Two new species of *Leptanilloides* Mann, 1823 (Formicidae: Dorylinae) from the Andes of southern Ecuador

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Abstract. Two new species of *Leptanilloides* are described: *L. copalinga* Delsinne & Donoso sp. nov. and *L. prometea* Delsinne & Donoso sp. nov., based on workers collected in the leaf litter and soil of the Andes of southern Ecuador. Both species belong to the *L. biconstricta* species-group (formally diagnosed here). The metatibial gland, considered a synapomorphy for Dorylinae, is observed in *L. prometea* sp. nov. but seems absent in *L. copalinga* sp. nov. We provide a COI DNA barcode for both species and a revised key for the worker caste of all known species in the genus. We also describe a single male identified as a potential new *Leptanilloides* species on the basis of morphology. Furthermore, its mitochondrial COI gene sequence does not match any previously barcoded species. However, we refrain from giving it a specific name because of our lack of knowledge about the worker caste. So far, half of the 14 *Leptanilloides* species have been discovered above 1500 m in the mountain forests or páramos of the Ecuadorian Andes, confirming, if needed, the biological significance of these threatened habitats.

Key words. Army ant syndrome, DNA barcoding, male description, metatibial gland, subterranean ants.


Introduction

The ant genus *Leptanilloides* Mann, 1923 belongs with *Amyrnx* Kusnezov, 1953 to a monophyletic clade on the basis of the molecular phylogeny of the recently reassessed subfamily Dorylinae Leach, 1815 (Brady *et al.* 2014). This clade was formerly recognized as a distinct subfamily, Leptanilloidinae Baroni Urbani *et al.*, 1992, and also comprises a third genus, *Asphinctanilloides* Brandão *et al.*, 1999, for which no DNA sequences are yet available.
Leptanilloides comprises 14 species of subterranean ants, including two species described as new in this paper, with morphological and behavioral traits distinctive of the army ant syndrome, including synchronization of larva production, brood carrying by workers under their mesosoma, nomadic habits, and a subdichthadiiform gyne (Brandão et al. 1999; Donoso et al. 2006; Borowiec & Longino 2011). They present an apparent disjunctive distribution with the majority of species infrequently collected from Mexico to Bolivia, at altitudes of between 440 and 3350 m (Brandão et al. 1999; Longino 2003; Donoso et al. 2006; Ward 2007; Ward & Brady 2009; Borowiec & Longino 2011), and a single species, L. atlantica Silva et al., 2013, only known from its type locality at 800 m in south-eastern Brazil, more than 2500 km away from the other species (Silva et al. 2013). However, this distribution pattern is suspected to be the result of the insufficient or inadequate collecting effort applied to the soil environment (Silva et al. 2013). Our knowledge concerning the diversity and distribution of this genus is indeed very low: most species are only known from their type locality, one species from just two sites (L. gracilis Borowiec & Longino, 2011 from Mexico and Guatemala; Borowiec & Longino 2011), and two species from three localities (L. mckennae Longino, 2003 in Costa Rica (AntWeb 2015a); L. biconstricta Mann, 1923 from Bolivia (Mann 1923), Colombia (Zabala et al. 2006) and Venezuela (AntWeb 2015b)).

Borowiec & Longino (2011) updated the diagnosis of the genus based on the worker caste. They described the state of 38 morphological variables and discussed the variability observed for some of them within the genus. In particular, they noted the presence, albeit reduced, of a metatibial gland in two species (L. erinys Borowiec & Longino, 2011 and L. nubecula Donoso et al., 2006). This observation was interesting because this gland is considered to be a synapomorphy of Dorylinae in relation to Formicidae (Bolton 1990; 2003) but was estimated to be secondarily absent in Leptanilloides (Brandão et al. 1999; Bolton 2003; Longino 2003; Brady & Ward 2005; Donoso et al. 2006).

Morphological data seem to support the segregation of Leptanilloides into two natural species-groups (Borowiec & Longino 2011): the Leptanilloides biconstricta species-group, easily recognizable for the completely unfused promesonotal connection and the postpetiole nearly as high as the following abdominal segment, and the Leptanilloides legionaria species-group, with ants that have a partially to completely fused promesonotal connection and a reduced and isolated postpetiole. Importantly, the phylogeny of Brady et al. (2014) inferred from 11 nuclear DNA markers also supports this distinction (but only five Leptanilloides species were included in their work). In this paper, we provide a formal diagnosis of both species-groups.

The male caste has only been described for two species of the L. legionaria species-group, namely L. mckennae (Ward 2007) and L. nubecula (Donoso et al. 2006). In addition, Borowiec & Longino (2011) described three males without associating them with workers or giving them specific names. The first male (called ‘Male 1’ in Borowiec & Longino 2011) was suspected to belong to L. gracilis, a species from the L. biconstricta species-group, because of sympatric distribution, similar abundance, small size and the presence of two simple spurs on mid and hind tibia in both ‘Male 1’ and workers of L. gracilis. However, because several Leptanilloides species can occur in sympatry (Donoso et al. 2006; Borowiec & Longino 2011), the association was cautiously not formalized without further evidence. No hypotheses were proposed concerning the identities of the two other males. It is noteworthy that males are also known for Amyrmex (Ward & Brady 2009), but not for Asphictanilloides.

Boudinot (2015) indicated that most males of Dorylinae are uniquely identified by the bidentate or pronged ninth abdominal sternite (subgenital plate), the lack of pygostyles, and the poorly developed clypeus. Males of Leptanilloides and Amyrmex are highly derived and are identifiable by the following combination of characters: antennal torulus abutting or very nearly abutting anterior clypeal margin, oblique mesopleural sulcus absent, four closed cells present on forewing, and cinctus between abdominal pre- and postsclerites IV absent (Boudinot 2015; see also Ward & Brady 2009). Moreover, the posterior
Here we describe the worker caste of two new *Leptanilloides* species belonging to the *L. biconstricta* species-group. A male with a unique mandible shape is also described, although we do not intend to formally name it until further information regarding its associated worker caste has been acquired. We also provide COI barcodes for the two new species and for this male. The key of Borowiec & Longino (2011) to the worker caste is updated to include our two new species as well as the recently described *L. atlantica* Silva *et al.*, 2013. Finally, we provide evidence of a metatibial gland in one of our species and note the hitherto unnoticed occurrence of a pectinate spur on the midtibia in some species.

**Material and methods**

**Molecular analyses**

One hindleg was sampled from 4 specimens (specimen codes of *L. prometea* sp. nov. workers: 4052302, 4052314; code of *L. copalinga* sp. nov. worker: 4006302; code of the unassociated male: 4870101), and DNA was extracted with the NucleoSpin Tissue Kit (Macherey-Nagel, Germany) following the manufacturer’s protocol. Moreover, the total genomic DNA was isolated from the complete body of 4 workers of *L. prometea* sp. nov. (specimen codes 4052311, 4052312, 4060602, and 4060603) also using the NucleoSpin Tissue Kit. After DNA extraction, these 4 specimens were preserved as vouchers in 96% ethanol. PCR amplification of the 5’ end of the cytochrome c oxidase subunit I (COI) marker (standard DNA barcode region) was performed with the primers LCO1490 and HCO2198 (Folmer *et al.* 1994). The wingless wnt-1 gene (Wg) was amplified using primers Wg578F (Ward & Downie 2005) and Wg1032R (Abouheif & Wray 2002). PCR conditions were the same as in Delsinne *et al.* (2012). Amplicons were directly sequenced in both directions and aligned with all COI and Wg sequences of *Leptanilloides* and *Cylindromyrmex* Mayr, 1870 (as outgroup) available in GenBank and in BOLD, the Barcode of Life Data System (Ratnasingham & Hebert 2007). We obtained COI sequences for 6 specimens of *L. prometea* sp. nov., 1 of *L. copalinga* sp. nov., and 1 of the unnamed *Leptanilloides* male. Unfortunately, we obtained only one Wg sequence for *L. prometea* sp. nov. (specimen code 4052311; Appendix 1). Mega ver. 6.06 (Tamura *et al.* 2013) was used to perform alignments with the Clustal W algorithm (Thompson *et al.* 1994), calculate pairwise uncorrected p-distances and construct neighbor-joining trees with bootstrapping (1000 replicates).

All COI and Wg sequences were deposited in GenBank with accession numbers KT601697–KT601704 (COI) and KT750331 (Wg). Process IDs of specimens in BOLD are from LEPEC001-15 to LEPEC007-15 and LEPEC009-15. Further phylogenetic analyses were performed on the basis of the Wg DNA dataset (Appendix 1), which included sequences representing more species than the COI dataset. Parsimony trees were calculated in R using the packages ape (Paradis *et al.* 2004) and phangorn (Schliep 2011) with 1000 bootstrap replicates (non-parametric bootstrapping) (Appendix 2). Best partitioning schemes and best-fit substitution models were estimated using PartitionFinder ver. 1.0 (Lanfear *et al.* 2012) in order to perform maximum likelihood analysis and Bayesian inference of phylogeny. Two partitions (one partition for the third codon position and one partition for the two other positions) were selected, with best-fit substitution models TIM+Gamma (Posada 2003) and JC (Jukes & Cantor 1969), respectively (using the Bayesian information criterion). Maximum likelihood analysis was conducted using GARLI ver. 2.0 (Zwickl 2006) with a bootstrapping test (500 replicates) (Appendix 3). Bayesian inference was performed with MrBayes ver. 3.2.3 (Ronquist *et al.* 2012; Appendix 1). Two parallel runs, with four chains each, were run for ten million generations, with unlinked nucleotide substitution parameters for each data partition. Every 1000th generation was sampled, convergence of the Markov chains was monitored, average standard deviation of split frequencies decreased below 0.01, and traces of the parameters were checked using Tracer ver. 1.6 (Rambaut *et al.* 2014) following
the recommendations of Ronquist et al. (2011), Lesaffre & Lawson (2012) and SAS Institute (2009). Finally, the first 25% of the trees were discarded (“burn-in”) to summarize the tree samples.

Images

High resolution digital images were taken using either a Leica DFC290 camera attached to a Leica Z6APO stereo microscope or a Leica MC170 camera attached to a Leica S8APO stereo microscope. A series of images was taken by focusing the sharpness on different levels of the specimen, using the Leica Application Suite ver. 38 (2003–2011) and combined with the stacking software Combine ZP (Hadley 2010). Final editing of the images was done in Adobe Photoshop CS5. SEM photographs of the specimens (spm 4006301 and spm 4052301 for *L. copalinga* sp. nov. and *L. prometea* sp. nov., respectively) were taken at the Royal Belgian Institute of Natural Sciences using a FEI Quanta 200 scanning electron microscope (SEM).

Measurements

Measurements were made using an Olympus SZ61 stereo microscope at a magnification of 45× with a micrometer. All the measurements are presented in millimeters. The following abbreviations are used:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>HL</td>
<td>Head length, measured in full face view, from anterior edge of frontal lobes (therefore without including clypeal lamella) to posterior border of head</td>
</tr>
<tr>
<td>HW</td>
<td>Head width, maximum width of head measured in full face view; HW for males includes eyes (workers eyeless)</td>
</tr>
<tr>
<td>SL</td>
<td>Scape length, excluding basal condyle and neck</td>
</tr>
<tr>
<td>LAII, LAIII, LAIV, LAXIII</td>
<td>Length of second, third, fourth and terminal (13th) antennal segments, respectively (male only)</td>
</tr>
<tr>
<td>EL</td>
<td>Eye length, measured in full face view, maximum length of eye parallel to midline (male only)</td>
</tr>
<tr>
<td>MaL</td>
<td>Mandible length, measured in full face view, length of line running medially from clypeal anterior margin to mandible apex (male only)</td>
</tr>
<tr>
<td>PrW</td>
<td>Pronotal width, maximum width in dorsal view</td>
</tr>
<tr>
<td>WL</td>
<td>Weber’s length, measured from anterior edge of pronotum to posterior edge of metapleural lobe; this measure called Mesosoma Length (ML) in Borowiec &amp; Longino (2011) and Silva et al. (2013)</td>
</tr>
<tr>
<td>MH</td>
<td>Mesosoma height: in lateral view, maximum height measured from lowermost point of mesopleuron (in front of middle coxa) to dorsal edge of mesosoma, measured perpendicularly to long axis of mesosoma</td>
</tr>
<tr>
<td>PL</td>
<td>Petiole length, maximum length of the petiole measured in dorsal view, starting at base of anterior face and ending at base of posterior edge</td>
</tr>
<tr>
<td>PW</td>
<td>Petiole width, maximum width of petiole measured in dorsal view</td>
</tr>
<tr>
<td>PPL</td>
<td>Postpetiole (= third abdominal segment) length, maximum length of node measured in dorsal view</td>
</tr>
<tr>
<td>PPW</td>
<td>Postpetiole width, maximum width of node measured in dorsal view</td>
</tr>
<tr>
<td>AivL</td>
<td>Length of fourth abdominal segment, in dorsal view</td>
</tr>
<tr>
<td>AivW</td>
<td>Maximum width of fourth abdominal segment, in dorsal view</td>
</tr>
<tr>
<td>FFeL</td>
<td>Maximum length of fore femur, measured along extensor (outer) surface</td>
</tr>
<tr>
<td>FFeW</td>
<td>Maximum width in lateral view of fore femur</td>
</tr>
<tr>
<td>HFeL</td>
<td>Length of hind femur, measured along extensor (outer) surface</td>
</tr>
<tr>
<td>HTL</td>
<td>Length of hind tibia, measured along extensor (outer) surface</td>
</tr>
<tr>
<td>CI</td>
<td>Cephalic index, HW/HL × 100</td>
</tr>
<tr>
<td>SI</td>
<td>Scape index, SL/HL × 100</td>
</tr>
<tr>
<td>PI</td>
<td>Petiolar index, PW/PL × 100</td>
</tr>
</tbody>
</table>
MI = Mesosomal index, MH/WL × 100
sc = specimen code(s) following the Maurice Leponce database at the Royal Belgian Institute of Natural Sciences

For consistency, we follow Borowiec & Longino (2011) for terminology, in particular for wing venation and genitalia.

Deposition of material

Holotypes and paratypes have been deposited at:

BMNH = British Museum of Natural History (Natural History Museum), London, UK
CASC = California Academy of Sciences, San Francisco, CA, USA
ICN = Insect Collection, Instituto de Ciencias Naturales, Universidad Nacional de Colombia, Bogotá D.C., Colombia
JTLC = John T. Longino, personal collection, University of Utah, Salt Lake City, UT, USA
MCZC = Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts, USA
MUTPL = Museo de Colecciones Biológicas de la Universidad Técnicas Particular de Loja, Loja, Ecuador
MZSP = Museu de Zoologia da Universidade de São Paulo, São Paulo, Brazil
QCAZ = Museum of Zoology of the Pontificia Universidad Católica del Ecuador, Quito, Ecuador
RBINS = Royal Belgian Institute of Natural Sciences, Brussels, Belgium

Results

Class Hexapoda Blainville, 1816
Order Hymenoptera Linnaeus, 1758
Suborder Apocrita Latreille, 1810
Infraorder Aculeata Latreille, 1802
Superfamily Vespoidea Latreille, 1802
Family Formicidae Latreille, 1809
Subfamily Dorylinae Leach, 1815

Genus *Leptanilloides* Mann, 1823

Diagnosis of the *Leptanilloides biconstricta* species-group

Species from the *Leptanilloides biconstricta* species-group possess the typical characters of the genus (see diagnosis in Borowiec & Longino 2011) but can be grouped by the fact that they share the following criteria: promesonotal connection completely unfused and flexible; lateroclypeal tooth (called genal tooth in Brandão et al. 1999, Longino 2003 and Donoso et al. 2006) well-developed (although apparently lacking in *L. caracola* Donoso et al., 2006, a species known only by its holotype; because this criterion may be difficult to observe with a stereo microscope, even at high magnification, the absence of a lateroclypeal tooth in *L. caracola* should be confirmed in SEM); abdominal segment III (postpetiole) nearly as high as abdominal segment IV; and postpetiolar spiracle situated proximately to the anterior margin of the tergite.

Moreover, species from this group are generally smaller than species of the *L. legionaria* species-group (Fig. 1).

This group includes the ten following species:
*L. atlantica* Silva et al., 2013
*L. biconstricta* Mann, 1923
Diagnosis of the *Leptanilloides legionaria* species-group

Species from the *Leptanilloides legionaria* species-group possess the characters of the genus (see diagnosis in Borowiec & Longino 2011) but are grouped because they share the following criteria: promesonotal connection at least partially fused; lateroclypeal tooth absent or reduced; postpetiolar reduced and isolated; and postpetiolar spiracle shifted posteriad on anteromedian side of the tergite.

This group includes the following four species:
*L. legionaria* Brandão et al., 1999
*L. mckennae* Longino, 2003

Key to workers of *Leptanilloides*

1. Abdominal segment III (postpetiole) in lateral view much smaller than adjoining fourth abdominal segment. Spiracle of segment III shifted posteriad on anteromedian side of tergite. Body size relatively large, HL 0.68–0.75 (Fig. 1) ………………2 (*L. legionaria* species-group)
   - Abdominal segment III (postpetiole) in lateral view nearly as high as abdominal segment IV. Spiracle of segment III situated forward on the tergite. Body size relatively small, HL 0.31–0.62 (Fig. 1) ………………………………………..5 (*L. biconstricta* species-group)

2. Abdominal segment III (postpetiole) in lateral view nearly as high as abdominal segment IV. Spiracle of segment III shifted posteriad on anteromedian side of tergite. Body size relatively large, HL 0.68–0.75 (Fig. 1) ………………2 (*L. legionaria* species-group)
   - Abdominal segment III (postpetiole) in lateral view nearly as high as abdominal segment IV. Spiracle of segment III situated forward on the tergite. Body size relatively small, HL 0.31–0.62 (Fig. 1) ………………………………………..5 (*L. biconstricta* species-group)

3. Head subquadrate, CI 85–88; lateral margin nearly straight and parallel. Propodeal declivity short and vertical, propodeum with dorsal and posterior faces clearly differentiated (Ecuador) ……… ………………………………………………………………………….. *L. nomada* Donoso et al., 2006
   - Head subrectangular, CI 75–83; lateral margin convex. Propodeal declivity usually rounded without clear distinction between dorsal and posterior faces ………………………………………..3

4. Legs shorter, HW/HTL×100 > 78. Hypostomal tooth present (Ecuador) ………………………………………………………………………………………………………………………………………………………… *L. nubecula* Donoso et al., 2006
   - Legs longer, HW/HTL×100 < 78. Hypostomal tooth absent (Costa Rica) ……………………………………………………………………………………………………………………………………….. *L. mckennae* Longino, 2003

5. Lateroclypeal tooth absent. Masticatory margin of mandible edentate (Ecuador) ……………………………………………………………………………………………………………………………………………………………….. *L. caracola* Donoso et al., 2006
   - Lateroclypeal tooth present. Masticatory margin of mandible dentate (at least minute denticles present) ………………………………………………………………………………………………………………………………………………………………..6

6. In lateral view, abdominal segment IV narrowly attached to preceding segment III and broadly to succeeding segment V, with contrast between widths of anterior and posterior articulations of segment IV in lateral view ……………………………………………………………………………………………………………………………………………………………….. *L. atlantica* Silva et al., 2013
   - In lateral view, abdominal segment IV relatively broadly attached to preceding segment III, with little contrast between widths of anterior and posterior articulations of segment IV in lateral view ………………………………………………………………………………………………………………………………………………………………..8

7. In lateral view, sternite of abdominal segment III evenly rounded, making sternal and tergal portions subequal. Petiole as long as postpetiole. Reticulation on mesopleuron and metapleuron uninterrupted. Head densely foveolate, with c. 20–30 foveolae covering a straight transverse line at head midlength. HL ≤ 0.36 (Brazil) ……………………………………………………………………………………………………………………………………………………………….. *L. biconstricta* Mann, 1823
   - In lateral view, sternite of abdominal segment III (postpetiole) distinctly bulging anteriorly, making sternal portion of segment deeper than tergite (Borowiec & Longino 2011: fig. 2d). Petiole longer than postpetiole (Borowiec & Longino 2011: fig. 2d). Reticulation on mesopleuron and metapleuron superficial and interrupted. Head sculpture less dense, with only 10–20 foveolae covering a straight transverse line at head midlength. HL = 0.49 in only measured specimen (Bolivia, Colombia, Venezuela) ……………………………………………………………………………………………………………………………………………………………….. *L. biconstricta* Mann, 1823
8. In lateral view, petiolar sternite distinctly bulging medially .................................9
   – In lateral view, petiolar sternite straight (not bulging) or bulging anteriorly ..............10

9. Hindtibia with two very small, simple spurs, without pectinate spur clearly visible under 50× magnification. Petiolar spiracle opening in an excavation distinctly larger than propodeal spiracle (Borowiec & Longino 2011: fig. 5g–h). Flange over metapleural gland opening sharply pointed posteriorly (Mexico, Guatemala) .........................L. gracilis Borowiec & Longino, 2011
   – Hindtibia with large pectinate spur, clearly discernable under 50× magnification. Petiolar spiracle not in excavation, similar and subequal to or smaller in diameter than propodeal spiracle (Borowiec & Longino 2011: fig. 4g–h). Flange over metapleural gland opening rounded posteriorly (Venezuela) ...........................................L. femoralis Borowiec & Longino, 2011

10. Head narrow, HL < 0.30, HW < 0.20, CI < 60. Head dorsum densely foveolate with foveolae separated by less than their diameter, often contiguous (Colombia) ..........................................................L. sculpturata Brandão et al., 1999
   – Head broader, HL > 0.30, HW > 0.20, CI > 65. Head dorsum less densely foveolate, with foveolae separated by about their diameter or more ..............................................................11

11. Size large, HL ≥ 0.50, HW ≥ 0.38. Flange over metapleural gland opening short, not surpassing propodeum declivity margin in lateral view, and rounded posteriorly .........................12
   – Smaller, HL < 0.45, HW < 0.35. Flange over metapleural gland opening long, surpassing propodeum declivity margin in lateral view, and sharply pointed posteriorly .................13

12. HL = 0.50 on single known specimen. Masticatory margin of mandible distinctly dentate, with regularly spaced and well-developed teeth (Brandão et al. 1999: fig. 16) (Ecuador) ......................L. improvisa Brandão et al., 1999
   – HL ≥ 0.58. Masticatory margin of mandible dentate but teeth extremely minute, blunt and irregularly shaped, difficult to discern even under magnifications of about 100× (Figs 4B, 6C) (Ecuador) ..........................................................L. prometea Delsinne & Donoso sp. nov.

13. Ventral margin of petiolar sternite forming an even convexity in lateral view. Reticulation on mesopleuron, metapleuron and lateral side of petiole uninterrupted. In dorsal view, petiole subquadrate, 67 ≤ PI ≤ 80 (Ecuador) .........................L. erinys Borowiec & Longino, 2011
   – Ventral margin of petiolar sternite relatively straight in lateral view (Fig. 3C). Reticulation on mesopleuron, metapleuron and lateral side of petiole superficial and interrupted. In dorsal view, petiole slender, rectangular, 52 ≤ PI ≤ 63 (Fig. 3D) (Ecuador) ..........................................................L. copalinga Delsinne & Donoso sp. nov.

Leptanilloides copalinga Delsinne & Donoso sp. nov.

Figs 2A–F, 3A–F

Diagnosis

Leptanilloides copalinga sp. nov. belongs to the L. biconstricta species-group. It can be distinguished from other species of this group by the combination of the following characters: masticatory margin of mandible feebly dentate, with teeth extremely minute, blunt and irregularly shaped (Fig. 3A); lateroclypeal tooth present (difficult to observe in stereo microscopy even at high magnification due to the small size of the species but obvious in SEM; Fig 3A); head with piligerous foveolae separated by smooth interspaces equaling 2–3 diameters; reticulation on mesopleuron, metapleuron and lateral side of petiole superficial and interrupted; flange over metapleural gland opening conspicuous and sharply
pointed posteriorly, resulting in a projection which surpasses propodeum declivity margin in lateral view (Fig. 3B); petiole slender in dorsal view, $52 \leq \text{PI} \leq 63$ (Fig. 3D); and subpetiolar process straight (not distinctly bulging), with a rounded projection anteriorly (Fig. 3C).

**Etymology**

Name in apposition which refers to the Copalinga Private Reserve, the type locality, in honour of its owners Catherine Vits and Boudewijn de Roover to acknowledge their efforts for the conservation of Ecuadorian biodiversity.

**Type material**

**Holotype**
ECUADOR: Worker, Zamora-Chinchipe Prov., Copalinga Private Reserve, 1510 m, 4°4’56.6” S, 78°58’5.71” W, 2 Apr. 2010, soil sample, coll. Thibaut Delsinne and Tania Milena Arias-Penna (QCAZ, sc 4006304).

**Paratypes**
ECUADOR: Same data as holotype: 1 worker, gold-coated for SEM (RBINS, sc 4006301); 1 pinned (MUTPL, sc 4006303); 1 worker, 96% ethanol (RBINS, sc 4006302, specimen with DNA data). GenBank accession number: KT601697.

**Measurements (in mm) and indices**

Holotype (paratype 4006303): HW 0.33 (0.34), HL 0.44 (0.44), SL 0.23 (0.21), PrW 0.2 (0.22), WL 0.56 (0.57), MH 0.17 (0.17), PL 0.17 (0.18), PW 0.09 (0.11), PPL 0.16 (0.18), PPW 0.16 (0.18), AivL 0.26 (0.26), AivW 0.3 (0.33), FFeL 0.23 (0.27), FFeW 0.09 (0.11), HFeL 0.29 (0.29), HTL 0.3 (0.29), CI 75 (76.25), SI 70 (60.66), PI 51.61 (62.5), MI 30 (30.39).

**Description**

**Worker**

With the characters typical of the genus (see diagnosis and description in Brandão et al. 1999 and Borowiec & Longino 2011) and of the *L. biconstricta* species-group (see diagnosis of the group above). Other characters or differences are as follows:

**HEAD.** Elongate and rectangular with lateral margin nearly straight and parallel. Posterior corner rounded. Posterior margin modestly convex, almost straight. Parafrontal ridge absent. Clypeal lamella strongly convex (Fig. 3A). Basal and masticatory margins of mandible dentate but teeth extremely minute, blunt and irregularly shaped, difficult to discern even under magnifications of about 100× (Fig. 3A). Basal and masticatory margins united by a broad convexity. Basal margin faintly crenulate. Labial and maxillary palps difficult to discern in situ but formula apparently 2,2. Hypostomal anterior border without distinct tooth. Scape when laid back failing to reach medial distance to posterior margin of head by nearly one maximum diameter.

**MESOSOMA.** Flange over metapleural gland opening conspicuous and sharply pointed posteriorly, forming a projection which surpasses propodeum declivity margin (Fig. 3B). Femur not conspicuously enlarged, relatively slender. Midtibia with one short (i.e., half size of foretibia strigil), pectinate spur (Fig. 3F), although pectination may be difficult to observe even at high magnification. Hindtibia with one broadly pectinate spur roughly as long as strigil. Metatibial gland absent or very reduced and not visible even in SEM.

**METASOMA.** Long and relatively slender. In dorsal view, petiole rectangular with lateral margin slightly convex, twice as long as wide (Fig. 3D), and as long as abdominal segment III (postpetiole). Anterior
face concave, posterior face straight. In lateral view, petiole height approximately ¼ smaller than height of abdominal segment III. Petiolar tergite dome-shaped, with short anterior and posterior faces, maximum height situated in posterior half. Short tubulated portion present posteriorly. Petiolar spiracle inconspicuous, not in excavation, set near anterior rim of tergite, similar in form and slightly smaller in diameter than propodeal spiracle. Subpetiolar process with ventral margin straight to slightly concave, not distinctly bulging, anteriorly forming rounded projection (Fig. 3C). Maximum height of petiolar sternite situated medially. In dorsal view, abdominal segment III (postpetiole) trapezoid, with straight, parallel anterior and posterior faces. Posterior face twice as long as anterior face. In lateral view, tergite evenly convex, without well-differentiated posterior face. Sternite evenly rounded. In dorsal view, abdominal segments IV–VI subequal in length.

PILOSITY AND SCULPTURE. Mandible smooth and shiny, with few, scattered piligerous punctures. Head with abundant deep piligerous foveolae and smooth interspaces on average equaling two or three puncture diameters. Mesosoma and abdomen more finely and sparsely punctate. Fine reticulate sculpture present laterally on lower pronotum, mesopleuron, propodeum and petiole. Dorsal part of pronotum, mesopleuron, propodeum and petiole smooth and shiny. Body and appendages with abundant, short and subdecumbent to suberect hairs. Body color yellow to reddish (Fig. 2), with head and mesosoma darker than petiole and gaster. Legs and antennae yellowish.

Fig. 3. *Leptanilloides copalinga* Delsinne & Donoso sp. nov. Paratype worker (specimen code 4006301). **A.** Anterior part of the head in full-face view, showing vertical and fused frontal carinæ, exposed antennal socket, lateroclypeal tooth, strongly convex clypeal lamella and subtriangular mandible with minute denticles along its inner margin. **B.** Lower part of propodeum in lateral view, showing the long and sharp flange over the metapleural gland opening. **C.** Petiole in lateral view. **D.** Propodeum and petiole in dorsal view. **E.** Abdominal segment III (postpetiole) in lateral view. **F.** Mesotibial spur, short and pectinate.
**Gyne**
Unknown.

**Male**
Unknown.

**Distribution**
Only known from the type locality.

**Biology**
Unknown. The type series was found by visual search in a 15×15×15 cm soil core inspected for ants during 20 person-minutes (the soil content was examined over a white plastic board using a headlamp to facilitate ant detection), which suggests subterranean habits as observed in other *Leptanilloides* species. The habitat is a relatively well-preserved evergreen lower montane forest. Soil texture is clay loam; proportion of sand, silt and clay is 44%, 22%, and 35%, respectively; mean pH = 4 ± 0.2 SD (n = 24 soil samples).

**Remarks**
This species belongs to the *L. biconstricta* species-group. The most similar species are *L. biconstricta* and *L. atlantica*, which also possess a long and sharply pointed flange over the metapleural gland opening and a straight (not bulging) petiolar sternite with a rounded anterior projection. However, these two species have a deeper constriction between the postpetiole and abdominal segment IV. Besides, *L. copalinga* sp. nov. has a petiole roughly as long as the postpetiole (longer in *L. biconstricta*) and its postpetiolar sternite is evenly rounded (distinctly bulging anteriorly in *L. biconstricta*). Finally, *L. copalinga* sp. nov. is larger than *L. atlantica* (Fig. 1). The other species of the *L. biconstricta* species-group are easily distinguished from *L. copalinga* sp. nov. by, among other characters, the shape of their petiolar sternite (bulging medially in *L. gracilis* and *L. femoralis*; bulging anteriorly in *L. erinys*), the shape of the flange over the metapleural gland opening (short and rounded posteriorly in *L. improvisa* and *L. prometea*), the head sculpture (more densely foveolate in *L. sculpturata*) and the lateroclypeal tooth (absent in *L. caracola*).

**Leptanilloides prometea** Delsinne & Donoso sp. nov.

Figs 4A–D, 5, 6A–F, 7A–C

**Diagnosis**
*Leptanilloides prometea* belongs to the *L. biconstricta* species-group and can be distinguished by the combination of the following characters: masticatory margin of mandible dentate but teeth extremely minute, blunt and irregularly shaped (Fig. 6C); lateroclypeal tooth present (Fig. 6C); head with piligerous foveolae separated by smooth interspaces equaling, on average, puncture diameters; fine, uninterrupted reticulation on mesopleuron, metapleuron and lateral side of petiole; flange over metapleural gland opening forming a short, blunt projection (Fig. 6D); and subpetiolar process relatively straight (not distinctly bulging), without posterior angle (Fig. 6E).

**Etymology**
Name in apposition, in honour of the Prometeo initiative of the Ecuadorian government that seeks to strengthen research and knowledge transference by getting national and international experts to work together. This project is aligned to the Ecuadorian “National Development Plan for Good Living”, which has the objective, among others, of guaranteeing the rights of Nature and promoting a healthy and
sustainable environment. Furthermore, the species is the largest of the *L. biconstricta* species-group and its name nicely reminds us of ‘Prometheus’, the Titan in Greek mythology who brought fire, a symbol of enlightenment, to mankind. We hope that this new ant species will symbolize the promise of a bright future for Ecuadorian biodiversity.

**Type material**

**Holotype**
ECUADOR: Worker, Zamora-Chinchipe Prov., Reserva Biológica San Francisco (RBSF), 2070 m, 3°58’ S, 79°05’ W, 13 May 2010, within 0.5 m² of leaf litter extracted with a mini-Winkler apparatus for 96 h, coll. Thibaut Delsinne and Tania Milena Arias-Penna (QCAZ, sc 4267803).

**Paratypes (n = 67)**
ECUADOR: Same data as holotype except that specimens were captured after 48 h of Winkler extraction: 30 workers, 96% ethanol (RBINS, General Inventory Number 33044: sc 4052302 [1 specimen with DNA data], sc 4052314 [1 specimen with DNA data], sc 4052316 [21 specimens without DNA data]; BMNH [1 specimen, sc 4052317]; CASC [1 specimen, sc 4052318]; ICN [1 specimen, sc 4052319]; JTLC [1 specimen, sc 4052320]; MCZC [1 specimen, sc 4052321]; MZSP [1 specimen, sc 4052322]; QCAZ [1 specimen, sc 4052323]); 3 workers, pinned (MUTPL, sc 4052310, 4052315; RBINS, sc 4052313);

**Fig. 4.** *Leptanilloides prometea* Delsinne & Donoso sp. nov. – **A–C.** Paratype worker (specimen code 4052301). **A.** Habitus, dorsal view. **B.** Head in full-face view. **C.** Habitus, lateral view. – **D.** Apex of hindtibia; reservoir of metatibial gland is visible under translucent cuticle at the base of the tibial spur (paratype, specimen code 4052313).
2 workers, 96% ethanol after non-destructive DNA extraction (RBINS, sc 4052311, 4052312); 1 worker, gold-coated for SEM (RBINS, sc 4052301). – Same data as holotype except 21 Mar. 2010, within upper 5 cm of organic layer of a core (Ø 5 cm) extracted by heat using a modified high gradient extractor for 4 days: 28 workers, 96% ethanol (RBINS, sc 4060601); 2 workers, 96% ethanol after non-destructive DNA extraction (RBINS, sc 4060602, 4060603); 1 worker, pinned (MUTPL, sc 4060604). GenBank accession numbers: KT601698–KT601703 and KT750331.

Measurements (in mm) and indices
Holotype (3 paratypes): HW 0.4 (0.4–0.44), HL 0.58 (0.61–0.62), SL 0.27 (0.29), PrW 0.27 (0.29), WL 0.71 (0.76–0.78), MH 0.23 (0.24), PL 0.22 (0.22–0.23), PW 0.11 (0.12), PPL 0.22 (0.22–0.23), PPW 0.2 (0.22–0.23), AivL 0.31 (0.31–0.33), AivW 0.37 (0.38–0.4), FFeL 0.38 (0.38–0.4), FFeW 0.11 (0.11), HFeL 0.44 (0.44–0.45), HTL 0.42 (0.44–0.46), CI 69.23 (65.45–71.43), SI 66.67 (66.25–72.22), PI 50.00 (52.5–53.66), MI 32.81 (31.43–32.36).

Description

Worker
With the characters typical of the genus (see diagnosis and description in Brandão et al. 1999 and Borowiec & Longino 2011) and of the L. biconstricta species-group (see diagnosis of the group above). Other characters or differences are as follows:

HEAD. Elongate and rectangular with lateral margin nearly straight and parallel. Posterior corner rounded. Posterior margin modestly convex, almost straight. Parafrontal ridge absent. Clypeal lamella

![Habitus in lateral view of paratype (specimen code 4060601) of Leptanilloides prometea Del-sinne & Donoso sp. nov. Note the rounded anterior projection of the subpetiolar process, which differs from the acute shape observed in the specimen on Figs 4C and 6E.](image)
strongly convex (Fig. 6C). Masticatory margin of mandible dentate but teeth extremely minute, blunt and irregularly shaped, difficult to discern even under magnifications of about 100× (Fig. 6C). Basal and masticatory margins united by a broad convexity. Basal margin faintly crenulate. Labial palp not readily visible but, at least, one-segmented, maxillary palp two-segmented. Hypostomal anterior border rounded and slightly projected outward but without forming a distinct tooth (Fig. 6A). Scape when laid back reaching about medial distance to posterior margin of head.

Fig. 6. Leptanilloides prometea Delsinne & Donoso sp. nov. Paratype worker (specimen code 4052301). A. Head, ventral view. B. Mesotibial pectinate spur. C. Anterior part of the head in full-face view, showing vertical and fused frontal carinae, exposed antennal socket, well-developed lateroclypeal tooth, strongly convex clypeal lamella and subtriangular mandible with minute denticles along its inner margin. D. Lower part of propodeum in lateral view, showing the flange over the metapleural gland opening. E. Petiole in lateral view. F. Simple claw (hindleg).
MESOSOMA. Flange over metapleural gland opening conspicuous and posteriorly forming short, blunt projection, not surpassing propodeum declivity margin (Fig. 6D). Femur not conspicuously enlarged, relatively slender. Midtibia with one short (i.e., half size of foretibia strigil), pectinate spur (Fig. 6B), although pectination may be difficult to observe even at high magnification. Hindtibia with one broadly pectinate spur roughly as long as strigil. Metatibial gland present, visible at high magnification and good lighting as translucent oval area at apex of tibia, behind spur insertion (Fig. 4D). Metatibial gland pore plate observable in SEM (Fig. 7A–C).

METASOMA. Long and relatively slender. In dorsal view, petiole uniformly rectangular, twice as long as wide, as long as abdominal segment III (postpetiole). Anterior face concave, posterior face straight. In lateral view, petiole height approximately ¼ smaller than height of abdominal segment III. Petiolar tergite dome-shaped, with short and poorly differentiated anterior and posterior faces, maximum height situated in posterior half. Short tubulated portion present posteriorly. Petiolar spiral inconspicuous, not in excavation, set near anterior rim of tergite, similar in form and slightly smaller in diameter than propodeal spiral. Subpetiolar process with ventral margin relatively straight (not distinctly bulging), and without posterior angle (Fig. 6E). Anterior projection of subpetiolar process variable in shape, acute to rounded (Figs 5, 6E). Maximum height of petiolar sternite situated in its anterior half. In dorsal view, abdominal segment III (postpetiole) trapezoid, with straight, parallel anterior and posterior faces. Posterior face almost twice as long as anterior face. In lateral view, tergite evenly convex, without well-differentiated posterior face. Sternite evenly rounded, slightly bulging anteriorly. In dorsal view, abdominal segments IV–VI subequal in length.

PILOSITY AND SCULPTURE. Mandible smooth and shiny, with few scattered piligerous punctures. Head with abundant deep piligerous punctures and smooth interspaces on average equaling puncture diameter, except on ventral side and front where punctures are sparser, separated by more than their diameter. Mesosoma and abdomen more finely and sparsely punctate. Fine reticulate sculpture present laterally on lower pronotum, entire mesopleuron, propodeum, and petiole. Body and appendages with abundant, short and subdecumbent to suberect hairs. Body color brownish to reddish, with head and mesosoma tending to be darker than petiole and gaster. Legs and antennae yellowish.

Gyne
Unknown.

Male
Unknown.

Distribution
Only known from the type locality.

Biology
Unknown. All the specimens were collected from a single reserve bordering the Podocarpus National Park on the eastern Andean slope of southern Ecuador. The habitat is an evergreen lower montane forest (Homeier et al. 2008) which is in nearly pristine condition. The forest harbours more than 300 tree species, with Lauraceae, Melastomataceae and Rubiaceae being the most species-rich families (Homeier et al. 2012). The mean annual temperature is c. 15 °C and mean annual precipitation is c. 2200 mm, with low seasonality (Bendix et al. 2008). Soils of the sampling area are cambisols, with a very thick (often > 50 cm) leaf litter layer (Homeier et al. 2012). Soil texture is sandy silt loam; proportion of sand, silt and clay is 41%, 52%, and 6%, respectively; mean pH = 3.2 ± 0.1 SD (n = 24 soil samples).
Remarks

*Leptanilloides prometea* sp. nov. is the largest species of the *L. biconstricta* species-group (Fig. 1). The most similar species in habitus and size is *L. improvisa*, but head of *L. prometea* sp. nov. is longer (Fig. 1).

**Fig. 7.** *Leptanilloides prometea* Delsinne & Denoso sp. nov. Scanning electron micrographs of the metatibial apex of paratype worker (specimen code 4052301), showing the metatibial gland pore plate (arrow) at different magnifications.

**Fig. 8.** *Leptanilloides* ‘Male 4’ (specimen code 4870101) from southern Ecuador. A. Head in full face view. B. Habitus in dorsal view. C. Habitus in lateral view.
and its mandible has only minute and irregular teeth along the masticatory margin (teeth are conspicuous and regularly spaced in *L. improvisa*). Moreover, *L. prometea* sp. nov. can easily be distinguished from other species in the group by: the presence of a conspicuous lateroclypeal tooth (absent in *L. caracola*), the evenly rounded sternite of abdominal segment III (bulging anteriorly in *L. biconstricta*), the petiole being as long as the postpetiole (petiole longer than postpetiole in *L. biconstricta*), the petiolar sternite being higher in its anterior half (bulging medially in *L. gracilis* and *L. femoralis*), the head dorsum with foveolae separated by about their diameter (more densely foveolate in *L. sculpturata*, with foveolae separated by less than their diameter), and the short, blunt flange over the metapleural gland opening (long and sharply pointed in *L. atlantica*, *L. copalinga* and *L. erinys*).

**Leptanilloides** sp. ‘Male 4’ (unassociated to workers)

Figs 8A–C, 9A–B, 10A–B

**Material examined**

**Measurements (in mm) and indices**

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**Description**

**Head.** Broader than long, with large convex eye that occupies anterior half of side of head. Mandible slender, falcate, twisted at apex, overlapping at closure (Fig. 8A). Basal and masticatory margins strongly concave, separated by well-developed tooth. External margin straight along basal and medial lengths but strongly bent at apex. Mandible longer than eye length. Lateroclypeal tooth and hypostomal tooth lacking, clypeus short and transverse, with narrow clypeal lamella (apron). Antennal socket horizontal and exposed, located at anterior clypeal margin, margin not projecting anteriorly beyond ventral articulation with labrum. Antenna 13-segmented, each segment longer than wide, with second segment shortest. Scape of moderate length, 1.5 times longer than length of ultimate antennal segment. Scape length nearly four times length of second antennal segment. Lateral ocellus separated from median ocellus by its diameter. Distance between lateral ocelli similar to distance between median and lateral ocellus; as a result, ocelli forming equilateral triangle.

**Mesosoma.** Pronotum U-shaped in dorsal view and reduced anteromedially to thin horizontal strip, set below level of dorsally protruding mesonotum and triangular in lateral view, with pointed posterior apex directed towards wing base. Mesoscutum lacking notaulus. Parapsidal line not discerned, apparently absent. Axillae depressed, not meeting medially, connected by narrow furrow; tegula very small and inconspicuous. Mesopleuron lacking oblique transverse sulcus and hence not divided into anepisternum and katepisternum. Transcutal sulcus deeply impressed. Mesoscutum and mesoscutellum prominently bulging, in lateral view. Metapleural gland not discernable. Propodeum rounded in profile, with dorsal and declivous faces only poorly differentiated; dorsal surface somewhat shorter than declivous. Propodeal spiral small, circular, positioned slightly above midheight of propodeum and slightly posterior to metanotum. Leg slender, mid tibia with one simple and hind tibia with one simple spur (observed under 50× magnification), pretarsal claw lacking preapical tooth.

**Wings** (Fig. 9A–B). With relatively well-developed venation (*for Leptanilloides*). Forewing with C present, tubular and pigmented. Sc+R approximated to wing margin, very narrow. Sc+R1 in line with
Fig. 9. Wings of *Leptanilloides* ‘Male 4’ (specimen code 4870101) and terminology used in the description. **A.** Forewing. **B.** Hindwing. Abbreviations: 2r-rs = second radial-radial sector crossvein, A = anal vein, C = costal vein, Cu = cubital vein, cu-a = cubital-anal crossvein, M = medial vein, M·f1 = first free abscissa of medial vein, M+Cu = fused medial-cubital veins, Pst = pterostigma, Rs+M = fused medial and radial sector veins, Rs = radial sector, Rs·f1 (to Rs·f5) = first (to fifth) free abscissa of radial sector vein, Sc+R = fused subcostal and radial veins, Sc+R1: fused subcostal and first radial veins.
Sc+R, tubular. Pterostigma well-marked. M+Cu tubular and pigmented, curved towards posterior wing margin before division. Rs·f1 nebulous. M·f1 pigmented, tubular. Rs+M, Rs·f2, and Rs·f3 all joined, tubular and pigmented. 1r-rs absent. 2r-rs present, tubular and pigmented. Rs·f4 and Rs·f5 joined and not differentiated in absence of 2rs-m. Rs·f4&Rs·f5 partly tubular, then nebulous, terminating before wing margin. M and Cu diverging at cu-a. Free abscissa of M partly nebulous, then spectral, very weakly visible, joining to Rs+M&Rs·f2&Rs·f3. Abscissa of Cu joined, nebulous throughout most of length and continuing as spectral. Vein A tubular, joining cu-a at obtuse angle and confluent with Rs+M, apparently absent beyond cu-a. Posterior margin of fore wing with fold where hamuli attach, narrow, conspicuous. Hindwing with C present, narrow and faint. Anterior hindwing margin with small differentiated pigmentation area located in distal half of wing. Three hamuli originate in pigmented region. Sc+R present, nebulous, almost reaching third of wing length. Sc+R1 absent. Rs·f1&Rs·f2 partly nebulous, then spectral. Jugal lobe absent.

METASOMA. Slender in lateral view, obovate in dorsal view, widest at abdominal segment V. Petiole (abdominal segment II) longer than high or wide, rectangular in dorsal view, with convex lateral margin, ovanate in lateral view, and weakly constricted posteriorly, helcium thus apparently quite broad. Petiolar spiracle located on anterior third of segment, near anterodorsal extremity. Abdominal segment III three times larger than petiole at its maximum width, not developed as postpetiole nor separated from abdominal segment IV by marked constriction. Abdominal spiracle III located on anterior third of tergite. Abdominal segments II and III with tergosternal fusion. Abdominal segment IV and succeeding segments lacking tergosternal fusion. Spiracle present on anterior third of tergite IV. Abdominal segments V and VI not separated from succeeding segments by constrictions. Abdominal spiracles V and VI indiscernible. Abdominal tergite VIII (pygidium) small and simple but visible dorsally, not wholly covered by abdominal tergite VII.

GENITALIA (Fig. 10A–B). Pygostyle absent. Abdominal sternite IX (subgenital plate) with posterior margin broadly and deeply concave but not bifurcate. Basal ring present, not hypertrophied. Paramere relatively small, harpago rounded at apex; paramere shorter than petiole length. Volsella a simple, broad lobe, lacking differentiated cuspis. Aedeagus subequal in length to paramere and volsella, simple, narrow, distally spatulate.

PILOSITY AND SCULPTURE. Integument mostly smooth and shiny, with scattered piligerous punctures. Pilosity common on most of body, suberect to decumbent. Color light brown, head and metasoma past abdominal segment III slightly darker. Appendages (antennae, mandibles, legs) lighter than body.

Distribution

Only known from the sampling locality.

Remarks

The male (specimen code 4870101) collected in southern Ecuador is easily distinguished from any other leptanilloid males (either of Leptanilloides or Amyrmex species, males of all species of Asphinctanilloides being unknown) by the unique shape of its mandible. We did not give it a specific name because it is not associated with workers. However, we called it ‘Male 4’ to indicate that it is different from the three males (‘Male 1’ to ‘Male 3’) described in Borowiec & Longino (2011), which are also without specific names.

The wing venation of ‘Male 4’ is relatively similar to that of ‘Male 3’ in Borowiec & Longino (2011), except that the free abscissa of M on the forewing joins Rs+M&Rs·f2&Rs·f3 (not joining in ‘Male 3’) and Sc+R1 is not visible on the hindwing (present as a short nebulous stub in ‘Male 3’). ‘Males 1 and 2’ of Borowiec & Longino (2011) have more reduced venation, as do the males of Amyrmex (Ward &
Fig. 10. Genitalia of *Leptanilloides* ‘Male 4’ (specimen code 4870101). A. In posterior view. B. In ventral view.
Brady 2009). The forewing venation of *L. mckennae* and *L. nubecula* is also closely similar but, in these species, veins M and Cu diverge distal to crossvein cu-a (diverging at cu-a in ‘Male 4’). Table 1 lists informative characters to separate leptanilloid males.

**DNA results**

The COI sequences of the two new species and ‘Male 4’ differ significantly from each other (p-distances ranging from 18.8 to 21.7%) (Fig. 11). They also diverge (18–21.8%) from the other *Leptanilloides* DNA barcodes available in BOLD (6 specimens, 2 species: *L. gracilis* from the *L. biconstricta* species-group and *L. nubecula* from the *L. legionaria* species-group).

Our phylogenetic analyses constructed on the basis of the nuclear gene Wg support one clade consisting of *L. prometea* sp. nov., *L. femoralis*, *L. gracilis* and *Amyrmex* sp. and another clade composed of *L. nubecula*, *L. nomada* and *L. mckennae*. Both clades are supported by bootstrap values of 98% and 99% in the neighbour-joining tree, 97% and 100% in the most parsimonious tree, 83% and 100% in the tree with maximum likelihood and by posterior probabilities of 1 in the tree built with Bayesian inference (Fig. 12).

**Discussion**

The discovery of two new *Leptanilloides* species and a distinct male, which potentially belongs to another species, confirms assertions such as “there are many more species to be found and we have no idea how species-rich the [leptanilloid clade] may be” (Longino 2003), “new species [of *Leptanilloides*] will continue to be discovered” (Borowiec & Longino 2011), and “the ants may be much more common and widespread than presently known” (Silva *et al.* 2013). The rate of species discovery within the leptanilloid clade is indeed rapidly increasing (Fig. 13), and should continue to do so, thanks to growing interest in the subterranean ant assemblages, considered as the “final frontier” in the study of the biodiversity of Formicidae (Ryder Wilkie *et al.* 2007) and linked with the development of new

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**Fig. 11.** Neighbour-joining tree showing p-distances among DNA sequences of the mitochondrial COI barcode fragment obtained for all specimens of *Leptanilloides* Mann, 1823 sequenced here and available in GenBank and BOLD. The tree was rooted with a COI sequence of *Cylindromyrmex striatus* Mayr, 1870 (GenBank accession number AY233723). Labels provide species identifications and field IDs (in bold) or GenBank or BOLD numbers. Values at nodes indicate bootstrap support only if it was above 80%.
Table 1. List of informative characters to separate the *Leptanilloides* male collected in southern Ecuador ('Male 4') from males of *L. mckennae* Longino, 2003 (Ward 2007), *L. nubecula* (Donoso et al. 2006), three males without specific names (Borowiec & Longino 2011), and from males of the genus *Amyrmex* Kusnezov, 1953 (Ward & Brady 2009).

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<th>Mandible shape</th>
<th>External margin of mandible</th>
<th>Clypeal apron</th>
<th>HL, HW, CI</th>
<th>Parapsidal line</th>
<th>Forewing venation</th>
<th>Forewing submarginal cell</th>
<th>Veins M and Cu</th>
<th>Paramere</th>
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<tr>
<td>'Male 4'</td>
<td>falcate, twisted at apex, basal and masticatory strongly concave, separated by a well-developed tooth</td>
<td>straight along its basal and medial lengths but strongly bent at apex</td>
<td>narrow</td>
<td>0.31, 0.34, 110</td>
<td>not discernible, apparently absent</td>
<td>relatively well-developed for <em>Leptanilloides</em> (e.g., Rs·f1 tubular, M+Cu tubular, Rs·f4&amp;f5 tubular, free abscissa of M partly nebulous and partly spectral, joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>relatively short, no more than three times longer than wide, not extending distad of stigma</td>
<td>diverging at crossvein cu-a</td>
<td>shorter than petiole length</td>
</tr>
<tr>
<td><em>L. mckennae</em></td>
<td>elongate-triangular, masticatory margin edentate and rounding into unarmad basal margin</td>
<td>weakly concave</td>
<td>prominent, subtriangular, bluntly pointed medially</td>
<td>0.59-0.64, 0.45-0.47, 131-135</td>
<td>present (long, according to observations made on pictures available on AntWeb 2015a)</td>
<td>relatively well-developed for <em>Leptanilloides</em> (e.g., according to fig. 3 in Ward 2007; Rs·f1 tubular, M+Cu tubular, Rs·f4&amp;f5 tubular, free abscissa of M nebulous, joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>relatively short, no more than three times longer than wide, not extending distad of stigma</td>
<td>diverging distal to cu-a by a distance greater than the length of the crossvein</td>
<td>longer (about 1.5×) than petiole length</td>
</tr>
<tr>
<td><em>L. nubecula</em></td>
<td>falcate, without differentiated masticatory margin, edentate</td>
<td>strongly and evenly curved</td>
<td>not specified, apparently absent</td>
<td>0.32, 0.32, 100 (perhaps erroneous; see Ward &amp; Brady 2009)</td>
<td>not specified</td>
<td>not observed but, according to fig. 26 in Donoso et al. (2006), relatively well-developed (e.g., Rs·f1 tubular, M+Cu tubular, Rs·f4&amp;f5 tubular, free abscissa of M nebulous, joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>relatively short, not extending distad of stigma</td>
<td>diverging distal to cu-a by a distance shorter than the length of the crossvein</td>
<td>not specified</td>
</tr>
<tr>
<td>'Male 1' of Borowiec &amp; Longino 2011 (good candidate for male caste of <em>L. gracilis</em>)</td>
<td>falcate, without differentiated masticatory margin, edentate</td>
<td>evenly curved</td>
<td>absent</td>
<td>0.18-0.24, 0.25-0.31, 115-130</td>
<td>not discernable</td>
<td>extremely reduced (e.g., Rs·f1 absent, M+Cu nebulous and inconspicuous, Rs·f4&amp;f5 nebulous and poorly visible, free abscissa of M absent or at least nebulous and not joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>no closed submarginal cell</td>
<td>diverging at cu-a</td>
<td>a little longer than petiole length</td>
</tr>
<tr>
<td>'Male 2' of Borowiec &amp; Longino 2011</td>
<td>falcate to linear, without differentiated masticatory margin, edentate</td>
<td>straight, slightly bent at apex</td>
<td>absent</td>
<td>0.3-0.31, 0.37-0.41, 125-127</td>
<td>weakly marked but long, running about one third of mesoscutum length</td>
<td>relatively well developed (intermediate with, e.g., Rs·f1 reduced, M-Cu nebulous but conspicuous, Rs·f4&amp;f5 partly tubular and partly nebulous, free abscissa of M nebuluous, joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>partially closed; relatively short, not extending distad of stigma</td>
<td>diverging at cu-a</td>
<td>a little longer than petiole length</td>
</tr>
<tr>
<td>'Male 3' of Borowiec &amp; Longino 2011</td>
<td>elongate-triangular to sublinear, masticatory margin edentate and weakly differentiated from the unarmad basal margin</td>
<td>straight, slightly bent at apex</td>
<td>narrow</td>
<td>0.32-0.33, 0.42-0.43, 127-136</td>
<td>long, running about one third of mesoscutum length</td>
<td>relatively well-developed (e.g., Rs·f1 nebulious, M+Cu tubular, Rs·f4&amp;f5 tubular, free abscissa of M nebulious, not joining Rs+M&amp;Rs·f2&amp;Rs·f3)</td>
<td>relatively short, less than four times longer than wide, not extending distad of stigma</td>
<td>diverging at cu-a</td>
<td>shorter than petiole length</td>
</tr>
<tr>
<td><em>Amyrmex</em> spp.</td>
<td>elongate-triangular to sublinear, masticatory margin edentate and weakly differentiated from the unarmad basal margin</td>
<td>curved basally, straight medially, and bent slightly mesad at apex</td>
<td>narrow</td>
<td>HL not specified, HW 0.32-0.41; CI 125-137</td>
<td>very weakly discerned, barely discernable</td>
<td>reduced (e.g., based on Eq. 6 in Ward &amp; Brady 2009; Rs·f1 tubular, M+Cu tubular, Rs·f4&amp;f5 tubular, free abscissa of M absent)</td>
<td>elongate, four times longer than wide, extending distad of stigma</td>
<td>diverging at cu-a</td>
<td>shorter (about 0.8×) than petiole length</td>
</tr>
</tbody>
</table>
techniques designed to specifically collect this subterranean ant fauna (Weissflog et al. 2000; Berghoff et al. 2003; Ryder Wilkie et al. 2007; Brandão et al. 2008; Schmidt & Solar 2010).

Workers of *L. copalinga* sp. nov. and *L. prometea* sp. nov. presented the characteristic morphology of species from the *L. biconstricta* species-group. Interestingly, our phylogenetic analyses, albeit based on a single nuclear gene, supported the existence of the *L. biconstricta* and *L. legionaria* species-groups, and placed *L. prometea* sp. nov. within the clade composed of the *L. biconstricta* species-group, supporting the morphology-based hypothesis. Our results also suggest that COI can be used as a molecular tool to identify species of *Leptanilloides*, since large divergences were observed at COI among specimens that were assigned to different species on the basis of morphology. Unfortunately, these taxa are currently known from a limited number of workers and are not all represented in the library of DNA barcodes. Therefore, we could not compare the new sequences with those of all other species in the genus. It remains necessary to accumulate information concerning the leptanilloid clade to facilitate male-worker caste association and to establish generic limits with confidence. This case study highlights the interest in building a comprehensive reference library of COI barcodes and other DNA markers.

The characters of the *L. legionaria* species-group are reminiscent of the three currently known species of *Asphinctanilloides* (Longino 2003; Donoso et al. 2006; Borowiec & Longino 2011) [two workers of a fourth species were recently discovered but have not yet been formally described (Silvestre et al. 2012; Silvestre pers. comm.)]. However, *Asphinctanilloides* does not exhibit constrictions (cincti) between the gastral segments, a condition considered as plesiomorphic within the leptanilloid clade (Brandão et al. 1999). In addition, four sting characters were identified as apomorphies for *Asphinctanilloides*, but because the sting apparatus of only one species of the *L. legionaria* species-group has been studied so far (*L. legionaria* in Brandão et al. 1999), it is not possible to assert that these traits are absent in all species of *Leptanilloides*.

The monotypic *Amyrmex* is only known from its male (Ward & Brady 2009). The genus was initially placed in Dolichoderinae Forel, 1878, but genetic studies, based on fragments of 7–11 nuclear genes, confirmed that it is nested within *Leptanilloides*, apparently as sister of the *L. biconstricta* species-group.

![Fig. 12](image-url)

Fig. 12. Neighbour-joining tree showing p-distances among DNA sequences of the wingless nuclear marker obtained for specimens of *Leptanilloides* Mann, 1823, *Amyrmex* Kusnezov, 1953 and *Cylindromyrmex* Mayr, 1870 (as outgroup), available in GenBank and sequenced here (specimen code 4052311). Labels provide species identifications and field IDs (in bold) or GenBank or BOLD numbers. Values at nodes correspond to the bootstrap values (%) and posterior probabilities obtained in the Neighbour-joining parsimony maximum likelihood/Bayesian inference analyses. Bootstrap values < 80% and posterior probabilities < 0.95 are not indicated.
(Ward & Brady 2009; Brady et al. 2014) or even nested within this species-group (this study). It is therefore clear that generic limits within the leptanilloid clade should be reassessed (Longino 2003; Ward & Brady 2009; Borowiec & Longino 2011). However, a broad set of uncertainties hampers that task, including (1) the unknown molecular-based phylogenetic position of Asphinctanilloides, (2) the unknown worker and male caste of Amyrmex and Asphinctanilloides, respectively, (3) the poor morphological distinction between the L. legionaria species-group and Asphinctanilloides, (4) the lack of collection series for nearly all species to correctly appreciate intra- and interspecific morphological variations, and (5) the availability of molecular data for only a limited number of species of Leptanilloides.

The morphology of the two new species of Leptanilloides corresponds to the diagnosis of the genus based on the worker caste offered by Borowiec & Longino (2011), except that these species have one pectinate spur on both mid- and hindtibiae, giving the spur formula 1p,1p. The midtibial spur is small, making the pectination difficult to observe with a stereo microscope even at high magnification; however, SEM examinations were conclusive. Hitherto, the spur formula in species of Leptanilloides has been considered to be either 1s,1p or 2s,2s (Borowiec & Longino 2011). We had the opportunity to study paratypes of L. nomada (QCAZ I 119692) and L. nubecula (QCAZ I 119693, QCAZ I 119694). The three specimens possess a single short, but pectinate midtibial spur. This character also seems to be present in L. atlantica (Silva et al. 2013: fig. 2d). It could be interesting to reassess this criterion in previously described species of Leptanilloides and Asphinctanilloides, in order to evaluate its taxonomic value.

A metatibial gland was detected in L. prometea sp. nov. (Fig. 7). In stereo microscopy, the gland reservoir is visible under the translucent cuticle at the base of the hindtibial spur, but observing it requires good

---

**Fig. 13.** Cumulative number of described species of Leptanilloides Mann, 1823 over time, since the description of L. biconstricta Mann, 1823.
lighting. In SEM, the gland area appears as an irregular patch of porous cuticle, contrasting with the smoother surrounding surface, as in *L. erinys* of the *L. biconstricta* species-group and *L. nubecula* of the *L. legionaria* species-group (Borowiec & Longino 2011). We were not able, however, to discern a metatibial gland in *L. copalinga* sp. nov.

As already noted by Donoso et al. (2006), the mandibles of *Leptanilloides* males vary greatly in shape, ranging from elongate-triangular to falcate (sickle-shaped), and with basal and masticatory margins ranging from undifferentiated to forming two distinct concavities. In *Eciton* Latreille, 1804 and *Dorylus* Fabricius, 1793, two other genera of Dorylinae, males use their sickle-shaped mandibles to grab the dichthadiigyne at her petiolar horns during copulation (Kronauer & Boomsma 2007). In *Leptanilloides*, the gyne caste is known for only two species, *L. nubecula* (Donoso et al. 2006) and *L. erinys* (Borowiec & Longino 2011). In both species, the gyne is subdichthadiiform with an enlarged petiole but without the dorsal or lateral horns typical of *Eciton* and *Dorylus* gynes. It is reasonable to expect that most males of *Leptanilloides* also use their elongate mandibles to grab the female behind the petiole during copulation. However, the variety of mandibular shapes also suggests that males may display different reproductive behaviors (Shik et al. 2013).

The recent discovery of *L. atlantica* in the Brazilian Atlantic Forest considerably expanded the geographic range of the genus (Silva et al. 2013). Nevertheless, it seems that the highest diversity of *Leptanilloides* occurs in the Andes, especially at high elevations. In this respect, the montane forests and páramos of Ecuador are noteworthy, since half of the currently known species of *Leptanilloides* (i.e., 7 out of 14) were collected in these habitats. Unfortunately, Andean ecosystems are currently being destroyed at an alarming rate (Goerner et al. 2007; Tejedor Garavito et al. 2012; Thies et al. 2014; Curatola Fernández et al. 2015). To determine whether the observed distribution of species of *Leptanilloides* is a mere sampling artifact or an actually existing pattern (Silva et al. 2013), it would be necessary to (1) carry out more inventories of the subterranean ant fauna in the Andes and elsewhere, and (2) secure the conservation of Andean habitats.

**Note added in press**

During the editing process, a 15th species of *Leptanilloides*, namely *L. chihuahuaensis* MacGown, Schiefer & Branstetter, 2015, was described based on six males from western Texas, USA. This species extends the range of the genus 2500 km to the north and indicates that it possesses a larger ecological niche than suspected, since it can also be found in xeric ecosystems. The COI sequence of *L. chihuahuaensis* is distant by more than 10 % from *L. copalinga* sp. nov., *L. prometea* sp. nov. and ‘Male 4’, providing convincing support that *L. chihuahuaensis* is neither the male of *L. copalinga* or *L. prometea*, nor the same species as ‘Male 4’. Moreover, the mandible shape of ‘Male 4’ remains unique within the genus.


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References


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Appendix 1

Nexus file with all aligned sequences of the wingless gene (Wg) used in this paper and Mr Bayes command block.

```
#NEXUS
begin data;
dimensions ntax=10 nchar=412;
format missing=? gap=- datatype=dna;
matrix
4052311_L_prometea
ACTACCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
FJ588490_Amyrmex_BR01
GCTGCCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
KJ523541_L_nubecula
GCTGCCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
KJ523540_L_gracilis
GCTGCCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
KJ523539_L_femoralis
GCTGCCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
KJ523545_L_gracilis
GCTGCCGAACTTCCGTTGTCGGCAACACCTGAAAGGATCTGGTTCGACGCGCGCTCC-
CGGTTAGTGATCCAGCAATTCGACCGCCCGCATCATCGCAACGGGATTACCC-
GAACCTCGGCGACAACTTCCGTCGACCGCCCGGCGTCTCGCACGCCGGCAGCGC-
TACAATTTCCAATTAACCCGTAAATTCGAGCAGCAGACCGCCCGGTCCAGGACCT-
CCTCTACGTGGAACCCGTACCGCCGGCTTCTGAGAAGAGAACCCGAAATTCCGATCCTCG-
CAGCGACCGGCGCGATTCGGCAGATGTCGACGCTGCGACCTGTGCTGCGGCAGGGACGACATG-
GTCGCGCGAGGCTACAAAGACCCAGAGGTGATTACGATGACGGCAGGGTACGCTGACCGCACC
```
TACAATTTCACAATCGAAGCCGACAACTCGAAGCCGACAGCAAGCAAGCACAGAGGAGGAGGACCTGC
GTGAGCTGAGCAGCAGGAGCAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGGAGG
DQ353106_L_nomada
GCTGCAGCAACTTCGCCTCAGTGGGCGACAAACCTGAAGGACCGGTTTCGACGGAGCCTGACG
AY867429_L_mckennae
GCTGCCGAACTTCCCGGTCTGCGGCGACAAACCTGAAGGACCGGTTTCGACGGAGCCTGACG
KJ523535_C_meinerti
GCTGCCCAACTTCGCCGTCTGCGGCGACAAACCTGAAGGACCGGTTTCGACGGAGCCTGACG
AY233640_C_striatus
ACTGCCGAACTTCCCGGTCTGCGGCGACAAACCTGAAGGACCGGTTTCGACGGAGCCTGACG
AY867426_C_striatus
ACTGCCGAACTTCCCGGTCTGCGGCGACAAACCTGAAGGACCGGTTTCGACGGAGCCTGACG
Appendix 2

R script used for the parsimony analysis.

```r
library(ape)
library(phangorn)
dna<-read.dna("input.fasta","fasta")
dnaphy<-as.phyDat(dna)
tre.ini<-nj(dist.dna(dna,model="raw"))
parsimony(tre.ini, dnaphy)
tre.pars<-optim.parsimony(tre.ini, dnaphy)
root(tre.pars,out=10)
tre.Ratchet<-pratchet(dnaphy, trace = 0)
tres.pars<-bootstrap.phyDat(dnaphy, pratchet, trace=0)
tre.Ratchet<-acctran(tre.Ratchet, dnaphy)
plotBS(tre.Ratchet, tres.pars)->treB.pars
write.tree(treB.pars, "RParsimony.phy")
```
Appendix 3

Configuration file used for the maximum likelihood analysis using GARLI ver. 2.0 (Zwickl 2006).

[generic]

datafname = wghok.nex
constraintfile = none
streefname = stepwise
attachesperX = 50
ofprefix = MLgarlibest
randseed = -1
availablememory = 512
logevery = 10
saveevery = 100
refinestart = 1
outputeachbettertopology = 0
outputcurrentbesttopology = 0
enforcertermsconditions = 1
genthreshforpoterm = 5000
scorethreshforpoterm = 0.001
significanttopochange = 0.01
outputphylibptree = 0
outputmostlyuselessfiles = 0
writecheckpoints = 0
restart = 0
outgroup = 10
resampleproportion = 1.0
insoftfprobs = 0
outputsitelikelihoods = 0
outputinputonly = 0
collapsebranches = 1

[searchreps = 5]
bootstrapreps = 500
linkmodels = 0
subsetspecificrates = 1

[model1]
datatype = nucleotide
ratematrix = ( 0 0 0 0 0 0 )
statefrequencies = estimate
ratehetmodel = none
numratecats = 1
invariantsites = none

[model2]
datatype = nucleotide
ratematrix = ( 0 0 0 0 0 0 )
statefrequencies = estimate
ratehetmodel = none
numratecats = 1
invariantsites = none

[generic]

datafname = wghok.nex
constraintfile = none
streefname = stepwise
attachesperX = 50
ofprefix = MLgarlibest
randseed = -1
availablememory = 512
logevery = 10
saveevery = 100
refinestart = 1
outputeachbettertopology = 0
outputcurrentbesttopology = 0
enforcertermsconditions = 1
genthreshforpoterm = 5000
scorethreshforpoterm = 0.001
significanttopochange = 0.01
outputphylibptree = 0
outputmostlyuselessfiles = 0
writecheckpoints = 0
restart = 0
outgroup = 10
resampleproportion = 1.0
insoftfprobs = 0
outputsitelikelihoods = 0
outputinputonly = 0
collapsebranches = 1

[searchreps = 5]
bootstrapreps = 500
linkmodels = 0
subsetspecificrates = 1

[model1]
datatype = nucleotide
ratematrix = ( 0 0 0 0 0 0 )
statefrequencies = estimate
ratehetmodel = none
numratecats = 1
invariantsites = none

[model2]
datatype = nucleotide
ratematrix = ( 0 0 0 0 0 0 )
statefrequencies = estimate
ratehetmodel = none
numratecats = 1
invariantsites = none

[startoptprec = 0.5]
minoptprec = 0.01
numberofprecreductions = 10
treerejectionthreshold = 50.0
topweight = 1
modweight = 0.05
brlenweight = 0.2
randnniweight = 0.1
randsprweight = 0.3
limsprweight = 0.6
intervallength = 100
intervalstostore = 5
meanbrlennmuts = 5
gammashapebrlen = 1000
gammashapemodel = 1000
uniqueswapbias = 0.1
distanceswapbias = 1.0