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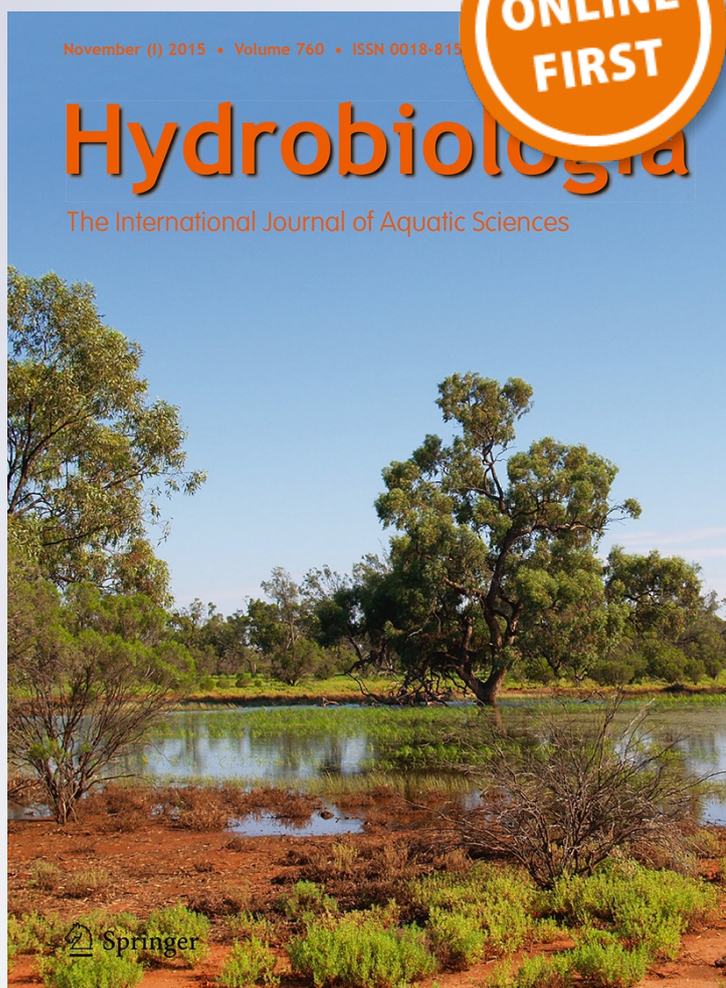
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Antarctic bdelloid rotifers: diversity, endemism and evolution

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Abstract Antarctica is an isolated continent whose conditions challenge the survival of living organisms. High levels of endemism are now known in many Antarctic organisms, including algae, tardigrades, nematodes and microarthropods. Bdelloid rotifers are a key, widespread and abundant group of Antarctic microscopic invertebrates. However, their diversity, regional distribution and endemism have received little attention until recently. We provide the first authoritative review on Antarctic Bdelloidea, based on published data and new collections. Our analysis

reveals the extreme levels of bdelloid endemism in Antarctica. Sixty-six bdelloid morphospecies are now confirmed from the continent, and 83–91 putative species are identified using molecular approaches (depending on the delimitation method used). Twelve previously unknown species are described based on both morphology and molecular analyses. Molecular analyses indicate that only two putative species found in Antarctica proved to be truly cosmopolitan. The level of endemism based on the available data set (95%) is higher than that in any other continent, with many bdelloid species occurring only in maritime or continental Antarctica. These findings are consistent with the long-term presence of Bdelloidea in Antarctica, with their considerable isolation facilitating intraregional radiation, providing further evidence that does not support the microbial global ubiquity hypothesis that “everything is everywhere.”

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Introduction

Antarctica's ecosystems are characterized by the challenges of extreme environmental stresses, including low temperatures, desiccation and high levels of solar radiation, all of which have led to the evolution and expression of well-developed stress tolerance features in the native terrestrial biota (Convey, 1996; Peck et al., 2006). The availability of liquid water, as well as its predictability, is considered to be the most important driver of biological and biodiversity processes in the terrestrial environments of Antarctica (Block et al., 2009; Convey et al., 2014). Antarctica's extreme conditions and isolation combined with the over-running of many, but importantly not all, terrestrial and freshwater habitats by ice during glacial cycles underlie the low overall levels of diversity that characterize the contemporary faunal, floral and microbial communities of the continent (Convey, 2013). Nevertheless, in recent years it has become increasingly clear that these communities contain many, if not a majority of species that have survived multiple glacial cycles over many millions of years and undergone evolutionary radiation on the continent itself rather than recolonizing from extra-continental refugia (Convey & Stevens, 2007; Convey et al., 2008; Fraser et al., 2014). With this background, high levels of endemism characterize the majority of groups that dominate the Antarctic terrestrial fauna, including in

particular Acari, Collembola, Nematoda and Tardigrada (Pugh & Convey, 2008; Convey et al., 2012).

The continent of Antarctica is ice-bound, and surrounded and isolated from the other Southern Hemisphere landmasses by the vastness of the Southern Ocean. The 1000-km Drake Passage separates it from South America and distances of 4–5000 km from Australia/New Zealand and South Africa. Terrestrial ecosystems reach their greatest development in the coastal regions, where most of the continent's biodiversity is found, most evidently along the Antarctic Peninsula and parts of the coastline of East Antarctica. Terrestrial communities are also present on isolated nunataks and the major mountain ranges inland, as well as in the 'dry valleys' of southern Victoria Land, which are the single largest ice-free areas of the continent (Convey, 2013). However, most ice-free areas are small and isolated by tens to hundreds of kilometers from neighboring areas.

Bdelloids, microscopic water-dwelling invertebrates belonging to the Subclass Bdelloidea of the Phylum Rotifera, account for 11–100% of all rotifer species recorded in Antarctic waterbodies and for 40–100% of species from terrestrial habitats (e.g., Dougherty & Harris, 1963; Sudzuki, 1964; Everitt, 1981; Sohlenius et al., 1996; Smykla et al., 2010). The evolutionary success of Bdelloidea in the extreme Antarctic environment is underlain by their parthenogenetic mode of reproduction and their ability to survive drying and/or freezing in an anabiotic state (cryptobiosis). Populations of bdelloids usually consist of a mix of reproductively isolated clonal lineages, often apparently morphologically uniform, but which are genetically distinguishable evolutionary entities (Birky et al., 2005). At least some clonal lineages can be identified by detailed examination of external morphological characteristics (Birky et al., 2011) and/or by the size and shape of hard parts of the masticatory apparatus (Fontaneto et al., 2007). To date only seven bdelloid morphospecies have been recognized as being endemic to the Antarctic and sub-Antarctic (Segers, 2007), although a recent preliminary molecular analysis has suggested that this number should be considerably greater (Velasco-Castrillón et al., 2014a). Unfortunately, many studies (including recent) use only superficial identification of rotifers, often incomplete or misleading when based on identification keys (Donner, 1965; Kutikova, 2005) for mostly European fauna. Much of the early

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literature on Antarctic Bdelloidea is inevitably in journals with limited access, and hence much relevant information is not easily accessible to contemporary researchers.

With this background, the aims of this study are: (1) to review contemporary knowledge of aspects of diversity, ecology and reproductive biology of Antarctic bdelloid rotifers, (2) to advance knowledge of morphological and molecular diversity of Bdelloidea in Antarctica and (3) to evaluate the level of endemism of Antarctic bdelloids.

Current state of knowledge of bdelloid diversity and biology in Antarctica

Early studies

The history of bdelloid research in Antarctica and the sub-Antarctic dates back more than a 100 years. Early records of Bdelloidea date to the start of the twentieth century, obtained from material collected by the First German Antarctic (1901–1903), Swedish (1901–1904), British (1907–1909) and Second French (1908–1910) Antarctic Expeditions. Richters (1907, 1908) was the first to record bdelloids from terrestrial mosses. However, the only two species unequivocally recognizable from his records, *Callidina angusticollis* (= *Habrotrocha angusticollis* Murray, 1905) and *C. longirostris* [= *Rotaria sordida* (Western, 1893)], were found further north, between 35° and 40° S (St. Paul and Amsterdam islands). The remaining 13 bdelloids, also attributed to the genus *Callidina*, are now unidentifiable to species. The illustrations available, depicting contracted bodies, jaws and foot appendages, suggest that these rotifers are most likely correctly referred to the genera *Habrotrocha* and/or *Macrotrachela*.

Scottish biologist, microscopist and polar explorer J. Murray was the first to describe new species of Antarctic Bdelloidea. In the excellently illustrated report on the British Expedition, Murray (1910) listed 12 bdelloid species from mosses and pools of Ross Island and one species [*Callidina tridens* = *H. tridens* (Milne, 1886)] from terrestrial moss from the Stranded Moraines of McMurdo Sound. Five species of the 12 found were previously unknown: *Philodina gregaria* Murray, 1910, *Ph. antarctica* Murray, 1910, *Ph. alata*

Murray, 1910, *Habrotrocha* (as *Callidina*) *angularis* (Murray, 1910) and *Adineta grandis* Murray, 1910. Four species that Murray identified as cosmopolitan, *A. barbata* Janson, 1893, *A. longicornis* Murray, 1906, *Callidina constricta* [= *Habrotrocha constricta* (Dujardin, 1841)] and *C. habita* [= *Macrotrachela habita* (Bryce, 1894)], were noted to have morphological differences from the original descriptions of these species as found in Europe. One further species, *Philodina* sp., while apparently new to science, was not further described.

Murray (1910) also discussed the tolerance of bdelloids to desiccation, salinity and extreme temperatures, their habitat and possible feeding preferences, presumed cosmopolitanism and possible dispersal mechanisms, and the origin of the Antarctic rotifer fauna. He noted the predominance of Bdelloidea over other rotifers in the habitats examined and the remarkably high proportion of species that appeared to be known only from Antarctica, which were fully adapted to the conditions of the Antarctic environment. He noted that the two most abundant species, *A. grandis* and *Ph. gregaria*, were both viviparous (possibly a means of increasing progeny survival under extreme conditions), although the only exclusively viviparous bdelloid genus, *Rotaria* Scopoli, 1777, would not be found in Antarctica for some time yet. Murray (1910) considered wind to be the main vector of bdelloid dispersal, also noting that the characteristics of air currents around the Antarctic continent made transportation of rotifers from sources to the north impossible. Waterbirds, along with wind, were also considered as dispersal vectors on the local scale, between different water bodies on Ross Island. Murray (1910) also included the first report of “watermelon snow,” a phenomenon caused by aggregations of *Ph. gregaria*, a large bdelloid rotifer with a bright-red colored stomach.

Early taxonomic studies of the Bdelloidea of the maritime Antarctic and sub-Antarctic islands were carried out by de Beauchamp (1913, 1940), who investigated terrestrial habitats of Jenny Island and Îles Kerguelen. However, the description of a new viviparous bdelloid *Philodina* (?) *jeanelli* Beauchamp, 1940, from Kerguelen was based only on contracted individuals and has possibly hampered identification of this species by subsequent researchers.

Further taxonomic studies

More recent taxonomic studies were made by Donner (1972a, 1980) using collections made by Dougherty and Harris (1963) on Ross Island and by Jennings (1976a) on Signy Island (South Orkney Islands; material initially erroneously attributed to the Falkland Islands). The Ross Island material allowed redescription of three species previously found by Murray—*A. grandis*, *Ph. gregaria* and *M. insolita* var., the latter apparently being identical with *M. habita* as described by Murray (1910). The Signy Island material included the previously undescribed species *Mniobia ostensa* Donner, 1980, and 11 other bdelloid species thought to be cosmopolitan. Suzuki (1964), examining material from Langhovde on the continental Antarctic coastline, depicted 11 unidentified bdelloids from the genera *Adineta*, *Habrotracha*, *Macrotrachela* and *Mniobia*, which cannot now be reliably attributed to any known species since many important characters (corona, trophi shape, oviparity/viviparity) were missing in the images presented. Dartnall (1983, 1995a, b) and Dartnall & Hollowday (1985) reported a total of 32 bdelloid species, depicting and redescribing 15 species from the maritime Antarctic and the continent (Princess Elizabeth Land), among which were nine previously unknown representatives of *Adineta*, *Habrotracha*, *Macrotrachela* and *Philodina*. Notwithstanding some uncertainty over details of the corona, most of the specimens described were clearly different from known species, while specimens identified as *A. gracilis* and the viviparous *Rotaria rotatoria* (Pallas, 1766) showed morphological inconsistencies with the original descriptions of non-Antarctic material. *Macrotrachela* (= *Callidina*) *papillosa* (Thompson, 1892) was erroneously listed as *Habrotracha papillosa* and *M. insolita* de Koning, 1947, as *M. insolata*.

Life cycle

Several studies have described the life cycles of endemic Antarctic bdelloids. Dougherty (1964) investigated reproductive features of *Ph. gregaria* cultivated in the laboratory and found its maturation time to be 28–110 days (in a laboratory refrigerator). This suggests a life span considerably longer than in any other cultivated bdelloid, including another Antarctic endemic, *A. grandis* (Dartnall, 1992; Ricci, 2001). The

fecundity of the viviparous *Ph. gregaria* (up to 24 offspring; Dougherty, 1964), was also much lower than those of various oviparous bdelloid species in cultures maintained at room temperature (Ricci & Caprioli, 2005). Dartnall & Hollowday (1985) recorded that *Ph. gregaria* could produce up to 32 young per female, a number close to that of many oviparous bdelloids but still lower than others. Dougherty (1964) stated that most *Ph. gregaria* offspring started to reproduce 27–90 days after birth. Dartnall (1992) confirmed the unusually long life span of *Ph. gregaria*—up to 89 days at 4°C, and twice that of *A. grandis* at the same temperature (40–50 days). Ruttner-Kolisko & Kronsteiner (1979, cited in Dartnall, 1992) reported that at 6°C *Ph. gregaria* lived longer than at 10°C (60 days vs 26) and produced more offspring (15 vs. 7). Also, Dartnall (1992) found the age at the first reproduction to be 36–37 days for *Ph. gregaria*, about ten times more than typical oviparous non-Antarctic bdelloids cultivated at room temperature (Ricci & Caprioli, 2005).

Ecology

A number of studies have described the interactions of Antarctic bdelloids with their substrata (moss, algal mats and soil), comparing these with other terrestrial microinvertebrates, while others have addressed seasonal changes in their populations. Davis (1981) evaluated the role of bdelloids in bryophyte communities of Signy Island by estimating their dry biomass from data presented by Jennings (1976b, 1979) on density and species composition. Both the average and maximum biomasses of Bdelloidea were comparable with or higher than those of Nematoda, though considerably lower than those of Tardigrada. The biomass of bdelloids in mosses could reach up to 29.5 mg dry mass m⁻²—fourfold greater than that of monogonont rotifers, with about a half of the bdelloid biomass being attributed to *Adineta* species. Davis' (1981) data on feeding preferences suggested that the diet of bdelloids consisted entirely of dead organic matter, contrasting with Dougherty (1964), who stressed the importance of unicellular algae in the diet of *Ph. gregaria*. However, members of the genus *Adineta* are also known to feed predominantly on dead organic matter elsewhere (Örstan, 1992). Everitt (1981) observed cyclical changes in abundance throughout the year in the bdelloid population of a

saline continental lake in the Vestfold Hills. Rotifers overwintered in a cryptobiotic state and during the summer reproduced with abundance peaks occurring at 3-week intervals. Dougherty (1964) and Dartnall (1992) reported that the time between recovery from cryptobiosis, or birth, and the first reproduction in *Ph. gregaria* was at least 1 month in the laboratory, but that it could be shorter in the natural environment. In the relatively stable lake environment, the abundance peaks observed could be successive new generations, especially as Bdelloidea, unlike another rotifer group, Monogononta, do not possess specific larval or programmed dormant stages. However, the largest abundance peak described by Everitt (1981) corresponded to a massive inflow of N and P compounds into the lake, indicating that environmental influences are also important. In the more unstable (in terms of water availability) terrestrial habitats environmental factors seem to be the major driver of bdelloid abundance dynamics (Iakovenko, 2004). Priddle & Dartnall (1978), investigating the microflora and microfauna of aquatic moss and algal communities in lakes of Signy Island, observed three- to seven-fold decreases in the abundance of *Philodina* sp. during winter compared to summer. They also reported that two non-sessile bdelloid species showed distinctive space distribution patterns inside moss cushions, dominating in different zones of stems and leaves. Cathey et al. (1981) found *Ph. gregaria* and *Ph. alata* to be able to colonize artificial substrata (polyurethane foam) in eight lakes of southern Victoria Land, the former being present in all the lakes and the latter in only three.

Based on recent studies, most or all rotifer species that have been recorded in Antarctic soil communities are bdelloids (Smykla et al., 2010). Even in soils of the McMurdo Dry Valleys, one of the driest places in Antarctica, rotifers were present in all sampled locations (Courtright et al., 2001). Confirming Murray's (1910) speculations of almost a century ago, Nkem et al. (2006) concluded that wind plays an important role in the dispersal of soil rotifers, and this has been proposed as the primary mechanism behind their colonization of remote ice-free areas such as isolated valleys and nunataks, where they can reach abundances of up to 135 ind g⁻¹ dry substrate (Sohlenius et al., 1996). In some soil types at Edmondson Point, Victoria Land, Smykla et al.

(2010, 2012) found bdelloid rotifers to be the dominant group of microinvertebrates, reaching over 8000 ind 100 g⁻¹ fresh soil. Smykla et al. (2010, 2012) also reported that bdelloids reached high abundances in wet soils under moss and algal and cyanobacterial mats, while being absent in both barren fellfields and heavily nutrient-enriched penguin colonies. In contrast, Porazinska et al. (2002) reported rotifers to be present and even dominant in terms of abundance (>4000 ind kg⁻¹ dry soil) in ornithogenic soils collected on Ross Island. Sohlenius & Boström (2008) similarly noted that rotifers were the most frequently encountered and abundant group of invertebrates in both ornithogenic soils and fellfields of Dronning Maud Land. Velasco-Castrillón et al. (2014b) reported bdelloid rotifers to be the most widespread and abundant taxon in soils from multiple locations in East Antarctica, being present in 87% of sampled sites and reaching 44 ind g⁻¹ dry soil. In this latter study, bdelloids were present in soils with widely varying particle size composition (from fine to coarse), both with and without vegetation, and with a broad variety of abiotic and geochemical parameters, all observations that are consistent with the high tolerance of this group toward extreme conditions.

As also noted in the Arctic (De Smet & Van Rompu, 1994), Bdelloidea play an important role in Antarctic cryoconite communities. In cryoconites on glaciers of the McMurdo Dry Valleys, rotifers were dominant, reaching over 3500 ind 100 g⁻¹ dry sediment, although abundance decreased with elevation and was also influenced by pH, nutrient concentrations and cryoconite area (Porazinska et al., 2004). Rotifer and tardigrade abundances were also positively correlated in these cryoconites.

“Watermelon snow” and similar phenomena on the surface of water, ice or algal mats resulting from the massive accumulation of red-coloured *Ph. gregaria* was originally described by Murray (1910) and later addressed briefly by Dougherty & Harris (1963), Dougherty (1964) and in more detail by Dartnall (1992). These accumulations can create very noticeable red patches on the surface of such substrata, ranging from a few centimeters to many meters in diameter (Dartnall, 1983). For a patch to grow to a size of about 10 m may take only a week, with the abundance of *Ph. gregaria* reaching up to over 20 million ind. m⁻².

Diversity and endemism

Studies that have included or provided compilations of the rotifer fauna of Antarctica and the sub-Antarctic, in particular terrestrial Bdelloidea, have been published by Dartnall (1983), Dartnall & Hollowday (1985), Sudzuki (1988), Adams et al. (2006), Segers (2007), Velasco-Castrillón et al. (Velasco-Castrillón et al. 2014a, b, c) and Fontaneto et al. (2015). These sources should be referred to for details on species diversity of particular regions. A few studies have been carried out at the same location over time. For example, Dougherty & Harris (1963), investigating Ross Island and the McMurdo Dry Valleys, found virtually the same species as recorded by Murray (1910). A number of previously unrecorded bdelloid species have been reported from Antarctica and sub-Antarctic by Jennings (1976a), Sudzuki (1979), Everitt (1981), Sohlenius et al. (1996) and Sohlenius & Boström (2005). These studies reported, along with the indigenous Antarctic bdelloids, some 20 morphospecies similar to species first described from Europe, thus considering Antarctic bdelloid fauna to include many cosmopolitan species. No Antarctic endemic bdelloid families or genera have been reported.

Velasco-Castrillón et al. (2014a) consider that the known Antarctic Bdelloidea diversity comprises 36 morphospecies. However, this figure does not include three species-level taxa identified by Murray (1910), Jennings (1976a) and Cathey et al. (1981) or ten further undescribed species reported by Dartnall & Hollowday (1985), Dartnall (1995a,b) and Sohlenius et al. (1996): *Adineta vaga minor* Bryce, 1893, *Ceratotrocha cornigera* (Bryce, 1893), *Philodinus* sp., *A.* sp., *Habrotrocha* sp., *Macrotrachela* sp. "A", *Macr.* sp. 1, *Macr.* sp. 2., *Mniobia* sp. N, *Philodina* sp. "A", *Ph.* sp. "B", *Ph.* sp. 1. and *Ph.* sp. 2. Including these taxa, in total 49 bdelloid morphospecies have been recorded in Antarctica and the sub-Antarctic over the last century.

Based on classical taxonomy, only seven endemic bdelloids [five described by Murray (1910), plus *Ph. jeanelli* and *Mn. ostensa*] have been reported for Antarctica, with the remainder being cosmopolitan and previously known from other continents including Europe (Donner, 1965; Segers, 2007). In contrast, the application of contemporary molecular approaches (Velasco-Castrillón et al., 2014a) suggests that the bdelloid fauna of Antarctica comprises mostly endemic species, or at least species not yet recorded from any other continent.

Molecular approaches

A number of genomic and molecular phylogeographic studies have been performed during the last decade on various groups of Antarctic microfauna, mainly microarthropods and nematodes (Stevens et al., 2006; Stevens & Hogg, 2006; McGaughan et al., 2008, 2010; Velasco-Castrillón & Stevens, 2014) as well as various microbial groups (see Vyverman et al., 2010) and mosses (Pisa et al., 2014). However, the application of such studies to bdelloid rotifers in Antarctica remains at an early stage. Fragmentary sequence data on Antarctic bdelloids have been published in studies of the evolution and global biogeography of Bdelloidea (Barraclough et al., 2007; Fontaneto et al., 2008, 2012). Velasco-Castrillón et al. (2014a) recently evaluated molecular diversity of Antarctic and sub-Antarctic bdelloids across a wide area. Their study identified 47 putative species, counting both sequence clusters, and singletons (entities with only one sequence obtained). All of the putative species were designated as Antarctic or Tierra del Fuego endemics based on percentage sequence similarities in comparison with representatives of eight bdelloid genera from other continents. The study also indicated that the true number of taxa in the genera *Adineta* and *Philodina* determined from the sequence data analyzed using the Poisson tree processes (PTP) model (Zhang et al., 2013) must be considerably higher than can apparently be determined by morphological approaches alone.

The current study shows that when appropriately analyzed, the morphological diversity of Antarctic bdelloids is sufficient to reveal most diversity detected by contemporary molecular markers. We also reevaluate previously published data on Antarctic bdelloid rotifer endemism and determine the ratio of endemic to cosmopolitan bdelloid species in comparison with such from other continents.

Materials and methods

Sampling and extraction of rotifers

New samples included in this study were obtained from both maritime and continental regions at sites between 63°60'S and 77°55'S (Fig. 1; Table 1). In continental Antarctica 11 sampling locations were

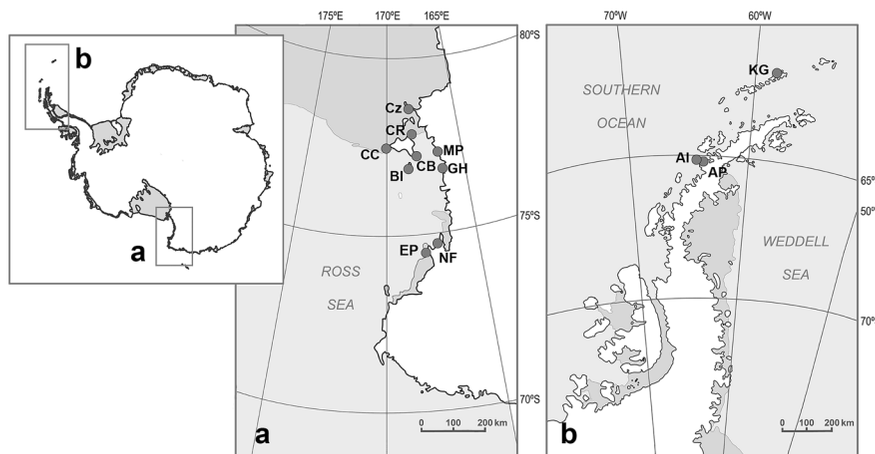


Fig. 1 Map indicating sampling locations in Antarctica. **a** Ross Sea area; **b** Antarctic Peninsula and adjacent islands. *BI* Beaufort Island. Ross Island: *CB* Cape Bird, *CC* Cape Crozier, *CR* Cape Royds. Coastal zone of Victoria Land: *Cz* Cape

Chocolate, *GH* Granite Harbor, *MP* Marble Point, *NF* Northern Foothills. Maritime Antarctica: *AI* Argentine Islands archipelago, *KG* King George Island, *AP* Mount Demaria, Kiev Peninsula, Graham Land

Table 1 Locations sampled in the Antarctic, including altitudes and the number of collected samples

Area	Locality ^a	Latitudes	Longitudes	Altitudes, m	Habitats
Antarctic Peninsula	AP	65°12'–65°17'S	64°06'–64°08'W	15–146	Soil, moss
Argentine Islands	AI	65°09'–65°56'S	64°03'–66°08'W	3–59	Soil, moss, lichens
King George Island	KG	63°60'41'–44''S	60°42'05'–29''W	5–11	Soil, waterbodies
Victoria Land	BI	76°55'–76°59'S	166°54'–166°56'E	6–141	Soil, moss, algal mats
–“–	CB	77°12'–77°15'S	166°22'–166°27'E	5–100	Soil, moss, algal mats
–“–	CC	77°27'21'–39''S	169°13'–169°15'E	61–201	Soil, moss
–“–	CR	77°32'–77°34'S	166°08'–166°10'E	6–28	Soil, moss, algal mats
–“–	Cz	77°56'21'–26''S	164°30'–164°32'E	20	Soil, moss, algal mats
–“–	EP	74°19'45'–60''S	165°07'–165°09'E	15–24	Soil
–“–	GH	77°00'25'–60''S	162°28'–162°32'E	10	Soil
–“–	MP	77°25'–77°27'S	163°40'–163°51'E	14–38	Soil, moss, algal mats
–“–	NF	74°42'25'–46''S	164°06'11'–54E	34–113	Soil

^a See Fig. 1

AP Antarctic Peninsula, *AI* Argentine Island Archipelago, *KG* King George Island, *BI* Beaufort Island, *CB* Cape Bird, *CC* Cape Crozier, *CR* Cape Royds, *Cz* Chocolate Point, *EP* Edmondson Point, *GH* Granite Harbour, *MP* Marble Point, *NF* Northern Foothills

visited in the Ross Sea area, including the Victoria Land coast, Beaufort and Ross Islands (Fig. 1a, Supplementary file I). The fieldwork and sampling in the Ross Sea area were conducted during five austral summer seasons between 2003/04 and 2011/12 within the project of J. Smykla (Smykla et al., 2010, 2011, 2012). Soil and moss were collected at all localities. The soil samples were obtained from barren fellfields, bryophyte communities, wetlands with algal and cyanobacterial mats, and the vicinity of active and

relict penguin colonies. Algal mats were collected in coastal areas and on Ross Island. Most of the collected material was stored frozen (–20°C), but some terrestrial mosses were dried and stored at room temperature; details of collection methods and primary sample processing are given in Smykla et al. (2010, 2012, 2015).

In the maritime Antarctic 237 samples were obtained during the summers of 2004/2005, 2006/2007 and 2009/2010 from the Argentine Islands

archipelago, King George Island and coastal areas of the Antarctic Peninsula under the projects of K. Janko, I. Kozeretka and V. Trokhymets. These included 50 soil and 183 moss samples, one sediment sample from a pool on King George Island and three lichen samples from the Argentine Islands archipelago (Fig. 1b; Table 1; Supplementary file I). Methods of collection, storage and rotifer extraction were as used for the continental samples, except for mosses, which were washed directly along with wet sieving and sugar gradient centrifugation as described by Freckman & Virginia (1993).

Alpha taxonomy procedures

Detailed procedures of rotifer sorting, identification, digital imaging and the preparation of type material (glycerin jelly slides and SEM mounts) are described in Iakovenko et al. (2013). We used the keys of Donner (1965) and Kutikova (2005) as a primary guide for identification, but detailed taxonomic analysis was based on the first descriptions (cited in Donner (1965) and further specific studies (Haigh, 1965, 1966; Donner, 1972a, b, 1980; Örstan, 1995; Koste, 1996a; Ricci et al., 2001, 2003; Birky et al., 2011).

Rotifer trophi (hard parts of the mastax) were extracted using Savo[®] Perex bleach and prepared for SEM according to De Smet (1998). Trophi measurements (ramus length and trophi width) were made as described by Iakovenko et al. (2013). Type material for newly described species (holotypes, paratypes and additional specimens) are deposited in the collections of the Schmalhausen Institute of Zoology, National Academy of Sciences of Ukraine, Kiev, Ukraine.

Additionally, we investigated and described rotifers depicted in photographs in Velasco-Castrillón et al. (2014a). To reliably distinguish, both morphologically and genetically, between several similar European and Antarctic species, we used material from our collections in Bulgaria, Czech Republic, Germany and Poland (Supplementary file I). Previously unpublished data on the morphology and morphometry of *A. barbata* Janson, 1893, *A. gracilis* Janson, 1893, *A. vaga* (Davis, 1873) and *Habrotrocha thienemanni* Hauer, 1924, from these collections, as well as COX1 sequences of the voucher specimens from which the morphometric data were taken, were used for these analyses.

Morphometric analyses

External rotifer body dimensions were taken on screenshots from digital videos, and trophi were measured on SEM photos, as described in Iakovenko et al. (2013). Total length (TL) in the case of adinetid rotifers was taken as the distance between the middle of the anterior rim of the head and the posterior rim of the spur pseudosegment, i.e., not including the rostrum, as it was usually bent under the head (Fig. 2).

To distinguish some Antarctic species from morphologically similar European ones, we measured specimens from clonal cultures, from which we subsequently obtained some COX1 sequences: 113 specimens of Antarctic *Adineta*, 69 specimens of European *Adineta* and 16 specimens of Antarctic *Habrotrocha*. We used the Linear Mixed Effects Model (LME) and Principal Components Analysis (PCA) to compare body and trophi measurements. The results of PCA were visualized as the two first principle components of variation plotted against each other. All statistical analyses were performed in R 2.15.1 following Crawley (2007).

DNA taxonomy procedures

The DNA extraction protocol followed Fontaneto et al. (2007) and Iakovenko et al. (2013). The target locus of the mitochondrial COX1 gene (355 bp in length) was amplified and sequenced from 194 bdelloid specimens (Supplementary file III) using universal primers LCO1490 and HCO2198 (Folmer et al., 1994) with the subsequent reamplification to increase the outcome of the product, using bdelloid-specific primers Bdel_CO1_FW (5'-CGTACWGAGTTAGGAATRGTA-3') and Bdel_CO1_Rev (5'-CCAAAATTWCGATC TAAYA-3') (Robeson et al., 2011).

To construct phylogenies, we downloaded available sequences of the taxonomically assigned bdelloid species from GenBank, available from all continents except South America (977 COX1 sequences, their detailed descriptions are given in the Supplementary file II). We used EMBL online version of MAFFT software (Kato et al., 2002) to construct one total alignment of both newly obtained sequences and those downloaded from GenBank and four separate alignments for four genus-specific data sets (*Adineta* Hudson and Gosse, 1886, *Habrotrocha* Bryce, 1910, *Macrotrachela* Milne, 1886, *Philodina* Ehrenberg,

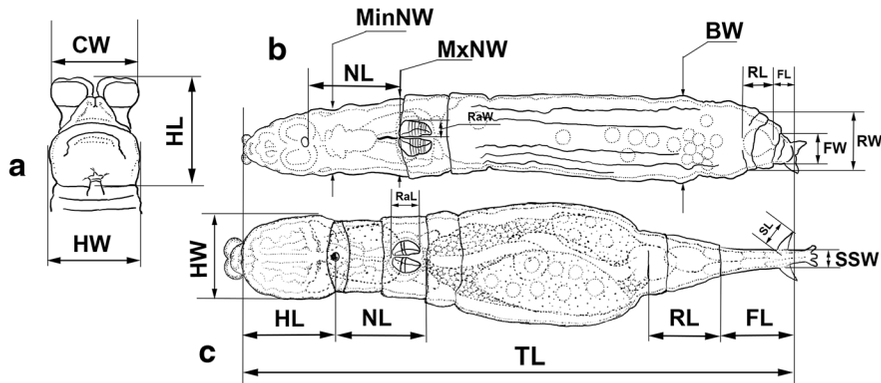


Fig. 2 Measurements of bdelloids of the families Habrotrochi-
dae and Philodinidae (a, b) and Adinetidae (c). *BW* body width,
CW corona width, *FL* foot length, *FW* foot width, *HL* head
length, *HW* head width, *MinNW* minimal neck width, *MxNW*

maximal neck width, *NL* neck length, *RaL* ramus length, *RaW*
ramus width, *RL* rump length, *RW* rump width, *SL* spur length,
SSW spur pseudosegment width, *TL* total length

1830). The monogonont rotifer *Brachionus calyciflorus* Pallas, 1755, was used as outgroup in each of these alignments, and each genus-specific data set also contained a member of another bdelloid genus as an additional outgroup: *Bradyscela clauda* (Bryce, 1893) for *Adineta*, *M. ehrenbergii* (Janson, 1893) for *Habrotrocha* and *H. constricta* (Dujardin, 1841) for *Macrotrachela* and *Philodina*.

We constructed phylogenetic trees in MrBayes 3.2.3 (Ronquist et al., 2012), running 8–20 million generations and sampling every 1000 generations. The optimal nucleotide substitution model (GTR+I+G) was chosen for each data set in jModelTest 2.1.6 (Darriba et al., 2012). The analysis was stopped when the standard deviation of split frequencies was below 0.01, with the PSRF being 1.00 for all the parameters. Effective sample size (ESS) sufficiency for the model parameters, process stationarity and the number of burn-in trees were checked using both MrBayes and Tracer 1.6 software (Rambaut et al., 2013). The resulting consensus trees constructed in MrBayes were visualized using FigTree 1.4.2 (Rambaut, 2012), and the full-size Bayesian trees are included in Supplementary file IV.

Three independent approaches were used for species delimitation based on DNA sequence data: 4× rule (Birky et al., 2005; Birky & Barraclough, 2009), Generalized Mixed Yule Coalescent Approach (GMYC; Fujisawa & Barraclough, 2013) and PTP with Bayesian support (bPTP; Zhang et al., 2013). The 4× rule identifies as putative species those monophyletic

clades whose genetic distances (K) to other sequences on the phylogenetic tree are larger than four times the intra-clade divergence (θ). To assess this, we constructed matrices of mean pairwise correlated sequence distances for each clade in the Bayesian trees in MEGA6 software (Tamura et al., 2013), calculating θ and estimating the K/θ ratio within and between the clades (Supplementary file V).

The GMYC method likewise identifies species as independently evolving entities represented by a number of clades on a phylogenetic tree. However, each clade is delimited by optimizing the tree nodes indicating transitions between inter- and intraspecific evolutionary processes. The maximum likelihood optimum is found between models of species diversification (based on the Yule model) and branching events within species (based on the neutral coalescent model). The initial tree should be time-calibrated (ultrametric), unrooted and not contain polytomies or zero-length branches. We used a single-threshold version of the method implemented in GMYC species delimitation software available online (<http://species.h-its.org/gmyc/>). The uploaded coalescent trees were produced from Bayesian unrooted trees in R 3.1.2 (<http://www.r-project.org/>) using the *chronopl* function of the “ape” package. This function utilizes a semiparametric method based on penalized likelihood (Sanderson, 2002) to estimate the tree node ages through a trade-off between contiguous and non-contiguous branch rates.

Unlike GMYC, the bPTP method does not require a time-calibrated and unrooted tree as input. In this method, the number of substitutions κ between intra- and interspecific events is used instead of time as a tree-calibrating parameter. Assuming that each substitution (which is independent of other substitutions) has a probability ρ of generating a speciation event, κ substitutions generate η speciations in a continuous process, and in a population of the size η the number of substitutions is sufficient; the process proceeds at the rate $\rho \times \eta$ and follows a Poisson distribution. The number of substitutions is calculated from the branch lengths of the input tree. We used online implementation of bPTP (<http://species.h-its.org/ptp/>) and the trees produced in MrBayes as the input.

Results

In total, we identified 60 morphospecies, including 20 taxa currently identified to the generic level only and still under investigation, and ten listed as “conformis” that show minor morphological differences from known species. Only 13 of the morphospecies found occurred in both maritime and continental Antarctica. The material examined included six of the seven known Antarctic endemics: *A. grandis*, *H. angularis*, *Mn. ostensa*, *Ph. alata*, *Ph. jeanelli* and *Ph. gregaria*. We have identified ten morphospecies reported by other researchers from Antarctica as *A. barbata* Janson, 1893, *A. vaga* (Davis, 1873), *H. gulosa* Milne, 1916, *H. vicina* Donner, 1980, *Macr. ambigua* Donner, 1965, *Macr. concinna* (Bryce, 1912), *Macr. habita* (Bryce, 1894), *Macr. musculosa* (Milne, 1886), *Macr. nixa* Donner, 1962, and *Rotaria rotatoria* (Pallas, 1766). These species are considered cosmopolitan, or at least are known from locations other than Antarctica. However, of these ten species, those resembling *A. barbata* and *A. vaga* s. str. are shown to be distinct new taxa and therefore currently endemic to Antarctica, based on both minor but consistent morphological differences and molecular analyses.

In *Alpha taxonomy*, below, we describe 12 new for science Antarctic bdelloid species. For some of them we also provide statistical analysis of morphometric data confirming their delimitation from morphologically similar described species occurring in Europe (*Morphometric analyses*). New records for the Antarctic, yet to be verified by molecular analyses belonging

to already described cosmopolitan species, included *H. angusticollis* (Murray, 1905), *Macr. nana* (Bryce, 1912), *Mniobia incrassata* (Murray, 1905), *Mn. scabrosa* Murray, 1911 and *Pleuretra lineata* Donner, 1962. The genus *Scepanotrocha* (*S.* cf. *semitecta* Donner, 1951) is reported from Antarctica for the first time. The list of known Antarctic bdelloids is therefore extended to 66 *morphospecies* (49 already known and reported in the existing literature, 12 new for science and 5 new for Antarctica).

Phylogenetic trees constructed using 194 original COX1 sequences and 977 sequences downloaded from GenBank gave similar results on the delimitation of independently evolving entities (IEE) according to the 4× rule, GMYC and bPTP models. These results are discussed in detail below (*DNA taxonomy* subsection). The 4× rule gave 140 IEEs: 44 of *Adineta*, 22 of *Habrotrocha*, 26 of *Macrotrachela* and 48 of *Philodina*. In total, 132 IEEs were identified by GMYC: 44 of *Adineta*, 20 of *Habrotrocha*, 18 of *Macrotrachela* and 50 of *Philodina*. Finally, bPTP generated a somewhat higher number of IEEs (160): 47 of *Adineta*, 26 of *Habrotrocha*, 29 of *Macrotrachela* and 58 of *Philodina*. Most of the IEEs identified by GMYC and bPTP were confirmed by the 4× rule. Delimitation according to the GMYC approach gave the best correspondence with rotifer morphology, considering both major and minor external features and morphometric data. Geographical distribution of the identified IEEs is discussed in *Biogeography*.

The integrity of most species identified by morphology, with the exception of *A. grandis*, *Ph. gregaria* and two new species of *Habrotrocha*, was confirmed by molecular analyses (*DNA taxonomy*). According to the molecular data, *A. grandis* consists of at least two cryptic species, one of which is described below as new for science. Ten putative species (IEEs) were identified from molecular data only, obtained from both the new material examined in this study and COI sequences downloaded from GenBank.

Alpha taxonomy

The list of locations is given after both the literature sources (cited in the Introduction) and our data (marked with *). Full descriptions of the examined samples, mentioned in Type material and Additional material below (as sample codes), are given in the Supplementary file I.

Abbreviations BW, body width; HL, head length; HW, head width; NL, neck length; MinNW, minimal neck width; MxNW, maximal neck width; RL, rump length; RW, rump width; FL, foot length; FW, foot width; SL, spur length; SSW, spur pseudosegment width; TL, total length. The abbreviations of the localities are explained in the Fig. 1 and Table 1, with the exception of the data from literature: DM, Dronning Maud Land; EB, Enderby; FI, Francis Island; HI, Haswell Island; LH, Langhovde; MM, McMurdo Sound; QM, Queen Mary's Land; SI, Signy Island; TF, Tierra del Fuego; WK, Wilkes Land.

Phylum Rotifera Cuvier, 1817
Class Eurotatoria De Ridder, 1957

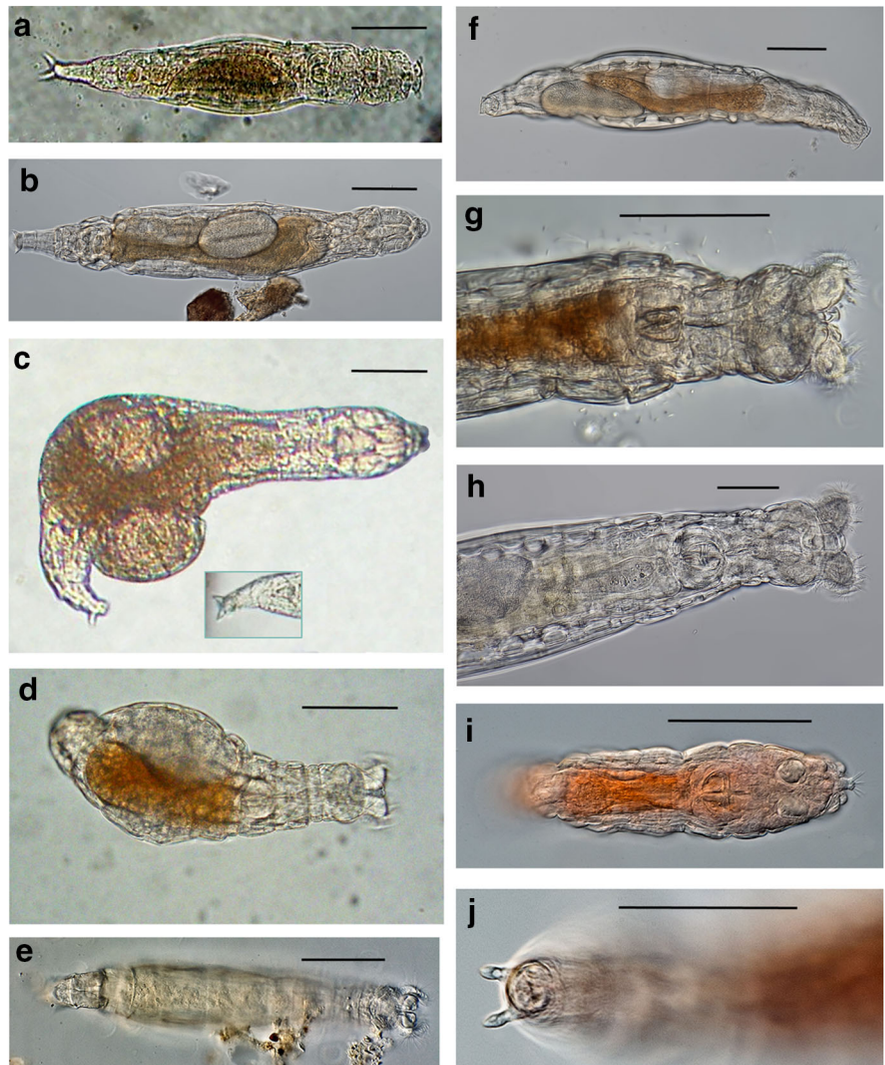
Subclass Bdelloidea Hudson, 1884
Order Philodinida Melone & Ricci, 2005
Family Adinetidae Hudson & Gosse, 1889
Genus *Adineta* Hudson and Gosse, 1886

***Adineta coatsae* sp. nov. (Figs. 3a, 4)**

Murray, 1910 (*A. barbata*?), pp. 53–54, Pl. XII Fig. 9a–c. Dartnall & Hollowday, 1985 (*A. barbata*), p. 30, Fig. 24a, b. Velasco-Castrillón et al., 2014a (*A. sp. Bd24*), p. 8 (main text), 2, Fig. 6 (Annex S1).

Type locality Chocolate Point (Victoria Land), 20 m asl., S77°56.400', E164°30.693'.

Fig. 3 New species of Antarctic bdelloids (photographs M. Plewka and N. Iakovenko): **a** *Adineta coatsae* sp. nov., holotype, habitus, dorsal view; **b** *A. editae* sp. nov., habitus, dorsal view; **c** *A. grandis*, habitus, dorsal view; **d** *H. antarctica* sp. nov., holotype, habitus, feeding, dorsal view; **e** *H. vernadskii* sp. nov., habitus, feeding, ventral view; **f** *M. jankoi* sp. nov., habitus, creeping, ventral view; **g** same, head, feeding, ventral view; **f** *M. ioannae* sp. nov., habitus, feeding, dorsal view; **i** *Ph. dartnallis* sp. nov., habitus, creeping, dorsl view; **j** same, spurs. Scale bar 50 µm



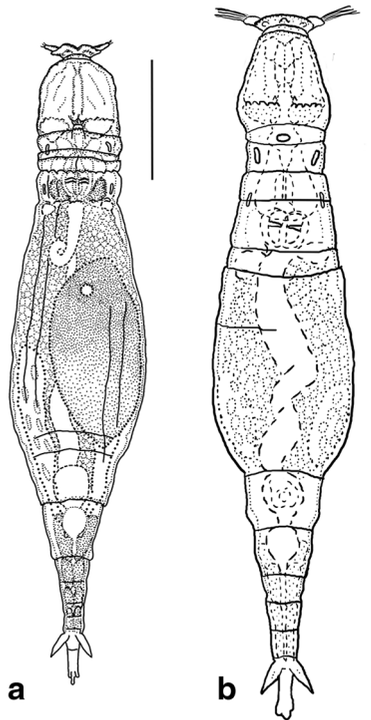


Fig. 4 *Adineta coatsae* sp. nov. (Antarctica): **a** holotype, habitus, dorsal view. *A. barbata* (Europe, BG0715): **b** habitus, dorsal view. Scale bar 50 μ m

Type habitat Algal and cyanobacterial mats.

Type material Holotype: SIZ 55.1 (CzM3NC-matAC1, 23.1.2010, Leg. J. Smykla), mounted in glycerin jelly. Paratypes: SIZ 55.2–55.3 (CzM3NC-matAC2–3), trophi mounted for SEM. *Additional material*. SIZ 55.4–10 (CzM3AS1–2, MPM4-mossAC1, V10AC1–2, KG2AC1, V10AC1), digital photos and videos.

Etymology Named in honor of a mountaineer and Antarctic researcher, Dr. Larry Coats, who assisted in the fieldwork done in the Ross Sea area.

Barcodes GenBank ID KJ543629–30.

ZooBank LSID. urn:lsid:zoobank.org:act:22DAD23A-DD71-4FB7-828C-59DCD677EAB7.

Diagnosis Similar to *A. barbata* (Fig. 4b) by flat laterally widened rostrum with two protrusions ending with a bundle of long thin sensory bristles and long sword-like spurs. However, the protrusions are leaf-like, while in *A. barbata* they are tubular. Frontal rim of the rostrum is concave and has a notch in the middle, while in *A. barbata* the notch is absent and the frontal rostral rim is prominently convex (Fig. 4b). Spurs gradually tapering from their base to the points,

shorter than in *A. barbata*. The new species seems to have substantially smaller body (TL 220–292 μ m) than *A. barbata* (TL 280–400 μ m according to Donner, 1965 and 223–374 μ m according to our data).

Description Body of moderate size, not very wide, flattened dorsoventrally, transparent, stomach usually of brown-yellow color. Darnall & Hollowday (1985) report the color of this rotifer as grayish-brown. Integument smooth, thin, without sculpturation, spines, knobs or bolsters. Head trapezoid, wider in the posterior part, HL is 15–19% of TL, HW is 76–94% of HL. Distal rostral pseudosegment flat, lobe-like widened, with a V-shaped shallow and wide notch in the middle. Rostral lamella shaped as two lateral leaf-like narrow protrusions with a bundle of long sensory bristles under each protrusion. Eight rectangular teeth in each rake. Neck of moderate length and width, NL is 12–17% of TL, antenna about 1/4–1/5 of bearing pseudosegment. Trunk oval, BW 17–28% of TL. Rump conical, first pseudosegment slightly swollen, RL is 12–17% of TL, RW is 75–92% of RL. Slim foot of five pseudosegments, of moderate length, FL is 28–35% of TL, FW is 40–62% of FL. Spurs sword-like, long, gradually tapering from the base to the points; SL is 115–181% of SSW. Three short unsegmented toes. No eyespots. Throat and straight esophagus of moderate size. Trophi small, round, 11–12 μ m long and 13–15 μ m wide; 2/2 major teeth and 26/26 minor teeth in unci. Oviparous; egg oval, 101 \times 46 μ m, smooth, 1–6 round knobs on both poles and the sides.

Measurements See Table 2. Body length 120 μ m (possibly in contracted state) according to Velasco-Castrillón et al. (2014a), and 325 μ m according to Darnall & Hollowday (1985).

Distribution Maritime Antarctica: AI*, KG*, SI. Continental Antarctica: EB, VL (CR, CH, GH*, MP*), possibly also DM and MM (Dougherty & Harris, 1963; Sohlenius et al., 1996). *Habitat* Algal and cyanobacterial mats in wetlands; terrestrial moss, soil.

Adineta editae sp. nov. Iakovenko (Figs. 3b, 5)

Darnall & Hollowday, 1985 (*A. gracilis*), p. 31, Fig. 24c. Fontaneto et al., 2008 (*A. gracilis*), p. 3139. Velasco-Castrillón et al., 2014a (*A. cf. gracilis* Bd8), p. 8 (main text); 1, Figs. 2–5 (Annex S1).

Table 2 Body dimensions (measured from light microscope photographs) and trophi dimensions (measured from SEM photographs) of the described bdelloid species

Species	Measurements (μm)						
	TL ^a	BW	HL	HW	CW	NL	
<i>Adineta coatsae</i> sp. nov.	220–292, 257 \pm 27 ^b	39–62, 52 \pm 9	35–55, 42 \pm 7	28–46, 36 \pm 6	–	30–49, 40 \pm 7	
Holotype	248	56	44	38	–	40	
<i>A. editae</i> sp. nov.	206–382, 285 \pm 41	40–90, 58 \pm 12	39–56, 45 \pm 4	30–47	–	34–63, 45 \pm 9	
Holotype	340	90	56	35 \pm 5	–	55	
<i>A. emsleitzi</i> sp. nov.	206–352, 294 \pm 44	40–87, 69 \pm 15	32–61, 49 \pm 8	27–47, 39 \pm 5	–	34–72, 50 \pm 10	
Holotype	332	84	50	38	–	47	
<i>A. grandis</i>	304–505, 414 \pm 61	60–152, 94 \pm 22	57–92, 70 \pm 9	48–71, 56 \pm 7	–	43–118, 82 \pm 19	
Type locality	500–505	81–90	73–92	64–71	–	84–103	
<i>A. fontanetoi</i> sp. nov.	471–509	81–98	67	51–61	–	79–87	
Holotype	471	98	67	51	–	79	
<i>Habrotracha antarctica</i> sp. nov.	225–299, 270 \pm 25	35–58, 48 \pm 8	29–36, 33 \pm 2	28–37, 33 \pm 3	25–32, 29 \pm 2	37–54, 46 \pm 6	
Holotype	249	40	31	31	27	88	
<i>H. devetteri</i> sp. nov.	209–282, 252 \pm 26	38–60, 48 \pm 7	22–31, 27 \pm 6	29–34, 32 \pm 4	30–41, 35 \pm 7	37–56, 46 \pm 7	
Holotype	216	38	22	29	30	42	
<i>H. vermadskii</i> sp. nov.	216–298, 256 \pm 17	29–62, 43 \pm 8	32–35, 33 \pm 1	28–38, 33 \pm 2	22–28, 25 \pm 1	38–61, 50 \pm 6	
Holotype	258	41	32	28	22	47	
<i>Macrotrachela donneri</i> sp. nov.	252–415, 325 \pm 44	39–87, 59 \pm 13	33–48, 41 \pm 6	41–52, 46 \pm 4	46–58, 52 \pm 5	38–72, 53 \pm 11	
Holotype	306	61	43	43	48	47	
<i>M. ioannae</i> sp. nov.	284–488, 409 \pm 80	38–85, 61 \pm 18	55–59	59–60, 59 \pm 1	60–64, 61 \pm 2	50–83, 64 \pm 11	
Holotype	283	38	59	60	60	50	
<i>M. jankoi</i> sp. nov.	245–570, 327 \pm 88	43–107, 61 \pm 18	34–54, 43 \pm 7	33–64, 45 \pm 9	40–73, 52 \pm 10	52–91, 61 \pm 13	
holotype	265	56	37	41	55	55	
<i>Philodina darnnalis</i> sp. nov.	226–349, 275 \pm 44	45–87, 57 \pm 12	27–38, 33 \pm 4	21–31, 26 \pm 25	37–55, 45 \pm 8	39–62, 48 \pm 7	
Holotype	226	49	36	42	50	45	
<i>Ph. shackletoni</i> sp. nov.	360–507, 451 \pm 64	66–148, 94 \pm 37	63–87, 75 \pm 17	71–120, 96 \pm 35	83–131, 107 \pm 33	60–118, 87 \pm 30	
Holotype	474	71	63	71	83	84	

Table 2 continued

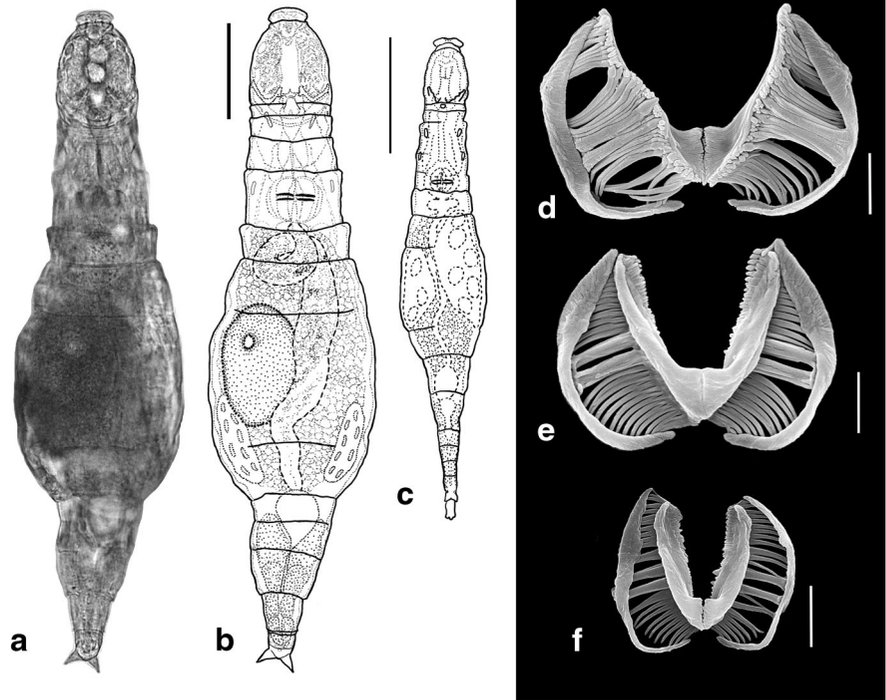
Species	Measurements (μm)						
	RL	RW	FL	SL	SSW	RaL	RaL
<i>Adineta coatsae</i> sp. nov.	29–47, 37 \pm 7	25–38, 31 \pm 5	28–35, 31 \pm 2	9–16, 12 \pm 2	7–11, 8 \pm 1	11.1–12.3	–
Holotype	31	28	30	13	7	–	–
<i>A. editae</i> sp. nov.	30–58, 40 \pm 8	22–43, 33 \pm 6	28–48, 36 \pm 6	6–10, 7 \pm 1	7–13, 10 \pm 1	15.0–18.6, 16.7 \pm 1	–
Holotype	52	44	35	10	13	–	–
<i>A. emsleitzi</i> sp. nov.	27–55, 41 \pm 8	23–48, 35 \pm 7	27–46, 36 \pm 5	6–9, 7 \pm 1	8–12, 9 \pm 1	14.7–19.3, 15.7 \pm 1	–
Holotype	49	48	39	8	9	–	–
<i>A. grandis</i>	36–72, 52 \pm 10	27–64, 46 \pm 10	28–69, 51 \pm 12	7–14, 11 \pm 2	9–17, 13 \pm 2	23.2–31.0, 25.4 \pm 1	–
Type locality	53–68	54–64	61–65	10–12	16–17	24.8–31.0	–
<i>A. fontanetoi</i> sp. nov.	56–59	51–58	49–63	8–10	13–14	24.1–28.3, 26.8 \pm 1	–
Holotype	59	58	63	8	13	–	–
<i>Habrobrocha antarctica</i> sp. nov.	29–39, 34 \pm 4	27–37, 32 \pm 4	22–37, 30 \pm 5	6–9, 7 \pm 1	8–15, 11 \pm 2	14.5–19.3	–
Holotype	29	31	29	7	11	–	–
<i>H. devetteri</i> sp. nov.	27–42, 35 \pm 5	27–37, 32 \pm 4	23–33, 26 \pm 3	4–9, 6 \pm 1	6–13, 10 \pm 2	16.0–17.0	–
Holotype	37	34	30	5	12	–	–
<i>H. vernadskii</i> sp. nov.	26–46, 37 \pm 4	24–43, 32 \pm 5	19–34, 24 \pm 4	6–11, 9 \pm 1	10–17, 13 \pm 2	15.0–17.4	–
Holotype	39	30	23	10	15	–	–
<i>Macrotrachela donneri</i> sp. nov.	34–51, 42 \pm 6	27–46, 37 \pm 7	24–38, 30 \pm 4	9–14, 11 \pm 1	9–13, 11 \pm 1	21.2–24.8, 23.3 \pm 1	–
Holotype	40	39	30	11	10	–	–
<i>M. ioannae</i> sp. nov.	41–72, 55 \pm 11	33–64, 55 \pm 11	31–37, 35 \pm 2	10–14, 13 \pm 2	13–21, 19 \pm 3	25.9	–
Holotype	41	33	33	14	21	–	–
<i>M. jankoi</i> sp. nov.	32–74, 46 \pm 11	28–54, 39 \pm 8	19–46, 28 \pm 8	5–8, 7 \pm 1	9–15, 12 \pm 2	18.1–21.9, 19.6 \pm 1	–
holotype	55	45	32	5	9	–	–
<i>Philodina darnnalis</i> sp. nov.	32–63, 40 \pm 10	26–47, 34 \pm 6	21–47, 30 \pm 10	5–10, 7 \pm 1	9–16, 12 \pm 2	24.7	–
Holotype	32	26	21	6	9	–	–
<i>Ph. shackletoni</i> sp. nov.	60–78, 69 \pm 9	46–66, 55 \pm 10	39–76, 58 \pm 15	17–24, 15 \pm 3	11–21, 15 \pm 4	24.2	–
Holotype	69	53	59	19	14	–	–

^a See explanations in “Materials and methods”

TL total length, BW body width, HL head length, HW head width, CW corona width, NL neck length, RL rump length, RW rump width, FL foot length, SL spur length, SSW spur pseudosegment width, RaL ramus length

^b Min–max, mean \pm SD

Fig. 5 *Adineta editae* sp. nov. (Antarctica): **a**, **b** holotype, habitus, dorsal view; **d** paratype, trophi, cephalic view; **e** paratype, trophi, caudal view. *A. gracilis* (Europe, PL0924): **c** habitus, dorsal view; **f** trophi, caudal view. Scale bar 50 μ m (**a–c**) or 5 μ m (**d–f**)



Type locality Rocka Islands (Argentine archipelago), 15 m asl, S65°10.738', W64°29.522'

Type habitat Soil.

Type material Holotype: SIZ 53.1 (MRockaAED1a, 15.02.2010, Leg. K. Janko), mounted in glycerin jelly. Paratypes: SIZ 53.2-7 (MRockaAED2-7), in glycerin jelly on a separate slide; SIZ 53.8-21 (MRockaAED1b-e, MRockaAED8-18), trophi mounted for SEM. *Additional material* SIZ 53.22-25 (870_1AED1, V12AED1-3), trophi mounted for SEM; SIZ 53.26-33 (870_1AED2, VRA01AED2-3, VS03AED1-3), digital photos and videos.

Etymology Named after the Czech biologist Dr. Edita Drdová-Janková, wife of the collector and project leader Dr. Karel Janko.

Barcodes Gen Bank ID EF173189-91, EF173193, KJ543598-600, see also Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:21EBEA58-A1F3-4D91-9249-CA4E6607986B.

Diagnosis Resembles *A. gracilis* s. str. Janson, 1893 (Fig. 5c, f) and *A. bartosi* Wulfert, 1960 known from Europe, by the short narrow rostrum, the arcuate rostral lamella not divided into lobes and without long sensory bristles underneath and the structure of rakes. Unlike other *Adineta*, all three mentioned species have

rod-like, V-shaped rakes with only two claw-like teeth in each rake directed toward each other. The new species differs from *A. gracilis* and *A. bartosi* by the shape of the head and spurs. The first head pseudosegment not bubble-like swollen as in *A. bartosi*. Head not elongated in the proximal part and not widened (hexagonal) in the distal part as in *A. gracilis* s. str. The head of the new species is larger and wider than in *A. gracilis* s. str. Differently from *A. gracilis* s. str., the new species has somewhat larger and stouter body. Spurs of the new species narrow conical, pointed, with short interspace, while *A. gracilis* s. str. has isocetes triangular spurs without an interspace, and the spurs of *A. bartosi* are narrow, peg-like and without an interspace. Trophi much larger than in *A. gracilis* s. str. with larger number of minor teeth in unci.

Description Body of moderate size, dorsoventrally flattened, transparent, colorless except the yellow-brown stomach. Integument smooth, thin, without knobs, spines or bolsters. Rostrum very short, of moderate width; its lamella wide, semicircular, not divided into lobes. Two claw-like sharp teeth pointing toward each other in each thin rod-like rake. Head wide oval, of regular shape or slightly narrowed toward rostrum, HL is 13–19% of TL. Neck rather short and

wide, NL 34–62% of TL, antenna about 1/3 of the bearing pseudosegment width. Trunk wide, oval. Rump somewhat swollen in the middle part, RL is 11–16% of TL. Foot of moderate length, five pseudosegments, FL is 10–16% of TL. Spurs short, conical, pointed, divergent, with tiny interspace, SL is 60–83% of SSW. Three short unsegmented toes. No eyespots. Trophi round, 15–19 μm long and 16–20 μm wide; 2/2 major and 28–34 minor teeth in unci. Oviparous. Eggs oval, smooth without knobs or spines. Egg size 71–89 \times 45–61 μm by our data and 70 \times 50 μm as reported by Dartnall & Hollowday (1985).

Measurements See Table 2. TL 300 μm by Dartnall & Hollowday (1985) and 220–300 μm according to Velasco-Castrillón et al. (2014a).

Distribution Maritime Antarctica: AI*, SI, AP*. Continental Antarctica: DM, EB, LH, MM, VL (Cz*, CR*), WK.

Habitat Soil, terrestrial moss and lichens, pools.

Comments Most likely all the researchers, except Murray (1910), have been reporting this species under *A. gracilis*—which, in spite of presumed cosmopolitanism, is very unlikely to inhabit dry and cold Antarctic, being a strict acidophile most common in sphagnum bogs (Bērziņš, 1987).

The head of the new species is 45 ± 4 μm long and 34 ± 4 μm wide, HW/HL is 69–90%. According to our data, *A. gracilis* s. str. has the head 40 ± 7 μm long and 29 ± 4 μm wide ($N = 42$), HW/HL is 53–70%. By our data, *A. gracilis* s. str. has TL 247 ± 45 μm ,

BW/TL 13–23%, RW/RL 54–82%, FW/FL 27–40% ($N = 42$). The new species TL is 286 ± 41 μm , BW/TL is 13–23%, RW/RL is 67–99%, and FW/FL is 38–58%. By our data, the trophi of *A. gracilis* 11.1 ± 0.4 μm long, 13.7 ± 0.9 μm wide ($N = 14$), 20–24 minor teeth in each unci. The new species has trophi of 16.6 ± 1 μm long and 18.4 ± 0.7 μm wide, with 28–34 minor teeth in each unci.

Adineta emslii sp. nov. Fig. 6a, b, d, e

Dartnall, 1995a (*A. sp.*), p. 13, Fig. 7a. Velasco-Castrillón et al., 2014a (*A. sp.* Bd1), p. 8.

Type locality Cape Royds (Ross Island), 27 m asl, S77°32.500', E166°8.933'.

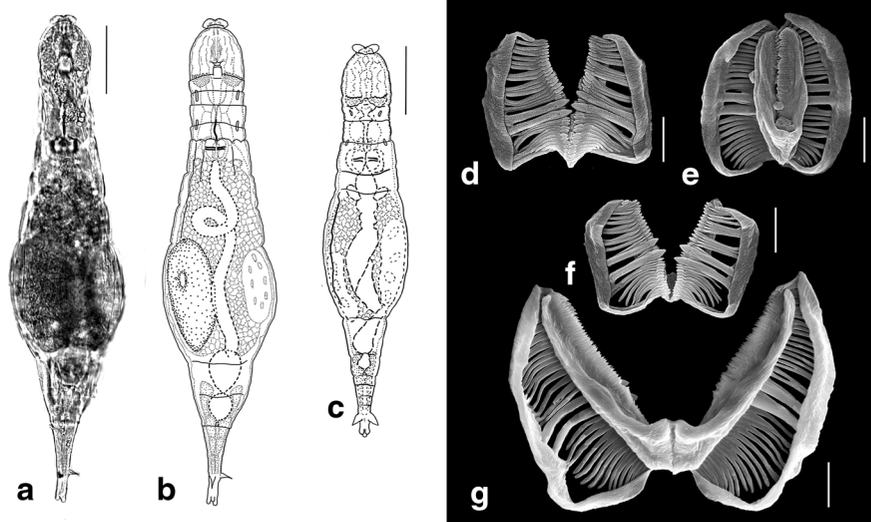
Type habitat Cyanobacterial mats in wetlands.

Type material Holotype SIZ 52.1 (CR23matAE1a, 14.1.2010, Leg. J. Smykla) mounted in glycerin jelly. Paratypes: SIZ 52.2–52.16 (CR23matAE2–16) in glycerin jelly on a separate slide; SIZ 52.16–17 (CR23matAE1b–c), trophi mounted for SEM. **Additional material** SIZ 52.18–52.22 (CR23matAE1e–j, CBM 2AE1), trophi mounted for SEM; SIZ 52.23–31 (CR23matAE17–21, CBM2matAE1–4), digital photos and videos.

Barcodes Gen Bank ID KJ543570–80, see also Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:45FA650B-0086-4E3C-BCCD-A3F228E987F3.

Fig. 6 *Adineta emslii* sp. nov. (Antarctica): **a**, **b** holotype, habitus, dorsal view; **d** paratype, trophi, cephalic view; **e** paratype, trophi, caudal view. *A. vaga* (Europe, PL0838): **c** habitus, dorsal view; **f** trophi, cephalic view. *A. grandis* Murray, 1910 (Antarctica): **g** trophi, caudal view. Scale bar 50 μm (a–c) or 5 μm (d–g)



Etymology Named in honor of the leading Antarctic researcher Dr. Steven D. Emslie for his invaluable support in the Ross Sea project.

Diagnosis Resembles *A. grandis* by the bright orange body color, but it is smaller and not viviparous. By our data, the new species is larger than the similar oviparous species *A. vaga* s. str. (Davis, 1873) (Fig. 6c). Trophi size is intermediate between *A. vaga* s. str. (Fig. 6f) and *A. grandis* (Fig. 6g). Spurs are needle-like with bulb-like swollen bases, while *A. vaga* s. str. has straight triangular spurs. From *A. vaga major* Bryce, 1893, and *A. vaga minor* Bryce, 1893, the new species differs by the shape of the spurs, and the intermediate head size (it is larger than *A. vaga minor*, but smaller than *A. vaga major*). From *A. vaga* s. lat. the new species differs by the orange body (*A. vaga* s. lat. is colorless inclusive stomach).

Description Body of moderate size, wide, flattened, of bright orange color. Integument smooth, thin, transparent, without knobs, spines, bolsters or other appendages. Rostrum short, sickle-like, distal rostral pseudosegment not plate-like flattened. Two short semicircular rostrum lobes, no stiff sensory bristles, only short cilia under the lobes. Wide-oval head of a moderate size, HL is 13–18% of TL, HW is 71–94% of HL. Six thin peg-like teeth in each massive scoop-like rake. Neck of moderate length and width, slightly contracted behind the head, NL is 14–21% of TL, antenna about 1/3 of the bearing pseudosegment width. Trunk oval, wide, BW is 19–27% of TL. Rump conical, somewhat swollen in the middle, RL is 11–16% of TL, RW is 74–98% of RL. Relatively short slim foot of five pseudosegments, FL is 10–16% of TL, FW is 29–45% of FL. Spurs short (SL 60–94% of SSW), pointed, needle-shaped with bulb-like swollen bases, divided by straight interspace of ~ 2 spur widths. Three short unsegmented toes. No eyespots. Trophi ramate, round, 15–18 μm long and 14–18 μm wide. Rami massive, the region of articulation is straight, protruding backwards, without incisure. Interior margins of rami with long numerous peg-like scleropili. Manubria thin, sickle-like. Two major teeth and 29–33 minor teeth in each uncus. Throat small, esophagus short, straight. Stomach glands of moderate size. Eight nuclei (3–7 according to Murray) in each germovitellarium. Oviparous. Eggs oval, 60–70 \times 39–44 μm , shell smooth, without knobs or spines.

Measurements See Table 2. TL 350 μm according to Dartnall (1995a, b).

Distribution Maritime Antarctica: AI*. Continental Antarctica: EB, HI, VL (CR*, CB*, MP*), WK.

Habitat Cyanobacterial mats wetlands, terrestrial moss, soil.

Comments According to our data, the new species has TL $294 \pm 44 \mu\text{m}$, while TL is $414 \pm 61 \mu\text{m}$ in *A. grandis* ($N = 20$) and $274 \pm 14 \mu\text{m}$ in *A. vaga* s. str. ($N = 15$). The new species has trophi $15.7 \pm 1.1 \mu\text{m}$ long with 29–32 minor teeth in each uncus, while *A. grandis* has trophi $25.4 \pm 1.4 \mu\text{m}$ long with 36–44 minor teeth ($N = 53$), and *A. vaga* s. str. has it $13 \pm 0.7 \mu\text{m}$ long with 25–27 minor teeth ($N = 14$).

Adineta grandis Murray, 1910 (Figs. 3c, 6g, 7a)

Murray (1910, pp. 51–53, Pl. XII Fig. 10). Voigt (1956–1957, p. 71, Taf. 5 Abb. 24, Taf. 8 Abb. 19, Taf. 14 Abb. 16). Donner (1965, p. 273, Fig. 200a). Donner (1972a, p. 252, Abb. 1). Koste (1996b) (as *A. grandis*, but most likely sibling species), p. 243, Abb. 5. Dartnall & Hollowday (1985, p. 31, Fig. 24d–f). Kutikova (2005, p. 275, Ris. 299). Velasco-Castrillón et al. (2014a) (*A. sp.* Bd2): 8 (main text); 2, Fig. 8 (Annex S1).

Type locality Cape Royds

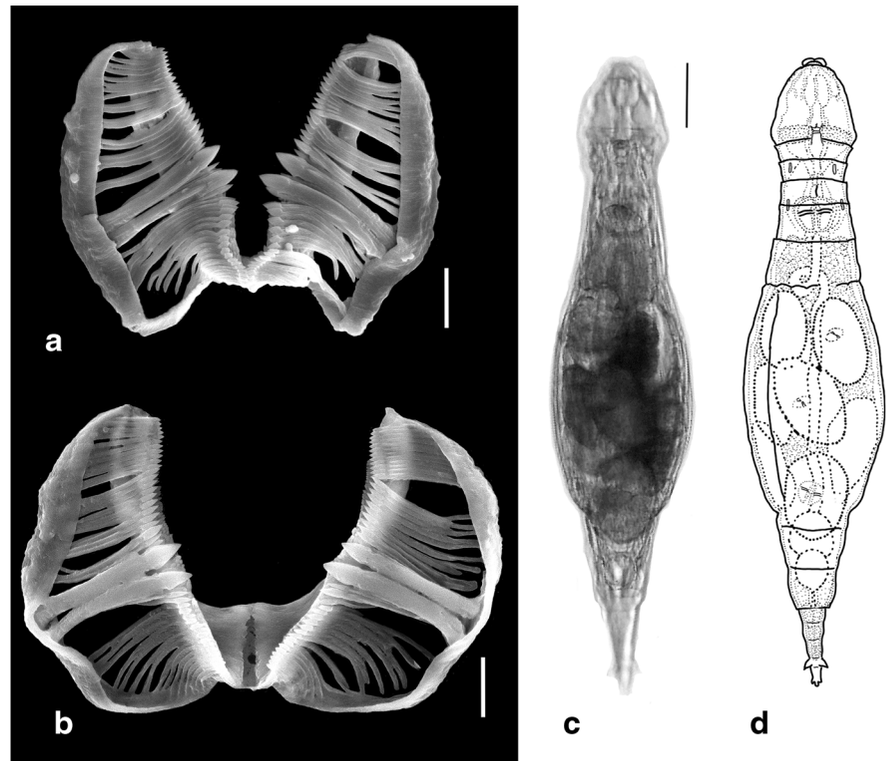
Type habitat “Brown vegetation” (algae?) in lake.

Barcodes GenBank ID KJ543581-88, see also Supplementary file III.

Material examined BI11, 1 ind.; BI23, 10 ind.; CBM1CYmat, 6 ind.; CBM2mat, 4 ind.; CBC1mat, 5 ind.; CBPc2mat, 1 ind.; CRL21, 2 ind.; CR24, 2 ind.; CRL24, 1 ind.; CzM2CYmat, 8 ind.; CzM3CYmat, 9 ind.; EPL23, 11 ind.; MPM3, 1 ind.; MPM5, 34 ind.; MPM5CYmat, 10 ind.

Description The largest species of the genus, and the only known viviparous one. Reported TL is 306–750 μm (Murray, 1910; Donner, 1965; Dartnall & Hollowday 1985) and 304–505 μm according to our data. Its foot is shorter than in other species of *Adineta*. Trophi length 23–29 μm (our data). Body pale orange or brownish yellow, sometimes reddish (“light brown or yellowish, darker in the alimentary tract” according to Murray). Integument smooth, thin, transparent, without knobs, spines or other appendages. Rostrum short, of moderate width, distal rostral pseudosegment not strongly widened or flattened. Rostral lamella divided into two small semicircular lobes. No stiff sensory bristles under rostrum lobes, only short soft

Fig. 7 *Adineta grandis*: **a** trophi, cephalic view. *A. fontanetoi* sp. nov.: **b** paratype, trophi, cephalic view; **c, d** holotype, habitus, dorsal view. Scale bar 50 μm (**c, d**) or 5 μm (**a, b**)



cilia. Head not large (HL is 13–19% of TL), wide oval (“ovate” by Murray), tapering toward rostrum, HW is 66–97% of HL. 6–10 teeth in each massive scoop-like rake. Neck massive, long (NL is 11–27% of TL). Dorsal antenna thick, about 1/4 of width of the antennal pseudosegment. Trunk wide (its width depends on the number of embryos inside), BW is 16–31% of TL. Rump conical, with both pseudosegments somewhat swollen laterally (in some specimens the lateral swellings look like knobs), gradually tapering into a very short narrow foot. RL is 7–16% of TL, RW is 74–103% of RL. Foot short, of 5 pseudosegments, FL is 6–15% of TL, FW is 40–56% of FL. Spurs conical, widened at the base (according to Murray, “short broad cones,” “stout and subacute”), pointed, narrow, divergent, divided by the straight interspace equal to 1–2 spur widths, SL is 60–98% of SSW. Three short unsegmented toes. No eyespots. Trophi ramate, large, round or elongate. Rami massive, interior margin with numerous peg-like scleropili. Articulation protruding to the ventral part, straight and without incisure. Manubria wide, flat, crescent-shaped. Major uncinal teeth thick, dental formula 2/2; 38–41 minor teeth. Trophi unusually large for *Adineta*: 30 μm long

according to Donner (1965), 24–31 μm long and width is equal to the length, according to our data. Throat voluminous, esophagus short, straight. Stomach glands large. Eight nuclei in each of germovitelaria. Viviparous, up to 4 embryos with developed trophi can be seen inside trunk.

Measurements See Table 2. TL up to 750 μm according to Murray (1910).

Distribution Maritime Antarctica: SI. Continental Antarctica: EB, HI, MM, VL (BI*, CB*, CC*, CR, Cz*, EP*, MP*). Africa (questionable): Madagascar (Koste, 1996a).

Habitat Algal mats and sediment in pools and seepages, soil, terrestrial moss.

Comments Velasco-Castrillón et al. (2014a) erroneously attributed this rotifer to “wheel-bearers” (although *A. grandis* has no trochi), and described it as “ovoviviparous” although the species is viviparous.

Adineta fontanetoi sp. nov. Fig. 7b–d

Type locality Beaufort Island, 9 m asl, S76°58.147', E166°54.217'.

Type habitat Soil.

Type material Holotype: SIZ 54.1 (BI27AG1a, 29.1.2010, Leg. J. Smykla), mounted in glycerin jelly. Paratypes: SIZ 54.2-4 (BI27AG1b-d), SIZ 54.5-22 (BI27AG2-19), trophi mounted for SEM.

Etymology The species is named after colleague rotiferologist Dr. Diego Fontaneto who first sequenced this species (as *A. grandis*).

Barcodes GenBank ID EF173184-85, KP869896.

ZooBank LSID urn:lsid:zoobank.org:act:15A138A9-A20D-41BE-A7E6-0EC4BF2F59B0.

Diagnosis By external morphology the new species does not differ from *A. grandis* (see the description above) and may be easily confused with the latter under the light microscope. However, it has somewhat larger trophi (ramus length mean \pm SD $26.8 \pm 1.2 \mu\text{m}$ in *A. fontanetoi* sp. nov. and $25.4 \pm 1.4 \mu\text{m}$ in *A. grandis*).

Description Viviparous. 8 teeth in each rake. Trophi 24–28 μm long and 27–28 μm wide; 2/2 major uncinial teeth, 38–43 minor teeth in the left uncus and 39–42 in the right one.

Measurements See Table 2.

Distribution Maritime Antarctica: SI. Continental Antarctica: BI*.

Habitat Soil.

Order Philodinida Melone & Ricci, 2005

Family Habrotrochidae Bryce, 1910

Genus *Habrotrocha* Bryce, 1910

***Habrotrocha antarctica* sp. nov. (Figs. 3d, 8)**

Murray, 1910 (*Callidina constricta*), pp. 48–49, Pl. XII Fig. 13a, b; Dartnall & Hollowday, 1985 (*H. constricta*), p. 32; Fig. 25a–c. Velasco-Castrillón et al., 2014c (Bd12), p. 8.

Type locality Cape Royds, 18 m asl, S77°32.532', E166°8.855'.

Type habitat Soil.

Type material Holotype: SIZ 56.1 (CRL23HE1a, 14.01.2010, Leg. J. Smykla), mounted in glycerine jelly, encircled in green ink. Paratypes: SIZ 56.2-4 (CRL23HE1b-d), on the same slide as holotype, encircled in black ink; SIZ 56. 5-11 (CRL23HE1e-k), trophi mounted for SEM. *Additional material* SIZ 56.12-23 (EPL24M51-5, CzL4CYmatHE1-7), digital videos and photos; SIZ 56.24 (CzL4CYmatHE8), trophi mounted for SEM.

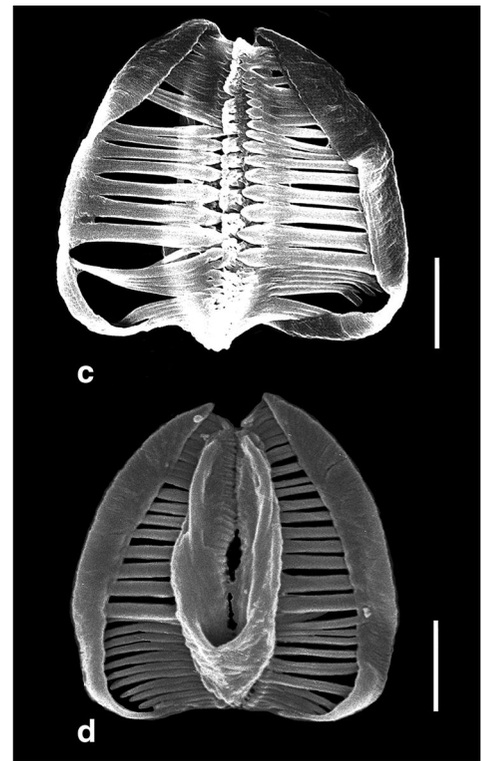
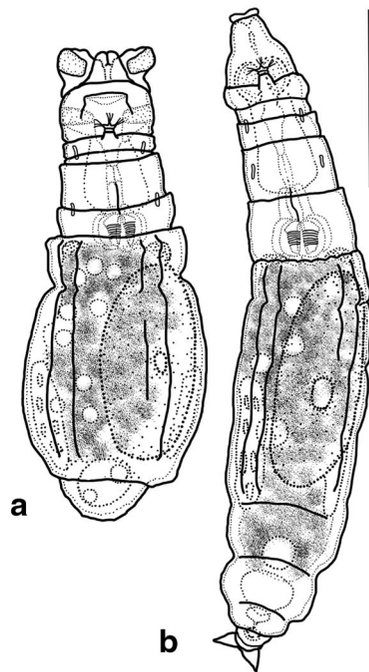
Etymology Named after the Antarctic continent where it was first found.

Barcodes GenBank ID EF650588-90, KJ543609-11, see also Supplementary file III. *ZooBank LSID*. urn:lsid:zoobank.org:pub:C3618A2A-F710-4318-B8EA-58C285EC6CDA.

Diagnosis Very similar to *H. elusa* s. lat. Milne, 1916, except of the integument sculpturation, rump shape and trophi structure. The foot is much wider and the spurs thicker and longer than in *H. elusa vegeta* Milne, 1916. Differently from *H. elusa* s. str. Milne, 1916, it has no lateral knobs on the first rump pseudosegment. The integument on the trunk and rump is not dotted or granulated, unlike it is reported for *H. elusa* s. str. (Milne, 1916; Donner, 1965). The same as in *H. elusa* s. str., corona width of the new species is almost equal to the cingulum, while in *H. elusa vegeta* it is substantially narrower. Similar to *H. constricta* by the size and body shape, however distinguished by the upper lip with a notch in the middle (so that the tip is divided into two small lobes), while in *H. constricta* the tip is whole. Dental formula 7/7 major uncinial teeth (the last 2–3 thinner than the rest), while in *H. elusa* s. str. it is reported to be 6/6. In *H. elusa vegeta* it is 3 + 4/3 + 4 and in *H. constricta* usually has 6/6 major teeth in unci and rarely 7/7 or 8/8 (Donner, 1965). Murray (1910) reports this species to have 4/4 thicker teeth, succeeded by several finer ones, what can be sometimes observed in our specimens.

Description Body of moderate size, transparent, spindle-shaped, colorless but usually with yellow-brown or bright orange stomach. Integument smooth, thin, without knobs, ribs or spines. No knob on the first foot pseudosegment. Rostrum short, lamella divided into two small semicircular lobes. Corona narrower than the oval head base, CW/HW 79–91%, HW is 96–100% of HL. Pedicels short, straight, divided by a narrow sulcus without membrane or ligula. Trochal discs kidney shaped in apical view. No papillae or sensory bristles on trochi. Upper lip triangular, reaching plane of trochal discs, upper rim thickened by cuticular bolster, tip divided by a notch into two small rounded lobes. Lower lip not wide, not projecting laterally. Cingulum bolster very narrow. Neck of moderate length and width, NL is 15–35% of TL. Trunk slim, BW is 15–20% of TL. Rump conical, first pseudosegment swollen, RL is 11–15% of TL, RW is 80–110% of RL. Foot very short, 4 pseudosegments,

Fig. 8 *Habrotrocha antarctica* sp. nov.: **a** holotype, habitus, feeding, dorsal view; **b** same, creeping, dorsal view; **c** paratype, trophi, cephalic view; **d** paratype, trophi, caudal view. Scale bar 50 μm (**a**, **b**) or 5 μm (**c**, **d**)



FL is 10–12% of TL, FW is 49–74% of FL. Spurs short, triangular with elongated narrow tips and slightly swollen middle part, divergent, without interspace, SL is 55–74% of SSW. Three short unsegmented toes. No eyespots. Throat narrow, esophagus short, straight. Stomach glands small, round. Food pellets rounded, small. Trophi ramate, heart-shaped, 15–19 μm long and 17–20 μm wide. Rami thick, with numerous short scleropili along the inner rims. Articulation straight, wide, without incisure. Manubria narrow, sickle-like. Dental formula 7/7 or 4 + 3/4 + 3, with 24–25 minor teeth in each uncus. Oviparous. Eggs oval, 65–70 \times 33–41 μm , shell smooth, without knobs or spines.

Measurements See Table 2. TL 250 μm by Murray (1910) or 375 μm (Dartnall & Hollowday, 1985).

Distribution Maritime Antarctica: SI. Continental Antarctica: EB, MM, VL (CR, EP*, Cz*).

Habitat Soil, algal mats, sediment in pools.

Comments Murray (1910) first depicted this species, but erroneously identified it as *C. (=H.) constricta*. In Murray's image the notch in the middle of the upper lip (absent in *H. constricta*) is clearly visible, and the dental formula seems to be 7/7 or 8/8

(though in the description Murray mentions only 4/4 major teeth). *H. antarctica* sp. nov. was identified as *H. constricta* by Dartnall & Hollowday (1985), but the specimen depicted by these authors has a two-lobed upper lip, while *H. constricta* has only one lobe.

***Habrotrocha devetteri* sp. nov. Iakovenko (Fig. 9a–d)**

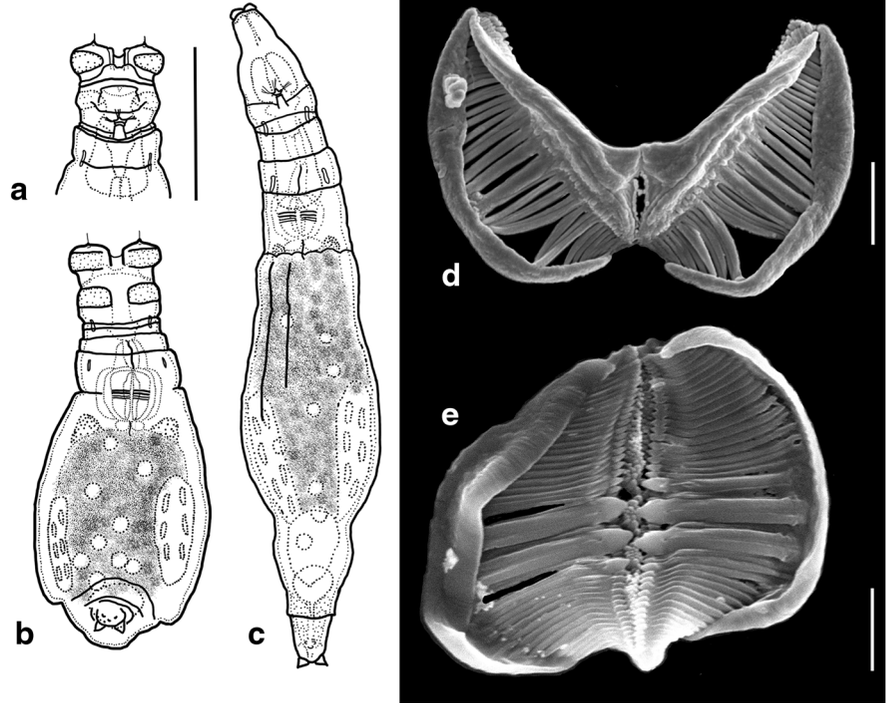
Velasco-Castrillón et al., 2014a (Bd42), p. 8 (main text); 5, Figs. 21–22 (Annex S1).

Type locality Cape Bird (Ross Island), 77 m asl, S77°13.207', E166°26.568'.

Type habitat: Soil.

Type material Holotype: SIZ 57.1 (CBM2HD2a, 19.01.2010, Leg. J. Smykla), mounted in glycerin jelly, incircled with green ink. Paratypes: SIZ 57.2–5 (CBM2HD1, CBM2HD3–5), mounted on the same slide as the holotype, incircled with black ink; SIZ 57.6 (CBM2HT2b), trophi mounted for SEM. **Additional material** SIZ 57.7–14 (CBC4HD1, CBM2HD6–9, CBM2matHD1–3), digital photos and videos; SIZ 57.15 (CzL4CymatHD2), trophi mounted for SEM.

Fig. 9 *Habrotrocha devetteri* sp. nov. (Antarctica): **a** holotype, head, feeding, dorsal view; **b** same, habitus, feeding, ventral view; **c** habitus, creeping, dorsal view; **d** paratype, trophi, caudal view. *H. thienemanni* (Europe): **e** trophi, cephalic view. Scale bar 50 μ m (**a–c**) or 5 μ m (**d, e**)



Etymology Named after colleague rotiferologist Dr. Miloslav Devetter participating in this study.

Barcodes GenBank ID KJ543668-74, see also Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:C39F702E-3A94-4529-879F-7093793196D2.

Diagnosis Similar to *H. thienemanni* s. lat. by the shape and size of the corona, trunk and spurs. Like *H. thienemanni* s. lat., its upper lip has two small lobes; however, the lobes are rounded and divided by a broad interspace, while in *H. thienemanni* s. lat. the lobes are often pointed and divided by a notch. The new species has $2 + 2/2 + 2$ major teeth in the unci, while *H. thienemanni* s. lat. has $2 + 1/2 + 1$ major teeth (Fig. 9d, e). It differs from *H. crassa* Donner, 1949, another species with a two-lobed upper lip, by the corona being wider than the head base, the smooth integument, body outline and dental formula ($4/4$ in *H. crassa*). It differs from *H. tranquilla* Milne, 1916, by its smaller size. The TL of the new species is 209–282 μ m, while for *H. tranquilla* it is 340–402 μ m. The lower lobes of the upper lip are divided by an interspace, while in *H. tranquilla* they are higher and divided by a notch. Dental formula is not $7/7-9/9$ as in *H. tranquilla*.

Description Body of moderate size, spindle-shaped, colorless, transparent. Integument thin, smooth, without knobs, spines or bolsters. Rostrum short, lamella with two small semicircular lobes. Corona wider than the oval head base, CW is 103–118% of HW, HL is 22–31% of TL. Pedicels short, straight. Sulcus very narrow, half-covered with membrane. Trochal discs with papillae and sensory bristles. Upper lip goes up to a half of the pedicels, it is arcuate with two small semicircular lobes divided by an interspace. Lower lip slightly protruding laterally. Cingulum narrow. Neck of moderate length, NL is 15–26% of TL. The length of the antenna is about $1/3$ of the bearing pseudosegment width. Trunk plump, BW is 17–30% of TL. Rump conical, RL is 12–17% of TL, RW is 81–98% of RL. Foot short, slim, 4 pseudosegments, FL is 8–14% of TL, FW is 51–88% of TL. Spurs short, triangular, divided by interspace as broad as one spur width, SL is 44–76% of SSW. Three short unsegmented toes. No eyespots. Throat and esophagus short, food pellets round, of moderate size. Stomach glands of medium size. Trophi ramate, heart-shaped, 16–19 μ m long and 16–19 μ m wide. Rami thin, with numerous short scleropili along the inner rim, articulation straight and without incisure. Manubria thin, sickle-like. $2 + 2/$

2 + 2 major teeth, 30–33 minor teeth in each uncus. Oviparous, egg oval, $67 \times 39 \mu\text{m}$, shell smooth, without knobs or spines.

Measurements See Table 2. TL 250–360 μm (Velasco-Castrillón et al., 2014a).

Distribution Continental Antarctica: EB, VL (CC*, CB*), WK.

Habitat Soil, algal mats.

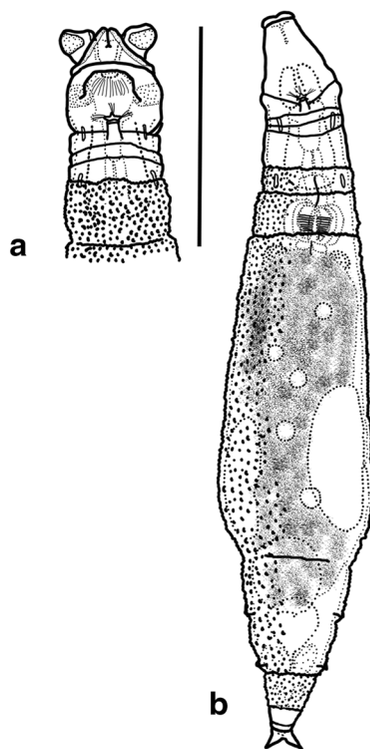
Habrotrocha vernadskii sp. nov. (Figs. 3e, 10)

Type locality Galindez Island (Argentine Islands archipelago), 4 m asl, S65°15.060', W64°14.558'.

Type habitat Soil.

Type material Holotype: SIZ 58.1 (V12HE2a, 1.03.2010, Leg. K. Janko), mounted in glycerin jelly, encircled in green ink. Paratypes: SIZ 58.2-4 (V12HE2a-c), on the same slide as holotype, encircled in black ink; SIZ 58. 5-8 (V12HE4a-b, V12HE13a-b), trophi mounted for SEM. **Additional material** SIZ 58. 9-15 (CCA2HE1-2, CCA4HE1-5), digital photos and videos.

Fig. 10 *Habrotrocha vernadskii* sp. nov.: **a** holotype, head, feeding, dorsal view; **b** same, habitus, creeping, dorsal view; **c** paratype, trophi, cephalic view; **d** paratype, trophi, caudal view. Scale bar 50 μm (**a**, **b**) or 5 μm (**c**, **d**)

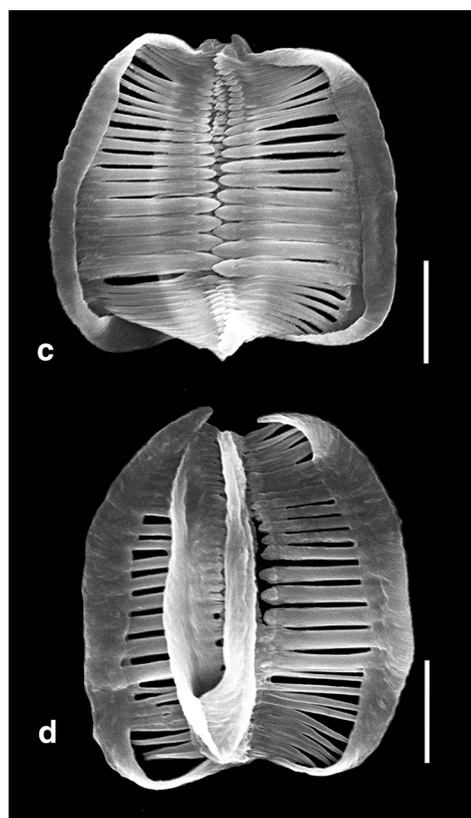


Etymology Named after the Ukrainian polar research base “Academician Vernadsky” in the vicinity of which it was found.

Barcodes GenBank ID—see Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:6FF7FEA-F-BC8C-46FB-AA46-F09ADF2D2147.

Diagnosis Resembles *H. elusa* s. str. Milne, 1916 by the yellowish body with thicker integument on trunk and rump, the triangular upper lip with the tip divided by a notch into two rounded lobes, by lateral knobs on the rump and by the short triangular spurs. However, the knobs seem to be smaller and sharper than in *H. elusa* s. str., and their number is 4, not 6. Milne (1916) states that *H. elusa* s. str. has “thick, leathery but smooth skin” that is stippled, but not granulated, on the trunk, rump and foot excluding spurs. Contradictory to this, the new species has a trunk, rump and first foot pseudosegment covered with minute granulae, as in “*H. elusa* s. str.” found by Donner (1965). Similarly to *H. elusa vegeta*, which lacks granulated cuticle, corona of the new species is distinctively narrower than the head base, CW/HW is 71–84%. Milne reports *H. elusa*'s s. str. corona to be



equal to or slightly wider than the head base (CW/HW is 100–111%). Dental formula 9/9 major uncinal teeth (the last two almost as thin as minor teeth), while in *H. elusa* s. str. it is 6/6, and in *H. elusa vegeta* is 3+4/3+4 (Donner, 1965). Differs from *H. crenata* s. lat. by the shape of the upper lip (in *H. crenata* s. lat. the tip of the upper lip is not divided into lobes) and by the pattern of sculpturation. In *H. crenata* s. str. the whole foot and spurs are granulated, and the first foot pseudosegment has a rounded knob absent in the new species. Dental formula of *H. crenata* is 7/7 or 8/8 (Donner, 1965). Differs from *H. antarctica* sp. nov. by the granulated integument, lateral knobs on rump, narrower corona, and slightly longer and narrower spurs. Also, it has a larger number of major and minor teeth in the unci (9/9) than *H. antarctica* sp. nov. (7/7 or 4+3/4+3).

Description Body slim, spindle-shaped, yellowish. Integument granulated on the last neck pseudosegment, trunk, rump and the first foot pseudosegment, and smooth on the rest of the body. Four small pointed lateral knobs on rump (2 on the distal rim of the first and 2 on the second pseudosegment). No knobs on foot. Rostrum very short, lamella with two small semicircular lobes. Corona narrower than rectangular head base, HL is 11–15% of TL. Upper lip triangular, reaches plane of trochal discs, tip divided by a notch into two small semicircular lobes. Trochi without papillae and sensory bristles. Pedicels short, straight. Sulcus narrow, partly covered by prominent retractors of trochi. Lower lip not protruding laterally. Neck rather long, of moderate width, NL is 38–61% of TL. Antenna is 1/3–1/4 of the bearing pseudosegment's width. Trunk narrow, BW is 29–62% of TL depending on the amount of eggs in a female. The first rump pseudosegment swollen, RL is 10–15% of TL, RW is 71–99% of RL. Foot short, 4 pseudosegments, FL is 8–11% of TL, FW is 54–85% of FL. Spurs of moderate length, narrow triangular, bases merged but seem to form short interspace, SL is 56–84% of SSW. Three short unsegmented toes. No eyespots. Throat small, esophagus short, straight. Food pellets small, of irregular shape. Oviparous, eggs oval, shell smooth, without knobs or spines. Trophi ramate, heart shaped, 15–17 µm long and 14–16 µm wide. Articulation straight, without incisure. Numerous short scleropili on inner rims of rami. Manubria narrow, sickle-like. Unci with 9/9 major teeth, gradually diminishing in

thickness, the last ones hardly distinguishable from minor teeth (26–27 in each uncus).

Measurements See Table 2.

Distribution Maritime Antarctica: AI*. Continental Antarctica: CC*.

Habitat Soil, terrestrial moss.

Family Philodinidae Ehrenberg, 1838

Genus *Macrotrachela* Milne, 1886

***Macrotrachela donneri* sp. nov. Fig. 11**

Murray, 1910 (*Callidina habita*): Pl. IX Fig. 3, Pl. XI Fig. 8a. Donner, 1965 (*Macr. insolita* var. 3), p. 132, Fig. 961, m. Donner, 1972a (*Macr. insolita* var.), p. 252, Abb. 2. Iakovenko & Tyshenko, 2006 (*Macr. hewitti*), p. 2, Ris. 2.

Type locality Marble Point (Victoria Land), 4 m asl, S77°25.597', E163°45.148'.

Type habitat Soil.

Type material Holotype: SIZ 59.1 (MPM5MD1a, 25.01.2010, Leg. J. Smykla), on slide in glycerine jelly, encircled with green ink. Paratypes: SIZ 59.2-6 (MPM5MD2-6), in glycerine jelly on the same slide as the holotype, encircled with black ink; SIZ 59.7-9 (MPM5MD1b-d), trophi mounted for SEM. **Additional material** SIZ 59.10-15 (KG1MD1, EPL25MD1, MPL1MD1, MPL3MD1-3), digital photos and videos; SIZ 59.15-18 (MPL3MD1-4), trophi mounted for SEM.

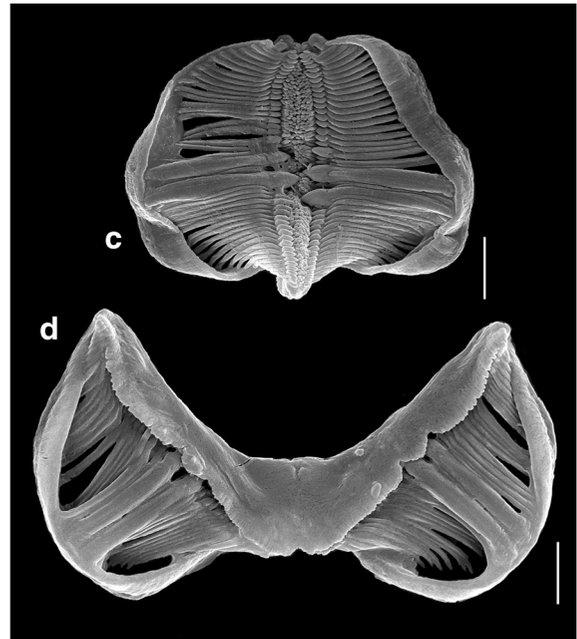
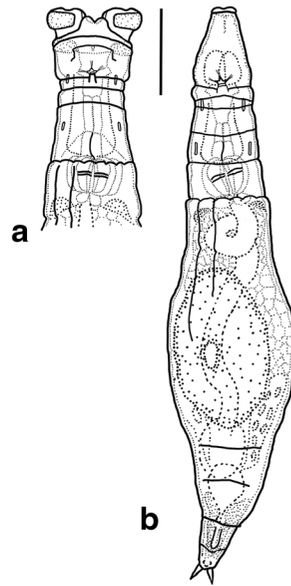
Barcodes GenBank ID KP869998.

ZooBank LSID urn:lsid:zoobank.org:act:86AC6997-0625-4B56-BEC9-FECCF2379EC5.

Etymology Named after the Austrian rotiferologist Dr. J. Donner who first depicted this species as *M. insolita* var.

Diagnosis Similar to *Macr. concinna* (Bryce, 1912), *Macr. habita* (Bryce, 1894), *Macr. hewitti* (Murray, 1911), *Macr. insolita* de Koning, 1947 and *M. plicata* s. str. (Bryce, 1892) by the upper lip with two rounded lobes. Alike *Macr. habita*, *Macr. hewitti* and *Macr. insolita*, the new species has a knob on the first foot pseudosegment, which is lacking in *Macr. concinna* and *Macr. plicata*. Ligula in the sulcus and knobs on rump typical for *Macr. plicata* are absent in the new species. The integument is smooth, while in *M. habita* it is very finely stippled (Bryce, 1894). The foot is rather stout (what distinguishes all three species

Fig. 11 *Macrotrachela donneri* sp. nov.: **a** holotype, head, feeding, dorsal view; **b** same, habitus, creeping, dorsal view; **c** paratype, trophi, cephalic view; **d** paratype, trophi, caudal view. Scale bar 50 μ m (**a**, **b**) or 5 μ m (**c**, **d**)



from *Macr. plicata* s. lat.). Similarly to *Macr. insolita*, the dental formula of the new species is 2/2, while in *Macr. habita* it is 2+1/1+2 (with an additional thinner teeth), and in *Macr. hewitti* it is 4/4 or 5+1/5+1 (Donner, 1965). It is also distinguished by the peg-like or narrow conical spurs with broad interspace—in *Macr. habita* and *Macr. insolita* the spurs are triangular, wide, and flat, the interspace is shorter; in *Macr. hewitti* the interspace is lacking (Donner, 1965; Murray, 1911). Larger than *Macr. insolita* (TL 325 \pm 44 μ m, TL of *Macr. insolita* is about 250 μ m). CW/HW ratio (107–119%) is intermediate between *Macr. habita* (120–125%) and *Macr. insolita* (100–103%), and the head seems to be shorter and wider than in *Macr. insolita*. The upper lip is shorter, and the lobes are more separated from each other than in *Macr. concinna* and *Macr. insolita*. The head base is rectangular, while it seems to be trapezoid in *Macr. habita*, and wide-oval in *Macr. insolita*.

Description Body large, transparent, colorless, but often with bright-orange stomach. Integument smooth, without spines or bolsters, no knobs except a large longitudinal knob on the first foot pseudosegment. Rostrum short, thick, lamella with two semicircular lobes. Head wide, corona wider than head base, CW is 107–119% of HW, HL is 10–15% of TL. Pedicels short, straight. Sulcus as wide as 1/2 of a trochus, covered with protruding trochi

retractors. No papillae or sensory bristles on trochi. Head base rectangular, shorter than its width. Upper lip arcuate with two large semicircular lobes not divided by interspace, reaching 1/2 of the pedicels' height. Lower lip protrudes laterally. Neck of moderate length and width, NL is 15–21% of TL. Length of antenna is about 1/4 of bearing pseudosegment width. Trunk thick, BW is 15–25% of TL. Rump large, swollen, RL is 12–15% of TL, RW is 65–97% of RL. Foot short, of 4 pseudosegments, first pseudosegment with dorsal elongated knob. Spurs rather long, rod-like, pointed, gradually tapering from the base to tips, interspace equal to 2 spur widths, SL is 86–111% of SSW. Three thick unsegmented toes. No eyespots. Throat voluminous, esophagus short, straight. Lumen long, thick, often with a loop. Stomach glands large, round. Trophi ramate, large, heart-shaped, 21–25 μ m long and 24–28 μ m wide. Rami massive, inner rim with numerous scleropili. Articulation long, flat, straight, without incisure. Manubria long, wide, crescent-like. Dental formula 2/2, with 42–43 minor teeth in each uncus. Oviparous. Egg lemon-shaped, shell smooth with two round knobs on each pole.

Measurements See Table 2. TL up to 570 μ m, CW 95 μ m in Murray (1910). According to Donner (1965), TL 410 μ m, CW 66 μ m, SL 29 μ m, trophi 30 μ m long.

Distribution Maritime Antarctica: AI*, KG*, SI. Continental Antarctica: VL (CB*, CC*, CR, Cz*, BI*, EP*, MP*).

Habitat Soil, algal mats in seepages, terrestrial moss.

Comments Murray (1910) first depicted this rotifer from Cape Royds, though he apparently described two different species under the name *C. habita*.

***Macrotrachela ioannae* sp. nov. Iakovenko (Figs. 3h, 12)**

Type locality Rocka Islands (Argentine Islands Archipelago), 15 m asl, S65°10.738', W64°29.522'.

Type habitat Soil.

Type material Holotype: SIZ 60.1 (MRockaMI1a, 15.02.2010, Leg. K. Janko), mounted in glycerin jelly, encircled with green ink. Paratypes: SIZ 60.2-3 (V12MI1-2), on the same slide with holotype, encircled with black ink. **Additional material** SIZ 60.4-6 (MRockaMI2-3, CrulsBMI1), digital photos.

Etymology Named after Mgr. Ioanna Vaňkova, a friend and a specialist in linguistics, who gave much advice on creating Latin names for the new rotifer species.

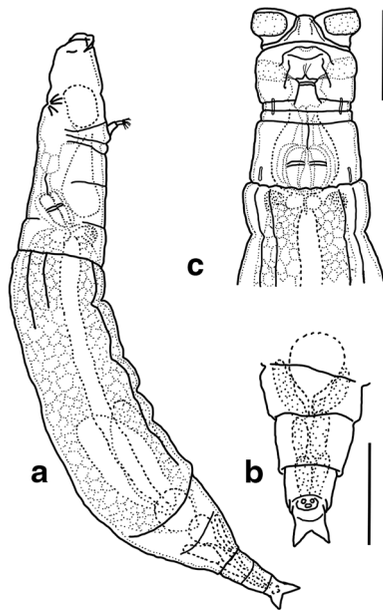


Fig. 12 *Macrotrachela ioannae* sp. nov.: **a** holotype, habitus, creeping, lateral view; **b** same, head, feeding, dorsal view; **c** same, foot, ventral view. Scale bar 50 μ m

Barcodes GenBank ID KP869995-97.

ZooBank LSID urn:lsid:zoobank.org:act:05D141A6-F494-45F3-B144-755BBCAAB31F.

Diagnosis Similar to *Macr. ehrenbergii* (Janson, 1893), *Macr. timida* s. lat., *Macr. induta* Donner, 1951, and *Macr. allani* (Murray, 1911). Differs from all these species by the shape of spurs with papillae-like tips, and dental formula (additional thinner tooth in each uncus). The head base is shorter and wider than in *Macr. ehrenbergii*. The corona is almost equal to the head base, while in *Macr. timida* and *Macr. allani* it is substantially wider. Spurs shorter than the bearing pseudosegment width, while in these species they are, on the contrary, longer. Unlike in *Macr. timida* s. lat., no knobs on foot or rump. Differs from *Macr. induta* also by the shape of the upper lip, which in *Macr. induta* is wide arcuate, with a low rounded lobe in the middle. The new species has a narrow arcuate upper lip with a high trapezoid lobe with a rounded tip in the middle, bearing a thin bolster along the upper rim.

Description Body large, colorless, transparent. Integument smooth, without knobs, spines or bolsters. Rostrum thick, of moderate length, lamella with two semicircular lobes. Corona not wide, CW is 100–108% of HW, HL is 11–21% of TL. Pedicels short, straight. Sulcus of moderate width, without ligula, not covered with membrane or trochi retractors. Head base rectangular, its width larger than height. Upper lip arcuate with single trapezoid lobe, its rounded tip has a bolster along the upper rim. Lower lip not protruding laterally. Neck of moderate length and width, NL is 12–18% of TL. Length of antenna is 1/3 of the bearing pseudosegment's width. Trunk cylindrical, BW is 13–17% of TL. First rump pseudosegment swollen, RL is 12–15% of TL, RW is 78–96% of RL. Foot short, 4 pseudosegments, FL is 8–11% of TL, FW is 52–77 of FL. Spurs short, flat, triangular, divergent, with bases merged and tips separated as small papillae. Three unsegmented toes. No eyespots. Throat voluminous, esophagus short, straight. Lumen wide, with a loop. Stomach glands round, not large. Trophi ramate, heart-shaped, 26 μ m long and 20–21 μ m wide. Dental formula 2 + 1/1 + 2 major teeth (with an additional thinner teeth) and about 30 minor teeth in each uncus. Oviparous. Eggs oval, 124 \times 66 μ m. Egg shell smooth, without knobs or spines.

Measurements See Table 2.

Distribution Maritime Antarctica: AI*.

Habitat Soil, terrestrial moss.

***Macrotrachela jankoi* sp. nov. Iakovenko (Figs. 3f, g, 13)**

Velasco-Castrillón et al., 2014a (Bd7), p. 8.

Type locality Squa Island (Argentine Islands archipelago), 20 m asl, S65°25.117', W64°26.583'.

Type habitat Soil.

Type material Holotype: SIZ 61.1 (VS02MJ1a, 15.03.2010, Leg. K. Janko), mounted in glycerine jelly. Paratypes: SIZ 61.2-3 (VS02MJ2, VS02MJ3), mounted in glycerine jelly; SIZ 61.4 (VS02MJ4), trophi mounted for SEM. *Additional material* SIZ 61.5-17 (V08MJ1, V11MJ1, V12MJ1-2, VRA01MJ1-7, VRA02MJ1), digital photos and videos; SIZ 61.18-26 (V12MJ3, MRockaMJ1-8), trophi mounted for SEM.

Barcodes GenBank ID KJ543594-97, KJ543597, KP869999, KP870000.

ZooBank LSID urn:lsid:zoobank.org:act:6E2BA135-65A8-4256-9B82-54E86D84865B.

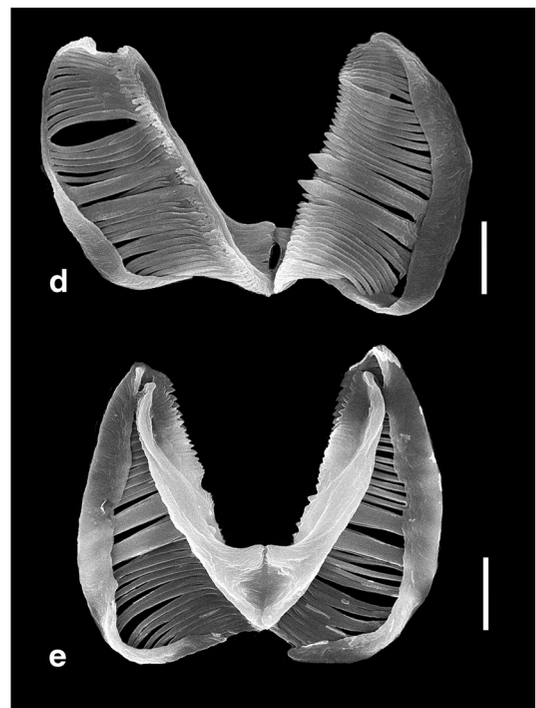
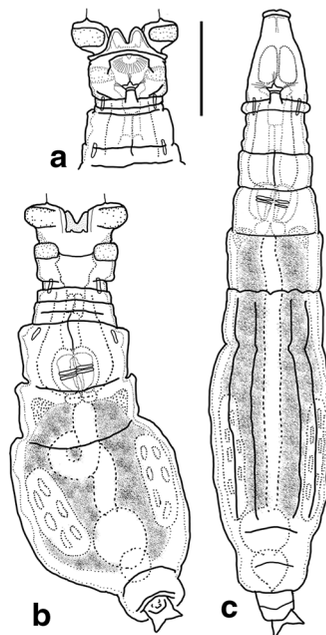
Etymology Named after Dr. Karel Janko, the leader of the project conducted on Vernadsky Base and collector of the material.

Diagnosis Resembles *Macr. insulana* Donner, 1962, by the shape of the corona and upper lip, the

characteristic sulcus with two denticles on the dorsal side, and the absence of a knob on the first foot pseudosegment. Differs by the flat and short triangular spurs, which are longer and peg-like in *Macr. insulana*. Dental formula of the new species is 2/2, while in *Macr. insulana* it is 1 + 2/2 + 1.

Description Body large, transparent, stomach of bright red or orange color. Integument smooth, thin, without knobs, spines or bolsters. Rostrum stout, of moderate length, lamella with two large rounded lobes. Corona wider than oval head base, CW is 100–133% of HW, HL is 35–54% of TL. Pedicels short, straight. Sulcus wide, not covered with membrane. On dorsal side of head two short denticles divided by interspace visible in sulcus. Trochi large, with papillae and sensory bristles. Upper lip with two triangular lobes reaching about a half of the sulcus and divided by interspace. Lower lip not protruding laterally. Neck stout, of moderate length, NL is 16–21% of TL. Antenna about 1/3 of the bearing pseudosegment. Trunk plump, BW is 15–21% of TL. Both pseudosegments of the rump swollen, RL is 12–21% of TL, RW is 73–93% of RL. Foot short, stout, 4 pseudosegments, without a dorsal knob, FL is 10–12% of TL, FW is 50–82% of FL. Spurs small,

Fig. 13 *Macrotrachela jankoi* sp. nov.: **a** holotype, head, feeding, dorsal view; **b** same, habitus, feeding, ventral view; **c** same, habitus, creeping, dorsal view; **d** paratype, trophi, cephalic view; **e** paratype, trophi, caudal view. Scale bar 50 μ m (a–c) or 5 μ m (d, e)



short, isosceles triangular, divided by an interspace almost equal to spur width, SL is 44–79% of SSW. Three short unsegmented toes. No eyespots. Throat voluminous, esophagus short, straight. Stomach glands round. Trophi ramate, 18–22 μm long and wide. Rami massive, with numerous short scleropili along the inner rim. Articulation somewhat concaved in the middle, but without incisure. Manubria thin, sickle-like. Dental formula 2/2, 39–41 minor teeth in each uncus. Oviparous. Eggs oval, shell smooth, without knobs or spines.

Measurements See Table 2.

Distribution Maritime Antarctica: AI*, AP*, KG*. Continental Antarctica: EB, VL (Cz*).

Habitat Soil, terrestrial moss.

Genus *Philodina* Ehrenberg, 1830

Philodina dartnallis sp. nov. (Figs. 3i, j, 14)

Priddle & Dartnall, 1978 (? *Philodina*), p. 475. Dartnall & Hollowday, 1985 (*Philodina* sp. 'A'), p. 24, Fig. 27a–e. Velasco-Castrillón et al., 2014a (*Ph.* sp. Bd46), p. 8.

Type locality Cape Bird, 77 m asl, S77°13.207', E166°26.568'.

Type habitat Soil.

Type material Holotype: SIZ 63.1 (CBM2PHD1a, 19.01.2010, Leg. J. Smykla), mounted in glycerin jelly, encircled with green ink. Paratypes: SIZ 63.2–4

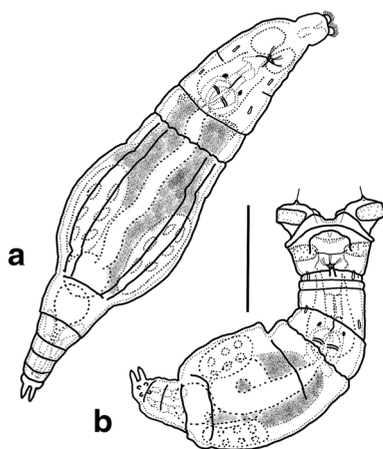


Fig. 14 *Philodina dartnallis* sp. nov.: **a** holotype, habitus, creeping, dorsal view; **b** same, feeding, dorsal view. Scale bar 50 μm

(CBM2PHD1b–d), on the same slide with holotype, encircled with black ink. **Additional material** SIZ 63.5–12 (CzM4PHD1–5, CzM4matPHD1, VDM2 PHD1–2), digital photos and videos.

Barcodes GenBank ID KJ543683–86, see also Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:6706ED3E-BBFF-4390-A602-4595C122986E.

Etymology Named after Antarctic researcher Dr. H. Dartnall who first depicted this species as *Ph.* sp. 'A'.

Diagnosis Similar to *Ph. flaviceps* Bryce, 1906, *Ph. australis* Murray, 1911, and some specimens of *Ph. brevipes* Murray, 1902. Resembles *Ph. flaviceps* by the shape of corona and spurs; however, the spurs of the new species are longer and with broader interspace. Differs from *Ph. australis* by the spur shape, which in the new species is peg-like with a broad interspace, but narrow triangular without an interspace in *Ph. australis*. Differs from *Ph. brevipes* by the shape of the upper lip (rounded lobes in the new species, pointed in *Ph. brevipes*) and by the absence of long sensory cilia in the rostrum. The foot seems to be shorter than in *Ph. brevipes*.

Description Body of moderate size, colorless, transparent. Integument smooth, thin, without knobs, spines or bolsters. Rostrum of moderate length, thick, lamella with two very small semicircular lobes. Corona wider than head base, CW is 102–119% of HW, HL is 27–38% of TL. Upper lip wide, arcuate, with two large low rounded lobes not reaching the plane of the trochal discs and divided by a broad interspace. Pedicels short, trochi large, with papillae and sensory bristles, retractors visible. Sulcus wide, not covered with membrane, without ligula. Lower lip not protruding laterally. Neck wide, of moderate length. Antenna 1/3 of bearing pseudosegment width. Trunk plump, BW is 14–25% of TL. Rump conical, first pseudosegment slightly swollen, without protrusions, RL/TL is 10–18%, RW/RL is 74–96%. Foot of moderate length, stout, 5 pseudosegments, without knobs or protrusions, FL is 8–15% of TL, FW is 13–22% of FL. Spurs peg-like, parallel to each other, divided by very narrow interspace, SL is 51–72% of SSW. Four thick unsegmented toes. Two cerebral orange or bright-red eyespots. Throat and esophagus of moderate length. Trophi ramate, round, 3/2. Stomach bright red, lumen wide. Egg oval, with

rounded knob on one pole, shell without spines or sculpturation, egg size 50–59 × 34–42 µm.

Distribution Maritime Antarctica: AI*, AP*. Continental Antarctica: EB, VL (CB*, Cz*), WK.

Habitat Soil, terrestrial moss, lakes.

Philodina shackletoni sp. nov. (Fig. 15)

Velasco-Castrillón et al., 2014a (*Ph. sp.* Bd45), p. 8 (main text); 5, Figs. 23–27 (Annex S1).

Type locality Cape Royds, 18 m asl, S77°32.532', E166°08.855'.

Type habitat Soil.

Type material Holotype: SIZ 62.1 (CRL25PHC1a, 14.01.2010, Leg. J. Smykla), mounted in glycerine jelly. Paratypes: SIZ 62.2-3 (CRL25PHC2-3), the same. **Additional material** SIZ 62.4 (CRL21PHC1), digital photos.

Barcodes GenBank ID KJ543677-86, see also Supplementary file III.

ZooBank LSID urn:lsid:zoobank.org:act:B2616FE6-B7B9-45D0-9F3A-DF065BE96A4F.

Etymology Named in honor of the leading Antarctic explorer, Sir Ernest Henry Shackleton, who in 1909 established his base on Cape Royds where the species was discovered.

Diagnosis Most closely resembles *Ph. flaviceps* Murray, 1906, by the shape of spurs and corona, however lacking eyespots. Spurs are longer than in *Ph. flaviceps* and divided by broader interspace.

Description Body large, spindle-shaped, colorless with yellow–brown stomach. Integument smooth, thin, without knobs, spines or bolsters. Rostrum of moderate size, with crescent-like lamella. Corona wider than trapezoid head base, CW is 109–117% of HW, HL is 13–17% of TL. Pedicels short, slightly bent inwards. Sulcus wider than diameter of a trochus, not covered with membrane. Trochi with papillae and sensory bristles. Upper lip very low, arcuate, with two small rounded lobes divided by interspace. Lower lip not protruding laterally. Neck of moderate length and width, NL is 18–23% of TL. Antenna long, almost equal to the bearing pseudosegment width. Trunk slim, BW is 15–20% of TL. Rump large, swollen, RL is 15–17% of TL, RW is 11–13 of RL. Foot long, slim, 5 pseudosegments, FL is 10–15% of TL, FW is 27–32% of FL. Spurs long, needle-like, SL is 115–164% of SSW. Four unsegmented toes. No eyespots. Throat small, esophagus short. Lumen wide. Stomach glands small, round. Trophi ramate, 24 µm long and wide, dental formula 2/2. Oviparous. Eggs oval, shell smooth, without knobs or spines.

Measurements See Table 2. TL 400 µm (Velasco-Castrillón et al., 2014a).

Distribution Maritime Antarctica: SI. Continental Antarctica: EB, VL (CR*), WK.

Habitat Soil, terrestrial moss, pools.

Morphometric analyses

Adineta editae sp. nov. differed from the similar European species *A. gracilis* by larger size of body and mastax, and longer spurs. LME demonstrates that the variation of body measurements between species represented over 60% of total variation for the parameters describing the width along the rotifer body (HW, MinNW, MxNW, FW, SSW) and for the spur length. The variation between localities and individuals was not significant for FW, SSW and SL

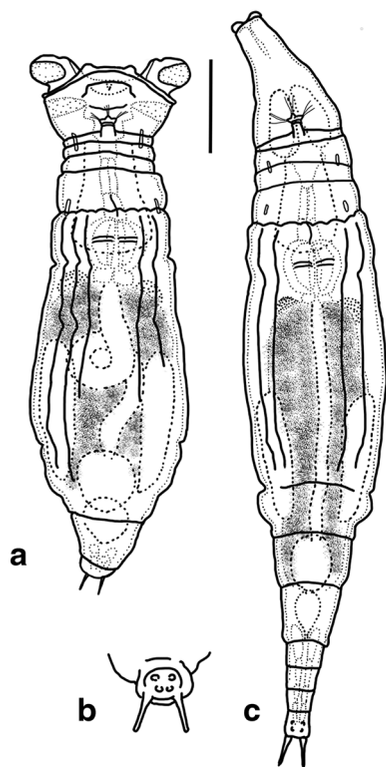


Fig. 15 *Philodina shackletoni* sp. nov.: **a** holotype, habitus, feeding, dorsal view; **b** foot, ventral view; **c** holotype, habitus, creeping, dorsal view. Scale bar 50 µm

(ANOVA on LME output: LR = 11.1 to 13.1, $P > 0.1$). This was in correspondence with our visual observation that *A. editae* sp. nov. had a distinctively stouter body than *A. gracilis* s. str. In the PCA plot (Fig. 16a) external measurements of the two species did not overlap along PC1 (correlating with all measurements) and PC2 (correlating with HW and SL).

The difference in trophi measurements between *A. editae* sp. nov. and *A. gracilis* represented over 90% of total variation in the number of minor teeth and trophi length, and over 80% in the case of trophi width. The variation between localities and individuals was insignificant for all measurements (LR = 0.2 to 2.1, $P > 0.5$). The trophi measurements of these species were completely separated on the PCA plot (Fig. 16b) and did not overlap along PC1 (correlates with all

measurements) and PC2 (correlates with the number of minor teeth and the unci width).

The body and trophi of the Antarctic species *A. grandis* and *A. fontanetoi* sp. nov. are indistinguishable by external morphology and did not differ significantly in any of the measured parameters. The Antarctic *A. emsliei* sp. nov. and the European *A. vaga* s. str. noticeably differed by at least one trophi measurement (the number of minor teeth in unci). The difference by this parameter consisted of over 80% total variation, with the variation between localities and individuals being insignificant (LR = 0.7 to 1.5, $P > 0.5$). The antarctic species *A. grandis* and *A. emsliei* sp. nov. were distinguished by all trophi measurements, the difference between species being over 90% of total variation. The variation between localities and individuals was not

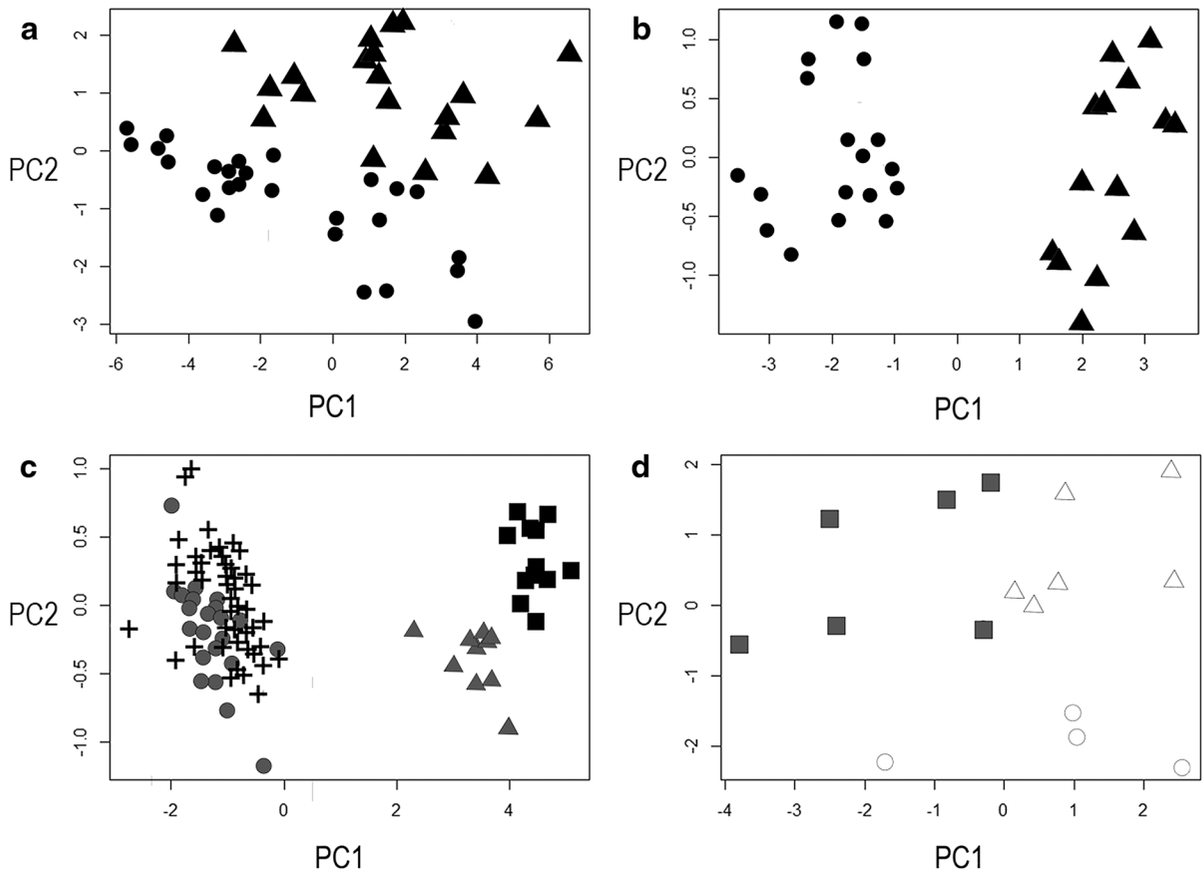


Fig. 16 Principal components analysis of rotifer body and trophi measurements: **a** *A. gracilis* (circles) and *A. editae* sp. nov. (triangles), body dimensions; **b** same, trophi dimensions; **c** *A. grandis* (crosses), *A. fontanetoi* sp. nov. (circles), *A. emsliei*

sp. nov. (squares) and *A. vaga* (triangles), trophi dimensions; **d** *H. antarctica* sp. nov. (squares), *H. vernadskii* sp. nov. (circles) and *H. sp. 4* (triangles), trophi dimensions

significant except for rami length (LR = 0.3 to 4.5, $P > 0.1$). In the PCA plot (Fig. 16c) the samples of trophi of *A. grandis* and *A. fontanetoi* sp. nov. overlapped completely on both PC1 (correlating with all trophi measurements) and PC2 (correlating with the number of minor teeth and trophi width), but the samples of *A. emsliei* sp. nov. did not overlap with any of the other species.

The Antarctic *H. antarctica* sp. nov. and *H. vernadskii* sp. nov. did not show any significant difference in trophi length and width, but could be distinguished by the number of minor teeth of the unci. Variation between the species on the latter measurement represented over 70%, the variation between localities and individuals being insignificant (LR = 3.6, $P > 0.1$). *Habrotrocha* sp. 4 is indistinguishable from *H. antarctica* sp. nov. by the external morphology, but has an intermediate trophi size between *H. antarctica* sp. nov. and *H. vernadskii* sp. nov., with the variation between species by all trophi parameters representing 50% or less of total variation. On the PCA plot of the samples of trophi measurements (Fig. 16d) *H. antarctica* sp. nov. and *H. vernadskii* sp. nov. did not overlap with each other on any axis. *Habrotrocha* sp. 4 did not overlap with either of the two other species.

DNA taxonomy

On the phylogenetic tree of *Adineta* (Fig. 17a) consisting of 46 IEEs and 28 singletons according to GMYC, the sequences of Antarctic rotifers grouped into 1 singleton and 8 independently evolving entities (IEE) identified by the GMYC and 4× rule approaches (32 singletons and 6 IEEs according to bPTP). Twenty-five IEEs of *Adineta* identified by GMYC contained sequences only from Europe, 6 IEEs—sequences from North America, 3 IEEs—sequences from each of Africa, Australia and New Zealand, and 2 IEEs—sequences from Asia. None of the Antarctic IEEs contained rotifers originating from any other continents, while three cosmopolitan IEEs (all attributed to the *A. vaga* species complex according to morphology) were identified from outside Antarctica. Two cosmopolitan IEEs had European-North American distributions and one occurred in Europe, Africa and New Zealand. The phylogeny shows that the Antarctic endemic *A. grandis* consists of at least two morphologically indistinguishable entities (one of

which was described above as *A. fontanetoi* sp. nov.). For three IEEs the morphology has not been described; therefore, these are listed as *A. sp.* 1–3. Finally, three IEEs that proved to be both genetically distinct and morphologically distinguishable are described above as *A. editae* sp. nov., *A. emsliei* sp. nov. and *A. coatsae* sp. nov. Molecular analysis confirmed that these species occur only in Antarctica, in spite of being previously confused with the cosmopolitan species *A. gracilis*, *A. vaga* s. str. and *A. barbata*. The integrity of *A. coatsae* sp. nov. as a single IEE was supported by the GMYC model, but not by the bPTP and 4× rule models. Both the GMYC and 4× rule, but not the bPTP model, confirmed the integrity of *A. fontanetoi* sp. nov. as a separate IEE.

The tree of *Philodina* consisted of 31 singletons and 42 IEEs identified by GMYC, mostly confirmed as IEEs by the 4× rule and bPTP (Fig. 17b). Nineteen singletons and ten IEEs contained sequences of Antarctic bdelloids. According to GMYC, 18 IEEs of *Philodina* had a European distribution, 17 were from North America and 4 from Asia, and 3 IEEs had cosmopolitan distributions. *Philodina* is the only one of the four investigated genera with cosmopolitan IEEs occurring in the Antarctic, those being *Ph. sp.* 4 (two sequences, from the Antarctic and the USA) and *Ph. sp.* 7 (25 sequences from the USA and one from Antarctica). Unfortunately, for both of these IEEs no data on morphology are available, and none of the sequences were obtained from vouchers of already known species. The remaining IEEs did not contain individuals from continents other than Antarctica. For six of them, listed here as *Ph. sp.* 1–6, no morphological data are available. *Philodina gregaria* appeared as one large pan-Antarctic IEE, three singletons and one IEE with atypical morphology, containing only two sequences. Two IEEs proved to be well distinguishable both morphologically and by the means of DNA taxonomy, both from *Ph. gregaria* and the morphologically similar non-Antarctic *Ph. acuticornis* Murray 1902, *Ph. flaviceps* Murray 1906, and *Ph. roseola* Ehrenberg, 1832. These two species are described above as the new Antarctic endemics: *Ph. shackletoni* sp. nov. and *Ph. dartnallis* sp. nov. The integrity of the *Ph. dartnallis* sp. nov. clade was supported by two of the three delimitation methods.

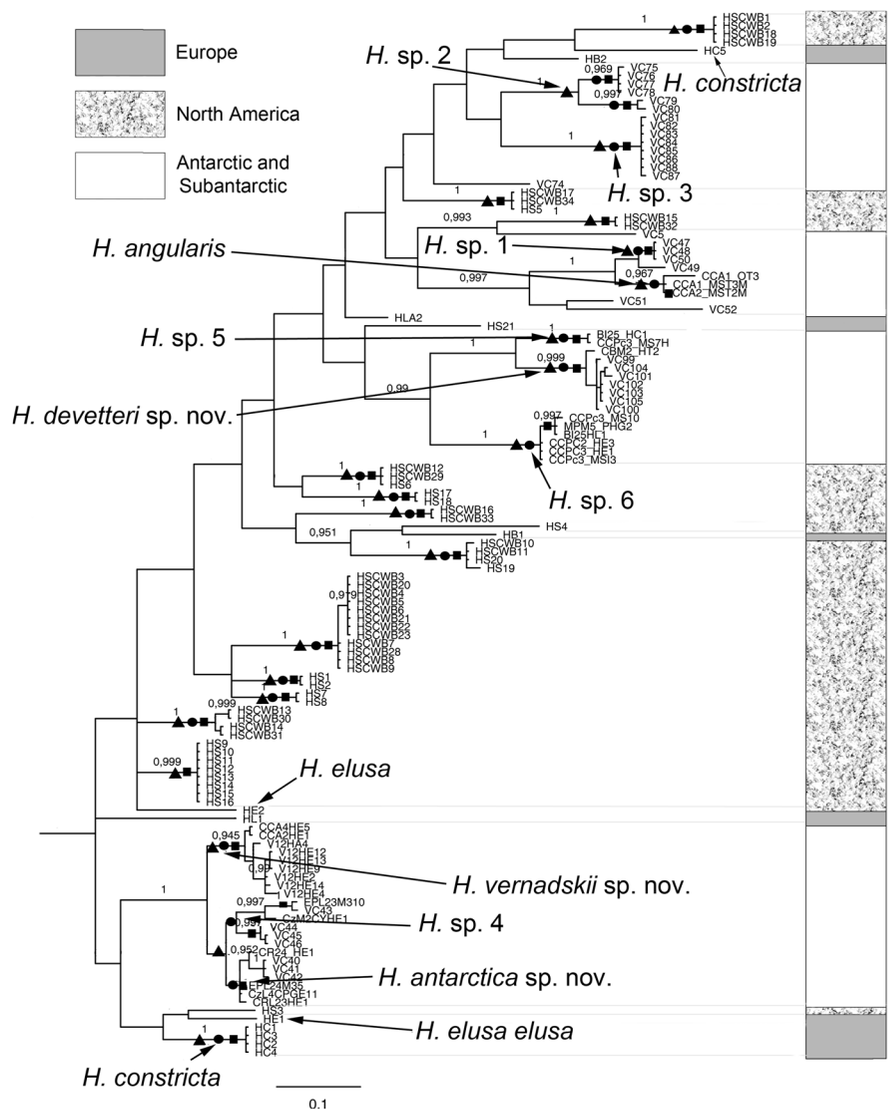
Putative species delimitation of *Habrotrocha* according to GMYC gave 22 IEEs and 1 singleton. Eleven IEEs and four singletons had strictly Antarctic

distribution. Ten IEEs contained sequences from North America and one was from Europe (Fig. 18). No cosmopolitan IEEs were detected. For three IEEs (*H. sp. 1*, 2 and 3) morphological data are not available. Three IEEs are described above as *H. antarctica* sp. nov., *H. devetteri* sp. nov. and *H. vernadskii* sp. nov. based on both morphological and molecular differences from the similar non-Antarctic *H. constricta*, *H. crenata* s. lat., *H. elusa* s. lat. and *H. thienemanni* s. lat. *Habrotracha* sp. 4 was identified as an IEE by the GMYC model, but not confirmed as a single entity by the 4× rule, being less than four times separated from both *H. antarctica* sp. nov. and *H. vernadskii* sp. nov. (which are genetically and

morphologically sufficiently separate to be good species). *H. sp. 4* and *H. sp. 5* are morphologically identical to *H. devetteri* sp. nov., but were delimited as separate IEEs by all three molecular delimitation methods. *H. angularis* has been confirmed as a separate species with a strictly Antarctic distribution.

Sixteen IEEs and 20 singletons were identified in *Macrotrachela* using GMYC, and this delimitation in most, but not all cases was confirmed by the two other approaches (Fig. 19). Four IEEs and three singletons were distributed in Antarctica, with eight IEEs identified from Europe, four from Asia, three from North America and one from Africa. No cosmopolitan IEEs were detected by any of the delimitation methods. For

Fig. 18 Phylogenetic relationships in the genus *Habrotracha* (consensus Bayesian tree, COX1 mt DNA data set). Putative species are delimited according to the 4× rule (triangles), GMYC (circles) and bPTP (squares). Wider distributions of rotifers are shown as in boxes on the right. Individual labels on branches are given in larger resolution in Supplementary file IV



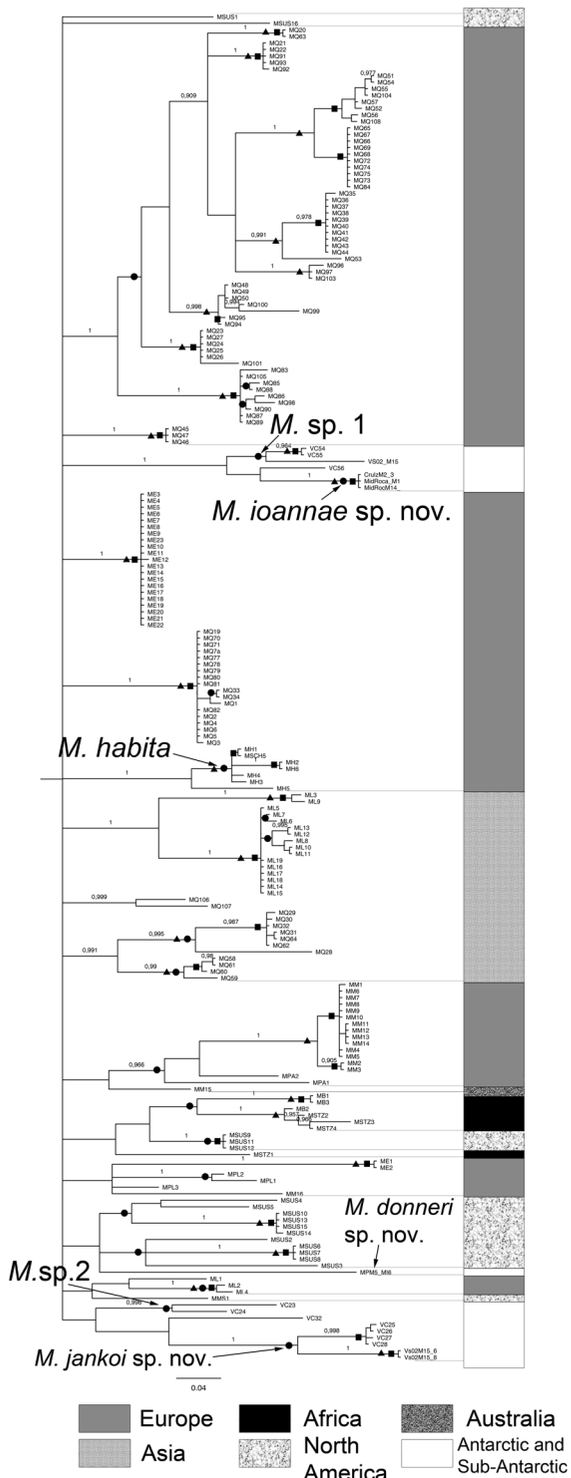


Fig. 19 Phylogenetic relationships in the genus *Macrotrachela* (consensus Bayesian tree, COX1 mt DNA data set). Putative species are delimited according to the 4× rule (triangles), GMYC (circles) and bPTP (squares). Wider distributions of rotifers are shown as in the boxes on the right. Individual labels on branches are given in larger resolution in Supplementary file IV

insolita or *Macr. cf. hewitti* (Donner, 1972) by Yakovenko & Tyshchenko (2006), was represented by a single sequence, which did not cluster with any non-Antarctic rotifers including the morphologically similar *Macr. habita*. For *Macr. sp. 1* more morphological data are required. *Macrotrachela ioannae* sp. nov. and *Macr. jankoi* sp. nov. were confirmed by GMYC as good species occurring in Antarctic, but the bPTP approach revealed *Macr. jankoi* sp. nov. as two IEEs. The integrity of *Macr. sp. 2*, identified using GMYC, was not confirmed by the other two approaches.

The identification of IEEs based on our combined set of sequences, including both new sequences and the previously published data, was the same as presented in Velasco-Castrillón et al. (2014a) with the exception of three IEEs. According to the results of our analysis, the putative species Bd15 and Bd16 were identified as the single entity *H. sp.1* by all three models. Bd31 and Bd32 were identified as *H. sp. 2* by the 4× rule, but as separate entities by the GMYC and bPTP models. Bd5 and Bd6 were identified as *Macr. sp. 2* according to GMYC, but this was not confirmed by the two other models. All the sequences published by Barraclough et al. (2007) and Fontaneto et al. (2008) were identified as members of *Adineta* (*A. editae* sp. nov., *A. fontaneto*i sp. nov., *A. sp. 2*, *A. sp. 3*). All 16 sequences published by Fontaneto et al. (2012) were confirmed as *Philodina* (*Ph. sp. 3–13*).

Biogeography

Of the 60 morphospecies found in the new material examined in this study, at least 17 can be considered true Antarctic and/or sub-Antarctic endemics, with 6 of these being already known and 11 newly recognized. A further ten morphospecies that closely resemble known and presumed cosmopolitan bdelloids, possessing only minor but consistent morphological differences from the original descriptions, require further detailed analysis. No molecular data are available for 13 morphospecies showing no discernible differences from known cosmopolitan

one Antarctic IEE, *Macr. sp. 2*, no information is yet available on morphology. *Macrotrachela donneri* sp. nov., previously reported in Antarctica as *Macr. cf*

species, as well as for 20 identified only to the generic level, and these cannot currently be attributed with confidence to any classification. That gives at least 28 and possibly up to 45% endemism (17 or up to 27 morphospecies out of 60) on the basis of classical taxonomy alone.

Based on molecular data obtained both from our new samples and GenBank, we identified 33 IEEs and 37 singletons from Antarctica and the sub-Antarctic using GMYC (largely confirmed by the two other approaches used), belonging to the genera *Adineta*, *Habrotrocha*, *Philodina* and *Macrotrachela*. Including a further five IEEs and eight singletons apparently representing other genera, a total of 38 IEEs and 45 singletons of Bdelloidea are now identified from this region. Only two of the IEEs occurring in Antarctica have been confirmed as having wider non-Antarctic distributions by molecular methods. At least 13 of the strictly Antarctic IEEs are clearly distinguishable by morphology, while at least four IEEs appear to represent cryptic species.

The molecular data obtained suggest there is a very high level of endemism among Antarctic bdelloids—36 out of 38 IEEs (95%). GMYC also identified considerable apparent endemism in the rotifer faunas (IEEs) of other major global regions: 55 of 59 found in Eurasia (93%), 65 of 70 (93%) in North America, and 3 of 4 (75%) in both Australia and New Zealand, and Africa. In the genus *Adineta*, 93.5% of the IEEs throughout the world were found to be endemic at the level of a continent (with no cosmopolitan IEEs found so far in Antarctica). In the genus *Philodina*, the percentage of endemic IEEs was lower (85%), with at least two cosmopolitan IEEs occurring in the Antarctic continent.

Sixteen IEEs were found only within continental Antarctic (5 representatives of *Adineta*, 4 of *Philodina* and 7 of *Habrotrocha*), three IEEs (2 of *Adineta* and 1 of *Macrotrachela*) were found only in the maritime Antarctic, and the distribution of 8 IEEs was either sub-Antarctic or unknown. Only six IEEs (1 of *Adineta*, 2 of *Habrotrocha* and 3 of *Philodina*), or 18% of their total number, occurred in both the maritime and continental regions of Antarctica.

Discussion

The hypothesis “Everything is everywhere, but the environment selects” (EiE hypothesis) was initially

proposed by Beijerinck and Baas Becking in the early twentieth century (Baas Becking, 1934 in Williams, 2011) and recently gained a renewed interest of biogeographers. It presumes that organisms less than 1–2 mm in length tend to be cosmopolitan, having no “true endemics” because of the high dispersal capacities and large population sizes, and that their diversity is driven by ecological factors rather than historical processes (Hillebrand & Azovsky, 2001; Williams, 2011 and references therein). While this seems to be true for some microscopic organisms (Fenchel & Finlay, 2004), including some but not all bdelloid rotifers (Fontaneto et al., 2008), increasing number of studies are providing evidence that microbes can have distinct distribution patterns and show endemism on a global scale (Hughes Martiny et al., 2006; Garey et al., 2008; Ganter, 2011; Lacap et al., 2011). Some bdelloid morphospecies appear to be restricted at least to a particular continent, and the largest number of such taxa is found in Antarctica (Segers, 2007).

After a 100-year period of research, the Antarctic Bdelloidea still remain obscure and knowledge sparse in terms of their diversity, distribution and origin. The current study is only the fourth to report previously undescribed Antarctic bdelloid rotifers, with all 12 new taxa being currently known only from the continent. The study is the first to provide a robust combination of detailed, morphological, morphometric and molecular approaches, which are being utilized in the description of these new Antarctic taxa. This study is also the second to apply molecular approaches in describing the diversity and biogeography of Antarctic Bdelloidea carried out, like in Velasco-Castrillón et al. (2014a), through COX1 sequencing. To date, 15 morphospecies of 60 (25%) found in this study have been barcoded successfully. Of these, we have (1) described 12 species new to science using both classical and DNA taxonomy approaches, (2) revealed the presence of a number of cryptic species that are apparently morphologically identical to *A. grandis*, *H. antarctica* sp. nov. and *H. devetteri* sp. nov., (3) linked the molecular data provided by Velasco-Castrillón et al. (2014a) with morphospecies, and (4) provided evidence of generally very high levels of endemism of bdelloid rotifers in Antarctica, with the exception of the finding of two cosmopolitan species of *Philodina*, not identified in previous studies.

The difficulty of distinguishing bdelloid species based on morphology alone, due to their generally

highly conserved body morphology and structure, and ambiguity in defining specific characters have led to a prevailing misconception that the contemporary Antarctic fauna includes a large proportion of cosmopolitan bdelloids. Thus, Donner (1965, 1972a) considered minor differences in the size and shape of the bdelloid body and its appendages as only representing intraspecific variability. Previous studies have often attributed Antarctic specimens to species already known from elsewhere (mostly Europe) if most of the external morphological characters matched the original descriptions or they were identified through keys based on the latter. Our morphometric and molecular data support the conclusions of Fontaneto et al. (2007), who proved that careful morphometric measurement of hard parts of the mastax (trophi) can differentiate several morphologically distinct entities within one “classical” species, corresponding with IEEs determined by molecular analysis.

All three models applied here for the delimitation of IEEs based on molecular sequence data showed good correspondence with rotifer morphology. GMYC gave the best correspondence with the species-specific morphology, and in most cases the results were supported by two other approaches. However, the bPTP model, used similarly by Velasco-Castrillón et al. (2014a), tended to give finer subdivision of IEEs, creating an excessive number of entities unidentifiable at the morphological level. Our data confirmed the integrity of all but three of the IEEs identified by Velasco-Castrillón et al. (2014a) and also confirmed the attribution of some sequences to *Adineta* and *Philodina* as proposed by Barraclough et al. (2007) and Fontaneto et al. (2008, 2012).

Our data demonstrate that, in spite of their acknowledged high dispersal capacities, Bdelloidea have distinctive patterns of distribution on the global scale. It has been considered that rotifer species, including bdelloids, are generally widespread with endemism occurring at the continental level (Dumont, 1983; Ricci, 1987; Segers, 2007; Fontaneto et al., 2008). Fontaneto et al. (2008) found that, within the bdelloid genera *Adineta* and *Rotaria*, IEEs were widely distributed over Europe, some even occurring in different continents (*A. vaga* found in UK, Tanzania and New Zealand, see Fig. 17). According to our data, not only is the distribution of most bdelloid IEEs limited by continents, with the highest levels of endemic IEEs in Antarctica, but even within the Antarctic region there

are substantial differences within local bdelloid faunas (maritime and continental Antarctica).

Conclusions

A striking feature of the data obtained in the current study is that of the extremely high levels of endemism to the Antarctic and sub-Antarctic region apparent in the bdelloid fauna. Clearly, consideration of the concept of endemism is itself limited by the quality and extent of the data available, from both the Antarctic and other regions. However, both the current study and that of Velasco-Castrillón et al. (2014a) are consistent in identifying (1) that considerably greater diversity in terms of divergence to the ‘species level’ is apparent in analyses of molecular (COX1) data than was the case in previous classical taxonomic studies of the group and (2) that Antarctic lineages are distinct from those of bdelloids from other continents available today in GenBank. In addition to the previous studies, we show that careful morphological analysis using morphometrics and SEM in many cases allows the detection of endemic Antarctic species of Bdelloidea even without molecular analysis.

Implicit in the assessment of considerable levels of endemism at the continental level is the conclusion that it is indicative of an extended history (long-term presence) allowing evolutionary divergence in situ in the Antarctic. This is consistent with a range of studies over the last 1 to 2 decades that have used both classical and molecular approaches to confirm high levels of endemism and long evolutionary histories in representatives of all the main terrestrial invertebrate groups occurring in Antarctica, including Tardigrada (Convey & McInnes, 2005), Nematoda (Andrássy, 1998; Maslen & Convey, 2006), Collembola (Green-slade, 1995; McGaughan et al., 2010; Torricelli et al., 2010), Acari (Pugh, 1993; Stevens & Hogg, 2006) and Diptera (Allegrucci et al., 2012). Wider reviews of this subject are provided by Convey et al. (2008) and Pugh & Convey (2008). Similar conclusions are increasingly being drawn from studies of some microbial groups (De Wever et al., 2009; Strunecký et al., 2012), most recently, mosses (Pisa et al., 2014).

The outcomes of the current study highlight the need for considerably greater survey efforts to be applied to groups of microscopic Antarctic fauna rich in cryptic species such as rotifers. Data obtained in the current

study suggest that at least some species of bdelloid rotifers are limited to particular parts of the Antarctic or sub-Antarctic. This again is consistent with recent findings in terrestrial biota (Convey et al. 2008; Pugh & Convey 2008; Velasco-Castrillón et al., 2014a) as well as the recent analysis of Terauds et al. (2012), which identified no less than 15 ‘Antarctic Conservation Biogeographic Regions’ across the Antarctic continent alone. Thus, further targeted research among the bdelloid rotifers of Antarctica, integrating classical, morphometric and molecular biological approaches, should identify considerably greater levels of diversity and both continental and intracontinental regional endemism than are currently appreciated.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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