Sudhausia aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp. (Nematoda: Diplogastridae): viviparous new species with precocious gonad development

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Summary – Two new species of diplogastrid nematodes, Sudhausia aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp., were isolated from dung beetles (Coleoptera: Scarabaeidae) in Ghana and South Africa, respectively. Sudhausia n. gen. is circumscribed by stomatal morphology, namely the presence of a tube-shaped gymnostom, a pair of conical subventral denticles in the metastegostom, and a disk-like telostegostom bearing minute conical denticles. Other characters distinguishing the new genus are the presence of a bursa in the male and a vulva lined anteriorly and posteriorly by columns of sac-like cells. The two new species, both of which are hermaphroditic, are distinguished from each other by spicule and gubernaculum morphology, male papillae arrangement, bursal shape, vaginal morphology, and phasmid position. A suite of unusual developmental traits that distinguishes Sudhausia n. spp. includes maturation of the gonad and development of juvenile progeny before molting to adulthood, a two-fold increase in embryo size during development, and constitutive vivipary. A phylogeny inferred from 11 ribosomal protein-coding genes and a fragment of the small subunit rRNA gene show Sudhausia n. gen. to be divergent from other sequenced diplogastrid taxa, including those characterised by a tube-like stoma. The two new species represent useful new reference points for the study of feeding-structure evolution in Diplogastridae.

Keywords – androdioecy, description, dung beetles, Ghana, hermaphrodite, molecular, morphology, morphometrics, new genus, new species, Onthophagus, phylogeny, Scarabaeidae, Scarabaeus, South Africa, stoma, taxonomy.

The nematode family Diplogastridae Micoletzky, 1922 currently includes 31 genera (Sudhaus & Fürst von Lieven, 2003; Kanzaki et al., 2009, 2012a; Mayer et al., 2009; Fürst von Lieven et al., 2011; Susoy & Herrmann, 2012), many of which are associated with insects. Consequently, intensive sampling of potential insect hosts has recently revealed diverse new taxa in the family (Kanzaki et al., 2009, 2012a). Scarab beetles (Coleoptera: Scarabaeidae) in particular are hosts to many diplogastrid species, especially in Pristionchus Kreis, 1932 (Herrmann et al., 2006; Mayer et al., 2007, 2009; Rae et al., 2008; Kanzaki et al., 2012a, 2013), but also in Diplogasteroides de Man, 1912, Koerneria Meyl, 1960, Mononchoides Rahm, 1928, Rhabditoides Rahm, 1928, and Tylopharynx de Man, 1876 (Mayer et al., 2009).

Given the diversity of feeding structures in diplogastrids (Fürst von Lieven & Sudhaus, 2000; Sudhaus & Fürst von Lieven, 2003), the isolation of new higher order taxa in the family is particularly valuable for studies of the evolution of form. In addition to the presence of detailed references for diplogastrid stomatal anatomy (Baldwin et al., 1997; Bumbarger et al., unpubl.), the established model system Pristionchus pacificus Sommer, Carta, Kim & Sternberg, 1996 allows genetic and developmental mechanistic studies of morphology (Bento et al., 2010; Sommer & McGaughran, 2013). The potential to extend studies in Pristionchus to other species of diplogastrids, especially those amenable to laboratory culture, is made more powerful by the discovery of new forms to include in a comparative context. Here, we describe a
new genus of Diplogastridae, Sudhausia n. gen., based on its unique stomatal morphology, sex-specific characters, and a suite of unusual developmental traits. Reflecting the frequent association of diplogastrids with scarab beetles, both of the new species, S. aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp., were recorded from dung beetles in Ghana and South Africa, respectively.

**Materials and methods**

**Nematode isolation and cultivation**

*Sudhausia aristotokia* n. gen., n. sp. was isolated from adult dung beetles as detailed for individual species below. Host beetles were dissected on a 2.0% agar plate, after which the plate was kept at room temperature for several weeks. Nematodes proliferated on bacteria associated with the host beetle cadavers. Individuals were thereafter transferred to nematode growth medium (NGM) agar plates seeded with the original microbes from the beetle carcass, and they have been since kept in laboratory culture on this medium.

**Morphological observation and preparation of type material**

One- to two-week-old cultures of *S. aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. provided material for morphological observation. Observations by light microscopy (LM) and differential interference contrast (DIC) microscopy were conducted using live nematodes, which were hand-picked from culture plates. Because of the clarity and integrity of its morphology, live material was used for morphometrics. For line drawings, specimens were mounted into water on slides and then relaxed by applying gentle heat. For morphometrics, specimens were mounted on slides with pads of 5% noble agar and 0.15% sodium azide and were additionally relaxed by heat when necessary. To prepare type material, nematodes were isolated from type strain cultures, rinsed in distilled water to remove bacteria, heat-killed at 65°C, fixed in TAF to a final concentration of 5% formalin and 1.5% triethanolamine, and processed through a glycerin and ethanol series using Seinhorst’s method (see Hooper, 1986). Alternatively, some paratypes were fixed in 5% formalin after heat killing at 65°C. Nomarski micrographs were taken using a Zeiss Axio Imager Z.1 microscope and a Spot RT-SE camera supported by the program MetaMorph v.7.1.3 (Molecular Devices).

**Scanning electron microscopy**

Nematodes were prepared for scanning electron microscopy (SEM) by fixation in 2.5% glutaraldehyde in M9 buffer and then post-fixed with 1% osmium tetroxide. After several rinses with water, samples were dehydrated through a graded ethanol series, followed by critical point drying using carbon dioxide. Specimens were mounted on polylysine-coated cover slips, sputter-coated with 20 nm gold/palladium, and then imaged with a Hitachi S-800 field emission scanning electron microscope operating at 20 kV.

**Molecular characterisation and phylogeny**

For species diagnosis and phylogenetic analysis, we amplified and sequenced an approximately 500-bp fragment of the small subunit (SSU) rRNA gene of the two new species using the primers SSU18A (5′-AAAGATTA AGCCATGCATG-3′) and SSU9R (5′-AGCTGGAATTA CCGCGGCTG-3′) (Floyd et al., 2002). Partial SSU rRNA sequences of *S. aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. have been deposited in the GenBank database under accession numbers KC831555 and KC831556, respectively. Additionally, we amplified and sequenced 11 ribosomal protein-coding genes for the two new species: *rpl-2*, *rpl-6*, *rpl-9*, *rpl-10*, *rpl-14*, *rpl-23*, *rps-7*, *rps-14*, *rps-27* and *rps-28*. All information regarding genes, primers and PCR conditions is given in Mayer et al. (2009). Except for *rps-28*, the sequence of which was less than 200 bp, all ribosomal protein gene sequences for *S. aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. have been deposited in GenBank under accession numbers KC831557-KC831566 and KC831567-KC831576, respectively. Phylogenetic analysis included species from all diplogastrid genera for which homologous sequences have been published (Mayer et al., 2009; Kanzaki et al., 2012a).

The concatenated dataset of ribosomal protein genes and the partial SSU rRNA gene was aligned using MUSCLE (Edgar, 2004), followed by manual alignment in MEGA5.10 (Tamura et al., 2011), which included deletion of ambiguous alignment positions. The final alignment comprised 5963 positions. The alignment was partitioned into four subsets: one for the partial SSU rRNA gene and three according to codon position for the concatenated set of ribosomal protein genes. Analyses by maximum likelihood (ML) and Bayesian inference, as implemented in RAxML v.7.2.8 (Stamatakis, 2006) and MrBayes 3.2 (Ronquist et al., 2012), respectively,
were used to infer the phylogeny. Fifty independent runs were performed for the ML analysis, in which trees were inferred under a general time reversible model with a gamma-shaped distribution of rates across sites. Bootstrap support was calculated by 1000 pseudoreplicates on the most likely tree among all runs. Bayesian analyses were initiated with random starting trees and were run with four chains for $4 \times 10^6$ generations. Markov chains were sampled at intervals of 100 generations. Two independent runs were performed for the analysis. After confirming convergence of runs and discarding the first $2 \times 10^6$ generations as burn-in, remaining topologies were used to generate a 50% majority-rule consensus tree with clade credibility values given as posterior probabilities (PP). Bayesian analysis invoked a mixed model of substitution with a gamma-shaped distribution across sites and, as informed by previously studies (Mayer et al., 2009; van Megen et al., 2009), specified Rhabditoides inermis (Schneider, 1866) Dougherty, 1955 as outgroup. Model parameters were unlinked across character partitions in both ML and Bayesian analyses.

**Results**

*Sudhausia* *n. gen.*

**DESCRIPTION**

Diplogastridae. Body cylindrical, stout. Cuticle with fine annulation, fine striations observed by LM but not appearing on surface, and one fine lateral line appearing on surface on each side, only visible by SEM. Lips fused in pairs (one dorsal, two subventral) and separated by thin rims of labial cuticle, resulting in an ypsiliform stomatal opening. Six short, papilliform labial sensilla present in both males and morphological females (only hermaphrodites are known) and four papilliform cephalic papillae present in males, as typical for diplogastrid nematodes. Labial and cephalic papillae visible by LM. Cheilostom triradiate, anterior rim of which punctuated by grooves corresponding to separation of lips. Gymnostom tube-shaped, twice as long as wide, divided into two distinct regions: i) an offset, short anterior region of smooth cuticle with a barely distinguishable midline separating an anterior rim that is slightly thicker than the posterior rim; and ii) a longer posterior region with cuticle refracting to give appearance of a rough texture. Stegostom comprising: i) a transversely flat rim of cuticle (pro-/mesostegostom) at anterior margin of muscular pharynx; ii) a cylindrical metastegostom bearing a moveable dorsal tooth, dorsoventrally narrowed for its entire length and transversely triangular, with a clear gland duct and orifice and, in each of subventral sectors, a pair of narrow, conical, equally sized and axially oriented denticles, all borne from a level slightly posterior to dorsal tooth; and iii) a disk-like telostegostom with an anterior rim and bearing minute, conical, axially oriented denticles arranged in three pairs (one dorsal, two subventral). Apodemes absent. Procorpus and metacorpus muscular. Metacorpus similar to, or slightly wider than, procorpus in diam. Isthmus narrow, not muscular. Basal bulb glandular. Pharyngo-intestinal junction clearly observed, well developed. Morphological females amphidelphic, ovotestes reflexed. Vagina surrounded anteriorly and posteriorly by columns of sac- or pouch-like cells, largest being those closest to ventral body surface. Bursa present. Male with 8-9 pairs of genital papillae, those on bursa forming rays, and one small, unpaired, ventral genital papilla on anterior cloacal lip.

**BIONOMICS**

In addition to the above morphological characters, which definitively circumscribe *Sudhausia* *n. gen.*, the following biological traits distinguish and are considered tentatively diagnostic of the genus. Species are androdioecious, consisting of males and self-fertile hermaphrodites. Somatic gonad of hermaphrodites develops precociously. Developing embryos occur inside the gonad during the early J4 stage of the mother, *i.e.*, when the vulva is beginning to invaginate. Gonads in older hermaphrodites appear disorganised due to the presence of growing embryos and hatching juveniles. Embryos lack a discernible eggshell, are irregular in shape, and grow two-fold in size from the zygote to the comma stage. Dauer juveniles are often present in the intra-uterine brood. Primary branches of gonads in mature adults are therefore filled with a brood of mixed developmental stages that together occupy most of the body volume of the living, active mother. Internal non-dauer juveniles have been seen to feed on vesicles of unknown origin and substance. Juveniles are born constitutively by vivipary, even under well-fed laboratory conditions.

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* Named in honour of Prof. Walter Sudhaus (Freie Universität Berlin, Berlin, Germany), in recognition of a career of outstandingly detailed work in rhabditid and diplogastrid morphology and systematics.
**TYPE SPECIES**

*Sudhausia aristotokia* n. gen., n. sp.

**OTHER SPECIES**

*Sudhausia crassa* n. gen., n. sp.

**RELATIONSHIPS**

*Sudhausia* n. gen. is distinguished from all other Diplogastridae by the combined presence of a tube-shaped (i.e., ‘rhabditiform’) gymnostom and a pair of conical denticles in the subventral metastegostom and in all sectors of the telostegostom (Fürst von Lieven & Sudhaus, 2000; Sudhaus & Fürst von Lieven, 2003). The new genus is therefore distinguished from all other diplogastrid genera with a tube-like gymnostom, namely *Demaniella* Steiner, 1914, *Diplogasteroides*, *Diplogastrellus* Para-monov, 1952, *Goffartia* Hirschmann, 1952, *Mehdinema* Farooqui, 1967, *Pseudodiplogasteroides* Körner, 1954, *Rhabditidoides*, and *Rhabditolaimus* Fuchs, 1914, by the presence of this armature. Also unique to *Sudhausia* n. gen. is a vagina lined anteriorly and posteriorly by columns of sac-like cells. The similar diameter of the procorpus and metacorpus distinguishes *Sudhausia* n. gen. from other diplogastrids except for *Rhabditolaimus* and *Odontopharynx* de Man, 1912, the latter being distinguished from *Sudhausia* n. gen. by its barrel-shaped and heavily cuticularised gymnostom and large, dagger-like dorsal tooth. The presence of a bursa distinguishes *Sudhausia* n. gen. from other diplogastrids except for *Heteropleuronema* Andrássy, 1970 and some species of *Neodiplogaster* Cobb, 1924 and *Rhabditolaimus*, all of which are clearly distinguished from *Sudhausia* n. gen. by stomatal and pharyngeal morphology. Finally, heterochronic gonad development and growing embryos both distinguish *Sudhausia* n. gen. from all other diplogastrid genera.

**MOLECULAR CHARACTERISATION AND PHYLOGENY**

The concatenated alignment comprised 2342 parsimony-informative sites. Phylogenetic analysis of sequences of the SSU rRNA and 11 ribosomal protein loci revealed divergence of *S. aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. from each other and from all other diplogastrid genera for which both SSU rRNA and orthologous ribosomal protein gene sequences were available (Figs 1, 2).

The diagnostic 450 bp fragment of SSU rRNA in particular differed by ten nucleotide positions, supporting their identity as two unique species.

Topologies inferred by ML (Fig. 1) and Bayesian (Fig. 2) analyses differed in several deep, poorly supported nodes, but the trees were congruent in several respects. Firstly, *S. aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. were fully supported as sister taxa (100% BS, PP). Secondly, *Sudhausia* n. gen. was shown to be a sister group to a clade comprising *Tylopharynx* sp. + *Neodiplogaster* sp. + *Mononchoides* sp., albeit with variable support (62% BS, 100% PP). Thirdly, a separate clade including *Diplogastrellus gracilis* Bütschli, 1876 + *Rhabditidoides* sp. and *Pseudodiplogasteroides* sp. + *Diplogasteroides* sp. was moderately or well supported as monophyletic (83% BS, 100% PP). Finally, *Sudhausia* n. gen. was nested within a well supported clade (100% BS, PP) that included all analysed diplogastrids other than *Kerneria* sp. Differing between the two analyses was the position of *Rhabditolaimus* sp. + *Oigolaimella attenuata* Fürst von Lieven & Sudhaus, 2008, which in both cases was inferred to have a negligibly supported relationship to their sister group (<50% BS, 65% PP). The relationship of a clade comprising *Sudhausia* n. gen., *Tylopharynx* sp., *Neodiplogaster* sp., and *Mononchoides* sp. to other lineages was also unresolved, being poorly supported and differing between analyses.

***Sudhausia aristotokia**** n. gen., n. sp.**

(Figs 3-6; 7A-C, E; 8A-C)

**MEASUREMENTS**

See Table 1.

**DESCRIPTION**

**Adults**

Labial region slightly offset, lips squared or acutely angular at junction of anterior and outer lateral walls. Amphidial apertures elliptical, located subdorsally at level of posterior cheilostom. Dorsal pharyngeal gland clearly observed, penetrating dorsal tooth to gland opening. Procorpus occupying half to two-thirds of corresponding body diam. Metacorpus similar to, or slightly wider than,

*Derived from the Greek ἄριτος (‘best’) + τόκος (‘birth’) and refers to the vivipary of this species.*
**Phylogenetic relationships of Diplogastridae, including Sudhausia aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp.**

The tree with the highest log-likelihood ($-55811.647527$) is shown. The percentage of trees in which the associated taxa clustered together in 1000 bootstrap pseudoreplicates is shown next to the nodes (left value). Bootstrap support values above 50% are shown. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site.

**Male**

Spontaneous (= fatherless) males rare in culture (<1%). Testis single, located along ventral side. Spermatogonia arranged in ca three rows in reflected part, in two to three rows in anterior half of main branch, then mature amoeboid spermatids arranged in multiple rows in remaining, proximal part of gonad. *Vas deferens* not clearly separated from other parts of gonad. Spicules paired, separate. Spicules long, slender, appearing pliable (bending inconsistently) in ventral view, adjacent to each other for distal fifth of their length. Spicule in lateral view smoothly and ventrally arcuate, giving spicule curvature of ca $135^\circ$. Manubrium squared, 1.5-2.0 times as long as wide, twice as wide as spicule blade (= lamina/calomus complex) in ventral view. Spicule blade only slightly expanded at one fourth of spicule length from anterior end. Gubernaculum half of spicule length long, slender, with medially split, knob-like anterior tip, anterior half not recurved with respect to posterior half, posterior half offset from anterior half and forming a tube enveloping spicules. Cloacal opening dome-shaped in ventral view. Three cloacal glands, two subventral and one dorsal, present but not obvious. Two pairs of genital papillae ventral (‘vn’) to alae, six pairs forming bursal rays, ending distinctly on dorsal side (‘Rnd’), ventral side (‘Rnv’), or margin (‘Rn’) of bursa (nomenclature is original, as homologies with papillae of other diplogastrids are ambiguous). Pairs of papillae...
and phasmids arranged from anterior to posterior as \( \langle R_1, R_2, R_3, (R_4d, v_1), C, v_2, (R_5d, Ph, R_6, R_7) \rangle \), whereby \( R_1 \) is far (almost one cloacal body diam.) anterior to \( R_2 \), \( R_2v-R_4d \) are close and regularly spaced, \( R_4d \) is at same level as \( v_1, C \) is equidistant from \( v_1 \) and \( v_2 \), single papillae of \( v_1 \) are as close to each other as are single papillae of \( v_2 \), and \( R_6 \) and \( R_7 \) separated as rays but emerging as papillae from a common opening or fusing at tip. Tail bluntly conical. Bursa peloderan, wide, such that alae extend 0.5 cloacal body diam. from body, terminating in a single posterior point.

**Hermaphrodite**

Each genital system arranged from vulva/vagina as uterus, oviduct, and ovotestis. In J4 or very young adults, uterus and oviduct extending ventrally and anteriorly on right of intestine, as typical for diplogastrids, but in most adults gonads expanded, possibly tangled, and filled with developing brood, such that positions are unclear. Ovotestis totally reflexed, extending dorsally. Dorsal flexure of gonad short, not usually extending to vulva, remaining short even after extension of main gonad branch during J4 and adult stages, in which oocytes are arranged in 2-3 rows, often terminating in one large oocyte just proximal to gonad flexure. At proximal end of main gonad branch, oviduct serving as spermatheca and filled with self sperm. Distal to spermatheca-like region, oviduct of young hermaphrodites composed of flattened, rectangular cells. Distal, remaining part of oviduct (= uterus) containing developing embryos from an early stage, even as early as J4 stage, before eruption of vulva and moulting to adulthood. Developing embryos and juveniles often arranged serially along body of mother. Vaginal glands present but obscure, observed in ventral view. Vagina perpendicular to body surface. Sac-like cells lining vulva anteriorly and posteriorly with thin cytoplasm, such that
Sudhausia aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp.

Type Host (Carrier) and Locality

The type strain was established from an individual nematode isolated from the body of a dung beetle of the genus *Onthophagus* Latreille, 1802 (Coleoptera: Scarabaeidae) in Borbiri, Ghana. Only one of eight beetles examined was positive for the nematode.

Type Material and Strain Culture

Holotype male, two paratype males, and six paratype hermaphrodites deposited in the University of California Riverside Nematode Collection (UCRNC), Riverside, CA, USA. Four paratype hermaphrodites deposited in the Natural History Museum Karlsruhe, Karlsruhe, Germany. The type strain is available in living culture from the Department of Evolutionary Biology, Max Planck Institute (MPI) for Developmental Biology, Tübingen, Germany, and can be provided to other researchers upon request. Attempts to achieve frozen stocks of this species have been unsuccessful.

Diagnosis and Relationships

*Sudhausia aristotokia* n. gen., n. sp. is diagnosed by the long and thin spicules with no ventral expansion of the spicule blade (*i.e.*, calomus is not offset from lamina), a straight and narrow gubernaculum, the male genital papillae arranged as \(R1, R2v, R3v, (R4d, v1), C, v2, (R5d, Ph, R6, R7)\), whereby R6 and R7 emerge from a common opening, and an offset labial region with squared or acutely angular lips.

The species is distinguished from its only known congener, *S. crassa* n. gen., n. sp., by the following characters: spicules that are long, thin with a manubrium twice as wide as blade transversely vs spicules that are robust with a manubrium only 1.5 times wide as blade transversely; a straight and narrow vs stout and recurved gubernaculum; a bursa with alae extending 0.5 cloacal body diam. vs less than 0.33 cloacal body diam. from the...
Fig. 4. *Sudhausia aristotokia* n. gen., n. sp. A: Neck region of hermaphrodite, right lateral view; B: Lip region of male, left lateral view; C: Stomatal region of hermaphrodite, right lateral view; D: Stomatal region of hermaphrodite, ventral view; E: Spicule and gubernaculum, left lateral view; F: Male tail, left lateral view; G: Male tail, ventral view. Abbreviations: R1-R7 = bursal ray papillae; d = dorsal; v = ventral.
Sudhausia aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp.

Fig. 5. Schematic representation of the stoma of Sudhausia aristotokia n. gen., n. sp. Abbreviations: cheilo = cheilostom; gymno = gymnostom; pro/meso stego = pro-/mesostegostom; meta stego = metastegostom; telostego = telostegostom.

Body; male genital papillae arranged as \((R_1, R_2, R_3, (R_4, v_1), C, v_2, (R_5, Ph, R_6, R_7)) vs (R_1, R_2, R_3, (v_1, C), v_2, (Ph, R_4) (R_5d, R_6v))\); by \(R_1\) far (almost one cloacal body diam.) anterior to \(R_2\), which is close to \(R_3\) vs \(R_1\) closer to \(R_2\) than is \(R_3\); \(C\) equidistant from \(v_1\) and \(v_2\) vs at same level as \(v_1\); single papillae of \(v_1\) as close to each other as are single papillae of \(v_2\) vs further from each other than single papillae of \(v_2\); \(R_7\) fused to or emerging from a common opening with \(R_6\) vs \(R_7\) absent; and a corpus that is twice as long vs 1.5 times as long as the postcorpus. Hermaphrodites of \(S. aristotokia\) n. gen., n. sp. are distinguished from those of \(S. crassa\) n. gen., n. sp. by an offset labial region with squared or acutely angular lips vs a labial region that is rounded and not offset. Although body shape can vary substantially with age and among individuals within species, \(S. aristotokia\) n. gen. n. sp. has a stout, vs fat, very stout, body and by a longer (4-5 anal body diam.) and narrowly conical vs shorter (3-4 anal body diam.), widely conical tail. Finally, each species is diagnosed by a unique sequence in a 450 bp fragment of SSU rRNA, which differs by ten positions between the two species.

**Sudhausia crassa** *n. gen., n. sp.*
(Figs 7D-H; 8D, F-H; 9, 10)

**Measurements**

See Table 1.

**Description**

**Adults**

Labial region not offset, lips rounded at junction of anterior and outer lateral walls. Amphidial apertures elliptical, located subdorsally at level of posterior cheilostom. Dorsal pharyngeal gland clearly observed, penetrating dorsal tooth to gland opening. Procorpus occupying half to two-thirds of corresponding body diam. Metacorpus similar in diam. to, or slightly wider than, procorpus. Corpus 1.5 times as long as postcorpus. Nerve ring surrounding anterior to mid-isthmus. Excretory pore located at level of mid-isthmus to anterior part of basal bulb. Hemizonid not clearly observed. Deirid posterior to pharyngo-intestinal valve.

**Male**

Spontaneous males rare in culture (<1%). Body stout. Testis single, located along ventral side, anterior part reflexed to either right or left side. Spermatogonia arranged in ca three rows in reflexed part, in 2-3 rows in anterior half of main branch, then mature amoeboid spermatids arranged in multiple rows in remaining, proximal part of gonad. Vas deferens not clearly separated from other parts of gonad. Spicules paired, separate. Spicules robust in ventral view, adjacent to each other for distal fifth of their

*S* Latin adjective meaning ‘fat, stout’ and referring to the general body shape of mature adults, especially hermaphrodites.
Fig. 6. Young hermaphrodites of *Sudhausia aristotokia* n. gen., n. sp. A-C: Variation in reproductive tract among individuals. A: Left lateral view; B: Right lateral view; C: Right lateral view; D: Anus, ventral view; E: Tail region, right lateral view; F: Vulva, ventral view.
length. Spicule in lateral view smoothly and ventrally ar-
cuate, giving a spicule curvature of ca 135°. Manubrium
ovoid, less than 1.5 times as long as wide, only slightly
wider than spicule blade in ventral view. Spicule blade
distinctly expanded at one-third of spicule length from an-
terior end. Gubernaculum 0.5 spicule lengths long, robust,
broad anteriorly such that dorsal wall slightly recurved
and dorsal and ventral walls separate at a 60° angle at pos-
terior end, posterior half not offset from anterior half, for-
making a tube enveloping spicules. Cloacal opening dome-
shaped in ventral view. Three cloacal glands, two subven-
tral and one dorsal, present but not obvious. Two pairs
of genital papillae ventral (vn) to alae, six pairs forming
bursal rays, ending distinctly on dorsal side (Rnd), ven-
tral side (Rnv), or margin (Rn) of bursa. Pairs of papillae
and phasmids arranged from anterior to posterior as (R1,
R2, R3d, (v1, C), v2, (Ph, R4d) (R5d, R6v)), whereby R2
is anterior and closer to R1 than to R3d, R2-R4d are close
and regularly spaced, v1 at same level as C, single papillae
of v1 further from each other than single papillae of v2,
Ph slightly anterior or at same level as R4d, and R5d and
R6v sharing same opening. Tail bluntly conical. Bursa pe-
loidenan, narrow, such that alae extend less than one-third
cloacal body diam. from body, terminating in two poste-
rior points.

Hermaphrodite

Body very stout. Each genital system arranged from
vulva/vagina as uterus, oviduct, and ovotestis. In J4 or
very young adults, uterus and oviduct extending ven-
trally and anteriorly on right of intestine, as typical for
diplogastrids, but in most adults gonads are expanded,
possibly tangled, and filled with developing brood, such
that positions of gonad arms are unclear. Ovotestis to-
tally reflexed, extending dorsally. Dorsal flexure of go-
nad short, not usually extending to vulva, remaining short
even after extension of main gonad branch during J4
and adult stages in which oocytes are arranged in 2-3
rows, often terminating in one large oocyte just proxim-
al to gonad flexure. At proximal end of main gonad
branch, oviduct serving as spermatheca and filled with
self sperm. Distal to spermatheca-like region, oviduct of
young hermaphrodites composed of flattened, rectangular
cells. Remaining distal part of oviduct containing de-
veloping embryos from an early stage, even as early as J4,
before eruption of vulva and moulting to adulthood. De-
veloping embryos and juveniles seemingly arranged haphaz-
dardly along body of mature mother. Vaginal glands present
but obscure, observed in ventral view. Vagina perpendic-
ular to body surface. Sac-like cells lining vulva anteriorly
and posteriorly with dense, refractive cytoplasm such that
sacs appear thickened. Vulva slightly to strongly protu-
berant in lateral view, pore-like in ventral view. Rectum
ca 1 anal body diam. long, intestinal-rectal junction sur-
rrounded by well developed sphincter muscle. Three rec-
tal glands, two subventral and one dorsal, present. Anus
in form of dome-shaped slit, posterior anal lip slightly
protuberant. Phasmid conspicuous, located ca one-third
anal body diam. anterior to anus. Tail thick and coni-
cal, ca 3-4 anal body diam. in length, with conical ter-
minus.

Type host (carrier) and locality

The type strain was established from an individual ne-
motode isolated from dung beetles of the tribe Canthonini
and from Onthophagus (Coleoptera: Scarabaeidae) on a
farm near Pretoria, South Africa.

Type material and strain culture

Holotype male, two paratype males, and six paratype
hermaphrodites deposited in the UCRNC, Riverside, CA,
USA. Four paratype hermaphrodites deposited in the Nat-
ural History Museum Karlsruhe, Karlsruhe, Germany.
The type strain is available in living culture from the
Department of Evolutionary Biology, MPI for Develop-
mental Biology, Tübingen, Germany and can be provided
to other researchers upon request. Attempts to achieve
frozen stocks of this species have been unsuccessful.

Other carrier and locality

In addition to its collection from the type host (carrier)
and locality, the species has been isolated from dung
beetles of Scarabaeus L. 1758 that were collected on
Corsica by Prof. H. Schultz-Key.

Diagnosis and relationships

Sudhausia crassa n. gen., n. sp. is diagnosed by a very
stout body shape, stout and robust spicules with a long
lamina (one-third of spicule length), male genital papillae
including only eight pairs and arranged as (R1, R2, R3d,
(v1, C), v2, (Ph, R4d) (R5d, R6v)), and the phasmid of the
hermaphrodite opening anterior to the anus.

It is distinguished from the type and only other species
in the genus as described thereunder.
Discussion

*Sudhausia* n. gen. is described according to a novel combination of traits. Whereas some individual characters, including a tube-shaped gymnostom, subventral stegotomosomal armature, and a developed bursa, independently circumscribe other genera of Diplogastridae, their shared presence in *Sudhausia* n. gen. clearly distinguishes the new genus from other lineages in the family. The composition of stomatal traits in particular raises questions about the evolution and possible convergence of the architecture of those structures.

Stomatal characters most clearly diagnose *Sudhausia* n. gen. from other diplogastrids. Almost all sequenced diplogastrid genera characterised by a tube-shaped gymnostom, namely *Demaniella*, *Diplogasteroides*, *Diplogastrellus*, *Pseudodiplogasteroides* and *Rhabditiloides*, putatively constitute a monophyletic clade (Mayer et al., 2009; Kiontke & Fitch, 2010; Figs 1, 2), with the possible exclusion of the highly autapomorphic genus *Rhabditolaimus*, formerly of a separate family, Cylindrocoporidae (Goodey, 1939; Andrásy, 1984). The position of *Sudhausia* n. sp. was variably supported as falling outside of this group, suggesting the possibility that a tube-shaped gymnostom has evolved independently in *Sudhausia* n. gen. and other diplogastrids. The presence of a pair of metastegomosomal denticles, presumably homologous with the subventral tooth and armature of diplogastrids with opposing, movable teeth (e.g., *Koerneria*, *Pristionchus*, *Mononchoides*), clearly sets *Sudhausia* n. gen. apart from other genera with a tube-shaped gymnostom. A phylogeny with broader taxon representation is needed to test whether the stoma of *Sudhausia* n. gen. represents an intermediate state in the loss of teeth or is due to convergent evolution of a tubular stoma. Tracking the loss of predatory feeding structures, and the presumptive advantages they entail (Kiontke & Fitch, 2010), holds important implications for the evolution of feeding strategies.

Male sexual characters are also distinct for *Sudhausia* n. gen., which is in part circumscribed by the presence of a male bursa and which shows divergent arrangements of genital papillae. A bursa has apparently arisen independently in different lineages of Diplogastridae, for example, in both *Neodiplogaster* and *Rhabditolaimus* (Figs 1, 2), where this character is even variable among congeners. In known diplogastrid species with a developed bursa, the arrangement of the nine pairs of papillae and rays is relatively conserved, such that their homologies are reliably discernible (Sudhaus & Fürst von Lieven, 2003). In *Sudhausia* n. gen., the positions of rays in particular are atypical for diplogastrids, making homologies difficult to interpret in this genus. The anterior and posterior pairs of papillae closest to the cloacal aperture in *Sudhausia* n. gen. are most likely the homologues of the ventral v2/v3 and v4 papillae *sensu* Sudhaus & Fürst von Lieven (2003), respectively. Furthermore, the most anterior ray is probably homologous with v1 in other diplogastrids. It is possible that at least ray R5v in *S. crassa* n. gen., n. sp. and rays R6 and R7 in *S. aristotokia* n. gen., n. sp. are homologous with one or two of the small ventral triplets (v5-v7), respectively, in other diplogastrids. The derived condition of only eight pairs of papillae in *S. crassa* n. gen., n. sp. is apparently due to the loss of the homologue of either R6 or R7 present in *S. aristotokia* n. gen., n. sp. The proclivity to lose one or two pairs of these papillae has also been observed in *Parasitodiplogaster* Poinar, 1979 and *Tera- todiplogaster* Kanzaki, Giblin-Davis, Davies, Ye, Center & Thomas, 2009 (Giblin-Davis et al., 2006; Kanzaki et al., 2012b). Identities of the other rays are unclear, and plasticity of rays even among species of *Sudhausia* n. gen. indicates that the rays are likely to be divergent with respect to outgroups.

Unusual developmental phenomena distinguish *Sudhausia* n. gen. from other similarly studied nematode

Fig. 7. Scanning electron micrographs of *Sudhausia aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. A: Entire body of mature hermaphrodite; B: Male tail, right subventral view. Nomenclature of genital papillae follows text. Asterisk indicates phasmid; C: Gubernaculum and thin tips of spicules extruded from cloacal opening, right subventral view. D-H: *S. crassa* n. gen., n. sp. D: Lip region of hermaphrodite, oblique *en face* and left subdorsal lateral view. Plugged amphid (am) is also apparent; E: Neck region of hermaphrodite, including excretory pore, ventral view; F: Vulva, ventral view; G: Male tail, left ventral view. Extruded, robust spicule tips also apparent; H: Anus, oblique left subventral view. Left phasmid is also apparent. Abbreviations: R1-R7 = bursal ray papillae; d = dorsal, v = ventral, V1, V2 = ventral papillae.
**Table 1.** Morphometrics of male holotypes (in glycerin) and male and hermaphrodite specimens of *Sudhausia aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. (temporary water mounts). All measurements are in μm and in the form: mean ± sd (range).

<table>
<thead>
<tr>
<th>Character</th>
<th><em>S. aristotokia</em> n. gen., n. sp.</th>
<th><em>S. crassa</em> n. gen., n. sp.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male Temporary water mounts</td>
<td>Hermaphrodite Temporary water mounts</td>
</tr>
<tr>
<td>n</td>
<td>– 5</td>
<td>10</td>
</tr>
<tr>
<td>L</td>
<td>479 ± 61 (412-573)</td>
<td>917 ± 117 (680-1086)</td>
</tr>
<tr>
<td>L′</td>
<td>461 ± 49 (390-519)</td>
<td>827 ± 111 (604-989)</td>
</tr>
<tr>
<td>a</td>
<td>11 ± 1.1 (10-13)</td>
<td>13 ± 1.6 (10-15)</td>
</tr>
<tr>
<td>a′</td>
<td>10.5 ± 1.0 (9.2-12)</td>
<td>12 ± 1.4 (9-14)</td>
</tr>
<tr>
<td>b</td>
<td>3.8 ± 0.4 (3.5-4.3)</td>
<td>6.3 ± 0.6 (5.2-7.1)</td>
</tr>
<tr>
<td>c</td>
<td>15 ± 3.3 (10.6-19)</td>
<td>10 ± 0.9 (8.9-12)</td>
</tr>
<tr>
<td>c′</td>
<td>3.5 ± 0.5 (2.4-4.4)</td>
<td>1.8 ± 0.7 (1.1-3.0)</td>
</tr>
<tr>
<td>V</td>
<td>– 58 ± 4.0 (51-63)</td>
<td>– 58 ± 4.0 (51-63)</td>
</tr>
<tr>
<td>Max. body diam.</td>
<td>32 ± 4.9 (38-50)</td>
<td>72 ± 15 (48-99)</td>
</tr>
<tr>
<td>Lip region diam.</td>
<td>8.0 ± 1.0 (7.8-10)</td>
<td>12 ± 0.7 (11-13)</td>
</tr>
<tr>
<td>Stoma width</td>
<td>3.3 ± 0.7 (3.0-4.8)</td>
<td>4.7 ± 0.5 (3.7-5.4)</td>
</tr>
<tr>
<td>Stoma height</td>
<td>13 ± 0.8 (12-14)</td>
<td>15 ± 0.9 (14-17)</td>
</tr>
<tr>
<td>Corpus length</td>
<td>64 ± 5.2 (75-89)</td>
<td>97 ± 4.5 (88-102)</td>
</tr>
<tr>
<td>Corpus/pharynx ratio</td>
<td>60 ± 3.3 (63-71)</td>
<td>67 ± 2.7 (62-72)</td>
</tr>
<tr>
<td>Median bulb diam.</td>
<td>12 ± 0.9 (10-13)</td>
<td>15 ± 1.5 (13-18)</td>
</tr>
</tbody>
</table>

**Fig. 8.** Nomarski micrographs of *Sudhausia aristotokia* n. gen., n. sp. and *S. crassa* n. gen., n. sp. A: Stomatal region of hermaphrodite of *S. aristotokia* n. gen., n. sp., sagittal plane, left ventral view. Short arrow indicates offset anterior part of gymnostom; asterisk indicates dorsal tooth; long arrows indicate pair of telostegostomatal denticles; B: Same individual shown in (A), plane left of sagittal, left lateral view. Arrow indicates one of the pair of left subventral metastegostomatal denticles (both are shown); C: Vagina of *S. aristotokia* n. gen., n. sp., sagittal plane, right lateral view. Vagina is lined anteriorly and posteriorly by series of sac-like cells with a thin cytoplasm and membranous appearance; D: Vagina of *S. crassa* n. gen., n. sp., showing vaginal sac-like cells with a refractive cytoplasm and thickened appearance; E: Mature hermaphrodite of *S. aristotokia* n. gen., n. sp., with a series of developing embryos and juveniles in oviduct; F: Oviduct of hermaphrodite of *S. crassa* n. gen., n. sp. with growing embryos; G: Dauer (D) and directly developing J3 (J3) juveniles within the oviduct of *S. crassa* n. gen., n. sp.; H: J4 of *S. crassa* n. gen., n. sp., gravid with developing embryos, located near developing vulva. (Scale bars = 10 μm unless otherwise indicated.)
Table 1. (Continued.)

<table>
<thead>
<tr>
<th>Character</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>S. aristotokia n. gen., n. sp.</td>
<td>S. crassa n. gen., n. sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Male Holotype</td>
<td>Male Temporary water mounts</td>
<td>Hermaphrodite Temporary water mounts</td>
<td>Male Holotype Temporary water mounts</td>
<td>Hermaphrodite Temporary water mounts</td>
</tr>
<tr>
<td>Basal bulb diam.</td>
<td>12</td>
<td>14 ± 1.3 (13-16)</td>
<td>18 ± 1.8 (16-21)</td>
<td>10</td>
</tr>
<tr>
<td>Excretory pore from ant. end</td>
<td>106</td>
<td>97 ± 6.2 (90-103)</td>
<td>112 ± 12 (95-127)</td>
<td>72</td>
</tr>
<tr>
<td>Nerve ring from ant. end</td>
<td>88</td>
<td>91 ± 5.0 (87-99)</td>
<td>107 ± 7.8 (96-121)</td>
<td>57</td>
</tr>
<tr>
<td>Vulva from ant. end</td>
<td>–</td>
<td>–</td>
<td>479 ± 60 (365-563)</td>
<td>–</td>
</tr>
<tr>
<td>Cloacal or anal body diam.</td>
<td>14</td>
<td>20 ± 5.9 (13-27)</td>
<td>26 ± 4.2 (21-35)</td>
<td>15</td>
</tr>
<tr>
<td>Tail length</td>
<td>18</td>
<td>35 ± 13 (22-54)</td>
<td>90 ± 8.1 (77-102)</td>
<td>21</td>
</tr>
<tr>
<td>Spicule length (curve)</td>
<td>31</td>
<td>30 ± 3.3 (27-35)</td>
<td>–</td>
<td>24</td>
</tr>
<tr>
<td>Gubernaculum length</td>
<td>10</td>
<td>9.4 ± 1.2 (8.2-11)</td>
<td>–</td>
<td>14</td>
</tr>
</tbody>
</table>

systems. Perhaps most remarkable is the precocious gonad development in Sudhausia n. gen., whereby fertilisation and offspring development occur before the mother reaches the adult stage, a phenomenon made possible by self-sufficient (hermaphroditic) reproduction. The twofold size increase of embryos in Sudhausia n. gen. is also unique, to our knowledge, among nematodes and other animals with an eggshell. Although the nematode eggshell is not expected to harden until after being passed through uterine or vaginal sphincters (e.g., Bird et al., 1993), embryonic development as currently perceived proceeds by cell division without an increase in embryo size (Sulston et al., 1983; Schulze & Schierenberg, 2011). Finally, the demonstrated capability for vivipary tentatively distinguishes Sudhausia n. gen. from most other diplogastrid species, except for Koeneria paramata (Schneider, 1938) Sudhaus & Fürst von Lieven, 2003 (Fürst von Lieven, 2005) and Mehedinema alii Farooqui, 1967 (Luong et al., 1999), although more detailed examination of other diplogastrids exhibiting intra-uterine birth (usually endotokia matricida) may reveal further cases of vivipary. However, whereas endotokia matricida in particular is commonly a facultative response to a stressful environment (Johnigk & Ehlers, 1999; Chen & Caswell-Chen, 2004), S. aristotokia n. gen., n. sp. and S. crassa n. gen., n. sp. are constitutively viviparous, even under well-fed laboratory conditions. We hypothesise that the unique vaginal morphology of Sudhausia n. gen., namely the anterior and posterior columns of sac-like cells lining the vagina, is correlated with this life-history trait, although presently the function of these structures is unknown.

The isolation of new higher-order taxa such as Sudhausia n. gen. from insect associates suggests that this general habitat is still far from being adequately sampled. The collection of new species from beetles in particular promises to reveal further cases of novel trait combinations and expand the known spectrum of morphological and biological diversity in Diplogastridae. The presently unclear relationship of Sudhausia n. gen. (+ Tylopharynx, Neodiplogaster and Mononchoides) to other major lineages of the family suggests that ongoing sampling will be necessary to provide greater resolution of deeper relationships among them. The collection of new or understudied taxa from other insect hosts (Herrmann et al., unpubl.; Kanzaki, unpubl.) is likely to help in this regard, and it will be essential for refining the framework needed for ongoing macroevolutionary studies in the group.
**Fig. 9.** Mature adults of *Sudhausia crassa* n. gen., n. sp. A: Hermaphrodite, right lateral view; B: Male, left lateral view; C: Neck region of hermaphrodite, left lateral view; D: Lip region of male, left subdorsal view; E: Stomatal region of hermaphrodite, left lateral view; F: Stomatal region of hermaphrodite, ventral view.
Fig. 10. Adults of *Sudhausia crassa* n. gen., n. sp. A: Reproductive tract of young hermaphrodite, right lateral view; B: Vulva, ventral view; C: Tail region of hermaphrodite, left lateral view; D: Anus, ventral view; E: Male tail, ventral view; F: Spicule and gubernaculum, left lateral view; G: Male tail, left lateral view. Abbreviations: R1-R6 = bursal ray papillae; d = dorsal; v = ventral.

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References


