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NEW GENUS AND SPECIES OF APOROCOTYLIDAE (DIGENEA) FROM A BASAL ACTINOPTERYGIAN, THE AMERICAN PADDLEFISH, POLYODON SPATHULA, (ACIPENSERIFORMES: POLYODONTIDAE) FROM THE MISSISSIPPI DELTA

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ABSTRACT: Acipensericola petersoni n. gen., n. sp. (Digenea: Aporocotylidae) infects the heart of the American paddlefish Polyodon spathula (Walbaum, 1792) in the Mississippi Delta. It has robust, spike-like body spines arranged in ventrolateral transverse rows; a bowl-shaped anterior sucker centered on the mouth and having minute spines on the inner anterodorsal surface only; a pharynx; an inverse U-shaped ceca extending to near the posterior body end; intercelcal testes comprising a pre-ovarian testicular column plus a single testis posteriorly; an extensively lobed ovary located medially and immediately posterior to the testicular column; a spherical ootype that is intercelcal and post-ovarian; a Laurer’s canal; and a common genital pore. The new species is the first-named aporocotylid collected from a basal actinopterygian. It resembles the chondrichthyian aporocotylids *Chimaerohemecus trondheimensis*, *Orchispirium heterovitellatum*, and *Hyperandrotrema cetorhini* in having an inverse U-shaped ceca, but it is morphologically most similar to the anguilliform aporocotylid *Paracardicoloides yamagutii* in having that feature plus a comparable anterior sucker, a single testis posteriorly, an intertesticular ovary, and a common genital pore. Sequence data for the complete small subunit ribosomal DNA (18S) do not refute its membership within Aporocotylidae nor its affinity to 1 of those aforementioned aporocotylids: *A. petersoni* was basal to the few teleost aporocotylids analyzed, and *C. trondheimensis* was the only taxon basal to *A. petersoni*. We regard the specimens of *Spirochis* sp. previously reported from the shortnose sturgeon *Acipenser brevisrostrum* Lesueur, 1818 as congeneric with the new species.

Adult blood flukes (Digenea: Schistosomatoida) infect jawed vertebrates (Gnathostomata) and historically have been grouped into 3 families correlating to the broad phylogenetic affiliations of their definitive host groups: Aporocotylidae Odhner, 1912 for those blood flukes that infect non-tetrapod gnathostomes, i.e., fishes (Smith, 1997a, 1997b, 2002; Spirorchidae Stunkard, 1921 for those of turtles (Platt, 2002); and Schistosomatidae Stiles and Hassall, 1898 for those of birds and mammals (Khalil, 2002). We concur with Stunkard (1921) who stated that, “In my opinion, the Aporocotylidae of fishes, the Spirorchidae of turtles, and the Schistosomatidae of birds and mammals constitute a well-defined group with inherent natural relationships.” At present, Aporocotylidae includes only 5 nominal species that infect cartilaginous fishes (Chondrichthy-es) (Bullard et al., 2006), plus >100 species that infect bony fishes (Actinopterygii: Teleostei) (see Smith, 1997a, 1997b). As such, aporocotylids infect definitive hosts allocated to widely separated gnathostome lineages, whereas all adult spirorchids and schistosomatids are reportedly restricted to members of Tetrapoda (Gnathostomata: Sarcopterygii). Although Aporocotylidae is the most diverse blood fluke family, with respect to the number of named species and accepted genera, at least tens of aporocotylid species remain unnamed (S. Bullard, unpubl.), and many potential host lineages seem vastly under-explored for the presence of aporocotylid infections. Fewer than 200 (Smith, 1997b) of the nearly 28,000 valid fish species (Nelson, 2006) are reported as hosts, and most fish orders (46 of 63) reportedly lack infections. This void of information is a barrier to understanding the relationship between host ancestry and the evolution of fish blood flukes. Most notable among those under-explored host lineages are the lower (basal) actinopterygians (sensu Grande and Bemis, 1996), which form a non-monophyletic group of convenience that includes extant members allocated to 2 distinct lineages, i.e., Acipenseriformes (including paddle-fishes [Polyodontidae] and sturgeons [Acipenseridae]) and Polypteriformes (including bichirs [Polypteridae] only) (e.g., Grande and Bemis, 1996; Nelson, 2006). Herein, we provide the first name and description for an aporocotylid that infects a basal actinopterygian, the American paddlefish *Polyodon spathula* (Walbaum, 1792) (Acipenseriformes: Polyodontidae) and propose a new genus to accommodate this new species.

MATERIALS AND METHODS

American paddlefish from the Mississippi Delta were captured with gill nets in April of 2004 and 2006. All fish were killed by spinal severance, and immediately afterwards the heart was extracted, placed in a sample bag, bisected to expose its lumen, immersed in an anticoagulant solution of ~5.0 gm NaCl and ~2.0 gm NaCl-citrate/L of distilled water, and kept in a cooler with a small amount of ice for several hours. Upon returning to the laboratory, the contents of the bag were examined with the aid of a dissecting microscope. Observations of living flukes were made with the aid of dissecting and compound microscopes. Flukes intended as whole mounts were killed with heat from an ethanol-burner flame, under little or no coverslip pressure, and transferred to a vial of 5% neutral buffered formalin (n.b.f.). Whole mounts were stained in Van Cleave’s hematoxylin with several additional drops of Ehrlich’s hematoxylin, made basic at 70% ethanol with lithium carbonate and butyl-amine, dehydrated, cleared in clove oil, and mounted in Canada balsam. Two specimens were embedded in paraffin, serially-sectioned at 4 μm, and stained with Gill’s hematoxylin and eosin. Three specimens for scanning electron microscopy (SEM) were dehydrated, immersed in hexamethyldisilazane for 30 min, air-dried for 45 min, and sputter-coated with gold-palladium. Drawings were made with the aid of a drawing tube and facilitated by differential interference contrast (DIC) optical components. Measurements are reported in μm and given as ranges with the sample size in parentheses. The specimen of “*Spirochis* sp.” of Appy and Dadswell (1978) (USNPC No. 73138) was lost and presumably destroyed during Hurricane Katrina on 29 August 2005.

Fish classification and higher level taxon names used herein follow primarily Grande and Bemis (1996) and Nelson (2006). Because “fishes” comprise a paraphyletic assemblage if one excludes Tetrapoda, we herein use that term not as a taxonomic rank but rather, as stated by Nelson (2006), “as a matter of convenience, essentially to describe...
those vertebrates studied by ichthyologists and covered in ichthyological courses."

The specimens intended for molecular analyses were fixed directly in 95% EtOH, and genomic DNA was extracted using a DNeasy tissue kit (QIAGEN, Valencia, California) according to the manufacturer’s instructions. Polymerase chain reaction (PCR) was used to amplify SSU rDNA with the forward primer 18SE (5'-CGG ATT TCG TCG ACA ACC TGG TTT ATC CTG CCA GT) and the reverse primer WORMB (5'-CAG AGT CTG GAA GAC GAT C) with long posterior ceca only and no anterior cecum, smooth, lacking diverticula or secondary rami, extending posteriorid to near body end. Testes intercecal, non-contiguous, an anterior testicular column plus 1 testis posteriorly, with extensively lobed margins. Cirrus sac clearly delineated from surrounding tissue, enveloping seminal vesicle, prostatic gland cells, and cirrus; cirrus evertting dorsally near dextral body margin; prostatic gland cells spherical, encircling ejaculatory duct at base of cirrus and proximal to common genital pore. Ovary single, medial, intercecal, separating anterior column of testes from single posterior testis, immediately posterior to testicular column, deeply lobed, located within posterior 1/4 of body. Oviduct sinistral, functioning as oviducal seminal receptacle. Vitellaria an extensive network of narrow, interconnecting branching bands, situated both dorsal and ventral to gonads and ceaca, not extending laterally beyond ventrolateral nerve cords. Laurer’s canal (sensu Snyder, 2004) and provisionally aligned using Clustal W (Thompson et al., 1994), and included Alaria alata (AY22098) was associated, Sunderland, Massachusetts) (Maddison and Maddison, 2003). Positions for which alignment was ambiguous were removed before analysis. Maximum parsimony analysis of these data was performed using the “branch and bound search,” “random sequence addition,” and “TBR branch-swapping” options of PAUP* (v. 4.0b10, Sinaur and Associates) (Swoford, 2001). Gaps were treated as missing data, and characters were unordered with equal weight. Nodal support was assessed using bootstrap resampling (Felsenstein, 1985) (1,000 bootstrap replicates, 100 heuristic searches/replicate).

**DESCRIPTION**

*Acipensericoloria* n. gen. (Figs. 1–28)

**Diagnosis:** Body flat, ventrally concave, elongate. <4 times longer than wide, with anterior and posterior ends tapering approximately equally, spined; tegumental body spines robust, spike-like, lacking recurved tip, in ventrolateral transverse rows. Rosethorn-shaped spines absent. Lateral nerve cord extending nearly entire body length, appearing subterminal in anterior body end, with commissure anteriorly. Dorso-lateral nerve cords indistinct. Sensory papillae abundant, occupying ventrolateral body surface between lateral nerve cord and body margin. Anterior sucker bowl-shaped, centered on mouth, demarcated from the anterior body end by peduncle, having minute spines on inner anteroverentral surface only. Pharynx between anterior sucker and nerve commissure, highly muscular, intensely basophilic. Esophagus medial, straight, ventral to anterior nerve commissure, extending posteriad approximately 1/6 of body length; posterior esophageal swelling present immediately anterior to cecal bifurcation. Anterior testicular column and posterior testis immediately posterior to testicular column, deeply lobed, located within posterior 1/4 of body. Oviduct sinistral, functioning as oviducal seminal receptacle. Vitellaria an extensive network of narrow, interconnecting branching bands, situated both dorsal and ventral to gonads and ceaca, not extending laterally beyond ventrolateral nerve cords. Laurer’s canal (sensu Snyder, 2004) and provisionally aligned using Clustal W (Thompson et al., 1994), and included Alaria alata (AY22098) was associated, Sunderland, Massachusetts) (Maddison and Maddison, 2003). Positions for which alignment was ambiguous were removed before analysis. Maximum parsimony analysis of these data was performed using the “branch and bound search,” “random sequence addition,” and “TBR branch-swapping” options of PAUP* (v. 4.0b10, Sinaur and Associates) (Swoford, 2001). Gaps were treated as missing data, and characters were unordered with equal weight. Nodal support was assessed using bootstrap resampling (Felsenstein, 1985) (1,000 bootstrap replicates, 100 heuristic searches/replicate).

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Acipensericoloides petersoni from the heart of Polyodon spathula, ventral view. (1) Body of holotype showing anterior sucker (as), pharynx (ph), nerve commissure (nc), esophagus (es), cecal bifurcation (cb), testes 1–6 (t1–t6), ovary (o), and excretory pore (ep). Bar = 500 μm. (2) Transverse rows of ventrolateral body spines, paratype. Bar = 50 μm. (3) Juvenile, body showing anterior sucker (as), pharynx (ph), esophagus (es), and intestinal anlagen (ia). Bar = 200 μm. (4) Genitalia, composite, ventral view, showing posterior-most testis of testicular column (t5), uterus (u), uterine eggs (ue), anterior trunk of vasa efferentia (ave), vas deferens (vd), ovary (o), everted cirrus (ec), seminal vesicle (sv), primary vitelline collecting duct (vt), Laurer’s canal (lc), ootype (oo), posterior trunk of vasa efferentia (pve), oviduct (ov) with sperm and serving as oviducal seminal receptacle, and posterior-most testis (t6). Bar = 200 μm.

Prepared specimens: Body 2,405–4,026 (10) long, 810–1,325 (10) wide, 2.9–3.2 times longer than wide (Fig. 1); dorsum with honeycomb-like surface features (Figs. 21, 22, 28); ventral surface relatively smooth medially (Fig. 18). Body spines 7–12 (8) long (Figs. 2, 11), nearly indistinct in some whole mounted specimens; proximal end broadly rounded (Fig. 2); distal end with sharp tip protruding only slightly from tegument (Fig. 24). Spine rows 10–12 (7) long, numbering 110–135 (3) per side, indistinct in posterior region of some specimens, not contiguous posteriorly. Ventrolateral nerve cord becoming confluent with paired cord 75–233 (5) or 4–6% of body length from posterior body.
FIGURES 5–6. *Acipenserericola petersoni* from the heart of *Polyodon spathula*, ventral view. (5) Posterior body end of paratype showing posteriormost testis (t6), mononucleate cells (mc) of cecal wall, distal end of dextral cecum (c), nerve cord (nc), excretory vesicle (ev), and dorsal, sub-terminal excretory pore (ep). Bar = 200 μm. (6) Paratype showing junction of Laurer’s canal (Lc), vitelline duct (vt), and oviduct (ov). Ootype (oo), glandular ducts (gd), and proximal portion of uterus (ut). Bar = 100 μm.

end (Figs. 1, 5); ventrolateral nerve commissure 139–273 (9) or 5–8% of body length from anterior body end, 64–140 (9) across width of worm, 20–35 (7) in diameter, perpendicular to midline of body (Fig. 1). Ventral sensory papillae 5–12 (10) in diameter, appearing volcano-shaped with light microscopy (Fig. 11) and nipple-like with SEM (Figs. 18, 20, 23), arranged in ventrolateral bands, delimited by approximate track of ventrolateral nerve cord and lateral body margin (Figs. 18, 23), indistinct or absent along midline of ventral and dorsal body surface. Anterior sucker 75–159 (9) in diameter or 9–13% of body width (Figs. 1, 7, 12, 18, 19, 21, 22, 25); anterior sucker spines about 3–5 long, conical, directed posteriorly, clustered, not occurring in clearly delineated rows (Figs. 24, 26, 27). Pharynx 139–199 (10) long or 38–53% of esophagus length, 109–144 (10) wide or 4–11 times esophagus width, with muscular wall 30–55 (10) thick, nearly as thick as anterior body end (Figs. 1, 7, 12). Esophagus 273–497 (10) long or 10–14% of body length, 10–25 (10) wide, with wall 5–10 (10) thick at level immediately posterior to pharynx; posterior swelling 40–100 (10) wide or 2–7 times width of anterior esophagus at level immediately posterior to pharynx, 104–224 (10) long or 31–67% of esophagus length, with wall 15–35 (10) thick (Figs. 1, 13). Esophageal gland indistinct surrounding swelling. Ceca bifurcating immediately posterior to esophageal swelling, 388–647 (10) or 14–19% of body length from anterior body end, extending posteriad in parallel 1,792–2,995 (10) or 72–79% of body length, ending 259–408 (10) or 9–13% of body length from posterior body end, 40–135 (10) wide, not extending laterally beyond ventrolateral nerve cord (Figs. 1, 3), with wall having cuboid basophilic mononucleate cells (Figs. 5, 14), containing granular material within lumen in some individuals; granular material blackish, probably comprising hematin and other compounds formed in decomposition of hemoglobin and erythrocytes, nearly filling cecal lumen in some specimens (Figs. 8, 10).

Testes spheroid, 6 in number, each approximately equal in diameter, comprising 5 testes (t1–t5) oriented in a single testicular column plus 1 separate testis (t6) posteriorly; testicular column intercecal, 933–1,743 (10) long or 35–50% of body length, with anterior-most testis in column 221–565 (10) or 9–21% of body length from cecal bifurcation, 173–393 (10) wide or 20–32% of body width at widest level; posterior-most testis intercecal, between distal ends of ceca, 259–512 (10) long or 9–13% of body length, 114–299 (10) wide or 11–24% of body width, 313–597 (10) or 11–16% of body length from posterior margin of an-

FIGURES 7–10. *Acipenserericola petersoni* from the heart of *Polyodon spathula*, adult, longitudinal sections. (7) Anterior region showing anterior sucker (as) and pharynx (ph). Dorsal (d) and ventral (v) surfaces. Bar = 50 μm. (8) Blackish residue, probably partially-digested blood components, occupying the cecal lumen (cl). Bar = 50 μm. (9) Posterior end showing excretory vesicle (ev) and subterminal excretory pore (ep) opening on dorsal surface. Bar = 50 μm. (10) Terminal genitalia showing common genital pore (arrow), cirrus (c), and cirrus sac (cs). Bar = 25 μm.
Laurer's canal extending 74–136 (7) anteromedially from distal region containing sperm for entire length, 25–99 (7) in maximum width (Fig. 4). Margin of t6, extending anteriad while curving toward midline, extending posteriad while curving mediad, recurving dorsally at anterior long or 17–24% of body length (Fig. 4). Oviduct sinuous, ultimately to vas deferens and seminal vesicle; post-ovarian space 465–870 (10) wide or 26–41% of body width, 0.9–1.5 times wider than long, dorsal lacking spines; post-cirrus sac space 475–811 (10) long or 17–23% of dextral, 65–99 (2) long, 30–50 (5) wide, directing anteriad, finger-like, body margin, containing sperm in all specimens; extruded cirrus dorsal, 131 (9) wide or 1.7–2.4 times longer than wide, orienting toward dextral laterally beyond dextral ventrolateral nerve cord, 99–273 (9) long, 59–35 in diameter. Pharynx 60 long, 40 wide. Esophagus 102 long or 15% of body length, 8 wide immediately posterior to pharynx, extending posterior loop of body length (Figs. 1, 3). Vasa efferentia an interconnecting meshwork of fine ducts entwining throughout testicular tissue, containing sperm in all specimens, with anterior and posterior trunks linking testicular column and posterior-most testis (Figs. 1, 4), difficult to trace in fixed specimens; anterior trunk of vasa efferentia 14–174 (4) long, extending posterior from posterior margin of t5 and curving dextrad before uniting with posterior trunk, 10–17 (4) wide (Fig. 4); posterior trunk of vasa efferentia extending 224–373 (4) or 7–12% of body length from anterior margin of t6 before uniting with anterior trunk and forming vas deferens, 5–12 (4) wide (Fig. 4); vas deferens dextral, 40–99 (6) long, 5–27 (6) wide, extending a short distance dorsolaterally before joining with cirrus sac and seminal vesicle, containing sperm in all specimens (Fig. 4). Cirrus sac thin-walled, enclosing seminal vesicle and glandular cells (Figs. 4, 10); seminal vesicle ovoid, between t5 and t6, not extending laterally beyond dextral ventrolateral nerve cord, 99–273 (9) long, 59–131 (9) wide or 1.7–2.4 times longer than wide, orienting toward dextral body margin, containing sperm in all specimens; extruded cirrus dorsal, dextral, 65–99 (2) long, 30–50 (5) wide, directing anterior, finger-like, lacking spines; post-cirrus sac space 475–811 (10) long or 17–23% of body length.

Ovary 189–448 (10) long or 8–10% of body length, 213–472 (10) wide or 26–41% of body width, 0.9–1.5 times wider than long, dorsal to vas deferens and seminal vesicle; post-ovarian space 465–870 (10) long or 17–24% of body length (Fig. 4). Oviduct sinuous, ultimately extending posterior while curving mediad, recurving dorsally at anterior margin of t6, extending anterior while curving toward midline, containing sperm for entire length, 25–99 (7) in maximum width (Fig. 4). Laurer’s canal extending 74–136 (7) anteromedially from distal region of oviduct, immediately before union of vitelline duct and oviduct, opening on dorsal surface, 7–10 (7) wide, highly glandular. Primary vitelline collecting duct sinistral, ventral, extending posterior and following a track between or ventral to testicular column and intestinal cecum, extending 174–448 (9) posterior from margin of t5, uniting with oviduct immediately before joining ootype and short distance distal to where Laurer’s canal joined (Figs. 1, 4, 6). Ootype 45–69 (9) in diameter, medial, at level of cirrus sac, anterior to major posterior loop of oviduct, dorsal to vasa efferentia (Figs. 1, 5, 17); Mehlis’ gland diffuse, surrounding ootype, with many hair-like ducts (Fig. 17). Uterus primarily dextral for most of length, running anterior a short distance along midline from ootype, ventral to ovary, dorsal to vas deferens, becoming somewhat convoluted lateral to ovary, looping anterior before running posterior, 20–37 (10) in maximum width (Figs. 1, 4). Uterine eggs 35–50 (7) long, large relative to many other aporocotylids, curved, crenulated, or collapsed completely in many specimens, lacking apparent operculum, with thin membranous shell when in distal portion of uterus (Fig. 4); ejected eggs seemingly more regular in shape (Fig. 15); mature egg in gill of definitive host not observed.

Excretory vesicle oblong, 71–146 (9) long, 10–32 (9) wide; excretory pore subterminal, dorsal, 42–128 (9) or 2–3% of body length from posterior body end (Figs. 1, 5, 9, 16); system difficult to observe in fixed specimens.

**Diagnosis of juvenile (based on 1 whole mounted specimen from heart):** Body 663 long, 98 wide; ventrolateral body spines indistinct. Nervous system indistinct. Sensory papillae not evident. Anterior sucker 35 in diameter. Pharynx 60 long, 40 wide. Esophagus 102 long or 15% of body length, 8 wide immediately posterior to pharynx, extending directly posterior; posterior esophageal swelling 28 long, 12 wide.
Acipensericola petersoni from the heart of Polyodon spathula, adult specimens, scanning electron micrographs. (18) Body, ventral view. Bar = 100 μm. (19) Ventral aspect of anterior sucker, lateral view. Bar = 10 μm. (20) Sensory papillae (circled) on ventral surface of body. Bar = 5 μm. (21) Body, dorsal view. Anterior sucker (as). Bar = 100 μm. (22) Anterior sucker (as), dorsal view. Bar = 100 μm. (23) Ventrolateral surface of body showing dispersion of sensory papillae (sp). Bar = 10 μm. (24) Spine row showing protruding tips of ventrolateral body spines (s). Bar = 0.5 μm. (25) Anterior sucker (as) and mouth (m), ventral view. Bar = 10 μm. (26) Higher magnification of Figure 25 showing anterior sucker spines (s). Bar = 4 μm. (27) Higher magnification of Figure 26 showing cluster of spines on the inner anterovelateral surface of the anterior sucker. Bar = 2 μm. (28) Tegument of body, dorsal view. Bar = 10 μm.

Esophageal gland indistinct. Cecal anlagen appearing as a sac-like medially-positioned mass (Fig. 3). Terminal genitalia, gonads, and excretory system not evident.

**Taxonomic summary**

**Type and only known host:** Polyodon spathula (Walbaum, 1792) (Acipenseriformes: Polyodontidae), the American paddlefish.

**Sites:** Adults and juvenile in atrium, ventricle, and bulbous arteriosus of heart.

**Type locality:** Six Mile Lake (33°41′14″N, 90°12′39″W), an oxbow of the Tallahatchie River near Greenwood, Mississippi. **Other locality:** Lower Lake (34°24′26″N, 89°43′12″W), a tailrace comprising the upper portion of the Little Tallahatchie River exiting Sardis Lake Reservoir, near Batesville, Mississippi.

**Specimens deposited:** Holotype USNPC No. 100676. Paratypes USNPC Nos. 100677–100678.

**Prevalence of infection:** Nine of 11 (82%) from Six Mile Lake and 6 of 6 (100%) from Lower Lake.

**Etymology:** The specific name ‘petersoni’ honors Jody Lee Peterson (Parasitology Section, Gulf Coast Research Laboratory) for his intuition and skill as a fisherman and for his invaluable field assistance to SAB during 1997 through 2007.

**Remarks**

Live specimens of Acipensericola petersoni used their flat, ventrally-concave surface and their ventrolateral body spines for attachment and locomotion. Like some other crawling aporocotylids that have been observed (Bullard and Overstreet, 2002, 2003, 2004), these flukes adhere to the walls of the heart, as well as to glass and plastic surfaces, by using the lateral body margin, presumably as a gasket that creates and maintains an internal negative pressure between the fluke’s body and the attachment surface. Specimens crawled by repeated, wave-like un-
dulations of the lateral body margins. As with adults of some species of Cardiola, as well as with Elaphroastes ecueti Bullard and Overstreet, 2003, the transverse rows of ventrolateral body spines of A. petersoni probably enhance grip and traction for attaching to, and crawling over, uneven fleshy surfaces. However, these spines apparently are not required for initial attachment and crawling since adults adhered to, and crawled over, impervious surfaces, e.g., glass and plastic, and the juvenile specimen lacked spines altogether (Fig. 3). Regarding the function of the anterior sucker, a few live specimens applied their anterior sucker to the surface of the plastic sample bag and remained anchored there, even after the bag was shaken vigorously. These specimens, however, could be dislodged by inserting the bristles of an artist’s brush beneath the rim of the anterior sucker.

Parsimony analysis of SSU data (Fig. 29) derived from a single specimen of A. petersoni, as well as from several other aporocotylids, produced a single most-parsimonious tree with nodal support ranging from 74–100 and a tree length of 1,311 (224 of the 1,815 total base pairs sequenced were informative). The tree topology showed that C. trondheimensis, a chondrichthyian blood fluke, was the most basal ingroup taxon in the tree, with A. petersoni basal to Sanguinicola cf. inermis, both of which were basal to 3 euteleost (Euteleostei) aporocotylids included in the analysis, i.e., Aporocotyle spinosicanalis, Plethorchis acanthus, and Neoparacardicola nasonis.

**DISCUSSION**

At least 1 species of the new genus infects a sturgeon in North America. This reported (Appy and Dadswell, 1978), previously collected, but unnamed aporocotylid infects the mesenteric vessels of shortnose sturgeon (Acipenser brevirostrum Lesueur, 1818 [Acipenseridae]) in the Saint John River Estuary, New Brunswick, Canada. Appy and Dadswell (1978) detailed 9 immature specimens and identified the specimens as Spirorchis sp.; however, we regard the unnamed species as congeneric with A. petersoni because it has: (1) a prominent, bowl-shaped anterior sucker that is centered on the mouth; (2) a clearly delineated pharynx that is located immediately posterior to the anterior sucker; (3) an inverse U-shaped ceca that extends to near the posterior body end; (4) a testicular column; and (5) ventrolateral body spines. Although ventrolateral body spines were neither described nor illustrated by Appy and Dadswell (1978), we examined a voucher specimen (USNPC No. 73138) and confirmed the presence of ventrolateral spines. However, since we could not discern ventrolateral body spines in the juvenile specimen of A. petersoni in our collection (Fig. 3), perhaps some of the younger specimens collected by Appy and Dadswell also lacked them. In addition, ventrolateral body spines can be overlooked quite easily because of their small size and, in poorly fixed specimens, these spines may be lost because the thin margins of the body seem to be especially vulnerable to autolysis, which can result in the subsequent detachment of the spines. The specimens of Appy and Dadswell (1978) from the sturgeon remain unidentified, and not yet ready to be described, because no adult aporocotylid material from a shortnose sturgeon has been reported or examined by us. We suspect that it represents a species distinct from A. petersoni.

As previously stated, the morphological features of the species certainly indicate that it belongs in Acipensericola, but its host’s phylogenetic affiliation (Acipenseridae rather than Polyodontidae) and geographic distribution (New Brunswick River draining to northwestern Atlantic Ocean rather than the Mississippi River drainage) suggest that it represents a new species. Addressing this taxonomic problem should require detailed necropsies of freshly killed, wild-caught shortnose sturgeons. Unfortunately, such an opportunity is rare and has proven logistically difficult for us to arrange because the shortnose sturgeon, like nearly all sturgeon species, is presently protected throughout its range.

Collectively, aporocotylids exhibit several different morphological types of suckers associated with the mouth (Bullard and Overstreet, 2003), and we think that the fine features associated with these various types of suckers help elucidate evolutionary relationships within the group. Hereinafter, we refer to them as

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**FIGURE 29.** Phylogram based on maximum parsimony analysis of small subunit ribosomal DNA. Nodal support based on bootstrap resampling is to the left of each ingroup node. Note that the aporocotylids infecting Chondrichthyes (Chimaerohemecus trondheimensis), Acipenseriformes (Acipensericola petersoni), and Ostariophysi (Sanguinicola cf. inermis) are basal to those aporocotylids that infect Euteleostei (Aporocotyle spinosicanalis, Plethorchis acanthus, and Neoparacardicola nasonis).
“anterior suckers.” The shape of the sucker, the spines of the sucker, and the location of the mouth are diagnostic for at least some genera, e.g., Elaphrobates Bullard and Overstreet, 2003. Homology of these various anterior suckers presently is uncertain for some taxa, which represents a barrier to understanding the phylogenetic interrelationships among aporocotylid genera. Demonstrating homology of these various suckers is beyond the scope of the present paper because it requires a complete phylogenetic analysis of the family; however, we regard the anterior suckers of A. petersoni and P. yamagutii as 2 slightly different variations of the same homologous sucker type, and we herein delineate that type from those of other aporocotylids by its general shape and the relative position of the mouth. In Acipenserica and Paracardicolaoides, unlike all other accepted aporocotylid genera, the anterior sucker is bowl-like, centered on the mouth, and demarcated from the anterior body end by a short trunk or peduncle that supports the sucker and can direct it anteroventrally (Figs. 1, 25; Fig. 3 of Martin [1974]). Despite these general similarities, the pharynx and spination associated with the anterior sucker differs among the species of Acipenserica and Paracardicolaoides. The anterior sucker of A. petersoni has spines on its inner anteroventral surface only and is accompanied by a muscular pharynx, whereas that of P. yamagutii reportedly lacks exposed spines and an associated pharynx. The bowl-shaped anterior sucker of Acipenserica and Paracardicolaoides is superficially like that of a spirorchiid because it is relatively large and centered on the mouth, but these aporocotylid genera have a peduncle associated with the sucker. In contrast, the spirorchiids studied by 1 of us (S.A.B.) have a peduncle associated with the sucker. It is relatively large and centered on the mouth, but these aporocotylid genera have a peduncle associated with the sucker and can direct it anteroventrally (Figs. 1, 25; Fig. 3 of Martin [1974]). Despite these general similarities, the pharynx and spination associated with the anterior sucker differs among the species of Acipenserica and Paracardicolaoides. Our understanding of the phylogenetic interrelationships of aporocotylids remains unclear because no clade-based phylogenetic hypothesis involving morphological or molecular sequence data for the majority of aporocotylid genera has been published. Two obvious obstacles to completing such a task are that (1) several of the most specious aporocotylid genera need revision, e.g., Aporocotyle Odhner, 1900, Cardicola Short, 1953, and Sanguinicola Plehn, 1905, and (2) type material for many of the named species in those genera are in poor condition or not available to borrow. Although a taxonomic revision and phylogenetic analysis of Aporocotylidae are in preparation by 1 of us (S.A.B.), additional descriptions of new aporocotylid species that infect fishes belonging to previously undocumented host lineages promise to further advance our knowledge of how these flukes evolved among various lineages of “fish.” The results of the present study offer some preliminary insight into the potential interrelationships among particular aporocotylid genera and their host groups and provide a framework from which to test future hypotheses about aporocotylid-fish co-phylogeny. For example, the morphological similarities we observed between A. petersoni and those aporocotylids that infect non-euteleost fishes indicate a strong phylogenetic affiliation among these genera. Further, the available SSU data indicate that non-euteleost aporocotylids are basal to those that infect euteleosts (Fig. 29). This preliminary result contradicts the notion that fish blood flukes lack a detectable level of phylogenetic host specificity to their definitive hosts.

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