Systematics of the Neotropical caddisfly genus
Notidobiella Schmid (Trichoptera, Sericostomatidae),
with the description of 3 new species

Ralph W. Holzenthal†, Roger J. Blahnik‡

Department of Entomology, University of Minnesota, 1980 Folwell Ave. 219 Hodson Hall, St. Paul, Minnesota, 55108, U.S.A.

† urn:lsid:zoobank.org:author:C27A1C49-F372-4232-A49C-B8B5F753CC70
‡ urn:lsid:zoobank.org:author:8E40859B-B83A-4C45-9B03-B3F56E861DE1

Corresponding author: Ralph W. Holzenthal (holze001@umn.edu)

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Abstract
Three new species of Notidobiella Schmid (Insecta: Trichoptera) are described from South America: Notidobiella amazoniana sp. n. (Brazil), N. brasiliana sp. n. (Brazil), and N. ecuadorensis sp. n. (Ecuador). In addition, the 3 previously described species in the genus, N. chacayana Schmid, N. inermis Flint, and N. parallelipipeda Schmid, all endemic to southern Chile, are redescribed and illustrated, including the females of each species for the first time, and a key to males of the species in the genus is provided. The occurrence of Notidobiella in Brazil and Ecuador represents a significant extension of the range of the genus beyond southern Chile where it previously was thought to be endemic. The biogeography of Sericostomatidae and other austral South American Trichoptera is reviewed. The presence of the family in South America may not be part of a “transantarctic” exchange, but instead may represent an earlier occurrence in the region. The distribution of Notidobiella in tropical South America likely represents recent dispersal from southern South America to the north.

Keywords
caddisfly, Neotropics, transantarctic, new species, biogeography, South America, taxonomy
Introduction

The caddisfly family Sericostomatidae occurs in all biogeographic regions, except the Australasian, but its species diversity is very unevenly distributed across these regions (Morse 2010). Nineteen genera and 100 species have been described worldwide, with half of the species occurring in the western Palearctic (50 species in 5 genera: *Cerasma* McLachlan, *Notidobia* Stephens, *Oecismus* McLachlan, *Schizopelex* McLachlan, *Sericostoma* Latreille) (Holzenthal et al. 2007b). South Africa harbors 12 species in 5 endemic genera (*Aclasma* Morse, *Aselas* Barnard, *Cheimacheramus* Barnard, *Petroplax* Barnard, *Rhoizema* Barnard), but no species are found in tropical Africa. The genus *Agarodes* Banks contains 12 species confined largely to the southeastern United States, where a second monotypic genus, *Fattigia* Ross, also occurs. There is a single species known from India, *Asahaya asambaddha* Schmid, and the 6 species in the genus *Gumaga* Tsuda occur in Mexico (Baja California) and the western United States (3 species) and in southern and eastern Asia (3 species). In the Neotropics, the family is represented in the Chilean subregion by 4 genera, *Chiloecia* Navás (1 species, *nomen dubium*), *Myotrichia* Schmid (1), *Notidobiella* Schmid (3), and *Parasericostoma* Schmid (10) (Flint et al. 1999b). Until now, only a single sericostomatid species, *Grumicha grumicha* (Vallot), was known from the Brazilian subregion (Flint et al. 1999a).

In this paper, we describe 3 new species of *Notidobiella*, 1 from the Amazon basin, Brazil, 1 from southeastern Brazil, and 1 from Ecuador, thus extending the range of this genus well beyond its Chilean representation. In addition, we provide illustrations and diagnoses of males and females (the latter for the first time) of the 3 previously described species of *Notidobiella*, *N. chacayana* Schmid, *N. inermis* Flint, and *N. parallelipipeda* Schmid, and a key to males of species in the genus. The Neotropical species of Sericostomatidae, including those in the genus *Notidobiella*, appear to be members of a southern Gondwana fauna (de Moor and Ivanov 2008).

Material and methods

Techniques and procedures used in the preparation and examination of specimens are those outlined by Blahnik and Holzenthal (2004) and Blahnik et al. (2007). Terminology for genitalia and wing venation follows that presented by Holzenthal et al. (2007b) and morphological structures are labeled in Figs 3–5. The species are presented in alphabetical order, except for the type species of the genus, *N. parallelipipeda*, which is presented first, after the generic diagnosis. Material examined and types are deposited in the collections of the University of Minnesota Insect Collection, St. Paul, Minnesota, USA (UMSP), the Museu de Zoologia, Universidade de São Paulo, São Paulo, Brazil (MZUSP), the Instituto Nacional de Pesquisas da Amazonia, Manaus, Brazil (INPA), and the National Museum of Natural History, Smithsonian Institution, Washington, DC, USA (NMNH). UMSP barcode accession label numbers for holotypes are included in the list of material examined, but not for paratypes.
Systematics

Genus Notidobiella Schmid

Notidobiella Schmid 1955: 152 [Type species: Notidobiella parallelipipeda Schmid 1955, original designation].

Diagnosis: (modified from Schmid 1955; characters pertain to the male of the type species, except where noted): Head short, but broad, with large, projecting eyes, with conspicuous setae between ommatidia (Fig. 1); in most species interocular distance equal to or slightly less than diameter of eye, when viewed frontally (eyes of females smaller, interocular distance about 2× diameter of eye). According to Schmid (1955) bordering the eyes medially is an elongate, convex cephalic tubercle, but this structure was not evident in the material of the type species examined by us. Head dorsally with prominent, elongate occipital and retrooccipital warts; ocellar, antennal, frontal, anterolateral, and hypomedial setal warts absent (terminology of Ivanov 1990) (although females with small anterolateral setal wart). Antennal scape much shorter than head and about as wide as long, with oval setal warts on both its dorsal and ventral surfaces; scapes almost touch medially. Maxillary palp very short, composed of enlarged, heavily setose basal article held against the face and sclerotized on its lateral side only, medial side membranous, with small, oval, sclerotized, setose apical article (Fig. 2) (female maxillary palp 5-segmented, unmodified). Labial palp 3-segmented, unmodified. Pronotum with pair of elongate setal warts (Fig. 1). Mesoscutum with very deep median fissure and pair of small, oval scutal warts; mesoscutellum with pair of large scutellar warts (Fig. 1). Each leg with pair of apical spurs (tibial spur formula 2-2-2). Wings are broad, forewing longer and broader than hind wing; wing venation uniform among the species. Forewing (Fig. 3A) with forks I, II, III, and V present; cross vein r between R1 and R2; discoidal cell present and short, forks II and III sessile; crossveins r-m and m-cu present (in N. amazoniana sp. n., and N. ecuadorensis sp. n., crossveins r and s absent or very weak and difficult to discern in the material examined; the absence of s leaves the discoidal cell open); Cu2 not attaining wing margin, merged apically with Cu1b; A1 and A2 each attaining wing margin, A3 absent. Hind wing (Fig. 3B) with discoidal cell open; forks I, II, V present; fork I petiolate; fork II sessile; M with single branch; fork V short. Male genitalia (Figs 4, 8): Sternum VII with posteromesal process. Segment IX with short or long ventral setose lobes or only single setose process (N. amazoniana sp. n.). Preanal appendages short, ovate, setose. Tergum X simple, triangular to subquadrate in shape. Inferior appendage narrow basally and broadly spatulate apically, with short to long mesal process on ventromesal margin (in N. amazoniana sp. n., inferior appendage uniformly narrow throughout its length and without mesal process on the ventromesal margin). Phallus simple, elongate, tubular, with prominent endophallic membranes. Female genitalia (Figs 5, 9): Tergum IX heavily setose, posterolateral lobe, with lateral, microsetose, elevated ridge (all species except N. brasiliiana sp. n.). Internal vaginal sclerites complex with no discernable differences
among the species. Bursa copulatix subspherical and semisclerotized; pair of small, oval sclerites lying in membranes above vaginal sclerites (position variable).

Flint (1967) described the immature stages of *N. chacayana*. Larvae construct slightly tapered and curved cases of small mineral fragments embedded in silk and occur on the bottoms of small streams. The 3 previously known species, *N. chacayana* Schmid, 1957, *N. inermis* Flint, 1983, and *N. parallelipipeda* Schmid, 1955, are endemic to southern Chile.

**Notidobiella parallelipipeda** Schmid

Figs 1–5

*Notidobiella parallelipipeda* Schmid, 1955: 152 [Type locality: Chile, Ñuble, Recinto; NMNH; male]. Flint 1974: 91 [distribution].

**Description.** Of the species in the genus with broad, spatulate inferior appendages (all species except *N. amazoniana*), the type species is the most distinctive based on the parallel-sided inferior appendages with their prominent, mesally directed, mesal processes.

Adult. Forewing length 7.8–8.0 mm male (n=2); 8.8–9.0 mm female (n=2). Color light brown, palps and legs stramineous; forewings light brown, with scattered golden setae. Sternum VII of male with broad, fingernail-like, postermesal process.

Male genitalia (Fig. 4). Segment IX with anterior margin broadly produced midlaterally; tergum IX narrow; sternum IX with pair of very short, postermesal processes, bearing long apical setae. Tergum X simple, subquadrate in lateral view, with slight apicolateral elevation, with slight dorsomesal excavation, setose apically. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, spatulate, dorsal and ventral edges parallel, narrow basally, with prominent, elongate mesal process on mesal margin; strongly directed mesally in ventral view; apex exposed in lateral view. Phallic apparatus simple, tubular, slightly curved from base to apex; endophalic membranes prominent, but simple; phalotremal sclerite not apparent.

Female genitalia (Fig. 5). Tergum VIII quadrate; pleural membranes extensive, highly folded; sternum VIII broad, anterior margin with apodemal ridge, extending dorsolaterally; posterolateral corners rounded, heavily setose, especially posteriorly. Tergum IX with heavily setose, posterolateral lobes, rounded in lateral view, triangular in dorsal view; with lateral, microsetose, elevated ridge; sternum IX highly membranous, the membranes with parallel pleats or folds; tergum IX semimembranous dorsally. Tergum X with short setose projection. Internal vaginal sclerites complex (no discernable differences among the species); bursa copulatix subspherical, semisclerotized; pair of small, oval sclerites lying in membranes above vaginal sclerites (position variable).

**Material Examined:** CHILE: Ñuble, Recinto, 4–6.iii.1968, Flint and Peña, 1 male, 1 female (pinned) (NMNH); Linares, El Castillo, Malcho, E Parral, 750 m, 8–10.i.1988, L.E. Peña, 1 male, 1 female (pinned) (NMNH).
Figures 1–3. Notidobiella parallelepiped Schmid. 1 Male head and thorax A head, dorsal B pro- and mesonota, dorsal. 2. Notidobiella parallelepiped Schmid. Maxillary palp, male, frontal view. 3. Notidobiella parallelepiped Schmid. Male wings A forewing B hind wing. Abbreviations: DC = discoidal cell, TC = thyridial cell.
Notidobiella amazoniana Holzenthal & Blahnik, sp. n.
urn:lsid:zoobank.org:act:6203A2A2-CB58-418F-A731-8C31A2243E7D
Figs 6–7

Description. This is the smallest species in the genus. Its wings are short and broad with venation typical for the genus except forewing crossveins $r$ and $s$ are absent, leaving the discoidal cell open (Fig. 7). Its genitalia are the most atypical in the genus in that the inferior appendages are not broadly spatulate, but sinuous in shape and uniform in width.
Tergum IX bears a short triangular, posteromesal process, rather than short, paired processes, as found in the other species. Otherwise, the genitalia are typical for the genus.

Adult. Forewing length 4.5–5.0 mm male (n=8). Color faded, overall pale stramineous (specimens in alcohol); forewings colorless, almost transparent, denuded. Sternum VII of male with broad, fingernail-like, posteromesal process.

Male genitalia (Fig. 6). Segment IX with anterior margin acutely produced ventrolaterally; tergum IX narrow, ridge-like; sternum IX with short, triangular, posteromesal process, bearing apical setae. Tergum X simple, subquadrate in lateral view, with slight dorsomesal excavation, setose. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, elongate, narrow throughout length, without mesal process on ventromesal margin; in ventral view, apex acute, slightly incurved. Phallic

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**Figure 6.** *Notidobiella amazoniana*, sp. n. Male genitalia **A** segments IX, X, inferior appendages, lateral **B** segments IX, X, dorsal **C** segment IX, inferior appendages ventral **D** phallus, lateral **E** apex of phallus, dorsal.
apparatus simple, tubular, slightly curved from base to apex; endophalic membranes prominent, with paired apical membranous lobes; elongate, lightly sclerotized band internally (perhaps the phallostremal sclerite).

**Female.** Unknown.

**Holotype male:** BRAZIL: Amazonas: AM 010, km 246, 15–16.vii.1979, J. Arias (alcohol) (UMSP000131226) (INPA).

**Paratypes:** BRAZIL: Amazonas: same data as holotype, 3 males (alcohol) (UMSP), 4 males (alcohol) (NMNH).

**Etymology:** Named for the state and region where the specimens were collected, which represents a significant northern extension of the range of the genus.

*Notidobiella brasiliana* Holzenthal & Blahnik, sp. n.

urn:lsid:zoobank.org:act:3365A48E-060D-4925-AC4B-B8AD7E74944F

Figs 8–10

**Description.** This new species is most similar to *N. chacayana* in the overall shape and structure of the inferior appendages. Both species possess an elongate mesal process on the ventromesal margin of the inferior appendage. In *N. ecuadorensis* sp. n., the ventromesal process is also present, but is shorter and broader in ventral view; in the other 2 Chilean species, *N. inermis* and *N. parallelipipeda*, the ventromesal processes are either very reduced (*N. inermis*) or long (*N. parallelipipeda*), but not nearly as long.
as in *N. brasiliana* sp. n. Setting *N. brasiliana* sp. n., apart from all of its congeners is the pair of elongate posteromesal processes on sternum IX; in all other species these processes are much shorter and broader. Furthermore, forewing crossveins *r* and *s* are absent, leaving the discoidal cell open (Fig. 10).

Adult. Forewing length 7.0 mm male (*n*=1); 7.9–8.2 mm female (*n*=4). Color medium to dark brown, palps and legs light brown; forewings dark brown with scattered golden hairs, pale golden spot on anal margin at about midlength. Sternum VII of male with broad, fingernail-like, posteromesal process.

Male genitalia (Fig. 8). Segment IX with anterior margin broadly produced mesolaterally; tergum IX narrow, elevated, mound-like; sternum IX with pair of prominent, elongate, posteromesal processes, bearing long apical setae. Tergum X simple, triangu-
Figure 9. *Notidobiella brasiliana*, sp. n. Female genitalia **A** segments VIII-X, lateral **B** segments IX, X, dorsal **C** segments VIII-X, ventral.

Figure 10. *Notidobiella brasiliana*, sp. n. Wings **A** forewing **B** hind wing.
lar in lateral view, with slight dorsomesal excavation, setose. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, broadly spatulate, narrow basally, with elongate mesal process on ventromesal margin. Phallic apparatus simple, tubular, slightly curved from base to apex; endophalic membranes prominent, but simple; phalotremal sclerite not apparent.

Female genitalia (Fig. 9). Tergum VIII quadrate; pleural membranes extensive, highly folded; sternum VIII broad, anterior margin with apodemal ridge, extending dorsolaterally; posterolateral corners rounded, heavily setose, especially posteriorly. Tergum IX with heavily setose, posterolateral lobes, rounded in dorsal and lateral views; without lateral ridge; sternum IX highly membranous, the membranes with parallel pleats or folds; tergum IX semimembranous dorsally. Tergum X with short, bifurcate, setose projection. Internal vaginal sclerites complex (no discernable differences among the species); bursa copulatrix subspherical, semiclerotized; pair of small, oval sclerites lying in membranes above vaginal sclerites (position variable).

**Holotype male: BRAZIL: São Paulo:** Parque Estadual de Campos do Jordão, 1st order trib. to Rio Galharada, 22°41.662'S, 45°27.783'W, el 1530 m, 14–16.ix.2002, Blahnik, Prather, Huamantinco (pinned) (UMSP000086351) (MZUSP).

**Paratypes: BRAZIL: São Paulo:** Parque Estadual de Campos do Jordão, Rio Galharada, 22°41.662'S, 45°27.783'W, el 1530 m, 13–15.ix.2002, Blahnik, Prather, Melo, Huamantinco, 2 females (alcohol) (MZUSP); same data as holotype, 2 females (pinned) (UMSP).

**Etymology:** Named for Brazil, the country of the type specimens, which represents a significant northeastward extension of the range of the genus.

**Notidobiella chacayana** Schmid

Figs 11–12

*Notidobiella chacayana* Schmid 1957: 392 [Type locality: Chile, Maule, Chacay; NMNH; male]. Flint 1967:63 [larva, pupa]; 1974:91 [distribution].

**Description.** This Chilean species is most similar to *N. brasiliana* because of the similarly shaped inferior appendages, with their similar elongate mesal processes. It differs from that species in the much shorter posteromesal processes of sternum IX.

Adult. Forewing length 6.8–7.5 mm male (n=3); 7.2–9.0 mm female (n=3). Color brown, palps and legs stramineous; forewings brown, with scattered golden setae. Sternum VII of male with broad, fingernail-like, posteromesal process.

Male genitalia (Fig. 11). Segment IX with anterior margin produced ventrolaterally; tergum IX slightly elevated, mound-like; sternum IX with pair of short, posteromesal processes, bearing long apical setae. Tergum X simple, triangular in lateral view, with dorsomesal excavation, setose. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, broadly spatulate, narrow basally, with elongate
mesal process on ventromesal margin. Phallic apparatus simple, tubular, curved from base to apex; endophalic membranes prominent, but simple; phallotremal sclerite not apparent.

Female genitalia (Fig. 12). Tergum VIII quadrate; pleural membranes extensive, highly folded; sternum VIII broad, anterior margin with apodemal ridge, extending dorsolaterally; posterolateral corners rounded, heavily setose, especially posteriorly. Tergum IX with heavily setose, posterolateral lobes, rounded to subtriangular in dorsal view; with lateral, microsetose, elevated ridge; sternum IX highly membranous, the membranes with parallel pleats or folds; dorsally tergum
IX semimembranous. Tergum X with short setose projection. Internal vaginal sclerites complex (no discernable differences among the species); bursa copulatrix subspherical, semiscerotized; pair of small, oval sclerites lying in membranes above vaginal sclerites (position variable).

**Material Examined: CHILE:** Cauquenes, Tregualeme, 35°56’S, 72°43’W, 11–12. xii.1993, C. and O. Flint Jr., 1 male, 1 female (pinned) (NMNH); X Región de los Lagos, Isla de Chiloé, Río Verde, 1.9 km W Puntra, 42°07.078’S, 73°50.364’W, el. 40 m, 3.ii.2005, Holzenthal, Blanhik, Chamorro, 2 males, 2 females (pinned) (UMSP); XIV Región de los Ríos, Monumento Nacional Alerce Costero, unnamed trib., trail to Alerce Milenario, 40°11.874’S, 73°26.217’W, el. 895 m, 5.ii.2008, Holzenthal, Pauls, Mendez, 1 male (pinned) (UMSP).

**Figure 13.** Notidobiella ecuadorensis, sp. n. Male genitalia A segments IX, X, inferior appendages, lateral B segments IX, X, dorsal C segment IX, inferior appendages ventral D inferior appendage, dorsal E phal- lus, lateral F phallus apex, dorsal G sternum VII posteromesal process, ventral.
Notidobiella ecuadorensis Holzenthal & Blahnik, sp. n.

URN:lsid:zoobank.org:act:20CB7DC2-73DF-4D4D-9593-0A87C589BABB

Figs 13–14

Description. The combination of broadly spatulate inferior appendage, thumb-like mesal process on the ventromesal margin of the inferior appendage, and short posteromesal processes on sternum IX separate this species from its congeners. The wing venation (Fig. 14) is similar to that of the type species.

Adult. Forewing length 6.2 mm (n=1). Color faded, overall yellowish-brown (specimen in alcohol); forewings stramineous, denuded. Sternum VII of male with broad, fingernail-like, posteromesal process.

Male genitalia (Fig. 13). Segment IX with anterior margin broadly produced midlaterally; tergum IX narrow, elevated, mound-like; sternum IX with pair of short, triangular, posteromesal processes, bearing very long apical setae. Tergum X simple, triangular in lateral view, with slight dorsomesal excavation, setose apically. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, very broadly spatulate, narrow basally, with short, thumb-like mesal process on ventromesal margin. Phallic apparatus simple, tubular, relatively straight from base to apex; endophalic membranes prominent, with paired apical membranous lobes; elongate, lightly sclerotized band internally (perhaps the phallotremal sclerite).

Female: Unknown.

Holotype: male, ECUADOR: Pastaza: Puyo, 1–7.ii.1976, Spangler et al. (alcohol) (UMSP000208470) (NMNH).
**Etymology:** Named for Ecuador, the country of the holotype, which represents a significant northern extension of the range of the genus.

**Notidobiella inermis** Flint

Figs 15–16

*Notidobiella inermis* Flint 1983: 90 [Type locality: Chile, Pcia Cautín, near Pucón; NMNH; male].

**Description.** *Notidobiella inermis* shares with *N. ecuadorensis* broadly spatulate inferior appendages with short, thumb-like mesal processes, but differs in details of the shape.
of the inferior appendages, as illustrated, and in the possession of a narrow posteromesal process on sternum VII.

Adult. Forewing length 6.0–6.5 mm male (n=2); 8.0 mm female (n=2). Color brown, palps and legs stramineous; forewings brown, with scattered golden setae. Sternum VII of male with narrow, fingernail-like, posteromesal process.

Male genitalia (Fig. 15). Segment IX with anterior margin produced ventrolaterally; tergum IX narrow; sternum IX with pair of short, posteromesal processes, bearing long apical setae. Tergum X simple, subquadrate in lateral view, with slight dorsomesal excavation, setose. Preanal appendage short, ovate, setose. Inferior appendage prominent, heavily setose, very broadly spatulate, narrow basally, with short, thumb-like mesal process on ventromesal margin. Phallic apparatus simple, tubular, relatively straight from base to apex; endophallic membranes prominent, but simple; phallotremal sclerite not apparent.

Female genitalia (Fig. 16). Tergum VIII quadrate; pleural membranes extensive, highly folded; sternum VIII broad, anterior margin with apodemal ridge, extending dorsolaterally; posterolateral corners rounded, heavily setose, especially posteriorly. Tergum IX with heavily setose, posterolateral lobes, subovate, small, bilobed in dorsal and lateral views; with lateral, microsetose, elevated ridge; sternum IX highly membranous, the membranes with parallel pleats or folds; dorsally tergum IX with sclerotized ridge. Tergum X with broad heavily setose projection. Internal vaginal sclerites complex, no discernable differences among the species; bursa copulatrix subspherical, semisclerotized; pair of small, oval sclerites lying in membranes above vaginal sclerites (position variable).

**Material Examined:** CHILE: Llanquihue, Salto Chamiza, Correntosa, 100 m, 19.i.1987, C.M. and O.S. Flint Jr., 1 male, 2 females (pinned) (NMNH); Llanquihue, El Chinque, N Correntosa (S Volcán Calbuco), 300 m, 20–25.i.1980, 1 male paratype (pinned) (NMNH).

**Key to males of Neotropical Notidobiella**

1 Inferior appendage narrow basally, broadly spatulate apically, with mesal process on ventromesal margin (Figs 4A, 8A); abdominal sternum IX with pair of posteromesal processes (Figs 4C, 8C); forewing length 6–8 mm ...............2

2 Inferior appendage elongate, narrow throughout length, without mesal process on ventromesal margin (Figs 6A, C); abdominal sternum IX with single, short, triangular, posteromesal process (Fig. 6C); forewing length 4.5–5 mm ...............2

.... Notidobiella amazoniana sp. n.

3 Inferior appendage mesal process elongate (Figs 4C, 8C) ..................3

4 Inferior appendage mesal process short, thumb-like (Figs 13C, 15C) .......5

5 Posteromesal processes of abdominal sternum IX short (Figs 6C, 13C); distribution: Chilean subregion (Chile) ...... Notidobiella brasiliana sp. n.

6 Posteromesal processes of abdominal sternum IX elongate (Fig. 8C); distribution: Brazilian subregion (southeastern Brazil) ....
### 4(3) Spatulate apex of inferior appendage broadly ovate (Fig. 11A)  

- Notidobiella chacayana Schmid

### 5(2) Abdominal tergum IX highly elevated, mound-like (Fig. 13A); ventromesal process of abdominal sternum VII broad (Fig. 13G); distribution: Brazilian subregion (Ecuador)  

- Notidobiella ecuadorensis sp. n.

- Abdominal tergum IX not elevated (Fig. 15A); ventromesal process of abdominal sternum VII narrow (Fig. 15F); distribution: Chilean subregion (Chile)  

- Notidobiella inermis Flint

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### Biogeographic considerations

As defined most recently by de Moor and Ivanov (2008), the Trichoptera fauna of southern Chile and adjacent patagonian Argentina exhibits a strong biogeographical affinity to Australia, New Zealand, and other southern Pacific islands (e.g., New Caledonia). This “Temperate Gondwana” (de Moor and Ivanov 2008) or “transantarctic” pattern (e.g., Brundin 1966) also includes the temperate, southernmost part of Africa and Madagascar. In the Neotropical Trichoptera, the affinity to the southern African fauna is very weak, perhaps exhibited only at the family level within Sericostomatoida (de Moor and Ivanov 2008) and potentially among genera within Sericostomidae (although relationships among genera within this family are yet to be inferred). On the other hand, the biogeographical affinity between southern South America and Australasia is strong. Within South America, nearly all species in Chile and adjacent Argentina are endemic, prompting Flint (1976) to divide the Neotropics into 2 distinct subregions, the Chilean (southern Chile/Argentina) and the Brazilian (the rest of the Neotropics as defined by Wallace 1876); these regions are equivalent to the “Patagonian” and “Neotropical” (sensu stricto) Trichoptera regions of de Moor and Ivanov (2008).

Four Trichoptera families are representative of a temperate Gondwanan pattern: Helicophidae, Kokiriidae, Philorheithridae, and Tasimiidae, each family with genera endemic to Australia/New Zealand/New Caledonia, southern South America, or Madagascar (no genera are shared) (Neboiss 1986, Flint et al. 1999b, Holzenthal et al. 2007b, Weaver et al. 2008) (Table 1). Sukatcheva and Jarzembowski (2001) questionably placed a fossil (fragment of a forewing) from the early Cretaceous of southern England in the Helicophidae.

While not wholly endemic to the region, other caddisfly families contain a diverse temperate Gondwana fauna including, most notably, Hydrobiosididae (reviewed by Schmid 1989, Ward et al. 2004), Limnephilidae: Dicosmoecinae (Wiggins 2002) and Leptoceridae: Triplectidinae (Holzenthal 1986a, b, Morse and Holzenthal 1987, de Moor 1997, Holzenthal and Pes 2004, Calor et al. 2006, Calor and Holzenthal 2008, Malm and Johanson 2008). Other extant caddisfly taxa conforming to a tem-
Table 1. Genera (number of included species) in the families Helicophidae, Kokiriidae, Philorheithridae, and Tasimiidae and their regional distributions, including references to recent works inferring or discussing phylogenetic relationships among genera.

| Family, genus (# species) | Distribution |
|--------------------------|--------------|
| Helicophidae (Flint 1992, 2002, Henderson and Ward 2007, Johanson 2003a, Johanson and Keijsner 2008, Johanson and Ward 2002, Neboiss 2002) | |
| *Alloecella* Banks (3) | SE Australia, Tasmania |
| *Alloecentrella* Wise (4) | New Zealand |
| *Alloecentrellodes* Flint (2) | Chile |
| *Austrocentrus* Schmid (3) | Chile, Argentina |
| *Eosericostoma* Schmid (2) | Chile, Argentina |
| *Helicopha* Mosely (21) | Australia, Tasmania, New Caledonia |
| *Helococcus* Neboiss (1) | E Australia |
| *Microthremma* Schmid (8) | Chile |
| *Pseudosericostoma* Schmid (1) | Chile |
| *Zelolessica* McFarlane (2) | New Zealand (incl. Stewart Island) |
| Kokiriidae (Johanson 2003b) | |
| *Kokiria* McFarlane (1) | New Zealand |
| *Mecynostomella* Kimmins (7) | New Caledonia |
| *Pangulia* Navás (2) | Chile |
| *Tanjistomella* Neboiss (1) | SE Australia |
| *Taskiria* Neboiss (3) | SE Australia, Tasmania |
| *Taskirosyche* Neboiss (1) | Tasmania |
| Philorheithridae (Henderson and Ward 2006, Weaver et al. 2008) | |
| *Aforheithrus* Weaver, Gibon, and Chvojka (3) | Madagascar |
| *Aphilorheithrus* Mosely (4) | SE Australia, Tasmania |
| *Australheithrus* Mosely (3) | SE Australia, Tasmania |
| *Kosrheithrus* Mosely (3) | SE, SW Australia, Tasmania |
| *Mystacopsyche* Schmid (2) | Chile, Argentina |
| *Philorheithrus* Hare (6) | New Zealand |
| *Psilopsyche* Ulmer (3) | Chile, Argentina |
| *Ramiteithrus* Neboiss (2) | SE Australia, Tasmania |
| *Tasmanthrus* Mosely (3) | Tasmania |
| Tasimiidae (no phylogenetic assessment available) | |
| *Charadropsyche* Flint (1) | Chile |
| *Tasiagma* Neboiss (2) | SE Australia, Tasmania, Lord Howe Island |
| *Tasimia* Mosely (5) | SE Australia, Tasmania |
| *Trichovespula* Schmid (1) | Chile |

perate Gondwanan pattern between the Neotropics (*sensu* Wallace) and Australasia include closely related or purported sister genera in otherwise more widely distributed or cosmopolitan families, including genera in Ecnomidae (Flint 1973, Li and Morse 1997, Cartwright 2009, Johanson and Espeland 2009), Hydropsychidae: Smicrideinae, Macronematinae (Neboiss 1984, Schefter 1996, Geraci et al. 2005), Philopotamidae (Blahnik 2005), and possibly Stenopsychidae. Other Trichoptera taxa endemic to
the Patagonian region (as defined by de Moor and Ivanov 2008), but that apparently either do not pertain to a transantarctic pattern or have unknown biogeographic affinities include genera in Anomalopsychidae (Holzenthal and Flint 1995, Holzenthal and Robertson 2006), Glossosomatidae (Robertson and Holzenthal 2005), Leptoceridae: Leptocerinae (Holzenthal 1986c), and Hydroptilidae (Harris and Armitage 1997, Harris and Flint 1993).

The family Sericostomatidae contains temperate Gondwanan components, including 5 endemic South African/Malagasy genera (Aclosma, Aselas, Cheimacheramus, Petroplax, Rhoizema) and 4 endemic South American genera (Grumicha, Myotrichia, Notidobiella, Parasericostoma, excluding Chiloecia, nomen dubium). As indicated above, the family includes other genera endemic to the Nearctic and West Palearctic regions. Sericostomatids are absent from the Australasian region (all Australasian species previously assigned to Sericostomatidae have been transferred to other families, see Holzenthal et al. 2007b for a historical review). Phylogenetic relationships among families and genera of Sericostomatoidea are largely unresolved (Holzenthal et al. 2007a) making it impossible to construct an area cladogram to test for congruence with the prevailing hypotheses of the geological sequence of the breakup of Pangea or the subsequent breakup of southern Gondwana (Sanmartín and Ronquist 2004, fig. 1).

The presence of Helicophidae and Hydrobiosidae in Eocene Baltic amber (Botosaneanu and Wichard 1983) and of fossil Plectrotarsidae (extant taxa endemic to Australasia) and a putative helicophilid from late Cretaceous deposits in England (Sukatsheva and Jarzembowski 2001) suggests, as hypothesized by de Moor and Ivanov (2008), that certain southern temperate Trichoptera may be relicts of a more widespread fauna which included now extinct (but still extant in Sericostomatidae) north temperate elements. Two species of Triplectides in Baltic amber (Ulmer 1912) suggests the same scenario for this southern Gondwanan genus. In addition, the putative triplectidine larva from South Africa (de Moor 1997) and the recent discovery of Philorheithridae in Madagascar (Weaver et al. 2008) suggest at least a Gondwanan origin for these taxa (category 3 of Amorin et al. 2009) and, by inference, other endemic austral Trichoptera.

Evidence suggests that the contemporary distribution of the Patagonian and Australasian temperate Gondwanan Trichoptera fauna reflects a past dispersal corridor between Australia and southern South America via Antarctica (Sanmartín and Ronquist 2004). This “transantarctic exchange” pattern also has been demonstrated for other insects, including aquatic taxa (Cranston and Edward 1999, Amorin et al. 2009, Daugeron et al. 2009). However, it may be that the current distribution of other southern Gondwana caddisflies, such as Sericostomatidae, reflects an older dispersal prior to the breakup of southern Gondwana, now represented by relict Southern Hemisphere distributions. The southeast Brazilian sericostomatid Grumicha grumicha might represent evidence to support the relict hypothesis.

The now widespread occurrence of Notidobiella in temperate southern Chile and tropical South America (Ecuador, southeast Brazil, Amazonian Brazil) suggests a more recent dispersal of the genus to northern tropical South America from Patagonia and...
its subsequent diversification. The data from insects analyzed by Sanmartín and Ronquist (2004, table 4) found a significantly higher frequency of dispersal from southern South America to northern South America than from the other direction. This may be true for other caddisflies with both Patagonian and Neotropical (sensu de Moor and Ivanov) distributions, including *Antarctoecia* (Huamantinco and Nessimian 2003), *Atopsyche* and *Cailoma* (Hydrobiosidae) (Ross and King 1952, Flint 1974, Schmid 1989 [although *Atopsyche* is absent from Patagonia, its putative sister genus is Patagonian]), *Contulma* (Anomalopsychidae) (Holzenthal and Flint 1995), *Smicridea* (*Smicridea*) (Hydropsychidae) (Flint 1989), *Tolhuaca* (Glossosomatidae) (Robertson and Holzenthal 2005), and *Triplectides* (Leptoceridae).

As confirmed by Crisci et al. (1991), Sanmartín and Ronquist (2004), and Amorin et al. (2009) the historical biogeography of southern South America is complex. The distribution of the austral South American caddisflies support this conclusion, with a fauna pertaining strongly to a transantarctic pattern, but with perhaps older Gondwana elements, as exhibited by Sericostomatidae. The current weight of evidence described above, however, supports a more recent dispersal of this southern fauna to northern South America.

Other than in a few studies, phylogenetic hypotheses are lacking for most of the taxa reviewed above. Phylogenies of Southern Hemisphere caddisfly taxa inferred from molecular data are even fewer (e.g., Johanson and Keijsner 2008) and only one (Johanson et al. 2009) has used events-based models (e.g., Sanmartín at al. 2001, Ronquist and Sanmartín 2004) or divergence time estimates (but see Amorin et al. 2009 for a critique of molecular dating) to address historical biogeography. Given the current ease with which molecular sequence data can be obtained and with the availability of newer analytical methods (e.g., Ronquist 1997, Sanderson 2002, Zaldivar-Riverón et al. 2008), there is a wealth of hypotheses that can be tested regarding the historical biogeography of the austral caddisfly fauna once phylogenetic information is available (Santos and Amorin 2007).

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