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Research article

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Integrative redescription of *Hypsibius pallidoides* Pilato *et al.*, 2011 (Eutardigrada: Hypsibioidea) with the erection of a new genus and discussion on the phylogeny of Hypsibiidae

Denis V. TUMANOV 1,2

¹Department of Invertebrate Zoology, Faculty of Biology, Saint-Petersburg State University, 199034, Universitetskaya nab. 7/9, Saint Petersburg, Russia.

²The Laboratory of the Marine Research, Zoological Institute of Russian Academy of Sciences, 199034, Universitetskaja nab. 1, Saint Petersburg, Russia.

Email: d.tumanov@spbu.ru

urn:lsid:zoobank.org:author:49E22A70-C27B-485B-941E-577666EE65F9

Abstract. An integrative redescription of *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011 was undertaken following a reexamination of the type material and new material using high-quality light microscopy, scanning electron microscopy and methods of molecular taxonomy. Detailed morphological investigations revealed a unique complex of characters that precluded the attribution of this species to the genus *Hypsibius* Ehrenberg, 1848. Furthermore, phylogenetic analyses indicated the affinity of this species within the subfamily Pilatobiinae (Hypsibiidae). *Notahypsibius* gen. nov. is erected for *H. pallidoides* and two putatively related species: *H. scaber* Maucci, 1987 and *Ramazzottius arcticus* (Murray, 1907). An emended diagnosis for the genus *Pilatobius* is given, while the subfamily Pilatobiinae lacks a cohesive morphological diagnosis despite representing, at the same time, a well-supported molecular clade. Obvious controversy between the results of the morphological and molecular analyses of the phylogeny of Hypsibioidea is discussed. The distribution of morphological characters such as the claw type, organization of the bucco-pharyngeal apparatus, and egg shell sculpture type within Eutardigrada is analyzed and their phylogenetic significance discussed.

Keywords. Tardigrada, morphology, trait evolution, Pilatobiinae, molecular taxonomy.

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Introduction

Phylum Tardigrada Doyère, 1840 is a group of widely distributed microscopical multicellular animals. Currently there are ca 1300 known species (Degma *et al.* 2019), but this is very likely an underrepresentation of the actual number of taxa, as the global diversity of tardigrades is considered poorly investigated (Bartels *et al.* 2016). Along with the description of new taxa, the redescription of known species using the integrative approach, i.e., combining a morphological analysis with methods of molecular taxonomy and phylogeny, is a promising way to improve our understanding of the real

taxonomical richness of this group (Bertolani *et al.* 2011a; Gąsiorek *et al.* 2016, 2018; Kaczmarek *et al.* 2018; Stec *et al.* 2018; Guidetti *et al.* 2019a). Molecular methods can also help to infer phylogenetic relationships within Tardigrada when a morphological analysis alone is insufficient due to the paucity of the morphological variation and the prevalence of evolutionary parallelism in taxonomically important structures (Kiehl *et al.* 2007; Sands *et al.* 2008; Bertolani *et al.* 2014; Cesari *et al.* 2016; Guil *et al.* 2019; Gąsiorek *et al.* 2019a, 2019b, 2019c).

Hypsibius Ehrenberg, 1848 is the type genus, and, with 40 described species (Degma et al. 2019, with corrections according to Dastych 2019), also the largest genus of the family Hypsibiidae Pilato, 1969. Intrageneric morphological heterogeneity and phylogenetic clues both suggest a polyphyletic nature of this taxon (Guil & Giribet 2012; Gąsiorek et al. 2018). However, a molecular phylogenetic analysis of most of the morphologically divergent forms previously revealed within the genus Hypsybius sensu lato (i.e., genera Borealibius Pilato, Guidetti, Rebecchi, Lisi, Hansen & Bertolani, 2006, Cryobiotus Dastych, 2019, and complex of species similar to H. scabropygus Cuénot, 1929) demonstrated their close affinity to the 'typical' species of Hypsibius, so the subfamily Hypsibiinae Pilato, 1969 seems to be monophyletic (Bertolani et al. 2014; Gąsiorek et al. 2018). Previously, only the genera Acutuncus Pilato & Binda, 1997 and Mixibius Pilato, 1992, both separated from the genus Hypsibius on the base of a morphological analysis (Pilato & Binda 1997; Pilato 1992), were demonstrated as lineages phylogenetically distant from the subfamily Hypsibiinae (Kiehl et al. 2007; Sands et al. 2008; Marley et al. 2011).

In 2016, I found a single adult tardigrade specimen and an exuvium containing eggs from a moss sample collected in St Petersburg, Russia, and an additional adult specimen from a moss sample collected in Golubinjak Forest Park, Croatia. I attributed both specimens preliminarily to the species *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011, which was described based on hypsibiid individuals from Kherson Oblast, South Ukraine (Pilato *et al.* 2011). In 2018, I obtained numerous specimens of *H. pallidoides* from moss collected in Carinthia, Austria. This finding made it possible to establish a laboratory culture of the species to be used for morphological and molecular analyses. I also reviewed some specimens in my collection that had previously been identified as *Hypsibius pallidus* Thulin, 1911 and found that these were actually attributable to *H. pallidoides*.

The initial investigations of the above mentioned material of *H. pallidoides* revealed some subtle differences to the original species description. This necessitated an examination of the type series. During my visit to Catania in August 2019, I had the opportunity to compare my specimens with the holotype and paratypes of *H. pallidoides* in the Binda and Pilato collection (Museum of the Department of Animal Biology "Marcello La Greca", University of Catania, Italy). Further comparisons were made with paratypes obtained from Y. Kiosya (Kharkiv, Ukraine). Studies of the type material and additional specimens revealed some inaccuracies in the original description of *H. pallidoides*. In this paper, I give a redescription of this species based on the type material and on my own specimens. Scanning Electron Microscopy (SEM) investigation provided an opportunity to explore details of the structures of this species which were previously indiscernible with the use of only Light Microscopy (LM). Sequencing and analyses of the phylogenetically significant genes (COI, 18S rRNA, 28S rRNA and ITS-2) led to the clarification of the phylogenetic position of *H. pallidoides* and to the institution of a new genus.

Material and methods

Sampling and culturing

Tardigrades were extracted from rehydrated samples using a standard technique of washing them through two sieves (Tumanov 2018). The content of the fine sieve was examined under a Leica M205C stereo microscope.

A laboratory culture of *H. pallidoides* was established from several living specimens extracted from a sample collected in Austria. Animals were kept in plastic Petri dishes with a mixture of distilled and filtered tap water (3:1) and *Chlorella* sp., a unicellular freshwater alga (received from Core Facilities Center "Culture Collection of Microorganisms" of St Petersburg State University). To aid tardigrade locomotion, the Petri dish bottom was scratched with fine sandpaper as recommended by Kosztyła *et al.* (2016). The culture was maintained at 16°C.

Microscopy and imaging

The tardigrades found in moss samples or acquired from the laboratory culture were fixed with acetic acid or relaxed by incubating live individuals at 60°C for 30 min (Morek *et al.* 2016) and mounted on slides in Hoyer's medium. Permanent slides were examined under a Leica DM2500 microscope equipped with phase contrast (PhC) and differential interference contrast (DIC). Photographs were made using a Nikon DS-Fi3 digital camera with NIS software.

For scanning electron microscopy (SEM) specimens were thermally relaxed at 60°C (Morek *et al.* 2016), dehydrated in an ascending ethyl alcohol series (10%, 20%, 30%, 50%, 70%, 96%) and acetone, critical-point dried in CO₂, mounted on stubs and coated with gold. The bucco-pharyngeal apparatus was prepared for SEM investigation following the protocol of Eibye-Jacobsen (2001) as modified by Gąsiorek *et al.* (2016). A Tescan MIRA3 LMU Scanning Electron Microscope was used for observations (Centre for Molecular and Cell Technologies, St Petersburg State University).

Morphometrics

The sample size for morphometrics was chosen following the recommendations of Stec *et al.* (2016a). All measurements are given in micrometres (µm). Structures were measured only if their orientations were suitable. Body length was measured from the anterior end of the body to the posterior end, excluding the hind legs. The bucco-pharyngeal tube was measured from the anterior margin of the stylet sheaths to the caudal end of the buccal tube, not including the buccal apophyses. Terminology for the structures within the bucco-pharyngeal apparatus and for the claws follows that of Pilato & Binda (2010). Elements of the buccal apparatus were measured according to Kaczmarek & Michalczyk (2017). Claws were measured following Beasley *et al.* (2008), but the total length of the claws was also measured (according to Pilato *et al.* 2002) to maintain compatibility with the initial description. The *pt* index used is the percentage ratio between the length of a structure and the length of the buccal tube (Pilato 1981) and is presented here in italics. Morphometric data were handled using ver. 1.6 of the "Parachela" template, which is available from the Tardigrada Register (Michalczyk & Kaczmarek 2013) with total length of the claws added.

Genotyping

DNA was extracted from 15 individual animals using QuickExtractTM DNA Extraction Solution (Lucigen Corporation, USA) using the following protocol (kindly provided by Torbjørn Ekrem, Norwegian University of Science and Technology).

- 1) Tardigrades were sorted in water and specimens were rinsed individually in ddH₂O.
- 2) Each individual specimen was transferred by pipette into a PCR-tube containing 70 μl QuickExtractTM.
- 3) PCR-tubes were vortexed well, spun down (5 min at 3500 RPM), then kept at room temperature ($\approx 25^{\circ}$ C) for 2 hrs.
- 4) PCR-tubes were incubated at 65°C for 15 min (in a PCR-machine), vortexed every 5 min and spun down.
- 5) PCR-tubes were incubated at 98°C for 2 min.

- 6) 60 μl of the extract supernatant were transferred into a new, sterile PCR tube. The supernatant was collected in order to avoid the exoskeleton remaining at the bottom. The PCR-tubes containing extract were then stored at -20°C for later use in PCR.
- 7) 70 µl ddH₂O were added to the tube with the exuvium and mixed well with the pipette to wash the exoskeleton.
- 8) Water and exoskeleton were transferred to a glass staining block with ddH₂O. The exoskeleton was collected and mounted on a microscope slide in Hoyer's medium and retained as the hologenophore (Pleijel *et al.* 2008).

Four genes were sequenced: a small ribosome subunit (18S rRNA) gene, a large ribosome subunit (28S rRNA) gene, internal transcribed spacer (ITS-2), and the cytochrome oxidase subunit I (COI) gene. Both 18S rRNA and 28S rRNA are nuclear markers used in phylogenetic analyses to investigate high taxonomic levels (Jørgensen et al. 2010, 2011; Guil & Giribet 2012; Bertolani et al. 2014; Guil et al. 2019; Gasiorek et al. 2019b, 2019c). COI is a protein-coding mitochondrial marker that is widely used as a standard barcode gene of intermediate to high effective mutation rate (Bertolani et al. 2011b). ITS-2 is a non-coding nuclear fragment with high evolution rates used for both intra-specific comparisons and comparisons between closely related species (Gasiorek et al. 2016, 2018; Stec et al. 2018). A complete 18S rRNA gene was amplified in several overlapping fragments using primer pairs: SSU F 04 and SSU R 26, 18Sfw and rev960, fw390 and rev18S, 5F and 9R (for primer details see Table 1). These products were sequenced with PCR primers and the internal primers fw1230 and rev1460. A fragment of the 28S rRNA gene was amplified and sequenced using primers 28 F0001 and 28S R1800. PCR reactions included 2 µl template DNA, 1 µl of each primer, 1 µl DNTP, 5 µl Taq Buffer (10X) (-Mg), 4 μl 25 mM MgCl₂ and 0.2 μl Taq DNA Polymerase (Thermo ScientificTM) in a final volume of 50 μl. The PCR cycling profile for the 18S and 28S genes was as follows: initial denaturation at 95°C for 5 min, then 35 cycles of 95°C for 1 min, 50°C for 1 min, 72°C for 2 min, and final elongation at 72°C for 10 min. ITS-2 was amplified and sequenced using primers ITS2 Eutar Ff and ITS2 Eutar Rr (Stec et al. 2018). The PCR cycling profile for ITS-2 was as follows: initial denaturation at 95°C for 3 min, followed by 30 cycles of denaturation at 95°C for 1 min, annealing at 55°C for 2 min, and elongation at 72°C for 2 min, and a final elongation step at 72°C lasting 10 min. A fragment of the COI mtDNA gene was amplified and sequenced using primers LCO1490 and HCO2198 (Folmer et al. 1994). The PCR cycling profile for the COI gene was as follows: initial denaturation at 94°C for 5 min, followed by five cycles of denaturation 1 min at 94°C, annealing at 42°C for 1.5 min and amplification at 72°C for 1.5 min; then 35 cycles of 94°C for 1 min, 50°C for 1.5 min, 72°C for 1 min, and final elongation at 72°C for 5 min. COI sequences were translated to amino acids by using the invertebrate mitochondrial code implemented in MEGA7 (Kumar et al. 2016) in order to check for the presence of stop codons and therefore of pseudogenes.

PCR products were visualized in 1.5% agarose gel stained with Ethidium bromide. All amplicons were sequenced directly using ABI PRISM Big Dye Terminator Cycle Sequencing Kit (Applied Biosystems, Foster City, CA, USA) on ABI Prism 310 Genetic Analyzer. Sequences were edited and assembled using ChromasPro software (Technelysium, USA).

Phylogenetic analyses

All sequences of Hypsibioidea Guil, Jørgensen & Kristensen, 2019 of appropriate length available in GenBank at the time of the analysis were downloaded and those originating from published works with reliable attribution of the investigated taxa were selected (see Supplementary file SM.01). Because of the low number of species of Hypsibioidea where both 18S and 28S sequences are available, these genes were analysed separately, also the analysis of concatenated 18S+28S sequences alignment was performed. Four species of *Macrobiotus C.A.S.* Schultze, 1834 (Macrobiotoidea Guil, Jørgensen & Kristensen, 2019) were used as outgroup. First, sequences were automatically aligned using the Muscle

Table 1. Primers used for amplification of the four DNA fragments sequenced in the study.

DNA fragment	Primer name	Primer direction	Primer sequence (5'-3')	Primer source
18S rRNA	SSU_F_04	forward	GCTTGTCTCAAAGATTAAGCC	Kiehl et al. 2007
	SSU_R_26	reverse	CGAAAGCATTTGCCAAGAATG	Kiehl et al. 2007
	18Sfw	forward	CTTGTCTCAAAGATTAAGCCATGCA	Dabert et al. 2014
	rev960	reverse	GACGGTCCAAGAATTTCAC	Dabert et al. 2014
	fw390	forward	AATCAGGGTTCGATTCCGGAGA	Dabert et al. 2014
	rew18S	reverse	TGATCCTTCCGCAGGTTCACCT	Dabert et al. 2014
	5F	forward	GCGAAAGCATTTGCCAAGAA	Giribet et al. 1996
	9R	reverse	GATCCTTCCGCAGGTTCACCTAC	Giribet et al. 1996
	fw1230	forward	TGAAACTTAAAGGAATTGACG	Dabert et al. 2014
	rev1460	reverse	CATCACAGACCTGTTATTGC	Dabert et al. 2014
28S rRNA	28SF0001	forward	ACCCVCYNAATTTAAGCATAT	Dabert et al. 2014
	28SR1800	reverse	GTTCACATGGAACCCTTCT	Dabert et al. 2014
COI	LCO1490	forward	GGTCAACAAATCATAAAGATATTGG	Folmer et al. 1994
	HCO2198	reverse	TAAACTTCAGGGTGACCAAAAAATCA	Folmer et al. 1994
ITS-2	ITS2_Eutar_Ff	forward	CGTAACGTGAATTGCAGGAC	Stec et al. 2018
	ITS2_Eutar_Rr	reverse	TGATATGCTTAAGTTCAGCGG	Stec et al. 2018

algorithm (Edgar 2004) as implemented in SeaView 4.0 (Gouy *et al.* 2010); the alignment was later refined manually. Final align lengths were 1685 bp for 18S gene and 2207 bp for 28S gene. Best fitting model evaluations for each analysis were performed using jModeltest 2.1.10 (Darriba *et al.* 2012) resulting in the GTR+Gamma+I model to be most suitable for all the datasets.

Maximum-likelihood (ML) topologies were constructed using the RaxML 8.2.10 program (Stamatakis 2014) with GTR+ γ +I model; the number of invariant sites, alpha parameter and tree topology were optimized by RAxML, 1000 bootstrap pseudoreplicates were used. Bayesian analysis of the same datasets was performed using MrBayes ver. 3.2.6, GTR model with gamma correction for intersite rate variation (8 categories) and the covarion model (Ronquist & Huelsenbeck 2003). The analyses were run as two separate chains (default heating parameters) for 10 million generations, by which time they had ceased converging (final average standard deviation of the split frequencies was less than 0.01). The quality of chains was estimated using built-in MrBayes tools. jModeltest, RaxML and MrBayes programs were run at the Cipres ver. 3.3 web-site (Miller *et al.* 2010). Uncorrected pairwise distances were calculated using MEGA7 (Kumar *et al.* 2016) with gaps/missing data treatment set to "complete deletion".

Institutional acronyms

Specimens from the following institutions and collections were examined (curator in parenthesis).

KNU = V.N. Karazin Kharkiv National University, Ukraine, School of Biology (Yevgen Kiosya)

MCVR = Museo Civico di Storia Naturale, Verona, Italy (Roberta Salmaso)

SPbU = St Petersburg University, Russia, Faculty of Biology, Department of Invertebrate Zoology

(Denis Tumanov)

UNICT = Università degli Studi di Catania, Italy, Museum of the Department of Animal Biology

"Marcello La Greca", Binda and Pilato collection (Giovanni Pilato)

Results

Redescription of Hypsibius pallidoides

Phylum Tardigrada Doyère, 1840 Class Eutardigrada Richters, 1926 Order Hypsibioidea Guil, Jørgensen & Kristensen, 2019 Family Hypsibiidae Pilato, 1969

Hypsibius pallidoides Pilato, Kiosya, Lisi, Inshina & Biserov, 2011 Figs 1–7

Material examined

Holotype

UKRAINE • Kherson Oblast, Ivano-Rybalchansky district of Chernomorsky biosphere reserve; 46°27′25″ N, 32°8′56″ E; Jun. 2008; D.A. Korolesova leg.; moss on wood; UNICT 5430.

Paratypes

UKRAINE • 1 spec. + 1 exuvium with eggs; same collection data as for holotype; UNICT 5430 • 4 specs + 2 exuviae with eggs; same collection data as for holotype; KNU 4ep-9 II.

Other material

AUSTRIA • 78 specs + 35 exuvia with eggs; Carinthia; 46.817818° N, 13.859837° E; 20 Aug. 2017; A. Smirnov leg.; moss on soil; GenBank: MK973069, MN912103, MK967961 to MK967964, MN927181, MN927182, MN919385, MN915220, MN915221, MK967241, MN918533; SPbU 251(1–13), 251(28).

CROATIA • 1 spec.; Park Šuma Golubinjak [Golubinjak Forest Park], Primorje-Gorski Kotar County; 45.35216° N, 14.76557° E; 10 Sep. 2005; O. Orlova leg.; moss on stone; SPbU 228(30).

RUSSIA – **St Petersburg** • 1 spec. + 1 exuvium; Puskin City; 59.72537° N, 30.39147° E; 15 May 2016; D. Tumanov leg.; moss on tree trunk; SPbU 234(10). – **Karelia** • 3 specs + 1 exuvium; vicinity of Akkaharju village; 61.49584° N, 29.84775° E; 11 May 1994; D. Tumanov leg.; mosses and leaf litter from the overgrown lake; SPbU 113(2).

Morphological redescription

MEASUREMENTS. Body elongated, of uniform width on the entire body length (Fig. 1), with a blunt snout (morphometrics Tables 2–3).

COLOUR. Body uncoloured or whitish with green gut content. Most specimens with eyespots, usually well-discernible after slide mounting (Fig. 1A) but absent in some specimens.

CUTICULAR SCULPTURE. Dorsal cuticle sculpture consists of a system of transverse folds with smaller irregular folds between (Figs 1B, 2A–D). Cuticle sculpture better visible in the caudal region of the body

(Figs 2C–D, 3A–B), well developed even in juveniles (Fig. 4E). Ventral surface with poorly developed foldings, visible in SEM only (Fig. 1C).

CEPHALIC SENSORY STRUCTURES. Cephalic body portion with a pair of elliptical sensory organs developed in the form of flat porous areas, separated from the body surface with a oval cuticular groove. These structures are scarcely visible in LM, but are well-discernible in SEM (Fig. 3C–D, black arrowheads). Two indistinctly demarcated porous areas are also developed in the fronto-lateral region of the head, on the each side of the mouth opening (visible in SEM only; Fig. 4A–B, white arrowheads). Central concavity on the dorsal surface of the head (Fig. 3C, white arrowhead) seems to be similar to the structures present in some Isohypsibioidea (see Gąsiorek *et al.* (2019c: 91, fig. 4b, d) and in *Cryobiotus roswithae* (Dastych 2019).

MOUTH. Opening antero-ventral, surrounded by six peribuccal lobes (visible in SEM only; Fig. 4C). In large specimens a line of elliptical structures is visible in LM around the mouth opening (Fig. 4G–H, black arrowheads), similar to those described for *Acutuncus antarcticus* (Richters, 1904) and *Hypsibius murrayi* (Richters, 1907) (Dastych 1991, 2018). These structures are possibly the compressed peribuccal lobes.

Bucco-pharyngeal apparatus. Hypsibiinae model sensu Pilato & Binda 2010 (Fig. 4F). Oral cavity armature with a ring of small teeth located it its anterior part followed by the second band of larger,

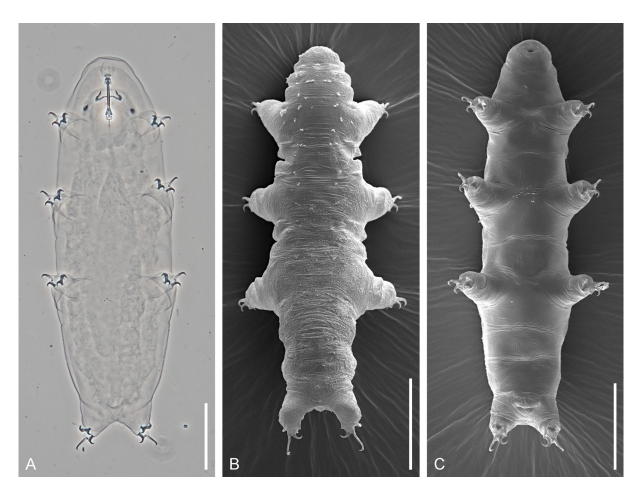


Fig. 1. *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011, total view. **A.** Phase contrast (SPbU 251(82)). **B.** Dorsal view in SEM. **C.** Ventral view in SEM. Scale bars = $50 \mu m$.

irregular teeth (visible in SEM only; Fig. 4D). Dorsal and ventral apophyses for the insertion of the stylet muscles (AISM) are evidently dissimilar. Dorsal AISM are shorter and higher than ventral, with thickened anterior margin (Fig. 5A–C). A short thickening of the buccal tube wall is present posteriorly to both these apophyses (the ventral poorly visible; Fig. 5A, black arrowheads). Buccal tube rigid, bent ventrally in its caudal part (Fig. 5A). Stylet furcae typically shaped (*sensu* Pilato & Binda 2010) (Fig. 4F). Pharyngeal bulb spherical (Fig. 5D, black arrowhead), with well-developed apophyses, two elongated macroplacoids, and a small dot-like structure interpreted here as a septulum (following Pilato *et al.* 2011) (Fig. 5A, white arrow), connected to the second macroplacoid with a thin cuticular line (often scarcely visible) (Fig. 5D, black arrow). No microplacoids. Posteriorly to the septulum, an indistinct thickening of the cuticular lining similar to "pseudoseptulum" described in *Diphascon mirabilis* Dastych, 1984 and *Hypsibius iskandarovi* Tumanov, 1997 is present (Dastych 1984; Tumanov 1997) (Fig. 5D, white arrowhead). First macroplacoid longer than second with a slight constriction in the middle (Fig. 5E–F, black arrowhead). Second macroplacoid can also have a poorly developed subterminal constriction (visible in SEM only) (Fig. 5F, black arrow).

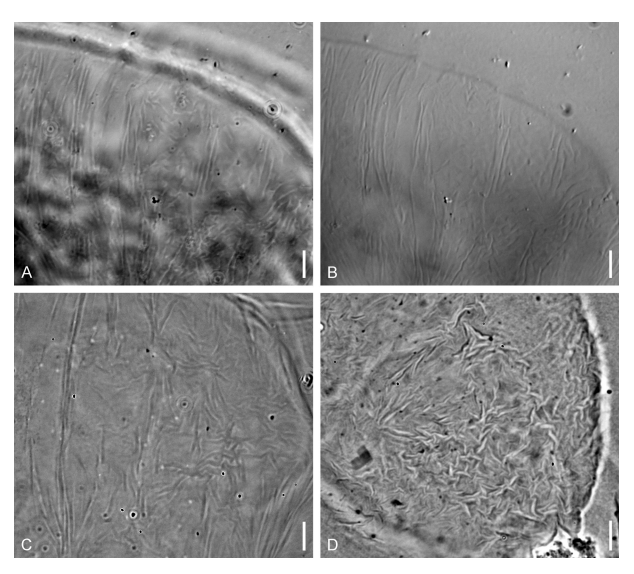


Fig. 2. Hypsibius pallidoides Pilato, Kiosya, Lisi, Inshina & Biserov, 2011, dorsal sculpture. **A.** Type series specimen (KNU Чер-9 II), PhC. **B.** Type series specimen (KNU Чер-9 II), DIC. **C.** Specimen from Austria (SPbU 251(82)), PhC. **D.** Specimen from Russia (SPbU 113(3)). Scale bars = 5 μm.

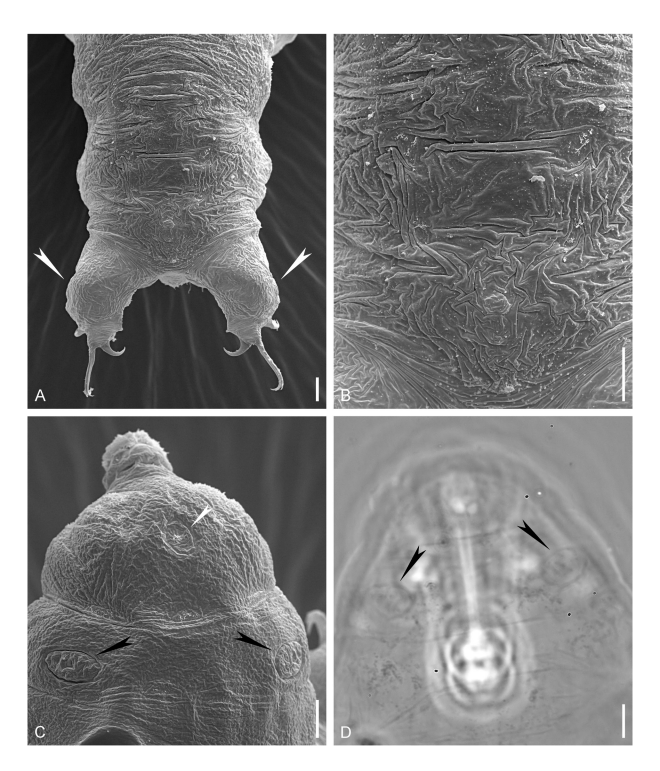


Fig. 3. *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011. **A.** Dorsal view of the caudal body end, white arrowheads indicate the dorsal inflations of the hind legs, SEM. **B.** Dorsal sculpture of the caudal body region, SEM. **C.** Dorsal surface of the head region, black arrowheads indicate the elliptical sensory organs, white arrowhead indicates the central concavity on the dorsal surface of the head, SEM. **D.** Dorsal surface of the head region, black arrowheads indicate the elliptical sensory organs (SPbU 251(89)), PhC. Scale bars = 5 μ m.

Table 2 (continued on the next page). Summary of morphometric data for *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011 (type series specimens (n = 4) and additional specimens from Croatia and Russia). Measurements are given in μm , pt values in % (the pt index is the percentage ratio between the length of a structure and the length of the buccal tube).

	type s	eries 1	type	series 2	type series 3		type series 4		Croatia		Russia 1		Russia 2	
CHARACTER	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt
Body length	258	985	235	957	201	951			220	895	288	1194	247	1030
Buccopharyngeal tube														
Buccal tube length	26.1	_	24.6	_	21.2	_	26.4	-	24.5	_	24.1	-	24.0	_
Stylet support insertion point	14.7	56.2	14.3	58.0	12.2	57.6	15.5	58.6	14.7	59.8	13.7	56.8	14.0	58.4
Buccal tube external width	1.8	6.9	1.8	7.2	1.5	7.1	1.9	7.1	1.8	7.3	2.0	8.4	1.7	7.1
Buccal tube internal width	1.0	3.7	0.8	3.4	0.6	2.8	0.9	3.4	0.9	3.5	1.1	4.4	0.7	3.1
Placoid lengths														
Macroplacoid 1	3.5	13.3	2.9	11.8	2.2	10.6	3.1	11.6	2.5	10.2	3.4	14.0	2.5	10.5
Macroplacoid 2	2.3	8.9	2.1	8.6	2.0	9.2	2.3	8.6	2.2	9.1	2.6	10.7	2.3	9.5
Macroplacoid row	6.6	25.1	5.7	23.2	4.7	22.1	6.2	23.6	5.6	22.9	6.5	26.8	5.8	24.2
Placoid row	8.8	33.7	8.0	32.6	6.6	31.0	8.5	32.0	7.7	31.4	8.8	36.2	8.2	34.1
Claw 1 heights														
External base	_	-	4.6	18.5	3.2	15.2	4.6	17.4	3.9	15.9	4.4	18.0	4.3	18.0
External primary branch	8.6	33.0	8.9	36.2	6.9	32.6	8.1	30.7	7.8	31.8	9.4	38.9	7.8	32.4
External secondary branch	4.5	17.3	4.2	17.1	4.1	19.5	5.0	19.1	4.4	18.0	4.3	17.8	4.6	19.0
External total	12.9	49.2	13.1	53.3	_	_	12.6	47.9	12.0	48.8	14.1	58.5	11.9	49.4
Internal base	3.6	13.9	3.9	16.0		_	3.4	12.7	_	_	4.4	18.1	3.5	14.4
Internal primary branch	4.7	18.1	5.7	23.0	_	_	5.1	19.2	_	_	4.9	20.1	4.7	19.7
Internal secondary branch	3.2	12.2			_	_	3.4	12.7	_	_	3.8	15.7	3.4	14.0
Internal total	8.7	33.1	8.8	35.6	_	_	8.4	31.7	_	_	8.3	34.2	7.8	32.4
Claw 2 heights														
External base	5.1	19.5	5.6	23.0	3.4	15.9	4.5	16.9	4.2	17.0	5.1	21.2	4.9	20.5
External primary branch	8.9	34.1	9.8	40.0	7.7	36.3	8.5	32.0	7.7	31.6	9.8	40.4	8.6	35.7
External secondary branch	4.7	17.9	5.4	22.0	3.7	17.7	4.9	18.5	4.4	18.0	4.5	18.8	4.7	19.5
External total	12.8	48.9	15.4	62.8	_	_	12.9	48.9	12.2	49.8	15.4	63.6	13.3	55.4
Internal base	4.8	18.2	_	_	_	_	4.1	15.6	3.7	15.0	4.5	18.8	4.3	17.7
Internal primary branch	4.2	16.0	_	_	_	_	4.7	17.9	4.4	17.9	5.7	23.7	4.5	18.7
Internal secondary branch	_	_	_	_	_	_	4.1	15.6	3.5	14.4	4.2	17.5	3.6	14.9
Internal total	_	_	_	_	_	_	8.5	32.3	7.9	32.2	9.3	38.4	8.2	34.0
Claw 3 heights														
External base	5.0	19.2	5.3	21.6	3.1	14.5	5.3	20.1	4.7	19.0	4.7	19.3	4.9	20.2
External primary branch	8.7	33.1	9.6	39.2	7.9	37.2	8.2	31.0	7.6	30.8	9.7	40.3	8.6	35.6
External secondary branch	4.3	16.5	5.9	24.2	3.8	17.9	4.9	18.5	4.4	17.8	5.2	21.7	4.8	20.0
External total	13.1	50.1	14.4	58.7	_	_	13.3	50.3	12.2	49.9	14.3	59.4	13.2	54.9
Internal base	4.2	16.0	_	_	_	_	_	_	_	_	4.8	19.8	_	_
Internal primary branch	-	-	_	_	_	_	_	_	_	_	4.9	20.4	5.2	21.8
Internal secondary branch	4.1	15.8	_	_	_	_	_	_	_	_	4.4	18.2	3.7	15.5
Internal total	_	-	_	_	_	_	8.5	32.1	_	_	9.8	40.7	8.5	35.3
Claw 4 lengths											0			
Anterior base	4.0	15.4	_		_		_		3.4	14.0			3.8	15.7
Anterior primary branch	5.4	20.6	_	_	_	_	_	_	4.8	19.6	5.8	24.0	5.4	22.4

Table 2 (continued).

	type s	type series 1 typ		pe series 2 type		series 3	ries 3 type series 4		Croatia		Russia 1		Rus	sia 2
CHARACTER	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt	μm	pt
Anterior secondary branch	-	_	_	_	-	-	_	_	3.5	14.3	4.9	20.2	4.8	20.0
Anterior total	9.0	34.4	-	-	_	_	_	_	8.0	32.6	10.3	42.7	9.3	38.5
Posterior base	_	_	_	_	_	_	_	_	4.7	19.2	5.0	20.7	5.4	22.6
Posterior primary branch	_	_	_	_	_	_	_	_	10.7	43.6	11.4	47.3	12.1	50.4
Posterior secondary branch	_	_	_	_	_	_	_	_	4.7	19.3	5.8	23.9	5.9	24.6
Posterior total	13.9	53.3	_	_	11.1	52.5	_	_	15.8	64.5	16.6	68.8	17.8	74.3

LEGS AND CLAWS. All legs with well-developed claws, increasing in size from legs I to legs IV (Fig. 6A–I). Legs IV evidently swollen dorsally, above the claws (Fig. 3A, white arrowheads). Claws similar to the Ramazzottius-type claws (sensu Pilato & Binda 2010 and Guidetti et al. 2019b) with external and internal claws of each leg strongly dissimilar. External claws with massive base+secondary branch complex, where the base is at least as long as the secondary branch and only slightly curved, while the secondary branch is thinner than the base and connected with it at a nearly right angle without forming a smooth ark (Fig. 6A–B, I). Thin and long primary branch connected to the base+secondary branch complex far from the claw's basal point (length of the claw base is equal or slightly exceeds the length of the secondary branch). The connection point is shifted laterally and located near the evident crest developed on the lateral surface of the claw base (Fig. 6G, I, white arrowheads). Basal part of the primary branch flexible, with thinned walls (Fig. 6A-B, black arrowheads). External and posterior claws of *H. pallidoides* differ from typical *Ramazzottius*-type claws only in having primary branches wider with less pronounced differentiation between rigid distal and soft basal parts. Primary branches are connected with the base by a filamentous structure (not always visible in LM, Fig. 6D, black arrow), but no distinct light-refracting unit is present. Internal claws much shorter than external ones, without flexible parts, with developed internal structure, consisting of the system of cavities and septae (Fig. 6A, D-E). All claws with developed accessory points and widened smooth bases (Fig. 6A-I). Claws of legs I–III with very poorly developed smooth lunules (or pseudolunules, according Gasiorek et al. 2017) (Fig. 6A, white arrow), usually not discernible on the external claws. Claws of legs IV with welldeveloped wide lunules (Fig. 6E, I). Posterior claws with thickened region on the lunule margin, visible in LM as a dark line, which can create the impression of the presence of a cuticular bar between the bases of the anterior and posterior claws (Fig. 6E-F, black arrows). Legs I-III without cuticular bars near the claw bases, but with an elongated bulge located near the base of the internal claw. In SEM, the pulvinus is similar in appearance to the typical cuticular bar of Hypsibioidea, but in LM no zone of thickened cuticle is visible (Fig. 6C, H, white arrows). Also, poorly developed pulvini are visible on the inner side of the legs (Fig. 6C, H, white asterisks)

EGGs. One to six white subspherical eggs are laid in the exuvium (Fig. 7A), 59.4–71.9 µm in diameter (65.96 ± 3.71; N = 20). Egg shell in LM appears sculptured with minuscule granules, visible only with PhC or DIC in high magnification (Fig. 7B–C, E–F). In fact, these granules are inner pillar-like structures in the egg shell (Fig. 7D).

DNA sequences

Sequences of good quality for the 4 aforementioned molecular markers were obtained from five specimens: 2 paragenophores and 3 hologenophores (voucher specimens 251(09), 251(10) and 251(87)). Each gene was represented by single haplotype.

COI sequence (GenBank: MK967241), 688 bp long.

18S rRNA sequence (GenBank: MN912103), 1777 bp long:

28S rRNA sequence (GenBank: MK967961), 1618 bp long:

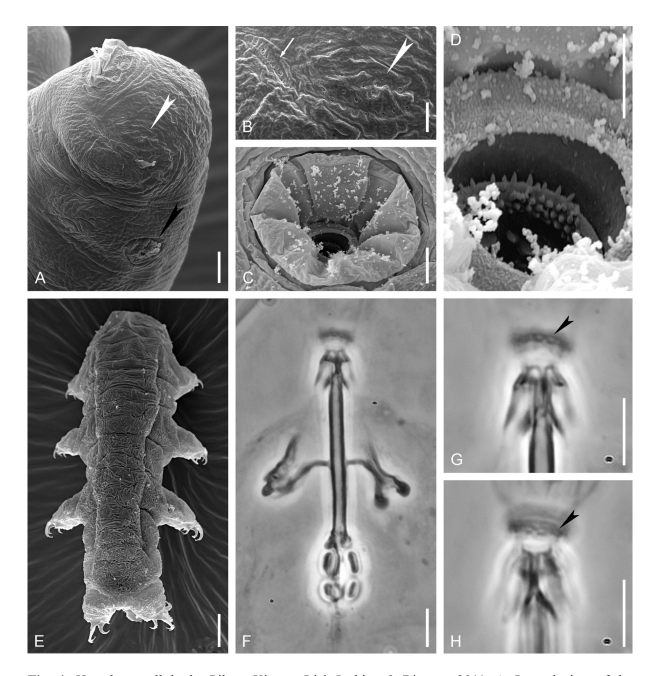


Fig. 4. *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011. **A.** Lateral view of the head region, white arrowhead indicates the anterior porous area, black arrowhead indicates the elliptical sensory organ, SEM. **B.** Enlarged view of the lateral surface of the head, white arrowhead indicates the anterior porous area, white arrow indicates the muscle attachment zone, SEM. **C.** Mouth opening with peribuccal lobes, SEM. **D.** Mouth opening with anterior ring of teeth, SEM. **E.** Dorsal sculpture of the juvenile, SEM. **F.** Bucco-pharyngeal apparatus (SPbU 251(82)), PhC. **G.** Dorsal view of the buccal cavity, black arrowhead indicates the circumoral elliptical structures (SPbU 251(82)), PhC. **H.** Ventral view of the buccal cavity, black arrowhead indicates the circumoral elliptical structures (SPbU 251(82)), PhC. Scale bars: A, F–H = 5 μm; B–C = 2 μm; D = 1 μm; E = 10 μm.

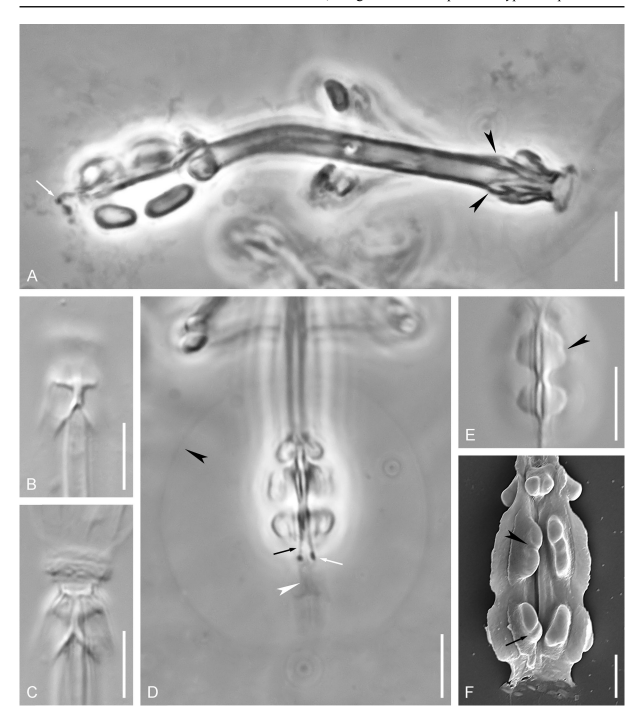


Fig. 5. *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011, bucco-pharyngeal apparatus. **A.** Lateral view of the bucco-pharyngeal apparatus, black arrowheads indicate the thickenings of the buccal tube, white arrow indicates the septulum (SPbU 251(109)), PhC. **B.** Dorsal AISM (SPbU 251(82)), DIC. **C.** Ventral AISM (SPbU 251(82)), DIC. **D.** Pharyngeal bulb, black arrowhead indicates the pharynx outline, black arrow indicates the cuticular line connecting second macroplacoid with septulum, white arrow indicates the septulum, white arrowhead indicates the "pseudoseptulum" (SPbU 251(89)), PhC. **E.** Macroplacoids, black arrowhead indicates the constriction of the first macroplacoid (SPbU 251(89)), DIC. **F.** Cuticular structures of the pharynx, black arrowhead indicates the constriction of the first macroplacoid, black arrow indicates the constriction of the second macroplacoid, SEM. Scale bars: $A-E=5 \mu m$; $F=2 \mu m$.

Table 3 (continued on the next page). Summary of morphometric data for *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011 (Austrian population). Measurements are given in μm , pt values in % (the pt index is the percentage ratio between the length of a structure and the length of the buccal tube).

CHARACTER	N			RA		MI	EAN	SD			
CHARACTER	11	μт				pt		μm	pt	μm	pt
Body length	30	132	-	292	695	-	1223	240	1016	39	120
Buccopharyngeal tube											
Buccal tube length	30	19.0	_	26.1		-		23.5	_	1.7	_
Stylet support insertion point	30	11.4	_	16.2	56.9	_	63.3	14.1	60.3	1.1	1.3
Buccal tube external width	30	1.4	_	2.1	6.8	-	8.6	1.8	7.8	0.2	0.4
Buccal tube internal width	30	0.7	_	1.3	3.2	_	5.1	1.0	4.1	0.1	0.4
Placoid lengths											
Macroplacoid 1	30	2.2	_	3.8	11.2	_	15.9	3.0	12.6	0.3	0.9
Macroplacoid 2	30	1.9	_	3.2	8.7	_	13.3	2.4	10.2	0.3	0.9
Macroplacoid row	30	4.5	_	7.2	23.1	_	30.0	6.0	25.7	0.6	1.3
Placoid row	30	5.9	_	9.6	30.8	_	40.2	7.9	33.8	0.7	1.7
Claw 1 heights											
External base	30	2.4	_	5.3	12.6	-	21.8	4.2	17.7	0.7	2.2
External primary branch	30	5.5	_	9.6	25.8	_	40.4	7.7	33.0	0.9	3.4
External secondary branch	30	2.8	_	4.9	14.6	_	20.4	4.2	18.0	0.5	1.5
External total	30	8.3	_	15.2	40.9	_	63.8	12.1	51.5	1.5	4.9
Internal base	28	2.9	_	4.4	13.3	_	17.6	3.7	15.6	0.4	1.1
Internal primary branch	27	4.0	_	5.7	18.0	_	24.0	4.9	21.0	0.4	1.6
Internal secondary branch	27	2.7	_	5.1	11.8	_	20.0	3.7	15.5	0.6	2.0
Internal total	27	6.7	_	9.5	29.6	_	39.6	7.9	33.7	0.6	2.2
Claw 2 heights											
External base	30	2.7	_	6.0	14.4	_	25.1	4.6	19.5	0.8	2.6
External primary branch	30	5.7	_	11.3	29.3	_	47.2	8.6	36.7	1.0	3.4
External secondary branch	30	2.8	_	5.8	14.6	_	23.8	4.5	19.3	0.7	2.0
External total	29	8.7	_	17.0	43.6	_	71.3	13.4	57.1	1.7	5.2
Internal base	27	2.8	_	4.7	12.7	_	18.8	3.9	16.4	0.5	1.4
Internal primary branch	26	3.7	_	6.7	17.5	_	25.8	5.2	22.2	0.6	1.9
Internal secondary branch	26	2.1	_	5.4	11.1	-	21.4	4.0	16.7	0.7	2.1
Internal total	26	6.2	_	10.2	29.8	-	42.5	8.4	35.8	0.9	2.6
Claw 3 heights											
External base	26	2.7	-	6.3	14.2	-	25.3	4.6	19.8	0.8	2.6
External primary branch	26	6.1	_	11.7	28.1	_	48.8	8.4	35.8	1.2	4.4
External secondary branch	26	2.8	_	5.9	14.9	_	24.2	4.7	19.9	0.7	2.0
External total	25	9.0	_	18.1	42.2	_	75.7	13.1	56.2	2.0	6.6
Internal base	21	2.6	_	4.9	13.9	_	19.2	3.9	16.8	0.5	1.2
Internal primary branch	19	4.0	_	6.6	18.3	_	27.6	5.1	22.1	0.6	2.0
Internal secondary branch	21	2.6	_	5.4	13.7	_	21.3	4.0	16.9	0.7	2.2
Internal total	20	6.4	-	10.4	31.7	-	43.3	8.4	36.2	0.9	2.8
Claw 4 lengths											
Anterior base	24	2.7	_	4.8	13.0	_	19.3	4.0	16.7	0.5	1.4

Table 3 (continued).

	N			RA	NGE	MF	EAN	SD			
	N		μm		pt			μm	pt	μm	pt
Anterior primary branch	24	4.6	_	6.4	19.9	_	27.0	5.4	23.1	0.4	1.6
Anterior secondary branch	24	2.8	_	5.2	14.7	_	20.9	4.4	18.7	0.6	1.6
Anterior total	24	7.1	_	11.0	31.3	_	46.1	9.0	38.2	0.9	2.8
Posterior base	28	3.1	_	6.1	15.0	_	24.4	5.2	21.8	0.7	2.3
Posterior primary branch	28	8.0	_	14.5	37.7	_	60.6	11.5	48.7	1.3	4.0
Posterior secondary branch	28	3.8	_	6.4	15.8	_	25.7	5.4	23.0	0.7	2.5
Posterior total	28	11.4	_	20.0	52.9	_	83.8	16.8	71.3	2.0	5.7

ITS-2 sequence (GenBank: MN927181), 486 bp long:

All obtained sequences were deposited in GenBank (https://www.ncbi.nlm.nih.gov/genbank/) under the following accession numbers: MK967241, MN915220, MN915221, MN918533, MN919385 (COI), MK973069, MN912103 (18S rRNA); MK967961, MK967962, MK967963, MK967964 (28S rRNA); MN927181, MN927182 (ITS-2).

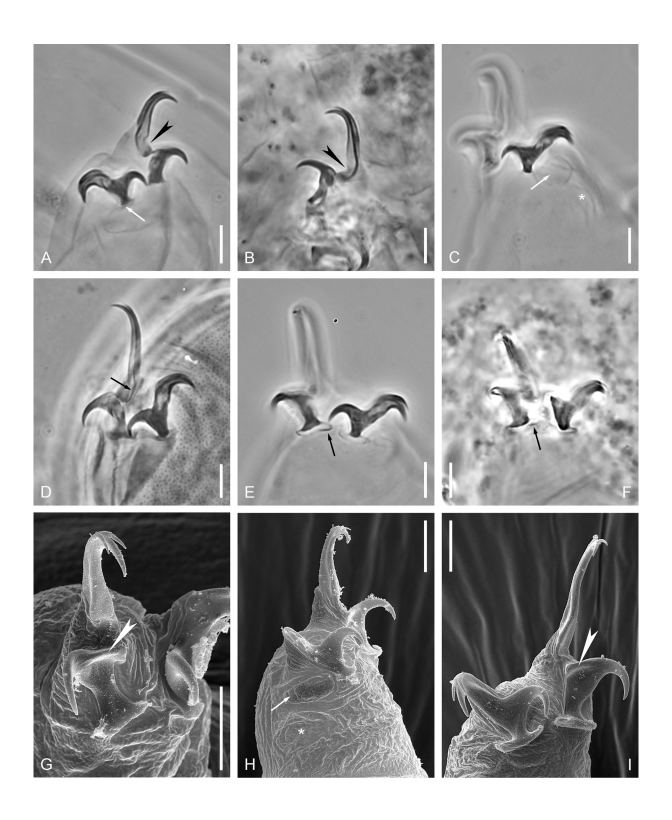
Phylogenetic analysis

In the 18S DNA phylogenetic analysis, the order Hypsibioidea was highly supported and divided into two well-supported clades: clade I, embracing the family Ramazzottiidae, Marley, McInnes & Sands, 2011, and clade II, comprised of taxa currently attributed to the families Hypsibiidae Pilato, 1969, Calohypsibiidae Pilato, 1969 and Microhypsibiidae Pilato, 1998 (Fig. 8). Clade II was further divided into two subclades well-supported with Bayesian analysis, but weakly supported or unsupported with ML analysis. The first subclade included the families Microhypsibiidae s. str. (genus *Microhypsibius* Thulin, 1928) and Calohypsibiidae s. str. (genus *Calohypsibius* Thulin, 1928) (see Gąsiorek *et al.* 2019a for a discussion on the taxonomic composition of these two families), together with two genera of unclear taxonomic position, *Acutuncus* and *Mixibius*. The second subclade was divided into three subclades with unclearly resolved phylogenetic relationships. The first of these subclades included the species representing the subfamily Pilatobiinae Bertolani, Guidetti, Marchioro, Altiero, Rebecchi & Cesari, 2014, *Hypsbius pallidoides*, and the species attributed to *Hypsibius convergens* (Urbanowicz, 1925) by Guil & Giribet (2012). The second one included the species of the subfamily Itaquasconinae Rudescu, 1964 and the third one was comprised of two well-supported lineages, the subfamilies Diphasconinae Dastych, 1992 and Hypsibiinae Pilato, 1969.

Analyses of the concatenated 18S + 28S sequences resulted in a phylogeny with the same tree configuration, but with slightly weaker support of the clades (see Supplementary file SM.02). This weakened support is possibly a consequence of the small number of sequences available for such analysis.

Comparison with the original description

Morphometry of specimens from all analysed populations (including the type series) corresponds well with the data from the original description (Pilato *et al.* 2011). Small differences in the values of the stylet supports insertion point *pt* index (54.2–55.2 in the original description vs 56.9–63.3 in the material investigated) and the length of the first macroplacoid (3.8–4.2 μ m (*pt* 15.5–17.0) vs 2.2–3.8 μ m (11.2–15.9) respectively) should be considered as the result of some differences in the measuring process, taking into account that my own measurements of the type series specimens are concordant with those of the specimens from the other populations (see Table 2).



It was stated in the original description (Pilato *et al.* 2011) that *H. pallidoides* had a smooth cuticle, but high quality LM and SEM observations revealed the presence of a cuticular sculpture (Figs 2A–D, 3A–B). It is poorly visible in the type series specimens because of the intensive staining of soft tissues with acetocarmine during the slide preparation.

Contrary to the absence of lunules in *H. pallidoides* stated by Pilato *et al.* (2011), my investigation determined that scarcely visible lunules on the claws of legs I–III and well-developed wide lunules on the claws of legs IV are present (Fig. 6A, E). In the original description of the species, Pilato *et al.* (2011) indicated the absence of a cuticular bar between the claw bases of legs IV, but considered this as unconfirmed. My observations revealed the presence of a thickened zone of the posterior claw lunule, located between the anterior and the posterior claw bases (Fig. 6E–F). This thickening can give the impression of a cuticular bar in the case when the main part of the lunule is not discernible.

While Pilato *et al.* (2011) described the eggs of *H. pallidoides* as being smooth, further scrutiny ascertained the presence of a granular pattern formed by the system of internal pillars in the egg shell of this species (Fig. 7B–G).

New phenotypic differential diagnosis

Hypsibius pallidoides is similar to the species of the genera Ramazzottius Binda & Pilato, 1986 and Cryoconicus Zawierucha, Stec, Lochowska-Cierlik, Takeuchi, Li & Michalczyk, 2018 in having claws of the Ramazzottius type; AISM asymmetrical with respect to the frontal plane; cephalic elliptical sensory organs and in laying ornamented eggs. It clearly differs from all species of those genera by having wider primary branches of the external and posterior claws, with less pronounced differentiation between rigid distal and soft basal parts; the dorsal AISM raised and thickened in its anterior margin, and eggs laid in the exuvium without external processes, but with pillars inside the egg shell only.

Hypsibius pallidoides is similar to the species of the genus Mixibius in having AISM asymmetrical with respect to the frontal plane, where the ventral apophysis is similar, but not identical, to the "semilunar hook" of Hypsibius; dorsal apophysis more stumpy with a blunt and swollen caudal apex. Also a short median cuticular thickening caudal to both these apophyses is present (the ventral one slightly visible) (Pilato & Binda 2010). It clearly differs from all species of this genus by having: cephalic elliptical sensory organs and Ramazzottius-like claws (external claws with elongated primary branches and less developed secondary branches).

The type of egg shell sculpture of *Hypsibius pallidoides* is similar to that of *Acutuncus antarcticus*, from the Antarctic region (see Dastych 1991 for a review of the old records) in that the sculpture,

Fig. 6 (opposite page). *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011, claws. Specimens SPbU 251(82) (A, C, E) and KNU Yep-9 II (B, D, F). **A**. Claws of leg I, black arrowhead indicates the flexible part of the external claw main branch, black arrows indicates the lunule of the internal claw, PhC. **B**. Claws of leg III, type series specimen, black arrowhead indicates the flexible part of the external claw main branch, PhC. **C**. Claws of leg II, white arrow indicates the elongated bulge near the base of the internal claw, white asterisk – pulvilla on the inner side of the leg, PhC. **D**. Claws of leg IV, type series specimen, black arrow indicates the filamentous structure connecting the external claw main branch with the claw base, PhC. **E**. Claws of leg IV, black arrow indicates the thickened region on the lunule margin, PhC. **F**. Claws of leg IV, type series specimen, black arrow indicates the thickened region on the lunule margin, PhC. **G**. Claws of leg II, white arrowhead indicates the lateral crest of the claw base, SEM. **H**. Claws of leg III, white arrow indicates the elongated bulge near the base of the internal claw, white asterisk – pulvilla on the inner side of the leg, SEM. **I**. Claws of leg IV, white arrowhead indicates the lateral crest of the claw base, SEM.

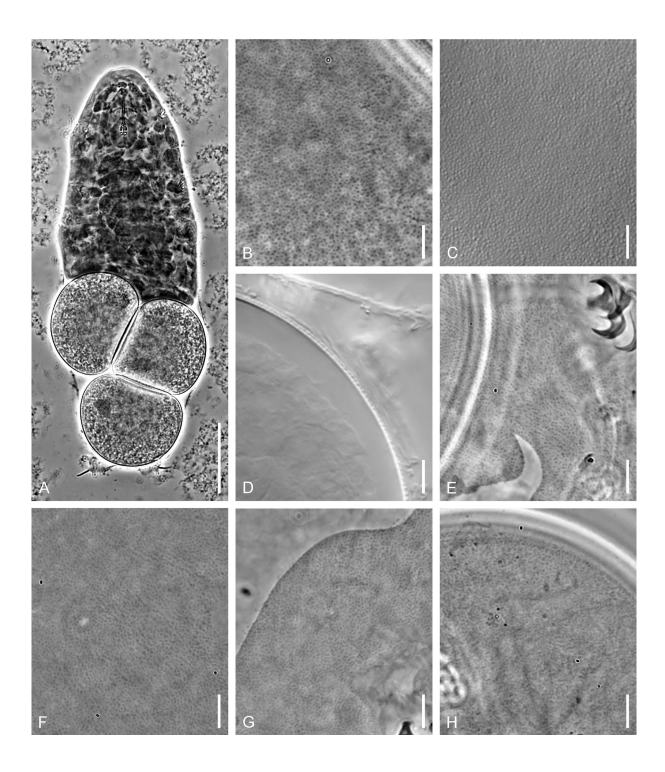


Fig. 7. A–G. *Hypsibius pallidoides* Pilato, Kiosya, Lisi, Inshina & Biserov, 2011 eggs. A–D, specimen KNU Чер-9 II. **A**. Type series specimen mounted during the egg laying process, PhC. **B**. Type series egg shell, PhC. **C**. Type series egg shell, DIC. **D**. Type series egg shell structure with numerous internal pillars visible, DIC. **E**. Austrian population egg shell (SPbU 251(3)), PhC. **F**. Karelian population egg shell (SPbU 113(2)), PhC. **G**. Pushkin population egg shell (SPbU 235(28)), PhC. – **H**. *Pilatobius recamieri* (Richters, 1911), egg shell (SPbU 203(7)), PhC. Scale bars: A = 50 μm; B–H = 5 μm.

formed by the pillars within the egg shell, presents as a dot-like pattern when observed in LM. *Hypsibius pallidoides* differs from *A. antarcticus* by having the *Ramazzottius*-type claws; AISM asymmetrical with respect to the frontal plane; a sculptured cuticle and a small dot-like septulum. The precise nature of the latter structure requires further investigation as its small size prevents it from being undoubtedly interpreted as microplacoid or septulum.

Hypsibius pallidoides is similar to the following species of the genus Hypsibius: Hypsibius allisoni Horning, Schuster & Grigarick, 1978 (known from New Zealand and South America (Horning et al. 1978; Maucci 1988; Pilato et al. 2003)); H. murrayi (=H. heardensis Miller, McInnes & Bergstrom, 2005; known from Antarctica (Dastych 2018)) and H. pachyunguis Maucci, 1996 (known from Greenland).

Hypsibius pallidoides clearly differs from the above mentioned species by having the Ramazzottius-like claws and by having cuticular sculpture. Additionally, H. pallidoides differs from:

Hypsibius allisoni by having a thinner buccal tube (external width up to 2.1 μm in H. pallidoides vs 4 μm in H. allisoni holotype) (Horning et al. 1978).

Hypsibius murrayi by the absence of cuticular bars near the claw bases of legs I–III, by having a dot-like pattern of the egg shell, a smaller body length (up to 292 μm in *H. pallidoides* vs 338.0–603.0 μm in *H. murrayi*) (Dastych 2018).

Hypsibius pachyunguis by having less elongated macroplacoids (see Maucci 1996: 196, fig. 1; Tumanov 2018: 440, fig. 4a–b).

Two species of the genus *Hypsibius* are known as laying eggs with granulated chorion in exuvium, *Hypsibius roanensis* Nelson & McGlothlin, 1993 (Guidetti *et al.* 1999) and *H.* cf. *scabropygus* (Guidetti & Bertolani 2001). *Hypsibius pallidoides* clearly differs from both of these species by having a septulum, the *Ramazzottius*-like claws, and a different cuticular sculpture.

Genotypic differential diagnosis

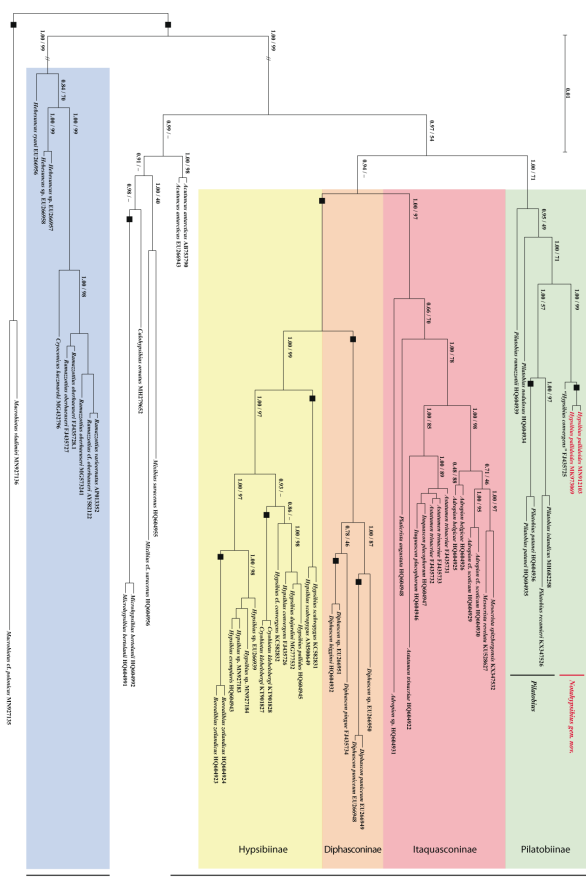
The ranges of uncorrected genetic *p*-distances between the studied population of *Hypsibius pallidoides* and species of the order Hypsibioidea for which sequences are available from GenBank (see Supplementary file SM.01) are as follows:

COI: 20.9%–26.7% (mean 23.0%), with the most similar being *Pilatobius recamieri* (Richters, 1911) (KX347530, Gąsiorek *et al.* 2017), and the least similar being *Diphascon puniceum* (Jennings, 1976) (KP013612, Velasco-Castrillón *et al.* 2015).

18S rRNA: 2.0%–8.7% (mean 6.2%), with the most similar being *Pilatobius recamieri* (KX347526, Gąsiorek *et al.* 2017) and *P. islandicus* Buda, Olszanowski, Wierzgoń & Zawierucha, 2018 (MH682258, Buda *et al.* 2018), and the least similar being *Diphascon puniceum* (EU266948, Sands *et al.* 2008).

28S rRNA: 5.9%–18.7% (mean 11.2%), with the most similar being *Mesocrista revelata* Gąsiorek, Stec, Morek, Zawierucha, Kaczmarek, Lachowska-Cierlik & Michalczyk, 2016 (KX347536, Gąsiorek *et al.* 2016), and the least similar being *Ramazzottius varieornatus* Bertolani & Kinchin, 1993 (MG432818, Zawierucha *et al.* 2018).

ITS-2: 19.6%–45.1% (mean 38.2%), with the most similar being *Pilatobius recamieri* (KX347528, Gąsiorek *et al.* 2017), and the least similar being *Ramazzottius subanomalus* (Biserov, 1985) (KU900019, Stec *et al.* 2016b).



Ramazzottiidae Hypsibiidae

Full matrices with p-distances are provided in the Supplementary file SM.03.

Sequences of the 18S and 28S rRNA genes, attributed to the species "Hypsibius convergens" by Guil & Giribet (2012) are nearly identical to those of Hypsibius pallidoides (p-distances 0.0% and 1.1% respectively).

Phylogeny of Hypsibioidea and phylogenetic position of Hypsibius pallidoides

The results of phylogenetic analysis presented herein correspond well with the molecular phylogenies of Tardigrada reconstructed in the recent works of other researchers (Guil & Giribet 2012; Bertolani et al. 2014; Guil et al. 2019), being most similar to the results of Bertolani et al. (2014). In comparison with the results of Guil et al. (2019), some differences in the tree topology may be attributable to a different approach taken in the selection of the compared sequences. In my opinion, the position of some sequences attributed to the species Acutuncus antarcticus within the cluster of species of Hypsibius (Guil et al. 2019: fig. 2) and the unclear differentiation of the Itaquasconinae and Hypsibiinae lineages are artefacts, caused by the inclusion of sequences derived from pooled samples which could contain multiple species (Sands et al. 2008), or by misidentifications of the sequenced specimens (see below).

My phylogenetic analysis confirmed the presence of the weakly supported but distinct basal clade within Hypsibiidae that includes the genera Acutuncus, Mixibius, Calohypsibius and Microhypsibius. With the addition of recently published data for two species of the genus *Pilatobius* Bertolani, Guidetti, Marchioro, Altiero, Rebecchi & Cesari, 2014 (Gasiorek et al. 2017; Buda et al. 2018), the Pilatobiinae clade, recognized in the analysis of Bertolani et al. (2014), became better supported in my analysis. Surprisingly, Hypsibius pallidoides (and a species attributed to H. convergens by Guil & Giribet (2012)) were distinctly placed within the Pilatobiinae clade, and even more interestingly within the genus Pilatobiotus itself, forming a cluster with the species P. patanei (Binda & Pilato, 1971)/P. islandicus/P recamieri, while the species P. ramazzottii (Robotti, 1970) and P. nodulosus (Ramazzotti, 1957) formed a separate paraphyletic group. Grouping of the species attributed to *H. convergens* with *Pilatobius recamieri* was obtained by Guil et al. (2019), but this result was not discussed by the authors. In an earlier publication (Guil & Giribet 2012), the taxon misidentified with H. convergens was joined with Astatumen trinacriae (Arcidiacono, 1962), but this result is likely an artefact because no species of *Pilatobius* were used in the analysis. In my opinion, extreme similarity of the 18S and 28S sequences of this species to the sequences of H. pallidoides (p-distances 0.0% and 1.1% respectively) should be considered as evidence of their identity on the genus level. Hypsibius pallidoides is morphologically similar to H. convergens and could be misidentified with this species, especially when temporary slides were used for the identification (Guil & Giribet 2012), because of the poor visibility of the cuticular sculpture and septulum in living specimens.

As a result, in the case of *H. pallidoides* we have a distinct contradiction between the morphological and molecular taxonomical approaches. Analysis of the morphological traits of this species reveals similarities with Ramazzottiidae (i.e., presence of the cephalic elliptical organs, the *Ramazzottius*-like claws, asymmetry of the AISM), but, according to the analysis of the gene sequences, this species should be attributed to the subfamily Pilatobiinae. Its position in the obtained phylogenetic tree also supports the presumably paraphyletic nature of the genus *Pilatobius*, also inferred by Gasiorek *et al.* (2018). To my knowledge, this is the first occurrence of such a distinct controversy between morphological and molecular taxonomy within Tardigrada. Previously, genetic analyses have supported

Fig. 8 (opposite page). The phylogeny of Hypsibioidea based on 18S rRNA sequences. Numbers at nodes indicate Bayesian posterior probability values (BI, first values) and bootstrap values (ML, second values). Black dots indicate the nodes supported by values of 1.0/100% with both methods. Scale bar and branch lengths refer to the Bayesian analysis.

the erection of taxa recognized by traditional morphological analysis (e.g., genera *Paramacrobiotus* Guidetti, Schill, Bertolani, Dandekar & Wolf, 2009, *Mesobiotus* Vecchi, Cesari, Bertolani, Jönsson, Rebecchi & Guidetti, 2016, *Acantechiniscus* Vecchi, Cesari, Bertolani, Jönsson, Rebecchi & Guidetti, 2016, family Ramazzottiidae, order Isohypsibioidea) (Guidetti *et al.* 2009; Vecchi *et al.* 2016; Sands *et al.* 2008) or provided an opportunity to resolve the phylogeny of a group when morphological data were insufficient (e.g., the clarification of the phylogenetic position of the genera *Apodibius* Dastych, 1983 and *Haplomacrobiotus* May, 1948, revisions of Isohypsibioidea and *Echiniscus* C.A.S. Schultze, 1840) (Dabert *et al.* 2014; Cesari *et al.* 2016; Gąsiorek *et al.* 2019b, 2019c). The presence of such controversy is a problem that has been acknowledged in current zoology since molecular methods began to be widely used (Hillis 1987; Osawa *et al.* 2004; Smirnov *et al.* 2005; Cohen 2018). Various authors who have analysed this problem (Hedges & Sibley 1994; Scotland *et al.* 2003; Osawa *et al.* 2004; Wiens 2004; Smith & Turner 2005) came to the conclusion that the best (if not the only) way to align the conflicting morphological and molecular phylogenies is to improve the morphological data by involving new characters in the analysis and by re-evaluating some characters already in use.

Taking into account the unique combination of the morphological features and the phylogenetic position of *Hypsibius pallidoides* distant from the remaining species of *Hypsibius*, as demonstrated by the analysis of the molecular data, the erection of the new genus *Notahypsibius* gen. nov. for the species *H. pallidoides* is proposed.

Taxonomic account

Genus *Notahypsibius* gen. nov. urn:lsid:zoobank.org:act:EDAD9932-04DD-4926-BEAA-369E0AA074C5

Type species

Hypsibius pallidoides Pilato, Kiosya, Lisi, Inshina & Biserov, 2011

Diagnosis

Hypsibiidae with *Ramazzottius*-like claws and completely rigid buccal tube. Apophyses for the insertion of the stylet muscles asymmetrical, dorsal AISM shorter and higher than ventral, with thickened anterior margin. Pharynx with two elongated macroplacoids and minute dot-like septulum. Cephalic elliptical organs present. Rugose cuticular sculpture. Eggs laid within the exuvium (or freely?), chorion with developed pillar-like internal structure visible in LM.

Etymology

The name refers to the phylogenetic position of the new genus, the type species of which was originally described as belonging to genus *Hypsibius*, but according to the phylogenetic analysis, definitely is "not a *Hypsibius*".

Genus composition (three species)

Notahypsibius pallidoides (Pilato, Kiosya, Lisi, Inshina & Biserov, 2011) gen. et comb. nov. Figs 1–7, Tab. 2, 3

Hypsibius pallidoides Pilato, Kiosya, Lisi, Inshina & Biserov, 2011: 13–15, fig. 7a–d (description).

Geographical distribution

This species was described from Kherson Oblast, South Ukraine (Pilato *et al.* 2011). Later it was recorded for the Minsk Oblast, Central Belarus (Pilato *et al.* 2012) and Sicily (Lisi 2015). My observations extend the distribution of this species to North-West Russia (St Petersburg and Karelia), Croatia and Austria (Carinthia). It should be noted that Dastych (1988) observed a configuration of the bucco-pharyngeal apparatus similar to *N. pallidoides* gen. et comb. nov. in some Polish specimens attributed by him to *H. convergens* (Dastych 1988: 147, pl. XXIa, c). Also, some of the microphotographs of the claws of Dastych's specimens of *H. convergens* show a similarity to those described for *N. pallidoides* gen. et comb. nov. (Dastych 1988: pl. XXIi). So, it is very likely that the latter species is present among the tardigrade fauna of Poland. The species attributed to as "*H. convergens*" by Guil & Giribet (2012) is nearly identical to *N. pallidoides* gen. et comb. nov. in 18S and 28S gene sequences (see Genotypic differential diagnosis). In my opinion, this is evidence for the presence of *N. pallidoides* gen. et comb. nov. in Spain, but it was recently shown (Guidetti *et al.* 2019a) that closely related species can share an identical 18S rRNA haplotype. Thus, without analyses of the more sensitive barcode genes (particularly COI and ITS-2), and in the absence of morphological data, the possibility of the presence of another species similar to *N. pallidoides* gen. et comb. nov. in Spain cannot be excluded.

Notahypsibius scaber (Maucci, 1987) gen. et comb. nov. Fig. 9A–I

Hypsibius scaber Maucci, 1987: 200, figs 11-12 (description).

Material examined

Holotype

USA • Yellowstone Park, near Undina Falls; Aug. 1984; W. Maucci leg.; moss on tree trunk; MCVR C.T. 12289.

Paratype

USA • 1 spec.; same collection data as for holotype; MCVR C.T. 12288.

Notes

The species described as *Hypsibius scaber Maucci*, 1987 (known from North America only; Maucci 1987) is very similar to N. pallidoides gen. et comb. nov. in having cuticular sculpture consisting of irregular ridges (Fig. 9A-B, D-E), highly differentiated external and internal claws that closely resemble the Ramazzottius-type claws (Fig. 9H–I), and a similar bucco-pharyngeal apparatus with a thin buccal tube and minute dot-like septulum (Fig. 9C, F). In my opinion, it should be transferred to the genus Notahypsibius gen. nov. as Notahypsibius scaber gen. et comb. nov. It seems also that N. scaber gen. et comb. nov. has cephalic elliptical organs (Fig. 9G), but their presence requires further confirmation because of the difficulties in observing the dorsal surface of the type specimens (R. Guidetti, pers. com.). Notahypsibius pallidoides gen. et comb. nov. differs from N. scaber gen. et comb. nov. in having a less developed cuticular sculpture, especially on the ventral side of the body (Fig. 9D-E), and in having external claws with the common base thinner and longer in relation to the secondary branch. It should be noted that the comparison of N. pallidoides gen. et comb. nov. with N. scaber gen. et comb. nov. cannot be considered to have definitively resolved the possible synonymy of these species. The latter species description was based on only two specimens, both of which have most of their claws in positions that obstruct observation and correct measurement. Also, the eggs of N. scaber gen. et comb. nov. are unknown.

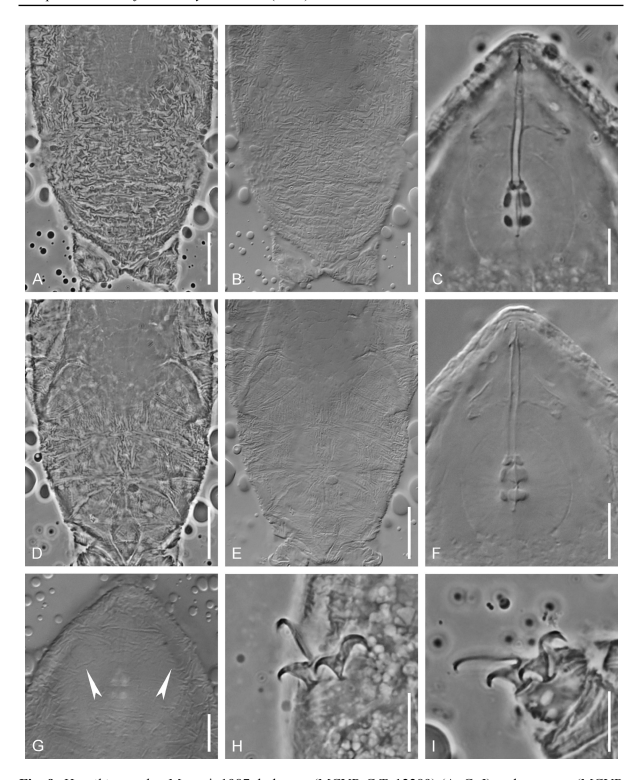


Fig. 9. *Hypsibius scaber* Maucci, 1987, holotype (MCVR C.T. 12289) (A–G, I) and paratype (MCVR C.T. 12288) (H). **A.** Dorsal view of the caudal body end, PhC. **B.** Dorsal view of the caudal body end, DIC. **C.** Bucco-pharyngeal apparatus, PhC. **D.** Ventral view of the caudal body end, PhC. **E.** Ventral view of the caudal body end, DIC. **F.** Bucco-pharyngeal apparatus, DIC. **G.** Dorsal surface of the head region, white arrowheads indicate the presumable elliptical sensory organs, DIC. **H.** Claws of leg III, PhC. **I.** Claws of leg IV, PhC. Scale bars: A–B, D–E = 20 μ m; C, F–I = 10 μ m. Photo G presented by R. Guidetti, University of Modena and Reggio Emilia, Italy.

Notahypsibius arcticus (Murray, 1907) gen. et comb. nov.

Macrobiotus arcticus Murray, 1907b: 677, pl. 1, fig. 5a–f (description). *Macrobiotus heinisi* Richters, 1911: 15, fig. 15.

Hypsibius arcticus – Thulin 1911: 27. — Marcus 1930: 380. — Dastych 1991: 141–159 (taxonomical notes).

Ramazzottius arcticus Gasiorek et al. 2018: 52.

Type locality

Franz Joseph Land (Murray 1907b).

Notes

The species Hypsibius arcticus (Murray, 1907) was recently transferred by Gasiorek et al. (2018) to the genus Ramazzottius on the basis of having Ramazzottius-like claws and freely laid eggs. In my opinion, the type of the chorion ornamentation in this species, consisting of the internal pillars, is definitely different from the external processes that are typical of the genus *Ramazzottius* (see Discussion). The combination of the *Ramazzottius*-like claws and eggs with developed internal pillars in the egg chorion makes this species more similar to N. pallidoides gen. et comb. nov. It should, therefore, be transferred to the new genus as Notahypsibius arcticus gen. et comb. nov. This species differs from N. pallidoides gen. et comb. nov. by having better developed pillars in the egg chorion and by laying free eggs. However, the latter trait requires confirmation as it is assumed upon the basis of a single observation (Murray 1907b), especially taking into consideration that *Macrobiotus heinisi* (Richters, 1911) – a similar species described from the same locality (Franz Joseph Land) and later synonymized with H. arcticus (Marcus 1930) – has eggs with a similar chorion structure laid within the exuvium (Richters 1911). Also, Acutuncus antarcticus, which shares an eggshell structure of a similar appearance, is known to lay eggs both within the exuvium and freely (Dastych 1991; pers. obs.). Other key characteristics, such as the presence of the minute septulum and cuticular sculpture, may have been overlooked by Murray (1907b) in his original description, as visualisation of these structures requires the use of high quality optics unavailable at that time. The specimen and the egg from Scotland, which were described and figured by Murray (1907a: 658, pl. IV, fig. 27a-d) as "Macrobiotus oberhäuseri Doy. ?", could not be attributed to N. arcticus gen. et comb. nov. because of the evident differencies in the claw structure (claws similar to the Cryoconicus type), the egg chorion appearance (much shorter pillars), and the significant difference in the value of the pt index for the stylet support insertion point (57% in Scottish specimen vs 70% in N. arcticus gen. et comb. nov., measurements taken from the original Murray's drawings). This material possibly represents an undescribed species of the genus *Cryoconicus*.

Discussion

Phylogenetic significance of some morphological characters

Ramazzottius-like claws

Morphology of the claws is one of the most important characters used in the taxonomy of Eutardigrada (Pilato 1969; Schuster *et al.* 1980; Pilato & Binda 2010; Gąsiorek *et al.* 2019c). *Ramazzottius* type claws were recognized as a separate morphotype (and denominated as "*oberhaeuseri* type" claws) by Binda & Pilato (1986) with the simultaneous erection of the genus *Ramazzottius* and were recently reanalysed by Guidetti *et al.* (2019b). Two other genera, *Ramajendas* Pilato & Binda, 1990 and *Thalerius* Dastych, 2009, were recognized as having a similar claw morphology (Pilato & Binda 1990; Dastych 2009). The phylogenetic and taxonomic position of these two genera is currently the subject of debate because of the evident controversy in their morphology and lack of DNA sequences. Being initially placed

within Isohypsibiidae (Marley et al. 2011; Guil et al. 2013), both genera were later attributed to the family Ramazzottiidae by Bertolani et al. (2014) on the basis of claw morphology. Zawierucha et al. (2018), taking into account the simple ridge-like form of the apophyses for the insertion of the stylet muscles and the deposition of smooth eggs in the exuvium known in the genus Ramajendas, proposed to place both of these genera back within Isohypsibiidae, suggesting the independent evolution of the Ramazzottius-like claws. Thus, for clarity, I use here the term "Ramazzottius-like claws" in order to distinguish the claws of Ramajendas, Thalerius and Notahypsibius gen. nov. from the Ramazzottius-type claws of Ramazzottiidae (see Guidetti et al. 2019b for a discussion). In a recent revision of the order Isohypsibioidea, Ramajendas and Thalerius are considered as incertae sedis pending molecular verification of their taxonomic positions (Gąsiorek et al. 2019c). In this situation, the obtained data showing the independent evolution of the Ramazzottius-like claws within the Pilatobiinae clade should be considered as an argument in favour of the hypothesis that Ramajendas and Thalerius are positioned phylogenetically distant from Ramazzottiidae.

Cephalic elliptical organs

Binda & Pilato (1986) pointed out the presence of these structures in the genus Ramazzottius and compared them with the papillae in the cephalic region of Calohypsibius ornatus (Richters, 1900). Since the genus Fractonotus Pilato, 1998, in which these structures are also known, was revised and transferred to the family Isohypsibiidae (Gasiorek et al. 2019a, 2019c) the cephalic elliptical organs are currently known within two Eutardigrada orders, Hypsibioidea (Calohypsibius and Notahypsibius gen. nov.) and Isohypsibioidea (Fractonotus). These structures should not be confused with 'cephalic papillae' or 'frontal lobes' known in several Isohypsibiidae genera (Halobiotus Kristensen, 1982, Apodibius, Ursulinius Gasiorek, Stec, Morek & Michalczyk, 2019 and Paradiphascon Dastych, 1992) (Gasiorek et al. 2019c) regarding the much more rostral position of the latter structures (within the anteriormost cephalic pseudosegment). Cephalic elliptical organs are located more caudally on the second pseudosegment following the cephalic one. In my opinion, only fronto-lateral porous areas of N. pallidoides (Fig. 4A–B) can be matched with the 'cephalic papillae' or 'frontal lobes' of Isohypsibiidae. Taking into account that cuticular sensory structures of Eutardigrada are very likely homologous to the cephalic sensory structures of Heterotardigrada (Zantke et al. 2008), the presence of such organs should be considered a plesiomorphic state and so supports the hypothesis of the basal phylogenetic position of Hypsibioidea and Isohypsibioidea (Marley et al. 2011; Gasiorek et al. 2019c).

Chorion structure

Although egg shell structure has been considered a valuable taxonomic character within Eutardigrada from the early years of its investigation (Marcus 1929, 1936; Ramazzotti & Maucci 1983), the phylogenetic significance of this trait was revealed considerably later (Bertolani *et al.* 1996). In their analysis, Bertolani and colleagues identified two types of the organization of the egg chorion – "smooth" and "ornamented" – and attributed *A. antarcticus* eggs to the ornamented type. In my opinion, the boundary between these two egg shell morphotypes within Tardigrada is not so obvious and the delimitation should be different.

Following the transfer of *R. arcticus* to the genus *Notahypsibius* gen. nov., there are only three genera known within Hypsibioidea with an egg chorion internal structure consisting of numerous pillars that connect the outer and inner layers of the shell: *Acutuncus* (Dastych 1991), *Notahypsibius* gen. nov. (present paper), and *Pilatobius* (present paper, see below). Additionally, two species of the genus *Hypsibius* (*H. roanensis* Nelson & McGlothlin, 1993 and *H.* cf. *scabropygus*) have finely granulated egg shells (Guidetti *et al.* 1999; Guidetti & Bertolani 2001). The similarity to the visible external pattern of the egg shell in *Acutuncus* and *Notahypsibius* gen. nov. makes it possible to suppose the same structure of the egg chorion for these two species. Ultrastructural investigations (Eibye-Jacobsen 1997; Poprawa 2011; Janelt *et al.* 2019) of egg development in *Halobiotus crispae* Kristensen, 1982,

Grevenius granulifer (Thulin, 1928) and Thulinius ruffoi (Bertolani, 1982) revealed the presence of the distinct pillars connecting the inner and outer layers of the chorion in the species with the egg shell usually considered to be smooth. As a result, the only difference between the typical 'smooth' eggs of most of the Hypsibioidea and Isohypsibioidea and the eggs with visible pillars within the shell (Acutuncus, Notahypsibius gen. nov., H. roanensis and H. cf. scabropygus) is the degree of the pillars' development making them visible in LM. In my opinion, this trait could have often been omitted in older observations of the eggs of other species due to the insufficient quality of the optics and the prevailing opinion that eggs laid in the exuvium are always smooth. For example, a careful investigation of the eggs of Pilatobius recamieri revealed the presence of the same type structure of the egg chorion (Fig. 7H).

The same structure of the egg shell was also described in Macrobiotidae Thulin, 1928 (Poprawa et al. 2015). Using the Transmission Electron Microscopy, the presence of the pillars connecting the outer and inner layers of the egg shell was revealed in *Macrobiotus polonicus* Pilato, Kaczmarek, Michalczyk & Lisi, 2003. This species has a continuous external layer of the egg shell, while in other species of the *Macrobiotus hufelandi* group this layer is modified to a mesh-like structure, supported by the pillars (Fig. 10A–B, white arrowheads). It also seems that such internal pillars form the dot-like pattern often visible in LM between the egg processes in some species of Macrobiotidae. For example, eggs of *Tenuibiotus voronkovi* (Tumanov, 2006) have a distinct dot-like pattern visible in LM, between the processes bases (Fig. 10C, black arrowhead; not mentioned in the original description), while SEM shows the absence of any granulation on the egg surface (Fig. 10D).

In my opinion, the presence of egg shell pillars visible in LM should be considered as a state, poorly delimited from the completely smooth egg shell (with pillars visible in EM only), and clearly different from the presence of true ornamentation consisting of the external processes. The development of the external processes does not exclude the presence of the internal pillars-like structures in the shell, but often these structures undergo progressive development, forming a mesh-like system of trabecules denoted as the "labyrinthine layer" (Węglarska 1982; Poprawa 2005).

It should also be noted that some tardigrade species have eggs with large, protruding pillar-like structures in the shell enclosed within the thin outer membrane. Egg shells of this type are known within both Heterotardigrada Marcus, 1927 (*Oreella* Murray, 1910) and Eutardigrada (Macrobiotidae, Murrayidae Guidetti, Rebecchi & Bertolani, 2000, Eohypsibiidae Bertolani & Kristensen, 1987) (Bertolani *et al.* 1996; Dastych *et al.* 1998). In my opinion, this type of egg shell structure could be derived from the primitive three-layered shell as a result of the progressive development of the internal pillars. This hypothesis can explain the emergence of similar-looking structures in several phylogenetically distant groups and possibly can partially resolve the known paradox in the tardigrade systematics formulated by Guidetti *et al.* (2006): "...there are closely related species, which share a very similar morphology of the animals but clearly differ in their egg morphology. Conversely, there are species belonging to different evolutionary lines that have similar eggs, but very different adult morphology". Surely, wide comparative TEM investigation of the egg chorion of different tardigrade species is needed to check this hypothesis.

It is interesting to note that, while within Eohypsibiidae the type of the egg shell ornamentation is genus-specific (pillar-like structures in *Eohypsibius* Kristensen, 1982 and external processes with labyrinthine layer in *Bertolanius* Özdikmen, 2008 and *Austeruseus* Trygvadóttir & Kristensen, 2011 (Trygvadóttir & Kristensen 2011; Hansen *et al.* 2017)), in Macrobiotidae and Murrayidae two genera include species with both types of the egg shell – *Murrayon* Bertolani & Pilato, 1988 and *Minibiotus* Schuster, 1980. The polyphyly of the genus *Murrayon* was previously demonstrated via molecular analyses (Bertolani *et al.* 2014; Guidetti *et al.* 2016). It incorporates at least two clades – one with the pillar-like structures

of the egg shell (*M. dianeae* (Kristensen, 1982) and related) and the other with the external conical processes (*M. pullari* (Murray, 1907) and related). Unfortunately, no molecular data are available on the species *M. ovoglabellus* (Biserov, 1988), which is known to lay smooth eggs. The genus *Minibiotus* is also suspected to be polyphyletic (Guidetti *et al.* 2007; Bertolani *et al.* 2014; Stec *et al.* 2015), but new molecular data are needed to test this hypothesis.

Bucco-pharyngeal apparatus of the Diphascon model

In the revision of the genus *Diphascon*, Pilato (1987) accepted the hypothesis of several independent origins of the long buccal tube with a flexible caudal part within Hypsibiidae as the most likely. He considered as a less likely alternative, the hypothesis of the presence of an independent monophyletic group within Hypsibiidae with a *Diphascon*-like buccal tube, where the shape of the apophyses for the insertion of the stylet muscles became identical to that of some species with a rigid buccal tube.

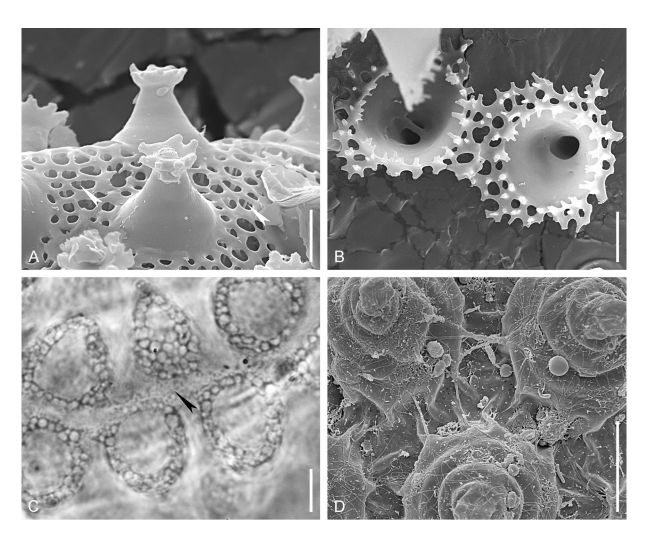


Fig. 10. Details of the egg shell structure in Macrobiotidae. **A.** Egg shell surface of *Macrobiotus* cf. *sottilei* Pilato, Kiosya, Lisi & Sabella, 2012, white arrowheads indicate the internal pillars of the egg shell, SEM. **B.** Fragment of the outer shell layer of *M.* cf. *sottilei* egg visible from inside, with numerous pillars, deriving from the lower surface of the reticulated outer layer, SEM. **C.** Fragment of the egg surface of *Tenuibiotus voronkovi* (Tumanov, 2006) with granulation-like pattern visible between the egg processes (SPbU 205(4)), PhC. **D.** Fragment of the egg shell of *T. voronkovi* in SEM, note the absence of the surface granulation. Scale bars: A–B = 2 μm; C–D = 5 μm.

The presence of the bucco-pharyngeal apparatus of the Diphascon model within Macrobiotidae and Eohypsibildae was considered as a confirmation of the possibility of the independent evolution of this trait. But after this work was published, a much wider distribution of the Diphascon-like buccal tube within Eutardigrada sensu lato (including the order Apochela Schuster, Nelson, Grigarick & Christenberry, 1980) has been revealed. Now, it is also known in three of four genera of Milnesiidae Ramazzotti, 1962 (Dastych 2011) and in the possibly closely-related Carphania Binda, 1978 (Binda & Kristensen 1986); in all major clades of Hypsibioidea: Ramazzottiidae, Diphasconinae, Itaquasconinae, and Pilatobiinae (Bertolani et al. 2014); in several genera of Macrobiotidae, possibly presenting different phylogenetic lines (Guidetti & Pilato 2003); in one of three known genera of Eohypsibiidae (Kristensen 1982) and, possibly, in Isohypsibioidea, if Paradiphascon is treated as belonging to that order (Gasiorek et al. 2019c). Consequently, a new hypothesis of the presence of the flexible pharyngeal tube as a plesiomorphy of the whole Eutardigrada sensu lato, was suggested by Bertolani et al. (2014). In my opinion, the position of the genus *Notahypsibius* gen. nov. on the obtained phylogram within the morphological genus Pilatobius can be considered as evidence of a possible reduction of the caudal flexible part of the buccal tube within the taxon with an initially Diphascon-like buccal tube (such a reduction was also recently hypothesized by Gasiorek & Michalczyk (2020) for the subfamilies Hypsibiinae and Itaquasconinae). As so, it strongly supports the hypothesis of the initially bipartite construction of the buccal tube within Eutardigrada sensu lato and independent reduction of the caudal flexible part in different phylogenetic lines.

Phylogeny and taxonomy of Pilatobiinae

The subfamily Pilatobiinae was established by Bertolani *et al.* (2014) when the phylogenetic analysis of 18S and 28S gene markers revealed that several species, previously attributed to the genus *Diphascon* (*D. nodulosum*, *D. patanei* and *D. ramazzottii*) form a separate clade within Hypsibiidae. These species, together with morphologically similar species of the genus *Diphascon*, were moved to a newly established genus *Pilatobius*. Morphological diagnosis of the subfamily Pilatobiinae, given by Bertolani *et al.* (2014), was based on the characters of the genus *Pilatobius* as it was the only genus within this clade at that time.

The phylogenetic analysis shown herein involved two additional *Pilatobius* species (*P. recamieri* and *P. islandicus*) with recently obtained gene sequences (Gąsiorek *et al.* 2017; Buda *et al.* 2018). The presence of the separate clade forming the subfamily Pilatobiinae was confirmed, but the analysis showed that the genus *Notahypsibius* gen. nov. was positioned within this clade and, moreover, within the genus *Pilatobius*. In this situation, the subfamily Pilatobiinae is still valid in terms of being a well-supported clade, but in lack of a suitable morphological diagnosis. No morphological characters can be pinpointed as an autapomorphy of this taxon. The only character possibly shared by *Notahypsibius* gen. nov. and *Pilatobius* is the presence of the pillars of the egg shell, visible in LM (known only for one species of *Pilatobius*), but this trait should not be considered significant because it is known to be present in other species of the family Hypsibiidae, belonging to the different clades.

The genus *Pilatobius* appears to be paraphyletic, as it consists of the monophyletic clade *P. islandicus/P. recamieri/P. patanei/N. pallidoides* and two species (*P. nodulosus* and *P. ramazzottii*) being sister to this species complex. Taking into account the small number of species of the genus *Pilatobius* with known gene sequences available for inclusion in the phylogenetic analysis (5 of 26 species), I prefer not to change the taxonomical status of this genus, and instead leave it with the diagnosis given to Pilatobiinae by Bertolani *et al.* (2014) with the following redaction: "*Genus* Pilatobius *Bertolani* et al. *2014. Buccal tube followed by an annulated pharyngeal tube, with a drop-like thickening between them; pharyngeal bulb roundish or slightly oval, always containing 2 macroplacoids similar in length and in rows that look as parentheses, and a septulum. Claws of the Hypsibius type."*

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Supplementary files

SM.01. Complete list of species analyzed, with their GenBank accession numbers.

SM.02. The phylogeny of Hypsibioidea based on concatenated 18S + 28S sequences.

SM.03. Uncorrected pairwise distances.