under ultraviolet light. Samples that produced a faint band were subjected to another round of PCR, done as above, except the template was the product from the initial PCR. Products were purified using ExoSAP-IT (Affymetrix, Santa Clara, California, USA) or by gel extraction using a QIAquick Gel Extraction kit (Qiagen), following the manufacturer's protocol for both methods and then sent to Eurofins MWG Operon LLC (Louisville, Kentucky, USA) for bi-directional Sanger sequencing. Complementary sequences were assembled, and their chromatograms were assessed and edited by eye using Sequencher version 5.3 (Gene Codes Corp., Ann Arbor, Michigan, USA). Resulting sequences were compared with sequences available in the National Center for Biotechnology

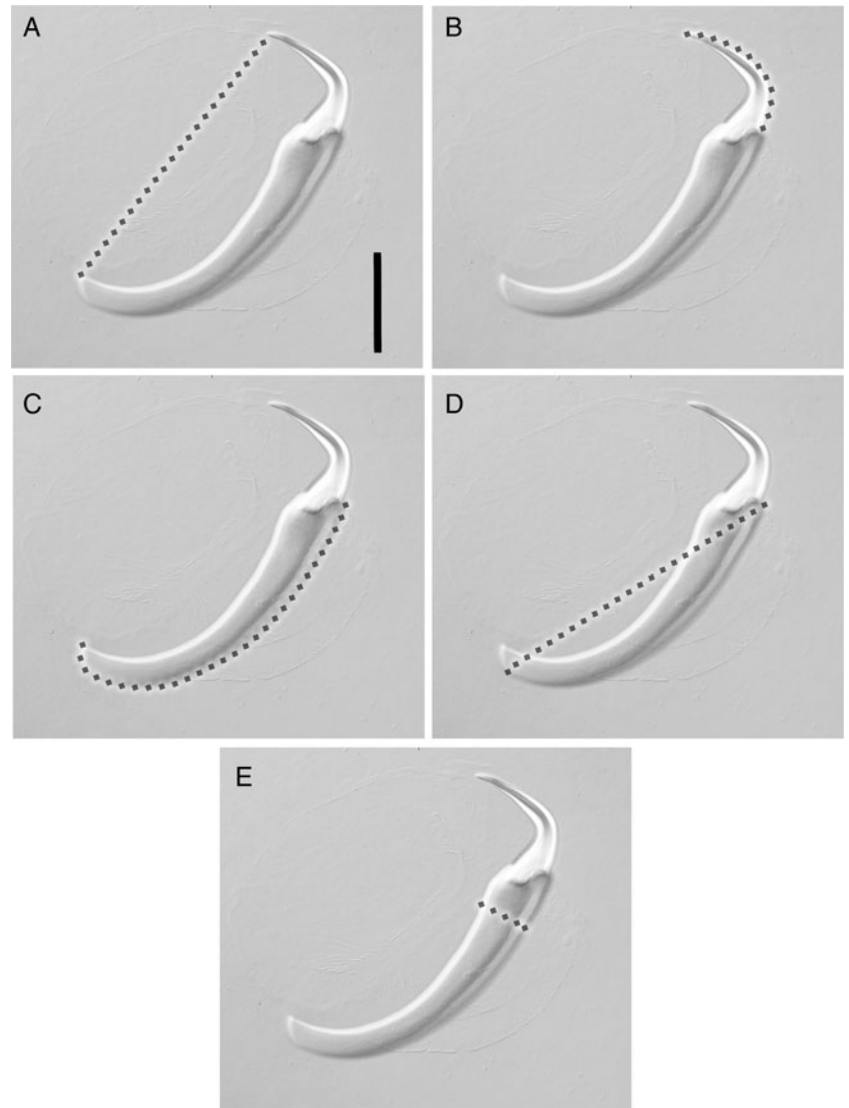


Fig. 1. Measurements of sclerite curvature were taken using ImageJ software (www.nih.org): (A) tip-to-tip (MacCallum, 1931), (B) perimeter hook length (Euzet and Maillard, 1967), (C) perimeter shaft length (Euzet and Maillard, 1967), (D) shaft length (Bullard and Dippenaar, 2003) and (E) max shaft width (Bullard and Dippenaar, 2003). Scale bar 80 μm .

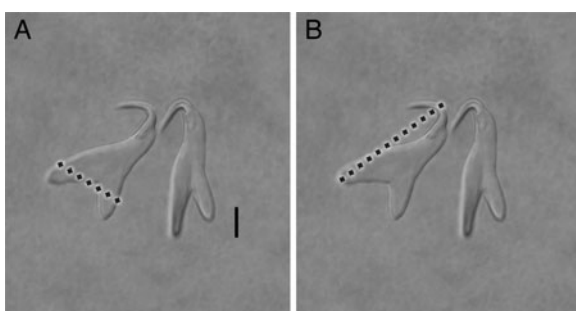


Fig. 2. Width (A) and length (B) measurements of anchors of the haptoral appendix. Scale bar 13 μm .

Information (NCBI) GenBank database using the Basic Local Alignment Search Tool (BLAST – Altschul *et al.*, 1990) and were deposited into GenBank (Table 1).

COI sequences were translated to confirm the absence of premature stop codons, which if present were corrected using Open Reading Frame Finder (RRID:SCR_016643) by NCBI and Geneious software version 4.8.5 (Kearse *et al.*, 2012). Sequences produced in this study and sequences of species of Hexabothriidae and Monocotylidae obtained from GenBank (Table 1) were aligned by hand in MEGA X (Kumar *et al.*, 2018), and distance analyses

were done using the same program. The maximum composite likelihood method (Tamura *et al.*, 2004) was used to calculate the number of base substitutions per site between pairs of sequences (*p*-distance), and phylograms were generated using the neighbour-joining (NJ) method (Saitou and Nei, 1987), implementing pairwise deletions and support values obtained by 1000 bootstrap replicates (Felsenstein, 1985). Resulting trees represent intraspecific and interspecific distances between sequences but not phylogenetic relationships (Figs 3 and 4).

A maximum likelihood phylogenetic tree for Monocotylidae was constructed using 28S sequences obtained herein and from GenBank (see Table 1 for accession numbers) – the only fragment available for the group at the moment for species previously allocated in Loimoidae. Sequences were aligned using GUIDANCE 2 (Penn *et al.*, 2010a, 2010b) using the multiple sequence alignment algorithm MAFFT (Katoh *et al.*, 2002) and set to 100 alternative guide trees. The phylogeny was generated using raxmlGUI 2.0 (version 2.0.6) (Silvestro and Michalak, 2012; Stamatakis, 2014) with the general time reversible model with gamma rates (GTR+G) for 1000 bootstrap repetitions. The final tree was opened and edited in MEGA X (Fig. 5).

Genomic DNA was extracted from hosts using a Mag-Bind Blood and Tissue DNA Kit (Omega Bio-Tek, Norcross, Georgia, USA). Double-digest restriction site-associated DNA sequencing libraries were prepared following the methods of Barker *et al.* (2019) and sequenced on an Illumina HiSeq 4000

Table 1. Sequences generated and used in the present study from families Monocotylidae and Hexabothriidae

	GenBank no.		Reference(s)	Notes
	COI	28S		
Monocotylidae Taschenberg, 1879				
<i>Calicotyle affinis</i> Scott, 1911	–	AF382061	Chisholm <i>et al.</i> (2001a)	
<i>Calicotyle hydrolagi</i> Ñacari <i>et al.</i> , 2019	–	MK659587	Ñacari <i>et al.</i> (2020)	
<i>Calicotyle japonica</i> Kitamura <i>et al.</i> , 2010	–	AB485996	Kitamura <i>et al.</i> (2010)	
<i>Calicotyle kroyeri</i> Diesing, 1850	–	AF279748 AF279744 AF279745 AF279746 AF279747, MW892410	Chisholm <i>et al.</i> (2001a); Bullard <i>et al.</i> (2021)	
<i>Calicotyle palombi</i> Euzet and Williams, 1960	–	AF279749 AF131709	Chisholm <i>et al.</i> (2001a); Mollaret <i>et al.</i> (2000)	
<i>Calicotyle</i> sp.	–	AF279750 FJ971978	Chisholm <i>et al.</i> (2001a); Perkins <i>et al.</i> (2009)	
<i>Calicotyle stossichi</i> Braun, 1899	–	AF279751	Chisholm <i>et al.</i> (2001a)	
<i>Calicotyle urolophi</i> Chisholm <i>et al.</i> , 1991	–	AF279753 AF279752	Chisholm <i>et al.</i> (2001a)	
<i>Cathariotrema selachii</i> (MacCallum, 1916)	–	MW892407 MW892406 MW892405 MW892404	Bullard <i>et al.</i> (2021)	
<i>Clemacotyle australis</i> Young, 1967	–	AF348350	Chisholm <i>et al.</i> (2001b)	
<i>Decacotyle floridana</i> (Pratt, 1910)	–	AF348357	Chisholm <i>et al.</i> (2001b)	
<i>Decacotyle lymmae</i> Young, 1967	–	AF348359	Chisholm <i>et al.</i> (2001b)	
<i>Decacotyle tetrakordyle</i> Young, 1967	–	AF348358	Chisholm <i>et al.</i> (2001b)	
<i>Dendromonocotyle ardea</i> Chisholm and Whittington, 1995	–	AF348351	Chisholm <i>et al.</i> (2001b)	
<i>Dendromonocotyle bradsmithi</i> Chisholm <i>et al.</i> , 1995	–	FJ971986	Perkins <i>et al.</i> (2009)	
<i>Dendromonocotyle octodiscus</i> Hargis, 1955	–	AF348352	Chisholm <i>et al.</i> (2001b)	
<i>Dictyocotyle coeliaca</i> Nybelin, 1941	–	AF279754 AF382062, AY157171	Chisholm <i>et al.</i> (2001b); Lockyer <i>et al.</i> (2003)	
<i>Electrocotyle whittingtoni</i> Vaughan <i>et al.</i> , 2016	–	KT735369 KT735368	Vaughan <i>et al.</i> (2016)	
<i>Empruthotrema aoneken</i> Irigoitia <i>et al.</i> , 2019	MN190708, MN190709	MN190270 MN190271 MN190272	Irigoitia <i>et al.</i> (2019)	
<i>Empruthotrema dasyatidis</i> Whittington and Kearn, 1992	–	AF348345	Chisholm <i>et al.</i> (2001b)	
<i>Empruthotrema doriae</i> Irigoitia <i>et al.</i> , 2019	MN190712, MN190711	MN190274 MN190273	Irigoitia <i>et al.</i> (2019)	
<i>Empruthotrema longipenis</i> Kritsky <i>et al.</i> , 2017	–	MW892409	Bullard <i>et al.</i> (2021)	
<i>Empruthotrema orashken</i> Irigoitia <i>et al.</i> , 2019	MN190702, MN190704, MN190705	MN190265 MN190266 MN190269 MN190268 MN190267 MN190264	Irigoitia <i>et al.</i> (2019)	
<i>Empruthotrema quindecima</i> Chisholm and Whittington, 1999	–	AF348346	Chisholm <i>et al.</i> (2001b)	
<i>Heterocotyle capricornensis</i> Chisholm and Whittington, 1996	–	AF348360	Chisholm <i>et al.</i> (2001b)	
<i>Loimopapillosum pascuali</i> Chero <i>et al.</i> , 2021	–	MZ367714 MZ367713	Chero <i>et al.</i> (2021)	

(Continued)

Table 1. (Continued.)

	GenBank no.		Reference(s)	Notes
	COI	28S		
<i>Loimosina wilsoni</i> Manter, 1944	OP342748, OP342749, OP342750, OP342751, OP342752, OP342753	OP348870, OP348871, OP348872, OP348873	Present study	
<i>Loimosina</i> sp.	–	KF908848	Boeger et al. (2014)	
<i>Thaumatoctyle australensis</i> Beverley-Burton and Williams, 1989	–	AF348348	Chisholm et al. (2001b)	Recorded in GenBank as <i>Merizocotyle australensis</i>
<i>Mycteranastes icopae</i> (Beverley-Burton and Williams, 1989)	–	AF026113, AF348349	Mollaret et al. (1997); Chisholm et al. (2001b)	Recorded in GenBank as <i>Merizocotyle icopae</i>
<i>Merizocotyle sinensis</i> Timofeeva, 1984	–	FJ514075	Unpublished	
<i>Thaumatoctyle urolophi</i> (Chisholm and Whittington, 1999)	–	AF348347	Chisholm et al. (2001b)	Recorded in GenBank as <i>Merizocotyle urolophi</i>
<i>Monocotyle corali</i> Chisholm, 1998	–	AF348353	Chisholm et al. (2001b)	
<i>Monocotyle helicophallus</i> Measures, Beverley-Burton and Williams, 1990	–	AF348355	Chisholm et al. (2001b)	
<i>Monocotyle multiparous</i> Measures, Beverley-Burton and Williams, 1990	–	AF348356	Chisholm et al. (2001b)	
<i>Monocotyle</i> sp.	–	AF387511	Chisholm et al. (2001b)	
<i>Monocotyle spiremae</i> Measures, Beverley-Burton and Williams, 1990	–	AF348354	Chisholm et al. (2001b)	
<i>Neoheterocotyle quadrispinata</i> Nitta, 2019	–	LC428038	Nitta (2019)	
<i>Neoheterocotyle rhinobatidis</i> (Young, 1967)	–	AF026107, AF348361 AF348362	Mollaret et al. (1997); Chisholm et al. (2001b)	
<i>Neoheterocotyle rhynchobatis</i> (Tripathi, 1959)	–	AF348363	Chisholm et al. (2001b)	
<i>Potamostrygonocotyle aramasae</i> (Young, 1967)	–	FJ755806 FJ755804 FJ755805, JN379514	Unpublished; Fehlauer-Ale and Littlewood (2011)	
<i>Potamostrygonocotyle chisholmae</i> Domingues and Marques, 2007	–	JN379519 JN379516 JN379515	Fehlauer-Ale and Littlewood (2011)	
<i>Potamostrygonocotyle dromedarius</i> Domingues and Marques, 2007	–	JN379518 JN379517	Fehlauer-Ale and Littlewood (2011)	
<i>Potamostrygonocotyle quadracotyle</i> Domingues, Pancera and Marques, 2007	–	FJ755807	Unpublished	
<i>Potamostrygonocotyle rara</i> Domingues, Pancera and Marques, 2007	–	FJ755809	Unpublished	Recorded as <i>Potamostrygonocotyle</i> <i>raram</i> in GenBank
<i>Potamostrygonocotyle rionegrensis</i> Domingues, Pancera and Marques, 2007	–	FJ755810	Unpublished	Recorded as <i>Potamostrygonocotyle</i> <i>rionegrense</i> in GenBank
<i>Potamostrygonocotyle tsalickisi</i> Mayes, Brooks and Thorson, 1981	–	JN379513	Fehlauer-Ale and Littlewood (2011)	
<i>Potamostrygonocotyle umbella</i> Domingues, Pancera and Marques, 2007	–	FJ755808	Unpublished	
<i>Thaumatoctyle</i> sp.	–	MW892408	Bullard et al. (2021)	
<i>Triloculotrema</i> sp.	–	AF387512	Unpublished	Boudaya and Neifar, 2016 reported this sequence corresponded to their new species <i>T. euzeti</i> Boudaya and Neifar, 2016

(Continued)

Table 1. (Continued.)

	GenBank no.		Reference(s)	Notes
	COI	28S		
<i>Troglocephalus rhinobatidis</i> Young, 1967	–	AF026110, AF348364	Mollaret <i>et al.</i> (1997); Chisholm <i>et al.</i> (2001b)	
Gyrodactylidae Cobbold, 1864				
<i>Bothitrema bothi</i> (MacCallum, 1913)	–	AF387508	Justine <i>et al.</i> (2002)	
<i>Gyrodactylus ticuchi</i> Pinacho-Pinacho <i>et al.</i> , 2021	–	MT879662.1	Pinacho-Pinacho <i>et al.</i> (2021)	
<i>Gyrodactylus tobala</i> Pinacho-Pinacho <i>et al.</i> , 2021	–	MT879660, MT879661	Pinacho-Pinacho <i>et al.</i> (2021)	
<i>Tetraonchus monenteron</i> (Wagener, 1857)	–	MK881304	Unpublished	
Hexabothriidae Price, 1942				
<i>Dasyonchocotyle</i> sp. Hargis, 1955	MT890380	–	Unpublished	
<i>Erpocotyle microstoma</i> (Brooks, 1934)	OP342755, OP342756, OP342757, OP342758, OP342759, OP342760, OP342761, OP342762, OP342763, OP342764, OP342765, OP342766, OP342767, OP342768, OP342769, OP342770, OP342771	–	Present study	
<i>Erpocotyle sphyrnae</i> (MacCallum, 1931)	OP342754, OP342755	–	Present study	
<i>Hexabothrium</i> sp. (Kuhn, 1829)	MT890381, MT890382	–	Unpublished	
<i>Narcinecotyle longifilamentus</i> Torres-Carrera <i>et al.</i> , 2020	MN367806, MN367807	–	Torres-Carrera <i>et al.</i> (2020)	
<i>Squalonchocotyle euzeti</i> Kheddam <i>et al.</i> , 2016	KX389260, KX389261, KX389262	–	Kheddam <i>et al.</i> (2016)	

COI sequences were used for distance analysis (species delimitation) and 28S sequences were used for phylogenetic analysis of the family Monocotylidae.

DNA sequencer. Subsequent data processing and species identification was performed following the methods of Barker *et al.* (2021). Briefly, the dDocent pipeline (Puritz *et al.*, 2014) was used to trim reads, map reads and call single-nucleotide polymorphisms (SNPs). Hosts were identified as *S. lewini* or *S. gilberti* using a panel of 1491 diagnostic SNPs, and a match of at least 95% to 1 species was required for identification. The program NewHybrids (Anderson and Thompson, 2002) was used to determine if ambiguous individuals could be assigned into a hybrid category [first-generation hybrid (F1), *S. lewini* backcross (BX), *S. gilberti* backcross].

Prevalence determination

Upon receipt of shark identifications, prevalence of infection per parasite species, including coinfections, as well as per family, was calculated. Coinfections included instances where specimens of multiple species were found to infect the same host individual. Monogenoid prevalence is the per cent of infected sharks, regardless of the parasite

species; Hexabothriidae and Monocotylidae prevalence includes all species from their respective families. In some instances, only fragments of specimens were collected but could not be identified to species or even family level, in which case these were included within the category of highest taxonomic classification possible.

Results

Species determination and prevalence of infection

Of the 87 neonates examined, 44 were determined molecularly as *S. gilberti*, 20 as *S. lewini* and 15 as hybrids of the 2 species (6 F1 hybrids, 5 *S. lewini* BX and 4 *S. gilberti* BX). Eight sharks could not be determined at the species level (i.e. no DNA or morphology available could ascertain *Sphyrna* species, as only the head of these specimens was provided and attempts to sequence were unsuccessful). Thus, only the remaining 79 neonates with confirmed species identities were used to report prevalence (Table 2). In total, 3 species of monogenoids were determined as *E. microstoma*, *E. sphyrnae*, and *L. wilsoni* (see

redescriptions below). Both species of *Erpocotyle* were found to infect both species and hybrids of the studied hammerhead sharks, while *L. wilsoni* was found only on specimens of *S. lewini* and hybrids.

Molecular species identity

In total, 22 COI and 4 28S sequences were generated (accession numbers in Table 1; sequence composition in Table 3). The 28S sequences from the Monocotylidae specimens ($n=4$) were 100% identical to that of a *Loimosina* sp. (Table 1; 99% query coverage).

There is no GenBank sequence available for *Erpocotyle*; however, NJ analysis using COI sequences obtained herein supported the finding of 2 distinct species of *Erpocotyle* in the sharks examined (Fig. 3). *Erpocotyle sphyrynae* and *E. microstoma* sequences formed 2 distinct groups each with 100% bootstrap support. Likewise, the 2 available sequences of species of *Loimosina* formed a clade (100% bootstrap support) distinct from all available Monocotylidae sequences from GenBank (Fig. 4). Intraspecific variation for *E. sphyrynae* was 0.00–0.02 and 0.01–0.04 for *E. microstoma*; for *L. wilsoni*, these values were 0.01–0.02 (Tables 4 and 5, respectively).

The maximum likelihood phylogeny using 28S sequences of Monocotylidae supports the proximity of *L. wilsoni* and *Loimosina* sp. (of Boeger et al., 2014), which comprise a monophyletic clade to the paraphyletic *Neoheterocotyle* and its sister *Troglocephalus* (Fig. 5). *Cathariotrema* (another monocotylid from hammerheads) is much further distanced in the tree, forming a sister clade to *Triloculotrema*.

Taxonomy

Subclass Heteronchoinea Boeger and Kritsky, 2001

Hexabothriidae Price, 1942

Erpocotyle microstoma (Brooks, 1934)

Figure 6A–H

Redescription (based on 9 whole- and 9 partially mounted specimens, including hologenophores, 2 syntypes; measurements based on specimens collected in present study): Body 3623–7425 (5562; $n=7$) long, 800–1575 (1172; $n=10$) at greatest width, tapering anteriorly (Fig. 5A). Oral sucker 160–283 (224; $n=9$) long, 200–375 (278; $n=10$) wide, multiple papillae present within mouth. Pharynx bulbous, 73–150 (98; $n=7$) long, 63–150 (88; $n=7$) wide. Oesophagus short. Caeca double, diverticulated, fusing just anterior to haptor, extends as non-diverticulated branches into haptor and haptoral appendix. Haptor symmetrical, squared-oval shaped 950–1550 (1257; $n=7$) long, 800–1075 (938; $n=6$) wide, armed with 2 parallel symmetrical rows of bell-shaped suckers, and haptoral appendix; haptoral suckers in 3 pairs; pairs 1–3 measuring 380–550 (464; $n=7$), 425–550 (493; $n=6$) and 375–500 (451; $n=6$) in diameters, respectively; hook-shaped sclerites embedded within haptoral suckers (see Fig. 6E–H), composed by long slightly curved shaft, short recurved hook, measurements are included in Table 6; haptoral appendix 1013–1875 (1515; $n=8$) long, 255–500 (407; $n=10$) wide, bearing 2 distal bell-shaped suckers composed of a small proximal and distal bulb and pair of anchors (Fig. 5D): proximal bulb 36–65 (51; $n=8$) long, 65–85 (73; $n=8$) wide; distal bulb 95–225 (164; $n=8$) long, 125–185 (151; $n=8$) wide; anchors 50–75 (62; $n=9$) in total length, 25–40 (31; $n=3$) wide. Testes numerous (40–55; $n=3$), irregular-shaped, located in posterior third of the body; vas deferens winding from testes to base of male copulatory organ (MCO). Germarium (ovary) J-shaped (Fig. 6C), 554–670 (612; $n=2$) long, 268–430 (349; $n=2$) wide, pre-testicular, proximally poorly lobate, descending branch straight, ascending

branch straight. Ootype smooth (see Fig. 6C); seminal receptacle present, 225–340 (278; $n=3$) long, 105–125 (117; $n=3$) wide; uterus ventral to vas deferens, containing few eggs; eggs 138–190 (168; $n=4$) long, 40–65 (53; $n=4$) wide, with polar filaments 70–120 (98; $n=3$). Common genital pore posterior to caeca bifurcation. MCO unarmed (Fig. 6B), distally bulbous, ovate, 18–63 (35; $n=5$) in diameter; prostatic region 33–120 (77; $n=5$) long, 13–65 (36; $n=5$) wide; distal portion of vas deferens with thick walls. Two latero-ventral vaginal apertures; vagina parallel, proximally connected to vitelline commissure; distal vaginal duct expanded with glands along entire length.

Taxonomic summary:

Type host and locality: *Sphyryna zygaena* Beaufort, NC, USA (western North Atlantic Ocean)

Present hosts and localities: *Sphyryna lewini*, *S. gilberti*, hybrid (*S. lewini*/*S. gilberti*) Bulls Bay, Awendaw, SC, USA (western North Atlantic Ocean)

Other reported localities: *Sphyryna mokarran* Panama Canal, Panama (Pacific Ocean); *Sphyryna tudes* Punta del Este, Uruguay (western South Atlantic Ocean)

Site of infection: gill filaments

Specimens studied: syntypes USNM 132155, HWML 1437; vouchers: USNM 1666647, USNM 1666648, USNM 1666649, USNM 1666651, USNM 1666652, USNM 1666653, USNM 1666654, HWML 216856, HWML 216857, HWML 216858, HWML 216859, HWML 216860, HWML 216861, HWML 216863, HWML 216864; hologenophores: USNM 1666642, USNM 1666643, USNM 1666644, USNM 1666645, USNM 1666646, USNM 1666650, HWML 216851, HWML 216852, HWML 216853, HWML 216854, HWML 216855, HWML 216861, HWML 216862, HWML 216863, HWML 216864, HWML 216865, HWML 216866

Representative sequences:	COI	–	Genbank
OP342755, OP342756, OP342757, OP342760, OP342761, OP342765, OP342766, OP342770, OP342771	OP342757, OP342761, OP342766, OP342767		OP342758, OP342759, OP342763, OP342764, OP342768, OP342769

Remarks: Brooks (1934) indicated the absence of the MCO in his original description of this species. It thus far has been either considered missed or assumed absent in all other redescriptions (Price, 1942; Caballero et al., 1956; Suriano and Labriola, 1998). However, the MCO is clearly present in available syntypes and specimens collected in the present study, and it corresponds to the morphology of MCO of *Erpocotyle* species as diagnosed by Boeger and Kritsky (1989), except for the relatively expanded prostatic region, which can represent either an artefact or an autapomorphic feature of the species. The vaginae, which were never previously described in detail (see Price, 1942; Caballero et al., 1956; Suriano and Labriola, 1998), also conform with the general morphology for species of the genus as diagnosed by Boeger and Kritsky (1989) – the vaginae are parallel, non-muscular, differentiated into 2 segments and distally expanded with basal glands. The generic diagnosis for *Erpocotyle* spp. includes glando-muscular terminal vaginae, a homologous character. However, compared to those from USNM, our specimens displayed a more dilated distal vagina in *E. microstoma* than in *E. sphyrynae*. This feature, along with the shape and size of the sucker sclerites, may potentially be used to distinguish *E. microstoma* from the remaining species of the genus.

Available sequences for hexabothriids are limited (only 9 COI, 14 Cytb, 13 28S and 7 18S). The 5 COI sequences of *E. microstoma* used in this analysis form a group in the NJ tree, with high bootstrap support (Fig. 3; Table 5). Conspecific distances among sequences of *E. microstoma* did not exceed 0.04 while interspecific distances varied between 0.29 and 0.40. Although sequences generated were short (~340 bp), such short sequences

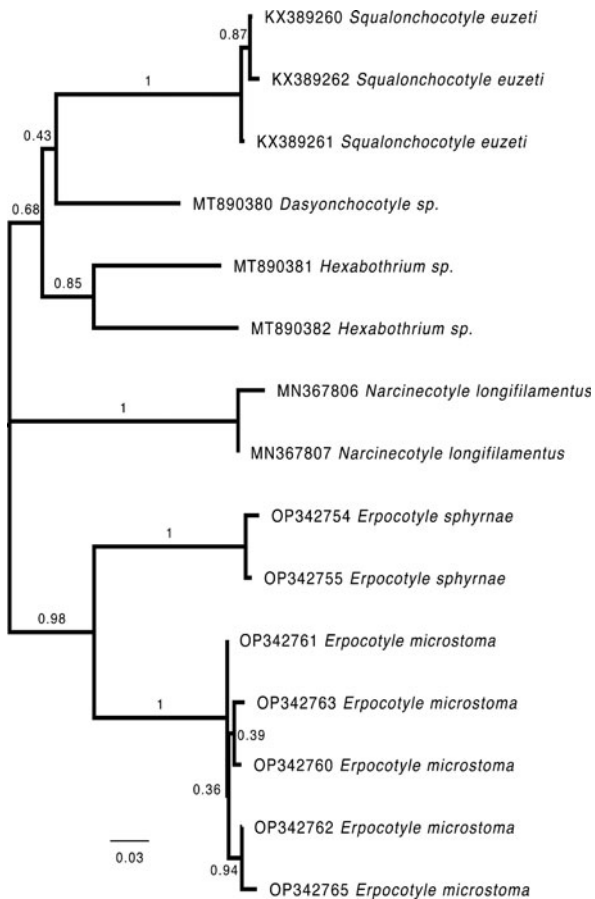


Fig. 3. Neighbour-joining tree based on Tamura and Nei (1993) distances for Hexabothriidae COI sequences from this study and from the GenBank database conducted in MEGA X (Kumar *et al.*, 2018). Sum of branch length = 1.11 and bootstrap support is shown, for 15 nucleotide sequences representing 6 species with a total of 821 positions in the final dataset. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances (number of base substitutions per site, see Table 4). Codon positions included were 1st + 2nd + 3rd.

have been demonstrated to still be useful in species delimitation of other organisms (Hajibabaei *et al.*, 2006). The distances (0.23–0.25; Table 4) between sequences of the 2 morphologically distinguishable species observed in this study support the delimitation between *E. microstoma* and *E. sphyrnae*.

Erpocotyle sphyrnae (MacCallum, 1931)

Figure 7A–H

Redescription (based on 7 whole- and 5 partially mounted specimens, including hologenophores, 5 syntypes; measurements are from specimens collected in the present study): Body fusiform, 1425–4550 (2719; $n = 7$) long, 263–890 (521; $n = 9$) wide at posterior third of body (see Fig. 7A). Oral sucker 140–445 (258; $n = 9$) long, 190–550 (350; $n = 8$) wide, multiple papillae present within the mouth. Pharynx bulbous, 48–80 (62; $n = 6$) in diameter. Oesophagus short. Caeca double, diverticulated, fusing just anterior to haptor, extend as non-diverticulated branches into haptor and haptoral appendix. Vitellaria extending posteriolaterally, branching into haptor and haptoral appendix. Haptor symmetrical, in the shape of a squared-oval 480–970 (704; $n = 8$) long and 375–562 (478; $n = 4$) wide, armed with 2 parallel symmetrical rows of bell-shaped suckers. Haptoral appendix projecting marginally between first sucker–sclerite pair; hook-shaped sclerites embedded within haptoral suckers, comprises long shaft and long recurved hook (see Fig. 7E–H). Haptoral suckers 1–3 measure 180–325 (252; $n = 7$), 180–330 (258; $n = 7$) and 180–325 (250; $n = 7$) in diameter measurements of sucker sclerites in Table 7. Haptoral appendix 410–1030 (740; $n = 8$) long, 140–320 (222; $n = 8$) wide with 2

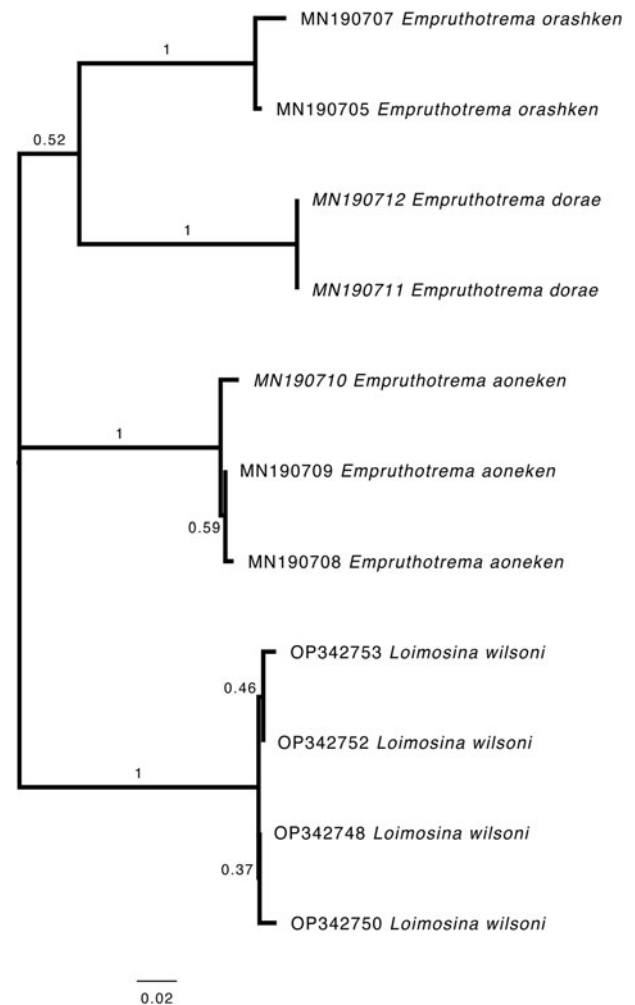


Fig. 4. Neighbour-joining tree based on Tamura and Nei (1993) distances for Monocotylidae COI sequences from this study and from GenBank conducted in MEGA X (Kumar *et al.*, 2018). Sum of branch length = 0.532 and bootstrap support is shown, for 14 nucleotide sequences representing 4 species with a total of 348 positions in the final dataset. The tree is drawn to scale, with branch lengths in the same units as those of the evolutionary distances (number of base substitutions per site, see Table 5). Codon positions included were 1st + 2nd + 3rd.

terminal bell-shaped suckers, each containing a muscular proximal bulb 13–53 (31; $n = 7$) long, 25–63 (46; $n = 7$) wide, and a distal bulb, 48–145 (107; $n = 8$) long, 58–148 (107; $n = 8$) wide and a pair of anchors between suckers (Fig. 7D), 45–58 (52; $n = 10$) long and 13–33 (22; $n = 5$) wide. Genital pore common. Testes multiple (60–98; $n = 3$), post-germarium, each subovate. Vas deferens distally wide with thick walls, winding from anterior testes to base of MCO; MCO composed by a distal unarmed, ovate muscular bulb 25–50 (40; $n = 4$) in diameter and a proximal elongate prostatic region 36–100 (79; $n = 4$) long and 29–45 (37; $n = 4$) wide (Fig. 7B). Two parallel vaginae (see Fig. 7B), located on either side of the MCO, differentiated into 2 segments: distal segment expanded with glands along entire length and a thin muscular layer on the most distal segment. Vaginal pores opening at level of genital pore, with each vaginal duct connecting proximally to the vitelline reservoir/commissure. Ootype smooth (see Fig. 7C); uterus with up to 3 eggs; eggs fusiform, 125–235 (197; $n = 3$) long, 35–50 (42; $n = 3$) wide, with 2 polar filaments 25–95 (53; $n = 3$) long. Germarium J-shaped, 180–648 (357; $n = 5$) long by 95–348 (185; $n = 5$) wide, adjacent to testes to the left of the medial line; proximal germarium lobate, straight descending and ascending germarium branches. Seminal receptacle ovate, 85–188 (139; $n = 4$) long and 40–90 (70; $n = 4$) wide.

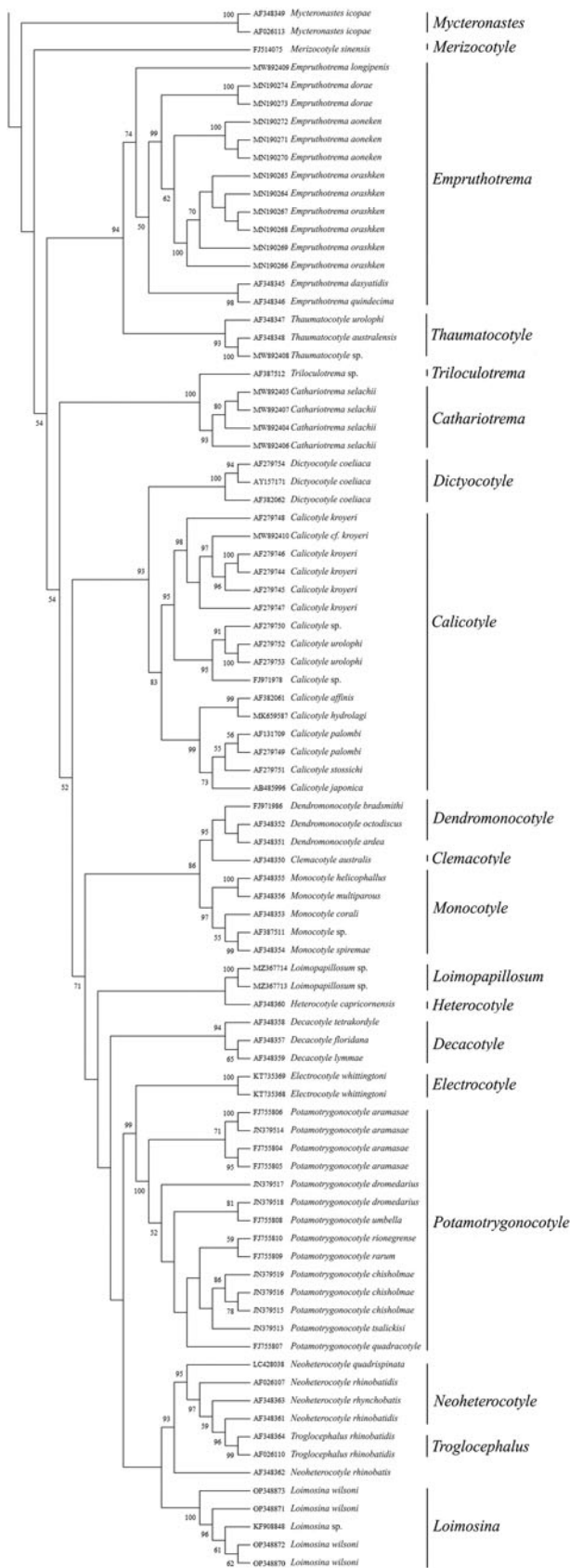


Fig. 5. Maximum likelihood tree (GTR+G substitution model, 1000 bootstraps) for Monocotylidae 28S sequences from this study and GenBank conducted in raxmlGUI 2.0 (version 2.0.6) (Silvestro and Michalak, 2012; Stamatakis, 2014) using 92 sequences representing 54 species with a total of 4022 positions in the final dataset. The consensus tree is shown with bootstraps as support; values lower than 50% have been removed.

Taxonomic summary:

Type host and locality: *Sphyrna zygaena* Woods Hole, MA, USA (western North Atlantic Ocean)

Present host and localities: *Sphyrna lewini*, *S. gilberti*, hybrid (*S. lewini*/*S. gilberti*) Bulls Bay, Awendaw, SC, USA (western North Atlantic Ocean)

Other hosts and localities: *Sphyrna zygaena*, *S. lewini* Dakar, Senegal (eastern North Atlantic Ocean); *S. lewini* Hawaii, USA (Pacific Ocean); *S. mokarran* Nueweiba, Egypt (Gulf of Aqaba)

Site of infection: gill filaments

Specimens studied: syntypes USNM 1320884, USNM 1320885; vouchers: USNM 1666656, USNM 1666657, USNM 1666659, USNM 1666660, USNM 1666661, HWML 216867, HWML 216868, HWML 216869; hologenophores: USNM 1666655, USNM 1666658, HWML 216870, HWML 216871

Representative sequences: COI_Genbank OP342754, OP342755

Remarks: Yamaguti (1968) presented the latest redescription of this species, which provides its morphology in detail, but indicated the presence of 6 irregular muscular pits, 2 median and 4 submedian, within the anterior sucker. These structures were not present in any of the specimens examined herein nor were they noted in any of the previous descriptions/redescriptions (MacCallum, 1931; Price, 1942; Euzet and Maillard, 1967).

Compared to the syntype specimens, our specimens depicted a protruding bulge on one of the proximal ends of the anchors (Fig. 7D). However, this feature was not visible in the museum specimens likely because their anchors were not properly flattened. Anchors have been shown to have high morphological variation among hexabothriids (Teo et al., 2013; Khang et al., 2016); thus, the presence of this bulge may be a possible differential diagnostic between *E. sphyrnae* and *E. microstoma*.

The 2 COI sequences generated for *E. sphyrnae* formed a group in the NJ tree (Fig. 3). This group is adjacent to the group composed by *E. microstoma* sequences, presenting a distance of 0.23–0.25 among these species, compared to distances between 0.35 and 0.41 with other, non-*Erpocotyle* Hexabothriidae sequences (Table 4).

Subclass Polyonchoinea Bychowsky, 1937

Monocotylidae Taschenberg, 1879

Loimosina wilsoni (Manter, 1944)

Figure 8A–F

Redescription (based on 9 whole- and 2 partially mounted specimens, including hologenophores and 3 syntypes; measurements are from specimens collected in present study): |Body 1415–3226 (2089; *n* = 4) long by 229–506 (371; *n* = 5) wide at mid-length (see Fig. 8A). Head tapering anteriorly bearing 3 pairs of symmetrical pits. Mouth subterminal, ventral with prepharynx; large bulbous pharynx, 104–221 (166; *n* = 8) long, 103–255 (155; *n* = 8) wide, large excretory vesicle on each side (see Fig. 8D); oesophagus inconspicuous; caeca lacking diverticulae, extending posterolaterally, terminating blind just anterior to haptor. Haptor subcircular, 184–393 (281; *n* = 8) long, 378–673 (504; *n* = 8) wide, with an external row of small and often inconspicuous loculi, a single latero-posterior pair of anchors (Fig. 8F), 7 pairs of smaller hooks organized evenly and symmetrically around the rim of the haptor (Fig. 8E). Anchors 27–60 (44; *n* = 10) long, 9–16 (12; *n* = 5) wide, evenly curved point and shaft, truncated superficial root, elongate deep root 16–37 (28; *n* = 7). Hooks similar in shape and size, 8 (*n* = 2) long, with short, recurved point, slightly curved shaft, erected truncate thumb, shank about as long as shaft; filamentous hook loop reaching ½ of shank. Common reproductive pore overlaps pharynx, common genital atrium expanded. Single, large, heavily lobed testis, 55–105 (85; *n* = 3) long, 21–28 (24; *n* = 3) wide, post-germarium, occupying middle third of body; vas deferens intercaecal, convoluted, several distal loops expanding into seminal vesicle; ejaculatory bulb pear-shaped with short, cone-shape sclerotized MCO (Fig. 8B). Germarium strongly lobate (especially in larger organisms) (see

Table 2. Summary of prevalence of infection and coinfection (%) per monogenoid species (*L. wilsoni*, *E. microstoma* and *E. sphyrnae*) and per monogenoid family (Hexabothriidae is represented by the combined prevalence of both *Erpocotyle* species; *L. wilsoni* was the only monocotylid found, thus its prevalence is representative of the family in the present study) successfully identified shark individuals ($n = 79$)

	Number of specimens per species	<i>L. wilsoni</i>	<i>E. microstoma</i>	<i>E. sphyrnae</i>	Coinfections	Hexabothriid	Monogenoid
<i>S. gilberti</i>	44	0	27	14	2	45	50
<i>S. lewini</i>	20	60	30	15	30	40	75
F1 hybrid	6	33	0	17	0	17	50
<i>S. lewini</i> BX	5	40	20	20	20	40	80
<i>S. gilberti</i> BX	4	0	25	0	0	50	50
Total prevalence		22	25	14	10	42	58

BX, backcross.

The shaded boxes represent the numbers of each species of host sampled.

Fig. 8D); uterus expanded; vaginal opening ventro-lateral; vagina sinuous duct distally expanded, glandular. Vitellaria follicular, coextensive with caeca; vitelline commissure overlapping with branch of germarium; ootype, seminal receptacle not observed. Egg ovate, 67–108 (94; $n = 3$) long, 46–87 (71; $n = 3$) wide, with a short proximal filament (Fig. 8C).

Taxonomic summary:

Type host and locality: *Sphyrna zygaena* Montego Bay, Jamaica (Caribbean Sea)

Present hosts and localities: *Sphyrna lewini* and hybrids (*S. lewini*/*S. gilberti*) Bulls Bay, Awendaw, SC, USA (western North Atlantic Ocean)

Other hosts and localities: *Sphyrna lewini* Dakar, Senegal (eastern North Atlantic Ocean)

Site of infection: gill filaments

Specimens studied: syntypes USNM 1337561, USNM 1337564, HWML 1425; vouchers: USNM 1666662, USNM 1666663, USNM 1666664, USNM 1666665, USNM 1666666, USNM 1666668, USNM 1666669, USNM 1666670, HWML 216873, HWML 216874, HWML 216875, HWML 216876, HWML 216878, HWML 216879, HWML 216880, HWML 216881; hologenophores: USNM 1666667, HWML 216872, HWML 216877

Representative sequences: COI-Genbank OP342748, OP342749, OP342750, OP342751, OP342752, OP342753; 28S-Genbank OP348870, OP348871, OP348872, OP348873

Remarks: Bravo-Hollis (1970) noted that the main differences between *L. parawilsoni* and *L. wilsoni* were geographical location (*L. parawilsoni* described from *S. lewini* in the Pacific Ocean), and few morphological features. *Loimosina parawilsoni* depicts a sub-circular haptor with hooks embedded in the papillae, a 'different [sic]' shape of the anchors, the presence of a common genital pore and a MCO with sclerotized, blade-like lateral walls. Manter (1944) interpreted the MCO of *L. wilsoni* as 'rudimentary,

consisting of a very short, very thinly chitinized tube near the male pore [sic]', uncertain of its position within or external to the ejaculatory bulb. Euzet and Maillard (1967) did not observe an MCO, instead considered this as the sclerotized distal wall of the ejaculatory bulb, protruding slightly ventrally. Syntype specimens from Manter (1944) were not mounted flat; structures were difficult to study and no MCO was visible in the syntypes. Unfortunately, there is no specimen from Euzet and Maillard (1967) in museum collections. Our specimens presented a short conical, sclerotized MCO at the distal extremity of the ejaculatory bulb (Fig. 8B) – thus, the presence of a sclerotized MCO projecting from the ejaculatory bulb is not a distinguishable diagnostic between the 2 species of *Loimosina*. Likewise, Bravo-Hollis (1970) mentions a difference in the shape of the anchors, but a difference between those of the syntype of *L. parawilsoni* (CNHE 154) and our specimens was not evident. Molecular sequences from specimens of *Loimosina* collected from *S. lewini* from the Pacific Ocean would allow clarification of this classification. However, as morphological descriptions stand, specimens of both *Loimosina* species are greatly similar which leaves the only differential criterion to be the shape of the haptor. However, the shape of non-sclerotized structures is often affected by fixation and mounting. Hence, given that both species are described from the same host, we strongly suggest that *L. parawilsoni* is a junior synonym of *L. wilsoni*. However, a definitive decision should await adequate restudy of specimens of *L. parawilsoni* and, preferably, a molecular delimitation of the species.

The same nucleotide composition of 28S from species of *Loimosina* redescribed in this study and that of Boeger *et al.* (2014) also strongly supports that *L. wilsoni* is present in at least 1 species of *Sphyrna* in the South Atlantic Ocean (see discussion).

Table 3. Sequence compositions, with no gaps, per monogenoid species collected from gills of *Sphyrna* spp.: length in base pairs (bp), reported as a range; variable sites (V); parsimony-informative sites (PI); per cent composition of each nucleotide

	Length (bp)	V	PI	A%	T%	C%	G%
COI							
<i>L. wilsoni</i>	347–370	9	1	28.5	44.2	7.5	19.9
<i>E. microstoma</i>	332–347	71	63	28.8	35.2	14.8	21.3
<i>E. sphyrnae</i>	332–364			27.7	35.0	14.6	22.7
28s							
<i>L. wilsoni</i>	642–950	0	0	28.4	24.9	28.3	18.5

Variable and PI sites are compared between both species of *Erpocotyle*. *Loimosina wilsoni* was the only monocotylid identified in the present study, and was not compared to the other species of Hexabothriidae.

Table 4. Tamura and Nei (1993) distances between Hexabothriidae COI sequences from this study (shaded) and from GenBank based on an 821 base pair alignment

	OP342754 <i>Erpocotyle sphyrrae</i>	OP342755 <i>Erpocotyle sphyrrae</i>	OP342763 <i>Erpocotyle microstoma</i>	OP342762 <i>Erpocotyle microstoma</i>	OP342760 <i>Erpocotyle microstoma</i>	OP342759 <i>Erpocotyle microstoma</i>	OP342761 <i>Erpocotyle microstoma</i>	KX389260 <i>Squalonchocotyle euzeti</i>	KX389261 <i>Squalonchocotyle euzeti</i>	KX389262 <i>Squalonchocotyle euzeti</i>	MN367806 <i>Narcinecotyle longifilamentus</i>	MN367807 <i>Narcinecotyle longifilamentus</i>	MT890380 <i>Dasyonchocotyle sp.</i>	MT890381 <i>Hexabothrium sp.</i>	MT890382 <i>Hexabothrium sp.</i>
OP342754 <i>Erpocotyle sphyrrae</i>		0.01	0.22	0.22	0.22	0.22	0.22	0.32	0.31	0.32	0.30	0.30	0.34	0.30	0.29
OP342755 <i>Erpocotyle sphyrrae</i>	0.02		0.22	0.22	0.22	0.21	0.22	0.32	0.32	0.32	0.30	0.30	0.33	0.30	0.31
OP342763 <i>Erpocotyle microstoma</i>	0.25	0.24		0.01	0.01	0.01	0.01	0.33	0.32	0.33	0.33	0.32	0.25	0.30	0.32
OP342762 <i>Erpocotyle microstoma</i>	0.25	0.25	0.03		0.01	0.01	0.01	0.31	0.29	0.31	0.30	0.32	0.25	0.29	0.31
OP342760 <i>Erpocotyle microstoma</i>	0.25	0.25	0.01	0.02		0.01	0.01	0.33	0.32	0.33	0.33	0.32	0.25	0.29	0.32
OP342759 <i>Erpocotyle microstoma</i>	0.25	0.24	0.04	0.01	0.03		0.01	0.35	0.35	0.36	0.33	0.32	0.25	0.36	0.40
OP342761 <i>Erpocotyle microstoma</i>	0.24	0.23	0.01	0.01	0.01	0.02		0.32	0.32	0.32	0.32	0.32	0.25	0.29	0.31
KX389260 <i>Squalonchocotyle euzeti</i>	0.39	0.40	0.38	0.36	0.38	0.43	0.37		0.01	0.00	0.34	0.25	0.22	0.25	0.25
KX389261 <i>Squalonchocotyle euzeti</i>	0.38	0.39	0.36	0.35	0.37	0.42	0.36	0.01		0.01	0.34	0.25	0.21	0.25	0.25
KX389262 <i>Squalonchocotyle euzeti</i>	0.39	0.39	0.39	0.37	0.39	0.44	0.38	0.01	0.01		0.34	0.25	0.22	0.25	0.25
MN367806 <i>Narcinecotyle longifilamentus</i>	0.37	0.36	0.38	0.36	0.38	0.39	0.35	0.41	0.40	0.41		0.00	0.30	0.36	0.31
MN367807 <i>Narcinecotyle longifilamentus</i>	0.36	0.35	0.36	0.37	0.37	0.37	0.35	0.30	0.30	0.31	0.00		0.30	0.31	0.26
MT890380 <i>Dasyonchocotyle sp.</i>	0.41	0.39	0.29	0.30	0.31	0.31	0.31	0.26	0.24	0.26	0.30	0.30		0.25	0.24
MT890381 <i>Hexabothrium sp.</i>	0.35	0.36	0.32	0.32	0.32	0.40	0.32	0.30	0.31	0.31	0.43	0.35	0.29		0.17
MT890382 <i>Hexabothrium sp.</i>	0.37	0.37	0.35	0.36	0.36	0.47	0.35	0.28	0.29	0.29	0.39	0.31	0.26	0.22	

The number of base substitutions per site between sequences was calculated in MEGA X (Kumar et al., 2018) using the maximum composite likelihood method (Tamura et al., 2004). Standard error estimates are above the diagonal. This analysis involved 6 species with a total of 15 sequences. Codon positions included were 1st + 2nd + 3rd.

Table 5. Tamura and Nei (1993) distances between Monocotylidae COI sequences from this study (shaded) and from the GenBank database based on a 348 base pair alignment

	OP342753 <i>Loimosina wilsoni</i>	OP342748 <i>Loimosina wilsoni</i>	OP342752 <i>Loimosina wilsoni</i>	OP342750 <i>Loimosina wilsoni</i>	MN190712 <i>Empruthotrema doraе</i>	MN190711 <i>Empruthotrema doraе</i>	MN190710 <i>Empruthotrema aoneken</i>	MN190709 <i>Empruthotrema aoneken</i>	MN190708 <i>Empruthotrema aoneken</i>	MN190707 <i>Empruthotrema orashken</i>	MN190706 <i>Empruthotrema orashken</i>	MN190705 <i>Empruthotrema orashken</i>	MN190704 <i>Empruthotrema orashken</i>	MN190703 <i>Empruthotrema orashken</i>
OP342753 <i>Loimosina wilsoni</i>		0.00	0.00	0.01	0.29	0.29	0.21	0.21	0.21	0.25	0.25	0.23	0.24	0.24
OP342748 <i>Loimosina wilsoni</i>	0.01		0.00	0.01	0.30	0.30	0.21	0.21	0.21	0.24	0.24	0.22	0.24	0.24
OP342752 <i>Loimosina wilsoni</i>	0.01	0.01		0.01	0.29	0.29	0.21	0.20	0.20	0.24	0.24	0.22	0.24	0.24
OP342750 <i>Loimosina wilsoni</i>	0.01	0.01	0.01		0.30	0.30	0.21	0.21	0.21	0.24	0.24	0.22	0.24	0.24
MN190712 <i>Empruthotrema doraе</i>	0.32	0.32	0.32	0.32		0.00	0.19	0.19	0.19	0.19	0.19	0.20	0.19	0.19
MN190711 <i>Empruthotrema doraе</i>	0.32	0.32	0.32	0.32	0.00		0.19	0.19	0.19	0.19	0.19	0.20	0.19	0.19
MN190710 <i>Empruthotrema aoneken</i>	0.24	0.24	0.24	0.25	0.21	0.21		0.01	0.01	0.21	0.21	0.20	0.21	0.21
MN190709 <i>Empruthotrema aoneken</i>	0.23	0.23	0.22	0.24	0.20	0.20	0.01		0.00	0.21	0.21	0.20	0.21	0.21
MN190708 <i>Empruthotrema aoneken</i>	0.24	0.24	0.23	0.25	0.20	0.20	0.01	0.00		0.21	0.21	0.20	0.21	0.21
MN190707 <i>Empruthotrema orashken</i>	0.28	0.26	0.27	0.27	0.22	0.22	0.26	0.26	0.26		0.00	0.01	0.01	0.01
MN190706 <i>Empruthotrema orashken</i>	0.28	0.26	0.27	0.27	0.22	0.22	0.26	0.26	0.26	0.00		0.01	0.01	0.01
MN190705 <i>Empruthotrema orashken</i>	0.24	0.22	0.22	0.23	0.21	0.21	0.25	0.24	0.25	0.03	0.03		0.01	0.01
MN190704 <i>Empruthotrema orashken</i>	0.26	0.25	0.25	0.26	0.22	0.22	0.26	0.25	0.26	0.02	0.02	0.01		0.00
MN190703 <i>Empruthotrema orashken</i>	0.26	0.25	0.25	0.26	0.22	0.22	0.26	0.25	0.26	0.03	0.02	0.02	0.00	

The number of base substitutions per site between sequences is shown and was calculated in MEGA X (Kumar *et al.*, 2018) using the maximum composite likelihood method (Tamura *et al.*, 2004). Standard error estimates are above the diagonal. This analysis involved 4 species with a total of 14 sequences. Codon positions included were 1st + 2nd + 3rd.

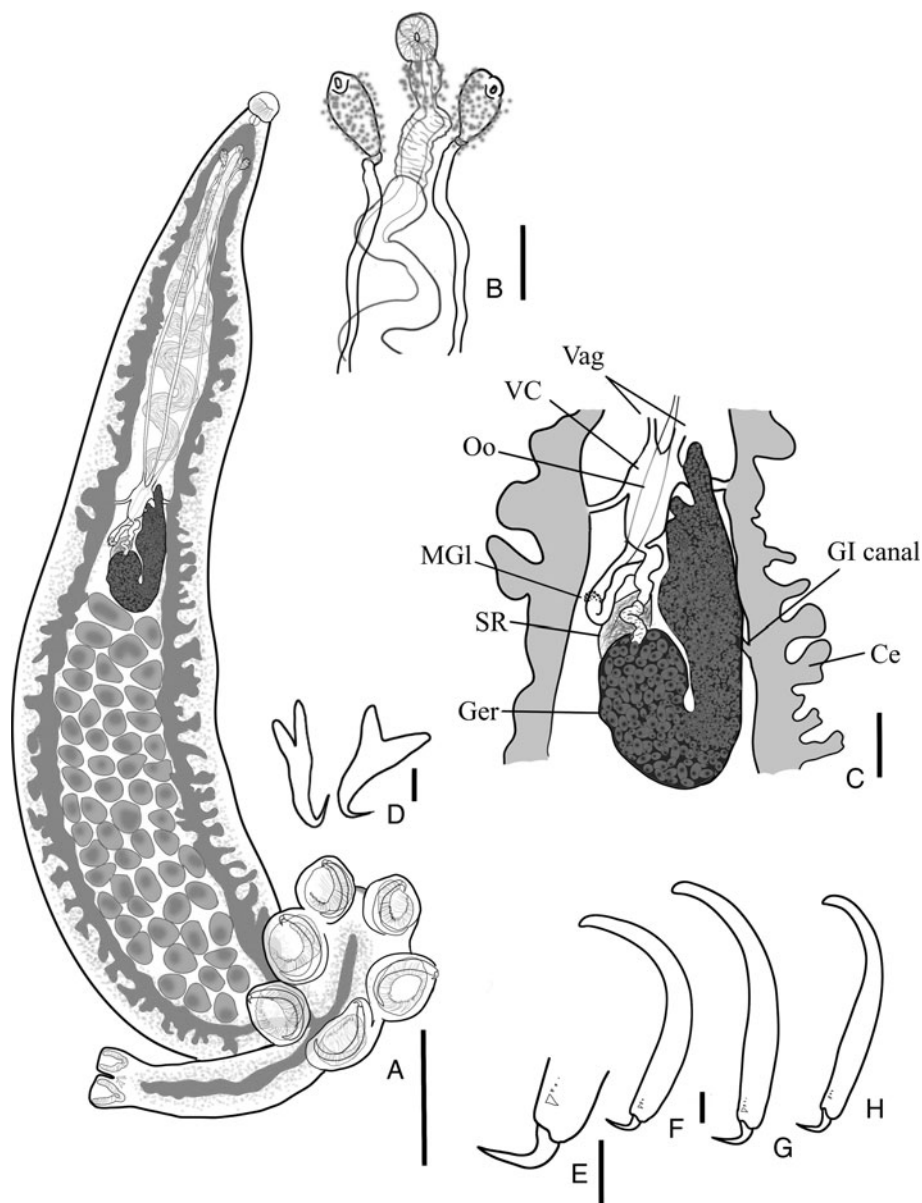


Fig. 6. *Erpocotyle microstoma* (A) whole body, scale bar 1000 μm; (B) male copulatory organ (MCO) and parallel vaginae, scale bar 50 μm; (C) midsection of specimen containing the Ce (caeca), GI (gastrointestinal) canal, reproductive organs Ger (germarium), MGI (Mehlis' gland), Oo (ootype), SR (seminal receptacle) and Vag (vaginae), and VC (vitelline canal), scale bar 100 μm; (D) anchors of the haptor appendix, scale bar 60 μm; (E) curvature of the terminal hook of a sucker sclerite, scale bar 60 μm; (F–H) sucker sclerites 1–3, respectively, scale bar 60 μm.

Table 6. *Erpocotyle microstoma* sclerite measurements in micrometres

Sclerite	A	B	C	D	E
1	230–710 (482; n = 11)	60–91 (71; n = 7)	518–1006 (802; n = 7)	345–720 (538; n = 10)	50–110 (75; n = 11)
2	260–650 (477; n = 11)	66–91 (81; n = 7)	554–980 (824; n = 7)	410–680 (547; n = 11)	60–120 (81; n = 11)
3	240–610 (439; n = 9)	65–80 (75; n = 6)	491–803 (693; n = 6)	370–630 (488; n = 9)	50–95 (72; n = 10)

Sclerites are numbered 1 through 3, with 1 being closest to the point of attachment of the haptor appendix. Ranges are given, with averages in parentheses and number of specimens measured. See Fig. 1 for images of measurements: (A) tip-to-tip (MacCallum, 1931), (B) perimeter hook length (Euzet and Maillard, 1967), (C) perimeter shaft length (Euzet and Maillard, 1967), (D) shaft length (Bullard and Dippenaar, 2003) and (E) max shaft width (Bullard and Dippenaar, 2003).

Available partial sequences of the monocotylids *Empruthotrema orashken* Irigoitia et al., 2019, *Empruthotrema dorae* Irigoitia et al., 2019 and *Empruthotrema aoneken* Irigoitia et al., 2019 of the COI mitochondrial DNA gene grouped with the sequences produced herein. Sequences of *L. wilsoni* cluster together with significant bootstrap support and present short intra-specific distances (0.01–0.02, see Table 5) and significantly greater interspecific distances (0.22–0.32; Fig. 4).

Discussion

Age-dependent parameters can produce high variability when averaging measurements (or providing ranges), and it would aid

in the validation of measurements and resolve diagnostic issues related to the genera of Hexabothriidae if these variables were identified (see Vaughan and Christison, 2012). For example, the extension of the vitellaria into the haptor appendix is disputed in the literature as to whether it is a reliable character of a species or genera, or if it is age-dependent for individuals (Price, 1942; Euzet and Maillard, 1967). Originally used by Price (1942) as a diagnostic in the genera of Hexabothriidae, the extension of the vitellaria was noted to vary with the maturity of the specimen by Cerfontaine (1899), Sproston (1946) and by Euzet and Maillard (1967) in their respective redescrptions of *E. sphyrnae*. Accordingly, Boeger and Kritsky (1989) did not mention this aspect of the vitellaria distribution in their revision of

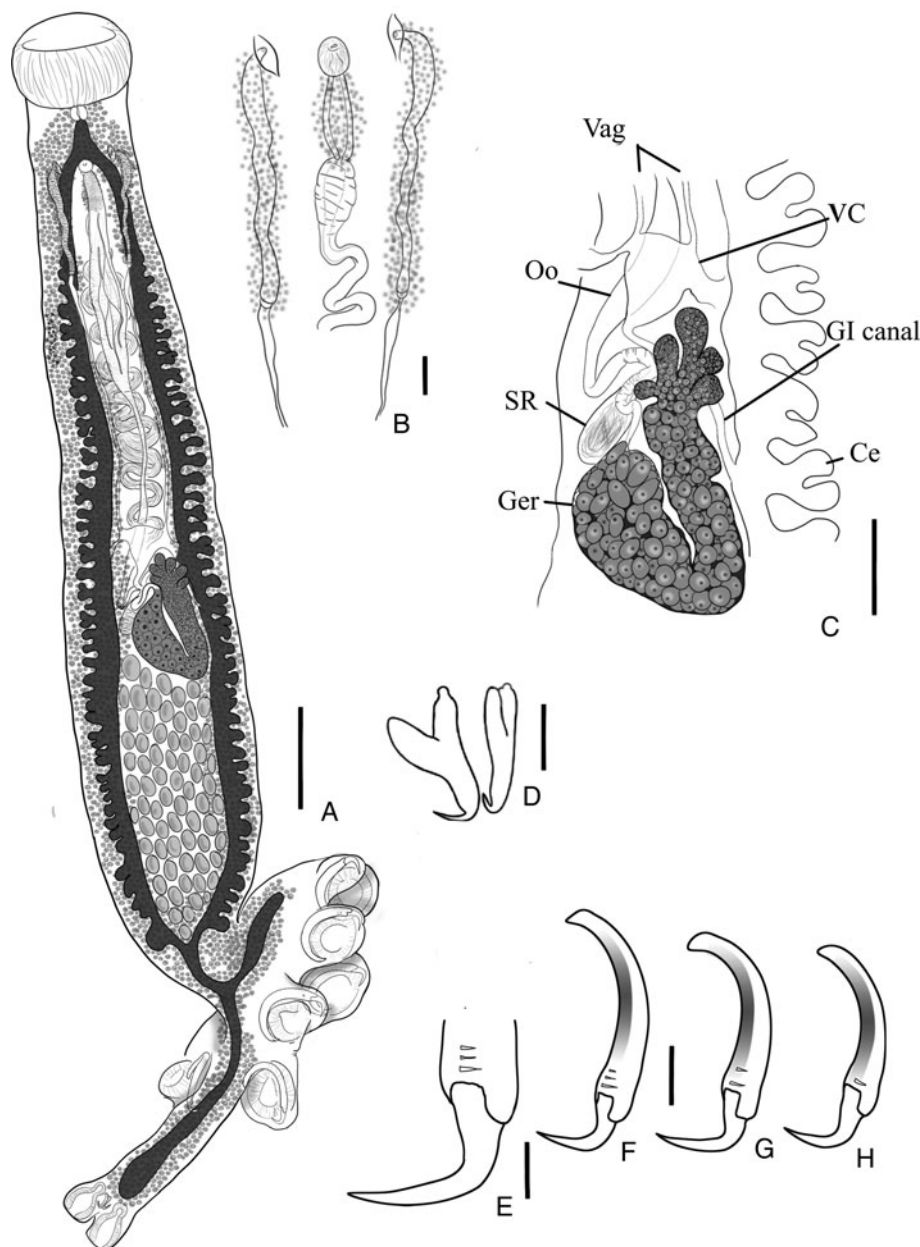


Fig. 7. *Erpocotyle sphyrnae* (A) whole body, scale bar 500 μm ; (B) male copulatory organ (MCO) and parallel vaginae, scale bar 60 μm ; (C) midsection of specimen (abbreviations as in Fig. 6), scale bar 200 μm ; (D) anchors of the haptor, scale bar 25 μm ; (E) curvature of the terminal hook of a sucker sclerite, scale bar 30 μm ; (F–H) sucker sclerites 1–3, respectively, scale bar 60 μm .

Table 7. *Erpocotyle sphyrnae* sclerite measurements in micrometres

Sclerite	A	B	C	D	E
1	82.5–257 (164; $n=10$)	81–166 (122; $n=5$)	386–444 (424; $n=5$)	150–347 (253; $n=9$)	25–50 (38; $n=9$)
2	100–240 (177; $n=10$)	87–172 (128; $n=6$)	373–530 (425; $n=6$)	170–310 (258; $n=10$)	25–50 (41; $n=10$)
3	75–225 (167; $n=10$)	80–141 (124; $n=5$)	287–508 (375; $n=5$)	145–305 (238; $n=10$)	20–475 (80; $n=10$)

Sclerites are numbered 1 through 3, with 1 being closest to the point of attachment of the haptor appendix. Ranges are given with averages in parentheses and number of measured specimens. See Fig. 1 for images of measurements: (A) tip-to-tip (MacCallum, 1931), (B) perimeter hook length (Euzet and Maillard, 1967), (C) perimeter shaft length (Euzet and Maillard, 1967), (D) shaft length (Bullard and Dippenaar, 2003) and (E) max shaft width (Bullard and Dippenaar, 2003).

Hexabothriidae. In our specimens, the vitellaria visibly extended into the haptor and the haptor appendix in all intact mature specimens of both species of *Erpocotyle* ($n=16$), but was not observed to do so in 3 juvenile representatives of *E. sphyrnae*, indicating that this character is age-dependent and not a reliable generic or specific diagnosis.

This study also revealed differences in the host repertoires of monogenoids – indicating different abilities to infect the spectrum of available hosts. Monogenoids typically demonstrate narrow host repertoires being either monoxenous – infecting a single host

species – or stenoxenous – infecting a few related host species (Euzet and Combes, 1980; Rohde, 1994; Whittington *et al.*, 2000). In this study, *L. wilsoni* presented a narrow host repertoire, as it was found only on *S. lewini* and the hybrids and did not infect any of the 44 *S. gilberti* examined. Host repertoire is typically determined by the opportunity of encounter and the compatibility of the involved host/parasite species (Combes, 2001; Araujo *et al.*, 2015; Brooks *et al.*, 2019). Factors involved in the success of the establishment of a parasite may depend on biological and behavioural aspects of the host, such as season of reproduction, stage of

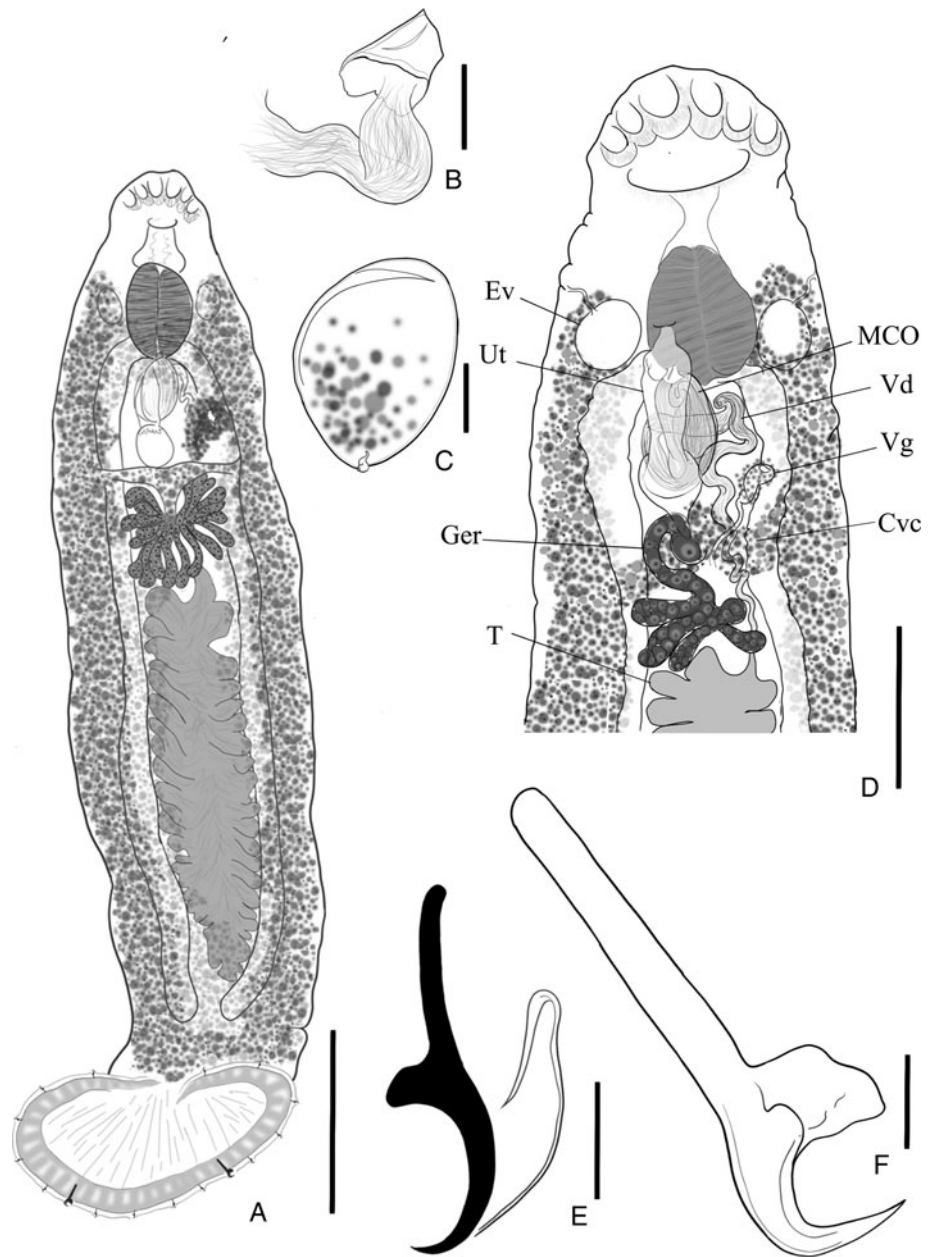


Fig. 8. *Loimosina wilsoni* (A) whole body, scale bar 500 μm ; (B) sclerotized, cone-shaped male copulatory organ (MCO), scale bar 16 μm ; (C) egg, scale bar 25 μm ; (D) mid- to terminal portion of specimen containing the Cvc (common vitelline commissure), Ev (excretory vesicle), Ger (germarium), MCO, T (teste), Ut (uterus), Vd (vas deferens) and Vg (vagina), scale bar 150 μm ; (E) haptoral hook, scale bar 10 μm ; (F) haptoral anchor, scale bar 10 μm .

maturation, physiology, feeding behaviour and immunity, to list a few (MacDonald, 1975; Tinsley and Jackson, 2002; Glennon et al., 2006; Ohashi et al., 2007; Whittington and Kearn, 2011).

Hosts for *L. wilsoni* have been identified as *S. zygaena* and *S. lewini* [syns: *Sphyrna diplana* (Springer, 1941) and *Sphyrna couardi* (Cadenat, 1950)] (Manter, 1944; Hargis, 1955; Euzet and Maillard, 1967). However, there remains some uncertainty in the literature over host identity. Hargis (1955) first suggested that the type host, *S. zygaena*, was possibly misidentified by Manter (1944), a doubt also mentioned by Euzet and Maillard (1967), and that the host could have been *S. diplana* (syn. *S. lewini*). This is supported by the fact that Manter (1944) never observed first-hand the shark specimens as they were collected and provided by Dr C. B. Wilson (for whom *L. wilsoni* was named). Euzet and Maillard (1967) also indicated the morphological similarities of *S. couardi* and *S. lewini* in their own report, which have been later synonymized in McEachran and Séret (1987). Paradoxically, as advances in molecular research have improved species delimitation, cryptic host species are often uncovered, such as the case of *S. gilberti* and *S. lewini*. Thus, it becomes even more intriguing that in this study that *L. wilsoni* remained specific to *S. lewini* and admixed

individuals with at least 50% *S. lewini* DNA (*S. lewini* F1 and BX) and that no *S. gilberti* or *S. gilberti* backcrosses (*S. gilberti* BX) were infected by *L. wilsoni*. The exclusion of *S. gilberti* BX from the host repertoire of *L. wilsoni* remains to be verified, however, given the small number of specimens of *S. gilberti* BX ($n = 4$). The exclusion of species *S. gilberti*, in contrast, from the host repertoire of *L. wilsoni* appears concrete given the large number of specimens of this species ($n = 44$).

Boeger et al. (2014) reported a specimen of *Loimosina* sp. from an unidentified hammerhead individual *Sphyrna* sp. off the coast of southeast Brazil. Although these authors were unable to identify this parasite to the species level based on morphology, its 28S rDNA sequence (GB Accession KF908848) was 98–100% identical to our specimens, suggesting that it was *L. wilsoni*. As *S. gilberti* appears not to be in the host repertoire of *L. wilsoni*, but only in *S. lewini* and hybrids, it is likely that the *Sphyrna* host species from Brazil was *S. lewini*, which is known from those coasts (Compagno, 1984), or yet another species, such as *Sphyrna media* Springer, 1940, *S. mokarran* (Rüppell, 1837), *Sphyrna tiburo* (Linnaeus, 1758), *S. tudes* (Valenciennes, 1822) or *S. zygaena*, all reported from south Brazil (Compagno, 1984). Whereas it is not

known if *L. wilsoni* can infect other *Sphyrna* species, its presence on hybrids of *S. lewini* and *S. gilberti* indicates the possibility of it being stenoxenous and thus potentially using any of these hammerheads as host. Previous studies of the molecular phylogeny of family Sphyrnidae has found the globally distributed *S. lewini* to be more closely related to smaller, range-restricted (eastern Pacific and western Atlantic) sharks *S. media*, *S. tudes* and *S. tiburo* than to other globally distributed large sharks *S. mokarran* and *S. zygaena* (Lim *et al.*, 2010). This is further supported by Pinhal *et al.* (2012), which showed the cryptic Atlantic lineage, now known as *S. gilberti*, to be more closely related to *S. lewini* than the other smaller sharks. This further draws doubt to *S. zygaena* being included in the host repertoire of this parasite.

The hexabothriids *E. microstoma* and *E. sphyrnae* have wider host ranges than *L. wilsoni*. Both have been originally described from specimens of *S. zygaena* from the western North Atlantic along the US coast (Woods Hole, MA and Beaufort, NC, respectively) and included in their repertoire are other hammerheads: *E. sphyrnae* has been reported in *S. zygaena* and *S. lewini* from Dakar, Senegal (Euzet and Maillard, 1967), *S. lewini* in Hawaii (Yamaguti, 1968), *S. mokarran* from Nuweiba, Egypt (Maillard and Paperna, 1978) and now in *S. lewini*, *S. gilberti* and their hybrids from the SC coast of the USA (present study). *Erpocotyle microstoma* has been reported also in *S. mokarran* but from the Panama Canal (Pacific Ocean; Caballero *et al.*, 1956), *S. tudes* from Punta del Este, Uruguay (Suriano and Labriola, 1998) and now in *S. lewini*, *S. gilberti* and their hybrids from the SC coast of the USA (present study). Maillard and Paperna (1978) were the first to recognize the wider repertoires of both these *Erpocotyle* species, and the vast geographic range of *E. sphyrnae*. Here, we further expand the host repertoire and geographic ranges of these 2 species.

Bullard *et al.* (2021) updated the list of monocotyliids in *S. lewini* to include *C. selachii*, which they found infecting the olfactory cavities of *S. lewini* in the northern Gulf of Mexico. The present study did not investigate the olfactory cavities of the specimens of *S. lewini*, *S. gilberti* and hybrids collected from the western North Atlantic. However, *C. selachii*, the only described species and thus the type for the genus, is noted as having a very broad host repertoire; it has been found to infect several *Sphyrna* spp., including *S. lewini*, as well as species of *Carcharhinus* Blainville, 1816, *Rhizoprionodon* Whitley, 1929 (both Carcharhinidae) and *Alopias* Rafinesque, 1810 (Alopiidae), in the Gulf of Mexico and in the western North Atlantic. Thus, it is entirely possible that *C. selachii* infects *S. lewini* and/or *S. gilberti* in the western North Atlantic.

In conclusion, this is the first study to compare monogenoid fauna between cryptic *Sphyrna* species and their hybrids in the western North Atlantic Ocean, and the first report of monogenoids collected from the gills of individuals of *S. gilberti* and hybrids with *S. lewini*. Genetic sequencing using nuclear ribosomal ITS2 sequences showed that species *S. lewini* and *S. gilberti* diverged around 4.5 million years ago (Pinhal *et al.*, 2012) yet can be found sympatrically and can produce reproductive and viable hybrids. This study demonstrated that these 2 hammerheads have differences in monogenoid fauna of the gills, which in addition to the number of vertebrae, further supports a distinction between species.

Lastly, Loimoidae or Loimoinae Price, 1936 – a subfamily of Monocotyliidae – was previously composed of species of 3 genera: *Loimos* McCallum, 1917, *Loimosina* and *Loimopapillosum* Hargis, 1955. The more comprehensive morphological analysis of the specimens provided in this study and the recognition that the sequence of the 28S rDNA fragment of *L. wilsoni* presented herein is identical to that of *Loimosina* sp. published by Boeger *et al.* (2014). This corroborates the decision that the species of these

genera are allocated within the Monocotyliidae and do not compose a monophyletic assemblage within the family, following the suggestion of Boeger *et al.* (2014) and Chero *et al.* (2021). Although there is no sequence currently available for the species of *Loimos*, molecular phylogenetic analyses using sequences of *Loimosina* from the present study and from Boeger *et al.* (2014), and *Loimopapillosum* (see Chero *et al.*, 2021), strongly indicate that these are members of distinct clades within the Monocotyliidae. The classification of *Loimos* and a more robust decision on the validity of Loimoidae/Loimoinae, however, should await sequencing of corresponding DNA fragments and phylogenetic analysis. Likewise, sequences of *Loimosina* specimens from the Pacific Ocean (in combination with morphological examination) would clarify whether *L. parawilsoni* is in fact a synonym for *L. wilsoni*, as suggested in the present study.

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Data availability. All sequences and representative specimens are available from respective repositories, as stated in the manuscript.

Author's contribution. I. d. B. conceived the idea of the initial study, which was carried out by K. M. D.; sharks were captured and provided by B. S. F. and A. S. G.; all molecular sequencing was done by A. B. and K. M. H.-S.; D. S. P. provided assistance in data manipulation and in writing; W. A. B. assisted in taxonomy, data analysis and in writing.

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Conflict of interest. None.

Ethical standards. All sharks were collected and sampled by authorized staff under official permits or scientific exemptions of US state government agencies (SCDNR Scientific Permit no. 2212).

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