RESEARCH ARTICLE



Redescription of two Pennellids (Copepoda, Siphonostomatoida) from Korea with a key to species of *Peniculus* von Nordmann, 1832

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Abstract

Redescriptions of two pennellid copepods, *Peniculus minuticaudae* Shiino, 1956 and *P. truncatus* Shiino, 1956, are provided, based on postmetamorphic adult females collected from marine ranched fishes captured at Tongyeong marine living resources research & conservation center, Korea. *Peniculus minuticaudae* was collected from the soft fin rays of black scraper *Thamnaconus modestus*. It can be distinguished from the other two closely related congeners *P. ostraciontis* Yamaguti, 1939 and *P. truncatus* by having a well developed triangular-shaped abdomen; the abdomen is rudimentary in other two species. This is the first report of the occurrence of *P. minuticaudae* in Korea. *Peniculus truncatus* was collected from the dorsal fin of Korean rockfish *Sebastes schlegelii*. It can be distinguished from *P. minuticaudae* by the combination of a rudimentary abdomen, long neck and setae on leg 1 and from *P. ostraciontis* by the long neck, slender trunk, and setae on leg 1. It is also shown that *P. truncatus* captured from the same host in Korea was misidentified as *P. ostraciontis* and hence, this is the second record of the occurrence of *P. truncatus* in Korea. A key is provided for the 14 nominal species of *Peniculus*.

Keywords

Copepod, pennellid, parasite, redescription, black scraper, rockfish, fins, identification, key

Introduction

The genus *Peniculus* von Nordmann, 1832 belongs to the family Pennellidae Burmeister, 1835 and contains 14 nominal species (Boxshall and Halsey 2004). Pennellids are highly transformed, often elongated copepods parasitic on marine fishes and cetaceans (Kabata 1979). Some of pennellids are ectoparasitic (e.g. *Exopenna* Boxshall, 1986; *Parinia* Kazachenko & Avdeev, 1977) but many are deeply inserted into the body of their host. The insertion can take place in the gills, the skin or in the musculature of the host without any particular preference, as is the case for the genus *Pennella* Oken, 1816 (Kabata 1981; Boxshall 1986).

Two species of *Peniculus* are redescribed from Korea in this study. They are *P. minuticaudae* Shiino, 1956 and *P. truncatus* Shiino, 1956. In Asia, nine species of *Peniculus* have so far been reported including six from India and three from Japan. The species reported from Japan are *P. minuticaudae*, *P. truncatus* and *P. ostraciontis* Yamaguti, 1939 (Shiino 1956, 1959; Yamaguti 1939, 1963). One of these three pennellids, *P. ostraciontis*, was redescribed from Korea by Choi et al. (1996) but we reveal here that theirs was a misidentification of *P. truncatus*.

Shiino (1956) described *P. minuticaudae* based on females collected from the fins of threadsail filefish *Stephanolepis cirrhifer* (Temminck and Schlegel, 1850) (= *Monacanthus cirrhifer*), from Shirahama, Wakayama Prefecture, Japan. Recently, infection of *P. minuticaudae* on two cultured fish hosts, *S. cirrhifer* and the black scraper *Thamnaconus modestus* (Günther, 1877), was reported from Oita Prefecture, Japan (Nagasawa et al. 2011), after Fukuda (1999) reported the same species from the same locality as an unidentified *Peniculus* sp.

Peniculus truncatus was also identified and described by Shiino (1956) based on a single female found on the fin ray of oblong rockfish *Sebastes oblongus* Günther (1877) [= *Sebastichthys mitsukurii*] collected off Wagu, Mie Prefecture, Japan. A third species, *P. ostraciontis*, was described based on females collected from the head of Humpback turretfish *Tetrosomus gibbosus* (Linnaeus, 1758) [= *Ostracion gibbosum*] on the Pacific coast of Japan (Yamaguti 1939). It was reported again from the triangular boxfish *Tetrosomus concatenatus* (Bloch, 1785) [= *Rhinesomus concatenatus*] from Sagami Bay by Shiino (1959) (Table 1). All three *Peniculus* species are in need of redescription and here we undertake the redescription of two of them.

The host *T. modestus* have been cultured at a few localities along the southern coastal regions of Korea. At Tongyeong marine living resources research & conservation center (TMRC), several commercially important fishes were ranched under the marine ranching program in Korea by Korea Institute of Ocean Science & Technology (KIOST) from 1998 (MOMAF 2007). Recently, we studied the symbiotic organisms associated with ranched fishes and their life cycles at TMRC (Venmathi Maran et al. 2012). The black scraper is one of the fishes that have been transferred into cages for the purpose of experimentally studying its feeding activities within this marine ranching program. The second host, *S. cirrhifer*, is uncommon in culture in Korea because of its small size

Pennellid	Host	Infected site	Host order: family	Locality	Reference
Peniculus minuticaudae Shiino, 1956	Stephanolepis cirrhifer (Temminck and Schlegel, 1850) [= Monacanthus cirrhifer]	Fins	Tetraodontiformes: Monocanthidae	Shirahama, Wakayama Prefecture, Japan	Shiino 1956
	Stephanolepis cirrhifer	Fins	Monocanthidae	Oita Prefecture, Japan	Nagasawa et al. 2011
	<i>Thamnaconus modestus</i> (Günther, 1877)	Fins	Monocanthidae	Oita Prefecture, Japan	Nagasawa et al. 2011
	Thamnaconus modestus	Fins	Monocanthidae	Tongyeong, Gyeongsangnam- do, Korea	Present study
Peniculus ostraciontis Yamaguti, 1939	Tetrosomus gibbosus (Linnaeus, 1758) [= Ostracion gibbosum]	Head	Tetraodontiformes: Ostraciidae	Pacific Ocean, Aziro, Kanagawa Prefecture, Japan	Yamaguti 1939
	<i>Tetrosomus concatenatus</i> (Bloch, 1785) [= <i>Rhinesomus concatenatus</i>]	Head	Ostraciidae	Sagami Bay, Japan	Shiino 1959
Peniculus truncatus Shiino, 1956	Sebastes oblongus (Günther, 1877) [= Sebastichthys mitsukurii]	Fins	Scorpaeniformes: Sebastidae	Off Wagu, Mie Prefecture, Japan	Shiino 1956
	Sebastes schlegelii Hilgendorf, 1880	Fins	Sebastidae	Haklim fish farm, Kamak Bay, Jeollanam- do, Korea	Choi et al. 1996
	Sebastes schlegelii	Dorsal Fin	Sebastidae	Tongyeong, Gyeongsangnam- do, Korea	Present study

Table 1. Hosts and localities of collections of Pennellids (Copepoda: Siphonostomatoida) from Korea and Japan.

and low growth rate, in contrast to Japan (Fukuda 1999). The Korean rockfish *Sebastes schlegelii* Hilgendorf, 1880 has been cultured at several localities around the southern coastal region of Korea due to its high commercial value (MOMAF 2007). Despite the increasing threat of parasites in aquaculture, information on parasites and diseases are largely lacking from farmed fishes in Korea. The redescription of *P. minuticaudae* and *P. truncatus* is necessary to reveal previously omitted or overlooked features of both species and also to correct the misidentification by Choi et al. (1996) in Korea. In addition, a key is provided for all 14 nominal species of *Peniculus*.

Materials and methods

The pennellids were carefully removed from the fin rays of the marine ranched *T. modestus* and *S. schlegelii* at TMRC, Tongyeong, Gyeongsangnam-do, Korea

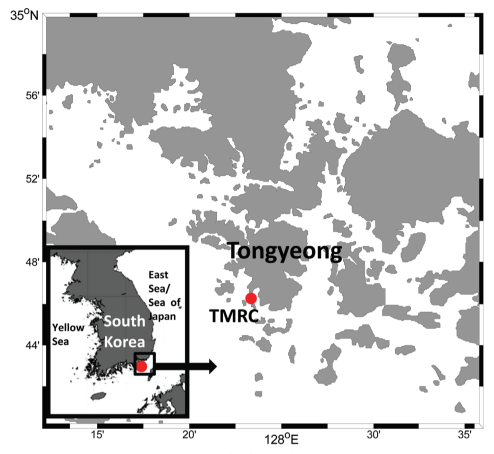


Figure 1. Map showing the marine ranched fish farming facility, Tongyeong marine living resources research & conservation center (TMRC), Tongyeong, Gyeongsangnam-do, Korea

(Figure 1) and they were preserved in 70% ethanol. Preserved copepods were cleared in a drop of 85% lactic acid or lactophenol prior to examination using an Olympus BX51 phase contrast microscope. Selected specimens were measured intact using an ocular micrometer and/or dissected and examined according to the wooden slide procedure of Humes and Gooding (1964). Measurements given are the mean followed by the range in parentheses. Drawings were made with the aid of a drawing tube. The descriptive terminology follows Kabata (1979) and the common and scientific names of host fishes follow FishBase (Froese and Pauly 2012). Voucher specimens are deposited at the National Institute of Biological Resources (NIBR), Incheon and Marine Biodiversity Institute of Korea (MABIK), Seocheon, Korea.

Results

Order Siphonostomatoida Burmeister, 1835 Family Pennellidae Burmeister, 1835 Genus *Peniculus* von Nordmann, 1832

Peniculus minuticaudae Shiino, 1956 http://species-id.net/wiki/Peniculus_minuticaudae Figures 2, 3

Peniculus minuticaudae Shiino, 1956: 593; Nagasawa et al. 2011: 43; Yamaguti 1963: 1104. *Peniculus* sp. Fukuda 1999: 57.

Material examined. 10 \bigcirc \bigcirc (NIBRIV0000245080) and 2 \bigcirc \bigcirc (MABIK CR00178439) from *Thamnaconus modestus*, Tongyeong, Gyeongsangnam-do, Korea, 20 September 2011.

Description. *Postmetamorphic adult female.* Body (Figure 2A), 2.42 (2.12–2.73) mm long (n=10) comprising oval head, slender neck, large trunk and reduced abdomen. Head (cephalothorax) ovoid, longer than wide, with blunt pointed apex (Figure 2B,C). Short slender neck (Figure 2C) consisting of three somites bearing legs 1, 2 and 3. Fourth pedigerous somite incorporated into trunk. Trunk large, cylindrical, longer than wide, bearing leg 4 proximally (Figure 2C). Abdomen slightly triangular-shaped (Figure 2D, E) long with subterminal caudal rami on ventral surface and projecting posterior tip with anal indentation. Egg sacs long and uniseriate with 33–40 eggs (Figure 2F). Caudal rami (Figure 2G) bearing 2 long, 3 medium sized subequal, 1 small setae. Antennule not observed. Antenna (Figure 2H) 2-segmented, chelate; proximal segment consisting of 2 pointed projections overlapping each other; terminal segment claw-like, acutely pointed with minute seta at base.

Mandible (Figure 3A) broad with 10 teeth terminally. Maxillule (Figure 3B) with 2 lobes having one and two long setae. Maxilla (Figure 3C) 2-segmented; proximal segment broad with spiniform small process, 2 rows of setules distally; distal segment blunt and curved with transverse striations and rows of spinules. Maxilliped absent. Legs 1 to 4 (Figure 3D–G) all represented by broad plate-like structures derived from the protopodal segments, without rami or seta. Leg 5 absent.

Variability. Some females showed variation on posterior end of trunk and abdomen (Figure 3H–J).

Attachment site. All fins of host fish.

Remarks. Careful comparison between our material and the original description of *P. minuticaudae* provided by Shiino (1956) revealed some differences: (1) the abdomen was described as trapezoid and rhomboid; (2) the striation and fine setulose ornamentation of the maxilla was not shown. The mandible was not described. Our redescription revealed that the abdomen of *P. minuticaudae* is triangular and protrudes, however, the two closely related congeners *P. ostraciontis* and *P. truncatus* both have a rudimentary abdomen. We also noted some variation in the posterior end of trunk and abdomen

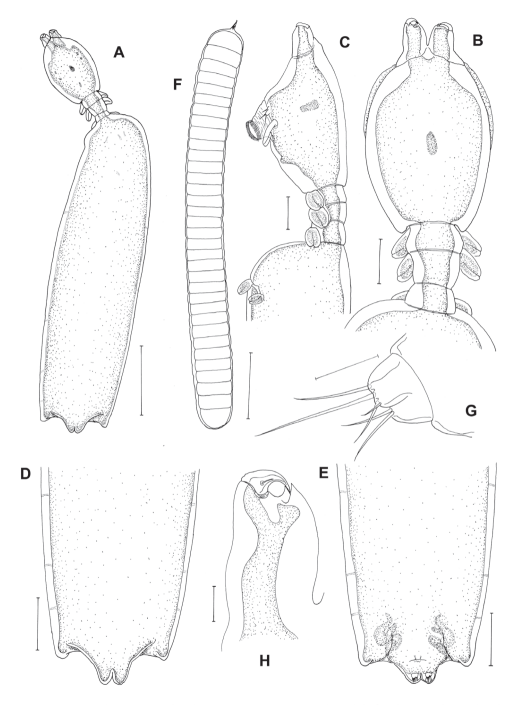


Figure 2. *Peniculus minuticaudae* Shiino, 1956. Postmetamorphic adult female. **A** Habitus, dorsal **B** Cephalothorax and free thoracic somites, dorsal **C** Cephalothorax and free thoracic somites, lateral **D** Posterior end of trunk with abdomen, dorsal **E** Posterior end of trunk with abdomen, ventral **F** Egg sac **G** Caudal ramus **H** Antenna, dorsal. Scale bars: $A=500 \mu m$; $B-F=200 \mu m$; $G=25 \mu m$; $H=50 \mu m$.

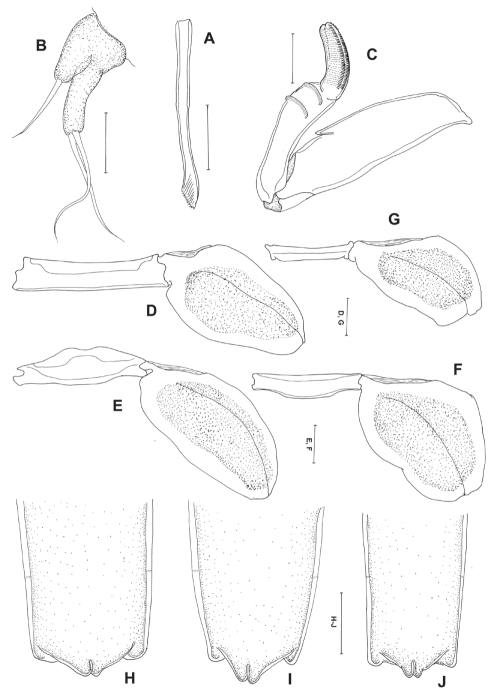


Figure 3. *Peniculus minuticaudae* Shiino, 1956. Postmetamorphic adult female. **A** Mandible, ventral **B** Maxillule, ventral **C** Maxilla, dorsal **D** Leg 1, ventral **E** Leg 2, ventral **F** Leg 3, ventral **G** Leg 4, ventral **H–J** variations of posterior end of trunk with abdomen, dorsal. Scale bars: **A–C=**25 μ m; **D–G=**50 μ m; **H–J=**200 μ m.

(Figure 3H–J). In the maxilla, fine striations and rows of setulose were found on the distal segment. In addition, the trunk is long and narrow in *P. minuticaudae* and there is no major gap between cephalothorax and trunk so it has a short neck, where legs 1 to 3 are located (Figure 2C). Leg 4 (Figure 2C) is embedded on the anterior part of the trunk. In comparison, the closely related congener *P. ostraciontis* has a stout trunk and short neck (Yamaguti 1939) while *P. truncatus* has a long trunk and neck, and leg 1 has minute setal structure which are not present in *P. minuticaudae* and *P. ostraciontis*.

Peniculus truncatus Shiino, 1956

http://species-id.net/wiki/Peniculus_truncatus Figures 4, 5

Peniculus truncatus Shiino, 1956: 593; Yamaguti 1963: 1104. *Peniculus ostraciontis*: Choi et al. 1996: 117.

Material examined. $4 \bigcirc \bigcirc$ (NIBRIV0000252624) and $1 \bigcirc$ (MABIK CR00178440) from *Sebastes schlegelii*, Tongyeong, Gyeongsangnam-do, Korea, 15 February 2012.

Description. *Postmetamorphic adult female.* Body (Figure 4A), 4.59 (4.14–5.41) mm long (n=4) comprising oval head, long slender neck, large trunk and reduced abdomen. Head (cephalothorax) ovoid, flattened dorsally but convex ventrally with pair of rounded swellings anteriorly bearing antennae (Figure 4B,C). Mouth tube prominent, directed posteroventrally (Figure 4C). Neck long (0.47–0.55 mm) (Figure 4B, C), slender, comprising about one sixth of trunk length; consisting of three somites bearing legs 1, 2 and 3 (Figure 4B, C). Fourth pedigerous somite incorporated into trunk. Trunk slender, cylindrical, longer than wide, 6 times longer than neck, bearing leg 4 proximally. Abdomen (Figure 4D), reduced with subterminal caudal rami on ventral surface. Caudal rami (Figure 4E) bearing 6 setae. Egg sacs long and uniseriate with 30-37 eggs. Antennule not observed. Antenna (Figure 4F) 2-segmented, chelate; proximal segment bearing 2 pointed projections overlapping each other; terminal segment claw-like, acutely pointed with minute seta at base. Mandible (Figure 4G) moderate-sized, broad, provided with 10 teeth terminally.

Maxillule (Figure 5A) with 2 lobes having one short and two long setae. Maxilla (Figure 5B) 2-segmented; proximal segment broad with robust spiniform process, projecting laterally, 2 rows of setules distally; distal segment blunt and curved with transverse striations and rows of spinules. Maxilliped absent. Leg 1 (Figure 5C) forming blunt plate-like structure derived from protopodal segments, with 2 minute setae laterally. Legs 2–4 (Figure 5D–F) as for leg 1, but without seta. Leg 5 absent.

Attachment site. Only on dorsal fin-rays.

Remarks. Comparison between our material and the original description of *P. truncatus* provided by Shiino (1956) revealed some omissions in that the antennae and mandibles were not shown, and possible differences, since the striation of setules on maxilla was not shown. The characteristic features of *P. truncatus* are: (1) the rudimentary abdomen; (2) the long neck (more than half as long as cephalothorax); (3) the maxilla with

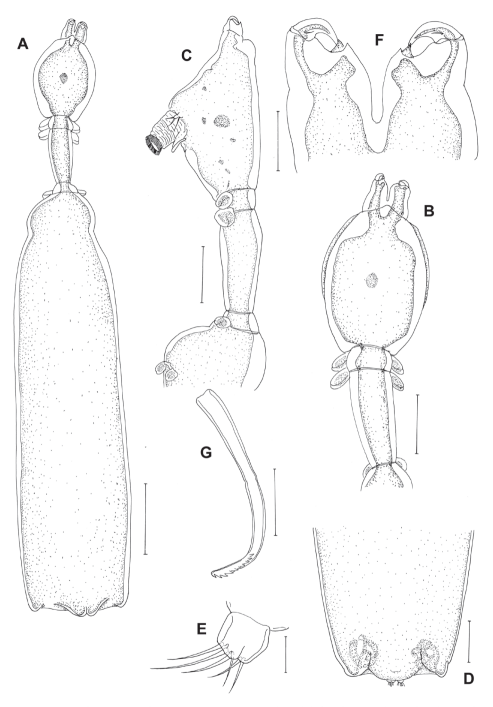


Figure 4. *Peniculus truncatus* Shiino, 1956. Postmetamorphic adult female. **A**. Habitus, dorsal **B** Cephalothorax and free thoracic somites, lateral **C** Cephalothorax and free thoracic somites, dorsal **D** Posterior end of trunk with abdomen, ventral **E** Caudal ramus, ventral **F** Antenna, dorsal **G** Mandible, ventral. Scale bars: **A**=500, μm; **B**–**D**=200 μm; **E**, **G**=25 μm; **F**=50 μm.

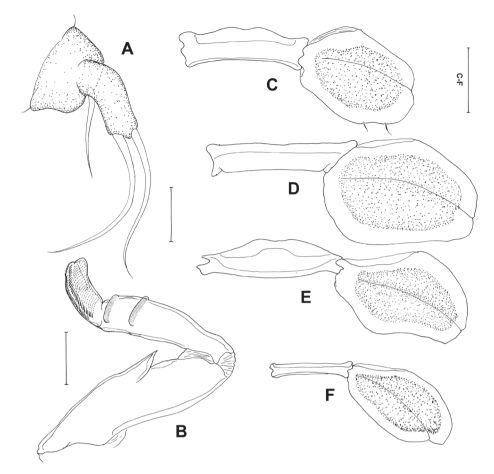


Figure 5. *Peniculus truncatus* Shiino, 1956. Postmetamorphic adult female. **A** Maxillule, dorsal **B** Maxilla, dorsal **C** Leg 1, ventral **D** Leg 2, ventral **E** Leg 3, ventral **F** Leg 4, ventral. Scale bars: **A**, **B**=25 μm; **C**-**F**=50 μm.

transverse striations of setules and rows of spinules on the distal segment; (4) the leg 1 is tipped with 2 minute setae laterally. *Peniculus truncatus* differs from *P. minuticaudae* in its rudimentary abdomen (vs. well developed abdomen); long neck (vs. short neck); and in the presence of setae on leg 1 (vs. absence of seta). It differs from *P. ostraciontis* in its moderately slender trunk (vs. stout trunk); long neck, ie: neck more than half as long as cephalothorax (vs. short neck, ie: neck less than half as long as cephalothorax); and in the presence of setae on leg 1 (vs. absence of setae) (Yamaguti 1939; Shiino 1956).

Choi et al. (1996) reported the same pennellid collected from the fins of *S. schlegelii* as *P. ostraciontis*. We compared our material with their illustrations (specimens were not deposited in the museum). It showed the features of *P. truncatus*: (1) long neck; (2) slender trunk [not as stout as like *P. ostraciontis* illustrated by Yamaguti (1939)] and the host was *S. schlegelii* (Choi et al. 1996), as in the present study.

Discussion

According to Boxshall and Halsey (2004), there are 14 species on the genus *Peniculus*: *P. asinus* Kabata & Wilkes, 1977; *P. clavatus* (Müller, 1779); *P. communis* Leigh-Sharpe, 1934; *P. elegans* Leigh-Sharpe, 1934; *P. elongatus* Boxshall, 1986; *P. fistula* Nordmann, 1832; *P. furcatus* Krøyer, 1863; *P. minuticaudae* Shiino, 1956; *P. ostraciontis* Yamaguti, 1939; *P. scomberi* Gnanamuthu, 1951; *P. stromatei* Gnanamuthu, 1951; *P. theraponi* Gnanamuthu, 1951; *P. trichuri* Gnanamuthu, 1951; and *P. truncatus* Shiino, 1956. Seven of these are reported from Asian countries.

Alexander (1983) reported *Peniculus haemuloni* from Brazil, however, it was subsequently treated as a separate genus, *Metapeniculus* Castro-Romero & Baeza-Kuroki, 1985 based on the presence of only 3 pairs of swimming legs (vs. 4 pairs for *Peniculus*) (Boxshall, 1986). According to Kabata (1979), two species of *Peniculus*, *P. calamus* Nordmann, 1864 and *P. fissipes* Wilson, 1917 should be regarded as *species inquirendae*, and in addition we treat *P. sciaenae* Gnanamuthu, 1951 as *species inquirenda* since it is also reported with 3 pairs of swimming legs (Gnanamuthu 1951a; Alexander 1983). Thus there are 14 species considered valid and a key is provided for nominal species following Alexander (1983). Most *Peniculus* species were not described adequately by modern standards and most need to be redescribed. In Asia, all species are poorly described and detailed studies are necessary for the five species reported from India (Gnanamuthu 1951a; 1951b; Pillai 1985) and for the three from Japan (Yamaguti 1939; Shiino 1956).

The mean body length of *P. minuticaudae* was 2.42 mm. It corresponds well to the body length (2.48 mm) of *P. minuticaudae* reported from Oita Prefecture, Japan (Nagasawa et al. 2011). The morphological features (Figures 2, 3) agree with the original description of *P. minuticaudae* (Shiino 1956). The present collection represents the first record of *P. minuticaudae* from ranched *T. modestus* in Korea. Thus, it is the third documented record of pennellid copepod from commercially cultured fishes.

Peniculus truncatus was originally reported from *S. oblongus* in Japan (Shiino 1956). This parasite is shown here to utilize a second host species, *S. schlegelii*, of the same host genus, although it was initially misidentified as *P. ostraciontis* by Choi et al. (1996). The misidentification was revealed by comparison between Choi's descriptions, our material and Yamaguti (1939) illustrations of *P. ostraciontis*. We collected *P. truncatus* from the same host species *S. schlegelii* cultured in Korea. The host for *P. ostraciontis* is *T. gibbosus* (Table 1). In Choi et al. (1996) redescription, they overlooked the third seta on the maxillule and the setules on the maxilla, in addition to the minute setal structures on leg 1.

Peniculus truncatus has so far been reported from two species of the genus *Sebastes*, *S. schlegelii and S. oblongus* and this pennellid appears to be host specific to rockfish (Table 1). *Peniculus minuticaudae* and *P. ostraciontis* might be specific to file fish and puffer hosts, respectively (Yamaguti 1939; Shiino 1956; 1959; Nagasawa et al. 2011; present study). A key is provided for all 14 valid species below.

Key to the species of Peniculus

(Modifi	ed from Alexander 1983)
1	Cephalothorax with 4 large holdfast processes
_	Cephalothorax without such processes
2	Cephalothorax with rounded swelling on ventral surface posterior to mouth
	tube
_	Cephalothorax without posterior swelling on ventral surface
3	Swimming legs apparently absent P. scomberi Gnanamuthu
-	Swimming legs with 4 pairs4
4	Trunk about 11 times longer than wide P. trichuri Gnanamuthu
-	Trunk about 8 times longer than wide P. stromatei Gnanamuthu
5	Legs 3 and 4 closer together than legs 1 and 2 P. communis Leigh-Sharpe
_	Legs 3 and 4 further apart than legs 1 and 2
6	Trunk conical-shaped
-	Trunk between 3 and 4.5 times longer than wide7
7	Mouth tube forming a massive posteriorly-directed proboscis
-	Mouth tube not forming a massive posteriorly-directed proboscis
8	Cephalothorax ovoid
-	Cephalothorax cylindrical
_	Cephalothorax widest near posterior margin and tapering anteriorly
0	<i>P. elegans</i> Leigh-Sharpe
9	Abdomen well developed; trunk longer than wide P. minuticaudae Shiino
-	Abdomen well developed; trunk longer than wide; with swelling on the
	head
-	Abdomen reduced; posterior margin of trunk more or less straight
10	High degree of ventral swelling on the head; neck constricted
_	Low degree of ventral swelling on the head; neck constricted
11	
11	rax
	C C
_	Trunk 3.3 times longer than wide; neck more than half as long as cephalotho- rax
	Ida F. Uruncutus Shiino

Conflict of interest statement

All authors declare that they do not have any conflict of interest.

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RESEARCH ARTICLE



A new genus and new species of Agathotanaidae (Crustacea, Tanaidacea) from West Australia

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Abstract

A new genus of Tanaidacea – *Bunburia*, collected from the region of Ningaloo in the vicinity of Bunbury (Western Australia), is erected to accommodate the new species – *B. prima* **sp. n.** This genus is classified in the family Agathotanaidae and it can be distinguished from the other members of the family by having a combination of antennulae covered with minute setae, reduced uropods and unusual setation of the propodus of pereopods 4 to 6. *B. prima* is the second species of Agathotanaidae known so far from Australia.

Keywords

Bunburia, NW Australia, Tanaidacea, Agathotanaidae

Introduction

The family Agathotanaidae erected by Lang (1971a) is represented by 41 species in four genera, namely *Agathotanais* Hansen, 1913, *Metagathotanais* Bird and Holdich, 1988, *Paragathotanais* Lang, 1971 and *Paranarthrura* Hansen, 1913 (Anderson 2012). The fifth genus, *Paranarthrurella* Lang, 1971, which had been previously considered as a member of this family (Larsen and Wilson 2002, Larsen 2005) was recently removed from Agathotanaidae (Jóźwiak et al. 2009).

The diagnosis of Agathotanaidae, modified by Larsen (2005), includes mandibles having a pointed or reduced molar, the absence of pleopods in the females and the uropod endopod that consists of one or two articles. The uropod exopods of Agathotanaidae is reduced to a blunt spur and only the presence of distal and middle setae suggests it is a fused exopod rather than a basis process. Another character that Larsen (op. cit.) has pinpointed as diagnostic for Agathotanaidae is cheliped attached directly to the cephalothorax or via pseudocoxa, however it was later questioned by Larsen (2007) and Bird (2010).

The genus *Agathotanais* is distinguishable from the other agathotanaid genera by its finely setulose body, 3-articled antennulae, reduced antennae, cheliped attached directly to the cephalothorax and uropodal endopod fused to the basis (Hansen 1913, Larsen 1999, 2005). *Metagathotanais* is the genus with strongly-reduced uropods (endopod fused with basis), but in contrast to *Agathotanais* it has fully-developed antennae that consist of six articles and 4-articled antennulae (Guerrero-Kommritz 2003). An unique feature of *Metagathotanais* is the fusion of the pleonites with the pleotelson in females, although the males retain complete pleonite segmentation. Members of *Paragathotanais* have both antennulae and antennae well-developed and the pleon with five distinct pleonites, while the uropods endopod is separated from the basis (Lang 1971b, Bird and Holdich 1988, Larsen 2005). *Paranarthrura* is the only genus of the family with uropod supported by a visible projection. The number of articles in the uropodal endopod of *Paranarthrura* can be either one or two.

The collection of Tanaidacea from the shelf and continental margin of Western Australia (WA), taken from on board the FRV *Southern Surveyor* in 2005 and 2007 has represented close to three hundred species new for science (Poore et al. in press). About 60% of the taxa are represented by single or few specimens only, while 82% of the taxa occurred just twice in the series of over two hundred samples. This scarcity of the material, together with the minute size of the specimens, which are often no longer than 1 mm, impede the formal taxonomical descriptions of the species.

One of the few species represented by more than one specimen was found to be a representative of a new genus of the family Agathotanaidae. The present paper presents the formal description of this species and the definition of the new genus that has been erected to accommodate it.

Material and methods

The material was collected in 2005 during the voyage of the FRV *Southern Surveyor* organized under the aegis of CSIRO (*Commonwealth Scientific and Industrial Research Organization*). The 14 specimens studied by us were recorded at ten of two hundred grab samples taken along the west coast of Australia from Dampier in the north to Albany in the south (from 21.0084°S, 114.381°E to 35.384°S, 118.316°E).

Appendages were dissected using chemically-sharpened tungsten-wire needles, stained with chlorazol black and mounted in glycerine. Drawings were prepared us-

ing a microscope combined with a *camera lucida* and redrawn on a digital tablet as proposed by Coleman (2003). The morphological terminology follows that proposed by Błażewicz-Paszkowycz and Bamber (2007). The body-length to width ratio was calculated using measurements from tip of the rostrum to end of pleotelson and of the widest part of cephalothorax. The ratio of particular articles was measured along their central axis. Abbreviations used in the morphological description: A1 – antennule, A2 – antenna, Mxp – maxilliped, P1-P6 – pereopods from first to sixth pair respectively.

The type material is deposited at Museum Victoria, Melbourne.

Systematics

Order Tanaidacea Dana, 1849 Suborder Tanaidomorpha Sieg, 1980 Family Agathotanaidae Lang, 1971

Bunburia gen. n.

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Diagnosis. Pleon with five free pleonites; antennulae 4-articled, with first article covered by numerous minute setae; antenna 6-articled, article 3 with dense setation; mandibles molar reduced; labium without outer or medial, setose process; maxillipedal bases unfused distally, endites unfused; epignath elongated and naked; cheliped sclerites unfused ventromedially, carpus stout (1.4 times as long as wide), chela with keel; pereopods with coxa; P1 propodus with elongate ventral seta, P1-P3 merus with long serrated seta, P4-P6 propodus with two long, serrated setae ventrodistally and three short setae dorsodistally, P5 and P6 with propodus shorter than carpus, dactylus/unguis of these pereopods setulated ventrally; ischium of all pereopods with only one seta; pleopods absent in female and well developed in male; uropod short, not projecting beyond pleotelson, with basis terminated with small projection, endopod short, one-articled.

Type species. *B. prima* sp. n. – by monotypy.

Etymology. The name refers to Bunbury, a port city near the type locality of *B. prima* sp. n.

Remarks. At first glance *Bunburia* gen. n., with its short uropods that not protrude the pleotelson, resembles the members of *Paragathotanais*. The new genus can be distinguish however from *Paragathotanais* by presence of dense setation on the proximal article of the antennulae and the fourth article of the antennae and by unusual chetotaxy of propodus of last three pereopods, which consists of three short setae dorsodistally and two long setae ventrodistally. The setation of propodus P4-P6 is variable in members of *Paragathotanais*. For example *P. abyssorum* Larsen, 2007, *P. insolitus* Guerrero-Kommritz, 2003 and *P. ipy* Jóźwiak i Błażewicz-Paszkowycz, 2011 have three long, distal setae, while five other species: *P. gracilis* Bird and Holdich, 1988, *P.* *nanus* Bird and Holdich, 1988, *P. robustus* Bird and Holdich, 1988, *P. typicus* Lang, 1971 and *P. vikingus* Bird, 2010 have three long setae in propodus of pereopods 4 and 5, but four setae in pereopod 6. Another species – *P. macrocephalus* Kudinova-Pasternak, 1986 lack of setae on propodus of pereopod 4, but it has three setae in pereopod 6. In *P. medius* Larsen, 2002 there are four long, distal setae at propodus of P4-P6. Beside the pereopods setation *Bunburia* can be separated from *Paragathotanais* by lack of medial process on the labium and bases of maxilliped unfused distally.

Larsen (2007) has pointed out that size of uropods and theirs position on the pleotelson distinguish *Paragathotanais* from *Paranarthrura. Bunburia* gen. n., with uropods similar to those observed in *Paragathotanais*, can be distinguished from *Paranarthrura* by short uropods, that are not reaching over pleotelson and are inserted more ventrally. The 4-articled antennula and the 6-articled antenna distinguish *Bunburia* from *Agathotanais*, that has 3-articled antennula and antenna reduced to one short article (Larsen 1999, 2005). An evident is also the difference between females of *Bunburia* and *Metagathotanais*, which have all pleonites fused with pleotelson (Bird and Holdich 1988, Guerrero-Kommritz 2003). Males of *Metagathotanais* have pleotelson with five distinct pleonites, but they differs from males of *Bunburia* in propodus P4-P6 chetotaxy. In *M. insulcatus* Bird and Holdich, 1988 propodus of these pereopods bears one short and three long setae distally and in *M. loerzae* Guerrero-Kommritz, 2003 there are only three long setae.

Bunburia represented by only one species is the second taxon of Agathotanaidae known so far from Australia, after *Agathotanais spinipoda* Larsen, 1999.

Bunburia prima sp. n.

urn:lsid:zoobank.org:act:ECDC9C27-D624-479B-9276-6E13A322B702 http://species-id.net/wiki/Bunburia_prima Figs 1–4

Etymology. The Latin ordinal number '*prima*' denotes the fact that the species described herein is the first member of genus *Bunburia*.

Material examined. Holotype, non-ovigerous female, J62967, 2.5 mm long, St. SS07/2005, 153, Bunbury, 33.0003°S, 114.579°E, depth 399 m, 07 August 2005.

Paratypes. 1 female dissected on slides, J62968, St. SS07/2005, 152, Bunbury, 32.9987°S, 114.576°E, depth 417 m, 2005.

1 male partially dissected, J62969, St. SS07/2005, 85, Zyutdorp, 27.1676°S, 112.778°E, depth 375 m, 29 July 2005.

1 specimen, J62974, St. SS07/2005, 8, Ningaloo, 22.0796°S, 113.797°E, depth 205 m, 2005; 1 specimen, J62973, St. SS07/2005, 23, Ningaloo, 22.0629°S, 113.723°E, depth 715 m, 2005; 2 specimens, J62971, St. SS07/2005, 24, 22.0631°S, 113.724°E, depth 713 m, 2005; 1 specimen, J62972, St. SS07/2005, 68, Point Cloates, 22.859°S, 113.328°E, depth 448 m, 2005; 1 specimen, J62970, St. SS07/2005, 75, Carnarvon, 24.5875°S, 112.253°E, depth 405 m, 2005; 2 specimens, J63690, St. SS07/2005,

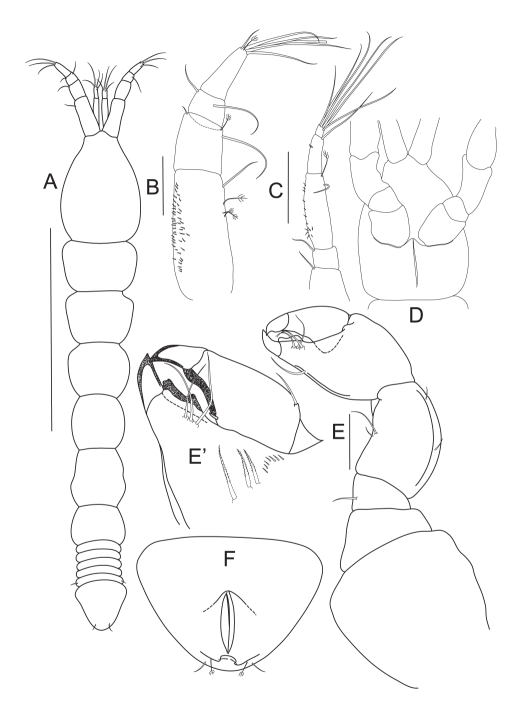


Figure 1. *Bunburia prima* sp. n., holotype female. **A** body, dorsal view; paratype female **B** antennule **C** antenna (proximal article not shown) **D** cephalothorax, ventral view **E** cheliped **E'** details of fixed finger and dactylus **F** pleotelson, ventral view. Scale lines = 1 mm for **A** and 0.1 mm for **B**–**E**.

76, Carnarvon, 24.5863°S, 112.254°E, depth 405 m, 2005; 3 specimens, J62975, St. SS07/2005, 126, Jurian Bay, 29.8604°S, 114.372°E, depth 499 m, 2005.

Type locality. near Bunbury, 33.0003°S, 114.579°E, depth 399 m.

Diagnosis. as for the genus.

Description of female. Habitus (Figs 1A, F): body 2.5 mm long, 6.2 times as long as wide. Carapace 23% of total body length, 1.5 times as long as wide. Length/ width ratios of pereonites 1 to 6: 0.7, 0.8, 0.9, 1.0, 1.1 and 0.7 respectively. Pleon about 18% of total body length; pleonites equal in length; fifth pleonite with lateral simple seta. Pleotelson (Fig. 1F) with a pair of bipinnate setae and two pairs of simple setae distally.

Antennule (Fig. 1B) 4-articled; article 1 longest, with three bipinnate setae and one simple seta on outer margin, inner margin covered with numerous minute setae; article 2 about 0.4 times as long as article 1, with one bipinnate and two simple setae distally; article 3 trapezoidal, wider than long, with two distal setae; article 4 twice as long as article 3, distally with one bipinnate, one short simple, five long simple setae and one aesthetasc.

Antenna (Fig. 1C) 6-articled; article 1 broken; article 2 with one simple seta distally; article 3 square, with one simple, distal seta; article 4 longest, four times as long as article 3, with one simple and one bipinnate setae distally and row of small spines laterally; article 5 half as long as article 4, with minute lateral setation and one long seta distally; last article very short, distally with six long setae.

Mouthparts: labrum (Fig. 2A) covered with dense setation; mandibles (Figs 2B, C) molar bent downward and tapering distally; right mandible incisor with four denticulations, left mandible with four denticulations on incisor and small lacinia mobilis with dorsal tooth. Maxillule (Figs 2D, D') ventrally with combs of short, simple setae, distally with six spines, two simple setae and minute setation, palp lost during dissection; maxilla (Fig. 2E) ovate. Labium (Fig. 2F) bilobed, with minute setation distally. Maxilliped (Fig. 2G) bases unfused distally, endites with pair of distal, simple setae and one tubercle; palp article 1 naked; article 2 with three inner setae; article 3 with three setae on inner margin, outer margin setulated; last article with five long spiniform setae and minute setation. Epignath (Fig. 2H) elongated, strap-like and naked.

Cheliped (Figs 1D–E') pseudocoxa massive, about as long as wide, naked, incompletely fused on midline of cephalothorax ventrum (Fig. 1D); basis trapezoidal and naked; merus triangular, with one seta ventrally; carpus 1.4 times as long as wide, with pair of setae on both dorsal and ventral margins; chela larger than carpus, propodus with one seta on ventral margin, inner comb of three serrated setae and row of minute spines; fixed finger with three setae on inner margin, and well calcified, inner teeth, ventrally with keel; dactylus with one small spiniform seta on inner margin and one seta dorsally.

Pereopod 1 (Figs 3A, A') coxa with seta; basis four times as long as wide, naked; ischium with simple seta; merus with one serrated seta; carpus as long as merus, with two serrated setae and one spiniform seta distally; propodus elongate, 1.5 times as long as merus, with three spiniform setae distally; dactylus 0.6 times as long as unguis, both together longer than carpus.

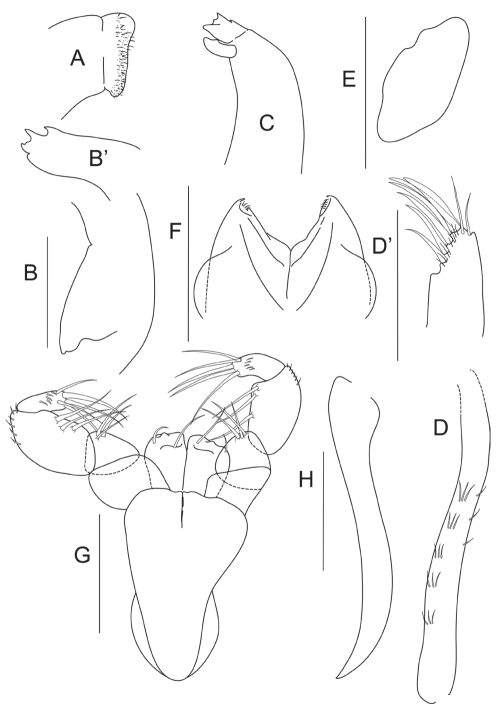


Figure 2. *Bunburia prima* sp. n., paratype female. **A** labrum **B** mandible molar **B'** incisor of right mandible **C** left mandible **D** maxillule endite **D** details of distal part of maxillule **E** maxilla **F** labium **G** maxilliped **H** epignath. Scale line = 0.1 mm.

Pereopod 2 (Fig. 3B) coxa with simple seta; basis five times as long as wide, with one bipinnate seta; ischium with single seta; merus with one serrated seta; carpus as long as merus, with two serrated setae and one spiniform seta distally; propodus elongate, 1.2 times as long as merus, with two spiniform setae distally, combs of small spines present; dactylus little shorter than unguis, with proximal seta.

Pereopod 3 (Fig. 3C) coxa with simple seta; basis four times as long as wide; ischium with one seta; merus with serrated seta ventrodistally; carpus longer than merus, with one spiniform and two serrated setae distally; propodus elongate, almost twice as long as merus, with one spiniform seta ventrally, combs of minute spines present; dactylus 0.6 times as long as unguis.

Pereopod 4 (Fig. 3D) coxa with one seta; basis 4.6 times as long as wide, with one bipinnate seta ventrally; ischium with single seta; merus with two serrated setae distally and row of minute setation; carpus longer than merus, with three serrated, strong setae ventrally, minute setation present on ventral margin; propodus little longer than carpus, with two serrated and three spiniform setae distally, ventral margin with rows of minute setation; dactylus twice as long as unguis, with numerous minute spines ventrally.

Pereopod 5 (Fig. 3E) basis with one bipinnate seta; ischium with single seta; merus with two serrated setae ventrally; carpus longer than merus, with one simple and three serrated setae distally, minute setation on ventral margin; propodus clearly shorter than carpus, with two serrated long setae and three short spiniform setae, ventrally with rows of minute setation; dactylus almost twice as long as unguis, dorsal margin with minute setation.

Pereopod 6 (Figs 3F, F') similar to pereopod 5, but dactylus/unguis slightly shorter. Pleopods absent.

Uropod (Fig. 3G) basis with small projection bearing one long and one short, simple distal setae; endopod one-articled, as long as basal article, with four long, simple setae distally and two bipinnate setae at midlength of article.

Male. Habitus (Fig. 4A): body 2.7 mm long, 6.7 times as long as wide. Carapace 20% of total body length, about 1.5 times as long as wide. Lateral margins of pereonites covered by small papillae; length/width ratios of pereonites 1 to 6: 0.7, 0.8, 0.8, 1.0, 1.0 and 0.8 respectively. Pleon about 18% of total body length; pleonites equal in length.

Antennule (Fig. 4B) stouter than that of female; 4-articled; article 1 longest and naked, 1.8 times as long as wide; article 2 0.4 times as long as first article, with four bipinnate setae distally; third article trapezoidal, about half as long as second article, with one outer and one inner simple setae; last article elongated, about half as long as first article; terminally with two bipinnate and five long, simple setae.

Cheliped (Fig. 4C) similar to that of female; merus with single seta; carpus stout – 1.3 times as long as wide, with single outer seta and pair of ventral setae (in figure only one). Propodus about as long as wide, with three setae near dactylus insertion, fixed finger with one ventral seta and three setae on inner (cutting) margin, ventral margin with keel. Dactylus as long as propodus, with single outer seta.

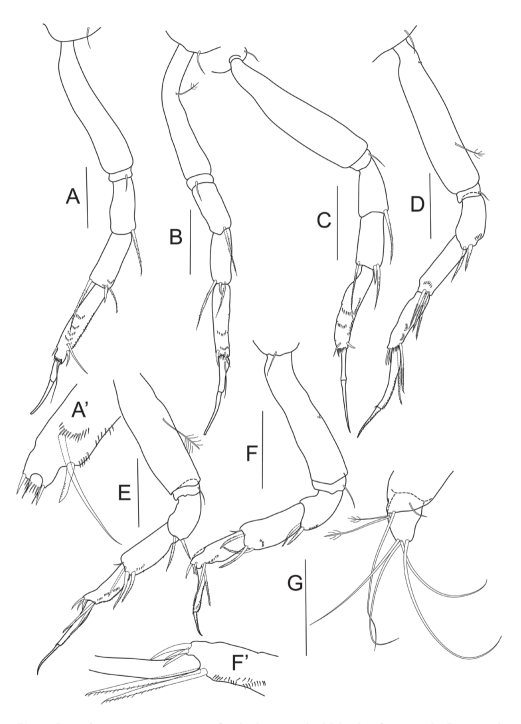


Figure 3. *Bunburia prima* sp. n., paratype female. **A** pereopod 1 **A'** detailes of P1 propodus **B** pereopod 2 **C** pereopod 3 **D** pereopod 4 **E** pereopod 5 **F** pereopod 6 **F'** details of P6 propodus **G** uropod. Scale line = 0.1 mm.

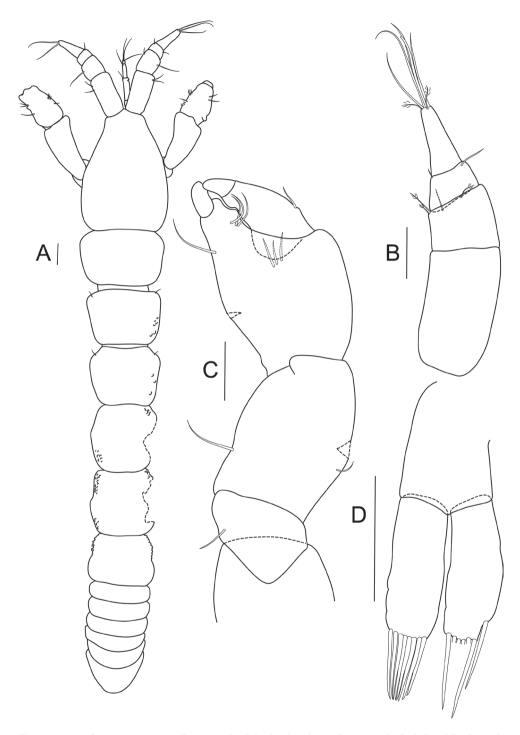


Figure 4. *Bunburia prima* sp. n., allotype male. **A** body, dorsal view **B** antennule **C** cheliped **D** pleopod. Scale line = 0.1 mm.

Pleopods (Fig. 4D) basis 0.8 times as long as each ramus, naked. Rami subequal, exopod terminating in eight strong, simple setae, endopod with one seta subdistally and six setae distally.

Distribution. The species is known from Western Australia and was recorded between Ningaloo and Bunbury City in a depth range from 205 to 715 m.

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New neotropical species of Opiinae (Hymenoptera, Braconidae) reared from fruit-infesting and leafmining Tephritidae (Diptera) with comments on the Diachasmimorpha mexicana species group and the genera Lorenzopius and Tubiformopius

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Abstract

Four new species of opiine Braconidae are described from Mexico. These are *Diachasmimorpha martinalujai* Wharton reared from *Rhagoletis* infesting fruits of *Crataegus* spp., *Diachasmimorpha norrbomi* Wharton reared from *Euphranta mexicana* infesting fruits of *Ribes pringlei*, *Eurytenes* (*Stigmatopoea*) *norrbomi* Wharton reared from *Trypeta concolor* mining leaves of *Barkleyanthus salicifolia* and *Eurytenes* (*Stigmatopoea*) *maya* Wharton reared from *Rhagoletis pomonella* infesting apples and fruits of *Crataegus* spp. Morphological features of the first metasomal segment and occipital carina, useful for placement of these species, are discussed relative to the genera *Diachasmimorpha, Eurytenes, Lorenzopius, Tubiformopius*, and *Opius* s.l. Descriptions and diagnoses are referenced to the Hymenoptera Anatomy Ontology. The following represent new combinations: *Diachasmimorpha hildagensis, Lorenzopius euryteniformis*, and *Tubiformopius tubibasis*. Revised diagnoses are provided for *D. hildagensis, D. mexicana, D. sanguinea, Eurytenes* (*Stigmatopoea*), *Lorenzopius, L. euryteniformis, Tubiformopius, T. tubigaster, T. tubibasis, Opius incoligma*, and *Opius rugicoxis*. Two species groups are delineated within *Lorenzopius* and a key to species of *Diachasmimorpha* occurring in the New World is provided.

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Keywords

Parasitoid, classification, Rhagoletis, HAO, Opius

Introduction

The subfamily Opiinae is a diverse assemblage of relatively small braconids that develop as koinobiont endoparasitoids of various cyclorrhaphous Diptera, emerging from the puparium of their hosts. Opiines have long been recognized as a distinct taxon within the Braconidae (Wharton and van Achterberg 2000), but specific features suitable for characterizing them as monophyletic relative to the Alysiinae have proven elusive (Wharton 1988, Quicke and van Achterberg 1990, Wharton et al. 2006). Koinobiont endoparasitism of cyclorrhaphous Diptera, with emergence from the puparium of the host, defines Opiinae+Alysiinae. Alysiines are readily characterized by the presence of exodont mandibles (non-overlapping, with teeth pointing outwardly) and an associated median sulcus on the back of the head (Wharton et al. 2006). Exothecines routinely appear as the sister group to Opiinae+Alysiinae (e.g. Wharton et al. 2006) and the labrum is flattened in Opiinae relative to exothecines (and other cyclostomes). The labrum is reduced in Alysiinae relative to Opiinae and cyclostomes in general. Molecular analyses published to date have provided evidence of monophyly for both Opiinae and Alysiinae when only 2-5 taxa are included in each (Dowton et al. 1998, Belshaw et al. 2000, Dowton et al. 2002) but have yet to resolve the problem completely when significantly more taxa are included (Gimeno et al. 1997; Wharton et al. 2006). There are over 1800 valid species in the Opiinae (Yu et al. 2005) and 116 genus group names (84 of these currently treated as valid by one or more authors) have been applied to various combinations of these species.

Fischer (1972, 1977, 1987) monographed the Opiinae on a world basis. This made the group more accessible for study, and this in turn led to numerous changes in the classification. Fischer (1972) initially recognized 23 genera (excluding the Gnamptodontinae, widely accepted subsequently as a separate subfamily). The number of genera currently accepted as valid varies from 17 (Wharton 1997) to about 24 (Fischer 1987, 1999) to 31 (van Achterberg and Maeto 1990, van Achterberg and Salvo 1997, van Achterberg 2004a, b, 2005). The primary purpose of the present study is to describe new species reared from fruit-infesting and leaf-mining Tephritidae from Mexico in order to broaden our understanding of host relationships within Opiinae. The search for the most appropriate genus group name for two of these species and the discovery of previously misplaced species revealed the need for re-characterization of certain genus-group taxa, and this is a secondary goal of the study.

Materials and methods

Specimens. Reared material of several species, including the four newly described below, was kindly sent for study to the senior author by Martin Aluja and Juan Rull

(Instituto de Ecologia, Xalapa, Mexico), Robert Jones (Universidad de Autónoma de Querétaro, Querétaro, Mexico), and Allen Norrbom (USDA Systematic Laboratory, Washington, D. C.). Other specimens used in this study, including type material of previously described species, were borrowed from or examined at the following institutions: American Entomological Institute, Gainesville, Florida, USA (AEIC), Canadian National Collection, Ottawa, Ontario, Canada (CNC), Hungarian Natural History Museum, Budapest, Hungary (HNHM), National Museum of Natural History, Leiden, The Netherlands (RMNH), Naturhistorisches Museum Wien, Vienna, Austria (NHNW), Texas A&M University Insect Collection, College Station, Texas, USA (TAMU), The Natural History Museum, London, England (BMNH), and U. S. National Museum of Natural History, Washington, D. C., USA (USNM).

In the material examined section under each species description, we record label data for the holotype exactly as they appear on the labels. We use a more standardized format for paratypes, additional specimens examined, and published data for other specimens.

Figures. Images were acquired digitally using Syncroscopy's AutoMontage[®] software, in combination with a ProgRes 3008 digital camera mounted on a Leica MZ APO dissecting microscope. All images were further processed using various minor adjustment levels in Adobe Photoshop[®] such as image cropping and rotation, adjustment of contrast and brightness levels, color saturation, and background enhancement. Automontage images are available in color and high resolution at http://peet.tamu.edu/projects/8/public/site/wharton_lab/home.

Database management, digital dissemination, and ontology reference. Illustrations and free-text diagnoses for morphospecies were assembled in mx, a web-based content management system that facilitates data management and dissemination for taxonomic and phylogenetic works (e.g. Yoder et al. 2006). The mx project is open source, with code and further documentation available at http://sourceforge.net/projects/mx-database/. Data pertinent to this work, including specimen-level data, images, diagnoses, and descriptions, are available at http://peet.tamu.edu/projects/8/ public/site/wharton_lab/home.

Morphological terms used in this revision were matched to the Hymenoptera Anatomy Ontology (HAO, Yoder et al. 2010) (Appendix). Identifiers (URIs) in the format http://purl.obolibrary.org/obo/HAO_XXXXXX represent anatomical concepts in HAO version http://purl.obolibrary.org/obo/hao/2011-05-18/hao. owl. They are provided to enable readers to confirm their understanding of the anatomical structures being referenced. To find out more about a given structure, including images, references, and other metadata, use the identifier as a web-link, or use the HAO:XXXXXXX (note colon replaces underscore) as a search term at http://glossary.hymao.org. For published examples see Wharton et al. (2010) and especially Talamas et al. (2011).

Terminology and measurements. Terminology as linked through the HAO (Appendix) largely follows Sharkey and Wharton (1997), with a few additions from Walker and Wharton (2011). For the first metasomal segment (sometimes referred to as the petiole), T1 is the median tergite and S1 is the sternite: the well-sclerotized basal

portion of the sternum. S1 is often greatly reduced in opiines but well developed in several of the species treated here. A tendon originates in the propodeum and inserts at the base of T1 medially. The point of insertion, which we have called the dorsal tendon attachment, serves as a convenient point of reference for orientation. The propodeum medially has at least a partial areola in most of the species treated here. Morphologically, this areola may not be strictly homologous with the areola as defined, for example, by Townes (1969) for Ichneumonidae or Sharkey and Wharton (1997, Fig. 8) for Braconidae since in these opiines there is no distinct petiolar area posteriorly. On the mesoscutum, some of the species treated here have a mesoscutal humeral sulcus extending along the lateral margin from the base of the notaulus. When present, it is usually carinately margined laterally, and we have referred to this as the supra-marginal carina in the text below. Wing cells are indicated in Fig. 36; abbreviations for wing veins are indicated in Fig. 16, both following Sharkey and Wharton (1997).

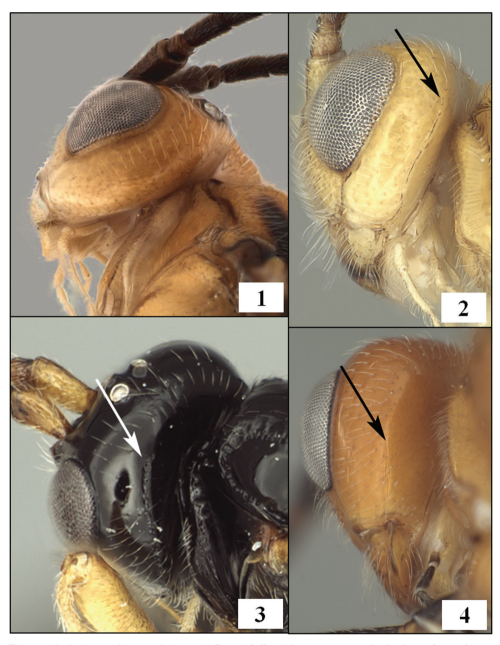
Quantitative data in descriptions are based on 5 individuals of each sex, when available. Measurements largely follow Walker and Wharton (2011). Mesosomal width is the distance across the mesoscutum between the tegula. Width of clypeus was measured at the lateral margin rather than at the anterior tentorial pit. The eye/ temple ratio is an important species-level characteristic, but is notoriously difficult to measure consistently because slight repositioning may result in significantly different ratios across this curved surface. The measurements are therefore provided to illustrate relative difference among species, and less emphasis should be placed on the absolute values. In the descriptions below, we have indicated whether eye/temple ratios were calculated from measurements made in dorsal view, lateral view, or both.

Results and discussion

Generic placement

The new species described below are placed in the genera *Diachasmimorpha* Viereck and *Eurytenes* Foerster. The basis for these placements, with particular reference to the nature of the occipital carina, characteristics of the first metasomal segment, and tephritid parasitism, are discussed in this section. Diagnoses of relevant taxa and descriptions of the new species follow in the next section, alphabetically by genus.

The occipital carina varies from completely present to completely absent in the Opiinae with most species having the carina broadly absent mid-dorsally but well-developed laterally (Figs 1–4). Fischer (1972) created the tribe Desmiostomatini for all species known to him in which the occipital carina was completely lost or apparently so (Fig. 1). Wharton (1983, 1987a, 1988) subsequently discovered that loss of the carina occurred in several other groups as well and hypothesized multiple independent losses within the subfamily. The opiine parasitoids of fruit-infesting Tephritidae are distributed among several genera (Wharton 1997), most of which have at least some species lacking an occipital carina. The New World endemics *Doryctobracon* Enderlein

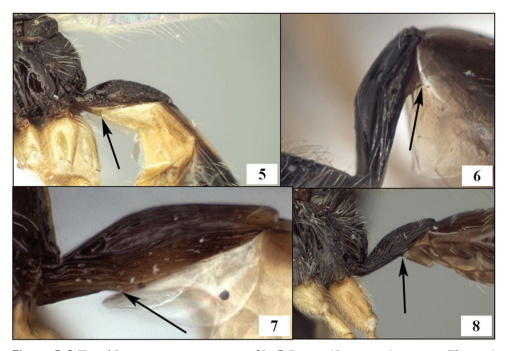


Figures 1–4. Occipital carina. I *Opius (Bellopius) bellus* Gahan, carina completely absent 2 *Diachasmi-morpha mellea* (Gahan), arrow at dorsal end of carina 3 *Lorenzopius tubulatus* (Fischer), holotype female, arrow at dorsal end of carina 4 *Diachasmimorpha sanguinea* (Ashmead), arrow at dorsal end of weak carina.

and *Bellopius* Wharton (the latter presently placed as a subgenus of *Opius* Wesmael s.l.) are thus far known only from tephritid hosts and all species lack the occipital carina. The Old World endemics *Psyttalia* Walker and *Fopius* Wharton are also known only

as parasitoids of Tephritidae but only a small percentage of the known species have the occipital carina completely lacking. The cosmopolitan Utetes Foerster, which may also be restricted to tephritid hosts, contains a group of New World endemics in which the occipital carina is completely lacking. This New World tropical and subtropical group was formerly treated as Bracanastrepha Brèthes (Fischer 1977, Wharton 1988). When Bracanastrepha was synonymized under Utetes (Wharton 1988), and Utetes restricted to species with a distinctive hind tibial carina, all of the remaining Bracanastrepha that lacked an occipital carina, including those species previously placed in the subgenera Thiemanastrepha Fischer and Buckanastrepha Fischer, were transferred to Opius s.l. (Wharton 1988). Hosts are unknown for nearly all of these, but at least one of the species is recorded from tephritids (Costa Lima 1938). Most species of *Diachasmimorpha*, another group of tephritid parasitoids, retain the lateral portion of the occipital carina, but there are parallel losses of the carina within Old and New World species groups that have caused confusion in the placement of a few species. Parasteres Fischer, for example, was defined solely on the basis of the loss of the occipital carina relative to other species with a short second submarginal cell. Parasteres originally included two species, each described from a single male specimen (Fischer 1964, 1967a). The type species of *Parasteres* was subsequently discovered to be the male of the Old World species D. tryoni (Cameron), with the holotype collected during a recovery program in Puerto Rico where D. tryoni had been released for control of tephritid pests. The second species originally included in *Parasteres* is treated below and belongs to the *Dia*chasmimorpha mexicana species group, endemic to the New World. The members of the *mexicana* species group are difficult to place because the occipital carina is present as a very short spur ventrally but the spur is easy to overlook and is often obscured by other body parts. Members of the mexicana species group have proven challenging to identify because two of the three previously described species were based on single male specimens and female ovipositor length is an important diagnostic feature. Diachasmimorpha was not recognized as valid until after publication of Fischer's (1972, 1977, 1987) monographs of the World Opiinae. Thus, a number of species undoubtedly remain incorrectly placed in other genera and no comprehensive key to species is available (but see Wharton and Yoder 2012).

The first metasomal segment, often referred to as the petiole (Sharkey and Wharton 1997), consists of a heavily sclerotized tergite (T1) and sclerotized sternite (S1) of varying length (Figs 5–8). In several New World species of Opiinae, the petiole is long and more or less parallel-sided. At least two genus-group names have been proposed for species with this characteristic: *Lorenzopius* van Achterberg and Salvo, 1997 and *Tubiformopius* Fischer, 1998. The relationships of the four explicitly included species to others in the Opiinae have not been discussed previously, nor is it clear that the feature used to define these two taxa (an elongate, tube-shaped T1) is sufficiently characterized to enable assessment of homology across the various species with an elongate petiole. Walker and Wharton (2011), for example, described a new species of *Eurytenes* s.s. with an exceptionally long, tubular petiole and Wharton (1988) placed *Opius macrocerus* Thomson in *Eurytenes* partly on the basis of a narrow, parallel-sided petiole.



Figures 5–8. T1 and S1, arrows at posterior margin of S1. **5** *Eurytenes (Stigmatopoea) macrocerus* (Thomson) **6** *Lorenzopius calicomyzae* van Achterberg and Salvo, holotype female **7** *Eurytenes (Stigmatopoea) maya* Wharton sp. n., paratype female **8** *Tubiformopius tubigaster* (Fischer), holotype male.

Van Achterberg and Salvo (1997) described *Lorenzopius* and characterized it on the basis of the tube-shaped petiole (Fig. 6) with at least the basal half of the tergite closed ventrally and with a midpit on the mesoscutum posteriorly. Three species were originally included: *Opius tubulatus* Fischer, 1979, *O. sanlorenzensis* Fischer, 1964, and the type species, *L. calycomyzae* van Achterberg and Salvo. *Opius tubibasis* Fischer, 1978 was also mentioned as a potential member of the newly described genus. Almost concurrently, Fischer (1998) described *Tubiformopius*, which he later (Fischer 1999) treated as a synonym of *Lorenzopius*. However, the type species of *Tubiformopius* (*Opius tubigaster* Fischer, 1968), while possessing a tubular petiole (Fig. 8), differs from *L. calycomyzae*, *L. tubulatus*, and *L. sanlorenzensis* in several important aspects. In *T. tubigaster*, there is no midpit on the mesoscutum, fore wing m-cu is widely antefurcal, the first subdiscal cell is broadly open distally, and the mandible has a distinct basal lobe (= basal tooth). Given these differences, I retain *Tubiformopius* as valid, at least for the present, and also include *Tubiformopius tubigaster*.

Neither van Achterberg and Salvo (1997) nor Fischer (1998) mentioned the sternite in their descriptions, focusing instead on the tubular tergite, closed ventrally. What is most distinctive about *Lorenzopius* and *Tubiformopius*, however, is the length of S1 and its apparent fusion with T1. The presence of a prominent S1 is an unusual feature in the Opiinae and it is therefore not surprising that two genus group names have been proposed for species with this characteristic. In the vast majority of opiine species S1 is present as a very short basal sclerite, clearly separated by membrane from the tergite, but S1 is often overlooked because it is difficult to see without removal of at least one hind leg. The type species of *Tubiformopius* and *Lorenzopius* have sternites illustrating different positions along the morphocline of an increasingly elongate S1 that appears fused to the tergite. S1 in T. tubigaster is 0.5–0.6 times the length of T1 (Fig. 8) while S1 in L. calycomyzae and L. tubulatus extends nearly the full length of T1 (Fig. 6). The other differences noted above between the type species of *Tubiformopius* and *Lorenzopius* make it relatively easy to place tubibasis in Tubiformopius rather than Lorenzopius, but other species with a narrow T1 and an elongate S1 are more challenging to place. Two such examples, O. incoligma Fischer and O. rugicoxis Fischer (Figs 74-83) are diagnosed below under Opius to highlight the problems in placing such species when focusing only on the presence of an elongate S1. Eurytenes is also problematic since several species have long, narrow petioles. The species of Lorenzopius are similar in many respects to Eurytenes macrocerus (Thomson), the type species of Eurytenes (Stigmatopoea). Both have an exposed labrum with sharp ventral margin to the clypeus, relatively well-developed notauli, a distinct midpit on the mesoscutum, relatively large scuto-scutellar sulcus, and similar venation, most notably the parallel-sided stigma. In Stigmatopoea, however, the dorsope is retained and S1, though longer than in most Opiinae, is short relative to Lorenzopius and clearly separated from T1 (as in Fig. 5). The number of shared features may indicate that Lorenzopius represents a distinct lineage derived from a Stigmatopoea-like ancestor. Otherwise, the exceptional morphological diversity in the species that share an elongate S1 suggests homoplasy, with possibly multiple derivations of an elongate S1. Until the relationships among the many Neotropical species with an elongate S1 are better understood, this feature will remain useful for characterizing opiine species, but must be used cautiously and in combination with other characters for defining genera.

Taxonomy

Diachasmimorpha Viereck

http://species-id.net/wiki/Diachasmimorpha

- *Diachasmimorpha* Viereck, 1913: 641. Type species: *Diachasmimorpha comperei* Viereck, 1913 [a junior subjective synonym of *Diachasmimorpha longicaudata* (Ashmead, 1905)]. Monobasic and original designation.
- *Biosteres (Parasteres)* Fischer, 1967a: 3. Type species: *Biosteres (Parasteres) acidusae* Fischer, 1967a [a junior subjective synonym of *Diachasmimorpha tryoni* (Cameron, 1911)]. Original designation.
- *Parasteres*: Fischer 1971: 33 (change in rank). Synonymized under *Biosteres* by Wharton and Marsh (1978:154) and under *Diachasmimorpha* by Wharton (1987a: 62).

Diagnosis. Mandible without basal lobe ventrally. Labrum concealed. Occipital carina broadly absent dorsally, present or absent laterally. Propleuron ventral-laterally

without oblique carina. Notauli deep, unsculptured or nearly so, well developed anteriorly, varying posteriorly from absent to deep and complete to midpit; midpit always present. Fore wing stigma short, broad, discrete posteriorly, r1 arising at or distad its midpoint; second submarginal cell short; m-cu arising from second submarginal cell. Hind wing RS absent basally, sometimes present as a weakly pigmented crease distally; 2M distinctly pigmented nearly to wing margin; m-cu present, well-developed. Dorsope absent.

The species of *Diachasmimorpha* are most readily recognized by the pattern of fore and hind wing venation (Figs 9, 16) in combination with the concealed labrum (Fig. 12), unsculptured notauli (Figs 11, 14, 19, 20), and lack of oblique carina on the propleuron (Fig. 23). The species of *Doryctobracon* Enderlein, endemic to the New World, are similar but have the fore wing m-cu interstitial or arising from the first submarginal cell and the labrum is partially exposed. *Fopius* Wharton, an Old World genus with species that have been introduced to the New World, is also similar. The species of *Fopius* differ by the presence of completely sculptured notauli and the presence of an oblique carina on the propleuron (Fig. 24).

Remarks. Both New and Old World species groups of *Diachasmimorpha* occur in Mexico. *Diachasmimorpha longicaudata* (Ashmead) and *D. tryoni*, both representatives of the Old World *longicaudata* species group (Wharton 1997), were established in various parts of Mexico during biological control programs directed against tephritid pests primarily in the genus *Anastrepha*. Females of the Old World species are readily distinguished from New World *Diachasmimorpha* because of the sinuate ovipositor (Fig. 28). The notauli are also more deeply incised posteriorly in the *longicaudata* species group (Fig. 19, in contrast to Fig. 20), which facilitates identification of males in biological control and other tephritid pest management programs. The name *Parasteres* continues to be used by some authors, for example as a subgenus of *Diachasmimorpha* (Yu et al. 2012), but we continue to treat *D. tryoni* and *D. longicaudata* in the same species group based in part on ovipositor morphology. We therefore do not treat *Parasteres* as valid, nor do we recognize subgenera under *Diachasmimorpha* at this time.

New World species have previously been referred to as the *mexicana* species group (Wharton 1997), a use we continue here. Wharton (1997) noted, however, that there were two subgroups distinguished in part on the basis of relative loss of the occipital carina. Further examination and discovery of additional species provides support for the two subgroups. One of these subgroups consists of *D. juglandis* (Muesebeck), *D. mellea* (Gahan), and *D. sublaevis* (Wharton). The occipital carina is generally better developed in this subgroup (usually readily visible laterally as in Fig. 2), the wings are hyaline, and the body is yellowish. As in the *longicaudata* species group, the anterior margin of the pronotum ventral-laterally is sharply excavated (Fig. 17). The second subgroup contains *D. mexicana* (Cameron), *D. sanguinea* (Ashmead), *D. hildagensis* (Fischer), new combination, and the new species described below. In all of these species, the occipital carina is greatly reduced, present only as a short spur ventrally near the mandible (maximum extent shown in

Fig. 4). These species also have infumate wings (Fig. 16) and the body tends to be orange rather than yellow. The anterior margin of the pronotum ventral-laterally is also more sinuate than abruptly excavated (Fig. 18). Detailed diagnoses are provided below for the three previously described species in this second subgroup, to facilitate comparison with the newly described species.

Key to species of Diachasmimorpha known from U.S. and Mexico

1	Female (ovipositor clearly visible, extending well beyond apex of metasoma)2
-	Male
2 (1)	Ovipositor distinctly sinuate subapically (Fig. 28)
-	Ovipositor straight or nearly so subapically (Fig. 29)
3 (2)	Metasomal tergum 2 distinctly striate medially (Fig. 21). Occipital carina
	well developed laterally, extending from base of mandible at least to mid eye
	height
-	Metasomal tergum 2 without striae or other sculpture (Fig. 22). Occipital ca-
	rina poorly developed to absent, not extending dorsally to lower eye margin
4 (2)	Head dark, at least on dorsal half (Fig. 9)
-	Head pale (Figs 2, 4), yellow or orange except sometimes ocellar field dark7
5 (4)	Ovipositor (total length) about 2.5 times longer than mesosoma. Notaulus
	extending anteriorly to margin of mesoscutum (Figs 10, 27)6
-	Ovipositor (total length) less than 2.0 times longer than mesosoma. Notaulus
	rarely extending anteriorly to margin of mesoscutum, usually terminating
	just before reaching margin (Fig. 32)D. norrbomi, sp. n.
6 (5)	Eye smaller than in Fig. 32, about $1.5-1.6 \times \text{longer than temple in lateral}$
	viewD. hildagensis (Fischer)
_	Eye larger, $2.1-2.9 \times \text{longer}$ than temple in lateral view (Fig. 33)
	D. martinalujai, sp. n.
7 (4)	Wings darkly infumate (as in Figs 16, 36). Occipital carina represented at
	most as in Fig 4, usually present as a short spur near mandible, otherwise
	absent
	Note: mexicana (Cameron) also keys here but is known only from the male,
	which has a much smaller eye than that of <i>sanguinea</i> .
_	Wings hyaline (Fig. 20). Occipital carina present laterally at least to lower
	margin of eye, usually as in Fig. 2
8 (7, 14)	Metasomal tergum 2 distinctly striate medially (as in Fig. 21)9
_	Metasomal tergum 2 without striae or other sculpture (as in Fig. 22)
	D. juglandis (Muesebeck)
9 (8)	Precoxal sulcus distinctly impressed, usually broad but very weakly sculp-
× /	tured, nearly smooth (as in Fig. 38). Hosts are walnut husk flies in species of
	Juglans

-	Precoxal sulcus distinctly impressed, broad, heavily sculptured: crenulate to
	foveolate (as in Fig. 17). Hosts are other species of <i>Rhagoletis</i> in other fruits
10(1)	Head black at least over dorsal half11
-	Head pale, yellow to orange except ocellar triangle sometimes black
11 (10)	Eye in dorsal view as long as temple; eye in lateral view 1.3–1.4 × longer than
	temple
_	Eye slightly larger, in dorsal view eye $1.4-1.9 \times longer$ than temple, in lateral
	view $1.7-2.4 \times \text{longer than temple}$ 12
12 (11)	Notaulus extending anteriorly to margin of mesoscutum (Fig. 27)
	D. martinalujai, sp. n.
-	Notaulus rarely extending anteriorly to margin of mesoscutum, usually ter-
	minating just before reaching margin (Fig. 32)D. norrbomi, sp. n.
13 (10)	Metasomal tergum 2 striate medially (Fig. 21)14
_	Metasomal tergum 2 without striae or other sculpture (Fig. 22)15
14 (13)	Notauli deep posteriorly as it nears midpit (Fig. 19)
_	Notauli more shallow posteriorly as it nears midpit (Fig. 20)8
15 (13)	Metasomal terga mostly black (Fig. 22)
_	Metasoma with at least terga 3–5 pale: yellow to orange
16 (15)	Wings hyaline
_	Wings darkly infumate
17 (16)	Eye larger, about $1.3-1.5 \times longer$ than temple in lateral view
	D. sanguinea (Ashmead)
_	Eye smaller, subequal to temple in lateral view (Fig. 35)
	D. mexicana (Cameron)

Diachasmimorpha hildagensis (Fischer), comb. n.

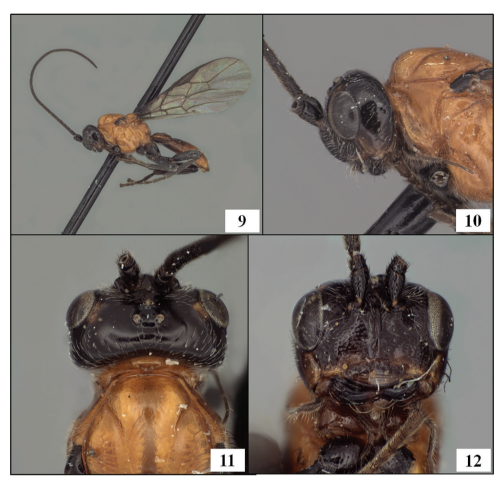
http://species-id.net/wiki/Diachasmimorpha_hildagensis Figs 9–12, 13–16

Opius (Biosteres) hildagensis Fischer, 1964: 12, 20–22. Holotype male in AEIC (examined). *Biosteres (Parasteres) hildagensis*: Fischer 1967a: 5 (generic transfer). *Parasteres hildagensis*: Fischer 1971: 33 (generic transfer); Fischer 1977: 880–883 (key,

redescription).

Type locality: Mexico, State of Mexico, Hidalgo National Park.

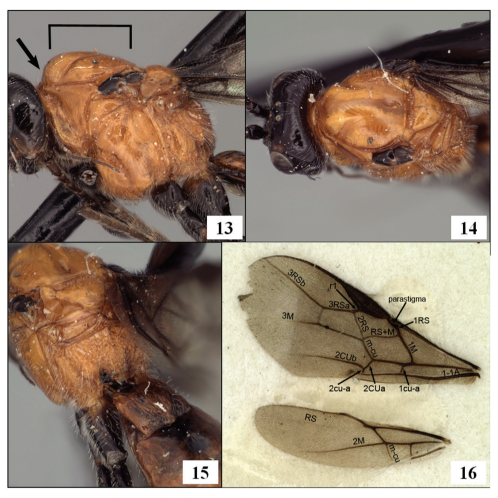
Type material. Holotype male (AEIC), first label, first line: Hidalgo Natl. Pk. second line: State of Mex., Mex. third line: x.12.62 3000 m. fourth line: H. & M. Townes Second label [purple]: Holotype Third label: Opius hildagensis [male symbol] sp. n. det. Fischer Fourth label: Type No. 336



Figures 9–12. *Diachasmimorpha hildagensis* (Fischer), holotype male. 9 habitus 10 head and base of notaulus, lateral view 11 head, pronope, and base of notaulus, dorsal view 12 face.

Other specimens examined: 2 females, 1 male, Mexico, Mexico, Rt 890, km 9, 6 km W Lago Zempoala, 2.x.1991, A.L. Norrbom, reared from *Oedicarina latifrons* infesting fruits of *Solanum brachycarpum* (91M14B) (TAMU, USNM).

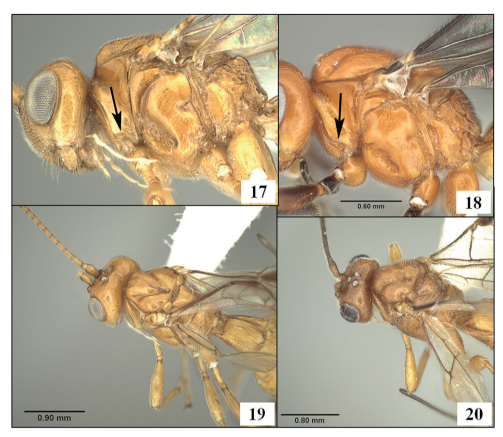
Diagnosis. Holotype male. Eye in dorsal view as long as temple, temples neither receding nor expanded beyond eyes; eye in lateral view 1.3 × longer than temple. Frons irregularly rugulose along midline between antenna and median ocellus. Clypeus 2.8 × wider than high. Occipital carina distinct near base of mandible, short, not extending dorsally to ventral margin of eye. Antenna with 46 flagellomeres; first flagellomere 1.25 × longer than wide. Pronope deep, large, interrupting posterior crenulate groove middorsally. Notauli deep anteriorly, reaching anterior-lateral margin of mesoscutum and extending posteriorly about 0.5 × distance to deep, elongate midpit. Precoxal sulcus distinctly crenulate throughout, nearly extending to anterior margin of mesopleuron. Propodeum rugose, areola extending over posterior 0.6 but largely obscured



Figures 13–16. *Diachasmimorpha hildagensis* (Fischer), holotype male. **13** mesosoma, lateral view, arrow showing anterior declivity of mesoscutum, bracket showing mesoscutal disc **14** head and mesonotum, dorsal view **15** propodeal sculpture **16** left fore and hind wings illustrating wing vein terminology.

by sculpture. Fore wing 2RS 0.95 × length of 3RSa; m-cu distinctly postfurcal. T1 with dorsal carinae weakly converging, widely separated at posterior margin, gradually weakening posteriorly. Meso- and metasoma orange, tegula black, head dark brown to black except narrow yellow-orange band along epistomal sulcus extending to and through malar sulcus and small orange spot on vertex adjacent eye; legs black except extreme base of hind coxa irregularly orange, joint between femora and trochantelli reddish orange, mid and hind tarsi dark brown. Body length about 4.3 mm, fore wing length 4.5 mm, mesosoma length 1.8 mm.

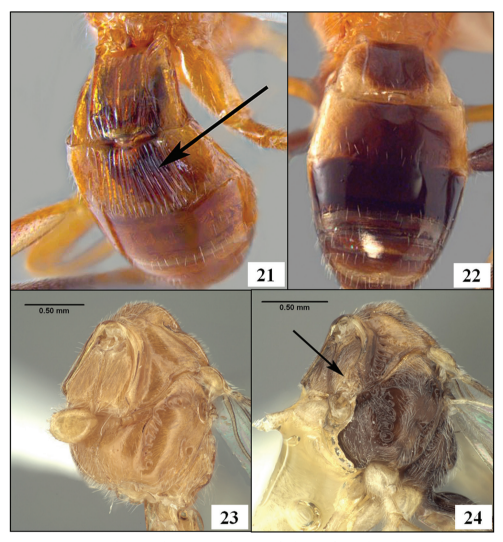
Specimens reared from *Oedicarena latifrons* (Wulp) vary as follows relative to the holotype: clypeus length/height ratio 2.6–2.8; eye/temple ratio, lateral view, 1.3–1.4 (males), 1.55 (female); antenna with 46–48 flagellomeres; 2RS/3RS ratio 0.95–1.0;



Figures 17–20. *Diachasmimorpha* spp. 17 *D. longicaudata* (Ashmead), arrow showing sharply indented margin of pronotum laterally 18 *D. sanguinea* (Ashmead), arrow showing less sharply indented margin of pronotum laterally 19 *D. longicaudata*, dorsal view 20 *D. mellea* (Gahan), dorsal view.

ovipositor sheath 2.5 times longer than the mesosoma; mesosoma length 1.85–1.9 mm (male), 2.0 mm (female); one male with T1 dorsal carinae absent over posterior 0.5 and mandible, clypeus, face, and hind coxa more extensively orange; female with outer surface of hind coxa completely pale (dark medially), mandible, clypeus and lower part of face more extensively pale than in holotype.

This species is slightly larger and has a smaller eye than both of the similarly-colored species described below, *D. martinalujai*, sp. n. and *D. norrbomi*, sp. n. Based on the single female reared from *O. latifrons*, *D. hildagensis* also has a much longer ovipositor than *D. norrbomi*. The ovipositors of *D. hildagensis* and *D. martinalujai* are similar in length. In *D. hildagensis* and *D. martinalujai*, the notaulus consistently extends anteriorly to the margin of the mesoscutum whereas in *D. norrbomi*, the notaulus usually does not. Color variation in the specimens reared from *O. latifrons* is similar to that in the paratype series of *D. martinalujai* and *D. norrbomi*. Both *D. hildagensis* and the two newly described species are similar in having the head mostly dark in contrast to the orange heads of *D. mexicana* and *D. sanguinea*, the other two members of this

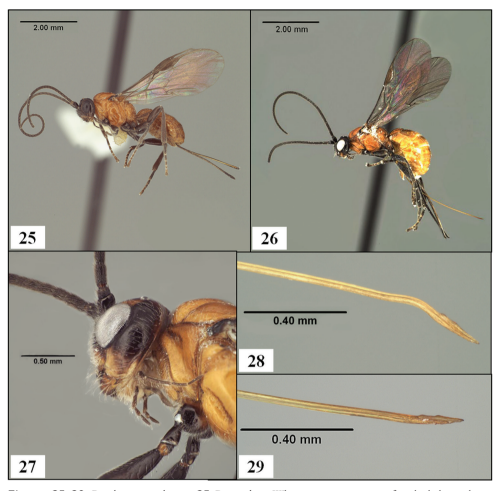


Figures 21–24. Propleuron and T2. 21 *Diachasmimorpha longicaudata* (Ashmead), T2 with striae (arrow) 22 *Diachasmimorpha tryoni* (Cameron), T2 without sulpture 23 *D. longicaudata*, propleuron without oblique carina 24 *Fopius arisanus* (Sonan), propleuron with oblique carina (arrow).

species group. The holotype of *D. hildagensis* exhibits subsurface discoloration on the metasoma, but the tergites are all entirely orange.

Biology. There is no biological information associated with the holotype. The non-type material listed above was reared from the tephritid *Oedicarina latifrons* infesting fruits of *Solanum brachycarpum* Correll. Collection data and host information can be found in Norrbom et al. (1988).

Remarks. The name *hildagensis* is based on a misreading of the locality label on the holotype, which is correctly written as Hidalgo Nat. Park, not "Hildago Nat. Park" as given by Fischer (1964) in the original description. In the original description, *hilda*-



Figures 25–29. *Diachasmimorpha* spp. 25 *D. norrbomi* Wharton sp. n., paratype female, habitus showing relatively shorter ovipositor 26 *D. martinalujai* Wharton sp. n., holotype, habitus showing relatively longer ovipositor 27 *D. martinalujai* paratype male, base of notaulus 28 *D. tryoni* (Cameron) apex of ovipositor showing subapical sinuation 29 *D. norrbomi*, paratype female, apex of ovipositor.

gensis is included in a key to the subgenus *Biosteres*, but the subgeneric name was not included in the heading for the species description. This species is here transferred to *Diachasmimorpha*, as diagnosed above, on the basis of fore and hind wing venation (Fig. 16), the morphology of the labrum, clypeus, and mandible (Fig. 12), and the well-developed notaulus and midpit (Figs 13–15). A detailed description of *Diachasmimorpha* is provided in Wharton (1997). Inclusion of *D. hildagensis* in the *mexicana* species group is based on the greatly reduced occipital carina, sinuate anterior margin of the pronotum ventral-laterally, and the body and wing coloration.

Both *D. hildagensis* and *D. mexicana* were described from single male specimens collected in the state of Mexico and the Distrito Federal, respectively, and unassociated with either hosts or host plants. Both have relatively small eyes, but are readily

separated from one another on the basis of head coloration. Associating the name *hildagensis* with the many dark-headed specimens available for study, however, has been considerably more challenging. Reared material, representing over 50 specimens kindly made available to us by Allen Norrbom, Martin Aluja, and Juan Rull, provides clear evidence of sexual dimorphism in eye size as well as variation in ovipositor length associated with different hosts and host plants. This material has been especially critical for understanding color patterns and associating males with females. Based primarily on eye size and body size, the holotype of *D. hildagensis* is closest to the series of three specimens listed above under "other specimens examined," that emerged from puparia of *O. latifrons* infesting fruits of *S. brachycarpum*. From the remaining reared material, we describe two closely similar species below.

Diachasmimorpha martinalujai Wharton, sp. n.

urn:lsid:zoobank.org:act:9E85B215-4032-4CEC-ADF4-6F068E83C029 http://species-id.net/wiki/Diachasmimorpha_martinalujai Figs 26, 27, 31, 33

Type locality. Mexico, Distrito Federal.

Type material. Holotype. Female (UNAM), first and only data label, first line: Mexico, D. F. second line: Host = R. pomonella third line: Host plant=Crataegus sp. fourth line: Common name=Tejocote fifth line: 7.xi.2007 J. Rull

Paratypes: 1 male, same data as holotype (TAMU). 1 male, Mexico, Hidalgo, Atotonilco, 4.xi.2002, J. Rull, key 30, reared from *Rhagoletis* nr. *pomonella* infesting fruit of *Crataegus* spp. (TAMU). 1 male, Mexico, Puebla, San Martin, 24.xi.2003, M. Pale key 69, reared from *Rhagoletis* nr. *pomonella* infesting fruit of *Crataegus mexicana* (TAMU).

Description. *Female.* Head in dorsal view $1.30 \times$ broader than mesoscutum, $1.65 \times$ broader than face; eye in dorsal view $2.0 \times$ longer than temple, temples not receding, but width at eyes greater than width at temples; eye in lateral view $2.05 \times$ longer than temple. Discrete facial midridge ending dorsally as a distinct elevation at base of antennae, continuing between antennae onto frons as low, sharp, bifurcating ridges. Frons irregularly rugulose along midline between bifurcating arms, otherwise polished, with moderately dense patch of decumbent, laterally-directed, white setae on either side of midline; bare on either side of ocellar field; width of ocellar field $0.95 \times$ distance from ocellar field to eye. Face $2.2 \times$ wider than high; uniformly setose (as in Figs 31, 33), distinctly punctate, punctures separated by about $1 \times$ their diameter or slightly less. Malar sulcus deep, complete; malar space about $1.1 \times$ basal width of mandible, $0.35 \times$ eye height. Clypeus $2.65 \times$ wider than high; very weakly convex, nearly flat. Occipital carina weak, difficult to discern near base of mandible, short, extending dorsally to ventral margin of eye. Hypostomal carina extending as short but distinct flange below mandible. Antenna with 45 flagellomeres; first flagellomere $1.3 \times$ longer than second; $1.8 \times$ longer than wide.

Mesosoma $1.4 \times \text{longer}$ than high; $1.9 \times \text{longer}$ than wide; $1.35 \times \text{higher}$ than wide. Pronotum not visible dorsally; crenulae extending over dorsal 0.3-0.4 of prono-

tum laterally within narrow, shallow groove; groove not margined anteriorly by carina; anterior margin of pronotum laterally sinuate, not abruptly excavated. Notauli deep anteriorly, ending abruptly posteriorly, short, not quite extending posteriorly to level of anterior margin of tegula, not reaching long, narrow midpit, anterior end extending to anterior-lateral margin of scutum; mesoscutum without supra-marginal carina adjacent margin of mesoscutum between base of notaulus and tegula. Scuto-scutellar sulcus rectangular or nearly so; 4.75 × wider than midlength; crenulate-foveolate. Propodeum rugose, areola extending over posterior 0.8 but partially obscured by sculpture. Precoxal sulcus crenulate, distinctly separated from anterior margin of mesopleuron.

Wings. Fore wing stigma short, broad, discrete distally, $3.5 \times$ longer than wide; r1 arising from midlength of stigma; 1RS (excluding parastigma) $0.30 \times$ length of 1M; m-cu postfurcal by $0.25 \times$ length of m-cu; second submarginal cell converging distally; 2RS $0.9 \times$ length of 3RSa; 2CUa about $1.7 \times$ longer than 2cu-a; 1cu-a distad 1M by about $1.0 \times$ its length.

Metasoma not distinctly petiolate; head $1.8 \times$ wider than apex of T1. T1 $1.05 \times$ as long as apical width; strongly diverging apically, with apex $2.1 \times$ wider than base; surface smooth; dorsal carinae parallel-sided, widely separated posteriorly, distinctly elevated over anterior 0.6, weaker and becoming indistinct posteriorly; lateral carina weaker than dorsal carina basally, extending distinctly ventrad spiracle, rounded and barely distinguishable posteriorad spiracle; spiracle at midlength of T1; dorsope absent but lateral and dorsal carinae elevated at junction, giving appearance of a slight depression; laterope deep; S1 very short. T2 unsculptured, with sharp lateral margins. Ovipositor sheath 2.4 × longer than mesosoma, densely setose over apical half, with 4–5 irregular rows of setae, the setae longer than sheath width, more sparsely setose basally.

Color (Fig. 26). Very similar to *D. hildagensis*. Meso- and metasoma orange, except tegula black; head dorsally black except for small orange spot on vertex adjacent eye; lower gena and most of occiput yellow-orange; narrow bands dorsad epistomal sulcus, along ventral margin of clypeus and vertically through middle of mandible orange; legs black to dark reddish brown except basal 0.5 of hind coxa orange, joint between femora and trochantelli reddish orange.

Male. Largely as in female with variation as follows: head in dorsal view $1.35-1.45 \times broader$ than mesoscutum, $1.6-1.7 \times broader$ than face; eye in dorsal view $1.6-1.85 \times longer$ than temple, in lateral view $1.7-1.95 \times longer$ than temple; face $1.95-2.1 \times wider$ than high; malar space $0.3-0.45 \times eye$ height; clypeus $2.6-2.8 \times wider$ than high; antenna with 39–47 flagellomeres; first flagellomere $1.1-1.2 \times longer$ than second, $2.0-2.1 \times longer$ than wide; mesosoma $1.25-1.35 \times longer$ than high; $1.85-1.95 \times longer$ than wide; $1.4-1.5 \times higher$ than wide; pronope deep, moderately large but not interrupting posterior crenulate groove middorsally; crenulae extending over dorsal 0.2-0.4 of pronotum laterally; scuto-scutellar sulcus $4.0-5.0 \times wider$ than midlength; areola of propodeum variably obscured, short and triangular rather than pentagonal in topotypic paratype; precoxal sulcus occasionally extending to anterior margin of mesopleuron; fore wing stigma $3.3-3.8 \times longer$ than wide; $1RS 0.2-0.25 \times length$ of 1M; m-cu postfurcal by $0.15-2.0 \times length$ of m-cu; 2RS $0.8-1.05 \times length$

of 3RSa; head $1.85-2.2 \times$ wider than apex of T1; T1 $0.95-1.05 \times$ as long as apical width, apex $2.1-2.25 \times$ wider than base; surface of T1 between dorsal carinae weakly rugulose; dorsal carinae weakly sinuate, weakly converging at posterior margin of T1; S1 extending posteriorly only to level of dorsal tendon attachment; head varying from darker as in female to more extensively pale (as in Fig. 31) with ventral 0.5 of face orange, outer surface of mandible entirely dark orange and clypeus reddish brown; hind coxa varying from almost entirely orange to almost entirely black; hind femur and tibia varying from black to reddish brown.

Body length 4.9 mm (female), 3.1–4.7 mm (male), fore wing length 4.0 mm (female), 2.7–4.1 mm (male), mesosomal length 1.55 mm (female), 1.0–1.7 mm (male).

Diagnosis. This species is nearly identical to *D. hildagensis* based on the similarly long ovipositor and the notaulus that consistently extends all the way to the anterior margin of the mesoscutum. The eye is distinctly larger in *D. martinalujai* than in *D. hildagensis. Diachasmimorpha norrbomi* is also similar, but has a shorter ovipositor and the notaulus only rarely extends anteriorly to the margin of the mesoscutum.

Biology. This is the species that has been referred to as *Diachasmimorpha mexicana* (vide Wharton) in previous publications on parasitoids of *Rhagoletis* Loew in Mexico (e.g. Rull et al. 2009). The holotype and paratypes were all reared from Mexican populations of *Rhagoletis pomonella* infesting fruits of various species of *Crataegus*, including *C. mexicana* DC., as characterized by Xie et al. (2007).

Etymology. This species is named after Martin Aluja in recognition of his many contributions to tephritid biology, particularly in Mexico.

Remarks. The male paratypes, though only three in number, are remarkably variable in size, with larger individuals closely approaching the size of *D. hildagensis*. Quantitative measures are also highly variable, which is not surprising given the variation in size.

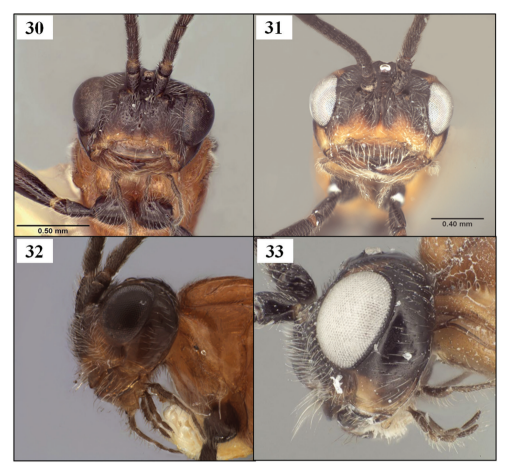
Detailed assessment of the available reared material suggests the presence of a diverse assemblage of *Diachasmimorpha* species in Mexico, associated with different hosts and host plants. The relatively small morphological differences between *D. hildagensis* and *D. martinalujai* are consistent among the available material and the differences in host and host plant associations lend support to the recognition of these as separate species.

Diachasmimorpha norrbomi Wharton, sp. n.

urn:lsid:zoobank.org:act:4900256F-3E99-41FC-8CCF-3A42E06D2033 http://species-id.net/wiki/Diachasmimorpha_norrbomi Figs 25, 29, 30, 32

Type locality. Mexico, State of Mexico, Parque Lago de Zempoala.

Type material. Holotype. Female (UNAM), first label, first line: Mexico, Parque second line: Lag. de Zempoala, path third line: along L. Zempoala, 10–11. fourth line: VIII.1989, A.L.Norrbom Second label, first line: reared ex. Euphranta second line: mexicana (Tephritidae) third line: ex. fruit of Ribes fourth line: pringlei Rose (89M13)



Figures 30–33. *Diachasmimorpha* spp., heads. 30 *D. norrbomi* Wharton, sp. n., paratype female, face 31 *D. martinalujai* Wharton, sp. n., paratype male, face 32 *D. norrbomi* paratype female, lateral view 33 *D. martinalujai*, holotype female, face.

Paratypes: 27 females, 20 males, same data as holotype, one of these with an additional ALN 31 label and a Biosteres sp. 1 det P. Marsh label (TAMU, UNAM, USNM).

Other specimens examined (not paratypes): 1 female, 1 male, Mexico, D.F., Delegacion Tlapan, Fracc. Tlapuente, 19.ix.2003, M. Aluja #50, reared from fruit of Granadilla (TAMU).

Description. *Female.* Head in dorsal view $1.25-1.30 \times$ broader than mesoscutum, $1.80-1.85 \times$ broader than face; eye in dorsal view $1.7-2.0 \times$ longer than temple, temples not receding, but width at eyes greater than width at temples; eye in lateral view $2.1-2.9 \times$ longer than temple. Facial midridge ending dorsally in short, very weak bifurcation between antennae. Frons irregularly rugulose along midline near bifurcation, otherwise polished, with moderately dense patch of decumbent, laterally-directed, white setae on either side of midline; bare on either side of ocellar field; width of ocellar field $1.0-1.2 \times$ distance from ocellar field to eye. Face $1.80-1.95 \times$ wider than

high; uniformly setose (as in Figs 30, 32), distinctly punctate, punctures separated by at least 1 × their diameter. Malar sulcus deep, complete; malar space about $0.9-1.0 \times$ basal width of mandible, $0.30-0.35 \times$ eye height. Clypeus $2.8-3.2 \times$ wider than high; very weakly convex, nearly flat. Occipital carina weak but distinct near base of mandible, short, extending dorsally to ventral margin of eye and often slightly beyond, not reaching mid eye height. Hypostomal carina extending as short but distinct flange below mandible. Antenna with 41-47 flagellomeres; first flagellomere $1.05-1.2 \times$ longer than second; $1.8-2.0 \times$ longer than wide.

Mesosoma 1.35–1.45 × longer than high; 1.85–1.95 × longer than wide; 1.35–1.40 × higher than wide. Pronope deep, large, interrupting posterior crenulate groove middorsally; crenulae extending along dorsal 0.2 of pronotum laterally within narrow, shallow groove; groove not margined anteriorly by carina; anterior margin of pronotum laterally sinuate, not abruptly excavated. Notauli deep anteriorly, gradually weakening posteriorly, extending posteriorly to level of tegula, not reaching long, narrow midpit, anterior end usually just short of and only rarely reaching anterior-lateral margin of scutum; mesoscutum usually without supra-marginal carina between base of notaulus and tegula, rarely with short, weak trace of a carina. Scuto-scutellar sulcus nearly rectangular, a little narrower medially; 4.2–4.8 × wider than midlength; crenulate-foveolate. Propodeum rugose, areola extending over posterior 0.8 but largely obscured by sculpture. Precoxal sulcus crenulate, widely separated from anterior margin of mesopleuron.

Wings. Fore wing stigma short, broad, discrete distally, $3.15-3.30 \times \text{longer}$ than wide; r1 arising from midlength of stigma; 1RS (excluding parastigma) $0.30-0.35 \times \text{length}$ of 1M; m-cu postfurcal by $0.2-0.3 \times \text{length}$ of m-cu; second submarginal cell distinctly converging distally; 2RS $1.0-1.2 \times \text{longer}$ than 3RSa; 2CUa $1.6-1.8 \times \text{longer}$ than 2cu-a; 1cu-a distad 1M by about $1.0 \times \text{its}$ length.

Metasoma not distinctly petiolate; head $1.6-1.9 \times$ wider than apex of T1. T1 $0.95-1.05 \times$ as long as apical width; strongly diverging apically, with apex $2.0-2.5 \times$ wider than base; surface smooth to weakly strigose posterior-medially, almost completed smooth laterally; dorsal carinae weakly converging, widely separated at posterior margin, strongly elevated over anterior 0.5, gradually weakening posteriorly; lateral carina weaker, extending distinctly ventrad spiracle, rounded and barely distinguishable posteriorad spiracle; spiracle at midlength of T1; dorsope absent but lateral and dorsal carinae elevated at junction, giving appearance of a slight depression; laterope deep; S1 very short, extending posteriorad to level of dorsal tendon attachment. T2 unsculptured, with sharp lateral margins. Ovipositor sheath $1.7-1.8 \times$ longer than mesosoma, setal pattern about as in *D. martinalujai*, with slightly greater density basally.

Color (Fig. 25). Very similar to *D. hildagensis*. Meso- and metasoma orange, except tegula black; head dorsally dark brown to black except for small orange spot on vertex adjacent eye, lower occiput mostly yellow-orange, similar in color to broad band extending through epistomal sulcus, clypeus, lower gena (often), and mandibles; clypeus usually with narrow, transverse brown band, mandible with apical teeth dark, rarely with entire mandible brownish; legs black except extreme base and most or all of dorsal side of hind coxa orange, joint between femora and trochantelli reddish orange.

Male as in female except head in dorsal view $1.3-1.4 \times broader$ than mesoscutum, $1.70-1.75 \times broader$ than face; eye slightly smaller, in dorsal view eye $1.45-1.60 \times longer$ than temple, in lateral view $1.9-2.4 \times longer$ than temple; antenna with 41-43 flagellomeres, first flagellomere $0.95-1.2 \times longer$ than second. Mesosoma slightly narrower, $1.95-2.05 \times longer$ than wide; $1.4-1.5 \times higher$ than wide; scuto-scutellar sulcus somewhat more variable in size, $4.0-5.5 \times wider$ than midlength. Fore wing stigma $3.1-3.4 \times longer$ than wide. T1 slightly smaller, head $1.9-2.2 \times wider$ than apex of T1, T1 $1.75-1.90 \times wider$ at apex than at base.

Body length 3.3-4.3 mm, fore wing length 3.5-4.1 mm, mesosoma length 1.15-1.65 mm.

Diagnosis. This species is similar in coloration to *D. hildagensis* and *D. martinalujai* but the ovipositor (with sheath $1.7-1.8 \times$ longer than mesosoma) is slightly but distinctly shorter and the notaulus only rarely extends all the way to the anterior margin. The notaulus always reaches the anterior margin in the other two species. *Diachasmimorpha norrbomi* is smaller and has a larger eye than *D. hildagensis*, and 2RS tends to be longer (relative to 3Ra) in *D. norrbomi* than in *D. hildagensis* and *D. martinalujai*.

Biology. The type series of *D. norrbomi* was reared from *Euphranta mexicana* Norrbom infesting fruits of *Ribes pringlei* Rose (Norrbom 1993). Two additional specimens that fit within the morphological limits of this species were reared from an unknown tephritid infesting *Passiflora ligularis* Juss.

Etymology. This species is named for Allen Norrbom, who reared many Opiinae from various fruit, stem, and flower-infesting tephritids in Mexico and Central America.

Remarks. Size variation in this species is similar to that exhibited by *D. martinalujai*, with males dominating the small end of the range.

Diachasmimorpha mexicana (Cameron)

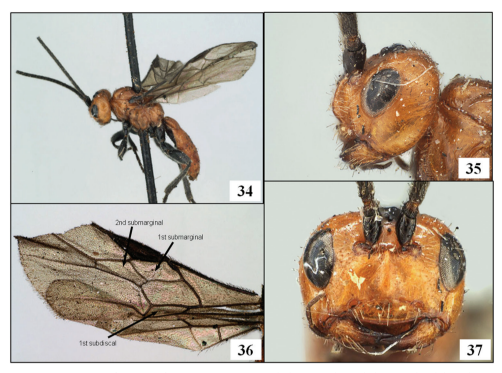
http://species-id.net/wiki/Diachasmimorpha_mexicana Figs 34–38

Opius mexicanus Cameron, 1887: 409–410. Holotype male in BMNH (examined). Desmiostoma mexicana: Fischer 1967b: 63–64 (redescription, generic transfer); Fischer 1977: 849, 872–873 (key, redescription).

Diachasmimorpha mexicana: Wharton 1997: 14 (generic transfer).

Type locality. Mexico, D. F., Chapultepec.

Type material. Holotype male (BMNH), first label [round, white with red margin], first line: Type second line: H. T. Second label, first line: B. M. TYPE second line: HYM third line: 3.°705 Third label, first line: B.C.A. Hymen. I. second line: Opius third line: mexicanus fourth line: Cam. Fourth label, first line: Opius second line: mexicanus third line: Cam. Type fourth line: BCA ii 409 Fifth label, first line: Bilimek second line: Mexico third line: 1871. fourth line: Chapul fifth line: tepek.



Figures 34–37. *Diachasmimorpha mexicana* (Cameron), holotype male. 34 habitus 35 head, lateral view 36 wings, showing names of cells used in descriptions 37 face.

Diagnosis. Holotype male. Eye in dorsal view shorter than temple, temples weakly expanded beyond eyes; eye in lateral view 0.95 × length of temple. Frons unsculptured along midline between antenna and median ocellus. Clypeus $3.4 \times$ wider than high. Occipital carina distinct near base of mandible, short, not extending dorsally to ventral margin of eye. Antenna broken. Pronope deep, large, interrupting posterior crenulate groove middorsally. Notauli deep anteriorly, reaching margin of mesoscutum anteriorly, apparently extending about half distance from anterior-lateral margin to elongate midpit but pin obliterates midpit and surrounding area of mesonotum. Precoxal sulcus very weakly crenulate, nearly smooth, short, not extending close to anterior margin of mesopleuron. Propodeum largely smooth, with rugulose sculpture largely confined to midline, especially around apex, and along border of metapleuron. Fore wing 2RS 0.8 × 3RSa; m-cu distinctly postfurcal. T1 with dorsal carinae widely separated, short, barely extending to level of spiracle, T1 otherwise unsculptured. Head, meso- and metasoma orange, tegula black; legs black as in holotype of D. hildagensis. Body length about 4.0 mm. This species has a much smaller eye (Figs 35, 37) than the similarly-colored D. sanguinea (Fig. 41) and is also less heavily sculptured. Females are unknown.

Biology. Unknown.

Remarks. The body of the *D. mexicana* holotype is remarkably smooth relative to that of other species in the *mexicana* species group. The precoxal sulcus, for example,

is very weakly crenulate, the propodeum is very weakly sculptured in general but completely smooth and polished anterior-laterally, and T1 is unsculptured except for the very short dorsal carinae. Sculpture is variable to some extent in other species of this species group, and thus it would be useful to obtain additional specimens of the true *D. mexicana* to determine the extent of sculptural variation in this species and ascertain whether reduction in sculpture is a useful diagnostic feature.

Fischer (1967b) noted that the specimen labeled as the type in BMNH is a male, but Cameron (1887) indicated in his original description that he was describing a female. The excellent figure in Cameron (1887) matches the type specimen, providing additional evidence of Cameron's error (either misinterpretation of the male genitalia as an ovipositor or, more likely given the general quality of Cameron's early work, a typographical error). The holotype was collected by D. Bilimek in Chapultepec and I have interpreted this as the large park that is now within Mexico City. Fischer (1967b) recorded the type label as type no. 3.c.505, but this is an inadvertent error. The type number for this specimens is 3.c.705.

See additional remarks under *D. hildagensis* above.

Diachasmimorpha sanguinea (Ashmead)

http://species-id.net/wiki/Diachasmimorpha_sanguinea Figs 4, 18, 39–41

- *Phaedrotoma (?) sanguinea* Ashmead, 1889: 655. Holotype female in USNM (examined). Marshall 1891: 47 (relationship to a European species of *Opius*).
- Opius sanguineus: Gahan 1915: 69, 74 (key, synonymy, expanded distribution and host); Muesebeck and Walkley 1951: 157 (synonymy, new distribution and host); Muesebeck 1967: 54 (catalog).

Opius (Biosteres) sanguineus: Fischer 1965: 116, 138-139 (key, redescription).

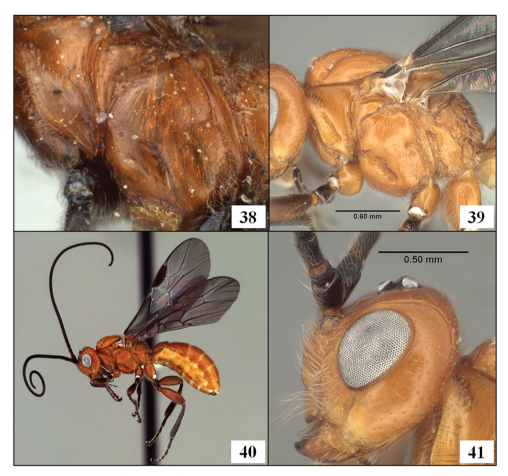
Biosteres sanguineus: Fischer 1971: 30 (catalog, change in rank); Wharton and Marsh 1978: 152, 156 (key, diagnosis, distribution, biology); Marsh 1979: 201 (catalog).

Biosteres (Chilotrichia) sanguineus: Fischer 1977: 804, 819–821 (key, redescription). *Diachasmimorpha sanguinea*: Wharton 1997: 14 (generic transfer).

Type locality. USA, Washington, D. C.

Type material. Syntype female (USNM), first label, first line: 3737[±] second line: Oct. 3. 85 Second label (red with black print), first line: Type second line: No2989 third line: U.S.N.M. Third label, first line: Phaedrotoma second line: sanguinea third line: Ashm ms. Syntype male, with same label data as syntype female except Third label = first line: Opius second line: sanguineus third line: Gahan Ashm Syntype male with first label, first line: 3737[±] second line: Aug. 5. 86 Second label: same as other two syntypes, no third label.

Other specimens examined. USA, Texas, 1 female, 1 male, Brazos Co., Yancey, xi.2010, emerged 9.iv & 3.v.2011, L. Ward, reared from *Zonosemata vittigera* infesting fruits of *Solanum eleagnifolium* (TAMU); 1 female, Hidalgo Co., Bentsen Rio Grande



Figures 38–41. *Diachasmimorpha* spp. 38 *D. mexicana* (Cameron) holotype male, mesopleuron 39 *D. sanguinea* (Ashmead), male mesosoma, lateral view 40 *D. sanguinea* habitus 41 *D. sanguinea*, male head, lateral view.

Valley State Park, 10.?.1978, C. Porter (TAMU); 5 females, 1 male, Hidalgo Co., Donna, J. W. Monk, reared from *Zonosemata vittigera*; 5 females, 1 male, Jeff Davis Co., 14 mi. S. Ft. Davis, 16–19.viii.1985, L. E. Carroll, reared from *Zonosemata* infesting fruits of *Solanum*; 5 males, Jeff Davis Co., Davis Mts. State Park, 12.vii.1995, R. Wharton; 1 female, Swisher Co., Happy, 17.viii.1977, W. F. Chamberlin.

Diagnosis. Male. Eye in dorsal view $1.1-1.3 \times \text{longer}$ than temple, temples not expanded beyond eyes; eye in lateral view $1.3-1.5 \times \text{longer}$ than temple. Frons between short, low, bifurcating ridges varying from unsculptured to irregularly strigose, frons otherwise smooth, polished. Clypeus $2.5-2.8 \times \text{wider}$ than high. Occipital carina distinct near base of mandible, short, not extending dorsally to ventral margin of eye. Antenna with 38–48 flagellomeres. Pronope deep, large, interrupting posterior crenulate groove middorsally. Notauli deep anteriorly, reaching margin of mesoscutum anteriorly, extending about half distance from anterior-lateral margin to elongate midpit. Precoxal

sulcus heavily sculptured, crenulate to foveolate, usually extending to or nearly to anterior margin of mesopleuron. Propodeum rugose, areola, when partially visible, extending over posterior 0.6-0.7 but frequently completely obscured by sculpture. Fore wing 2RS $0.9-1.05 \times$ length of 3RSa; m-cu distinctly postfurcal. T1 with dorsal carinae weakly converging, widely separated at posterior margin, gradually weakening posteriorly, T1 smooth to strigose between carinae. Head, meso- and metasoma orange; tegula orange to brown, legs varying from black except hind coxa mottled black and orange to more extensively orange. Female about as in male except eye in lateral view $1.2-1.6 \times$ longer than temple. Ovipositor sheath $1.6-1.75 \times$ longer than mesosoma. Body length 3.6-5.3 mm, fore wing length 3.3-4.6 mm, mesosoma length 1.2-1.9 mm. This species has a larger eye than the similarly-colored *D. mexicana* and is generally more heavily sculptured.

Biology. This species was originally described from several specimens reared from a tephritid infesting fruits of *Solanum carolinense* L. (Ashmead 1889). The tephritid host was later identified as *Zonosemata electa* (Say) (Gahan 1915). Muesebeck and Walkley (1951) added *Z. vittigera* (Coquillett) as a host and Cazier (1962) published on the biology of *Z. vittigera* with notes on parasitization by *D. sanguinea*. The only known host of *Z. vittigera* is *Solanum eleagnifolium* Cav. (Foote et al. 1993) and this is the host plant from which we have reared *D. sanguinea* in central and western Texas. Adult *D. sanguinea* are active in summer and fall in Texas, overwinter in the host puparium, and emerge the following year, over a period of several months.

Remarks. The diagnosis is based on the material from Texas listed in the other material examined section. Ashmead (1889) described this species from a single series of reared material, without designation of a type. The specimen in the type collection of the USNM is therefore a syntype, as are the remaining two specimens from this series in the general collection. There is no compelling reason to designate a lectotype, and we have therefore not done so. The original series is currently represented by 2 males and 1 female in the USNM collection. The syntypes agree in all essential details with the material from Texas, though the eye/temple ratio is at the smaller end of the range given above.

The sculpture is somewhat variable in this species, with smaller individuals having a tendency towards rugulose rather than rugose sculpture on the propodeum. The precoxal sulcus is always heavily sculptured, however, never approaching the reduction in sculpture seen in the holotype of *D. mexicana* (Fig. 39 vs. Fig. 38). The syntypes from Washington, D. C. are as variable in sculpture of the propodeum and T1 as are the specimens from Texas. Specimens from Texas, even within the same reared series, are exceptionally variable in leg coloration. The syntypes from Washington, D. C. have black legs with mostly orange hind coxa. Some specimens from Jeff Davis Co., Texas also have this pattern while in others only the tarsi are dark with the remaining parts orange. Similarly, the tegula is usually orange, but varies from orange to brown even within the same reared series.

Diachasmimorpha sanguinea is nearly identical to *D. mexicana* and additional material from the type locality of the latter is needed for a better understanding of the relationship between these two nominal species.

Eurytenes Wesmael

http://species-id.net/wiki/Eurytenes

Eurytenes (Stigmatopoea Fischer)

Opius (Stigmatopoea Fischer, 1986: 609–611). Type species: *Opius macrocerus* Thomson, 1895. Original designation.

Eurytenes (*Stigmatopoea*): Wharton 1988: 357 (revised status); Fischer 1998: 21–25 (subgeneric keys, diagnoses); Walker and Wharton 2011: 24 (review of classification).

Xynobius (Stigmatopoea): van Achterberg 2004: 314–315 (revised status, subgeneric keys). *Eurytenes (Xynobius)*: Wharton 2006: 330–333 (revised status, relationships).

Diagnosis. Mandible without basal lobe ventrally. Labrum broadly exposed. Occipital carina broadly absent dorsally, present laterally. Propleuron ventral-laterally without oblique carina. Notauli deep, well developed anteriorly, varying posteriorly from large-ly absent to deep and extending to scuto-scutellar sulcus or nearly so; midpit present. Fore wing stigma long, narrow, parallel-sided, discrete posteriorly, r1 arising distinctly basad its midpoint; second submarginal cell with 2RS shorter than 3RSb; 2CUb arising above middle of hind margin of first subdiscal cell. Dorsope present; S1 0.2–0.3 × length of T1, never fused to T1.

Remarks. The new species described below have been placed in *Eurytenes (Stig-matopoea)* based on the relative length of S1 (Figs. 5, 7) and the specific characteristics of T1 (Figs 5, 7, 54, 56, 57), wing venation (Fig. 64), mesoscutal sculpture (Figs 44, 48, 49), clypeus (Figs 50–53), and mandibles (Figs 50, 51) listed in the diagnosis. The wing venation is similar to that in *Lorenzopius* but in *Lorenzopius*, the dorsope is absent and S1 is longer and apparently fused to T1 (Fig. 6). We follow Wharton (1988, 2006) and Fischer (1998) in treating *Stigmatopoea* as a subgenus of *Eurytenes*. Wharton (2006) provides a detailed explanation of the morphological basis for this treatment as well as a discussion of alternative classifications.

Aulonotus Ashmead has usually been characterized on the basis of well-developed notauli (Fischer 1972, 1998), similar to the condition found in the species described below. *Aulonotus* shares other similarities with *Stigmatopoea*, including the presence of a dorsope, but the petiole is broader, S1 is very poorly developed, the stigma is not parallel-sided, and the precoxal sulcus is distinctly sculptured. Both the type species of *Stigmatopoea* and the two species described here will key to *Opius* (*Nosopoea* Foerster) in Fischer's classification of Opiinae (Fischer 1972, 1977) because the precoxal sulcus is unsculptured in nearly all individuals (as in Figs 43, 44). Difficulties in interpreting the variable nature of sculpture in the precoxal sulcus, and the emphasis placed on this character in existing keys to Opiinae, make it possible for relatively closely related species to become widely separated in current classifications.

Eurytenes (Stigmatopoea) maya Wharton, sp. n.

urn:lsid:zoobank.org:act:A5E2449E-78E5-48A3-B4CD-B4FC77A410A4 http://species-id.net/wiki/Eurytenes_maya Figs 7, 42, 44, 46, 48, 50, 52, 56, 59, 64

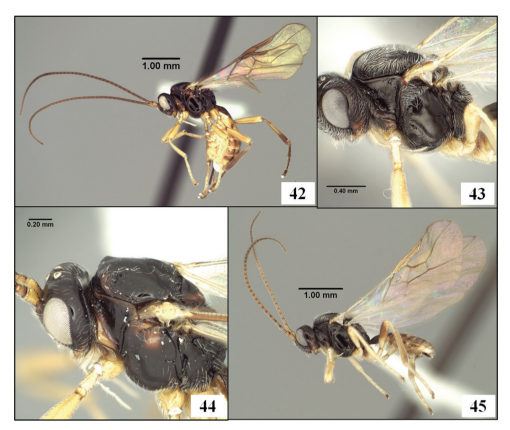
Type locality. Mexico, Chiapas, San Cristobal de las Casas.

Type material. Holotype. Female (TAMU), first label, first line: MEXICO: Chiapas second line: San Cristobal de las third line: Casas, xi.2001, #37A fourth line: J. Marquez, M. Aluja Second label, first line: host: Rhagoletis second line: pomonella third line: ex fruit of: fourth line: Crataegus mexicana

Paratypes: 2 females, same data as holotype but collected 26.xi.2001, #35A (TAMU); 1 female, same locality, 14.xi.2001, M. Aluja, Key 30A, host: *Rhagoletis* sp. on tejocote, manzanita (TAMU); 1 female, same locality, 14.xi.2001, J. Marquez, ex: *R. pomonella* on *Crataegus* sp., #27 (TAMU); 1 female, Chiapas, Rancho Nuevo, 5 km to San Cristobal de las Casas-freeway 190, 15.xi.2002, J. L. Marquez, M. Aluja, # 42, host: *Rhagoletis pomonella* ex fruit of *Crataegus mexicana* (TAMU); 2 males, Chiapas, 3 km E. San Cristobal, 15.xi.1994, R. Jones, ex pupa of *Rhagoletis pomonella* (TAMU); 3 females, Chiapas, Huixtan, 15.ix.2002, J. Marquez, Key 34, host: *R. pomonella* ex fruit of *Crataegus* sp. (TAMU); 1 male, 1 female, Chiapas, Cruz Quemada, 15.xi.2002, host: *Rhagoletis pomonella* ex fruit of *Malus* sp., J. Marquez, Key 35, and J. L. Marquez, M. Aluja, #45 (TAMU); 1 male, 1? (abdomen missing), Chiapas, Teopisca, 26.xi.2001, J. L. Marquez, ex: *R. pomonella* on *Crataegus* sp. #26 (TAMU).

Other specimens examined (not paratype): 1 male, Mexico: San Luis Potosi, Rio Verde, 7.x.2003, M. Pale, Key 71, *Rhagoletis* nr. *pomonella* on *Crataegus parrayana* (TAMU) [sequenced].

Description. *Female.* Head in dorsal view 1.25–1.30 × broader than mesoscutum, $1.80-1.95 \times$ broader than face; eye in dorsal view $2.5-3.2 \times$ longer than temple, temples distinctly receding behind eyes. Frons and vertex highly polished, unsculptured except for shallow, median depression between toruli; frons bare, vertex and occiput with a few, short, scattered setae; width of ocellar field $1.05-1.3 \times distance$ from ocellar field to eye. Face 1.55-1.70 × wider than high; slightly less polished than frons; uniformly setose (as in Figs 50, 52), with very fine punctures, these separated by at least 2 × their diameter. Frons and face delimited by slight change in sculpture resulting in weak, shallow sulcus between torulus and eye; distance between antennal toruli equal to distance from torulus to eye, eye not distinctly emarginate in region of antenna. Malar sulcus deep, complete; malar space about 0.5 × basal width of mandible, 0.2 × eye height. Face weakly convex, bulging slightly medially along the low midridge. Epistomal sulcus weak mid-dorsally, more distinct laterally. Clypeus 2.2-2.5 × wider than high; weakly convex, slightly protruding in profile; ventral margin sharp, truncate to very weakly concave in frontal view. Labrum broadly exposed, gap between ventral margin of clypeus and dorsal margin of mandible varying from $0.5-1.0 \times$ height of clypeus, depending on how tightly closed the mandibles are. Occipital carina distinctly curved medially at dorsal end, broadly absent mid-dorsally, the space where the carina



Figures 42–45. *Eurytenes (Stigmatopoea)* spp. 42 *E. (S.) maya* Wharton sp. n., paratype female, habitus 43 *E. (S.) norrbomi* Wharton sp. n., holotype female, mesosoma 44 *E. (S.) maya*, paratype female, head and mesosoma, dorsal-lateral view 45 *E. (S.) norrbomi*, holotype female, habitus.

is absent distinctly wider than width of ocellar field; occipital and hypostomal carinae widely separated at base of mandible, the latter extending as a flange beneath about basal 0.2 of mandible. Mandible without basal lobe ventrally; bidentate apically, lower tooth much smaller than dorsal tooth and slightly twisted beneath dorsal tooth; ventral margin carinate throughout. Antenna $1.35-1.45 \times longer$ than fore wing, with 39-43 flagellomeres; first flagellomere $1.1-1.3 \times longer$ than second, $1.2-1.3 \times longer$ than third; flagellomeres $2.3-2.7 \times longer$ than wide basally, twice longer than wide apically. Maxillary palps a little longer than head height; fifth and sixth segments equal in length or nearly so, fourth segment $1.1-1.15 \times longer$ than both fifth and sixth.

Mesosoma $1.4 \times \text{longer}$ than high; $1.9 \times \text{longer}$ than wide; $1.35-1.40 \times \text{higher}$ than wide. Pronotum dorsally a narrow, polished, smooth band with crenulate groove along posterior margin; rarely with discernible, slightly enlarged pit in middle of crenulate groove; crenulae extending in narrow, shallow groove onto pronotum laterally, but only covering dorsal 0.2-0.4; groove margined anteriorly by sharp carina that continues ventrally along full length of pronotum. Anterior declivity of mesoscutum completely vertical, bare or nearly so; anterior-lateral corners of mesoscutum at upper edge

of declivity elevated, rounded, sparsely setose; notauli extending 0.4 × distance from anterior declivity to scuto-scutellar sulcus, extending posteriorly from lateral side of elevated anterior-lateral corners, not extending to mesoscutal margin anteriorly, very weakly converging posteriorly; narrow, crenulate throughout; mesoscutum with distinct supra-marginal carina extending from elevated anterior-lateral corner to tegula. Lateral and median mesoscutal lobes bare except scattered setae along notauli; midpit deep, round to somewhat elongate, never extending to notauli. Scuto-scutellar sulcus nearly rectangular, a little narrower medially; $3.75-4.25 \times$ wider than midlength; crenulate-foveolate, with 7 ridges; all sides vertical, clearly delineated. Scutellum very weakly convex, nearly flat, not strongly elevated; bare except for scattered setae posteriorly; unsculptured, even along posterior margin. Propodeum with median carina over anterior 0.3, bifurcating at this point to form an inverted v-shaped transverse carina extending to pleural carina just posteriad spiracle; pleural carina complete from base to apex though sometimes partly obscured by sculpture posteriad spiracle; lateral longitudinal carina parallel to and narrowly separated from pleural carina anteriad spiracle, more medially displaced when visible posteriad transverse carinae, forming part of broad areola; area between pleural and lateral longitudinal carinae rugose and sparsely setose anteriorly; lateral propodeal areas anteriorly on either side of median carina smooth, bare, unsculptured; areola broad, varying from distinct (with surface irregularly, weakly rugulose) to indistinct (surface rugose, disrupting carinate margin of areola); lateral propodeal areas posteriorly varying from nearly unsculptured and distinct to rugose and indistinct; propodeum largely bare medially, with a few scattered setae. Mesopleuron largely bare, with sparse setae in unsculptured subalar region and a small patch of setae dorsad mid coxa; posterior margin unsculptured. Precoxal sulcus weakly impressed but distinct; unsculptured. Metapleuron bare on dorsal half except for small patch below wing, with a few long setae medially, and patches of setae among rugulose sculpture along ventral margin and in groove on ventral half of anterior margin; otherwise unsculptured.

Metasoma distinctly petiolate; head 3.5–3.8 × wider than apex of T1. T1 2.15–2.35 × longer than apical width; nearly parallel-sided, with apex 1.20–1.35 × wider than base;



Figures 46–49. *Eurytenes (Stigmatopoea)* spp. **46** *E. (S.) maya* Wharton sp. n., paratype female, head, lateral view **47** *E. (S.) norrbomi* Wharton sp. n., holotype female, head, lateral view **48** *E. (S.) maya*, paratype female, head and mesosoma, dorsal view **49** *E. (S.) norrbomi*, holotype female, dorsal view.

surface striate throughout, above and below lateral carina; one or two very shallow, subapical depressions usually present dorsally; dorsope distinct, deep; laterope completely absent; dorsal carina present only at base, lateral carina usually distinct throughout; spiracle positioned 0.6 × length of T1 from the base; S1 extending about 0.25–0.30 × length of T1; dorsal surface of petiole in profile evenly convex from base to apex. T2 and following without sharp lateral margins; spiracle of second metasomal tergum laterally displaced, not visible in dorsal view. Ovipositor as long as mesosoma; ovipositor sheath 0.6–0.7 × length of mesosoma, with 2–3 irregular rows of long setae along its length.

Color: head, including antenna, mesosoma, petiole and ovipositor sheath dark brown except scape yellow; mandible, lower gena, ventral portion of clypeus, pedicel (occasionally), face adjacent antennal base, propleuron, anterior margin of pronotum, spot on mesopleuron below wing and a smaller spot above mid coxa, two streaks on either side of midpit on mesoscutum, posterior margins of scutellum and metapleuron, and petiole laterally (occasionally) dark yellow to orange; palps pale yellow, nearly white. Legs and metasoma beyond T1 yellow except hind tibia, hind tarsi, lateral margin of metasomal terga 2 + 3 and often anterior half of terga 4–6 brown, the hind tibia often paler medially.

Male. As in female except antenna with 41-45 flagellomeres, head $4.0-4.6 \times$ wider than apex of T1 and T1 2.5–2.9 × longer than apical width. Body somewhat darker in color, with metasomal terga 6, 7, and most or all of 5 dark brown.

Body length 3.2–4.3 mm; wing length 3.5–4.2 mm.

Diagnosis. This species runs to *Opius (Nosopoea)* in Fischer (1972, 1977) on the basis of the exposed labrum, distinct midpit on the mesoscutum, and absence of sculpture within the precoxal sulcus. It differs from described species placed in the subgenus *Nosopoea* by the combination of larger size, more numerous flagellomeres, relatively well-developed notauli (Fig. 44), parallel-sided T1 (Fig. 56), and parallel-sided stigma (Fig. 64), all characters which it shares with the type species of *Stigmatopoea, Eurytenes (Stigmatopoea) macrocerus.* In *E. maya* the anterior declivity of the mesoscutum is more vertical and the anterior-lateral corners of the mesoscutal disc are distinctly elevated (Fig. 44) in comparison to *E. macrocerus. Eurytenes maya* differs from the other species described below, *E. norrbomi*, sp. n., by the possession of a relatively longer ovipositor (Fig. 42 vs. Fig. 45) and a less densely setose mesoscutum (Fig. 44 vs. 43).

Biology. All specimens were reared from Mexican populations of *Rhagoletis pomonella* (Walsh) infesting either hawthorns (species of *Crataegus* L.) or apples (*Malus domestica* Borkh.).

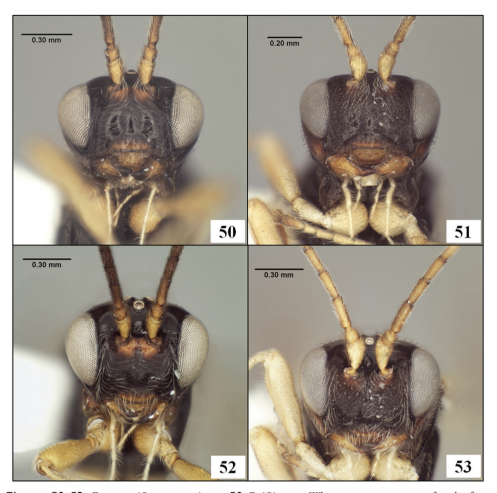
Etymology. The species name is in reference the Mayan Indians of this region.

Remarks. This species is similar in general appearance to members of the genus *Lorenzopius*, but T1 is not distinctly tubular as it is in the latter genus (see discussion below under *Lorenzopius*). The overall resemblance to *Lorenzopius* is enhanced by the presence of weak depressions on T1 that are similar in position in *E. maya* and *L. calycomyzae* van Achterberg and Salvo (Figs 55, 56). The depressions are variable within members of the same reared series of *E. maya*: being absent, for example, in the holotype, but well developed in some of the paratypes.

The limited information on hosts suggests that species with a more tubular petiole, such as those in *Lorenzopius*, are parasitoids of leaf-mining Agromyzidae while the species of *Stigmatopoea* attack both leaf-mining and fruit-infesting tephritids.

Eurytenes (Stigmatopoea) norrbomi Wharton, sp. n. urn:lsid:zoobank.org:act:FC57AAB3-6290-4076-BB44-6289F8ABF2EC http://species-id.net/wiki/Eurytenes_norrbomi Figs 43, 45, 47, 49, 51, 53, 57

Type locality. Mexico, Morelos, Km. 9–10 between Huitzilac and Lago Zempoala.



Figures 50–53. *Eurytenes (Stigmatopoea)* spp. 50 *E. (S.) maya* Wharton sp. n., paratype female, face, frontal view 51 *E. (S.) norrbomi* Wharton sp. n., holotype female, face, frontal view 52 *E. (S.) maya*, paratype female, face, slightly deflected 53 *E. (S.) norrbomi*, holotype female, face, slightly deflected.

Type material. Holotype. Female (UNAM), first label, first line: MEXICO: Morelos second line: Km 9–10, btw. Huitzilac third line: & Lago Zempoala fourth line: roadside, 22–24.ix.1991 fifth line: A. L. Norrbom #42

Paratypes: Mexico, 4 females, same data as holotype (TAMU, USNM); 1 female, Mexico, Rt. 890, Km 9 area, 6 km W Lago Zempoala 2.x.1991, Norrbom, #43, reared ex. *Trypeta concolor* ex. leafmines on *Barkleyanthus salicifolius* (91M1D) (TAMU). 3 males, Distrito Federal, Rt. 95 (libre), Km 42–43, 1 km N. La Cima, 20–26.ix.1991 A. L. Norrbom, #41, reared ex. *Trypeta concolor* ex. leafmines on *Barkleyanthus salicifolius* (91M1) (TAMU, USNM).

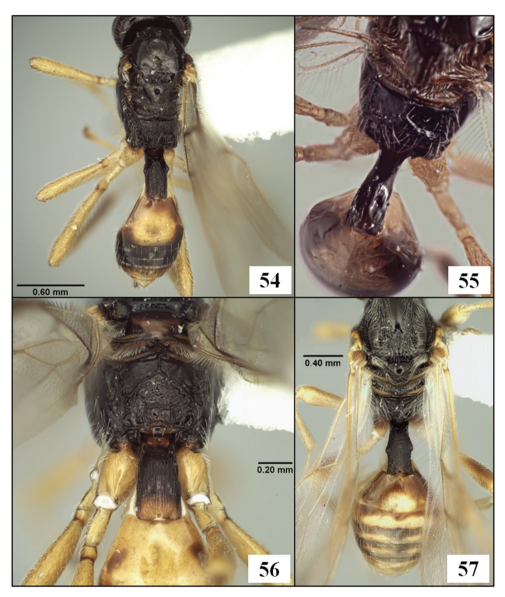
Description. *Female.* Head in dorsal view $1.2-1.3 \times$ broader than mesoscutum, $1.75-1.85 \times$ broader than face; eye in dorsal view $1.2-1.5 \times$ longer than temple, temples weakly receding behind eyes. Frons and vertex as in *E. maya* except vertex and outer part

of occiput densely covered with long, decumbent setae; width of ocellar field $1.20-1.35 \times$ distance from ocellar field to eye. Face $1.75-1.85 \times$ wider than high; slightly less polished than frons; uniformly setose (as in Figs 51, 53), distinctly punctate, the punctures separated by about 1 × their diameter. Frons and face delimited by a slightly more distinct change in sculpture in area between torulus and eye. Malar space about $0.6 \times$ basal width of mandible, $0.25 \times$ eye height. Clypeus $3.0-3.4 \times$ wider than high; protruding in profile. Occipital carina distinctly curved medially at dorsal end, absent mid-dorsally, the space where the carina is absent approximating width of ocellar field. Antenna $1.15-1.30 \times$ longer than fore wing, with 31-33 flagellomeres; first flagellomere $1.05-1.10 \times$ longer than second, $1.05-1.20 \times$ longer than third; flagellomeres $3.1-4.1 \times$ longer than wide basally, 2.3-2.7 longer than wide apically. Head otherwise as described for *E. maya*.

Mesosoma 1.35-1.45 × longer than high; 1.8-1.9 × longer than wide; 1.3-1.4 × higher than wide. Pronotum dorsally as in E. maya but with slightly enlarged pit in middle of crenulate groove consistently present; crenulae extending in shallow groove onto pronotum laterally, covering dorsal 0.2-0.6; groove margined anteriorly as in E. maya. Anterior declivity of mesoscutum completely vertical, densely covered with white, decumbent setae except for bare median band extending posteriorly to midpit; anterior-lateral corners of mesoscutum at upper edge of declivity elevated, rounded, densely setose, the setal pattern extending in broad bands all along notauli and laterally from anterior declivity to tegula; notauli complete, extending from anterior margin to scuto-scutellar sulcus, weakly converging posteriorly alongside but not into tear-drop shaped midpit; crenulate throughout, with sculpture extending laterally around margin to tegula, sculpture largely obscured by dense setae; lateral lobes of mesoscutum bare posterior-medially. Scuto-scutellar sulcus $4-5 \times$ wider than midlength, lateral margins difficult to discern due to setal density; with low midridge and indistinct crenulae on either side; otherwise as in E. maya. Scutellum as in E. maya except with long marginal setae extending medially to cover most of posterior 0.5. Propodeum extensively rugulose, obscuring nearly all traces of carinae; pleural carina weak, often indistinct, very short median carina often present basally; transverse carina rarely weakly indicated across middle; propodeum uniformly setose anteriorly, with a few scattered setae posteriorly. Mesopleuron as in E. maya except subalar region densely setose and groove below subalar ridge varying from nearly smooth to weakly rugulose. Precoxal sulcus distinctly impressed, unsculptured. Metapleuron a little more extensively setose but otherwise as in E. maya.

Wings. Fore wing stigma parallel-sided, discrete posteriorly, $6.3-6.6 \times longer$ than wide; r1 arising from basal 0.35; 1RS (excluding parastigma) 0.25-0.35 × length of 1M; RS+M weakly sinuate; 3RSa 1.05-1.30 × longer than 2RS; 2RS 2.6-3.1 × longer than r; 2RS and 3RSb straight; 3M variable, but often pigmented and sclerotized for most of its length; 2CUa 0.8-0.9 × length of 2cu-a, 2CUb arising slightly above middle of first subdiscal cell; position of m-cu, 1cu-a, and 1-1A, shape of second submarginal cell, and angle between r1 and 2RS as in *E. maya*. Hind wing as in *E. maya*.

Metasoma distinctly petiolate; head $3.75-4.10 \times$ wider than apex of T1. T1 2.2– 2.5 × longer than apical width; nearly parallel-sided, with apex $1.20-1.35 \times$ wider than base; surface granular coriaceous throughout; completely without subapical depres-



Figures 54–57. *Eurytenes (Stigmatopoea)* and *Lorenzopius*. 54 *E. (S.) macrocerus* (Thomson), mesosoma and metasoma, dorsal view 55 *L. calycomyzae* van Achterberg and Salvo, holotype female, T1, dorsal view 56 *E. (S.) maya* Wharton sp. n., paratype female, propodeum and T1, dorsal view 57 *E. (S.) norrbomi* Wharton sp. n., holotype female, mesosoma and metasoma, dorsal view.

sions dorsally; dorsope, laterope, dorsal carinae, dorsal surface of T1 in profile, as in *E. maya*; lateral carina at least partially present but difficult to distinguish from surrounding sculpture. S1 extending about $0.25-0.30 \times$ length of T1; T2 and following without sharp lateral margins; spiracle of second metasomal terga laterally displaced, only partially visible in dorsal view. Ovipositor shorter than mesosoma, base not visible

in type series, but total length approximately $0.6-0.7 \times \text{length of mesosoma}$; ovipositor sheath $0.30-0.35 \times \text{length of mesosoma}$, with setal pattern as in *E. maya*.

Color: Mesosoma, T1, S1, ovipositor sheath, and most of head dark brown to black; antenna yellow basally, apical 0.3 brown; mandibles yellow; palps white; lower gena adjacent malar sulcus brown to brownish red; ventral 0.3–0.4 of clypeus yellow to brownish red. Tegula reddish brown with yellow margin. Legs yellow to pale yellow except most of hind coxa, apical 0.6–0.7 of hind femur, and fifth tarsomere of all legs brown; hind tibia varying from weakly infumate to light brown, basal 0.2 nearly always pale yellow. T2 mostly brownish red with median yellow blotch posteriorly; T3 yellow with anterior and lateral margins brownish red; T4-T6 yellow with anterior and lateral margins dark brown; visible parts of remaining terga yellow.

Male. As in female except antenna with 37 flagellomeres; eye in dorsal view $1.55-1.75 \times longer$ than temple; width of ocellar field $1.05-1.10 \times longer$ from ocellar field to eye. Color same except visible parts of apical terga dark brown.

Body length 2.8–3.5 mm; wing length 3.2–3.6 mm.

Diagnosis. This species shares with *E. maya* and *E. macrocerus* the diagnostic features noted above for *Stigmatopoea*. *Eurytenes norrbomi* is most readily differentiated from *E. maya* on the basis of the more densely setose head and body (Figs 43, 47, 49), particularly the vertex, occiput, and mesoscutum, and the more extensively rugose propodeum. It also has a shorter ovipositor than *E. maya* (Fig. 45 vs. Fig. 42). The setal pattern on the mesoscutum also differentiates *E. norrbomi* from *E. macrocerus*. The latter has shorter setae that are more sparsely distributed laterally (Fig. 54).

Biology. Four of the specimens from the type series were reared from puparia of *Trypeta concolor* (Wulp) (Tephritidae) mining leaves of *Barkleyanthus salicifolius* (H.B.K.) H. Robins & Brett (Asteraceae). The remaining specimens were collected from flowers of this same plant together with *T. concolor* and *T. reducta* Han and Norrbom. See Han and Norrbom (2005) for details on the hosts and the collecting localities.

Etymology. This species is named after the collector, Allen Norrbom, who has provided many valuable host records for tephritid parasitoids.

Remarks. This species attacks leaf-mining tephritids, as does *E. macrocerus*, while *E. maya* attacks fruit-infesting tephritids. Despite the difference in host habitat, all three species share many morphological features, and readily fit the characterization of *Eurytenes (Stigmatopoea)* as given above.

Lorenzopius van Achterberg & Salvo

http://species-id.net/wiki/Lorenzopius

Lorenzopius van Achterberg & Salvo, 1997: 190–192. Type species: Lorenzopius calycomyzae van Achterberg & Salvo, 1997. Original designation.

Diagnosis. Mandible distinctly narrowed from base to apex, without basal lobe ventrally. Labrum exposed. Clypeus relatively flat, not distinctly protruding in profile; ventral margin sharp, truncate to weakly concave. Malar sulcus a sharp, weakly curved groove. Occipital carina broadly absent dorsally, present laterally; widely separated from hypostomal carina ventrally. First flagellomere longer than second. Propleuron ventrallaterally without oblique carina; pronotum dorsally without pronope or otherwise enlarged pit, posterior margin transversely rugulose. Notauli deep, narrow, well developed anteriorly, usually extending onto disc posteriorly; midpit present. Precoxal sulcus distinctly impressed. Propodeum with large areola, posterior portion often obscured by rugose sculpture. Fore wing stigma long, narrow, parallel-sided, discrete posteriorly, r1 arising distinctly basad its midpoint but not from extreme base; m-cu entering base of second submarginal cell; second submarginal cell with 2RS shorter than 3RSb; 2CUb arising above middle of hind margin of first subdiscal cell. Dorsope and laterope of T1 absent; S1 at least 0.7 × length of T1 in females, slightly shorter in males, apparently fused to T1; T1 long and narrow throughout; T2 and following terga unsculptured. Ovipositor tapering evenly to a fine point, without dorsal nodes or ridges.

Remarks. Lorenzopius and Tubiformopius are both characterized by having a tubular petiole with a long S1 which appears fused to T1 (Figs 6, 8). In the material available, S1 is longer in Lorenzopius than in Tubiformopius but there are more significant differences in the shape of the mandible, wing venation, and mesoscutal sculpture, as noted above in the section discussing genus group characters. Lorenzopius also shares many features with Eurytenes (Stigmatopoea), but the petiole is less tubular in the latter, with a distinctly shorter S1 that is clearly separated by membrane from T1 (Fig. 5).

The shape of the stigma has been proposed as a useful feature for assessing relationships among opiines (Wharton 1988), and both *Lorenzopius* and *Stigmatopoea* have the stigma more or less parallel-sided or slightly expanded distally. Unfortunately, the stigma often curls as specimens dry after death, and this feature then not only becomes difficult to assess properly, but is often illustrated in the curled position giving a misleading impression of the true form. For example, the shape of the stigma is difficult to discern on the holotype of *L. calycomyzae* (Fig. 62). However, the shape is more readily discernible in the holotype of *L. tubulatus* (Fig. 68) and in several other specimens of *Lorenzopius* available for examination (from CNC and TAMU), and these clearly show a parallel-sided stigma.

We recognize two distinct species groups within *Lorenzopius*: the *calycomyzae* species group containing the orginially included species *L. calycomyzae*, *L. tubulatus*, and *L. sanlorenzensis* and a second group typified by *L. euryteniformis* (Fischer), new combination. All have same basic wing venation and petiole. The precoxal sulcus is distinctly sculptured in the *calycomyzae* species group (Fig. 69) but the distinctly impressed sulcus is unsculptured or nearly so in the *euryteniformis* species group (Fig. 66). The smallest specimens of the *calycomyzae* species group examined during this study are slightly larger than the largest available specimens of the *euryteniformis* species group and perhaps as a consequence they tend to have slightly longer notauli and more sculpture bordering the supra-marginal carina extending from the base of the notaulus to the tegula. Most of the species we have examined from the *euryteniformis* species group have reduced propodeal sculpture with the areola clearly visible (Figs 72,

73). In addition to holotypes of *L. tubulatus* and *L. sanlorenzensis* and the holotype and paratypes of *L. calycomyzae*, we have seen two additional specimens from Argentina (TAMU), and one specimen each from Peru and Costa Rica (both CNC) representing the *calycomyzae* species group. RAW has examined 17 specimens representing the *euryteniformis* species group in addition to the holotype of *L. euryteniformis*. The material examined includes specimens housed in TAMU and CNC collected in Bolivia, Colombia, Costa Rica, Dominican Republic, Guatemala, and Mexico (as far north as Monterrey in Nuevo Leon).

Lengthy descriptions (Fischer 1963, 1964, 1979, van Achterberg and Salvo 1997) and some redescriptions (Fischer 1977) are available for the described species of *Lorenzopius* and van Achterberg and Salvo (1997) provide a useful key to the species of the *calycomyzae* species group. Species pages for *L. calycomyzae* (Figs 6, 55, 58, 60–62), *L. tubulatus* (Figs 3, 68, 69) and *L. euryteniformis* (Figs 66, 67, 70–73) can be found at http://peet.tamu.edu/projects/8/public/site/wharton_lab/home. The described species are readily differentiated. T1 is exceptionally long and narrow in *L. tubulatus* (at least 4 × longer than apical width) and this species has darker legs than the others, with most of the hind femur dark brown. T1 is about 3 × longer than apical width in the other two species of the *calycomyzae* species group and the hind femora are yellow. The presence of a pair of pits on T1 is thus far a unique feature of *L. calycomyzae* within *Lorenzopius* and this species is also characterized by orange markings dorsally in the middle of the mesosoma. The metasoma is darker in *L. sanlorenzensis*, with T2+3 dark brown in this species and largely yellow in the other two members of the *calycomyzae* species group. *Lorenzopius euryteniformis* lacks sculpture within the depression of the precoxal sulcus.

The type species of *Lorenzopius* was described from specimens reared from *Caly-comyza mikaniae* Spencer, a leafminer in the family Agromyzidae. RAW has also seen specimens from Colombia of a species nearly identical to *L. euryteniformis* that was also reared from an agromyzid leafminer. No other host records are known for this genus but given the general similarity of the habitus and the length and shape of the ovipositor, we predict that other species will also prove to be agromyzid leafminer parasitoids.

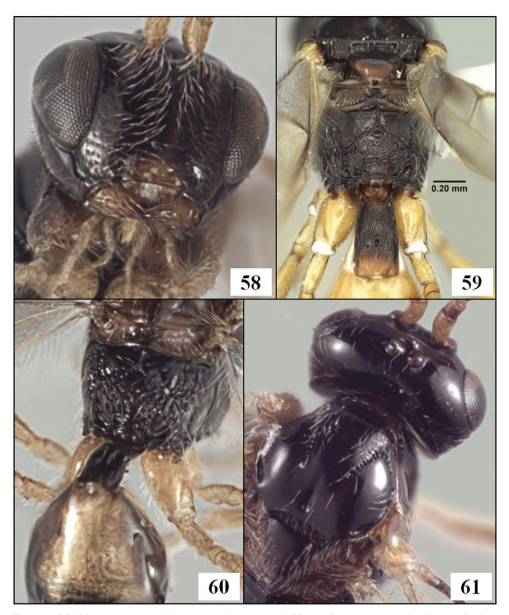
Lorenzopius euryteniformis Fischer, 1963, comb. n.

http://species-id.net/wiki/Lorenzopius_euryteniformis Figs 66, 67, 70–73

Opius euryteniformis Fischer, 1963: 288–290. Holotype male NHMW (examined). *Opius (Nosopoea) euryteniformis*: Fischer 1977: 195, 206–208.

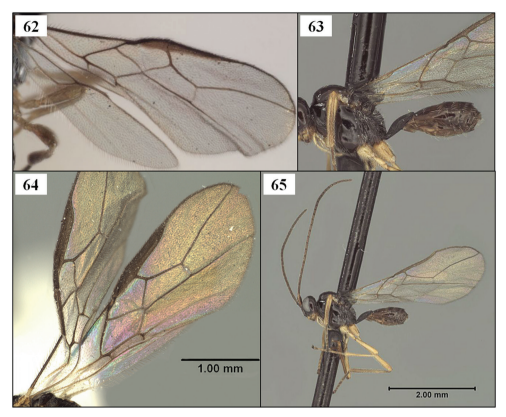
Type locality. Costa Rica, Mount Irazu, 2200–2300 m.

Type material. Holotype. Male (NHMW), first label, first line: Costa Rica, Irazu, second line: 2200–2300 m, 21–28. third line: V.'30. Reimoser Second label, first line: Opius second line: euryteniformis third line: sp. n. fourth line: det. Fischer Third label: Holotype [purple], Fourth label: NHMW



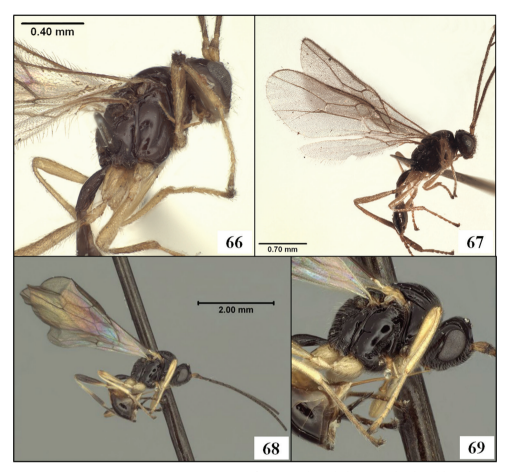
Figures 58–61. *Lorenzopius* and *Eurytenes (Stigmatopoea)*. 58 *L. calycomyzae* van Achterberg and Salvo, holotype female, face 59 *E. (S.) maya* Wharton sp. n., propodeum and T1 60 *L. calycomyzae*, propodeum 61 *L. calycomyzae*, head and mesoscutum, dorsal view.

Diagnosis. Holotype male. Head in dorsal view with temples neither receding nor expanded beyond eyes; in lateral view, eye about 1.6 × longer than temple. Labrum partly exposed between clypeus and mandibles (Fig. 70); clypeus about twice as wide as tall, flat or nearly so, not distinctly protruding in profile, ventral margin truncate to very weakly concave. Mandible without basal lobe. Malar space well developed,



Figures 62–65. Opiinae. **62** *Lorenzopius calycomyzae* van Achterberg and Salvo, holotype female, wings **63** *Tubiformopius tubigaster* (Fischer), holotype male, lateral view showing T1, S1, and wing base **64** *Eurytenes* (*Stigmatopoea*) *maya* Wharton, sp. n., paratype male, fore wing **65** *T. tubigaster*, holotype male, habitus.

longer than basal width of mandible; malar sulcus deeply impressed. Antenna with 27 flagellomeres. Pronotum dorsally not visible in holotype. Disc of mesoscutum nearly bare, with scattered setae along margin of anterior declivity and a single pair of setae arising about midlength of notauli; notaulus extending posteriorly along anterior 0.3 of disc, less than half distance to small, deep, round midpit; supra-marginal carina distinct anteriorly, not extending to level of tegula. Scuto-scutellar sulcus relatively narrow (Fig. 71), densely crenulate throughout. Precoxal sulcus distinctly impressed, long, narrow, completely unsculptured. Propodeum largely smooth with broad, pentagonal areola on posterior 0.65, anterior 0.35 with median carina. Fore wing stigma long, narrow, with some postmortem curling, but at least 4.5×1000 km width at r1; r1 arising from basal 0.3; second submarginal cell long, weakly converging distally, 3RSa 1.7 × longer than 2RS; 1RS 0.2 × length of 1M; m-cu postfurcal; 2CUb arising a little above middle of hind margin of first subdiscal cell, 2cu-a present, tubular. T1 long, narrow, apparently fused ventrally with S1 for most of its length, 4x longer than apical width, apex as wide as base; surface completely striate. T2 and following smooth, polished.

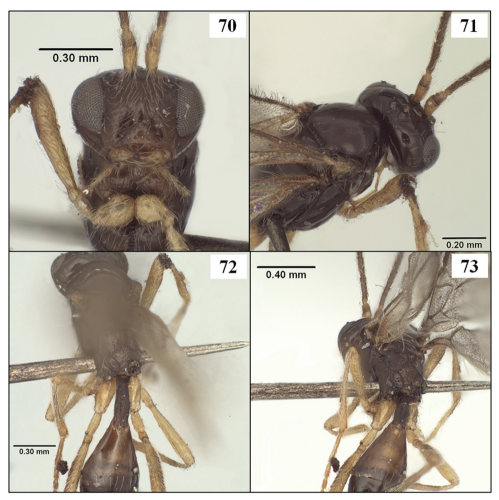


Figures 66–69. *Lorenzopius* spp. 66 *L. euryteniformis* (Fischer), holotype male, dorsal-lateral view showing unsculptured precoxal sulcus 67 *L. euryteniformis*, holotype male, habitus 68 *L. tubulatus* (Fischer), holotype female, habitus 69 *L. tubulatus*, holotype female, mesosoma, lateral view showing sculptured precoxal sulcus.

Biology. Unknown.

Remarks. Placement of this species in *Lorenzopius* is based on the wing venation and long S1, which is $0.65 \times$ length of T1 in the male holotype; S1 appears fused to T1. See additional comments on species groups under the remarks section for the genus.

The holotype bears a single data label containing the information given above. However, the label data listed in the original description are as follows: "Costa Rica, La Caja bei San José, H. Schmidt". As this species was described from a single male specimen, and the specimen from Irazu labeled as the holotype matches the original description, it is likely that the locality data in the original publication is an inadvertent error. The new species described immediately before *euryteniformis* in the same publication is from the La Caja locality. The type locality should therefore be Irazu (a mountain in Costa Rica), somewhere in the 2200–2300 m range in elevation.



Figures 70–73. *Lorenzopius euryteniformis*, holotype male. 70 face 71 mesoscutum and head, dorsallateral view 72 propodeum and T1–3 73 propodeum and metasoma.

Opius Wesmael

http://species-id.net/wiki/Opius

Opius Wesmael, 1835: 115. Type species: *Opius pallipes* Wesmael, 1835. Subsequent designation (Wharton 1987b, ICZN 1988).

Remarks. Van Achterberg and Salvo (1997) restricted the name *Opius* to species with a basal lobe on the mandible, referring to a classification in press that has yet to be published. A major concern in this regard is that the type species of *Biosteres* Foerster, another large genus within the Opiinae, also has a basal mandibular lobe. Until a more complete classification is offered, we prefer to treat *Opius* in a much broader sense as

a repository for the bulk of the Opiinae whose relationships remain uncertain, largely following the approach of Fischer (1972) and Wharton (1997).

Diagnoses are presented below for two species that represent a fairly diverse group of neotropical Opiinae that differ from both *Lorenzopius* and *Tubiformopius* in several features. These species all have a narrow, parallel-sided T1 and distinctly visible S1, though S1 is never as long as in *Lorenzopius*, and seldom as long as in *Tubiformopius*. Ultimately, the relationships of genus group taxa such as *Eurytenes* s.l., *Lorenzopius*, and *Tubiformopius* will have to be carefully considered in order to place the many neotropical species with a distinct S1.

Opius incoligma Fischer

http://species-id.net/wiki/Opius_incoligma Figs 74–77, 81

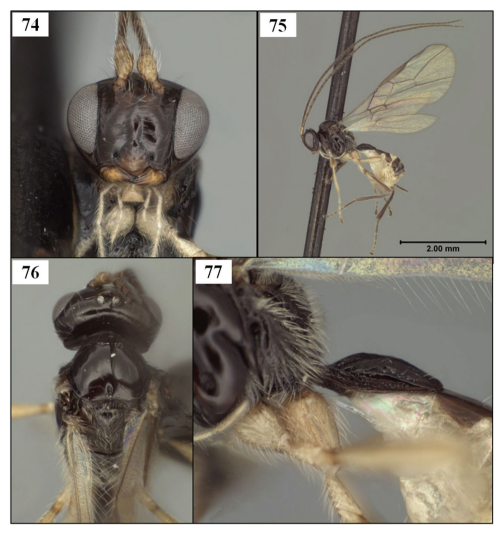
Opius (Nosopaeopius) incoligma Fischer, 1979: 274–276. Holotype female AEIC (examined).

Opius (Nosopaeopius) incoligma: Yu et al. 2005, 2012 (electronic catalogs).

Type locality. Colombia, Magdalena, 41 km south of Sta. Marta, 7000 ft.

Type material. Holotype. Female (AEIC), first label, first line: 41Km S.St. Marta second line: Magd., Colombia third line: V.6.1973 7000 ft. fourth line: Howden&Campbell second label [red]: Holotype third label, first line: [female symbol] Opius second line: incoligma third line: Holotype fourth line: det Fischer sp. n.

Diagnosis. Holotype female. Labrum completely concealed by mandibles; clypeus nearly as tall as wide, flat, not protruding, ventral margin convex. Mandible without basal lobe, distinctly narrowing apically to narrow, bifid tooth. Malar space distinct, malar sulcus deep, distinct. Antenna with 33 flagellomeres. Pronotum dorsally without pronope or distinct pit, mostly unsculptured, crenulate posterior margin broadly interrupted medially. Disc of mesoscutum nearly bare, with a few setae along traces of notauli; midpit small, distinct, narrowly elongate; notauli weak, present as very short, weakly sculptured grooves directed posterior-medially from and along edge of anterior declivity, not extending posteriorly onto disc of mesoscutum; distinct supra-marginal carina extending laterally from base of notaulus to tegula. Scuto-scutellar sulcus narrow (about $6-7 \times$ wider than long but difficult to measure), crenulate throughout. Precoxal sulcus distinct, moderately deep, long, completely unsculptured, somewhat vertically oriented as in Lorenzopius. Propodeum granular rugose, with very short median carina anteriorly, densely setose throughout. Fore wing stigma parallel-sided to weakly expanded apically; r1 longer than stigma width; second submarginal cell long, weakly narrowing distally; m-cu weakly postfurcal; 2CUb arising distinctly above middle of first subdiscal cell, 2CUa nearly absent. Hind coxa smooth; hind femur slender, weakly bilobed. T1 weakly strigose,



Figures 74–77. *Opius incoligma* Fischer, holotype female. 74 face 75 habitus 76 head and mesonotum, dorsal view 77 T1, lateral view.

irregularly sculptured with smooth patches; dorsal carina short but distinct; lateral carina very well developed, extending from junction with dorsal carina to apex, passing ventrad spiracle; dorsope shallow, indistinct, laterope shallow, weakly indicated by a long, narrow groove; T1 spiracle situated slightly posteriad midlength of T1; T1 narrow, parallel-sided, $2.6 \times longer$ than apical width; no visible membrane between S1 and T1, though lateral margin between the two clearly visible; S1 0.35 \times length of T1.

Remarks. The venation (Fig. 75) and features of the first metasomal segment (Figs 77, 81) suggest a relationship to *Eurytenes* (*Stigmatopoea*), but this species dif-

fers most remarkably by the completely concealed labrum (Fig. 74). Also, unlike the other species of *Eurytenes, Lorenzopius*, and *Tubiformopius* treated here, the individual flagellomeres are long throughout in *O. incoligma* but notably decreasing in length in the other species.

Opius rugicoxis Fischer

http://species-id.net/wiki/Opius_rugicoxis Figs 78–80, 82, 83

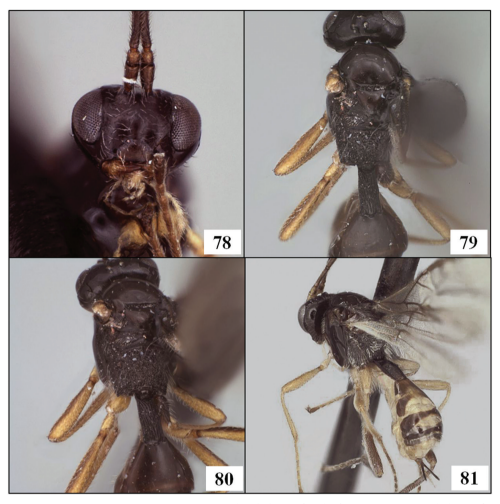
Opius rugicoxis Fischer, 1969: 251–254. Holotype female in AEIC (examined).
Opius (Stomosema) rugicoxis: Fischer 1977: 223, 248–249 (key, redescription); Yu et al. 2005, 2012 (electronic catalogs).

Type locality. Ecuador, Troya, 2900 m.

Type material. Holotype. Female (AEIC), first label, first line: Troya, Ecuador second line: VI. 10–13. 65 2900m. third line: Luis Pena second label [purple]: Holotype third label, first line: Opius [female symbol] second line: rugicoxis third line: det Fischer sp. n. fourth label, first line: Type no. second line: 659

Diagnosis. Holotype female. Labrum completely concealed by mandibles (Fig. 78); clypeus tall, flat, not protruding, ventral margin truncate. Mandible with broad, discrete basal lobe, apical half narrow, nearly parallel-sided. Malar space distinct; malar sulcus weak but present. Antenna with 25 flagellomeres. Pronotum not visible dorsally. Disc of mesoscutum (Figs 79, 80) bare, midpit small, round; notauli weak, present as very short, weakly sculptured grooves directed posterior-medially from and along edge of anterior declivity, not extending posteriorly onto disc of mesoscutum; weak supra-marginal carina extending laterally from base of notaulus nearly to tegula. Scuto-scutellar sulcus narrow $(5-6 \times \text{wider than long})$, crenulate throughout. Precoxal sulcus absent, thus unsculptured (Fig. 83). Propodeum (Fig. 80) completely granular rugose, without carinae, very sparsely setose. Fore wing (Fig. 82) with stigma folded, shape not readily discernible; r1 shorter than stigma width; second submarginal cell long, distinctly narrowing distally; m-cu distinctly postfurcal; 2CUb arising below middle of first subdiscal cell. Hind coxa granular-rugose, hence the species name; hind femur slender, distinctly bilobed. T1 (Figs 79, 80) completely striate, the striae curving medially from basal-lateral area adjacent dorsal tendon attachment, obscuring dorsal and lateral carinae; dorsope absent, laterope not apparent; T1 spiracle indistinct, situated posteriad midlength of T1; T1 nearly parallel-sided, 2.25 × longer than apical width; S1 appears fused to T1; S1 $0.3 \times \text{length of T1}$.

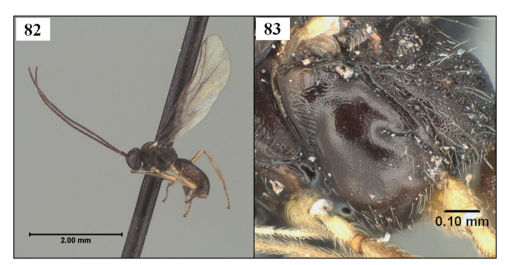
Remarks. Fischer (1977) placed this species in his subgenus *Opius (Stomosema)*, which he earlier (Fischer 1972) characterized on the basis of three features: a concealed labrum, absence of a mesoscutal midpit, and presence of sculpture in the precoxal sulcus. Unfortunately, the holotype has a small, shallow, but distinct mid-



Figures 78–81. *Opius* spp. 78 *O. rugicoxis* Fischer, holotype female, face 79 *O. rugicoxis*, holotype female, dorsal view 80 *O. rugicoxis*, holotype female, propodeum and T1 81 *O. incoligma* Fischer, holotype female, dorsal-lateral view.

pit (Figs 79, 80) and lacks a precoxal sulcus (Fig. 83). This species would therefore key to *Opius (Nosopaeopius)* in Fischer (1972) and Fischer (1999). Regardless of subgeneric assignment, this species falls within *Opius* in the classifications of Fischer (1977, 1999), van Achterberg and Salvo (1997), and Wharton (1997). The shape and sculpture of the first metasomal segment and the relatively long S1 suggest a relationship to *Tubiformopius*, but I exlude this species from *Tubiformopius* for the present time primarily on the basis of wing venation and from *Lorenzopius* on the basis of the form of the mandible.

The hind coxa is smooth to weakly punctate in other species treated here.



Figures 82-83. Opius rugicoxis Fischer, holotype female. 82 habitus 83 mesopleuron.

Tubiformopius Fischer, stat. rev.

http://species-id.net/wiki/Tubiformopius

Tubiformopius Fischer, 1998: 26. Type species: *Opius tubigaster* Fischer, 1968. Original designation.

Diagnosis. Mandible very weakly narrowing, nearly parallel-sided over distal 0.5, more abruptly widening basally, with weak to distinct basal lobe. Labrum narrowly exposed to concealed. Clypeus relatively weakly but distinctly protruding in profile; ventral margin truncate. Malar sulcus absent or represented only by a short, weak indentation adjacent eye; malar space distinct, at least as long as basal width of mandible. Occipital carina broadly absent dorsally, present laterally, distinctly separate from hypostomal carina ventrally. First flagellomere much longer than second. Propleuron ventral-laterally without oblique carina. Notauli short, shallow, narrow, confined to anterior declivity, not extending onto disc posteriorly; distinct midpit absent. Precoxal sulcus broad, very weakly impressed, unsculptured. Propodeum granular rugose, without areola. Fore wing stigma long, narrow, curled in holotypes of both species treated below, but not as discrete distally as in Lorenzopius and Stigmatopoea; r1 arising distinctly basad midpoint of stigma but not from extreme base; m-cu entering first submarginal cell, widely separated from second submarginal cell; second submarginal cell with 2RS much shorter than 3RSb; 2CUb arising near middle of hind margin of first subdiscal cell, the posterior-distal corner of the latter broadly open. Dorsope and laterope of T1 absent; S1 about $0.5-0.6 \times$ length of T1, apparently fused to T1; T1 long and narrow throughout; T2 and following terga unsculptured. Ovipositor not tapering evenly to a fine point.

Remarks. The diagnosis above is based on the holotypes of *T. tubigaster* (Fischer) and *T. tubibasis* (Fischer), new combination.

Fischer's (1998) original description of *Tubiformopius* was very brief since it was only included in a key to the eight genera he treated in the *Opius* genus group. Although two species are indicated in the relevant couplet, only one, designated as the type species, is specifically named. As noted above under the general discussion of genus-group characters, Fischer (1999), without discussion, treated *Tubiformopius* as a synonym of *Lorenzopius*. Aside from the original descriptions and Fischer's (1999) subsequent synonymy, neither *Tubiformopius* nor *Lorenzopius* has been further treated until now. We retain *Tubiformopius* as a valid genus distinct from *Lorenzopius* primarily on the basis of strong differences in the shape of the mandible (Fig. 85), fore wing venation (Figs 63, 65), and the notauli (Figs 86, 87). Fischer (1978) originally placed *T. tubibasis* in *Opius* s.s. Differences in venation and the first metasomal segment (especially the long and apparently fused S1) separate *Tubiformopius* from *Opius* s.s. Fischer (1977) placed *O. tubigaster* in the subgenus *Allophlebus* Fischer, 1972 but the type species of *Allophlebus* has T1 distinctly broadening apically with a very short, clearly separated S1, a distinct laterope, and the fore wing m-cu is postfurcal.

There is as yet no host data for either of the species currently included in *Tubiformopius*.

Tubiformopius tubigaster Fischer, stat. rev.

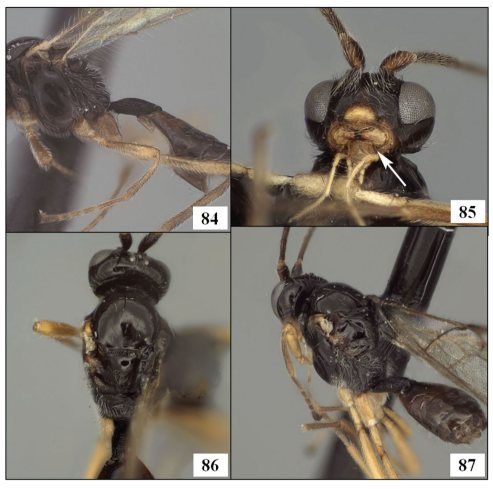
http://species-id.net/wiki/Tubiformopius_tubigaster Figs 8, 63, 65, 85–87

Opius tubigaster Fischer, 1968: 463–464, 483–485. Holotype male AEIC. *Opius (Allophlebus) tubigaster*: Fischer 1977: 223, 248–249 (key, redescription). *Tubiformopius tubigaster*: Fischer 1998: 26. *Lorenzopius tubigaster*: Fischer 1999: 282; Yu et al. 2005, 2012 (electronic catalogs).

Type locality. Ecuador, Cerro Tinajillas, 3200 m.

Type material. Holotype. Male (AEIC), first label, first line: Cerro Tinajillas second line: 3200m Ecuador third line: III. 18–21. 65 fourth line: Luis Peña second label [purple]: Holotype third label, first line: Opius [male symbol] second line: tubigaster third line: det Fischer sp. n. fourth label: first line: Type no. second line: 589

Diagnosis. Holotype male. Labrum partly concealed by mandibles (Fig. 85); clypeus nearly twice as wide as tall, protruding in profile, ventral margin truncate to very weakly concave. Mandible with basal lobe, apically nearly parallel-sided. Malar space distinct, malar sulcus not evident except as a small impression adjacent eye. Antenna with 26 flagellomeres. Pronotum dorsally not readily visible in holotype. Disc of mesoscutum nearly bare, with a sparse row of setae between notauli and transscutal articulation; midpit completely absent; notauli weak, present as very short, unsculptured grooves on anterior declivity, not extending posteriorly onto disc of mesoscutum; supra-marginal carina between base of notaulus and tegula absent. Scuto-scutellar sulcus



Figures 84–87. Tubiformopius spp. **84** T. tubibasis (Fischer), holotype female, T1, lateral view **85** T. tubigaster (Fischer), holotype male, face, arrow showing basal lobe of mandible **86** T. tubigaster, holotype male, head and mesoscutum, dorsal view **87** T. tubigaster, holotype male, dorsal-posterior view.

relatively narrow (Figs 86, 87), crenulate throughout. Precoxal sulcus indistinct, short, broad, very shallow, completely unsculptured. Propodeum granular rugose, without median carina anteriorly, moderately setose. Fore wing stigma long, curled in holotype, but appears to be very gradually tapered distally; r1 equal to or slightly longer than stigma width; second submarginal cell long, distinctly narrowing distally; m-cu widely antefurcal (Fig. 63, 65); 2CUb arising about middle of hind margin of first subdiscal cell, 2cu-a absent, first subdiscal cell broadly open at posterior-distal corner. Hind coxa smooth; hind femur very long, slender, weakly bilobed. T1 (Figs 63, 65, 86, 87) completely striate, the striae curving medially from basal-lateral area adjacent dorsal tendon attachment, completely obscuring dorsal and lateral carinae; dorsope and laterope absent; T1 spiracle indistinct, situated posteriad midlength of T1; T1 nearly parallel-sided, 2.1 × longer than apical width; S1 appears fused to T1; S1 0.5 × length of T1.

Remarks. This species is very similar to *T. tubibasis*, but differs in having a little more of the labrum exposed between the apex of the clypeus and the tightly closed mandibles. The hind coxae are yellow in *T. tubigaster* and distinctly infumate in *T. tubibasis*. Both species were described from Ecuador.

Tubiformopius tubibasis Fischer, comb. n.

http://species-id.net/wiki/Tubiformopius_tubibasis Fig. 84

Opius (Opius) tubibasis Fischer, 1978: 163–165. Holotype female in AEIC. *Opius (Opius) tubibasis*: Yu et al. 2005, 2012 (electronic catalogs).

Type locality. Ecuador, Cañar, Naupán, 3200 m.

Type material. Holotype. Female (AEIC), first label, first line: W. Naupán(Cañar) second line: 3200m. Ecuador third line: XII. 10. 70 fourth line: Luis Peña second label [red]: Holotype third label, first line: [female symbol] Opius second line: tubibasis third line: Holotype sp. n. fourth line: det. Fischer fourth label [yellow] Type 1195

Diagnosis. Holotype female. Labrum completely concealed by mandibles; clypeus tall, narrow, weakly protruding in profile, ventral margin truncate. Mandible with weak basal lobe, apically nearly parallel-sided. Malar space distinct, malar sulcus not evident except as a small impression adjacent eye. Antenna with 24 flagellomeres. Pronotum dorsally not readily visible in holotype. Disc of mesoscutum nearly bare, with a very sparse row of setae between notauli and transscutal articulation; midpit absent or nearly so, with faint indication of a depression when viewed in certain angles; notauli weak, present as short, weakly sculptured grooves on anterior declivity, not extending posteriorly onto disc of mesoscutum; supra-marginal carina between base of notaulus and tegula absent. Scuto-scutellar sulcus relatively narrow as in O. tubigaster, crenulate throughout. Precoxal sulcus barely visible as a short, broad, very shallow, completely unsculptured indentation. Propodeum granular rugose, without median carina anteriorly, moderately setose. Fore wing with stigma long, curled in holotype, but very gradually tapered distally; r1 equal to or slightly longer than stigma width; second submarginal cell long, distinctly narrowing distally; m-cu widely antefurcal; 2CUb arising slightly below middle of hind margin of first subdiscal cell, 2cu-a absent, first subdiscal cell broadly open at posterior-distal corner. Hind coxa smooth; hind femur very long, slender, weakly bilobed. T1 completely striate, the striae curving medially from basal-lateral area adjacent dorsal tendon attachment, completely obscuring dorsal and lateral carinae; dorsope and laterope absent; T1 spiracle indistinct, situated at 0.65 length of T1; T1 parallel-sided, $2.5 \times$ longer than apical width; S1 appears fused to T1; S1 $0.6 \times \text{length of T1}$.

Remarks. Van Achterberg and Salvo (1997) suggested the possibility that *tubibasis* might belong in *Lorenzopius* despite the absence of a midpit on the mesoscutum. The subsequently described *Tubiformopius* is a better fit because *tubibasis* is nearly identical to the type species of *Tubiformopius*, especially with respect to critical features of mesosomal sculpture and fore wing venation in addition to the shape of the mandible.

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Appendix

Hymenoptera Anatomy Ontology Table. (doi: 10.3897/zookeys.243.3990.app) File format: Microsoft Exsel file (xls).

Explanation note: Morphological terms used in text, referenced to the Hymenoptera Anatomy Ontology (HAO).

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RESEARCH ARTICLE



Two new species and new records of the genus Spinolyprops Pic, 1917 from the Oriental Region (Coleoptera, Tenebrionidae, Lupropini)*

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Abstract

Two new species of the genus *Spinolyprops* Pic, 1917 (Tenebrionidae, subfamily Lagriinae Latreille, 1825) are described from Thailand and China (*S. cribricollis* **sp. n.**, *S. thailandicus* **sp. n.**). The species characters within the genus are discussed, photographs of all seven Oriental species are added, a species key for the Oriental species is compiled, and a map with the distributional patterns is provided.

Keywords

Tenebrionidae, Lagriinae, Lupropini, Spinolyprops, new species, Oriental region, distribution, map, species key

Introduction

The genus *Spinolyprops* Pic, 1917 (Tenebrionidae, subfamily Lagriinae Latreille, 1825, tribe Lupropini Ardoin, 1958) was based on the type species *Spinolyprops rufithorax* Pic, 1917 from Zanzibar (Pic 1917). Kulzer (1954) published the first species from the Oriental Region (Sri Lanka). Later Kaszab (1965) and Schawaller (1994, 1996) described additional Oriental species. The purpose of the present paper is the description of two further species from the Oriental region (Thailand, China), to discuss the

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^{*} Contributions to Tenebrionidae no. 105. – For no. 104 see: Annales Zoologici 62, 2012.

species characters, to present for the first time photographs of all seven Oriental species (Figs 2–13), to provide a key for all Oriental species, and finally to add new faunistic data including a map with the distributional patterns (Fig. 1).

The separation of the genera *Pseudolyprops* Fairmaire, 1882, *Sphingocorse* Gebien, 1921, and *Spinolyprops* Pic, 1917 within the tribe Lupropini is still in a preliminary state and not yet based on discriminating characters. At present, the species with spine-like posterior corners of the pronotum, and with elytral colour pattern, are assigned to *Spinolyprops*. Congeners of all three genera live in Africa or in the Oriental/Papuan regions, thus zoogeographical aspects should also be considered during a future revision.

All species of this group are soil dwellers and are adapted also to extreme dry conditions (personal observations). All known species have fully developed wings and thus possess a high ability for dispersal. Specimens are usually collected by sifting litter and similar substrates, and are also attracted by light.

Depositories

BMNH	The Natural History Museum, London (Max Barclay)
CRGT	Collection Dr. Roland Grimm, Neuenbürg
HNHM	Hungarian Natural History Museum, Budapest (Dr. Ottó Merkl)
MHNG	Muséum d'Histoire Naturelle, Genève (Dr. Giulio Cuccodoro)
MNB	Museum für Naturkunde, Berlin (Dr. Manfred Uhlig)
MSNF	Museo di Storia Naturale, Firenze (Dr. Luca Bartolozzi)
NHMB	Naturhistorisches Museum, Basel (Dr. Michel Brancucci †)
NMPC	National Museum (Natural History), Prague (Dr. Jiří Hájek)
SMNK	Staatliches Museum für Naturkunde, Karlsruhe (Dr. Alexander Riedel)
SMNS	Staatliches Museum für Naturkunde, Stuttgart

Species characters

All species have a similar dorsal colour pattern (Figs 1–13), which is quite variable and helpful only in a low extent for species separation. On the contrary, the combination of the following characters are considered as diagnostic for species. Dorsal punctation of pronotum and elytra either fine (Figs 3–6) or rough (Figs 2, 7–13). The shape of pronotum with the lateral parts broadly (Figs 2, 7, 10–13) or narrowly (Figs 3–6) separated from disc; and with the anterior margin feebly (Figs 2–6) or deeply (Figs 10–12) excavated. Frons between eyes narrower (Figs 2, 7) or wider (Figs 3–6) than dorsal eye diameter. Antennomeres 8–10 longer than broad (Figs 2, 7) or as long as broad (Figs 3–6, 8–9). Sexual dimorphism of middle tibia present in one species (*S. pakistanicus*), absent in all other species. Specific shape of the apicale of aedeagus, considering a certain variability (Figs 14–25). Males and females can be separated only by dissection. For separation of the species see key below (suitable only for males).

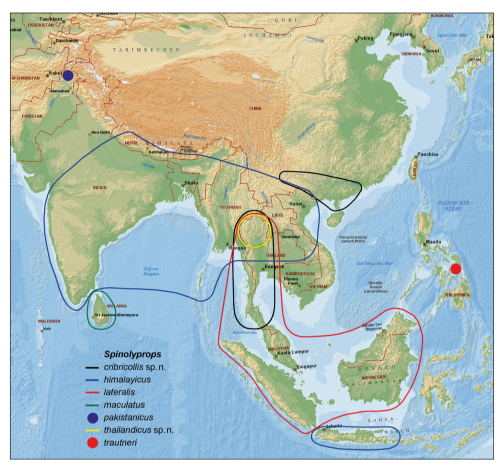


Figure 1. Idealized distributional patterns of the *Spinolyprops* species in the Oriental Region (modified Microsoft Encarta map).

Key to the species of *Spinolyprops* from the Oriental Region (\Diamond)

1	Frons between eyes wider than dorsal eye diameter, dorsal punctation of pro-
	notum and elytra fine, pronotum with lateral parts narrowly separated from
	disc (Figs 3–6, 16–19) himalayicus
_	Frons between eyes narrower than eye diameter, dorsal punctation of prono-
	tum rough, pronotum with lateral parts broadly separated from disc2
2	Pronotum with anterior margin feebly excavated (compare figures)
_	Pronotum with anterior margin deeply excavated
3	Elytral colour pattern apically with an arrow-shaped dark element, aedea-
	gus with long and broad triangular apicale with straight sides, antennomeres
	8–10 as long as broad (Figs 8, 22) maculatus
_	Elytral colour pattern apically with an narrowing pointed dark element, ae-
	deagus with short and narrow triangular apicale with rounded sides, anten-
	nomeres 8–10 longer than broad (Figs 2, 14–15) cribricollis sp. n.

Middle tibia of males on inner side with about five distinct spines (Figs 9,
23)pakistanicus
All tibiae unarmed
Separated lateral parts of pronotum extremely broad, aedeagus with broad
spade-like apicale (Figs 10–12, 25) thailandicus sp. n.
Separated lateral parts of pronotum narrower, aedeagus with apicale pentago-
nal6
Dorsal setation of pronotum and elytra short, body shape narrower (elytra
1.4× longer than broad) (Figs 13, 24) trautneri
Dorsal setation of pronotum and elytra long, body shape broader (elytra 1.3×
longer than broad) (Figs 7, 20-21)lateralis

The species

Spinolyprops cribricollis sp. n.

urn:lsid:zoobank.org:act:19A94C99-40A0-4D72-9CEC-716EEA19213C http://species-id.net/wiki/Spinolyprops_cribricollis Figs 2, 14–15

Type specimens. Holotype male: S Thailand, Island Ko Chang, western side, 1999 (without detailed data), leg. A. Schulz & K. Vock, SMNS. – Paratypes: N Thailand, Chiang Mai Prov., Doi Inthanon, 1800 m, 14.V.2006, leg. R. Grimm, 4 ex. CRGT, 1 ex. SMNS. – NW Thailand, Doi Pui, 1600–1685 m, 7.–9.V.2004, leg. R. Grimm, 4 ex. CRGT. – NW Thailand, Doi Pui, 1600–1685 m, 22.–23.V.2006, leg. R. Grimm, 1 ex. SMNS. – China, Yunnan, 22 km NE Dali, NE bank of Er Hai Lake, 2010 m, 12.VI.2007, leg. M. Schülke, 1 ex. MNB, 1 ex. SMNS. – China, S Yunnan, Mengyang NR, 500 m, 12.IX.1994, leg. S. Kurbatov, 2 ex. HNHM. – China, NE Guangxi, 15 km N Longsheng, 1000 m, 15.–22.VI.1995, leg. S. Kurbatov, 1 ex. HNHM.

Diagnosis. *S. cribricollis* sp. n. shares with *S. lateralis* the rough dorsal punctation of pronotum and elytra, the shape of the pronotum with lateral parts broadly separated from disc and bent up, the frons between eyes smaller than dorsal eye diameter, and the antennomeres 8–10 longer than broad. Both can be separated mainly by the anterior margin of the pronotum with feeble (*S. cribricollis* sp. n.) or deep excavation (*S. lateralis*), and by different shape of the aedeagus (in *S. lateralis* the apicale is pentagonal, compare Figs 20–21). Additionally, *S. lateralis* is somewhat larger in the average (5.0–6.0 mm), and the elytra are slightly more rounded. *S. maculatus* has a similar shape of the pronotum with feebly excavated anterior margin, but the aedeagus has the apicale of the aedeagus different triangular with straight sides. *S. trautneri* has also a different aedeagus with broad pentagonal apicale (Fig. 24). See also the species key above.

Description. Body length 4.5–5.0 mm. Dorsal and ventral surfaces and all appendages brown without metallic shine, head and pronotum slightly darker, elytra bicoloured with darker and lighter parts (see Fig. 2); dorsal surface roughly punctate, punc-



Figures 2–7. Dorsal view of *Spinolyprops* species from the Oriental Region. 2 *S. cribricollis* sp. n., paratype Thailand CRGT 3 *S. himalayicus*, non-type Thailand SMNS 4 *S. himalayicus*, non-type Bali SMNS
5 *S. himalayicus*, non-type W India SMNS 6 *S. himalayicus*, non-type Thailand SMNS 7 *S. lateralis*, non-type Sabah SMNS

tures with long erect setae, surface between punctures shining. Head with punctation similar as on pronotum; genae distinctly swollen, clypeal suture somewhat indistinct by rough punctation, clypeus with punctation as on frons, anterior margin of clypeus straight; eyes reniform, frons between eyes smaller than dorsal eye diameter, temples impunctate; maxillary palps with large securiform terminal palpomere; shape of antennomeres see Fig. 2, antennomere 3 not elongate, terminal three antennomeres not forming club. Pronotum widest in middle, anterior and posterior margins unbordered, lateral margins unbordered but distinctly crenulate, anterior corner rounded, posterior corners acute, surface flat with irregular rough and partly confluent punctation, lateral parts broadly separated from disc and bent up; propleura with sparser and smaller punctation and shorter setation as on pronotum, prosternal process not prominent; metaventrite with punctation as on propleura. Scutellum visible, shining, without punctation. Hind wings present. Elytra elongate oval, widest in middle, lateral margin distinctly dentate in humeral region, margin completely visible from above; surface with rough punctation as on pronotum, but not confluent, punctation as on elytral disc, similar as on propleura. Ventrites with fine and widely separate punctation, terminal ventrite unbordered, intersegmental membranes exposed between ventrites 3–5. Legs without particular modifications, tibiae without external keels, tibial spurs short. Aedeagus with triangular apicale with acute tip (Figs 14–15). No distinct external sexual dimorphism.

Remarks. I hope not to fail in assigning the (so far disjunct) Chinese specimens from Yunnan and Guangxi to the same species. Shape and punctation of the pronotum, elytral colour and shape of aedeagus are not distinctly different from the specimens from Thailand. The type locality lies in a lowland habitat (Island Ko Chang), and the paratypes from Thailand were collected in higher altitudes (Doi Pui and Doi Inthanon). Obviously, this species has a wide ecological range.

Etymology. The name refers to the rough punctation of the pronotum.

Spinolyprops himalayicus Kaszab, 1965

http://species-id.net/wiki/Spinolyprops_himalayicus Figs 3–6, 16–19

Type specimens examined. India, Darjeeling (labelled as West Bengal), Peshok, 710 ft., 19.IX.1959, leg. F. Schmid, holotype HNHM.

New material. Nepal, Gorkha Distr., Arughat Bazar, 600 m, 26.VII.1983, leg. J. Martens & W. Schawaller, 1 ex. SMNS. – Nepal, Surkhet Distr., Bheri Khola Bridge, 500 m, 24.–25.V.1998, leg. W. Schawaller, 1 ex. SMNS. – Nepal, Chitwan NP, Rampur, V.2005, leg. D. Ahrens, 1 ex. SMNS. – N India, Darjeeling, Sukna, 180 m, 21.–23.V.1980, leg. G. Topál, 1 ex. HNHM. – N India, Uttar Pradesh, Mussorie, 1300 m, 10.VII.1989, leg. A. Riedel, 1 ex. SMNS. – NE India, Meghalaya, 1 km E Tura, 500–600 m, 13.–18.V.2002, leg. M. Trýzna & P. Benda, 1 ex. SMNS. – W India, Maharasthra State, 70 km S Pune, Wai, 3.–6.X.2005, leg. F. & L. Kantner, 1 ex. SMNS. – W India, Maharasthra State, 40 km W Pune, Mulshi, 7.–11.X.2005, leg. F. & L. Kantner, 1 ex. SMNS. – S India, Anamalai Hills, Cinchona, 3500 ft., 1959, no collector labelled, 9 ex. NHMB (Frey collection, det Kulzer). – Bhutan, Samchi, 300 m, 7.–11.V.1972, Basel Expedition, 2 ex. HNHM. – Andaman Islands, Havelock Island, village no. 7, 22.IV.–14.V.1998, leg. K. & S. Majer, 2 ex. NHMB, 1 ex. SMNS. – N Thailand, Chiang Mai, Doi Pui, 1500 m, 19.XII.1988, leg. K. Geigenmüller & J. Trautner, 1 ex. SMNS. – N Thailand, Chiang Dao, 70 km N Chiang

Mai, 26.–28.IV.2003, leg. O. Šafránek, 2 ex. SMNS. – N Thailand, Chiang Mai Prov., Ban San Pakia, 1700 m, 25.IV.–7.V.1996, leg. S. Bilý, 7 ex. NHMB, 1 ex. SMNS. – N

Ban San Pakia, 1700 m, 25.IV.-7.V.1996, leg. S. Bilý, 7 ex. NHMB, 1 ex. SMNS. - N Thailand, Nan, 22.–24.V.1999, leg. R. Grimm, 16 ex. CRGT, 1 ex. SMNS. – N Thailand, Nan, 2.-4.V.2003, leg. R. Grimm, 2 ex. CRGT. - NW Thailand, Doi Pui, 1600-1685 m, 15.-16.IV.2004, leg. R. Grimm, 1 ex. CRGT. - N Thailand, Chiang Mai, Doi Pui, 1600 m, 15.–16.IV.2004, leg. W. Schawaller, 2 ex. SMNS. – NW Thailand, Chiang Dao, 700-800 m, 4.V.2004, leg. R. Grimm, 3 ex. CRGT. - NW Thailand, Mae Hong Son, 5.V.2004, leg. R. Grimm, 2 ex. CRGT. – NW Thailand, 5 km E Pai, 700 m, 19.IV.2004, leg. W. Schawaller, 1 ex. SMNS. - NW Thailand, Soppong, 700 m, 23.IV.2004, leg. W. Schawaller, 9 ex. SMNS. – NW Thailand, Soppong (Pangmapa), 17.–18.V.2006, leg. R. Grimm, 5 ex. CRGT. – Thailand, Chumphon Prov., Pha To, 27.III.–14.IV.1996, leg. K. Majer, 3 ex. NHMB. - Thailand, Thanon Thong Chai, Palong, 750 m, 26.-28.V.1991, leg. V. Kubán, 3 ex. NHMB. – Thailand, Prachin Buri Prov., Sakaerat Ecology Research Institute, 4.VI.2001, leg. E. Horváth & G. Sziráki, 5 ex. HNHM. – Burma, N Shan State, Namhsan, 1500-1900 m, 18.-28.II.1996, leg. S. Kasantsev, 1 ex. NHMB. - C Laos, Khammouan Prov., Ban Khoun Ngeun, 200 m, 19.-31.V.2001, leg. L. Dembický, 5 ex. SMNS. – C Laos, Khammouan Prov., Ban Khoun Ngeun, 17.V.–6.VI.2007, leg. M. Strba, 1 ex. SMNS. – CE Laos, Boli Kham Xai Prov., 8 km NE Ban Nape, 600 m, 1.– 18.V.2001, leg. L. Dembický, 6 ex. SMNS. – Laos, Champassak Prov., Bolavens Plateau, 3 km SE Ban Lak, 1070 m, 9.V.2010, leg. J. Hájek, 1 ex. NMPC. – Laos, Champassak Prov., Ban Nong Luang, 12 km S Paksong, 800 m, 6.IV.1998, leg. O. Merkl & G. Csorba, 1 ex. HNHM. – Laos, Phongsaly Prov., Phongsaly, 1500 m, 28.V.–20.VI.2003, leg. M. Brancucci, 2 ex. NHMB. – Laos, Phongsaly Prov., Phongsaly, 1500 m, 6.–17.V.2004, leg. M. Brancucci, 1 ex. SMNS. – Vietnam, Daklak Prov., Buon Ma Thuot, Dak Linn, 500 m, 28.–29.IV.1986, leg. S. Golovatch & L. Medvedev, 7 ex. SMNS. – Vietnam, Bac Kan Prov., Ba Be NP, 350 m, 3.–8.VI.2011, leg. L. Bartolozzi et al., 3 ex. MSNF. – Java, Batavia (now Jakarta), III.1921, no further data, 1 ex. HNHM. – C Bali, Bedugul, Tamlingan, 1210 m, 6.XI.2007, leg. A. Riedel, 10 ex. SMNK, 4 ex. SMNS.

Distribution. India (type locality Peshok/Darjeeling), Nepal, Bhutan, N Thailand, Vietnam (Schawaller 1996); Andaman Islands, Burma, Laos, Java, Bali (new records).

Spinolyprops lateralis Pic, 1917

http://species-id.net/wiki/Spinolyprops_lateralis Figs 7, 20–21

Spinolyprops rufithorax var. lateralis Pic, 1917

New material. NE Sumatra, Tebing-Tinggi, 1 ex. NHMB (Frey collection). – E Sumatra, Lampung, Bawang, Pedada Bay, Gn. Tanggang, 660 m, 9.VIII.2006, leg. A. Riedel, 1 ex. SMNS. – Borneo, Brunei, Temburong Distr., ridge NE Kuala Belalong, 300 m, X.1992, leg. J. H. Martin, 1 ex. BMNH. – Borneo, Sabah, Crocker Range, Tenom, Kalang waterfall, 17.VI.1998, leg. J. Kodada & F. Čiampor, 5 ex. SMNS. – Borneo, Sabah, Sapulut, Batu Pungull, 24.–26.VI.1998, leg. J. Kodada & F. Čiampor, 1 ex. SMNS. - Borneo, Sabah, Poring, 650 m, 15.V.2005, leg. R. Grimm, 1 ex. CRGT. -Borneo, Sarawak, Gunung Santubong, 10–200 m, 4.–8.IV.2009, leg. R. Grimm, 1 ex. CRGT. - Borneo, Sarawak, Gunung Santubong, 30-200 m, 30.XI.-5.XII.2010, leg. R. Grimm, 4 ex. CRGT. – Borneo, Sarawak, Gunung Gading NP, 100–300 m, 31.III.-4. IV.2009, leg. R. Grimm, 1 ex. CRGT. - Borneo, Sarawak, Gunung Gading NP, 50-200 m, 8.–10.XII.2010, leg. R. Grimm, 2 ex. CRGT. – NW Thailand, Mae Hong Son Prov., 32 km NNE Mae Hong Son, 5.V.2004, leg. R. Grimm, 2 ex. CRGT. - S Thailand, Khao Lak NP, Thone Chong Fa Waterfall, 100-300 m, 6.-15.I.1998, leg. A. Schulz & K. Vock, 3 ex. SMNS, 1 ex. MNB. – W Malaysia, Perak, 25 km NE Ipoh, Banjaran Titi Wangsa Mts., Mt. Korbu, 1200 m, 6.–12.V.2001, leg. P. Čechovský, 1 ex. SMNS.

Distribution. Sumatra (type locality); Borneo, Thailand, W Malaysia (new records).

Spinolyprops maculatus Kulzer, 1954

http://species-id.net/wiki/Spinolyprops_maculatus Figs 8, 22

Type specimens examined. Sri Lanka, Colombo, III.1953, leg. G. Frey, holotype and 1 paratype NHMB (Frey collection), 2 paratypes HNHM.

New material. Sri Lanka, Uva, Diyaluma Falls, 400 m, 23.I.1970, leg. C. Besuchet, I. Löbl & R. Mussard, 3 ex. MHNG, 1 ex. SMNS. – Sri Lanka, Uva, Monaragala, 300 m, 13.II.1970, leg. C. Besuchet, I. Löbl & R. Mussard, 1 ex. HNHM. – Sri Lanka, Uva, S Wellawaya, 300 m, 25.I.1970, leg. C. Besuchet, I. Löbl & R. Mussard, 1ex. HNHM. - Sri Lanka, Periyapullumalai, 11.II.1970, leg. C. Besuchet, I. Löbl & R. Mussard, 1 ex. HNHM. - Sri Lanka, Kandy, 18.III.1973, leg. G. Zimmermann, 1 ex. SMNS. - S Burma (labelled as Tenasserim), no additional data, 1 ex. HNHM (det. Kaszab).

Remarks. The specimen from Tenasserim was already published by Kaszab (1965). This specimen clearly belongs to S. maculatus and shares with the specimens from Sri Lanka the elytral colour pattern apically with an arrow-shaped dark element (Fig. 8). It is the only specimen of *S. maculatus* out of Sri Lanka, so probably it was mislabelled (and is not mapped herein).

Distribution. Sri Lanka (type locality), ? S Burma (Kaszab 1965).

Spinolyprops pakistanicus Schawaller, 1996

http://species-id.net/wiki/Spinolyprops_pakistanicus Figs 9, 23

Type specimens examined. Pakistan, Hazara, Malkandi, 1500 m, 3.VI.1983, leg. C. Besuchet & I. Löbl, 2 paratypes SMNS. – Pakistan, Swat, Madyan, 1400 m, 16.V.1983, leg. C. Besuchet & I. Löbl, 1 paratype SMNS.

Distribution. Northern Pakistan in Hazara and Swat.

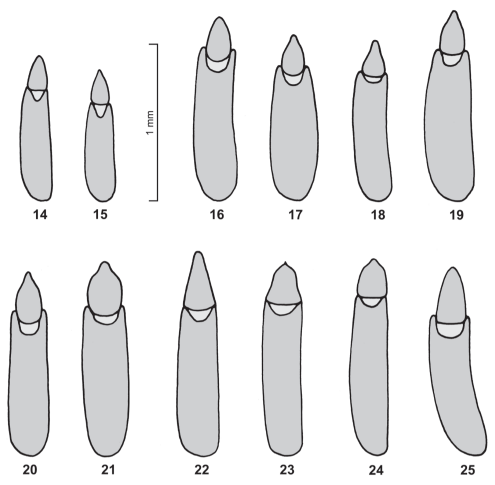


Figure 8–13. 8 *S. maculatus*, non-type Sri Lanka SMNS 9 *S. pakistanicus*, paratype Pakistan SMNS 10 *S. thailandicus* sp. n., holotype Thailand SMNS 11 *S. thailandicus* sp. n., paratype Thailand CRGT 12 *S. thailandicus* sp. n., paratype Thailand SMNS 13 *S. trautneri*, holotype Philippines SMNS.

Spinolyprops thailandicus sp. n.

urn:lsid:zoobank.org:act:E51CF1A3-852E-4A33-8196-46C799657133 http://species-id.net/wiki/Spinolyprops_thailandicus Figs 10–12, 25

Type specimens. Holotype male: N Thailand, Chiang Mai, Doi Pui, 1600 m, 15.–16. IV.2004, leg. W. Schawaller, SMNS. – Paratypes: same data as holotype, 3 ex. SMNS. – N Thailand, Chiang Mai, Doi Pui, 1600-1685 m, 23.IV.–12.V.2003, leg. R. Grimm,



Figures 14–25. Aedeagus of *Spinolyprops* species in the Oriental Region. 14 *S. cribricollis* sp. n., holotype Thailand/Ko Chang SMNS 15 *S. cribricollis* sp. n., paratype Thailand/Doi Inthanon SMNS 16 *S. himalayicus*, non-type Nepal SMNS 17 *S. himalayicus*, non-type Bali SMNS 18 *S. himalayicus*, non-type Thailand SMNS 19 *S. himalayicus*, non-type W India SMNS 20 *S. lateralis*, non-type Borneo SMNS 21 *S. lateralis*, non-type Thailand SMNS 22 *S. maculatus*, non-type Sri Lanka SMNS 23 *S. pakistanicus*, paratype Pakistan SMNS 24 *S. trautneri*, holotype Philippines SMNS 25 *S. thailandicus* sp. n., holotype Thailand SMNS.

4 ex. CRGT. – N Thailand, Chiang Mai, Doi Pui, 1600-1685 m, 7.–9.V.2004, leg. R. Grimm, 6 ex. CRGT. – N Thailand, Chiang Mai, Doi Pui, 1600-1685 m, 12.– 13.V.2006, leg. R. Grimm, 7 ex. CRGT, 4 ex. SMNS. – N Thailand, Chiang Mai, Doi Pui, 1600-1685 m, 22.–23.V.2006, leg. R. Grimm, 6 ex. CRGT, 3 ex. HNHM.

Diagnosis. *S. thailandicus* sp. n. is characterized by the shape of the pronotum with deeply excavated anterior margin and with the lateral parts broadly separated from disc and bent up, in combination with rough dorsal punctation of pronotum and elytra and the frons between eyes smaller than dorsal eye diameter. The aedeagus of *S. thailandicus* sp. n.

is similar as in *S. cribricollis* sp. n. (compare Figs 14–15), but in this species the lateral parts of the pronotum are not so widely separated from the disc as in *S. thailandicus* sp. n., and the anterior margin of the pronotum is only feebly excavated. See also species key above.

Description. Body length 4.3–5.3 mm. Dorsal and ventral surfaces and all appendages brown without metallic shine, elytra bicoloured with darker and lighter parts in different variation (Figs 10–12); dorsal surface roughly punctate, punctures with long erect setae, surface between punctures shining. Head with punctation similar as on pronotum; genae distinctly swollen, clypeal suture somewhat indistinct by rough punctation, clypeus with punctation as on frons, anterior margin of clypeus straight; eyes reniform, frons between eyes as broad as dorsal eye diameter, temples impunctate; maxillary palps with large securiform terminal palpomere; shape of antennomeres see Figs 10-12, antennomere 3 not elongate, terminal three antennomeres not forming club. Pronotum widest slightly before middle, anterior and posterior margins unbordered, lateral margins unbordered but distinctly crenulate, anterior corner rounded, posterior corners acute, surface flat with irregular rough and partly confluent punctation, lateral parts broadly separated from disc and bent up; propleura with sparser and smaller punctation and shorter setation as on pronotum, prosternal process slightly prominent; metaventrite with punctation as on propleura. Scutellum visible, shining, without punctation. Hind wings present. Elytra elongate oval, widest in middle, lateral margin distinctly dentate in humeral region, margin completely visible from above; surface with rough punctation as on pronotum, but not confluent, punctation irregular and not arranged in rows or striae; epipleura with sparser and smaller punctation as on elytral disc, similar as on propleura. Ventrites with fine and widely separate punctation, terminal ventrite unbordered, intersegmental membranes exposed between ventrites 3-5. Legs without particular modifications, tibiae without external keels, tibial spurs short. Aedeagus with broad spade-like apicale with rounded tip (Fig. 25). No distinct external sexual dimorphism.

Etymology. Named after the type locality in Thailand.

Spinolyprops trautneri Schawaller, 1994

http://species-id.net/wiki/Spinolyprops_trautneri Figs 13, 24

Type specimen examined. Philippines, Leyte, Lake Danao, 500 m, 19.II.–18. III.1991, leg. K. Geigenmüller, W. Schawaller & J. Trautner, male holotype SMNS. **Distribution.** Philippines (type locality Leyte Island).

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SHORT COMMUNICATION



Rhabdorrhynchus echii (Brahm, 1790), a "forgotten" name (Coleoptera, Curculionidae, Lixinae)

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Abstract

The application of the name *Curculio echii* Brahm, 1790 is discussed. Based on the description it is evident that it should be applied to a German species of the genus *Rhabdorrhynchus*, and that it has priority over the name currently applied to the species, *R. seriegranosus* Chevrolat, 1873. The new combination *Rhab-dorrhynchus echii* (Brahm, 1790) is proposed. As there is a lack of any type material of *C. echii* a neotype is designated. Based on the study of the type specimen, *R. seriegranosus* is restored as a valid species.

Keywords

Neotype, new combination, nomenclature, European fauna, Palaearctic weevils

Introduction

In the course of the preparation of the Catalogue of the Palaearctic Coleoptera: Lixinae: Cleonini several nomenclatural questions had to be resolved, and several previously undetected cases of priority came to light. Some of these concerned taxa seldom, if ever, cited in the literature. However, in some cases the "lost" names were applied to taxa more broadly known.

Brahm (1790) published an "Insect Calendar", in which he mentioned, month by month, the species that he had seen or collected in the surroundings of Mainz (Germany). In most cases he referred them to previously named taxa, but some of the

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species were described as new. One of these is *Curculio echii* Brahm, 1790: 175. The description is rather accurate, cites the plant where Brahm collected one specimen (*Echium vulgare* L.) and emphasizes its key traits. This description allows identification of *C. echii* as belonging to the genus *Rhabdorrhynchus* Motschulsky, 1860. Only one species of the genus is known to occur in Germany, thus the application of the name is undisputed. The comb. n. *Rhabdorrhynchus echii* (Brahm, 1790) is here proposed, based on *Curculio echii* Brahm, 1790: 175.

However, even though this was the first name to have been assigned to this species, it was born under an unlucky star, and it soon became forgotten. The name *Curculio echii*, published as it was in a book that did not have any influence in nomenclature, was not used in subsequent works. It was occasionally cited in nomenclatural checklists, both older ones (Sherborn 1902) and recent on-line name indexes (i.e., ION 2012), but it was never positively applied to any taxon and no transfer to other genera was ever proposed.

In 1795 Herbst described *Curculio varius* Herbst, 1795, from Europe. Regardless to the fact that this name is a junior homonym of *Curculio varius* Fabricius, 1775, and several other senior homonyms, it was continuously applied to the central European species, firstly (Schoenherr 1826) as *Pachycerus varius*, and subsequently, after publication of Chevrolat (1873), as *Rhabdorrhynchus varius*. Chevrolat (1873) also described *Rhabdorrhynchus seriegranosus* Chevrolat, 1873, from Algeria. This name was placed under synonymy of *Rhabdorrhynchus varius* by Faust (1904).

Throughout the 20th century the species, in Faust's concept, was named *Rhab-dorrhynchus varius* (Csiki 1934, as in all other papers which cited the species, including Ter-Minasyan 1988). It ranges in central-southern Europe, northern Africa and western Asia. Eventually, Alonso-Zarazaga and Lyal (1999) discovered the homonymy of *Curculio varius* Herbst, 1795 and its only synonym, *R. seriegranosus* Chevrolat, 1873, became the valid name and it was applied to specimens from Europe, Northern Africa and Western Asia.

However, *R. echii* is the first available name to be applied to the taxon and must be used as the correct name for this species. *R. seriegranosus* was seldom used since 1999 and article 23.9.1.2 ICZN cannot be applied.

It should be added that the taxonomy of the genus was never thoroughly revised, and there is still uncertainty regarding the validity of some species and their range. The type specimen of *Rhabdorrhynchus seriegranosus*, recently examined by one of the authors (M.M.), differs quite significantly from the European taxon (Figs 1-4), and attribution of specimens from the southern Mediterranean coasts to the central European species, as originally proposed by Faust (1904) and never subsequently discussed, seems questionable. The synonymy *Rhabdorrhynchus echii* (Brahm, 1790) = *Rhabdorrhynchus seriegranosus* Chevrolat, 1873 is here rejected and *R. seriegranosus* is restored as a valid species.

This act also allows maintainance as valid the first epithet attributed to the Algerian *Rhabdorrhynchus*. Two more species were in fact subsequently described from Algeria, namely, *R. curvirostris* Solari, 1950 and *R. longicollis* Solari, 1950, both based on a

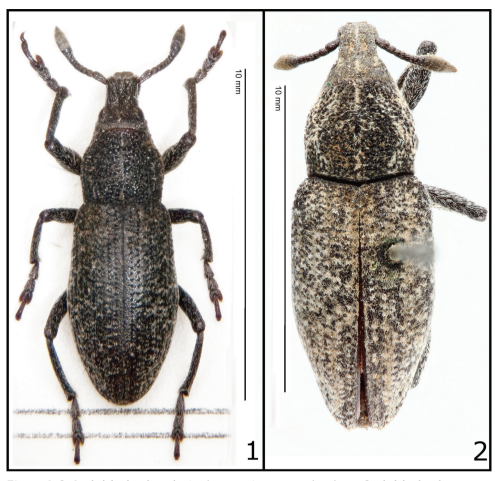


Figure 1–2. l *Rhabdorrhynchus echii* (Brahm, 1790). Neotype, dorsal view 2 *Rhabdorrhynchus seriegranosus* Chevrolat, 1873. Type specimen, dorsal view. Conserved at the Stockholm Museum of Natural History, Chevrolat colletion.

single specimen, never recollected anywhere else and thus apparently endemic to the country – which is at least unlikely. The status of these species, and their relations with *R. seriegranosus* and the other north-african taxa of the genus, were never discussed after their description.

It is evident that a typification of *Curculio echii* Brahm is required. Nikolaus Joseph Brahm (1751–1821) was a German zoologist, but there is no information about the fate of his collection, which was never cited in the literature. Horn et al. (1990) do not even report the name. Thus we consider the type specimen of *Curculio echii* to be lost.

We hereby designate a neotype of *Curculio echii* Brahm, using a specimen from southern Germany. This act is done with the intent to fix once and for all the meaning of the name and to stabilize nomenclature (under Art. 75.3 of the International Code of Zoological Nomenclature), with particular regard to the central European taxon.

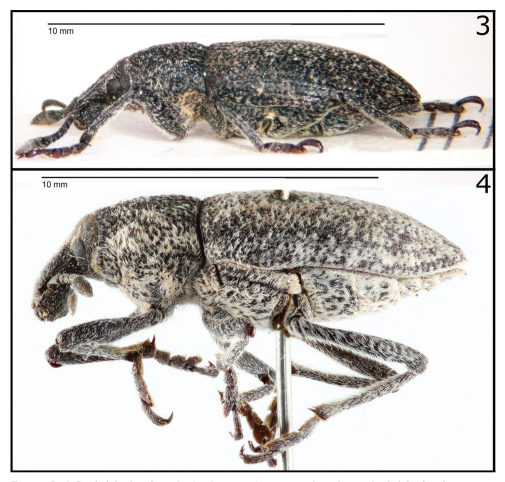


Figure 3–4. 3 *Rhabdorrhynchus echii* (Brahm, 1790). Neotype, lateral view 4 *Rhabdorrhynchus seriegranosus* Chevrolat, 1873. Type specimen. Same specimen as Fig. 2.

A specimen conserved at the Staatliches Museum für Naturkunde, Stuttgart, Germany, collected in Baden-Württenberg, southern Germany – thus not far from Mainz – is appropriate to be used as the neotype of *R. echii*. An appropriate description for the central-European populations was given by Dieckmann (1983), under *R. varius* Herbst.

Curculio echii Brahm 1790. NEOTYPE (here designated): A male specimen so labeled: 1. Germany, Ba- / den–Württ. / Grißheim / 6.Aug.1994 / leg. Kasper // Rheinaue // Rhabdorhyn- / chus varius / Herbst / det. Kasper // *Curculio echii* Brahm 1790 / NEOTYPUS / 2012 Meregalli & Alonso-Z. des. [red]

The range of the species is here considered to include only the forms morphologically referable to *R. echii*, mainly present in Central and central-eastern Europe. The populations from southern Europe, northern Africa and Western Asia are referred to *R. seriegranosus*.

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