A new species of *Notogynaphallia* (Platyhelminthes, Geoplanidae) extends the known distribution of land planarians in Chacoan province (Chacoan subregion), South America

Lisandro Negrete¹,², Ana Maria Leal-Zanchet¹ and Francisco Brusa¹,²*

**Abstract**

**Background:** The subfamily Geoplaninae (Geoplanidae) includes land planarian species of the Neotropical Region. In Argentina, the knowledge about land planarian diversity is still incipient, although this has recently increased mainly in the Atlantic Forest ecosystem. However, other regions like Chacoan forests remain virtually unexplored.

**Results:** In this paper, we describe a new species of the genus *Notogynaphallia* of the Chacoan subregion. This species is characterized by a black pigmentation on the dorsum and a dark grey ventral surface. The eyes with clear halos extend to the dorsal surface. The pharynx is cylindrical. The main features of the reproductive system involve testes anterior to the ovaries, prostatic vesicle intrabulbar (with a tubular proximal portion and a globose distal portion) opening broadly in a richly folded male atrium, common glandular ovovitelline duct and female genital canal dorso-anteriorly flexed constituting a “C”, female atrium tubular proximally and widening distally.

**Conclusions:** This is the first report of the genus *Notogynaphallia* in Argentina (Chacoan subregion, Neotropical Region) which increases its geographic distribution in South America. Also, as a consequence of features observed in species of the genus, we propose an emendation of the generic diagnosis.

**Keywords:** Land flatworms; *Notogynaphallia*; Geoplaninae; Argentina; Chacoan subregion; Neotropical Region

**Background**

Land planarians are free-living flatworms that live in humid environments. They cannot endure desiccation since they have not developed mechanisms for water conservation (Kawaguti 1932). Therefore, they hide from the sunlight (under fallen logs and leaf litter) and exhibit greater activity at night, predating on soil invertebrates such as earthworms, snails, slugs, insects and arachnids (Negrete et al. 2014a; Prasniski and Leal-Zanchet 2009). Due to their habitat requirements and position in the food chain, land planarians are good indicator taxa in biodiversity and conservation studies (Sluys 1998).

The genus *Notogynaphallia* Ogren & Kawakatsu, 1990, as originally proposed, included 23 species which could clearly be distinguished in at least two groups (Leal-Zanchet and Froehlich 2001, 2006; Froehlich and Leal-Zanchet 2003). Carbayo (2010) removed six species, which have striped dorsum, dorsal eyes and an extrabulbar prostatic vesicle to a new genus named *Luteostriata* Carbayo, 2010. Later, based on morphological and molecular data for some species, the genus *Notogynaphallia* was re-defined, including now seven species, besides four morphologically poorly known species (*Notogynaphallia* incertae sedis) (Carbayo et al. 2013). Species of *Notogynaphallia* so far have been described for Brazil (over 70 % of the species) and also...
for Colombia, Panama, Paraguay and Peru (Tyler et al. 2006–2015; Carbayo et al. 2013). This genus groups Geoplaninae species of small and medium-sized body (16–70 mm in length), with a reproductive system that includes a dilated intrabulbar prostatic vesicle opening broadly into a richly folded male atrium, in which folds form an eversible penis. Also, the distal ascending portions of the oovivitelline ducts are arranged laterally to the posterior portion of the female atrium, joining each other behind it. Additionally, the female genital canal is dorso-anteriorly flexed, arising from the posterior region of the female atrium (Ogren and Kawakatsu 1990; Carbayo et al. 2013).

In Argentina, where the land planarian diversity is still incipiently known, there are no records of this genus. Recently, the research about land flatworms has increased, mainly in the Atlantic Forest ecosystem (Negrete and Brusa 2012; Negrete et al. 2014a, b). However, other regions, such as the Chacoan forests, remain virtually unexplored. There are few old records of land planarians in the Chacoan province (Neotropical Region) in Asunción, Paraguay (Graff 1894, 1899). Unfortunately, the original landscape has suffered changes since then, mainly by deforestation and population growth, and there are no recent records of land planarians in this region.

In this paper, we describe a new species of land planarian, which represents the first land planarian species from this ecosystem in Argentina and the first record of Notogynaphallia in this country. Also, as a result of new features observed in species of the genus, we propose an emendation of its diagnosis.

**Methods**

Land planarians were collected between 2007 and 2012 in native forests within La Marcela farm (26°17′35″S, 59°06′67″W) in Formosa province, north-eastern Argentina. The ecosystem belongs to the Chacoan province, Chacoan subregion (Neotropical Region), which extends through southern Bolivia, western Paraguay, a small portion of southern Brazil and north central and eastern Argentina (Morrone 2000, 2014). It is characterized by savannahs, halophyllum steppes and xerophyllous caducifolious forests (Morrone 2000). However, in this region, the native forests are highly fragmented and reduced to small patches due to farming. The specimens were manually collected during the day beneath fallen logs in these forest fragments. Some animals were directly fixed in 10 % formaldehyde, and others were killed in boiling water and then fixed in 10 % formaldehyde and conserved in 70 % ethanol. Land planarians were sectioned in fragments, dehydrated in an ascending series of ethanol and embedded in Paraplast®. Sagittal and transverse serial sections (6–8 μm thick) of the anterior region, transverse sections of the pre-pharyngeal region (6–8 μm thick) and sagittal serial sections of the pharynx and copulatory apparatus (6–8 μm thick) were performed with a microtome and stained with Masson’s trichrome and haematoxylin–eosin methods (Bancroft and Gamble 2008). The cutaneous musculature height to body height ratio (cutaneous muscular index, CMI) was calculated according to Froehlich (1952a) from transverse sections at the pre-pharyngeal level, and the parenchymatic musculature height to body height ratio (parenchymatic muscular index, PMI) was also calculated at the same level according to Winsor (1983).

The type specimens were deposited in the Invertebrate Collection at Museo de La Plata (MLP), Argentina.

**Results**

**Taxonomy**

Family Geoplanidae Stimpson, 1857

Subfamily Geoplaninae Stimpson, 1857

Genus Notogynaphallia Ogren & Kawakatsu, 1990

Emended diagnosis of Notogynaphallia

Geoplaninae of small-to-medium-sized body (16–70 mm in length); slender body with margins nearly parallel; dorso-ventral sides slightly convex. Eyes monolobulated marginally arranged along the body, sometimes spreading dorsally. No cephalic specializations. Cutaneous musculature not insunk into parenchyma. Parenchymatic longitudinal musculature absent. Pharynx cylindrical or bell-shaped. Prostatic vesicle intrabulbar, dilated, broadly communicated with the richly folded male atrium; penis papilla eversible. Ascending portion of oovivitelline ducts lateral to posterior region of the female atrium and joining each other behind it; female genital canal dorso-antiorily flexed, arising from the posterior region of the female atrium; female atrium irregular and narrow. Adenodactyls or glandulo-muscular organs absent.

**Notogynaphallia nawei** sp. nov. urn:lsid:zoobank.org:act:B36A4C5F-452-4502-8AF5-774460DCF61A

**Type series**

Holotype: MLP-He 6807. Formosa, Argentina, 17 September 2012; anterior region 1: transverse sections on 14 slides (6–8 μm thick); anterior region 2: sagittal sections on 32 slides (8 μm thick); pre-pharyngeal region: transverse sections on 6 slides (8 μm thick); pharynx and copulatory apparatus: sagittal sections on 47 slides (7 μm thick).

Paratype 1: MLP-He 6808. Formosa, Argentina, 17 September 2012; anterior region 1: transverse sections on 12 slides (6 μm thick); anterior region 2: sagittal sections on 42 slides (8 μm thick); pre-pharyngeal region: transverse sections on 6 slides (8 μm thick); pharynx and copulatory apparatus: sagittal sections on 56 slides (7 μm thick).
Paratype 2: MLP-He 6808. Formosa, Argentina, 17 September 2012; anterior region: sagittal sections on 29 slides (8 μm thick); pre-pharyngeal region: transverse sections on 7 slides (8 μm thick); pharynx and copulatory apparatus: sagittal sections on 25 slides (6 μm thick).

Paratype 3: MLP-He 6809. Formosa, Argentina, 3 October 2007; anterior region: sagittal sections on 18 slides (8 μm thick); pre-pharyngeal region: transverse sections on 8 slides (8 μm thick); pharynx and copulatory apparatus: sagittal sections on 24 slides (8 μm thick).

Paratype 4: MLP-He 6809. Formosa, Argentina, 3 October 2007; anterior region: sagittal sections on 18 slides (8 μm thick); pre-pharyngeal region: transverse sections on 8 slides (8 μm thick); pharynx and copulatory apparatus: sagittal sections on 24 slides (8 μm thick).

Paratype 5: MLP-He 6809. Formosa, Argentina, 3 October 2007; pre-pharyngeal region: transverse sections on 23 slides (6 μm thick); pharynx and copulatory apparatus: sagittal sections on 68 slides (6 μm thick).

Paratype 6: MLP-He 6809. Formosa, Argentina, 3 October 2007; anterior region 1: transverse sections on 22 slides (6 μm thick); anterior region 2: sagittal sections on 16 slides (6 μm thick); pre-pharyngeal region: transverse sections on 21 slides (6 μm thick); pharynx and copulatory apparatus: sagittal sections on 26 slides (6 μm thick).

Paratype 7: MLP-He 6809. Formosa, Argentina, 5 October 2007; pre-pharyngeal region: transverse sections on 6 slides (7 μm thick); pharynx and copulatory apparatus: sagittal sections on 23 slides (7 μm thick).

Paratype 8: MLP-He 6809. Formosa, Argentina, 5 October 2007; anterior region 1: transverse sections on 16 slides (6 μm thick); anterior region 2: sagittal sections on 18 slides (6 μm thick); pre-pharyngeal region: transverse sections on 5 slides (6 μm thick); pharynx and copulatory apparatus: sagittal sections on 18 slides (6 μm thick).

Diagnosis
Species of Notogynaphallia with dorsal surface black and ventral surface dark grey with a thin longitudinal whitish line along the body; eyes dorsal, with clear halos; glandular margin absent; prostatic vesicle intrabulbar, with two regions: a tubular proximal portion and a globose distal portion with narrow lumen; common glandular ovovitel-line duct and female genital canal constituting a “C”; female atrium tubular proximally and wide distally.

Type locality. La Marcela farm, Pirané Department, Formosa Province, Argentina.

Habitat. The flatworms were found on the ground under fallen logs and palm leaves in environments with native vegetation.

Etymology. The specific name derives from the language of Toba indigenous people that inhabit the Chacoan region; nawe means black, which refers to the colour of the dorsal surface of this species.

Description

External morphology
The body is elongated with parallel margins. In live specimens, the anterior end is pointed, becoming rounded after fixation. The posterior end narrows abruptly, ending sharp-edged. The colour pattern of the dorsal surface of live specimens is black (Fig. 1). The ventral surface is dark grey with a whitish and thin median longitudinal line along the body (Fig. 2a), which becomes weaker behind the gonopore. The eyes, with small clear halos, surround the anterior tip uniserially in an irregular row, and they extend toward both body margins for 2–5 mm in two and three longitudinal rows. Further back, they spread to both sides of the dorsal surface, forming various rows (15–20) at the end of the first third of body. At the level of the pharynx, the eyes are still dorsal (6–8 rows of eyes). They become less numerous in the copulatory apparatus region, and a few reach the posterior end (Fig. 2b).

After fixation, body length of the specimens ranged between 19 and 42 mm. The maximum width was 2.4–8.3 mm and maximum height 1.2–2.4 mm. The mouth was located at a distance of 54–62 % from the anterior tip and the gonopore at 69–79 % (Table 1).

Internal morphology

Cephalic region
There are no musculo-glandular specializations. The dorsal epidermis is ciliated from the anterior tip until 1.9–2.5 mm (Fig. 3a). The creeping sole on the ventral epidermis is narrow (30–40 % of body width) and progressively widens...
The epidermis receives abundant secretions from xanthophil glands of two types (amorphous and fine granular secretions) and erythrophil and cyanophil glands with granular secretions. Also, rhabditogen secretions are abundant both in dorsal (rhammites) and ventral (small rhabdites) epidermis (Fig. 2a). Sensory pits, as simple invaginations (25–40 μm deep), surround the anterior tip (Fig. 3b, c). They spread along the ventral body margins in a single row forming the sensory border until ~6 mm from the anterior tip (Fig. 2a).

Epidermis and musculature at pre-pharyngeal region
The dorsal epidermis is 20–40 μm high, and ventrally, it is 25–40 μm high. The ventral epidermis is ciliated on the creeping sole, which is approximately 90–95 % of the body width (Fig. 2a, Table 1). Rhabditogen cell bodies are located in the parenchyma, below the cutaneous musculature (Fig. 3d). Their glandular secretions are abundant in the dorsal epidermis and the body margins (rhammites) and in the ventral epidermis partially occupy its height (small rhabdites) (Figs. 3d, e and 4). Numerous glands with fine and coarse granular erythrophil secretion and glands with fine granular and amorphous cyanophil secretion, whose cell bodies are located into the parenchyma, open through the entire epidermis. The creeping sole receives openings of abundant cells with erythrophil coarse granular secretion and cyanophil amorphous secretion and less numerous cells with erythrophil fine granular secretion and erythrophil amorphous secretion. There is no glandular margin.

The cutaneous musculature is composed of a subepithelial circular layer followed by a diagonal layer with decussate fibres and a longitudinal layer arranged in bundles. Cutaneous muscular index (CMI) varies between 3 and 7 % (Table 2). The parenchymatic musculature is composed by ill-defined supra-intestinal and subintestinal transverse layers and dorso-ventral fibres located between the intestinal branches (Fig. 4). The thickness of the parenchymatic musculature (PMI) represents 1.5–2 % of the body height (Table 2).

Digestive system
The pharynx (1.5–5.1 mm in length) is cylindrical, with the dorsal insertion posteriorly displaced (0.5–1.1 mm) (Fig. 5a–d). The mouth is located in the middle of the pharyngeal pouch (2.1–5.7 mm in length). The epithelial lining of the outer surface of the pharynx is cuboidal and ciliated. The outer pharyngeal musculature is arranged in a thin longitudinal subepithelial layer (2.5–5 μm thick) followed by a thicker circular layer (7.5–20 μm thick). The pharyngeal lumen is lined with a columnar and ciliated epithelium. The inner pharyngeal musculature consists of a thick layer of circular fibres (85–200 μm thick) and a subjacent longitudinal thin layer (5–10 μm thick). The pharynx receives secretion from abundant glands, the cell bodies of which are located both laterally.
and anteriorly to the pharynx, and their cell necks extend
to the pharyngeal epithelium. Four types of pharyngeal
glands occur: abundant glands with (1) erythrophil, fine
granules and (2) xanthophil amorphous secretion, as well
as less numerous glands with (3) erythrophil and (4)
cyanophil amorphous secretion. The oesophagus (150
– 350 μm in length) is lined with a columnar and ciliated
epithelium. Its subjacent musculature (40–125 μm thick)
is continuous with the internal pharyngeal musculature,
but it is thinner than the latter. The oesophaguspharynx
ratio varies from 5 to 10 %.

Male reproductive system
The testes are arranged in two or three irregular rows
on each side of the body, situated dorsal to the intestinal
branches and just below the supra-intestinal transverse
parenchymatic layer (Figs. 3d and 4). They begin before
the ovaries and extend to the pre-pharyngeal region,
slightly before the pharyngeal root. They are located at a
distance between 10–17 % and 47–54 % of the body
length from the anterior end (Table 3). Sperm ducts run
among the fibres of subintestinal parenchymatic muscle
layer, located slightly dorsal, medially displaced, to ovovi-
telline ducts (Fig. 4). Their distal portions are expanded
with their lumen full of spermatozoa. They bend slightly
toward the dorsum and the sagittal plane, going through
the common muscle coat, and open, close to each other,
into the proximal portion of the prostatic vesicle (Fig. 6).
The intrabulbar prostatic vesicle is composed by a tubu-
lar and sinuous proximal portion (~1 mm in length on
average) followed by a globose part with narrow lumen
(Figs. 6 and 7a). The prostatic vesicle broadly commu-
nicates with the male atrium. The ejaculatory duct is
absent. The male atrium possesses richly folded walls,
and thus, the communication with the female atrium
occurs through a narrow lumen (Figs. 6 and 7a). The
male atrium is longer than the female atrium (1.8 times
on average).

Sperm ducts are lined with a ciliated squamous epithe-
lium, and they are enveloped by a thin circular muscle
layer (2.5 μm thick). The prostatic vesicle, both the sinuous
portion as the globose one, is lined with a ciliated column-
nar epithelium traversed by erythrophil fine and coarse
granular secretion and scarce erythrophil amorphous
secretion arising from cell bodies located in the vicinity
of the prostatic vesicle and in the surrounding parenchyma.
These glandular secretions are more abundant and
strongly stained in the globose portion (Fig. 7a–c). A
thin muscular layer composed by interwoven circular,
longitudinal and oblique fibres (5–15 μm thick) sur-
round the prostatic vesicle. The male atrium is lined
with a columnar epithelium, taller in the distal portion
of the male atrium than in its proximal portion. The
epithelium of the male atrium is only ciliated in its
proximal portion, in the transition with the prostatic
vesicle (Fig. 6). Its muscularis is composed by a thin
subepithelial circular layer (5–20 μm thick) followed
by a thicker longitudinal subjacent layer (15–40 μm
thick) (Fig. 7d); it is thicker distally than proximally.
The proximal part of the male atrium receives granu-
lar secretion from abundant cyanophil glands and less
abundant erythrophil glands. The distal part receives
openings from erythrophil glands with granular secre-
tion and less abundant cyanophil glands with amorph-
ous and coarse granular secretions. The cell bodies of
these glands are located internally to the common muscle
coret. The lumen of prostatic vesicle and male atrium of
some specimens present spermatozoa. Around the male
organs, the common muscular coat is mainly composed
by longitudinal fibres, with some circular and oblique
fibres, thicker dorsally (30–40 μm thick) than ventrally
(20 μm thick).
Female reproductive system

The ovaries are located at a distance of 15–22 % of the body length from the anterior end (Table 3). They are ventral, situated just below the intestine, and ovoid in shape (Fig. 7e). The ovovitelline ducts emerge dorsally from the middle of the ovaries and pass along their dorsal side. The proximal portions of the ovovitelline ducts are full of spermatozoa (Fig. 7e). The ovovitelline ducts run backward, and at the level of the proximal portion of the female atrium, they ascend almost vertically, bending a short track toward the sagittal plane to join above the female genital canal and form the common glandular ovovitelline duct (Fig. 6). The common glandular ovovitelline duct is short (100–200 μm in length) which, together with the female genital canal (100–300 μm in length), constitutes a “C” (Fig. 7f). The proximal portion of the female atrium is tubular, and its lumen progressively widens to form a cavity with scarcely folded walls (Figs. 6 and 7a, f).

Fig. 3 Transverse sections of Notogynaphallia nawei sp. nov. a Detail of the cephalic region (paratype 6). b, c Anterior tip (paratype 1). d, e Pre-pharyngeal region, in dorsal (d) and ventral (e) view (paratype 4). The arrow indicates the dorsal ciliated epidermis and the head arrows indicate rhabditogen secretions.
The ovovitelline ducts are lined with a ciliated cuboidal epithelium, and they are enveloped by a circular muscle layer (2.5–5 μm thick). The distal ascending portions of the ovovitelline ducts receive scarce secretion of the shell glands. The lining epithelium of the common glandular ovovitelline duct is ciliated and columnar, receiving abundant secretion of the shell glands (Figs. 6 and 7f). The musculature of the common glandular ovovitelline duct is formed by intermingled circular and longitudinal fibres (5–15 μm thick). The female genital canal is lined with a ciliated columnar epithelium followed by a subjacent musculature composed by intermingled circular and longitudinal fibres (10–30 μm thick). The epithelium of the female genital canal receives erythrophil granular secretion and cyanophil amorphous secretion from glands located below the musculature. The female atrium is lined with a columnar epithelium, and the subjacent muscularis is composed by two layers, a circular subepithelial (10–25 μm thick) and a longitudinal subjacent (15–40 μm thick). Abundant erythrophil granular secretion and scarce cyanophil amorphous secretion, from glands situated outward of its muscularis, open into the female atrium. Spermatozoa were observed in the lumen of the female genital canal and female atrium of the holotype (Fig. 7f) and in other specimens. Around the female organs, the common muscular coat is composed by intermingled fibres of longitudinal, circular and oblique orientation, thicker dorsally (30–40 μm thick) than ventrally (15–20 μm thick).

Vitellaria are well developed and highly abundant along the body. In the anterior and pre-pharyngeal regions, vitellaria occupy the spaces between intestinal branches (Figs. 3d, e, 4 and 7e).

Some comments about specimen fixation
It is widely well known that a rapid fixation is necessary to preserve tissues and maintain its morphological features, minimizing the enzymatic destruction of cellular and extracellular molecules by autolysis processes and microorganisms (Bancroft and Gamble 2008). The fixatives sometimes produce some artefacts in the appearance of tissues. However, in the case of flatworms, another source of artefacts arises from the methodology used to kill them. Over the years, the use of boiling water as a method to rapidly killing land flatworms was established to minimize the contortion and rupture of these animals (Lincon and Sheals 1979; Winsor 1991). This procedure should be done when the specimen is creeping, with an extended body. As quickly as possible, the water should be replaced by the fixative (usually 10 % formaldehyde), trying not to touch the specimen due to its stickiness. If this protocol is not properly followed (e.g. performing the fixation directly in aqueous formaldehyde), the animals sometimes do not die instantly, suffering contractions and distortions of the

| Table 2 Measurements of the thickness of the musculature of Notogynaphallia nawei sp. nov. |
|-----------------------------------------------|----------------|----------------|----------------|--------|
| Holotype | Paratypes | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
| CM dorsal | | | | | | | | | |
| Circular | 5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 |
| Diagonal | 10 | 10 | 10 | 5 | 7.5 | 5 | 5 | 5 | 5 |
| Longitudinal | 25 | 30 | 40 | 17.5 | 30 | 25 | 15 | 25 | 25 |
| CM ventral | | | | | | | | | |
| Circular | 5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 | 2.5 |
| Diagonal | 10 | 10 | 10 | 7.5 | 7.5 | 5 | 5 | 15 | 10 |
| Longitudinal | 25 | 35 | 35 | 17.5 | 37.5 | 25 | 25 | 50 | 25 |
| CMI (%) | 6 | 7 | 5 | 4 | 7 | 3 | 5 | 4 | 5 |
| PM supra-intestinal | 10 | 10 | 10 | 10 | 15 | 15 | 10 | 25 | 10 |
| PM subintestinal | 10 | 15 | 25 | 15 | 15 | 10 | 30 | 10 | |
| PMI (%) | 2 | 2 | 2 | 2 | 2 | 1.5 | 2 | 2 | 1.5 |

CM cutaneous musculature (μm), PM parenchymatic musculature (μm), CMI cutaneous muscular index, PMI parenchymatic muscular index

Fig 4 Schematic drawing of a transversal section of the pre-pharyngeal region of Notogynaphallia nawei sp. nov. (holotype). For simplicity, only the cutaneous longitudinal muscle layer was drawn.
body (commonly with the pharynx and the penis papilla protruded) making it difficult its study or causing a mis-
interpretation of the anatomy.

The holotype of *N. nawei* sp. nov. shows no signs of 
contraction or distortion because it was properly fixed. 
The paratypes show artefacts since the method of rapidly 
killing them was not followed properly. Some comments 
mainly about variations in the morphology of the copula-
tory apparatus (compared to the holotype) could be helpful 
for comparison with other species that exhibit similar 
artefacts.

In paratype 1, the distal portion of the sperm ducts 
has been contorted so that they ascend and open dor-
sally into the prostatic vesicle (Fig. 8a). With slight varia-
tions, in all specimens, the tubular portion of the 
prostatic vesicle maintains similar appearance to the 
holotype, but the distal globose portion shows alter-
ations in its form, with the exception of paratype 3 
(Fig. 8c). In paratype 1, the distal portion of the prostatic 
vesicle is poorly differentiated and somewhat contracted 
(Fig. 8a); in paratypes 2, 6 and 8, there are signs of con-
traction, and it is higher than longer (Fig. 8b, e and f), 
while in the paratype 5 it is longer than higher (Fig. 8d). 
However, all specimens show the same histological 
features observed in the holotype, distinguishing both 
regions of the prostatic vesicle according to their 
staining pattern with the globose portion more strongly 
stained than the tubular portion (Fig. 7a–c). The male 
atrium of paratypes shows contraction, evidenced by 
tighter folds than the holotype, resulting in a male 
aviulum with narrower lumen, remarkably evident in 
paratype 1 (Fig. 8a).

Regarding the female atrium, some alterations were 
observed in comparison with the holotype. In paratypes 
1, 3, 5 and 8, the atrium bent toward the dorsum and 
adopted an oblique position, while in paratypes 2 and 6 
the female atrium suffered contraction, especially in the 
latter (Fig. 8).

Fig. 5 Sagittal sections of the pharynx of *Notogynaphallia nawei* sp. nov. a Holotype. b Paratype 2. c Paratype 3. d Paratype 5. The head arrow 
indicates the outer pharyngeal musculature and the arrow indicates the inner pharyngeal musculature.
Additional evidence of artefacts was observed in the pharynx. Although in the holotype there is no evidence of contraction or distortion (Fig. 5a), the pharynx of the remaining specimens suffered some contraction (Fig. 5b, c). In paratype 5, the pharynx was slightly protruded and contracted, with a distortion of the pharyngeal pouch (Fig. 5d).

Discussion
General comments on the genus Notogynaphallia and on its emended diagnosis
After a phylogenetic analysis of the subfamily Geoplaninae based on molecular data, Carbayo et al. (2013) proposed to restrict the genus to six species of the group originally proposed by Ogren and Kawakatsu (1990), i.e., Notogynaphallia sexstriata (Graff, 1899), Notogynaphallia modesta (Graff, 1899), Notogynaphallia mourei (C.G. Froehlich, 1956) Notogynaphallia parca (E.M. Froehlich, 1955), Notogynaphallia plumbea (C.G. Froehlich, 1956) and Notogynaphallia froehlichae Ogren & Kawakatsu, 1990. Carbayo et al. (2013) also proposed the inclusion of Geoplanis biseminalis Riester, 1938 into this group. Four other species were considered to be morphologically not well known (Notogynaphallia incertae sedis), namely Notogynaphallia atra (Schultze & Müller, 1857), Notogynaphallia garua (Du Bois-Reymond Marcus, 1951), Notogynaphallia quinquestrata (Hyman, 1962) and Notogynaphallia andina (Hyman, 1962). From the seven remaining species in Notogynaphallia, three were analysed by Carbayo et al. (2013)

Table 3 Measurements of the reproductive system of Notogynaphallia nawei sp. nov.

<table>
<thead>
<tr>
<th>Holotype</th>
<th>1</th>
<th>2</th>
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<tr>
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<td>0.8</td>
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<td>0.15–0.25</td>
<td>0.4–0.9</td>
<td>0.4–0.5</td>
<td>0.5–0.7</td>
<td>0.7–0.4</td>
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<td>(15%)</td>
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<td></td>
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<tr>
<td>LCGD</td>
<td>0.15</td>
<td>0.15</td>
<td>0.2</td>
<td>0.125</td>
<td>0.125</td>
<td>0.2</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>LFG</td>
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<td>0.15</td>
<td>0.2</td>
<td>0.2</td>
<td>0.25</td>
<td>0.3</td>
<td>0.1</td>
<td>0.15</td>
</tr>
<tr>
<td>LFA</td>
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<td>0.8</td>
<td>0.9</td>
<td>0.65</td>
<td>0.8</td>
<td>0.65</td>
<td>0.6</td>
<td>0.8</td>
</tr>
</tbody>
</table>

The numbers given in parentheses represent the position relative to body length.
LPV = height of prostatic vesicle, LCGD = length of common glandular ovovitelline duct, LFA = length of male atrium, LFG = length of female genital canal, LMA = length of male atrium, LPV = length of prostatic vesicle, T:BH = ratio of the height of testes to the height of the body.

Fig. 6 Sagittal schematic reconstruction of the copulatory apparatus of Notogynaphallia nawei sp. nov. (holotype)
in the molecular phylogeny of Geoplaninae. Although it is not the aim of the present paper to make a revision of Notogynaphallia, our comparison of the new species with the other species of the genus indicated the need of some comments on the whole genus and a proposal for altering its generic diagnosis.

Regarding the external morphology, there are species of Notogynaphallia with four to six stripes over the dorsum,
such as *N. modesta*, *N. sexstriata* and *N. froehlichae* (Graff 1899; Riester 1938), or with two broad lateral bands over the dorsum (*N. mourei* and *N. parca*) (Froehlich 1955b; Froehlich, 1956). *N. modesta* shows a brown pigmented dorsum with two darker medial as well as two lateral stripes (Graff 1899), whereas *N. sexstriata* and *N. froehlichae* have dark stripes over a yellowish ground. The type species, *N. plumbea*, has a homogeneous dorsal pattern (Froehlich 1956). From *Noto- gynaphallia biseminalis*, known just by a body fragment at the level of the copulatory apparatus of a single specimen (Riester 1938), there is no information about its general external morphology.

In relation to eye pattern, most species of *Notogynaphallia* have exclusively marginal eyes, but *N. modesta* and *N. froehlichae* have eyes spreading over the dorsum (Riester 1938; Carbayo 2003). Thus, we propose to modify the generic diagnosis to include the variation in eye pattern in species of the group.

Regarding pharyngeal morphology, most species of the genus present a cylindrical pharynx, but *N. modesta* has a cylindrical to bell-shaped pharynx (Carbayo 2003). In addition, at least *N. parca* has a cylindrical pharynx with a dorsal insertion shifted posteriorly. In fact, there is just a slight difference between a cylindrical pharynx with a dorsal insertion shifted posteriorly and a bell-shaped pharynx. Therefore, in view of these arguments, we propose an alteration to the generic diagnosis to consider this variation in the pharyngeal morphology.

In relation to the copulatory apparatus, all species of *Notogynaphallia* share a compact copulatory apparatus with a highly folded male atrium, which probably forms a temporary penis papilla during copulation. Such an eversible penis is characterized by strong atrial musculature (Winsor 1998). Also, the prostatic vesicle, exclusively intrabulbar, opens broadly into the male atrium, without an ejaculatory duct. Furthermore, in species of *Notogynaphallia*, the ascending portion of the female ducts is lateral to the posterior region of the female atrium and joins each other behind it. The genital canal, arising from the posterior region of the female atrium, is dorso-anteriorly flexed (Carbayo et al. 2013). Among species of the genus, some
variation is observed in the prostatic vesicle, which can be globose (N. mourei, N. parca, N. plumbea and N. sexstriata) or tubular (N. biseminalis, N. froehlichae and N. modesta), and may have two different regions, as occurs in N. modesta (Carbayo 2003). In addition, adenodactyls or glandulo-muscular organs have not been observed according to the original descriptions (Graff 1899; Riester 1938; Froehlich 1955b; Froehlich 1956).

From the four species which were considered as Notogynaphallia incertae sedis, N. atra has an almost homogeneous coloured dorsum (Froehlich 1956), whereas N. garua, N. andina and N. quinquestriata have a striped dorsal surface (Du Bois-Reymond Marcus 1951; Hyman 1962). Eyes spread over the dorsal surface in N. atra, N. garua and N. andina. The pharynx is cylindrical in N. garua, N. quinquestriata and N. andina (Du Bois-Reymond Marcus 1951; Hyman 1962; Carbayo 2003) and seems to vary from cylindrical to bell-shaped in N. atra (Froehlich 1956; Carbayo 2003). Two of these species, N. atra and N. andina, have some peculiarities in their genital anatomy in relation to other species of the genus, as already commented by Froehlich and Leal-Zanchet (2003). The genital anatomy is unknown in N. quinquestriata, since the single studied specimen was in an initial stage of maturation (Hyman 1962; Carbayo 2003). N. garua has a general genital anatomy similar to that of other species of the genus, but their type specimens were not fully mature (Du Bois-Reymond Marcus 1951; Carbayo 2003).

It should be noted that other additional characteristics need to be considered for inclusion in the diagnosis of the genus, namely the cutaneous and parenchymatic musculatures in the anterior and pre-pharyngeal body regions. These musculatures provide diagnostic characters at the genus level (Froehlich 1955a; Carbayo and Leal-Zanchet 2003); thus, it is justified to make some comments. According to Carbayo (2003), the type species of the genus, N. plumbea, together with N. froehlichae, N. modesta, N. mourei, N. parca and N. sexstriata have the usual tripartite cutaneous musculature of Geoplaninae in the pre-pharyngeal region. The arrangement of the cutaneous muscle layers remains the same toward the anterior region, without cephalic specializations, and the cutaneous musculature is not sunk into the parenchyma in the species that constitute the genus Notogynaphallia sensu Carbayo et al. (2013). In N. froehlichae, however, the cutaneous musculature of the anterior body region is unknown because the holotype is damaged. Carbayo (2003) also indicated for N. modesta, N. mourei, N. parca, N. plumbea and N. sexstriata that the parenchymatic musculature is composed of three main layers: a dorsal diagonal with decussate fibres and two layers with transverse fibres, namely a supra-intestinal and a subintestinal.

In N. froehlichae, only the dorsal diagonal and suprintestinal parenchymatic muscle layers have been observed. Parenchymatic longitudinal fibres are absent in species of Notogynaphallia sensu Carbayo et al. (2013) with the exception of N. sexstriata, in which loose longitudinal fibres intermingled with the subintestinal parenchymatic layer were observed (Carbayo 2003). For N. biseminalis, the arrangement of both the cutaneous and parenchymatic muscle layers is unknown (Riester 1938). Thus, the diverse characteristics which were observed in the parenchymal musculatures of N. froehlichae and N. sexstriata should be profoundly addressed, as well as the description of the cutaneous and parenchymatic musculatures of N. biseminalis. These actions should be taken in the light of a phylogenetic analysis, improving the definition of the genus.

Comparative discussion of N. nawei sp. nov. in the genus Notogynaphallia

Specimens of N. nawei sp. nov. have a medium-sized, elongated body with parallel margins, similarly to other species of the genus. The new species shows eyes spreading over the dorsal surface, similarly to N. froehlichae and N. modesta. Regarding pharyngeal anatomy, N. nawei is similar to most species of the genus, presenting a cylindrical pharynx. Considering that these features of the external morphology and pharyngeal anatomy can also occur in other genera of Geoplaninae, we found in its genital anatomy the main reasons to propose the inclusion of the new species into the genus Notogynaphallia, namely an intrabulbar prostatic vesicle broadly communicating with the folded male atrium, an eversible penis, female ducts lateral to the posterior region of the female atrium and joining each other behind it and a dorso-anteriorly flexed female genital canal, arising from the posterior region of the female atrium. In relation to the colour pattern, N. nawei sp. nov., with a homogeneous and dark dorsal surface, can be distinguished from the majority of its congeners (N. froehlichae, N. modesta, N. mourei, N. parca and N. sexstriata) and resembles N. plumbea. Notogynaphallia plumbea exhibits a dark grey to dark brown pigmentation on the back and greyish ventral surface. However, no clear halos occur surrounding the eyes, which are situated only along body margins (Froehlich 1956; Carbayo 2003). In the new species, the eyes extend toward the dorsal surface and present small clear halos that are better distinguished in live specimens. In addition, regarding genital anatomy, the prostatic vesicle is ovoid in N. plumbea (Froehlich 1956; Carbayo 2003), whereas in N. nawei sp. nov. it has a proximal tubular portion, absent in N. plumbea.

Although the external aspect of N. modesta differs from that of the new species, the copulatory apparatus of N. nawei sp. nov. resembles N. modesta since in both
species the sperm ducts pass through the common muscular coat and immediately open into an intrabulbar prostatic vesicle, which anatomically and histologically is differentiated in two portions: tubular and sinuous proximally and globose with folded walls distally. However, in *N. modesta*, sperm ducts join each other at the point where they traverse the common muscle coat, thus being a single opening in the prostatic vesicle, while in the new species sperm ducts open into the proximal portion of the prostatic vesicle close to each other, but without forming a common portion. In both species, the male atrium is richly folded, but in *N. modesta*, the folds are almost restricted to the ventral wall, while in *N. nawei* sp. nov. they occur both dorsally and ventrally. Both species also share the morphology of the common glandular ovovitelline duct and the female genital canal, being C-shaped.

Regarding the species of *Notogynaphallia* incertae sedis, *N. nawei* sp. nov. can be distinguished from the striped specimens of *N. andina*, *N. quinquestrata* and *N. garua*. The dorsal pattern of *N. nawei* sp. nov. is similar to that of *N. atra*, which has a black pigmentation on the back and greyish ventral surface. However, *N. atra* has a bell-shaped pharynx, according to Carbayo (2003), and the male reproductive system presents an extrabulbar, ovoid and vertically disposed prostatic vesicle (Froehlich 1956; Carbayo 2003) and testes extending behind the pharynx (Carbayo 2003). Thus, *N. nawei* sp. nov., having a cylindrical pharynx, an intrabulbar and tubular prostatic vesicle and pre-pharyngeal testes, can be easily distinguished from *N. atra*.

In summary, *N. nawei* sp. nov. can be differentiated from other species of *Notogynaphallia* by the following combination of characters: homogeneous coloured dorsum, eyes spreading over the dorsal surface of the body, intrabulbar prostatic vesicle with two distinct portions and common glandular ovovitelline duct and female genital canal constituting a “C”.

**Conclusions**

Considering the known distribution of the genus *Notogynaphallia* mainly in southern Brazil, besides the occurrence of species of *Notogynaphallia* incertae sedis in Colombia, Panama and Peru, the finding of a new species of *Notogynaphallia* for the first time in the Chacoan province in Argentina extends the distribution range of the genus. In addition, the present work increases the knowledge of the morphology of the genus, proposing an emendation to its diagnosis.

**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<td>cm</td>
<td>common muscular coat;</td>
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</table>

**References**


Froehlich CG (1957) Notogynaphallia and common glandular ovovitelline duct and female genital canal constituting a “C”.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

LN sampled part of the specimens, made part of the histological preparations, performed morphological analyses and figures and drafted the manuscript. FB sampled part of the specimens, conceived the study, participated in its design and helped to draft the manuscript. AMLZ participated in the design of the study and helped to draft the manuscript. All authors contributed to the specific identification of the new taxon, as well as read and approved the final manuscript.

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