RESEARCH ARTICLE



# A new species-group of Camptothlipsis (Braconidae, Agathidinae) from South Africa, with notes on the evolution of long mouthparts

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### Abstract

A new species-group of *Camptothlipsis* (Hymenoptera: Braconidae: Agathidinae), Lingualongis, endemic to the Western Cape of South Africa is proposed and three new species are described, i.e., *Camptothlipsis lingualongis, Camptothlipsis aagota* and *Camptothlipsis inertusursus*. Phylogenetic analyses support this as a distinct monophyletic group within *Camptothlipsis*. The phenomenon of elongate mouthparts in Hymenoptera and the unusual multiple evolution of such within the agathidines is briefly discussed.

### Keywords

Agathidinae, lingualongis, long mouthparts, systematics, phylogeny, endemism.

# Introduction

Agathidinae (Hymenoptera: Braconidae) is an unusual subfamily of hymenopterans. It is found all over the world in most terrestrial habitats, has widely variable life history traits, and has been used several times for biological control (Sharkey et al. 2006). What makes this subfamily most unusual, however, is multiple, independent evolution of elongate mouthparts. Adult mouthparts have multiple functions within Hymenoptera including: emergence facilitation, phoretic copulation, defense, host handling, and nest excavation and construction (Jervis 1998). However, the most common use of hymenopteran mouthparts is to feed on nectar (Jervis 1998, Krenn et al. 2005). Long mouthparts, used for nectar extraction, only occur in a handful of families within Hymenoptera outside of Aculeata, i.e., Tenthredinidae, Pergidae, Braconidae, and Ichneumonidae (Jervis 1998, Krenn et al. 2005). Within Braconidae, elongate mouthparts occur in few subfamilies other than Agathidinae, i.e., Braconinae, Cardiochilinae, and Cheloninae (Jervis 1998, Krenn et al. 2005). As uncommon as the trait of long mouthparts appears to be, it is remarkable that agathidines have independently evolved elongate mouthparts at least five times including *Disophrys*, *Agathis*, *Lytopylus* (formerly part of *Bassus*), *Agathirsia* and Cremnoptini (Sharkey et al. 2006, Jervis 1998).

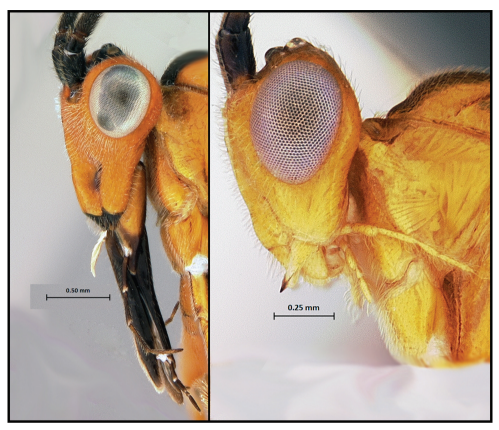
Here we report, a new species-group of *Camptothlipsis* with elongate mouthparts, the Lingualongis species-group, which is endemic to the Western Cape of South Africa (Fig. 1). Members of the Lingualongis species-group have mouthparts (both maxilla and labium) that are 1 to 1.5 times as long as their heads are high. We suggest two hypotheses in this paper to explain the extreme elongation of mouthpart in this group: 1) they are highly specialized nectar feeders and dependent on the flower of the plant on which their host lepidopterans feed or 2) that the long mouthparts facilitate the extraction of nectar from a wide variety of flowers with both shallow and deep nectaries.

### Methods

The data from the specimens used to produce the cladogram in Fig. 2, with the exception of the new species-group specimen data, are from the data sets of Sharkey et al. (2006) and Sharkey et al. (2011). The *Camptothlipsis* specimens in this analysis are a subset of the total available sequences, representing all parts of the genus, in an effort to keep the data set small and manageable for this project.

Most of the material for this study was collected in Malaise traps set up throughout the Western Cape of South Africa, along with a few specimens from sweep samples, with dates ranging from 1997 to 2009. Specimens were sent to the Sharkey Lab in Lexington, Kentucky where they were treated in hexamethyldisilazane (HMDS), mounted, and labeled. Morphological terminology follows Sharkey and Wharton (1997). Images for this study were taken with an automontage imaging system, Archimed 5.2.2, mounted on a Leica Z16 ZP0 stereomicroscope and edited using Adobe Photoshop CS4.

Regions D2–D3 of 28S rDNA were sequenced using the following primers: 28SD2hymF 5' – AGAGAGAGATTCAAGAGTACGTG – 3' and 28SD3hymR 5' – TAGTTCACCATCTTTCGGGTC – 3'. Sequences were edited using Geneious Pro v4.7.5 (Drummond et al. 2009) and aligned based on a secondary structure model for Ichneumonoidea developed by Yoder and Gillespie (2004-present) and Gillespie et al. (2005). Regions of expansion and contraction (RECs), regions of slipped-strand

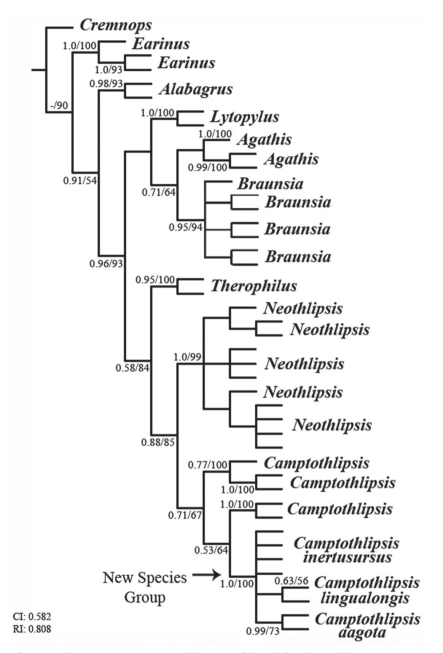


**Figure 1.** Mouthparts of *Camptothlipsis lingualongis*, a member of the species-group Lingualongis (left) compared with the mouthparts of a representative of the rest of the genus (right).

compensation (RSCs), and short regions of alignment ambiguity were further aligned/ corrected by eye (see supplementary file 1).

Phylogenetic trees were constructed using maximum parsimony and Bayesian analyses. Maximum parsimony analyses were performed using PAUP (Swofford 2002). A standard search of 1,000 replicates with 25% of trees saved was performed. The Bayesian analysis was performed using MrBayes v3.1.2 (Ronquist and Huelsenbeck 2003). Best-fitting DNA substitution models were determined using MrModeltest2.2 (Nylander 2004). The general time reversible model of evolution with a parameter for invariant sites and rate heterogeneity modeled under a gamma distribution (GTR+I+G) was determined as the best-fitting model.

Each Bayesian analysis consisted of two independent Bayesian runs initiated from different random starting trees. The analysis ran for 2,000,000 generations, reaching a topological similarity criterion of 0.005; trees were sampled every 200 generations. 25% of the trees from each run were removed as burn-in upon topological convergence.



**Figure 2.** A partial maximum parsimony cladogram of the subfamily Agathidinae. Consistency (**CI**) and retention (**RI**) index values are given at bottom left; branch support values are posterior probabilities/ bootstrap values.

See Appendix 1 for the data used to perform this analyses.

Specimen depository codes: HIC: Hymenoptera Institute Collection, University of Kentucky. SAMC: Natural History Department, Iziko South African Museum.

## Results

The maximum parsimony and Bayesian analyses produced almost identical cladograms (the *Braunsia* clade varying with more or less polytomies). Branch support values (bsv) for the phylogenetic analyses are given as Bayesian posterior probabilities and parsimony bootstrap values, respectively, in Fig. 2, e.g., 63/56. Within the Lingualongis species-group, three different species are indicated in the molecular analysis, i.e. *Camptothlipsis inertusursus, Camptothlipsis lingualongis*, and *Camptothlipsis aagota. C. aagota* is strongly supported molecularly (bsv 0.99/73), and morphological characters (deep, foveolate no-taulus, long foveolate sternaulus, deeply punctuate propodeum and granulate first tergite) suggest further evidence for the species. *C. inertusursus* and *C. lingualongis* are more weakly supported as species molecularly (bsv -/- and 0.63/56 respectively), but distinct morphological characters (very long malar space, foveolate sternaulus, rugose propodeum, and smooth first metasomal tergite, vs. short malar space, smooth notauli, smooth sternaulus, and weak propodeal texturing, respectively) suggest independent species.

The clade containing the Lingualongis species-group is strongly supported molecularly (bsv 1.0/100) and morphological characters (distinctive long mouthparts) suggest further evidence for the species-group. Phylogenetic analyses place the new species-group within the genus *Camptothlipsis* (Fig. 2). However, since the support values that suggest this placement are rather weak (bsv 0.53/64), more molecular markers are necessary to falsify the hypothesis that the Lingualongis species-group does not deserve generic status as the sister clade to *Camptothlipsis*.

While almost 60 specimens were tested for viable DNA, only eight 28S sequences were obtained. The degradation of viable DNA could be due to unfavorable conditions, e.g. extreme heat, and/or improper storage conditions before specimens were obtained by the Sharkey Lab. Although not much can be done about the former, if the later is the problem, this can be avoided in the future by proper training and instruction to those tasked with storing specimens before distribution. Due to this inability to obtain viable DNA, and thus 28S sequences, from a larger range of specimens and the high variability of morphological characteristics, only three species are described in this paper. All other potential species (we expect there are approximately 10 of these) are lumped as "undescribed species" in the key.

### Taxonomy

# Lingualongis new species-group

Figs 3.a–3.e

**Description.** *Head* (Figs 3.c - 3.e) - Lateral carina of frons absent; lateral ocellus raised; interantennal space raised, converging into a single distinct anteromedial carina; antennal depression deep; scape simple; 20–35 flagellomeres, apical flagellomere slightly rounded; gena rounded posteroventrally; malar length slightly short-

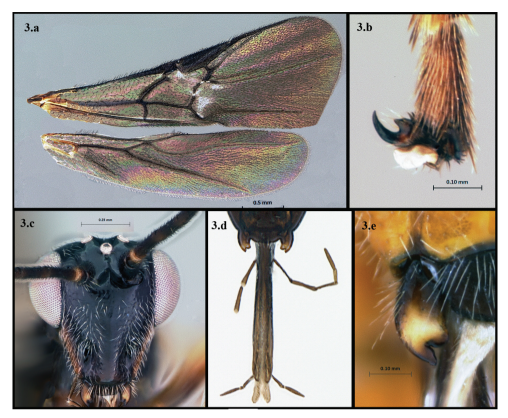


Figure 3. Lingualongis species-group a wings b hind claw c anterior head d mouthparts e mandible.

er to 1½ times as long as the height of the compound eye in lateral profile; anterior tentorial pits deep and distinct; mandible with two teeth; galea 1.0–1.5 times as long as height of head; maxillary palpus five segmented, the fourth segment somewhat reduced, length of palp as long or longer than galea; labium slightly longer than galea, and slightly longer than head to 1.5 times longer; labial palpus four segmented, third segment highly reduced.

*Mesosoma* - Subpronopes separate, and shallow; notauli present; scutellar sulcus present, with numerous longitudinal carinae; metanotum smooth or with carinae; propodeum lightly granulate to deeply pitted to rugose; mesopleuron with mesopleural groove (sternaulus) present; metapleuron slightly rugose ventrally with a rounded point anteroventrally.

*Metasoma* - Median tergite 1 lacking pair of longitudinal carinae, distal width roughly 1 to 2 times basal width; median syntergum 2 + 3 smooth, with transverse depression separating terga 2 and 3; ovipositor length as long as metasoma + most of mesosoma to as long as entire body.

*Wings* (Fig. 3.a) - Fore wing: second submarginal cell absent; m-cu broken; 3RSb straight to slightly sinuate. Hind wing: R1 and r-m present; RS nebulous or spectral and sinuate; cu-a not tubular throughout, broken; CUb present as a nebulous or spectral vein.

*Legs* (Fig. 3.b) - Tarsal claws simple with a pointed basal lobe and 3–5 basal pectines; mid tibia with apical and sometimes subapical spines; hind tibia with apical spines.

**Hosts/Biology.** Unknown. Presumably they are koinobiont endoparasitoids of lepidopteran larvae. More specifically, caterpillars of the family Gelechiidae, many of which are agricultural pests, are good candidates for this group's hosts because, to date, the only host records for members of the genus *Camptothlipsis* are on Gelechiidae larvae (Sharkey et al. 2009). Judging from the long length of their ovipositors, the group's hosts, which are probably also endemic, are most likely concealed deeply within the host plant tissue.

Distribution. South Africa, Western Cape.

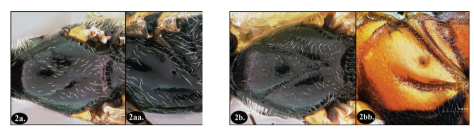
**Etymology.** From the Latin, *lingua* (tongue), and *longis* (long). The name refers to the very long mouthparts of this species-group.

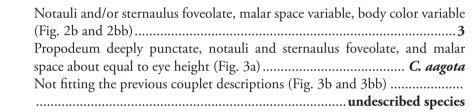
**Comments.** Within this species-group the morphology and color are highly variable. The malar length range from slightly shorter to much longer than the height of the compound eye; the anterior face of the head ranges from wider than long to longer than wide; the notauli ranges from V-shaped to Y-shaped, distinctly foveolate to completely smooth; the scutellar sulcus is partitioned into 5 to about 9 sections. These can be almost rectangular to round; the lateral metanotum ranges from smooth to ridged; the sternaulus ranges from ½ to 3⁄4 the length of the mesopleuron, straight to wavy or curved, and can be foveolate to completely smooth; the propodeal sculpture ranges from almost smooth with some slight granulation, to deeply pitted, to rugose; the first metasomal tergite ranges from square-shaped to distinctly elongate, and although the sculpture is usually granulate, it can also be slightly rugose with some specimens being completely smooth. Body length ranges from 2mm to 8mm. Color ranges from completely black to completely orange-yellow with every combination in-between. Three species are described in this paper; however, there are most likely many more species in this species-group that remain to be described.

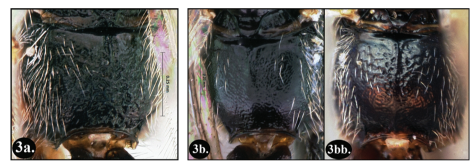
#### Key to species of the Lingualongis Species-Group

1 First metasomal tergite smooth, malar space distinctly longer than eye height; body mostly orange (Fig. 1a and 1aa)......*C. lingualongis* 









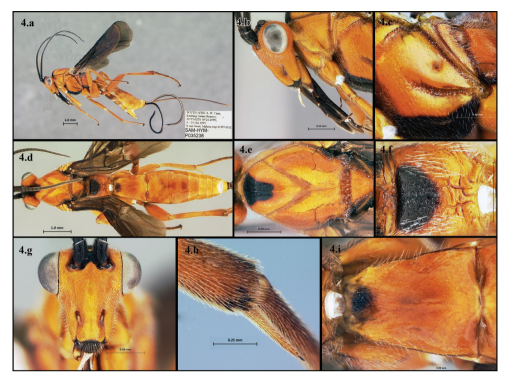
# **Species Descriptions**

Descriptions are of the holotype, with species' variation given within parentheses, e.g., 33 (31–35).

*Camptothlipsis lingualongis* Tucker & Sharkey, sp. n. urn:lsid:zoobank.org:act:D9B7511B-DC36-453A-BCE3-CD6AB0BA3BFD http://species-id.net/wiki/Camptothlipsis\_lingualongis Figs 4.a-4.i

**Diagnosis.** This is the largest and most distinctive species in the Lingualongis species-group. It is easily distinguished by its very long malar space, foveolate sternaulus, rugose propodeum, and its smooth or almost smooth first metasomal tergite

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**Figure 4.** *Camptothlipsis lingualongis* **a** lateral habitus **b** lateral head **c** mesopleuron **d** dorsal habitus **e** dorsal thorax **f** propodeum **g** anterior head **h** hind tibial spines **i** first median tergite of metasoma.

(Figs 4.a - 4.i). It is also one of the few members of the group that is all or almost all orange.

**Description.** Holotype female; length: 7.5mm (4.7–7.5mm).

*Head* (Figs 4.b, 4.g) – 33 (28–33) flagellomeres; malar length about  $1.1 \times$  eye height; glossa as long as head height (3.3 vs. 3.3mm); anterior face about as wide as high; ventral-most anterior edge of face shorter than greatest interantennal space; lightly covered with pale setae.

*Mesosoma* (Figs 4.c, 4.e, 4.f) - Median mesonotal lobe smooth except for faint beginnings of a carina anteromedially; notauli smooth, slightly extended posteriorly after joining medially; scutellar sulcus partitioned into 7 sections by longitudinal carinae; metanotum almost smooth, very faintly rugose; propodeum distinctly rugose, becoming almost areolate rugose posteromedially; sternaulus about <sup>3</sup>/<sub>4</sub> length of mesopleuron, foveolate and slightly sinuate; metapleuron with some carinae, but mostly smooth, and lightly to densely covered with pale setae.

*Metasoma* (Fig. 4.i) - First median tergite slightly longer than wide (1.2mm vs. 0.96mm), apical width 2× basal width, sculpture smooth with some very faint beginnings of rugosity basally; ovipositor as long as mesosoma + metasoma.

*Legs* (Fig. 4.h) - Mid leg with 2 apical spines; hind femur length about  $3.5 \times$  its width at widest point, hind tibia with 7 (7–10) apical spines, entire leg covered with pale setae.

*Color* - Orange-yellow except black as follows: median vertex of head, median mesonotal lobe apicomedially, posterolateral edges of mesoscutum, most of metanotum, apicomedial part (to all) of propodeum, 1<sup>st</sup> metasomal tergite basomedially, apical most part of last tergite, labrum and mouthparts, mesosternum and ovipositor sheath; antenna and wings brown, wing veins slightly darker brown.

**Etymology.** From the Latin, *lingua* (tongue), and *longis* (long). The name refers to the long mouthparts.

GenBank accession numbers. JN564494; JN564495.

**Material Examined.** HOLOTYPE: female, South Africa: Western Cape: Koeberg Nature Reserve, 33°37.62'S, 18°24.26'E, 3–31 Oct. 1997, S. van Noort, Malaise trap KO97-M12 (SAMC). PARATYPES: 1 female, South Africa: Western Cape: Kogelberge Nature Reserve, 34°16.48'S, 19°01.03'E, 16 Nov – 16 Dec 1999, S. van Noort, Malaise trap, KO98-M48 (HIC). 1 female, 2 males, South Africa: Western Cape: Mt. Rochelle, 5km S. of Franschoek, sweep, 10.i.2006, M. Buffington (HIC and SAMC) (accession number JN564495).

#### Camptothlipsis aagota Tucker & Sharkey, sp. n.

urn:lsid:zoobank.org:act:5515E4C5-1A8E-4F03-9A53-E9E716340FA8 http://species-id.net/wiki/Camptothlipsis\_aagota Figs 5.a–5.i

**Diagnosis.** This species is distinguished by its deep, foveolate notaulus, long, foveolate sternaulus, deeply punctate propodeum, and granulate first median tergite (Figs 5.a - 5.i).

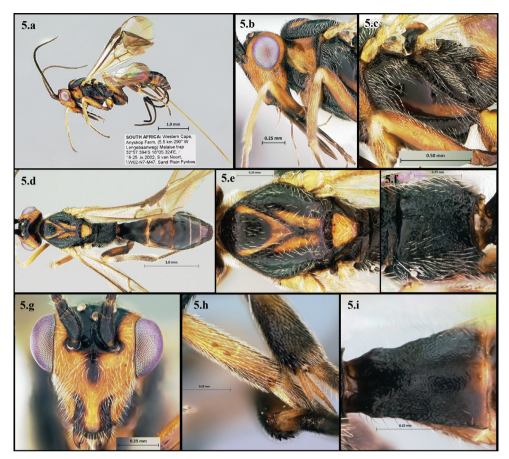
Description. Holotype female; length: 3.7mm (3.5–4.5mm)

*Head* (Figs 5.b, 5.g) – 23 (23–26) flagellomeres; malar length about 1x eye height; glossa slightly longer than head height (0.87 vs. 0.88mm); anterior face about as wide as high or slightly wider (0.83 vs. 0.80mm); ventral-most anterior edge of face shorter than greatest interantennal space; lightly covered with pale setae.

*Mesosoma* (Figs 5.c, 5.e, 5.f) - Median mesonotal lobe smooth with very faint, shallow pitting; notauli deep and foveolate (sometimes slightly extending posteriorly after joining medially); scutellar sulcus partitioned into 6 (5–9) sections by longitudinal carinae; metanotum very faintly rugose; propodeum moderately to deeply punctate; sternaulus about  $\frac{4}{5}$  ( $\frac{4}{5}$  - $\frac{3}{4}$ ) length of mesopleuron, foveolate and slightly sinuate; metapleuron  $\frac{1}{2}$  ( $\frac{1}{2}$ - $\frac{3}{4}$ ) punctate, somewhat densely covered by pale setae.

*Metasoma* (Fig. 5.i) - First median tergite slightly longer than wide (0.55 vs. 0.46mm), apical width 2x basal width, sculpture granulate; ovipositor as long as entire body.

*Legs* (Fig. 5.h) - Mid leg with 2 apical and 2 subapical spines; hind femur length about  $3 \times$  as long as wide, hind tibia with 5 (5–8) apical spines, entire leg covered with pale setae.



**Figure 5.** *Camptothlipsis aagota* **a** lateral habitus **b** lateral head **c** mesopleuron **d** dorsal habitus **e** dorsal thorax **f** propodeum **g** anterior head **h** mid and hind tibial spines **i** first median tergite of metasoma.

*Color - Head -* Orange-yellow except black as follows: vertex, spot on dorsomedial face, clypeal pit area, labrum, and mouthparts. Antenna and mandible brown. *Meso-soma -* Black except orange-yellow as follows: pronotum, edges of notauli, two thick medial longitudinal bands on mesoscutum, posteromedial scutellum, longitudinal band above sternaulus, dorsomedial spot on metapleuron. *Metasoma -* Terga mostly black with some orange-yellow highlights, ovipositor yellow, ovipositor sheath black. *Wings -* Translucent to pale yellow-brown, veins brown to light brown. *Legs -* Yellow, except brown as follows: trochanter, basal patch on hind femur, apical tibia and tarsal segments. Hind coxa mostly black.

**Etymology.** The species is named after the first author's mother, Ågot Marga. **GenBank accession number.** JN564488; JN564489.

**Material Examined.** HOLOTYPE: female, South Africa: Western Cape: Anyskop Farm, (5.5 km 290°W Langebaanweg), Malaise trap 32°57.39'S 18°05.32'E, 8–25. ix.2002, S. van Noort, LW02-N7-M47, Sand Plain Fynbos (SAMC) (accession num-

ber JN564489). PARATYPES: 2 females, same locality data as holotype with dates 9–16.x.2002 and code LW02-N7-M98 (HIC and SAMC).

# Camptothlipsis inertusursus Tucker & Sharkey, sp. n.

urn:lsid:zoobank.org:act:746C4EC3-62DD-4432-ABCC-FDDF29C77FC5 http://species-id.net/wiki/Camptothlipsis\_inertusursus Figs 6.a–6.i

**Diagnosis.** This species is distinguished by its short malar space, smooth notauli, smooth or almost smooth sternaulus, weak propodeal sculpture, and mostly black coloration (Figs 6.a - 6.i).

**Description.** Holotype female; length: 3.6mm (3.5–4.5mm).

*Head* (Figs 6.b, 6.g) - 22 (22-27) flagellomeres; malar length shorter than eye height (0.26 vs. 3.6mm); glossa longer than head height (0.80 vs. 0.70mm); anterior face about as wide as high; ventral-most anterior edge of face about same length as greatest interantennal space; lightly covered with pale setae.

*Mesosoma* (Figs 6.c, 6.e, 6.f) - Median mesonotal lobe smooth; notauli smooth, ending posteriorly where joined medially; scutellar sulcus partitioned into 7 (5–8) sections by longitudinal carinae; metanotum weakly to very weakly pitted; propodeum very weakly rugose, often appearing almost smooth, with a weak anteromedial carina; sternaulus about <sup>3</sup>/<sub>4</sub> length of mesopleuron, smooth to very weakly foveolate; metapleuron mostly smooth with some weak rugosity around the ventral edges, moderately covered with pale setae.

*Metasoma* (Fig. 6.i) - First median tergite longer than wide (0.45 vs. 0.34mm), apical width slightly less than  $2 \times$  basal width, sculpture granulate; ovipositor slightly longer than mesosoma + metasoma.

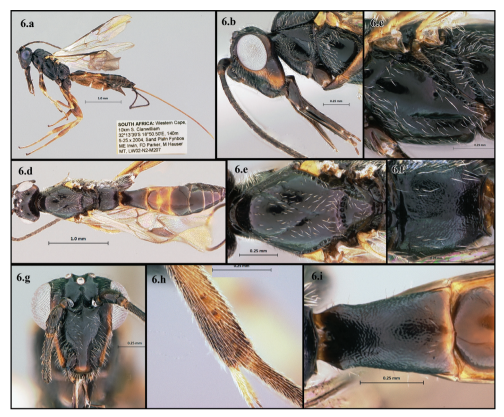
*Legs* (Fig. 6.h) - Mid leg with 3 (3–5) apical and 2 (2–3) subapical spines; hind femur length about 3x as long as wide, hind tibia with 10 (6–12) apical spines (occasionally 1 subapical spine), entire leg covered with pale setae.

*Color* - Black except yellow to orange as follows: longitudinal band on malar space, apex of antennal pedicel, base of first flagellomere, labrum, mandible, tegula, legs,  $2^{nd}$  and  $3^{rd}$  metasomal terga and ovipositor, coxa brown-black; wings translucent with brown to light brown veins.

**Etymology.** From the Latin *inertus* (lazy), and *ursus* (bear). The species is named for the first author's father, whom she thought would be amused by the name.

GenBank accession numbers. JN564490; JN564491; JN564492; JN565593.

Material Examined. HOLOTYPE: female, South Africa: Western Cape, 10km S. Clanwilliam, 32°13.39'S 18°50.50'E, 140m, 5–25.X.2004, sand plain fynbos, ME Irwin, FD Parker, M Hauser, MT, LW02-N2-M207 (SAMC). PARATYPES: 2 females, same locality and date data as holotype (HIC and SAMC) (accession number JN565593).



**Figure 6.** *Camptothlipsis inertusursus* **a** lateral habitus **b** lateral head **c** mesopleuron **d** dorsal habitus **e** dorsal thorax **f** propodeum **g** anterior head **h** mid tibial spines **i** first median tergite of metasoma.

# Discussion

This species-group has type 1 mouthparts, i.e., the glossa extracts nectar and then in retracted within the galeae, which close around it and allow the liquid to be sucked into the alimentary canal (Pucci and Sharkey 2004). The extreme elongation of mouthparts in this group suggests two possibilities: 1) they are highly specialized nectar feeders and dependent on the flower of the plant on which their host lepidopterans feed or 2) that the long mouthparts facilitate the extraction of nectar from a wide variety of flowers with both shallow and deep nectaries. The first hypothesis is supported by the biology of *Agathis malvacearum* Latreille, adults of which commonly feed on the same plant as their hosts *Metzneria lappella* Linnaeus (Lepidoptera: Gelechiidae) i.e., burdock, *Arctium minus* Bernh (Juhala 1967). The second hypothesis is supported by bees such as *Apis mellifera* Linnaeus that feed on a wide variety of plants with multiple nectary morphologies.

Our preferred hypothesis is the first and we suggest that, like *A. malvacearum*, members of the Lingualongis species-group may have an obligate and symbiotic rela-

tionship with their host flowers, in that the host plant provides nectar and host lepidopterans, and the wasp pollinates and protects the plant by keeping host lepidopteran populations in check.

The Western Cape of South Africa is a biodiversity hotspot and its long term environmental stability allows for the survival of large concentrations of relict species, such as the stag beetle *Colophon montisatris* Endrödy-Younga, and encourages organisms to radiate and differentiate morphologically (Fjeldsa and Lovett 1997, National World Heritage Serial Site 2005). South Africa also has a huge number of smaller endemic organisms including 46 invertebrate families, 150 species of Apoidea, and over 6,000 species of plants (Picker and Samways 1996, Kuhlmann 2009, National World Heritage Serial Site 2005). Due to its endemism, the species-group is undoubtedly parasitic on an endemic group of moths, which most likely feed on host plants that are also endemic.

#### Acknowledgements

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## References

- Drummond AJ, Ashton B, Cheung M, Heled J, Kearse M, Moir R, Stones-Havas S, Thierer T, Wilson A (2009) Geneious v4.7. http://www.geneious.com
- Fjeldsa J, Lovett JC (1997) Geographical patterns of an old and young species in African forest biota: the significance of specific montane areas as evolutionary centres. Biodiversity and Conservation 6: 325–346. doi: 10.1023/A:1018356506390
- Gillespie JJ, Munro JB, Heraty JM, Yoder MJ, Owen AK, Carmichael AE (2005) Secondary structural model of the 28S rDNA expansion segments D2 and D3 for chalcidoid wasps (Hymenoptera: Chalcidoidea). Molecular Biology and Evolution 22: 1593–1608. doi: 10.1093/molbev/msi152
- Jervis M (1998) Functional and evolutionary aspects of mouthpart structure in parasitoid wasps. Biological Journal of the Linnaean Society 63: 461–493. doi: 10.1111/j.1095-8312.1998.tb00326.x
- Juhala C (1967) Notes on parasitic Hymenoptera associated with a Gelechiid moth, *Metzneria lappella*, in the Common Burdock, and description of a new species of *Agathis* (Braconidae). Annals of the Entomological Society of America 60(1): 95–97.
- Krenn HW, Plant JD, Szucsich NU (2005) Mouthparts of flower-visiting insects. Arthropod Structure & Development 34: 1–40. doi: 10.1016/j.asd.2004.10.002
- Kuhlmann M (2009) Patterns of diversity, endemism and distribution of bees (Insecta: Hymenoptera: Anthophila) in southern Africa. South African Journal of Botany 75: 726–738. doi: 10.1016/j.sajb.2009.06.016

- National World Heritage Serial Site (2005) Cape floral region protected areas Western Cape & Eastern Cape provinces South Africa. United Nations Environmental Programme World Conservation Monitoring Centre.
- Nylander JAA (2004) MrModeltest v2. Program distributed by the author. Evolutionary Biology Centre, Uppsala University.
- Picker MD, Samways MJ (1996) Faunal diversity and endemicity of the cape peninsula, South Africa – a first assessment. Biodiversity and Conservation 5: 591–606. doi: 10.1007/ BF00137611
- Pucci T, Sharkey M (2004) A revision of *Agathirsia* Westwood (Hymenoptera: Braconidae: Agathidinae) with notes on mouthpart evolution. Journal of Hymenoptera Research 13: 64–107.
- Ronquist F, Huelsenbeck JP (2003) MrBayes 3: Bayesian Phylogenetic inference under mixed models. Bioinformatics 19: 1572–1574. doi: 10.1093/bioinformatics/btg180
- Sharkey MJ, Laurenne NM, Sharanowski B, Quicke DLJ, Murray D (2006) Revisions of the Agathidinae (Hymenoptera: Braconidae) with comparisons of static and dynamic alignments. Cladistics 22: 546–567. doi: 10.1111/j.1096-0031.2006.00121.x
- Sharkey MJ, Parys KA, Clutts SA (2011) A new genus of Agathidinae with the description of a new species parasitic on *Samea multiplicalis* (Guenee). Journal of Hymenoptera Research (submitted). doi: 10.3897/jhr.23.1100
- Sharkey M, Yu D, van Noort S, Seltmann K, Penev L (2009) Revision of the Oriental genera of Agathidinae (Hymenoptera, Braconidae) with an emphasis on Thailand and interactive keys to genera published in three different formats. ZooKeys 21: 19–54. doi: 10.3897/ zookeys.21.271
- Sharkey MJ, Wharton RA (1997) Morphology and Terminology. In: Wharton RA, Marsh PM, Sharkey MJ (Eds) Manual of the new world genera of Braconidae (Hymenoptera). Special publication of the International Society of Hymenopterists, Washington, DC, 19–37
- Swofford DL (2002) PAUP version 4.0b10 for Macintosh. Sinauer Associates, Inc. Publishers, Sunderland, Massachusetts.
- Yoder M, Gillespie J (2004-present) jRNA Exploring insect phylogeny using RNA secondary structure. http://hymenoptera.tamu.edu/rna

# Appendix I

Data used to perform the analyses. (doi: 10.3897/JHR.24.1909.app1). File format: NEXUS file (NEX).

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