**Lanfrediella amphicirrus gen. nov. sp. nov. Nematotaeniidae** (Cestoda: Cyclophyllidea), a tapeworm parasite of *Rhinella marina* (Linnaeus, 1758) (Amphibia: Bufonidae)

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The family Nematotaeniidae, tapeworms commonly found in the small intestines of amphibians and reptiles, includes 27 recognised species distributed among four genera: Bitegmen Jones, Cylindrotaenia Jewell, Distoichometra Dickey and Nematotaenia Lühe. The taxonomy of these cestodes is poorly defined, due in part to the difficulties of observing many anatomical traits. This study presents and describes a new genus and species of nematotaeniid parasite found in cane toads (*Rhinella marina*) from eastern Brazilian Amazonia. The cestodes were collected during the necropsy of 20 hosts captured in the urban area of Belém, Pará. The specimens were fixed and processed for light microscopy, scanning electron microscopy (SEM) and three-dimensional (3D) reconstruction. Samples were also collected for molecular analyses. The specimens presented a cylindrical body, two testes and paruterine organs. However, they could not be allocated to any of the four existing nematotaeniid genera due to the presence of two each of dorsal compact medullary testes, cirri, cirrus pouches, genital pores, ovaries and vitelline glands per mature segment. Lanfrediella amphicirrus gen. nov. sp. nov. is the first nematotaeniid studied using Historesin analysis, SEM and 3D reconstruction, and it is the second taxon for which molecular data have been deposited in GenBank.

Key words: Cestoda - Nematotaeniidae - 3D reconstruction - *Rhinella marina*

Nematotaeniidae Lühe 1910 is a family of cyclophyllidean cestodes characterised by their small size, the presence of paruterine organs and a cylindrical body with external segmentation restricted to the terminal region of the body. They also have two compact testes, ovaries and vitellaria located in the medulla (Jewell 1916, Douglas 1958, Yamaguti 1959, Jones 1987).

Adult specimens of these worms are differentiated by a number of morphological traits that are of considerable importance for the diagnosis of genera and species, including the position and number of paruterine capsules, as well as the terminal genitalia morphology. Although the taxonomy of the group is unclear, due in part to the difficulties of observing some anatomical characteristics, Jones (1987) divided the nematotaenids into four genera: the monospecifics Bitegmen Jones, 1987, and Distoichometra Dickey, 1921, Nematotaenia Lühe, 1910, with seven species, and Cylindrotaenia Jewell, 1916, with a total of 18 species.

Nematotaeniids are found in the small intestines of reptiles and amphibians, such as the cane toad *Rhinella marina* (syn. *Bufo marinus*) (Linnaeus, 1758). The natural range of *R. marina* extends from southern Texas to central Brazil, although the species has also been introduced in Florida, a number of Caribbean and Pacific islands, New Guinea and northeastern Australia (Barton 1997). The known helminth parasites of this species include 55 nematodes, 41 digenea, eight cestodes, one monogenea and an acanthocephalan (Espinoza-Jiménez et al. 2007, Espínola-Novelo & Guillén-Hernandez 2008, Santos et al. 2008, 2011).

The present study identified and describes a new nematotaeniid parasite of *R. marina* from eastern Brazilian Amazonia. This nematotaeniid represents a new genus and the diagnosis of the new taxon was based in part on analysis of the scolex and segments using scanning electron microscopy (SEM), three-dimensional (3D) reconstruction and histology. The description of the new taxon is also supported by molecular data.

**MATERIALS AND METHODS**

A total of 20 *R. marina* toads were collected from urban areas of Belém, state of Pará (PA), Brazil, between October 2007-September 2008. The cestodes found in each specimen were fixed in 2% glacial acetic acid and 3% formaldehyde in 95% ethanol for analysis using both light microscopy and SEM. Some of the specimens were fixed with 2.5% glutaraldehyde in 0.1 M sodium cacody-
late buffer (pH 7.2) and then embedded in paraffin and hydroxyethyl methacrylate resin (Leica Historesin Embedding Kit) for histological analysis.

The cestodes were stained with carmin, dehydrated in ethanol, cleared in methyl salicylate and mounted on glass slides with Entellan. Illustrations were made with the aid of a camera lucida linked to an Olympus BX41 microscope. All measurements, means and standard deviations are given in micrometers unless otherwise indicated, and the ranges are given in parentheses.

Mature segments were embedded in paraffin and sectioned transversely into 8 µm-thick serial sections, which were then stained with hematoxylin-eosin. Serial, 3 µm-thick, transversal and longitudinal sections of specimens embedded in Historesin were stained with 1% toluidine blue and used in the morphometric analyses.

Specimens prepared for SEM were post-fixed in 1% OsO4, dehydrated, mounted on stubs, sputter-coated with gold and examined using a JEOL 5310 microscope.

For the 3D reconstructions, serial longitudinal sections of Historesin-embedded samples of strobilae were photographed. The alignments and 3D reconstructions were conducted using the RECONSTRUCT™ software. Images from stained and cleared specimens were captured on different focal planes using a Nikon Eclipse E600 microscope equipped with differential interference contrast and a single image with the best-adjusted focus was obtained using the CombineZP software.

Genomic DNA (gDNA) was obtained from whole worms using the ChargeSwitch gDNA Mini Tissue Kit (Invitrogen Life Technologies, Gaithersburg, MD, USA). The 18S small subunit ribosomal RNA (18S rRNA) gene was partially amplified via polymerase chain reaction (PCR) in two overlapping fragments using the primers Ces1/Ces2 and 2880/B and following the procedure of Skeriková et al. (2001). The PCRs were performed in 25 µL final volume, containing 5-10 ng of DNA, 50 mM KCl, 2 mM MgCl2, 10 mM Tris-HCl, 50 µM of each dNTP, 0.5 µM of each oligonucleotide and one unit of Taq DNA polymerase (Invitrogen). Reactions containing the Ces1/Ces2 primers were denatured for 4 min at 94ºC followed by 35 cycles of 30 s at 94ºC for denaturing, 30 s at 60ºC for annealing and a 30 s extension at 72ºC with a final 5 min extension at 72ºC. Reactions containing the 2880/B primers were amplified following the same procedure, except for an annealing temperature of 62ºC. The amplified fragments were cycle-sequenced in a 3130 Genetic Analyzer (Applied Biosystems) according to the manufacturer’s specifications. Both strands of individual DNA fragments were sequenced to confirm ambiguous base calls. The sequences were merged using BioEdit (Hall 1999).

To evaluate the phylogenetic signal of the 18S rRNA gene, sequences were compared with a 465 bp sequence from Distoichoemotera bufonis (GenBank accession Z98377), a closely-related parasite of bufonids.

Ethics - The present study was approved by the Animal Research Ethics Committee of the Federal University of Pará (UFPA) through authorisation CEPAE-UFPN: BIO010-10.

RESULTS

Twelve of the 20 *R. marina* specimens were infected with cestodes (a prevalence of 60%) with five to 160 parasites per host.

*Lanfreidiella* gen. nov.

(Figs 1-33)

**Diagnosis** - Strobila aplolytic becoming thinner in the terminal portion. Segments numerous, acraspedote, with external segmentation apparent in gravid and pre-gravid segments. Scolex with four suckers, simple, unarmed. Large ventral osmoregulatory canals. Double reproductive structures, including the following: bilateral genital atrium, two compact medullary dorsal testes, two lateral piniform cirrus pouches, two ovaries and two vitelline glands. Genital atrium lightly muscular. Cirri pouches thin not divided by a septum. Cirri with terminal spines. Vas deferens arise from testis loops twice, enters in medially to the cirrus pouch. External seminal vesicle absent. Paruterine organs paired. Two paruterine capsules, united basally, not surrounded by an outer envelope structure. Eggs with three membranes.

**Etymology** - The genus has been named after the late Reinalda M Lanfredi, PhD, in recognition of her valuable contributions to Brazilian helminthology and helminth systematics.

**Emended key to the genera of the Nematotaeniidae Lühe, 1910.**

1. Single group of reproductive organs (1 testis, 1 cirrus, 1 ovary and 1 vitelline gland) ........................................... 3
2. Double group of reproductive organs (2 testes, 2 cirri, 2 ovaries and 2 vitelline glands) ...... *Lanfreidiella*

3a. Two paruterine capsules or organs per segment … 4
3b. More than two paruterine capsules or organs per segment .......................................................... 5

4a. Paruterine capsules surrounded by a second envelope .............................................. *Bitegmen* Jones, 1987
4b. Paruterine capsules not surrounded by a second envelope .............................................. *Cylindrotaenia* Jewell, 1916

5a. Paruterine capsules in row of two-six pairs of organs; four-12 paruterine capsules clustered in the posterior half of the segment ...... *Distoichoemotera* Dickey, 1921
5b. Paruterine organs not paired; five-150 paruterine capsules or paruterine organs distributed randomly throughout the segment ...... *Nematotaenia* Lühe, 1899

*Lanfreidiella amphicirrus* gen. nov. sp. nov.

(Figs 1-33)

**Description** - Body elongated, 70 mm ± 1.75 (4.0-9.0) (n = 20) long, scolex 541.46 ± 100.70 (373.33-706.67) in diameter, four oval acetabula, with no rostelum or apical organ. Suckers 151.10 ± 16.31 (128.67-183.00) × 110.94 ± 6.81 (99.33-122.00) (Fig. 1); excretory canal ventral, 19.45 ± 3.63 (13.30-26.60) in diameter; mature segments 38.66 ± 2.98 (33.00-44.00) × 333.83 ± 10.14 (322.66-354.00); testes oval, 88.12 ± 27.15 (84.40-116.00) × 31.52 ± 4.17 (25.90-38.90) (Figs 2-4); two
thin-walled piriform, cirrus pouches, not divided by a septum, 40.05 ± 1.50 (36.84-41.57) × 18.00 ± 1.05 (15.78-19.47) (Figs 4, 5, 11-13); genital atrium lightly muscular, cirri straight, with terminal spines, 16.54 ± 0.97 (15.52-18.15) in length (Figs 24, 26, 27); two ovaries, spherical to oval, situated in the middle of the segment, 48.24 ± 4.56 (38.90-55.19) × 38.47 ± 6.30 (30.51-50.00); two vitelline glands, oval, dorsolateral to the ovaries, 30.38 ± 4.74 (24.00-38.90) in diameter (Fig. 11); uterus dorsal or dorsolateral to the ovaries (Figs 4, 5, 11); pregravid segments 80.76 ± 8.42 (76.13-98.73) × 853.41 ± 36.63 (789.42-894.73); paruterine complex 96.05 ± 15.76 (65.78-110.52) × 34.29 ± 5.79 (28.31-47.36); gravid segments 259.67 ± 65.08 (191.56-374.68) × 251.95 ± 63.16 (200.00-388.96); two paruterine capsules, oval to spherical, 67.20 ± 8.29 (51.95-77.92) × 34.29 ± 5.79 (28.31-47.36) (Figs 6-9); seven-10 oncospheres per capsule and 15-20 per segment; oncospheres with an oval outer envelope 30.15 ± 3.18 (26.31-33.94) × 16.45 ± 1.61 (13.94-16.68), embryophore 18.83 ± 5.88 (17.30-22.36) × 13.73 ± 1.73 (12.76-15.52) and internal hooks, 7.85 ± 1.07 (6.57-9.89) in length (Fig. 10).

The arrangement of the cirrus pouches was reconstructed from the 116 serial longitudinal dorsoventral sections of mature strobila segments. This 3D reconstruction revealed the equidistant distribution of the cirrus pouches that line each lateral margin of the mature segment of the strobila, between opposing pouches. This reconstruction confirms the presence of two sets of reproductive structures in each proglottid (Figs 14, 15).

Figs 1-10: light microscopy of *Lanfrediella amphicirrus* gen. nov. sp. nov. 1: dorsoventral view of the scolex (Bar = 100 µm); 2: immature segments in a dorsoventral view (Bar = 50 µm); 3: mature segments (Bar = 50 µm); 4: cross section of mature segment showing two groups of reproductive organs and two cirrus pouches (Bar = 50 µm); 5: reconstruction of a cirrus pouch with cirri showing the genital atrium, deferent channel and vaginal channel (Bar = 50 µm); 6: pregravid strobila segments showing the initial region of segmentation (Bar = 100 µm); 7: gravid segments showing the maturation of the paruterine capsules and visible segmentation (Bar = 100 µm); 8: posterior extremity showing gravid segments (Bar = 100 µm); 9: detail of a gravid segment where it is possible to observe two paruterine capsules per segment (Bar = 50 µm); 10: details of eggs showing the outer envelope, embryophore, and the oncosphere (Bar = 30 µm).
Type host - *R. marina* (Linnaeus, 1758) (Amphibia: Bufonidae), Cane Toad, Giant Toad.

Site of infection - Small intestine.

Type-locality - Belém, PA (01º28'03”S 48º20'18”W).

Type data and depository - Holotype, deposited in the Helminthological Collection of the Oswaldo Cruz Institute (CHIOC) of the Oswaldo Cruz Foundation in Rio de Janeiro, Brazil, under catalogues CHIOC 37314a (holotype) and CHIOC 37314b, 35706 (paratype).

Molecular data - A partial sequence (1,253 nucleotides) of the 18S rRNA gene was obtained from *L. amphicirrus* gen. nov. sp. nov. and added to GenBank under accession HM185494. A comparison with *D. bufonis* identified a number of sites with potential for phylogenetic differentiation, including 18 substitutions and 15 indels, restricted to the 5’ half of the alignment (not shown).

SEM of *L. amphicirrus* gen. nov. sp. nov. (Figs 16-26, 28-33) - SEM revealed that the scolex has simple suckers and no apical organ (Figs 16-19). The microtriches observed throughout the surface of the parasite are delicate, very short and filiform (Figs 20-22). The body presents distinct segmentation only in its most posterior portion. Two genital pores were observed in the mature segments.
Some mature segments have partially extrverted, conical cirri with spine-like structures located distally (Figs 24, 26, 28). Genital pores could not be observed in the gravid segments (Figs 30–33).

Host-parasite data - Prevalence: 45%.

Etymology - The species name is derived from the Greek prefix amphi-, meaning both, and thus refers to the double cirri and other reproductive structures.

DISCUSSION

The new genus *Lanfrediella* was allocated to the Nematotaeniidae family due to the presence of diagnostic morphological traits described for this family by Jewell (1916), Hsü (1935), Douglas (1958), Jones (1987) and Khalil et al. (1994). These traits include few testes per segment (2 or rarely 3), a cylindrical form, acraspedote segments with segmentation evident only in the posterior region of the body, the presence of conical paruterine organs immediately adjacent to the uterus, a distinct pattern of paruterine capsules (which are thin-walled that bear eggs formation developing at the anterior surface of uterus) and cylindrical strobila. All nematotaeniids are found in amphibian and reptile hosts.

Khalil et al. (1994) proposed a key to the family Nematotaeniidae that uses characteristics that are visible in the gravid and pregravid segments to distinguish the genera *Nematotaenia*, *Cylindrotaenia*, *Distoichometra* and *Bitegmen*. These genera can be differentiated primarily by the number and arrangement of the paruterine organs. *Lanfrediella* gen. nov. can be differentiated easily from *Nematotaenia* and *Distoichometra*. While the...
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A new genus has only two paruterine capsules per gravid segment, *Nematotaenia* and *Distoichometra* have more than two capsules (Hsü 1935, Douglas 1958, Yamaguti 1959, Jones 1987, Khalil et al. 1994).

*Bitegmen* has only two paruterine capsules per segment, although each capsule is surrounded by an outer enveloping structure (Jones 1987, Khalil et al. 1994). The sperm duct does not loop within the cirrus pouch, the anterior portion of the genital atrium is surrounded by muscular tissue and a septum separates the distal portions of the cirrus pouch. In contrast, mature and gravid segments of *Lanfrediella* gen. nov. lack all of these traits.

According to Jones (1987) and Khalil et al. (1994), the mature segments of *Cylindrotaenia* (Jewell 1916) have two paired paruterine organs that are united basally and gravid segments with two paruterine capsules, a feature also seen in *Lanfrediella* gen. nov. However, the new genus differs conspicuously from *Cylindrotaenia* in that it has two of each reproductive structure (cirri, cirrus pouches, testes, ovaries and vitelline glands) in each segment.

*L. amphicirrus* gen. nov. sp. nov. is the first cyclophyllidean tapeworm to be found in Amazonian *R. marina*. Specimens of *Cylindrotaenia americana* have been collected from *Rhinella* toads in southern Brazil, including the states of Paraná (Stumpf 1982), São Paulo (Jones 1987) and Rio Grande do Sul (Santos & Amato 2010).

The comparison of *Lanfrediella* gen. nov. with other nematotaeniid genera using classical techniques was
hampered by the difficulties of observing the internal organs as described by Jones (1987), showing the necessity for the use of different tools to improve the classification. This is the first study of cestodes that used Historesin, which simplified the diagnosis and observation of internal structures, such as testes, ovaries, vitelline glands, sperm ducts and cirrus pouches. Based on these results, this technique is recommended for use in future studies of these organisms. Embedding in Historesin preserved the tegument and other structures, which become retracted in specimens embedded in paraffin.

Reconstruction of complex internal structures from serial histological sections is a complex and time-consuming task (Fiala 2005); however, in the present study, 3D reconstruction facilitated the visualisation of these structures in the helminths, which are difficult to observe using standard techniques. As the segmentation of the body is difficult to observe in mature segments, this technique may become an extremely useful tool for the analysis of internal structures in nematotaeniids and the development of a more systematic approach to the taxonomy of these helminths.

This is also the first study of a nematotaeniid parasite using SEM, which revealed morphological details of structures such as the scolex, suckers, genital pores, cirri, microtriches and segmentation of the strobila, which are representative of the genus. The use of SEM improved upon the results obtained with light microscopy in these analyses. The use of SEM also permitted the detection of two genital pores per segment and thus appears to be a valuable tool for analysing the external morphology of cestodes.

Figs 29-33: scanning electron microscopy of mature and gravid segments of *Lanfrediella amphicirrus* gen. nov. sp. nov. 29: mature segments of parasite showing the segmentation (arrows) (Bar = 100 µm); 30: gravid segment view in strobila where is possible to observe the segmentation (arrows) (Bar = 100 µm); 31: posterior end of a strobila showing the segmentation of gravid segments (arrows) (Bar = 400 µm); 32: end portion of the parasite showing the constrictions between gravid segments (arrows) (Bar = 100 µm); 33: isolated gravid segment. Note the elliptical morphology, characteristic of this stage, and the absence of genital pores (Bar = 50 µm).
The molecular database for cestodes is still very limited and most of the nucleotide sequences available in GenBank as of December 2010 were from species of medical or veterinary interest. The only available sequences from nematotaeniids are short fragments of the 18S rRNA gene of *D. bufonis* obtained by Mariaux (1998). Our molecular analysis results represent an important complement to the data available for the Nematotaeniidae, which will be valuable for future phylogenetic analyses and species classifications through the addition of new taxa and genes.

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