# Redescription of Milnesium alpigenum Ehrenberg, 1853 (Tardigrada: Apochela) and a description of Milnesium inceptum sp. nov., a tardigrade laboratory model 

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#### Abstract

Intra- and interspecific variability, being at the very core of alpha taxonomy, has been a long-standing topic of debate among tardigrade taxonomists. Early studies tended to assume that tardigrades exhibit wide intraspecific variation. However, with more careful morphological studies, especially those incorporating molecular tools that allow for an independent verification of species identifications based on phenotypic traits, we now recognise that ranges of tardigrade intraspecific variability are narrower, and that differences between species may be more subtle than previously assumed. The taxonomic history of the genus Milnesium, and more specifically that of the nominal species, M. tardigradum described by Doyère in 1840, is a good illustration of the evolution of views on intraspecific variability in tardigrades. The assumption of wide intraspecific variability in claw morphology led Marcus (1928) to synonymise two species with different claw configurations, M. alpigenum and M. quadrifidum, with M. tardigradum. Currently claw configuration is recognised as one of the key diagnostic traits in the genus Milnesium, and the two species suppressed by Marcus have recently been suggested to be valid. In this study, we clarify the taxonomic status of M. alpigenum, a species that for nearly a century was considered invalid. We redescribe M. alpigenum, using a population collected from the locus typicus, by the means of integrative taxonomy, i.e. including light microscopy, scanning electron microscopy, ontogenetic observations, and genetic barcoding. Moreover, the redescription of M. alpigenum allowed us to verify the uncertain taxonomic status of two popular laboratory models that were originally considered to be M. tardigradum; though one was recently reidentified as $M$. cf. alpigenum. Our analysis showed that both laboratory strains, despite being morphologically and morphometrically nearly identical to M. alpigenum, in fact represent a new species, M. inceptum sp. nov. The two species, being disnguishable only by statistical morphometry and/or DNA sequences, are the first example of pseudocryptic species in tardigrades.


Key words: barcoding, cryptic species, integrative taxonomy, M. tardigradum s.s., phylogeny, pseudocryptic species

## Introduction

Tardigrades, also known as water bears, are a phylum of microscopic invertebrates that dwell in marine, freshwater and terrestrial ecosystems (Nelson et al. 2015). The first formal descriptions of tardigrade species were published in the first half of the XIX century. Among them was Milnesium tardigradum Doyère 1840, described by a French zoologist Louis Michel François Doyère. The species was established as the nominal taxon for the genus Milnesium Doyère, 1840, family Milnesiidae Ramazzotti, 1962, and the order Apochela Schuster et al., 1980. In the following decades, two further species of this genus were described: M. alpigenum Ehrenberg, 1853 from Monte Rosa (Italy/Switzerland) and M. quadrifidum Nederström, 1919 from Utsjoki, Finland. Both species were differentiated from M. tardigradum based on claw morphology (Morek et al. 2016a), a trait currently termed the claw configuration (CC). However, soon after M. quadrifidum was described, Marcus (1928), in his influential monograph on tardigrade biology, expressed an opinion that claw morphology was only a manifestation of
intraspecific variability, making it unsuitable for the differentiation of Milnesium species. Therefore, he synonymised both M. alpigenum and M. quadrifidum with M. tardigradum. This resulted in a widespread conviction that M. tardigradum exhibited considerable morphological variability and throughout the following decades researchers used to classify any Milnesium species as M. tardigradum, no matter what geographic origin of specimens or morphological variation (e.g. see Ramazzotti \& Maucci 1983; Dastych 1988).

Thus, with numerous records throughout the globe, M. tardigradum inevitably became recognised as cosmopolitan and for decades remained the only species in the genus Milnesium. This started to change only at the end of the XX century, when several new Milnesium species descriptions, although based on traits other than claw configuration, were published (Ramazzotti, 1962; Binda \& Pilato, 1990; Maucci, 1991; Pilato \& Binda, 1991). Later, Tumanov (2006) described additional five species and proposed to standardise some of the key morphometric measurements. However, the greatest increase in Milnesium species descriptions occurred after the redescription of M. tardigradum (Michalczyk et al. 2012a, b), when $46 \%$ of all known species of the genus were described within only six years (i.e. from 2012 to 2017). Based on their observations, Michalczyk et al. (2012a, b) assumed that CC is stable at the species level and they emphasised the value of this trait in Milnesium spp. differentiation. However, recently Morek et al. (2016a), using an experimental approach, discovered that some Milnesium species may undergo ontogenetic CC change. In fact, the latest integrative study on intraspecific variability of M. tardigradum by Morek et al. (2019) showed that the nominal species also exhibited developmental variability in CC, which had gone undetected in the redescription by Michalczyk et al. (2012a, b). Importantly, however, the discovery of developmental variability in CC does not undermine the taxonomic value of CC itself, because the developmental pattern seems to be species-specific Morek et al. (2019). Moreover, the pattern of CC ontogenetic variability may constitute an additional set of traits for species delimitation in the genus Milnesium.

In other words, modern advances in Milnesium taxonomy showed that the opinion of Marcus (1928) was incorrect, and M. alpigenum and M. quadrifidum are good species and are now pending integrative redescriptions (Morek et al. 2016a). These redescriptions are particularly important because with the simplistic original descriptions, it is impossible to differentiate M. alpigenum and M. quadrifidum from congeners that differ solely in quantitative (morphometric) and/or molecular traits. This, in turn, may prevent the identification of new species and lead to underestimation of species diversity and overestimation of species geographic ranges. Milnesium quadrifidum is the only known Milnesium species with the [4-4]-[4-4] CC. Therefore, describing new species that exhibit this claw configuration and morphology before M. quadrifidum is redescribed involves a risk of taxonomic inflation (Morek et al. 2016a). On the other hand, M. alpigenum, with the [3-3]-[3-3] CC, was the first described member of the largest group of Milnesium species defined by the CC. Thus, the redescription should verify whether any of the eighteen reported species with the [3-3]-[3-3] CC and smooth cuticle require synonymising with $M$. alpigenum (Morek et al. 2016a). Moreover, the two most studied Milnesium laboratory strains, one from Japan (Suzuki 2003) and the other from Germany (Schill et al. 2004), fit the description of M. alpigenum. Both laboratory strains were originally considered to be "M. tardigradum"; though the German strain was recently tentatively reidentified as "M. cf. alpigenum" (see Michalczyk et al. 2012a, Morek et al. 2016a, and Morek et al. 2019 for details). Thus, it is vital to verify whether the laboratory strains represent M. alpigenum, or a new species of the [3-3]-[3-3] CC group (Morek et al. 2016a). These laboratory strains have been used in studies on cryptobiosis (e.g. Hengherr et al., 2008a, b, Hengherr et al., 2009a), astrobiology (e.g. Jönsson et al. 2016), cell biology (e.g. Beisser, et al. 2012, Schokraie et al. 2012, Grohme et al. 2013), physiology (e.g. Reuner et al. 2010a, Förster et al. 2012), developmental biology (e.g. Suzuki 2003, Suzuki 2006), experimental taxonomy (e.g. Kosztyła et al. 2016, Morek et al. 2016b, Stec et al. 2016), and ethology (Shcherbakov et al. 2010), thus pinpointing their identity is of great importance.

In this paper, we aim to clarify the taxonomy within the genus Milnesium and to verify the taxonomic status of the popular laboratory models. To achieve this, we integratively analysed Milnesium individuals, with the [3-3]-[33] CC and unsculptured cuticle, collected from the M. alpigenum type locality in northern Italy, and compared them with the two M. cf. alpigenum laboratory cultures (from Japan and Germany). Additionally, we analysed two similar populations, from Switzerland and Bulgaria, to test whether these represented either M. alpigenum or the species used to establish the laboratory strains. In either case, these additional populations could extend the genetic and phenotypic variability as well as the geographic range of the species in question.

## Materials and methods

Nomenclature. Claw configuration (abbreviated throughout the text as "CC") is denoted according to Michalczyk et al. (2012a, b), i.e. as a string of bracketed numbers that represent the number of points on the secondary branches on external and internal claws I-III, and on anterior and posterior claws IV: formula [e-i]-[a-p]. The developmental
terminology follows Morek et al. (2019), i.e. immature individuals: the first instar $=$ hatchling, and the second instar $=$ juvenile; mature individuals from the third instar onwards = adults.

Sampling and specimen isolation. Detailed collection data are provided in Table 1. All moss samples were collected and processed according to standard methods (e.g. Stec et al. 2015). Tardigrades were cultured following the protocol in Kosztyła et al. (2016), i.e. they were fed rotifers, Lecane inermis (Bryce, 1892), and kept on plastic Petri dishes with scratched bottoms, at a stable temperature ( 8 or $16^{\circ} \mathrm{C}$ ), and in complete darkness. Individuals isolated from samples and/or cultures were split into 3-4 analysis groups: (i) development tracking, (ii) imaging and morphometry in phase contrast light microscope (PCM), (iii) imaging in scanning electron microscopy (SEM), and (iv) DNA sequencing; see Table 1 for details.

Milnesium alpigenum was originally described from the Monte Rosa massif, thus we sampled this locality in order to find a population that could be designated as the neotype series. Since the original description is very limited and the type material no longer exists, any Milnesium with unsculptured cuticle and the [3-3]-[3-3] CC found in the vicinity of the type locality could be designated as neotype M. alpigenum. Our colleagues from the Adam Mickiewicz University (Poland) kindly provided us with a moss sample with a candidate population collected in the Monte Rosa massif, from which we isolated five live females (see Table 1 for details). The small number of individuals was insufficient for all the required analyses, so we cultured the five females separately (to control for potential multiple species exhibiting similar morphology and dwelling in a single moss cushion). After the isolines have perpetuated for several generations, producing sufficient numbers of individuals, we sequenced animals from all isolines (see below for details). DNA sequencing confirmed that all five isolines represent a single species, thus all individuals were pooled and used for the planned analyses (Table 1).

Alongside the Italian M. alpigenum neotype material, we also analysed a further four Milnesium populations, which conformed to the original description of M. alpigenum, to test whether they represent M. alpigenum or new species (see Table 1 for details). These were:

- a German laboratory strain established by R.O. Schill at the University of Tübingen in 2003 with specimens collected in the Bebenhausen forest and subsequently maintained at the University of Stuttgart (sample code DE.001; "Tübingen strain");
- a Japanese laboratory strain established by A.C. Suzuki at the University of Keio in 2002 with specimens collected in Hiyoshi (sample code JP.010; "Hiyoshi H-1 strain");
- a wild population from Switzerland (sample CH.002);
- a wild population from Bulgaria (sample BG.058).

Microscopy and imaging. A total of 47 individuals of the neotype M. alpigenum population (IT.057), 98 from the Tübingen laboratory strain (DE.001), 15 from the Japanese laboratory strain (JP. 010), 9 from the Swiss population (CH.002), and 69 from the Bulgarian population (BG.058) were used for the PCM analysis. All were mounted in Hoyer's medium on microscope slides according to the recipe and protocol described by Morek et al. (2016b). Photographs and measurements were taken using Nikon Eclipse 50i PCM associated with a Nikon Digital Sight DS-L2 digital camera (termed here as "a standard class/quality PCM") and with Olympus BX53 PCM associated with Olympus DP74 digital camera ("high class/quality PCM"). For structures that could not be focused in a single photograph, a series of up to eight pictures were taken and merged into one deep-focus image using Corel Photo-Paint X8.

Additionally, 20 individuals of the neotype population of M. alpigenum (IT.057), 15 from the German laboratory strain (DE.001), and 18 from the Bulgarian population (BG.058) were processed for SEM imaging, following the protocol described in Stec et al. (2015), and then examined under high vacuum in Versa 3D DualBeam SEM at the ATOMIN facility of the Jagiellonian University.

Ontogenetic variability detection. In order to test for ontogenetic variability, developmental tracking according to Morek et al. (2016a) was employed. In brief, exuviae with eggs were incubated individually and emerging hatchlings were split into three subsets: (i) mounted on permanent microscope slides in Hoyer's medium

| Sample code | Locality | Coordinates, altitude | Sample type | Collection date | Collector | Species |  | Analyses erformed |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| IT. 057 | Italy, Monte Rosa, lower chair-lift station of Macugnaga | $\begin{aligned} & 45^{\circ} 58^{\prime} 13^{\prime \prime N} \\ & 07^{\circ} 57^{\prime} 07^{\prime \prime} \mathrm{E} \\ & 1370 \mathrm{~m} \text { asl } \end{aligned}$ | moss on roof | 27.06.2015 | Łukasz Kaczmarek, Milena Roszkowska, Weronika Erdmann, Krzysztof Zawierucha | M. alpigenum Ehrenberg, 1853 | 1 | ii iii iv |
| DE. 001 | Germany, <br> Tübingen, Bebenhausen, Schönbuch Nature Reserve, mixed forest | $\begin{aligned} & 48^{\circ} 33^{\prime} 42^{\prime \prime N} \\ & 09^{\circ} 03^{\prime} 48^{\prime \prime} \mathrm{E} \\ & 377 \mathrm{~m} \text { asl } \end{aligned}$ | moss on soil | 10.2002 | Ralph O. Schill | M. inceptum sp. nov. | i | ii iii iv |
| JP. 010 | Japan, Yokohama, Hiyoshi | $\begin{aligned} & 35^{\circ} 33^{\prime} 10.6^{\prime \prime} \mathrm{N} \\ & 139^{\circ} 39^{\prime} 01^{\prime \prime} \mathrm{E} \\ & 35 \mathrm{~m} \text { asl } \end{aligned}$ | $\begin{aligned} & \text { moss } \begin{array}{l} \text { (Bryum } \\ \text { argenteum }) \\ \text { concrete wall } \end{array} \end{aligned}$ | 03.04.2000 | Atsushi C. Suzuki | M. inceptum sp. nov. | i | ii iv |
| CH. 002 | Switzerland, <br> Zürich, corner of Rämistrasse and Schmelzbergstrasse | $\begin{aligned} & 47^{\circ} 22^{\prime} 38^{\prime \prime} \mathrm{N} \\ & 08^{\circ} 32^{\prime} 56^{\prime \prime} \mathrm{E} \\ & 470 \mathrm{~m} \text { asl } \end{aligned}$ | moss on concrete wall | 10.12.2015 | Łukasz Michalczyk, Grzegorz Kwiatkowski | M. inceptum sp. nov. | i | ii iv |
| BG. 058 | Bulgaria, <br> Shanovo, <br> Kazanlak Valley, near the north slope of Sarnena Mt. | $\begin{aligned} & 42^{\circ} 333^{\prime 2} 7^{\prime N} \mathrm{~N} \\ & 25^{\circ} 37^{\prime} 51^{\prime \prime} \mathrm{E} \\ & 300 \mathrm{~m} \text { asl } \end{aligned}$ | moss (Grimmia sp.) on brick wall | $25.08 .2015$ | Dilian Georgiev, Maria Yankova | M. inceptum sp. nov. | i | ii iii iv |

one day after hatching, (ii) reared to the juvenile stage and mounted one day after moulting, and (iii) cultured to the first adult instar and mounted one day after moulting. A comparison of the morphology between the first three instars permitted observation of any ontogenetic variability in taxonomically important traits such as the CC, cuticle morphology and the presence/appearance of cuticular bars under claws I-III.

Morphometrics. The number of measured specimens follow recommendations by Stec et al. (2016). Specimens were measured according to Tumanov (2006) and Michalczyk et al. (2012a). The pt ratio is the ratio of the length of a given structure to the length of the buccal tube, expressed as a percentage (Pilato 1981), in the text the $p t$ values are given in italics.

Morphometric species delineation. Despite molecular analyses indicating clear genetic differences between the two species addressed in this study, the morphometric ranges (both absolute and relative values) overlapped (see below for details). Therefore, in order to test for statistical differences between the species, we used Principle Component Analysis (PCA) followed by a series of Student $t$-tests with $\alpha$-level adjusted with the BenjaminiHochberg correction. The PCA allows a reduction of the original dataset components while retaining the maximum possible variation of data. The analysis was performed in R 3.4.2. (R Core Team 2015) by prcomp function, using only the $p t$ values as ratios to reduce allometric effects. The dataset had $c a .18 \%$ of missing measurements, which were replaced by median values of each variable in a given population to avoid losing statistical power. Afterwards, the variables were scaled to unit variance and zero mean to minimise the effect of different scales of measured traits. The results were visualised using package ggfortify (version 0.4.1, Tang et al. 2016).

Genotyping. First, we tested whether the five isolines of the M. alpigenum type locality (IT.057) and populations from Germany (DE.001), Japan (JP.010), Switzerland (CH.002), and Bulgaria (BG.058) represent a single or multiple species. This we achieved by sequencing two variable barcodes, the nuclear Internal Transcribed Spacer 2 (ITS-2), and the mitochondrial Cytochrome Oxidase C subunit I (COI), for eight individuals per population. We established that all Italian isolines represented a single species, and that all the remaining populations represented another species. With this knowledge, we further sequenced two conservative nuclear markers, small ribosomal subunit (18S rRNA), large ribosomal subunit ( 28 rRNA ), for four individuals from each species (specifically, from populations IT. 057 and DE. 001 ).

DNA was extracted from individual tardigrades following the protocol of Chelex ${ }^{(8)} 100$ resin (Bio-Rad), extraction method by Casquet et al. (2012) with modifications by Stec et al. (2015). Primer sequences and sources as well as references for PCR programs are listed in Table 2. All sequences were handled in BioEdit ver. 7.2.5 (Hall 1999). COI sequences were translated into amino acids to test for potential pseudogenes. Additionally the uncorrected p-distances were calculated utilising MEGA 7 (Kumar et al. 2016) for comparisons between species and populations. All sequences were aligned using the default settings of MAFFT version 7 (Katoh et al. 2002; Katoh \& Toh 2008). The obtained alignments were edited and checked manually in BioEdit and then trimmed to 585 bp (ITS-2) and 509 bp (COI).

TABLE 2. Primers and references for specific protocols for amplification of the four DNA fragments sequenced in the study.

| DNA <br> fragment | Primer name | Primer direction | Primer sequence ( $5^{\prime}-3$ ') | Primer source | PCR programme |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 18S rRNA | $\begin{aligned} & \text { SSU01_F } \\ & \text { SSU82_R } \end{aligned}$ | forward reverse | AACCTGGTTGATCCTGCCAGT TGATCCTTCTGCAGGTTCACCTAC | $\begin{aligned} & \text { Sands et al. } \\ & \text { (2008) } \end{aligned}$ | Zeller (2010) |
| 28 S rRNA | $\begin{aligned} & 28 \mathrm{SF} 0001 \\ & 28 \mathrm{SR} 0990 \end{aligned}$ | forward reverse | ACCCVCYNAATTTAAGCATAT CCTTGGTCCGTGTTTCAAGAC | Mironov et al. (2012) | Mironov et al. (2012) |
| ITS-2 | $\begin{aligned} & \text { ITS2_Eutar_Ff } \\ & \text { ITS2_Eutar_Rr } \end{aligned}$ | forward reverse | GCATCGATGAAGAACGCAGC TCCTCCGCTTATTGATATGC | Stec et al. (2018) | Stec et al. (2018) |
| COI | $\begin{aligned} & \text { COI_Mil.tar_Ff } \\ & \text { COI_Mil.tar_Rr } \end{aligned}$ | forward reverse | TATTTTATTTTTGGTATTTGATGTGC ССТСССССTGCAGGATC | Morek et al. (2019) | Morek et al. (2019) |

Phylogenetic analysis. In order to visualise evolutionary relationships between the four Milnesium populations analysed in this study, phylogenetic trees using all available Milnesium ITS-2 and COI sequences were constructed. Thus, in addition to new sequences, the data set comprised the ITS-2 sequences for the following
species: M. berladnicorum Ciobanu, Zawierucha, Moglan \& Kaczmarek, 2014; (KT951662 from Morek et al. 2016a); M. dornensis Ciobanu, Roszkowska \& Kaczmarek, 2015; (MG923557 from Morek et al. 2019); M. variefidum Morek, Gąsiorek, Stec, Blagden \& Michalczyk, 2016a; (KT951666-7 from Morek et al. 2016a); "M. tardigradum"; (GQ403681-2 from Schill, unpublished); M. tardigradum sensu stricto Doyère, 1840; (MG9235515 from Morek et al. 2019). Additional COI sequences for the following species were also included: $M$. berladnicorum (KT951659 from Morek et al. 2016a); M. dornensis (MG923566 from Morek et al. 2019); M. variefidum (KT951663 from Morek et al. 2016a); M. tardigradum s.s. (MG923558-65 from Morek et al. 2019); M. cf. alpigenum (KU51342 from Kosztyła et al. 2016); "M. tardigradum" (EU244603-4 from Schill, unpublished); "M. tardigradum" (FJ435810 from Guil \& Giribet 2012); "M. tardigradum" (JX683822-5 from Vicente et al. 2013); M. sp. (EF632553 from Sands et al., unpublished); M. sp. (KX306950 from Fox et al., unpublished); M. spp. (KJ857001-2, KP013598, KP013601 and KP013613 from Velasco-Castrillón et al. 2015). For the outgroup, sequences of Diploechiniscus oihonnae Richters, 1903 (MG063724 from Gąsiorek et al. 2017a and MG923556 from Morek et al. 2019) were used. Concatenation was run in SequenceMatrix (Vaidya et al. 2010) and the final ITS-2+COI alignment was 1094 bp long.

Using PartitionFinder version 2.1.1 (Lanfear et al. 2016), under the Akaike Information Criterion (AIC), the best substitution model was chosen for posterior phylogenetic analysis. As COI is a protein-coding gene, the alignment was divided into three data blocks representing three separated codon positions. As best-fit partitioning scheme, PartitionFinder suggested to retain three predefined partitions for the COI data set and four predefined partitions for the concatenated data set separately. As RAxML (Stamatakis 2014) allows for only a single model of rate heterogeneity (from the GTR family) in partitioned analyses, each data set was analysed twice: first to test all possible models implemented in the program (for Bayesian Inference, BI), and then for models from the GTR family (for Maximum Likelihood analysis, ML). The best fit-models for four partitions for BI were: TRN+I+G for the first codon position, K81UF+G for the second and the third codon position, and GTR+G for the ITS-2 partition. For ML, the best model was GTR+G.

BI marginal posterior probabilities were calculated using MrBayes v3.2 (Ronquist \& Huelsenbeck 2003). Random starting trees were used and the analysis was run for ten million generations, sampling the Markov chain every 1000 generations. An average standard deviation of split frequencies of $<0.01$ was used as a guide to confirm that the two independent analyses had converged. The program Tracer v1.3 (Rambaut et al. 2014) was then used to ensure Markov chains had reached stationarity and to determine the correct "burn-in" for the analysis, which was the first $10 \%$ of generations. The consensus tree was obtained after summarising the resulting topologies and discarding the "burn-in". In the BI consensus tree, clades recovered with posterior probability (PP) between 0.95 and 1.00 were considered well supported, those with PP between 0.90 and 0.94 were considered moderately supported and those with lower PP were considered unsupported. The consensus tree was viewed and visualised by FigTree v.1.4.3, available from http://tree.bio.ed.ac.uk/software/figtree. ML topologies were constructed using RAxML v8.0.19 (Stamatakis 2014). The strength of support for internal nodes of ML construction was measured using 1000 rapid bootstrap replicates. Bootstrap (BS) support values $\geq 70 \%$ on the final tree were regarded as significant statistical support.

Data deposition. Raw morphometric data are deposited in the Tardigrada Register (Michalczyk \& Kaczmarek 2013) under http://tardigrada.net/register/0056.htm (M. alpigenum), http://tardigrada.net/register/0057.htm (M. inceptum sp. nov.) as well as in Supplementary Materials SM.1-5. The sequences of all haplotypes were uploaded to GenBank (accession numbers are listed under species redescription/description and in the Appendix 1).

## Results

## Taxonomic accounts

## Phylum Tardigrada Doyère, 1840

Class Eutardigrada Richters, 1926
Order Apochela Schuster et al., 1980

## Family Milnesiidae Ramazzotti, 1962

## Genus Milnesium Doyère, 1840

## Milnesium alpigenum Ehrenberg, 1853

Fig. 1, Table 3
M. tardigradum Marcus (1928)

Material examined: The neotype series consisting of the neotype and 37 neoparatypes (see Table 1 and "Type repositories" below for details).

Integrative redescription. Females (morphometrics in Table 3): Body slightly yellowish, rather slender as for a Milnesium (Fig. 1A). Eyes present in live specimens, quickly dissolving after fixation in Hoyer's medium. Cuticle smooth in SEM and with minute pseudopores visible on the caudo-dorsal part only under high quality PCM (Fig. 1C). Weakly outlined pseudoplates on the caudo-dorsal cuticle visible in some specimens only under SEM (Fig. 1B and D). Six peribuccal papillae present, with the ventral being the smallest. Six triangular peribuccal lamellae of unequal size; the two lateral being slightly smaller than the dorsal and ventral lamellae, i.e. with the $4+2$ configuration (identifiable only in SEM; Fig. 1F). Two lateral papillae present. Buccal tube funnel-shaped (Fig. 1E). Claws slender, primary branches with tiny accessory points, more visible on claws IV. All secondary branches with three points, i.e. with the [3-3]-[3-3] CC (Fig. 1G-H). Spurs on secondary branches long and slender, especially on internal and anterior claws. Cuticular bars under claws I-III present.

Males: No males were found in the sample or culture, confirming that the neotype population is parthenogenetic (at least facultatively).

Juveniles: Morphologically identical to adults, except for the lack of pseudopores.
Hatchlings: Morphologically identical to adults, except for the lack of pseudopores and the absence of cuticular bars under claws I-III in the majority of examined specimens ( $7 / 8$ specimens $=88 \%$ ).

Ontogenetic variability: No developmental variability in the CC. Pseudopores visible only in adults. Cuticular bars under claws I-III mostly absent in hatchlings but always present in juveniles and adults.

Eggs: Oval, yellow, smooth and laid in exuviae. In the culture, up to 12 eggs were recorded in a single clutch.
DNA markers: All sequences were of a very good quality and every marker was represented by a single haplotype: 18S rRNA (1054 bp, MG996146); 28S rRNA (809 bp, MH000384); ITS-2 (530 bp, MH000382); and COI (560 bp, MH000380). Sequences are provided in Appendix 1.

Neotype locality: $45^{\circ} 58^{\prime} 13^{\prime \prime} \mathrm{N}, 07^{\circ} 57^{\prime} 07^{\prime \prime} \mathrm{E} ; 1370 \mathrm{~m}$ asl: Italy, Monte Rosa massif, lower chair-lift station of Macugnaga; moss on roof.

Etymology: Ehrenberg (1853) did not explain the choice of the species name; however, it seems reasonable to assume that Christian Ehrenberg named the species after the Alps, the mountain chain in which the type locality, the Monte Rosa massif, is located.

Type repositories: The neotype series consist of the neotype (slide IT.057.17) and 37 "neoparatypes" (IT.057.01-16; 18-38). The neotype and 15 neoparatypes (IT.057.01-12; 45-47) are preserved at the Institute of Zoology and Biomedical Research, Jagiellonian University, Gronostajowa 9, 30-387 Kraków, Poland), further 14 neoparatypes (IT.057.13-16; 18-26) are deposited in Department of Animal Taxonomy and Ecology, Adam Mickiewicz University in Poznań, Umultowska 89, 61-614 Poznań, Poland, 10 neoparatypes (IT.057.27-38) are stored in the Marine Biology \& Ecology Research Centre, University of Plymouth, Drake Circus, Plymouth, PL4 8AA, United Kingdom, one paratype (IT.057.47) is deposited in Natural History Museum, Cromwell Road, London SW7 5BD, United Kingdom, and the remaining 6 neoparatypes (IT.057.39-44) are deposited in the collection of Binda \& Pilato, Museum of the Department of Biological, Geological and Environmental Sciences, Section of Animal Biology "Marcello La Greca", University of Catania, Italy.

Phenotypic differential diagnosis: Milnesium alpigenum has the [3-3]-[3-3] CC and "smooth" cuticle (i.e. cuticle smooth in SEM and with minute pseudopores visible only under high quality PCM, but with no sculpturing, such as reticulation, on cuticle surface). This places it in the largest group of Milnesium species that share these characteristics (20 species; Morek et al. 2016a; Pilato \& Lisi 2016; Young et al. 2016; Pilato et al. 2016; Pilato et al. 2017; Schlabach et al. 2018). Nevertheless, M. alpigenum differs specifically from:
TABLE 3. Measurements (in $\mu \mathrm{m}$ ) and the pt values of selected morphological structures of 30 specimens (neotype and neoparatypes) of Milnesium alpigenum Ehrenberg, 1853 available life stages as possible.

| CHARACTER | N | RANGE |  |  |  |  | MEAN |  | SD |  | Neotype |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ |  |  | $p t$ |  | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | $p t$ |
| Body length | 30 | 383 | - | 983 | 1164 | - 1509 | 614 | 1292 | 166 | 84 | 751 | 1322 |
| Peribuccal papillae length | 23 | 5.8 | - | 11.7 | 15.1 | - 22.0 | 8.7 | 17.8 | 1.7 | 1.5 | 10.2 | 18.0 |
| Lateral papillae length | 28 | 5.6 | - | 11.7 | 14.1 | - 20.4 | 8.6 | 18.0 | 2.1 | 1.5 | 10.4 | 18.3 |
| Buccal tube |  |  |  |  |  |  |  |  |  |  |  |  |
| Length | 30 | 32.3 | - | 67.5 |  | - | 47.1 | - | 10.5 | - | 56.8 | - |
| Stylet support insertion point | 30 | 21.8 | - | 43.3 | 61.1 | - 70.3 | 30.4 | 64.9 | 6.2 | 2.1 | 34.7 | 61.1 |
| Anterior width | 29 | 11.5 | - | 28.6 | 29.6 | - 44.8 | 17.9 | 38.1 | 4.6 | 3.5 | 22.1 | 38.9 |
| Standard width | 30 | 8.6 | - | 23.6 | 21.0 | - 38.4 | 14.8 | 31.4 | 4.0 | 3.7 | 19.7 | 34.7 |
| Posterior width | 30 | 9.7 | - | 22.3 | 23.7 | - 39.4 | 15.6 | 33.0 | 4.0 | 3.8 | 20.9 | 36.8 |
| Standard width/length ratio | 30 | 21\% | - | 38\% |  | - | 31\% | - | 4\% | - | 35\% | - |
| Posterior/anterior width ratio | 29 | 75\% | - | 97\% |  | - | 86\% | - | 6\% | - | 95\% | - |
| Claw 1 lengths |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 30 | 13.7 | - | 27.0 | 36.7 | - 47.9 | 19.1 | 40.8 | 3.9 | 2.9 | 21.6 | 38.0 |
| External base + secondary branch | 23 | 10.1 | - | 19.7 | 26.8 | - 34.4 | 13.9 | 29.6 | 2.9 | 2.1 | ? | ? |
| External spur | 17 | 4.0 | - | 9.1 | 11.3 | - 17.8 | 6.4 | 13.3 | 1.4 | 1.5 | ? | ? |
| External branches length ratio | 23 | 66\% | - | 86\% |  | - | 73\% | - | 5\% | - | ? | - |
| Internal primary branch | 29 | 13.2 | - | 26.6 | 36.0 | - 44.8 | 18.5 | 39.2 | 3.6 | 2.4 | 21.0 | 37.0 |
| Internal base + secondary branch | 18 | 9.9 | - | 20.8 | 25.1 | - 32.7 | 13.4 | 29.2 | 3.0 | 2.1 | 15.8 | 27.8 |
| Internal spur | 23 | 3.7 | - | 10.1 | 11.2 | - 16.9 | 6.8 | 14.2 | 1.7 | 1.2 | 8.4 | 14.8 |
| Internal branches length ratio | 17 | 66\% | - | 84\% |  | - | 74\% | - | 5\% | - | 75\% | - |
| Claw 2 lengths |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 29 | 15.3 | - | 29.9 | 40.0 | - 50.2 | 21.2 | 44.8 | 4.3 | 2.9 | 24.4 | 43.0 |
| External base + secondary branch | 25 | 10.4 | - | 23.0 | 27.8 | - 34.2 | 15.0 | 31.5 | 3.6 | 1.9 | 18.6 | 32.7 |
| External spur | 20 | 4.5 | - | 10.7 | 12.7 | - 17.5 | 7.6 | 15.3 | 1.8 | 1.4 | 8.5 | 15.0 |

TABLE 3. (Continued)

| CHARACTER | N | RANGE |  |  |  |  | MEAN |  | SD |  | Neotype |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ |  |  | $p t$ |  | $\mu \mathrm{m}$ | $p t$ | $\mu \mathrm{m}$ | $p t$ | $\mu \mathrm{m}$ | $p t$ |
| External branches length ratio | 24 | 62\% | - | 80\% |  | - | 71\% | - | 4\% | - | 76\% | - |
| Internal primary branch | 27 | 15.2 | - | 29.0 | 39.2 | - 49.8 | 21.0 | 44.2 | 4.1 | 3.0 | 25.0 | 44.0 |
| Internal base + secondary branch | 14 | 10.4 | - | 21.1 | 28.1 | - 34.4 | 13.9 | 31.8 | 3.5 | 1.5 | 18.0 | 31.7 |
| Internal spur | 24 | 4.4 | - | 10.5 | 12.9 | - 18.4 | 8.1 | 16.3 | 1.9 | 1.4 | 9.7 | 17.1 |
| Internal branches length ratio | 12 | 65\% | - | 88\% |  | - | 71\% | - | 6\% | - | 72\% | - |
| Claw 3 lengths |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 28 | 15.9 | - | 32.4 | 41.1 | - 51.4 | 22.0 | 45.8 | 4.5 | 2.7 | 24.4 | 43.0 |
| External base + secondary branch | 21 | 11.1 | - | 23.0 | 29.9 | - 35.7 | 16.1 | 32.7 | 3.5 | 1.8 | 18.4 | 32.4 |
| External spur | 20 | 4.4 | - | 12.0 | 12.4 | - 17.8 | 7.5 | 15.2 | 1.8 | 1.4 | 8.2 | 14.4 |
| External branches length ratio | 20 | 65\% | - | 82\% |  | - | 72\% | - | 4\% | - | 75\% | - |
| Internal primary branch | 26 | 15.3 | - | 29.1 | 39.7 | - 51.7 | 21.5 | 45.6 | 4.6 | 2.7 | 24.8 | 43.7 |
| Internal base + secondary branch | 13 | 10.5 | - | 18.8 | 29.8 | - 36.0 | 13.6 | 32.5 | 3.0 | 1.7 | 18.7 | 32.9 |
| Internal spur | 21 | 4.5 | - | 9.8 | 13.1 | - 18.7 | 7.6 | 16.3 | 1.9 | 1.7 | 9.5 | 16.7 |
| Internal branches length ratio | 12 | 62\% | - | 80\% |  | - | 71\% | - | 5\% | - | 75\% | - |
| Claw 4 lengths |  |  |  |  |  |  |  |  |  |  |  |  |
| Anterior primary branch | 30 | 19.6 | - | 37.9 | 50.9 | - 63.2 | 26.6 | 56.9 | 5.4 | 3.2 | 30.7 | 54.0 |
| Anterior base + secondary branch | 29 | 11.2 | - | 25.5 | 32.2 | - 39.1 | 16.3 | 34.9 | 3.8 | 1.9 | 18.4 | 32.4 |
| Anterior spur | 28 | 4.3 | - | 9.6 | 12.5 | - 17.0 | 7.2 | 15.0 | 1.6 | 1.1 | 8.7 | 15.3 |
| Anterior branches length ratio | 29 | 57\% | - | 72\% |  | - | 61\% | - | 4\% | - | 60\% | - |
| Posterior primary branch | 29 | 18.2 | - | 35.0 | 46.9 | - 60.8 | 25.1 | 53.8 | 5.1 | 3.8 | 28.4 | 50.0 |
| Posterior base + secondary branch | 25 | 10.9 | - | 25.7 | 30.7 | - 39.8 | 16.4 | 35.1 | 3.8 | 2.3 | 18.1 | 31.9 |
| Posterior spur | 27 | 4.2 |  | 10.8 | 12.8 | - 19.9 | 8.0 | 16.5 | 2.1 | 1.9 | 8.9 | 15.7 |
| Posterior branches length ratio | 24 | 58\% | - | 73\% |  | - | 65\% | - | 4\% | - | 64\% | - |



FIGURE 1. Milnesium alpigenum Ehrenberg, 1853. A-habitus, ventral view (neotype, PCM). B-habitus, dorsal view (SEM). C-dorsal cuticle, the arrow indicates area with visible pseudopores (neotype, PCM). D-dorsal cuticle with the barely visible outline of a pseudoplate (neoparatype, SEM). E-buccal apparatus (neotype, PCM). F-six peribuccal lamellae with the $4+2$ configuration (neoparatype, SEM). G-claws III with the cuticular bar below (neoparatype, PCM). H-claws IV (neotype, PCM). All scale bars in $\mu \mathrm{m}$.

- M. antarcticum Tumanov, 2006, only recorded from the Antarctic (Smykla et al. 2012), by the maximal length of the buccal tube ( $\leq 68 \mu \mathrm{~m}$ in $M$. alpigenum $v s>68 \mu \mathrm{~m}$ in $M$. antarcticum), a smaller buccal tube standard width (8.6-23.6 $\mu \mathrm{m}$ in M. alpigenum vs $25.9-31.8 \mu \mathrm{~m}$ in M. antarcticum), and by a statistically lower pt of the stylet support insertion point (61.1-70.3, on average 64.9 in $M$. alpigenum vs $70.0-73.7$, on average 71.5 in $M$. antarcticum; $t_{38}=16.708, \mathrm{p}<0.001$ ).
- M. argentinum Roszkowska, Ostrowska \& Kaczmarek, 2015, reported from Argentina, by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the dorsal cuticle in $M$. alpigenum $v s$ well-visible pseudopores in M. argentinum on the entire dorsum with a standard PCM), and by the lower pt of the primary branches IV (46.9-63.2 in M. alpigenum vs 28.4-36.4 in M. argentinum).
- M. asiaticum Tumanov, 2006, recorded from Kirghizstan (type locality), China (Beasley \& Miller 2007), Estonia (Zawierucha et al. 2014) and the Svalbard archipelago (Kaczmarek et al. 2012), by a statistically lower pt of primary branches III (39.7-51.7, on average 45.8 in M. alpigenum vs 51.5-58.3, on average 55.3 in M. asiaticum; $t_{36}=15.385, \mathrm{p}<0.001$ ) and by a lower pt of primary branches IV (46.9-63.2 in M. alpigenum vs 63.9-76.0 in M. asiaticum).
- M. barbadosense Meyer \& Hinton, 2012, only reported from the type locality in Barbados, by a lower pt of the stylet support insertion point (61.1-70.3 in M. alpigenum vs 71.6-82.1 in M. barbadosense) and by a higher pt of the primary branches IV (46.9-63.2 in M. alpigenum vs 28.4-42.2 in M. barbadosense).
- M. beatae Roszkowska, Ostrowska \& Kaczmarek, 2015, only reported from the type locality in Argentina, by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the dorsal cuticle in M. alpigenum vs well-visible pseudopores in M. beatae on the entire dorsum with a standard PCM), by a more slender buccal tube (standard width/length ratio 21-38\% in M. alpigenum $v s$ standard width/ length ratio 58-66\% in M. beatae).
- M. bohleberi Bartels, Nelson, Kaczmarek \& Michalczyk, 2014, recorded from North Carolina and Tennessee, USA, by a more slender buccal tube (standard width/length ratio $21-38 \%$ in M. alpigenum $v s$ standard width/ length ratio 54-64\% in M. bohleberi).
- M. brachyungue Binda \& Pilato, 1990, recorded from the type locality in Chile and south Argentina (Roszkowska et al. 2016), by a higher pt of primary branches of all claws (36.0-63.2 in M. alpigenum vs 22.933.1 in M. brachyungue).
- M. burgessi Schlabach, Donaldson, Hobelman, Miller \& Lowman, 2018, reported from Kansas, USA, by a higher $p t$ of the buccal tube standard width (21.0-38.4 in M. alpigenum vs 52.9-68.5 in $M$. burgessi) and by a lower pt of primary branches IV (46.9-63.2 in M. alpigenum vs 66.6-96.2. in M. burgessi).
- M. dornensis Ciobanu, Roszkowska \& Kaczmarek, 2015, recorded from Romania (type locality), Poland (Kaczmarek et al. 2018) and Tunisia (Gąsiorek et al. 2017b), by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the dorsal cuticle in M. alpigenum vs well-visible pseudopores in $M$. dornensis on the entire dorsum with a standard PCM) and by a statistically lower $p t$ of the buccal tube standard width (21.0-38.4, on average 31.4 in $M$. alpigenum vs $37.8-51.6$, on average 44.1 in $M$. dornensis; $t_{43}=10.473, \mathrm{p}<0.001$ ).
- M. eurystomum Maucci, 1991, recorded from Greenland (type locality), Chile and Argentina (Maucci 1996), and Mongolia (Kaczmarek \& Michalczyk 2006), by a more slender buccal tube (standard width/length ratio $21-38 \%$ in M. alpigenum vs standard width/length ratio $62-65 \%$ in M. eurystomum).
- M. longiungue Tumanov, 2006, reported from the type locality in the Himalayas (India) and China (Beasley \& Miller 2007), by the presence of accessory points on primary branches, a lower pt of primary branches III (39.7-51.7 in M. alpigenum vs 57.1-73.5 in M. longiungue), and by a lower pt of primary branches IV (46.963.2 in M. alpigenum vs 81.8-92.4 in M. longiungue).
- M. minutum Pilato \& Lisi, 2016, only reported from the type locality in Sicily, by a lower pt of the buccal tube standard width (21.0-38.4 in M. alpigenum vs 38.6-42.4 in M. minutum).
- M. sandrae Pilato \& Lisi, 2016, only reported from the type locality in Hawaii, by a higher pt of the stylet support insertion point (61.1-70.3 in M. alpigenum vs 58.0-60.5 in M. sandrae) and by a lower pt of the buccal tube standard width (21.0-38.4 in M. alpigenum vs 44.9-48.0 in M. sandrae).
- M. shilohae Meyer, 2015, only reported from the type locality in Hawaii, by a lower pt of the stylet support insertion point ( $61.1-70.3$ in M. alpigenum vs $75.5-77.5$ in M. shilohae), a lower pt of the buccal tube standard width (21.0-38.4 in M. alpigenum vs 47.1-55.9 in M. shilohae), and by a higher pt of external spurs I-III (11.3-17.8 in M. alpigenum vs 1.9-7.5 in M. shilohae).
- M. swansoni Young, Chappell, Miller \& Lowman, 2016, only reported from the type locality in USA, by a higher number of peribuccal lamellae (six in M. alpigenum vs four in M. swansoni), a lower pt of the buccal tube standard width (21.0-38.4 in M. alpigenum vs 39.2-42.2 in M. swansoni), a lower pt of the posterior buccal tube width (23.7-39.4 in M. alpigenum vs 39.9-42.2 in M. swansoni), and by a lower pt of primary branches I (36.0-47.9 in M. alpigenum vs 48.4-53.7 in M. swansoni). It should be noted that the number of peribuccal lamellae in M. swansoni was identified only with the use of PCM, thus until SEM observations are made, the number of lamellae should be treated as a working hypothesis.
- M. tumanovi Pilato, Sabella \& Lisi, 2016, only reported from the type locality in Crimea, by a higher pt of the stylet support insertion point (61.1-70.3 in M. alpigenum specimens $383-983 \mu \mathrm{~m}$ long vs ca. 52.3 in $M$. tumanovi in a specimen $774 \mu \mathrm{~m}$ long) and by a lower $p t$ of the buccal tube standard width (21.0-38.4 in M . alpigenum specimens $383-983 \mu \mathrm{~m}$ long vs ca. 55.1 in M. tumanovi in a specimen $774 \mu \mathrm{~m}$ long).
- M. validum Pilato, Sabella, D'Urso \& Lisi, 2017 only reported from the type locality in the Antarctic, according to the measurements presented in the description of $M$. validum all pt ranges overlap, but a comparison of specimens of a similar body length (414-509 $\mu \mathrm{m}$ in M. alpigenum and 424-482 $\mu \mathrm{m}$ in $M$. validum) shows that $M$. alpigenum has a shorter buccal tube ( $33.0-43.0 \mu \mathrm{~m}$ in $M$. alpigenum vs $44.1-55.6 \mu \mathrm{~m}$ in M. validum), moreover the two species differ in the shape of secondary branches (slender in M. alpigenum vs robust in M. validum, compare Fig. 1G-H here and Fig. 6B-D in Pilato et al. 2017), and in the shape of spurs (of typical width in M. alpigenum vs very thin in M. validum).
- M. inceptum sp. nov. (described below), recorded from Germany, Japan, Switzerland and Bulgaria-please see the section "Delineation of M. alpigenum and M. inceptum sp. nov." below for a detailed differential diagnosis between these two pseudocryptic species.
- M. zsalakoae Meyer \& Hinton, 2010, recorded from Arizona and New Mexico (USA), by the presence of accessory points on primary branches and by a lower pt of primary branches of all claws (36.0-63.2 in $M$. alpigenum vs 64.4-102.9 in M. zsalakoae).

Genotypic differential diagnosis: All sequences obtained for $M$. alpigenum were unique and distinct from the sequences deposited in GenBank. The ranges of the uncorrected p-distances between neotype M. alpigenum and sequences of other congeners are as follows:

- 18S rRNA: $1.1 \%-3.6 \%(2.6 \%$ on average $)$, with the most similar being $M$. inceptum sp. nov. from Europe (MH000383, present study) and the least similar being an undetermined species from Marion Island in the subAntarctic (EU266922, Sands et al. 2008).
- 28S rRNA: $4.5 \%-8.0 \%$ ( $6.1 \%$ on average), with the most similar being an undetermined species from the USA (JX888585-7, Adams et al, unpublished) and the least similar being M. tardigradum s.s. from Poland (KC138809, Zawierucha, unpublished).
- ITS-2: $20.4 \%-23.2 \%$ ( $20.2 \%$ on average), with the most similar being M. tardigradum s.s. from Germany (JF951049, Michalczyk et al. 2012a) and the least similar being M. tardigradum s.s. from France (MG923555, Morek et al. 2019).
- COI: $14.8 \%-25.8 \%$ ( $17.4 \%$ on average), with the most similar being M. variefidum from the UK (KT951663, Morek et al. 2016a) and an undetermined species from the USA (KX306950, Fox et al., unpublished), whereas the least similar being an undetermined species from the Antarctic (KP013598, Velasco-Castrillón et al. 2015).


## Milnesium inceptum sp. nov.

Figure 2, Tables 4-5
M. tardigradum: Suzuki 2003, Schill at al. 2004, Suzuki 2006, Pfannkuchen et al. 2007, Schill 2007, Schill \& Steinbruck 2007, Hengherr et al. 2008a, Hengherr et al. 2008b, Jönsson et al. 2008, Schill \& Fritz 2008, Suzuki 2008, Takahashi et al. 2008, Förster et al. 2009, Hengherr et al. 2009a, Hengherr et al. 2009b, Neumann et al. 2009, Hengherr et al. 2010, Mali et al. 2010, Reuner et al. 2010a, Reuner et al. 2010b, Schökraie et al. 2010, Shcherbakov et al. 2010, Grohme et al. 2011, Förster et al. 2011, Schökraie et al. 2011, Wełnicz, et al. 2011, Beisser, et al. 2012, Förster et al. 2012, Schökraie et al. 2012, Grohme et al. 2013, Wang et al. 2014, Jönsson et al. 2016.
M. cf. alpigenum strain Mil.alp_DE.001: Kosztyła et al. (2016), Morek et al. (2016ab), Stec et al. (2016).


FIGURE 2. Milnesium inceptum sp. nov. A-habitus, ventral view (holotype, PCM). B-dorsal cuticle (holotype, German population, PCM). C-dorsal cuticle with faint pseudopores (specimen from Bulgaria, the white arrowhead indicates the area where the pseudopores are more densely arranged, PCM). D-dorsal cuticle with the barely visible outline of a pseudoplate (paratype, SEM). E—pseudoplate surface (paratype, SEM). F-buccal apparatus (holotype, PCM). G-six peribuccal lamellae with the $4+2$ configuration. H-claws I with the cuticular bar below (paratype, PCM). I-claws IV (paratype, PCM). All scale bars in $\mu \mathrm{m}$.
TABLE 4. Measurements (in $\mu \mathrm{m}$ ) and the pt values of selected morphological structures of 30 specimens (holotype and paratypes) of Milnesium inceptum sp. nov. from the laboratory culture derived from the type locality, from Tui equal representation of all available life stages as possible.

| CHARACTER | N | RANGE |  |  |  |  |  | MEAN |  | SD |  | Holotype |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ |  |  | pt |  |  | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | $p t$ |
| Body length | 30 | 326 | - | 848 | 1136 | - | 1588 | 583 | 1381 | 164 | 122 | 743 | 1561 |
| Peribuccal papillae length | 13 | 4.5 | - | 11.1 | 17.2 | - | 21.6 | 8.6 | 19.1 | 2.0 | 1.4 | 10.3 | 21.6 |
| Lateral papillae length | 27 | 3.7 | - | 10.6 | 13.4 | - | 19.8 | 6.8 | 15.5 | 1.9 | 1.6 | 8.0 | 16.8 |
| Buccal tube |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Length | 30 | 25.8 | - | 53.5 |  | - |  | 41.8 | - | 9.7 | - | 47.6 | - |
| Stylet support insertion point | 30 | 17.1 | - | 34.3 | 61.4 | - | 69.4 | 27.1 | 65.0 | 5.9 | 1.8 | 30.9 | 64.9 |
| Anterior width | 30 | 8.4 | - | 20.9 | 30.8 | - | 42.9 | 15.4 | 36.5 | 4.4 | 3.1 | 20.4 | 42.9 |
| Standard width | 30 | 7.1 | - | 19.6 | 23.1 | - | 37.8 | 13.8 | 32.5 | 4.2 | 3.7 | 18.0 | 37.8 |
| Posterior width | 30 | 7.4 | - | 20.1 | 25.2 | - | 39.9 | 14.1 | 33.0 | 4.3 | 3.7 | 19.0 | 39.9 |
| Standard width/length ratio | 30 | 23\% | - | 38\% |  | - |  | 32\% | - | 4\% | - | 38\% | - |
| Posterior/anterior width ratio | 30 | 81\% | - | 99\% |  | - |  | 91\% | - | 4\% | - | 93\% | - |
| Claw 1 lengths |  |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 27 | 11.0 | - | 22.4 | 35.1 | - | 45.6 | 17.7 | 40.8 | 3.7 | 2.9 | 20.9 | 43.9 |
| External base + secondary branch | 28 | 8.3 | - | 17.1 | 26.8 | - | 35.2 | 12.6 | 30.6 | 2.9 | 2.4 | 15.6 | 32.8 |
| External spur | 16 | 2.1 | - | 6.6 | 7.6 | - | 14.8 | 4.6 | 11.0 | 1.4 | 1.9 | 5.0 | 10.5 |
| External branches length ratio | 25 | 67\% | - | 79\% |  | - |  | 74\% | - | 4\% | - | 75\% | - |
| Internal primary branch | 28 | 10.0 | - | 21.4 | 33.3 | - | 43.8 | 16.6 | 39.6 | 3.7 | 2.6 | 20.7 | 43.5 |
| Internal base + secondary branch | 21 | 8.2 | - | 17.4 | 24.4 | - | 32.7 | 11.9 | 29.7 | 3.0 | 2.6 | 14.7 | 30.9 |
| Internal spur | 19 | 3.4 | - |  | 11.5 | - | 15.7 | 5.8 | 13.6 | 1.6 | 1.4 | 6.7 | 14.1 |
| Internal branches length ratio | 19 | 68\% | - | 86\% |  | - |  | 74\% | - | 4\% | - | 71\% | - |
| Claw 2 lengths |  |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 29 | 10.8 | - | 25.0 | 37.5 | - | 50.0 | 18.3 | 43.3 | 4.2 | 3.1 | 21.6 | 45.4 |
| External base + secondary branch | 21 | 8.8 | - | 17.2 | 27.4 | - | 36.7 | 13.2 | 31.6 | 3.1 | 2.4 | ? | ? |
| External spur | 15 | 2.9 | - | 7.1 | 9.2 | - | 15.2 | 4.8 | 11.8 | 1.2 | 1.5 | ? | ? |

[^0]TABLE 4. (Continued)

| CHARACTER | N | RANGE |  |  |  |  |  | MEAN |  | SD |  | Holotype |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ |  |  | $p t$ |  |  | $\mu \mathrm{m}$ | $p t$ | $\mu \mathrm{m}$ | $p t$ | $\mu \mathrm{m}$ | $p t$ |
| External branches length ratio | 20 | 67\% | - | 81\% |  | - |  | 73\% | - | 4\% | - | ? | - |
| Internal primary branch | 28 | 10.6 | - | 22.9 | 38.8 | - | 46.3 | 17.4 | 42.4 | 4.0 | 2.3 | 20.0 | 42.0 |
| Internal base + secondary branch | 16 | 8.3 | - | 17.7 | 27.3 | - | 33.6 | 12.3 | 30.7 | 3.3 | 2.1 | 15.3 | 32.1 |
| Internal spur | 16 | 3.6 | - | 9.1 | 12.1 | - | 18.0 | 6.3 | 14.8 | 1.8 | 1.7 | 6.2 | 13.0 |
| Internal branches length ratio | 14 | 68\% | - | 77\% |  | - |  | 71\% | - | 2\% | - | 77\% | - |
| Claw 3 lengths |  |  |  |  |  |  |  |  |  |  |  |  |  |
| External primary branch | 30 | 11.7 | - | 24.0 | 38.8 | - | 51.2 | 18.4 | 44.2 | 4.1 | 2.8 | 21.7 | 45.6 |
| External base + secondary branch | 26 | 8.5 | - | 18.2 | 27.6 | - | 36.3 | 13.5 | 32.0 | 3.2 | 2.2 | 15.7 | 33.0 |
| External spur | 15 | 3.3 | - | 7.9 | 8.9 | - | 14.8 | 5.0 | 12.0 | 1.5 | 1.9 | ? | ? |
| External branches length ratio | 26 | 67\% | - | 80\% |  | - |  | 72\% | - | 3\% | - | 72\% | - |
| Internal primary branch | 28 | 10.6 | - | 22.2 | 37.3 | - | 47.7 | 17.4 | 42.5 | 4.0 | 2.6 | 20.0 | 42.0 |
| Internal base + secondary branch | 11 | 8.3 | - | 14.9 | 25.9 | - | 33.8 | 10.6 | 30.4 | 2.4 | 2.3 | ? | ? |
| Internal spur | 14 | 3.7 | - | 9.0 | 13.1 | - | 17.8 | 5.9 | 15.1 | 1.7 | 1.5 | 6.5 | 13.7 |
| Internal branches length ratio | 11 | 62\% | - | 78\% |  | - |  | 70\% | - | 5\% | - | ? | - |
| Claw 4 lengths |  |  |  |  |  |  |  |  |  |  |  |  |  |
| Anterior primary branch | 29 | 13.3 | - | 27.9 | 45.9 | - | 59.5 | 22.2 | 52.7 | 4.9 | 3.3 | 24.7 | 51.9 |
| Anterior base + secondary branch | 29 | 8.8 | - | 19.9 | 32.8 | - | 40.2 | 15.2 | 35.7 | 3.7 | 2.0 | 17.6 | 37.0 |
| Anterior spur | 16 | 2.6 | - |  | 7.4 | - | 18.1 | 5.1 | 12.0 | 1.7 | 2.7 | ? | ? |
| Anterior branches length ratio | 28 | 61\% | - | 76\% |  | - |  | 68\% | - | 4\% | - | 71\% | - |
| Posterior primary branch | 27 | 12.1 | - | 28.9 | 44.1 | - | 58.4 | 20.9 | 49.8 | 5.0 | 3.2 | ? | ? |
| Posterior base + secondary branch | 26 | 8.5 | - | 20.1 | 29.7 |  | 40.3 | 14.3 | 34.4 | 3.6 | 2.6 | 17.4 | 36.6 |
| Posterior spur | 25 | 3.0 | - |  | 11.6 | - | 19.2 | 6.7 | 15.9 | 2.0 | 1.8 | 7.0 | 14.7 |
| Posterior branches length ratio | 23 | 61\% |  | 81\% |  | - |  | 68\% | - | 4\% | - | ? | - |

Material examined: Type series consisting of 96 specimens (population DE.001) and additional 93 specimens (15 from JP. 010 population, 9 from population CH. 002 , and 69 from BG. 058 population). See Table 1 and "Type repositories" below for details.

Integrative description. Females: Body yellowish. Eyes present in live specimens, dissolved after fixation in Hoyer's medium in $50 \%$ of specimens (remained visible in $1 / 30=3 \%$ specimens of the German type series, $15 / 15$ $=100 \%$ of the Japanese series, $9 / 9$ specimens $=100 \%$ of the Swiss series, and in $14 / 23$ specimens $=61 \%$ of the Bulgarian series). Cuticle with very small pseudopores ( $0.46 \pm 0.06 \mu \mathrm{~m}$, detectable only under a high quality PCM) in the German and the Swiss population and with slightly larger (but still small, $0.62 \pm 0.06 \mu \mathrm{~m}$ ) pseudopores in the Bulgarian and the Japanese population (detectable under a standard PCM). In the German and the Swiss population, the cuticle on the entire body appears smooth under PCM (Fig. 2B), but under SEM a weak outline of a single dorsal pseudoplate is visible in some specimens in the caudal part of the body (Fig. 2D-E). In the Bulgarian and Japanese populations, no pseudoplates were detected either under PCM or in SEM. Six peribuccal papillae present, with the ventral being the smallest. Six triangular peribuccal lamellae of unequal size, with the two lateral being noticeable smaller than the two dorsal and the two ventral, i.e. with the $4+2$ configuration (identifiable only in SEM; Fig. 2G). Two lateral papillae present. Buccal tube funnel-shaped (Fig. 2F). Primary branches with typically developed and clearly visible accessory points. All secondary branches with three points, i.e. with the [3-3]-[3-3] CC (Fig. 2H and I). Spurs on secondary branches of moderate length. Cuticular bars under claws I-III present in the majority of examined specimens ( $23 / 29$ specimens $=79 \%$ in the German type population, $15 / 15$ specimens $=100 \%$ in the Japanese population, $7 / 9$ specimens $=78 \%$ in the Swiss population, and in $11 / 16$ specimens $=69 \%$ in the Bulgarian population; Fig. 2H).

Males: No males were found in German, Swiss, or Bulgarian populations and culturing of isolated virgin females confirmed that the type population is parthenogenetic. However, males were found to appear spontaneously in an otherwise parthenogenetic culture of the Japanese strain (Suzuki 2008). This suggests that the species is facultatively parthenogenetic with males appearing only occasionally.

Juveniles: Morphologically identical to adults, except for the lack of the cuticular pseudopores.
Hatchlings: Morphologically identical to adults, except for the lack of cuticular bars under claws I-III in the majority of examined hatchlings ( $14 / 15$ specimens $=93 \%$ ), and the absence of cuticular pseudopores.

Ontogenetic variability: No developmental variability in the CC. Pseudopores visible only in adults. Cuticular bars under claws I-III mostly absent in hatchlings but usually present in juveniles and adults.

Eggs: Oval, yellow, smooth and laid in the exuviae, up to 18 in a single clutch were found in laboratory culture.

DNA markers: All sequences were of a very good quality. The 18 S rRNA and 28 S rRNA, sequenced only in the German type population, were 1070 bp (MH000383) and 817 bp (MH000385) long, respectively. In ITS-2, two haplotypes were found: H1 was shared by the German, the Japanese and the Swiss population ( 528 bp long, MH000386), whereas H 2 was found in the Bulgarian population 528 bp , MH000387). The p-distance between the two ITS-2 haplotypes was $0.8 \%$. The COI marker exhibited three haplotypes: H1 shared by the German and the Swiss population ( $658 \mathrm{bp}, \mathrm{KU} 513422$ ), H2 in the Japanese population ( 580 bp , MK628723), and H3 in the Bulgarian population ( 647 bp , MH000381). The p-distances between the COI haplotypes were as follows: $0.5 \%$ ( H 1 vs H 2 and H 1 vs H 3 ), and $0.3 \%(\mathrm{H} 2 v s \mathrm{H} 3)$. Sequences with marked differences are provided in Appendix 1.

Morphology and genetic markers: The sample size of four populations does not allow us to formulate strong conclusions on the relationship between genetic markers and animal morphology. Nevertheless, it should be noted that populations with COI H1 (DE. 001 and CH .002 ) exhibited statistically smaller pseudopores than populations with COI H2 (JP.010) and H3 (BG.058): $0.46 \pm 0.06 \mu \mathrm{~m}$ (DE.001) vs $0.62 \pm 0.06 \mu \mathrm{~m}$ (BG.058), $t_{28}=7.450, p<0.001$. No associations were observed between ITS-2 haplotypes and phenotypic taxonomic traits.

Type locality: $48^{\circ} 33^{\prime} 42^{\prime \prime} \mathrm{N}, 09^{\circ} 03^{\prime} 48^{\prime \prime} \mathrm{E} ; 377 \mathrm{~m}$ asl: Germany, Tübingen, Bebenhausen; forest; moss on soil.
Etymology: The name of the new species originates from the Latin "inceptor", meaning "an initiator", or "a pioneer", as this species was among the very first tardigrade laboratory models. Milnesium inceptum sp. nov. has been used in a number of studies, including first studies on molecular mechanisms underlying cryptobiosis.

Type repositories: The type series consist of the holotype (slide DE.001.34) and 96 paratypes representing hatchlings, juveniles and adult females (slides DE.001.01-33). The holotype (DE.001.34) with 14 paratypes (DE.001.04-07; 32-33) are preserved at the Institute of Zoology and Biomedical Research, Jagiellonian University, Gronostajowa 9, 30-387 Kraków, Poland; 18 paratypes (DE.001.08-13) are deposited in Department of

Animal Taxonomy and Ecology, Adam Mickiewicz University, Poznań, Umultowska 89, 61-614 Poznań, Poland; 18 paratypes are deposited in the Department of Zoology, Institute of Biomaterials and Biomolecular Systems, Stuttgart University, Germany (DE.001.14-19), 18 paratypes (DE.001.20-25) are stored in Marine Biology \& Ecology Research Centre, Plymouth University, Drakes Circus, Plymouth, PL4 8AA, United Kingdom, one paratype (DE.001.34) is deposited in Natural History Museum, Cromwell Road, London SW7 5BD, United Kingdom, 18 paratypes (DE.001.26-31) are deposited in Department of Ecology and Environmental Conservation, Faculty of Biology, University of Plovdiv, Tzar Assen 24, BG-4000 Plovdiv, Bulgaria and the remaining 9 (DE.001.01-03) are deposited in the collection of Binda \& Pilato, Museum of the Department of Biological, Geological and Environmental Sciences, Section of Animal Biology "Marcello La Greca", University of Catania, Italy.

TABLE 5. Measurements (in $\mu \mathrm{m}$ ) and the pt values of selected morphological structures of 75 specimens of Milnesium inceptum sp. nov. from the type locality in Tübingen (Germany) and the additional localities from Hiyoshi (Japan), Zürich (Switzerland), and Kazanlak Valley (Bulgaria) mounted in Hoyer's medium. Individuals were chosen to represent the entire body length range, with as equal representation of all available life stages as possible.

| CHARACTER | N | RANGE |  | MEAN |  | SD |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | $p t$ | $\mu \mathrm{m}$ | $p t$ |
| Body length | 75 | 326-998 | 1136-1841 | 628.4 | 1493 | 181 | 164 |
| Peribuccal papillae length | 48 | 4.4-13.1 | 14.9-24.0 | 8.2 | 19.0 | 2.1 | 1.7 |
| Lateral papillae length | 69 | 3.4-11.2 | 11.8-21.6 | 7.1 | 16.5 | 2.1 | 2.1 |
| Buccal tube |  |  |  |  |  |  |  |
| Length | 75 | 25.8-56.2 | - | 41.6 |  | 9.1 |  |
| Stylet support insertion point | 73 | 17.1-36.4 | 59.0-71.6 | 26.9 | 65.5 | 5.4 | 2.4 |
| Anterior width | 75 | 8.4-23.8 | 28.5-45.2 | 15.5 | 36.8 | 4.4 | 4.0 |
| Standard width | 73 | 7.1-21.2 | 23.1-41.7 | 13.5 | 32.2 | 4.2 | 4.5 |
| Posterior width | 75 | 7.4-22.1 | 25.2-42.7 | 13.8 | 32.6 | 4.1 | 4.4 |
| Standard width/length ratio | 73 | 23\%-42\% | - | 32\% |  | 5\% |  |
| Posterior/anterior width ratio | 75 | 74\%-101\% | - | 87\% |  | 7\% |  |
| Claw 1 lengths |  |  |  |  |  |  |  |
| External primary branch | 65 | 11.0-26.1 | 34.2-51.4 | 18.5 | 43.2 | 4.2 | 3.7 |
| External base + secondary branch | 61 | 8.3-19.7 | 26.8-38.0 | 12.8 | 32.0 | 3.1 | 2.8 |
| External spur | 31 | 2.1-6.8 | 7.6-14.8 | 4.4 | 11.3 | 1.3 | 1.8 |
| External branches length ratio | 53 | 67\%-81\% | - | 73\% |  | 4\% |  |
| Internal primary branch | 70 | 10.0-24.8 | 32.5-50.2 | 17.5 | 42.0 | 4.2 | 3.5 |
| Internal base + secondary branch | 57 | 8.2-21.2 | 24.4-38.8 | 12.7 | 31.5 | 3.5 | 3.1 |
| Internal spur | 43 | 3.0-8.5 | 10.8-16.8 | 5.7 | 13.8 | 1.7 | 1.5 |
| Internal branches length ratio | 54 | 68\%-88\% | - | 73\% |  | 5\% |  |
| Claw 2 lengths |  |  |  |  |  |  |  |
| External primary branch | 72 | 10.8-28.4 | 37.5-56.8 | 19.6 | 46.7 | 4.8 | 4.5 |
| External base + secondary branch | 59 | 8.8-22.8 | 27.4-41.8 | 13.7 | 33.5 | 3.4 | 2.9 |
| External spur | 36 | 2.9-8.2 | 9.2-16.6 | 5.2 | 12.9 | 1.6 | 2.0 |
| External branches length ratio | 57 | 64\%-83\% | - | 70\% |  | 4\% |  |
| Internal primary branch | 70 | 10.6-27.6 | 38.8-55.0 | 18.8 | 45.5 | 4.8 | 4.3 |
| Internal base + secondary branch | 52 | 8.3-21.0 | 27.3-40.7 | 13.3 | 32.9 | 3.7 | 2.9 |
| Internal spur | 42 | 3.2-20.1 | 11.8-36.8 | 6.8 | 16.4 | 2.8 | 4.0 |
| Internal branches length ratio | 49 | 59\%-81\% | - | 70\% |  | 5\% |  |

TABLE 5. (Continued)

| CHARACTER | N | RANGE |  | MEAN |  | SD |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | pt | $\mu \mathrm{m}$ | $p t$ |
| Claw 3 lengths |  |  |  |  |  |  |  |
| External primary branch | 69 | 11.7-27.7 | 38.8-54.5 | 19.6 | 46.9 | 4.7 | 4.3 |
| External base + secondary branch | 64 | 8.5-21.9 | 27.6-42.4 | 13.9 | 33.9 | 3.3 | 3.0 |
| External spur | 37 | 3.3-8.7 | 8.9-16.8 | 5.1 | 12.4 | 1.6 | 1.8 |
| External branches length ratio | 59 | 64\%-90\% | - | 71\% |  | 6\% |  |
| Internal primary branch | 70 | 10.6-27.2 | 37.3-55.7 | 18.7 | 45.5 | 4.6 | 4.4 |
| Internal base + secondary branch | 42 | 8.3-20.5 | 25.9-39.5 | 13.0 | 33.2 | 3.8 | 2.9 |
| Internal spur | 44 | 3.3-11.0 | 11.7-21.4 | 6.6 | 16.2 | 2.1 | 2.4 |
| Internal branches length ratio | 40 | 62\%-86\% | - | 71\% |  | 6\% |  |
| Claw 4 lengths |  |  |  |  |  |  |  |
| Anterior primary branch | 67 | 13.3-34.1 | 44.2-65.7 | 23.1 | 55.5 | 5.4 | 4.7 |
| Anterior base + secondary branch | 63 | 8.8-24.2 | 31.7-46.2 | 15.5 | 37.5 | 4.0 | 3.2 |
| Anterior spur | 45 | 2.6-11.1 | 7.4-22.8 | 6.0 | 14.1 | 2.3 | 3.9 |
| Anterior branches length ratio | 59 | 61\%-76\% | - | 67\% |  | 4\% |  |
| Posterior primary branch | 64 | 12.1-32.7 | 44.1-65.7 | 22.5 | 54.3 | 5.5 | 5.8 |
| Posterior base + secondary branch | 59 | 8.5-22.8 | 29.7-43.8 | 15.2 | 36.7 | 4.0 | 3.6 |
| Posterior spur | 48 | 3.0-10.7 | 11.6-20.6 | 6.9 | 16.4 | 1.9 | 2.4 |
| Posterior branches length ratio | 55 | 60\%-81\% | - | 66\% |  | 4\% |  |

Phenotypic differential diagnosis. Milnesium inceptum sp. nov. has the [3-3]-[3-3] CC and "smooth" cuticle (i.e. cuticle smooth in SEM and with minute pseudopores, but with no sculpturing, such as reticulation, on cuticle surface), which places it in the largest group of Milnesium species that share these characteristics (19 species). Nevertheless, M. inceptum sp. nov. differs from:

- M. alpigenum Ehrenberg, 1853 only reported from the type locality in Italy-please see the section "Delineation of $M$. alpigenum and $M$. inceptum sp. nov." below for a detailed differential diagnosis between these two pseudocryptic species.
- M. antarcticum Tumanov, 2006, only reported from the Antarctic (Smykla et al. 2012), by the maximal length of the buccal tube ( $\leq 57.0 \mu \mathrm{~m}$ in the new species $v s>67.0 \mu \mathrm{~m}$ in $M$. antarcticum), by a lower buccal tube standard width (7.1-21.2 $\mu \mathrm{m}$ in the new species $v s 25.9-31.8 \mu \mathrm{~m}$ in $M$. antarcticum), and by a statistically lower $p t$ of the stylet support insertion point (59.0-71.6, on average 65.5 in the new species $v s 70.0-73.7$, on average 71.5 in $M$. antarcticum; $t_{80}=20.590, \mathrm{p}<0.001$ ).
- M. argentinum Roszkowska, Ostrowska \& Kaczmarek, 2015, recorded from Argentina, by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the dorsal cuticle in the new species $v s$ well-visible pseudopores in M. argentinum on the entire dorsum with a standard PCM), the maximal length of the buccal tube (up to $57 \mu \mathrm{~m}$ in the new species $v s$ up to $74 \mu \mathrm{~m}$ in $M$. argentinum), and by a lower pt of the primary branches IV (44.1-65.7 in the new species vs 28.4-36.4 in M. argentinum).
- M. asiaticum Tumanov, 2006, recorded from Kirghizstan (type locality), China (Beasley \& Miller 2007), Estonia (Zawierucha et al. 2014), and the Svalbard archipelago (Kaczmarek et al. 2012), by a statistically lower pt of primary branches IV (44.1-65.7, on average 54.8 in the new species vs 63.9-76.0, on average 69.7 in M. asiaticum; $t_{54}=26.040, \mathrm{p}<0.001$ ).
- M. barbadosense Meyer \& Hinton, 2012, only reported from the type locality in Barbados, by a lower pt of the stylet support insertion point (59.0-71.6 in the new species vs 71.6-82.1 in M. barbadosense), and by a higher $p t$ of the primary branches IV (44.1-65.7 in the new species vs 28.4-42.2 in M. barbadosense).
- M. beatae Roszkowska, Ostrowska \& Kaczmarek, 2015, only reported from the type locality in Argentina, by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the
dorsal cuticle in the new species $v s$ well-visible pseudopores in $M$. argentinum on the entire dorsum with a standard PCM), and by more elongated buccal tube (standard width/length ratio $23-42 \%$ in the new species $v s$ standard width/length ratio $58-66 \%$ in $M$. beatae)
- M. bohleberi Bartels, Nelson, Kaczmarek \& Michalczyk, 2014, recorded from North Carolina and Tennessee, USA, by the more slender buccal tube (standard width/length ratio $23-42 \%$ in the new species $v s$ standard width/length ratio 54-64\% in M. bohleberi).
- M. brachyungue Binda \& Pilato, 1990, reported from the type locality in Chile and south Argentina (Roszkowska et al. 2016), by a higher pt of primary branches of claws I-III (32.5-56.8 in the new species vs 22.9-27.1 in M. brachyungue) and by the pt of primary branches IV (44.1-65.7 in the new species vs 33.1 in M. brachyungue).
- M. burgessi Schlabach, Donaldson, Hobelman, Miller \& Lowman, 2018, recorded from Kansas, USA, by a higher pt of the buccal tube standard width (23.1-41.7 in the new species vs 52.9-68.5 in M. burgessi) and by the lower pt of primary branches IV (44.1-65.7 in the new species vs 66.6-96.2. in M. burgessi).
- M. dornensis Ciobanu, Roszkowska \& Kaczmarek, 2015, recorded from Romania (type locality), Poland (Kaczmarek et al. 2018) and Tunisia (Gąsiorek et al. 2017b), by the appearance of cuticle (faint pseudopores visible only with a high quality PCM on the caudal part of the dorsal cuticle in the new species vs well-visible pseudopores in M. dornensis on the entire dorsum with a standard PCM), and by a statistically lower pt of buccal tube standard width (23.1-41.7, on average 32.2 in the new species vs $37.8-51.6$, on average 44.1 in $M$. dornensis; $t_{22}=10.686, \mathrm{p}<0.001$ ).
- M. eurystomum Maucci, 1991, recorded from Greenland (type locality), Argentina and Chile (Maucci 1996), and Mongolia (Kaczmarek \& Michalczyk 2006), by a more slender buccal tube (standard width/length ratio $23-42 \%$ in the new species $v s$ standard width/length ratio $62-65 \%$ in M. eurystomum).
- M. longiungue Tumanov, 2006, reported from the Himalayas (India, type locality) and China (Beasley \& Miller 2007), by the presence of accessory points on primary branches, a lower pt of primary branches III (37.3-55.7 in the new species vs 57.1-73.5 in M. longiungue), and by the lower pt of primary branches IV (44.1-65.7 in the new species vs 81.8-92.4 in M. longiungue).
- M. minutum Pilato \& Lisi, 2016, only reported from the type locality in Sicily, by a statistically lower pt of the buccal tube standard width (23.1-41.7, on average 32.2 in the new species vs 38.6-42.4, on average 41.1 in $M$. minutum; $t_{3}=7.990, \mathrm{p}=0.002$ ).
- M. sandrae Pilato \& Lisi, 2016, only reported from the type locality in Hawaii, by a higher pt of the stylet support insertion point (59.0-71.6, on average 65.5 in the new species vs $58.0-60.5$, on average 58.9 in $M$. sandrae; $t_{22}=8.506, \mathrm{p}<0.001$ ) and by a lower $p t$ of the buccal tube standard width (23.1-41.7 in the new species vs 44.9-48.0 in M. sandrae).
- M. shilohae Meyer, 2015, only reported from the type locality in Hawaii, by a lower pt of the stylet support insertion point (59.0-71.6 in the new species vs 75.5-77.5 in M. shilohae) and by a lower pt of the buccal tube standard width (23.1-41.7 in the new species vs 47.1-55.9 in M. shilohae).
- M. swansoni Young, Chappell, Miller \& Lowman, 2016, only reported from the type locality in the USA, by a higher number of peribuccal lamellae (six in the new species vs four in M. swansoni; but note that the number of peribuccal lamellae in M. swansoni was determined only with a PCM) and by a lower pt of the buccal tube standard width (23.1-41.7, on average 32.2 in the new species vs 39.2-42.2, on average 40.3 in M. swansoni; $\left.t_{12}=10.325, \mathrm{p}<0.001\right)$.
- M. tumanovi Pilato, Sabella \& Lisi, 2016, only reported from the type locality in Crimea, by a higher pt of the stylet support insertion point (59.0-71.6 in the new species specimens being 326-998 $\mu \mathrm{m}$ long vs 52.3 in M . tumanovi in a specimen $774 \mu \mathrm{~m}$ long) and by a lower $p t$ of the buccal tube standard width (23.1-41.7 in the new species in specimens $326-998 \mu \mathrm{~m}$ long vs 55.1 in $M$. tumanovi in a specimen $774 \mu \mathrm{~m}$ long).
- M. validum Pilato, Sabella, D’Urso \& Lisi, 2017, only reported from the type locality in the Antarctic; according to measurements presented in the description of $M$. validum all pt ranges overlap, but a comparison between specimens of similar body length (393-513 $\mu \mathrm{m}$ in the new species and 424-482 $\mu \mathrm{m}$ in M. validum) shows that $M$. inceptum $\mathbf{s p}$. nov. has a shorter buccal tube ( $27.1-39.0 \mu \mathrm{~m}$ in the new species $v s 44.1-55.6$ in $M$. validum), moreover the two species differ in the shape of the secondary branches (typical in the new species vs robust in M. validum, compare Fig. 2H-I here and Fig. 6B-D in Pilato et al. 2017), and in the shape of spurs (moderate length and of normal width in the new species vs long and very thin in M. validum).
- M. zsalakoae Meyer \& Hinton, 2010, recorded from Arizona and New Mexico (USA), by the presence of accessory points on primary branches, by a lower pt of primary branches I-III (32.5-56.8 in the new species $v s$ 64.4-88.6 in M. zsalakoae) and by a lower pt of primary branches IV (44.1-65.7 in the new species vs 94.8102.9 in M. zsalakoae).

Genotypic differential diagnosis: Four sequences deposited in GenBank prior to this publication, labelled as "M. tardigradum", in fact represent M. inceptum: two ITS-2 (GQ403681-2) and two COI (EU244603-4) (all by Schill, unpublished). The GQ403683 and EU244604 sequences originated from Germany and represent the same laboratory strain that was utilised herein to describe the new species. The sequences GQ403682 and EU244603 originated from Japan and represent the Japanese strain, also used in the present study.

The ranges of uncorrected p-distances between the new species and sequences of other congeners are as follows:

- 18S rRNA: $1.1 \%-3.9 \%$ ( $2.9 \%$ on average), with the most similar being M. alpigenum, (MG996146, present study) and the least similar being an undetermined species from the USA (GQ925696, Chen et al. unpublished) as well as an undetermined species from South Georgia in the sub-Antarctic (EU266922, Sands et al. 2008).
- 28S rRNA: $0.4 \%-8.8 \%$ ( $6.0 \%$ on average), with the most similar being an undetermined species from the USA (AY210826, Mallatt et al. unpublished) and another undetermined species also from the USA (JX888540-1, Adams et al. unpublished) and the least similar being an undetermined species from Spain (FJ435779-80, Guil \& Giribet 2012).
- ITS-2: $19.6 \%-22.8 \%$ ( $20.3 \%$ on average), with the most similar being M. tardigradum s.s. from Hungary and Poland (MG923553, Morek et al. 2019) and the least similar being M. tardigradum s.s. from France (MG923555, Morek et al. 2019).
- COI: $17.8 \%-25.8 \%$ ( $19.7 \%$ on average), with the most similar being $M$. dornensis from Romania (MG923566, Morek et al. 2019) and an undetermined species from the USA (KX306950, Fox et al., unpublished) whereas the least similar being two undetermined species from the Antarctic (KP013601 and KP013598, Velasco-Castrillón et al. 2015).


## Delineation of M. alpigenum and M. inceptum sp. nov.

The two species are genetically distinct but morphologically very similar, although not identical. Therefore, they could be classified as pseudocryptic species, i.e. species that can be differentiated morphologically but only with a detailed analysis; in this case-with the use of statistical testing of morphometric traits measured in a number of specimens, since the classical identification based on qualitative traits and non-overlapping morphometric ranges is not sufficient to tell the species apart.

The PCA analysis indicated three components comprising $59.3 \%$ of the total variance in the $p t$ ratios. The three factors were as follows: PC1: the pt of the primary and secondary branches of all claws (38.2\%), PC2: the pt of external and posterior spurs (11.3\%), and PC3: the pt of buccal tube widths and stylet support insertion point $(9.8 \%)$. The relationships between the principal components are shown in Fig. 3. The two species did not differ in PC 1 and PC3 but in contrast, they differed in PC2, thus comparisons of the first three principal components did not result in congruent conclusions. Specifically, when PC1 and PC3 were compared (Fig. 3B), ranges for the two species largely overlapped. In the PC1 vs PC2 comparison (Fig. 3A), the overlap between species was smaller. Finally, when PC2 and PC3 were compared, the ranges barely overlapped (Fig. 3C). Thus, we compared only the traits constituting PC2 (i.e. the pt values of claw spurs) with a series of $t$-tests and adjusted $\alpha$-levels. Student's $t$ tests revealed significant differences in three of the eight compared traits (mean values, $\pm \mathrm{SD}$, and [ranges] for $M$. alpigenum vs $M$. inceptum sp. nov.):

- spur on external claw I: $13.3 \pm 1.5[11.3-17.8]$ vs $11.3 \pm 1.3[7.6-14.8], t_{46}=4.002, p<0.001$;
- spur on external claw II: $15.3 \pm 1.4[12.7-17.5]$ vs $12.9 \pm 2.0[9.2-16.6], t_{54}=4.630, p<0.001$;
- $\quad$ spur on external claw III: $15.2 \pm 1.4[12.4-17.8]$ vs $12.4 \pm 1.8[8.9-16.8], t_{48}=5.758, p<0.001$.

In other words, the analysis showed that M. alpigenum has statistically longer (relatively to the buccal tube length) external spurs than M. inceptum sp. nov. (compare also Fig. 1G-H and 2H-I).

In contrast to subtle morphometric differences, the two species exhibit considerable genetic distances in all four analysed markers. Specifically, they differ by $1.0 \%$ in 18 S rRNA, $5.2 \%$ in 28 S rRNA, $21.6 \%$ in ITS-2, and by $18.1 \%$ in COI. Most importantly, however, the two species are not immediately related to each other (see Fig. 4 for the positions of both species on the Milnesium phylogenetic tree), which is the strongest evidence that the two species represent separate phylogenetic lineages.

To conclude, differences both in phenotypic and genetic traits unequivocally show that $M$. inceptum sp. nov. is a bona species. Nevertheless, an extreme care must be taken when identifying these species using solely phenotypic data.


FIGURE 3. Graphs illustrating the relationships between the first three principal components revealed by the PCA for the single population of M. alpigenum Ehrenberg, 1853 and the three pooled populations of M. inceptum sp. nov.

## Phylogenetic relationships

Phylogenetic trees obtained with BI and ML methods did not exhibit the same topologies. Importantly, however, majority of nodes were weakly supported in the ML tree, thus we considered it uninformative and we focused on the BI tree, which had good statistical support (Fig. 4). The first clade encompasses M. tardigradum s.s. and a sister clade composed of M. berladnicorum, M. dornensis and M. variefidum. This group is in a sister relationship to the clade comprising M. inceptum sp. nov. and an undescribed species ("Milnesium hisatsinomorum"; KX306950; Fox et al., unpublished). The abovementioned taxa are in polytomy with $M$. alpigenum and an unknown species (described as "M. cf. tardigradum"; JX683822-5; Vicente et al. 2013). The following three undescribed species: EF632553 (Sands et al., unpublished), KP013613 and KJ857002 (Velasco-Castrillón et al. 2015) formed a clade that, together with another undescribed species KJ857001 (Velasco-Castrillón et al. 2015), were in polytomy with the abovementioned species. Finally, an undescribed species represented by two sequences (KP013598 and KP013601; Velasco-Castrillón et al. 2015) was a sister group to all remaining Milnesium species. Thus, our analysis indicated that $M$. inceptum sp. nov. and M. alpigenum are not sister species, even though the relationships of M. alpigenum with other congeners were not fully resolved.

## Discussion

Nearly a century since the synonymisation of M. alpigenum with M. tardigradum by Marcus (1928), M. alpigenum is now redescribed and reinstated utilising the tools of integrative taxonomy. Moreover, the redescription of $M$. alpigenum allowed us to verify the taxonomic status of the commonly used German "Tübingen" and Japanese
"Hiyoshi H-1" Milnesium laboratory strains. Both strains were originally identified as "M. tardigradum" (Suzuki 2003; Schill et al. 2004), but the redescription of M. tardigradum by Michalczyk et al. (2012a, b) showed that the original identifications were incorrect (see also Morek et al. 2019 for an updated delineation of M. tardigradum). In fact, Michalczyk et al. (2012a) hypothesised that the "Tübingen" strain might represent a new species. More recently, Morek et al. (2016a) tentatively classified the "Tübingen" strain as " $M$. cf. alpigenum" since the morphology of the strain fit the limited original description of M. alpigenum. However, the present study has shown unambiguously that both laboratory strains indeed represent a distinct taxon, M. inceptum sp. nov.


FIGURE 4. The positions of M. alpigenum Ehrenberg, 1853 and M. inceptum sp. nov. on the Milnesium phylogenetic tree based on the concatenated data set of ITS-2 and COI sequences. Branch support values are BI posterior probabilities. Species names in square brackets and in grey font are GenBank labels that are incorrect species identifications, uncertain identifications, or invalid names (correct identifications are provided in black font, before the incorrect labels). Filled circles represent both ITS-2 and COI sequences, empty circles indicate COI sequences only. Scale bar shows the number of substitutions per site.

Interestingly, M. alpigenum and $M$. inceptum sp. nov. are barely distinguishable using standard morphometric traits, even though they are genetically distant (not even immediate kin; see Fig. 4). In fact, the two species would have been unrecognised using classical taxonomic methods alone and only the use of molecular markers allowed a post hoc identification of statistical phenotypic differences. Still, the two species differ morphometrically, which means that they are not truly cryptic. As the identified differences are minor, and can only be revealed by the use of statistical tests, M. alpigenum and M. inceptum sp. nov. should be termed "pseudocryptic", meaning that they are species that exhibit minor morphometric differences, which are not detectable using classical diagnostic keys that rely on the presence or absence of qualitative (morphological) traits and non-overlapping ranges of quantitative (morphometric) traits. This example shows explicitly how important it is to base new species descriptions on a proper sample size and on both phenotypic and genetic traits.

In our opinion, species described with a few measured individuals and without associated molecular data pose a serious threat to the development of tardigrade taxonomy. At the time of description (Ehrenberg 1853), M. alpigenum was easy to differentiate from the only congener, M. tardigradum, because there were no other known Milnesium species with the [3-3]-[3-3] CC. However, now there are 22 Milnesium species with this CC, with most
of them only differentiated by morphometrics. At the time of Doyère and Ehrenberg, the knowledge and available analytical methods were, of course, limited compared to the present time. However, despite the undeniable increase in knowledge of tardigrade biology and progress in taxonomic methodology, the same process may occur again and again if new species are continually described with classical methods and low numbers of individuals. Even if a new species is obviously new (e.g. because it exhibits a unique qualitative trait) but is not integratively described, it may hinder the detection of similar species in the future. When describing a new species with a unique trait, it cannot be predicted as to whether this is the only such species in the world or whether it is the first representative of a complex of cryptic, pseudocryptic, or even only roughly similar taxa. Therefore, if the species is the first representative of a group of similar species, many of those species may go unnoticed for decades, until the nominal species is integratively redescribed. Furthermore, if individuals collected from distant locations that represent morphologically similar species are erroneously identified as the nominal species, the geographic range of the nominal species may be highly overestimated.

The history of confusion with the taxonomic identity of the "Tübingen" Milnesium laboratory strain also underlines the vital importance of integrative redescriptions. As mentioned above, the taxonomic identification of the strain was possible only after the redescriptions of M. tardigradum and M. alpigenum. Importantly, however, it needs to be underlined that if M. alpigenum was redescribed classically, the Tübingen Milnesium laboratory strain would not have been recognised as a separate species, since statistical differences in a fraction of morphometric traits-even if they were detected-would not have been accepted by the taxonomic community as sufficient evidence to erect a new species. Recently, another popular laboratory strain, originally identified as "Hypsibius dujardini (Doyère, 1840)", was shown to represent a new species, Hypsibius exemplaris Gąsiorek et al., 2018. The reason for the misidentification of the Hypsibius strain was similar to M. inceptum sp. nov., i.e. the original description of $H$. dujardini was insufficient to identify other morphologically similar species such as $H$. exemplaris (Gąsiorek et al. 2018). As there are numerous species requiring integrative redescriptions (many of them being nominotypical taxa for genera and higher taxonomic ranks), we would like to urge taxonomists to prioritise redescribing existing taxa over describing new species (e.g. see Meier \& Dikow 2004). Only when species with older, limited descriptions are integratively redescribed and new taxa are described integratively, may we hope to reliably estimate tardigrade diversity, species geographic distributions, and identify evolutionary mechanisms underlying these phenomena.

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## APPENDIX 1. Neotype and type DNA sequences

## Milnesium alpigenum Ehrenberg, 1853

18S rRNA (1054 bp, MG996146):
TAGATCGTATCATCCTACATGGATAACTGTGGTAATTCTAGAGCTAATACATGCAAAAAGCCGTCTGGCCTCGTGTCAGCGGCG CAGTTATTAGATTAAAACCAATATAGGCTTTTCGGGTCTATTAAACTTGTGATGAATCTGAATAACCGAAGCAAAGCGCATGGT CTCGTACCGGCGCTAGATCTTTCAAGTGTCTGATCTATCAGCTTGTCGTTAGGTTATGTTCCTAACGTGGCTTCGACGGGTAAC GGGGTATCAGGGTCCGATACCGGAGAGGGAGCCTGAGAAATGGCTACCACATCCAAGGAAGGCAGCAGGCGCGCAAATTACCCA CTCCCAGTTCGGGGAGGTAGTGACGAAAAATAACGATGCGGGAGCATAATGCTTCCCGTAATCGGAATGAGTACACTTTAAATC CTTTAACGAGGATCCATTGGAGGGCAAGTCTGGTGCCAGCAGCCGCGGTAATTCCAGCTCCAATAGCGTATATTAAAGTTGCTG CGGTTAAAAAGCTCGTAGTTGGATCTGGGTGTTTCGATGAGTGGTGCATCTATTCGATGTCTACTACTCCATCGACACCACAAG CCAACCATGTCCTTCTATACCCTTCACTGGGCGTAGATAATGGGCGGTTGGAACGTTTACTTTGAAAAAATTAGAGTGCTCAAA GCAGGCGTATGGCCTTGAATAATGGTGCATGGAATAATGGAATAGGACCTCGGTTCTATTTGTTGGTTTTCAAGAGCTCGAGGT AATGATAAAGAGGAACAGACGGGGGCATTCGTATTGCGACGTTAGAGGTGAAATTCTTGGATCGTCGCAAGACGAACTACTGCG AAAGCATTTGCCAAGAATGTTTTCATTAATCAAGAACGAAAGTTAGAGGTTCGAAGGCGATCAGATACCGCCCTAGTTCTAACC ATAAACGATGCCAACCAGCGATCTGTCGGTGTTTATTTAACGACTCGACAGGCAGCTTCCGGGAAACCAAAGTGCTTAGGTTCC GGGGGAAGTATGGTTGCAAAGCTGAAACTAAAGGAATGACGAAGCC

28S rRNA (809 bp, MH000384):
TACTAAGCGGGGGAAAAGAAACCAACGGGGATTCTCCTAGTAACTGCGAGTGAACGGAGAAAAGCCCAGCGCTGAATCCTGTAG CTGGTAACGGTTATGGGAGCTGTAGCGTGAAGAAGGTGTACAACCATTGCAGTCAATACACGTAAGTCTCCCTGAGTGGAGCTC CATCCCAAGGAGGGTGCAAGGCCCGTATCGTGTTTGACGCGTGATGGTATAGCATCTTCAGAGAGTCGGGTTGTTTGGGATTGC AACCTAAAGCCGGTGGTAAACTCCATCGAAGGCTAAATATGACCACGAGTCCGATAGCGAACAAGTACCGTGAGGGAAAATTGA AAAGCACTTTGAAGAGAGAGCGAAATAGTGCGTGAAACCGCTTAGAGGCAAGCAGATGGGGCCTCGAAGGTAGAGCAGCGAATT

CAGCTTGCATTTCTGCTAGACTACTGTCGGCGTAGAGATCGTAAGACTCTTGTCGATGTAGGGTGTATAGTGGAATGTGAGTGC ACTTTCGCTGTTTGTACGCCACCGCTGTTAAGTGTGCATCCGCTGTGGTCTTGCGTGAGGCCTTGAGTGGCTTGCTGCTCAAGT CACCTACGCTTGGCTATTATGCAGCGCGTTTGCCTATTAACTGGACAAGTCATTCCTATGCCAGCATCGCTTCGGTGGTGTGAT GTCGAACACTGGCGTGTTTATTGCTGCTTCGTGGCAGTTGACGTGCTTGCACGGCTTCAGCTGCTGGTGGTATACTGCGTCGGC TCTACAGGCATAGTGTAGATTTGGTGGCGAGTAGATGGCTGCCCATCTAACCC

## ITS-2 (530 bp, MH000382):

CTTTATGAACGTTGTTTCTTCGAACGCAAATTGCGGCTATGGGTTGACCGTAGCCACGTCTGGTTGAGGGTCAAACGAAAAAAA AATGATAGCTACGTGTTTGCTATCGATTGTCTGTCATCCTATACTGGCCATCTCAGAGCCAGGCGAAGGCTGACAGATGAAGTA TCAACCCTTTGACGAGCGTATTCCTGGTCTGTAGCGGATCGGAAGCCTACGGGCGCGCACATATGCGTATATGTATGTACGGGC TGTGTGATGGCAGTAGGTTGGAGTCGCTGATAGGCTCTGTATCGCTTAGCTGGTTAATGCATGCGGCAGTTGTGCATATATAGA GCCGGCTACGAGCAGAGTTTTGACCAATCGAACGACAGCCCACTAAAGTACATACTCGCATACACAATGCGCTTGGCTTCGAGT ACAGATATCAGTACGCTGAATGGTCATAGGTGAAGCCATGTGTTTGTCGCTTGCGACATGCTCATACACATATACACTCATTAC GTTGACCTCAGCTCAGGCGAGATTAC

## COI (560 bp, MH000380):

GCCTTTTTGTAGGTTCAGCATTGAGTATATTAATTCGACTTGAATTATCTCAACCTAATTCTATACTTATAAGAGAAGATATTT ATAATGCTTTTATTACAAGTCATGCTTTAGTAATAATTTTTTTTTTTGTTATACCAGTGTTAATTGGAGGGTTTGGTAATTGAT TGGTTCCTTTAATAATTAGTTCTCCGGATATAGCTTTTCCTCGTATTAATAATGTGAGATTTTGAATGTTAATTGCTTCTTTTT TGTTATTAATTTTAAGAATATTTTCTGGAGCTGGTGTAGGAGCAGGTTGAACTTTGTATCCTCCTCTTACTAATATTTATGGTC ATAGAAATTCTTCAGTTGATTTTGCGATCTTATCATTGCATATTGCTGGTGCTTCTTCTGTGTTTAGAGCAATAAATTTTTTAA CTACAATTTTTAATATACATTATTTTGGTCTTCGAATAGATAAATTACCTTTGTTTGTATGATCAATTTTTATTACGGCAATCT TGTTAGTTTTGGCTTTGCCTGTTTTGGCTGGTGCTATTACAATATTAATTTTGATC

## Milnesium inceptum sp. nov.

18S rRNA (1070 bp, MH000383):
ATTAATCAGGTATGGGGTMCTAGATCGTATCATCCTACATGGATAACTGTGGTAATTCTAGAGCTAATACATGCAATGAAGCTG TCTGGCCTTGTGTCAGCAGCGCAGTTATTAGATTAAAACCAATATAGGCTTTCGGGTCTATTAACTTGTGATGAATCTGAATAA CCGAAGCAAAGCGCATGGTCTCGTACCGGCGCTAGATCTTTCAAGTGTCTGATCTATCAGCTTGTCGTTAGGTTATGTTCCTAA CGTGGCTTCGACGGGTAACGGGGTATCAGGGTCCGATACCGGAGAGGGAGCCTGAGAAATGGCTACCACATCCAAGGAAGGCAG CAGGCGCGCAAATTACCCACTCCCAGTTCGGGGAGGTAGTGACGAAAAATAACGATGCGGGAGCATAATGCTTCCCGTAATCGG AATGAGTACACTTTAAATCCTTTAACGAGGATCCATTGGAGGGCAAGTCTGGTGCCAGCAGCCGCGGTAATTCCAGCTCCAATA GCGTATATYAAAGTTGCTGCGGTTAAAAAGCTCGTAGTTGGATCTGGGTGTTTCGATGAGTGGTGCATCTATTCGATGTCTACT АСТССАTCGACACCACAAGCCAACCATGTCCTTCTATACCCTTCACTGGGCGTAGATAATGGGCGGTTGGAACGTTTACTTTGA AAAAATTAGAGTGCTCAAAGCAGGCGTATGGCCTTGAATAATGGTGCATGGAATAATGGAATAGGACCTCGGTTCTATTTGTTG GTTTTCAAGAGCTCGAGGTAATGATAAAGAGGAACAGACGGGGGCATTCGTATTGCGACGTTAGAGGTGAAATTCTTGGATCGT CGCAAGACGAACTACTGCGAAAGCATTTGCCAAGAATGTTTTCATTAATCAAGAACGAAAGTTAGAGGTTCGAAGGCGATCAGA TACCGCCCTAGTTCTAACCATAAACGATGCCAACCAGCGATCTGTCGGTGTTTATTTAACGACTCGACAGGCAGCTTCCGGGAA ACCAAAGTGCTTAGGTTCCGGGGGAAGTATGGTTGCAAAGCTGAACTAAAGGAATTGACGAG

## 28S rRNA (817 bp, MH000385):

TACTAAGCGGAGGAAAAGAAACCAACGGGGATTCTCCTAGTAACTGCGAGTGAACGGAGAAAAGCCCAGCGCTGAATCCTGTAG CTGGTAACGGTTACGGGAGCTGTAGCGTGAAGAAGGTGTACAACCATTGCAGTCAATACACGTAAGTCTCCCTGAGTGGAGCTC CATCCCAAGGAGGGTGCAAGGCCCGTATCGTGTTTGACGCGTGATGGTATAGCATCTTCAGAGAGTCGGGTTGTTTGGGATTGC AACCTAAAGCCGGTGGTAAACTCCATCGAAGGCTAAATATGACCACGAGTCCGATAGCGAACAAGTACCGTGAGGGAAAATTGA AAAGCACTTTGAAGAGAGAGCGAAATAGTGCGTGAAACCGCTTAGAGGCAAGCAGATGGGGCCTCGAAGGTAGAGCAGCGAATT CAGCTTGCATCTCTGCTGGACTACTGTTGGCGTAGAGATCGTAAGACTCTTGTCAATGTAGGCTGTCATAGTGGAATGTGAGTG CACTTTCGCTGTTTGTACGCCACCGCTGATAAATGTTTCTGCATCCGCTGTGGCCTTGTGTGAGGCCTTGAGTGGCTTGCTACT CAAGTAACCTACACTTGGCTATATACAGCGCGTTTGCCTTTTATCTGGTCGAGTCACATCCTATGCCGGCATTGCTTTACGGCG ATGCAGTGTAGATACTAGCGTGTTTATTGCTACTTCGCATTGCGGTTGACGTGCTTGCACGGCTGCTGCGGCTGGTGGTATACT GCGTTGGATCTACTGGTATAGTGTAGATTTGGTGGCGAGTAGACGGCTGCCCATCTAACCC

ITS-2/H1, Germany (DE.001), Japan (JP.010) and Switzerland (CH.002) (528 bp, MH000386):
CTTTATGAACGTTGTTTCTTCGAACGCAAATTGCGGCTATGGGTTGACCGTAGCCACGTCTGGTTGAGGGTCAAACGAAAAAAT CACTGATAGCTACGTGTTTGCTGTCGATTGTCTGTCAGTCTATACTGGCCATTTTAGTGCCAGGTCAAGGCTGACAGATGAAGT TTGGACCCTATGGCTAGCGTACTTTTGGTCTGTGACGGATCGGGAGCTGGCGCGTACATGCTCATATAGAGTTATACTGGGACT GTGTGATCACAGTAGGTTGGAGTCGCTGATGGGCTCTACCGTTTAGCTGACAATGCATGCGGCAGTTGTGCATAACTAGTCAGC GACGCGTAGACGCTTGGCCAACCGAACGACAGTCCACTCACGGTATACTCTGGCATGTTCAAGCGTACGGCTACCGAGTGCAGC TTCCAATACGCAGTGATATAGTCATAGGTTGATAAAGCGTGTGTCGTACGCTTCATGTGCGCGACGCATACACACTCATTACGT TGACCTCAGCTCAGGCGAGATTAC

ITS-2/H2, Bulgaria (BG.058) (528 bp, MH000387):
CTTTATGAACGTTGTTTCTTCGAACGCAAATTGCGGCTATGGGTTGACCGTAGCCACGTCTGGTTGAGGGTCAAACGAAAAAAT CACTGATAGCTACGTGTTTGCTATCGATTGTCTGTCAGTCTATACTGGCCATTTTAGTGCCAGGTCAAGGCTGACAGATGAAGT TTGGACCCTATGGCTAGCGTACTTTTGGTCTGTGACGGATCGGAAGCCGGCGCGTACATGCTCATATAGAGTTATACTGGGACT GTGTGATCACAGTAGGTTGGAGTCGCTGACGGGCTCTACCGTTTAGCTGACAATGCATGCGGCAGTTGTGCATAACTAGTCAGC GACGCGTAGACGCTTGGCCAACCGAACGACAGTCCACTCACGGTATACTCTGGCATGTTCAAGCGTACGGCTACCGAGTGCAGC TTCCAATACGCAGTGATATAGTCATAGGTTGATAAAGCGTGTGTCGTACGCTTCATGTGCGCGACGCATACACACTCATTACGT TGACCTCAGCTCAGGCGAGATTAC

COI/H1, Germany (DE.001) and Switzerland (CH.002) (658 bp, KU513422):
TATATTGTATTTTATTTTTGGTATTTGATGTGCTTTTGTAGGTTCAGGTTTAAGTGTGTTAATTCGTCTTGAATTATCTCAGCC TAACACAATATTAATAAGTGAAGATATTTATAATGCTTTTATTACAAGTCATGCTTTAGTAATGATTTTTTTTTTTGTTATACC TGTTTTAATTGGAGGTTTTGGAAATTGATTAGTTCCTCTTATAATTAGATCACCAGATATAGCTTTTCCTCGTATTAACAATGT AAGATTTTGAATATTAGTTGCTTCTTTTGGTTTGTTGCTTTTTAGAATATTTAGGGGTACAGGAGTAGGAGCTGGTTGAACACT TTATCCTCCGTTAACTAGGTATAATGGCCATAGCAGTCATGCTGTCGATTATGCAATTTTGTCTTTACATATTGCAGGAGCATC GTCAATTTTTAGTGCACTGAATTTTTTAACGACGATTATTAATATACACTATTTTGGAGTACGAATAGATAAATTACCGTTGTT TGTGTGATCGATTTTTATTACTGCTCTATTGTTAGTTTTGGCTTTACCAGTACTTGCTGGAGCAATTACAATATTAATTTCTGA TCGTAATTTTACTACTACATTTTTTGATCCGGCAGGGGGAGGAGATCCTGTTTTATTTCAACATTTATTT

COI/H2, Japan (JP.010) (580 bp, MK628723):
TTTTGTAGGTTCAGGTTTAAGTGTGTTAATTCGTCTTGAATTATCTCAGCCTAACACAATATTAATAAGTGAAGATATTTATAA TGCTTTTATTACAAGTCATGCTTTAGTAATGATTTTTTTTTTTGTTATACCTGTTTTAATTGGAGGTTTTGGAAATTGATTAGT TCCTCTTATAATTAGATCACCAGATATAGCTTTTCCTCGTATTAATAATGTAAGATTTTGAATATTAGTTGCTTCTTTTGGTTT GTTGCTTTTTAGAATATTTAGGGGTACAGGAGTAGGAGCTGGTTGAACACTTTATCCTCCGTTAACTAGGTATAATGGTCATAG CAGTCATGCTGTCGATTATGCAATTTTGTCTTTACATATTGCAGGAGCATCGTCAATTTTTAGTGCACTGAATTTTTTAACGAC GATTATTAATATACACTATTTTGGAGTACGAATAGATAAATTACCGTTGTTTGTGTGATCGATTTTTATTACTGCTCTATTGTT AGTTTTGGCTTTACCAGTACTTGCTGGAGCAATTACAATATTAATTTCTGATCGTAATTTCACTACTACATTTTTT

COI/H3, Bulgaria (BG.058) (647 bp, MH000381):
TTTTGTAGGTTCAGGTTTAAGTGTGTTAATTCGTCTTGAATTATCTCAGCCTAACACAATATTAATAAGTGAAGATATTTATAA TGCTTTTATTACAAGTCATGCTTTAGTAATGATTTTTTTTTTTGTTATACCTGTTTTAATTGGAGGTTTTGGAAATTGACTAGT TCCTCTTATAATTAGATCACCAGATATAGCTTTTCCTCGTATTAATAATGTAAGATTTTGAATATTAGTTGCTTCTTTTGGTTT GTTGCTTTTTAGAATATTTAGGGGTACAGGAGTAGGAGCTGGTTGAACACTTTATCCTCCGTTAACTAGGTATAATGGCCATAG CAGTCATGCTGTCGATTATGCAATTTTGTCTTTACATATTGCAGGAGCATCGTCAATTTTTAGTGCACTGAATTTTTTAACGAC GATTATTAATATACACTATTTTGGAGTACGAATAGATAAATTACCGTTGTTTGTGTGATCGATTTTTATTACTGCTCTATTGTT AGTTTTGGCTTTACCAGTACTTGCTGGAGCAATTACAATATTAATTTCTGATCGTAATTTCACTACTACATTTTTTGATCCGGC AGGGGGAGGAGATCCTGTTTTATTTCAACATTTATTTTGGTTTTTTGGGCATCCAGAAG

## Supplementary Materials

Supplementary Materials SM.01-Morphometric measurements of the neotype (Italy, IT.057) population of Milnesium alpigenum Ehrenberg, 1853.
Supplementary Materials SM.02-Morphometric measurements of type population (Germany, DE.001) of Milnesium inceptum sp. nov.
Supplementary Materials SM.03-Morphometric measurements of the Japan (JP.010) population of Milnesium inceptum sp. nov.
Supplementary Materials SM.04-Morphometric measurements of the Swiss (CH.002) population of Milnesium inceptum sp. nov.
Supplementary Materials SM.05-Morphometric measurements of the Bulgarian (BG.058) population of Milnesium inceptum sp. nov.


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