Morphology and Gene Sequence of *Levicoleps biwae* n. gen., n. sp. (Ciliophora, Prostomatida), a Proposed Endemic from the Ancient Lake Biwa, Japan

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**ABSTRACT.** *Levicoleps biwae* n. gen., n. sp. was discovered in organic mud on the shore of Lake Biwa, Japan. Its morphology and small subunit rRNA gene sequence were studied with standard methods. Further, we established a terminology for the colepid armour and selected four features for genus recognition: the number of armour tiers, the structure of the tier plates, the presence/absence of armour spines, and the number of adoral organelles (three or five). The Japanese colepid, a barrel-shaped ciliate with an average size of $75 \times 45$ µm, has six armour tiers and *hirtus*-type tier plates, but lacks armour spines, both in the environment and in laboratory culture. Thus, it is considered to represent a new genus. This rank is supported by the considerable genetic distance (7%) from the common *Coleps hirtus*. Although *L. biwae* looks quite similar to *C. hirtus* in vivo, it is very likely most closely related to *Coleps amphacanthus*, a species with conspicuous armour spines, as indicated by body size, the number of ciliary rows and, especially, the multiple caudal cilia. Lake Biwa is about four million years old and inhabited by many endemic organisms, ranging from algae to large fish. Thus, we suspect that *L. biwae* is restricted to Lake Biwa or, at least, to Asia. Based on literature data and the generic features established, we also propose the new genus *Reticoleps* for *Coleps remanei* Kahl, 1933, and resurrect the genus *Pinacocoleps* Diesing, 1865 to include *Coleps incurvus* Ehrenberg, 1833, *Coleps palcher* Spiegel, 1926, *Coleps tessalatus* Kahl, 1930 and, probably, *Baikalocoleps quadrata* Obolkina, 1995a. Nine colepid genera are diagnosed and dichotomously keyed.

**Key Words.** Biogeography, *Coleps*, endemic protists, *Pinacocoleps*, *Reticoleps*, *Levicoleps* n. gen.

CILIATES of the family Colepadae Ehrenberg, 1838 are characterized by an armour composed of complex, calcified plates (Corliss 1979). They occur in a conspicuous variety of habitats: in freshwater and the sea, in the benthos and plankton, and even in the limnetic and marine psammon (Dragesco and Dragesco-Kernéis 1991; Kahl 1930, 1935; Obolkina 1995a, b). Probably, many species wait to be discovered, especially in the poorly studied interstitial of large freshwater lakes and sandy marine shores.

Kahl (1930, 1935), the last reviser of the colepids, recognized 14 valid and two doubtful *Coleps* species. Since then, several new colepid genera and about 20 new species have been described, according to the Zoological Record, for instance, by Foissner (1983, 1984), Lepsi (1962), Noland (1937), and Vacelet (1961). Unfortunately, most descriptions are very incomplete, both in the past and present. For instance, plate details were studied mainly by Kahl (1930, 1935), and the reported lack of a circumoral kinety in *Planicoleps*, *Baikalocoleps*, *Kotinia*, *Macrocoleps*, and *Tiarinella* (Dragesco and Dragesco-Kernéis 1991; Obolkina 1995a) is very likely caused by insufficient preparations. Detailed investigations and redescriptions, including scanning electron microscopy (SEM) of individual armour plates, are available for only a few common freshwater species (Foissner 1984; Foissner, Berger, and Kohmann 1994; Foissner, Berger, and Schaumburg 1999; Huttelauch 1985, 1986, 1987; Wilbert and Schmall 1976).

The poor knowledge of colepids is unfortunate because they are, due to the complex armour plates, ideal for investigating the hotly discussed question of whether or not endemic, free-living micro-organisms exist (for a review, see Foissner 2006). At the present state of knowledge, *Planicoleps* occurs only in Lake Tanganyika (Africa), and the genera *Baikalocoleps*, *Kotinia*, *Macrocoleps*, and *Tiarinella* seem to be restricted to Lake Baikal. We investigated a third ancient freshwater lake, the 4-million-year-old Lake Biwa in Japan, and immediately recognized a special colepid in the shore mud. This new species, which also represents a new genus, is described here in great detail so that later researchers can reliably compare it with species from other biogeographic regions.

**MATERIALS AND METHODS**

**Materials.** *Levicoleps biwae* n. gen., n. sp. was discovered in a manually taken mud sample from the flat shore of Lake Biwa at the end of November, 2006. The site was very near to the Lake Biwa Museum and contained various filamentous algae and decaying water plants, especially *Nelumbo nucifera*. See Rossiter (2000) for a detailed description of the lake. *Levicoleps biwae*, which was moderately abundant above and in the mud, could not be found in two plankton samples taken with a fine-meshed net from the north and south basin of the lake. In the environmental sample, *L. biwae* fed on coccoid green algae, *Trachelomonas*, dinoflagellates, and even on rather large and highly mobile ciliates, such as *Urocentrum turbo* (Fig. 14). In the laboratory, *L. biwae* could be cultivated on *Eau de Volvic* enriched with some squashed wheat grains and a few milliliters of natural mud. Here, it engulfed heterotrophic flagellates, *Euplotes*, and even large starch grains from the squashed wheat kernels. *Levicoleps biwae* grew well in such raw cultures for some months, but then became smaller and smaller and declined, even in fresh medium. Pure cultures with some middle-sized ciliates and flagellates as a food source were not successful. When feeding on large prey, the circumoral tier and the anterior tier opened widely, and one has the impression that *L. biwae* nibbles at the prey (Fig. 14).

**Morphological methods.** Specimens from the environmental sample and the raw cultures were used for the investigations. Living cells were studied using a high-power oil immersion objective and differential interference contrast. Various silver impregnation methods and SEM of washed, air-dried cells were used to reveal the ciliary pattern, cytological details, and the calcified armour. All these methods are described in Foissner (1991). Counts and measurements on silvered specimens were performed at a magnification of 1,000X. In vivo measurements were conducted at magnifications of 100–1,000X. Illustrations of live specimens were based on free-hand sketches and micrographs; those of impregnated cells were made with a drawing device. *Coleps hirtus*, which occurred in the same sample, was studied for comparison.
Molecular methods. Single live cells of *L. biwae* were isolated under a light microscope and transferred to sterile Mili-Q water droplets 2 times to facilitate the removal of contaminants, suspended in 2 µl of sterile Mili-Q water, and placed in 0.2-ml thin-walled polymerase chain reaction (PCR) tubes. Samples were then frozen at ~20 °C until analysis.

PCR amplification. The first round of PCR amplification was carried out on single cells, using two sets of external primers, SR1 and SR12 (Nakayama et al. 1996) and the PCR protocol of Puitika et al. (2007).

Cloning, sequencing, and tree construction. PCR products were purified with Wizard® SV Gel and a PCR Clean-Up System (Promega, Madison, WI). Sequencing of three *L. biwae* specimens was performed on an ABI PRISM® 310 Genetic Analyzer (Applied Biosystems, Inc., Foster City, CA) for both DNA strands, using the primers SR1, SR3, SR6, and SR8–SR12 of Nakayama et al. (1996) and the ciliate-specific primer set CS 322 F and EU929R of Puitika et al. (2007).

The *L. biwae* sequences and reference sequences from the nucleotide sequence library (NCBI) were aligned with CLUSTAL X 1.83 (Thompson et al. 1997). Phylogenetic trees were generated using neighbour-joining (NJ), maximum parsimony (MP), and maximum likelihood (ML). Neighbour-joining analysis was conducted using the program package MEGA 4.0 (Tamura et al. 2007). Distances were estimated by the NJ method with the TrN model of substitution (Tamura and Nei 1993) and with the assumption of rate heterogeneity among sites. The γ-shaped parameters (γ) were estimated with eight categories from PUZZLE version 5.2 (Strimmer and von Haeseler 1996). The proportion of invariant sites was 2.237 for the data sets. The transition/transversion ratio of the Has-egawa, Kishino, and Yano (1985) model was estimated by maximizing the likelihood value for the NJ topology. The statistical significance of the tree branches was assessed by 1,000 bootstrap resamplings (Felsenstein 1985). Maximum parsimony analysis was conducted using default settings in MEGA (Tamura et al. 2007). Maximum likelihood analysis was conducted using version 4.0 of PAUP (Swofford 2002). The shape parameter of the γ distribution was the same as used in the NJ analysis.

Accession numbers. The accession numbers of the SSU rDNA nucleotide sequences used for the phylogenetic analysis are given in the phylogenetic tree.

Terminology. General ciliate terminology follows Corliss (1979). The colepid terminology, which is rather confused and shown in Fig. 1–3, is based on Kahl (1930) and Huttenlauch and Barden (1987). Of particular importance is the number of armour tiers: frequently, only those that can be easily seen are counted, while the circumoral and caudal tiers are neglected. Thus, tier number is often incorrect in the literature.

RESULTS

*Levicoleps biwae* n. gen., n. sp. (Table 1 and Fig. 4–16, 18–20, 22–24, 26–47, 49, 51)

Morphological description. The size in vivo is 60–90 × 30–50 µm, usually about 75 × 45 µm in the environmental sample, while about 60 × 40 µm after prolonged laboratory cultivation. Body width increases after silver nitrate impregnation by about 15%, so that the length:width ratio is 1.5 in preparations and 1.8 in vivo (Table 1). It is barrel shaped and inconspicuously asymmetrical, occasionally slightly narrowed in mid-body where the main armour plates abut. The anterior end is transversely truncate and crown-like due to the acute ends of the secondary tier plates, while the posterior end is moderately broadly rounded. Poorly nourished specimens are flattened laterally by up to 30%, while well-fed individuals are more or less distinctly ellipsoidal (Fig. 4, 5, 7, 8, 15, 16, 18, 19, 22–24). The macronucleus is in or near mid-body close...
Table 1. Morphometric data on *Levicoleps biwae* n. gen., n. sp.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>$\bar{X}$</th>
<th>M</th>
<th>SD</th>
<th>SE</th>
<th>CV</th>
<th>Minimum</th>
<th>Maximum</th>
<th>n</th>
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<tbody>
<tr>
<td>Body, length, $\mu$m</td>
<td>72.4</td>
<td>71.0</td>
<td>5.3</td>
<td>1.2</td>
<td>7.3</td>
<td>65.0</td>
<td>81.0</td>
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<tr>
<td>Body, width, $\mu$m</td>
<td>49.8</td>
<td>49.0</td>
<td>4.9</td>
<td>1.1</td>
<td>10.2</td>
<td>42.0</td>
<td>60.0</td>
<td>21</td>
</tr>
<tr>
<td>Body, length:width ratio</td>
<td>1.5</td>
<td>1.5</td>
<td>0.1</td>
<td>0.1</td>
<td>5.2</td>
<td>1.3</td>
<td>1.6</td>
<td>21</td>
</tr>
<tr>
<td>Body, length:width ratio in vivo</td>
<td>1.8</td>
<td>1.8</td>
<td>0.2</td>
<td>0.1</td>
<td>9.3</td>
<td>1.6</td>
<td>2.2</td>
<td>16</td>
</tr>
<tr>
<td>Anterior body end to macronucleus, distance, $\mu$m</td>
<td>36.4</td>
<td>36.0</td>
<td>5.7</td>
<td>1.2</td>
<td>15.5</td>
<td>24.0</td>
<td>48.0</td>
<td>21</td>
</tr>
<tr>
<td>Macronucleus, length, $\mu$m</td>
<td>12.4</td>
<td>12.0</td>
<td>1.7</td>
<td>0.4</td>
<td>13.4</td>
<td>10.0</td>
<td>15.0</td>
<td>21</td>
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<tr>
<td>Macronucleus, width, $\mu$m</td>
<td>9.7</td>
<td>10.0</td>
<td>1.3</td>
<td>0.3</td>
<td>13.1</td>
<td>8.0</td>
<td>13.0</td>
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<td>1.0</td>
<td>0.1</td>
<td>0.1</td>
<td>5.2</td>
<td>1.3</td>
<td>1.6</td>
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<td>Macronucleus, number (protargol impregnation)</td>
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<td>1.0</td>
<td>0.1</td>
<td>0.1</td>
<td>5.2</td>
<td>1.3</td>
<td>1.6</td>
<td>21</td>
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<tr>
<td>Circumoral kinety, diameter$^a$, $\mu$m</td>
<td>5.9</td>
<td>6.0</td>
<td>1.1</td>
<td>0.3</td>
<td>18.7</td>
<td>4.5</td>
<td>8.0</td>
<td>13</td>
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<td>Internal oral basket, length, $\mu$m</td>
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<td>8.0</td>
<td>0.9</td>
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<tr>
<td>Anterior main plate, width from ridge to ridge, $\mu$m</td>
<td>5.0</td>
<td>5.0</td>
<td>0.4</td>
<td>0.1</td>
<td>7.8</td>
<td>4.0</td>
<td>6.0</td>
<td>21</td>
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<tr>
<td>Anterior main plate, number of windows$^b$</td>
<td>7.1</td>
<td>7.0</td>
<td>1.2</td>
<td>0.3</td>
<td>17.7</td>
<td>5.0</td>
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<td>21</td>
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<tr>
<td>Anterior secondary plate, number of windows$^c$</td>
<td>2.5</td>
<td>3.0</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>2.0</td>
<td>3.0</td>
<td>21</td>
</tr>
<tr>
<td>Posterior main plate, number of windows$^c$</td>
<td>5.9</td>
<td>6.0</td>
<td>0.9</td>
<td>0.2</td>
<td>15.1</td>
<td>5.0</td>
<td>7.0</td>
<td>21</td>
</tr>
<tr>
<td>Posterior secondary plate, number of windows$^c$</td>
<td>2.2</td>
<td>2.0</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>2.0</td>
<td>3.0</td>
<td>21</td>
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<tr>
<td>Somatic kineties, number</td>
<td>24.3</td>
<td>25.0</td>
<td>1.7</td>
<td>0.4</td>
<td>6.9</td>
<td>20.0</td>
<td>27.0</td>
<td>21</td>
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<tr>
<td>Caudal cilia, number</td>
<td>7.6</td>
<td>7.0</td>
<td>0.7</td>
<td>0.2</td>
<td>11.4</td>
<td>6.0</td>
<td>9.0</td>
<td>21</td>
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</tbody>
</table>

$^a$Data based, if not mentioned otherwise, on mounted, silver nitrate-impregnated, randomly selected specimens from the environmental sample.

$^b$From specimens with closed mouth.

$^c$Identical with number of basal bodies.

Pals to the cell’s periphery and is globular to broadly ellipsoidal with an average length:width ratio of 1.3; it is about 15 $\mu$m across in vivo. The discoidal micronucleus, about 3 $\times$ 1 $\mu$m in vivo, is attached to the macronucleus (Fig. 4, 5, 29, 31). The contractile vacuole is ventrally, in line with the adoral organelles, and is distinctly subterminal with a conspicuous, posteriorly directed canal about 5 $\mu$m long, whose opening is recognizable both in silver and SEM preparations (Fig. 4, 5, 11, 13, 24, 26–28, 40, 41, 43). The cytoproct, which occurs posterior to the contractile vacuole pore, appears as a granular, thick silverline (Fig. 11, 24). Exusomesomes, which stain darkly with silver carbonate and protargol, are mainly in the oral opening, but are difficult to recognize in vivo because they are only 4 $\times$ 0.3 $\mu$m in size (Fig. 6, 12, 32, 34). The cortex is conspicuous because it is about 3 $\mu$m thick due to the armour plates, which are studded with small but distinct convexities caused by the sieve domes (Fig. 4, 10, 18, 19, 22, 23). The exusomesomes are colourless, usually containing some food vacuoles up to 20 $\mu$m wide and highly refractile lipid droplets 1–5 $\mu$m thick due to the armour (Fig. 3–5, 10, 18, 19, 22, 23). The cytoplasm is colourless, usually containing some food vacuoles up to 20 $\mu$m wide and highly refractile lipid droplets 1–5 $\mu$m thick due to the armour (Fig. 3–5, 10, 18, 19, 22, 23). The ciliate outlet in the armour plates is hemispherical and with the SEM (Fig. 3–5, 15, 16, 20, 44–47).

The somatic cilia are about 9 $\mu$m long in vivo and are very regularly arranged, forming an average of 18 transverse circles and 24 longitudinal rows (Table 1); small irregularities occur frequently. Each cilior row commences with two dikinetids, forming the perioral cilature; the anterior cilium of the perioral dikinetids is distinctly shortened. The kinetics are slightly shortened and their perioral portion is curved rightwards underneath the adoral organelles (Fig. 4, 5, 11, 24, 29, 30, 35, 42). Six to nine, but usually seven, caudal cilia about 15 $\mu$m long are arranged in a circle, which is more or less open to the right side of the cell (Fig. 4, 5, 13, 27, 28, 30, 31, 42). The somatic kineties are composed of an anterior, ciliated basal body (large granule in silver preparations) and a posterior, slightly obliquely oriented parasomal sac (small granule in silver preparations). The kinetodesmal fibre is short. The cilial outlet in the armour plates is hemispherical and broadly fusiform when seen obliquely (Fig. 5, 6, 24, 30, 32, 44–47).

The silverline pattern is of the “stria” type: the silverlines connect basal bodies longitudinally and converge on the posterior pole, forming a wide, rather irregular reticulum. The ends of the main tiers are marked by a thick, granular silverline (Fig. 11, 13, 24, 26–28).

The oral opening occupies the central half of the anterior pole. In vivo it is about 8 $\mu$m across, but may open widely when engulfing large prey (Fig. 4–9, 11, 14, 15, 19, 24, 32–37, 49). The circumoral kinety is circular and slightly disordered, but not interrupted at the site of the adoral organelles (Fig. 15, 20, 44, 49). The anterior secondary plates have an acute anterior end, thus forming a beautiful crown (Fig. 15, 20, 49). The anterior main tier, the posterior main tier, and the posterior secondary tier are without peculiarities. The caudal tier is recognizable only in oblique or posterior polar view, and the arrangement, shape, and fine structure of the caudal plates is slightly different in all specimens analysed. They may even form a stopper-like accumulation (Fig. 38). Rarely, and only in the largest specimens, are there some posterior spines up to 2 $\mu$m long (Fig. 8, 38–44). The fine structure of the armour plates is as shown in Fig. 2: the plate ridge, which is at the left margin, is smooth to slightly serrate, but never wing-like; the midbar, which separates the windows of a pair, is usually difficult to recognize both in vivo and with the SEM (Fig. 3–5, 15, 16, 20, 44–47).
made of bundles of nematodesmata originating from the circumoral dikinetids and extending to the second third of the body (Fig. 4, 6, 8, 10). Three minute, obliquely arranged adoral organelles lie in a small concavity posterior to the circumoral kinety; organelles 1 and 2 are each composed of two (di?) kinetids, while organelle 3 is composed of three (di?) kinetids. All adoral organelles are associated with about 7-μm-long fibres that contribute to the external oral basket (Fig. 5, 9, 11, 20, 24, 33–37, 49).

Gene sequence. The 18SSU rDNA sequence of *L. biwae* is 1,749 bp long and available under Accession number AB354737 of the DDBJ database. Comparing the *L. biwae* sequence with other ciliate SSU rDNA sequences identifies *Coleps* spp. as the closest relatives in all analyses. We present the NJ tree with bootstrap support values for the MP and ML trees (Fig. 5). The three specimens analysed have identical sequences. The base substitutions between *L. biwae* and *C. hirtus* is comparatively large, viz., 122 bp (~7%). Interestingly, *C. hirtus* seems to be a mixture of

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**Fig. 4–13.** *Leviceps biwae* n. gen., n. sp. from life (4, 7, 8, 10), after protargol impregnation (5, 6, 9, 12), and after silver nitrate impregnation (11, 13). 4. Broad side view of a representative specimen, length 75 μm. 5, 6. Ciliary pattern of ventral and dorsal side of holotype specimen, which has the mouth opened. 7. Schematic view at low magnification, showing the main armour tiers and the crenulate surface. 8. A 90-μm-long specimen with a minute spine (arrow). 9. Oral ciliary pattern of a squashed specimen; the outer oral basket is not shown; arrowheads mark the minute adoral organelles. 10. Optical section showing the cortex which is about 31 μm thick due to the endoskeletal, calcified armour plates. 11. Ventral view of holotype specimen, showing the silverline and ciliary pattern as well as the contractile vacuole apparatus. 12. Oral basket of a specimen with the mouth opened. 13. Posterior polar view, showing the arrangement of the basal bodies and silverlines. AMT, anterior main tier; AO, adoral organelles; AST, anterior secondary tier; BB, basal body; C, cilium; CA, canal of contractile vacuole; CC, caudal cilia; CK, circumoral kinety; CV, contractile vacuole; CY, cytopyge; E, extrusomes; EX, excretory pore of contractile vacuole; FV, food vacuole; IB, inner oral basket; MI, micronucleus; OB, outer oral basket; P, cortex with armour plates; PC, perioral ciliature; PMT, posterior main tier; PS, parasomal sac; PST, posterior secondary tier; SL, silverlines; W, windows. Scale bars: 30 μm (Fig. 4, 11), 20 μm (Fig. 5, 6), 10 μm (Fig. 9, 12, 13).
at least two species. Possibly, the U97109 population, which has been submitted by Hirt, Dyal, Embley, Esteban, and Finlay (unpubl.), represents the "green", symbiotic algae containing C. hirtus viridis Ehrenberg, 1831 (see Foissner et al. 1999 for a detailed redescription) because it is likely from Priest Pot, where Esteban and Finlay have been working and where a green C. hirtus is abundant in the hypolimnion (Guhl, Finlay, and Schink 2001).

**DISCUSSION**

**Family Colepidae.** The Colepidae Ehrenberg, 1838 include ciliates with a calcified armour and a prostome ciliary organization (Corliss 1979; Foissner 1984; Foissner et al. 1994, 1999; Huttenlauch 1986, 1987; Huttenlauch and Bardele 1987; Kahl 1930; Small and Lynn 1985; Wilbert and Schmall 1976). Although details vary considerably, the basic organization is the same in all genera and species (Fig. 1). Thus, the family is well circumscribed, except for the genus Plagiopogon, which Kahl (1930) and Corliss (1979) assigned to the haptorids while Small and Lynn (1985) classified it in the Colepidae. Unfortunately, details of the scales and the ciliary pattern of Plagiopogon are not known, and thus both classifications remain doubtful.

Briefly, the colepid armour is composed of several plate tiers that are distinct in most but not all genera (Macrocoleps). The tiers consist of oblong, calcified plates with a complex fine structure (Fig. 2). Depending on the species, there are about 15–60 monokinetal ciliary rows that extend meridionally, rarely spirally. Each ciliary row commences with two or more dkininetids, forming the perioral ciliature (Fig. 1, 9, 32). The oral apparatus consists of the oral basket, a dkininetid circumoral kinety, and 3–5 min adoral organelles in a more or less deep cavity close to the circumoral kinety (Fig. 1, 5, 6, 9, 12, 15, 20, 35).

We recognize nine genera with a total of about 40 species. An updated revision of the species is not available.

**Recognition of genera.** Diesing (1865), who synonymized Coleps Nitzsch, 1827 with Plagiopogon Stein, 1859, founded three colepid genera, using armour details as distinguishing features: Dictyocoleps, Pinacocoleps (resurrected in the diagnostic section), and Cricocoleps (for Coleps amphiacanthus), all considered as junior synonyms of Coleps by later authors (Corliss 1979; Kahl 1930; Kent 1881). Kahl (1930) and Corliss (1979) recognized only two genera, also using armour details as the main generic features: Coleps Nitzsch, 1827 and Tiarina Bergh, 1881. They considered Stappersia Meunier, 1910 as a junior synonym of Tiarina. No new colepid genus was established between 1882 and 1985, when Small and Lynn created Nolandia, using the different orientation of the adoral organelles to distinguish Nolandia from Coleps. A few years later, Dragesco and Dragesco-Kerneis (1991) discovered a new colepid in the psammion of Lake Tanganyika and established the genus Planicoleps, using the number of armour tiers, the lack of armour spines, and the presence of special extrusomes as generic features. Finally, Obolkina (1995a) established four new genera for colepids discovered in the coastal psammion of Lake Baikal, using body shape, the number of armour tiers, the presence vs. absence of armour spines and a circumoral kinety, the number of adoral organelles, and the habitat (freshwater vs. marine) as the main generic features: Baikalocoleps, Kotinia (for the homonym Alexandria, see Aescht 2001), Macrocoleps, and Tiarinella. Thus, altogether 12 generic names have been created for colepids, to which we add the new genera Levicoleps and Reticolesps (see "Diagnostic").

Based on the data reviewed above and the present investigations, we selected four features for genus recognition in the family Colepidae: the number of armour tiers, the presence vs. absence of armour spines and a circumoral kinety, the number of adoral organelles, and the habitat (freshwater vs. marine) as the main generic features: Baikalocoleps, Kotinia (for the homonym Alexandria, see Aescht 2001), Macrocoleps, and Tiarinella. Thus, altogether 12 generic names have been created for colepids, to which we add the new genera Levicoleps and Reticolesps (see "Diagnostic").

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Levicoleps biwae n. gen., n. sp. (15, 16, 18–20, 22, 23) and Coleps hirtus (17, 21) from life (18, 19, 22, 23) and in the scanning electron microscope (15–17, 20, 21). 15, 16. Lateral and dorsal view, showing the flattening of the cell and the main components of the calcified armour; note the absence of anterior and posterior spines. 17, 21. Lateral view of C. hirtus, which occurred together with L. biwae. These species are easily distinguished by the presence vs. absence of spines and the number of plate (ciliary) rows, which is significantly higher in L. biwae (cp. Fig. 15). 20. Oblique anterior polar view of a specimen with the mouth partially open. Note the central mouth opening and the site of the adoral organelles (arrow). 18, 22. Surface views at medium (250×) and low (100×) magnification. 19, 23. Optical sections showing the body shape and thick cortex (apposed triangles). AMT, anterior main tier; AS, anterior spines; AST, anterior secondary tier; AT, acute end of a plate of the anterior secondary tier; CP, circumoral plates; CT, circumoral tier; Oa, oral apparatus; OB, oral basket; PMT, posterior main tier; PST, posterior secondary tier. Scale bars: 40 mm (Fig. 18, 19, 22), 30 μm (Fig. 15–17, 23), 10 μm (Fig. 20, 21).
Coleps and Leviceps, which differ mainly by the presence vs. absence of armour spines, indicate that spines are an important character in colepids. Unfortunately, the adaptive value of the spines is not known, but they persist in laboratory cultures, at least in C. hirtus and L. biwae.

The structure and shape of the tier plates has been considered a species-specific feature by Kahl (1930) and others. However, scales are now widely considered both genus- and species-specific in many protists. Thus, we suggest to follow this trend, that is, to use the main colepid plate types for genus distinction (Fig. 3).

The fourth generic feature, the number of adoral organelles is at first glance also quantitative and thus not very strong; furthermore, the organelles are small, and their details thus difficult to recognize. However, the number of adoral organelles is not size-related, suggesting some generic significance. For instance, several large colepids (Planiroleps, Macrocoleps) have only three adoral organelles, as is typical for small species (Foissner et al. 1994, 1999), while the large Kotinia species have five (Obolkina 1995a).

Certainly, our selection of generic features is a matter of intuitive feeling rather than of solid knowledge, but it provides a testable hypothesis for further investigations, especially molecular phylogenies. Other features that might be of significance include body shape and flattening, the number of perioral dikinetids and caudal cilia, and the habitat (freshwater vs. marine). In contrast to Obolkina (1995a), we do not include the circumoral kinety because this is an ordinal character and its supposed absence in several genera is very likely the result of insufficient preparations.

Using the nomenclatural rules and the four generic features discussed above, the 14 colepid genera decrease to nine: Dictycoleps is an objective synonym of Coleps because it includes C. hirtus Nitzsch, 1827; Ericoleps and Stappersia are very probably junior synonyms of Coleps and Tiarina, respectively; Baikalocoleps is a subjective synonym of (likely) Pinacocoleps; and Tiarinella is a subjective synonym of Tiarina. As concerns Baikalocoleps and Tiarinella, there is a fair chance that their genetic distance from, respectively, Pinacocoleps and Tiarina is sufficiently high to be classified as distinct genera, similar to the case of Coleps and Leviceps. See the diagnostic section for the characterization of the other genera.

The adoral organelles and the genus Nolandia. The adoral organelles of the colepids are, with few exceptions, \( \approx 3 \mu m \) long, that is, are very small and close to the circumoral kinety. Thus, they are difficult to investigate and prone to misinterpretations. Fortunately, there are now detailed electron microscopic investigations available (Huttenlauch 1986, 1987) that give us the

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Fig. 24–28. Leviceps biwae n. gen., n. sp. (24, 26–28) and Coleps hirtus (25) after silver nitrate impregnation. All specimens are from the environmental sample, i.e. not from laboratory cultures. 24. Ventral view of holotype specimen, showing the silverline and ciliary pattern as well as the main armour tiers. 25. Coleps hirtus is easily distinguished from L. biwae by the significantly lower number of ciliary (plate) rows. 26–28. Posterior polar views showing the ciliary and silverline pattern as well as the circle of caudal cilia (triangles) and the pore of the contractile vacuole. AMT, anterior main tier; AO, adoral organelles; AST, anterior secondary tier; CAT, caudal tier; CY, cytopyge; EX, excretory pore of contractile vacuole; PC, perioral ciliature; PMT, posterior main tier; PST, posterior secondary tier; SL, silverlines. Scale bars: 40 \( \mu m \) (Fig. 24), 30 \( \mu m \) (Fig. 25), 15 \( \mu m \) (Fig. 26–28).
Fig. 29–37. *Levicoleps biwae* n. gen., n. sp., somatic and oral ciliary pattern and other organelles after silver carbonate impregnation. All specimens more or less flattened by coverslip pressure. 29, 30. Overview showing left and right side of same specimen; note the short kinetodesmal fibre at the right side of the kinetids. 31. Oblique, posterior polar view showing seven caudal cilia. 32–37. Oral area showing the perioral ciliature and the 3 min adoral organelles (asterisks) from various perspectives; the first and second adoral organelle each consist of two (di?) kinetids, while organelle 3 is composed of three (di?) kinetids; note that the circumoral kinety is not interrupted at the site of the adoral organelles. The arrows (33, 35) denote the two perioral dikinetids underneath adoral organelle 1 and the triangle (34) marks the fibres forming the inner oral basket. AO, adoral organelles; CC, caudal cilia; CK, circumoral kinety; E, extrusomes; IB, inner oral basket; KD, kinetodesmal fibres; MA, macronucleus; MI, micronucleus; N, nematodesmata forming the outer oral basket; OA, oral apparatus; PC, perioral dikinetids. Scale bars: 30 μm (Fig. 29, 30), 20 μm (Fig. 31), 10 μm (Fig. 32–37).
Levicoleps biwae n. gen., n. sp., posterior body region in the scanning electron microscope. All specimens seen have an individually distinct plate and ridge pattern in the caudal tier; a few specimens have minute spines (≤ 2 μm) associated with some caudal plates (40, 43, arrows). The plate ridge may be smooth (38, 41, 42) or slightly serrate (39, 40). Triangles (42) mark openings for the caudal cilia. CAP, caudal plates; CAT, caudal tier; EX, plate opening for the contractile vacuole; PMT, posterior main tier; PST, posterior secondary tier; R, plate ridge. Scale bars: 10 μm (Fig. 38–42), 5 μm (Fig. 43).
Fig. 44–49. *Levicolesps biwae* n. gen., n. sp. (44–47, 49) and *Coleps hirtus* (48) from life (44) and in the scanning electron microscope (45–49). 44. A squashed specimen showing the six armour tiers and individual plates in interference contrast; triangles denote sites where the typical, pretzel-shaped structure of the windows can be seen. 45–47. Plates at high magnification; arrows (46) mark the inconspicuous midbar. 48. The armour of *C. hirtus* has the same fine structure as that of *L. biwae*. 49. Oral area showing the pharyngeal opening and the circumoral tier, which has a slight irregularity at the site of the adoral organelles. AMT, anterior main tier; AO, adoral organelles; AST, anterior secondary tier; CAT, caudal tier; CO, ciliary outlet; CT, circumoral tier; D, sieve dome; PMT, posterior main tier; PR, plate processes; PST, posterior secondary tier; R, plate ridges; W, plate windows. Scale bars: 30 μm (Fig. 44), 10 μm (Fig. 45, 48), 5 μm (Fig. 46, 47, 49).
possibility to reinterpret earlier light microscopic descriptions. Such data show that the arrangement and structure of the adoral organelles is very similar in all colepids. The organelles are in a minute depression close to the circumoral kinety and obliquely arranged with respect to the meridional somatic ciliary rows, just as are the adoral membranelles of *Tetrahymena* (Fig. 1). Usually the individual organelles consist of 10 dikinetids forming short rows that slightly increase in length from anterior (organelle 1) to posterior (organelle 3). Thus, each organelle is composed of two rows of basal bodies of which those of the left row bear 2-μm-long cilia while those of the right row are barren. This might explain why the adoral organelles may appear to be composed of mono- or dikinetids in the light microscope, depending on the silver method used and the interpretation of the observer (Fig. 1, 9, 33–37).

Huttenlauch (1986, 1987) showed that the circumoral kinety is not interrupted by the adoral organelles, which matches the present data (Fig. 9, 33–37), but disagrees with previous light microscopic observations (Foissner 1984; Wilbert and Schmall 1976);
Table 2. Comparison of well-known colepids.

<table>
<thead>
<tr>
<th>Species</th>
<th>Methods*</th>
<th>Number of specimens studied</th>
<th>Length: width ratio</th>
<th>Number of tiers</th>
<th>Number of ciliary rows (extremes and median)</th>
<th>Number of caudal cilia</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coleps hirtus hirtus</td>
<td>CHL</td>
<td>15</td>
<td>1.9</td>
<td>6</td>
<td>15–16; 15</td>
<td>1</td>
<td>Foissner (1984)</td>
</tr>
<tr>
<td>Coleps hirtus viridii</td>
<td>IV, SC</td>
<td>≤ 10</td>
<td>1.8</td>
<td>6</td>
<td>14–19; 15</td>
<td>1</td>
<td>Foissner et al. (1999)</td>
</tr>
<tr>
<td>Coleps spetai</td>
<td>P</td>
<td>12</td>
<td>1.6</td>
<td>6</td>
<td>16–18; 17</td>
<td>1</td>
<td>Foissner (1984)</td>
</tr>
<tr>
<td>Coleps elongatus</td>
<td>IV, SC</td>
<td>17</td>
<td>2.3</td>
<td>6</td>
<td>16–18; 16</td>
<td>2</td>
<td>Foissner et al. (1999)</td>
</tr>
<tr>
<td>Coleps amphacanthus</td>
<td>P</td>
<td>10</td>
<td>1.8</td>
<td>6</td>
<td>22–28; 27</td>
<td>~ 10</td>
<td>Foissner and O’Donoghue (1990)</td>
</tr>
<tr>
<td>Levicoleps biwae</td>
<td>CHL</td>
<td>21</td>
<td>1.5</td>
<td>6</td>
<td>20–27; 25</td>
<td>7</td>
<td>Present paper</td>
</tr>
<tr>
<td>Nolandia nolandi</td>
<td>CHL, P</td>
<td>≤ 10</td>
<td>2.5</td>
<td>6</td>
<td>12–17; 13</td>
<td>1</td>
<td>Wilbert and Small (1976), Foissner et al. (1999)</td>
</tr>
<tr>
<td>Planicoleps psammophilus</td>
<td>P</td>
<td>≥ 15</td>
<td>2.2</td>
<td>8</td>
<td>32–38; 36</td>
<td>0</td>
<td>Dragesco and Dragesco-Kernéis (1991)</td>
</tr>
</tbody>
</table>

*aCHL, Chatton-Lwoff silver nitrate impregnation, as modified by Corliss (1953); IV, in vivo; P, protargol impregnation; SC, silver carbonate impregnation.

bInclude some unpublished data.

Obviously, the circumoral dkinetids neighbouring the adoral organelles were interpreted as belonging to the latter.

Most of these data were not known when Small and Lynn (1985) established the genus Nolandia: ‘’genus differs from Coleps with brosse files (= adoral organelles) parallel, rather than perpendicular, to body kinetics’’. Small and Lynn (1985) concluded this from a reinvestigation of Coleps nolandi by Wilbert and Schmall (1976) and original data from protargol-impregnated C. elongatus. However, the figure they provided is rather schematic and the observations were not confirmed in a more detailed study which shows, by clear micrographs, an oblique arrangement of the organelles in C. elongatus, highly similar to that described in C. hirtus and C. nolandi (Foissner et al. 1999; Wilbert and Schmall 1976). Thus, Nolandia is obviously based on a misobservation and thus should be synonymized with Coleps. On the other hand, C. nolandi, type and sole species of the genus Nolandia, has unique tier plates (Foissner et al. 1994; Huttenlauch 1985; Kuhl 1930), requiring generic separation. The International Code of Zoological Nomenclature does not rule on such ‘’taxonomic’’ problems, but requires that the name and its author(s) are conserved if the genus has been typified correctly. Thus, an ‘’nomic’’ problems, but requires that the name and its author(s) are conserved if the genus has been typified correctly. Thus, an

Levicoleps as a new genus. As discussed above, we base Levicoleps n. gen. on the absence of armour spines. Although this feature tends to vary in general, it is apparently very stable in colepids because few spineless populations have been described. More importantly, no spines developed in cultures held for up to 6 months. Further, there is 7% genetic divergence to C. hirtus and C. spetai, in spite of the identical fine structure of the armour plates, showing that L. biwae is a rather different species (Fig. 58).

Indeed N. nolandi, which has different armour plates, clusters in between the C. hirtus group and L. biwae, supporting our classification as a distinct genus. Further molecular studies are required to clarify whether the considerable genetic distance of Levicoleps is due to ancient isolation (endemism) or due to a rather different ancestor. Unfortunately, the sequence of the supposed closest relative, C. amphacanthus, is not known (Table 2).

Comparison of L. biwae with related species. Five colepids without armour spines have been described (Fig. 50–57): Coleps inermis Perty, 1852, a doubtful species, which could be a haptorid ciliate; C. striatus Smith, 1897; C. kenti Bhatia, 1936; C. trichotus Savi, 1913, which possibly represents a distinct genus; and Planicoleps psammophilus Dragesco and Dragesco-Kernéis, 1991. Of these, only the first three species resemble L. biwae, while C. trichotus is conspicuously fusiform (Fig. 57) and P. psammophilus is much larger (Table 2). The three species resembling L. biwae are smaller than 55 μm (vs. 75 μm) and have, according to the figures, fewer than 20 ciliary rows (vs. 25). Thus, they are more closely affiliated to C. hirtus than to L. biwae, which resembles a spineless C. amphacanthus in these respects (Table 2).

The armour and armour plates of L. biwae are virtually identical to those of C. hirtus and C. amphacanthus (Foissner et al. 1999; Huttenlauch 1986), except for the spines, which are lacking in L. biwae (cp. Fig. 1, 15, 38–45, 49 with Fig. 17, 21, 48). Spineless ‘’C. hirtus’’ have been found in England (Kent 1881), Switzerland (Perty 1852), the United States (Smith 1897), and in India (Bhatia 1936). Whether these populations belong to the same or different species is not known because the descriptions are too incomplete. However, these descriptions show the existence of one or more spineless Levicoleps in these regions, resembling C. hirtus in size and number of ciliary rows.

Biogeographic aspects. Ancient lakes are those with an uninterrupted history that dates back more than 100,000 yr (GORTHNER 1994). They are famous for their high biodiversity, including many endemic species. For instance, about 50% of the species known from Lake Baikal (Russia) and Lake Tanganyika (Africa) are endemic (MARTENS 1997). Such numbers mainly refer to multicellular organisms because protists are poorly explored in ancient lakes. Some endemic amoebae and ciliates have been reported: for instance, Litilimorpha viridis from Lake Baikal (OBOLKINA 1995b) and Diffugia biwae from Lake Biwa (MORI AND MIURA 1980; NISHINO AND WATANABE 2000).

Ancient lakes could decide the hotly discussed problem of whether or not endemic protists exist (for a review, see FOISSNER 2006). The data available support restricted distribution, for instance, the many endemic diatoms in Lake Tanganyika (COQUY 2000), a considerable number of endemic ciliates in Lake Baikal (OBOLKINA 1995b), and some endemic algae in Lake Biwa (MORI AND MIURA 1980; NISHINO AND WATANABE 2000).

Colepids with their complex armour plates are biogeographic flagspar excellence. Indeed, several new coelid genera and species were discovered in the psammon of Lake Baikal (OBOLKINA 1995a) and Lake Tanganyika (DRAGESCO AND DRAGESCO-KERNÉIS 1991). Now, a further new coelid has been discovered in Lake Biwa (Fig. 4). Are all these discoveries only a fortunate chance? We do not believe this but interpret it as an indication for restricted distribution. Otherwise they would not have been found in the few samples taken. Whether they are local, regional, or
continental endemics requires further investigation. Probably they are rather widely distributed, like *D. biwae*, formerly considered as a local endemic of Lake Biwa (Mori and Miura 1980), but which has been reported recently from three lakes in central China (Yang and Shen 2005). In Europe, North America, and India obviously occurs a different *Levicoleps*, which is closely similar to the common *C. hirtus* (see ‘‘Comparison of *L. biwae* with related species’’), while *L. biwae* resembles a spineless *C. amphicanthus*. Indeed, we found such a species recently in Germany. It highly resembles *C. hirtus*, except for the lack of distinct spines.

**DIAGNOSIS**

In the following compilation, mainly those species which have been carefully described or reinvestigated are assigned to a certain genus. All others remain in *Coleps* for now.

**Coleps** Nitzsch 1827

**Emended diagnosis.** Colepidae with spiny armour composed of six tiers with plates of the *hirtus* type (Fig. 3). Three adoral organelles.


**Etymology.** Unfortunately, Nitzsch (1827) did not provide the etymology of *Coleps*. At first glance, it seems to be a Greek noun with the meaning ‘‘bend of the knee’’. Obviously, this is senseless, which is supported by the gender: the Greek *Coleps* is feminine, while all authors have treated the *Coleps* of Nitzsch as masculine. Possibly, *Coleps* is derived from the Latin neuter noun ‘‘colum’’ (sieve); indeed, *Coleps* looks like a sieve at moderate magnification!). Nitzsch (1827) may have added ‘‘eps’’ as an artificial combination of letters. As long as the etymology has not been clarified, we suggest treating *Coleps* as masculine.

**Species assignable.** *Coleps amphicanthus* Ehrenberg, 1833 (redescribed by Huttenlauch 1986, 1987; Huttenlauch and Bardele 1987; Foissner and O’Donoghue 1990; Kreutz and Foissner 2006); *C. elongatus* Ehrenberg, 1831 (redescribed by Foissner et al. 1999; Small and Lynn 1985); *C. hirtus hirtus* (O. F. Müller 1786) Nitzsch, 1827 (redescribed by Foissner 1984; Foissner et al. 1999); *C. hirtus viridis* Ehrenberg, 1831 (redescribed by Foissner et al. 1999); *C. spetali* Foissner, 1984 (reviewed in Foissner et al. 1999).

**Kotinia** Obolkina in Aescht, 2001

**Emended diagnosis.** Colepidae with spiny armour composed of eight tiers with not yet specified plates. Five adoral organelles.


**Etymology.** Named after Bolshie Koty, the locus classicus in southern Baikal. Feminine gender.


**Remarks.** Originally, this genus was named *Alexandria* which is, however, a homonym; for details, see Aescht (2001). All species need redescription, especially of the fine structure of the armour plates.

**Levicoleps n. gen.**

**Diagnosis.** Colepidae with smooth armour composed of six tiers with plates of the *hirtus* type (Fig. 3). Three adoral organelles.

**Type species.** *Levicoleps biwae* n. sp.

**Etymology.** Composite of the Latin adjective ‘‘levis’’ (smooth) and the generic name *Coleps*. Masculine gender.

**Species assignable.** So far only the type species, *L. biwae*, can be assigned to the new genus (but see Discussion).

**Levicoleps biwae** n. sp.

**Diagnosis.** Size in vivo about 75 x 45 μm, barrel shaped. On average 24 ciliary rows, each composed of about 18 monokinetics and two perioral dikinetids; six to nine caudal cilia. Anterior and posterior main plates each with an average of seven and six windows, respectively; anterior and posterior secondary plates each with two to three windows.

**Type locality.** Japan, shore of Lake Biwa near the Lake Biwa Museum, Karasuma Peninsula, Shiga Prefecture 35°04’20.64”N 135°56’21.74”E.

**Type material.** Two holotype slides (hapantotypes; one protargol impregnated, the other silver nitrate impregnated) have been deposited in the Oberösterreichisches Landesmuseum in Linz (L). The para type slides have been deposited in the Lake Biwa Museum.

**Gene sequence.** Accession number of the SSU rDNA of *L. biwae* is AB354737 in the DDBJ database.

**Etymology.** Named after the site found; ‘‘biwae’’ is a noun in genitive case.

**Macrocoleps** Obolkina, 1995a

**Emended diagnosis.** Colepidae with spiny armour composed of about 12 irregular tiers with not yet specified plates. Three adoral organelles.

**Type species** (by original designation). *Macrocoleps caudatus* Obolkina, 1995a.

**Etymology.** Not given in the original description. Obviously, it is composite of the Greek adjective ‘‘macro’’ (large) and the generic name *Coleps*. Masculine gender.

**Species assignable.** *Macrocoleps caudatus* Obolkina, 1995a; *Macrocoleps aculeatus* Obolkina, 1995a.

**Remarks.** Both species need redescription, especially of the fine structure of the armour plates.

**Nolandia** Small and Lynn, 1985

**Emended diagnosis.** Colepidae with spiny armour composed of six tiers with plates of the *nolandi* type (Fig. 3). Three adoral organelles.


**Etymology.** Named in honour of Lowell Evan Noland (1896 – 1972), who provided a revision of the genus *Coleps* in 1925. Feminine gender.

**Species assignable.** So far only the type species can be assigned to this genus. *Nolandia nolandi* was redescribed by Foissner et al. (1994), Huttenlauch (1985), and Wilbert and Schmal (1976).

**Remarks.** For taxonomy, see Discussion.

**Pinacoleps** Diesing, 1865

**Emended diagnosis.** Colepidae with spiny armour composed of six tiers with plates of the *incurvus* type (Fig. 3). Number of adoral organelles not known.


**Etymology.** Not given in the original description. Possibly, it is a composite of the Greek ‘‘pinia’’ (tile) and of the generic name *Coleps*, referring to the appearance of the cell’s armour. Masculine gender.
**Species assignable.** The incurvus plate type occurs in a freshwater and two marine species (Kahl 1930), which are thus combined with Diesing’s genus: Pinacocoleps incurvus (Ehrenberg 1833) n. comb., basionym: Coleps incurvus Ehrenberg, 1833; P. tessalatus (Kahl 1930) n. comb., basionym: Coleps tessalatus Kahl, 1930; P. pulcher (Spiegel 1926) n. comb., basionym: Coleps pulcher Spiegel, 1926. None of these species has been investigated with modern methods. Possibly, each represents a distinct genus or subgenus because plate details look fairly different (Fig. 3). Likely, B. quadratus Obolkina, 1995a also belongs to Pinacocoleps. Obolkina (1995a) defined this genus mainly by the lack of a circumoral kinety; all other features match Coleps or Pinacocoleps. Unfortunately, plate details of B. quadratus were described too incompletely to be sure about the generic assignment. Thus, and because the species might be genetically highly distinct from other colepids—considering the special habitat, we neither formally synonymize the genus nor combine the species with another genus.

**Remarks.** We resurrect, with improved diagnosis, this almost forgotten genus created by Diesing in 1865. Interestingly, he already used organ/padetail details to define the genus.

**Planicoleps** Dragesco and Dragesco-Kerneis, 1991

**Emended diagnosis.** Colepids with smooth armour composed of eight tiers with not yet specified plates. Three adoral organelles.

**Type species** (by original designation). Planicoleps psammophilus Dragoesco and Dragoesco-Kerneis, 1991.

**Etymology.** Not given in the original description. Very likely, the name refers to the strong flattening (plane) of the type species. Masculine gender.

**Species assignable.** So far only the type species can be assigned to this genus (Fig. 56).

**Remarks.** Planicoleps differs from Levicoleps in having eight instead of six plate tiers and in adoral organelle 3, which is an irregular field of basal bodies in Planicoleps. Possibly, there are also differences in the fine structure of the plates which, unfortunately, Dragesco and Dragesco-Kerneis (1991) described rather superficially.

**Reticoleps n. gen.**

**Diagnosis.** Colepids with spiny armour composed of six tiers with plates of the remanei type (Fig. 3). Number of adoral organelles not known.

**Type species.** Reticoleps remanei (Kahl 1933) n. comb. Basionym: Coleps remanei Kahl, 1933.

**Etymology.** Composite of the Latin noun ‘‘reticulum’’ (fine net) and the generic name Coleps. Masculine gender.

**Species assignable.** So far only the type species can be assigned to the new genus. Other marine and psammophilic colepids do not have a reticular plate fine structure (Fig. 3).

**Remarks.** Like Nolandia, this new genus is based on the unique structure of the armour plates, as explained in the section “Recognition of genera”.

**Tiarina** Bergh, 1881

**Emended diagnosis.** Fusiform Colepids with spiny or smooth armour composed of six tiers with plates of the tiarina type (Fig. 3). Three adoral organelles.

**Type species** (by monotypy). Tiarina fusus (Claparede and Lachmann 1859) Bergh, 1881. Basionym: Coleps fusus Claparède and Lachmann, 1859.

**Etymology.** Not given in the original description. Obviously, the name is derived from the Latin tiarás (tiara = crown of oriental kings), referring to the complex fine structure of the armour. Feminine gender.

**Species assignable and remarks.** Possibly only the type species belongs to that genus because T. meunieri Kahl, 1930 has a smooth, rather different armour, suggesting resurrection of the genus Stappesia. The type species, briefly investigated by Small and Lynn (1985), has a colepid organization with, however, rather conspicuous adoral organelles, while Tiarinella gracilis Obolkina, 1995a has minute adoral organelles and conspicuous oral spines, suggesting that it could represent a distinct genus (see also “Discussion”). Unfortunately, the fine structure of the armour plates was insufficiently described (“like fuzzy squares”). Considering these problems, we do not formally synonymize Tiarinella with Tiarina.

**KEY TO THE COLEPID GENERA**

The key uses simple features recognizable without silver impregnation. Nonetheless, silver impregnation should be applied because not yet discovered genera might differ in features recognizable only in silver preparations.

1. Armour tiers regular, i.e. forming rings . . . 2
   – Armour tiers irregular, i.e. not forming rings . . . Macrocoleps

2. Six armour tiers . . . 4
   – Eight armour tiers . . . 3

3. Armour with spines . . .
   – Armour without spines . . .
   – Body fusiform . . . 4
   – Body barrel- or pillow-shaped . . .
   – Armour without spines . . .
   – Armour with spines . . . 3

   – Plate ridge in or left of plate midline . . .
   – Plate ridge at left margin, windows of hiria type . . . 7
   – Plate ridge at left margin, windows of hiria type . . . 8
   – Plate window of nolandi type (Fig. 3) . . .
   – Plate windows of incurvus type (Fig. 3) . . .
   – Plate windows of incurvus type (Fig. 3) . . .

5. Armour without spines . . .
   – Tier plates different . . .
   – Tier plates different . . .

6. Tier plates of remanei type, i.e. with reticular fine structure (Fig. 3) . . .
   – Tier plates different . . .

7. Plate ridge at left margin, windows of hiria type . . .

8. Plate window of nolandi type (Fig. 3) . . .

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