A new species of *Stenoptilia* Hübner (Lepidoptera: Pterophoridae) associated with *Neobartsia peruviana* (Orobanchaceae) in the Andes of northern Chile

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ABSTRACT

The plume moth genus *Stenoptilia* Hübner, [1825] (Lepidoptera: Pterophoridae) is recorded for the first time from Chile. Adults of *Stenoptilia socoromaensis* Vargas & Gielis sp. nov. from the northernmost part of the Chilean Andes are described and illustrated. The larvae of *S. socoromaensis* feed on buds, flowers and unripe fruits of the hemiparasitic plant *Neobartsia peruviana* (Walp.) Uribe-Convers & Tank (Orobanchaceae). Pairwise distances of a DNA barcode sequence of *S. socoromaensis* with congeneric species ranged from 9.1 to 12.6% (K2P).

Material and methods

Collecting, rearing and morphological study

The study site is located near Socoroma village (18°16'45"S; 69°35'22"W), Parinacota Province, at about 3300 m elevation in the Andes of northern Chile. It has a tropical xeric climate with seasonal rains between December and March (Luebert and Pliscoff, 2006) and a seasonal vegetation cover with higher levels shortly after

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the rains (Muñoz and Bonacic, 2006). Larvae were collected on the hemiparasitic plant Neobartsia peruviana (Walp.) Uribe-Convers & Tank (Orobanchaceae) in May 2017. They were feeding on buds, flowers and unripe fruits. The collected larvae were placed in plastic vials with pieces of the plant and paper towel at the bottom and brought to the laboratory. The vials were cleaned periodically and fresh leaves, flowers and unripe fruits were provided until the larvae finished feeding. Vials were observed regularly after pupation. One pupa was kept in ethanol 95% until DNA extraction. The adults obtained were mounted, their abdomens were removed, cleared in hot KOH 10% for a few minutes, stained with Eosin Y and Chlorazol black and slide-mounted with Euparal. Images were captured with a Sony CyberShot DSC-HX200V and Micropublisher 3.3 RTV-QImaging digital cameras attached to a Leica M125 stereomicroscope and an Olympus BX51 optical microscope, respectively.

**DNA extraction and phylogenetic analysis**

Genomic DNA was extracted from one pupa following the procedures described in Huanca-Mamani et al. (2015). Genomic DNA was sent to Macrogen Inc. (Seoul, South Korea) for purification, PCR amplification and sequencing of the DNA barcode fragment of the cytochrome oxidase subunit 1 (sensu Hebert et al., 2003a, 2003b) with the primers LCO-1490 and HCO-2198 (Folmer et al., 1994). Amplification procedures follow the PCR program described in Escobar-Suárez et al. (2017). Sequences of 658 base pair (bp) lengths of additional species of *Stenoptilia* and one sequence of *Stenoptilodes taprobanes* (Felder & Rogenhofer, 1875) (Table 1) were downloaded from BOLD Systems (Ratnasingham and Hebert, 2007). The sequences were aligned with ClustalW in the software MEGA7 (Kumar et al., 2016). No presence of stop codons or gaps was detected. To evaluate the interspecific evolutionary distance proposed for DNA barcode (Hebert et al., 2003a, 2003b), genetic distance was assessed using the Kimura-2-parameter (K2P) model. The presence of phylogenetic signal was assessed with a substitution saturation analysis using the Xia test (Xia et al., 2003) in the Dambe 7.2.1 program (Xia, 2018). Because different evolutionary rates may occur along the marker used (e.g. Pentinsaari et al., 2016), the phylogenetic tree was inferred using a Bayesian Markov-chain Monte Carlo (MCMC) framework with a general likelihood-based mixture model of gene sequence evolution (Pagel and Meade, 2004) using the BayesPhylogenies 1.1 software (University of Reading, 2019a). The MCMC analysis was run using 300,000,000 iterations, sampling every 10,000 trees, removing the first 15% as burn-in. Finally, the consensus phylogenetic tree was visualized in BayesTrees 1.3 (University of Reading, 2019b).

**Abbreviations of institutional collections**

MNHC: Museo Nacional de Historia Natural de Santiago, Santiago, Chile
IDEA: Colección Entomológica de la Universidad de Tarapacá, Arica, Chile

### Results

*Stenoptilia socoromaensis* Vargas & Gielis sp. nov.

**Figure 1** Male holotype of *Stenoptilia socoromaensis* Vargas & Gielis sp. nov. in dorsal view. Scale bar: 1 mm.

| Species                      | BOLD accession  | GenBank accession | Country  |
|------------------------------|-----------------|--------------------|----------|
| *Stenoptilia socoromaensis*  | PBLMS2319-09    | MN847778           | Chile    |
| *Stenoptilia annadactyla*    | PHLAB792-10     | GU706533           | Germany  |
| *Stenoptilia bigoti* Gibeaux | DEEU1018-16     | HQ968800           | Italy    |
| *Stenoptilia bipunctidactyla*| ABOLA790-15     | Austria            |
| *Stenoptilia coproductyla*   | LASTSS72-14     | Austria            |
| *Stenoptilia graphodactyla*  | LEFIA025-10     | HM396375           | Finland  |
| *Stenoptilia islandica*      | PHLAB763-10     | HQ968774           | Italy    |
| *Stenoptilia lutescens*      | PHLAB601-12     | KP253705           | Austria  |
| *Stenoptilia mariaeulai*     | GRAFP298-11     | KU373285           | Greenland|
| *Stenoptilia mimula* Gibeaux | PHLA348-11      | JN772760           | Spain    |
| *Stenoptilia nokleini*       | LEFI194-11      | KT782438           | Finland  |
| *Stenoptilia pelidnoda*      | LEFGC350-10     | HM876027           | Finland  |
| *Stenoptilia pneumonanthes*  | LEEUA261-1      | JN772713           | Denmark  |
| *Stenoptilia plagiodactyla*  | PHLABH757-12    | KP253229           | Austria  |
| *Stenoptilia pterodactyla*   | FBLMS218-09     | HM901993           | Germany  |
| *Stenoptilia stigmatodactyla*| LEATC462-13     | Australia          |
| *Stenoptilia veronicae*      | LEFIA1376-10    | GU828689           | Finland  |
| *Stenoptilia zaproductyla*   | ANICG159-10     | H922425            | Australia|

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ex-larva *Neobartsia peruviana*, collected May 2017, genitalia slide HAV-1319 (MNNC).

Paratypes: Adults preserved pinned and dried. One male, genitalia slide HAV-1074, two females, genitalia slides HAV-1073, HAV-1303; same data as holotype (MNNC). Two males, genitalia slides HAV-1302, HAV-1305, one female, genitalia slide HAV-1306, same data as holotype (IDEA).

**Diagnosis**

*Stenoptilia socoromaensis* is recognized by the morphology of the genitalia. The male has a short, narrow uncus whose apex surpasses the excavation of the tegumen, juxta somewhat semicircular with thickened dorsal margin and phallus with narrow cornutus in the vesica. The female has a cone-like antrum with slightly sinuous sides and broadly excavated postero-ventral margin.

**Description**

Male (Figs. 1 and 2). Forewing length: 11.3–11.8 mm (n = 4).

Head. Vertex and frons mostly brownish grey, laterally flanked by a creamy white stripe with a few brownish yellow scales anteriorly. Antenna filiform, shortly ciliated, mostly creamy white, brownish grey scales scattered. Labial palp; first segment mostly creamy white, brownish grey dorso-apically; second segment mostly brownish grey, creamy white ventrally; third segment mostly brownish grey, creamy white dorso-apically.

Thorax. Dorsally mostly brownish grey, a slightly differentiated creamy white transverse stripe in the posterior margin of mesothorax; metathorax brownish yellow, laterally mostly brownish yellow. Foreleg mostly creamy white with brownish grey scales scattered, excepting brownish yellow coxa. Midleg and hind leg mostly creamy white with brownish grey scales scattered, one and two pairs of tibial spurs, respectively. Forewing dorsal surface mostly brownish grey with creamy white and dark brown scales scattered; a dark brown circular spot before the base of cleft; a narrow longitudinal dark brown stripe on the basal two thirds of the first lobe; a somewhat broad longitudinal brownish yellow stripe on anal margin, well differentiated basally, gradually obscured towards the base of the second lobe; fringe with brownish grey, creamy white and dark brown scales, one small dark brown blotch on anal margin of first lobe, two on distal margin of second lobe; ventral surface mostly brownish grey with creamy white scales on distal portion of the two lobes. Hindwing dorsal surface brownish grey; fringe brownish grey; ventral surface mostly brownish grey, first lobe with abundant creamy white scales scattered, third lobe mostly creamy white; two longitudinal rows of reddish brown venous scales, the anterior row longer, without scales in the medial third.

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*Figure 2* Male paratype genitalia of *Stenoptilia socoromaensis* Vargas & Gielis sp. nov. A) Male genitalia in ventral view, phallus removed. B) Phallus in lateral view. C) Uncus in detail, ventral view. D) Lobe indicated by closed arrow in A). E) Detail of the cornutus, lateral view. F) Lobe indicated by open arrow in A). G) Apex of the processes of the anellus. Scale bars: 0.2 mm.
Abdomen. Mostly yellowish brown dorsally, narrow longitudinal creamy white stripe on lateral margins of terga I-VIII, slightly differentiated creamy white transversal stripe on distal margin of terga III-VII, two dark brown spots on distal margin of terga II-VII. Mostly brownish grey latero-ventrally, slightly differentiated creamy white longitudinal stripes, creamy white and dark brown scales scattered.

Male genitalia (Fig. 2)
Tegumen bilobed, excavated sub-terminally, posterior portion folded. Uncus triangular basally, cylindrical distally, a few short setae on the two parts, apex of uncus surpasses the excavation of the tegumen. Saccus narrow, slightly thickened ventrally. Juxta somewhat semicircular, ventral margin straight, dorsal margin thickened. Anellus slightly sclerotized, lateral margin slightly differentiated; anellus arm finger-like, about 1.5 times length of uncus, a few setae at apex. Valvae symmetrical; costa straight basally, down curved distally; long hair-like setae from a small lobe near base of costa; cucullus as a downward-curved projection with narrow, rounded apex; sacculus bilobed, basal lobe triangular, short hair-like setae near proximal margin, ventral margin convex, distal lobe semicircular, about a third of basal lobe length, a small blister-like lobe on distal half bearing a few setae. Phallus strongly curved, coecum small, cylindrical, width about half that of base of phallus, vesica with narrow cornutus, length about half that of phallus.

Female. Similar to male in coloration and size.

Female genitalia (Fig. 3)
Papillae anales slightly sclerotized, long hair-like setae mostly apically, short hair-like microtrichia basally. Posterior apophyses about four times the length of papillae anales. Anterior apophyses...
absent. Antrum cone-like, slightly sinuous laterally, length about
two thirds that of posterior apophyses, tubercle-like ornamentation,
postero-ventral margin broadly excavated, a sclerotized lateral tab on
each side of ostium. Ductus bursae membranous, slightly shorter than
antrum, narrow sclerite longitudinally on central third. Corpus bursae
membranous, sub-spherical, slightly elongated, finely scobinate, two
horn-like signa with mesal margin serrated.

Geographic distribution (Fig. 4)

*Stenoptilia socoromaensis* is known from the type locality in the
neighborhood of Socoroma, Parinacota Province, northern Chile.

Host plant (Fig. 4)
The larvae of *S. socoromaensis* feed on buds, flowers and unripe fruits
of *Neobartsia peruviana*(Walp.) Uribe-Convers & Tank (Orobanchaceae).

*Neobartsia* is a South American genus of hemiparasitic plants mainly
associated with the Andes highlands (Uribe-Convers and Tank, 2016).

**Etymology**
The specific epithet is derived from Socoroma, the type locality of
*S. socoromaensis*.

DNA barcodes and phylogeny (Fig. 5)
Pairwise distances of the DNA barcode sequence of *S. socoromaensis*
(GenBank accession MN847778) with congeneric species ranged
from 9.1 to 12.6% (K2P). The lowest interspecific divergence was
between *Stenoptilia mengeli* Fernald, 1898 and *Stenoptilia islandicus***

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**Figure 4** The habitat and host plant of *Stenoptilia socoromaensis* Vargas & Gielis sp. nov. A) Habitat of *S. socoromaensis* in its type locality, near Socoroma village, Parinacota Province, at about 3400 m elevation on the Andes of northern Chile. B) The host plant *Neobartsia peruviana* at the type locality. C) Flower in detail.
Figure 5 Bayesian tree of the sequences of the DNA barcode fragment (658 bp) of the cytochrome c oxidase subunit I (COI) gene of *Stenoptilia socoromaensis* Vargas & Gielis sp. nov. (highlighted in red) and congeneric species. Numbers indicate node support (posterior probability) of branches.

Although surveys for plume moths are generally focused on the adult stage, searching for their immature stages on plants provides interesting additional information about their natural history, enabling further research involving both the plume moth and its host plants (e.g. Matthews, 2006; Vargas et al., 2018). Host plant records are known for 23 species of *Stenoptilia*, many of which feed on plants of only one or two families; the most polyphagous species is the Palearctic *Stenoptilia bipunctidactyla* (Scopoli, 1763), whose larvae feed on plants of at least nine families (Matthews and Lott, 2005). Larvae of *S. socoromaensis* were found only on *N. peruviana* at the study site, although they were also searched for on native plants of Asteraceae, Fabaceae, Malvaceae, Solanaceae, Verbenaceae and Vivianiaceae without success, suggesting a narrow host range. Accordingly, larvae of *S. socoromaensis* should be searched for throughout the geographic range of *N. peruviana*, in the highlands of the Andes of southern Peru and northern Chile, as a first step to characterize adequately the geographic range of this plume moth. The discovery of *S. socoromaensis* represents the first record of herbivory by Lepidoptera on *N. peruviana*. As the vulnerable status was recently proposed for this plant in its narrow Chilean range (Gatica-Castro et al., 2015) and the host range of *S. socoromaensis* appears to be narrow, further studies are needed to understand better the interaction between this plume moth and its host plant.

The clustering of the DNA barcode sequence of *S. socoromaensis* with congeners agrees with the morphological evidence in recognizing this species as a member of *Stenoptilia*. Morphological and molecular analysis of a greater number of species of the genus, especially the Neotropical representatives, would certainly be needed to assess the evolutionary relationships of *S. socoromaensis*. The recent discovery of many plume moths in the Neotropical Region (e.g. Landry et al., 2004; Gielis, 2006, 2011a, 2012, 2013, 2014; Matthews et al., 2012, 2019), including *S. socoromaensis* and some other representatives of *Stenoptilia*, suggests that surveys in undersampled habitats could be extremely useful to characterize better the taxonomic diversity of the genus *Stenoptilia* and the complete family Pterophoridae in the Neotropics.

**Discussion**

*Stenoptilia socoromaensis* is the first species of the genus described from Chile. The morphology suggests that *S. socoromaensis* belongs to the group of *Stenoptilia tenuis* (Felder & Rogenerofer, 1875), from Colombia, Ecuador and Peru, and *Stenoptilia suprema* Meyrick, 1926, from Bolivia, Colombia, Ecuador and Peru (Gielis, 2006). However, *S. socoromaensis* shows more black scales in the second forewing lobe. In the male genitalia of this species the valva has a longer and slender apical part; the uncus reaches well over the tegumen, in contrast to the other two species; and the anellus arms are longer than in *S. tenuis* and slightly longer than in *S. suprema*. In the female genitalia of *S. socoromaensis* the ostium bursae is wider than in the other two species; the antrum of *S. suprema* is longer and more slender, while in *S. tenuis* the antrum is slightly more slender; and the signum is much longer and more pronounced in *S. socoromaensis* than in the other two species.

Only two species of *Stenoptilia* from the Neotropics have a long, narrow cornutus in the vesica like *S. socoromaensis*: the widespread *Stenoptilia zophodactyla* (Duponchel, 1838), originally described from France and also found in the New World (Gielis, 2006), and the Neotropical *S. neblina* (Gielis, 1995, only known from Venezuela (Gielis, 1995). However, these two species can be accurately separated from *S. socoromaensis* based on the fringe pattern of the first forewing lobe, which has a single dot near the anal angle in *S. socoromaensis* and a double dot in both other species; and in the morphology of the male genitalia, *S. zophodactyla* and *S. neblina* have lateral pointed extensions near the top of the tegumen, which are absent in *S. socoromaensis*. In addition, the juxta is wider in *S. zophodactyla* and *S. neblina* than in *S. socoromaensis*. In the female genitalia, the sclerite of the ductus bursae is relatively small in *S. socoromaensis*, being restricted to its central third, while the length of the sclerite is about a half of the ductus bursae in *S. zophodactyla* and comprises almost the complete length of the ductus bursae in *S. neblina*.

(Faudsing, 1857). The BMCMC analysis separated the sequences of the species of *Stenoptilia* into two groups; the sequence of *S. socoromaensis* was placed at the base of one of them.
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Conflicts of interest

The authors declare no conflicts of interest.

Compliance with ethical standards

The specimens were collected in accordance to national legislation of Chile. Type material is deposited in public scientific collections.

Author contribution statement

HAV conducted fieldwork and dissections. CG compared morphological structures with congeneric species. MVO conducted the laboratory procedures and molecular analysis. All authors conceived the research and wrote a part of the manuscript. All authors approved the final version of the article.

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