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Author for correspondence:

I. Blasco-Costa, E-mail: isa.blasco.costa@gmail.com

A new species of *Atriophallophorus* Deblock & Rosé, 1964 (Trematoda: Microphallidae) described from *in vitro*-grown adults and metacercariae from *Potamopyrgus* antipodarum (Gray, 1843) (Mollusca: Tateidae)

I. Blasco-Costa^{1,2} , K. Seppälä^{3,4,5}, F. Feijen^{3,4}, N. Zajac^{3,4}, K. Klappert^{3,4} and J. Jokela^{3,4}

¹Natural History Museum of Geneva, PO Box 6434, CH-1211 Geneva 6, Switzerland; ²Department of Arctic and Marine Biology, UiT The Arctic University of Norway, Langnes, PO Box 6050, 9037 Tromsø, Norway; ³Department of Aquatic Ecology, Swiss Federal Institute of Aquatic Science and Technology (EAWAG), Dübendorf, Switzerland; ⁴Institute of Integrative Biology, ETH-Zürich, Zürich, Switzerland and ⁵Research Department for Limnology, University of Innsbruck, 5310 Mondsee, Austria

Abstract

The adult and metacercaria life stages of a new species of the microphallid genus Atriophallophorus Deblock & Rosé, 1964 are described from specimens collected at Lake Alexandrina (South Island, New Zealand). In addition to molecular analyses of ribosomal and mitochondrial genes, metacercariae of Atriophallophorus winterbourni n. sp. from the snail host Potamopyrgus antipodarum (Gray) were grown in vitro to characterize internal and external morphology of adults using light and scanning electron microscopy and histological techniques. Atriophallophorus winterbourni n. sp. is readily distinguishable from Atriophallophorus coxiellae Smith, 1973 by having a different structure of the prostatic chamber, sub-circular and dorsal to genital atrium, rather than cylindrical, fibrous, elongate and placed between the seminal vesicle and the genital atrium. The new species is most similar to Atriophallophorus minutus (Price, 1934) with regards to the prostatic chamber and the morphometric data, but possesses elongate-oval testes and subtriangular ovary rather than oval and transversely oval in A. minutus. Phylogenetic analyses including sequence data for A. winterbourni n. sp. suggested a congeneric relationship of the new species to a hitherto undescribed metacercariae reported from Australia, both forming a strongly supported clade closely related to Microphallus and Levinseniella. In addition, we provide an amended diagnosis of Atriophallophorus to accommodate the new species and confirm the sinistral interruption of the outer rim of the ventral sucker caused by the protrusion of the dextral parietal atrial scale at the base of the phallus.

Introduction

A microphallid species reported as *Microphallus* sp. from New Zealand has been largely used as a model species in parasitology and evolutionary studies for more than three decades (e.g. Lively, 1987; Dybdahl & Lively, 1998; Lively & Dybdahl, 2000; Jokela *et al.*, 2009; Gibson *et al.*, 2016a). Broad knowledge has accumulated on the role of this parasite in coevolutionary dynamics with the host populations. Particularly, in this case the coevolutionary dynamics are suggested to favour sexual reproduction by the host (i.e. 'parasite hypothesis' or 'Red Queen hypothesis' for maintenance of sex; Jaenike (1978); Hamilton (1980); Lively (2016)). However, basic knowledge such as the identity of the species, its genetic diversity and phylogenetic affinities are still lacking, with the consequences that it may be difficult to judge the source of heterogeneity in infection experiments and field surveys.

Microphallids are a diverse and cosmopolitan group of very small worms, typically found in the intestine of birds. Second intermediate hosts tend to be crustaceans, but several species present abbreviated life cycles where the metacercariae encyst within the mollusc first intermediate host. Their metacercariae are usually identical to the adult stage in the definitive host. In the definitive host, microphallids tend to mature in a few days and live only for several weeks (Galaktionov & Dobrovolskij, 2003). This feature of microphallids makes them amenable to *in vitro* culture as a method to obtain the mature adults and be able to characterize and describe new species. Such an alternative is, nowadays, desirable, considering the limitations of obtaining permits to collect potential vertebrate definitive hosts, which are in many cases protected by national or international regulations (Blasco-Costa & Poulin, 2017).

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The aim of this study is to morphologically characterize and describe the adult and metacercaria life stages of this microphallid, which represents a new species of the genus *Atriophallophorus* Deblock & Rosé, 1964 instead of *Microphallus* Ward, 1901 as had been previously reported. Metacercariae of microphallid specimens from the snail host *Potamopyrgus antipodarum* (Gray) were grown *in vitro* and the adults were analysed with light microscopy, scanning electron microscopy (SEM) and histological techniques to characterize their internal and external morphology. In addition, this study provides an amended diagnosis of *Atriophallophorus*, molecular data for two nuclear and two mitochondrial markers of the new species and an evaluation of its relationship with other microphallids.

Material and methods

Specimens

Potamopyrgus antipodarum snails were collected using a kick net and snorkelling from two sites in Lake Alexandrina (South Island, New Zealand), site 'Swamp' (-43.962102, 170.441728) and site 'JMS' (-43.937199, 170.459495). Live metacercariae were dissected from four infected snails from each locality and allowed to excyst in Tyrode's salt solution (Sigma, Buchs, Switzerland) with pancreatin (3 mg/ml; Sigma) and penicillin-streptomycinneomycin solution (8% v/v; Sigma) at 39°C. Excysted juvenile worms were washed twice with Tyrodes's salt solution supplemented with penicillin-streptomycin-neomycin solution (8% v/v). Then, juveniles from the same locality were pooled and transferred to culture medium of RPMI 1640 (Gibco, ThermoFisher Scientific, Waltham (MA) U.S.) supplemented with horse serum (20% v/v, Gibco), penicillin-streptomycin-neomycin solution (8% v/v), HEPES buffer (25 mm; Gibco) and Amphotericin B solution (0.25 mg/ml; Sigma). Juveniles were incubated at 39°C and cultured for up to three days (72 h). Fresh culture media was changed daily. Metacercariae right after hatching and in vitro-grown adults after 24 h, 48 h and 72 h of culturing were fixed in hot saline and preserved in 75% ethanol for later morphological examination. A subsample of specimens was preserved in hot formalin for examination using SEM and histological analysis. Additionally, five metacercariae from each site (from a pool of four snails each) were preserved in 100% ethanol for molecular analyses.

Morphological data

Metacercariae and adult specimens grown in vitro were stained using iron acetocarmine, dehydrated through a graded ethanol series, cleared in dimethyl phtalate and examined as permanent mounts in Canada balsam. Figures were made using a drawing tube mounted on a Zeiss light compound microscope at ×1600 magnification. Measurements of the specimens were taken from drawings at ×640 magnification. Five specimens were dehydrated in a graded ethanol series, critical point-dried and sputter-coated with gold for SEM examination using a Zeiss DSM 940A (Zeiss AG, Oberkochen, Germany) at an accelerating voltage of 5 kV. Three specimens were used for histology after dehydration through a graded ethanol series, followed by propylene oxide and immersed in Epon resin. Histological sections of 1 µm were made with an ultramicrotome Reichert-Jung Ultracut E (Leica Microsystems, Heerbrugg, Switzerland), stained with toluidine blue 0.75% and examined using the light microscope mentioned previously. All measurements in the text are in micrometres unless otherwise stated, and are given as the range followed by the mean \pm standard deviation. Permanent mounts of the type material and SEM preparations are deposited in the Platyhelminthes collection of the Natural History Museum of Geneva, the Institute of Parasitology of the Academy of Sciences of the Czech Republic and the Otago Museum, Dunedin, New Zealand.

Molecular data

Four metacercariae isolates from Swamp and five from JMS originating from a pool of four infected P. antipodarum from each site were characterized molecularly. Due to the small size of these specimens, it was impossible to keep hologenophores of the sequenced specimens. Nonetheless, the specimens used for morphological description herein represent paragenophores and likely genetic clones of the sequenced material. Genomic DNA was extracted from ethanol-fixed metacercariae isolates in 200 µl of a 5% suspension of Chelex® in deionized water and containing 0.1 mg/ml proteinase K followed by incubation at 56°C for 5 h, boiling at 90°C for 8 min and centrifugation at 14,000 g for 10 min. Partial fragments were amplified of the large ribosomal subunit (28S) (1800 bp; primers U178F: 5'-GCA CCC GCT GAA YTT AAG-3' and L1642R: 5'-CCA GCG CCA TCC ATT TTC A-3'; Lockyer et al., 2003) and internal transcribed spacer 2 (ITS2) (500 bp; primers 3S: 5'-GTA CCG GTG GAT CAC GTG GCT AGT G-3' and ITS2.2: 5'-CCT GGT TAG TTT CTT TTC CTC CGC-3' (Morgan & Blair, 1995; Cribb et al., 1998)). Additionally, two fragments of mitochondrial genes (mt) were amplified: cytochrome oxidase subunit I (cox I) (1050 bp; primers JB3: 5'-TTTTTTGGGCATCCTGAGGTTTAT -3' and microph_rev: 5'-AAT CAT GAT GCA AAA GG-3' (Bowles et al., 1993; newly designed)) and nicotinamide adenine dinucleotide dehydrogenase subunit 5 (NADH5) (744 bp; primers F2_micND5: 5'-CTTCAACCTTGGTTGCTGCC-3' and R2_micND5: 5'-TCCC AACGAAACCTAAAACTGC-3' (newly designed)).

Polymerase chain reaction (PCR) amplifications were performed in 20 µl reactions containing 2 µl of extraction supernatant (~10-20 ng of template DNA), 2× MyFiTM Mix (Bioline France, France; containing DNA polymerase, dNTPs, MgCl₂ and enhancers at optimal concentrations) and 0.4 µM of each primer combination. Thermocycling conditions used for amplification of the rDNA regions followed Galaktionov et al. (2012). The following thermocycling profile was used for amplification of the mt cox I and NADH5 fragments: denaturation (95°C for 3 min); 38 cycles of amplification (94°C for 50 s, 52°C for 30 s and 72°C for 1 min); and 4 min extension step at 72°C. PCR amplicons were purified prior to sequencing using exonuclease I and shrimp alkaline phosphatase enzymes (Werle et al., 1994). Amplicons were cycle-sequenced from both strands using PCR primers and an internal primer for the 28S fragment (L1200R: 5'-GCA TAG TTC ACC ATC TTT CGG-3'; Littlewood et al., 2000) at the commercial facility Macrogen (Amsterdam, The Netherlands). Contiguous sequences were assembled and edited using Geneious® (v. 8.1 Biomatters Ltd., Auckland, New Zealand) and submitted to GenBank (see accession numbers in table 1).

Molecular analyses

Newly generated sequences for the 28S rDNA and the ITS2 fragments were aligned in two independent datasets, together with the published sequences of other microphallids from GenBank (see accession numbers in table 1). The sequences were aligned

 Table 1. List of taxa included in the phylogenetic analyses, GenBank accession numbers and references.

AF387801 AY220617 AF151918 MN342153- MN342154 MG783585 KX712084 AY220629 KT355818 KT880222 KJ144173	MN342155 MG783580 KX712086 HM584171 KT355824	Snyder & Tkach (2001) Tkach et al. (2003) Tkach et al. (2000) This study Galaktionov & Blasco-Costa (2018) Kudlai et al. (2016) Galaktionov et al. (2012) Kudlai et al. (2015)
AY220617 AF151918 MN342153- MN342154 MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	Tkach et al. (2003) Tkach et al. (2000) This study Galaktionov & Blasco-Costa (2018) Kudlai et al. (2016) Galaktionov et al. (2012)
AF151918 MN342153- MN342154 MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	Tkach et al. (2000) This study Galaktionov & Blasco-Costa (2018) Kudlai et al. (2016) Galaktionov et al. (2012)
AF151918 MN342153- MN342154 MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	Tkach et al. (2000) This study Galaktionov & Blasco-Costa (2018) Kudlai et al. (2016) Galaktionov et al. (2012)
MN342153- MN342154 MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	This study Galaktionov & Blasco-Costa (2018) Kudlai <i>et al.</i> (2016) Galaktionov <i>et al.</i> (2012)
MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	Galaktionov & Blasco-Costa (2018) Kudlai <i>et al.</i> (2016) Galaktionov <i>et al.</i> (2012)
MG783585 KX712084 AY220629 KT355818 KT880222	MG783580 KX712086 HM584171	Galaktionov & Blasco-Costa (2018) Kudlai <i>et al.</i> (2016) Galaktionov <i>et al.</i> (2012)
KX712084 AY220629 KT355818 KT880222	KX712086 HM584171	Kudlai <i>et al.</i> (2016) Galaktionov <i>et al.</i> (2012)
AY220629 KT355818 KT880222	HM584171	Galaktionov et al. (2012)
KT355818 KT880222		
KT880222	KT355824	Kudlai <i>et al</i> . (2015)
Κ I1 <u>44</u> 172		Hernandez-Orts et al. (2016)
I/DT44T12		Presswell et al. (2014)
JF826247	HQ650132	Al-Kandari et al. (2011)
AY220632		Tkach et al. (2003)
	KF575167	Diaz & Cremonte (2010)
AF151927		Tkach <i>et al.</i> (2000)
KJ144178	KJ540203	Presswell et al. (2014)
AY220630	HM584170	Tkach et al. (2003); Galaktionov et al. (20
KJ144175		Presswell et al. (2014)
AY220631		Tkach <i>et al.</i> (2003)
AF151926	HM584172	Tkach et al. (2000); Galaktionov et al. (20
	KC012521	Gilardoni et al. (2011)
	KC222022	Gilardoni et al. (2011)
KT355820	KT355826	Kudlai <i>et al.</i> (2015)
AB974360		Kakui (2014)
AY220633		Tkach et al. (2003)
AY220626	HM584173	Tkach et al. (2003); Galaktionov et al. (20
		Tkach et al. (2003)
HM584125	HM584183	Galaktionov et al. (2012)
HM584140	HM584185	Galaktionov et al. (2012)
KT355822	KT355828	Kudlai <i>et al</i> . (2015)
HM584142	HM584175	Galaktionov et al. (2012)
HM584122	HM584181	Galaktionov et al. (2012)
AY220627		Tkach <i>et al.</i> (2003)
HM584126	HM584198	Galaktionov et al. (2012)
HM584133	HM584190	Galaktionov et al. (2012)
HM584138	HM584178	Galaktionov et al. (2012)
HM584139	HM584195	Galaktionov et al. (2012)
KJ868216		O'Dwyer et al. (2014)
	GQ377842	Al-Kandari & Al-Bustan (2010)
	AY220632 AF151927 KJ144178 AY220630 KJ144175 AY220631 AF151926 KT355820 AB974360 AY220633 AY220626 AY220628 HM584125 HM584140 KT355822 HM584142 HM584142 HM584133 HM584133 HM584138 HM584139	AY220632 KF575167 AF151927 KJ144178 KJ540203 AY220630 HM584170 KJ144175 AY220631 AF151926 HM584172 KC012521 KC222022 KT355820 KT355820 KT355826 AB974360 AY220633 AY220626 HM584173 AY220628 HM584125 HM584183 HM584140 HM584155 KT355822 KT355822 KT355828 HM584142 HM584155 HM584160 HM584175 HM584181 AY220627 HM584133 HM584198 HM584138 HM584138 HM584138 HM584139 HM584195 KJ868216

(Continued)

Table 1. (Continued.)

Таха	28S rDNA	ITS2	References
Pleurogenidae			
Parabascus duboisi	AY220618		Tkach <i>et al.</i> (2003)
Pleurogenes claviger	AF151925		Tkach <i>et al</i> . (2000)
Prosthogonimidae			
Prosthogonimus ovatus	AF151928		Tkach <i>et al</i> . (2000)
Telorchiidae			
Telorchis assula	AF151915		Tkach <i>et al</i> . (2000)

using default parameters of MAFFT implemented in Geneious®, and the extremes of the alignment were trimmed to match the shortest sequences. The 28S dataset (1280 bp long) included 14 representative sequences of Microphallus spp., 12 of Maritrema spp., one each of Levinseniella, Longiductotrema and an unidentified microphallid of Kudlai et al. (2015) and a sequence labelled as Microphallus fusiformis (which should be disregarded as a species of Microphallus; see Kudlai et al., 2015) retrieved from GenBank (table 1). Additionally, five sequences of species belonging to sister families of the Microphallidae - i.e. Lecithodendriidae, Pleurogenidae and Prosthogonimidae in the Microphalloidea - and three sequences of species in the Plagiorchioidea were retrieved from GenBank and included as outgroups. The ITS2 dataset (401 bp long) included ten representative sequences of Microphallus spp.; nine sequences of Maritrema; one each of Levinseniella, Probolocoryphe and Longiductotrema; and one of an unidentified microphallid of Kudlai et al. (2015). The phylogenetic analyses were run on the two datasets individually under the maximum likelihood (ML) and Bayesian inference (BI) criteria, employing the nucleotide substitution model GTR+Γ. ML analyses were conducted using the program RAxML v. 8.2 (Stamatakis, 2014). All model parameters, bootstrap nodal support values (1000 repetitions) and an extended majority-rule consensus topology were estimated using RAxML. BI trees were constructed using MrBayes v. 3.2 (Ronquist et al., 2012), running two independent Markov Chain Monte Carlo runs of four chains with standard settings for 10⁷ generations and sampling tree topologies every 10³ generation. Burn-in periods were set automatically to 25% generations, ensuring the remaining trees were obtained after values for standard deviation of split frequencies were <0.01. A majority-rule consensus topology and nodal support estimated as posterior probability values (Huelsenbeck et al., 2001) were calculated from the remaining trees. All MrBayes and RAxML analyses were performed on the computational resource CIPRES (Miller et al., 2010). Genetic divergences amongst taxa were calculated as uncorrected p-distances for each gene region using MEGA v. X (Kumar et al., 2018).

Results

Family: Microphallidae Ward, 1901 Atriophallophorus Deblock & Rosé, 1964

Atriophallophorus winterbourni n. sp.

Taxonomic summary

Synonyms. Metacercaria A of Winterbourn (1974); Microphallus sp. of Lively (1987, 1989), Lively & McKenzie (1991), Jokela &

Lively (1995), Dybdahl & Lively (1996, 1998), Levri & Lively (1996), Lively & Jokela (1996), Levri (1999), Krist et al. (2000, 2004), Levri & Fisher (2000), Jokela et al. (2003, 2009), Dybdahl & Krist (2004), Lively et al. (2004, 2008), Osnas & Lively (2004, 2006, 2011), Levri et al. (2005), Fromme & Dybdahl (2006), Koskella & Lively (2007, 2009), Lagrue et al. (2007), Dybdahl et al. (2008), Lagrue & Poulin (2008), Lively et al. (2008), King et al. (2009, 2011a, b), Koskella et al. (2011), Vergara et al. (2013, 2014, 2017), Paczesniak et al. (2014, 2019), Gibson et al. (2016a, b, 2018), McKone et al. (2016), Bankers et al. (2017), Bankers & Neiman (2017); Microphallus sp. 'lively' of Hofmann et al. (2016); Microphallus sp. 'lively' of Hechinger (2012), Soper et al. (2014), Bankers & Neiman (2017); Microphallus livelyi (nomen nudum) of Bankers & Neiman (2017).

Type host. Potamopyrgus antipodarum (Gray) (Mollusca: Tateidae; first and second intermediate host).

Definitive host. Waterfowl (see Osnas & Lively, 2011).

Type locality. Lake Alexandrina, South Island, New Zealand (site 'Swamp': -43.962102, 170.441728).

Other localities. Lake Alexandrina, South Island, New Zealand (site 'JMS': -43.937199, 170.459495).

Site in host. Intermediate host: metacercaria encysted in gonads.

Type material. Holotype (MHNG-PLAT-129859), 22 paratypes (MHNG-PLAT-129860–MHNG-PLAT-129865) and seven metacercariae vouchers (MHNG-PLAT-129866–MHNG-PLAT-129867) deposited in the Platyhelminthes collection of the Natural History Museum of Geneva (NHMG), two paratypes (IPCAS D-803) deposited in the helminthological collection of the Institute of Parasitology, Česke Budějovice, Czech Republic (IPCAS) and four paratypes (OMNZ IV110293–OMNZ IV110295) and two voucher metacercariae (OMNZ IV110296) deposited at the Otago Museum (OM), Dunedin, New Zealand.

Representative DNA sequences. 28S rDNA, four sequences of specimens from Swamp and five of specimens from JMS (GenBank MN342153–MN342154); ITS2 rDNA, eight identical sequences (four of specimens from Swamp and four from JMS; GenBank MN342155); mt cox I, eight identical sequences (three of specimens from Swamp and five from JMS; GenBank MN342156); and mt NADH5, four sequences of specimens from Swamp and five of specimens from JMS; GenBank MN342157–MN342158).

ZooBank registration. To comply with the regulations set out in article 8.5 of the amended 2012 version of the International Code of

Zoological Nomenclature, details of the new species have been submitted to ZooBank. The Life Science Identifier for *A. winterbourni* n. sp. is urn:lsid:zoobank.org:pub:E0E7C4F3-430E-42BD-9FEC-2D 11A159DD12.

Etymology. The species is named after Professor Michael Winterbourn in recognition for his contribution to the field of freshwater ecology – in particular, our knowledge of freshwater invertebrates of New Zealand, including the first mention of this species.

Description of adult (figs 1a-c and 2a-h; table 2; supplementary table S1)

(Based on whole mounts of 29 gravid specimens grown *in vitro* culture for 24–72 h and SEM preparations. Measurements provided as range and mean ± standard deviation for the type series, variation associated with specimens from each age class (duration of culture) are provided separately in supplementary table S1.)

Body minute, triangular, often curved concave ventrally (body-width-to-length ratio 1:1.3-1.9 ($1:1.6\pm0.4$)) with maximum width at posterior level of testes, 145-200 (167 ± 12) × 85-115 (100 ± 8). Tegument bears spines, glands and sensory papillae. Spines palmate, smooth, 5-9 prongs; present in lateral margins and alongside midline, separated by two narrow ventro-lateral regions devoid of spines, sparser towards posterior extremity on dorsal side; anterior forebody spines width 1.5-1.8, inter-spine space 0.6-0.8; midforebody spines width 1.9-2.2, inter-spines space 0.6-1; anterior dorsal spines width 1.5-1.8, inter-spines space 1.0-1.5; mid-dorsal spines width 1.5-1.8, inter-spines space 2.1-4.1; lateral posterior dorsal spines width 1.6-1.8, inter-spines space 1.2-1.5. Forebody 81-125 (97 ± 9) long, representing 52-63% ($58\pm2\%$) of body length. Glands and sensory papillae at anterior extremity, surrounding oral sucker (>14) and lateral margins of body.

Oral sucker subterminal, spherical, 21-29 (25 ± 2) × 21-29 (25 ± 2). Ventral sucker at two-thirds of body length, subspherical, complete, 24-30 (27 ± 1) × 22-33 (28 ± 2); outer rim crescent, interrupted sinistrally by genital pore, bearing spines and nine glands; oral-sucker-to-ventral-sucker-length ratio 1: 0.9-1.2 ($1:1.1\pm0.2$), width ratio 1: 0.9-1.3 ($1:1.1\pm0.1$). Pre-pharynx absent. Pharynx small, oval 11-17 (14 ± 2) × 11-15 (13 ± 1). Pharynx-length-to-oral-sucker-length ratio 1:1.5-2.4 ($1:1.7\pm0.2$). Oesophagus 29-46 (36 ± 5) long. Intestinal bifurcation pre-equatorial, immediately anterior to seminal vesicle. Caeca as long as oesophagus, widely divergent, extend to anterior margin of testes.

Testes two, postovarian, symmetrical, lateral, somewhat diagonal, smooth, slightly elongate-oval, right testis $18-29 (25 \pm 4) \times 17-28$ (20 ± 3) ; left testis 18-33 $(25 \pm 5) \times 14-24$ (19 ± 3) . Seminal vesicle arcuate, transversely oval, intercaecal in mid-body, overlapping anterior margin of ventral sucker dorsally, $12-24 (17 \pm 3) \times 24-39$ (32 ± 3) . Seminal-vesicle-length-to-ventral-sucker-length 1:1.1–2.2 (1:1.7 \pm 0.3). Seminal proximal duct long, entering prostatic chamber (i.e. phallophorus). Prostatic chamber subspherical, with loose fibres, sinistral to ventral sucker, dorsal to genital atrium, reaching proximal part of seminal vesicle. Prostatic glands not observed. Ejaculatory duct sinuous, enters prostatic chamber, opens to small papillae dorso-sinistral in genital atrium, together with prostatic ducts running through to periphery of male duct. Phallus of 'microphalloid'-type, glabrous, turgid, evaginable, with ejaculatory duct in axis, with large dextral triangular distal scale at the base. Genital atrium surrounding phallus, 16-24 (20 ± 3), genital pore large, sinistral to ventral sucker.

Ovary dextral to ventral sucker, pre-testicular, ventral to caeca, adjacent to or slightly overlapping ventral sucker laterally,

subtriangular with large cells, 13-26 (21 ± 3) × 17-34 (25 ± 4). Oötype inter-testicular, slightly dextral posterior to ovary and ventral sucker. Mehlis' gland surrounding oötype posteriorly. Laurer's canal not observed. Uterus confined posterior to midlevel of ventral sucker, overlapping testes ventrally. Metraterm thin-walled, with widened opening into sinistral wall of genital atrium. Vitellarium in two compact clusters of follicles, disaggregated in older specimens (after 24 h cultured), para-, post-testicular or overlapping testes, converging into seminal receptacle next to oötype. Eggs few, large. Excretory vesicle obscured by vitellarium masses and eggs. Flame-cell formula not observed.

Description of metacercaria (fig 1d; table 2; supplementary table S1)

Overall form highly developed and consistent with adult anatomy, except for the absence of eggs.

(Measurements based on 18 encysted specimens; measurements provided as range followed by mean ± standard deviation.)

Metacercaria folded within small spherical, translucent cyst, $114-130\times106-120$ ($123\times113\pm4$). Cyst wall consisting of three or more hyaline layers, 5-6 (5 ± 0.5) thick. Metacercaria encysted in the gonads of *P. antipodarum*.

Genetic affinities

Two distinct genotypes for the 28S rRNA gene were obtained from the sequences of the nine specimens analysed. The two genotypes differed in two transitions at the nucleotide positions 281, a cytosine in the sequence of specimens originated from Swamp and a thymine in specimens from JMS and 1358, adenine and guanine, respectively. However, all specimens shared the same sequence for the ITS2 region and the mt cox I. Independent phylogenetic analyses of the 28S rDNA and ITS2 regions using BI and ML methods showed congruent results (figs 3 and 4), with the Microphallidae as monophyletic and A. winterbourni embedded within. A. winterbourni appeared as sister taxon to the sequence of an undescribed microphallid from Australia by Kudlai et al. (2015). In the 28S phylogenetic tree (fig. 3), the sequence of M. fusiformis appeared closely related to A. winterbourni and the undescribed microphallid with strong nodal support, although considerably divergent. In all trees, Atriophallophorus appears closely related to Microphallus and Levinseniella, but the sister relationship among these three genera could not be established due to the low support of an internal node within the clade (figs 3 and 4).

Genetic divergence between *A. winterbourni* and the unidentified microphallid from Australia was 1.2–1.3% for the 28S and 3.2% for the ITS2, which are within the lower limit of the range observed among congeneric species of *Microphallus* (28S: 0.8–9.1%; ITS2: 1.0–9.9%) and *Maritrema* (28S: 0.6–9.1%; ITS2: 0.3–12.3%). The 28S sequences of *A. winterbourni* and the unidentified microphallid diverged 10.5–10.8% from the sequence of *M. fusiformis*, which fell within range of intergeneric distances in the 28S region of microphallids (*Maritrema–Microphallus*: 7.5–12.6%; *Microphallus–Levinseniella*: 5.5–10.0%; *Maritrema–Levinseniella*: 9.1–11.4%; *Longiductotrema–Levinseniella*: 7.7%; *Longiductotrema–Maritrema*: 7.8–11.1%; *Longiductotrema–Microphallus*: 6.5–10.6%).

Sequence data for the COI marker resulted in identical sequences for all specimens (five from JMS and three from Swamp; one sample from swamp did not amplify and another one produced chromatograms with double peaks), whereas sequences for the NADH5 marker showed variability at the

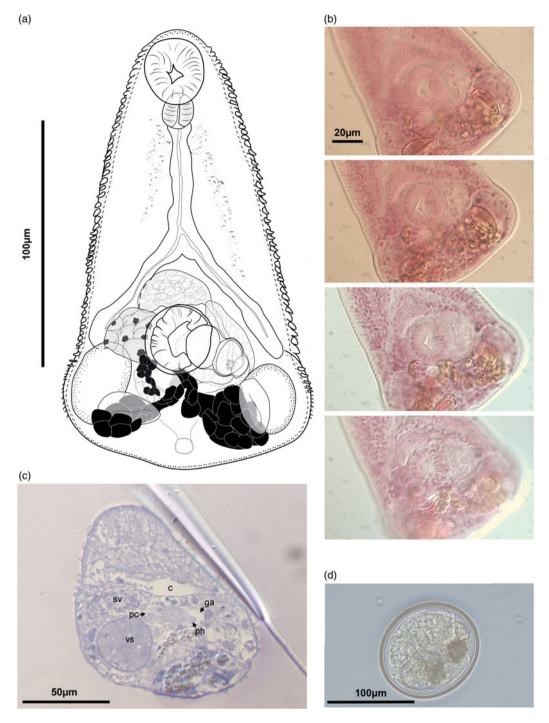


Fig. 1. Atriophallophorus winterbourni n. sp. (a) Illustration of the holotype, 24 h in vitro grown adult in ventral view. (b) Microphotographs of the terminal genitalia of the holotype using light microscopy. (c) Histological oblique section of a paratype at the level of the ventral sucker. (d) Microphotograph of an encysted metacercaria ex *Potamopyrgus antipodarum* (Gray). Abbreviations: c, caeca; ga, genital atrium; pc, prostatic chamber; ph, phallus; sv, seminal vesicle; vs, ventral sucker.

position 191, with specimens from Swamp having a thymine and specimens from JMS a cytosine.

Discussion

Species differentiation

Following the key to the Microphallidae provided by Deblock (2008), the new species described from *in vitro* grown adults

conforms to the general morphology of the Microphallidae in having a typically very small body, longer than broad, densely covered with squamous spines, suckers well-separated and subequal, short digestive tract with divergent caeca not extending posteriorly beyond ventral sucker. Ovary pretesticular, in the opposite side of the body to the genital pore. Two testes, lateral and symmetrical, with male terminal genitalia intercaecal and anterior to ventral sucker. These specimens fit to the diagnosis of the supersubfamily Microphallidi Ward, 1901 by having the

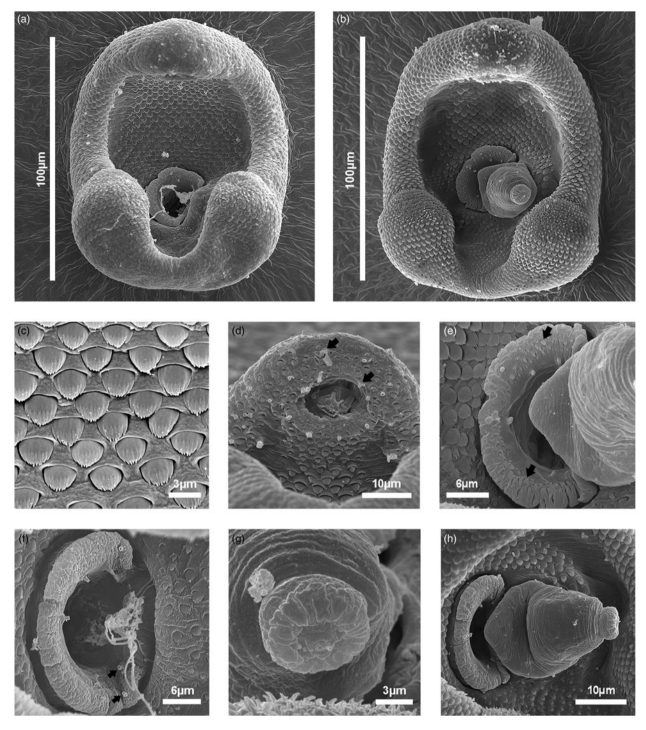


Fig. 2. Scanning electron micrographs of Atriophallophorus winterbourni n. sp. (a) Adult. (b) Adult with phallus protruded. (c) Palmate spines on the ventral surface of the body. (d) Detail of the oral sucker, arrows point at a gland opening and a sensory papilla surrounding the oral sucker. (e) Detail of the outer rim of the ventral sucker with spination and the parietal atrial scale at the basis of the phallus. (f) Outer rim of the ventral sucker interrupted sinistrally by the opening for the genital pore with arrows pointing at glands. (g) Tip of the phallus evaginated, which appears as a flower-like structure when invaginated. (h) Detail of the configuration of the protruded phallus and the sinistrally interrupted outer rim of the ventral sucker.

terminal genitalia free in the parenchyma, and the subfamily Microphallinae Ward, 1901 by presenting a genital atrium that closely envelops the phallus. However, the opening of the genital atrium and the presence of the scale at the base of the phallus of the new species were found to alter the outer rim of the ventral sucker. Whereas the latter feature is characteristic of the monotypic Endocotylinae, our specimens differ from the genus

Endocotyle in lacking a connection between the cavity of the sucker and the genital pore, and presenting a scale at the base of a phallus of the 'microphalloid-type'. Our specimens fall within the Microphallini tribe by having a fleshy and muscular phallus, and the opening of the metraterm on the sinistral wall of the genital atrium. The new species exhibits features consistent with the genus Atriophallophorus: body of triangular shape; male genital

 Table 2. Comparative metrical data for Atriophallophorus spp.

Species	Atı	Atriophallophorus winterbourni n. sp.			Atriophallophorus r	Atriophallophorus coxiellae Smith, 1973			
Stage	Adu	Adult In vitro cultured		Metacercaria Potamopyrgus antipodarum (Mollusca: Tateidae)		Adult Aythya affinis (Eyton) (Aves: Anatidae)	Adult and metacercaria Ecrobia truncata (Vanatta) and Amnicola limosa (Say) (Mollusca: Hydrobiidae) and mice (experimental)	Metacercaria	
Hosts	In vitro c							Coxiella bad (Johnston) (Tomichi Charadrius d Vieillot a ruficapillus T (Aves: Chara	(Mollusca: idae); cucullatus and C. Temminck
Distribution	New Zealand				North America, Caribbean Islands and Europe (France)			Tasmania	
Source	Present study		Present study		Deblock & Rosé, 1964	Price (1934)	Stunkard 1958	Smith, 1973	
	Range (<i>n</i> = 29)	Mean ± SD	Range (<i>n</i> = 9)	Mean ± SD	Range or mean		Range	Range	Mean
Body length	145-200 (28)	167 ± 12	165–193 (9)	175 ± 9	170-200	153-180	120-200	143-229	195
Body width	85–115 (28)	100 ± 8	98-118 (9)	106 ± 7	90-100	105–112	90-120	72-125	90
Forebody	81–125 (28)	97 ± 9	97-115 (9)	104 ± 5					
Oral sucker length	21-29 (29)	25 ± 2	23–28 (9)	25 ± 2	29 ^a		27 ^a		
Oral sucker width	21–29 (28)	25 ± 2	21–28 (9)	24 ± 2	22–25	23-25	23-30	26-34	29
Pharynx length	11–17 (26)	14 ± 2	12-14 (9)	14 ± 1	12ª		12ª	16-44	20
Pharynx width	11-15 (26)	13 ± 1	11-15 (9)	13 ± 1	12–15	10-18	10-18	15-20	17
Oesophagus	29-46 (22)	36 ± 5	40-50 (8)	44 ± 4		18-30	16???40		
Ventral sucker length	24–30 (28)	27 ± 1	28-35 (9)	31 ± 2	23 ^a		26 ^a		
Ventral sucker width	22-33 (28)	28 ± 2	24-31 (9)	28 ± 2	19-25	22-27	20-28	26-31	29
Right testis length	18-29 (13)	25 ± 4	26-34 (8)	30 ± 3					
Right testis width	17-28 (12)	20 ± 3	24-29 (9)	26 ± 2	20-30	28	12-29	36-40	39
Left testis length	18-33 (10)	25 ± 5	26-35 (7)	30 ± 3					
Left testis width	14-24 (10)	19 ± 3	23-29 (9)	26 ± 2	25-30				
Seminal vesicle length	12-24 (27)	17 ± 3	9–15 (2)	13 ± 3	13-19		14-20		
Seminal vesicle width	24–39 (28)	32 ± 3	35-44 (3)	40 ± 6	13-26		26 ^a	29-36	34
Ejaculatory duct length	21–34 (11)	27 ± 3			20-25				
Ejaculatory duct width	1.5-2.3 (13)	2.0 ± 0.4			2.5				
Genital atrium width	16-24 (10)	20 ± 3	22-23 (2)	22 ± 1	20-25		20-35		

Ovary length	13-26 (20)	21 ± 3	23-31 (9)	25 ± 2	19–25	18	15–20		
Ovary width	17-34 (20)	25 ± 4	17-31 (9)	26 ± 5	13-16	22	21–24	30-48	35
Right vitellarium length	15-26 (4)	20 ± 5	20-28 (9)	25 ± 3	20		5-6 follicles		
Right vitellarium width	23-35 (4)	29 ± 5	28-54 (8)	37 ± 9	30				
Left vitellarium length	15-28 (4)	21 ± 5	20-23 (9)	22 ± 1					
Left vitellarium width	23-37 (4)	28 ± 6	20-35 (9)	31 ± 5					
Egg number	1–12 (29)	5 ± 3			20-25		10-30		
Egg length × width	20-26 × 11-15				21-25 × 10-13	22 × 13	20-22 × 12-13		
Oral-to-ventral-sucker-length ratio	0.9-1.2	1.1 ± 0.2	1.1-1.4	1.3 ± 0.1	0.8 ^a		1.0 ^a	0.9 ^a	
Oral-to-ventral-sucker-width ratio	0.9-1.3	1.1 ± 0.1	0.9-1.3	1.1 ± 0.1	1.1		0.7 ^a	0.9 ^a	
Body-width-to-length ratio	1.3-1.9	1.6 ± 0.4	1.5-1.8	1.7 ± 0.1	1.8ª		-	1.6ª	
Seminal-vesicle-to-ventral-sucker-length ratio	1.1-2.2	1.7 ± 0.3	2.1-3.8	2.8 ± 0.9	0.7 ^a		1.0 ^a	1.9 ^a	
Pharynx-to-oral-sucker-length ratio	1.5-2.4	1.7 ± 0.2	1.7-2.0	1.8 ± 0.1	2.4 ^a		2.3 ^a	1.3ª	
Forebody as percentage of body length	52-63	58 ± 2	57-62	60 ± 1	62 ^a		-	63ª	
Metacercaria cyst length			114-130	123 ± 4					
Metacercaria cyst width			106-120	113 ± 4			≤100	88-125	
Metacercaria cyst wall			5-6	5 ± 0.5			8.0-10		

^aEstimated from the published illustration. *n*, number of specimens measured; SD, standard deviation of the mean.

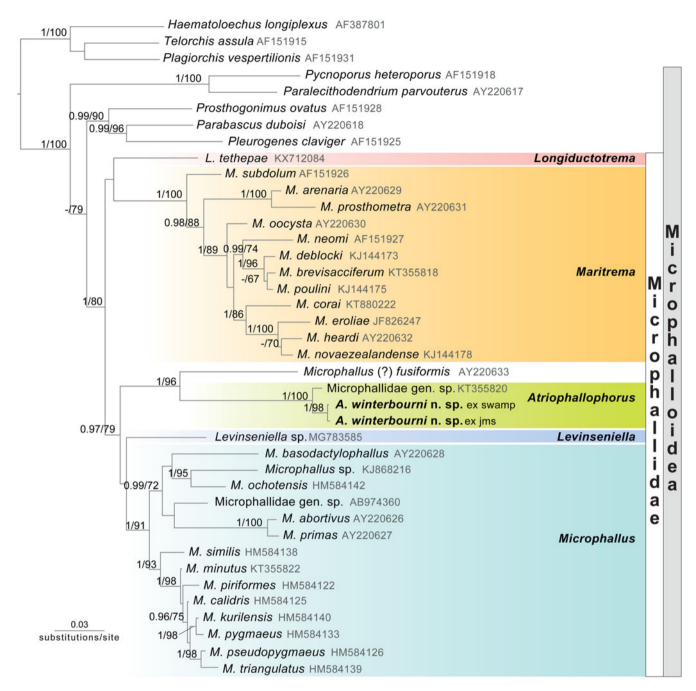


Fig. 3. Phylogenetic relationships for representatives of the family Microphallidae, inferred by maximum likelihood analysis of 28S rDNA sequence data. The newly generated sequences are indicated in bold. Values on the branches correspond to posterior probabilities >0.95 followed by bootstrap support >60. Values below these thresholds were not reported.

pouch absent; proximal ejaculatory ducts very long and entering a large prostatic chamber (described as 'phallophorus' by Deblock & Rosé (1964)), subcircular and similar in size to the ventral sucker and dorsal to the genital atrium, with long prostatic ducts running through the periphery of the male duct; genital atrium containing a phallus of the 'microphalloid-type' and, at its base, a large dextral parietal atrial scale often protruding from the genital pore; a metraterm that widens at its opening into the sinistral wall of the genital atrium; and few but large eggs.

Currently the genus contains only two species, A. minutus (Price, 1934) and A. coxiellae Smith, 1973. Atriophallophorus

coxiellae was described from metacercariae infecting Coxiella badgerensis in a freshwater lake in Tasmania (Smith, 1973). However, the metacercaria of A. winterbourni n. sp. is readily distinguishable from A. coxiellae by having a different structure of the prostatic chamber, sub-circular and dorsal to genital atrium, rather than cylindrical, fibrous, elongate and placed between the seminal vesicle and the genital atrium, as well as smaller pharynx and testes, and somewhat smaller oral sucker and ovary, which barely overlap the lower limit of the range of A. coxiellae.

The new species is most similar to A. minutus with regards to the prostatic chamber and the morphometric data. But A.

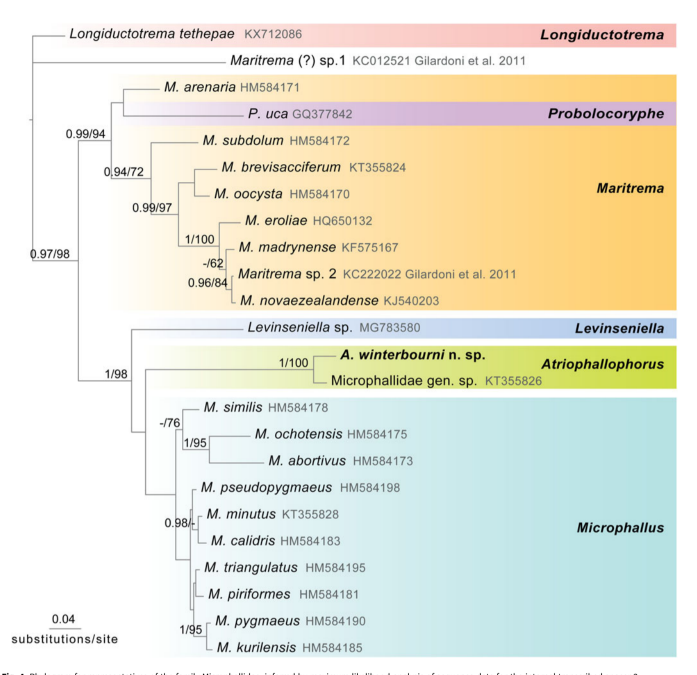


Fig. 4. Phylogram for representatives of the family Microphallidae, inferred by maximum likelihood analysis of sequence data for the internal transcribed spacer 2 of the rRNA genes. The newly generated sequences are indicated in bold. Values on the branches correspond to posterior probabilities >0.95 followed by bootstrap support >60. Values below these thresholds were not reported.

minutus possesses oval testes and a transversely oval ovary, versus elongate-oval and subtriangular, respectively, in A. winterbourni. In addition, the original description of A. minutus, based on adult specimens from Aythya affinis from the Caribbean (Price, 1934), reported A. minutus had a shorter oesophagus than the new species, although this feature might vary greatly depending on the position of the specimen. Morphometric data provided by Stunkard (1958) for the redescription of A. minutus were based on adults experimentally grown in white mice from metacercariae infecting Hydrobia minuta and Amnicola limosa along the east coast of the USA. However, Stunkard's specimens were measured alive, whereas our measurements were taken from fixed, stained and mounted worms. Therefore, the morphometric data are not directly comparable. Furthermore, Stunkard noted

that fixed specimens were slightly smaller than alive, which suggests that if the measurements of *A. minutus* were taken after mounting they would be slightly smaller than those of *A. winter-bourni*. Nonetheless, live specimens of *A. minutus* also show smaller dimensions for several metrical features (body length, ventral sucker width and testes width) extending outside the lower range for *A. winterbourni*, and eggs three times more numerous and with size overlapping the lower range of variation of the new species.

Metrical data of *A. minutus* described by Deblock & Rosé (1964) from France supports that *A. minutus* differs from *A. winterbourni*. Whereas the specimens examined by Deblock & Rosé (1964) were of comparable size to the largest in our sample, their ovary width is smaller than that of *A. winterbourni*, and the ventral sucker width

overlapped and extended below the lower range of variation of *A. winterbourni*. *Atriophallophorus minutus* also shows smaller values for the ratios oral sucker length to ventral sucker length and seminal vesicle length to ventral sucker length (estimated from the original illustrations) than *A. winterbourni*.

In vitro cultivation of the metacercaria and adult allowed us to observe developmental changes through time in the shape of the vitellarium, the seminal vesicle and testes (see also supplementary table S1). In metacercariae, the testes are well visible, the seminal vesicle is small, indistinct and quite empty, and the vitellarium forms two masses of tight follicles. After 24 h of growth, the adults are already gravid, testes become indistinct, the seminal vesicle is full and extended, and the follicles constituting the vitellarium masses start to disaggregate slightly. After 48 h or more, the testes are faintly visible, the seminal vesicle stays fully extended and the vitellarium follicles are drawn further apart in the posterior region of the body. When descriptions are based on specimens collected from the wild birds, this variation in the sample of specimens should be taken into account.

Previous researchers working on Atriophallophorus spp. have emphasized the difficulty of observing the features of specimens of such tiny size. Thus, it is highly recommended to describe new species with the support of genetic data and using a holistic biological approach (Blasco-Costa et al., 2016a). Despite A. winterbourni showing subtle morphological differences from A. minutus, we consider them sufficient to distinguish the new species given the intrinsic difficulties of this group. Molecular results showed, convincingly, that specimens of A. winterbourni belong to a different microphallid genus to those already represented by sequence data. The new species appeared sister to an unidentified microphallid recovered from Posticobia brazieri (Smith) (Gastropoda, Tateidae) and Caridina indistincta Calman (Decapoda, Atyidae) in Australia by Kudlai et al. (2015). These authors suggested that their specimens were likely closely related to (or even conspecific) with the material reported as Microphallus sp. 'livelyi' by Hechinger (2012), which is considered a junior synonym of A. winterbourni herein. Divergence between our sequences and those of the Australian microphallid of Kudlai et al. (2015) suggests that they represent two distinct but congeneric species, as the authors anticipated.

The examination of specimens of A. winterbourni with SEM has allowed us to confirm a sinistral interruption of the outer rim of the ventral sucker caused by the opening of the genital pore and protrusion of the dextral parietal atrial scale, while light microscopy and histological sections show that, internally, the ventral sucker is complete. Deblock & Rosé (1964) mentioned the possibility of the scale being united to the rim of the ventral sucker (p. 229). However, this feature went unnoticed and was never confirmed by a later redescription or new species description for the genus, nor was it mentioned in the most recent diagnosis provided by Deblock (2008). Since the three species known for the genus so far are characterized by the presence of the large atrial scale, it is likely that in all three cases it has resulted in the same modification of the ventral sucker. Thus, we consider this feature of diagnostic value for the genus and amend herein the generic diagnosis of Atriophallophorus as follows.

Atriophallophorus Deblock & Rosé, 1964

Generic diagnosis

Body piriform or triangular, small (150–200 μm). Resembles *Microphallus*. Ventral sucker postequatorial, outer rim

interrupted sinistrally by genital pore. Oesophagus medium or short. Caeca short, divergent, in mid-body. Testes symmetrical in hindbody. Male genital pouch absent. Seminal vesicle intercaecal in mid-body, ovoid; prostatic gland caps distal part of seminal vesicle; proximal ejaculatory duct very long, supported by envelope acting as large prostatic chamber, either: (i) subcircular with diameter of ventral sucker, dorsal to genital atrium (formation described as 'phallophorus' (apparently bearing phallus)) and with long prostatic duct running through to periphery of male duct inside phallus; or (ii) cylindrical, fibrous, elongate between seminal vesicle and genital atrium, adjacent to margin of ventral sucker; bundle of long prostatic ducts enter as far as mid-part of cylindrical prostatic chamber (not inside phallus). Phallus of 'microphalloid'-type, more or less turgid, with ejaculatory duct in axis. Genital atrium present, envelopes phallus, with enormous dextral parietal atrial scale; genital pore sinistral to ventral sucker. Ovary dextral to ventral sucker. Uterus postcaecal, with few coils around testes. Metraterm long, with widened opening into sinistral wall of genital atrium. Eggs not numerous, relatively large. Vitellarium formed of two clusters of follicles, paratesticular and post-testicular in hindbody; vitelline ducts short, arched, post-testicular. Excretory vesicle short, Y-shaped, post-testicular. In intestine of birds (Anseriformes, Charadriiformes); cosmopolitan.

Type species. Atriophallophorus minutus (Price, 1934).

Synonym. Atriophallophorus samarae Deblock & Rosé, 1964.

Synonym lapsus calami. Atriophallus samarae Deblock & Rosé, 1964 (Fig. 1).

Life cycle and putative definitive hosts

Compared to other microphallid genera, the known species diversity of *Atriophalophorus* is quite low. So far, the four members of this genus (including the yet undescribed but molecularly characterized lineage from Australia of Kudlai *et al.* (2015)) show an abbreviated life cycle with the absence of the cercarial stage. Furthermore, *A. winterbourni* also lacks daughter sporocyst parthenitae-bearing germ cells (or has a very reduced life span), so that germ balls and embryonic metacercariae are observed free in the visceral mass of the snail host and appear to develop directly into encysted metacercariae, as reported by Krist & Lively (1998).

Based on the experimental exposure of snails to the faeces of different waterfowl species from Lake Alexandrina, Osnas & Lively (2011) concluded that the likely definitive hosts of *A. winterbourni* are mallard ducks (*Anas platyrhynchos* L.), grey ducks (*Anas superciliosa* Gmelin), their hybrids and the New Zealand scaup (*Aythya novaeseelandiae* Gmelin). Their conclusion agrees with the known distribution of both the putative hosts and *A. winterbourni* throughout the South Island. Furthermore, recent studies of the parasite fauna of mallards have discovered new species to science of microphallids and strigeids in New Zealand (Presswell *et al.*, 2014; Blasco-Costa *et al.*, 2016b). Altogether, these results highlight the still scarce knowledge on the parasites of birds, the most diverse group of vertebrates native to New Zealand, and the need for more biodiversity studies to address this gap.

Supplementary material. To view supplementary material for this article, please visit https://doi.org/10.1017/S0022149X19000993.

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Conflicts of interest. None.

Ethical standards. The authors assert that all procedures contributing to this work comply with the ethical standards of Switzerland and New Zealand and our institutional guides on the care and use of wild invertebrate animals

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