## Tijdschrift voor

 EntomologieA journal of systematic and evolutionary entomology since 1858


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The abstract starts with a bibliographical reference (authors, year, tide), see recent issues for layout. Then the abstract follows, without the word 'abstract' and without indentation. The abstract, written in English, should be concise, yet cover the main results
of the paper, including new taxa and nomenclatorial changes. The name and address of one and only one of the authors follows, if needed preceded by the word 'Correspondence:'. This should also be the address for the galley proofs. The abstract ends with a list of key-words.

The text proper starts on a new page, the introduction (if any) starts without heading. Use a maximum of three categories of headings, all to be typed left, and using capitals and lower-case letters. The first type may also be typed in small capitals, the second and third type may be typed in bold. After the heading the paragraph follows without blank. A fourth category may be used in descriptions of species, etc. It is separated from the following paragraph by a long dash $(-)$ to be typed as double dash.

New paragraphs should not be indented in wordprocessors. Scientific names of genera and species should be typed in italics or underlined. No underlining or italics are allowed for any other text. Abbreviations of museums ('codens') etc. should be typed in SMALL CAPS, if available.

## References

In the text they are given as Lopes (1982a), (Lopes 1982) or (Brown \& White 1975: 24). All cited papers should be listed alphabetically at the end of the paper under the heading 'References', papers not cited in the text should be omitted from the list of references. Examples for format:

Boer, P. J. den, 1970. On the significance of dispersal power for populations of carabid-beetles (Coleoptera, Carabidae). - Oecologia 4: 1-28.
Karsholt, O. \& E. S. Nielsen, 1976. Systematisk fortegnelse over Danmarks sommerfugle. - Scandinavian Science press, Klampenborg, 128 pp.
Johansson, R. \& E. S. Nielsen, 1990. Tribus Nepticulini. In: Johansson, R. et al. The Nepticulidae and Opostegidae (Lepidoptera) of NW Europe. - Fauna entomologica scandinavica 23: 111-238, pls.

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In new taxa the type material should be listed immediately after the name. Only holotype, lectotype, neotype, paratype and paralectotype are allowed. Label data should not be quoted literally (except for primary types), but arranged in a standardized sequence. Material should be listed alphabetically or chronologically under the present day countries or other geographical units. Long lists of non-type material should be summarized. Geographical names should be written according to present day spelling, original spelling or label names may be given in brackets. Use standard transcription for non-latin scripts (e.g. Pinyin for chinese, BSI for cyrillic, etc.) or refer to the 'Times Atlas of the World'.

Abbreviations ('codens') for depositories preferably follow R. H. Arnett, Jr., G. A. Samuelson \& G. M. Nishida (1993: The insect and spider collections of the world. Second edition. Sandhill Crane Press, Gainesville). Otherwise, they should be listed under 'Material and methods' or in the introduction.

Data for primary types of previously described species follow directly the reference to the original description as:

Elachista subnitidella Duponchel, [1843]: 326, pl. 77: 8. Lectotype $\delta$ [designated by van Nieukerken \& Johansson 1987: 471]: [Austria, Vienna region], Duponchel coll., Genitalia slide EvN 2522 (mNHN) [examined].

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All illustrations, including photographs, graphs, maps, etc. should be serially numbered as figures. No subdivision with letters is recommended. Illustrations are to be reduced to column width $(65 \mathrm{~mm}), 1.5 \times$ column width ( 102.5 mm ) or text width ( 135 mm ). Line figures should be mounted in blocks, or are printed singly. When all figures are mounted in fullpage blocks (after reduction: $135 \times 195 \mathrm{~mm}$ includ-
ing caption), they may be printed after the text, otherwise the approximate place in the text should be indicated with pencil in the margin of the manuscript. Line-drawings are numbered with pre-stencilled or pre-printed figures, which should not be too large after reduction, preferably using a font like 'Garamond' or 'Times'. Photographs should be unmounted glossy prints. Numbering of photos should be left to the discretion of the editors. Captions should be typed on a separate sheet (or in a separate file), consult a recent issue for style. Colour plates will only be printed at the author's expense.

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# THE TAXONOMY AND BIOGEOGRAPHY OF THE LORIAE GROUP OF THE GENUS BAETURIA STÅL, 1866 (HOMOPTERA, TIBICINIDAE) 


#### Abstract

Boer, A. J. de, 1993. The taxonomy and biogeography of the loriae group of the genus Baeturia Stål, 1866 (Homoptera, Tibicinidae). - Tijdschrift voor Entomologie 137: 1-26, figs. 1-102, table 1. [ISSN 0040-7496]. Published 15 July 1994 The loriae group is proposed for a presumably monophyletic group of 9 species. Species of the group combine characters seemingly synapomorphic for the species of Baeturia Stål, 1866, with characters seemingly synapomorphic for the species of Gymnotympana Stàl, 1861. Though the generic relationships of the loriae group are not clear, it is included here in the genus Baeturia, on account of shared male genital characters. One species ( $B$. Loriae Distant, 1897) is redescribed and eight species (B. bemmeleni, B. daviesi, B. fortuini, B. hamiltoni, B. hartonoi B. Bigrami, B. silveri, and B. wegeneri) are described as new. B. loriae is transferred back from Gymnotympana to Baeturia. A key to the males is presented. A redescription of $B$. tenuispina Blöte, 1960 is included in this publication, since a character of the clasper indicates a relationship with some species of the loriae group. Nevertheless, the attribution of that species to the loriae group is far from certain. The loriae group is endemic to New Guinea, most species are restricted to the central and eastern mountain ranges of that island. A. J. de Boer, Institute for Systematics and Population Biology (Zoölogisch Museum), University of Amsterdam, PO Box 4766, 1009 AT Amsterdam, The Netherlands. Key words. - Baeturia, loriae group, taxonomy, biogeography, New Guinea.


Current studies of the tibicinid cicadas of eastern Malesia show, that a number of New Guinean and Australian genera form a monophyletic group: 'the Baeturia and related genera complex' (de Boer 1990, 1991). The largest genus of this group, Baeturia Stal, 1866, appears to be non-monophyletic in its present concept, but, within that genus, several monophyletic species groups have been recognized. Taxonomic and biogeographic revisions have been published so far for the B. nasuta group (de Boer 1982), the B. conviva group (de Boer 1986), the B. bloetei group (de Boer 1989) and the B. viridis group (de Boer 1992).
These pages present the revision of a presumably monophyletic group of nine species, for which the name Baeturia loriae group is proposed. This group is of particular interest from a taxonomic point of view, since its species combine several characters regarded apomorphic for Baeturia, with characters apomorphic for Gymnotympana Stàl, 1861. The distribution of these characters within the group is such, that any reconstruction of the in-group relationships will unavoidably show several conflicting characters. The $B$. loriae group is endemic to northern, eastern and central New Guinea.
B. tenuispina Blöte has a character of the clasper in common with some species of the $B$. loriae group, but
differs in most other respects. This species is not attributed to the $B$. loriae group, but described here, since it seems more related to this group than to any of the other species groups of Baeturia.

## Material and methods

The material used for this study comes from the following institutions:

AMS - Australian Museum, Sydney; BMNH

- Natural History Museum [formerly: British Museum (Natural History)], London; BPBM Bernice P. Bishop Museum, Honolulu; csiro Commonwealth Scientific and Industrial Research Organisation, Australian National Insects collection, Canberra; KBIN - Koninklijk Belgisch Instituut voor Natuurwetenschappen, Brussel; MSNG - Museo Civico di Storia Naturale 'G. Doria', Genova; MVM Museum of Victoria, Melbourne; RMNH - Nationaal Natuurhistorisch Museum (formerly: Rijksmuseum van Natuurlijke Historie), Leiden; SMN - Staatliches Museum für Naturkunde, Stuttgart; ZMA - Institute for Systematics and Population Biology (Zoölogisch Museum), Amsterdam.

The following sources have been used for tracing
the localities: 'Atlas van tropisch Nederland' (1938), 'The Times Arlas of the World (Comprehensive Edition)' (1968), 'Papua New Guinea a travel survival kit' by Tony Wheeler, 1988, Lonely Planet Publications, Victoria, Australia and a 'List of New Guinea localities' published by the Bernice P. Bishop Museum (1966).

To examine the male genitalia, the pygofer was pulled out, after overnight softening, with a sharp needle inserted berween the pygofer and the 8th abdominal segment. The aedeagus was pulled out at the same time, by inserting the needle berween the claspers. Body and tegmen lengths of all specimens were measured, other measurements are based on a maximum of ten specimens, when available.

## Phylogeny

## Classification of the $B$. Loriae group

The generic classification of the loriae group presents a major problem. As stated above, the species combine characters of Baeturia with characters of Gymnotympana. Some of these characters appear to be unique for either Baeturia or Gymnotympana (or part of these genera). The distribution of these characters within the loriae group is not homogeneous; some species have more in common with Baeturia, others more closely resemble Gymnotympana.

The following characters are found in the loriae group and indicate a relationship with Baeturia, the first four are presumed apomorphies for that genus.
(1) a strongly curved aedeagus, tapering to its apex, with bluntly rounded lateral lobes and oval aedeagus pore. This aedeagus shape is unique for Baeturia.
(2) a narrow vertex, with large ocelli close together (fig. 3). Two species of Gymnotympana have an even narrower vertex. In other species of the complex, the vertex is broader than in Baeturia.
(3) a broad and triangular middle spine on fore femur (fig. 19) is shared by all species of Baeturia.
(4) a fairly narrow and frontally bluntly rounded postclypeus, distinctly protruding beyond vertex lobes and, in most species, distinctly and angularly swollen ventrally (figs. 3-4). A similar swelling only occurs in some species of Baeturia, the narrow, protruding, and rounded shape of the postclypeus is possibly apomorphic for Baeturia.
(5) a racher stout caudodorsal beak, often angularly bent and curved over anal valves. Other species of the complex have a much shorter and erect beak. The curved caudodorsal beak is probably synapomorphic for several of the Baeturia species groups together.
(6) almost straight parallel claspers, separated at the
base, and with a broad and rounded ventral hollow in the apical part. Such separated claspers are unique for Baeturia and Scottotympana de Boer, 1991.
(7) a distinct and angular clasper heel (in B. bemmeleni, B. hamiltoni, and B. wegeneri). A clasper heel is found in many species of Baeturia, but not in other genera.
(8) rows of setae on veins of tegmina and wings (in B. hamiltoni and B. wegeneri). Similar rows of setae are found in many Baeturia species, but not in any of the related genera. However, such setae also occur, though more distinctly, in several genera of the Prasiini, a possible sister group of the 'Baeturia and related genera complex'.
(9) dense brown speckling on body (in B. daviesi and $B$. wegeneri). Such speckling, though strongly variable in density, only occurs within Baeturia.
(10) irregularly shaped brown spots on tegmina (in $B$. pigrami and $B$. silveri, fig. 59). Such spots are only found in some species of Baeturia (e.g. B. guttulinervis, B. guttulipennis and B. nasuta).
(11) a distinct crest between the aedeagal lobes. A similar crest occurs in some species of the $B$. conviva group and some species related to that group.

Other characters found in the B. loriae group do not occur in other species groups of Baeturia, but seem to indicate a relationship with Gymnotympana. The first two must possibly be regarded as apomorphies for Gymnotympana, others are either restricted to part of that genus, or occur in several other genera as well.
(1) sexual dimorphism in wing venation (males with very slender anal fields, compare figs. 59 and 84) is shared by all species of Gymnotympana, but does not occur in other genera of the complex. (Similarly slender anal fields are recorded for Prasia Stål and Plautilla Stål (Boulard 1975), however. In Prasia this character is restricted to the males, but the literature does not record such dimorphism for Plautilla).
(2) a sharp ventrolateral fold in tergites 4-7 (more or less distinct in all species) is shared by nearly all species of Gymnotympana, but not found in related genera.
(3) an enlarged distal part of male operculum. Both, the broadly rounded shape (as found in B. bemmeleni, $B$. bamiltoni, and $B$. wegeneri), as the triangular shape (as found in $B$. daviesi, $B$. fortuini, and B. pigrami), are found in Gymnotympana, but do not occur in related genera.
(4) a very short,. often rudimentary meracanthus
(found in all species with enlarged opercula) is shared by Venustria Goding \& Froggatt, 1904 and most species of Gymnotympana. This character might be coupled to the character described immediately above.
(5) a laterally elongate basal part of male operculum (in six species) is shared by nearly all Gymnotympana species and not found in other genera.
(6) a broad hyaline border along hind margin of tegmen (in B. fortuini and B. loriae, fig. 84) is shared by all species of Gymnotympana, Scottotympana, and Venustria.
(7) an inwards curving ventral corner of tymbal, with strongly inwards directed connecting bar between abdomen and tymbal (only found in $B$. fortuint is found in most species of Gymnotympana, but not in other related genera.
(8) furthermore, all species except $B$. daviesi and $B$. wegeneri have a more or less distinct colour pattern of dark streaks and spots on head, pronotum, and abdomen (figs. 8-9). This colour pattern is much more distinct than in other species of Baeturia. An even more distinct colour pattern is found in many Gymnotympana species, in Scottotympana and in two species of Chlorocysta Westwood, 1851. Since these patterns are very variable between the various species, it is not clear whether the patterns found in the loriae group can be considered homologue with those in Gymnotympana.

The loriae group thus, seems to combine some apparently convincing synapomorphies of Baeturia and Gymnotympana. At this point of investigation, with several groups of the 'Baeturia and related genera complex' still undescribed, and with only vague indications as to the possible sister groups of the complex, it is not possible to fully evaluate the meaning of the character distribution in the loriae group. Current phylogenetic analysis, involving all species of the 'Baeturia and related genera complex', however, shows that a monophyletic loriae group as part of the genus Baeturia, is the most parsimonious solution. Furthermore, this solution is in accordance with the distribution of all male genital characters.

Nevertheless, the remarkable number of conflicting characters concentrated in the loriae group is upsetting to any phylogenetic reconstruction. Whatever generic allocation of the loriae group is favoured, the parallel evolution and / or the successive gain and loss of several characters must be presumed. Biochemical study of the loriae group could be interesting as a test for a hybrid origin of that group.

## Monophyly of the $B$. loriae group

The above discussed combination of characters found in the loriae group involves all those characters
that would, by non-existence of that group, be used as apomorphies for either Baeturia or Gymnotympana. As a result neither Baeturia nor Gymnotympana can be properly defined by shared apomorphies in exclusion of the loriae group. But also for the loriae group itself, no unambiguous apomorphies could be found. The group can be best defined by the unique combination of two characters: sexual dimorphism in wing, and an S -curved aedeagus with lateral lobes at the base of the curvation. Furthermore, there are three characters shared by all, or nearly all, species of the group, and not widely distributed outside the group, that could indicate a close relationship between the species. These characters are: 1) a distinct crest between the lateral lobes of aedeagus, only shared by some species of the B. conviva group and two species related to that group; 2) an enlarged and angularly rounded ventral part of the lateral lobe of pygofer, shared by only a few species of Gymnotympana (e.g. G. stridens); and 3) a rather short and broad eighth apical area of tegmen, tending to be shorter than in Gymnotympana or other species of Baeturia. The latter character is somewhat variable within the species and cannot be used as a diagnostic character.

## Ingroup phylogeny

As long as the problems concerning the phylogenetic position of the loriae group, relative to other species groups of Baeturia and to the genus Gymnotympana, are not satisfactorily solved, characters that occur either in Baeturia or Gymnotympana and also in some species of the loriae group cannot be properly evaluated as plesiomorphous or apomorphous, and can therefore not be used in a phylogenetic reconstruction of the group. Only the distribution of characters that appear to be unique for some species of the loriae group allows some remarks on ingroup phylogeny.
B. bemmeleni, B. bamiltoni, and B. wegeneri almost certainly form a monophyletic group. These species share an almost identical clasper, characterized by a nearly rectangular dorsal corner (figs. 11, 26, and 30), which is regarded synapomorphic. Furthermore, these species share an almost identical male operculum; very broad, slightly domed and broadly rounded at its apex (figs. 14, 22, and 38). A very similar operculum was found in G. olivacea Distant, 1905, but in that species the lateral margin of operculum is almost continuous with the lateral crest of operculum base. Some undescribed Gymnotympana species, related to G. olivacea, have a similarly broad and curved, but more angular, square-shaped, operculum.
B. daviesi, B. fortuini, B. bartonoi, B. Loriae, B. pigrami, and $B$. silveri, most probably form a monophyletic group on account of a shared dorsal protrusion on the clasper. This protrusion is partly, or complete-


Fig. 1. Localities of Baeturia bemmeleni, B. hamiltoni, B. hartonoi, B. tenuispina, and B. wegeneri.
ly fused with the upwards curving proximal part of the dorsal margin of clasper, and merges with the clasper base, were in many other Baeturia species the clasper heel is formed. B. tenuispina Blöte has a similarly shaped protrusion at the corner of its clasper heel (fig. 97) and should possibly be included in this group.
In B. daviesi, B. pigrami, and B. silveri the dorsal protrusion is very distinct and finger-shaped (figs. 40, 50,64 ); in $B$. hartonoi, this protrusion is more angular and strongly curved mesiad (fig. 70); and in B. fortuini it is rudimentary and only recognizable in the specimens from Mt. Missim (fig. 79). The dorsal protrusion is believed to be lost in the lectotype of B. loriae and the remaining specimens of $B$. fortuini.

The clasper of these six species is very broad in lateral view; the dorsal margin of the clasper curves upwards from clasper base, so that no clasper heel is formed. This clasper-shape is probably also synapomorphous.
B. fortuini, B. loriae, and B. bartonoi possibly form a monophyletic group on account of the shared elongate lateral lobes of aedeagus (figs. 73, 82, 92), and the very broad, angular and laminiform clasper.
B. fortuini and B. Loriae are very closely related and probably sister species. They share triangular bronzed spots in the tegmina (fig. 84) and a very long and slender aedeagus (figs. 82, 92). Possibly these two
species are synonyms, and the lectotype of $B$. loriae is just an aberration.
The relationships between $B$. daviesi, B. pigrami, and $B$. silveriare less clear. These species share a triangular clasper, with convex dorsal margin (lateral view), which must probably be regarded more primitive than the squarely laminiform clasper of the three species mentioned above. Especially the claspers of $B$. daviesi and B. pigrami are very similar, sharing a broadly rounded and outcurving dorsal lobe, but the clasper of $B$. bartono $i$ has a very similar outcurving lobe. B. pigrami and B. silveri are conspicuous by the brown patches along veins of tegmina. But B. guttulinervis Blöte and B. guttulipennis Blöte, two otherwise very different and not closely related species, have similar patches.

Table 1. Altitudinal ranges of the species of the B. loriae group in meters above sea level.

| B. bemmeleni | $35-1350$ |
| :--- | :---: |
| B. fortuini | $1100-2100$ |
| B. bamiltoni | $50-1200$ |
| B. bartonoi | 100 |
| B. Loriae | 1300 |
| B. pigrami | $120-1260$ |
| B. silveri | $1800-2590(8500 \mathrm{ft})$ |
| B. wegeneri | 250 |



Fig. 2. Localities of Baeturia daviesi, B. fortuini, B. loriae, B. pigrami, and B. silveri.

## Biogeography

The B. loriae group is endemic to New Guinea, and distributed in the northeastern part of Irian Jaya, and in central and eastern Papua New Guinea. Many specimens come from, or close by, the central and eastern mountain ranges. The group seems to consist mainly of montane species, probably with very restricted areas of distribution, often reaching altitudes of over 1000 m (table 1).
It is remarkable, that the subdivision made above, between B. bemmeleni, B. hamiltoni, and B. wegeneri versus the remaining species, is a subdivision between a mainly western and a mainly eastern species group (figs. 1, 2).

## Taxonomy

## Description of the B. loriae group

The species of the $B$. loriae group are predominantly brown coloured, sometimes greenish tinged. $B$. wegeneri and $B$. daviesi are densely speckled with small brown spots, the other species are conspicuous by a colour pattern of regularly distributed dark markings. These species generally have two dark spots or longitudinal streaks between eyes and lateral ocelli and often darkened lateral corners of postclypeus; a light immaculate middorsal band on pronotum, someti-
mes bordered by almost black lines, and dark brown streaks in and along the oblique fissures. The dark markings on the mesonotum form two semi-circular paramedian spots, at pronotal collar, two blackened dents in front of cruciform elevation and lateral bands, converging from pronotal collar to edges of elevation. The cruciform elevation and a triangular area in front of the elevation is generally light ochraceous and unmarked. On the male abdomen, dark markings tend to form dorsal, and sometimes lateral bands, while a latero-ventral row of dark spots is often very clear. The dorsal markings tend to be interrupted middorsally by a narrow lighter coloured line. Ventral side of male abdomen generally light ochraceous, segmental hind margins often red. Female abdomen more irregularly dark spotted all over, though the light middorsal band is often more conspicuous.
Females are shorter than males (on average), but have a larger, more robust head and thorax and often longer tegmina.
Head in dorsal view (fig. 3): Postclypeus distinctly protruding beyond vertex lobes and broadly rounded at frontal margin. Postclypeus 1.2-2.2 $\times$ as broad as long and $0.5-0.7 \times$ as broad as distance between eyes, strongly varying within the species. Head 0.7-0.8× as wide as pronotal collar and only slightly narrower than anterior width of pronotum. Vertex narrow, ocelli large and close together. Distance between lateral ocelli less than $1.5 \times$ the width of frontal ocellus and $0.7-1.4 \times$ as long as distance between eye and lateral ocellus. Eye $0.6-0.8 \times$ as wide as distance between eyes. Head $0.9-1.2 \times$ as long and 2.2-2.6 $\times$ as wide as distance between eyes.
Legs: Ochraceous, sometimes with longitudinal brown stains on femora and tibia. Fore femur (fig. 19) with row of three sharply pointed spines, diminishing in length towards tibia. Proximal spine about as long as distance to middle spine. Middle spine broad, triangular.

Tegmina and wings: Hyaline, though tegmina in some species with brown markings. Venation ochraceous, costa often reddish. Tegmina with 8 , wings with 6 apical areas. Costal area of tegmen very narrow. Anal fields of wing in male distinctly narrower than in female (compare figs. 59 and 84).

Tymbal organ (fig. 6): Six or seven parallel transverse sclerotized ridges spanning the tymbal from dorsal to ventral margin. Short intercalary ridges forming a midlateral band across tymbal.

Opercula: Basal part of operculum vaulted, with distinct crest around rectangular distolateral corner. Distal part of male operculum either very long and domed, and completely covering tymbal cavity in ventral view; or short, oblong-shaped, laminiform and flat against body, only partly covering tymbal cavity. Distal part of operculum medially extending

 al view; 6, tymbal; 7, female operculum.


Fig. 8. Baeturia hamiltoni sp. n., habitus male.
beyond meracanthus. Distal part of female operculum short, sickle-shaped and erect, medially extending just to base of meracanthus. Basal part of female operculum forming narrow rim around base of meracanthus.

Abdomen: With tergites 4-7 of the male abdomen sharply folded the the ventrolateral edge of abdomen, though these folds are not always distinct an all specimens.

Male genitalia: Pygofer with stout and often angularly bent caudodorsal beak. Lateral lobes of pygofer with small and bluntly rounded protuberances, that do not extend beyond pygofer margin. Lateral lobes


Fig. 9. Baeturia hamiltoni sp. n., habitus female.
strongly folded inwards. Claspers parallel and directed posteriad, slightly bending down towards their apices. Apical part of clasper with small, rounded and sharply edged clasper hollow. Aedeagus strongly Scurved, with two narrowly elongate lateral lobes at base of S-curvation, and a distinct crest between lateral lobes. Aedeagus in lateral view broad at base, gradually narrowing to pointed apex.

Female genitalia: Caudodorsal beak sharply pointed at apex. Ovipositor sheats reaching just beyond apex of beak (fig. 5).

Etymology: This revision is to form part of an area cladistic study of the New Guinean region, in which
the general patterns of distribution of organisms are compared to the geological history of that island. The new species described in this publication are named in honour of geologists, who contributed to a paleogeographic reconstruction of the area. The species are named after R. W. van Bemmelen, H. L. Davies, A. R. Fortuin, W. B. Hamilton, H. M. S. Hartono, C. J. Pigram, E. A. Silver, and A. Wegener.

## Key to the males

1. Clasper with distinct angular clasper heel; apical part of clasper slender in lateral view, with rectangular dorsal corner (fig. 11). Distal part of operculum broad and long, weakly domed, and broadly rounded at apex (fig. 14)

- Clasper without clasper heel; apical part of clasper in lateral view broader than clasper base, often with dorsal protrusion, partly fused to proximal part of dorsal margin (fig. 40). Distal part of operculum either oblong and flat (fig. 41), or triangular, domed and pointed (fig. 51) at apex ....... 3

2. Body densely speckled all over, not forming an unspeckled or lighter coloured middorsal band ..
B. wegeneri

- Body not speckled, but with regularly distributed dark markings and an immaculate middorsal band over part or whole length of body $\qquad$

3. Body length $25.4-29.7 \mathrm{~mm}$. Caudodorsal beak broad and truncate at apex .............B. Bamiltoni

- Body length 17.1-21.3 mm. Caudodorsal beak narrowly truncate or pointed at apex $\qquad$ B. bemmeleni

4. Aedeagus very long, with elongate and slender lateral lobes (fig. 82). Clasper quadrangular in lateral view; dorsal part of clasper angular, laminiform (figs. 70, 79, 80, 89) $\qquad$ .. 5

- Aedeagus short and broad, lateral lobes not elongate (fig. 45). Clasper triangular in lateral view; dorsal part of clasper swollen, with weakly convex dorsal margin (figs 40, 50, 64)

5. Postclypeus not swollen in lateral view. Tegmen immaculate. Aedeagus angularly bent to apex (fig. 73). Clasper with dorso-lateral lobe (fig. 70) B. hartonoi

- Postclypeus distinctly swollen in lateral view. Tegmen with triangular brown spots in apical areas (fig. 84). Aedeagus not angularly bent to apex (fig. 81). Clasper without dorso-lateral lobe (fig. 79) ... 6

6. Distal part of operculum triangular, domed and narrowly rounded at apex (fig. 77). Margin of aedeagus pore concave at apex (fig. 83) . B. fortuini

- Distal part of operculum square-shaped, flat against body (fig. 91). Margin of aedeagus pore not concave at apex (fig. 93) ..................B. Loriae

7. Tegmina with brown markings along veins (fig. 59) .. 8

- Tegmina immaculate B. daviesi

8. Postclypeus angularly swollen in lateral view. Distal part of operculum triangular and slightly domed. Hind margin of tegmen with distinct hyaline border
.B. pigrami

- Postclypeus not swollen in lateral view. Distal part of operculum square-shaped and flat against body. Hind margin of tegmen with narrow hyaline border .B. silveri


## Description of the species

## Baeturia hamiltoni sp. n.

(figs. 1, 3-20)
Type material. - Holotype ठ: 'Neth. Ind.-American New Guinea Exped. Rattan camp, $1200 \mathrm{~m}, 6 . \mathrm{iiii}$ 1939, L. J. Toxopeus', rmNh. - Paratypes: inian jaya: new guinea (W): Araucaria camp, $800 \mathrm{~m}, 8 . \mathrm{iii} .1939$, L. J. Toxopeus,
 10.iii.1939, 3우; 11.iii.1939, 18; 12.iii.1939, 1 ㅇ;

 29.iii. 1939, $10^{\circ}$, all RMNH; same data but 19.iii. 1939, 2 年; 28.iii. 1939, $2 \delta^{\circ}$, all ZMA; Bernhard camp, 100 m , 11.iv. 1939, L. J. Toxopeus, 2 , R, RMnH.

Other material. - PAPUA: NEW GUINEA (NE): Ambunti, Sepik R., $50 \mathrm{~m}, 10$ v. 1963 , R. Straatman, 1 $\delta^{\circ}$, вРвм.
B. hamiltoni is a large brown coloured, species with a light middorsal band on head and thorax, sometimes continuing over abdomen. The species is easily recognized by a small and almost black medial spot on pronotum, at pronotal collar. Females have a glistering middorsal band of silvery setae on the abdomen, making the light and immaculate middorsal band more conspicuous than in males.

## Description

Body of males brown, abdomen sometimes reddish, with distinct colour pattern of darkened markings and an immaculate middorsal band on head and thorax, sometimes continuing over abdomen. Females on average slightly smaller than males, with more robust head and thorax, but shorter abdomen and considerably longer tegmina. Male abdomen 1.3$1.4 \times$ as long as head and thorax, of female 0.9-1.0×. Tegmina of males 1.1-1.2 $\times$ as long as total body length, of females 1.4-1.5 $\times$.

Head (fig. 3): Light brown, with traces of red around ocelli, longitudinal dark brown to black streaks on verrex lobes, between eyes and lateral ocelli, and often brown spots at margins of postclypeus. Postclypeus very stout, 1.3-1.9× as long as broad, distinctly protruding beyond vertex lobes and broadly rounded at anterior margin. Postclypeus in lateral view (fig. 4) strongly inflated, and angularly protruding; dorsal part of anterior margin (lateral view) al-


Figs. 10-20. Baeturia hamiltoni sp. n. - 10, pygofer in lateral view; 11, clasper in lateral view; 12, clasper in dorsal view; 13, male caudodorsal beak in dorsal view; 14, male operculum; 15, aedeagus from aslant; 16, aedeagus in lateral view; 17, aedeagus from behind; 18 , detail crest between lateral lobes of aedeagus; 19, fore femur; 20 , female caudodorsal beak in dorsal view. Lettering: $\mathrm{bp}=$ basal part of operculum; $\mathrm{c}=\mathrm{crest}$ between lateral lobes of aedeagus; $\mathrm{cb}=$ caudodorsal beak; $\mathrm{dm}=$ distomedial margin of operculum; $\mathrm{di}=$ distal margin of pygofer; $\mathrm{dc}=$ crest around distolateral corner of basal part of operculum; do = dorsal margin of pygofer; $\mathrm{dp}=$ distal part of operculum; he = clasper heel; ho = clasper hollow; la = lateral margin of operculum; $m=$ meracanthus; $m e=$ medial margin of operculum; $p=$ protuberance on lateral lobe of pygofer; ve $=$ ventral margin of pygofer.
most straight, ventral part strongly concave. Head very narrow between eyes, distance between lateral ocelli only slightly larger than width of frontal ocellus. Distance between lateral ocelli 0.8-1.2 $\times$ distance between eye and lateral ocellus. Head narrower than anterior part of pronotum.

Thorax: Pronotum brown, with broad dark brown streaks, in and along oblique fissures, and broad, immaculate medial band. Markings variable in intensity and shape, and sometimes completely absent. All specimens with a small dark brown or black middorsal spot at margin of pronotal collar. Mesonotum light brown, with two paramedian dark spots at pronotum margin, darkened lateral streaks converging from pronotal collar to corners of cruciform elevation and two black spots in front of elevation. Cruciform elevation, and a narrow triangular area in front of el-
evation, often reaching to pronotal collar, light ochraceous and immaculate.

Legs: Fore femur (fig.19) with row of three erect spines, diminishing in length towards tibiae.

Tegmina and wings: Hyaline, though slightly bronzed, venation ochraceous or reddish tinged, but costa often bright red. Veins densely set with short setae. Tegmina with very short apical areas, 8th area almost square-shaped, and with a very narrow hyaline border along hind margin. This border slightly broader in wings.

Tymbal organs (fig. 6: Tympanum not curving inwards; connecting bar between tymbal and abdomen quite short and almost parallel to body axis. Six transverse sclerotized ridges spanning the tymbal, from dorsal to ventral margin. A 7th ridge, close to proximal tymbal margin, almost reaching ventral margin.

Six short intercalary ridges seem to form a band across tymbal.

Opercula: Male operculum (fig. 14) ochraceous brown with darkened streaks. Basal part of operculum almost oblong-shaped, but at half-width slightly longer than at lateral or medial margin. Distal part of operculum very broad, shell-shaped, and almost flat against body. Operculum broadly rounded at apex and reaching to, or beyond, margin of abdominal segment 3. Distomedial margin and distal is of lateral margin convex, proximal $/ 6$ of lateral margin straight. Medial margin short and straight, recurving to base of meracanthus. Opercula well separated medially; 1st sternite slightly elevated between opercula, with distinct distal lobe. Meracanthus very short, reaching to about $1 /$ the operculum length. Female operculum (fig. 7) sickle-shaped and erect, with weak crest along distal margin; area between operculum and base of meracanthus broad and flat.

Abdomen: Male abdomen light brown, with some darker brown spots on lateral sides and latero-ventral rows of dark spots on segments 3-7. Segments 2-7 darkened dorsally as well, though these markings often interrupted by a narrow lighter middorsal band. Segment 8 almost entirely dark brown, but pygofer ochraceous. Ventral side of abdomen light brown, immaculate. Segmental hind margins often reddish. Anterior margin of 2nd tergite convex medially; anterolateral sides of 2 nd tergite distinctly swollen adjacent to tymbal. Female abdomen more irregularly spotted all over. Light middorsal band more conspicuous than in males, often glistering with short, silvery setae. Ovipositor sheaths reaching just beyond apex of caudodorsal beak (fig. 5). Female caudodorsal beak (fig. 20) stout, slightly erect and rounded at apex.

Male genitalia: Pygofer slender in lateral view as in fig. 10. Dorsal margin straight, angularly bending into stout, short and slightly erect caudodorsal beak. Distal margin straight angularly bending into margin of beak. Lateral lobes of pygofer strongly curved inwards, with bluntly rounded lateral protuberances. Ventral margin angularly convex, but concave near base of pygofer. Caudodorsal beak in dorsal view (fig. 13) broad and truncate at apex. Clasper in lateral view (fig. 11) broad at base, with very distinct and angular clasper heel, strongly bent down at half-length, forming almost right angled dorsal corner. Dorsal margin of clasper concave between clasper heel and this dorsal corner. Distinct crest running along $2 / 3$ of dorsal margin, and bending outwards around aedeagus to$\because$ ards clasper base. Clasper hollow short, broadly rounded at apex. Clasper in dorsal view (fig. 12), with dorsal margins of clasper bases very close together, leaving only a narrow gap for anal valves. Aedeagus in lateral view (fig. 16) strongly concave around aedeagus pore, lateral lobes long and slender. Distinct crest,
at base of S-curvation, between lateral lobes (figs. 15, 18). Aedeagus pore (fig. 16) large and oval-shaped.

Measurements: Body length ${ }^{\text {oै }}: 25.4-29.7 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ $27.6 \mathrm{~mm} \pm 1.3$ ), $\quad: \quad 23.5-28.7 \mathrm{~mm}(\overline{\mathrm{x}} 26.6 \mathrm{~mm} \pm$ 1.4); tegmen length $\delta: 28.0-35.0 \mathrm{~mm}$ ( $\overline{\mathrm{x}} 32.5 \mathrm{~mm} \pm$ $1.8), f: 32.1-40.5 \mathrm{~mm}(\vec{x} 37.7 \mathrm{~mm} \pm 2.3)$; head length $\bar{\delta}: 2.3-2.6 \mathrm{~mm}(\overline{\mathrm{x}} 2.4 \mathrm{~mm}), \circ: 2.4-3.1 \mathrm{~mm}(\overline{\mathrm{x}}$ 2.6 mm ); pronotum length $\delta$ : 3.1-3.8 mm ( $\overline{\mathrm{x}} 3.5$ $\mathrm{mm})$,,$:=3.5-4.3 \mathrm{~mm}(\overline{\mathrm{x}} 4.0 \mathrm{~mm})$; mesonotum length ठ: $5.8-6.6 \mathrm{~mm}(\bar{x} 6.3 \mathrm{~mm}), \quad \uparrow: 6.5-7.8 \mathrm{~mm}(\overline{\mathrm{x}} 7.1$ mm ); head width $\delta$ : $5.5-6.2 \mathrm{~mm}(\overline{\mathrm{x}} 5.8 \mathrm{~mm})$, if: 5.9-6.9 mm ( $\bar{x} 6.6 \mathrm{~mm}$ ); width of pronotal collar ठ: $7.1-8.5 \mathrm{~mm}(\overline{\mathrm{x}} 8.0 \mathrm{~mm}), \quad$, $: 8.3-9.9 \mathrm{~mm}(\overline{\mathrm{x}} 9.2$ mm ).

Distribution (fig. 1). - B. hamiltoni is known from a very restricted area in Irian Jaya and from one locality in NW Papua.
Etymology. - The species is named in honour of the geologist W. B. Hamilton.
Remark. - Three females from Araucaria camp that have been described as B. guttulinervis Blöte (allo- and paratypes) are very similar in size and colour pattern to the females of $B$. hamiltoni, but have darkened patches along the venation of tegmen, are slightly brown speckled on the pronotum and miss the dark middorsal spot at pronotal collar.

## Baeturia bemmeleni sp. n .

(figs. 1, 21-29)
Type material. - Holotype ठ': 'NEW GUINEA NETH. Bodem, $100 \mathrm{~m}, 11 \mathrm{~km}$ SE of Oerberfaren, July 7-17. 1959'; 'T. C. Maa Collector', BPBM. - Paratypes: IRIAN: NEW GUINEA (W): Boven Digul gebied [Upper Digul area], 400 km N. Merauke, Dr A. Kalthofen, legit 1926, $10^{\star}$, ZMA; Boven Sermowai riv., $\pm 400 \mathrm{~m}, 8$.iv.1911, K. G. [K. Gjellerup], 20, RMNH.

Other material. - PAPUA: NEW GUINEA (NE): Eliptamin valley, $1200-1500 \mathrm{~m}, 16-31 . v i i .1959$, W.W. Brandt, 1 f, bPBM; NEW GUINEA (SE): Kiunga, Fly riv., 35 m, viii. 1969, J. and M. Sedlacek, 1 ㅇ, BPвм.
B. bemmeleni closely resembles $B$. hamiltoni in shape of male operculum and clasper, but is considerably smaller and less intensely coloured. The female from Eliptamin Valley deviates somewhat in shape and colour pattern, but the shape of its 8th apical area of tegmen suggests that this specimen belongs to the $B$. loriae group and it presumably belongs to this species.

## Description

Body of males ochraceous or reddish brown, with light brown patches, smaller than in foregoing species. Distribution of patches resemble markings in $B$. hamiltoni; leaving a light middorsal band on head and


Figs. 21-30. 21-29: Baeturia bemmeleni sp. n. - 21, pygofer in lateral view; 22, male operculum; 23, female operculum; 24, male caudodorsal beak in dorsal view; 25, female caudodorsal beak in dorsal view; 26, clasper; 27, aedeagus from behind; 28 , aedeagus in lateral view; 29, detail crest between lateral lobes of aedeagus. - 30: Baeturia wegeneri sp . n ., clasper.
thorax immaculate. This band sometimes interrupted by a vaguely darkened middorsal spot, at margin of pronotal collar. Females greenish brown, specimen from Eliptamin Valley densely brown speckled. Both females show the light brown middorsal marking on pronotum. Male abdomen 1.3-1.6× as long as head and thorax, of females $1.1 \times$. Male tegmen 1.1-1.2× as long as total body length, of females $1.3 \times$.

Head: Ochraceous, with brown markings on vertex lobes and along sides of postclypeus, but densely speckled in female of Elipramin Valley. Postclypeus strongly protruding, almost triangular in dorsal view, $1.8-2.3 \times$ as wide as long. Postclypeus in lateral view, angularly inflated as in B. hamiltoni, though in some specimens more rounded at anterior margin. Head about as wide as anterior part of pronotum. Distance between lateral ocelli $0.7-0.9 \times$ as long as distance between eye and lateral ocellus.

Thorax: Pronotum light brown, with dark brown spots along oblique fissures and a light, but narrow, middorsal band. Female from Eliptamin Valley brown speckled between medial fissures and showing this band only on proximal half of pronotum. Slightly darkened middorsal spot, at margin of pronotal collar, marking end of immaculate band; this spot less clear than in B. hamiltoni. Spot U-shaped in one male from Sermowai riv. and the female from Kiunga riv.; connecting two dark lines, running partly along the immaculate middorsal band. Mesonotum variable in
colour pattern between individual specimens, though always with a light, immaculate cruciform elevation and, with exception of the densely speckled female from Eliptamin Valley, with a light middorsal band or narrow triangular area in front of elevation. Two dark spots in front of cruciform elevation.
Tegmina and wings: Hyaline, veins slightly reddened. Tegmina with short 8th apical area, and with narrow hyaline border along hind margin. Wings with slightly broader hyaline border.

Tymbal organs: As in B. hamiltoni, not curving inwards, but with connecting bar between tymbal and abdomen distinctly longer and more curved inwards. Six transverse sclerotized ridges spanning the tymbal, from dorsal to ventral margin. A 7th ridge, close to proximal tymbal margin, almost reaching ventral margin. Six short intercalary ridges seem to form a band across tymbal.

Opercula: Male operculum (fig. 22) closely resembling that of $B$. hamiltoni, broadly rounded, shellshaped, and ochraceous brown with some darkened streaks. Basal part of operculum almost oblong, but weakly tapering mesiad at half its width. Distal part weakly domed, almost flat against body, its rounded apex reaching well beyond margin of abdominal segment 3. Distomedial margin and distal $2 / 2$ of lateral margin weakly convex, proximal $1 / 3$ of lateral margin more strongly convex. Meracanthus very short, reaching to about $1 / 4$ the operculum length. Female opercu-


Figs. 31-38. Batturia wegeneri sp. n. - 31, pygofer in lateral view; 32, pygofer from aslant; 33, first sternite; 34, caudodorsal beak in dorsal view; 35 , aedeagus from behind; 36 , aedeagus in lateral view; 37, detail crest between lateral lobes of aedeagus; 38 , operculum.
lum (fig. 23) sickle-shaped and erect, with broad and flat area between base of meracanthus and medial edge of operculum.

Abdomen: Male abdomen ochraceous brown, stained with dark brown spots, concentrated in nearly continuous middorsal band. Sternites immaculate. A latero-ventral row of clear dark spots on segments 3-7. Segmental hind margins bright red. Anterior margin of 2nd tergite convex medially; sides of 2nd tergite distinctly swollen and adjacent to tymbal. Female abdomen darker brown with irregular brown markings. Latero-ventral row of dark spots less conspicuous than in males. Segmental hind margins ochraceous. Female caudodorsal beak (fig. 25) long, slender, and pointed at apex.

Male genitalia: Pygofer (fig. 21) closely resembling that of $B$. bamiltoni, slender in lateral view. Dorsal margin straight, angularly bending into straight, stout and slightly erect caudodorsal beak. Distal margin straight, angularly bending into margin of beak. Lateral lobes of pygofer strongly incurved, with welldeveloped, bluntly rounded lateral protuberances. Ventral margin angularly convex, but concave near base of pygofer. Caudodorsal beak (fig. 24) short, sharply pointed or narrowly truncare at apex. Clasper
in lateral view (fig. 26) almost identical to that of $B$. hamiltoni, with very distinct angular clasper heel, and rectangular dorsal corner. Dorsal margin of clasper concave between clasper heel and this dorsal corner. Distinct crest running along $2 / 3$ of dorsal margin, and bending outwards around aedeagus towards clasper base. Clasper hollow short, broadly rounded at apex. In dorsal view, dorsal margins of clasper bases wider apart than in B. hamiltoni. Aedeagus in lateral view (fig. 28) slender, strongly concave along margin of pore. Lateral lobes long and slender. Distinct crest at base of S-curvation, between lateral lobes (fig. 29). Aedeagus seen from behind (fig. 27) very slender. Aedeagus pore narrow and oval.

Measurements: Body length $\delta$ : $17.1-21.3 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ $18.4 \mathrm{~mm} \pm 1.7$ ), $.9: 19.3$ and 21.7 mm ; tegmen length ©: 19.8-20.4 mm, $:: 27.8$ and 28.6 mm ; head length $\delta: 1.5-1.8 \mathrm{~mm}(\overline{\mathrm{x}} 1.7 \mathrm{~mm}), ~ ㅇ, ~: 1.9$ and 2.2 mm ; pronotum length $\delta$ : $1.9-2.2 \mathrm{~mm}(\bar{x} 2.1 \mathrm{~mm}), ~ q: 2.8$ and 3.2 mm ; mesonotum length $\delta: 3.4-3.8 \mathrm{~mm}$ ( $\overline{\mathrm{x}} 3.7$ $\mathrm{mm}), ~ f: 5.0$ and 5.3 mm ; head width $\delta$ : $: 4.2-4.5 \mathrm{~mm}$ ( $\bar{x} 4.3 \mathrm{~mm}$ ), 우: 5.0 and 6.0 mm ; width of pronotal collar $\delta$ © : $5.2-5.4 \mathrm{~mm}(\bar{x} 5.3 \mathrm{~mm})$, $ㅇ: 9.6$ and 7.2 mm .

Distribution (fig. 1). - B. bemmeleni is known
from several localities in the northern and central parts of Irian Jaya.

Etymology. - The species is named in honour of the geologist R. W. van Bemmelen.

## Baeturia wegeneri sp. n.

(figs. 1, 30-38)
Type material. - Holotype đ': 'New guinea: NE, May R. Petrol sta., $250 \mathrm{~m}, 3 . \mathrm{vi} .1963$ ' [print]; 'Dry Forest' [print]; 'R. Straatman Collector BISHOP' [print], BPBM. - Paratype: same locality and collector as holotype but 100 m , 31.v. 1963,1 ठ), вРВМ.
B. wegeneri is an ochraceous brown coloured species, with densely brown speckling all over its body, as characteristic for many Baeturia species, and in this respect quite different from most other species of the B. loriae group. However, B. wegeneri is very similar to the two foregoing species in shape of male operculum and genitalia.

## Description

Body ochraceous brown, densely brown speckled all over. Abdomen strongly inflated, $1.7 \times$ as long as head and thorax. Tegmina about as long as body length.

Head: Ochraceous, brown speckled. Postclypeus broad and oblong in dorsal view, 1.4-1.6 $\times$ as broad as long. Postclypeus angularly inflated ventrally (as in fig. 4). Head narrower than anterior part of pronotum. Distance between lateral ocelli $0.8-1.0 \times$ distance between eye and lateral ocellus.

Thorax: Pronotum ochraceous, brown speckled in middorsal band and on pronotal collar. Mesonotum grey-brown, densely brown speckled. Two dark spots in front of speckled cruciform elevation, almost lost in speckling.

Tegmina and wings: Hyaline, veins ochraceous or slightly reddish and set with short setae. Tegmen with short and squarish 8th apical area, and narrow hyaline border along hind margin. Wing with slightly broader hyaline border.

Tymbal organs: Not curving inwards, connecting bar between tymbal and abdomen quite short and almost parallel to body axis. Six transverse sclerotized ridges spanning the tymbal, from dorsal to ventral margin. A 7th ridge, close to proximal tymbal margin, almost reaching ventral margin. Six short intercalary ridges seem to form a band across tymbal.

Operculum (fig. 38): Broad and rounded as in foregoing species, ochraceous with some darkened stains near lateral margin. Basal part of operculum almost oblong, but weakly tapering towards mesiad at half its width. Distal part slightly domed, almost flat against body, broadly rounded at apex and reaching
just beyond margin of abdominal segment 3 . Distomedial margin and distal $2 / 3$ of lateral margin convex, proximal $1 / 3$ of lateral margin more strongly convex. Opercula well separated medially, by distinctly smaller and anteriorly lobate first sternite (fig. 33). Meracanthus short, reaching to about $1 / 5$ of operculum length.
Abdomen: Ochraceous brown, tergites densely brown speckled all over, sternites unspeckled. Lateroventral row of dark spots almost inconspicuous, by speckling. Segmental hind margins bright red. Anterior margin of 2nd tergite convex middorsally; sides of 2nd tergite weakly swollen and adjacent to tymbal.

Genitalia: Pygofer in lateral view (fig. 31) resembling that of foregoing species. Dorsal margin straight, angularly bending into stout and slightly erect caudodorsal beak. Distal margin straight and more gradually bending into margin of beak. Lateral lobes of pygofer strongly curved inwards, with well-developed, bluntly rounded, lateral protuberances. Ventral margin straight, not concave to base; ventral margins converge to sharp angle at base of pygofer opening (fig. 32). Caudodorsal beak slightly longer than in B. bemmeleni, very slender in dorsal view (fig. 34) and sharply pointed apically. Clasper in lateral view (fig. 30) strongly resembling that of two foregoing species, with a very distinct angular clasper heel and rectangular dorsal corner. Clasper strongly bent down to apex at half-length. Distinct crest running along $2 / 3$ of dorsal margin. Clasper hollow short and broadly rounded at apex. Dorsal margins of clasper bases wider apart than in B. hamiltoni. Aedeagus in lateral view (fig. 36) concave along margin of pore, with broader basal lobes than in B. bemmeleni. Distinct crest at base of S-curvation, between basal lobes (fig. 37). Aedeagus from behind (fig. 35) broader than in B. bemmelent, with a short and broad oval pore.

Measurements: Body length: 26.0 and 27.3 mm ; tegmen length: 26.4 and 26.6 mm ; head length: 2.0 and 2.3 mm ; pronotum length: 2.6 and 2.7 mm ; mesonotum length: 5.3 mm ; head width: 4.7 and 4.8 mm ; width of pronotal collar: 6.4 and 6.5 mm .

Distribution (fig. 1). - B. wegeneri is only known from May river, in central west Papua New Guinea.

Etymology.- The species is named in honour of the geologist A. Wegener.

## Baeturia silveri sp. n.

(figs. 2, 39-48)
Type material. - Holotype ठ': 'NEW GUINEA: NE Purosa, 20-26 km SE Okapa, $1800-2020 \mathrm{~m}, 28$. viii. $1964^{\prime}$ [print]; 'J. and M. Sedlacek Collectors BISHOP' [print], BPBM. Paratypes: same data as holotype $1 \delta^{\star}, 1$, BPBM.

Other material. - papua: New guinea (NE): Kandep,


Fič. 39-48. Bacturiz silzerisp. n. -39 , pygofer in lateral view; 40, clasper; 41, male operculum, Kandep; 42, male caudodoral heak in doral view: 43 , female caudodorsal beak in dorsal view; 44, aedeagus from behind; 45, aedeagus in lateral view; 4 (3. detail crevt hetween lateral lobes of aedeagus; 47, male operculum, holotype; 48, female operculum. Lettering: $\mathrm{d}=$ distal margin of operculum; 1 = lateral margin of operculum; $m=$ medial margin of operculum.

Western Highlands, 8000 ft , 23.xii.1961-14.ii.1962, W. W. Brandt, $1 \delta^{\circ}$. Csiro.
B. silveri can be recognized by the brown speckled tegmina and short, oblong male operculum.

## Description

Body of males light brown or ochraceous, with a colour pattern of dark brown spots and a narrow, immaculate, middorsal band as characteristic for most species of the B. Loriae group. Female slightly reddish brown with irregularly distributed dark spots. Male abdomen $1.4-1.6 \times$ as long as head and thorax, of female $1.1 \times$. Tegmina of males $1.2-1.4 \times$ as long as total body length, of female $1.5 \times$.

Head: Ochraccous brown, with dark brown streaks on vertex lobes, almost black between eyes and lateral ocelli. Female with black markings between pronotum and ocelli. Postclypeus sometimes darkened towards lateral margins, 1.6-1.9 $\times$ as broad as long and broadly rounded at anterior margin. Postclypeus not, or only slightly, swollen ventrally, anterior margin (lateral view) straight or weakly convex. Head narrower than anterior part of pronotum. Distance between lateral ocelli $0.9-1.3 \times$ distance between eye and lateral ocellus.

Thorax: Pronotum light brown, darkened along fissures; with a lighter, ochraceous, middorsal band, sharply bordered by irregular black streaks. Middorsal band abruptly widening near pronotal collar, and ending at a vaguely darkened middorsal spot. Middorsal band reddish brown and brown speckled in the female, but clearly recognizable by its black lining. Pronotal collar red-brown, brown speckled. Mesonotum with two distinct semi-circular dark spots at pronotum margin and darkened, greenish tinged, broad lateral streaks, converging from pronotum margin to corners of cruciform elevation. Elevation and narrowly pointed triangular area in front of elevation, light ochraceous. Mesonotum of female, including cruciform elevation, reddish tinged and densely brown speckled.

Tegmina and wings: Hyaline, venation reddish ochraceous. Tegmen with irregular brown markings, predominantly along venation. Hyaline border along hind margin of tegmen broader than in foregoing species and almost as broad as in wing. The 8th apical area of tegmen slightly longer, more slender, than in other species of this group.

Tymbal organ: Not curving inwards, connecting bar between tymbal and abdomen quite short and al-
most parallel to body axis. Five transverse sclerotized ridges spanning the tymbal, from dorsal to ventral margin. A 6th ridge almost reaching ventral margin, and a 7 th, most proximal, ridge running from dorsal margin to about half the tymbal width. Six intercalary ridges seem to form a band across tymbal. Traces of red between tymbal ridges.

Opercula: Male operculum (fig. 47) quite different from that of the three foregoing species, more resembling the opercula found in other groups of Baeturia. Basal part of operculum oblong, equally long at lateral and medial margins. Distal part angularly oblong, wider than long and flat against the body, almost completely covering tymbal cavity. Lateral and medial margin almost straight, distal margin weakly convex. Distolateral corner broadly rounded, distomedial corner angular. Opercula wide apart medially, separated by broad and rounded 1st sternite. Meracantus almost as long as operculum. Operculum of Kandep specimen (fig.41) much longer and more directed mesiad, with longer lateral margin. Female operculum (fig. 48) sickle-shaped and erect. Distal margin convex.

Abdomen: Male abdomen light brown, ochraceous in Kandep specimen. Two parallel dorsal tows of dark spots along a narrow light middorsal band and a very clear latero-ventral row of dark spots on segments 37. Lateral and ventral sides of abdomen slightly brown stained. Abdomen of Kandep specimen with midlateral band of dark spots, as in B. fortuini and $B$. loriae. Anterior margin of 2nd tergite convex medially; sides of 2nd tergite distinctly swollen and adjacent to tymbal. Female abdomen irregularly brown spotted all over, though with unstained light middorsal band. Female caudodorsal beak (fig. 43) pointed at apex.

Male genitalia: Pygofer in lateral view rounded, with stout caudodorsal beak (fig. 39). Dorsal margin straight almost continuous with erect caudodorsal beak. Distal margin straight or slightly concave, broadly rounded into margin of beak. Lateral lobes of pygofer curved inwards, with well-developed, bluntly rounded protuberances, that curve slightly upwards towards pygofer margin. Ventral margin angularly convex, but concave near base of pygofer. Caudodorsal beak (fig. 42) rounded, almost truncate at apex. Clasper in lateral view (fig. 40) strongly rounded and quite different from the three foregoing species. Clasper gradually widening distad to clasper base, so that no clasper heel is formed. Dorsal margin of clasper rounded, forming an almost globular, slightly inwards curved, protrusion at about $1 / 4 \mathrm{its}$ length and, distad to this protrusion, convex to apex. Clasper slightly curved outwards, towards dorsal margin and around aedeagus. Clasper hollow narrow triangular, narrowly rounded at apex. Aedeagus longer
than in foregoing species, in lateral view (fig. 45) strongly concave along margin of pore, with long and slender basal lobes. Distinct crest at base of S-curvation between basal lobes (fig. 46). Aedeagus pore (fig. 44) large and oval-shaped, but truncate at apex.

Measurements: Body length ठ': $20.0-22.0 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ $20.9 \mathrm{~mm})$, 오: 18.8 mm ; tegmen length oै: $25.9-28.8$ $\mathrm{mm}(\overline{\mathrm{x}} 27.2 \mathrm{~mm})$, 우:28.6 mm; head length ot: 1.5 1.7 mm, ㅇ: : 1.8 mm ; pronotum length ot: 2.2-2.5 mm, 오: 2.5 mm ; mesonotum length of $4.2-4.6 \mathrm{~mm}$, 우: 5.0 mm ; head width $\delta: 3.8-4.3 \mathrm{~mm}$, ㅇ $: 4.4 \mathrm{~mm}$; width of pronotal collar $\delta: 5.4-6.2 \mathrm{~mm}, ~: ~: ~ 6.2 \mathrm{~mm}$.

Distribution (fig. 2). - B. silveri is known from only two localities, both in the central mountain ranges of Papua New Guinea.

Etymology. - The species is named in honour of the geologist E.A. Silver.

## Baeturia pigrami sp. n.

(figs. 2, 49-59)
Type material. - Holotype os: 'Museum Leiden Neth New Guinea Exp. Star Range, 1260 m, Sibil, 21.viii. 1959, op licht' [print], RMNH.

Other material. - PAPUA: NEW GUINEA (NE): Feramin, 150-120 m, 1-6.vi.1959, W.W. Brandt, 2 오, BPBM.

Of this species only one male is available, it resembles B. silveri in colour patterns of body and tegmina, and in clasper shape. However, B. pigrami is much larger, has very large triangular opercula, and an angularly swollen postclypeus. Two females, with similar colour patterns and from the same area, probably belong to this species.

## Description

Body of male dark reddish brown, with very distinct light middorsal band on head and thorax. This band much broader in females, and continuing over abdomen. Male abdomen strongly inflated and $1.5 \times$ as long as head and thorax of females $1.0 \times$. Tegmina of male $1.2 \times$ as long as total body length, of females 1.4-1.5×.

Head: Light brown with dark stains on vertex lobes, mainly between lateral ocelli and eyes. Vertex and postclypeus with long setae, more densely set than in foregoing speces. Postclypeus unstained, light ochraceous in females, slightly brown suffused in male. Male postclypeus $1.2 \times$ as broad as long, in females 1.5 and $2.2 \times$. Postclypeus angularly swollen, almost conical in lateral view (as in fig. 4). Vertex in male very narrow, with ocelli close together. Head narrower than anterior part of pronotum. Male head $1.2 \times$ as long as distance between the eyes, in female $0.9 \times$. Distance between lateral ocelli in male $0.7 \times$ distance between eye and lateral ocellus, in females 1.1-1.2×.


Figs. 49-58. Baeturia pigrami sp. n. -49 , pygofer in lateral view; 50, clasper; 51, male operculum; 52, aedeagus from behind; 53. aedeagus in lateral view; 54, detail crest between lateral lobes of aedeagus; 55 , male caudodorsal beak in dorsal view; 56, female operculum; 57, female caudodorsal beak in dorsal view; 58, tymbal.

Thorax: Pronotum dark brown with light ochraceous middorsal band, narrow and sharply defined in male. much broader and more diffuse in females. Pronotal collar slightly reddish and, especially in male, stained with brown. Male mesonotum dark brown, but light ochraceous in triangle-shaped area in front of reddish brown cruciform elevation. Female mesonotum with continuous broad ochraceous middorsal band. Two black spots in front of cruciform elevation in male and female.

Tegmina and wings (fig. 59): Hyaline, though tegmina with irregular brown markings, predominantly along veins, venation reddish brown. Eight apical area short and broad, especially in females. Tegmen and wing with very narrow hyaline border along hind margin.

Tymbal (fig. 58): Five transverse sclerotized ridges spanning the tymbal from dorsal to ventral tymbal margin. A Gth ridge almost reaching ventral tymbal margin, and a 7 th, most proximal, ridge running from dorsal margin to only about half the tymbal width. Six intercalary ridges seem to form a band
across tymbal. Traces of red between tymbal ridges.
Opercula: Male operculum (fig. 51) very large, reddish ochraceous with dark stains. Basal part of operculum with greatest length at $1 / 4$ its width, slightly tapering towards lateal margin, more strongly tapering to base of meracanthus. Distal part triangle-shaped and slightly domed, narrowing towards narrowly rounded, almost pointed apex and reaching to halflength 3rd abdominal segment. Lateral margin of male operculum convex near base, concave at halflength, and weakly convex towards apex. Distomedial margin concave near apex and convex to short and straight medial margin. Opercula close together medially, separated by narrow and weakly elevated 1st sternite. Meracanthus very short, reaching to about $1 / 6$ the operculum length. Female operculum (fig. 56) sickle-shaped, slightly angular at distal margin, and erect. Base of operculum forming narrow rim around base of meracanthus.

Abdomen: Male abdomen brown, dark brown stained dorsally and on sides of 7 th and 8 th segments, though lighter and reddish tinged, in very narrow


Fig. 59. Baeturia pigramin. sp., female tegmen and wing.
middorsal band. A latero-ventral row of slightly darkened patches, hardly conspicuous. Segmental hind margins slightly red. Anterior margin of 2nd tergite convex middorsally. Lateral parts of 2nd tergite, with auditory capsules, and anterolateral part of 3rd tergite, weakly incurving towards ventral corner of tymbal. Tergites 4-7 with sharp latero-ventral fold. Female abdomen ochraceous, densely stained with irregular brown spots laterally and ventrally, but with broad and immaculate middorsal band. Female caudodorsal beak very characteristic in dorsal view (fig. 57 ), with greatest width at about $2 / 3$ its length and sharply pointed at apex.

Male genitalia: Pygofer in lateral view (fig. 49). Dorsal margin of pygofer bending gradually into erect caudodorsal beak. Distal margin straight, angularly bending into margin of beak. Lateral lobes of pygofer strongly curving inwards, with well-developed, bluntly rounded protuberances. Ventral margin angularly convex, but strongly concave to base of pygofer. Caudodorsal beak in dorsal view (fig. 55) very slender, narrowly rounded, almost pointed at apex. Clasper (fig. 50) closely resembling that of B. silveri, not forming a clasper heel and slightly bent down to apex. Dorsal margin of clasper strongly bent upwards from clasper base, forming a long finger-shaped, slightly incurved protuberance. This protuberance distinctly longer than in B. silveri. Clasper strongly bent outwards distally of this protuberance, and forming a broadly rounded dorsal crest. Clasper hollow longer than in B. silveri. Aedeagus in lateral view (fig. 53) slightly concave along margin of pore, with narrow
and elongate lateral lobes. Aedeagus with distinct crest at base of S-curvation, between lateral lobes (fig. 54). Aedeagus pore broad and short, oval-shaped (fig. 52).

Measurements: Body length $\bar{\delta}: 27.0 \mathrm{~mm}, ~ ㅇ: 20.9$ and 22.4 mm ; tegmen length $\delta: 32.0 \mathrm{~mm}, \circ: 31.3$ and 31.8 mm ; head length $\delta^{*}: 2.6 \mathrm{~mm}, ~ ㅇ: 9: 2.1$ and 2.3 mm ; pronotum length $\left.\delta^{\circ}: 2.9 \mathrm{~mm}, ㅇ:\right\}: 3.5$ and 3.6 mm ; mesonotum length $\delta$ : $: 6.1 \mathrm{~mm}, ~ ㅇ: 6.1$ and 6.6 mm ; head width $\delta$ : $: 5.4 \mathrm{~mm}, 9: 5.4$ and 5.7 mm ; width of pronotal collar ot: $7.7 \mathrm{~mm}, 9: 7.8$ and 8.4 mm .

Distribution (fig. 2). - The only male known of this species comes from Sibil, Star Range, in the central mountain ranges of New Guinea. Two females, that probably belong to this species, come from nearby Feramin.

Etymology. - The species is named in honour of the geologist C. J. Pigram.

## Baeturia daviesi sp. n.

(figs. 2, 60-66)
Type material, - Holotype ठ̄: 'Papua, Mt Lamington, G. H. Muray, 1924' (written); 'W. W. Froggatt Collection' (print); 'CANB' (print), Csiro.

Of this species only one male is available. B. davie$s i$ is very similar to $B$. pigrami in the shapes of the male operculum and genitalia, but $B$. daviesi is distinguished by a distinctly smaller size, immaculate tegmina and the absence of distinct colour markings, characteristic to B. pigrami and most other species of


Figs. 60-67. - 60-66, Baeturia daviesin. sp., 60, pygofer in lateral view; 61, pygofer from aslant; 62, operculum; 63, caudodorsal beak in dorsal view; 64 , clasper; 65 , aedeagus from behind; 66 , aedeagus in lateral view. -67 , Baeturia hartonoi sp. n ., fore femur.
the $B$. loriae group, though its abdomen is densely brown speckled, as in $B$. wegeneri.

## Description

Body dull grey-brown with vaguely lighter coloured middorsal band, slightly more distinct on pronotum. Abdomen $1.4 \times$ as long as head and thorax. Tegmina $1.1 \times$ as long as body length.
Head: Greyish brown, with vaguely darkened spots between eyes and lateral ocelli. Postclypeus triangularly protruding, $1.5 \times$ as wide as long. Postclypeus angularly swollen ventrally, almost conically protruding in lateral view (as in fig. 4) Head narrower than anterior part of pronotum. Vertex very narrow, with ocelli close together; distance between lateral ocelli about as wide as frontal ocellus and $0.8 \times$ the distance between eye and lateral ocellus.

Thorax: Pronotum greyish brown with vaguely lighter coloured. middorsal band, most conspicuous over proximal half. Mesonotum brown with a greenish tinge, without special colour markings, apart from two dark spots in front of slightly reddish cruciform devation.

Legs: Proximal spine of fore femur rather short, about as long as distance to middle spine.

Tegmina and wings: Hyaline and immaculate, veins ochraceous. Tegmina with fairly short, square, 8th apical area, and very narrow hyaline border along hind margins. This border is slightly broader in wing.
Tymbal organ: Tymbal not curved inwards towards distal margin; connecting bar between tymbal and abdomen almost parallel to body axis. Six transverse sclerotized ridges spanning the tymbal from dorsal to ventral tymbal margin. Most proximal ridge narrowing considerably towards the ventral tymbal margin and only just reaching it. Six short intercalary ridges seem to form a band across the tymbal. Traces of bright red between the ridges, especially on dorsal half of tymbal.
Operculum (fig. 62): Very similar to that of B. pigrami, but smaller, with dark stains near apex. Basal part of operculum somewhat quintangular, with its greatest length at base of lateral margin of distal part, weakly tapering to distolateral corner and more strongly tapering to base of meracanthus. Distal part large, triangle-shaped and slightly domed, curved to abdomen near its almost pointed apex, and reaching to about half-lenght abdominal segment 3. Proximal $1 / 3$ of lateral margin slightly convex, distal $/ 2 / 3$ almost straight to apex. Distomedial margin concave near


Figs. 68-75. Baeturia hartonoi sp. n. -68 , pygofer in lateral view; 69, caudodorsal beak in dorsal view; 70, clasper; 71, postclypeus in lateral view; 72, operculum; 73, aedeagus in lateral view; 74, aedeagus from behind; 75 , detail crest between lateral lobes of aedeagus.
apex and convex to almost rectangular medial corner. Medial margin short and straight. Opercula close together medially, separated by narrow and distally lobate 1st sternite. Meracanthus very short, reaching to about $1 / 6$ the operculum length.

Abdomen: Light brown with brown speckling, darker brown dorsally and lateroventrally, with a slightly lighter coloured narrow middorsal line, though without a distinct row of latero-ventral spots. Segmental hind margins bright red. First tergite only slightly shorter than 2nd. Anterior margin of 2nd tergite weakly convex middorsally. Ventrolateral part of 2nd tergite, auditory capsules, and anterolateral corners of 3rd tergite, curved inwards towards ventral corner of tymbal. Anterolateral part of 2nd tergite adjacent to tymbal.

Genitalia: Pygofer in lateral view (fig. 60) slender. Dorsal margin straight, continuous with straight, slender and erect caudodorsal beak. Distal margin of pygofer straight, angularly bending into margin of beak. Lateral lobes of pygofer bent inwards, with small, bluntly rounded protuberances. Caudodorsal beak very short, pointed at apex (fig. 63). Ventral margin convex, but weakly concave to base of pygofer; ventral margins converging to sharp angle at base of pygofer opening (fig. 61). Clasper (fig. 64) much resembling that of B. pigrami, without clasper heel, but with dorsal margin strongly bent upwards from clasper base, forming a long finger-shaped, slightly incurved protuberance. Dorsal margin strongly bent outwards distad to protuberance, forming a broad
and distinctly laterally protruding crest. Clasper slightly bending down to rounded apex. Apical part of clasper with small clasper hollow. Aedeagus in lateral view (fig. 66) slightly concave along margin of pore, with narrow and elongate lateral lobes. Aedeagus with distinct crest at base of S-curvation, between lateral lobes. Aedeagus pore broad and short, oval-shaped (fig. 65).

Measurements: Body length: 23.9 mm ; tegmen length: 25.6 mm ; head length: 2.0 mm ; pronotum length: 2.8 mm ; mesonotum length: 5.1 mm ; head width: 4.8 mm ; width of pronotal collar: 6.5 mm .

Distribution (fig. 2). - The only specimen available comes from Mt. Lamington on the Papuan Peninsula.

Etymology. - The species is named in honour of the geologist H.L. Davies.

## Baeturia hartonoisp. n.

(figs. 1, 67-75)
Type material. - Holotype ô: 'PNG: NEW GUINEA: SE: S Highlands Distr.: Kutubu: Tugiri, 1000 m, 7-9.ii.1978' (print); 'J. L. Gressitt Collector BISHOP Museum' (print), ВРВМ.

Of this species only one male specimen is available. B. hartono $i$ is a small species, that can be recognized by an elongate apical part of aedeagus and a lateral protuberance on the clasper.

## Description

Body ochraceous brown, head and thorax slightly reddish. Abdomen with conspicuous dark dorsal and midlateral bands, and slightly inflated. Abdomen $1.3 \times$ as long as head and thorax. Tegmina $1.3 \times$ as long as body length.

Head: Brown, slightly darkened around ocelli, but not as distinctly marked as in some other species of the B. loriae group. Postclypeus distinctly protruding beyond vertex lobes, broadly rounded at anterior margin, and $1.8 \times$ as broad as long. Postclypeus only very slightly swollen ventrally, anterior margin (lateral view) weakly convex (fig. 71). Head narrower than anterior part of pronotum. Distance between lateral ocelli about $1.5 \times$ the width of frontal ocellus and $1.2 \times$ the distance between eye and lateral ocellus.

Thorax: Pronotum, plain brown with no special markings. Mesonotum almost plain grey-brown, cruciform elevation slightly lighter ochraceous. Two vaguely darkened spots in front of elevation hardly visible.

Legs: Proximal spine of fore femur strongly bent, almost adjacent to femur, and shorter than distance to middle spine (fig. 67).

Tegmina and wings: Hyaline and immaculate, venation ochraceous. Tegmen with narrow hyaline border along hind margin, wing with fairly broad border.

Tymbal organ: Tymbal not curved inwards towards distal margin; connecting bar between tymbal and abdomen almost parallel to body axis. Six sclerotized transverse ridges spanning the tymbal from dorsal to ventral tymbal margin. A 7th, most proximal, ridge almost reaching ventral tymbal margin. Seven short intercalary ridges seem to form a midlateral band across tymbal.

Operculum (fig. 72): Short and oblong shaped as in $B$. silveri. Basal part of operculum oblong; equally long at lateral and medial margins. Distal part broader than long, flat against the body. Operculum directed mesiad, so that connecting bar between tymbal and abdomen becomes partly visible in ventral view. Lateral margin very short, bending gradually into slightly convex distal margin. Distomedial corner angular. Medial margin almost straight. Medial part of operculum reaching beyond margin of abdominal segment 2 . Opercula widely separated medially, by broad and rounded lst sternite. Meracanthus reaching to about $3 / 4$ the operculum length.

Abdomen: Ochraceous, greenish tinged, darkened in dorsal and lateral bands, though less intensely in narrow middorsal line. Latero-ventral row of dark spots most clear on segments 3-5 on right side, and on segments $3-4$ on left side of body, but faded in successive segments. First tergite very short medially. Anterior margin of 2nd tergite weakly convex medially. Lateral parts of 2nd tergite not incurved.

Genitalia: Pygofer in lateral view (fig. 68). Dorsal margin straight and angularly bending into slender and erect caudodorsal beak. Distal margin broadly rounded into margin of beak. Lateral lobes of pygofer strongly bent inwards, with small and bluntly rounded protuberances. Ventral margin weakly convex, but distinctly concave towards base of pygofer. Caudodorsal beak in dorsal view (fig. 69) slender and pointed at apex. Clasper very broad, squarely rounded in lateral view (fig. 70), abruptly broadening dorsally, distally of clasper base, so that no clasper heel is formed. Dorsal margin angularly bending down at about half-lenght, forming a small and slightly incurving dorsal protrusion on clasper. This protrusion seems to function as a support or lock for the aedeagus, since it curves inwards, around aedeagus. This in contrast with the dorsal protrusions on the claspers of the three foregoing species, which lie proximally of the aedeagus. Dorsal margin of clasper bending into almost straight distal margin, at rounded, slightly inwards curved, distodorsal corner. Claspers slightly diverging towards pointed apices. Apical part of clasper directed downwards, with narrow clasper hollow. Lateral side of clasper forming a small and angular, laminiform lateral protrusion, unique for this species. Aedeagus elongate in apical part of its S-curvation (fig. 71), with narrowly elongate lateral lobes and truncate apex. Aedeagus with distinct crest between lateral lobes (fig. 75), crest very low mid-between basal lobes, and strongly concave (seen from behind (fig. 74). Aedeagus pore narrow and sharply incised, truncate at aedeagus apex.

Measurements: Body length: 19.0 mm ; tegmen length: 24.5 mm ; head length: 1.8 mm ; pronotum length: 2.5 mm ; mesonotum length: 4.3 mm ; head width: 4.3 mm ; width of pronotal collar: 5.8 mm .

Distribution (fig. 1), - The only specimen known comes from Lake Kutubu, in central Papua New Guinea, just south of the central mountain ranges.

Etymology. - The species is named in honour of the geologist H.M.S. Hartono.

## Baeturia fortuini sp. n.

(figs. 2, 76-87)
Type material. - Holotype ơ: 'Coll R.I. Sc. N.B. Papua New Guinea Morobe pr.: Anguaia, 1800 m, 21.v. 1988 (st. 050) J. van Stalle, I.G. no.: $27363^{\prime}$ (print, orange label), KBIN. - Paratypes: PAPUA: NEW GUINEA (NE): Bulldog rd., 60 km S Wau, $2070 \mathrm{~m}, 22-31 . v .1964$, J. Sedlacek, 1 §̃, BPBM; Dowalo, W Zenag, $2000 \mathrm{~m}, 4 . \mathrm{ii} .1971$, J.L. Gressitt, 2 if, вPBM; Vagau, Herzog Mts., 4000 ft, 4-17.i.1965, M.E. Bacchus, 1 O', BMNH; U. Watut SW, 1500 m, 3.v.1968, J.L. $_{\text {S. }}$ Gressitt, $1 \delta^{\top}, \mathrm{BPBM}$; same data but $1100-1600 \mathrm{~m}$, 30.iv. 1968, 10 , BPBM.

Other material. - Mt Missim, 2100 m, 15.iii.1968, P.


Figs. 76-83. Baeturia fortuini sp. n. - 76, pygofer in lateral view; 77, male operculum; 78, first sternite; 79, clasper, Mt. Missim; 80, idem, holotype; 81 , aedeagus from aslant; 82 , aedeagus in lateral view; 83 , aedeagus from behind.

Colman, $1 \delta^{*}, 1$ ㅇ, вPBM; Wau, ix,1965, J. Sedlacek, 1 우, BPBM; Wau, Morobe Dist, 1200-1300 m, 14-17.i.1963, J. Sedlacek, 1 ㅇ, вРвм; Wau, Morobe Dist., Mt. Missim, 2000 m, 1.v.1966, O. R. Wilkes, $1 \delta^{\star}$, вРВМ.

The most striking character of this species is the row of bronzed triangular spots along the hind margins of the tegmina. This characer is also found in $B$. loriae, described next. B. fortuini is very similar to $B$. loriae, but can be separated from that species by its larger and triangle-shaped male operculum.

## Description

Body of males light brown or greenish tinged, with dark markings, forming dorsal and lateral bands on abdomen, and with narrow and light middorsal band over whole length of body. Females provided with irregular dark patches, and a narrow and light middorsal band. Male abdomen 1.4-1.7 $x$ as long as head and thorax, of females $0.9-1.0 \times$. Tegmina of males 1.2-1.4× as long as total body length, of females 1.4$1.7 \times$.

Head: Greenish or brown. Vertex and postclypeus with very long setae, as in B. pigrami. Dark spots on vertex lobes, between eyes and ocelli, and on lateral parts of postclypeus. Postclypeus triangularly protruding and $1.4-2.2 \times$ as wide as long. Postclypeus angularly swollen ventrally (as in fig. 4). Head narrower than anterior part of pronotum. Distance between lateral ocelli 1.1-1.4× as long as distance between eye and lateral ocellus.

Thorax: Greenish, with light brown streaks, partly along oblique fissures, and with two pairs of paramedian black spots, at both ends of a broad, greenish tinged, and immaculate middorsal band. Long setae, predominantly on these black areas and on pronotal collar. Colour pattern on mesonotum very variable. Cruciform elevation and small triangle-shaped area in front of elevation light brown or bright green, lateral parts of mesonotum either completely, or only the distal half, dark brown to castaneous.

Legs: Ochraceous, fore femora with dark brown streaks. Proximal spine of fore femur shorter than distance to middle spine.


Fig. 84. Baeruria fortuini sp. n., male tegmen and wing.

Tegmina and wings (fig. 84): Hyaline, though tegmen with a regular row of seven triangular dark spots, close to hind margin, with one spot in each apical area from apical area 2 to 8 . The 8 th apical area of tegmen generally broad and almost square-shaped. Venation red. Tegmina and wings with fairly broad hyaline border along hind margins.

Tymbal organ: Tymbal somewhat incurved ventrally, connecting bar between abdomen and tymbal directed inwards. Six transverse sclerotized ridges spanning the tymbal, from dorsal to ventral tymbal margin. Most proximal ridge narrowing considerably towards ventral margin and only just reaching it. Some specimens with a 7 th ridge, hardly separated from proximal tymbal margin, running from dorsal margin to about half-width of tymbal. Six short intercalary ridges seem to form a midlateral band across rymbal.
Opercula: Male operculum (fig. 77) ochraceous with longitudinal dark streaks on distal part. Basal part of operculum almost oblong, slightly elongate at half-width and slightly tapering towards medial margin. Distal part very large, triangular, and slightly domed, narrowing towards narrowly rounded and almost pointed apex, and reaching to about half-length of abdominal segment 3. Lateral margin of operculum convex near operculum base, concave at halflength and convex again near apex. Operculum strongly curved to body along distomedial margin. Distomedial margin weakly convex. Medial margin short and straight. Opercula close together medially, separated by globularly protruding 1st sternite (fig. 78). Meracanthus very short, reaching to about $1 /$ the operculum length. Female operculum (fig. 85) sickle-
shaped and erect, with weak crest along its distal margin. Basal part of female operculum forming a narrow rim around base of meracanthus.
Abdomen: Male abdomen light brown or greenish. Dark brown markings forming two dorsal bands, separated by a narrow and lighter coloured middorsal line, and lateral bands over abdomen. These bands are interrupted by lighter, often reddish coloured, segmental hind margins. Latero-ventral row of dark spots on segments 3-7 very clear. Ventral side of male abdomen immaculate, with bright red segmental hind margins. First tergite quite long and weakly curved. Anterior margin of 2nd tergite medially straight. Lateral parts of 2nd tergite hardly inflated, forming a weak crest along anterolateral margin, auditory capsules somewhat removed from ventral corner of tymbal. Female abdomen more intensely and irregularly dark stained. Dark lateral bands less conspicuous than in males. Light middorsal line often very clear, broader than in males. Segmental hind margins ochraceous or reddened. Ventral side of female abdomen dark brown stained. Female caudodorsal beak (fig. 86) light ochraceous, long and slender, pointed at apex.

Male genitalia: Pygofer in lateral view (fig. 76) rather stout, with very long and slender caudodorsal beak. Dorsal margin concave, gradually bending into convexly bent beak. Distal margin straight, almost forming right angle with margin of beak. Lateral lobes of pygofer strongly curving inwards towards distal margin, with well-developed, angularly rounded, protuberances. Ventral margin weakly convex, distinctly concave near base of pygofer. Caudodorsal beak in dorsal view (fig. 87) long and slender, point-
ed at apex. Clasper in lateral view (fig. 80) squareshaped as in B. hartonoi, dorsally abruptly broadening, distally of clasper base, so that no clasper heel is formed. Dorsal part of clasper forming a broad and laminiform protrusion, broadly rounded at its proximal corner, and rectangular at its distal corner. Claspers very narrow and directed downwards towards weakly diverging apices, and with a narrow and rounded clasper hollow. Both specimens from Mt. Missim with quite different clasper (fig. 79), more resembling those of $B$. silveri and B. loriae, with broad and laminiform, but less angular, dorsal protrusion and forming a very small, slightly inwards curved, protrusion at its rectangular distal corner. Aedeagus in lateral view (fig. 82) very long and slender, with extremely long and slender lateral lobes, and a very distinct ridge between these lobes (fig. 81). Aedeagus pore slightly larger than in foregoing species. Margin of aedeagus pore distinctly concave at aedeagus apex (fig. 83).

Measurements: Body length $\delta$ : $22.3-24.5 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ $23.4 \mathrm{~mm} \pm 0.9$ ), $:$ tegmen length $\delta$ : $27.0-31.3 \mathrm{~mm}(\overline{\mathrm{x}} 29.8 \mathrm{~mm} \pm 1.5)$, 우: $31.4-36.6 \mathrm{~mm}(\overline{\mathrm{x}} 34.6 \mathrm{~mm} \pm 1.7)$; head length ot: $1.7-2.0 \mathrm{~mm}(\overline{\mathrm{x}} 1.9 \mathrm{~mm})$, , $: 2.2 .1-2.3 \mathrm{~mm}(\overline{\mathrm{x}} 2.2 \mathrm{~mm})$; pronotum length $\delta: 2.5-3.0 \mathrm{~mm}(\overline{\mathrm{x}} 2.8 \mathrm{~mm})$, , ㅇ: $: 3.2$ $3.5 \mathrm{~mm}(\overline{\mathrm{x}} 3.4 \mathrm{~mm}$ ); mesonotum length ot: 4.5-5.4 $\mathrm{mm}(\overline{\mathrm{x}} 4.9 \mathrm{~mm})$, $\quad$ : $: 5.7-6.4 \mathrm{~mm}(\overline{\mathrm{x}} 6.1 \mathrm{~mm})$; head width $\bar{\delta}: 4.6-5.2 \mathrm{~mm}(\bar{x} 5.0 \mathrm{~mm})$, $;: 5: 5-5.9 \mathrm{~mm}(\overline{\mathrm{x}}$ 5.7 mm ); width of pronotal collar $\bar{\delta}: 6.2-7.4 \mathrm{~mm}(\overline{\mathrm{x}}$ 6.8 mm ), $, 7: 7.6-8.2 \mathrm{~mm}(\overline{\mathrm{x}} 7.9 \mathrm{~mm})$.

Distribution (fig. 2). - B. fortuini seems restricted to a small area in Papua New Guinea, just south and west of the Huon Gulf.

Etymology. - The species is named in honour of the geologist A. R. Fortuin.

## Baeturia loriae Distant, 1897 stat. n.

(figs. 2, 88-93)
Baeturia loriae Distant, 1897: 382.
Gymnotympana loriae, Distant 1906: 158; Metcalf 1963: 153.

Identification of types: In his description Distant (1897) indicates that the type series of this species contains at least one male and one female, originating from Moroka and Paumomu river, and collected by Loria. Two males from Moroka could be traced: one with a type label was found in the BMNH, the other, with identification label: 'Baeturia loriae Dist.', was located in the MSNG. Females, or material from Paumomu river, that could possibly belong to the type series, were not found. The specimen from the BMNH is hereby designated lectotype, the other specimen belongs to a different species, possibly Baeturia
bicolorata Distant. The lectotype bears the following labels: 'Type' (round label, red margin); 'loriae Dist' (written); 'New Guinea SE Moroka, 1300 m, Loria, vii-xi.93' (print); 'Distant coll. 1911.383' (print).

Classification. - The reasons for the re-allocation of this species in the genus Baeturia are given in the paragraph on phylogeny of the $B$. loriae group

Only one male of this species is known. B. loriae closely resembles $B$. fortuini in body markings, and shares the triangular dark spots in apical areas of tegmina with that species. B. loriae can be separated from B. fortuini by its flat and broader, almost rectangular, operculum.

## Description

Head and thorax brown, abdomen olive green. Dark, almost black, markings forming regular dorsal and lateral bands on abdomen. A narrow and immaculate middorsal band over whole length of body. Abdomen $1.5 \times$ as long as head and thorax. Tegmen $1.2 \times$ as long as total body length.

Head: Grey-brown, with dark brown spots on vertex lobes and lateral parts of postclypeus. Postclypeus broadly protruding, $1.8 \times$ as wide as long, anterior margin broadly rounded. Postclypeus distinctly swollen and ventrally conically protruding, anterior margin (lateral view) broadly rounded. Head narrower than anterior part of pronotum. Distance between lateral ocelli $1.4 \times$ distance between eye and lateral ocellus.

Thorax: Pronotum ochraceous, with brown streaks in and along oblique fissures and two pairs of dark brown spots at both ends of immaculate and lighter coloured middorsal band. Mesonotum with two large and triangle-shaped dark spots, in front of cruciform elevation, and two brown straeks laterad of elevation.

Tegmina and wings: Hyaline, veins ochraceous. Tegmina, as in B. fortuini, with a regular row of seven triangular dark spots, close to hind margin, one spot in each apical area from area 2 to 8 . The 8th apical area of tegmen short, almost square-shaped. Tegmina and wings with a fairly broad hyaline border along hind margin.

Tymbal organ: Six transverse sclerotized ridges spanning the tymbal, from dorsal to ventral margin. Most proximal ridge narrowing considerably towards ventral tymbal margin and only just reaching it. A 7th ridge, hardly separated from proximal tymbal margin, running from dorsal margin to about half width of tymbal. Six short intercalary ridges seem to form a midlateral band across tymbal.

Operculum (fig. 91): Ochraceous. Basal part of operculum oblong. Distal part very large and angular, flat against body. Lateral margin straight, bending into convex medial margin, at almost rectangular distal corner. Medial margin bending angularly back to base of meracanthus, at medial corner. Meracanthus


Figs. 85-93. - 85-87, Baeturia fortuinisp. n., 85, female operculum; 86, female caudodorsal beak in dorsal view; 87, male caudodorsal beak in dorsal view. - 88-93, Baeturia loriae, 88 , pygofer in lateral view; 89 , clasper; 90 , caudodorsal beak in dorsal view; 91, operculum; 92, aedeagus in lateral view; 93, aedeagus from behind.
very short, reaching to about $1 / 4$ the operculum length. Abdomen: Greenish brown. Dark brown markings forming two dorsal bands, separated by a lighter narrow middorsal line, and lateral bands over abdomen. These bands are interrupted by lighter, often reddish coloured, segmental hind margins. Latero-ventral row of dark spots on segments 3-7 very clear. Ventral side of male abdomen immaculate, with bright red segmental hind margins. Anterior margin of 2nd tergite straight medially. Lateral parts of 2nd tergite hardly inflated, forming a weak crest along anterolateral margin, auditory capsules somewhat removed from ventral corner of tymbal.

Genitalia: Pygofer in lateral view (fig. 88) slender, with long and slender, slightly bent, caudodorsal beak. Dorsal margin of pygofer straight, angularly bending into beak. Distal margin straight and broadly rounded into margin of beak. Lateral lobes of pygofer strongly bent inwards, with bluntly rounded protuberances. Ventral margin weakly convex, distinctly concave near base of pygofer. Margins of caudodorsal beak in dorsal view (fig. 90) converging from base, but parallel in apical part of beak. Caudodorsal beak rounded at apex. Clasper in lateral view (fig. 89) angular, square-shaped, as in the Mt. Missim specimens of B. fortuini. Dorsal margin of clasper very slightly bending upwards, distally of
clasper base, so that no clasper heel is formed. Straight dorsal margin bending almost rectangularly into straight distal margin. Distal margin concavely incurved, just before reaching downwards directed apical part of clasper. Clasper hollow small and narrowly rounded at clasper apex. Aedeagus in lateral view (fig. 92) very long and slender, with extremely long and slender lateral lobes, and a very distinct ridge between lobes. Aedeagus pore oval (fig. 93).

Measurements: Body length: 24.1 mm ; tegmen length: 30.0 mm ; head length: 1.9 mm ; pronotum length: 2.9 mm ; mesonotum length: 5.0 mm ; head width: 5.2 mm ; width of pronotal collar: 6.9 mm .

Distribution (fig. 2). - The only specimen available comes from Moroka on the Papuan Peninsula.

## Baeturia tenuispina Blöte, 1960

(figs. 1, 94-102)
Baeturia tenuispina Blöte, 1960: 72, figs. 22-23. - Duffels \& Van der Laan 1985: 254.

Material examined. - PAPUA: NEW GUINEA (NE): Gewak, Salawaker Range, 1530 m, 6.ix. 1956, E.J. Ford Jr., 1 ठ̃, BPBM; Gurakor, 7.vii. 1965, H. Pyka, 1 ó, 7 ㅇ, Smi; Kokoda, 1200 ft , viii. 1933 , L.E. Cheesman, $1 \delta^{\text {o }}$ holotype Baeturia tenuispina Blöte, $30^{\hat{\circ}}$, BMNH; same data, $10^{\hat{}}$ paratype, RMNH; same


Figs. 94-102. Baeturia tenuispina Blöte. - 94, pygofer in lateral view; 95, pygofer from aslant; 96, male operculum; 97, clasper; 98, male caudodorsal beak in dorsal view; 99, aedeagus from behind; 100 , aedeagus in lateral view; 101, female operculum; 102, female caudodorsal beak in dorsal view.
data but vi-x.1933, 9 ס', 65 우, BMNH; Kokoda, xi.1957, 2 ㅇ, BMNH; Kokoda, 400 m, 15-20.xi.1965, J. and M. Sedlacek, 1õ, BPBM; Kokoda-Pitoki, 400 m, 25.iii.1956, J.L. Gressitt, $1 \delta^{\circ}$, BPBM; Sangeman Village nr Busu R., NE of Lae, 25 m , 30.viii.1957, D. Elmo Hardy, 3ठ, вРBM; Wau, Morobe Dist., 1200 m, 29-30.ix. 1963, J. Sedlacek, $1 \delta^{\top}, 1$ ㅇ, BPBM; PApua: new guinea (SE): Bori nr Sasambota, Popondetta Subdist., 31.x. 1963 , D.K. McAlpine, $1 \delta^{\top}$, AMS; Mt Lamington Dist., Northern Division, 1925, C.T. McNamara, 1 ㅇ, AMS; same data but vii.1927, 2 우; i-ii.1929, $2 \delta^{\star}, 1$ ㅇ, all AMS; Popondetta, $25 \mathrm{~m}, \mathrm{v} .1966$, ShanahanLippert, $1 \delta^{\text {§ }}$, BPBM; same data but vi.1966, 2ठ, BPBM; D'Entrecastaux Islands: GOodenough: Goodenough Id., x.1943, F/D, C. Ralph, 1 ठ $^{\star}$, MVM.

Males of $B$. tenuispina are easily recognized by the long spine-shaped caudodorsal beak and a small fin-ger-shaped protrusion on the clasper heel. This protrusion forms the only indication that the species might be related to the $B$. loriae group.

## Description

Body ochraceous to castaneous brown, covered with brown speckling. Females on average shorter than males, but with equally large head and thorax. Abdomen of males 1.3-1.5 $\times$ as long as head and thorax, of females 1.0-1.2×. Tegmen of males 1.1-1.2× as long as body length, of females 1.3-1.4×.

Head: Ochraceous, dark brown or blackish speckled. Postclypeus angularly protruding, $1.7-2.3 \times$ as
wide as long, anterior margin convex. Postclypeus slightly swollen ventrally, anterior margin (lateral view) weakly convex.

Thorax: Pronotum ochraceous brown speckled medially, between medial pair of oblique fissures. Mesonotum greyish brown, densely brown speckled but without any distinct black spots in front of cruciform elevation.

Tegmina and wings: Hyaline, venation ochraceous or reddish, often quite densely set with short setae. Tegmen with 8 apical areas, the 8 th apical area longer than in most species of the $B$. loriae group. No sexual dimorphism in wings. Tegmen with narrow hyaline border along hind margin, this border distinctly broader in wing.

Tymbal organs: Tymbal not curved inwards towards distal margin; connecting bar between tymbal and abdomen almost parallel to body axis. Seven sclerotized transverse parallel ridges spanning the tymbal from dorsal to ventral margin and an 8th, most proximal ridge nearly reaching ventral margin. Seven short intercalary ridges seem to form a midlateral band across tymbal. Tymbal reddish coloured at dorsal margin and dorsally between ridges.

Opercula: Male operculum (fig. 96) quite large and almost completely covering tymbal cavity in ventral view. Distal part of male operculum broad, angularly rounded, oblong and curved towards body. Lateral
margin short, convexly bending into long weakly convex distal margin. Distomedial and medial margins weakly convex. Medial corners rounded. Meracanthus reaching to about $1 / 2$ operculum length. Female operculum (fig. 101) with very short, sickle-shaped and erect distal part. Medial margin short and straight

Abdomen: Male abdomen light brown or ochraceous, densely brown speckled dorsally; ventral part, and sometimes lateral parts of segments $4-5$, unspeckled. Latero-ventral row of darkened spors generally very distinct on segments $3-4$, but often only vaguely visible on segments 5-7. Segmental hind margins bright red. First tergite quite short. Anterior margin of 2nd tergite medially convex. Lateral parts of 2nd tergite distinctly swollen adjacent to tymbal. Female abdomen greyish brown and densely speckled, lateroventral row of spors less distinct. Segmental hind margins often ochraceous. Ovipositor sheaths just reaching apex of caudodorsal beak. Female caudodorsal beak in dorsal view (fig. 102) very long and slender, triangle-shaped and sharply pointed at apex.
Male genitalia: Pygofer (figs. 94-95) very characteristic, with broadly rounded lateral lobes and long, slender and erect caudodorsal beak. Dorsal margin concave to base, but convexly bent into caudodorsal beak. Distal margin angularly convex, forming a narrowly rounded angle with straight margin of beak. Ventral margin convex. Lateral lobe of pygofer weakly curving inwards towards end of distal margin, forming a weakly swollen elongate and bluntly rounded protuberance. Caudodorsal beak in dorsal view (fig. 98) extremely long and slender, its parallel margins converge at about half-length of beak, towards pointed apex. Claspers conspicuous by slender finger-shaped protuberance on corner of clasper heel (fig. 97). Dorsal crest very stout and strongly outcurving, abruptly ending at distolateral corner of clasper. Apical part of clasper slightly curved down and with small clasper hollow. Aedeagus (fig. 100) very small with narrow lateral lobes, in lateral view weakly concave along its pore. Aedeagus pore oval (fig. 99).

Measurements: Body length of: $18.3-23.3 \mathrm{~mm}(\overline{\mathrm{x}}$ $20.3 \mathrm{~mm} \pm 1.4),: \circ: 16.7-20.0 \mathrm{~mm}(\bar{x} 18.5 \mathrm{~mm} \pm 1.2)$; tegmen length $\delta^{\hat{c}}: 20.0-24.9 \mathrm{~mm}(\overline{\mathrm{x}} 22.7 \mathrm{~mm} \pm 1.2$ ), ?: $23.6-26.5 \mathrm{~mm}(\bar{x} 24.9 \mathrm{~mm} \pm 1.2)$; head length $\delta$ : $1.6-2.0 \mathrm{~mm}(\bar{x} 1.8 \mathrm{~mm})$, 여: $1.6-1.9 \mathrm{~mm}(\bar{x} 1.8 \mathrm{~mm})$; pronotum length $\delta^{*}: 2.3-3.2 \mathrm{~mm}(\bar{x} 2.7 \mathrm{~mm})$, $8: 2.7-$ 3.0 mm ( $\overline{\mathrm{x}} 2.9 \mathrm{~mm}$ ); mesonotum length os: 4.2-5.4 $\mathrm{mm}(\bar{x} 4.6 \mathrm{~mm}), \quad \mp: 4.2-4.9 \mathrm{~mm}(\bar{x} 4.6 \mathrm{~mm})$; head width $\delta^{3}: 4.4-5.2 \mathrm{~mm}(\bar{x} 4.8 \mathrm{~mm})$, $\circ: 4.5-5.0 \mathrm{~mm}(\bar{x}$ 4.8 mm ); width of pronotal collar $\delta$ : $5.5-7.1 \mathrm{~mm}(\overline{\mathrm{x}}$ 6.3 mm ), $9: 6.2-6.9 \mathrm{~mm}(\bar{x} 6.6 \mathrm{~mm})$.

Distribution (fig. 1). - B. tenuispina is distributed in the eastern parts of Papua New Guinea and is also recorded from Goodenough Island of the D'Entrecasteaux islands.

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## References

Blöre, H. C., 1960. The genus Baeturia Stål as represented in New Guinea (Homoptera, Cicadidae). - Zoologische Mededelingen 37: 61-80.
Boer, A. J. de, 1982. The taxonomy and biogeography of the nasuta group of the genus Baeturia Stàl, 1866 (Homoptera, Tibicinidae). - Beaufortia 32: 57-78.
Boer, A. J. de, 1986. The taxonomy and biogeography of the conviva group of the genus Baeturia Stå, 1866 (Homoptera, Tibicinidae). - Beaufortia 36: 167-182.
Boer, A. J. de, 1989. The taxonomy and biogeography of the bloetei group of the genus Baeturia Stàl, 1866 (Homoptera, Tibicinidae). - Beaufortia 39: 1-43.
Boer, A. J. de, 1990. Aedeastria, a new cicada genus from New Guinea, its phylogeny and biogeography (Homoptera, Tibicinidae), preceded by a discussion on the taxonomy of New Guinean Tibicinidae. - Beaufortia 40: 63-72.
Boer, A. J. de, 1991. Scottotympana, a new cicad genus from New Guinea, with the description of three new species, their taxonomy and biogeography (Homoptera, Tibicinidae). - Beaufortia 42: 1-11.
Boer, A. J. de, 1992. The taxonomy and biogeography of the viridis group of the genus Baeturia Stål, 1866 (Homoptera, Tibicinidae). - Bijdragen tot de Dierkunde 61 (3): 163-183.
Distant, W. L., 1897. Viaggio di Lamberto Loria nella Papuasia orientale, xvii. Additions to our knowledge of the Cicadidae of New Guinea. - Annali del Museo civico di Storia naturale Giacomo Doria Genova (2) 17: 378-383.
Distant, W. L., 1906. A synonymic catalogue of Homoptera. Part 1. Cicadidae 1906: 1-207 (Trustees Indian Museum, London).
Duffels, J. P. \&\& P. A. van der Laan, 1985. Catalogue of the Cicadoidea (Homoptera, Auchenorhyncha) 1956-1980. - Series Entomologica 34: i-xvi, 1-414.

Metcalf, Z. P., 1963. General catalogue of the Homoptera, VIII. Part 2. Tibicinidae: i-iv, 1-492. North Carolina State College, Raleigh, N.C.

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# DIE PUPPEN DER SPANNER MITTELEUROPAS <br> (LEPIDOPTERA, GEOMETRIDAE): CHARAKTERISTIK <br> UND BESTIMMUNGSTABELLE DER GATTUNGEN 


#### Abstract

Patočka, J., 1994. Die Puppen der Spanner Mitteleuropas (Lepidoptera, Geometridae): Charakteristik und Bestimmungstabelle der Gattungen. - Tiijdschrift voor Entomologie, 137: 27-56, figs. 1-227. [ISSN 0040-7496]. Published 15 July 1994. Characteristics of the pupae of the family Geometridae (Lepidoptera) and an identification key for its 161 Central European genera are given. The present system of this family is discussed from the view of pupal characters. The species examined are listed in an appendix. Institut für Waldökologie der SAW, Śtúrova 2, 96053 Zvolen, Slovakia. Key words. - Geometridae; Pupae; Taxonomy; Key for genera.


Die Familie der Spanner, Geometridae (Lepidoptera) ist eine der größten und wichtigsten Schmetterlingsfamilien. Viele ihrer Vertreter sind ernste Schädlinge in Land- und Forstwirtschaft, einige sind sogar Vorratsschädlinge. Die Mehrzahl der Arten ist wegen ihrer Gebundenheit an spezifische Standorte auch vom ökologischen Gesichtspunkt wichtig und kommt als Indikatoren der bedrohten Umwelt in Betracht. Die Kenntnis ihrer jüngeren Entwicklugsstadien ist für viele Fragen unbedingt notwendig; z. B. für das Studium der Schädlinge, der Prädatoren und Parasitoide, der Bioindikatoren, Bodenfauna, verschiedener Synusien usw. Die bisherige Kenntnis der Puppen der mitteleuropäischen Geometridae ist lückenhaft. Speziell die Puppen dieser Familie behandelt Khotko (1977) und erwähnt 25 Gattungen. In der vorliegenden Arbeit werden zum Vergleich 161 mitteleuropäische Gattungen, also mehr als $90 \%$ ihres Bestandes berücksichtigt. An Khotko's Arbeit knüpfen die Veröffentlichungen des Verfassers an, die zahlreiche weitere Arten der Unterfamilie Larentiinae (Patočka 1980-1994) und mehrere Gruppen der Ennominae (Patočka 1978, 1985, 1986 a, b, 1992, 1993) behandeln. Auf diese Weise ist zwar die Mehrzahl der Arten dieser beiden Unterfamilien eingehend bearbeitet, es fehlt jedoch eine zusammenfassende Bestimmungstabelle der Gattungen. Diese Lücke soll die vorliegende Arbeit ausfüllen. Eingehendere Angaben über viele Geometridenpuppen finden sich bei Ljungdahl (1919) und Nordström et al. (1941), über die Arten an Obstbäumen Speyer (1958), an Tannen Patočka et al. (1960), an Eichen Patočka (1980) und über die Puppen in der Bodenstreu in Kiefernwäldern Koehler (1937). Einzelbeschreibungen von Geometridenpuppen sind in weiteren Litera-
turquellen verstreut. Die Handbücher über die Schmetterlinge Mitteleuropas, wie Forster \& Wohlfahrt (1981) geben nur kurze und für eine Bestimmung nicht ausreichende Puppenbeschreibungen. Die Puppen der kanadischen Geometridae wurden im Rahmen einer Monographie dieser Familie von McGuffin (1967-1981) bearbeitet.

Das hier benützte System und die Nomenklatur folgen grundsätzlich Leraut (1980) und Gustafsson (1987), die morphologische Terminologie richtet sich nach Mosher (1916) und McGuffin (19671981). Das untersuchte Material (die hier berücksichtigten Gattungen und Arten sind im Anhang aufgelistet) stammt großenteils aus der Sammlung des Verfassers, zu welcher viele Fachkollegen durch Überlassung von Puppen und Puppenexuvien beigetragen haben, wofür ihnen auch an dieser Stelle herzlich gedankt sei. Viele Arten konnte sich der Verfasser von Museen ausleihen, insbesondere von dem Museum für Naturkunde der Humboldt-Universität in Berlin (Dr. W. Mey), von dem Naturhistorischen Museum Wien (Dr. M. Lödl), von dem Zoologischen Museum Kopenhagen (O. Karsholt), von der Zoologischen Staatssammlung München (Dr. W. Dierl) und aus der Entomologischen Abteilung des Nationalmuseums in Prag (Dr. F. Krampl). Den genannten Institutionen und Herren dankt der Verfasser verbindlichst, ebenso wie auch der Alexander von Humboldt-Stiftung Bonn für die Erteilung eines Stipendiums, welches ihm das Studium des Materials in München, sowie der Hanskarl-Göttling-Stiftung der Bayerischen forstlichen Versuchs- und Forschungsanstalt, welche einen Aufenthalt in Berlin ermöglichten.

Die Puppen und ihre Exuvien wurden mit Hilfe eines Stereomikroskopes und z. T. eines Abbeschen Zeichenapparates untersucht und gezeichnet.

## Charakteristik der familie GeometridaE

Eine Übersicht der Puppenmorphologie dieser Familie geben die Abb. 1-6. Die Puppen der mitteleuropäischen Geometridae sind kleine bis mittelgroße (erwa $5-25 \mathrm{~mm}$ lang) Pupae obrectae. Sie sind schlank (Abb. 86) bis gedrungen (Abb. 94), meistens in der Mitte am breitesten, vorne abgerundet und hinten mehr zugespizt (Abb. 1, 2, 88, 94). Manche Formen (z.B. Cyclophora, Abb. 130) sind vorne durch einen Höcker an der Basis der Vorderflügel breiter und sehen in Ventralansicht eher keilartig aus. Die Mehrzahl der Puppen ist einfarbig gelb-, rot-, oder schwarzbraun bis fast schwarz gefärbt und $\pm$ glänzend. Es handelt sich hierbei um Arten die sich im Boden oder in der Bodenstreu, in einem Erdkokon bzw. einer Erdhöhle verpuppen. Die übrigen Puppen sind grün, gelblich oder sandfarben, $\pm$ glanzlos und oft mit dunklen Zeichnungen versehen. Das sind Arten. die sich oberirdisch bzw. am Boden in einem leichren Gespinst, oder nur mit Einzelfäden befestigt verpuppen. Eine Gattung (Abraxas) ist glänzend schwarz mit satt gelben Querstreifen am Abdomen, die sich ebenfalls oberirdisch verpuppt.

Die Oberfläche der Puppen ist fein bis grob skulpruriert. Meist ist der Kopf und Thorax $\pm$ gerunzelt oder gefurcht und das Abdomen (manchmal auch der Thorax, insbesondere das Metanotum) mit kleinen punktförmigen Grübchen, den sog. Punktgrübchen (Abb. 210) besetzt. Selten entspringen von diesen Microbörstchen (Apeira, Abb. 74). Bei wenigen Puppen (z. B. bei der Gattung Geometra, Abb. 78) kommt eine Stachelskulptur vor. Die Körperborsten sind kaum sichtbar bis relativ groß und stark (Abb. 31). Der Vertex wächst entweder mit der Frons ohne cine Epicranialnaht zusammen (Vertreter der Unterfamilie Ennominae und einige andere, Abb. 4), oder er ist durch eine deutliche Epicranialnaht abgetrennt (Mehrzahl der übrigen Unterfamilien, Abb. 3). Beim Schlüpfen löst sich der Vertex bei dieser Gruppe von der Frons ab. Die Frons ist in wenigen Fällen mit einem Paar Höcker (Abb. 8, 9, 86, 125) versehen, oder kegelförmig vorgezogen (Abb. 43), ansonst meist abgerundet. Zwei Paare von Frontal(Abb. 4) und ein Paar von Clypealborsten (Abb. 116) sind meist deutlich und groß ausgebildet. Von der Frontoclypealnaht sind nur Teile nahe der Antennen deutlich erkennbar. Die Komplexaugen sind groß und bestehen aus einem $\pm$ glatten, ventralen Streifen und einem oft mehr rauhen Dorsalteil (Abb. 95, 96). Die Mandibeln sind meist kaum differenziert (Abb. 112, 116), bei Theria jedoch stark hervortretend
(Abb. 22). Das Labrum erscheint meist halbkreis(Abb. 46), schaufel- (Abb. 117) oder trapezförmig (Abb. 47), gemeinsam mit dem schwach abgegrenzten Postclypeus oft fünfeckig (Abb. 116). Das Labium ist entweder ganz verdeckt (Abb. 87) oder klein (Abb. 209) bis groß (Abb. 101), als eine dreioder fünfeckige Platte sichtbar. Die Maxillar- und Labialpalpen sind stets verdeckt. Die Genae grenzen an die Vorderbeine; diese Grenze immer kürzer als die Grenze zwischen Auge und Vorderbein. Die Grenzen zwischen Vorderbeine und Antennen kurz oder lang (Abb. 19, 21). Bei den sonst ähnlichen Puppen der Herminiinae, Noctuidae, grenzen die Vorderbeine nicht an die Antennen, weil die Mittelbeine die Augen berühren. Die Vorder- und Mittelbeine sind meist relativ lang, die Vorderbeine zuweilen (Abb. 16), die Mittelbeine meistens ungefähr so lang wie die Antennen und Vorderflügel. Die Enden der Hinterbeine sind oft sichtbar. Die Vorderschenkel sind sichtbar, groß (Abb. 1, 125) oder klein, schmal (Abb. 94, 77, 165), oft aber ganz verdeckt (Abb. 16, 94a). Die Proboscis ist meist so lang oder wenig kürzer als die Vorderflügel, selten stark verkürzt und deutlich kürzer als die Mittelbeine, bzw. auch als die Vorderbeine, (Abb. 16, 98, 99). In wenigen Fällen überragt die Proboscis (Abb. 9, 12, $94 a, 123$ ) und noch seltener überragen auch die Mittelbeine und Antennen (Abb. 13) die Spitzen der Vorderflügel und den Kaudalrand des 4. Abdominalsegmentes. Bei Rhodometra (Abb. 123) verschiebt die vergrößerte Proboscis das Labrum bis nahe am Frontalende des Kopfes, was oft auch bei Sphingidae mit langer Proboscis vorkommt. Die Antennen sind meist ungefähr so lang wie die Vorderflügel (Abb. 1). Die Vorderflügel überragen den Kaudalrand des 4. Abdominalsegmentes nicht, sind voneinander getrennt (Ausnahme: Apeira, Abb. 76) und zeichnen sich zuweilen durch hervorstehende (Abb. 94) oder dunkel gestreifte bzw. helle Adern aus. Selten liegt an ihren Wurzeln je ein Höcker (Abb. 123, 125, 130). Die Hinterflügel säumen die Vorderflügel meist zu $1 / 4$ bis $2 / 3$ des 4 . Abdominalsegmentes, nicht aber bis zu deren Spitzen, wie es bei der Familie Drepanidae, die den Geometridae sonst ähnlich ist, vorkommt. Eine Ausnahme ist die Gattung Archiearis.

Die thorakalen Spiracula sind klein und spaltförmig. Oft bildet sich kaudal von ihnen am Mesonotum, ein auffälliger, tomentöser (Abb. 31, 91), rauh skulpturierter (Abb. 54) oder 'ohrartig' hervortretender, kantiger Höcker (Abb. 33, 35). Das Metanotum besitzt frontal in der Mitte meist einen mitteltiefen und abgerundeten Ausschnitt. Die Frontolaterale Lappen des Metanotums sind meist abgerundet (Abb. 102, 103), zuweilen spitz oder winkelig (Abb. 153), insbesondere bei den Vertretern der Unterfamilie Sterrhinae. Die Vorderecken des 1. Abdominalseg-
mentes sind meist abgerundet, Ihre Seiten $\pm$ parallel. Die Spiracula sind am 2.-3. Abdominalsegment sichtbar oder $\pm$ verdeckt. Die abdominalen Spiracula sind klein bis mittelgroß, meist schmal elliptisch, oft mit breiten elliptischen Höfen und nicht selten $\pm$ erhaben. Die 5.-7. Abdominalsegmente sind beweglich. An der Basis des 5. Segmentes frontal des Spiraculums finden sich oft Leisten, Gruben und verdichtete Punktgrübchen (Abb. 34, 38, 40, 45, 49-55, 138, 140-142), die gute Unterscheidungsmerkmale bieten. An der intakten Puppe sind sie jedoch durch den Kaudalrand des 4. Abdominalsegmentes oft verdeckt; dieser sollte deshalb, um eine Untersuchung zu ermöglichen, mit einer Nadel beseitigt werden. An der Puppenexuvie sind diese Gebilde auch ohne diesen Eingriff meist gut sichtbar. Bei der Gattung Rheumaptera gibt es Querleisten und Rinnen an der Basis der Dorsalseite des 5. Abdominalsegmentes (Abb. 139). Zwischen dem 9. und 10. Segment besitzt die Mehrzahl der Spannerpuppen einen $\pm$ tiefen, rinnenartigen Quereinschnitt, die sog. Dorsalrinne, welche an ihrer Kaudalseite meist einen oder mehrere Ausläufer aufweist, deren Ränder oft verdunkelt und $\pm$ tomentös sind. Seltener bildet die Dorsalrinne auch einen oder einige Ausläufer frontalwärts (Frontalausläufer, Abb. 6 ). An den Seiten entspringt von dieser Rinne oft noch ein dorso-kaudalwärts gerichteter Lateraleinschnitt (Abb. 5).

Dorsalseite des 10. Abdominalsegmentes ist verschieden lang (Abb. 70, 71), seine Ventralseite oft $\pm$ breit mit gewölbten Seiten, abgeflacht, und der manchmal erhabene frontolaterale Teil durch flache Furchen abgegrenzt. Die Analnaht ist meist lang, ihre oft erhabenen Seiten wallartig, manchmal auch mit Längs-, seltener Querrunzelung. Das sog. Analfeld groß und auffällig. Die Genitalnaht und ihr Genitalfeld, besonders bei den Männchen sind oft groß. Der Absturz zur Basis des Kremasters verläuft steil bis schräg (Abb. 137, 182). Der Kaudalausläufer des 10. Segmentes, der Kremaster, welcher bei der Puppe eine Haft- und Stützfunktion besitzt (Patočka 1978), ist bei den Puppen der Geometridae immer vorhanden und $\pm$ groß, seltener klein (Abb. 17) und mehr oder weniger scharf von dem 10. Abdominalsegment abgegrenzt bzw. abgesetzt. In der Dorsalsicht ist er oft haufen- (Abb. 119) oder zungenförmig (Abb. 80), dreieckig (Abb. 178), seltener trapez- (Abb. 18, 164) oder ankerförmig (Abb. 7). Ventral an seiner Basis gibt es oft ein Paar Basalhöcker, dazwischen eine Vertiefung bzw. Mulde. Bei der Unterfamilie Geometrinae ist der Kremaster ventral charakteristisch radial gefurcht (Abb. 75, 79), ansonst ist er $\pm$ glatt oder verschiedenartig skulpturiert (gerunzelt, gefurcht, gerippt; Abb. 120, 122). Der Kremaster trägt meist 4, seltener 1-3 oder 5-7 (bei Apeira eine Anzahl) Paare von Borsten, die oft am Ende hakenartig gekrümmt
sind. Die Bezeichnung von 4 Paaren von Primärborsten am Kremaster (nach McGuffin 1967-1981) zeigen die Abb. 5-6. Bei vielen Geomerriden-Puppen wachsen die starken Borsten $\mathrm{D}_{2}$ zu einem Stiel zusammen und nur ihre Spitzen bleiben als eine Gabelspitze frei (Abb. 32), die manchmal auch vermehrt sind (Abb. 28). Während bei vielen Vertretern der Unterfamilie Larentiinae die übrigen Borsten noch erhalten bleiben (Abb. 177, 178) verschwinden sie bei zahlreichen Ennominae vollständig (Abb. 71). Bei Bupalus bleibt nur eine einfache, stumpfe Spitze übrig (Abb. 42).

Die Puppen der Spanner findet man z. T. an der Nahrungspflanze, durch die Häkchen am Kremaster und einen Gürtelfaden befestigt (z.B. bei Cyclophora) oder in einem lockeren Gespinst (Abraxas, Ennomos, bei den Comibaenini in dem Sack der Raupe) direkt an der Nahrungspflanze oder auch am Boden. Die Mehrzahl der Puppen ruht jedoch in einem $\pm$ festen Erdkokon oder einer Erdhöhle in der Bodenstreu oder im Boden selbst. Die Puppen variieren in der Größe insbesondere nach Geschlecht, Generation oder relativer Abundanz. Die Raupen leben an verschiedenen Gruppen von Pflanzen, sogar an Sporenpflanzen bzw. an Detritus. Es gibt ebenso polyphage, wie relativ streng oligophage, an spezielle Pflanzengattungen gebundene Arten (z. B. Clematis, Actaea, Berberis, Rhamnus, Hypericum, Veratrum). Manche Gruppen (Eupitheciini, Perizomini) bevorzugen Blüten und Früchte. Viele Arten leben an Laubhölzern, an denen sie oft auch schädlich werden, aber auch an Nadelhölzern, und eine zahlreiche Gruppe von Larentiinae lebt an Galium.

Die Spanner bewohnen verschiedenartige Biotope, sehr viele Wälder und steppenartige, mehrere auch ruderale Standorte, Sümpfe bzw. Torfmoore, Felsen und Alpenwiesen im Hochgebirge. Viele Arten sind für bestimmte Biotope recht charakteristisch.

## Bestimmungstabelle der Gattungen

Einige Gattungen, wie Orthostixis Hübner, 1822, Costaconvexa Agenjo, 1949, Coenocalpe Hübner, 1825, Bichroma Gumppenberg, 1887, Eilicrinia Hübner, 1825, Mannia Prout, 1915, konnten infolge Materialmangels nicht berücksichtigt werden.

1 Kaudalhälfte des Kremasters in Dorsalsicht einfach zugespitzt (selten mit zwei Börstchen am Ende) (Abb. 42) Bupalus Leach

- Kaudalhälfte des Kremasters in Dorsalsicht mit 2-4 Dornen oder Zähnen, jedoch ohne Börstchen oder Häkchen (Abb. 7, 14, 20). Die Dornen oft gestielt (Abb. 32, 52), zuweilen auch verzweigt (Abb. 28).

2




- Kaudalhälfte des Kremasters mit 2-8 (seltènêr mehr) Häkchen oder Borsten (Abb. 6, 72, 80, 89, 136). Die Endborsten ( $\mathrm{D}_{2}$ ) zuweilen $\pm$ dornartig umgebildet (Abb. 89, 143, 218).
(1) Frons mit einem Paar kräfriger Höcker (Abb 8, 9) 3
- Frons ohne ein Höckerpaar, selten kegelförmig vorgezogen (Abb. 43), sonst abgerundet (Abb. 16) 4
3 (2) Proboscis überragt deutlich den Kaudalrand des 4. Abdominalsegmentes (Abb. 9)

Siona Duponchel

- Proboscis überragt den Kaudalrand des 4. Abdominalsegmentes nicht (vgl. Abb. 1) ...... ............................. Aethalura McDunnough
4 (2) Einzelne Abdominalsegmente dorsal mit großen, elliptischen, dunkel umrandeten Gruben (Abb. 11) .5
- Abdominalsegmente ohne solche Gruben (vgl. Abb. 2) .6
5 (4) Am 5.-7. Abdominalsegment dorsal fehlen die Gruben ...................... Lithostege Hübner
- Auch am 5.-7. Abdominalsegment gibt es dunkel umrandete Gruben (Abb. 11)

Chesias Treitschke
6 (4) Proboscis überragt den Kaudalrand des 4. Abdominalsegmentes (Abb. 12, 13)........... 7

- Proboscis überragt nicht den Kaudalrand des 4. Abdominalsegmentes (Abb. 1) $\qquad$
7 (6) Sowohl die Antennen wie auch Mittelbeine überragen den Kaudalrand des 4. Abdominalsegmentes 8
- Nur die Antennen, nicht jedoch die Mittelbeine überragen den Kaudalrand des 4. Abdominalsegmentes. $\qquad$ Odezia Boisduval
- Weder Antennen noch Mittelbeine überragen den Kaudalrand des 4. Abdominalsegmentes (Abb. 12) ........... Schistostege Hübner
8 (7) Kremaster länger als basal breit (Abb. 15) ..... .. Carsia Hübner
Kremaster kürzer als basal breit (Abb. 13, 14) Aplocera Stephens

9 (6) Kremaster ankerartig, stumpf abgerundet, an seinen Seiten je ein größerer und oft auch kleinerer Zahn (Abb. 7). Puppe schlank. Archiearis Hübner Kremaster am Ende mit gabelartigen, schräg kaudalwärts gerichteten Dornen (Abb. 25, 32, 52), diese zuweilen weiter verzweigt (Abb. 28). Wenn die Dornen $\pm$ gegeneinander stehen (Abb. 17, 20), nur in einem Paar und die Puppe ist gedrungen 10
10 (9) Weder Labium, noch Vorderschenkel sichtbar (Abb. 16, 39) 11

- Labium, Vorderschenkel, oder beides sichtbar (vgl. Abb. 1) ....................................... 23
11 (10) 8. Abdominalsegment dorsal mit einem Paar spitzer Höcker.................. Nychiodes Lederer
- 8. Abdominalsegment dorsal ohne Höcker ...
.12
12 (11) Kremaster mit zwei Paaren schlanker, wenig divergierender Dornen, außerdem in seinem Basalteil noch ein Paar Höcker (Abb. 23) .....
. Artiora Meyrick
Kremaster mit 1 Paar Dornen, die zuweilen noch verzweigt sein können, außerdem höchstens mit kurzen Zähnen an den Seiten (Abb. $17,20,25,26,28)$ 13
13 (12) Kremaster klein, kurz, seine Dornen stark divergierend (Abb. 17, 18, 20). 14
- Kremaster groß, am Ende mit langen Dornen oder mit einer gestielten Gabel (Abb. 25, 27, 30) 16
14 (13) Proboscis kürzer als die Vorderbeine (Abb. 16). $\qquad$ Operophtera Hübner
- $\quad$ Proboscis länger als die Vorderbeine (Abb. 1) .15
15 (14) Grenzen zwischen Vorderbeine und Antennen etwa $2 \times$ so lang wie die Grenzen zwischen Vorderbeine und Augen (Abb. 21). Mandibeln erhaben .............. Theria Hübner Grenzen zwischen Vorderbeine und Antennen höchstens wenig länger (oft kürzer) als die Grenze mit den Augen (Abb. 19). Mandibeln nicht erhaben $\qquad$ Alsophila Hübner

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16 (13) 9. Abdominalsegment mit einer Querreihe von Dornen (Abb. 24).. Eurrhanthis Hübner

- 9. Abdominalsegment ohne Dornen (Abb. 32)

17
17 (16) 10. Abdominalsegment mit je einem spitzen lateralen Fortsatz (manchmal ein weiterer an der Mitte des Kremasters, Abb. 25)

Lycia Hübner

- 10. Abdominalsegment ohne, Kremaster oft mit einem Paar von laterale Fortsätze (Abb. 28,32 ).

18
18 (17) Dorsalrinne vorhanden, Kaudalausläufer $\pm$ tomentös, Lateraleinschnitt deutlich (Abb. 26, 27, 28) 19

- Dorsalrinne und Lateraleinschnitt nicht vorhanden (Abb. 32) 21
19 (18) Erhebung kaudal des thorakalen Spiraculums flach, länglich elliptisch, stark tomentös (Abb. 31). Puppe 12-15 mm lang.

Erannis Hübner

- Erhebung kaudal des thorakalen Spiraculums 'ohrartig', kantig (Abb. 33, 35). Puppe über 17 mm lang

20
20 (19) Leiste frontal des Spiraculums am 5. Abdominalsegment doppelt (Abb. 34). Erhebung kaudal des thorakalen Spiraculums steiler frontal- als kaudalwärts (Abb. 33). Puppe 17 mm lang Apocheima Hübner

- Leiste frontal des Spiraculums am 5. Abdominalsegment einfach (Abb. 36). Erhebung kaudal des thorakalen Spiraculums weniger steil frontal- als kaudalwärts. (Abb. 35) Puppe $20-26 \mathrm{~mm}$ lang.

Biston Leach
21 (18) Puppe 17-21 mm lang.. Phigalia Duponchel

- Puppe $10-15 \mathrm{~mm}$ lang............................ 22

22 (21) Kremaster mit deutlichen lateralen Fortsätzen (Abb. 41). Vertiefung frontal des Spiraculums am 5. Abdominalsegment mit zahlreichen Punktgrübchen frontal der Querleiste (Abb. 40)....... Selidosema Hübner

- Kremaster ohne, bzw. nur mit angedeuteten lateralen Fortsätzen (Abb. 32). Vertiefung am 5. Abdominalsegment frontal des Spiraculums mit Querleisten, ohne Punktgrübchen (Abb. 37, 38) $\qquad$ Agriopis Hübner
23 (10) Dorsalrinne mit 1-2 großen Kaudalausläu-
fern (Abb. 26, 27, 29) 24
- Dorsalrinne mit mehreren Kaudalausläufern, oder nicht vorhanden (Abb. 56, 62, 70) .. 28
24 (23) Vorderschenkel nicht sichtbar (vgl. Abb. 82) 25
Vorderschenkel sichtbar (vgl. Abb. 1) ...... 26
25 (24) Dorsalrinne mit nur einem großen Kaudalausläufer. Kremaster ohne laterale Fortsätze (Abb. 26) $\qquad$ Chondrosoma Anker
- Dorsalrinne mit zwei großen Kaudalausläufern (Abb. 27)................. Lignyoptera Lederer
26 (24) Frons breit kegelförmig vorgezogen (Abb. 43). Dorsalrinne auch an der Frontalseite ausgebuchtet (Abb. 44) ....... Pelurga Hübner
- Frons nicht vorgezogen. Dorsalrinne nur an der Kaudalseite mit einem großen, spitzen Vorsprung (Abb. 29) ............................... 27
27 (26) Puppe mehr gedrungen, schwärzlich, Exuvie schwarzbraun. Puppenlänge $6-8 \mathrm{~mm}$.

Phibalapteryx Stephens

- Puppe schlanker, bräunlich, Exuvie gelbbraun. Puppenlänge $9-10 \mathrm{~mm}$.

Cataclysme Hübner
28 (23) Dornen am Kremaster voneinander entfernt entspringend (Abb. 30) 29

- Dornen am Kremaster gabelartig, gestielt (Abb. 28, 67, 202, 206) 30
29 (28) Kremaster an der Dorsalseite fein skulpturiert. Puppe nur $6-8 \mathrm{~mm}$ lang. $\qquad$ Pygmaena Boisduval
- Kremaster an der Dorsalseite grob skulpturiert. Puppe über 10 mm lang.

Gnophos Treitschke s. lat.
30 (28) Proboscis wesentlich kürzer als die Mittelbeine. Labium sehr klein. Puppe $7-9 \mathrm{~mm}$ lang.. Tephronia Hübner

- Proboscis nicht oder wenig kürzer als die Mittelbeine (im Zweifelsfall Labium Mittelgroß) 31
31 (30) Dornen am Kremaster $\pm$ länger als ihr Stiel (Abb. 102, 206). Puppen nur 6-8 lang... 152 Dornen am Kremaster $\pm$ kürzer als ihr Stiel (Abb. 52, 62, 67). Meist größere Puppen .....
.............................................................. 32

32 (30) Dorsalrinne entwickelt (Abb. 56, 58, 61-63)
33

Abb. 22-57. - 22, 39, Theria rupicapraria; 23, Artiora evonymaria; 24, Eurranthis plumistarias 25, Lycia zonarias 26, Chondrosoma fiduciaria; 27, Lignyoptera thaumastaria; 28, 31, Erannis defoliara, 29, Phibalapteryx virgata; 30, Gnophos obfussatus; 32, 38, Agriopis aurantiaria; 33, 34, Apocheima bispidaria; 35, 36, Biston stratariat, 37, Agriopis bajariat 40, 41, Selidosema plumaria, 42, Bupalus piniarius, 43, 44, Pelurga comitata, 45, 46, Ectropis crepuscularia, 47-49, Calospilos sylvata, 50, Cleora cinctaria; 51, Fagivorina arenaria, 52-54, Serraca punctinalis, 55, Ematurga atomaria, 56, 57, Chiasmia clatbrata 22, Labrum, Mandibulae; 23-30, 32, 41, 42, 44, 52, 56, Abdominalende in Dorsalsicht; 31, 33, 35, Erhebung kaudal des thorakalen Spiraculums; 34, 36, 38, Basis des 5. Abdominalsegmentes, Lateralsicht; 29, Kopf und Beine in Ventralsichr; 40, 45, 49-51, 53, 57, Basis des 5. Abdominalsegmentes, Lateralsicht; 43, Kopf in Ventralsicht; 46, 47, Labrum, Labium; 48, Abdominalende in Lateralsicht; 54, Erhebung kaudal des thorakalen Spiraculums.


- Dorsalrinne nicht entwickelt (Abb. 70, 71) 47
33 (32) Kaudal des thorakalen Spiraculums je eine relativ große, breite, sehr rauhe, erhöhte Fläche (Abb. 54). An der Basis des 5. Abdominalsegmentes lateral zwischen zwei dunklen Querleisten wenigstens 12 Zellen in einer Reihe (Abb. 53, 55) 34
- Kaudal des thorakalen Spiraculums eine kleine, längliche oder eine unauffällige Erhebung (vgl. Abb. 106, 111). Wenn diese jedoch größer und rauher ist (Abb. 227) und auch sonst, gibt es an der Basis des 5. Abdominalsegmentes lateral anders aussehende Bildungen (Abb. 51, 57, 65, 68, 69) oder nur etwa 5 Zellen (Abb. 50) 36
34 (33) Mesonotum ohne Punktgrübchen Hypomecis Hübner
- Mesonotum mit Punktgrübchen.............. 35

35 (34) Spiraculum am 5. Abdominalsegment liegt in der Nähe der 10.-15. Zelle (von der Ventralseite gezählt, Abb. 53). Puppe meist über 14 mm lang, eher glänzend ... Serraca Moore

- Spiraculum am 5. Abdominalsegment liegt in der Nähe der 2.-6. Zelle (Abb. 55). Puppe meist matter, unter 13 mm lang

Ematurga Lederer
36 (33) Abdomen ohne Punktgrübchen (vgl. Abb. 92) Lomaspilis Hübner

- Abdomen mit Punktgrübchen (vgl. Abb. 51) 37
37 (36) Am 5. Abdominalsegment frontal des Spiraculum eine länglich ovale, von einer dunklen Leiste umgebene Fläche (Abb. 45) 38
- Am 5. Abdominalsegment vor dem Spiraculum andere Bildungen: Zellen (Abb. 50), Leisten (Abb. 65, 68, 69) oder nur Punktgrübchen (Abb. 57) bzw. eine glatte Fläche (Abb. 51) 39
38 (37) Labium ganz klein (Abb. 46) bzw. fehlend. Vorderschenkel gut sichtbar (vgl. Abb. 1). Lateraleinschnitt undeutlich

Ectropis Hübner

- Labium (Abb. 47) und Lateraleinschnitt (Abb. 48) deutlich. Vorderschenkel verdeckt (vgl. Abb. 39) $\qquad$ Calospilos Hübner
39 (37) Frontal des Spiraculums am 5. Abdominal-
segment nur eine glatte Fläche, weder mit Leisten, noch Punktgrübchen (Abb. 51) Fagivorina Wehrli
- Frontal des Spiraculums am 5. Abdominalsegment eine dunkle Leiste und vor ihr etwa fünf Zellen (Abb. 50)............ Ascotis Hübner Frontal des Spiraculums am 5. Abdominalsegment Leisten und Punktgrübchen (Abb. 65, 69), oder nur Leisten (Abb. 68) oder Punktgrübchen (Abb. 57) 40
40 (39) 9. Abdominalsegment mit Punktgrübchen (Abb. 56, 61) 41
- 9. Abdominalsegment ohne Punktgrübchen (Abb. 62, 63).......................................... 43
41 (40) Frontal des Spiraculums am 5. Abdominalsegment keine glatte Fläche, nur Punktgrübchen (Abb. 57) $\qquad$ Chiasmia Hübner
- Frontal bzw. frontodorsal des Spiraculums am 5. Abdominalsegment eine glatte Fläche und davor liegen Punktgrübchen bzw. eine Querleiste (Abb. 59, 60) 42
42 (41) An der Basis des 5. Abdominalsegmentes lateral nur Punktgrübchen ohne eine dunkle Querleiste (Abb. 59). Stiel am Kremaster relativ kurz (Abb. 58)

Parectropis Sato

- An der Basis des 5. Abdominalsegmentes lateral eine dunkle Querleiste mit einer Reihe vòn dicht gedrängten Punktgrübchen (Abb. 60). Stiel des Kremasters lang (Abb. 61) .......

Tephrina Gueneé
43 (40) Dorsalrinne ohne Kaudalausläufer $\qquad$ Euconista Lederer Dorsalrinne mit Kaudalausläufern ........... 44
44 (43) Frontal des Spiraculums am 5. Abdominalsegment liegen nur Punktgrübchen (Abb. 57). 45

- Frontal des Spiraculum am 5. Abdominalsegment liegt eine glatte Fläche, dann Punktgrübchen (Abb. 59) 46 Frontal des Spiraculums am 5. Abdominalsegment liegt zuerst eine glatte Fläche, dann eine dicke Leiste und eine Vertiefung mit Leisten und Punktgrübchen (Abb. 226).

Odontognophos Wehrli
45 (44) Kaudal des thorakalen Spiraculums eine längliche tomentöse Erhebung (vgl. Abb. 31)

Isturgia Hübner

[^1]

- Kaudal des thorakalen Spiraculums keinẻ tởmentöse Erhebung, nur grobe Runzeln. $\qquad$ Lobophora Curtis
46 (44) Dorsalrinne mit 8-9 mittelgroßen kaudalen Ausbuchtungen (Abb. 62). Lateraleinschnitt ohne Runzeln um die Spitze. Ventralseite des Kremasters mit großen Basalhöckern $\qquad$ Deileptenia Hübner
- Dorsalrinne mit etwa 20 kleinen Ausbuchtungen (Abb. 63). Lateraleinschnitt mit starken Runzeln um die Spitze. Basalhöcker an der Ventralseite des Kremasters schwach oder fehlend Arichanna Moore
47 (32) 5. Abdominalsegment frontal des Spiraculums mit einer schlank ovalen Leiste. Auch der Kaudalrand des 4 . Segmentes dort verdickt (Abb. 64)

48

- 5. Abdominalsegment frontal des Spiraculums mit etwa fünf Zellen, die kaudal durch eine dicke Querleiste begrenzt sind (Abb. 50)

Cleora Curtis

- 5. Abdominalsegment frontal des Spiraculums mit Punktgrübchen und/oder Leisten (Abb. 57, 60, 65, 68, 69) 49
48 (47) Puppe $6-8 \times 2,5 \mathrm{~mm} . . . . . .$. Stegania Gueneé - Puppe $9-10 \times 3 \mathrm{~mm} . . . . . . . . .$. Lygdia Gueneé

49 (47) Frontal des Spiraculums am 5. Abdominalsegment eine fast skulpturlose, glatte Fläche und vor ihr Punktgrübchen bzw. Leisten (Abb. 59, 65) 50

- Frontal des Spiraculums am 5. Abdominalsegment Punktgrübchen und/oder Leisten, ohne eine glatte Fläche dazwischen (Abb. 57, 68,69 ) 52
50 (49) 9. Abdominalsegment mit Punktgrübchen (vgl. Abb. 61). Alcis Curtis
- 9. Abdominalsegment ohne Punktgrübchen (vgl. Abb. 71)

51
51 (50) Lateral an der Basis des 5. Abdominalsegmentes eine dicke Querleiste und vor ihn dichte Punktgrübchen (Abb. 65)

Peribatodes Wehrli Lateral an der Basis des 5. Abdominalsegmentes nur Punktgrübchen, keine Querleiste (Abb. 59) $\qquad$ Paradarsia Warren
52 (49) An der Basis des 5. Abdominalsegmentes lateral nur Punktgrübchen (Abb. 57) 53

An der Basis des 5. Abdominalsegmentes lateral Leisten und dazwischen manchmal Punktgrübchen (Abb. 68, 69) 54

53 (52) Stiel des Kremasters lang (Abb. 66). Ventralseite des Kremasters ohne deutliche Basalhöcker. Metanotum ohne Punktgrübchen (vgl. Abb. 102) $\qquad$ Semiothisa Hübner

- $\quad$ Stiel des Kremasters kurz (Abb. 70). Ventralseite des Kremasters mit deutlichen Basalhöckern. Metanotum mit Punktgrübchen (vgl. Abb. 210)

Narraga Walker
54 (52) Stiel des Kremasters lang und schlank (Abb. 71) Diastictis Hübner Stiel des Kremasters kurz und dick (Abb. 67)

Itame Hübner
55 (1) Puppe glänzend braunschwarz, Kaudalstreifen des 5.-8. Abdominalsegmentes leuchtend gelb. Häkchen am Kremaster gleich groß, dick, zerstreut, $\mathrm{Sd}_{1}$ entspringt nahe der Basis des Kremasters (Abb. 72) ....... Abraxas Leach Puppe nicht glänzend schwärzlich mit gelben Seitenstreifen. Wenn die Häkchen am Kremaster gleich groß und dick sind, entspringt $\mathrm{Sd}_{1}$ nicht nahe der Basis des Kremasters................................................... 56
56 (55) Mittelteil der Puppe stark verbreitert (Abb. 76) Thorax und Abdomen dorsal mit dichten winzigen Börstchen besetzt (Abb. 74) $\qquad$ Apeira Gistl

- Puppe nicht verbreitert, ohne Börstchen.......
$\qquad$
57 (56) Am 9. Abdominalsegment an Stelle der Borsten $\mathrm{D}_{1}$ (und manchmal auch $\mathrm{D}_{2}$ ) zahnartige Spitzen. 9. Abdominalsegment dorsal stark längsgefurcht (Abb. 73). Puppen etwa 15-22 mm lang, matt.... Ennomos Treitschke Am 9. Abdominalsegment $\mathrm{D}^{1}$ borstenartig oder rückgebildet. Puppe oft kleiner und/oder glänzend

58
58 (57) Kremaster ventral stark radial gefurcht, und/oder seine Borsten, auch $D_{1}$, entspringen fächerartig von den Seiten der Spitze, sind $\pm$ gleich groß ( $\mathrm{D}_{2}$ manchmal etwas stärker) und am Ende stark spiralig eingerollt (Abb. 75, 79,80 ). Vorderschenkel nicht sichtbar oder klein (Abb. 82, 86, 88). Puppen bunt, oft grün gefärbt und/oder mit Zeichnungen.......

Abb. 88-114. - 88, Chlorissa cloraria; 89, Epirrita christyi; 90, Larentia clavaria; 91, Selenia dentaria; 92, Hypoxystis pluviaria; 93, Plagodis pulveraria; 94, Opisthograptis luteolata; 94a, Therapis flavicaria; 95, 100, Angerona prunaria; 96, 101, Crocallis elinguaria; 97, Epione parallelaria; 98, Synopsia sociaria; 99, Aspilates gilvaria; 102, Epione repandaria; 103, 105-107, Dyscia conspersaria; 104, 108, Pseudopanthera macularia; 109, Petrophora chlorosata; 110-112, Epirrhanthis diversata, 113, Hylaea fasciaria; 114, Odontopera bidentata.
88, 94, 94a, Habitusbild in Ventralsicht; 89, 90, 105, 108, 109, 110, 114, Abdominalende in Dorsalsicht; 91, 106, 111, 113, Erhebung kaudal von dem thorakalen Spiraculum; 92, Basis des 5. Abdominalsegmentes in Lateralsicht; 93, 107 , Abdominalende in Lateralsicht; 95, 96, Augen und Umgebung; 97, Lateraleinschnitt und Dorsalrinne in Lateralsicht; 98101, Kopf und Beine in Ventralsicht; 102-104, Metanotum, 1. Abdominalsegment; 112, Labrum, Labium.

- Kremaster anders gestaltet, ventral meist nicht regelmäßig radial gefurcht. $D_{1}$ wenn vorhanden, entspringt von seiner Dorsalseite. Borsten oft ungleich groß oder lang. Im Zweifelsfall Vorderschenkel groß (Abb. 1), Puppen oft zeichnungslos und $\pm$ glänzend .... 67
59 (58) Proboscis so lang wie die Mittelbeine (Abb. 82, 86). Abdomen ohne stachelige Skulptur . 63
- Proboscis kürzer als die Mittelbeine, doch länger als die Hälfte dieser (Abb. 77). Wenigstens die mittleren Abdominalsegmente mit stacheliger Skulptur (Abb. 78) ....... 60
- Proboscis kürzer als die Hälfte der Mittelbeine. Abdominalsegmente ohne stachelige Skulptur

Aplasta Hübner
60 (59) Puppe einfarbig grün, Exuvie weißlich grüngelb. Hemistola Warren

- Puppe mit Zeichnungen und dunklen Pinnacula der Borsten. 61
61 (60) Puppe grün, am Rücken z. T. rotbraun, 1520 mm lang. Hinterbeine meist nicht sichtbar. Geometra Linnaeus
- Puppe nicht grün gefärbt, $11-14 \mathrm{~mm}$ lang. Spitze der Hinterbeine sichtbar, groß ...... 62
62 (61) Flügeladerung deutlich schwarz abgesetzt, Grundfärbung weißgräulich

Thetidia Boisduval

- Flügeladern kaum dunkler, Grundfärbung hell rostbraun ................ Comibaena Hübner
63 (59) Kremaster mittellang mit 4 Paaren von Häkchen (Abb. 80)

64

- Kremaster sehr lang mit 5-7 Paaren von Häkchen (Abb. 79) .................. Jodis Hübner
64 (63) Dorsalrinne und Lateraleinschnitt vorhanden, deutlich (Abb. 80, 81)
$P$ seudoterpna Hübner
- Dorsalrinne und Lateraleinschnitt nicht vorhanden (vgl. Abb. 93)

65
65 (64) Puppe recht schlank, mehr als 4 mal so lang wie breit. Frontalhöcker relativ groß (Abb. 86). Puppe eintönig sandfarben oder grünlich, nur der Rückenstreifen am 1.-8. Abdominalsegment dunkel. Spiracula hell .... Hemithea Duponchel Puppe $\pm$ gedrungen, Frontalhöcker klein (Abb. 82, 88). Puppe meist bunter gezeichnet, auch am Kopf und Thorax. Spiracula dunkel gefleckt (Abb. 83)

66
66 (65) Grenze zwischen Vorderbein und Antenne etwa $2 \times$ so lang wie die Grenze zwischen Vorderbein und Gena (Abb. 84). Kopf und Thorax ventral, auch die Adern der Vorderflügel sehr kontrastreich dunkel gezeichnet.

Puppe relativ schlanker, $13-15 \mathrm{~mm}$ lang....... Thalera Hübner

- Grenze zwischen Vorderbein und Antenne etwa $3 \times$ so lang wie die Grenze zwischen Vorderbein und Gena (Abb. 85). Kopf und Thorax ventral oft weniger kontrastreich gezeichnet. Puppe mehr gedrungen, $9-11 \mathrm{~mm}$ lang. Chlorissa Stephens
67 (58) Vorderschenkel nicht sichtbar (Abb. 99, 101) 68
- Vorderschenkel sichtbar (Abb. 1, 165, 166)

98
68 (67) $\mathrm{D}_{2}$ am Kremaster $\pm$ parallel, Spitze hakenförmig, wenn divergierend, dann erst im Kaudalteil (Abb. 109) oder Kremaster kurz und stumpf abgerundet (Abb. 119) ......... 72

- $\quad \mathrm{D}_{2}$ am Kremaster stark divergierend, Spitze oft mehr dorn- als hakenförmig, manchmal $\pm$ gestielt. Kremaster länger und/oder spitz (Abb. 89, 167, 168, 174)......................... 69
69 (68) Mittlerer Kaudalausläufer der Dorsalrinne vergrößert, tief (Abb. 89, 174)................. 70
- Mittlerer Kaudalausläufer der Dorsalrinne klein oder flach (Abb. 167, 168, 171) bzw. Dorsalrinne nicht entwickelt (Abb. 93) ... 73
70 (69) Labium nicht sichtbar (Abb. 87). Kremaster lang vorgezogen, $\mathrm{D}_{2}$ einen Winkel von fast $180^{\circ}$ bildend (Abb. 89) ...... Epirrita Hübner Labium groß (vgl. Abb. 135), Stiel des Kremasters kürzer, $\mathrm{D}_{2}$ einen spitzen Winkel bildend 71
71 (70) $\mathrm{Sd}_{1}$ vorhanden, entspringt nahe der Basis des Kremasters, Kaudalausläufer der Dorsalrinne spitz (Abb. 90) $\qquad$ Larentia Treitschke
- $\quad \mathrm{Sd}_{1}$ am Kremaster fehlt. Kaudalausläufer der Dorsalrinne stumpf abgerundet (Abb. 174)

131
72 (68) Proboscis überragt deutlich den Kaudalrand des 4. Abdominalsegmentes (Abb. 94 a)

Therapis Hübner

- Proboscis überragt den Kaudalrand des 4. Abdominalsegmentes nicht (Abb. 94)...... 73
73 (72) Dorsalrinne nicht entwickelt (Abb. 93) ... 74 Dorsalrinne entwickelt, mindestens stärker und tiefer als der Dorsaleinschnitt zwischen dem 8. und 9. Abdominalsegment (Abb. $105,109,110$ )

75
74 (73) Punktgrübchen am Abdomen fehlen (Abb. 92). Proboscis kürzer als die Vorderbeine (vgl. Abb. 98) $\qquad$ Hypoxystis Prout

- Punktgrübchen am Abdomen vorhanden (vgl. Abb. 102). Proboscis länger als die Vorderbeine (vgl. Abb. 94)....... Plagodis Hübner
75 (73) Kremaster mit 4 Paaren von Häkchen (Abb. $105,114) . \mathrm{D}_{2} \pm$ größer als die übrigen Borsten.

76

- Kremaster mit 2 Paaren von Häkchen. $\mathrm{D}_{2}$ lang, schlank. Puppe $10-11 \mathrm{~mm}$ lang. Pachycnemia Stephens
- Kremaster mit 2 Paaren von Häkchen. $\mathrm{D}_{2}$ kürzer, relativ dick (Abb. 164). Puppe 16-18 mm lang

Colotois Hübner
76 (75) Adern der Vorderflügel stark hervortretend. Vorderschenkel meist etwas sichtbar (Abb. 94). Puppe sandfarben

Opisthograptis Hübner

- Adern der Vorderflügel nicht oder kaum hervortretend 77
77 (76) Vorderbeine grenzen aneinander (Abb. 98, 99)......................................................... 78 Vorderbeine grenzen nicht aneinander (Abb. 94, 100)................................................. 79
78 (77) Mittelbeine grenzen aneinander (Abb. 98). Kremaster wenig länger als basal breit. $\qquad$ Synopsia Hübner
- Mittelbeine grenzen nicht aneinander (Abb. 99). Kremaster länger als basal breit (Abb. 105)

Aspilates Treitschke
79 (77) Mittelbeine grenzen aneinander (Abb. 100, 101)....................................................... 80

- Mittelbeine grenzen nicht aneinander (vgl. Abb. 94) ................................................. 82
80 (79) Kremaster in Lateralsicht stumpf. Dorsalrinne mit flachen, breiten Ausläufern. Puppe unter 13 mm lang Cleorodes Warren
- Kremaster in Lateralsicht eher spitz. Dorsalrinne mit tieferen Kaudalausläufern. Puppe über 15 mm lang 81
81 (80) Puppe fein skulpturiert, am Rücken glänzend, Augen glatt (Abb. 95). Antennen und Mittelbeine kaudal stumpf, abgerundet (Abb. 100) .................. Angerona Duponchel
- Puppe grob skulpturiert, am Rücken kaum glänzend. Augen dorsal grob skulpturiert (Abb. 96). Antennen und Mittelbeine kaudal spitz, keilförmig (Abb. 101).
..................................... Crocallis Treitschke
82 (79) Kremaster (in Dorsalsicht, ohne Häkchen) fast zweimal so lang als basal breit (Abb. 105, 107). Kaudal der thorakalen Spiracula je eine relativ große tomentöse Erhebung (Abb. 91, 106). 83
- Kremaster kaum länger oder kürzer als basal breit. Im Zweifelsfall der Höcker kaudal des thorakalen Spiraculum klein, unauffällig, kaum tomentös....................................... 85
83 (82) Dorsalrinne mit starken, abgerundeten Kaudalausläufern (vgl. Abb. 114)

Charaspilates Wehrli

- Dorsalrinne mit undeutlichen, kleinen bzw. mehr spitzen Kaudalausläufern (Abb. 105) ...

84 (83) 1. Abdominalsegment ohne Punktgrübchen (Abb. 103). Lateraleinschnitt groß, tief (Abb. 107).................................... Dyscia Hübner 1. Abdominalsegment mit starken Punktgrübchen (Abb. 102). Lateraleinschnitt schwach entwickelt oder klein (Abb. 97) ......

Epione Duponchel
85 (82) Metanotum mit Punktgrübchen (Abb. 104)
86

- Metanotum ohne Punktgrübchen (Abb. 102, 103)....................................................... 87

86 (85) Pro- und Mesonotum mit Punktgrübchen (Abb. 104). $\mathrm{D}_{2}$ am Kremaster hakenförmig (Abb. 108) .............. Pseudopanthera Hübner Pro- und Mesonotum ohne Punktgrübchen (vgl. Abb. 210). $\mathrm{D}_{2}$ am Kremaster schlank dornförmig, in der kaudalen Hälfte voneinander divergierend (Abb. 109)

Petrophora Hübner
87 (85) Thorakale Spiracula kaudal mit je einem großen, elliptischen, tomentösen Höcker (Abb. 91) Selenia Hübner

- Thorakale Spiracula kaudal ohne oder nur mit kleinen und schmalen, bzw. nicht tomentösen Höckern (Abb. 111, 113) ......... 88
88 (87) 1-2. Abdominalsegment ohne Punktgrübchen (vgl. Abb. 103). $\mathrm{D}_{2}$ am Kremaster meist $\pm$ weit voneinander entfernt (Abb. 120-122)

96

- 1-2. Abdominalsegment mit Punktgrübchen (vgl. Abb. 102). $\mathrm{D}_{2}$ am Kremaster entspringen nahe beieinander (Abb. 110, 114, 119)

89
89 (88) $\mathrm{D}_{2}$ am Kremaster in Dorsalsicht deutlich länger als der Kremaster selbst, parallel (Abb. 110). Labrum auffallend lang, abgerundet (Abb. 112)................... Epirrbanthis Hübner

- $\quad \mathrm{D}_{2}$ am Kremaster nicht länger als der Kremaster selbst oder distal stark divergierend (Abb. 114, 115, 119). Labrum meist kurz und breit (Abb. 116, 117)

90
90 (89) 10. Abdominalsegment (ohne Kremaster) in Dorsalsicht fast 2 mal länger als das 9. Abdominalsegment, mit einer gebogenen, feinen Querfurche (Abb. 114). Puppe $17-20 \mathrm{~mm}$ lang.

Odontopera Stephens

- 10. Abdominalsegment (ohne Kremaster) nicht oder wenig länger als das 9. Abdominalsegment, ohne durchgehende Querfurche. Puppen oft kürzer als 15 mm .................... 91
91 (90) Lateraleinschnitt groß, wenigstens bis zur Mitte des 10. Abdominalsegmentes reichend (Abb. 115)

92
Lateraleinschnitt klein und kurz (Abb. 118)
95
92 (91) Kaudal des thorakalen Spiraculums je eine

längliche, stark tomentöse Erhebung ( Abb . 113). Lateraleinschnitt distal schmal. Kremaster auch im distalen Bereich dorsal skulpturiert, $\mathrm{Sd}_{1}$ entspringen etwa in seiner Mitte

Hylaea Hübner

- Kaudal des thorakalen Spiraculums ebenfalls je eine längliche tomentöse Erhebung. Lateraleinschnitt distal $\pm$ breit. Kremaster dorsal auch im distalen Bereich skulpturiert, $\mathrm{Sd}_{\mathrm{i}}$ entspringt nahe seiner Basis


## Menophra Moore

Kaudal des thorakalen Spiraculum liegt keine oder eine ganz unauffällige und kaum tomentöse Erhebung (Abb. 111). Kremaster im distalen Bereich $\pm$ glatt. Sd $_{1}$ entspringt meist nicht so nahe der Kremasterbasis ............. 93
93 (92) Labrum mit wenig konvergierenden Seiten (Abb. 117). Puppen fein skulpturiert, relativ stark glänzend. Kremaster im Basalteil schwach gerunzelt ... Puengeleria Rougemont Labrum trapezförmig mit stark konvergierenden Seiten (Abb. 116). Puppe nur mäßig glänzend, $\pm$ gröber skulpturiert. Kremaster dorsal an der Basis stark gerunzelt. 94
94 (93) Puppe $12-20 \mathrm{~mm}$ lang. 8. Abdominalsegment ohne Punktgrübchen Campaea Lamarck

- $\quad$ Puppe 8-11 mm lang. 8. Abdominalsegment mit starken Punktgrübchen $\qquad$
95 (91) $\mathrm{Sd}_{1}$ entspringen nahe der Basis des Kremasters, dieser kurz, stumpf, abgerundet, $\mathrm{D}_{2}$ groß (Abb. 119)........... Lomographa Hübner
- $\quad \mathrm{Sd}_{1}$ entspringen etwa von der Mitte des Kremasters, dieser relativ lang, spitz (Abb. 118)

Cabera Treitschke
96 (88) $\mathrm{Sd}_{1}$ entspringen am Kremaster auf der Höhe von $D_{1}$ in der Mitte des Kremasters oder distal davon (Abb. 121, 122). Lateraleinschnitt schmal 97

- $\quad \mathrm{Sd}_{1}$ entspringen am Kremaster mehr frontal als $\mathrm{D}_{1}$, nahe der Basis (Abb. 120). Lateraleinschnitt kurz, breit, stumpf abgerundet

Sciadia Hübner
97 (96) $\mathrm{D}_{2}$ mehr als halb so lang wie der Kremaster, dieser kürzer als basal breit.

Dorsalrinne mit 2-3 Kaudalausläufern (Abb. 224). Labium sehr groß (Abb. 225) Perconia Hübner
$\mathrm{D}_{2}$ mehr als halb so lang wie der Kremaster, dieser kürzer als basal breit. Dorsalrinne mit mehreren Kaudalausläufern (Abb. 122). Labium groß

Gnophos Treitschke s. lat.
$\mathrm{D}_{2}$ weniger als halb so lang wie der Kremaster, dieser länger als basal breit. Dorsalrinne mit mehreren Kaudalausläufern (Abb. 121). Labium groß Crocota Hübner
98 (67) An der Basis der Vorderflügel je ein starker Höcker (Abb. 123, 125, 130) 99 An der Basis der Vorderflügel kein Höcker (Abb. 1, 2) 101
99 (98) Proboscis überragt den Kaudalrand des 4. Abdominalsegmentes (Abb. 123). Kremaster mit relativ langen Borsten (Abb. 124)........................ Rhodometra Meyrick Proboscis überragt den Kaudalrand des 4. Abdominalsegmentes nicht (Abb. 125, 130). Kremaster mit kurzen Borsten (Abb. 127, 128) 100
100 (99) Frons mit langem, zweispitzigem Fortsatz (Abb. 125). Die Borsten am Ende des Kremasters einzeln (Abb. 127)

Timandra Duponchel

- Frons ohne Fortsatz (Abb. 130). Die Borsten am Ende des Kremasters in einer Gruppe (Abb. 128) .... Cyclophora Hübner
101 (98) Puppe $\pm$ matt, grün oder leder- bzw. sandfarben, dann mit dunkler Sprenkelung oder anderer Zeichnung bzw. schwarzen Spiracula. Exuvie grünweiß, gelbweiß oder trüb bräunlichgelb 102
- Puppe $\pm$ glänzend, gelb-, rot- bis schwarzbraun, zeichnungslos.......................... 113
102 (101) Frontolaterale Lappen des Metanotums spitz (Abb. 153) 128
- Frontolaterale Lappen des Metanotums abgerundet (Abb. 195)....................... 165
103 (165) Adern der Vorderflügel stark hervortretend. 76 Adern der Vorderflügel nicht stark hervortretend ............................................. 104

Abb. 115-145. - 115, 117, Puengeleria capreolaria; 116, Campaea honoraria; 118, Cabera pusaria; 119, Lomographa temerata; 120, Sciadia tenebraria; 121, Crocota nivearia, 122, Gnophos dilucidaria; 123, 124, Rhodometra sacraria, 125-127, Timandra griseata; 128, 130, Cyclophora linearia; 129, Ourapteryx sambucaria; 131, Eulithis populatac 132, 133, Ecliptopera silaceata; 134, Electrophaes corylata; 135, Dysstroma truncata; 136, Cidaria fulvata; 137, Thera variata; 138, Rheumaptera cervinalis, 139, 140, R. undulata; 141, Pbilereme transversata; 142, Triphosa dubitata; 143, Pterapherapteryx sexalata, 144, Xanthorhoe designata; 145, Epirrhoe alternata.
115-118, 137, Abdominalende in Lateralsicht; 116, 117, 133, 135, Labrum, Labium; 119-122, 124, 127-129, 131, 132, 134, 136, 143, Abdominalende in Dorsalsicht; 123, 125, 130, Habitusbild in Ventralsicht; 126, Kopffortsatz in Lateralsicht (Umriss); 138, 140, 141, 142, Basis des 5. Abdominalsegmentes in Lateralsicht; 139, Basis des 5. Abdominalsegmentes, Dorsalsicht; 144, 145, Auge und Umgebung.


104 (103) $\mathrm{D}_{2}$ am Kremaster an einem kurzen Stieli, eine Winkel von fast $90^{\circ}$ bildend (Abb. 129). Puppe über 18 mm lang
. Ourapteryx Leach

- $\quad \mathrm{D}_{2}$ am Kremaster nicht oder kaum gestielt. Puppe unter 16 mm lang 105
105 (104) Kremaster mit 4 Paaren von Borsten (Abb. 131, 132) 106
- Kremaster mit 6 Paaren von Borsten (Abb. $136,137)$ 111
106 (105) 10. Abdominalsegment dorsal stark verkürzt und $\pm$ längsgefurcht oder längsgerippt (Abb. 131). Puppen über 12, oft über 13 mm lang $\qquad$ Eulithis Hübner
- 10. Abdominalsegment stark verkürzt, jedoch nicht längsgefurcht. Puppenlänge unter 12 mm ....... Xanthorhoe Hübner p.p ( $X$. munitata (Hübner))
- 10. Abdominalsegment nicht stark verkürzt, kaum längsgefurcht (Abb. 132).

107
107 (106) Labium nicht sichtbar oder ganz klein (Abb. 133, 209) ................................ 108

- Labium relativ groß (Abb. 135) .......... 110

108 (107) Dorsalrinne ohne Kaudalausläufer (Abb. 132) Ecliptopera Warren

- Dorsalrinne mit Kaudalausläufern, wenigstens der mittlere gut sichtbar (Abb. 134, 208)

109
109 (108) Dorsalrinne mit deutlichen spitzen tomentösen Ausläufern (Abb. 134). Puppe deutlich dunkel gefleckt ...... Electrophaes Prout

- Dorsalrinne mit undeutlichen Ausläufern, nur der mittlere etwas mehr ausgeprägt (Abb. 208). Puppe undeutlich oder nicht gefleckt 156
110 (107) Puppe dünnschalig, nicht gesprenkelt, nicht bereift, Exuvie matt gelblich $\qquad$ Dysstroma Hübner Puppe dickschalig, dunkel gesprenkelt, deutlich bereift, Exuvie hellbraun …........................... Chloroclysta Hübner gen, insbesondere an den Adern der Vorderflügel, nicht bereift, nur $7-8 \mathrm{~mm}$ lang 150

111 (105) Dorsalrinne und Lateraleinschnitt nicht sichtbar (Abb. 136) ..... Cidaria Treitschke Dorsalrinne und Lateraleinschnitt $\pm$ sichtbar (Abb. 137) 112
112 (111) Borsten am Kremaster satt braunrot, Exuvie weißlichgelb. Dorsalrinne schwach, meist nur mit einzelnen, kleinen Kaudalausläufern. $\qquad$ Plemyria Hübner

- Borsten am Kremaster bräunlich. Exuvie bräunlich weiß oder gelb. Dorsalrinne mit mehreren meist deutlichen Kaudalausläufern (Abb. 137)

Thera Stephens
113 (101) An der Basis des 5. Abdominalsegmentes, dorsal oder lateral, dunkle Leisten, die Querrinnen oder -vertiefungen begrenzen (Abb. 138-142). 114

- An der Basis des 5. Abdominalsegmentes keine Leisten, Querrinnen oder -vertiefungen, höchstens dichte Punktgrübchen (Abb. 221)

116
114 (113) An der Dorsalseite der Basis des 5. Abdominalsegmentes Querleisten und dazwischen eine Querrinne. Sie laufen lateral $\pm$ in die Nähe der Spiracula herab (Abb. 138-140) Rheumaptera Hübner Nur lateral an der Basis des 5. Abdominalsegmentes Querleisten und Vertiefungen (Abb. 141, 142) 115
115 (114) 1-2 laterale Vertiefungen am 5. Abdominalsegment ungeteilt (Abb 141)

Philereme Hübner

- Laterale Vertiefung an der Basis des 5. Abdominalsegmentes in mehrere Zellen unterteilt (Abb. 142) ........ Triphosa Stephens
116 (113) Am Kremaster - außer den Borsten D2, die starke Endhäkchen, Enddornen bzw. eine gestielte Endgabel bilden - gibt es nur ein Paar kleiner Borsten ( $\mathrm{L}_{1}$ oder $\mathrm{D}_{1}$ ) (Abb. $143,146,150$ ) 117
- Am Kremaster, außer $D_{2}$ noch 2-3 Paaren von Borsten (Abb. 167, 183, 201, 218) ....

123
117 (116) Kremaster mit einem Paar von lateralen Höckern. $\mathrm{D}_{2}$ zusammen gestielt, stark divergierend (Abb. 143)

Pterapherapteryx Curtis

[^2]Kremaster ohne Lateralhöcker. Enddornen
schlanker, nicht so lang gestielt, weniger
divergierend (Abb. 146, 150)............ 118
118 (117) Grenzen zwischen Vorderbeine und Antennen viel länger àls die Grenzen zwischen Vorderbeine und Augen. Grenzen zwischen Antennen und Augen kaum länger als die zwischen Antennen und Vorderbeine (Abb. 144)

119

- Grenzen zwischen Vorderbeine und Antennen nicht oder wenig länger als die Grenzen zwischen Vorderbeine und Augen. Grenzen zwischen Antennen und Augen deutlich länger als die zwischen Antennen und Vorderbeine (Abb. 145) .... 120
119 (118) Am Kremaster außer $\mathrm{D}_{2}$ noch $\mathrm{L}_{1}$ vorhanden, nahe bei $D_{2}$ (vgl. Abb. 148) $\qquad$ Xanthorhoe Hübner
- Am Kremaster außer $\mathrm{D}_{2}$ noch $\mathrm{D}_{1}$, von $\mathrm{D}_{2}$ entfernt 120
120 (118) Dorsalrinne nur mit einem großen Kau-dal- und Frontalausläufer in der Mitte (Abb. 146, 147) .......... Catarhoe Herbulot
- Dorsalrinne mit mehreren Kaudalausläufern (Abb. 148, 150, 152) 121
121 (120) $\mathrm{D}_{2}$ so lang wie oder länger als der Kremaster (Abb. 148)................ Epirrhoe Hübner
- $\quad D_{2}$ kürer als der Kremaster, dieser $\pm$ länger als basal breit (Abb. 150, 152)

122
122 (121) Frontolaterale Lappen des Metanotums spitz (Abb. 149). Dorsalrinne im Mittelteil wenig verbreitert (Abb. 150)

Nycterosea Hulst

- Frontolaterale Lappen des Metanotums abgerundet. Dorsalrinne im Mittelteil stark verbreitert (Abb. 152).
........................ Camptogramma Stephens
123 (116) Frontolaterale Lappen des Metanotums spitzwinkelig (Abb. 153, 160). (Wenn jedoch das 1. Abdominalsegment mit deutlichen Punktgrübchen versehen und die Skulptur relativ grob ist, suche weiter unter 143) 124
- Frontolaterale Lappen des Metanotums stumpf, abgerundet (Abb. 151, 181).. 129
124 (123) $D_{2}$ gleichlang und selten stärker als die übrigen Borsten des Kremasters (Abb. 155-157) 125
- $\quad \mathrm{D}_{2}$ deutlich länger als die übrigen Borsten des Kremasters (Abb. 154, 159, 161)

125 (124) Dorsalrinne nur mit einem mittelgroßen Vorsprung in der Mitte. Labium klein (Abb. 193). Lateraleinschnitt kurz, abgerundet............................................... 149

Dorsalrinne mit kleinen Vorsprüngen, der mittlere aber sehr lang, bis nahe zur Basis des Kremasters reichend. (Labium an der untersuchten Puppe nicht vorhanden)

Glossotrophia Prout

- Dorsalrinne mit mehreren Vorsprüngen, keiner von ihnen nähert sich der Basis des Kremasters (Abb. 155, 157). Wenn nur einer dorsal in der Mitte oder gar keiner vorhanden ist, dann fehlt das Labium und/oder der Lateraleinschnitt sehr groß

Idaea Treitschke
126 (124) Kremaster rauh skulpturiert, $\mathrm{D}_{2}$ entspringen weit voneinander entfernt. Vorderschenkel klein 97

- Kremaster $\pm$ fein skulpturiert. $D_{2}$ entspringen nahe beieinander, Vorderschenkel groß 127
127 (126) Sd, entspringen nahe der Basis des Kremasters (Abb. 161). Labium klein (vgl. Abb. 192)

Baptria Hübner

- $\quad \mathrm{Sd}_{1}$ (wenn vorhanden) entspringen weit von der Basis des Kremasters entfernt (Abb. 154, 159). Labium meist groß (vgl. Abb. 135)

128
128 (127) Kaudal des thorakalen Spiraculums je eine große, elliptische, stark tomentöse Erhebung (Abb. 158). Puppe sandfarben oder grün ...................... Rhodostrophia Hübner Kaudal des thorakalen Spiraculums eine längliche, weniger auffällige Erhebung (vgl. Abb. 113). Puppe rotbraun, glänzend Scopula Schrank
129 (123) Kremaster mit 4 Paaren steifer, gleich groBer Borsten, am Ende nicht hakenartig. $\mathrm{D}_{2}$ entspringen weit voneinander entfernt (Abb. 162). Punktgrübchen fehlen

Glacies Milliere, 1874
Wenigstens ein Teil der Borsten am Kremaster hakenartig. Punktgrübchen am Abdomen $\pm$ vorhanden

130
130 (129) Vorderschenkel klein, oft recht schmal. Labium klein oder fehlend (Abb. 165). $\mathrm{D}_{2}$ am Kremaster länger und stärker als die übrigen Borsten, $\pm$ divergierend (Abb. $167,168,171,174)$.

131
Vorderschenkel, Labium, oder beide mittelgroß bis groß (Abb. 1), sonst $\mathrm{D}_{2}$ am Kremaster nicht viel länger oder stärker als die übrigen Borsten. (Abb. 193)......... 135
131 (130) Dorsalrinne in der Mitte beiderseits, insbesondere kaudalwärts, mit abgerundeten Ausschnitt. Borsten am Enddrittel des Kremasters konzentriert (Abb. 174). Punktgrübchen an der Basis des 1.-3. Abdominalsegmentes stark vergrößert (Abb.
176)...................... Anticlea Stephens p.p. (A. badiata Denis et Schiffermüller) Dorsalrinne mit einem oder mehreren flachen Kaudalen Ausschnitten. Borsten auch an der Basalhälfte des Kremasters (Abb. 167, 168, 171). Punktgrübchen an der Basis des 1. -3. Abdominalsegmentes nicht vergrößert

132
132 (131) Metanotum mit Punktgrübchen (Abb, 181, 210) 133

- Metanotum ohne Punktgrübchen (Abb. 151)................................................. 134

133 (132)Kremaster in Dorsalsicht länger als basal breit. Borsten $\mathrm{Sd}_{1}$ entspringen nahe der Basis des Kremasters (Abb. 167) $\qquad$ . Nothacasis Prout

- Kremaster in Dorsalsicht kürzer als basal breit. $\mathrm{Sd}_{1}$ entspringen etwa in der Mitte des Kremasters (Abb. 168)

Acasis Duponchel
134 (132) Puppe ohne oder mit sehr kleinem Labium. Lateraleinschnitt undeutlich oder in der Form eines Grübchens (Abb. 169, 170). Trichopteryx Hübner

- Puppe mit deutlicherem mittelkleinem Labium. Lateraleinschnitt rinnenförmig, mit einem kurzen Kaudalausläufer (Abb. 172). $\qquad$ . Anticollix Prout 135 (130) Kremaster kürzer als basal breit, sehr grob skulpturiert (gefurcht, gerippt). $\mathrm{D}_{2}$ hakenförmig, kaum divergierend, stärker und länger als die übrigen Borsten, basal $\pm$ voneinander entfernt, der Kremaster dazwischen $\pm$ konkav (Abb. 163). Epicranialnaht zwischen Vertex und Frons nicht sichtbar (Abb. 4). Puppen über 11 mm
lang. 97
- Kremaster kürzer als breit, stumpf, grob skulpturiert. $\mathrm{D}_{2}$ stärker als die übrigen Borsten, basal nahe beieinander, ohne Einschnitt dazwischen, ziemlich parallel laufend. Epicranialnaht sichtbar. Puppenlänge $10-12 \mathrm{~mm}$ Epilobophora Inoue
- Kremaster oft länger als breit, nur schwach oder mäßig skulpturiert. $\mathrm{D}_{2}$ entspringen entweder nahe beieinander und divergieren, oder sind nicht größer als die übrigen Borsten des Kremasters. Puppen manchmal auch kürzer als 10 mm . Epicranialnaht relativ deutlich (Abb. 3) ..................... 136
136 (135) Kremaster mit 3 Paar Borsten (Abb. 178180)................................................. 137
- Kremaster mit 4 Paar Borsten (Abb. 190, 207, 213, 218).................................. 144
137 (136) Lateraleinschnitt nicht entwickelt, oder als einfache Rinne ohne einen größeren
Kaudalausläufer (Abb. 175, 182) ..... 138
Lateraleinschnitt deutlich entwickelt, miteinem starken Kaudalausläufer (Abb. 184)142

138 (137) Borsten am Kremaster gleich groß (Abb. 179) ......................... Gymnoscelis Mabille $\mathrm{D}_{2}$ viel größer als die übrigen Borsten am Kremaster (Abb. 180) ........................ 139
139 (138) Punktgrübchen an der Basis des 1.-4. Abdominalsegment stark vergrößert und vertieft (Abb. 176). 140

- Punktgrübchen an der Basis des 1-4. Abdominalsegmentes nicht vergrößert (Abb. 151) 141
140 (139) Kremaster am Ende abgerundet, schaufelförmig (Abb. 177). Puppe $7-8 \mathrm{~mm}$ lang.... Discoloxia Warren
- Kremaster am Ende spitz (Abb. 178). Puppe $8-9 \mathrm{~mm}$ lang........... Venusia Curtis
141 (139) Metanotum ohne Punktgrübchen (vgl. Abb. 194) ..................... Euchoeca Hübner Metanotum mit Punktgrübchen (Abb. 181) .. Hydrelia Hübner
142 (137) Kremaster in Dorsalsicht nicht länger als basal breit (Abb. 183)..... Euphyia Hübner
- Kremaster in Dorsalsicht länger als basal breit (Abb. 185)................................ 143
143 (142) Dorsalrinne nur mit drei Kaudalausläufern (Abb. 185)................... Mesoleuca Hübner Dorsalrinne mit mehreren Kaudalausläufern (Abb. 186) .................................. 146
144 (136) Puppenlänge über 11 mm . Punktgrübchen fehlen am Metanotum und meist auch am 1. Abdominalsegment (Abb. 187, 219); falls vorhanden, dann Kremaster recht lang und die Borsten am 9. Abdominalsegment deutlich (Abb. 186) 145
- Metanotum oder wenigstens das 1. Abdominalsegment mit deutlichen Punktgrübchen, oder Puppen unter 10 mm lang....... . 148
145 (144) $D_{1}$ am Kremaster entspringt weit ventral von Sd (Abb. 186, 220) .................... 146
- $\quad D_{1}$ entspringt etwa auf derselben Höhe wie Sd $_{1}$ (Abb. 188, 189) ........................... 147
146 (145) Kremaster lang zungenförmig (Abb. 186). Borsten am 9. Abdominalsegment relativ kräftig (Abb. 186) ...... Scotopteryx Hübner - Kremaster in Dorsalsicht dreieckig, nicht länger als basal breit (Abb. 220).

Pareulype Herbulot
147 (145) Grenzen zwischen Vorderbeinen und Antennen viel länger als zwischen Vorderbeinen und Augen; Grenzen zwischen Antennen und Augen kaum länger als zwischen Antennen und Vorderbeinen (Abb. 144) ..


## Xanthorhoe Hübner p.' p." <br> (X.montanata (Denis et Schiffermüller), X. incursata (Hübner))

 Grenzen zwischen Vorderbeinen und Antennen nicht oder kaum länger als zwischen Vorderbeinen und Augen; Grenzen zwischen Antennen und Augen viel länger als zwischen Antennen und Vorderbeinen (Abb. 145) Entephria Hübner 148 (144) Grenzen zwischen Vorderbeinen und Antennen deutlich länger als zwischen Vorderbeinen und Augen (etwa gleich lang wie die Grenzen mit Augen und Genae zusammen). Grenzen zwischen Antennen und Augen etwa gleich lang wie zwischen Antennen und Vorderbeine (Abb. 144, 191). Kleine Arten, unter 9, meist unter 8 mm Länge, Skulptur oft grob, $\mathrm{D}_{2}$ am Kremaster hakenförmig, nicht oder nur wenig divergierend. (Abb. 196-201) Lateraleinschnitt und Labium meist gut entwickelt 149 Grenzen zwischen Vorderbeinen und Antennen nicht oder kaum länger als zwischen Vorderbeinen und Augen; Grenzen zwischen Antennen und Augen deutlich länger als zwischen Antennen und Vorderbeinen (Abb. 145). Wenn die Puppen unter 8 mm lang, fehlt oft das Labium und der Lateraleinschnitt. $\mathrm{D}_{2}$ oft gabelartig divergierend (Abb. 203-205, 220) 151149 (148) Labium sehr klein (Abb. 192). Dorsalrinne mit einem einzigen, kleinen Kaudalausläufer (Abb. 193). Borsten am Kremaster $\pm$ gleich groß, nur im distalen Drittel (Abb. 193). Frontolaterale Lappen des Metanotums ziemlich spitz (Abb. 194)

## Calliclystis Dietze

Labium größer (Abb. 1), Dorsalrinne mit mehreren Ausläufern (Abb. 196, 199, 200 ), oder $\mathrm{D}_{2}$ deutlich größer als die übrigen Borsten des Kremasters (Abb. 198). Frontolaterale Lappen des Metanotums $\pm$ abgerundet (Abb. 210)

150
150 (149) Metanotum ohne, 1. Abdominalsegment ohne oder mit schwach angedeuteten Punktgrübchen (Abb.195). Dorsalrinne
mit kleinen Kaudalausläufern, Borsten am Kremaster fast gleich groß, nur in der distalen Hälfte (Abb. 196)
................................ Chloroclystis Hübner

- Skulptur $\pm$ grob, 1. Abdominalsegment und oft auch das Metanotum (bzw. ganze Thorax dorsal) mit deutlichen Punktgrübchen. Sonst meist entweder die Kaudalausläufer der Dorsalrinne groß (Abb. 201) oder $\mathrm{D}_{2}$ länger als die übrigen Borsten des Kremasters (Abb. 197, 198, 200) bzw. Borsten auch in der Basalhälfte des Kremasters $\qquad$ Eupithecia Curtis
151 (148) $\mathrm{D}_{2}$ am Kremaster deutlich, weit voneinander entfernt, einem stumpfen Winkel bildend (Abb. 190). Metanotum mit Punktgrübchen (vgl. Abb. 210)

Anticlea Stephens p. p. (Anticlea erivata (Denis et Schiffermüller)) $\mathrm{D}_{2}$ entspringen nahe beieinander (Abb. 216, 218)

152
152 (151) $\mathrm{D}_{2}$ in Dorsalsicht länger als der Kremaster, dornartig, divergierend und $\pm$ lang gestielt. Oft kleinere Arten, unter 9 mm Länge (Abb. 203-205) $\qquad$ Perizoma Hübner

- $\quad \mathrm{D}_{2}$ in Dorsalsicht nicht länger als der Kremaster, oft hakenartig und $\pm$ parallel laufend (Abb. 213, 216, 223), wenn divergierend (Abb. 218, 220), dann oft größere Arten, über 9 mm lang 153
153 (152) Labium fehlt, oder sehr klein (vgl. Abb. 87, 209). Dorsalrinne und Lateraleinschnitt $\pm$ schwach ausgebildet. $\mathrm{D}_{2}$ hakenartig, viel stärker als die übrigen Borsten (Abb. 207, 208, 211). Oft kleinere Arten, 6-8 mm lang..................................... 154 Labium mittelgroß, bis groß (Abb. 1, 135). Dorsalrinne und Lateraleinschnitt meist deutlich und tief. $\mathrm{D}_{2}$ stärker oder auch gleichstark wie die übrigen Kremasterborsten 157
154 (153) Metanotum mit Punktgrübchen (vgl. Abb. 210). Puppe nur etwa 6-7 lang........... 155 Metanotum ohne Punktgrübchen (vgl. Abb. 195). Puppe oft länger als $7 \mathrm{~mm} . . . .$. 156
155 (154) $\mathrm{D}_{1}$ am Kremaster viel näher an $\mathrm{L}_{1}$ als an $\mathrm{Sd}_{1}$

Abb. 179-211. - 179, Gymnoscelis rufifasciata; 180, 181, Hydrelia flameolaria; 182, Euchoeca nebulata; 183, Eyphyia; 185, Mesaleuca albicillata; 186, 187, Scotopteryx moeniata; 188, Xanthorhoe montanata; 189, Entephria caesiata; 190 Anticlea derivata; 191-194, Calliclystis v-ata; 195, 196, Chloroclystis rectangulata; 197, Eupithecia absinthiata; 198, E. pulchellata; 199, E. tenuiata; 200, E. abietaria; 201, E. pusillata; 202, Perizoma alchemillata; 203, P. bifasciata; 204, P. didymata; 205, P. perallelolineata; 206, P. blandiata; 207, P. incultaria; 208, 209, Eustroma reticulata; 210, 211, Asthena albulata.
$179,180,183,185,186,188-190,193,196-208,211$, Abdominalende in Dorsalsicht; 181, 187, 194, 195, 210, Metanotum, 1. bzw. 2. Abdominalsegment; 182, 184, Abdominalende in Lateralsicht; 191, Augen und Bereich; 192, 209, Labrum, Labium.


Athb. 212-225. 212. itmon murinata; 213. Cosmorhoe ocellata; 214, Nebula salicata; 215, N. nebulata; 216, Colostygia aptalit: $21^{-}$. Hirtrumena ruheratas: 218, Lampropteryx otregiata; 219-221, Pareulype berberata; 222, Spargania luctuata; 223, Melanthia procellata, 224, 225, Perconia strigillaria, 226, 227, Odontognophos dumetata.
212-218. 220. 222-224, Ahdominalende in Dorsalsicht; 219, Metanotum, 1-2. Abdominalsegment; 221, 226, Basis der 5. Abdrminnkencenta in 1 aterakicht: 225, Labrium, Labium; 227, Höcker kaudal von dem thorakalen Spiraculum.
entspringend (Abb. 212)
Minoa Treitschke
$\mathrm{D}_{1}$ am Kremaster etwa in der Mitte zwischen $\mathrm{Sd}_{1}$ und $\mathrm{L}_{1}$ entspringend (Abb. 211)
$\qquad$ Asthena Hübner
156 (154) $\mathrm{D}_{1}$ am Kremaster entspringen frontal von $\mathrm{Sd}_{1}$ (Abb. 207)....... Perizoma Hübner p.p. .................(P. incultaria Herrich-Schäffer)

- $\quad \mathrm{D}_{1}$ am Kremaster entspringen kaudal von $\mathrm{Sd}_{1}$ (Abb. 208). Eustroma Hübner
157 (153) Borsten am Kremaster fast gleich lang und gleich dick, schlank. $\mathrm{Sd}_{1}$ entspringt nahe der Basis des Kremasters (Abb. 213)
. Cosmorhoe Hübner
- Borsten am Kremaster nicht gleich lang und gleich groß ( $\mathrm{D}_{2}$ deutlich größer, Abb . $216,218)$. Wenn doch gleich groß, dann entspringt $\mathrm{Sd}_{1} \pm$ auf der Höhe von $\mathrm{D}_{1}$, weit von der Basis des Kremasters entfernt (Abb. 214) 158
158 (157) $\mathrm{D}_{1}$ entspringt am Kremaster frontal von $\mathrm{Sd}_{1}$ (Abb. 217) ......... Hydriomena Hübner
- $\quad \mathrm{D}_{1}$ entspringt $\pm$ auf der Höhe von $\mathrm{Sd}_{1}$ (Abb. 214, 216) 159 $\mathrm{D}_{1}$ entspringt kaudal von $\mathrm{Sd}_{1}$. (Abb. 220, 222, 223).......................................... 161
$159(158) \mathrm{D}_{2}$ nicht, oder nur wenig länger als die übrigen Borsten (Abb. 214, 216) ....... 160
- $\quad \mathrm{D}_{2}$ deutlich länger und stärker als die übrigen Borsten des Kremasters.

Orthonama Hübner
160 (159) $\mathrm{D}_{2}$ kaum länger, manchmal etwas stärker als die übrigen Borsten am Kremaster, dieser kaudal $\pm$ stumpf (Abb. 214, 215)

Nebula Bryard

- $\quad \mathrm{D}_{2}$ deutlich länger und stärker als die übrigen Borsten des Kremasters, dieser kaudal zugespitzt (Abb. 216) .. Calostygia Hübner
161 (158) Metanotum und 1. Abdominalsegment ohne Punktgrübchen (Abb. 219) ....... 146
- Metanotum und 1. Abdominalsegment mit Punktgrübchen (Abb. 210).......... 162
162 (161) Kremaster in Dorsalsicht länger als basal breit, $\mathrm{D}_{2}$ oft dornartig, divergierend (Abb. 218) $\qquad$ Lampropteryx Stephens
- $\quad$ Kremaster in Dorsalsicht nicht länger als basal breit, $\mathrm{D}_{2}$ hakenartig (Abb. 220, 223)

163
163 (162) Kremaster in Dorsalsicht kürzer als basal breit. $\mathrm{D}_{2}$ hakenartig (Abb 223) $\qquad$ ............................... Melathia Duponchel

- Kremaster in Dorsalsicht nicht oder kaum kürzer als basal breit, $\mathrm{D}_{2}$ mitunter dornartig (Abb. 5, 222) 164
164 (163) Dorsalrinne mit deutlichen Kaudalausläufern, auch oft ein Frontalausläufer in
der Mitte vorhanden (Abb. 6). $\mathrm{D}_{2} \pm$ hakenförmig ...................... Horisme Hübner Dorsalrinne mit relativ kleinen Kaudalund keinem Frontalausläufer. $D_{2}$ am Kremaster eher dornartig, gebogen (Abb. 222)

Spargania Guéneé, 1857
165 (102) Proboscis kürzer als die Vorderbeine. Vorderflügel grenzen kurz aneinander.

Lythria Hübner
Proboscis länger als die Vorderbeine. Vorderflügel grenzen nicht aneinander....

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## Diskussion

In der vorliegenden Arbeit wurde auf eine Bestimmungstabelle der in den derzeitigen Systemen üblichen Unterfamilien der Geometridae, sowie der Tribus nach Herbulot (1962-1963) verzichtet. Die bisherigen Tabellen zur Bestimmung der Unterfamilien von Khotko (1977), sowie von McGuffin (1967-1981) kann man für größere oder kleinere Zahl der Gattungen, jedoch nicht für alle erfolgreich benutzen. Die Tabelle von Khotko ist allzu kompliziert - zur Bestimmung von fünf Unterfamilien sind dort mehr als zwei Textseiten nötig. Die Tabelle von McGuffin ist einfacher, es gibt jedoch bei der Bestimmung mancher Puppen der mitteleuropäischen Fauna Schwierigkeiten, besonders was die Unterscheidung der Unterfamilien Sterrhinae und Larentiinae betriff. Noch schwieriger ist dies bei den Tribus im Sinne von Herbulot. Nur für einen Teil von ihnen gelang es mir anhand der Puppenmerkmale eine benutzbare Diagnose aufzustellen und auch dort ist diese nicht für alle Gattungen nach dem Herbulot'schen System gültig. Unter die von puppenmorphologischer Hinsicht uneinheitlichen Tribus gehören z. B. die Pseudoterpnini (Unterfamilie Geometrinae) und die Lythriini (Unterfamilie Larentiinae). Mehrere Gattungen sollten anhand der Puppenmorphologie in andere Tribus gehören als in welche sie von Herbulot gestellt wurden, wie z. B. : Calospilos, Lomaspilis, Lygdia und Stegania (siehe Patoc"ka 1986a), oder Odontognophos, Cleorodes, Crocota, Synopsia und Menophra (siehe Patoc ${ }^{\text {ka }}$ 1993).
Die Gattungen des Herbulot'schen Systems entsprechen dagegen in der Mehrzahl auch auf Grund der Puppenmerkmale sehr gut. Auf wenige Ausnahmen soll hier aufmerksam gemacht werden: Die Art obstipata weicht von vittata stark ab, es wird für sie deshalb der Gattungsname Nycterosea (im Sinne von Forster-Wohlfart 1981) benützt. Ebenso wurde es mit den Gattungen Dysstroma, Chiasmia, Diastictis und Phigalia gemacht. Außerdem ist auch die Gattung Xanthorhoe puppenmorphologisch uneinheitlich. Insbesondere die Art $X$. munitata entspricht
viel besser der gartung Eulithis. Die Arten Xanthorhoe montanata und incursata weichen von den übrigen stärker ab. Ähnlich ist es auch mit der Gattung Perizoma, wo besonders die Art incultaria den übrigen puppenmorphologisch unähnlich ist. Stark voneinander abweichende Arten enthält auch die Gattung Anticlea. Dagegen sind die größeren oder großen Gattungen Cyclophora, Sterrha, Idaea und Eupithecia hinsichtlich der Puppenmorphologie relativ einheitlich gebaut. Die Verhältnisse in der Gattung Gnophoss. lat. (vgl. Sauter 1990) werden in einer besonderen Arbeit (Patočka im Druck) besprochen.

## Literatur

Forster, W. \& T. A. Wohlfahrt, 1981. Die Schmetterlinge Mitteleuropas 5, Spanner (Geomerridae). - Franckh'sche Verlagshandlung, Stuttgare: 1-311.
Gustafsson, B. (ed), 1987. Catalogus Lepidopterorum Sueciae. - Riksmuseet, Stockholm: 1-140.
Herbulot, C., 1962-1963. Mise à jour de la liste des Geometridae de France. - Alexanor, 2: 117-124, 147154, 3: 17-24, 85-93.
Khotko, E. I., 1977. A key to the spanpupae (Lepidoptera, Geometridae). - Academia Scientiarum Beloruss. SSR, Minsk: 1-80.
Koehler, W., 1937. Beitrag zur Kennenis der, unter Streudecke der Kiefernbestände überwinternden Schmetterlingspuppen. - Institut des recherches des forêts dominiales Pologne, Traxaux et comptes rendus Warszawa, Ser. A. Nr. 29: 1-81
Leraut, P. 1980. Liste systématique et synonymique des Lépidoptères de France, Belgique et Corse. - Alexanor, Supplément: 1-334.
Ljungdahl, D., 1919. Några puppenskrivningar. - Entomologisk Tidskrift 40: 97-119.
McGuffin, W. C., 1967-1981. Guide to the Geometridae of Canada. - Memoirs of the Entomological Society of Canada 50: 1-103, 86: 1-159, 101: 1-191, 117: 1-153.
Mosher, E., 1916. A classification of the Lepidoptera based on characters of the pupa. - Bulletin of the Illinois State Laboratory of Natural History 12: 1-159.
Nordström, F., E. Wahlgren \& A. Tulgren, 1941. Svenska Fjärilar. - Nordisk Familjeboks Förlags Akriebolag Stockholm: 1-354.
Patočka, J., 1978. Zur Puppenmorphologie und -Taxonomie der Unterfamilie Ennominae insbesondere der Tribus Bistonini (Lepidoptera, Geometridae). - Vestnik Ceskoslovenské Společnosti Zoologické 42: 143-151.
Patockka, J., 1980. Die Raupen und Puppen der Eichenschmetterlinge Mitteleuropas. - Monografiën zur angewandten Entomologie, Verl. Paul Parey, Hamburg \& Berlin: 1-188.
Patočka, J. 1980-1983. Beitrag zur Puppentaxonomie der mitteleuropäischen Larentiinae (Lepidoptera, Geometridae). - Biológia, Bratislava, 35: 97-109, 36: 583-593, 37: 559-570, 38: 117-131.
Patocka, J., 1985. Beitrag zur Kenntnis der Puppen der Unterfamilie Ennominae (Lepidoptera, Geomerridae). Biológia, Bratislava, 40: 997-1012.
Patočka, J., 1986a. Zur Kenntnis der Puppen der mitteleu-
ropäischen Spanner aus der Tribus Abraxini und Semiothisini (Lepidoptera, Geometridae). - Biologgia, Bratislava, 41: 579-595.
Patočka, J., 1986b. Zur Kenntnis der Puppen der Tribus Boarmiini (Lepidoptera, Geometridae) in Mitteleuropa. - Acta Entomologica Bohemoslovaca 83: 301-315.

Patočka, J., 1992. Über einige Puppen der Spanner aus der Tribus Ennomini (Lepidoptera, Geometridae, Ennominae). - Entomologische Berichten, Amsterdam 52: 171176.

Patočka, J., 1993. Über einige Puppen der Spanner aus der Tribus Boarmini (Lepidoptera, Geometridae, Ennominae). - Entomologische Berichten, Amsterdam 53: 114-120.
Patočka, J., 1994. Die Puppen der mittel- und westeuropäischen Spanner aus der Tribus Asthenini (Lepidoptera, Geometridae, Larentinae). - Entomologische Berichten, Amsterdam 54: 13-19.
Patočka, J., im Druck. Die Puppen der Gnophos-Gruppe (Lepidoptera, Geometridae) Mitteleuropas. - Zeitschrift der Arbeitsgemeinschaft österreichischen Entomologen: im Druck.
Patočka, J., J. Burgan, M. Capek \& M. Stolina, 1960. Die Tannenschmetterlinge der Slowakei. - Ed. SAV Bratislava: 1-214.
Sauter, W., 1990. Zur Systematik der Gnophos-Gruppe (Lepidoptera, Geometridae). - Nota Lepidopterologica 12:328-343.
Speyer, W., 1958. Lepidopteren-Puppen an Obstgewächsen und in ihrer näheren Umgebung. - Mitteilungen aus der Biologischen Bundesanstalt für Land- und Forstwirtschaft 93: 1-40.

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## ANHANG

Übersicht der Gattungen und Arten der Familie Geometridae, deren Puppen als Unterlage für die Aufstellung dieser Gattungstabelle untersucht wurden.

Archiearis Hübner, 1823
parthenias (Linnaeus, 1761) (Slowakei, Holland)
notha (Hübner, 1803) (Slowakei)
puella (Esper, 1787) (Slowakei)
Alsophila Hübner, 1825
aescularia ([Denis \& Schiffermüller], 1775) (Slowakei)
quadripunctaria (Esper, 1787) (Slowakei)
Aplasta Hübner, 1823
ononaria (Fuessly, 1783) (Deutschland)
Pseudoterpna Hübner, 1823
pruinata (Hufnagel, 1767) (Slowakei)
Geometra Linnaeus, 1758
papilionaria (Linnaeus, 1758) (Böhmen)
Comibaena Hübner, 1823
bajularia ([Denis \& Schiffermüller], 1775) (Slowakei)

Thetidia Boisduval, 1840
smaragdaria (Fabricius, 1787) (Slowakei)
Hemithea Duponchel, 1829
aestivaria (Hübner, 1799) (Slowakei, Böhmen)
Chlorissa Stephens, 1831
viridata (Linnaeus, 1758) (Polen)
cloraria (Hübner, 1813) (Slowakei)
etruscaria (Zeller, 1848) (Slowakei)
Thalera Hübner, 1823
fimbrialis (Scopoli, 1763) (Österreich)
Hemistola Warren, 1893
biliosata (Villers, 1789) (Slowakei)

Jodis Hübner, 1823
lactearia (Linnaeus, 1758) (Slowakei)
putata (Linnaeus, 1758) (Böhmen, Österreich)
Cyclophora Hübner, 1822
pendularia (Clerck, 1759) (Slowakei, Böhmen)
albiocelaria (Hübner, 1789) (Slowakei) annulata (Schulze, 1775) (Slowakei) albipunctata (Hufnagel, 1767) (Slowakei, Böhmen) puppillaria (Hübner, 1799) (Frankreich) ruficiliaria (Herrich-Schäffer, 1855) (Slowakei) porata (Linnaeus, 1767) (Österreich) quercimontaria (Bastelberger, 1897) (Slowakei) punctaria (Linnaeus, 1758) (Slowakei)
suppunctaria (Zeller, 1847) (Slowakei)
linearia (Hübner, 1799) (Slowakei)
Timandra Duponchel, 1829
griseata W. Petersen, 1902 (Slowakei)
Scopula Schrank, 1802
immorata (Linnaeus, 1758) (Slowakei)
umbelaria (Hübner, 1813) (Österreich)
virgulata (Denis \& Schiffermüller, 175) (Slowakei)
ornata (Scopoli, 1763) (Slowakei)
rubiginata (Hufnagel, 1767) (Slowakei)
marginepunctata (Goeze, 1781) (Slowakei)
corrivalaria (Kretschmar, 1862) (Deutschland)
immutata (Linnaeus, 1758) (Slowakei)
ternata Schrank, 1802 (Slowakei)
Glossotrophia Prout, 1913
asellaria (Millière, 1868) (Italien)
Idaea Treitschke, 1825
rufaria (Hübner, 1799) (Österreich) serpentata (Hufnagel, 1767) (Slowakei)
aureolaria (Denis \& Schiffermüller, 1775) (Österreich)
extersaria (Herrich-Schäffer, 1848) (Italien)
typicata (Gueneé, 1857) (Italien)
filicata (Hübner, 1799) (Österreich)
laevigata (Hübner, 1799) (Deutschland)
obsoletaria (Rambur, 1833) (Italien)
biselata (Hufnagel, 1767) (Slowakei)
calunetaria (Staudinger, 1859) (Schweiz)
inquinata (Scopoli, 1763) (Deutschland)
dilutaria (Hübner, 1799) (Slowakei)
fuscovenosa (Goeze, 1781) (Slowakei)
bumiliata (Hufnagel, 1767) (Slowakei)
seriata (Schrank, 1802) (Ősterreich)
subsericeata (Haworth, 1809) (Italien)
emarginata (Linnaeus, 1758) (Deutschland)
aversata (Linnaeus, 1758) (Slowakei)
degeneraria (Hübner, 1799) (Slowakei)
deversaria (Herrich-Schäffer, 1847) (Slowakei)
Rhodostrophia Hübner, 1823
vibicaria (Clerck, 1759) (Slowakei)
Rhodometra Meyrick, 1892
sacraria (Linnaeus, 1767) (Bosnien)
Lythria Hübner, 1823
purpuraria (Linnaeus, 1758) (Slowakei)
rotaria (Fabricius, 1798) (Russland)
Cataclysme Hübner, 1825
riguata (Hübner, 1813) (Slowakei)
Phibalapteryx Stephens, 1829
virgata (Hufnagel, 1767) (Slowakei)
Scotopteryx Hübner, 1825
moeniata (Scopoli, 1763) (Österreich)
coarctaria ([Denis \& Schiffermüller], 1775) (Slowakei) chenopodiata (Linnaeus, 1758) (Slowakei, Dänemark) luridata (Hufnagel, 1767) (Slowakei)

Larentia Treitschke, 1825
clavaria (Haworth, 1809) (Polen)
Orthonama Hübner, 1825
vittata (Borkhausen, 1794) (Deutschland)
Nycterosea Hulst, 1896
obstipata (Fabricius, 1794) (Österreich)
Xanthorhoe Hübner, 1825
biriviata (Borkhausen, 1794) (Slowakei)
designata (Hufnagel, 1767) (Slowakei)
munitata (Hübner, 1809) (Schweiz)
spadicearia ([Denis \& Schiffermüller], 1775) (Slowakei)
ferrugata (Clerck, 1759) (Slowakei)
quadrifasciata (Clerck, 1759) (Slowakei)
montanata ([Denis \& Schiffermüller], 1775) (Slowakei)
fluctuata (Linnaeus, 1758) (Slowakei)
incursata (Hübner, 1813) (Schweiz)
Catarhoe Herbulot, 1951
rubidata ([Denis \& Schiffermüller], 1775) (Slowakei)
cuculata (Hufnagel, 1767) (Slowakei)
putridata (Herrich-Schäffer, 1852) (Frankreich)
Epirrhoe Hübner, 1825
hastulata (Hübner, 1813) (Slowakei)
pupillata (Thunberg, 1788) (Deutschland)
tristata (Linnaeus, 1758) (Slowakei)
alternata (Müller, 1764) (Slowakei)
rivata (Hübner, 1813) (Dänemark)
molluginata (Hübner, 1813) (Slowakei)
galiata ([Denis \& Schiffermüller], 1775) (Slowakei)
Camptogramma Stephens, 1831
bilineata (Linnaeus, 1758) (Dänemark, Slowakei)
Entephria Hübner, 1825
cyanata (Hübner, 1809) (Schweiz)
flavicinctata (Hübner, 1813) (Deutschland)
infidaria (La Harpe, 1852) (Schweiz)
caesiata (Denis \& Schiffermüller, 1775) (Slowakei)
Anticlea Stephens, 1831
badiata ([Denis \& Schiffermüller], 1775) (Slowakei)
derivata ([Denis \& Schiffermüller], 1775) (Slowakei)
Mesoleuca Hübner, 1825
albicillata (Linnaeus, 1758) (Slowakei)
Pelurga Hübner, 1825
comitata (Linnaeus, 1758) (Slowakei)
Lampropteryx Stephens, 1831
suffumata ([Denis \& Schiffermüller], 1775) (Slowakei) otregiata (Metcalfe, 1917) (Mähren)

Cosmorhoe Hübner, 1825
ocellata (Linnaeus, 1758) (Slowakei)
Nebula Bruand, 1846
salicata (Hübner, 1799) (Slowakei)
tophaceata (Denis \& Schiffermüller, 1775) (Mähren)
nebulata (Treitschke, 1828) (Österreich)
achromaria (La Harpe, 1852) (Slowakei)
Eulithis Hübner, 1821
prunata (Linnaeus, 1758) (Slowakei)
testata (Linnaeus, 1761) (Slowakei, Böhmen)
populata (Linnaeus, 1758) (Böhmen)
pyropata (Hübner, 1822) (Polen)
pyraliata ([Denis \& Schiffermüller], 1775) (Slowakei)
Ecliptopera Warren, 1894
silaceata ([Denis \& Schiffermüller], 1775) (Slowakei) capitata (Herrich-Schäffer, 1839) (Slowakei)

Chloroclysta Hübner, 1825
siterata (Hufnagel, 1767) (Slowakei)
miata (Linnaeus, 1758) (Slowakei, Deutschland)
Dysstroma Hübner, 1825
citrata (Linnaeus, 1761) (Deutschland, Slowakei)
truncata (Hufnagel, 1767) (Slowakei)
infuscata (Tengström, 1869) (Schweden)
Cidaria Treitschke, 1825
fulvata (Forster, 1771) (Slowakei)
Plemyria Hübner, 1825
rubiginata ([Denis \& Schiffermüller], 1775) (Slowakei)

## Thera Stephens, 1831

obeliscata (Hübner, 1787) (Slowakei)
variata ([Denis \& Schiffermüller], 1775) (Slowakei)
britannica (Turner, 1925) (Slowakei)
stragulata (Hübner, 1809) (Slowakei)
cognata (Thunberg, 1792) (Österreich)
juniperata (Linnaeus, 1758) (Slowakei)
cupressata (Geyer, 1831) (Frankreich)

Eustroma Hübner, 1825
reticulata (Denis \& Schiffermüller, 1775) (Slowakei)
Electrophaes Prout, 1923
corylata (Thunberg, 1792) (Slowakei)
Colostygia Hübner, 1825
aptata (Hübner, 1813) (Österreich)
austriacaria (Herrich-Schäffer, 1856) (Österreich)
tempestaria (Herrich-Schäffer, 1856) (Österreich)
multistrigaria (Haworth, 1809) (Schweiz)
Hydriomena Hübner, 1825
furcata (Thunberg, 1784) (Slowakei)
impluviata ([Denis \& Schiffermüller], 1775) (Slowakei)
ruberata (Freyer, 1831) (Österreich, Deutschland)
Horisme Hübner, 1825
vitalbata ([Denis \& Schiffermüller], 1775) (Slowakei, Österreich)
tersata ([Denis \& Schiffermüller], 1775) (Slowakei, Deutschland)
aemulata (Hübner, 1813) (Slowakei)
aquata (Hübner, 1813) (Deutschland)
corticata (Treitschke, 1835) (Österreich)
radicaria (La Harpe, 1855) (Deutschland)
Melanthia Duponchel, 1829
procellata ([Denis \& Schiffermüller], 1775) (Slowakei)
alaudaria (Freyer, 1846) (Slowakei)
Pareulype Herbulot, 1951
berberata ([Denis \& Schiffermüller], 1775) (Dänemark, Slowakei)

Spargania Guenee, 1857
luctuata ([Denis \& Schiffermüller], 1775) (Slowakei)
Rheumaptera Hübner, 1822
hastata (Linnaeus, 1758) (Slowakei)
cervinalis (Scopoli, 1763) (Slowakei)
undulata (Linnaeus, 1758) (Slowakei, Österreich)
Triphosa Stephens, 1829
dubitata (Linnaeus, 1758) (Slowakei)
sabaudiata (Duponchel, 1830) (Ö́sterreich)
Philereme Hübner, 1825
vetulata ([Denis \& Schiffermüller], 1775) (Slowakei)
transversata (Hufnagel, 1767) (Slowakei)
Euphyia Hübner, 1825
biangulata (Haworth, 1809) (Slowakei)
unangulata (Haworth, 1809) (Böhmen)
frustata (Treitschke, 1828) (Österreich)
Epirrita Hübner, 1822
dilutata ([Denis \& Schiffermüller], 1775) (Slowakei) christyi (Allen, 1906) (Slowakei)
autumnata (Borkhausen, 1794) (Slowakei)
Operóphtera Hübner, 1825
brumata (Linnaeus, 1758) (Slowakei)
fagata (Scharfenberg, 1805) (Slowakei)

Perizoma Hübner, 1825
affinitata (Stephens, 1831) (Polen)
alchemillata (Linnaeus, 1758) (Slowakei)
bydrata (Treitschke, 1829) (Schweiz)
blandiata ([Denis \& Schiffermüller], 1775) (Slowakei)
flavofasciata (Thunberg, 1792) (Slowakei)
didymata (Linnaeus, 1758) (Slowakei, Böhmen)
obsoletaria (Herrich-Schäffer, 1838) (Österreich, Schweiz)
sagittata (Fabricius, 1787) (Dänemark, Böhmen)
incultaria (Herrich-Schäffer, 1848) (Österreich)
parallelolineata (Retzius, 1783) (Österreich)
Baptria Hübner, 1825
tibiale (Esper, 1791) (Österreich)
Eupithecia Curtis, 1825
tenuiata (Hübner, 1813) (Slowakei)
inturbata (Hübner, 1817) (Schweiz, Deutschland)
haworthiata Doubleday, 1856 (Dänemark, Deutschland)
immundata (Lienig \& Zeller, 1846) (Slowakei, Deutschland)
plumbeolata (Haworth, 1809) (Deutschland)
abietaria (Goeze, 1781) (Deutschland)
analoga Djakonov, 1926 (Deutschland, Polen, Slowakei)
linariata ([Denis \& Schiffermüller], 1775) (Deutschland, Slowakei)
pulchellata Stephens, 1831 (Deutschland)
pyreneata Mabille, 1871 (Slowakei, Deutschland)
laquaearia Herrich-Schäffer, 1848 (Slowakei, Deutschland)
irriguata (Hübner, 1813) (Slowakei, Deutschland)
liguriata Millière, 1884 (Frankreich)
exiguata (Hübner, 1813) (Deutschland)
insigniata (Hübner, 1790) (Deutschland)
valerianata (Hübner, 1813) (Slowakei, Deutschland)
pygmaeata (Hübner, 1799) (Deutschland)
undata (Freyer, 1840) (Schweiz)
variostrigata Alpheraky, 1878 (Schweiz) silenata Assman, 1849 (Böhmen, Deutschland) carpophagata Staudinger, 1871 (Italien)
venosata (Fabricius, 1787) (Slowakei, Deutschland) schiefereri Bohatsch, 1893 (Italien)
silenicolata Mabille, 1866 (Italien)
alliaria Staudinger, 1870 (Slowakei)
egenaria Herrich-Schäffer, 1848 (Deutschland)
extraversaria Herrich-Schäffer, 1852 (Deutschland)
centaureata ([Denis \& Schiffermüller], 1775) (Slowakei, Deutschland)
queneeata Miller, 1862 (Österreich)
gratiosata Herrich-Schäffer, 1861 (Frankreich)
thalictrata (Püngeler, 1902) (Schweiz, Italien)
acteata Walderdorf, 1869 (Deutschland)
selinata Herrich-Schäffer, 1861 (Deutschland, Slowakei) trisignaria Herrich-Schäffer, 1848 (Slowakei, Deutschland) intricata (Zetterstedt, 1839) (Slowakei, Deutschland) veratraria Herrich-Schäffer, 1848 (Slowakei, Deutschland) cauchiata (Duponchel, 1830) (Deutschland) satyrata (Hübner, 1813) (Slowakei, Deutschland) cretaceata Packard, 1874 (Italien) pernotata Guenée, 1857 (Schweiz) absinthiata (Clerck, 1759) (Slowakei, Deutschland) goossensiata Mabille, 1869 (Dänemark, Deutschland) assimilata Doubleday, 1856 (Slowakei, Deutschland) expallidata Doubleday, 1856 (Deutschland) vulgata (Haworth, 1809) (Slowakei, Deutschland)
tripunctaria Herrich-Schäffer, 1852 (Slowakei, Deutschland)
denotata (Hübner, 1813) (Slowakei, Deutschland)
subfuscata (Haworth, 1809) (Slowakei, Deutschland)
icterata (Villers, 1789) (Deutschland)
succenturiata (Linnaeus, 1758) (Deutschland)
denticulata (Treitschke, 1828) (Deutschland)
impurata (Hübner, 1813) (Deutschland)
orphnata W. Petersen, 1909 (Slowakei, Deutschland)
subumbrata ([Denis \& Schiffermüller], 1775) (Deutschland)
semigraphata Bruand, 1851 (Deutschland)
millefoliata Rössler, 1866 (Deurschland)
santolinata Mabille, 1871 (Frankreich)
simpliciata (Haworth, 1809) (Slowakei, Deutschland)
sinuosaria Eversmann, 1848 (Slowakei, Polen, Deutschland)
druentiata Dietze, 1902 (Frankreich)
distinctaria (Herrich-Schäffer, 1848) (Slowakei, Deutschland)
graphata (Treitschke, 1828) (Schweiz, Italien, Österreich)
pimpinellata (Hübner, 1813) (Deutschland)
pauxillaria Boisduval, 1840 (Deutschland)
gelidata (Möschler, 1860) (Böhmen, Deutschland)
nanata (Hübner, 1813) (Dänemark, Deutschland)
innotata (Hufnagel, 1767) (Deutschland)
fraxinata Grewe, 1863 (Deutschland)
unedonata Mabille, 1868 (Spanien)
virgaureata Doubleday, 1861 (Slowakei, Deutschland)
abbreviata Stephens, 1831 (Deutschland)
dodoneata Guenee, 1857 (Deutschland)
pusillata ([Denis \& Schiffermüller], 1775) (Slowakei, Deutschland)
ericeata (Rambur, 1833) (Slowakei)
phoeniceata (Rambur, 1834) (Spanien)
scopariata (Rambur, 1833) (Frankreich)
oxycedrata (Rambur, 1833) (Italien)
lanceata (Hübner, 1825) (Slowakei, Deutschland)
lariciata (Freyer, 1842) (Slowakei, Deutschland)
tantillaria Boisduval, 1840 (Slowakei, Deutschland)
Gymnoscelis Mabille, 1868
rufifasciata (Haworth, 1809) (Slowakei, Deutschland)
Chloroclystis Hübner, 1825
v-ata (Haworth, 1809) (Slowakei, Deutschland)
chloerata (Mabille, 1870) (Slowakei, Deutschland, Dänemark)
rectangulata (Linnaeus, 1758) (Slowakei, Deutschland)
debiliata (Hübner, 1817) (Böhmen, Dänemark, Deutschland)

## Anticollix Prout, 1938

sparsata (Treitschke, 1828) (Dänemark, Deutschland)
Chesias Treitschke, 1825
legatella ([Denis \& Schiffermüller], 1775) (Mähren, Schweden)

## Carsia

sororiata (Linnaeus, 1758) (Schweden)
Aplocera Stephens, 1827
plagiata (Linnaeus, 1758) (Slowakei)
efformata (Guence, 1857) (Dänemark)
praeformata (Hübner, 1826) (Slowakei)

Odcais Boisduval. 1840
atraka (Linnaeus, 1758) (Deutschland)
Schistostege Hübner, 1825
treisschkei Kovács, 1957 (Ungarn)
decussata (Denis \& Schiffermüller, 1775) (Österreich)
Lithostege Hübner, 1825
griseata ([Denis \& Schiffermüller], 1775) (nach Khotko, 1977)

Disciloxia W'arren, 1895
blomeri (Curtis, 1832) (Slowakei)
Venusia Curtis, 1839
cambrica (Curtis, 1839) (Slowakei)
Euchoeca Hübner, 1823
nebulata (Scopoli, 1763) (Slowakei)
Asthena Hübner, 1825
albulata (Hufnagel, 1767) (Slowakei)
anseraria (Herrich-Schäffer, 1855) (Slowakei)
Hydrelia Hübner, 1825
flammeolaria (Hufnagel, 1767) (Slowakei)
Minoa Treitschke, 1825
murinata (Scopoli, 1763) (Slowakei)
Lobophora Curtis, 1825
halterata (Hufnagel, 1767) (Slowakei)
Trichopteryx Hübner, 1825
polycommata ([Denis \& Schiffermüller], 1775) (Slowakei)
carpinata (Borkhausen, 1794) (Slowakei)
Epilobophora Inoue, 1943
sabinata (Geyer, 1831) (Schweiz)
Nothocasis Prout, 1937
sertata (Hübner, 1817) (Slowakei)
Pterapherapteryx Curtis, 1825
sexalata (Retzius, 1783) (Slowakei)
Acasis Duponchel, 1845
appensata (Eversmann, 1842) (Slowakei)
Abraxas Leach, 1815
grossulariata (Linnaeus, 1758) (Slowakei)
Calospilos Hübner, 1825
sylvata (Scopoli, 1763) (Slowakei)
Lomaspilis Hübner, 1825
marginata (Linnaeus, 1758) (Slowakei)
Ligdia Guenéc, 1857
adustata ([Denis \& Schiffermüller], 1775) (Slowakei)
Stegania Gucnée, 1857
dilectaria (Hübner, 1790) (Slowakei, Frankreich)
Scmiothisa Hübner. 1818
notata (Linnaeus, 1758) (Slowakei)
alternaria (Hübner, 1809) (Slowakei)
signaria (Hübner, 1809) (Slowakei)
Chiasmia Hübner, 1823
clathrata (Linnaeus, 1758) (Slowakei)
glarearia (Brahm, 1791) (Slowakei)
Diastictis
artesiaria (Denis \& Schiffermüller, 1775) (Slowakei)
Isturgia Hübner, 1823
limbaria (Fabricius, 1775) (Slowakei)
Narraga Walker, 1861
fasciolaria (Hufnagel, 1767) (Slowakei)
Itame Hübner, 1823
wauaria (Linnaeus, 1758) (Slowakei)
brunneata (Thunberg, 1784) (Böhmen, Schweden)
Tephrina Guenée, 1845
arenacearia (Denis \& Schiffermüller, 1775) (Slowakei)
murinaria (Denis \& Schiffermüller, 1775) (Slowakei)
Enconista Lederer, 1853
miniosata (Duponchel, 1829) (Frankreich)
Pygmaena Boisduval, 1840
fusca (Thunberg, 1792) (Schweiz)
Cepphis Hübner, 1823
advenaria (Hübner, 1799) (Polen)
Petrophora Hübner, 1811
chlorosata (Scopoli, 1763) (Slowakei, Böhmen)
Plagodis Hübner, 1823
pulveraria (Linnaeus, 1758) (Slowakei)
dolabraria (Linnaeus, 1767) (Slowakei)
Pachycnemia Stephens, 1829
hippocastanaria (Hübner, 1799) (Holland)
Opisthograptis Hübner, 1823
luteolata (Linnaeus, 1758) (Slowakei)
Epione Duponchel, 1829
repandaria (Hufnagel, 1767) (Slowakei)
paralellaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Pseudopanthera Hübner, 1823
macularia (Linnaeus, 1758) (Slowakei)
Hypoxystis Prout, 1915
pluviaria (Fabricius, 1787) (Slowakei)
Therapis Hübner, 1823
flavicaria (Denis \& Schiffermüller, 1775) (Österreich)
Epirrhanthis Hübner, 1823
diversata (Denis \& Schiffermüller, 1775) (Böhmen)
Ennomos Treitschke, 1825
autumnaria (Werneburg, 1859) (Slowakei)
quercinaria (Hufnagel, 1767) (Slowakei) alniaria (Linnaeus, 1758) (Slowakei, Deutschland) fuscantaria (Haworth, 1809) (Österreich, Deutschland) erosaria ([Denis \& Schiffermüller], 1775) (Slowakei) quercaria (Hübner, 1813) (Italien)

Selenia Hübner, 1823
dentaria (Fabricius, 1775) (Slowakei)
lunularia (Hübner, 1788) (Slowakei, Böhmen)
tetralunaria (Hufnagel, 1767) (Slowakei)
Apeira Gistl, 1848
syringaria (Linnaeus, 1758) (Böhmen, Slowakei)
Artiora Meyrick, 1892
evonymaria (Denis \& Schiffermüller, 1775) (Slowakei)
Odontopera Stephens, 1831
bidentata (Clerck, 1759) (Slowakei)
Crocallis Treitschke, 1825
tusciaria (Borkhausen, 1793) (Deutschland)
elinguaria (Linnaeus, 1758) (Slowakei)
Ourapteryx Leach, 1814
sambucaria (Linnaeus, 1758) (Slowakei, Böhmen)
Colotois Hübner, 1823
pennaria (Linnaeus, 1761) (Slowakei)
Angerona Duponchel, 1829
prunaria (Linnaeus, 1758) (Slowakei)

## Chondrosoma

fiduciaria Anker, 1854 (Österreich)
Apocheima Hübner, 1825
hispidaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Phigalia Duponchel, 1829
pilosaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Lycia Hübner, 1825
hirtaria (Clerck, 1759) (Slowakei)
graecaria (Staudinger, 1870) (Bosnien)
isabellae (Harrison, 1914) (Böhmen)
pomonaria (Hübner, 1790) (Slowakei)
zonaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Biston Leach, 1815
strataria (Hufnagel, 1867) (Slowakei)
betularia (Linnaeus, 1758) (Slowakei)
Agriopis Hübner, 1825
leucophaearia ([Denis \& Schiffermüller], 1775) (Slowakei) bajaria ([Denis \& Schiffermüller], 1775) (Slowakei)
aurantiaria (Hübner, 1799) (Slowakei)
marginaria (Fabricius, 1776) (Slowakei)
Erannis Hübner, 1825
defoliaria (Clerck, 1759) (Slowakei)
Nychiodes Lederer, 1853
obscuraria (De Viller, 1789) (Frankreich)

Eurranthis Hübner, 1823
plumistaria (De Viller, 1789) (Frankreich)
Peribatodes Wehrli, 1943
rhomboidaria ([Denis \& Schiffermüller], 1775) (Slowakei)
umbraria (Hübner, 1809) (Italien)
secundaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Cleora Curtis, 1825
cinctaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Deileptenia Hübner, 1825
ribeata (Clerck, 1759) (Slowakei)
Alcis Curtis, 1826
repandata (Linnaeus, 1758) (Slowakei)
maculata (Staudinger, 1890) (Slowakei)
Arichanna Moore, 1868
melanaria (Linnaeus, 1758) (Slowakei, Böhmen)
Hypomecis Hübner, 1821
roboraria ([Denis \& Schiffermüller], 1775) (Slowakei)
viertlii (Bohatsch, 1883) (Slowakei)
Serraca Moore, 1887
punctinalis (Scopoli, 1763) (Slowakei)
Fagivorina Wehrli, 1943
arenaria (Hufnagel, 1767) (Slowakei)
Ascotis Hübner, 1825
selenaria (Denis \& Schiffermüller, 1775) (Slowakei)
Ectropis Hübner, 1825
crepuscularia ([Denis \& Schiffermüller], 1775) (Slowakei)
Paradarsia Warren, 1897
consonaria (Hübner, 1799) (Slowakei)
Parectropis Sato, 1980
similaria (Hufnagel, 1767) (Slowakei)
Aethalura Mc Dunnough, 1920
punctulata ([Denis \& Schiffermüller], 1775) (Slowakei)
Ematurga Lederer, 1853
atomaria (Linnaeus, 1758) (Slowakei)
Tephronia Hübner, 1825
sepiaria (Hufnagel, 1767) (Deutschland)
Odontognophos Wehrli, 1951
dumetata (Treitschke, 1827) (Slowakei)
Selidosema Hübner, 1823
brunnearia (Villers, 1789) (Böhmen)
plumaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Bupalus Leach, 1815
piniaria (Linnaeus, 1758) (Slowakei)
Cracota Hübner, 1823
lutearia (Fabricius, 1794) (Frankreich, Schweiz)
niveata (Scopoli, 1763) (Österreich)

Menophra Moore, 188 ?
nychemeraria (Geyer, 1831) (Deutschland)
Synopsia Hübner, 1825
sociariad (Hübner, 1799) (Slowakei)
Cleorodes W'arren. 1894
lichenaria (Hufnagel, 1767) (Deutschland)
Gnophos Treitschke, 1825
furuatus ([Denis \& Schiffermüller], 1775) (Slowakei) obfuscarus ([Denis \& Schiffermüller], 1775) (Österreich)
ambiguatus (Duponchel, 1838) (Österreich)
prollatus ([Denis \& Schiffermüller], 1775) (Österreich)
glaucinarius (Hübner, 1799) (Österreich)
variegarus (Duponchel, 1830) (Österreich, Schweiz)
intermedia Wehrli, 1917 (Slowakei)
dilucidaria ([Denis \& Schiffermüller], 1775) (Österreich, Schweiz)
serotinaria ([Denis \& Schiffermüller], 1775) (Österreich, Schweiz)
zelleraria (Freyer, 1836) (Österreich)
caelibaria (Herrich-Schäfer, 1852) (Österreich)
operaria (Hübner, 1813) (Österreich)
Sciadia Hübner, 1822
tenebraria (Esper, 1806) (Österreich, Schweiz)
Glacies Millière, 1874
alticolaria (Mann, 1853) (Österreich)
Siona Duponchel, 1829
lineata (Scopoli, 1763) (Slowakei)
Chariaspilates Wehrli, 1953
formosaria (Eversmann, 1837) (nach Khotko, 1977)
Aspitates Treitschke, 1825
gilvaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Dyscia Hübner, 1825
conspersaria (Fabricius, 1775) (Slowakei)
Perconia Hübner, 1823
strigillaria (Hübner, 1787) (Mähren)
Cabera Treitschke, 1825
pusaria (Linnaeus, 1758) (Slowakei)
exanthemata (Scopoli, 1763) (Slowakei)
Lomographa Hübner, 1825
bimaculata (Fabricius, 1775) (Slowakei)
temerata ([Denis \& Schiffermüller], 1775) (Slowakei)
Campaea Lamarck, 1816
margaritata (Linnaeus, 1767) (Slowakei)
honoraria ([Denis \& Schiffermüller], 1775) (Böhmen)

Hylaea Hübner, 1822
fasciaria (Linnaeus, 1758) (Slowakei)
Puengeleria Rougemont, 1903
capreolaria ([Denis \& Schiffermüller], 1775) (Slowakei)
Lignyoptera
thaumastaria Rebel, 1901 (Bosnien)
Theria Hübner, 1825
rupicapraria ([Denis \& Schiffermüller], 1775) (Slowakei)
primaria (Haworth, 1809) (Deutschland)

# FOUR NEW GENERA OF MICROVELIINAE (HETEROPTERA) FROM NEW GUINEA 


#### Abstract

Polhemus, J. T. \& D. A. Polhemus, 1994. Four new genera of Microveliinae (Heteroptera) from New Guinea. - Tijdschrift voor Entomologie 137: 57-74, figs. 1-30. [ISSN 0040-7496]. Published 15 July 1994. Four new genera of Microveliinae are described from New Guinea. These genera and their constituent species are as follows: Tanyvelia gen. n. containing type species T. missim sp. n. from Papua New Guinea; Aegilipsicola gen. n. containing type species A. rapida sp. n. from Papua New Guinea; Tarsovelia gen. n. containing type species T. alta sp. n. from Papua New Guinea, plus T. arfak sp. n. from Irian Jaya, and T. dani sp. n. from Irian Jaya; and Neusterinsifer gen. n. containing type species $N$. compacta sp. n. from Papua New Guinea, plus N. sepiksp. n. from Papua New Guinea, $N$. cyclops sp. n. from Papua New Guinea, N. nabire sp. n. from Irian Jaya, and $N$. gladius sp. n. from Irian Jaya. Habitus figures of the above new genera are provided, accompanied by illustrations of the male genital structures and distribution maps for all new species. Correspondence: Dr Dan A. Polhemus, Dept. of Natural Sciences, Bishop Museum, P.O. Box 19000-A, Honolulu, HI. 96817, U.S.A.


Key words. - Veliidae; Microveliinae; New Guinea; taxonomy; new genera; new species; keys; distribution.

The assemblages of Microveliinae occurring on the islands of the Malay Archipelago exhibit a profound shift in taxonomic composition as one progresses from west to east through the region. On the Greater Sunda Islands, Celebes and the Philippines such faunas are dominated by members of the genera Pseudovelia Hoberlandt and Microvelia Westwood, accompanied by several other more ecologically specialized and less speciose genera. On New Guinea and surrounding islands, by contrast, Pseudovelia is absent and there occurs instead a group of previously undescribed endemic genera which fill the typical Pseudovelia niches. In addition, New Guinea also supports numerous endemic species of in the genus Microvelia sensu lato, but as noted by Andersen (1982) this generic grouping is almost certainly polyphyletic. A detailed analysis on a world basis of the subgroups contained within it will be necessary before it can be certain that the Papuan and Australian species presently held in Microvelia are in fact congeneric with the Asian forms.

New Guinea thus represents a major center of diversification for the Microveliinae, with a fauna derived independently from that occurring in the Asian tropics. In the present report we describe four new genera of endemic Papuan Microveliinae, all of which are easily separable from Microvelia sensu lato on the basis of distinct apomorphies. The relationships among these genera are still unresolved, however, pending the completion of ongoing revisions of the
diverse Australian and New Guinea Microvelia faunas.

## Key to genera of New Guinea Microveliinae

1. Middle tarsi subequal in length to middle tibia (fig. 9); light markings on hemelytra restricted to basal angles; fore femur of male at least slightly modified for phoresy $\qquad$ Tarsovelia gen. n.

- Middle tarsi distinctly shorter than middle tibia (figs. 1, 5, 17); light markings on hemelytra may be restricted to basal angles, or may occur also on distal portions; fore femur may or may not be modified for phoresy (figs. 3, 6, 24-26)2

2. Claws extremely long; light hemelytral markings entirely bright greyish to light bluish pruinose; gula long, rostral cavity demarcated by strongly raised carina $\qquad$ Aegilipsicola gen. n.

- Claws relatively short; light hemelytral markings not bright pruinose; gula short, rostral cavity not demarcated by strongly raised carina

3
3. Antennae extremely long (fig. 1); comb on fore tibia 3/4 the length of the tibia (fig. 3); only micropterous morph known

Tanyvelia gen. n.

- Antennae not extremely long (fig. 17); comb on fore tibia less than $3 / 4$ length of tibia (except in Neuterinsifer compacta), if comb on fore tibia 3/4 length of tibia then male proctiger bearing elongate process and female abdominal tergites VII
and VIII forming an anal plate (see below); only aprerous and macropterous morphs known, latter with large hemelytral light markings basally, and often occurring in all closed cells $\qquad$ .. 4

4. Female abdominal tergites VII-VIII deflected ventrad forming an anal plate (fig. 23); male genitalia highly modified, bearing a long sinuate anreriorly directed sword-like process (figs. 18-22)

Neusterinsifer gen. n.

- Female abdominal tergites VII-VIII not deflected ventrad, not forming an anal plate; male genitalia may be modified, but without a long sword-like process $\qquad$ Microvelia Westwood

Tanyvelia gen. n .
(figs. 1-4, 27)

## Diagnosis

Size. - Micropterous form, length of males 2.732.91 mm , females $2.81-3.13 \mathrm{~mm}$; general body characteristics and size sexually dimorphic, males smaller, female abdomen more robust.

Colour. - Ground colour blackish brown, tinged with orange brown, without silvery pubescence; anterior pronotal lobe yellowish, pruinose, sharply demarcated from dark collar and posterior lobe. Micropterous wing pads brown, set with anterolateral light streaks.

Structural characters. - Only micropterous form known, shape elongate (fig. 1). Eyes globose, exserted but usually not beyond anterolateral pronotal angles, separated by about twice the width of an eye, appressed to anterior pronotal margin, with short ocular setae. Head declivant anteriorly, recessed into pronotum, posterior margin sloping caudo-dorsally, with usual three pairs of facial trichobothria; gular region moderately long, plainly visible, rostral cavity closed posteriorly. Rostrum reaching to middle of mesosternum, segment I short, enclosed in rostral cavity, segments I and IV subequal in length and about three times longer than II, segment III about 8 times as long as II. Antennae slender, very long, about 3/4 of body length.

Pronotum slightly raised medially, with weak median longitudinal carina; collar weakly formed, set off by a ragged row of small dark foveae, terminating under eyes laterally; anterior and posterior lobes set off by a transverse row of small foveae, evanescent medially; anterior lobe with scattered stiff erect dark setae; posterior lobe with numerous small foveae, humeri not prominent, broadly rounded posteriorly, not modified, covering metanotum. Thoracic venter not diagnostic, with weakly formed tubercles on either side of mesosternal midline on posterior margin opposing an unmodified metasternum. Metasternal scent gland opening (omphalium) small but visible,
marked by a small tubercle; scent channels prominent, curving slightly anterad to base of metacetabulae.

Abdomen without silvery setae or other adornment, except weak longitudinal pruinose line on midline of tergites IV-VII; with short paired longitudinal carinae on basal half of tergite II. Abdominal sternites set off from laterosternites by hair-free glabrous oval lacunae. Micropterous wing pads elongate.

Legs slender, long, hind legs very long; anterior femur set beneath with short light setae, unmodified in males; anterior tibia of male with a comb of minute black setae occupying $3 / 4$ of tibial length; middle femora set ventrally with $4-5$ very long slender setae; all tarsi long (fig. 3), claws moderately long; both upand down curving arolia large, evident.
Male genital segments moderately large, protruding, modified; proctiger unmodified (fig. 4); parameres small, slender symmetrical, acuminate (fig. 2). Female tergite VIII on same plane as VII, truncate posteriorly; first gonocoxae small, barely exposed, plate-like; tergite IX of both sexes triangular, protruding posteriorly.

## Type-species: Tanyvelia missim sp. n.

## Remarks

Comparative notes. - Separable from other genera of Papuan Microveliinae by the characters given in the key. Similar in general appearance and ecological habits to Baptista Distant, but distinguished by the presence of a long grasping comb that runs nearly the entire length of the male fore tibia (fig. 3), and the absence of modifications to abdominal sterna VI and VII. Among the Papuan genera, Tanyvelia appears most closely allied to Aegilipsicola, but the pronotal collar is not as pronounced, the legs, although long and slender, are slightly stouter, and the fore tibial grasping comb is shorter. The proctiger of Tanyvelia is broadly rounded distally rather than coming to a rounded point as in Aegilipsicola (compare figs. 4 and 7), and the paramere is slender and elongate rather than short and stout (compare figs. 2 and 8).
Etymology. - The generic name Tanyvelia is derived from tany- (Gr.), long, referring to the slender body shape, and Velia, the nominate genus of the family. Gender feminine.

Distribution. - New Guinea (fig. 27).

## Tanyvelia missim sp. n .

(figs. 1-4, 27)
Type material: Holotype, micropterous male: Papua New Guinea, Morobe Prov., upper Poverty Creek on Mt. Missim, nr. Wau, 1600 m., 18 Sept.


Figs. 1-4. Tanyvelia missim, gen. n., sp. n. . - 1. Male, dorsal habitus; 2. Male foreleg; 3. Male paramere; 4. Male proctiger. Figs. 5-8. Aegilipsicola rapida, gen. n., sp. n. - 5. Male, dorsal habitus; 6. Male foreleg; 7. Male proctiger; 8. Male paramere

1983, CL 1830, D. A. and J. T. Polhemus (BPBM), Paratypes: 23 micropterous males, 12 micropterous females, same data as holotype (JTPC).

## Diagnosis

Size. - Micropterous male, length 2.73-2.91 mm ( $\overline{\mathrm{x}}=2.81, \mathrm{n}=10$ ); width $0.79-0.90 \mathrm{~mm}(\overline{\mathrm{x}}=0.86, \mathrm{n}$ $=10$ ). Micropterous female, length $2 \cdot 81-3.13 \mathrm{~mm}(\overline{\mathrm{x}}$ $=2.96, \mathrm{n}=10)$; width $0.83-0.94 \mathrm{~mm},(\overline{\mathrm{x}}=0.88, \mathrm{n}=$ 10).

Colour. - Micropterous male: ground colour black, venter and connexiva tinged with brown. Head black, brown ventrally; rostrum luteous on basal three segments, piceous distally. Pronotum with anterior lobe entirely yellowish including propleura, except narrowly embrowned medially; disc and collar black. Abdomen black, tergites mostly pruinose, lighter ventrally. Legs and antennae luteous to yellow brown, distally darker.

Structural characters. - Micropterous male: head of moderate length, declivant anteriorly; length 0.47 ; width of eye/interocular space, $0.16 / 0.30$. Pronotum length:width, 0.74 : 0.81 . Abdominal tergites not shining; tergites, II-VI subequal in length (0.160.19 ), VII longer (0.30). Abdominal venter set with short appressed setae; ventrite VII with a short Vshaped depression on caudal half. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Legs unarmed.

Antennal formula, segments I : II : III : IV; 0.51 : $0.40: 0.70: 0.70$.

Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.79: 0.77: 0.30: 0.0$; of middle leg, $1.07: 1.05: 0.14: 0.28$; of hind leg, 1.16 : $1.47: 0.23: 0.30$.
First genital segment with an antero-ventrally directed tuft of stiff dark setae. Paramere small, long, triangular (fig. 2).

Micropterous female: Similar to male in most respects, but somewhat larger; connexiva vertical or reflexed over dorsum except in gravid specimens.

## Remarks

Comparative notes. - Recognized among the Pa puan Microveliinae by its elongate form with long legs and antennae (fig. 1), relatively unmodified male foreleg with a long tibial grasping comb (fig. 3), acuminate paramere (fig. 2), and elongate male proctiger (fig. 4).

Biological notes. - Upper Poverry Creek at the Mt. Missim type locality was a small, rocky headwater stream dropping steeply through a boulder-strewn bed heavily shaded by primary rain forest.

Etymology. - The name 'missim' is a noun in appo-
sition and refers to the Mt. Missim type locality. Distribution. - Eastern New Guinea (fig. 27).

## Aegilipsicola gen. n.

(figs. 5-8, 28)

## Diagnosis

Size. - Macropterous form, length of males 2.702.84 mm , females $3.16-3.31 \mathrm{~mm}$, general body characteristics not sexually dimorphic, but males slightly smaller.

Colour. - Macropterous form: ground colour blackish brown, tinged with yellowish brown; dorsally dark brown on head and ventrally, wings black, brown to black beneath; anterior pronotal lobe narrowly yellowish on either side of midline, laterally dark but covered with silvery pubescence except medially, demarcated from dark collar and posterior lobe. Hemelytra black, bearing six elongate bluishgrey pruinose streaks, with two confined within the basal cells, and four others on distal third (fig. 5)

Structural characters. - Only macropterous form known. Form elongate, slender (fig. 5); eyes globose, exserted beyond anterolateral pronotal angles, separated by about twice the width of an eye, removed from anterior pronotal margin, with short ocular setae. Head moderately declivant anteriorly, recessed into pronotum, posterior margin sloping somewhat caudo-dorsally, with usual three pairs of facial trichobothria; gular region long, plainly visible, rostral cavity closed posteriorly by a strong carinate ridge. Rostrum reaching to middle of mesosternum, segment I short, enclosed in rostral cavity, I and IV subequal in length and about three times longer than II, segment III about 8 times as long as II. Antennae very slender, very long, about $2 / 3$ of body length. Pronotum slightly raised medially, with weak median longitudinal carina marked by a line of golden pubescence; collar prominent, distinctly set off by a row of deep foveae, terminating laterally; near ventral eye margins; anterior and posterior lobes set off by a transverse row of foveae, evanescent medially; entire pronotum sparsely set with fine decumbent golden pubescence; posterior lobe with numerous obscure shallow foveae, humeri moderately prominent, almost triangular, narrowly rounded posteriorly. Thoracic venter not diagnostic, with weakly formed tubercles on either side of mesosternal midline along posterior margin opposing an unmodified metasternum. Metasternal scent gland opening (omphalium) not visible; scent channels prominent, curving anterad to base of metacetabulae.

Abdomen without silvery setae or other adornment; prominent paired longitudinal carinae present on basal half of tergite II in macropters (visible only after removal of wings). Abdominal sternites set off
from laterosternites by hair-free glabrous oval lacunae. Female connexiva expanded, plate like along tergite VII. Macropters with five closed cells in hemelytra, venation of fore and hind wings similar to figs. 312-313 in Andersen (1982) except distal cell closed; fore wings set with fine semi-recumbent stiff setae on basal half; micropterous form unknown.

Legs slender, long, hind legs very long; anterior femur set beneath with short light setae, with ventral tubercle at distal $2 / 3$ in males; anterior tibia of male with a fine line of minute black setae occupying $4 / 5$ of tibial length; femora ventrally without long setae; all tarsi long (fig. 6), claws very long; both up- and down curving arolia long, slender.

Male genital segments moderately large, protruding, modified; proctiger unmodified (fig. 7); parameres small, slender, symmetrical (fig. 8). Female tergite VIII on same plane as VII, protruding posteriorly over tergite IX; first gonocoxae small, barely exposed, plate-like; tergite IX of both sexes button-like, protruding posteriorly.

## Type-species: Aegilipsicola rapida sp. n.

## Remarks

Etymology. - The generic name Aegilipsicola is derived from Aegilips (Gr.), sheer, steep, referring to the habitat, and -cola, (L.), inhabitant. Gender feminine.

Comparative notes. - In general facies Aegilipsicola most closely resembles the Neotropical genus Aegilipsivelia, however this similarity is apparently due to convergence. Although the two genera share many similar structures adapted to life on steep hygropetric habitats, such as long legs and extremely long claws, Aegilipsivelia lacks the carinate rostral cavity on the gula, the bright pruinose hemelytral markings, and the carinate female connexiva seen in Aegilipsicola. In addition, the collar in Aegilipsivelia is set off by a sulcus rather than a row of deep foveae, and the comblike process on the male fore tibia is relatively short instead of almost as long as the tibia.

Among Papuan genera of Microveliinae Aegilipsicola is apparently most closely allied to Tanyvelia gen. n. (see discussion under Tanyvelia)

Distribution. - Eastern New Guinea (fig. 28).

## Aegilipsicola rapida sp. n.

(figs. 5-8, 28)
Type material: Holotype, macropterous male: Papua New Guinea, Morobe Prov., seeps along Mt. Kaindi road nr. Kunai Creek, above Wau, 17 Sept. 1983, CL 1823, D. A. and J. T. Polhemus (вРвм). Paratypes: 39 macropterous males, 37 macropterous females, 62 immatures, same data as holotype (JTPC);

4 macropterous males, 4 macropterous females, 2 im matures, Morobe Prov., Namie Creek, Mr. Kaindi, 17 Sept. 1983, CL 1829, J. T. \& D. A. Polhemus (JTPC); 1 macropterous female, Morobe Prov., Kunai River, 1500 m, 10 Oct. 1966, Illies (ZBSM).

## Diagnosis

Size. - Macropterous male, length $2.70-2.84 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=2.74, \mathrm{n}=10$ ); width $0.86-0.97 \mathrm{~mm}(\overline{\mathrm{x}}=0.94, \mathrm{n}$ $=10$ ). Macropterous female, length $3.06-3.31 \mathrm{~mm}(\overline{\mathrm{x}}$ $=3.16, \mathrm{n}=10)$; width $1.01-1.12 \mathrm{~mm}$, $(\overline{\mathrm{x}}=1.04, \mathrm{n}=$ 10).

Colour. - Macropterous male: ground colour black, tinged with yellowish brown ventrally. Head black; two longitudinal regions between eyes and medial furrow, posterior margin, brown. Pronotum with anterior lobe narrowly, transversely orange brown on either side of midline, extending to inner eye margins; posterior lobe uniformly black. Hemelytra blackish brown, long, extending beyond tip of abdomen. Abdomen brownish yellow ventrally, with a broad brown longitudinal stripe along sternal-laterosternal boundary. Legs yellowish, darker distally; antennae brown, segment I yellowish basally.

Structural characters. - Macropterous male: head of moderate length, declivant anteriorly; length 0.51 ; width of eye/interocular space, $0.14 / 0.30$. Pronotum long, humeri evident but not raised; length : width, $0.84: 0.91$.

Abdominal venter set with very short appressed fine setae; venter VII depressed on either side of midline; first genital segment excavated ventrally, with three small brown sclerotized denticles distally. Legs, antennae thickly clothed with short inconspicuous setae, without longer setae. Legs unarmed, except fore femur set near distal $2 / 3$ with patch of short stiff setae (fig. 6).
Antennal formula I : II : III : IV; $0.40: 0.28: 0.74$ : 0.60 .

Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.74: 0.60: 0.28: 0.0$; of middle leg, $1.00: 0.98: 0.12: 0.33$; of hind leg, 1.44 : $1.74: 0.09: 0.40$.

Paramere small, short, ovate (fig. 8).
Macropterous female: Similar to male in most respects, but somewhat larger. Connexiva vertical, strongly raised and plate-like along tergite VII, produced posteriorly along tergite VIII.

## Remarks

Comparative notes. - Easily recognized among the Papuan Microveliinae by its elongate form (fig. 5), broadly pointed proctiger (fig. 7), short and broad paramere (fig. 8), elongate fore tibial grasping comb


Figs. 9-16. Tarsovelia gen. n. - 9. Tarsovelia alta sp. n., micropterous female, dorsal habitus; 10. Tarsovelia danisp. n., male proctiger; 11-13. Male forclegs of Tarsovelia species, 11. Tarsovelia arfak sp. n.; 12. Tarsovelia dani sp. n.; 13. Tarsovelia alta sp. n.; 14-16. Male parameres of Tarsovelia species. 14. Tarsovelia dani sp. n.; 15. Tarsovelia alta sp. n.; 16. Tarsovelia arfak sp. n.
(fig. 6), and small tuft of black setae ventrally on the male fore femur (fig. 6).

Biological notes. - The type series was taken from wet seeping vertical rock faces formed by road cuts along the Mt. Kaindi road. The insects were abundant, and ran across the wet rocks and moss with great speed and agility.

Etymology. - The name 'rapida' refers to the speed with which these insects can move across vertical rock faces.

Distribution. - Eastern New Guinea (fig. 28).
Tarsovelia gen. n .
(figs. 9-16, 29)

## Diagnosis

Size. - Length of males 2.41-2.95 mm, females $2.77-3.45 \mathrm{~mm}$, general body characteristics and size sexually dimorphic, males smaller, female abdomen more robust.

Colour. - Ground colour brown to blackish brown, often blackish above, without silvery pubescence; winged forms dorsally black, wings black, brown to black beneath. Anterior pronotal lobe and collar usually entirely yellowish to orange brown, only in T. arfak dark laterally.

Structural characters. - Form robust (fig. 9); eyes globose, exserted but usually not beyond anterolateral pronotal angles, separated by at least twice the width of an eye, appressed to anterior pronotal margin, with long ocular setae. Head declivant anteriorly, slightly recessed into pronotum, with usual three pairs of facial trichobothria; gular region moderately long, plainly visible, rostral cavity closed posteriorly. Rostrum reaching almost to metasternum, segment I short, enclosed in rostral caviry, segments I and IV subequal in length and about three times longer than II, segment III about 8 times as long as II. Antennae slender, long, about $1 / 2$ of body length.

Pronotum of micropterous form slightly raised medially; collar weakly formed, set off by a ragged row of small dark foveae, terminating under eyes laterally; anterior and posterior lobes set off by a transverse row of small foveae in a weak transverse sulcus, evanescent medially; anterior lobe set with stiff erect setae; posterior lobe with numerous shallow depressions, humeri not prominent, broadly rounded posteriorly, not modified, covering metanotum. Pronotum of alate form longer, rounded posterior margin extending farther caudad, almost triangular, humeri more prominent. Thoracic venter not diagnostic, with weakly formed tubercles on either side of mesosternal midline on posterior margin opposing an unmodified metasternum. Metasternal scent gland opening (om-
phalium) not evident; scent channels prominent, angled slightly anterad to base of metacetabulae.

Abdomen without silvery setae or other adornment, except weak longitudinal pruinose line on midline of tergites IV-VII; prominent paired longitudinal carinae on tergite II of macropters (visible only after removal of wings), absent in micropterous form. Abdominal sternites set off from laterosternites by hair-free round or oval lacunae. Micropters with elongate white wing pads, often infuscated distally; macropters with four closed cells in dark hemelytra with white region basally; venation of fore and hind wings similar to figs. 312-313 in Andersen (1982); female fore wing Sc often terminating in a triangular thickened region, resulting in a slight to pronunced sinuosity of the wing margin; fore wings set with long dark stiff erect setae on basal half, much more pronounced in females.
Legs slender; anterior femur set beneath with short light setae, sometimes slightly modified in males (figs. 11-13); anterior tibia with a distal comb of many stiff setae; middle and anterior femora set ventrally with numerous long slender setae; middle tibia set with an evenly spaced row of about 10 erect setae beginning at basal third, very long basally, decreasing in length distally; middle tarsi extremely long (fig. 9); claws short; both up- and down curving arolia evident, but very slender.
Male genital segments small, not protruding, unmodified; proctiger unmodified; parameres small, slender symmetrical (figs. 14-16). Female tergite VIII on same plane as VII, truncate posteriorly; first gonocoxae small, barely exposed, plate-like; tergite IX small, button-like, protruding posteriorly.

Type-species: Tarsovelia alta sp. n.

## Remarks

Comparative notes. - Tarsovelia is similar in general facies to Microvelia, but is easily separated from this genus and all other Microveliinae by the elongate middle tarsi (fig. 9), which are adapted for rowing rather than running. In addition, the light colored markings on the hemelytra of Tarsovelia are restricted to the basal angles, plus occasionally a weak medial fascia, whereas in the type-species of Microvelia (M. pulchella Westwood) the hemelytra also bear ovate light spots in every closed cell, plus a large bright spot in the open distal cell.

Biological notes. - Tarsovelia species are found along the margins of flowing pools on rocky mountain streams. They are swift and agile skaters if disturbed, and will fly readily if captured in a net.

Etymology. - The generic name Tarsovelia is derived from tarsos (Gr.), referring to the extremely long
slender middle tarsi, and Velia, the nominate genus of the family. Gender feminine.

Distribution. - New Guinea (fig. 29).
Key to the species of Tarsovelia
Males

1. Fore femur with patch of short, stiff hairs on ventral surface (figs. 11, 12)

- Fore femur lacking patch of short, stiff hairs on ventral surface (fig. 13) ................. T. alta sp. n.

2. Fore femur with patch of short, stiff hairs near middle of ventral margin (fig. 11)
T. $\operatorname{arfak} \mathrm{sp} . \mathrm{n}$.

- Fore femur with patch of short, stiff hairs near base of ventral margin (fig. 12) ..... T. dani sp. n.


## Tarsovelia alta sp. n .

(figs. 9, 13, 15, 29)
Type material: Holotype, micropterous male: Papua New Guinea, Morobe Prov., bog nr. Mt. Kaindi summit, above Wau, 17 Oct. 1983, CL 1825, D. A. and J. T. Polhemus (BPBM). - Paratypes: $13 \mathrm{mi}-$ cropterous males, 14 micropterous females, 16 immatures, same data as holotype (ZMUC, JTPC, BPBM).
Additional material examined: 2 macropterous females, Papua New Guinea, Morobe Prov., Eddie Creek, 2050 m, 11 Oct. 1966, Illies leg. (ZBSM, JTPC); 1 macropterous female, Morobe Prov., Poverty Creek, Mt. Missim, 18 Sept. 1983, CL 1830, 1600 m, J. T. \& D. A. Polhemus (JTPC); 9 macropterous males, 6 macropterous females, 2 immatures, Western Highlands Prov., rocky stream 17 km N. of Mt. Hagen on Baiyer River road, 6 and 8 Sept. 1983, CL 1780, D. A. and J. T. Polhemus (JTPC).

## Diagnosis

Size. - Micropterous male, length 2.41-2.48 mm ( $\overline{\mathrm{x}}=2.44, \mathrm{n}=10$ ); width $0.83-0.89 \mathrm{~mm}(\overline{\mathrm{x}}=0.85, \mathrm{n}$ $=10$ ). Micropterous female, length $2.81-2.88 \mathrm{~mm}(\overline{\mathrm{x}}$ $=2.84, \mathrm{n}=4$ ); width $0.94-1.01 \mathrm{~mm}$, $(\bar{x}=0.99, n=4)$.
Macropterous male, length $2.77-2.95 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=$ $2.87, \mathrm{n}=9$ ); width $1.10-1.15 \mathrm{~mm}(\overline{\mathrm{x}}=1.14, \mathrm{n}=9)$. Macropterous female, length $3.31-3.45 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=$ 3.36, $n=6)$; width $1.19-1.33 \mathrm{~mm},(\overline{\mathrm{x}}=1.28, \mathrm{n}=6)$.

Colour. - Micropterous male: ground colour orange brown, venter slightly lighter. Head dark orange brown, lighter beneath; rostrum luteous medially, piceous distally. Pronotum with anterior transverse band orange brown, extending onto propleura; disc dark brown, mortled with lighter brown. Abdomen orange brown, tergal margins darker, lighter ventrally. Distal segements of legs, antennae brown, basal segments lighter.

Structural characters. - Micropterous male: head
of moderate length, declivant anteriorly; length 0.37 ; width of eye/interocular space, $0.14 / 0.30$. Pronotum long, covering metanotum, humeri evident but not raised; length:width, $0.84: 0.86$. Abdominal tergites not shining; lengths of tergites, II : 0.14; III : 0.16; IV-VI : 0.19; VII : 0.26 .

Abdominal venter not modified, set with short appressed setae. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Legs unarmed.

Antennal formula I : II : III : IV; $0.29: 0.33: 0.37$ : 0.42 .
Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.65: 0.63: 0.28: 0.0$; of middle leg, $0.98: 0.93: 0.42: 0.35$; of hind leg, 0.86 : $1.00: 0.19: 0.29$.
Paramere small, long, slender (fig. 15).
Micropterous female: Similar to male in most respects, but somewhat larger; connexiva reflexed over dorsum except in gravid specimens.

Macropterous male: Similar to micropterous form in most respects. Pronotum with anterior lobe entirely orange brown, extending onto propleura; posterior lobe uniformly dark. Hemelytra blackish brown, basally entirely white except slight infuscation of Sc, white region extending beyond tip of pronotum.

Macropterous female: Very similar to macropterous male, except lateral hemelytral margin slightly sinuate at distal thickening of Sc. Posterior lobe of pronotum with two $(1+1)$ tufts of erect stiff black setae on either side of midline at about middle. Connexival margin vertical.

## Remarks

Comparative notes. - T. alta sp. n. may be separated from T. dani sp. n. and T. arfaksp. n. by the unmodified male fore femur (fig. 13), brownish coloration in wingless forms, male paramere shape (fig.15), and length of the second segment of the hind tarsi, which is about 1.5 times as long as the first (versus only slightly longer in the other two species). In $T$. arfak sp. n. the anterior femora are dark distally, and the middle and posterior femora plus all the tibiae and tarsi are dark brown to black, while in T. alta sp. n . and $T$. dani sp. n. all femora are largely light brown, and the tibiae and tarsi are dark brown but not black. In the macropters of T. alta sp. n. and $T$. arfak sp. n . the entire basal angle of the hemelytra is white (including the veins, although these are occasionally slightly darkened), whereas in T. dani sp. n. the veins are dark and the light coloring is often a sordid white.

We originally considered the micropterous and macropterous specimens at hand from Papua New Guinea to represent different taxa. However with two
additional new species now available from Irian Jaya that show marked differences from the Papua New Guinea material, we have now concluded that the latter represent but a single, albeit somewhat variable, species. We have thus included all of the specimens from eastern New Guinea under T. alta sp. n., but the macropters are not named as paratypes. Macropterous specimens from Eddie Creek and Poverty Creek (not measured) have very long hemelytra, considerably exceeding the tip of the abdomen, and they are lighter in color than specimens from Mt. Hagen. Measurements of leg and antennal segments are not helpful, as they appear to vary randomly between populations, thus we defer the final decision about the specific distinctness of these populations from Papua New Guinea until more material, hopefully including both micropters and macropters in the same sample, is available.

Biological notes. - Near Mt. Hagen T. alta was found in a plunge pool below a small cascade, with individuals hiding in the crevices of a sheer rock wall. When disturbed the insects would skate rapidly over the water near the wall until disappearing into their hiding places once more. By contrast, the Mt. Kaindi specimens were taken from beneath a large, partly submerged $\log$ in a bog pool, where violent flushing was required to dislodge them. So far this species has been found only at altitudes above 1200 meters.

Etymology. - The name 'altd' refers to the high elevation at which this species was taken.

Distribution. - Eastern New Guinea (fig. 29).

## Tarsovelia arfaksp. n.

(figs. 11, 16, 29)
Type material: Holotype, macropterous male: Indonesia, Irian Jaya Prov., Saumarin River, nr. Warkomi, Arfak Mountains, 42 km S. of Manokwari, 90 m , water temp. $25^{\circ} \mathrm{C}$, 18 Oct. 1991, CL 2647, D. A. and J. T. Polhemus (USNM). Paratypes: 67 macropterous males, 48 macropterous females, 6 immatures, same data as holotype (JTPC).

## Diagnosis

Size. - Macropterous male, length $2.45-2.77 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ $=2.62, \mathrm{n}=10)$; width $1.01-1.12 \mathrm{~mm}(\overline{\mathrm{x}}=1.06, \mathrm{n}=$ 10). Macropterous female, length $2.81-3.09 \mathrm{~mm}(\overline{\mathrm{x}}=$ $2.98, \mathrm{n}=10)$; width $1.15-1.19 \mathrm{~mm},(\overline{\mathrm{x}}=1.17, \mathrm{n}=10)$.

Colour. - Macropterous male: ground colour black, tinged with brown. Head black. Pronotum with anterior lobe and collar medially orange brown, extending to inner eye margins; posterior lobe uniformly black. Hemelytra blackish brown, long, extending beyond tip of abdomen, basally with broad elongate rectangular white region along outer margin,
extending beyond tip of pronotum. Abdomen black, ventrally tinged with brown along connexival margins and caudally. Legs, antennae brown, fore femur yellowish except distally, mid and hind femora yellowish ventrally.

Structural characteristics. - Macropterous male: head of moderate length, declivant anteriorly; length 0.40 ; width of eye/interocular space, $0.12 / 0.33$. Pronotum long, humeri evident but not raised; length:width, $0.86: 1.00$.

Abdominal venter not modified, set with short appressed setae. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae; all tibia, middle and hind tarsi set with rows of long setae dorsally, in addition to long ventral setae. Legs unarmed, except fore femur ventrally slightly excavate basally, set near middle with patch of short stiff setae (fig. 11).

Antennal formula I : II : III : IV; $0.28: 0.33: 0.35$ : 0.40 .

Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.65: 0.51: 0.28: 0.0$; of middle leg, $0.88: 0.81: 0.49: 0.35$; of hind leg, 0.81 : $0.95: 0.23: 0.28$.

Paramere small, long, slender (fig. 16).
Macropterous female: Similar to male in most respects, but somewhat larger; hemelytra with lateral margin sinuate, thickened and set with long dense black setae at distal Sc. Connexiva vertical along tergites II-IV, slightly reflexed over tergites V-VII.

## Remarks

Comparative notes. - Recognized by the patch of short stiff black setae on the ventral margin of the male fore femur (fig. 11), and the shape of the male paramere (fig. 16). For additional comparative notes see key, and discussion under T. alta sp. n.

Biological notes. - The Saumarin River at the type locality was a moderate sized stream dropping steeply to the sea off the flanks of the Arfak Mountains through a bed of boulders, cobbles and sand, heavily shaded by primary rain forest. The stream profile was punctuated by numerous small waterfalls, and T. arfak was most abundant on the plunge pools below these falls.

Etymology. - The name ' $a r f a k$ ' is a noun in apposition and refers to the Arfak Mountain type area.

Distribution. - Western New Guinea (Vogelkop Peninsula) (fig. 29).

## Tarsovelia danisp. n.

(figs. 10, 12, 14, 29)
Type material: Holotype, macropterous male: Indonesia, Irian Jaya Prov., swift rocky stream in

upper Pass Valley, 52 km NE of Wamena, 2015 m .,' water temp. $14^{\circ} \mathrm{C}, 23$ Sept. 1991, CL 2614, D. A. and J. T. Polhemus (USNM). - Paratypes: 61 macropterous males, 26 macropterous females, 28 immatures, same data as holotype (JTPC).

## Diagnosis

Size. - Macropterous male, length $2.55-2.95 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=2.78, \mathrm{n}=10$ ); width $1.01-1.15 \mathrm{~mm}(\overline{\mathrm{x}}=1.09$, n $=10$ ). Macropterous female, length $2.99-3.45 \mathrm{~mm}(\overline{\mathrm{x}}$ $=3.12, \mathrm{n}=10$ ); width $1.15-1.37 \mathrm{~mm}$, $(\overline{\mathrm{x}}=1.22, \mathrm{n}=$ 10).

Colour. - Macropterous male: Ground colour orange brown, marked with black. Head blackish brown dorsally, ventrally lighter. Pronotum with anterior lobe and collar orange brown, extending onto propleura, collar marked only laterally, demarcated by a row of dark foveae; posterior lobe uniformly black. Hemelytra blackish brown, basally with narrow elongate white stripe along costal margin, extending beyond tip of pronotum, plus a shorter sordid white region basally along pronotum; with elongate grayish brown streak in middle, barely visible in some specimens; length variable, shortest reaching barely onto tergite VII, longest almost covering genital segments. Abdomen orange brown. Legs, antennae brown, fore femur yellowish except distally, mid and hind femora, tibia yellowish ventrally.
Structural characteristics. - Macropterous male: head of moderate length, declivant anteriorly; length 0.38 ; width of eye/interocular space, $0.14 / 0.30$. Pronotum long, humeri evident but not raised; length:width, $1.00: 1.12$.

Abdominal venter not modified, set with short appressed setae. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Legs unarmed, except fore femur flattened ventrally, curved, basally with a tuft of stiff setae (fig. 12).
Antennal formula I : II : III : IV; $0.30: 0.35: 0.42$ : 0.42 .
Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.74: 0.72: 0.30: 0.0$; of middle leg, $1.12: 1.02: 0.49: 0.35$; of hind leg, $0.98:$ $1.07: 0.26: 0.33$.
Paramere small, long, slender (fig. 14).
Macropterous female: Similar to male in most respects, but somewhat larger. Hemelytra length variable, shorter than in male, shortest not reaching tergite VII, longest reaching caudal margin of tergite

VII; with lateral margin weakly sinuate, narrowly thickened and set with stiff setae at distal Sc. Connexiva vertical along tergites II-III, IV in transition, strongly reflexed over tergites V-VII.

## Remarks

Comparative notes. - Recognized by the small tuft of black setae basally on the ventral surface of the weakly bowed male fore femur (fig. 12), and the shape of the male paramere (fig. 14). For additional comparative notes see key, and discussion under T. alta sp. n.

Biological notes. - The type series of T. dani was taken from pools along a very swift, cold mountain stream in a limestone bed bordered by Nothofagus and Rhododendron forest. The insects were most common along the pool margins, skating rapidly across the water if disturbed and flying readily if taken in a net.
Etymology. - The name 'dani' is a noun in apposition and refers to the native Dani people who inhabit the type area.
Distribution. - Central New Guinea (fig. 29).

## Neusterinsifer gen. n .

(figs. 17-26, 30)

## Diagnosis

Size. - Form stout (fig. 17), length of males 1.872.48 mm , females $1.91-2.77 \mathrm{~mm}$, general body characteristics and size sexually dimorphic, males smaller, female abdomen more robust.
Colour. - Ground color black, tinged with orange brown, heavily marked with silvery pubescence; anterior pronotal lobe orange brown on posterior half, with streak extending from near midline to middle of eyes, weakly demarcated from collar and posterior lobe.
Structural characteristics. - Both apterous and macropterous forms known. Eyes globose, exserted but usually not beyond anterolateral pronotal angles, separated by about twice the width of an eye, appressed to anterior pronotal margin, with moderately long ocular setae. Head strongly declivant anteriorly, recessed into pronotum, posterior margin almost vertical, with usual three pairs of facial trichobothria; gular region short, not visible, rostral cavity open posteriorly, with raised carinate bucculae. Rostrum reaching to middle of mesosternum, segment I short, enclosed in rostral cavity, I and IV subequal in length

Figs. 17-26. Neusterinsifer gen. n. - 17. Neusterinsifer compactus sp. n., micropterous male, dorsal habitus; 18-22. Processes arising from male proctigers of Neusterinsifer species, 18. Neusterinsifer compactus sp. n.; 19. Neusterinsifer cyclops sp. n.; 20. Neusterinsifer sepik sp. n.; 21. Neusterinsifer nabire sp. n.; 22. Neusterinsifer gladius sp. n.; 23. Neusterinsifer compactus sp. n., posterior view of female abdomen showing anal plate formed by tergites VII and VIII. 24-26. Male forelegs of Neusterinsifer species: 24. Neusterinsifer sepik sp. n.; 25. Neusterinsifer gladius sp. n.; 26. Neusterinsifer compactus sp. n.
and about two times longer than II, segment III about four times as long as II. Antennae slender, very long, about $2 / 3$ of body length.

Pronotum almost flat, without median carina; collar very weakly formed, set off by a ragged row of small dark foveae, terminating under eyes laterally; anterior and posterior lobes set off by a transverse row of foveae; anterior lobe thickly set with appressed silvery setae except triangular median area; posterior lobe with numerous small foveae, humeri not formed, almost straight posteriorly, not modified, covering mesonotum. Metanotum almost as long as pronotum. Thoracic venter not diagnostic, with weakly formed tubercles on either side of mesosternal midline on posterior margin opposing an unmodified metasternum. Metasternal scent gland opening (omphalium) not visible, but marked by a tiny tubercle; scent channels prominent, curving slightly anterad to base of metacetabulae.

Abdomen with silvery setae laterally on tergites II, VII, VIII, covering most of tergites II-VI in females, less extensive in males; prominent paired longitudinal carinae on tergite II of macropters (visible only after removal of wings), absent in micropterous form. Abdominal sternites set off from laterosternites by small hair-free glabrous oval lacunae; laterosternite VII of both sexes set with small round tuft of stiff posterolaterally directed setae, near connexival margin at basal $2 / 5$ in males, much closer to base in females.

Macropterous forms with four closed cells in dark hemelytra, occasionally with fifth distal cell closed; always with white elongate streak basally in outer basal cell, usually with faint to pronounced light areas in other cells; venation of fore and hind wings similar to figs. 312-313 in Andersen (1982).

Legs moderately stout, moderately long; anterior femur set beneath with short light setae, modified in males of some species, with a patch of stiff setae or denticles ventrally on basal half; anterior tibia of male with a comb of minute black setae occupying $2 / 5$ to 1/2 of tibial length; all femora set ventrally with 7-12 long slender setae; all tarsi moderately long, claws moderately long; both up- and down curving arolia long, slender.

Male abdominal ventrites VI, VII modified; genital segments very large, protruding, highly modified, proctiger bearing a large sword-like appendage arising from anterior margin, directed antero-laterally beneath abdomen (figs. 18-22); segment IX, proctiger rotated by about $90^{\circ}$; parameres vestigial or absent. Female tergite VIII on same plane as VII basally, distally sharply deflected ventrad closing abdomen posteriorly except for protruding tip of proctiger (fig. 23); first gonocoxae hidden.

Type-species: - Neusterinsifer compacta sp. n.

## Remarks

Comparative nores. - Very similar in general facies to Microvelia (fig. 17), but separated by sword-like projection on the male proctiger (figs. 18-22), the small tuft of stiff setae on laterotergite VII, and other characters as given in the generic key. Macropters differ from Microvelia and most other genera of Microveliinae in having a well defined long slender white streak basally in the outer cell (with only ill defined light areas in all other cells), and lacking the distal white spot that is characteristic of Microvelia and other closely related genera. The wing markings are similar to those of Tenagovelia Kirkaldy from Africa, but species in this latter genus are much larger and possess much longer legs.

Biological notes. - Species of Neusterinsifer are most commonly encountered in sheltered areas along the margins of stream pools, or on the surfaces of water filled potholes in bedrock exposures adjacent to flowing sections of streams.

Etymology. - The generic name Neusterinsifer is derived from neuster (Gr.), swimmer, and ensifer (L.), sword bearer. Gender masculine.

Distribution. - New Guinea (fig. 30).

## Key to species of Neusterinsifer

1. Male process on proctiger bifurcate, short (fig. 21). Female abdomen strongly narrowed posteriorly, boat shaped $\qquad$ . nabire sp. n.

- Male process on proctiger not bifurcate, longer (figs. 18-20, 22). Female abdomen not strongly narrowed posteriorly, not boat shaped $\qquad$

2. Small species, male 1.87 mm , female 1.91 mm long. Male process on proctiger relatively short, broad, narrowly triangular distally (fig. 19
cyclops sp. n.

- Larger species, male at least 2.19 mm , female at least 2.27 mm long. Male process on proctiger long, relatively narrow, not triangular distally (figs. 18, 20, 22)

3. Relatively small species, males $2.19-2.27 \mathrm{~mm}, \mathrm{fe}-$ males $2.27-2.34 \mathrm{~mm}$ long. Process of male proctiger long, broad, expanded medially and distally (fig. 22); male fore femur with a row of 5-10 black spinules basally (fig. 25). Female connexival segment VII forming a small acute angle at level of caudal margin of tergite VII ...... gladius sp. n.

- Larger species, males $2.27-2.48 \mathrm{~mm}$, females $2.37-2.77 \mathrm{~mm}$ long. Process of male proctiger long, narrow or broad, not expanded medially and distally (figs. 18, 20); male fore femur without a basal row of 5-10 black spinules, instead with either a basal row of many short stiff black setae, or unarmed. Caudal extreme of female con
nexival segment VII either forming an acute ờr obtuse angle, but without a small acute angle at level of caudal margin of tergite VII $\qquad$ 4

4. Process of male proctiger long, slender, sharp distally (fig. 18); male fore femur strongly bowed, bearing small black denticles basally (fig. 26). Caudal extreme of female connexival segment VII forming an obtuse angle ..... compactus sp. n.

- Process of male proctiger long, broad (fig. 20); male fore femur not bowed, with a patch of black spinules basally (fig. 24). Caudal extreme of female connexival segment VII slightly produced posteriorly, forming an acute angle ... sepik sp. n.


## Neusterinsifer compactus sp. n.

(figs. 17, 18, 23, 26, 30)
Type material: Holotype, apterous male: Papua New Guinea, Western Highlands Prov., Baiyer River, 53 km N. of Mt. Hagen, 6 and 8 Sept. 1983, D. A. and J. T. Polhemus (BPBM). - Paratypes: 19 apterous males, 12 apterous females, same data as holotype (JTPC); 1 apterous male, 3 apterous females, Western Highlands Prov., Baiyer River Bird Sanctuary, 8 Sept. 1983, CL 1792, J. T. \& D. A. Polhemus (JTPC); 1 apterous male, Morobe Prov., Bamboo Creek, nr. Wau, 16 Sept. 1983, CL 1817, J. T. \& D. A. Polhemus (JTPC); 4 apterous males, 1 apterous female, Morobe Prov., trib. to Bumbu River, N. of Lae, 20 Sept. 1983, CL 1836, J. T. \& D. A. Polhemus (JTPC).

## Diagnosis

Size. - Apterous male, length $2.37-2.48 \mathrm{~mm}(\overline{\mathrm{x}}=$ 2.43, $\mathrm{n}=10$ ); width $1.33-1.40 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=1.39, \mathrm{n}=$ 10). Apterous female, length $2.63-2.77 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=$ 2.70, $\mathrm{n}=10$ ); width $1.40-1.55 \mathrm{~mm},(\overline{\mathrm{x}}=1.50, \mathrm{n}=$ 10).

Colour. - Apterous male: Ground colour blackish brown, venter slightly lighter, entire dorsum, laterotergites covered with fine appressed golden pubescence. Head black, orange brown beneath; rostrum fuscous medially, piceous distally. Pronotum with anterior transverse orange brown band barely interrupted medially; disc blackish brown. Abdomen blackish brown, lighter ventrally, connexiva margined with orange brown. Antennae brown; anterior legs yellowish, distally darkened; middle, hind legs dark brown, basally lighter; coxae, trochanters luteous.
Structural characters. - Apterous male: head of moderate length, declivant anteriorly, with weak impressed median line; length 0.40 ; width of eye/interocular space, $0.15 / 0.30$. Pronotum long, covering metanotum, thickly set with obscure foveae, humeri depressed; length:width, 0.40 : 1.00. Metanotum length:width, $0.33: 1.00$. Abdominal tergites not
shining; tergites II-VI subequal in length ( $0.14-0.16$ ), VII longer (0.28).
Abdominal venter set with short appressed setae, and scattered longer setae caudally; venter VI posteriorly set with two $(1+1)$ tufts of dense dark erect setae on either side of midline; venter VII broadly depressed medially, carinate along posterior margin. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Middle and hind legs unarmed; fore femur strongly bowed, bearing a patch of black denticles basally, fore tibia with grasping comb extending $3 / 4$ the length of the tibia (fig. 26).
Antennal formula I : II : III : IV; $0.35: 0.28: 0.40$ $: 0.51$.

Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.65: 0.60: 0.28: 0.0$; of middle leg, $0.86: 0.86: 0.16: 0.30$; of hind leg, 0.93 : $1.07: 0.17: 0.30$.
Proctiger produced anteriorly into a long slender sinuate ensiform process (fig. 18). Parameres vestigial or absent.

Micropterous female: Similar to male in most respects, but somewhat larger; connexiva vertical, incurved and slightly refexed at caudal extreme, which forms an obtuse angle.

## Remarks

Comparative notes. - A relatively large species for the genus, recognized by the short, blunt process on the male proctiger (fig. 18), and the strongly bowed male foreleg with a patch of black denticles basally on the ventral surface (fig. 26). Similar in size and general facies to $N$. sepik sp. n., but this latter species possesses an unbowed foreleg with a large patch of black denticles (fig. 24), and a much larger and more pointed process on the male proctiger (fig. 20).

Biological notes. - The Baiyer River at the type locality was a swift, deep river flowing in a rocky bed with wet mossy banks, and partially shaded by disturbed upland rain forest. $N$. compactus was taken here skating on the surface of water filled potholes in an exposure of igneous rock along the river margin.
Etymology. - The name compactus (L.) refers to the shape of this species.

Distribution. - Eastern New Guinea (fig. 30).

## Neusterinsifer cyclops sp. n.

(figs. 19, 30)
Type material: Holotype, apterous male: INDONESIA, Irian Jaya Prov., small rocky stream and spring above Sentani, Cyclops Mountains, $325 \mathrm{~m}, 25$ Sept. 1991, CL 2618, J. T. \& D. A. Polhemus (JTPC, to be placed in USNM when more material is available).

## Diagnosis

Size. - Apterous male, length, $\overline{\mathrm{x}}=1.87, \mathrm{n}=1$; width, $\bar{x}=0.79, n=1$. Apterous female, length, $\bar{x}=$ 1.91, $\mathrm{n}=1$ : width, $\overline{\mathrm{x}}=0.90, \mathrm{n}=1$.

Colour. - Apterous male: Ground colour deep brown, marked with brown, venter slightly lighter, entire dorsum, laterotergites covered with fine appressed golden pubescence; entire dorsum, pleura, sides of abdomen covered with moderate length erect setae. Head brown, orange brown beneath; rostrum fuscous medially, piceous distally. Pronotum with anterior transverse orange brown band embrowned medially; disc blackish brown. Abdomen blackish brown, lighter ventrally, connexiva margined with orange brown. Antennae brown, base of segment I lighrer; legs yellowish, distally darkened; femora darker posteriorly; coxae, trochanters luteous.

Structural characters. - Apterous male: head of moderate length, declivant anteriorly, with weak impressed median line; length 0.33 ; width of eye/interocular space, $0.12 / 0.30$. Pronotum long, covering metanotum, thickly set with obscure foveae, humeri depressed; length:width, 0.28 : 0.79. Metanotum length:width, $0.26: 0.79$. Abdominal tergites not shining; tergites II-VI subequal in length (0.12-0.14), VII longer (0.26).

Abdominal venter set with short appressed setae; venter VI posteriorly set with two ( $1+1$ ) tufts of dense dark erect setae on either side of midline, between them a V-shaped depression open posteriorly; venter VII broadly depressed medially, sharply excavated on posterior half, carinate along anterior margin of excavation. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Middle and hind legs unarmed; foreleg lacking denticles or spinules, tibial comb extending for only $1 / 2$ the length of the tibia.

Antennal formula I : II : III : IV; $0.21: 0.19: 0.33$ : 0.40 .

Proportions of legs as follows: Femur, tibia, tarsal 1 , tarsal 2 of fore leg, $0.51: 0.44: 0.19: 0.0$; of middle leg, $0.63: 0.60: 0.12: 0.19$; of hind leg, 0.74 : $0.86: 0.12: 0.26$.

Proctiger produced anteriorly into a broad ensiform process (fig. 19). Parameres vestigial or absent.

Apterous female: Similar to male in most respects, but somewhat larger; connexiva vertical; caudal extreme gently curving ventrad.

Macropterous forms: unknown.

## Remarks

Comparative notes. - Separable from all other Neusterinsifer species by its small size, and short, pointed process on the male proctiger (fig. 19).

Biological notes. - The type of $N$. cyclops was taken from a small, clear, spring fed stream at the margin of the remaining primary rain forest on the slopes of the Cyclops Mountains above Sentani. The stream flowed in a bed of metamorphic rocks intermixed with coarse grey sand, and was partially shaded by trees felled for garden clearings.

Etymology. - The name cyclops, a noun in apposition, refers to the mountain range where the species was found.

Distribution. - Northern New Guinea (Cyclops Mountains) (fig. 30).

## Neusterinsifer gladius sp. n.

(fig. 22, 25, 30)
Type material: Holotype, apterous male: INDONEsia, Irian Jaya Prov., Nabire River and swift tributary, 5 km E. of Nabire, $60 \mathrm{~m}, 14$ Oct. 1991, CL 2642, J. T. \& D. A. Polhemus (USNM). - Paratypes: 2 apterous males, 2 apterous females, same data as holotype (JTPC).

## Diagnosis

Size. - Apterous male, length 2.19-2.27 mm ( $\overline{\mathrm{x}}=$ $2.24, \mathrm{n}=3$ ); width $0.90-0.94 \mathrm{~mm}(\overline{\mathrm{x}}=0.91, \mathrm{n}=3)$. Apterous female, length $2.27-2.34 \mathrm{~mm}(\overline{\mathrm{x}}=2.30, \mathrm{n}=$ 2); width $1.01-1.04 \mathrm{~mm}$, $(\bar{x}=1.03, \mathrm{n}=2)$.

Colour. - Apterous male: Ground colour blackish brown, venter slightly lighter, entire dorsum, laterotergites covered with fine appressed golden pubescence; entire body except thoracic venter covered with long erect setae. Head black, orange brown benearh; rostrum fuscous medially, piceous distally. Pronotum with anterior transverse orange brown band sometimes interrupted medially; disc blackish brown. Abdomen blackish brown, lighter ventrally, connexiva margined with orange brown. Antennae brown; anterior femora luteous, distally darkened; anterior tibia, tarsi brown, darker distally; middle, hind legs brown dorsally, ventrally yellowish, femora basally lighter; coxae, trochanters luteous, tinged with fuscous.

Structural characters.- Apterous male: head of moderate length, declivant anteriorly, with weak impressed median line; length 0.35 ; width of eye/interocular space, $0.15 / 0.28$. Pronotum long, covering metanotum, thickly set with obscure foveae, humeri depressed; length:width, 0.37 : 0.86. Metanotum length:width, $0.28: 0.86$. Abdominal tergites not shining; tergites II-VI subequal in length ( $0.12-0.14$ ), VII longer (0.28).

Abdominal venter set with short appressed setae; venter VI posteriorly set with two ( $1+1$ ) widely separated tufts of dense dark erect setae on either side of

Fig. 27. Distribution of Tanyvelia missim, gen. n., sp. п.


Fig. 28. Distribution of Aegilipsicola rapida, gen. n., sp. n.

midline, with shallow hair free sulcus between, extending the length of ventrites V-VI; venter VII broadly and strongly depressed medially. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Middle and hind legs unarmed; fore tibia bearing a row of 5-10 black spinules basally; fore femur with grasping comb extending $1 / 2$ the length of the tibia (fig. 25).
Antennal formula I : II : III : IV; $0.35: 0.28: 0.35$ : 0.47 .
Proportions of legs as follows: Femur, tibia, tarsal 1, tarsal 2 of fore leg, $0.60: 0.51: 0.28: 0.0$; of middle leg, $0.79: 0.74: 0.16: 0.30$; of hind leg, 0.84 : $1.02: 0.16: 0.30$.
Proctiger produced anteriorly into a long broad sinuate ensiform process, expanded medially and distally (fig. 22). Parameres vestigial or absent.
Apterous female: Similar to male in most respects, but somewhat larger; connexiva vertical; caudal extreme of VII forming an acute angle; VIII gently curving ventrad.
Macropterous forms: Unknown.

## Remarks

Comparative notes. - Similar in size to $N$. nabire
sp. n., but easily separable by the elongate process on the male proctiger (fig. 22), the patch of stiff, black spinules basally on the ventral surface of the male fore femur (fig. 25), and the relatively unmodified female abdomen which is not laterally compressed or boat-shaped.

Biological notes. - The Nabire River at the type locality was a swift, clear, moderate sized stream flowing in a rocky bed, and shaded by primary lowland rain forest. The type series of $N$. gladius was taken downstream of a limestone waterfall, where the banks of the river included numerous boulders and bedrock exposures. The insects were taken from the surfaces of water filled potholes and on still pools amid the rocks.
Etymology. - The name gladius (L.), refers to the ensiform process of the male proctiger.

Distribution. - Western New Guinea (fig. 30).

## Neusterinsifer nabire sp. n.

(figs. 21, 30)
Type material: Holotype, apterous male: INDONESIA, Irian Jaya Prov., Nabire River and swift tributary, 5 km E. of Nabire, 60 m , 14 Oct. 1991, CL 2642, J. T. \& D. A. Polhemus (USNM). - Paratypes: 98 apterous males, 13 macropterous males, 72 apterous fe-
males, 35 macropterous females, 32 immatures, same data as holotype (ITPC).

## Diagnosis

Size. - Apterous male, length 2.01-2.09 mm ( $\overline{\mathrm{x}}=$ $2.04, \mathrm{n}=10$ ); width $0.79-0.83 \mathrm{~mm}(\overline{\mathrm{x}}=0.81, \mathrm{n}=$ 10). Apterous female, length $2.19-2.41 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=$ $2.27, \mathrm{n}=10$ ); width $0.79-0.90 \mathrm{~mm}(\overline{\mathrm{x}}=0.83, \mathrm{n}=$ 10). Macropterous male, length $2.23-2.34 \mathrm{~mm}(\bar{x}=$ $2.29, \mathrm{n}=6)$; width $1.04-1.12 \mathrm{~mm}(\overline{\mathrm{x}}=1.10, \mathrm{n}=6)$. Macropterous female, length $2.41-2.55 \mathrm{~mm}$ ( $\overline{\mathrm{x}}=$ $2.46, \mathrm{n}=6)$; width $1.08-1.15 \mathrm{~mm},(\overline{\mathrm{x}}=1.12, \mathrm{n}=6)$.

Colour. - Apterous male: Ground colour blackish brown, venter slighty lighter, entire dorsum, laterotergites covered with fine appressed inconspicuous pubescence; dorsum, body laterally covered with long erect setae. Head black, orange brown beneath; rostrum fuscous medially, piceous distally. Pronotum with anterior transverse orange brown band broadly interrupted medially; disc blackish brown. Abdomen blackish brown, slightly lighter ventrally, connexiva margined with orange brown. Antennae brown; anterior femora yellowish, distally darkened, tibia brown; middle, hind legs dark brown to piceous, femora basally lighter; coxae, trochanters luteous, tinged with fuscous.

Structural characteristics. - Apterous male: head of moderate length, declivant anteriorly, with weak impressed median line; length 0.33 ; width of eye/interocular space, $0.14 / 0.28$. Pronotum long, covering metanotum, thickly set with obscure foveae, humeri depressed; length:width, 0.37 : 0.81. Metanotum length:width, $0.28: 0.79$. Abdominal tergites not shining; tergites II-VI subequal in length ( $0.12-0.16$ ), VII longer (0.30).

Abdominal venter set with short appressed setae; venter V posteriorly set with two $(1+1)$ tufts of dense dark erect setae on either side of midline; venter VI with caudal margin medially produced, set with a large quadrate ruft of dense dark erect serae; venter VII excavated medially along posterior margin. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Middle and hind legs unarmed; fore femur lacking denticles or spinules; fore tibia with grasping comb extending 1/2 the length of the tibia.

Antennal formula I : II : III : IV; $0.30: 0.21: 0.28$ $: 0.44$.

Proportions of legs as follows: Femur, tibia, tarsal 1, tarsal 2 of fore leg, $0.51: 0.47: 0.21: 0.0$; of middle leg, $0.70: 0.63: 0.12: 0.23$; of hind leg, 0.74 : $0.93: 0.16: 0.28$.

Proctiger produced anteriorly into a bifurcate process (fig. 21). Parameres vestigial or absent.

Apterous femalc: Similar to male in most respects,
but somewhat larger; abdomen narrowing posteriorly. Connexiva vertical, touching at caudal extreme; caudally rounded; thickly set along posterior margin with long stout dark setae; tuft of stiff posterolaterally directed setae near connexival margin of laterosternite VII of females pronounced, more so than in other species.

Macropterous male: Similar to apterous male in many respects, but broader, humeri more pronounced. Hemelytra always with light streaks in each of the two basal cells, often with light areas in all closed cells.

Macropterous female: Similar to apterous female in many respects, but broader, humeri more pronounced. Hemelytra with light areas in each of the four basal closed cells.

## Remarks

Comparative notes. - Easily recognized by the bifurcate process on the male proctiger (fig. 21) and the laterally compressed, boat-shaped abdomen in females. This species is similar in size to $N$. gladius sp. n., and larger than $N$. cyclops sp. n., but smaller than either $N$. compactus sp. n. or $N$. sepik sp. n.

Biological notes. - See discussion under N. gladius sp. n.

Etymology. - The name 'nabire', a noun in apposition, refers to the Nabire River type locality.

Distribution. - Northern New Guinea (fig. 30).

## Neusterinsifer sepik sp. n.

(figs. 20, 24, 30)
Type material: Holotype, apterous male: Papua New Guinea, East Sepik Prov., Yemogu Creek, 2 km W. Tring, 12 Sept. 1983, CL 1805, J. T. \& D. A. Polhemus (BPBM). - Paratypes: 11 apterous males, 6 apterous females, same data as holotype (JTPC); 1 apterous male, 4 apterous females, 1 macropterous female, East Sepik Prov., Mandi Creek, E. of Wewak, 10 Sept. 1983, CL 1797, J. T. \& D. A. Polhemus (JTPC); 4 apterous males, 1 apterous female, East Sepik Prov., Brandi Creek, nr. Wewak, 10 Sept. 1983, CL 1796, J. T. \& D. A. Polhemus (JTPC).

## Diagnosis

Size. - Apterous male, length $2.27-2.41 \mathrm{~mm}(\overline{\mathrm{x}}=$ $2.35, \mathrm{n}=10$ ); width $0.94-1.01 \mathrm{~mm}(\overline{\mathrm{x}}=0.99, \mathrm{n}=$ 10). Apterous female, length $2.37-2.63 \mathrm{~mm}(\overline{\mathrm{x}}=$ $2.48, \mathrm{n}=10)$; width $0.97-1.15 \mathrm{~mm},(\overline{\mathrm{x}}=1.04, \mathrm{n}=$ 10). Macropterous female, length, $\overline{\mathrm{x}}=2.62, \mathrm{n}=1$ ); width, $\overline{\mathrm{x}}=1.15, \mathrm{n}=1$ )

Colour. - Apterous male: Ground colour blackish brown, venter slightly lighter, entire dorsum and la-

Fig. 29. Distribution of species of Tarsovelia gen. n. - Square $=$ Tarsovelia dani sp. n.; Circles = Tarsovelia alta sp. n.; Triangle $=$ Tarsovelia arfak sp. n.


Figure 30. Distribution of species of Neusterinsifer gen. n. - Square = Neusterinsifer sepik sp. n.; Circles $=$ Neusterinsifer compactus sp. n.; Triangle = Neusterinsifer cyclops sp. n.; Diamond $=$ Neusterinsifer nabire sp. n.; Half filled square $=$ Neusterinsifer gladius sp. n.
terotergites covered with fine appressed golden pubescence; entire body except thoracic venter covered with long erect setae. Head black, orange brown beneath; rostrum fuscous medially, piceous distally. Pronotum with anterior transverse orange brown band barely interrupted medially; disc blackish brown. Abdomen blackish brown, lighter ventrally, connexiva margined with orange brown. Antennae brown; anterior legs yellowish, distally darkened; middle, hind legs dark brown to piceous dorsally, ventrally yellowish, femora basally lighter; coxae, trochanters luteous, tinged with fuscous.

Structural characteristics. - Apterous male: head of moderate length, with weak impressed median line; length 0.27 ; width of eye/interocular space, $0.14 / 0.30$. Pronotum long, covering metanotum, thickly set with obscure foveae, humeri depressed; length:width, $0.37: 0.95$. Metanotum length:width, $0.28: 0.95$. Abdominal tergites not shining; tergites II-VI subequal in length (0.14-0.16), VII longer (0.30).

Abdominal venter set with short appressed setae; venter $V$ I posteriorly set with two $(1+1)$ tufts of dense dark erect setae on either side of midline, with hair free sulcus between; venter VII broadly depressed me-
dially, weakly carinate along posterior margin. Legs, antennae thickly clothed with short to moderate length setae, with scattered longer setae. Middle and hind legs unarmed; fore femur bearing an elongate patch of stiff erect black spinules basally; fore tibia with a swelling distally bearing 5-7 erect short stiff setae (fig. 24).

Antennal formula I : II : III : IV; $0.33: 0.26: 0.37$ : 0.47 .

Proportions of legs as follows: femur, tibia, tarsal 1, tarsal 2 of fore leg, $0.65: 0.53: 0.28: 0.0$; of middle leg, $0.84: 0.79: 0.16: 0.28$; of hind leg, $0.93: 1.07$ : 0.19: 0.28.

Proctiger produced anteriorly into a long broad sinuate ensiform process (fig. 20). Parameres vestigial or absent.

Apterous female: Similar to male in most respects, but somewhat larger; connexiva vertical; caudal extreme forming an acute angle.

Macropterous female: Similar to apterous female in many respects, but broader, humeri more pronounced. Hemelytra with light areas in each of the four basal closed cells.

## Remarks

Comparative notes. - Recognized by the large, elongate process on the male proctiger that comes to a rounded point (fig. 20), and the elongate patch of black denticles on the inner surface of the male foreleg (fig. 24).

Biological notes. - Yemogu Creek at the type locality was a slow flowing stream running through a rock bed, and containing numerous large pools shaded by slightly disturbed primary rain forest. $N$. sepik was found here in sheltered areas along the margins of the stream pools, and in similar habitats at Brandi River and Mandi Creek, two lowland rainforest streams entering the sea east of Wewak.

Erymology. - The name sepik, a noun in apposition, refers to the district of Papua New Guinea from which the type series was taken.

Distribution. - Northern New Guinea (fig. 30).

## Acknowledgements

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Holorypes of all new species described herein are deposited in the Bishop Museum, Honolulu (BPBM), or the U.S. National Museum of Natural History, Washington, D. C. (USNM); paratypes are held in the J. T. Polhemus collection, Englewood, Colorado (JTPC), and where numbers permit will be distributed to the institutions noted above as well as the Museum Zoologicum Bogoriense, Bogor, Indonesia (MZB), and the Universitets Zoologisk Museum, Copenhagen (ZMUC). We also thank Dr. Martin Baehr of the Zoogische Sammlung des Bayersichen Staates, Munich (ZBSM) for allowing us to examine specimens held under his care.

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## References

Andersen, N. M. 1982. The semiaquatic bugs (Hemiptera, Gerromorpha). Phylogeny, adaptations, biogeography and classification. - Scandinavian Science Press, Klampenborg, Denmark, Entomonograph 3, 455 pp.
Andersen, N. M. 1989. The Old World Microveliinae (Hemiptera: Veliidae). II. Three new species of Baptista Distant and a new genus from the Oriental region. Entomologica Scandinavica, 19:363-380.

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# ADULTS AND LARVAE OF TWO PTECTICUS LOEW FROM PENINSULAR MALAYSIA (DIPTERA, STRATIOMYIDAE) 


#### Abstract

Rozkošný, R. \& D. Kovac, 1994. Adults and larvae of two Ptecticus Loew from Peninsular Malaysia, (Diptera, Stratiomyidae). - Tijdschrift voor Entomologie 137: 75-86, figs. 1-43. [ISSN 0040-7496]. Published 15 July 1994. Descriptions and redescriptions of larvae and adults of $P$. malayensis sp. n. and $P$. longipennis (Wiedemann) are given, based on material from Malaysia. The phylogenetic relationships of the two species and the systematic position of the genus based on larval and adult characters are briefly discussed, and biological notes on larvae are provided. The diagnostic characters of the larvae and imagines, including the male genitalia, are illustrated. A generic key to the larvae of the subfamily Sarginae is given. Correspondence: R. Rozkošný, Department of Environmental Studies, Faculty of Science, Masaryk University, Kotlăřská 2, 61137 Brno, Czech Republic. Key words. - Peninsular Malaysia; Stratiomyidae; larvae, key; Ptecticus, new species, life history.


The genus Ptecticus Loew belongs to the subfamily Sarginae and differs in the adult stage from the closely related genus Sargus Fabricius by the distinctly prominent pedicel on the inner side of the antenna. According to the recent catalogue by James (1975), 31 valid species have been described from the Oriental region but many of them are poorly known. The male genitalia as the most important criterion for species discrimination, for example, have not yet been examined in most Oriental species and the taxonomic status of many species needs a re-evaluation.

The junior author, while studying insects on bamboo (Gigantochloa scortechinii Gamble) in Peninsular Malaysia, found larvae of Stratiomyidae living inside bamboo internodes. Following our study on Camptopteromyia fractipennis de Meijere (Rozkošný \& Kovac 1991) the present paper represents the second modern contribution to the taxonomy, morphology and biology of the Oriental Stratiomyidae.

Two species are treated here, both reared from larvae found under semi-natural conditions. The first of them, P. longipennis (Wiedemann), is a well distinguishable species, but we decided on re-description in order to include important diagnostic characters not mentioned in the original description, including the male genitalia. The second species is described here as new; it differs from all known Oriental Ptecticus species. Larvae and puparia of both species are described for the first time, and their descriptions enable us to propose generic characters for Ptecticus in the larval stage.

## Ptecticus longipennis (Wiedemann, 1824)

(figs. 1-10, 26-32)

## Diagnosis

An easily recognizable, chiefly blue black and yellow species from the cingulatus-group with conspicuously long, and apically darkened wings.

Measurements: Body $8.2-11.5 \mathrm{~mm}$, wing 10.8 13.0 mm .

Male. - Head hemispherical, deeply concave posteriorly, bare eyes touching on frons (figs. 1-2). Upper part of frons elongate, triangular, subshining black and sparsely covered with fairly long brownish hairs. Ocellar triangle relatively narrow and low, barely visible in lateral aspect (fig. 2). Lower frons and face protuberant, white to pale yellow, white frontal callus slender, nose-shaped. Face with dense yellow hairs. Antennae yellow, arista brown, both basal segments yellow haired. Pedicel with prominent projection on inner side. Flagellum wider than long, about as long as pedicel. Postocular margin with a complete fringe of erect hairs. Proboscis pale yellow, somewhat swollen but relatively slender, with two-segmented labelliae.

Thorax bright yellow in ground colour, with extensive dark pattern. Mesonotum chiefly shining bluishblack, leaving large anterior corners and postalar calli yellow. Scutellum dark, with margin broadly brownish or yellow. Pleura yellow, with large brownish to bluish-black spots on greater part of anepisternum,


Figs. 1-6. Ptecticus longipennis, adult characters. - 1, male head, dorsal view; 2, the same, lateral view; 3, female head, dorsal view; 4, dorsal part of male terminalia; 5 , aedeagal complex; 6 , ventral part of male terminalia.
lower half of katepisternum and anterior parts of anepimeron and katepimeron. Laterotergite shining brown and mediotergite behind scutellum completely bluish-black. Narrow subnotopleural stripe, upper margin of katepisternum, and, especially, propleuron pale, often whitish yellow.

Wings slightly greyish infuscate, with yellow pterostigma and darkened apical part reaching almost half way berween apex and discal cell. Dense microtrichia absent in wing base, alula, subcostal cell and greater part of posterior cubital cell. Anterior crossvein situated just at beginning of $R_{3.5}, M_{3}$ shortened, not reaching wing margin. Narrow squamae white to yellow, pale yellow haired. Halteres yellow, with darkened knob.

Legs including coxae chiefly yellow, only hind pair partly darkened. All femora distinctly thickened in basal half. Hind coxa and femur yellow as in fore and middle legs, hind tibia and narrow base of basitarsus dark brown to black. Rest of basitarsus and all following tarsal segments white, with whitish hairs at least on dorsal side.

Abdomen shining bluish-brown to black, with yellow transverse stripes and pleural membrane. Transverse stripes broad on the posterior, and narrow on the anterior margin of each tergum of segments 2 4, apical terga completely black. Transverse stripes sometimes reduced to broad yellow sidemarkings on posterior corners of segments. Male genitalia (figs. 46) rotated by $90^{\circ}$ along longitudinal axis, so that epandrium is in right lateral position. Apart from this, hypopygium symmetrical. Epandrium almost round and compact, tightly connected with paraprocts posteroventrally. Epiproct of proctiger narrow, bearing slender, rodlike cerci. Synsternum deeply emarginate, with slender and bipartite medial process shielding short, symmetrical, partly reduced aedeagal complex from ventral side.

Female. - Eyes separated by shining black frontal stripe being anteriorly somewhat broader than ocellar triangle (fig. 3). White frontal callus bulbose, wider than in male. Scutellum dark, its posterior margin slightly yellowish. Dark pleural spots as in male but
partly reduced on katepisternum. Abdomen shining bluish black, with transverse yellow stripes at posterior margins of terga 2-4 extending to anterior margins of following terga. Yellow stripes about half as wide as dark stripes between them, somewhat broader than in male. Dark pattern of hind legs as in male. Cerci with two segments, slender and relatively long.

Larva (and puparium). - Subcylindrical, with prominent head, slightly arched lateral margins of body segments and distinctly angulate though generally
rounded anal segment. Cuticle with usual mosaic appearance, some cells somewhat larger and darker, forming almost complete double rows at boundary of body segments and groups in sublateral position. General colour yellowish brown, with diffuse brown pattern consisting of broad, longitudinal, undulate bands. Anal segment yellow, with large anterodorsal brownish spot. Ventral side paler, especially in median third where brownish spots are almost indistinct. Head (figs. 7-8, 26-27) moderately flat, subconical. Vestiges of mandibular-maxillary complexes about as


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Figs. 7-10. Ptecticus longipennis, larval characters. - 7, head and first thoracic segment, dorsal view (A, antenna; Ad, anterodorsal setae; Cf, clypeofrontal setae; D, dorsal setae; Bl, dorsolateral setae; E, eye prominence; L, lateral setae; Lb, labral setae; VL3, posterior ventrolateral setae); 8, the same, ventral view (SI, sublabral setae; V, ventral setae; VL1-3, ventrolateral setace; V3, posterior ventral setae); 9, last abdominal segments, dorsal view ( Sa, subapical setae); 10, the same, ventral view (Ap, apical setae; as, anal slit; L1 + L2, lateral setae; sp, sternal patch).


Figs 11-19. Ptecticus malayensis sp. n., adult characters. - 11, male head, dorsal view; 12, the same, lateral view; 13, mesonorum patrern. dnrcal view: 14. male rerminalia, lateral view; 15 , female abdomen, dorsal view; 16 , antenna, inner side; 17 , doral part of male terminalia; 18, aedeagal complex; 19, ventral part of male terminalia.
long as labrum in dorsal view, low and rounded antennae placed anteriorly, their apical segment about as high as broad at base. Eyes relatively little prominent, posterior part of head with two rounded protuberances on each side.

Three thoracic segments about same shape as abdominal segments $1-7$, first thoracic segment with lateral vestiges of prothoracic spiracles. Thoracic segment 3 and abdominal segments $1-7$ with small, round, dorsolateral, probably non-functional larval spiracles. In puparia, abdominal segments $2-5$ with very inconspicuous, short, rodlike pupal respiratory horns near lateral margin. Very slender and long midsternal patch on abdominal segment 6 consisting of 112-114 pale cells (figs. 10, 31). Rounded and much smaller sternal patch with 18-19 cells usually at posterior margin of abdominal segment 5 (fig. 32). Anal segment (figs. 9-10, 28) distinctly emarginate between insertions of marginal setae, with a pair of low protuberances distally. Opening of spiracular chamber with undulate posterior margin and short pennate setae (figs. 29-30). Anal slit on ventral side with somewhat swollen marginal lips, boundered proximally by usual transverse fold. Two low symmetrical protuberances developed around insertions of posterior ventral setae.

General scheme of chaetotaxy as in other known genera of Sarginae (cf. Rozkošný 1982). All setae on head short, labral setae not longer than others, spinelike sublabral setae well developed. Anterior clypeofrontal setae inserted almost at level of eye prominences. Setae on thoracic and abdominal segments arranged in transverse rows (3 pairs of dorsal and 2-3 pairs of ventral setae) and lateral groups ( $2-4$ setae). Thoracic segment 1 , on each half, with 2 anterodorsal, 3 dorsal, 1 dorsolateral and 1 ventrolateral seta. All 3 thoracic segments only with 2 pairs of ventral setae. Lateral group of setae on thoracic segments consisting of 2 setae only ( 1 dorsolateral and 1 ventrolateral).

On lateral side of abdominal segments, 4 setae ( 1 dorsolateral, 2 lateral and 1 ventrolateral) on abdominal segments 1 and $6-7$, but only 2 setae (probably 1 ventrolateral and 1 lateral) on abdominal segments 2 5. All marginal setae on anal segment of about same length, virtually not longer than setae on preceding abdominal segments. Five pairs of ventral setae on anal segment well visible, two distal pairs inserted closely together (fig. 10). Especially dorsal and ventral setae on body segments distinctly pubescent. Length 9.3-10.6 mm, maximum width $2.6-3.0 \mathrm{~mm}$ (based on 4 puparial exuviae).

Material examined. - Peninsular Malaysia, University of Malaya Field Studies Centre, Ulu Gombak, Selangor $\left(3^{\circ} 20^{\prime} \mathrm{N}, 101^{\circ} 45^{\prime} \mathrm{E}\right.$, altitude 250 m$)$, $1 \delta^{\text {º }}$ from 10 May, $2 \delta^{\star}$
from 12 May and 19 from 16 June 1991, all specimens ex puparia, D. Kovac leg. Material deposited in the Forschungsinstitut Senckenberg, Frankfurt (smfD) (2 $\mathbf{\delta}^{\circ}$ ), and in the Faculty of Science, Masaryk University, Brno (MUB) ( $10^{\circ}$ and 1 iq).

## Remarks

Distribution. - India, Malaya, Sumatra, Java, Borneo, Philippines.

Discussion. - According to Brunetti (1923) this slender species with long legs and wings belongs to the cingulatus species group. It differs from the other species of this group by the darkened wing tip, a diagnostic character shared only with $P$. sumatranus Enderlein. The latter species, however, possesses the normal, not elongate wings and its mesonotum is reddish yellow, not dark with a metallic shine. Of the few Ptecticus species in which the male genitalia have been illustrated, only P. okinawensis Ôuchi may belong to the same group, as shown by the compact epandrium and deeply emarginate synsternum (cf. Nagatomi 1975).

## Ptecticus malayensis sp.n.

(figs. 11-25, 33-39)
Type material. - Holotype, male: Peninsular Malaysia, University of Malaya Field Studies Centre, Ulu Gombak, Selangor ( $3^{\circ} 20^{\prime} \mathrm{N}, 101^{\circ} 45^{\prime} \mathrm{E}$, altitude 250 m ), ex puparium 15 November 1991, D. Kovac leg., SmFD. - Paratypes 3 すै 29: Same locality, 30 reared 15 November, 1 if 23
 mub.

## Diagnosis

A species of the cingulatus group with longitudinally striped mesonotum and a conspicuously spindleshaped abdomen in both sexes.

Measurements: Body 11.2-12.5 mm, wing 11.812.2 mm .

Male (holotype). - Head hemispherical, with conspicuously protuberant white frontal callus (figs. 1112). Eyes separated by yellow, sparsely black-haired frontal stripe being in its narrowest point about as wide as distance between outer margins of posterior ocelli. Postocular area narrowly protuberant in dorsal view. Postocular hairs fine and moderately long, in several rows. Antennae ochrous yellow, scape and pedicel black-haired. Pedicel with elongate process on inner side (fig. 16), annulation of flagellum barely visible. Arista in subapical position, bare. Bi-partite labellae of proboscis distinctly swollen, white in ground colour and with short white hairs. Posterior part of head deeply concave.

Thorax yellow in ground colour, with extensive


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Fig. 20-25. I'tecticus malaycusis sp.n., larval characters. - 20, head and first thoracic segment, dorsal view; 21, the same, ventrial view: 22 last abdominal segments, dorsal view; 23 , the same, ventral view; 24 , scheme of setae on lateral wall of 6 th abdominal segment ( I ), dorsolateral seta; st, larval spiracle; VI, ventrolateral seta); 25, scheme of setae on lateral wall of abdominal segment 3 (ph, pupal respiratory horn; st, larval spiracle).
brown whlackish pattern. Mesonotum (fig. 13) with 3 broad black stripes reaching its posterior margin. Pleura predominantly yellow with several brownish
to black spots in central part of anepisternum, anterior side and posterior corner of katepisternum, pasterior half of katepimeron and anterior third of latero-
tergite. Also posterior margin of mediotergite broadly brownish. Thoracic pubescence short and mostly adpressed, yellow on pleura and predominantly brown to black on mesonotum and scutellum.
Wings shaded with grey, veins along anterior margin including discal cell yellow, other veins brown except for bases. Anterior cross-vein placed at basis of radial fork $R_{3.5,}, M_{3}$ not reaching wing margin. Dense microtrichia covering virtually the whole wing membrane, only wing base, alula and posterior cubital cell mostly bare. Squamae very narrow and inconspicuous, with brownish margin and yellow hairs. Halteres with yellow stem and blackish knob.
Legs yellow with extensive blackish pattern on hind pair. Both anterior pairs mainly yellow, including coxae, only apical half of mid-femora somewhat brownish. Hind coxae brownish on anterior side, hind femur mainly dark except for both ends. Also hind tibia chiefly brown but this darkening more pronounced along dorsal side and in mid-ring. Hind tarsi black but basal two thirds of basitarsus and last tarsal segment pale yellow.
Abdomen spindle-shaped, broadest behind middle, yellow with brownish black transverse bands on terga (fig. 15). First tergum yellow with a transverse brown spot in middle and following terga with relatively broad posterior margin and narrow anterior margin yellow. Venter yellow but more brownish in apical half. Genitalia (figs. 14, 17-19) brown, distinctly asymmetrical, although dorsal part only slightly so. Cerci pointed, with tips oriented ventrally (figs. 14, 17). Synsternum strongly asymmetrical in proximal half, dististyli almost hoodlike in ventral view (fig. 19). Aedeagal complex almost $S$-shaped in lateral aspect (fig. 14), right paramere conspicuously reduced (fig. 18).

Female. - As male but with broader wings and somewhat wider abdomen in apical half. Frontal stripe only slightly broader than in male, particularly in anterior half, white frontal callus distinctly bipartite and lower, antennae relatively larger. Colours of thorax, legs and abdomen virtually the same, only darkening of hind legs more extensive, leaving only bases of femora and knees yellow. Hind tarsi coloured as in male. Terminalia dark brown, cerci with two narrow black segments (fig.15).

Larva (and puparium). - Larvae on average stouter and longer than those of $P$. longipennis, often predominantly reddish brown, though dark pattern consisting of dark undulate longitudinal bands, especially in puparia, mostly distinct. Integument rough and sclerotised, with usual mosaic surface structure. Some cuticular cells forming inconspicuous pattern as in preceding species. Head (figs. 20-21, 33-34) promi-
nent, strongly sclerotised, partly retracted into first thoracic segment, distinctly longer than wide and conspicuously flattened, rather subtriangular in dorsal view. Antennae very small and low, distinctly closer to eye prominences than in $P$. longipennis, apical segment barely prominent. Eyes conspicuously protuberant, posterolateral tubercles less distinct, not doubled as in P. longipennis.
Tracheal system as in larva of $P$. longipennis, including vestiges of larval spiracles and very small lateral pupal respiratory horns on abdominal segments $2-5$. Sternal patch in middle of abdominal segment 6 drop-like to elongate oval, consisting of 90-116 cells (fig 23). Similar but much smaller and rounded sternal patch at posterior margin of sternum 5 formed by 17-20 cells. Anal segment (figs. 22-23, 35) almost round posteriorly, with a small incision in middle of distal margin. Opening of posterior spiracular chamber with arched and finely fringed posterior margin (figs. 36-37). Anal slit distinctly emarginate, somewhat longer than one third of anal segment.

General scheme of chaetotaxy as in P. longipennis although setae relatively shorter and more pubescent, especially in marginal setae of anal segment. Anterior clypeofrontal setae inserted almost between eye prominence and tip of labrum (fig. 20). Configuration of setae on lateral wall of abdominal segments and ventral side of anal segment identical with those of preceding species and probably confirming generic characters in this way: 4 setae on lateral wall of abdominal segments 1 and 6-7, 2 setae on abdominal segments 2-5 (figs. 24-25) and 5 pairs of ventral setae on anal segment present (fig. 23). Length 10.0-13.4 mm , maximum width $3.6-5.0 \mathrm{~mm}$ (based on 2 larvae, 5 puparia and 3 puparial exuviae).

## Discussion

The new species presumably belongs to the cingulatus species group as defined by Brunetti (1923). Three species of this group have dark longitudinal stripes on the thorax: P. brevipennis Rondani, P. intensivus Brunetti and $P$. kambangensis de Meijere. The first was redescribed by van der Wulp (1885) and Brunetti (1923). It is, in contrast to $P$. malayensis sp. n ., characterized by the shining blue black frontal stripe and by much narrower and shorter stripes on the mesonotum, the black tibiae and bases of the basitarsi of the hind leg. $P$. intensivus is, according to the original description by Brunetti (1927), closely related to $P$. brevipennis and like that species should have a black frontal stripe. Moreover, it also differs in the coloration of the leg (the fore tarsi are partly black, only the basal third of the hind femur is darkened and the hind tarsi are completely black). P. kambangensis distinctly differs by the pleura being shining blackish



brown with a yellow band, and by the hind basitarsus, which is completely whitish as is the basis of the following tarsal segment.

From the other Oriental Ptecticus species only $P$. tricolor van der Wulp displays the longitudinally striped mesonotum. Brunetti (1923) considered this species to be an intermediate form with uncertain relationships. According to the original description and illustration (van der Wulp in de Meijere 1904) P. tricolor differs from the new species by the colouring of the legs (hind tibia and tarsi completely black) and abdomen (segments 2-4 entirely red).

The colour of the leg and partly also the male genitalia of the new species resemble the East Palaearctic $P$. sinchangensis Ouchi redescribed and figured recently by Nagatomi (1975). However, this species does not have a striped mesonotum, and its male genitalia, including the well developed aedeagal complex, are symmetrical.

## Biological notes

The larvae of Ptecticus longipennis were found in felled bamboo culms whose internodes - now lying in the horizontal position on the ground - had been sawn in such a way that the upper half formed a hinged lid that could easily be opened for inspection and than closed again. Two lateral slits, about 2 mm wide, provided access to the interior. The internodes were partly filled with water, and the larvae stayed in the moist part above the water surface (fig. 43). They appeared about two months after the culms had been felled. During the six-months study period larvae of Ptecticus longipennis were found only in three out of over one hundred internodes prepared in the described manner; moreover, these three internodes were adjacent to each other, i.e. the eggs were probably laid by the same female. This indicates that bamboo internodes are only an occasional habitat for the larvae of Ptecticus longipennis.

Ptecticus malayensis, in contrast, was found in bamboo more frequently. The larvae of this species were restricted to bamboo shoots that had died recently (fig. 40). Judging from the traces of insect attack found in the internodes, death of the shoots was probably caused by a weevil, Cyrtotrachelus sp. (Coleoptera, Curculionidae), adults of which had been observed on bamboo shoots. The dead shoots were 2.5 m to 6 m high. At the time of felling the lower internodes were still green while the upper part
had turned dark brown and begun to decay. In the lower part the rotten section (c. 1 m from the top) in some cases a small hole (diameter c. $1-2 \mathrm{~mm}$ ) was found which led into the interior of the culm. Beginning at this hole a groove in the inner wall led upwards, piercing the node septa (usually c. 10-15) and gradually becoming wider and deeper (fig. 42). At several points along this depression there were holes through the internode wall to the outside which were covered by the culm sheets and therefore hidden from outside view. Just below the shoot tip there was no segmentation into internodes. In this section, which was moist and gave off a putrid smell, the culm sheets had been penetrated with a circular opening $c$. 1.5 cm in diameter. In one case 2 grooves and circular openings were found in the same shoot.
Apparently the Cyrtotrachelus larva had worked itself up all the way to the nutritious apical meristem from the small entrance hole bored by the female, feeding and grooving along the way. After feasting on the apical meristem the fully grown larva then probably broke to the outside tu pupate in the soil, as described by Witt (in Beeson 1941: 209) for Cyrtotrachelus longipes F.

Ptecticus malayensis probably uses a fresh Cyrtotrachelus emergence hole to enter the bamboo shoot. Although all shoots attacked by Cyrtotrachelus died, there was probably still some limited growth after departure of the weevils as indicated by a shift of the holes in the different culm sheets relative to each other (fig. 41). Consequently, at the time of felling there was no longer a contiguous opening all the way to the interior of the culm. The larvae of Ptecticus malyensis were found in all internodes connected by holes. In the lower internodes with punctured septa water usually accumulated, and dead stratiomyid larvae were found there on several occasions. Pupation of Ptecticus malayensis took place in the upper section of the shoot, behind the Cyrtotrachelus escape hole (fig. 41). Probably the freshly eclosed flies had to sqeeze themselves past the sheets to escape from the bamboo. The adult flies left before the rotten shoot tips fell to the ground. Five out of twelve investigated bamboo shoots were occupied by Ptecticus malayensis, with a maximum number of 22 larvae per shoot. In two shoots larvae of Solva completa de Meijere (Xylomyidae, Diptera) were found together with Ptecticus malayensis.

Only a few literature accounts on larvae of Ptecticus are available. Lindner (1928), who superficially de-

Figs. 26-32. Ptecticus longipennis, larval structures (SEM-photographs). -26 , head and fore margin of first thoracic segment, dorsal view; 27 , the same, lateral view; 28 , last abdominal segments, ventral view; 29, opening to spiracular chamber, posterior view; 30, short pennate setae on ventral lip of spiracular chamber opening; 31 , sternal patch on abdominal segment $6 ; 32$, sternal patch on abdominal segment 5 .

. wis sp.n.. larval structures (SEM-photographs). -33 , head and fore margin of first thoracic segid and two thoracic segments in lateral view; 35, last abdominal segments, ventral view; 36, opennterior view: 37, pennate setae on lips of spiracular chamber opening; 38, sternal patches on ab1 patch on abdominal segment 5 .


Figs. 40-43. Habitat of the larvae of Ptecticus malyensis sp.n. and P. longipennis. - 40, dead shoot tip of Gigantochloa scortechinii. The arrow points to the emergence hole of the weevil larva, Cyrtotrachelus sp., which was probably used by $P$. malayensis to enter the hollow interior of the shoot; 41, emergence hole of Cyrtotrachelus sp. after removal of several culm sheets. Two puparia of $P$. malayensis are visible (arrows). As can be seen, the openings of the culm sheets have shifted relative to each other; 42, the hollow interior of the shoot. Arrow points to a groove made by the larva of Cyrtotrachelus sp.; 43, larva of Ptecticus longipennis on the moist interior wall of an internode laying on the ground.
scribed a larva of $P$. testaceus Fabricius from South America, pointed out that larvae were found in decaying cactuses, oranges and lemons. Engel \& Cuthbertson (1939), who examined a larva of African P. posticus Wiedemann, found the larvae in the semi-liquid mass of decaying Conopharyngia johnstoni Stapf. (Apocynaceae) fruits together with other dipterous larvae.

Larvae of North-American Ptecticus trivittatus (Say) were collected from a corn compost pile, decaying tomatoes, garbage and fruit, from watermelon rinds, rotting paper and a fungus (McFadden 1967).

Our observations suggest that $P$. longipennis and $P$. malayensis are unspecialized phytosaprophages. These findings conform to the previously reported larval mode of life within the subfamily Sarginae: saprophagy with some tendency to coprophagy as known in many European species.

## Relationships of Ptecticus Loew

Only one reliable character separates the genus Ptecticus Loew from the related Sargus Fabricius, viz. the shape of the pedicel projecting into the elongate
process on the inner side of the antenna. However, this character may be indistinct in some species, and its generic relevance therefore seems disputable. A comparison of the structure of the male genitalia is not possible for all species of the two genera because it has not been included in all descriptions. It may nevertheless be stated that the differences found among some Ptecticus species seem to be more pronounced than those among the European genera of the Sarginae. This fact indicates the necessity of a revision of the generic concept of Ptecticus.

Judging from the description of the larva of NorthAmerican P. trivittatus (Say) by McFadden (1967) and the descriptions of the two larvae presented here, Ptecticus appears, on the other hand, well separated from the genera of Sarginae in the larval stage. First of all, in addition to a long and slender sternal patch on abdominal segment 6, a small and rounded sternal patch on abdominal segment 5 is developed. Relatively short setae on the head as well as on the anal segment, the configuration of setae on the lateral wall of abdominal segments and the presence of five pairs of ventral setae on the anal segment are cosid-
ered diagnostic generic characters. So, the known larvae of Sarginae may be distinguished according to the following identification key:

1. Setae on anal segment shorter than length of penultimate segment

- At least some setae on anal segment longer than penultimate segment $\qquad$ .4

2. Eye prominence anterior to protruding lateral tubercle (according to McFadden 1967) .................

Merosargus Loew

- Eye prominence on protruding lateral tubercle . 3

3. Abdominal segment 5 without sternal patch, four pairs of ventral setae on anal segment, 2-4 setae on lateral wall of abdominal segment $1,6-7$, and always one seta less or equal number of setae on abdominal segments 2-5 $\qquad$ Sargus Fabricius

- Abdominal segment 5 with a small but distinct sternal patch in addition to sternal patch on segment 6, five pairs of ventral setae on anal segment, four setae on lateral wall of abdominal segments $1,6-7$, and only two setae on abdominal segments 2-5 (i.e. two setae less) ... Ptecticus Loew

4. Anterior labral setae short, apical lobes on anal segment prominent $\qquad$ Chloromyia Duncan

- Anterior labral setae strikingly long, apical lobes on anal segment less distinct .. Microchrysa Loew


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## References

Beeson, C.F.C., 1941. The ecology and control of the forest insects of India and the neighbouring countries. - Vasant Press, Dehra Dun, 1007 pp. (Reprinted 1961).
Brunetti, E., 1923. Second revision of the Oriental Stratiomyidae. - Records of the Indian Museum 25: 45180.

Brunetti, E., 1927: Notes on Malayan Diptera, with descriptions of new species. - Journal of the Federal Malay States Museums 13: 281-309.
Engel, E.O. \& A. Cuthbertson, 1939. Systematic and biological notes on some brachycerous Diptera of southern Rhodesia. - Journal of the Entomological Society of South Africa 2: 181-195.
James, M.T., 1975. Family Stratiomyidae. In Delfinado, M.D. \& D.E. Hardy (eds.), A catalog of the Diptera of the Oriental Region. Vol.II. The University Press of Hawaii, Honolulu, 459 pp.
Lindner, E. 1928. Dr. L. Zürchers Dipteren-Ausbeute aus Paraguay: Stratiomyiiden. - Archiv für Naturgeschichte (A) 22 (12): 94-103

McFadden, M. W., 1967. Soldier fly larvae in America north of Mexico. - Proceedings of the United States National Museum 121: 1-72.
Meijere, J. C. H. de, 1904. Neue und bekannte SüdAsiatische Dipteren. - Bijdragen tot de Dierkunde 17-18: 83-118.
Nagatomi, A., 1975. The Sarginae and Pachygasterinae of Japan (Diptera: Stratiomyidae). - Transactions of the Royal Entomological Society of London 126: 305-421.
Rozkošný, R., 1982/83. A biosystematic study of the European Stratiomyidae (Dipteta). - Dr. W. Junk Publishers, The Hague, Vol. 1: 1-401, Vol. 2: 1-431.
Rozkošný, R. \& D. Kovac, 1991. First description of the male and the larva of Camptopteromyia fractipennis de Meijere from Malaysia (Diptera: Stratiomyidae). Entomologica scandinavica 22: 297-304.
Wulp, F. M. van der, 1885. On exotic Diptera. - Notes from the Leyden Museum 7: 1-15.

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# THE ODONATA OF SULAWESI AND ADJACENT ISLANDS 

Part 3. The genus Macromia Rambur (Corduliidae)'


#### Abstract

Tol, J. van, 1994. The Odonata of Sulawesi and adjacent islands. Part 3. The genus Macromia Rambur (Corduliidae). - Tijdschrift voor Entomologie 137: 87-94, figs. 1-25, table 1 [ISSN 0040-7496]. Published 15 July 1994. The Macromia species of Sulawesi (Celebes) are detailed, and a key to the species is provided. One new species, Macromia celebica sp, n., and the hitherto unknown female of M. irina Lieftinck are described. The occurrence on Sulawesi of M. moorei fumata Krüger is considered doubtful. The phylogenetic relationships of the Indo-Australian species of Macromia are analysed. The Sulawesi species are shown to have a sister group relationship to the monophyletic group of the Papuan species. M. chalciope from Halmahera is considered the most recently evolved taxon. These results are compared with recent reconstructions of the geological history of the region. J. van Tol, National Museum of Natural History, P. O. Box 9517, 2300 RA Leiden, The Netherlands. Key words. - Odonata; Corduliidae; Macromia; new species; phylogeny; biogeography; Indonesia; Sulawesi; New Guinea.


' Part 2: Zoologische Mededelingen [1987] 61 (13): 160-176.

Hitherto, only two species of the rheophilic dragonfly genus Macromia Rambur have been mentioned from Sulawesi (Celebes), viz. M. moorei fumata Krüger and M. irina Lieftinck (Lieftinck 1950: 714716). Specimens of Macromia that were recently collected in the southern part of Central Sulawesi, appear to belong to a species as yet undescribed. The number of species reliably recorded from this island, however, remains two, since the only record of Macromia moorei fumata must be regarded doubtful (see below).

The dragonflies of the virtually cosmopolitan genus Macromia are remarkable insects in many respects. With more than 120 species, several of which polytypic, Macromia is one of the largest genera of the Anisoptera. They are especially speciose in the tropical areas of the Old World. In Europe (one species), North America (nine species) and Australia (two species), they are far less numerous. All other species are confined to Africa and Southeast Asia.

Our present knowledge of the Macromia species of Southeast Asia is mainly based on the studies of Krüger (1899) and especially Lieftinck (1929, 1950, 197 la; see also 1954).

Due to their elusive behaviour, most species are poorly represented in collections. In fact, many are only known from the type series, often consisting of
one specimen only. Most species are fierce flyers. The larvae seem to be quite stenotopic, so that larvae as well as adults can only be found in very restricted parts of streams. Notwithstanding their strong flying capabilities, many species have small ranges, apparently due to the fact that they seldom fly far from their breeding sites.

In the present paper I describe the new species from Sulawesi, provide some new observations on $M$. irina, and give a key to species recorded from Sulawesi. Finally, after having analysed the affinities of the Sulawesi species to those of the Oriental and the Papuan regions, I propose a hypothesis for a scenario of the biogeographical history of this genus in the Indo-Australian region.

## Systematic part

## Key to the Macromia species of Sulawesi

1. Outer margin of superior appendage of male with well-developed tooth-like tubercle approximately halfway (cf. fig. 18); vertex bilobed (cf. figs. 22, 23); thorax green with metallic lustre $\qquad$ M. moorei fumata
[Occurrence on Sulawesi doubtful].

- Outer margin of superior appendage of male


Figs. 1-7. Macromia celebica sp. n. - 1, Head in frontal view; 2, Anal appendages of male, dorsal view; 3, Idem, left lateral view; 4 , Base of right hind wing, male; 5 , Pterostigma of right fore wing; 6 , Accessory genitalia of male, left lateral view; 7, Fourth segment of penis, left lateral view.
somewhat curved inwards, in some specimens with very small teeth along the margin, but especially in dorsal view not a conspicuous extero-lateral tooth (figs. 2, 9); vertex dorsally smoothly rounded, without a paired tubercle (figs. 1, 8); thorax green or brown $\qquad$2
2. Appendix inferior distinctly shorter than superiors (figs. 2, 3); coloration at least on synthorax green with greenish blue metallic shine; frontal side of head brownish black .... M. celebica sp. n.

- Appendix inferior approximately as long as superiors (figs. 9, 10); coloration without any indication of metallic lustre, but body generally orange-brown; head orange-brown with some middle-brown parts, but without conspicuous brownish black frons M. irina


## Macromia celebica sp. n.

(figs. 1-7)
Type material. - Holotype ơ. Indonesia. Southern Central Sulawesi. Puncak Palopo. $600-1000 \mathrm{~m}$.

October 1993. Leg. Yohan R. (JvT 6007) (MBBJ). Paratypes: 20 ${ }^{\text {on }}$, same data (JvT 5930-5931) (RMNH).

Description. - A medium-sized Macromia, with green thorax and dark abdomen.
Male. Head with labium, labrum and clypeus orange-brown, especially dypeus without yellow markings, labrum with outer corners darker. Pyramidal processes of frons high and subacute (fig. 1); frontal side very flat, more or less triangular, with very fine transverse striae from upper-inner to lowerouter corner of each side; frontal and upper parts of frons shiny black with blue metallic lustre; frons against eyes orange-brown as lower part of face; vertex black with fine punctation, the paired dorsal tubercles hardly discernable, the lateral angles tapering into small projecting cones on top of which the lateral ocelli are situated (shape as in irina; westwoodii has somewhat more pronounced dorsal tubercles), covered with thin setae. Occiput shiny black with very fine punctation. Antennae brownish black.

Thorax. Brilliant metallic green, densely covered
with long and slender creamish white setae; antehumeral stripe yellow and distinct, somewhat tapering dorsally, extending dorsally to ca. three-seventh of mesepisternum, ventrally continuing over mesinfraepisternum; yellow stripe over spiracle tather narrow, tapering in width dorsally and nearly reaching the upper margin of episternum. Pale stripe on lateroventral part of metepimeron covering anterior anterior half (stripe absent in chalciope). Legs long and slender, hind femur 10 mm , reaching to base of posterior hamulus, colour brown, keel on fore leg very short, ca. 1 mm , not reaching the apex, middle tibia without keel, hind tibia with long keel from ca. 2 mm from base of tibia to less than 1 mm from its apex.

Wings relatively short, fore wing with 17-18 Ax, 8 Px ; hind wing with 11 Ax, 11-13 Px; fore wing with 7 Cux, hind wing with 5 Cux (incl. subtriangle); anal loop 7-9 cells without central cell; anal triangle relatively short (fig. 4; cf. M. chalciope, fig. 19); discoidal triangle of hind wing very narrow; pterostigma (fig. 5) reddish brown, anterior side ca. 2.0 mm .

Abdomen brownish black with some blue-green metallic lustre especially on first few segments; yellow spot on anterior side of segment 7; general shape anteriorly slender, especially segment 7-9 significantly widened; tergites ventro-anteriorly somewhat paler, dark or middle-brown. Spot on segment 7 consisting of a complete ring covering anterior one-fourth of segment, medio-dorsally extending posteriorly for another one-eighth segment length and ca. one-third the segment's width. Segment 10 rather short, middorsal process not spine-like, but consisting of a low triangular-shaped subterminal tubercle. Appendages (figs. 2-3) with superiors in dorsal view rather slender, significantly longer than inferior, basal two-fifths rather straight, then innerside slightly tapering posteriorly and outerside strongly tapering, caudalmost part somewhat curved outwards; no trace of a lateral tubercle on superior appendage.

Secondary genitalia: anterior side margin of tergite 2 with dense setae, a mixture of strong and soft setae, longer than in irina; also tuft of setae on ventral tu-


Figs. 8-15. Macromia irina Lieftinck. - 8, Head in frontal view; 9, Anal appendages of male, dorsal view; 10, Idem, left lateral view; 11, Base of right hind wing, male; 12 , Pterostigma of right fore wing; 13. Accessory genitalia of male, left lateral view: 14, Fourth segment of penis, left lateral view; 15, Valvula vulvae, female.
bercle of tergite longer; genital lobe directed posteriorly, elongate, and tapering towards the top, but tip not acute; posterior hamulus glossy brown, basally coloured brown, the straight outer part pale brown, and the endhook reddish brown, curved strongly $\left(90^{\circ}\right)$ inwards, endhooks in ventral view slightly diverging, the tip reaching ca. two-thirds of genital lobe.

Measurements. - Abdomen, incl. appendages 48 (paratypes $46-48$ ) mm, hind wing 42 (paratypes 42 43) mm

Female. - Unknown.
Differential diagnosis. - Differs clearly from the only other Sulawesi Macromia by the presence of green metallic lustre, the black markings on the frons, the presence of an antehumeral stripe; structurally the short inferior appendage, and shape of abdominal segment 10 are diagnostic. Both Sulawesi species can be distinguished from other Indo-Australian Macromia by the intermediate sized pterostigma, the very narrow discoidal triangle of the hind wing, and especially the rounded (not bilobed) vertex. Also, the superior appendages lack an extero-lateral tooth, but are not particularly slender. The Papuan Macromia have extremely slender superiors with or without an extero-lateral tooth (fig. 20), while the Oriental species have rather solid appendages with a more or less conspicuous extero-lateral tooth (fig. 18).
Remarks. - Although this species differs in many respects from $M$. irina, I consider it the sister species of M. irina. This view is based on shared, presumably apomorphous, character states in the structure of vertex and superior appendage of male (see below).
Distribution. - Only the southwestern part of Central Sulawesi. Collected between $600-900 \mathrm{~m}$.

## Macromia irina Lieftinck

(figs. 8-15)
Macromia irina Lieftinck, 1950: 714-716. Holotype male 'S Celebes, foot of Mt Lompobatang, Borong Rapao, 800 $\mathrm{m}, 12$.viii. 1949, leg. A. Diakonoff in RMNH [examined]. - Lieftinck 1971b: 94 [note on type]; Askew et al. 1989: 118 [records Northern Sulawesi, Dumoga Bone National Park].

Diagnosis. - Unmistakable among Indo-Australian Macromia: head, thorax and abdomen brown without any metallic lustre. Structural characters: wing base with anal angle sharp, but not angulated (fig. 11); the anal triangle rather short (long in moorei fumata), membranula ending circa halfway between anal crossvein and base of anal triangle; prerostigma rather short (fig. 12), circa twice as long as high (more than three times in moorei fumata); anal appendages of male with superiors with rather smooth outer margin, without any indication of tooth (figs. 9-10) (cf. moo-
rei fumata and westwoodit); inferior appendage approximately as long as superiors (clearly shorter in celebica).

Description. - Little can be added to the extensive description of the male in Lieftinck (1950). Live colour of eye dark green. Since Lieftinck did not provide any illustration, I have added here drawings of the head, wing base, anal appendages and secondary genitalia (figs. 8-14). The up to now undescribed female was collected in Southeast Sulawesi.

Female. - General coloration as the male, and structurally very similar to the male, although more robust. Head and thorax not differing from male. Wings somewhat broader, especially at base; membranula fully opaque white or only greyish in most narrow part; rusty brown base of wings more extensive than in male, in costal and subcostal area of fore and hind wing reaching to first Ax or even somewhat beyond; some less extensive rusty brown at base of medio-anal space; Arculus at Ax2 in fore wing and just beyond Ax2 in hind wing; fore wing 16-17 Ax, 9 Px, 7 Cux; hind wing 10-12 Ax, 11 Px, 5 Cux. Abdomen of the coloration ('Mikado-brown') as male; segment 1 dorsum dark brown and lateral parts brown; segment 2 brown, somewhat darker above than laterally, as the male with two narrow transverse yellow streaks, one over transverse carina halfway abdominal segment and the other pair more oblique running just before posterior margin; segment 3 anteriorly brown to transverse carina, somewhat darker above, posteriorly dark brown except for space just before segment 4 ; segments $4-7$ similar to segment 3 , but dark patch anterior to transverse carina gradually increasing in size, and posterior paler brown ring also increasing from segment 4 to 7 ; segment 8 brown with dorsal carina black and some darkening against dorsal carina; segment brown, dorsal carina dark; segment 10 brown; appendages middle brown, rather short, acute. Valvula vulvae short, approximately four times as wide as long, with wide triangular emargination, the tips just bifid, somewhat curved inwards (fig. 15).

Other material examined. - Sulawesi Tenggara. S of Sanggona: Gunung Watuwila. Sunggai Lalonduwasi near Centipede camp. c. 1050 m. 2-4 Nov 1989 (J. van Tol) $3 \delta^{\text {ot }} 1$ ㅇ (JvT 5932-5935); Sulawesi Tengah, ca. 10 km WNW Palopo near Tojambu. $800-1000 \mathrm{~m}, \mathrm{c} .2^{\circ} 56^{\prime} \mathrm{S} 120^{\circ} 07^{\prime} \mathrm{E}, \mathrm{Jul} /$ Aug 1991 (Yohan R.), 1 if (JvT 5936), all in RMNH. I have also studied one of the specimens collected by Askew et al. (1989): $1 \delta^{\star}$, in RSME.

Remarks. - I collected this species myself along a small, shaded stream in a steep valley in the Mengkoko mountain area. The insects appeared immediately after sunrise, and were active for approxi-


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Figs. 16-23, Macromia species. 16-18, 22, M. westwoodii; 19-21, 23, M. chalciope - 16,19 , Base of right hind wing, male; 27, 21, Pterostigma of right fore wing; 18, 20, Anal appendages of male, dorsal view; 22-23, Vertex in frontal view (fig. 23 dorsal-most part only).
mately one hour. Males as well as females flew low over the water up and down the stream, more or less in a straight line. With a rather slow and small wing beat, they were not extremely fast. The specimens could be collected at a site where a big log of more than one meter diameter was laying in the stream. Macromia specimens crossed the log at just a few centimeters height to return immediately to the water surface. The female was caught on a dark day at 16.30 h, i.e. 1.5 hours before sunset.

## Macromia moorei fumata Krüger

Macromia fumata Krüger, 1899: 333-335 (Java) [not examined].
Macromia moorei fumata. - Lieftinck 1950: 678 (key, mentioned Celebes); Lieftinck 1954: 119, 185 (references, geographical distribution); Lieftinck 1971a: 29 (geographical distribution).

Remarks. - I have been able to check the specimen (in SMFD) at which Lieftinck's notice of this taxon on

Sulawesi was based. Although this specimen can be assigned to $M$. moorei fumata without doubt, I consider its occurrence on Sulawesi still doubtful. The 'Lompo Battau' labels of Fruhstorfer have on many occasions proved to be incorrect (see van Tol 1987: 163). M. moorei fumata is rather common on Java, where Fruhstorfer also collected during his long journey through the former Dutch Indies.

Material examined. - Lompa Battau, S. Celebes, III. 1896 (H. Fruhstorfer) '16764' (in green ink) 10 [label in Ris' hand] [identified by M.A. Lieftinck as Macromia moorei fumata, Jan. 1928].

## Phylogenetic relationships

## Introduction

The so-called Papuasian representatives of Macromia share at least four characters (Lieftinck 1952, 1971a), viz., an exceptionally small discoidal triangle of the hind wing, a minute pterostigma of ca. one mm , a sharply acute anal angle of the hind wing in

Table 1 Character matrix for taxa of Macromia of the Indo-Australian archipelago.

the male, and long and slender genital hamules in the male. Lieftinck (1971a: 30-32) distinguishes three groups among the Papuan Macromia species, viz. M. terpsichore, M. melpomene, and M. chalciope, with the last one consisting of one species only.

The species assigned to these groups, which the exception of viridescens described from Cape York (Australia), are confined to the islands of New Guinea (astarte Lieftinck, celaeno Lieftinck, eurynome Lieftinck, melpomene Ris, terpsichore Foerster, viridescens Tillyard (= amymone Lieftinck)), the Bismarck Archipelago (lachesis Lieftinck), Waigeu (euphrosyne Lieftinck, sophrosyne Lieftinck) and Misool (hermione Lieftinck). M. chalciope Lieftinck is restricted to Schouten Is, and the Moluccan islands of Halmahera and Bacan (Lieftinck 1971a). No species of Macromia have been mentioned from any of the other Moluccan islands up to now. According to Lieftinck (1952: 460) M. chalciope can be considered 'an immediate derivative of the eastern cluster', i.e. that the sister-species of chalciope could be discovered among the species of New Guinea and adjacent islands.

The Malay peninsula and the Greater Sunda Islands are inhabited by 15 species of Macromia (see Lieftinck 1971a: 29). Although Lieftinck (1929) attempted to distinguish species groups in the western Malesian species, this author (1950:716) had to admit that the groups had an artificial character. Variation in morphology among the species of west-
ern Malesia is more prominent than that among the Papuan representatives.
Finally, the Philippine islands are inhabited by at least one endemic species, viz. M. negrito Needham \& Gyger from Luzon. The widespread M. cincta is reported from Palawan (Lieftinck 1929: 92), and Tawi Tawi (Lieftinck 1974), while also M. gerstaeckeri may


Fig. 24. Tree for Macromia species of the Indo-Australian archipelago.


Fig. 25. Phylogenetic relationships of Macromia species (groups) represented on the map of Malesia.
occur in the Philippines (Lieftinck 1971a: 29).
I present here the results of a preliminary investigation of the phylogeny of the Macromia of the IndoAustralian region, mainly based on the characters used by Lieftinck to establish his Papuan species groups, or to distinguish the Oriental and Papuan groups. As outgroups I have added $M$. moorei fumata Krüger and M. westwoodii Selys, both considered closely allied to M. irina (Lieftinck 1950). For the present study only a limited number of the Papuan species have been included. Unfortunately, M. negrito Needham \& Gyger from Luzon, was not available for study. A more extensive phylogenetic study, involving more of the Oriental as well as the Papuan species, has to wait for the future.

## Methods and results

The matrix (table 1) was analysed with PAUP, version 3.1.1 (Swofford 1993), using 'exhaustive search'. Macromia moorei fumata was designated as outgroup, but the results appeared to be similar when $M$. westwoodii was selected as such. The analysis resulted in one tree of 20 steps ( $\mathrm{CI}=0.800, \mathrm{RI}=0.818$ ).

This tree (fig. 24) shows that all Papuan species form a monophyletic group, with the Sulawesi species as sister group. The Papuan groups as defined by Lieftinck (1971a) are not corroborated by the present analysis. It is of interest to notice the sistergroup relationship of M. terpsichore from NE New Guinea, and M. chalciope from Halmahera.

Homoplasies in the present tree expecially occur in
two non-structural characters, viz. character 7 (antehumeral stripe) and to a lesser extent character 6 (postclypeus).

## Discussion and biogeography

The results of the phylogenetic analysis partly confirm earlier suggestions by Lieftinck on relationships among Indo-Australian Macromia species. Lieftinck (1950) considered M. irina from Sulawesi closely related to $M$. moorei and $M$. westwoodiz, both rather widespread in Southeast Asia, including Borneo. The rather solid build of the superior appendage, in combination with the absence of an extero-lateral tooth, ranks the Sulawesi representatives in an intermediate position between the Oriental and the Papuan species. This intermediate position can also be recognized in the shape of the pterostigma.

The rather distant relationships of the Sulawesi and Halmahera species are of interest in a biogeographical context. Although the geological history of Wallacea and Halmahera is still under debate (e.g. Daly et al. 1991, Honza 1991), there is general agreement that there is no shared history for the North Moluccas (Halmahera) and Sulawesi. Reconstructions prior to 10-15 million years ago (Ma) are highly uncertain because of the very complicated plate movements. In the reconstruction of Daly et al. (1991) (see also Nichols \& Hall 1991) Halmahera was formed at the margin of the Pacific and the Philippine plates, north of New Guinea, with a westward progression from c. 5 Ma to its present position. At least some parts of

Sulawesi have a much longer ( 65 million years) history above sea level on a far more western position. In other words, from approximately 60 to 5 Ma , the Oriental region + Sulawesi have shared (in some respects) a common history, while Halmahera shares its history mainly with New Guinea. The close contact of the Oriental and Papuan region is more recent than 5 Ma .

Given these phylogenetic relationships, and our present knowledge of the geological history of the area, the following historical scenario may be outlined. All Macromia species in the Indo-Australian region originate from an Oriental ancestor. The ancestor of the Sulawesi + Papuan species groups possibly inhabited Sulawesi, while the ancestor of the Halmahera M. chalciope + New Guinea M. terpsicho$r e$ species groups apparently lived in north(eastern) New Guinea. The occurrence of two relatively remotely allied species on Sulawesi and Halmahera can be attributed to the westward movement of Halmahera during the last ten million years.

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## References

Askew, R.R., G.G. Cleland, D.A.L. Davies \& T.W. Harman, 1989. A report on a collection of Odonata from North Sulawesi, Indonesia. - Tijdschrift voor Entomologie 132: 115-121, fig. 1, tables 1-2.
Daly, M.C., M.A. Cooper, I. Wilson, D.G. Smith \& B.G.D. Hooper, 1991. Cenozoic plate tectonics and ba-
$\sin$ evolution in Indonesia. - Marine and Petroleum Geology 8: 2-21, figs. 1-15.
Honza, E., 1991. The Tertiary arc chain in the Western Pacific. - Tectonophysics 187: 285-303, figs. 1-5
Krüger, L., 1899. Die Odonaten von Sumatra. III. Theil. a. Familie Libelluliden (with supplement: Neue Cordulinen von Java und Borneo. - Stettiner Entomologische Zeitung 60: 321-338.
Lieftinck, M. A., 1929. A revision of the known malaysian dragonflies of the genus Macromia Rambur, with comparative notes on species from neighbouring countries and descriptions of new species. - Tijdschrift voor Entomologie 72: 59-108, figs. 1-22.
Lieftinck, M. A., 1950. Further studies on Southeast Asiatic species of Macromia Rambur, with notes on their ecology , habits and life history, and with descriptions of larvae and two new species (Odon., Epophthalmiinae). Treubia 20: 657-716.
Lieftinck, M. A., 1952. On the Papuasian representatives of the genus Macromia Ramburm with descriptions of five new species and some larval forms (Odon.). - Treubia 21: 437-468, p. 1-22.
Lieftinck, M. A., 1954. Handlist of Malaysian Odonata. Treubia 22 (Supplement): i-xiii + 1-202, map.
Lieftinck, M. A., 1971a. Studies in Oriental Corduliidae (Odonata), I. - Tijdschrift voor Entomologie 114 (1): 163, figs. 1-50.
Lieftinck, M. A., 1971b. A catalogue of the type-specimens of Odonata preserved in The Netherlands, with a supplementary list of the Odonata types described by Dutch scientists deposited in foreign institutional collections. Tijdschrift voor Entomologie 114 (2): 65-139 + plate 17.

Lieftinck, M. A., 1974. Dragonflies collected by the Noona Dan Expedition in the southwestern Philippine Islands (Insecta, Odonata). - Steenstrupia 3: 111-147, figs. 1-26.
Nichols, G. J. \& R. Hall, 1991. Basin formation and Neogene sedimentation in a backarc setting, Halmahera, eastern Indonesia. - Marin and Petroleum Geology 8 (1): 50-61, figs. 1-9.
Swofford, D. L., 1993. Paup: Phylogenetic Analysis Using Parsimony. Version 3.1.1. - Illinois Natural History Survey, Champaign.
Tol, J. van, 1987. The Odonata of Sulawesi and adjacent islands. Part 2. The genus Diplacina Brauer on Sulawesi. Zoölogische Mededelingen 61: 160-176, figs. 14-39.

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## THE GENUS DIPSEUDOPSIS WALKER FROM ASIA (TRICHOPTERA: DIPSEUDOPSIDAE)


#### Abstract

Weaver, J. S., III \& H. Malicky, 1994. The genus Dipseudopsis Walker from Asia (Trichoptera: Dipseudopsidae). - Tijdschrift voor Entomologie 137: 95-142, figs. 1-55. [ISSN 0040-7496]. Published 15 July 1994.

The genus Dipseudopsis Walker from Asia is revised, and a phylogeny of Hydropsychoidea is provided showing the relationships of its family taxa and the genera of Dipseudopsidae. Presently 36 Asian species of Dipseudopsis are recognized, having a distribution including Pakistan, India, Sri Lanka, Nepal, Bangladesh, Burma, Thailand, Cambodia, Vietnam, China, Japan, Philippines, Malaysia and Indonesia, including Sumatra, Java, Borneo, and Sulawesi. Six new species are described, Dipseudopsis adiaturix sp. n., D. finti sp. n., D. lucasi sp. n., D. malaisei sp. n., $D$. marrynovi sp. n., and D. schmidisp. n., and one new subspecies, D. robustior andamanensis ssp. n. The identities of several problematic species are resolved, with 16 species names recognized as new junior synonyms (cf. species checklist p.102). D. nervosella Ulmer stat. n. is elevated from former subspecies status. New descriptions are provided for all species, including figures of the male hind mesoapical tibial spur (the modified spur), the male genitalia, and other notable characteristics of the head, thorax and wings, except two species recognized as nomina dubia, D. onychophora Navás, and D. orientalis (Navás). New lectotypes are designated for $D$. contorta Banks, $D$. morosa Banks and D. triclavata Martynov. A pictorial atlas is provided for rapid species identification. The genus Eodipseudopsis Marlier is removed from the Dipseudopsidae. Correspondence: J. S. Weaver III, Department of Entomology, University of New Hampshire, Durham, NH, 03824, U. S. A.


Key words. - Asia, Dipseudopsis, Dipseudopsidae, Hydropsychoidea, phylogeny, systematics, Trichoptera.

The genus Dipseudopsis Walker exhibits a number of characteristics that are quite remarkable among the Trichoptera. The adults are the only members of the suborder Annulipalpia Martynov having siphoning mouthparts, with the lacinia forming a proboscis, a parallel modification of the galea in the higher Lepidoptera. Sexual dimorphism is conspicuous in the genus, with the males having highly contrasting colour patterns in their wings and bodies, and the females less striking in comparison. Also, males have the mesoapical spur of the hind tibia modified, often twisted and/or branched. The function of the modified spur is unknown, but it has been used almost exclusively for species determinations, to the extent that customary descriptions of the male genitalia often have been omitted from species descriptions. The convention of using the modified spur as the basis for species descriptions was established by Ulmer (1907b: 180): 'Bei Dipseudopsis sind die Genitalien aller Species ganz gleich' (i.e: In Dipseudopsis the genitalia of all species are totally alike). Hence, the mod-
ified spur became the primary characteristic used to recognize species; e.g. this is the main distinguishing character in every couplet of Ulmer's (1951) key to eleven species of Dipseudopsis of the Sunda Islands.

The precise orientation of the modified spur must be made apparent for species identification, because slight rotations of the tibia will show different perspectives of the spur. Some of the previous species descriptions based on the modified spur were ambiguous, because the figure of the spur did not show the remainder of the leg, and hence the actual orientation of the spur was uncertain. This problem accounts for several of the synonyms that we discovered in the course of this study. The figures of the spur herein are usually of the left leg and include the adjacent apicolateral spur and a portion of the tibia and tarsus to provide proper comparison of other specimens in the same perspective.

It seems somewhat ironic that the taxonomy of this genus, which exhibits so many fascinating character-

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    Adult dorsal onelli prevent \(=0\); onelli absent \(=1(\) cf. Schmid 1980)
    sgment \(I X\) completely selcrotized eylinder \(=0 ; I X\) jawlike with articulation between tergum \(\& x\) sternum \(=1\) (cf.
    Shmad lusu
    esement \(X\) without lateral papillae \(=0 ;\) X with lateral papillae \(=1(\) cf. Schmid 1980).
    sgment IX normal = 0: IX long completely sclerotized cylinder = 1 (cf. Schmid 1980).
    I an a spunneret short \(=0\); spinneret longer than other mouthparts \(=1\) (cf. Wiggins 1977).
    jsernum \III normal. undivided \(=0\); sternum VIII divided into a pair of lateral lobes \(=1\) (cf. Schmid 1980).
    \(\because\) infertor appendages 2 -segmented \(=0 ; 1\)-segmented \(=1(c f\). Schmid 1980).
    Adult baves of antermace distant \(=0\); bases close together \(=1\) (cf. Crichton 1957).
    Lina tani nearly calindrical, pretarsal claws long \(=0\); tarsi flattened, claws short \(=1\) (Ulmer 1957, Gibbs 1968, Wiggins
    \(1)^{--}\). Wells \& (artwright 1943).
    ianal retreat differs from the following \(=0\); larva with branched tube-dwelling and filter-feeding behaviour as in
    P/omloicntropm (Wallace et al. 1976, Wiggins 1977).
11. Adule labial palps present \(=0\); absent \(=1\) (Ulmer 1951).
12. Adulr pronorum normal \(=0\); pronorum enlarged and with deep median fissure \(=1\).
13. Aduls with head and thorax setous \(=0\); head and thorax glabrous \(=1\).
14.. Adult regula knoblike \(=0\); tegula fartened like epaulets \(=1\).
15. Adult with galca shore, vestigial \(=0\); modified into proboscis \(=1\).
16. Lana labrum expanded and more membranous = 1, autapomorphy for Philopotamidae (cf. Wiggins 1977).
\(1^{-}\). Lana trochantin broad, either with blunt apical angle or truncate \(=1\), autapomorphy for Psychomyiidae (cf. Wiggins
    (1)
18. Lana with mesopleural lobes = 1, autapomorphy for Xiphocentronidae (cf. Wiggins 1977).
19. I ana with cibiac and carsi of legs fused \(=1\), autapomorphy for Xiphocentronidae (cf. Wiggins 1977).
20. Lana with ventral thoracic gills whorled \(=1\), autapomorphy for Hydropsychidae (cf. Wiggins 1977).
21. Lana cpidermis with modified setae, especially on sternum VIII \& IX and tergum \(=1\), autapomorphy for
    Hudropsychidac (of. Wiggins 1977).
22. Lana head with ventrolateral bands of transverse ridges used in stridulation = 1, autapomorphy for Hydropsychidae (cf.
    Wigeins 19"-
23. Lanal head and body flattened dorsoventrally = 1, autapomorphy for Ecnomidae (cf. Lepneva 1964).
24. Laraa lateral fringe dense \(=1\), autapomorphy for Ecnomidae (cf. Lepneva 1964).
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istics, should have become so confused at the species level. This is chiefly because Banks, Martynov, Navás, and (Vlmer from 1905 to 1937 proposed descriptions of 25 different species from Asia and the species deecriptions by: Banks and Navás are poor. Another facfor mighe be that specimens of Dipseudopsis are not parricularly abundant in muscum collections, even shough adults are attracted to light traps.

The primary objective of this work is to clarify the iavonomy of the Asian species of Dipseudopsis. To accomplish this we have examined the types of most of the species as well as other specimens from several muscums and private collections. The species descriptitins have been expanded beyond the characteristics of the modified spur to include colouration of the hody and wings and illuserations of the male genitalia. Hence, most of the redescriptions of species herein provide the first illuserations of the male genitalia. We recognize 36 species of Dipseudopsis from Asia, 30 previously descrited species and six new species, and one new subspecies. W'e also recognize the names of 16 species and one subspecies as new junior synonyms. Two additional Asian species, D. onychophora Navis and $D$ ) orrensalis (Xavás), are recognized as nomina dubia. because both were poorly described and their types could not be located for examination.

## DEPOSITORIES

The depositories of material are abbreviated as follows: BMNH, Natural History Museum (formerly the British Museum of Natural History), London; BPBM, Bernice Pauahi Bishop Museum, Honolulu; CASC, California Academy of Sciences, San Francisco; CNCI, Canadian National Collection, Ottawa; CLDD, collection of D. G. Denning (now deposited at CASC); CLHM, collection of H. Malicky; CLJW, collection of J. S. Weaver III; McZC, Museum of Comparative Zoology, Harvard University, Cambridge; MNHN, Muséum National d'Histoire Naturelle, Paris; MZBS, Museu de Zoologia, Barcelona; MzLU, Museum for Zoology, University of Lund; NHMW, Naturhistorisches Museum Wien; NHRS, Naturhistoriska Riksmuseet, Stockholm; NZSI, National Zoological Collection, Zoological Survey of India, Calcutta; RMNH, Nationaal Natuurhistorisch Museum (formerly Rijksmuseum van Natuurlijke Historie), Leiden; SOFM, National Museum of Natural History, Sofia; UOPJ, Entomological Laboratory, University of Osaka Prefecture, Sakai; USNM, Smithsonian Institution, United States National Museum of Natural History, Washington; ZRAS, Zoological Institute of the Russian Academy of Sciences, St. Petersburg; ZMAN, Zoo-
logisch Museum, Amsterdam; ZMHB, Zoologisches Museum der Humboldt-Universität Berlin; ZMUC, Zoologisk Museum København; ZMUH, Zoologisches Museum der Universität Hamburg; and ZSMC, Zoologische Staatssammlung München.

## METHODS

## Taxonomy

The morphological terminology of wing venation and genitalic characters mostly follows that of Schmid (1980), and Schuster \& Hamilton (1984). The species synonymies are complete listing all original species descriptions, but subsequent references listed in Fischer's Trichopterorum Catalogus $(1962,1972)$ are not included. Locality names in current use are provided for specimens examined, where old names taken from labels are in quotes. For the type data, the term 'holotype' is used when the type specimen has actually been labelled or cited as such, otherwise it is listed as 'rype'. An asterisk $\left({ }^{*}\right)$ designates types of previously described species examined by the authors.

## Phylogenetics

Hennig86 (Farris 1988) is used for phylogenetic analysis to determine the relationship of Dipseudopsidae within the Hydropsychoidea s. str. Weaver, and the relationships of the proposed dipseudopsid genera. Twenty four character states (table 1) are selected, and with polarity determined for twelve taxa, using Philopotamidae Stephens as an outgroup, the data matrix (table 2) is produced. These data are analyzed using the 'mhennig*' command of Hennig86 for calculating trees. All characters have default values, weight $=1$, active and additive, except weight $=2$ for character 6 .

## DIPSEUDOPSIDAE ULMER

The family Dipseudopsidae is included in the suborder Annulipalpia Martynov, in the infraorder Curvipalpia Weaver (Weaver 1984, Weaver \& Morse 1986). The family name, Dipseudopsinae Ulmer (1904b), including only the nominative genus, was first proposed as a subfamily of the Hydropsychidae Curtis, but was later placed within the Polycentropodidae Ulmer (1906), as it has been recognized by several others (Tsuda 1942, Marlier 1962, Ross \& Gibbs 1973, Wiggins 1977, Schuster \& Hamilton 1984). However, some workers in the past have recognized Polycentropodinae and the Dipseudopsinae as subfamilies of Psychomyiidae Curtis (Ross 1956, Ross \& Kingsolver 1959), and more recently, some still recognize the Polycentropodinae as such (Flint 1991). The Dipseudopsidae were first given full family status by Ross (1967), and later concurred by Gibbs (1968), Wiggins (1982), Weaver (1984), Scott (1985), Weaver \& Morse (1986), and Wells \& Cartwright (1993).

The genus Protodipseudopsis Ulmer (1909) from the Afrotropical Region was the first addition to the subfamily, a relationship later confirmed by Ross \& Kingsolver (1959). Two additional genera from the Afrotropical Region were placed in the Dipseudopsinae, Eodipseudopsis Marlier (1959) by original designation, and Limnoecetis Marlier (1955) originally placed in the Leptoceridae Leach, but later moved to the Dipseudopsinae (Marlier 1961). Ross \& Gibbs (1973) transferred Phylocentropus Banks (1907) from Polycentropodinae to Dipseudopsinae, based predominantly on synapomorphic larval characteristics. They recognized only three genera in the Dipseudopsinae, Dipseudopsis, Phylocentropus, and Protodipseudopsis, and did not consider the status of Eodipseudopsis and Limnoecetis. However, Schmid (1980) later

Table 2. Data matrix for the families of Hydropsychoidea and genera of Dipseudopsidae, including outgroup Philopotamidae.

|  | Character |  |  |  |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
|  |  |  |  |  |  |
|  |  | 1 | 11111 | 1112 | 2222 |
| Taxon | 12345 | 67890 | 12345 | 67890 | 1234 |
| Philopotamidae | 00000 | 00000 | 00000 | 10000 | 0000 |
| Psychomyiidae | 11011 | 00000 | 00000 | 01000 | 0000 |
| Xiphocentronidae | 11011 | 00000 | 00000 | 00110 | 0000 |
| Hydropsychidae | 11100 | 10000 | 00000 | 00001 | 1100 |
| Polycentropodidae | 11100 | 11000 | 00000 | 00000 | 0000 |
| Ecnomidae | 11100 | 11000 | 00000 | 00000 | 0011 |
| Dipseudopsis | 11101 | 01111 | 01111 | 00000 | 0000 |
| Protodipseudopsis | 11101 | 01111 | 01110 | 00000 | 0000 |
| Phylocentropus | 11101 | 01111 | 00000 | 00000 | 0000 |
| Limnoecetis | $1110 ?$ | $011 ? ?$ | 01110 | $? ? ? ?$ | $? ? ? ?$ |
| Hyalopsyche | 11101 | 01111 | 10000 | 00000 | 0000 |
| Hyalopsychella | $11 ? ? ?$ | $? 1 ? ? ?$ | 10000 | $? ? ? ? ?$ | $? ? ? ?$ |

made the novel assignment of placing P/ylocentropus in the family Hyalopsychidae Lestage, on the basis of adule characteristics shared with the genus Hyalopsyche UTmer (19) (0)b). W'ells \& Carmsright (1993) recently concurred with the position of Ross and Gibbs, and based on the morphology of the larva and female of $H$ ozlopsorthe, added this genus to the Dipseudopsidac, and accordingly proposed the suppression of the family group name Hyalopsychidae. Our phylogeneric analusis tests the classifications of Ross \& Gibbs and Wells \& Cartwright, versus that of Schmid.

## Results

Phylogeny: - The analysis of the data matrix via Hennig86 generated three trees with different topologies, having length 28, CI 0.89, RI 0.91 (fig. 55). All three trees depict Dipseudopsidae as a monophyletic group, comprising six genera. The first tree has the fewest number of ancestors, having 18 nodes, and is also identical to the strict consensus tree (fig. 56) of the three trees. The consensus tree depicts (Psychomyiidae, Xiphocentronidae Ross) ((Hydropsychidae, Ecnomidac Ulmer, Polycentropodidae) Dipseudopsidac)), thus supporting monophyly of the Hydropsychoidea. The clade comprising Hydropsychidae, Polycentropodidae, Ecnomidae is recognized as the sister group of the Dipscudopsidae. A few differences are noted among the three original trees: 1) trees 0 and 2 depict the trichotomy (Hydropsychidae, Ecnomidac. Polycentropodidae), which is resolved as (Hydropsychidac (Ecnomidae, Polycentropodidae)) in utee 1; 2) trees 0 and 1 each have a basal and a distal trichotomy among the genera of Dipseudopsidae, both of which are resolved in tree 2 .

[^3]that the antennal scapes of Dipseudopsidae are not merely close together but rather are so enlarged that they occupy most of the space between the eyes along the anterodorsal margin of the head (fig. 13).

Conclusions. - Monophyly of the Dipseudopsidae, including Dipseudopsis, Hyalopsyche, Hyalopsychella Ulmer (1930), Limnoecetis, Phylocentropus, and Protodipseudopsis, is unequivocal. Two clades are recognized among these genera, (Hyalopsyche + Hyalopsychella) supported by synapomorphy 11 (cf. table 1), and (Dipseudopsis + Protodipseudopsis + Limnoecetis) supported by synapomorphies 12-14. Hence, the sister group of the genus Dipseudopsis is unclear. The adult of Limnoecetis, a monotypic genus endemic to Lake Tanganyika, are quite remarkable, having unique adaptations to walk on water. The striking autapomorphies of Limnoecetis present a formidable challenge in resolving the trichotomy (Dipseudopsis + Protodipseudopsis + Limnoecetis). One obvious prediction based on our phylogeny, is that the unknown larval forms of Hyalopsychella and Limnoecetis probably share many similarities with the dipseudopsid larvae known. The phylogeny also provides evidence for removing Eodipseudopsis from the Dipseudopsidae (cf. final section of text).

The sister group relationship of the genus Phylocentropus remains uncertain. Phylocentropus phenetically agrees closely with Hyalopsyche, but the similarities of these two genera are predominantly plesiomorphic. Thus, the phylogenetic analysis above does not provide supporting evidence for Hyalopsychidae s. str. Schmid (1980) and in fact, infers that this taxon might be paraphyletic. Therefore we concur with Ross and Gibbs for the most part on the classification of Phylocentropus, and Wells and Cartwright on the placement of Hyalopsyche and the suppression of the Hyalopsychidae, a taxon that presently serves no useful purpose in phylogenetics.

## Keys to genera of Dipseudopsidae

## Larvae

The larvae of Hyalopsychella Ulmer and Limnoecetis Marlier are unknown.

1. Mandibles with apicolateral teeth; head capsule short, with lateral margins nearly parallel and its total length less than its maximum width (Ulmer 1957, Gibbs 1968) 2
Mandibles without apicolateral teeth; head capsule long, with lateral margins tapering anteriad, and with its length slightly greater than its maximum width (Wiggins 1977, Wells \& Cartwright 1993) 3
2. Frontoclypeal suture V-shaped with lateral lines tapering irregularly posteriad toward the dorsal coronal suture (Ulmer 1957)

Dipseudopsis Walker

- Frontoclypeal suture with the posterior portion U-shaped, and the anterolateral lines nearly parallel between the eyes and then curving mesad to meet the coronal suture posteriad (Gibbs 1968)

Protodipseudopsis Ulmer
3. Frontoclypeal suture sessile in respect to posterior margin, posterior $1 / 3$ nearly regular $V$-shaped and without hourglass-shaped constriction (Wells \& Cartwright 1993) .. Hyalopsyche Ulmer

- Frontoclypeal suture with short petiole joining posterior margin, posterior $1 / 3$ more irregular $V$ shaped and with hourglass-shaped constriction (Wiggins 1977) $\qquad$ Phylocentropus Banks


## Adults

1. Pronotum with setaceous warts conspicuous and without deep median fissure
.. 2

- Pronotum with setaceous warts inconspicuous and with deep median fissure $\qquad$

2. Labial palps present; forewing with fork I long and sessile $\qquad$ . Phylocentropus Banks

- Labial palps absent; forewing with fork I short and petiolate or absent $\qquad$ 3

3. Fore and hind wings with fork I short and petiolate $\qquad$ Hyalopsyche Ulmer

- Fore and hind wings with fork I absent $\qquad$
Hyalopsychella Ulmer

4. Tibial spurs 1, 2, 2; female with maxillary palps three-segmented; one species endemic to Lake Tanganyika $\qquad$ Limnoecetis Marlier

- Tibial spurs 3, 4, 4; female with maxillary palps five-segmented; Africa and Asia. $\qquad$

5. Mouthparts with maxilla modified into a proboscis; hindwing with median cell closed; male hind mesoapical tibial spur modified, often twisted and/or branched; Asia and Africa

Dipseudopsis Walker

- Mouthparts without a proboscis; hindwing with median cell open; male hind mesoapical tibial spur normal; Africa ...... Protodipseudopsis Ulmer


## Systematic part

## Dipseudopsis Walker, 1852

Dipseudopsis Walker, 1852: 91. - Type species: Dipseudopsis capensis Walker (monobasic).
Nesopsyche McLachlan 1866: 168. - Type species: Neopsyche flavisignata McLachlan (monobasic).
Esperona Navás 1915: 397. - Type species: Esperona orientalis Navás (monobasic).
Bathytinodes Iwata 1927: 235. - Type species: Bathytinodes albus Iwata (monobasic).

Dipseudopsodes Lestage 1936: 170 (invalid, type species not designated).

The genus Dipseudopsis was proposed by Walker (1852) with the description of D. capensis from South Africa and placed in the family Sericostomatidae Stephens, a family designation which was common for caddisflies, exhibiting bizarre characteristics, that did not agree with the diagnoses of other families. Later when McLachlan (1863) added a second species to the genus, $D$. collaris from China, he placed $D i$ pseudopsis in the family Rhyacophilidae. Brauer (1868) placed it in the Hydropsychidae s. lat., near Cyrnus Stephens. However, after examination of the adult mouthparts, Ulmer (1904a, b) placed the genus in the Polycentropodidae, subfamily Dipseudopsinae. Accordingly, most workers have placed the genus in Polycentropodidae (Navás 1913, Martynov 1935, Schmid 1949). However, Banks (1931b) maintained a conservative approach, having never recognized the full family status of Polycentropodidae, and followed Brauer's placement of the genus in the Hydropsychidae s. lat.

Etymology. - Dipseudopsis: Greek di, two; pseudos, false; ops, eye; feminine. The actual etymology is uncertain. The basis of this name is uncertain also, since there are no noticeable characters such as wing spots present in the type species of the genus, $D$. capensis. Perhaps it was based on the appearance of the pronotum, having the setal areas expanded and divided by a deep median fissure.

## Adult

Head and thorax (figs. 12A-B, 13, 14): The head has a sparse number of short inconspicuous setae and is nearly glabrous; sometimes it is glossy and very glabrous. The setal areas of both the head and thorax are not raised like typical setaceous warts, but are flush with other surrounding sclerites. The setae of the body and wings are mostly short sparse and inconspicuous, giving the imago a glabrous appearance. The pronotum is raised to the same level as the head and mesonotum and is divided by a deep median furrow, probably resulting from modifications of the lateral setal areas, which have remained separate, but have become both enlarged and glabrous. The tegula are flat and sit like epaulets on the bases of the forewings. Legs with tibial spur formula complete: 3, 4, 4. The head, thorax, and sometimes the anterior segments of the abdomen have patterns of yellowish or orangish brown contrasting with dark brown. The forewing is narrow, length ranging from 10 to 20 mm , usually mostly brown with a hyaline spot at m -cu and the arculus, and sometimes with a pattern of translucent
bands or spors against a dark background, or with a striped pattern of dark veins against lighter translucene membranes; fork 1 is either short and petiolate or absent, II and IV are long arid sessile, III and V are long and periolate, and the discal and medial cells are closed. The hindwing is shorter and sometimes broader than the forewing, being somewhat triangular and lighter in colour, sometimes nearly translucent; forks II and V are long and sessile, forks I, III, IV absent, the discal cell closed, and unlike other dipseudopsid genera described the median cell is closed. The scent glands of sternum $V$ are present in the female and absent in the male, their openings are rather inconspicuous and lack exterior projections. This condition differs in Phylocentropus placidus (Banks) examined, having the scent gland absent in both sexes.

The mouthparts (fig. $12 \mathrm{~B}, 13,14$ ) have been described by Ulmer (1904a), Cummings (1913) and Crichoon (1957: fig. 53), and are quite unique, having the lacinia modified into a proboscis, functioning as a siphoning tube. The two processes which form the proboscis are not held together in preserved specimens, but curl away from each other distally. Each process is shaped like half of a long slender straw, with its longitudinal edges fringed with numerous short scalelike setae, having many transverse ridges along its shaft (thus being semiannulate), and having several mesal dentations at its base. The proboscis is at least as long as the first $2-4$ segments of the maxillary palps, or usually about the same length as the labium, including base plus palps. The haustellum is slender, about as long as the base of the labium, and its apex is bilobed. The frontoclypeus is large and slightly bulbous, apparently well-developed to support a pumping organ. Protodipseudopsis differs noticeably from Dipseudopsis by lacking a well-developed proboscis (fig. 12C); its frontoclypeus is shorter and less bulbous and apparently the pumping organ is reduced.

Male. - The male is dark brown, sometimes with contrasting patterns in the wings and body, and more striking than in the female. The mesoapical spur of the hind tibia is modified, often twisted like a corkscrew and sometimes branched into two or more processes. It is interesting to note that the same spur is also modified in Phylocentropus auriceps (Banks), (Schuster \& Hamilton 1984), being exceptional among the other dipseudopsids.

Genitalia (fig. 17): The male genitalia are blunt, generally lacking in conspicuous characteristics, and the processes of the genitalia are retracted and obscured by the large blunt preanal appendages. This blunt appearance of the male terminalia, especially in dry material, probably is why males sometimes have been mistaken as females, e.g. the types of $D$. flavisig-
nata (McLachlan, 1866) and D. benardi Navás (1930). Segment IX with articulation between tergite and sternite, resembling an open jaw in lateral view; the tergum and sternum also articulate with the preanal appendages. The tergum of segment LX is much smaller than the sternum and usually overhangs segment X , and its apical margin may be triangular, rounded, or bilobed in dorsal view. The sternum of segment IX is heavily sclerotized and is the most massive feature of genitalia, providing support for the inferior appendages, the phallus, and the ventral articulation of the preanal appendages. The sternum of segment IX is a simple structure except for the mesosuperior processes that, along with segment $X$, form a hood that hangs over the dorsal portion of the phallus which is mostly membranous. Segment X is inferior to segment IX, and subequal in length to the phallus. The phallus is mostly membranous dorsad and heavily sclerotized ventrad. Hence, segment $X$ and the phallus appear to function in concert, with segment $X$ providing dorsal coverage and support to the phallus. The inferior appendages are thumblike, with apices directly dorsad, perhaps modified to hold segment X and the phallus together, as well as clasping the female during copulation.

Female. - Females are usually lighter in colour than the males, and as a result they are more dull and have fewer contrasting colour patterns in their wings and bodies. However, the contrasting colouration of an unidentified female specimen (fig. 2) may be exceptional. In several species, such as $D$. nervosa (figs. 3, 4), where the males have brown wings with large translucent spots, the females have striped forewings with dark veins against a translucent background. Sometimes females have forewings with faded indistinct spots that are remotely similar to translucent spots in the forewings of males. However, the female forewings are most often dull brown and show no striking colour patterns.

Genitalia (figs. 15-16): Sternum VIII forms a single plate, not divided into lateral lobes as in the Polycentropodidae. Segment IX has a well-sclerotized tergite, as in some species such as D. recta (fig. 15), but is more membranous in others (fig. 16); the sternum is mostly membranous. The vaginal apparatus and the vulval scale are attached to the sternum of segment IX by sclerotized bands and membranous folds. Both the vaginal apparatus and the vulval scale are sclerotized. The length of the vaginal apparatus is variable, being long in $D$. recta, where it extends anteriad reaching the anterior margin of sternite VIII (fig. 15), but is much shorter in other species (fig. 16). The genitalia are generally similar to those of Phylocentropus (Schmid 1980, Schuster and Hamilton 1984), except that segment IX is more heavily
sclerotized. This diagnosis is provisional, because the females of most species remain undescribed. We observe that the females of some species can be distinguished from others by colour patterns of the body and wings, and characteristics of the genitalia.

## Larva

The larva of Dipseudopsis was first described by Iwata (1927), although under a different name. A key separating the larvae of Dipseudopsis and Protodipseudopsis was provided by Ulmer (1957), and a similar key was provided by Marlier (1962), whereby the larvae of Dipseudopsis and Protodipseudopsis were distinguished by the gular sclerite, mandibles, and hind tarsal claw. Protodipseudopsis and Dipseudopsis are unique in having mandibles with apicolateral teeth, a probable synapomorphy (Ulmer 1957: f. 401; Gibbs 1968: fig. 1; Marlier 1962: fig. 50; Marlier 1979: fig. 5). The known larvae of Dipseudopsis known can be distinguished from those of other dipseudopsids by having head capsule short, lateral margins nearly parallel with total length less than its maximum width, frontoclypeal suture $V$-shaped and the lateral lines tapering irregularly and posteriad toward the dorsal coronal suture, and mandibles with apicolateral teeth.

Larval behaviour and habitat. - The larval tubedwelling and feeding-behaviour have been described for Protodipseudopsis (Gibbs 1968) and Phylocentropus (Wallace et al. 1976, Wiggins 1977). The larvae of these genera live in the bottom substrates of streams or lakes and construct a somewhat Y-shaped silken tube composed of sand grains and detritus. The ends of the two upper branches, usually one vertical and the other inclined, are open and exposed to the stream current, the rest of the tube is located within the bottom substrate, and the end of the bottom branch is closed. The opening of the vertical branch is an intake and the inclined branch an outlet, through which a directional flow is established via rapid oscillations of the larva. This forced flow enables a fine capture net, within an enlarged chamber of the outlet branch, to collect fine organic food particles for the larva to eat. The larval tubes of Hyalopsyche are branched and appear to be similar to those of Phylocentropus (Wells \& Cartwright 1993), as are the tube-dwellings of Dipseudopsis. Therefore, since branched tube-dwellings are characteristic for the larvae of Dipseudopsis, Hyalopsyche, Protodipseudopsis, and Phylocentropus, this type of larval retreat and its associated mode of feeding-behaviour is a synapomorphy of Dipseudopsidae. The larva retreat of the hydropsychid genus Macrostemum Kolenati (Wiggins 1977, as Macronema Pictet) is generally similar, but differs in many details and is certainly a homoplasy.

Ulmer's (1957: 303) description of the larval retreat and biology of Dipseudopsis is translated as follows: 'Larvae and pupae live in calm pools of rivers, also in lakes and lake-outflows, in the mud or fine sandy substrates; the larvae not free-living there, but construct tubes out of secretions, covered with mud etc., in the depths; these tube-dwellings are more or less bent and twisted, elastic and flexible, but nevertheless fairly resistant; sometimes they are branched into 2 or 3 tubes, but only one of the branches reaches the [substrate] surface and there it is closed, while the other lower ends are open.' It seems obvious that Ulmer's orientation of the tube-dwelling is upside down; his description should depict the two open ends reaching the surface, and the one closed branch completely buried in the substrate. This would agree more with the tube-dwellings described in detail for Phylocentropus and Protodipseudopsis, and the description by Scott (1985: 334) for Dipseudopsis in southern Africa: 'The larvae are bottom dwellers in standing or slow-flowing water where they construct U-shaped silk-lined tubes with a transverse net in mud or silt or on submerged branches, filtering out algae and organic particles on which they feed.'
The larvae and pupae of Dipseudopsis inhabit a range of freshwater habitats, including the sandy depositional zones of lakes and the pool areas of rivers and streams. Marlier $(1953,1962)$ reports that immatures in Africa were found in shallow water, while Tsuda (1939) reported that larvae in Japan were discovered at a depth of 90 m . The collections of $D$. diehli Malicky \& Weaver, at elevations of $1,400 \mathrm{~m}$ in Sumatra, devoid of large rivers and lakes, suggests that some species also live in mountain streams.

## Distribution

Dipseudopsis is known to occur generally throughout the old world tropics of Africa and Asia. The African fauna recorded from the northern Nile River Valley, including Egypt, Sudan, and throughout most of the Afrotropical Region, from Ethiopia and Kenya to Nigeria, Ghana and Sierra Leone, and to Zaire, Mozambique, South Africa, and Madagascar. The genus is represented by 42 described species, 23 from the African continent, and 19 from Madagascar. However, in light of the problems discovered in this work, we suspect that the African species are in need of a thorough taxonomic revision. The Asian fauna with 36 recognized species is known throughout most of the Oriental Region and part of the southeastern Palearctic Region, including Pakistan, India, Sri Lanka, Nepal, Bangladesh, Burma, Thailand, Cambodia, Vietnam, China, Japan, Philippines, Malaysia, and Indonesia, including Sumatra, Java, Borneo and Sulawesi.

## Checklist of Dipsewdopsis from Asia

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*adsafuerix sp. n.
benarrdi Navas, 1930.
    Langazht Navás, 1930 syn. n.
    arctulata Navás, 1931 syn. n.
    ubmeriSchmid & Denning, 1979 syn. n.
bisolorata Martynov, }1935
collaris McLachlan, }1863
    srelluse McLachlan, 1875 syn. n.
    a/b,a (Iwata, 1927).
    bakeri Banks, }1916\mathrm{ syn. n.
    discors Navás, 1924 syn. n.
ronforta Banks, 1931b.
dichli Malicky & Weaver, }1988
digirala Ulmer, 1907a.
doehleri Ulmer, 1929 (döhlers).
clongasa Banks, }1920
fluzisignaza(McLachlan, 1866)
flintisp. n.
immaculata Ulmer, 1905.
    moeska Banks,1931b syn. n.
indica McLachlan, 1875.
    buddha Banks, 1913 syn. n.
    i. sindica Martynov, }1935\mathrm{ syn. n.
infuscata McLachlan, }1875
knappiSchmid & Denning, 1979.
lamellata Martynov, 1935.
lucasisp. n.
maculata Ulmer, 1907a.
malaisei sp. n.
martynozi sp. n.
modesta Banks,1911.
    pallida Martynov,1935 syn. n.
morosa Banks,1924.
nebulosa Albarda, 1881.
nervosa Brauer, 1868.
    luctuosa Banks, }1913
    venosa Navás, 1931 syn. n.
nervosellat Ulmer, 1951 stat. n.
p. }12
nieuwenhuisi Ulmer, 1909.
p. }13
notaka(Fabricius, 1781).
    hormi U'lmer, 1915 syn. n.
recha Martynov, }1935
p. }10
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p. 127
luctuosa Banks. 1913.
venosa Navás, 1931 syn. n.
nervosella Ulmer, 1951 stat. n.
p. 128
notata (Fabricius, 1781).
p. 130
p. 130
recta Martynov, 1935.
p. 131
bombayana Martynov, 1935 syn. n.
pobustior robustior Ülmer, 1929 stat. \(\mathbf{n}\).
p. 132 akhila Schmid \& Denning, 1979 syn. n. -hatandica Schmid \& Denning, 1979 syn. n. iunki Marlier. 1979 syn. a.
\begin{tabular}{|c|c|}
\hline \multirow[t]{3}{*}{\begin{tabular}{l}
robustior andamanensss s.sp. n. tmidisp. n. \\
- . bilis Banks. 1931a.
\end{tabular}} & p. 134 \\
\hline & p. 134 \\
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\hline -.batenses Malicky \& Weaver, 1988. & p. 135 \\
\hline : mkemensss Navás. 1921. & p. 135 \\
\hline - ata Martyov, 1935. & p. 137 \\
\hline 29. & p. 138 \\
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    petersorum Schmid \& Denning, 1979 syn. n.
    voluta Ulmer, 1906.
p. 138 voluta Ulmer, 1906.
p. 138 onychophora Navas, 1935 nomen dubium
p. 139 orientalis (Navás, 1913) nomen dubium
p. 139

## Dipseudopsis adiaturix sp. n.

(fig. 18)
Type material. - Holotype ơ: INDONESIA: Sumatra, Huta Padang, 26.I.1990, Diehl (CLHM).

Male. - Head with dorsum and frontoclypeus dark brown, almost black, but posterior dorsal warts yellowish brown. Basal segments of antennae brown (type with distal parts of antennae and mouthparts missing due to ant attack). Thorax, tergites and sternites dark brown. Coxae of fore and mid legs brown, coxa of hind legs yellowish brown. Basal $1 / 2$ of fore femora dark brown, femora of mid and hind legs with basal $1 / 2$ brownish; distal $1 / 2$ of all femora, tibia of fore and mid legs, and all tarsi yellow; distal $1 / 2$ of hind tibia brown. Modified spur (fig. 18) long and slender, more than $1 / 2$ as long as adjacent tarsal segment, more than $2 \times$ longer than outer spur, distal claw about $1 / 8$ as long as spur and bent obliquely laterally, and with short lateral thorn at $2 / 3$ length from base. Forewing 16 mm , slender, dark brown, but with nervation darker, having hyaline lunula at $\mathrm{m}-\mathrm{cu}$, and indistinct translucent spots between $\mathrm{M}_{4}$ and $\mathrm{Cu}_{1}$ and near apex of 1 A and 2 A . Hindwing 10 mm , dark brown, with hyaline lunula at m-cu. Abdomen yellowish brown. Genitalia (18A-D): Tergum IX broad and bilobate with median sized cavity below in dorsal view; sternum IX with mesosuperior process fingerlike in lateral view. Dorsal edge of segment X with strong incision in lateral view, but apex not incised. Preanal appendages long and triangular in lateral view. Inferior appendages curved dorsad in lateral view, with distal part slightly clavate and basolateral mesal margins squarish in caudal view. Phallus calyx-like in ventral view, about $2 / 3$ as long as segment X .

Etymology. - A noun in apposition: named after a Celt who lived in Austria 2000 years ago.

Distribution. - Indonesia: Sumatra.
Remarks. - This species is only known from the unique type. It has a modified spur similar to $D$. flavisignata, being long slender, acuminate, slightly sinuate and bearing a short subapical lateral thorn. However, it differs in having the male modified spur with the lateral thorn closer to its apex, rather than near the base, the male forewings mostly dark brown and without a striking pattern of translucent stripes and by characteristics of the male genitalia.

Pictorial atlas (p. 103-106)
A pictorial atlas is provided for quick species determination. The figures of the following male characters are provided for each species: 1) dorsal aspect of segment IX and X, 2) shape and relative size of the left modified spur, and 3) caudal aspect of the right infe-

## D. nebulosa Albarda



Burma, Thailand, Sumatra, Malay Pen.

D. immaculata Ulmer


Sumatra, Borneo, Malay Pen.

D. schmidi Weaver and Malicky


India, Bangladesh
rior appendage (but for subspecies, only figures of the modified spur). The distributional data are listed below the figures. Possible identifications should be confirmed with the complete species descriptions and figures in the text.


Sulawesi
D. lamellata Martynov
D. varians Ulmer
D. doehleri Ulmer

Assam, Burma, Thailand


Thailand, Malay Pen.


D. collaris McLachlan

China, Japan, Phllippines
D. robustior Ulmer


Burma. Thailand, Vietnam, Cambodia, Malay Pen.
D. r. andamanensis Weaver and Malicky Andaman Is

D. infuscata McLachlan


Sumatra, Borneo, Java, Sulawesi
D. diehli Malicky and Weaver

D. knappi Schmid and Denning

D. nervosa Brauer

D. notata (Fab.)

India. Sri Lanka

Dipseudopsis berardi Navás, 1930
f1es. 10.191
"5 steluatia - L'Imer 1905: 96, fig. 75 b [misdet]. isenards へavas, 1930: 141, Type $0^{*}$ ", IIFINAM Phu I ang Thunng. 1909, det. P. Navás S. J.,
, same data ( M : $\mathrm{H}: \mathbf{\circ}$ ). - Fischer

1962: 5; Fischer 1972: 3.
Dipseudopsis langana Navás, 1930: 141, Type ㅇ*, vietnam $^{*}$ 'Tonkin': Phu Lang Thuong, 1909, det. P. Navás S. J., G. Benard (MNHN). - Fischer 1962: 10; Fischer 1972: 5. Syn. n.
Dipseudopsis arculata Navás, 1931: 85-86, Holotype o*, China: Guangfong: 'Cong-tcheou' Canton (MZBS). Fischer 1962: 5; Fischer 1972: 2. Syn. n.

## D. morosa Banks


D. nervosella Ulmer

D. maculata Ulmer


Borneo

D. digitata Ulmer


Philippines


Dipseudopsis moesta. - Ulmer 1951: 128 [misdet].
Dipseudopsis ulmeri Schmid \& Denning, 1979: 243, fig. 1ae, Holotype on $^{*}$, Thailand: Chiang Mai Prov., E fork Mae Ping, 56 km N of Chiang Mai, 1300ft, at light, 24.XI. 1964, Peters (CldD). Syn. n.

Specimens examined. - CAMBODIA: $10^{\star} 2$ 아, 'Cambodge'

## D. bicolorata Martynov


D. contorta Banks


Sulawesi, Borneo, Java, Malay Pen.
D. martynovi Weaver and Malicky

D. tonkinensis Navás


Vietnam, Hainan


Erratum: D. contorta, read Sumatra for Sulawesi.
Pavie, 1886 (MNHN). - CHINA: Fujian: I © , 'Foochow' Fuchou, 1936-37, det. D. stellata by Mosely, M. S. Yang (bMNH). - THAiland: 2 ô, Chiang Mai, 19.V.1952, D. \& E. Thurman (USNM). 10 , Doi Sutep, 1952, D. \& E. Thurman (MCZC). 1才, Bangkok, 5.X.1929, H. Smith (MCZC). 1才, Ping River, Chiang Mai, 9.IV.1989, Malicky (Clhm). 20 , Ping River near Lampun, 20.XII.1989, Malicky (CLHM).
D. stabatensis Malicky and Weaver

D. flinti Weaver and Malicky

D. malaisei Weaver and Malicky

D. benardi Navás


Thailand, Vietnam. Cambodia, China

## D. recta Martynov



[^4]/(te). light trap. 24-31.X.1988, (11:9). 22. Kanchanaburi. 14. V. 1988: Allen (co HMi).

1 montly dark brown but pos-

## D. modesta Banks


D. triclavata Martynov


India, Sri Lanka

D. lucasi Weaver and Malicky

terior warts light brown; frontoclypeus light brown but dorsal portion darker; antennal scape and pedicel light brown, flagellum dark. Pronotum, mesonotum and tegula dark brown. Forewing (fig. 10): 13.5-16.5 mm , mostly dark brown, but with translucent spots: distinct small translucent spot just distal of r ; small


Figures 1-2. - Fig. 1, Dipseudopsis bicolorata Martynov, male, dorsal. Fig. 2, Dipseudopsis sp. female, dorsal, India, Manipur, Nungha.
faded spots just distal of transverse cord, hyaline lunula at $\mathrm{m}-\mathrm{cu}$, large translucent spot at arculus. Legs brown. Modified spur (fig. 19): basal $3 / 4$ unmodified and apical $1 / 1 /$ bifid with two short acuminate points curved, spiraling opposite each other distally, each point slightly longer than width of main shaft. Genitalia (fig. 19): Tergum IX with posterior margin curved in dorsal view, with posterior $1 / 2$ extending over segment X in lateral view; sternum IX having mesosuperior process with short dorsal triangular point and long slender distal point, extending above basal $1 / 2$ of phallus in lateral view. Segment X tapering distally into slender lobe in lateral view, extending posteriorly distal to preanal appendages about as far as inferior appendages, apex bilobed. Preanal appendages with posterior margin not distinctly incised in lateral view, having blunt dorsal point, dorsal and ventral margins converging to truncate posterior margin. Inferior appendages thumblike with obtuse midventral angle in lateral view.

Distribution. - Cambodia; China: Guangdong, Fujian; Thailand; Vietnam.
Remarks. - This species can be recognized by having the modified spur with two short apical points
spiraling opposite from each other, and the forewing longer and darker brown than most other species in southern China and Indochina. The types of $D$. benardi, D. arculata, D. langana, and D. ulmeri were examined, and the latter three are recognized as junior synonyms of $D$. benardi. The type of $D$. benardi is a male, not a female as previously reported by Navás (1930). The type of D. langana is recognized as a female of $D$. benardi, because it is has an identical locality label as the type of $D$. benardi and is similar in size and general appearance.

Ulmer (1905: fig 75b) provided a figure of the male modified spur of an undetermined specimen of Dipseudopsis from 'Cambodscha, Museum Paris' and Ulmer (1951: 128) later identified this figure as $D$. moesta (syn. n. of D. immaculata). Ulmer's figure is more similar to the spur of $D$. benardi, and is probably not $D$. immaculata because this species is unknown from Cambodia. Also, we have examined a male specimen of $D$. benardi from Cambodia in the MNHN, which might be the same specimen figured by Ulmer.

Dipseudopsis bicolorata Martynov, 1935 (figs. 1, 20)
Dipseudopsis bicolorata Martynov, 1935: 165-166, figs. 68ac, 69a-b, Type series: IndiA: ' 2 §', ㅇ. Castle Rock, North Kanara district, Bombay Pres. October 1916. S. W. Kemp ... ס. Talewadi, Castle Rock, N. Kanara district, Bombay Pres. 3-10.1916. S. W. Kemp'. The type material present in the collection of the Zoological Survey of India is in poor condition and too delicate to send by mail (pers. comm. Dr. K. Rai). Syntype 1 $^{*}$ : det. Martynov, Karnataka, N. Kanara dist., Castle Rock, in alcohol, X.1916, S. Kemp (ZRAS). - Fischer 1962: 5; Higler 1992: 80.

Specimens examined. - INDIA: Karnataka: $1 \delta^{\hat{N}}$, Shimoga Dist, Agumbe Ghat 2000ft, V.1974, Lucas (Clhm). Tamil Nadu: 4ず, Kattalaimala, 25-26.XII.1961, F. Schmid (CNCI). 1 or', Ottakada, 5.I. 1962 , F. Schmid (CNCI). $_{\text {. }}$

Male. - Body and wings (fig. 1). Head dorsum and pronotum orangish brown, frontoclypeus mostly dark brown but dorsal margin orangish brown, genae yellowish, antennal scape orangish, pedicel and flagellum brown, palps dark brown, but stipes orangish brown, maxillary palps 3.2 mm , proboscis 1.7 mm , as long as first three segments of maxillary palps, labial palps 1.4 mm ; cervical sclerites and mesonotum dark brown. Forewing $16-21 \mathrm{~mm}$, almost unicoloured dark brown, but darker along anterior margin between C and R and posterior margin, only hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Front leg brown, middle and hind legs with coxae, trochanters and femora orangish brown, distal segments brown. Modified spur (fig. 20) having one apical point, curving in

-psis nervosa Braucr: Fig. 3, male. Fig. 4, female. Fig. 5, D. doehleri Ulmer, male. Fig. 6, D. D. nebulosa Albarda, male. Fig. 8, D. digitata Ulmer, male. Fig. 9, D. spectabilis Banks, male malc forewing. Fig. 11, D. flinti, n. sp., male forewing.

Figures 12A-C. - Fig. 12A, Dipseudopsis nebulosa AIbarda, male, head and thorax, dorsal. Fig. 12B, $D$. doehleri Ulmer, male, head with mouthparts, ventral. Fig. 12C, Protodipseudopsis sjoestedti Ulmer, male, head with mouthparts, anterior.

plane perpendicular to longitudinal axis of spur.
Genitalia (fig. 20): Tergum IX triangular with blunt apex in dorsal view, posterior margin only slightly overhanging segment X in lateral view. Segment X ellipsoidal with length $11 / 2 \times$ basal width in dorsal view, tapered distad and apex incised. Preanal appendages with dorsal margin irregular in lateral view, directed dorsad with apex obtuse or sometimes rounded and ventral margin broadly rounded. Inferior appendages with massive dorsal extension that doubles height at middle as compared to base in lateral view, with distinct apicomesal tooth in ventral view, and with curved apicolateral flanges.

## Distribution. - India: Karnataka, Tamil Nadu.

Remarks. - This is the largest species of Dipseudopsis known from India and is easily recognized by its colouration of the males which superficially resemble ctenuchid moths. The orangish brown dorsum of the head and pronotum contrast strikingly with the unicoloured, dark brown mesonotum and forewings. The modified spur bears a single curved apical point that lies in a plane perpendicular to the longitudinal axis of the spur. The specimen in fig. 2 is possibly a female of this species.

## Dipseudopsis collaris McLachlan, 1863

(fig. 21)
Dipseudopsis collaris McLachlan, 1863: 496, pl. 19: fig. 6, Lectotype đ*, selected by Kimmins (1957: 101), Hong Kong: (вмNн). Cotype $1 \mathbf{\delta}^{\boldsymbol{*}}$ : Hong Kong: Soochow (MCZC 11080). Weidner (1964): Paratype if, ChINA: Hong Kong (ZMUH). - Fischer 1962: 7; Weidner 1964: 73; Fischer 1972: 3.
Dipseudopsis stellata McLachlan, 1875: 16-17, p1. 2: fig. 11, Holotype ơ (Kimmins, 1957: 101): [CHINA: Zhejiang:] Shanghai, McL. coll. (BMNH); $3 \delta^{*}$ [possible syntypes]: Shanghai, McL. coll. (BMNH); 'Paratype' 1 O*: 'N China' McL. coll (вмNH). - Fischer 1962: 15. Fischer 1972: 7-8. Syn. n.
nec Dipseudopsis stellata. - Ulmer 1915: 73 (Sri Lanka); Lestage 1925: 65, 115 (Java); Banks 1931b: 401 (stellatus, Thailand); Tsuda 1942: 266 (Thailand); Schmid 1958: 35, 89 (Sri Lanka).
Dipseudopsis nebulosa, - Banks 1914: 178 (Luzon) [misdet.].
Dipseudopsis bakeri Banks, 1916: 215-216, pl. 2: figs. 20, 21, Type ${ }^{\text {ot }}$, philuppines: Luzon, Laguna, Mt Maquiling and Los Baños,Baker (MCZC 11768). Only $1 \delta^{\text {o }}$ at the mCZC labelled as type, although several other specimens, not labelled as types, at the MCZC and USNM are part of the type series. - Fischer 1962: 5; Fischer 1972: 3. Syn. n.
Dipseudopsis discors Navás, 1924: 205. - Fischer 1962: 8. Syn. n.
Dipseudopsis morosa. - Banks 1924:450 (pro parte). Syntype $1 \delta^{*}$ is actually D. collaris: Luzon: Mt Maquiling, Baker (MCZC 14826).
Bathytinodes alba Iwata, 1927: 209-210, pl. 4: fig. 87-91.
Dipseudopsis alba (Iwata, 1927). - Fischer 1962: 217; Fischer 1972: 2.


13－14．－Fig．13．Dipseudopsis sp．，male head，anteri－ merdified after（Crichton 1957）．Fig．14，Dipseudopsis onehteri L＇Imer，male head and prothorax，lateral．

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\begin{aligned}
& \text { 'peumens examined. - (111NA: Guangdong: } 30^{\circ} \text {, }
\end{aligned}
$$

19（ts）as D．stellata．＇Kiang－si＇，1869，A．David
Hong Kong：18．Foochow（mezc）．Jiangsu： $60^{\circ}$ ，
Fulm．18．Vitit．198．Tian（cthms）． 8819 ，Jinhu，
＇hek ang Hangreheou＇．Hangrhou，1925，A．Pichon
thanghai．det．McLachlan as D．stellata，
vhumn．20．V］．195－，Tian（（IHM）．APAN： $10^{0}$ ．
－Mivis：Luzon： 8319 Los
Wr M）．If（BMNHI）．10，Manila
Wt Maquiling．Baker（ ©N：M）， 20
wings and legs with colour－ mon strmulat to it mbustior．Forcwing $13-15 \mathrm{~mm}$ ， ．int small translucent spots
around discal cell，having most spots distal to cord， and spot at arculus adjoining hyaline area at apex of $\mathrm{Cu}_{2}$ ．Modified spur（fig．21）apical $1 / 3$ bifid，crescent－ shaped，with two subequal curved points．Genitalia （fig．21）：Tergum IX triangular in dorsal view，slight－ ly overhanging segment X in lateral view；sternum IX mesosuperior process ellipsoidal with blunt apex pointed posteriad，extending above basal $2 / 3$ of phallus． Preanal appendages with posterior margin slightly in－ cised in lateral view，dorsal margin at right angle to posterior margin，ventral margin at obtuse angle， sometimes varied with ventral margin continuous with posterior margin．Inferior appendages fingerlike with short basoventral shelf in lateral view．

Distribution．－China：Guangdong，Hong Kong， Jiangsu，Jiangxi，Zhejiang；Philippines：Luzon；Japan： Honshu．

Remarks．－This species is similar to D．robustior， but differs by having inferior appendages more broad and fingerlike in lateral view（although this is vari－ able）and modified spur with two apical points both crescent－shaped and subequal in length．In D．robust－ ior the inferior appendage is more slender and angled， its modified spur bears two points，one point slender－ er，more curved and more than $2 x$ longer than other． Previous records of $D$ ．collaris，as $D$ ．stellata，from Sri Lanka，Java and Thailand are probably in error．

## Dipseudopsis contorta Banks， 1931

（fig．22）
Dipseudopsis sp．－Ulmer，1905：96，fig．75c（not fig．75e as listed by Fischer 1962）．
Dipseudopsis infuscata．－Ulmer 1905：96，fig．75d；Ulmer 1951： 128 ［prob．misder．］．
Dipseudopsis contorta Banks，1931b：400－401，fig．20，lecto－ type by present designation ot $^{*}$ ：MALAYsiA：West Ma－ laysia：Selangor，Kuala Lumpur，29．I．1924，at light，H． M．Pendlebury（BMNH）；paralectorype 10＊，without hind legs：Pahang，Kuala Tahan，300ft at light，23．XI．1921，H． M．Pendlebury（MCZC 16480）．－Fischer 1962：8； Malicky \＆Weaver 1988：3－4，fig．3a－e．

Specimens examined．$-20^{\circ}$ ，＇Borneo＇det．Ulmer 1905 as D．infuscata（MNHN）．IndONESiA：Kalimantan： 10 ， Pontianak，VII．1907，F．Muir（USNM）．Java：1才゙，［Muller？］ （RMinh）．Sumatra： $1 \delta^{\star}$ ，Dolok Merangir，19．VII－20．VI－ II． 1971 ，Dichl（Clhm）．10 ${ }^{\text {a }}$ ，same data，V－X．1980，Diehl （CI．hm）．10 O Ost－Aceh 26－30．VIII．1972，Diehl（CLHM）． 1才，NE Sumatra，16．I－18．II．1979，Diehl（CLHM）．

Male．－Head dorsum dark brown，except posteri－ or warts orangish brown，frontoclypeus dark brown， antennae orangish brown；cervical sclerites dark brown，pronotum light orangish brown and mesono－ tum dark brown．Maxillary palps 2.2 mm，brown；

Figures 15-16. - Fig. 15, Dipseudopsis recta Martynov, female: 15 A , genitalia, lateral; 15 B , genitalia, ventral; 15C, abdominal segment VII, lateral. Fig. 16, Dipseudopsis sp. female, 'Manipur Nungha' (CNC): 16 A , genitalia, lateral; 16B, genitalia, ventral.


16A

i
proboscis 1.6 mm , as long as first two segments of maxillary palps. Forewing $10-12.5 \mathrm{~mm}$, brown with row of lighter spots from stigma to $\mathrm{Cu}_{1}$ just distal to cord, and large translucent spot at arculus, hyaline spot at $\mathrm{m}-\mathrm{cu}$. Legs mostly light brown, except front and middle coxae dark brown. Modified spur (fig. 22) distal $1 / 3$ branched into two subequal curved points, with points diverging basally but converging distally, sometimes with rounded subapical shelf. Genitalia (fig. 22): Tergum IX semicircular in dorsal view, with posterior margin having almost no extension overhanging segment $X$ in lateral view; sternum IX with mesosuperior process short, slender, acuminate having apex curved dorsad in lateral view. Segment X ellipsoidal in dorsal view, long trapezoidal in lateral view, extended distad beyond apex of inferi-
or appendage. Preanal appendages variable, either Dshaped with lobe extending distally as far as inferior appendages in lateral view, or with posterior margin incised and with short dorsoapical blunt point. Inferior appendages with basoventral notch and apicoventral obtuse angle in lateral view, apical $1 / 2$ variable, either equal in height to base and truncate, or sometimes more expanded and clavate; with apex and apicolateral margin curved in ventral view, but bearing subapicomesal obtuse tooth.

Distribution. - Indonesia: Kalimantan, Sumatra, Java; Malaysia: West Malaysia.

Remarks. - This species is similar to $D$. collaris, but differs by having the inferior appendage capitate, with


Figure 17．－Fig．17．Dipseudopsis sp．，male genitalia，lateral．
a basodorsal notch and an apicoventral obtuse angle in lateral view，and the modified spur with apical points directed more laterally．Slight variation is no－ ticcable in the male modified spur and genitalia（cf． Malicky \＆Weaver 1988）．

## Dipseudopsis diehli Malicky \＆Weaver， 1988

（fig．23）
Dipseudopsis diehli Malicky \＆Weaver，1988：8，fig．6a－f． Holotype $\delta^{\circ}$ ：INDONESIA：Sumatra：Prapat， 1400 m ， ［）ichl（CI．HM）：paratypes $27 \mathbf{\delta}^{\text {：}}$ ：Prapat，Bukit Tinggi，Aek「arum．Sitahoan，Dolok Merangir（clhm，CLJW）．

[^5]Male．－Head dorsum yellowish brown with dark semicircular spot at anterior margin．Frontoclypeus brown．Scape yellowish brown，distal portion of an－ tenna brown．Maxillary palps 3.0 mm ，proboscis 2.4 mm ．extending as far as $41 / 2$ segment of maxillary palp． Cervical sclerites and pronotum yellowish brown． Mesonotum and regulac brown，with scutellum yello－ wish brown．Forcwing $16-20 \mathrm{~mm}$ ，brown with four translucent arcas：1）long stripe adjacent to costal margin：2）ellipsoidal spor between stems of $R$ and $M$ ； 3）transverse band distal to cord，between $\mathrm{R}_{1}$ and $\mathrm{M}_{4}$ ， with two $V$－shaped incisions along distal margin，and having narrow connection with anterior margin；4） －riangular spot at arculus．Hindwing unicoloured hrown，but with hyaline comma at $\mathrm{m}-\mathrm{cu}$ ．Legs yello－ vish brown，with light brownish tibiae，apparently
caused by more dense covering of fine brown setae． First segment of hind tarsi uniquely bent mesad about $1 / 3$ of its length to permit free motion of broad modified spur．Modified spur（fig．23）with apex bi－ fid with two sinuate points nearly parallel and slight－ ly unequal in length，both curved mesad，but apices curved distad．Genitalia（fig．23）：tergum IX with ap－ ical $1 / 3$ hanging over base of segment X in lateral view； broad and short with posterior margin bilobed or slightly triangular in dorsal view，anterior margin slightly bilobed．Sternum IX mesosuperior process extending to end of phallus．Segment X ellipsoidal in dorsal view．Preanal appendages triangular in lateral view with dorsal margin slightly inclined dorsad and ventroposterior margin curved steeply dorsad． Inferior appendages fingerlike and with ventral mar－ gin slightly concave in lateral view，but dorsal margin serrate in middle；with U－shaped basomesal notch in ventral view，distal arms slightly diverging apically and each tapering into small apical lobe．

Distribution．－Indonesia：Sumatra．
Remarks．－This species is easily recognized by hav－ ing the male modified spur extremely broad and bi－ furcate，with points curved a wide whorl，and by hav－ ing forewing with large translucent spots．Some specimens are darker brown，having brown parts that are usually yellowish brown in other species．

## Dipseudopsis digitata Ulmer， 1907

（figs．8，24）
Dipseudopsis digitata Ulmer，1907a：39－40，figs．56－57， Holotype $\delta^{*} *$ ：MALAYsia：Banguey Is，Staudinger（ZMHB）． －Fischer 1962：8；Fischer 1972： 3.

Specimens examined．－philippines：Balabac： $20^{\star}$ ， Dalawan Bay，5．X．1961；1才，9．X．1961；3ठ̊，12．X． 1961 （ZMUC）．Busuanga： $50^{\pi}, 4 \mathrm{~km} \mathrm{~N}$ San Nicolas， $20-$ 29．V．1962，H．Holemann（BPBM）．Culion： $6 \delta^{t} 19,6 \mathrm{~km}$ W Culion，7－11．VI．1962，light trap，H．Holtmann（BPBM）． Palawan： 10 ． ，Brooke＇s Point，Macagua， 75 m, 1－4．IV． 1962 （вPBM）． $19 \delta^{\circ} 1$ ㅇ，Tarumpitao Point，H．E．Milliron（BPBM）． $17 \delta^{7} 7$ ㅇ，Irawan， 14 km W Puerto Princessa，5m，D．R． Davis（UsNm）． $1 \delta^{\top}$ ，Litso，Amoyan $\mathrm{Ck} ., 57 \mathrm{~km}$ N Puerto Princessa， $10 \mathrm{~m}, 10 . \mathrm{XII} .1965$ ，D．R．Davis（UsNm）．10， Mainit， 11 km NW Brooke＇s Point，18．XI．1965，D．R． Davis（USNM）． $10^{*}$ ，Chromite Mine， 28 km W Puerto Princessa， $400 \mathrm{~m}, 1-7$. XII．1965，D．R．Davis（USNM）． $300^{\star}$ ， Brooke＇s Point，Uring Uring，14－23．VIII． 1961 （ZMUC）．

Male．－Body and wings（fig．8）．Head，thorax and forewing with pattern of light and dark bands．Head dorsum with dark brown median band widening an－ teriad and narrowing posteriad，lateral areas light brown and posterior warts lighter．Frontoclypeus mostly dark brown，but with yellowish V－shaped band pointed dorsad．Antennae light brown． Maxillary palps 2.3 mm ，proboscis 1.4 mm ，as long as


Figures 18-21. - Dipseudopsis, A-H, male genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior appendages, ventral; D, phallus, lateral; E, phallus, ventral; F, segments IX, X, dorsal; G, ventral; H, left inferior appendage, ventral. I-J, male modified apical spurs of hind tibia, ventral. Fig. 18, D. adiaturix, n. sp. Fig. 19, D. benardi Navás. Fig. 20, D. bicolorata Martynov. Fig. 21, D. collaris McLachlan, 21H, type of D. stellata McLachlan.


[^6]first three segments of maxillary palp. Cervical sclerites dark brown ventrally, light brown dorsally. Pronotum with raised areas light brown, median furrow and lateral sides dark brown. Mesonotum and tegula dark brown. Forewing $10-14 \mathrm{~mm}$, with striped pattern of dark brown veins and lighter cells, noticeably darker along thicker veins, $\mathrm{R}_{1}$ and $\mathrm{Cu}_{1}$. Legs with coxae dark brown and trochanters and distal segments light brown, except front coxa bicoloured, dark brown ventrad and lighter dorsad. Modified spur (fig. 24) with three minute apical points and one long acuminate subapical point curved laterally, all points distal to blunt subapical shelf. Genitalia (fig. 24): Tergum IX short and truncate in lateral view, posterior margin not overhanging segment X , rectangular in dorsal view; sternum IX small, with massive mesosuperior process having large broad dorsal lobe and slender posteroventral process extending above basal $2 / 3$ of phallus. Segment X with pair of minute basodorsal lateral lobes, dorsal margin angled obliquely ventrad in lateral view, but with apex turned dorsad, ventral margin straight but also turned dorsad apically; about $1 / 2$ as wide as tergum IX and about twice as long as wide in dorsal view, lateral sides parallel but tapered into triangular apex, and apex slightly incised. Preanal appendages broad and not incised in lateral view, somewhat triangular with dorsal margin and ventral margin curved and converging at blunt dorsal apex. Inferior appendages capitate in lateral view, dorsal margin concave, ventral margin straight with short basal shelf, apex truncate with corners slightly curved, extended dorsad.

Distribution. - Malaysia: Banguey Is.; Philippines: Balabac, Busuanga, Culion, Palawan.
Remarks. - This species is easily recognized, having the male segment X with a pair of minute dorsolateral lobes at its base, and modified spur with three minute apical points, one long acuminate subapical point curved laterally, and all points distal to a squarish subapical shelf.

## Dipseudopsis doehleri Ulmer, 1929

(figs. 5, 12B, 14, 25)
Dipseudopsis döbleri Ulmer, 1929: 189-190, fig. 27, Holotype $\delta^{*}$ : IndiA: Meghalaya: Khasi Hills, McL. coll (BMNH). - Fischer 1962: 8.
Dipseudopsis doehleri Ulmer. - Fischer 1972: 4; Higler 1992: 80

Specimens examined. - BURMA: 10 , Washaung, 20 km E of Myitkyina, $200 \mathrm{~m}, ~ 4 . V I I .34$, Malaise (NHRS). $10^{\text {T, }}$ Myitkyina, 175 m , 1934, Malaise (BMNH). INDIA: Assam: 1 §', 8 mi E Ledo, $125 \mathrm{~m}, 27.34 \mathrm{~N} 95.34 \mathrm{E}$, 13.X.1961, Ross \& Cavagnaro (CASC). 2 ㅇ, Abhoypur For., Naphuk, 360m, 12.X.1961, Ross \& Cavagnaro (CASC). THAILAND: 90,

Nakhon Nayok Prov., Khao Yai National Park, 700 m , 29.LX-6.X. 1984, Karsholt, Lomholdt \& Nielsen (ZMUC). $7 \mathbf{\sigma}^{\circ}$, Nam Nao, Pet Cha Boon, 5.V.1989, Allen (ClHM). 20 ${ }^{\circ}$, Puk Hieo, Chayapoom, 6.V.1989, Allen (ClHM).

Male. - Body and wings (fig. 5). Head and all of its appendages dark brown, except proboscis yellowish. Mouthparts (fig. 12B, 14): Maxillary palps 3.6 mm ; proboscis 2.7 mm , extended to midlength of apical segment of maxillary palp; haustellum 0.8 mm ; labial palps 1.7 mm . Cervical sclerites and prothorax ivory. Mesonotum, tegula and metanotum dark brown. Forewing 16-17 mm, dark brown with three distinct translucent spots: 1) basal triangle between $\mathrm{R}_{1}$ and $\mathrm{Cu}_{1}$ from base to fork I of Rs, 2) subapical spot distal to cord with apical margin incised at forks II, and III) ellipsoidal spot at arculus, similar as in $D$. nebulosa except for translucent spot at arculus. Legs dark brown, except front coxa with lateral side ivory. Hindwing with large translucent triangle as in front wing, apical region brown, anal region translucent, $\mathrm{Cu}_{1}$ dark brown. Modified spur (fig. 25) similar to D. immaculata, unbranched, slightly longer than adjacent spur, basal $2 / 3$ unmodified, apical $1 / 3$ twisted $180^{\circ}$ into short spiral, apical point directed mesad. Abdomen with anterior sternites II-V ivory, giving translucent triangles of forewings a white background so that contrasting pattern is displayed when wings are folded, sternites VI-VIII dark brown. Genitalia (fig. 25): Tergum IX triangular with posterior margin rounded in dorsal view, apical $1 /$ overhanging segment X in lateral view. Segment X long slender in lateral view, with short base having parallel sides in dorsal view, and apical portion cordate and apex deeply incised. Preanal appendages abbreviated with slender dorsoapical point in lateral view, but truncate and triangular in dorsal view. Inferior appendages long and thumblike, without incisions or points, curved slightly dorsad in lateral view.

Distribution. - Burma; India: Assam, Meghalaya; Thailand.

Remarks. - This species is most similar to $D$. nebulosa, but differs by having the head entirely dark brown and pronotum ivory, forewing with translucent round spot at the arculus, and modified spur relatively shorter, with less than $1 / 2$ of apical portion twisted. It also differs by having male genitalia with segment X and the preanal appendages more slender in lateral view, and segment X also bears a unique pair of basolateral humps in dorsal view. A teneral male specimen from Hainan Island, possibly an undescribed species near $D$. doehleri, was observed in the collection at the MCZC.

## Dipseudopsis elongata Banks, 1920

(fig. 26)
Dipscudopsis elongata Banks, 1920: 361-362. pl. 1: fig. 13, Tipe 8 $^{\circ}$ : MALAISIA: Sabah, Sandakan, Baker (MCZC 10882). - Fischer 1962: 8-9: Fischer 1972: 4.

Specimens examined. - Malaysla: Sabah: 10 º, Sandakan Bay: NW' Sepilok For. Res., 1-10m, 26.X.1957, J. L. Gressitt (APBM). $18^{\circ}$. 'North Bornco' McL. coll (BMiNH).

Male. - Head dorsum glossy dark brown, but posterior warts light brown, frontoclypeus glossy dark brown, genae yellowish. Cervical sclerites, pronotum, mesonotum tegula dull dark brown. Antenna brown, but scape light brown. Palps brown, maxillary palps 3.0 mm ; proboscis 2.0 mm , as long as first $31 / 2$ segments of maxillaty palp. Forewing 16 mm , uniformly dark brown, with small hyaline spots at nygma and m -cu, and large translucent spot at arculus. Front and middle leg with coxa, trochanter and basal $1 / 2$ of femur dark brown, and distal portion of leg light brown; hind leg light brown. Modified spur (fig. 26) long and truncate, about 3 x longer than adjacent spur, with apex more chan 2 X as wide as base, and minute apical points curved mesad. Genitalia (fig. 26): Tergum IX short and slightly overhanging segment X in lateral view, posterior margin rectangular with lateral edges curved in dorsal view; sternum IX with elongate mesosuperior process, extending above $3 / 1 /$ of phallus in lateral view. Segment X acuminate with blunt apex in lateral view; broad ellipsoidal with apex slightly incised in dorsal view. Preanal appendages triangular with corners rounded and dorsal margin slightly concave in lateral view, ventroposterior margin curved dorsad, apex broad and blunt. Inferior appendages capitate with curved truncate apex in lateral vicw.

Distribution. - Malaysia: Sabah.
Remarks. - This species was previously only known from the unique type from Sabah. Two additional male specimens were found in the material borrowed from BPBM and BMNH. The male is distinguished by having modified spur long and truncate, about $3 x$ longer than adjacent spur, with apex more than $2 x$ wider than base, and with minute apical points curving mesad.

Dipseudopsis flavisignata (McLachlan, 1866) (fig. 27)

[^7](1933), one slide with wings mounted dry, and the second slide with body mounted in balsam, but prothorax absent.
Dipseudopsis flavisignata (McLachlan, 1866). - Mosely 1933: 499-502, figs. 1-7. Fischer 1962: 9. Fischer 1972: 4.

Male. - Head dorsum dark brown, yellowish brown between antenna, frontoclypeus dark brown. Antenna base of scape yellowish brown and remainder dark brown. Maxillary palps 2.3 mm dark brown, labial palps 0.8 mm yellowish. Proboscis 1.7 mm about as long as first four segments of maxillary palp. Mesothorax and abdomen black. Forewing 12 mm , similar to that of D. spectabilis (fig. 9), dark brown background with large yellowish spots: 1) irregular band on anterior margin tapering toward base, 2) subapical C-shaped spot having distal side open and filled with dark narrow stripe along $\mathrm{M}_{1}$ (but in description by Mosely (1933: fig. 1) the latter is depicted as an irregular O-shaped spot), 3) semicircular irregular band through anal region, running from base, distad through anal loops and thyridial cell to arculus. Front leg with coxa, trochanter and basal portion of femur dark, apical portion yellowish; Middle and hind leg with similar colour pattern but lighter. Modified spur (fig. 27) nearly as long as first tarsal segment, basal $1 / 1$ unmodified, distal slender twisted blade, bearing a small tooth at base, a ridge at midlength and apical $1 / 2$ flattened like a blade. Genitalia (fig. 27): Tergum IX posterior margin broad at base in dorsal view, but abruptly tapering into slender fingerlike extension, apical $1 / 2$ overhanging segment X in lateral view. Segment X long ovoid projection tapering slightly posteriad in dorsal view. Preanal appendages with dorsal margin nearly horizontal, meeting curved ventroposterior margin at small dorsoapical lobe in lateral view. Inferior appendages similar as in D. spectabilis, heavily sclerotized, and darker brown than other appendages, basal $1 / 3$ angled dorsally, and apical $2 / 3$ fingerlike and angled more posteriorly in lateral view.

## Distribution. - Indonesia: Sulawesi.

Remarks. - This species is only known from the unique type. It is a sibling species to $D$. spectabilis, but differs by having the forewing with a yellowish Cshaped subapical spot, almost closed except for a narrow dark strip along $M_{1}$, and by having the modified spur with one short subapical point distant from apical point.

## Dipseudopsis flinti sp. n.

(figs. 11, 28)
Type material. - Holotype ő: MALAYSIA: West Malaysia:


Figures 26-29. - Dipseudopsis, A-G, male genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior appendages, ventral; D, phallus, lateral; E, phallus, ventral; F, segments IX, X, dorsal; G, ventral. I-J, male modified apical spurs of hind tibia, ventral. Fig. 26, D. elongata Banks. Fig. 27, D. flavisignata (McLachlan). Fig. 28, D. flinti, n. sp. Fig. 29, D. immaculata Ulmer.

Selangor, Kuala lumpur, 21.IV',1931, H. M. Pendlebury, (-


Make. - Head completely glossy dark brown, almose black, antennae and mouthparts dark brown. Maxillary palps 2.3 mm , proboscis 1.3 mm , as long as firse chree segments of maxillary palp. Pronotum dull dark brown. except anterior margin light, mesonorum and tegula dark brown and metanotum light brown. Forewing 12 mm , dark brown except for slender lighe brown stripe along posterior margin, running from cell 3A and expanded at arculus; cells below M: and Cu : light brown distally. Front leg with coxa. trochanter and basal z/ of femur dark brown, distal portions of leg light brown; middle leg with coxa dark brown and distal segments light brown; hind leg light brown. Modified spur (fig. 28) about $2 \times$ longer than adjacent spur, having less than apical $1 / 1$ bifid with two short apical points parallel and angled distad toward tarsus. Genitalia (fig. 28): Tergum IX posterior margin broadly bilobed in dorsal view; IX sternum base rectangular in lateral view, mesosuperior process with blunt dorsal lobe and slender posterior projection with apex curved anteriad. Preanal appendages rrapezoidal with dorsal margin slightly inclined dorsad toward blunt distal apex in lateral view, ventral margin curved gradually into posterior margin. Segment X about $2 / 3$ as wide as tergum IX in dorsal view, somewhat rectangular with posterior margin broadly bilobed. Inferior appendages capitate with apical 1/ellipsoidal and about 3 X as thick as base in lateral view, lateral arms slender and fingerlike with truncate apex in ventral view.

Etymology. - Named after Oliver S. Flint, Jr.
Distribution. - Malaysia: West Malaysia.
Remarks. - This species is only known from the unique type. The modified spur of the male is similar to that of $D$. sfabatensis, but it differs by having two apical points parallel and angled toward the tarsus, the adjacent spur is relatively shorter, about $1 / 2$ as long as the modified spur; in $D$. stabatensis the apical poines of the modified spur are not parallel but are directed in opposite directions, and the adjacent spur is relatisely longer, \% as long as the modified spur. Also, 1. flinti differs by having the inferior appendage capcate in lateral view, with the apical $1 / 2$ ellipsoidal and it $3 x$ as broad as the base.

## Dipseudopsis immaculata Ulmer, 1905

[^8]9; Weidner 1964: 73; Fischer 1972: 4; Malicky \& Weaver 1988: 4, fig. la-e.
Dipseudopsis moesta Banks, 1931b: 400, 401, fig. 9, Type ठ*: MALAYSIA: West Malaysia: Johor, Lubok, Kedondong, NW Gunung Ledang (Mt Ophir), Lalang \& Belukar, H. C. Abraham (MCZC 16481). - Fischer 1962: 11. Syn. n.
nec Dipseudopsis moesta. - Ulmer 1951: 128 [reidentification of Ulmer 1905, fig. 75 b sub nom. D. stellata $][=D$. benardi].

Specimens examined. - INDONESIA: Sumatra: 2 ot, Karny, $^{\circ}$, Pakan Baroe, at light, $23 . X .1925$ (RMNH). 10 , Huta Padang, 20.I.1990, Diehl (Clhm). 1 ơ, Dolok Merangir, 15.VI.82-4.I.1983, Diehl (Clhm). MalaySla: Sarawak: 1 ठे, $^{\text {h }}$ Merirai Valley near Kapft, 180 m , 28.VII.-6.VIII.1958, T. C. Maa (BPBM). West Malaysia: $1 \delta^{*} 1$ 우, Pahang, Rompin Mining Co., railway track, 31 km , Petoh Swamp, 11.XII. 1960 (ВРВм).

Male. - Head dorsum, cervical sclerite and pronotum light orangish brown; frontoclypeus mostly dark brown, but lateral margins yellowish brown; genae yellowish brown except dark brown ventral spike. Antennae light brown. Mouthparts brown, maxillary palps 2.2 mm ; proboscis 1.2 mm , about as long as first three segments of maxillary palps. Mesonotum and tegula dark brown, about as dark as eyes. Legs light brown. Modified spur (fig. 29) slightly longer than adjacent spur, apical $1 / 2$ modified into twisted point with setose basal pocket; apex almost truncate, but lateral side curved and other side with short point directed toward tarsus. Tegula dark brown. Forewing 11-12 mm, almost uniformly brown, except for hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Genitalia (fig. 29): Tergum IX triangular in dorsal view, with about $3 / 4$ of posterior overhanging segment X in lateral view. Segment X thumblike in lateral view, extending posteriad as far as inferior appendage; broadly cordate in dorsal view with apex slightly incised. Preanal appendages rhomboidal or sometimes simply rounded in lateral view. Inferior appendages oblong and truncate in lateral view, ellipsoidal in ventral view, but apicomesal margin incised with blunt subapicomesal tooth.

Distribution. - Indonesia: Sumatra; Malaysia: Sarawak, West Malaysia.

Remarks. - The male of this species has a modified spur which is similar to that of $D$. doehleri and $D$. nebulosa, but can be distinguished from these by its almost uniformly dark brown forewing. It also differs by having the male genitalia with more than $2 / 3$ of tergum IX overhanging segment X , and the inferior appendage with a blunt subapicomesal tooth in ventral view. A lighter phase is noted in male specimens from Borneo, having apical region of the forewing brown, remainder nearly translucent except for several dark


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Figures 30-33. - Dipseudopsis, A-G, male genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior appendages, ventral; D, phallus, lateral; E, phallus, ventral; F, segments IX, X, dorsal; G, ventral. I-J, male modified spurs apital spurs of hind tibia, ventral. Fig. 30, D. indica McLachlan. Fig. 31, D. infuscata McLachlan. Fig. 32, D. knapp Schmid and Denning. Fig. 33, D. lamellata Martynov.
veins. A slight variation also is observed in males from Sumatra, having forewing with narrow lighter strip along anterior margin and indistinct translucent area in outer s of wing. and preanal appendage more noticably incised distally. Regarding specimen determined as D. serellara by Ulmer (1905: fig. 75b) and later determined as D. moesta by Ulmer (1951), cf. 'Remarks' of D. benardi above.

## Dipseudopsis indica McLachlan, 1875

(fig. 30 )
Drpscudopsis indica MicLachlan, 1875: 18-20, pl. 2: figs. 1314 (indicous). Type 'Habitus in India orientali.' Lectorype vinta, designated by Kimmins (1957: 101). - Fischer 19)(2: 10; Fischer 1972: 4: Higler 1992: 80.
(7ipscudopsis buddhaz Banks. 1913: 239, pl. 23: fig. 8, pl. 24: fig. 19. Type $\delta^{\circ}$ : indla: West Bengal, Chapra, Mackenzic, no dare (mczc 11757). - Fischer 1962: 5; Higler 1992: 80. Syn. n.
Drpsendopsis indica sindica Marrynov, 1935: 159, 206, fig. ${ }^{59}$ a-b. Type: Pakistan: " ${ }^{\circ}$. Bubak, Sind, from pools. 12.X1.1927. B. Prashad and B. N. Chopra.' - Fischer 1962: 10; Fischer 1972: 4. Syn. n.

Specimens examined. - BANGLADESH: $180^{\circ} 36$ 名, Chicknagul, 10 mi N Sylhet, $10 \mathrm{~m}, 30 . \mathrm{IX} .61$, Ross \& Cavagnaro (casc). $50^{\circ} 69$, Cormilla, 50ft, 23.IX.61, Ross \&. Cavagnaro (CASC). INDIA: Bihar: $1 \delta^{\circ}$, Pusa, at light, 22.1711.08 (MczC). $18^{\circ}$, Pusa, IV.1919, M. A. Husain (BMNH). 12 . Pusa, 2. 1TII. 1924 , Mukerjee (BMNH). Orissa: 20, det. Martynov. Balighai, near Puri, 16-20.VIII. 11 (ZRas). $1 \delta^{\hat{\prime}}$, Cuttack. 6.III.1944, at light (BMNH). 20., Bhubaneswar, 11.1983. J. Olảh (clhm). West Bengal: 10, Chapra, Mackenzic (mcz.C). $6 \delta^{\circ} 32$ ? , Malda $110 \mathrm{~m}, 28 . X .61$, Ross \& Cavagnaro (CASC). $2 \delta^{\circ} 2 \%, 10 \mathrm{mi}$ SE Asansol 175 m , 6.XI.61. Ross \& Cavagnaro (CASC). Pakistan: Sind: $10^{\circ}$, Makli, near Thatta, 22.IX. 1976, Hevel \& Dietz (USNM). 2 oै 1 U. Makli, near Hyderabad, 24.IX.1976, Hevel \& Dietz (-

Malc. - Colour dull brown, with eyes dark brown. Pronotum lighter brown than head and mesonotum. Maxillary: palps 2.0 mm ; proboscis 0.8 mm , as long as firse 3* segments of maxillary palp. Legs light brown. Modified spur (fig. 30) with apical $1 / 2$ modified, subdisided into two acuminate branches, first branch bifid with apical points unequal and both directed distally, econd branch curved, directed laterally with minute point at its base. Forewing $11-14 \mathrm{~mm}$, mostly uni-
dull hrown, but costal margin lighter brown, with are of faint translucent spots distad of cord, cenpot at m -cu. and two translucent spots at arcu(ienitalia (fig. 30): Tergum IX only slightly rhanging segment $X$ in lateral view, posterior mardorsal view, but middle slightly . .. rnum IX short, with mesosuperior in lateral view, extending above regment $X$ dorsal margin inclined rad with blunt apex lobiform in lateral view, el-
lipsoidal and long in dorsal view. Preanal appendages with dorsal margin straight and horizontal in lateral view, ventroposterior margin curved apicodorsal angle squarish. Inferior appendages capitate in lateral view, with short ventrobasal shelf, narrow middle section and broad squarish apex; lateral arms with apicomesal tooth in ventral view.

Distribution. - Bangladesh; India: Bihar, Orissa, West Bengal; Pakistan: Sind.

Remarks. - This species is easily recognized by the complex male modified spur, having two branches, one branch directed distally and bifid, and the other branch curved laterally and acuminate, but with a minute point at its base. Examination of the type of $D$. buddha revealed that it is identical to $D$. indica. Based on its original description, the subspecies D. i. sindica is recognized as a junior synonym of $D$. indica.

## Dipseudopsis infuscata McLachlan, 1875

(fig. 31)
Dipseudopsis infuscata McLachlan, 1875: 15-16, pl. 2: figs. 6-10, Holotype on: $^{*}$ indonesia: Sulawesi, Ujung Pandang 'Piepers Celebes Macassar' (RMNH). - Fischer 1962: 10; Fischer 1972: 4; Malicky \& Weaver 1988: 3, fig. $4 \mathrm{a}-\mathrm{e}$.
nec Dipseudopsis infuscata. - Ulmer 1905: 96, fig. 75d;
Ulmer 1951: 128; Malicky 1984: 214, 216
Specimens examined. - Indonesia: Java, Jakarta 'Bata-
 10, I.1908, Jacobson (ZMAN). Kalimantan: $1 \delta^{\text {t }} 3$ ? , Telang X. 1881 (MCZC). 10 ${ }^{\text {® }}$, West Java, [Piepers?], McL. coll (BMNH). Sumatra: 20 , Palembang, 13.V.1972, Diehl (CLHM).

Male. - Head and antennae brown. Maxillary palps $1.6-1.8 \mathrm{~mm}$, proboscis $0.5-1.0 \mathrm{~mm}$ as long as first two or three segments of maxillary palp. Antenna yellowish brown with fine brown setae that become less dense apically so that distal part appears more yellowish. Forewing 12 mm , brown with small translucent spot distal to r , hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Modified spur (fig. 31) slightly longer than adjacent spur, bifid with two acuminate apical points, longer point straight directed distally and almost $1 / 2$ as long as long as spur, shorter point curved with apex directed mesad. Genitalia (fig. 31): Tergum IX with posterior margin rounded in dorsal view, with slight protrusion in middle, short and not extended over segment $X$ in lateral view. Segment $X$ with basal $2 / 3$ of dorsal margin inclined ventrad apical portion directed posteriad, having apex truncate, in lateral view; dorsal and ventral margins narrowing toward apex, with apex truncate and slightly emarginate in dorsal
view. Preanal appendages with posterior margin incised in lateral view. Inferior appendages with dorsal and ventral margins nearly parallel and curved dorsad in lateral view, having thumblike apex; base with mesal margin squarish in ventral view, distal portion long with mesal margin slightly convex, apicolateral margin rounded, and with apicomesal point.

Distribution. - Indonesia: Java, Kalimantan, Sulawesi, Sumatra.

Remarks. - This species is similar to D. robustior, but differs by having the two apical points of the modified spur with the longer point straight and slightly shorter than the unmodified base of the spur, and the shorter point curved with its apex directed mesad. All specimens determined as $D$. infuscata by Malicky (1984) from South Andaman Island are now considered to be $D$. robustior andamanensis Weaver \& Malicky, n. ssp. The specimen referred to by Ulmer (1905: fig, 75d, 1951) is D. contorta, not D. infuscata (cf. D. contorta above).

## Dipseudopsis knappi Schmid \& Denning, 1979 (fig. 32)

Dipseudopsis knappi Schmid \& Denning, 1979: 246, 248, fig. 5a-e, Holotype ot: Thailand: Tha Phra, Khom Kaen, 17.V.1968, blacklight trap, F. W. Knapp (CASC).

Male. - Body brown. Forewing 13 mm , brown with scattered translucent spots. Modified spur (fig. 32) trifid with long curved acuminate point and slightly shorter fingerlike lobe bearing minute subapical point. Genitalia (fig. 32): Tergum IX broadly rounded in dorsal view, short in lateral view; sternum IX with mesosuperior process triangular in lateral view. Segment X slender with apex distinctly bilobed in dorsal view, lateral margins sinuate, base of dorsal margin slightly inclined dorsally distad in lateral view, apical $1 / 2$ horizontal with ventral margin concave, apex slender and fingerlike. Preanal appendages trapezoidal with corners rounded and dorsal margin inclined dorsad in lateral view, having blunt apicodorsal lobe. Inferior appendages slightly clavate with basodorsal shelf in lateral view, middle portion constricted, and having clavate apex.

## Distribution. - Thailand.

Remarks. - This species is only known from the type series from northeastern Thailand. The modified spur of $D$. knappi is trifid and somewhat similar to that of $D$. spectabilis, but differs by having second longest process of the spur lobiform, rather than pointed. The male genitalia differ by having segment

X with a bilobed apex. The diagnosis of the male given above is based on the original species description (Schmid \& Denning 1979).

## Dipseudopsis lamellata Martynov, 1935

(fig. 33)
Dipseudopsis lamellata Martynov, 1935: 160-161, figs. 60ab, 61, Type series: burma: Myitkyina Dist.: $1 \delta^{\circ} 12$ ㅇ, Lonton, W shores Indawgyi Lake, U. Burma, 1831.X.1926, B. N. Chopra; Syntype 1 O*: Kamaing, U. Burma, X.1926, B. N. Chopra (NzSI); indiA: Assam: Mangaldai, $26.26 \mathrm{~N} 92.02 \mathrm{E}, 16-18 . \mathrm{X} .1910$, S. W. Kemp. Fischer 1962: 10; Higler 1992: 80.

Specimens examined. - BANGLADESH: $1 \delta^{7}$, Chicknagul, 10 mi N Sylhet, $10 \mathrm{~m}, 30 . \mathrm{IX} .61$, Ross \& Cavagnaro (CASC). $20^{\star} 4$, Harbang For. 6mi N Chiringa, 50ft, 18.IX.61, Ross \& Cavagnaro (CASC). 10 , Umgebung Dacca, 115.V.1976, Dietz (Clhm). indiA: Assam: 2 ở 3 우, Kohara, $_{\text {, }}$ Kaziranga, $110 \mathrm{~m}, 26.36 \mathrm{~N}$ 93.28E, 7-16.X.61, Ross \& Cavagnaro (CASC). Orissa: 20, Bhubaneswar, II.1983, J. Oláh (CLHM).

Male. - Colour brown. Forewing 13-15 mm, colouration similar as in D. indica, mostly brown, but with arc of five faint translucent spots distad of cord, central hyaline spot at $\mathrm{m}-\mathrm{cu}$, and broad translucent spot at arculus. Modified spur (fig. 33) $2 x$ longer than adjacent spur, bearing two distal processes, longer process slender twisted and fingerlike, shorter process slender acuminate and bent laterad at right angle. Genitalia (fig. 33): Tergum IX broad triangular in dorsal view, apex slightly overhanging segment X in lateral view; sternum IX short, with mesosuperior process having blunt dorsal lobe and slender ventroposterior process in lateral view. Segment X dorsal margin concave, ventral margin convex, apical part narrow and curved dorsad in lateral view; lateral margins slightly convergent at basal $1 / 3$, distal portion elongate ellipsoidal, with apex slightly incised in dorsal view. Preanal appendages slender and somewhat trapezoidal in lateral view, with posterior margin strongly incised, bearing two blunt apical points. Inferior appendages capitate in lateral view, with base constricted, having short slender basal projection and widening into rhomboid apex; basomesal margin with deep curved incision in ventral view, distal portion widened, having apex with lateral margin curved and with short apicomesal point.

Distribution. - Bangladesh; Burma; India: Assam, Orissa.

Remarks. - This species differs from other species of Dipseudopsis by having a unique modified spur, bifid, with a long slender fingerlike twisted process, and with an acuminate process angled laterad.

c genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior E. phallus, ventral; F, segments IX, X, dorsal; G, ventral. I-J, male modified spurs apV. Iucasi, n. sp. Fig. 35, D. maculata Ulmer. Fig. 36, D. malaisei, n. sp. Fig. 37, D.

## Dipseudopsis lucasi sp. n.

(fig. 34)
Type material. - Holotype ô: INDIA: Karnataka: Shimoga Dist., Agumbe Ghat, 2000ft, 1974, Lucas (CLHM). Paratypes: $7 \mathrm{o}^{\text {on }} 3$ 오, Someshwar, 27.I.1959, F. Schmid (CNCI). 1 ${ }^{\circ}$, Nagodi, 28.I.1959, F. Schmid (CNCI). $1 \delta^{\hat{}}$, Kerala: Sangalipalam, 20-21.XII.1961, F. Schmid (CNCI).

Male. - Head brown, eyes darker brown, frontoclypeus brown, antennae brown but ventral sides lighter brown, palps brown, maxillary palps 2.8 mm , labial palps 1.0 mm , proboscis 1.2 mm , almost as long as first three segments of maxillary palp. Pronotum dark brown, but lateral sides lighter, mesonotum and tegula dark brown. Forewing $14-17 \mathrm{~mm}$ unicoloured dark brown except small hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Legs with coxae dark brown, distal segments light brown. Modified spur (fig. 34) slightly longer than adjacent spur, apical $1 / 2$ modified with short curved subapical shelf, and long slender acuminate curved point directed distad. Genitalia (fig. 34): Tergum IX with posterior $1 / 4 /$ overhanging X in lateral view, posterior margin broadly rounded in dorsal view; sternum IX mesosuperior process with middle constricted and apex curved abruptly dorsad in lateral view. Segment X thumblike and directly posteriad in lateral view; elongate ovoid, with apex slightly incised in dorsal view. Preanal appendages short, extending to middle of segment X in lateral view, posterior margin broadly rounded, almost semicircular but with blunt apicodorsal angle. Inferior appendages basal $1 / 2$ thick and broadened toward middle in lateral view, apical $1 / 2$ tapered to truncate apex; lateral margin rounded and mesal margin almost straight in ventral view, but with obtuse apicomesal angle and minute basomesal angle.

Etymology. - Named after the collector J.A.W. Lucas.

Distribution. - India: Karnataka, Kerala.
Remarks. - The male genitalia of $D$. lucasi are similar to those of $D$. doehleri and $D$. nebulosa, having abbreviated preanal appendages. However, D. lucasi differs by having forewing almost unicoloured dark brown, and lacking the contrasting pattern, as in the forewings of $D$. doebleri and $D$. nebulosa (figs. 5, 7).

## Dipseudopsis maculata Ulmer, 1907

(fig. 35)
Dipseudopsis maculata Ulmer, 1907a: 37-39, fig. 55a-b, Type of*: Malaysia: Sabah: Sandakan (BMnH). Fischer 1962: 11. Fischer 1972: 5.

Specimens examined. - malaysia: Sabah: $1 \delta^{7}$, Tawau, Quoin Hill, Cocoa Res. Sta., 1.X.1962, Y. Hirashima (вРвм). 10, Tawau, Quoin Hill, 15-20.VII.1962, Y. Hirashima (BPBM). $10^{\hat{7}}$, Sepilok For. Res., Sandakan Bay, 110m, 27.X. 1957, J. L. Gressitt (BPBM).

Male. - Head dorsum with anterior $1 / 2$ glossy dark brown, almost black, and posterior warts light brown, frontoclypeus glossy dark brown; antennal scape and pedicel light brown, flagellum darker, palps dark brown, maxillary palps 3.6 mm , proboscis 2.2 mm , as long as first four segments of maxillary palp. Pronotum, mesonotum and tegula dark brown, but metathorax and abdomen yellowish brown. Forewing 17-18 mm, veins dark brown against brown and translucent background, large translucent triangular spot between $\mathrm{R}_{1}$ and $\mathrm{Cu}_{1}$, also translucent spots at arculus, discal cell, and area distal to cord, and dark band distal to hyaline spot at m -cu. Fore leg coxa to basal $1 / 2$ of femur dark brown, distal $1 / 2$ of femur and other distal leg segments yellowish brown; mid leg coxa dark brown, distal leg segments yellowish brown; hind leg yellowish brown. Modified spur (fig. 35) $2 x$ longer than adjacent spur, apical $1 / 3$ modified, into two unequal short nearly parallel points curved laterally away from tarsus. Genitalia (fig. 35): Tergum IX distal $1 / 3$ overhanging segment X and with broad apex in lateral view, posterior margin broadly rounded and bilobed in dorsal view. Sternum IX mesosuperior process triangular with blunt dorsal angle and acute posterior angle in lateral view. Segment X ventral margin straight and horizontal, dorsal margin slightly concave and inclined ventrad, apex rounded; ellipsoidal about $1 / 3$ as wide as tergum IX in dorsal view. Preanal appendages broad and triangular with corners rounded in lateral view, dorsal and ventroposterior margins slightly irregular, apicodorsal apex blunt. Inferior appendages inclined dorsad and widening apically, having apex truncate with rounded corners in lateral view, dorsal margin straight and longer than ventral margin; ventral margin with basal $1 / 3$ having protruding shelf extended as short free lobe, oblique angle at midlength, and distal $1 / 2$ inclined dorsad; distal process thumblike in ventral view, mesal margin with deep narrow curved basolateral notch, and apex rounded.

## Distribution. - Malaysia: Sabah.

Remarks. - The male of this species has a unique modified spur with a prominent apical point extending laterally from a blunt apical shelf, and a shorter subapical point curved laterally, having both points curved distad. $D$. maculata and $D$. nieuwenhuisi appear to be closely related based on similar colouration in the male forewing (cf. fig. 6 of D. nieuwenhuist), and similar characteristics in the male genitalia. $D$.
maculura differs by being smaller in size，and by hav－ ing male modified spur with only two apical points， rather than having three as in D．nieuwenhuisi．

## Dipseudopsis malaisei sp． n ． （fig．36）

［ype material．－Holorype đ＇：BURMA：Washaung，600ft， 13．＇II．1934，Malaise（NHRS）．Pararypes： $30^{\circ}$ ，same data， Malaise（NHRS）．

Male．－Head dorsum and frontoclypeus dark brown，nearly black，posterior warts brown． Antennae，mouthparts，legs and sclerotized parts of thorax unicoloured brown．Maxillary palps 2．2－2．5 mm ；proboscis $1.2-1.8 \mathrm{~mm}$ ，as long as first four seg－ ments of maxillary palps．Forewing length 13－15 mm ，dark brown except for small hyaline comma at m －cu．Modified spur（fig．36）slender，slightly curved and longer than adjacent spur，with two small spiral－ ling apical points．Abdomen yellowish brown． Genitalia（fig．36）：Tergum IX broadly rounded in dorsal view，with distal $2 / 3$ overhanging segment X in lateral view；sternum IX with mesosuperior process fingerlike and $1 / 2$ as long as segment X in lateral view． Segment X long and fingerlike，tapering slightly dis－ tad and apex incised in dorsal view．Preanal appenda－ ges oblong，with posterodorsal margin ellipsoidal in lateral view．Inferior appendages curved dorsad and broad in lateral view；broad with lateral and mesal margins parallel，and apices slightly bent mesad in ventral view．

Etymology．－Named after the collector R．Malaise．
Distribution．－Burma．
Remarks．－This species is only known from the type series．The male has segment X long and slender and modified spur with two small apical spiralling points similar as in D．benardi．However，D．malaisei differs by having male with preanal appendages ob－ long and ellipsoidal in lateral view，inferior appenda－ ges broad and flat，and dark brown forewings without large translucent spots，similar to D．benardi（fig．10）．

[^9]M\｛ale．－Head dark brown，but posterior warts light
brown；frontoclypeus with ventral $1 / 2$ light brown，dor－ sal $1 / 2$ dark brown；antennae light brown，palps light brown，maxillary palps 2.3 mm ，proboscis 1.2 mm ，as long as first three segments of maxillary palp． Pronotum light brown．Mesonotum and tegula dark brown．Forewing 13 mm ，dark brown with C and small light brown stigma and small hyaline spors at m －cu and arculus．Legs with coxae dark brown，distal segments light brown．Modified spur（fig．37）short and stout，about 4 X ，as long as basal width apical 2／3 modified with three broad points clustered together． Genitalia（fig．37）：Tergum IX posterior $1 / 3$ extended over segment X in lateral view，broad and triangular in dorsal view；sternum IX mesosuperior process straight rodlike with apex curved dorsad in lateral view．Segment X dorsal margin slightly concave and inclined ventrad in lateral view，ventral margin curved ventrad at middle and curved dorsad distad， apex blunt；ellipsoidal and about $1 / 2$ as wide as tergum IX in dorsal view．Preanal appendages slightly shorter than segment X in lateral view，dorsal margin slightly sinuate at base，with broad dorsoapical curve，ventral margin nearly straight inclined dorsad，posterior mar－ gin with shallow incision．Inferior appendages finger－ like and inclined dorsad in lateral view，with narrow basoventral shelf，squarish with distal arms parallel and straight in ventral view，lateral apex rounded，ba－ somesal margin with square－angled basal shelf，and lateral notch and apicomesal angle both square．

Etymology．－Named after Andrei Vasilievich Martynov．

Distribution．－Philippines：Sulu，Tawitawi．
Remarks．－This species is superficially similar to D．recta and D．triclavata，having modified spur tri－ fid．However，it differs by having the spur short and stout，about $4 \times$ longer than basal width，with apical $2 / 3$ of spur modified，and inferior appendage fingerlike in lateral view．

## Dipseudopsis modesta Banks， 1911

（fig．38）
Dipseudopsis modesta Banks，1911：105－106，pl．6：fig．2， Type ठ＊：IndiA：Bihar：Pusa，21．IX．08，T．N．J．（MCZC 11756）．－Fischer 1962：11；Fischer 1972：5；Higler 1992： 80.
Dipseudopsis pallida Martynov，1935：166－167，fig．70a－b， Type series： 1 ot $^{1}$ 워 in copula，Pusa，31．III．1911－4．IV，F． H．Gravely（probably NZSI）．－Fischer 1962：14；Higler 1992：80．Syn．n．

Specimens examined．－IndiA：Assam： $3 \delta^{\circ} 3$ ，Kohara， Kaziranga， $110 \mathrm{~m}, 26.36 \mathrm{~N}$ 93．28E，16．X．61，Ross \＆ Cavagnaro（CASC）． $1 \delta^{\star} 3$ 우， 13 mi SW Kochugaon， 50 m ，

17．X．61，Ross \＆Cavagnaro（CASC）． 2 o $^{\text {o }} 2$ 오，Meleng For．， Mariani， $110 \mathrm{~m}, 26.39 \mathrm{~N} 94.18 \mathrm{E}, 10 . \mathrm{X} .61$ ，Ross \＆ Cavagnaro（CASC）． $2 \delta^{\circ} 3$ ㅇ，Garampani Res． 10 mi S Gologhat， $110 \mathrm{~m}, 9 . \mathrm{X} .61$ ，Ross \＆Cavagnaro（CASC）．Assam： 1 or $^{\text {2 }}$ 우，＇Doom Dooma＇Dum Duma，27．34N 95．34E， VI．1943，E．Hardy（USNM）．＇U．D．M．N．C．H．＇［United District of Mikir and North Cachae Hills］：1才，Kalanga，1．V．1960， F．Schmid（CNCI）．Bihar： $40^{\top} 29$ ，Pusa（MCZC）．2 $\delta^{\circ}$ ，Pusa （BMNH）．Orissa：2ơ，Bhubaneswar，21．II．1985，J．Oláh （CLHM）．West Bengal： $3 \delta^{\circ}$ ，Chapra，Mackenzie（MCZC）．

Male．－Head glossy dark brown；frontoclypeus，an－ tennae and palps dark brown；maxillary palps 2.6 mm ， proboscis 1.0 mm ，almost as long as first three seg－ ments of maxillary palp．Pronotum，mesonotum and tegula dark brown．Forewing 12－15 mm，dark brown， except for hyaline lunula at m － cu and small translu－ cent spot at arculus．Legs light brown slightly darker basad．Modified spur（fig．38）slightly shorter than ad－ jacent spur with two minute apical points（cf． ＇Remarks＇below）．Genitalia（fig．38）：Tergum IX tri－ angular with acute apex slightly inclined dorsad and distal $1 / 3$ overhanging segment $X$ in lateral view；poste－ rior margin broadly rounded in dorsal view；sternum IX mesosuperior process with short broad dorsal trian－ gle，posterior process narrow having dorsal margin ir－ regular and slightly concave，and ventral margin curved dorsad in lateral view，extending above basal $1 / 20$ phallus．Segment X long slender and apex bilobed with deep mesal notch in dorsal view；fingerlike and slightly tapered in middle，extending posteriad distal to inferior appendages in lateral view．Preanal appen－ dages trapezoidal，posterior margin slightly incised in lateral view．Inferior appendages slender and long in lateral view，extending dorsad toward apex of preanal appendage，dorsal margin curved and concave，ventral margin irregular；lateral margins curved and apicome－ sal margin straight with apical point in ventral view， mesal margin with curved notch in basolateral corner．

> Distribution. - India: Assam, Bihar, Orissa, West Bengal.

Remarks．－This species can be distinguished by its unique male modified spur，having a short inconspic－ uous apical point directed distally and a long slender point curving halfway around the apex；the straight point is minute and lightly sclerotized，and without clear magnification it appears to be a small lobe．This species is similar to $D$ ．benardi，but it differs by hav－ ing male modified spur with smaller apical points and inferior appendages narrower in lateral view．We have observed some variation in the shape of the preanal appendages．Comparison of the type of $D$ ．modesta with the original description of D．pallida，reveals that these species are conspecific，and thus the latter is rec－ ognized as a junior synonym．

## Dipseudopsis morosa Banks， 1924

（fig．39）
Dipseudopsis morosa Banks，1924：450，pl．3：fig．31．－ Fischer 1962：11；Fischer 1972： 5.
nec Dipseudopsis morosa．－Schmid 1958：13，24，27，89，pl． 15：figs．12－13．
Lectotype designation．－お＊：PHILIPPINES： Mindanao：Dapitan，Baker（MCZC 14826）．1才＊： Basilan：probably syntype but without type label， Baker（USNM）．The type series： $3 \delta^{\sigma}$（MCZC）is mixed and comprises three different species；in addition to the lectotype，the o＊syntype from Luzon is D．colla－ ris，and the $\delta^{*}$ syntype from＇Borneo＇Kalimantan is D．stabatensis．

Specimens examined．－philippines：Mindanao： $10^{\top}$ ， Lanao， 4.8 km E of Dansalan， $750 \mathrm{~m}, 11 . \mathrm{VI} .1958$ ，jungle along stream，H．E．Milliron（вРВм）．
Male．－Head brown with black triangular median stripe，broad along anterior margin and tapering to middle of posterior margin，frontoclypeus glossy black and genae brown，antennae and palps brown， maxillary palps 2.3 mm ，proboscis 1.0 mm ，almost as long as first three segments of maxillary palp． Pronotum black with posterior dorsal margin brown， mesonotum and tegula almost black．Forewing 9－10 mm ，uniformly dark brown，but small hyaline spots at m －cu and arculus．Legs with coxae of fore and mid legs dark brown and distal segments yellowish，hind leg yellowish．Modified spur（fig．39） $11 / 2 x$ longer than adjacent spur，apex truncate with two apical points， both points curved and one point nearly in plane per－ pendicular to longitudinal axis of spur，and other longer point with apex directed more distad． Genitalia（fig．39）：Tergum IX not extended over base of segment X in lateral view，slightly inclined dorsad； sternum IX mesosuperior process short and blunt in lateral view，with square dorsoanterior angle，posteri－ or lobe inclined dorsad and extended above basal $1 / 3$ of phallus．Segment X dorsal and ventral margins nearly parallel and inclined ventrad in lateral view，but api－ cal portion of ventral margin horizontal，apex round－ ed；triangular and with rounded posterior apex in dorsal view．Preanal appendages broad and somewhat D－shaped with apex extended dorsoposteriad in late－ ral view．Inferior appendages oblong with distinct subapical ventral tooth in lateral view，distal portion capitate in ventral view with apex and basomesal notch curved．

Distribution．－Philippines：Basilan，Mindanao．
Remarks．－$D$ ．morosa has been confused with $D$ ． triclavata，due to similarities of the male modified spurs．However，these species are not closely related and are distinguished from each other by characteris－

tics of the male genitalia，$D$ ．morosa having the inferi－ or appendage broader and with a distinct blunt ven－ tral tooth in lateral view，and by having modified spur with only two apical points．The type series of $D$ ．mo－ rosa at the MCZC is mixed，comprising three syntypes， each a different species．Records of $D$ ．morosa from Sri Lanka by Schmid（1958）are actually D．triclava－ ta．

## Dipseudopsis nebulosa Albarda， 1881

（figs．7，12A，40）
Dipseudopsis nebulosa Albarda，1881：19，pl．5：fig．4，Type O，Indonesia：＇in insula Sumatra＇．－Fischer 1962：11－
12；Fischer 1972：6；Malicky \＆Weaver 1988：4，fig．2a－ f．
nec Dipseudopsis nebulosa．－Banks 1914： 178 （Luzon）．
Specimens examined．－burma：2 $\widehat{\delta}$ ，Tenasserim，Sukli， 75 km E of Moulmein， $600 \mathrm{~m}, 27-31 . X .1934$ ，Malaise （BMNH）．10＇，Mekane， 20 km W Myawaddy，Amherst Dist， 700ft，3．XI．1934，Malaise（NHRS）．INDONESIA：Sumatra：
 （RMNH）．1 $\widehat{\text { th}}$ ，Loeboek Sikaping， 450 m （RMNH）． $1 \delta^{\text {º }}$ ， Pematang Siantar，1．VI－12．XII．1985，Diehl（CLHM）．12 $\widehat{\text { h }}$ Huta Padang，Diehl（Clhm）．malaysia：West Malaysia： Pahang：30，Kuala Tahan，at light，Pendlebury（вмnh）． 10̊，Sungei Tembeling，18．XI．1922，Pendlebury（MCZC）． $1^{10}$ ，Kuala Teku，500ft，5．XII．1921，Pendlebury（MCZC）． Selangor： 12 § 6 영，Kuala Lumpur，15－31．XII．1958，Quate （BBBM）．thailand：38＇，upper Pran River，13－15．IV． 1926 （MCZC $1 \delta^{\circ}$ ，BMNH $2 \delta^{\circ}$ ）． $2 \delta^{\circ}$ ，det．Mosely as D．doehleri，same data，15．IV．1927，Landell（BMNH）．10，Trang，26．VI．1924， Evans（MCZC）． $2 \delta^{\pi} 1$ 오，Trang Prov．，Khaophappha Khaochang， $200-400 \mathrm{~m}, 12-13 . \mathrm{I}$ ．1964，Samuelson（BPBM）． $5{ }^{\text {on}}$, Krachong For．near Trang， $100 \mathrm{~m}, 2$. VII．62，Ross \＆ Cavagnaro（CASC）． 1 万人， 20 km E of Krabi，10－20．II．1962， Friedel（clhm）．10 trang，Banchang，16．V．1924，at light， Evans（BMNH）．

Male．－Body and wings（fig．7）．Head dorsum，an－ tennae，cervical sclerites and pronotum orangish brown，and contrasting with dark brown mesono－ tum，tegula and forewings（fig．12A）．Frontoclypeus and mouthparts brown．Fore and mid legs with coxae and trochanters brown，femora with basal portions dark brown and apical portions orangish brown，tib－ iae and tarsi orangish；hind leg dark brown．Forewing 12－14 mm，dark brown with translucent spots：long triangular spot below $\mathrm{R}_{1}$ above $\mathrm{Cu}_{1}$ from base of $R$ s to hyaline lunula at $\mathrm{m}-\mathrm{cu}$ ，translucent oblong spot distal to cord with apical margin deeply incised，and small translucent spot at arculus．Modified spur（fig．40） with twisted glabrous apical point，about $2 / 3$ as long as spur．Genitalia（fig．40）similar to D．schmidi． Tergum IX with posterior margin triangular with apex curved and lateral sides slightly depressed in dor－ sal view，apical $/ 2$ of dorsum overhanging segment X in lateral view．Sternum IX extended anteriad，with mesosuperior process globular in lateral view．

Segment X short stout and ellipsoidal in lateral view， dorsal and ventral margins nearly parallel and hori－ zontal，apex bluntly rounded，broad and cordate in dorsal view，length and width subequal，with apex slightly incised．Preanal appendages abbreviated in length，with posterior margin incised．Inferior appen－ dages broad，somewhat ellipsoidal in lateral view，but dorsal margin slightly concave．

Distribution．－Burma；Indonesia：Java，Sumatra； Malaysia：West Malaysia；Thailand．

Remarks．－This species is similar to $D$ ．doebleri， but differs by having a translucent spot of forewing with faded edges，rather than a well defined oval－ shaped spot at the arculus，and genitalia with sternum IX，segment X ，and preanal appendages slightly broader in lateral view．The modified spur of $D$ ．neb－ ulosa is similar to that of $D$ ．immaculata and $D$ ．doeh－ leri，with a twisted glabrous apical point，having a se－ tose pocket at its base，but it differs by having a longer glabrous point，about $2 / 3$ as long as spur．We have ac－ cepted the description of the male provided by Ulmer （1951），who associated it with the female type from middle Sumatra．We have observed that a male spec－ imen from Luzon was misidentified by Banks（1914） and actually is $D$ ．collaris．

## Dipseudopsis nervosa Brauer， 1868

（figs．3，4，41）
Dipseudopsis nervosa Brauer，1868：267－268，Type 오：PHILI－ pPines：（Semper）［cf．＇Remarks＇below］．－Fischer 1962： 12；Fischer 1972： 6.
Dipseudopsis luctuosa Banks，1913：178，pl．8：fig．3，Type か＊：Luzon：Los Baños，P．L．Baker（Mczc 11769）．－ Ulmer 1930：380，as syn．－Fischer 1962：12；Fischer 1972： 6.
Dipseudopsis venosa Navás，1931：86，Type 우＇Filipinas： Mindanao，Kolambogan，11．VI．1914．Col．m．＇［We were unable to locate the female type of $D$ ．venosa for examina－ tion，but are relatively certain that this species is synony－ mous with $D$ ．nervosa，because only two species of Dipseudopsis are known from Mindanao，D．modesta and $D$ ．nervosa，and the size of the female type of $D$ ．venosa only agrees with the female of $D$ ．nervosa．］－Fischer 1962：16．Syn．n．

Specimens examined．－philippines：Leyte：30̃，Palo（－
 MCZC）． $2 \delta^{\star} 5$ ㅇ，Los Baños（USNM）． $4 \delta^{\circ} 3$ 오，Mt Maquiling （CNCI，USNM）． $1 \delta^{\hat{}}$ ，det．Ulmer（1905），bez．Bilneao，prov． Laguna，1876，Laglaize（MNHN）．Mindanao： $10^{\top}$ ， Kolambugan（BMNH）． 1 of（＇Paratypus＇see discussion above）
 19，Agusan Esperanza（BPBM）． 30 ，Agusan Jabonga Mayogda（MCZC）．2す，Bukidnon，Dalongdong， 40 km NW Maramag，Talakag，Urwaldrand， $800 \mathrm{~m}, 7^{\circ} 53^{\circ} \mathrm{N} 124^{\circ} 40^{\circ} \mathrm{E}(-$ ZMHB）． $1 \delta^{\text {t }} 2$ ㅇ，Butuan（MCZC，USNM）． $1 \delta$ ，Cotabato， Kalaong（ВРBM）．2才，Sapamoro，Curnan dist．（ZMUC）．

Mindoro: $80^{*}$. Laguna de Nanjan Soldanski (Clhm). Negros: 2038 . L. Balinsasayao (BPBMM). $1 \delta^{*}$. Sibulan
 (Capiz. Lithazo (/SMC). Sibưyan: 10, MagdiwangTampayan, Ga-ong. Pawala River (clema).

Malc. - Body and wings (fig. 3). Head dorsum similar to $D$. morosa having dark brown median acute triangle, tapering posteriad, and lateral sides ycllowish brown; frontoclypeus with dark brown median stripe tapering dorsad and lateral sides yellowish brown; antennae and palps brown, maxillary palps 2.8 mm , proboscis 1.7 mm , as long as first $31 / 2 \mathrm{seg}$ ments of maxillary palp. Cervical sclerites mostly dark brown, but dorsal part yellowish brown. Pronotum yellowish brown, median furrow and lateral sides dark brown; propleuron yellowish brown. . Mesonotum, metanotum and tegula dark brown. Mesothorax. metarhorax and abdomen brown dorsad and yellowish brown ventrad. Forewing $14-17 \mathrm{~mm}$, dark brown with three translucent spots: 1) basal stripe between $\mathrm{R}_{1}$ and $\mathrm{Cu}_{t}$, 2) triangular spot distal to cord, and 3) round spot at arculus. Fore leg coxa bicoloured, basal $1 / 2$ dark brown and distal $1 / 2$ yellowish brown. Pleuron bicoloured, cervical sclerites and side of thorax with lower portion dark brown and upper part yellowish brown. Modified spur (fig. 41) trifid, with long apical point spiraling distad, minute apical point, and subapical tobe. Genitalia (fig. 41): Tergum IX inclined dorsad in lateral view, minute portion of apex overhanging segment $X$, posterior margin broadly curved in dorsal view; sternum IX mesosuperior process oblong in lateral view, dorsal margin curved. Segment X dorsal margin slightly concave, strongly inclined ventrad to blunt apex in lateral view; base broad and narrowing toward oval-shaped apex in dorsal view. Preanal appendages irregular triangle, with dorsal margin irregular and curved in lateral view, ventroposterior margin steeply inclined dorsad, dorsoposterior apex squarish. Inferior appendages fingerlike and curved dorsad in lateral view, lateral margin curved in ventral view, extended arms fingerlike with middle of mesal margin having small semicircular incision.

Fermale. - Body and wings (fig. 4).
Distribution. - Philippines: Leyte, Luzon, .Mindanan, Mindoro, Negros, Panay, Sibuyan.

Remarks. - This species is similar to D. digitata, but differs by having unique colouration of the male head and forewings, and by characteristics of the male hind spur and genitalia.

Brauer (1868) described this species after a female from the Philippines, and the species epithet, nervosa, was probably derived in references to the colouration
pattern of the female forewing, having dark veins against a translucent background. The type which should be at the NHMW is missing. Instead a male specimen with a hand-written type label reading 'Paratypus' is present, but this specimen was not mentioned by Brauer and thus cannot be regarded as a syntype. The red 'paratype' label was probably added to this specimen after the species was described. This specimen cannot be the type because the type description was clearly based on a female specimen; also the forewing length of the type should be 19 mm , but that of the male 'paratype' is only 17 mm . Therefore, we conclude that the type is missing. We accept the male association of this species by Ulmer (1929). Furthermore, we are reasonably sure that the identity of this species is correct because there are no other species as large as $D$. nervosa known from the Philippines., i.e. the other four Philippine species, $D$. collaris, D. digitata, D. martynovi, and D. morosa, are much smaller.

## Dipseudopsis nervosella Ulmer, 1905 stat. $\mathbf{n}$.

 (fig. 42)Dipseudopsis sp.- Ulmer, 1905: 96, fig. 75e.
Dipseudopsis morosa. - Ulmer 1951: 128 [misdet.].
Dipseudopsis morosa nervosella Ulmer, 1951: 141-143, pl. 7: figs. 140-142, Type $\delta^{* *}$ : 'Borneo' 1886, F. Baczes (NHMW). - Fischer 1972: 5.

Specimens examined. - MALAYSLA: Sarawak: 10 , at foot of Mt Dulit, junction of rivers Tinjar \& Lejok, 19.VIII. 1932 (BMNH). 1 ? same data, 30.VIII. 1932 (BMNH). $10^{\text {t. }}$ R. Kapah trib. of R. Tinjar, 25.X.1932, Oxford Univ. Exp. (BMNH).

Male. - Dorsum of head with curved M-shaped pattern marked by division between glossy dark brown, almost black, anterior triangle pointed posteriad and dark brown lateral margins, and yellowish brown area, filling in two triangles of ' M ' and with acute angle of each triangle pointed toward base of antenna. Frontoclypeus glossy, almost black, antennae yellowish brown, cervical sclerites dark brown. Maxillary palps 3.0 mm , dark brown; proboscis brown, 1.8 mm , almost a long as first four segments of maxillary palp; labial palps 1.2 mm . Pronotum dark brown with slender yellowish brown posterior edges along median fissure and lateral margins, propleuron light brown; mesonotum almost black, mesopleuron dark brown except epimeron light brown; metathorax and abdomen yellowish brown. Forewing $15-16 \mathrm{~mm}$, dark brown with golden hairs, and three large translucent spots, similar to $D$. nervosa and $D$. nebulosa (cf. figs. 3, 7): 1) long band between $\mathrm{R}_{1}$ and $\mathrm{Cu}_{1}$, 2) equilateral triangle at arculus, and 3) subapical area distal to cord with six translucent gaps between


Figures 42-45. - Dipseudopsis, A-G, male genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior appendages, ventral; D, phallus, lateral; E, phallus, ventral; F, segments IX, X, dorsal; G, ventral. I-J, male modified spurs apical spurs of hind tibia, ventral. Fig. 42, D. nervosella Ulmer. Fig. 43, D. nieuwenhuisi Ulmer. Fig. 44, D. notata (Fabricius). Fig. 45, D. recta Martynov.
dark apical veins, and in discal ceil. Fore leg and middle leg with coxace, trochanter and basal portion of femur dark brown, distal portion of leg yellowish; hind . cg vellowish. Modified spur (fig. 42) 1/x longer than didiacent spur, apex eruncare with two apical points, toeth poines curved in plane perpendicular to longitudinal axis of spur (cf. 'Remarks' below). Genitalia (fig. 42 ): Tergum IX with posterior $1 / 8$ overhanging segment X in lateral view, posterior margin bilobed with two blune lateral angles separated by a obtuse mesal notch in dorsal view; sternum IX dorsomesal process slonder horizontal, with pointed apex curved dorsad in lateral view: Segment X dorsal margin slightly concave in lateral view, distal portion narrow; ellipsoidal and with apex slightly incised in dorsal view. Preanal appendages large lobate and extended dorsoposteriad in lateral view. Inferior appendages thumblike and inclined dorsad in lateral view, ventrobasal shelf with short frec apical projection; distal arms ellipsoidal in veneral view, without sharp mesal tooth, mesal margin with decp narrow curved basolateral norch.

## Distribution. - Malaysia: Sarawak.

Remarks. - This species was originally described as a subspecies of $D$. morosa. Comparison of the male type of $D$. neriosella with that of $D$. morosa reveals that each is a distinct species. The modified spur in $D$. nervosella is similar to that of D. morosa but more robust and actually a mirror image of the other, having apical points curved in opposite directions. This species also differs by having the male forewing with a more contrasting colour pattern, and inferior appendage having mesal margin with deep curved basolateral notch in ventral view: Also, $D$. nervosella is only known from Sarawak, and $D$. morosa is only known from the islands of Mindanao and Basilan of the Philippines.

## Dipseudopsis nieuwenhuisi Ulmer, 1909

fies. 6. 431
()ipsendopses nieuwenhuisi Ulmer, 1909: 128-129, fig.4,
Type ? : Borneo Exp., Dr. Nicuwenhuis, L. Dengay,
1894 (RM:NH). - Fischer 1962: 12; Fischer 1972: 6.
xamined. - Mabaysia: Sarawak, Mt Dulit,
., at light, Oxford Univ. Exp., B. M
1-.IX. 1932

posterior lateral corners light brown, propleuron light brown, mesonotum and tegula dark brown, metathorax light brown. Forewing $19-23 \mathrm{~mm}$, dark brown, veins dark brown with several translucent stripes between veins, similar to that $D$. digitata, having three translucent areas, basal stripe in thyridial cell, round marginal spot at arculus and subapical translucent stripes distal to cord. Front and middle legs with coxae dark brown, trochanters to apex yellowish brown; hind leg mostly yellowish. Modified spur (fig. 43) 2 X as long as adjacent spur and 2 X as thick, with three apical points, one minute point, and two large points parallel and curved, one large point subapical. Genitalia (fig. 43): Tergum IX posterior margin broadly bilobed, having shallow mesal notch in dorsal view. Segment X originating from relatively shallow cavity of tergum IX in lateral view, apex blunt, ventral margin slightly concave; ellipsoidal and posterior margin broadly bilobed in dorsal view. Preanal appendages triangular and rounded with dorsal margin horizontal, broadly curved and slightly emarginate in lateral view, ventroposterior margin broadly curved and strongly inclined dorsad. Inferior appendages oblong with ventrobasal shelf in lateral view; angulate with distal arms thumblike in ventral view, mesal margin with deep narrow curved basolateral notch.

Distribution. - Malaysia: Sarawak.
Remarks. - This is the largest Asian species of Dipseudopsis known. It appears to be closely related to D. digitata and D. nervosella, but differs by its larger size and by characteristics of the male modified spur and genitalia.

## Dipseudopsis notata (Fabricius, 1781)

(fig. 44)
Phryganea notata Fabricius, 1781: 390, Type ठ*: 'N. America' [mislabelled, cf. 'Remarks' below], (BMNH). Fischer 1962: 13.
Dipseudopsis horni Ulmer, 1915: 46-47, fig. 13. Type series: SRI LANKA: '2ठ', [N. Cen. Prov.: Mahl Iluppallama] Maha Illupalama, Ceylon, Juli 1912, J. C. F. Fryer leg., im Museum Cambridge [MCZC]; ferner 9 Exemplare ( $(0, q)$, [Cen. Prov.]: Kandy, resp. [E. Prov.: Kantalai?] Kanthaley, resp. [Cen. Prov.: Matale] Matala, Ceylon, Dr. W. Horn leg., im Deutsch. Entom. Museum BerlinDahlem' [ZMUH]. - Weidner (1964: 73): 'Ceylon, Matala, Paratypoid' of D. horni, (ZMUH). - Fischer 1962: 9; Weidner 1964: 73; Fischer 1972: 4. Syn. n.

Specimens examined. - INDIA: Karnataka: $80^{\star}$, Bhadravati (MCZC). $2 \delta^{\circ}$, Shimoga, R. Tunga 1865 ft (MCZC). Tamil Nadu: $3 \delta^{\top}$, Tanjore dist. 'Thanjavur', Nedungadu (mczC). SRI Lanka: N. Prov.: $20^{\circ}$, Marichchukkaddi (BMNH). N. Cen. Prov.: 60, Moradan Maduwa, Wilpattu Natl. Park,

23 mi W of Anuradhapura， 80 m （MZLU）． 10 ，Kontalai， 60 m （MZLU）．Anuradhapura Dist： $7 \delta 8$ 只，Wildlife Soc． Bungalow，Hunuwilagama，Wilpattu Natl．Park，200ft（－ USNM）． $10{ }^{\text {® }} 10$ ㅇ，Irrigation Bungalow，［Padawiya Tank］ ＇Padaviya＇，180ft（USNM）． $80^{\star} 2$ ㅇ，Padaviya， 180 ft （USNM）． Polonnaruwa Dist： $2 \delta^{\text {t }} 7$ 아，Pinburettawa， 13 mi S Mannampitiya（USNM）．10＊，Polonnaruwa（SOFM）．N．W．
 Puttalam，5m（Mzlu）．S．Prov．： 1 §，Hikkaduwa， 11 mi NW of Galle， 10 m （MZLU）． $2 \delta^{\star} 2$ 오，Badagiriya Tank， 6.5 mi N Hambantota， 75 ft （ClJw）． $1 \delta^{\text {t }} 1$ ㅇ，Hambantota（BMNH）． $1 \delta^{\star}$ ，Tangalla（MCZC）． $1 \delta^{\star} 1$ ㅇ，Weligama（BMNH）．W． Prov．： $100^{\star} 11$ ㅇ，Colombo（BMNH）． $5 \delta^{\star}, 18 \mathrm{mi}$ NE of Colombo，Dambuwa Estate， 30 m （mZLU）．

Male．－Head dorsum mostly brown，posterior margin lighter，frontoclypeus，antennae，and palps brown．Maxillary palps 2.3 mm ；proboscis 0.9 mm ， as long as first $21 / 2$ segments of maxillary palp． Pronotum brown，slightly lighter in colour than head and mesonotum．Mesonotum and tegula brown． Forewing 12－14 mm，mostly brown，similar to that of D．robustior，with arc of small translucent spots distad of cord，hyaline spot at $\mathrm{m}-\mathrm{cu}$ and big spot at arculus． Legs with coxae brown and distal segments lighter． Modified spur（fig．44）slightly longer than adjacent spur，bifid with two unequal sinuate points directed distally，longer point almost $1 / 2$ as long as spur． Genitalia（fig．44）：Tergum IX apical $1 / 4$ overhanging segment X in lateral view，posterior margin with broad obtuse angle in dorsal view；sternum IX meso－ superior process triangular and extended above basal $2 / 3$ of phallus in lateral view．Segment X dorsal margin slightly concave with blunt apex in lateral view，oval－ shaped almost $1 / 2$ as wide as IX in dorsal view．Preanal appendages trapezoidal with dorsal margin horizontal in lateral view，ventral margin inclined dorsad and posterior margin slightly incised．Inferior appendages angled dorsad in lateral view，dorsal margin concave and nearly parallel with ventral margin，middle of ventral margin with obtuse angle，apex rounded；dis－ tal arms diverging in ventral view，lateral margins with base and apex rounded，mesal margin with squa－ rish basomesal angle，and short apicomesal point．

Distribution．－India：Karnataka，Tamil Nadu；Sri Lanka．

Remarks．－This species has male modified spur bi－ fid and somewhat similar to that of $D$ ．robustior，but differs by having spur with the shorter point nearly straight and directed distad．Until now the identity of D．notata has been enigmatic，because the locality stated by Fabricius（1781）as＇America boreali＇was very doubtful．McLachlan（1864）placed this species in Dipseudopsis，and suggested that the original loca－ lity information by Fabricius was in error，since at the time the genus was known only from South Africa
and China．Our examination of the male type of $D$ ． notata reveals that it is identical to Ulmer＇s（1915） original description of D．horni，only known from southern India and Sri Lanka，and we therefore con－ cur with McLachlan that the type of $D$ ．notata is mis－ labelled．

## Dipseudopsis recta Martynov， 1935

（figs．15，45）
Dipseudopsis recta Martynov，1935：163－164，fig．65a－b， Type，IndiA：‘ô．Sanjai River，Chakradharpur，Chota Nagpur．8－10．ii．1918．N．Annandale and F．H．Gravely （probably NZSI）．－Fischer 1962：14；Higler 192： 80.
Dipseudopsis bombayana Martynov，1935：164－165，figs． $66 a-b, 67$ ，Type，india：＇$\delta$＇．Gokak，Belgaum district， Bombay Presidency（at light）．June 1928 B．Prashad and H．S．Rao．＇［Type（NZSI）in poor condition；not able to borrow for examination，pers．comm．Dr．K．Rai］．－ Fischer 1962：5；Higler 1992：80．Syn．n．

Specimens examined．－india：Andhra Pradesh：10ㅈ， Lamnasinghi Ghats， 75 km SW Vishak， $850 \mathrm{~m}, 10 . \mathrm{VII} .1982$, D．R．Davis（USNM）．Bihar： $1 \delta^{\text {§ }} 5$ ，Namkum Ranchi， 675m，8．XI．61，Ross \＆Cavagnaro（CASC）．Karnataka： $12 \delta$ 1\％．Haveri，2．XII．1961，F．Schmid（CNCI）．18＇，Jannapura， 22．I．1959，F．Schmid（CNCI）．2§＇，Kakankote，10．I．1959，F． Schmid（CNCI）． $4 \mathbf{\delta}^{\$} 8$ ㅇ，Yellapur，30．I．1959，F．Schmid （CNCI）． $2 \delta^{\circ}$ ，Shimoga，R．Tunga， 1865 ft ，III－12．VIII，P．S． Nathan（MCZC）．Madhya Pradesh：Satanwara， 37 万 17 여，
 Phalghat，6．II．1959，F．Schmid（CNCI）．Maharashtra： $3 \delta^{\circ}$ 1 ㅇ，（Bombay）Sykes，5．II．1959，F．Schmid（cnci）．Tamil Nadu：3才，Varaiyattu Tittu，21．XII．1958，F．Schmid （CNCI）．26，Maraiyur，19．XII．1958，F．Schmid（CNCI）． $1 \delta^{\top}$ ， Sathuparai，1．XII．1958，F．Schmid（CNCI）．Nepal： $1 \begin{gathered}\text { oे，}\end{gathered}$ Chitwan Dist．，Sauraha S．I．Camp，31．X．1985，W．N． Mathis（USNM）．

Male．－Head brown，occipital margin light brown， frontoclypeus light brown except dorsal margin with dark band，antennae with dorsal portions brown，and ventral areas light brown，palps light brown，maxil－ lary palps 3.7 mm ，proboscis 1.5 mm ，almost as long as first three segments of maxillary palp．Pronotum brown，hind margin light brown，lateral sides light brown；mesonotum brown，with tegula and forew－ ings light brown．Forewing $10-12 \mathrm{~mm}$ long，faded ar－ ea distal to cord，small hyaline spots at m－cu and ar－ culus．Legs light brown．Modified spur（fig．45）about 1.5 x long as adjacent spur，apical $1 / 6$ with crown of three minute points，having two larger exterior points spiraling around minute middle point，middle point slightly curved，directed distad，inconspicuous and more lightly sclerotized than other points．Genitalia （fig．45）：IX tergum with apical $1 / 2$ overhanging $X$ in lateral view，posterior margin slightly incised in dorsal view；IX sternum mesosuperior process with blunt dorsal extension and slender posterior lobe overhan－ ging basal $1 / 3$ of phallus in lateral view．Segment $X$
slighely longer than preanal appendage, with lateral margins parallel and apex decply bilobed in dorsal vicw. Preanal appendages dorsal margin straight in lateral view, ventral margin curved dorsad, posterior margin truncate, apicodorsal angle square, ventroposterior angle obruse. Inferior appendages similar as in D. trichatlata, basal a twice as thick as apical portion in lateral view; with dorsal margin almost straight, ventral margin having distal $1 /$ incised, distal $1 /$ fingerlike but slightly acuminate; basal shelves separated by Vshaped notch in ventral view, posterior arms with blunt apicomesal point, lateral margin straight with curved apical and basal angles.

Female. - Tergite of abdominal segment VIII (fig. 15) with ventral submarginal recessed groove. Genitalia (fig. 15).

Distribution. - India: Andhra Pradesh, Bihar, Karnataka, Madhya Pradesh, Maharashtra, Tamil Nadu: Nepal.

Remarks. - D. recta is a sibling species to D. trichurara, but differs by having male modified spur with smaller apical points. The specific differences shown in the original species descriptions of $D$. recta and $D$. bombayana can be observed by rotating the modified spur. Hence, $D$. bombayana is considered to be a junior synonym of $D$. recta.

Dipseudopsis robustior robustior Ulmer, 1929 stat. n. (fig. 46)
$D_{\text {ipseudopsis robustior Ulmer, 1929: 185-186, fig. 22, Type }}$ scrics: Malaysta: West Malaysia: 'Material in Coll. :McLachlan: 1 8. Perak (etwas verletzt).' Cotype o *: abdomen missing, Perak (BMNH). - Fischer 1962: 14.
Dipseudopsis akhila Schmid \& Denning, 1979: 243-244, fig. 2. Holorype $\delta^{*}$ : Thalland: Chiang Mai Prov., Mae Ping. Chiang Mai, 1000f, 10.XI.1964, Peters (CLDD); P'aratypes". Tha Phra, Khon Kaen. Syn. n.
Dipscudopst chailandica Schmid \& Denning, 1979: 246, fig. 4. Holotype " ': Tha Phra, Khon Kaen, 24.XII.1968, Knapp; Paratypes". Nong Pora Kit bamroog, Roi-et, and Kalasin. Syn. n.
iepseudopss junki Marlier, 1979: 1-10, figs. 1-7, Type serics: $20^{\circ} 78,4$ larvae, 1 pupa, Bung Borapet, lake in central Thailand. 5.IX. 1971, Dr. W. Junk. Syn. n.

[^10]Lampam (bMnh). $8 \delta^{\circ}$, Tha Phra, Khon Kaen (Cidd). $7 \delta$, Chiang Mai, River Ping (Clhm, ZMUC). $10{ }^{\circ}$, Chiang Mai Zoo, light trap (ClHM). 27 ${ }^{\circ}$, Tap-Tan (Clhm). 10 , Chantaburi, Khao Soi Dao, 400 m (UOPJ). 18 , Kanchanaburi Dist., Sai Yok Nati. Park, 400 m (CLHM). 1 ठु, Nam Nao Natl. Park (сlhm). 30', Puk Hieo, Chayapoom (CLHM). $50^{\circ}$, Ayutthaya (CLHM). 20ㅇ, Bangsaen (CLHM). $1 \delta^{\circ}$, Tham Than Lod Natl. Park (clum). 18., Phuker, Kao Prataew Natl. Park (clhm). vietnam: 10, Long Xuyen (MNHN). 28 ${ }^{\circ}$, Hue (MNHN).

Male. - Head mostly brown, posterior warts light brown, frontoclypeus with ventral $1 / 2$ light brown and dorsal part brown, antennae mostly yellowish brown but dorsal surface dark brown, palps yellowish brown, maxillary palps 2.1 mm , proboscis 1.1 mm , as long as first three segments of maxillary palp. Pronotum and tegula light brown, mesonotum dark brown. Forewing 11-12 mm, similar as in D. collaris, with 57 small translucent spots in circular pattern around discal cell, having most spots distal to cord. Legs with coxae dark brown, distal segments yellowish brown. Modified spur (fig. 46) bifid, with two apical points, shorter point stout and angular, longer point slender, semicircular, and distant from other point. Genitalia (fig. 46): Tergum IX with short apical lobe extending over X in lateral view, posterior margin triangular in dorsal view; sternum IX mesosuperior process with short rounded dorsal lobe and long slender fingerlike posterior lobe in lateral view. Segment X dorsum inclined steeply ventrad toward blunt acuminate apex in lateral view, apex directed posteriad, having middle of ventral margin curved ventrad, and apicoventral margin slanted dorsad, ellipsoidal in dorsal view. Preanal appendages trapezoidal in lateral view, dorsal margin straight, ventral margin slanted dorsoposteriad, apicodorsal angle obtuse, apicoventral apex curved and extended more posteriad than apicodorsal angle. Inferior appendages with dorsal and ventral margins parallel and apical $1 / 2$ bent dorsad in lateral view, basal $1 / 2$ with thin basal shelf, and apical $1 / 2$ trapezoidal with apex slighted rounded, distal arms almost fingerlike in ventral view, lateral margins with apical and basal ends curved, basoventral shelf short, and mesal margin with squarish basal angle.

Distribution. - Burma; Cambodia; Malaysia: West Malaysia; Thailand; Vietnam.

Remarks. - D. robustior is the most common species of Dipseudopsis in Thailand, and is somewhat difficult to distinguish from $D$. collaris and D. infuscata. Possibly these species are merely variants of a widely distributed species.


Figures 46-50. - Dipseudopsis, A-G, male genitalia: A, lateral; B, preanal appendages, segments IX and X, dorsal; C, inferior appendages, ventral; D, phallus, lateral; E, phallus, ventral; F, segments IX, X, dorsal; G, ventral. H, see specific descriptions below. I-J, male modified apical spurs of hind tibia, ventral. Fig. 46, D. robustior Ulmer. Fig. 47, D. robustor andamanensis, n. ssp. 47 I, modified spur, isolated spur depicted at left. Fig. 48, D. schmidi, n. sp., 48 H . left inferior appendage, ventral, variation with apicomesal tooth. Fig. 49, D. spectabilis Banks. Fig. 50, D. stabatensis Malicky \& Weaver; 50H, segment IX, dorsal.

Dipseudopsis robustior andamanensis ssp. $\mathbf{n}$. (fig. 47)

Dipscudopsis infiuctata. - Malicky 1984: 214, 216 [misder.].
Type data. Holorype ठ : INDIA: Sourh Andaman Island: Mongelutonge, 20.X11.1976, Starmühlner (ClHM). Paranyes: $1 \delta^{\circ} 5$ ?, same data (CLHM); $2 \delta^{\circ} 15$ ㅇ, Bimblton, 22.xil. 1976. Starmühlner (сlemi).

Male. - Head, body, wings and genitalia similar to D. robustior. Modified spur (fig. 47) with two apical subequal in length and size.

Remarks. - This subspecies of D. robustior has unique, pincher-like modified spur (fig. 47) with two subequal points, but in D. robustior (fig. 46) the apical points of the modified spur are unequal, having a long slender curved point and a short broad point. It differs from $D$. infuiscata (fig. 41) which has the modified spur with a long straight point directed distad and a shorter curved point. In D. collaris the points are shorter, only $1 /$ as long as the spur. This subspecies represents the only member of the genus known from the Andaman Islands. All previous records of D. infiscata from South Andaman Island (Malicky 1984) are now recognized as this subspecies.

Etymology. - This subspecies is named after the Andaman Islands.

Distribution. - India: South Andaman Island.

## Dipseudopsis schmidisp. n.

(fig. 48)
Type material. - Holotype ठ': IndiA: Manipur: Kaiphundai, 20.V.1960, F. Schmid (CNCI). Paratypes: $90^{\circ}$, same data, F. Schmid (CNCI). $1 \delta^{\star}$, Kambiron, 24.V.1960, F. Schmid (cNCI). Assam: $60^{\circ} 2$ 2 , Garampani Res., 10 mi S Golaghat, $110 \mathrm{~m}, ~ 9 . X .61$, E. S. Ross \& D. Cavagnaro (-
C). $13^{3}$, Kaziranga, $75 \mathrm{~m}, 26.36 \mathrm{~N} 93.28 \mathrm{E}, 7-9 . \mathrm{V} .1976$, Wittmer (clemm). 'U.D.M.N.C.H' [United District of Mikir and North Cachac Hills]: 1才, Langtrang, 30.IV. 1960 , F. Schmid (CNCI). BANGLADESH: $10^{7} 2$, Lawa Chera For., Srimangal, $110 \mathrm{~m}, 27 . \mathrm{IX} .6 \mathrm{I}, ~ E . ~ S . ~ R o s s ~ \& ~ D . ~$ C.avagnaro (CASC).

Male. - Head dorsum, pronotum and cervical sclerites light orangish brown, but head slightly darker. Frontoclypeus dark brown except for small orangish brown triangular area between bases of antennae; genae yellowish brown except ventral sides dark brown. Scape mostly orangish brown, but apical portion and remainder of antenna dark brown. Mouthparts, with palps dark brown, maxillary palps ? 5 mm , proboscis 1.5 mm , as long as first four seg... .. of maxillary palp. Mesonotum and tegula dark .... F.....ing 11-13 mm, dark brown with indis-
tinct translucent spots, large basal triangular translucent spot bordered by $\mathrm{Sc}, \mathrm{Cu}_{1}$ and first branches of Rs and M , another translucent spot distal to cord. Hindwing lighter brown than forewing, with translucent basal triangular area between Sc and $\mathrm{Cu}_{1}$. Legs with colouration similar as in D. nebulosa, mostly dark brown, but apical $1 / 2$ of front and middle femora lighter brown, as well as tibiae and tarsi. Hind leg mostly dark brown, tarsi slightly lighter. Modified spur (fig. 48) with basal $2 / 3$ unmodified, distal $1 / 3$ with setose pocket at base of straight acuminate apex pointed distad. Genitalia (fig. 48): Tergum IX broad and triangular in dorsal view, long with apical $1 / 3$ overhanging segment X in lateral view. Segment X broad, extending distad slightly further than inferior appendages in lateral view, almost twice as long as broad and somewhat ellipsoidal, with dorsal and ventral margins inclined ventrad but extended posteriad at apex, apex broadly curved; elongate and cordate in dorsal view, with lateral margins tapered distad and apex slightly incised. Preanal appendages broad and rhomboid in lateral view. Inferior appendages broad and somewhat rectangular with curved ventral margin in lateral view, dorsal margin with short broad basal hump, maximum height about $2 / 3$ total length.

Etymology. - Named after its collector Fernand Schmid.

Distribution. - Bangladesh; India: Assam, Manipur.

Remarks. - This species is easily recognized by having the male modified spur acuminate and nearly straight. The male is somewhat similar to $D$. nebulo$s a$, but differs by having light areas of forewing with faded borders and less distinct, and sternum IX with mesosuperior process not short and globular, but long and slender, extending above basal $2 / 3$ of phallus.

## Dipseudopsis spectabilis Banks, 1931

(figs. 9, 49)
Dipseudopsis spectabilis Banks, 1931a: 69, pl. 5: fig. 1, Type $\delta^{*}$ : MALAYSIA: Sabah: Sandakan, Baker (mCzC 16421). Fischer 1962: 15.

Male. - Head dorsum with anterior portion dark brown, posterior warts and lateral margins yellowish brown, frontoclypeus dark brown, genae and antennal scape yellowish brown, but flagellum dark brown, palps dark brown; maxillary palps 1.8 mm , labial palps 0.6 mm , proboscis 1.4 mm , longer than first four segments of maxillary palp. Pronotum, mesonotum and tegula dark brown. Wings (fig. 9): Forewing 11 mm , fork I absent, conspicuous pattern of lighter
bands on dark brown background, similar as in $D$. flavisignata, with five translucent spots: 1) long slender stripe along anterior margin, $2 / 3$ as long as $S c ; 2$ ) small oval-shaped spot at base of Rs; 3) cordate spot in discal and medial cells; 4) truncate $V$-shaped band running basad from subapex to crossvein $\mathrm{r}-\mathrm{m}$, then distal to cord to base of $\mathrm{M}_{4}$, and to submargin through fork IV including $M_{4,}$ 5) long irregular semicircular band running from base of anal margin, tapering at $\mathrm{m}-\mathrm{cu}$, and widening toward margin at arculus. Modified spur (fig. 49): trifid, with three apical points, first point longest, slightly curved at base and $1 / 2$ as long as spur, second point $1 / 2$ as long as spur, and third point minute and adjacent to second point and located in curved notch between first and second points. Genitalia (fig. 49): Tergum IX with posterior margin slightly extended over segment X in lateral view, posterior margin broadly rounded in dorsal view; sternum IX mesosuperior process acuminate, dorsal margin irregular, ventral margin nearly straight, extending above basal $3 / 4$ of phallus. Segment X dorsal and ventral margins nearly parallel in lateral view; thumblike with lateral margins parallel in dorsal view. Preanal appendages trapezoidal in lateral view, posterior margin slightly incised and lateral side with ridge parallel to anterior margin. Inferior appendages heavily sclerotized, fingerlike with dorsal and ventral margins tapering slightly distad in lateral view, having short ventrobasal shelf, and dorsal margin with some minute bumps; squarish in ventral view, with lateral arms fingerlike with lateral margins converging slightly distad and mesal margin finely serrate.

Distribution. - Malaysia: Sabah.
Remarks. - This species is only known from the unique type; it is a sibling species to $D$. flavisignata and differs from it in having the male forewing with a translucent subapical truncated V-shaped band. It also differs by characteristics of the male modified spur and genitalia.

## Dipseudopsis stabatensis Malicky \& Weaver, 1988

 (fig. 50)Dipseudopsis stabatensis Malicky \& Weaver, 1988: 4-5, fig. Sa-e, Holotype ठ** $^{*}$ indonesiA: Sumatra: Dolok Merangir, 2-18.V.1971, Diehl (ClHM). Paratype ©*: Stabat 23.VI. 1974 (ClJw).
Dipseudopsis morosa, - Banks 1924: 450 (pro parte) [misdet. syntype 1 '**: Indonesia, Kalimantan, 'Borneo', Telang, X. 1881 (MCZC)].

Male. - Head dorsum with dark brown median stripe tapering posteriad, lateral sides and posterior warts yellowish brown, frontoclypeus dark brown, genae light brown, antennae light brown, palps light
brown, maxillary palps 2.4 mm , proboscis 1.2 mm , as long as first three segments of maxillary palp. Pronotum with raised areas yellowish brown, but deep median furrow and lateral sides dark brown. Mesonotum and tegula dark brown. Metanotum light brown. Forewing 13 mm , brown with translucent apical and anal spots, hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Fore and mid legs with coxae dark brown and distal segments yellowish brown, hind leg yellowish brown. Modified spur (fig. 50) 11/3x longer than adjacent spur, apex bifid with two minute curved apical points, each about $1 / 3$ length of spur. Genitalia (fig. 50): Tergum IX posterior margin bilobed, with lobes separated by squarish mesal notch in dorsal view, minute portion of posterior margin overhanging segment X in lateral view; sternum IX mesosuperior process similar to that of D. flinti, but slightly slenderer in lateral view. Segment X thumblike with lateral sides parallel and sinuate in dorsal view. Preanal appendages ellipsoidal and extended dorsoposteriad in lateral view. Inferior appendages capitate with trapezoid apex extended dorsad in lateral view; lateral arms fingerlike with lateral margins straight and parallel in ventral view, and mesal margins with broad curved basolateral incisions.

## Distribution. - Indonesia: Kalimantan, Sumatra.

Remarks. - This species is similar to $D$. flinti, but differs by having inferior appendage slenderer in lateral view, and the modified spur with apical points shorter.

## Dipseudopsis tonkinensis Navás, 1921

(fig. 51)
Dipseudopsis tonkinensis Navás, 1921: 79-80, Holotype ơ*: VIetnam: 'Tonkin, Chogahn, 1919, P. de Joannis' (MZBS). - Fischer 1962: 16; Fischer 1972: 8.

Specimens examined. - CHINA: Hainan: 10, Ta Hau, 4.VII.35, J. L. Gressitt (MCZC).

Male. - Head dorsum brown, posterior margin lighter brown, frontoclypeus brown, antennae brown but ventral side lighter brown, palps brown, maxillary palps 2.2 mm , proboscis 1.0 mm , as long as first three segments of maxillary palp. Pronotum brown, but posterior margin and lateral sides yellowish brown. Mesonotum and tegula brown. Forewing 12.5 mm , faded brown with small hyaline spots at m-cu and arculus. Legs with coxae brown and distal segments yellowish. Modified spur (fig. 51) bifid, with two robust acuminate points, first point directed obliquely distad and second point curved mesad. Genitalia (fig. 51): Tergum IX posterior margin only slightly overhanging segment X in lateral view, posterior margin

broad triangular in dorsal view; sternum IX mesosuperior process in lateral view with blunt dorsal projection and acuminate posterior process extended above basal $1 / 2$ of phallus. Segment $X$ dorsal margin slightly concave and slanted ventrad having blunt apex in lateral view, ellipsoidal with apex slightly incised in dorsal view. Preanal appendages dorsal margin straight and ventral margin inclined dorsad distad with posterior margin broadly incised in lateral view. Inferior appendages, clavate, being slightly broadened in lateral view, bent dorsad with oblique angle in middle, dorsal and ventral margins parallel, apex truncate; apical arms with apicolateral margins rounded and slightly broadened in ventral view, apicomesal angles squarish, mesal margins with square basolateral notch and square basomesal shelf.

## Distribution. - China: Hainan; Vietnam.

Remarks. - This species is similar to $D$. collaris, but differs by having male modified spur with apical points shorter, and by having the male genitalia with segment X broader in dorsal view.

## Dipseudopsis triclavata Martynov, 1935

## (fig. 52)

Dipseudopsis triclavata Martynov, 1935: 161-162, figs. 62ab, 63a-b, 64. New Lectotype ot $^{*}$ : INDIA: Maharashtra: Khanapur, Belgaum dist., Bombay Presidency, 1819.XI.1928, B. P. \& H. S. R. (NzsI). Remainder of type series: 1 ㅇ, same data as lectotype. $1{ }^{\text {đ }}$ [doubtful det.], Meghalaya: The Peak, Shillong, $8,400 \mathrm{ft}, 25.34 \mathrm{~N} 91.53$, 12.X.1914. S. W. Kemp. - Fischer 1962: 16; Higler 1992: 80.
Dipseudopsis morosa. - Schmid 1958: 13, 24, 27, 89, pl. 15: figs. 12-13 [misdet.].

Specimens examined. - IndiA: Karnataka: 3才, Uppinangadi (CNCI). $1 \delta$, Patan (CNCI). $14 \mathbf{x}^{\circ} 49$, Shimoga, R. Tunga, 1865 ff (MCZC). 2 す 10 ? , Bhadravati (MCZC). 8 đ 49 , Kakankote (CNCI). SRI LANKA: E. Prov.: Amparai Dist.: $1 \delta^{\circ}$, Inginiyagala, 250 ft (USNM). Cen. Prov.: $10^{\circ}$ det. Ulmer as D. stellata, Peradeniya (BMNH). Kandy Dist.: $110^{\circ} 69$, Hasalaka, Dam Spillway, 300ft (USNM). Matale Dist.: 2 ठ 27, Sigiriya, 800 ft (USNM). Nuwara Eliya Dist.: $2 \delta^{\delta}$, Milk Board Dairy, 4.2mi SW Nuwara Eliya, 6200ft (UsNM). N. Prov.: Vavuniya Dist.: $8{ }^{\circ} 29$, Parayanalankulam Irrigation Canal, 25 mi NW of Medawachchiya, 100 ft (USNM). N. Cen. Prov.: Anuradhapura Dist.: $4 \$^{\circ} 29$, Wildlife Soc. Bungalow, Hunuwilagama, Wilpattu Natl. Park, 200ft (USNM). NW. Prov.: Puttulam Dist.: 1 ${ }^{\text {T}}$, Wilpattu Natl. Park, 100 ft , Tala Wila (USNM). Sabaragamuwa Prov.: $1 \delta^{\top}$, Niriella (USNM). Ratnapura Dist.: $15 \delta^{\circ}$, Walawe Ganga, Embilipitiya, 100 ft (USNM). S. Prov.: Galle Dist.: $1 \delta^{\text {to }} 19$, Hiniduma (USNM). Hambantota Dist.: $1 \delta^{\delta}$, Palatupana, 10 ft (UsNM). Uva Prov.: Badulla Dist.: 1 © 1 iq, Dunhinda Falls, 1300 ft (USNM). Monaragala Dist.: 903, Sella Karagama, Menik Ganga, 150 ff (USNM). W. Prov.: Colombo Dist.: 1 §ै, Malewana (USNM).

Male. - Head dorsum brown, except narrow yellowish brown posterior margin; frontoclypeus mostly yellowish brown with brown stripe along dorsal margin, antennae dorsum brown and ventral sides yellowish brown; palps light brown, maxillary palps 2.5 mm , proboscis 1.3 mm , as long as first three segments of maxillary palp. Pronotum mostly brown, but lateral sides yellowish brown, propleuron and cervical sclerites yellowish brown. Mesonotum and tegula brown. Forewing 11-14 mm, dark brown with translucent stigma and hyaline spots at m -cu and arculus. Fore and mid legs with coxae to mid femora dark brown and apical $1 / 2$ of femora and distal leg segments light brown; hind leg yellowish brown. Modified spur (fig. 52) trifid, apex of spur appears truncate, but with three distinct apical points spiraling in plane perpendicular to longitudinal axis of spur, having one minute point and two longer points, long points each $1 / 1$ as long as spur. Genitalia (fig. 52): Tergum IX with minute portion of apex overhanging segment $X$ in lateral view, posterior margin truncate with broadly rounded lateral corners in dorsal view. Sternum IX mesosuperior process with rounded triangular dorsal process and smaller posterior lobe extended above ba$\mathrm{sal} 2 / 3$ of phallus in lateral view. Segment X dorsal margin slightly concave and inclined steeply ventrad in lateral view, apex acuminate with rounded end, ellipsoidal with apex incised in dorsal view. Preanal appendages trapezoidal with dorsoposterior apex broadly rounded and dorsal and posterior margins concave in lateral view. Inferior appendages similar to $D$. rec$t a$, base 2 x as thick as fingerlike apical $1 / 2$ in lateral view, ventral margin of apical portion incised; basal shelves with broad V-shaped medial notch in ventral view, apical arms with middle of mesal margin having oblique incision, lateral apical margin rounded, basolateral margin squaxish and apicomesal angle squarish with short blunt tooth.

Distribution. - India: Karnataka, Maharashtra; Sri Lanka.

Remarks. - This species is closely related to D. rec$t a$, but it differs by having male modified spur with longer apical points. There has been some confusion regarding the identity of this species, due to an erroneous redescription of ' D . morosa' by Schmid (1958), based on a specimen from Sri Lanka that was actually $D$. triclavata. D. morosa is only known from the southern islands of the Philippines. The male modified spurs of these species are similar, however the spur of $D$. triclavata (fig. 52) differs by being trifid, but that of $D$. morosa (fig. 39) is bifid. The record of the male syntype from Shillong, Meghalaya by Martynov (1935) is doubtful and should be confirmed.

Dipseudopsis varians Ulmer, 1929
rig. 531
Dipsesudopsis zarrians Ulmer, 1929: 191-192, figs. 28-30, Type series: Malaysla: West Malaysia: 'Material in Coll. Mitachlan: $38^{\circ}$. Perak.' Type $1 \delta^{\circ}$ : Perak (BMNH), condition poor, abdomen and hind legs absent, no other type matcrial available at Mus. (pers, comm. P. C. Barnard), Fischer 1962: 16.
Dipseudopss perersorum Schmid \& Denning, 1979: 345, fig. 3a-e. Holorype ō, thalland: Chiang Mai Prov., east lork. Mace Ping at junction of small stream, 59 km north Chiang Mai, 1350ft, at light, 30.XI.1964, W. L. \& J. G. Peters. Syn. n.

Specimens examined. - BURMA: Tenasserim: $10^{\hat{0}}$, Mekane, 90 km E Moulmein, 200 m (NHRS). MALAYSIA: West Malaysia: $10^{\circ} 2$ ??, Melaka, det. Ulmer 1905 as Dipseudopsis sp., 'de Malacca' Presquile, 1899, Errington de la Croix et P. Chapé (MNHN). $10^{\circ}$, Johor: Kota Tinggi (BMNH). $10^{\star}$, Negeri Sembilan: Kuala Pilah, at light (BMNH). $30^{7}$, Pahang: Kuala Tahan, at light, (MCZC, BMNH, BPBM). 20', Kedah: Bumbong Lima, light trap (USNM). 10*, Selangor: Kuala Lumpur (bMNH). Thalland: $10^{*}$, Trang (MCZC). $10^{\star} 1$ if, 40 mi SE Prachuab, 75 m (CASC). $1 \delta^{\circ}$, det. Denning as $D$. perensonum, Krabinburi, Thung Pho, 5.X.1970, Balmer ((ClDD). $1 \delta^{\circ}$. Phuket, Tonesai waterfall (SOFM). $1 \delta^{\text {on }}$, Khao Soy Dan (Clhm). $1 \delta^{\top}$, Chantaburi, Phliu, 100 m (UOPJ). 20. Loei, Phu Rua, 800 m (UOPJ). $10^{\star}$, Nakorn Nayok Prov.. Khao Yai Natl. Park, 700 m (ZMUC). $23 \delta^{\top} 7$ ㅇ, Chiang Mai Zoo, light trap (CLHM).

Malc. - Head, cervical sclerites, and pronotum orangish brown. Mesonorum, tegula, frontoclypeus, palps and antennae dark brown, but genae lighter brown. Maxillary palps 2.4 mm , proboscis 1.3 mm , as long as first three segments of maxillary palps. Forewing 12 mm , colouration similar to that of $D$. nebulosa but borders of translucent spots more indistince, dark brown background, large triangular spot berween bases of $\mathrm{R}_{1}$ and $\mathrm{Cu}_{1}$ and translucent subapical spot proximal to fork I, similar to bilobed spot of D. nebulosa but second lobe below $\mathrm{R}_{\mathrm{i}}$ less distinct. Legs with coxae dark brown, distal segments light brown. Modified spur (fig. 53) shorter than adjacent spur, with modified apical portion $1 / 2$ as long as spur, having spiraling point with tuft of setae in basal pocket. Genitalia (fig. 53): Tergum IX with apical $3 / 4$ or -nore overhanging segment $X$ in lateral view, triangu-
with postcrior margin rounded in dorsal view; ternum [X mesosuperior process slender and acumiin lateral view. Segment X oblong with dorsal nd ventral mareins parallel in lateral view, apex - $\quad$ : rounded $\quad$ ith venter slightly incised; cordate I with lateral sides tapering distad in dorsal lunt and slightly incised. Preanal appen-D-shaned ; th obruse dorsal angle in lateral r appendages oblong with irregular dorI margins in lateral view, ventral margin al bump, dorsal margin irregu-
lar with conspicuous tooth in middle, distal arms fingerlike in ventral view, with lateral margins diverging slightly distad and basal shelves broad.

## Distribution. - Burma; Malaysia: West Malaysia;

 Thailand.Remarks. - This species is somewhat similar to $D$. immaculata, but differs by having male with modified spur shorter and twisted like a short corkscrew, and by having the male genitalia with inferior appendages with a mid dorsal tooth in lateral view and without an apicomesal tooth in ventral view.

## Dipseudopsis voluta Ulmer, 1906

(fig. 54)
Dipseudopsis voluta Ulmer, 1906: 87-88, 110, fig. 89, Type $\delta^{*}$ : INDONESIA: Sulawesi, 'Makassar', McL. coll, without abdomen and hind legs (BMNH). - Fischer 1962: 17; Fischer 1972: 8.

Specimens examined. - INDONESIA: Sulawesi: 10 , without hind legs, 'Makassar', McL. coll (BMNH). Tenggara: 1 §̂, Desa Aopa, 50m, 27.X.1989, at light, R. de Jong \& J. Huisman (RMNH). $1 \delta^{\text {ºn }}$, Moramo, Sg Sena, 50 m , 15.XI.1989, at light, R. de Jong \& J. Huisman (RMNH).

Male. - Head dorsum mostly dark brown, posterior warts light brown; frontoclypeus brown with dorsal portion dark brown, genae light brown, antennal scape dark dorsad and light brown ventrad. Maxillary palps 2.3 mm , apical segments $1-3$ yellowish brown, but basal segments 4-5 dark brown; proboscis 1.1 mm , as long as first three segments of maxillary palps, labial palps 0.5 mm , second and third segments minute. Pronotum mostly brown, posterior margin light brown with a few darker muscle scars, and propleuron yellowish brown. Mesothorax with notum and tegula dark brown, epimeron dark brown, episternum and sternum yellowish brown. Forewing 14-15 mm, dark brown with golden setae, and some translucent spots between $\mathrm{R}_{2+3}$ and $\mathrm{R}_{4+5}, \mathrm{M}_{1+2}$ and $\mathrm{M}_{3}$, and hyaline spots at $\mathrm{m}-\mathrm{cu}$ and arculus. Front leg with anterior portion of coxa dark brown, remainder of leg yellowish brown, mid and hind legs yellowish brown, but hind tibia with brown ring at subapical spur. Modified spur (fig. 54) 2 longer than adjacent spur, bifid with long subapical point spiraling around longer straight point, about $2 / 3$ as long as spur. Genitalia (fig. 54): Tergum IX broad with posterior margin blunt triangular in dorsal view; sternum IX dorsomesal process with blunt posterior lobe in lateral view. Segment $X$ similar as in D. tonkinensis, but base of ventral margin nearly straight and apex curved dorsad in lateral view. Inferior appendages basal $1 / 3$ with slender squarish ventral shelf, distal $2 / 3$ of ventral margin
bent dorsad, dorsal margin slightly concave, apex broadened slightly and obtuse.

Distribution. - Indonesia: Sulawesi.
Remarks. - This species is easily recognized by having the modified spur with a long straight apical point and another more slender point spiraling completely around its base. The genitalia of D. voluta are similar as in $D$. tonkinensis, but differs by having inferior appendages with short obtuse subapical mesal tooth in ventral view, and with apical portion slightly tapered in lateral view. This species was previously only known from the male type which lacks the abdomen and is in rather poor condition. Two male specimens were collected recently, making it possible to provide a description of the male genitalia for the first time.

## Nomina dubia

## Dipseudopsis onychophora Navás, 1935

nomen dubium
Dipseudopsis onychophora Navás, 1935: 104-105, Type ठै: india: 'Lonawla (Bombay), 9-X-1934. H. Benavent, S. J.' [location of type unknown]. - Fischer 1962: 14; Higler 1992: 80.

Remarks. - The distributional data of other species of Dipseudopsis in India and the figure provided in the original description of $D$. onychophora by Navás (1935) of the modified spur of the male type, suggests that this species is probably synonymous with $D$. triclavata. However, since we are unable to locate and examine the type, the status of this species remains uncertain.

## Dipseudopsis orientalis (Navás, 1913) nomen dubium

Esperona orientalis Navás, 1913: 12-13, fig. 2a-c, Type, sex unknown: vietnam: 'Hanoï (Tonkin). Un échantillon envoyé et cédé généreusement pour ma collection par M. Laboissière.' [location of type unknown]. - Fischer 1962: 14.

Remarks. - We were not able to locate and examine the type of this species, hence its identity remains uncertain. D. orientalis could be synonymous with one of the three species of Dipseudopsis known from Vietnam, D. benardi, D. robustior, and D. tonkinensis. According to the original species description (Navás 1913) the forewing of the type of $D$. orientalis is 12.4 mm , suggesting that this species is probably synonymous with either D. robustior or D. tonkinensis.

## POLYCENTROPODIDAE ULMER

Eodipseudopsis Marlier 1959 nomen dubium
Eodipseudopsis Marlier 1959: 117. Type species: Eodipseudopsis tomensis Marlier (original designation).

The genus Eodipseudopsis is removed from the Dipseudopsidae and is provisionally placed in the Polycentropodidae. Three of its characteristics support its removal from Dipseudopsidae: 1) Female genitalia as figured in the original description have sternum VIII bearing a pair of ventrolateral processes (Marlier 1957: fig. 10A). 2) The antennae (cf. Marlier 1959: fig. 48A) are slender with their bases distant from each other. 3) The long slender lobe of the sent gland of sternum $V$ (Marlier 1962: fig 10B) is uncharacteristic of dipseudopsid females examined, i.e. Dipseudopsis, Hyalopsyche, Phylocentropus.


Figure 55. Total tree output calculated by Hennig 86 for Hydropsychoidea, including outgroup and genera of Dipseudopsidae.


Figute 56．Consensus tree of the families of the Hydro－ psychoidea s．sir．Weaver，showing phylogenetic relation－ ships of the genera of Dipseudopsidae．

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## References

Albarda，H．，1881．Neuroptera．－In：J．P．Veth，ed．， Midden－Sumatra 4 （2），Natuurlijke Historie．5：1－22， pls．1－6，Leiden．
Banks，N．，1907．Descriptions of new Trichoptera．－ Proceedings of the entomological Society of Washington 8：117－33，pls．8－9．
Banks，N．，1911．Notes on Indian neuropteroid insects．－ Proceedings of the entomological Society of Washington 13：99－106，pl． 6.
Banks，N．，1913．Synopsis and descriptions of exotic Neuroptera．－Transactions of the American entomolog－ ical Society 39：201－42，pls．23－6．
Banks，N．， 1914 （1913）．On a collection of neuropteroid in－ sects from the Philippine Islands．－Proceedings of the en－ tomological Society of Washington $15: 170-80$ ，pls．8－9． （22 January 1914）．
Banks，N．，1916．Neuropteroid insects of the Philippine Islands．－Philippine Journal of Science 11：195－217，pls． 1－2．
Banks，N．，1920．New neuropteroid insects．－Bulletin of the Museum of Comparative Zoology 64：299－362，pls． 1－7．
Banks，N．，1924．Descriptions of new neuropteroid insects． －Bulletin of the Museum of Comparative Zoology 65： 421－55，pls．1－4．
Banks，N．，1931a．Some Oriental neuropteroid insects．－ Psyche 38：56－70，pl． 5.
Banks，N．，1931b．Some neuropteroid insects from the Malay Peninsula．－Journal of the Federated Malay Museums 16：377－409．
Brauer，F．，1868．Neue von Herrn Dr．G．Semper gesa－ mmelte Neuropteren．－Verhandlungen der zoologisch－ botanischen Gesellschaft in Wien 18：263－268， 2 pls．
Crichton，M．I．，1957．The structure and function of the mouth parts of adult caddis flies（Trichoptera）．－ Philosophical Transactions of the Royal Society of London（Series B）241：45－91．
Cummings，B．F．，1913．Apropos of the first maxillae in the genus Dipseudopsis Walk．（Trichoptera）．－Annals and Magazine of Natural History 11：308－312．
Fabricius，J．C．，1781．Species Insectorum．Vol．I， 552 p．－ Hamburg．
Farris，J．S．，1988．Hennig86，version 1．5．Program and doc－ umentation．Port Jefferson，New York．
Fischer，F．C．J．，1962．Polycentropodidae and Psychomyi－ dae．Trichopterorum Catalogus．Nederlandsche Entomo－ logische Vereeniging，Amsterdam 3： 236 p．
Fischer，F．C．J．，1972．Supplement to volumes III and IV． Trichopterorum Catalogus．Nederlandsche Entomologi－ sche Vereniging，Amsterdam 13： 172 p．
Flint，O．S．，Jr．1991．Studies of Neotropical caddisflies，

XLV: The taxonomy, phylogeny, and faunistics of the Trichoptera of Antioquia, Colombia. - Smithsonian Contributions to Zoology, no. 520: 1-113.
Gibbs, D. G., 1968. The larva, dwelling-tube and feeding of a species of Protodipseudopsis (Trichoptera: Dipseudopsidae). - Proceedings of the Royal entomological Society London (Series A) 43: 73-79.
Higler, L. W. G., 1992. A check-list of the Trichoptera recorded from India and a larval key to the families. Oriental Insects 26: 67-106.
Iwata, M., 1927. Trichopterous larvae from Japan. Annotationes zoologicae japonenses 11: 203-233.
Kimmins, D. E. 1957. Lectotypes of Trichoptera from the McLachlan Collection now in the British Museum (Natural History). - Bulletin of the British Museum (Natural History) 6(4): 93-126.
Lepneva, S. G., 1964. Trichoptera (Rucheiniki), Larvae and pupae of Annulipalpia. In: Pavlovskii, E. N., ed., Fauna of the U.S. S. R., 2 (1): 560 p. (Translated into English, by Program for Scientific Translations, Jerusalem 1970).
Lestage, J. A., 1925. Notes trichoptérologiques (7me Note).

- Bulletin et Annales de la Société royal entomologique de Belgique 65: 35-44.
Lestage, J. A., 1936. Notes trichoptérologiques. XIV. - Les composantes de la faune sud-africaine et la dispersion transafricaine de quelques espèces. - Bulletin et Annales de la Société royal entomologique de Belgique 76: 165192.

McLachlan, R., 1863. On Anisocentropus, a new genus of exotic Trichoptera, with descriptions of five species, and of a new species of Dipseudopsis. - Transactions of the entomological Society of London 1(3): 492-6, pl. 19.
McLachlan, R., 1864. On the types of Phryganidae described by Fabricius from the Banksian Collection. Transactions of the entomological Society of London 1: 656-659.
McLachlan, R., 1866. Descriptions of new or little-known genera and species of exotic Trichoptera, with observations on certain species described by Mr. F. Walker. Transactions of the Royal entomological Society of London 5: 247-75, pls. 17-9.
McLachlan, R., 1875. Descriptions de Plusieurs Névro-ptères-Planipennes et Trichoptères nouveaux de l'île de Célèbes et de quelques espèces nouvelles de Dipseudopsis avec considérations sur ce genre. - Tijdschrift voor Entomologie 18: 1-21, pls. 1, 2.
Malicky, H., 1984. The caddisflies (Trichoptera) collected by the Austrian-Indian Mission in 1976 on the Andaman Islands. - Annalen des Naturhistorischen Museums in Wien 86: 213-8.
Malicky, H. \& J. S. Weaver III, 1988. Die von Dr. E. W. Diehl in Sumatra gesammelten Dipseudopsis-Arten (Trichoptera: Dipseudopsidae). - Opuscula Zoologica Fluminensia 24: 1-9.
Marlier, G., 1953 (1952). Études hydrobiologiques dans les rivières du Congo oriental. B. Trichoptères - Polycentropodidae (Dipseudopsinae). - Annales du Musée royal du Congo belge, Série $8^{\circ}$ (Sciences Zoologiques) 21: 51-65.
Marlier, G., 1955. Un Trichoptère pélagique du Lac Tanganika. - Revue de zoologie et de botanique africaines 52: 150-155.
Marlier, G., 1959. Mission du Muséum dans les Iles du Golfe de Guinée. Entomologie XII (1). Trichoptères de Sao Tomé. - Revue Française d'entomologie 63: 108120.

Marlier, G., 1961. La nymphe et la position systematique de Limnoecetis tanganicae Marlier (Trichoptera). - Bulletin Institut royal des Sciences naturelles de Belgique 38: 1-6.
Marlier, G., 1962. Genera des Trichoptères de l'Afrique. Annales du Musée Royal de l'Afrique Centrale, Série 80 (Sciences Zoologiques) no. 109: 1-261.
Marlier, G., 1979. Une nouvelle espèce de Dipseudopsis de Thaillande (Trichoptères: Polycentropidae). - Bulletin Institut royal des Sciences naturelles de Belgique 51: 1 11.

Martynov, A. [V.], 1935. On a collection of Trichoptera from the Indian Museum. - Records of the Indian Museum 37: 93-209.
Mosely, M. E., 1933. Synonyms in the order Trichoptera. Annals and Magazine of Natural History (series 10) 12: 497-503.
Navás, L., 1913. Espèces nouvelles de Néveroptères exotiques. - Annales de l'Assacociation des naturalistes de Levallois-Perret 19: 10-13.
Navás, L., 1915. Neurópteros nuevos o poco conocidos (cuarta serie). - Memorias de la Real Academia de ciencias y artes de Barcelona 11(23): 373-98.
Navás, L., 1921. Insectos exóticos neuvos, críticos o poco conocidos. - Revista de la Academia de Ciencias exactas físico-químicas y naturales de Zaragoza 6: 61-81.
Navás, L., 1924. Neue Trichopteren. - Konowia, Wien 3: 204-9.
Navás, L., 1930. Insectos del Museo de Paríe ( $6^{2}$ serie). Brotéria, Série zoológia 26(3): 120-44.
Navás, L., 1931. Decadas de insectos neuvos. - Revista de la Academia de ciencias exactas físico-químicas y naturales de Madrid 26: 60-86.
Navás, L., 1935. Décadas de insectos nuevos. - Brotéria, Série Ciencias naturais 4(31): 97-107.
Ross, H. H., 1956. Evolution and classification of the mountain caddisflies. Univ. Illinois Press, Urbana, 213 p.
Ross, H. H., 1967. The evolution and past dispersal of the Trichoptera. - Annual Review of Entomology 12: 169206.

Ross, H. H. \& J. M. Kingsolver, 1959. The Madagascan fauna of Dipseudopsis with speculations about the phylogeny and dispersal of the genus (Trichoptera). - Mémoires de l'Institut Scientifique de Madagascar, Série E 11: 103128.

Ross, H. H. \& D. G. Gibbs, 1973. The subfamily relationship of the Dipseudopsinae (Trichoptera, Polycentropodidae). - Journal of the Georgia entomological Sociery 8: 312-6.
Schmid, F., 1949. Les Trichoptères de la collection Navás. Eos, Madrid 25: 305-426.
Schmid, F., 1958. Trichoptères de Ceylan. - Archiv für Hydrobiologie 54: 1-173, 34 pls.
Schmid, F., 1980. Les insectes et arachinides du Canada, partie 7: Genera des Trichoptères du Canada et des États adjacents. Agriculture Canada. pub. 1692. Ottawa. 296 p.

Schmid, F. \& D. G. Denning, 1979. Descriptions of new Annulipalpia (Trichoptera) from southeastern Asia. Canadian Entomologist 111: 243-9.
Schuster, G. A. \& S. W. Hamilton, 1984. The genus Phylocentropus in North America (Trichoptera: Polycentropodidae). - Proceedings International Symposium on Trichoptera 4 (Series Entomologica 30): 347-62.
Scott, K. M. F., 1985. Order Trichoptera (caddis flies). In: C.H. Scholtz \& E. Holm, eds., Insects of Southern

Mirka. p. 32--40. Buttenworths, Durban.
Iisuda. M.. 1434. Metamorphose von drei Köcherfliegen, Whannaz fikkitha C'lmer. Tinedes satuteri Ulmer und Depersdopses sellatad Machachtan. - Annotationes Zoolofivac laponenses 18(3): 20--212.
Tivia. M.. 1942. Japanische Trichoperen, pt. 1. swematik. - Memoirs of the College of Science, Kyoto Imperal Universiey, Series B, 17(1): 239-339.
Clmer. (i.. 1904,3. U'her cinige Trichopteren mit rüsselforamigen Koptanhängen. - Zoologischer Anzeiger 28: 5654.
(Vmer. (i.. ( $\left.{ }^{( }\right)()+b$. Uber westafrikanische Trichopteren. Zoulugischer Anzeiger 28: 353-359.
('lmer. (i.. 1905. Neue und wenig bekannte außereuropaische Trichopteren, hauptsächlich aus dem Wiener Muscum, - Annalen des Naturhistorischen Museums in Wien 20: 59-98.
Llmer. G.. 1906. Neuer Beitrag zur Kenntnis aussereuropaischer Trichopteren. - Notes from the Leyden Muscum 28: 1-116.
Clmer, G... $190^{-}$a. Neue Trichopteren. - Notes from the Levden Muscum 29: 1-53.
Limer. G.. 190h. Trichoptera. In: P. Wytsman, ed., Genera Insectorum. V'erteneuil \& Desmer, Bruxelles, Fasc. 60, 254 p.. 41 pls.
('lmer. G.. 1909. Einige neue exotische Trichopteren. Notes from the Levden Museum 31: 125-42.
('lmer, G.. 1915. Trichopteren des Ostens, besonders von Ceylon und Neu-Guinea. - Deutsche entomologische Zcitschrift. Berlin pp. 41-75.
("imer. G.. 1929. Über einige, hauptsächlich asiatische, Pphemeropteren und Trichopteren aus der Sammlung R. Mclachlan. - Deutsche entomologische Zeitschrift (Ber(in) 3:161-195.
L"mer, G.. 1930. Trichopteren von den Philippinen und von den Sunda-Inseln. - Treubia 11:373-498.

Ulmer, G., 1951. Köcherfliegen (Trichopteren) von den Sunda-Inseln. Teil I. - Archiv für Hydrobiologie. Suppl. 19: 1-528, 28 pls.
Ulmer, G., 1957. Köcherfliegen (Trichopteren) von den Sunda-Inseln. Teil III. - Archiv für Hydrobiologie. Suppl. 23: 109-470, 31 pls.
Walker, F., 1852. Catalogue of the specimens of neuropterous insects in the collections of the British Museum, pt. 1, 135 p .
Wallace, J. B., W. R. Woodall \& A. A. Staats, 1976. The larval dwelling-tube, capture net and food of Phylocentropus placidus (Trichoptera: Polycentropodidae). - Annals of the entomological Sociery of America 69: 149-154.
Weaver, J. S., III., 1984. The evolution and classification of Trichoptera, part I: The groundplan of Trichoptera. Proceedings International Symposium Trichoptera 4 (Series Entomology 30): 413-419.
Weaver, J. S., III \& J. C. Morse., 1986. Evolution of feeding and case-making behavior in Trichoptera. - Journal of the North American Benthological Society 5: 150-8.
Weidner, H., 1964. Die entomologischen Sammlungen des Zoologischen Staatsinstituts und Zoologischen Museums Hamburg. - Mitteilungen aus dem Hamburgischen zoologischen Museum und Institut 62: 55-100.
Wells, A. \& D. Cartwright, 1993. Females and immatures of the Australian caddisfly Hyalopsyche disjuncta Neboiss (Trichoptera), and a new family placement. Transactions of the Royal Society of South Australia 117: 97-104.
Wiggins, G. B., 1977. Larvae of the North American Caddisfly genera (Trichoptera). Univ. Toronto Press, 401 p.
Wiggins, G. B., 1982. Trichoptera. - In: S. P. Parker, ed., Synopsis and classification of living organisms. McGrawHill, New York, vol. 2, p. 599-612.

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# REVIEW OF THE SOUTHERN PALAEARCTIC AND <br> PALAEOTROPICAL LEAFHOPPER GENUS HENGCHUNIA VILBASTE (HOMOPTERA, CICADELLIDAE) 


#### Abstract

Asche, M. \& M. D. Webb, 1994. Review of the southern Palaearctic and Palaeotropical leafhopper genus Hengchunia Vilbaste (Homoptera, Cicadellidae). - Tijdschrift voor Entomologie 137: 143-154, figs. 1-63 (ISSN 0040-7496). Published 15 December 1994. The leafhopper genus Pseupalus Remane \& Asche (Deltocephalinae: Paralimnini) is synonymized with Hengchunia Vilbaste syn. n. Four new species are described: H. helleri sp. n. from Cameroun and Zaire, H. indica sp. n. from India, H. pakistanica sp. n. from Pakistan and H. javana sp. n. from Indonesia: Java. Hengchunia now contains nine species which are probably all associated with the grass genus Imperata. The zoogeographical distribution of the species and their potential phylogenetic relationships are briefly discussed. Additions and amendments to Webb \& Heller's (1990) check-list of Afrotropical and Oriental Paralimnini are made and include Paralimnellus Emeljanov (1972), oldest available name for Bubulcus Dlabola (1961) (junior homonym of Bubulcus Bonaparte 1854, Aves) and Paralimnellus vittata (Matsumura) comb. n. from Henschia. Correspondence: Dr Manfred Asche, Am Schwanhof 2A, D-35037 Marburg, Germany. Key words. - Homoptera; Auchenorrhyncha; Cicadellidae; Paralimnini; Hengchunia; Pseupalus; review; new species; South Palaearctic and Palaeotropical Region.


This paper concerns a group of paralimnine leafhoppers (Cicadellidae: Deltocephalinae) apparently associated with the Palaeotropical grass Imperata, especially the species I. cylindrica. This association was reported by Remane \& Asche (1980) who erected the genus Pseupalus for a single paralimnine species, $P$. graecanarus from the Canary Islands (type locality) and Greece. Recently, Pseupalus and related taxa were reviewed by Webb \& Heller (1990) who added three more species: P. gaiseri from Ivory Coast, P. digitus from Thailand, and $P$. murtus from the Philippines: Luzon. Webb \& Heller (1.c.) discussed the validity of the tribe Paralimnini sensu Emeljanov (1962), Remane \& Asche (1980), Ossiannilsson (1983), and Emeljanov \& Kirilova (1989) and Hamilton's 'Deltocephalina' (Hamilton 1975), and confirmed the tribal placement of Pseupalus. In addition, Webb \& Heller (l.c.) provided a check-list of Afrotropical and Oriental paralimnine genera. Since then, the second author has examined specimens of a species from Taiwan, described as Thamnotettix koshunensis Matsumura, for which Vilbaste (1969) had established the genus Hengchunid. A comparison revealed not only the placement of Hengchunia in the Oriental Paralimnini, but also its generic synonymy with

Pseupalus. Moreover, four new species of this genus were discovered in samples from Africa, India, and Indonesia, described below, together with an unidentifiable female specimen from Australia. The genus Hengchunia now contains nine species: one in the Southwestern Palaearctic, two in the Afrotropical Region, and 6 in the Oriental Region.
The following amendments and additions apply to the Afrotropical and Oriental paralimnine check-list (Webb \& Heller 1990), noted above: Paralimnellus Emeljanov (1972) (type species: cingulatus Dlabola) oldest available name for Bubulcus Dlabola (1961) (type species: cingulatus Dlabola) (junior homonym of Bubulcus Bonaparte, 1854, Aves); Paralimnellus vittata Matsumura comb. n. (Henschia); Pteropyx acutus (Singh) (Jassargus) Emeljanov, 1972: 66; Pteropyx maculatus Emeljanov, 1972: 67 (Mali); Khasia prima Rao, 1989: 82 (India).

## Depositories

Abbreviations for depositories of material used in this paper: M. Asche \& H. Hoch, Marburg, private collection (AH); Agricultural University Wageningen, The Netherlands (auwn); The Natural History Museum, London, United Kingdom (bmnh); Musée


X"thb \& Hellerl, paratype of from Ivory Coast; Figs. 6, 10: ot from Sierra Leone. - 1, male bital plate, dorsal view; 3, same, dorsolateral view; 4, subgenital plates, ventral view; nd connective, ventrocaudal view; 8, aedeagus, left lateral view; 9,10 , same, left la$: 0.1 \mathrm{~mm}$.

Royal de l'Afrique Centrale, Tervuren, Belgium (mrac); Institut Royal des Sciences Naturelles de Belgique, Brussels, Belgium ( ISNB ); Entomological Institute, Hokkaido University, Sapporo, Japan (eiHu); Staatliches Museum für Naturkunde Stuttgart, Germany (smns); University of Agricultural Sciences, Bangalore, India (UASB).

## Taxonomy

## Hengchunia Vilbaste

Hengchunia Vilbaste, 1969: 8. - Type species by original designation: Thamnotettix koshunensis Matsumura.
Pseupalus Remane \& Asche, 1980: 88, syn. n. - Type species by original designation: Pseupalus graecanarus Remane \& Asche.

For relationship and diagnosis of Hengchunia see the corresponding paragraphs for Pseupalus in Webb \& Heller (1990: 2, 3).

## Key to the species of Hengchunia (males)

(after Webb \& Heller 1990, modified and supplemented)

1. Preatrium of aedeagus with a pair of long and slender processes (e.g., figs. 7, 8) 2

- Preatrium of aedeagus without a pair of processes (fig. 46)
H. pakistanica sp. n.

2. Aedeagal shaft with four spinose processes apically (fig. 63) H. koshunensis (Matsumura)

- Aedeagal shaft with two or less spinose processes apically or subapically (e.g., fig. 7) 3

3. Aedeagal shaft with a single spinose process or lateral flanges subapically 4

- Aedeagal shaft with two spinose processes apically or subapically .5

4. Aedeagal shaft with a horn-shaped subapical process directed ventrad, dorsal side of shaft at midlength with a bifurcate process and shortly basad of it with a single short process (figs. 32-34) ......
H. indica sp.n.

- Aedeagal shaft with a subapical flange on each side, devoid of spinose processes (see Webb \& Heller, 1990: 5, fig. 13) $\qquad$
$\qquad$ H. graecanarus (Remane \& Asche)

5. Subgenital plates with distal margin slightly concave (fig. 14); aedeagal shaft medially on dorsal side with a group of 3 short spines (fig. 18) .... 6

- Subgenital plates with distal margin convex (e.g., figs. 2, 22); aedeagal shaft medially on dorsal side devoid of short spines (e.g., figs. 8, 26, 61) ..... 8

7. Aedeagal shaft with apical processes almost directed laterad; mediodorsal processes comparatively distant from each other; style almost evenly tapering to apex without a subapical lobe (see

Webb \& Heller, 1990: 6 figs. 14, 15)
H. digitus Webb \& Heller

- Aedeagal shaft with apical processes directed ventrobasad; mediodorsal processes near to each other (fig. 18); style with a subapical lobe (fig. 16) ..
.H. murtus Webb \& Heller

8. Aedeagal shaft with processes apically, directed dorsocaudad, not reflected (figs. 60-62) ............ . H. javana sp. n.

- Aedeagal shaft with processes subapically or medially, directed laterobasad (e.g., figs. 7, 25) .... 9

9. Aedeagal shaft with processes located subapically (figs. 7-10) ............... H. gaiseri Webb \& Heller

- Aedeagal shaft with processes located mediolateral (figs. 25, 26)
H. belleri sp. n.


## 1. Hengchunia koshunensis (Matsumura)

(fig. 60)
Thamnotettix koshunensis Matsumura, 1914: 178. Lectotype
$\delta^{\pi}$, Taiwan (EIHU), here designated [examined].
Hengchunia koshunensis: Vilbaste 1969: 8, fig. 6: A-J.
Material examined. - Lectotype ठ', 'Formosa/Matsumura' and '5.vii.1906/Koshun' on reverse; 1 it, 1 ? (sex unknown). - Paralectotypes, originally on same mount as lectotype (re-mounted) (EIHU).


Fig. 11. Hengchunia murtus (Webb \& Heller), ot from Sulawesi. - 11, habitus. Scale: 1.0 mm .


[^11]Remarks．－H．koshunensis can be separated from the other Hengchunia species by the crown－shaped apex of the aedeagal shaft consisting of two pairs of spines：a pair of longer spines directed ventrad and a pair of shorter spines directed dorsad（not vice versa as noted in Vilbaste 1969：8）．
Distribution．－This species is only known from Taiwan（Matsumura 1914；Vilbaste 1969）．

## 2．Hengchunia graecanarus（Remane \＆Asche） comb．n．

Pseupalus graecanarus Remane \＆Asche，1980：90，fig． 16 a－ l．Holotype ô，Canary Is，（Remane coll．）［examined］．－ Webb \＆Heller 1990：5，fig． 13.

Material examined（additional to the type material of Remane \＆Asche 1980）．－20才，5우，S Greece， Peloponnessos，Skala nr Gythion，on Imperata cylindrica， 6．x．1981，M．Asche \＆H．Hoch．－ $90^{\text {® }}, 26$ 年，NW Greece， Corfu I．，Aghios Stephanos，coastal biotopes with Imperata cylindrica，14．x．1981，M．Asche \＆H．Hoch．－ $3 \delta^{\circ}, 1$ O O ，NW Greece， 27 km S．Igoumenitsa，S．Kastri，on Imperata cylin－ drica， 10 ．viii． 1983, M．Asche \＆H．Hoch（AH）．

Remarks．－Within the genus，H．graecanarus dis－ plays a unique configuration of the aedeagus with subapical flanges at the shaft instead of spinose pro－ cesses．
Distribution．－This species is known from the type－locality in the Canary Islands：Gran Canaria， and from several localities in Greece（Remane \＆ Asche 1980）．According to the distribution of its host plant，Imperata cylindrica，a wider distribution at least in the Mediterranean Region is possible．

## 3．Hengchunia digitus（Webb \＆Heller）comb．n．

Pseupalus digitus Webb \＆Heller，1990：4，figs．14－18． Holotype $\delta \overline{\text { on }}$ ，Thailand（вмлн）［examined］．

Remarks．－No material additional to the type ma－ terial has been examined．
$H$ ．digitus closely resembles $H$ ．murtus in the shape of the subgenital plates（including the toothed area） and aedeagus but the aedeagal shaft is straighter and has the three mediodorsal spines less closely adjacent and the style apex narrower．
Distribution．－This species has been reported on－ ly from Thailand（Webb \＆Heller 1990）．

## 4．Hengchunia gaiseri（Webb \＆Heller）comb．n． （figs．1－10）

Pseupalus gaiseri Webb \＆Heller，1990：4，figs．1－12． Holotype ${ }^{\text {on }}$ ，Ivory Coast（smns）［examined］．

Remarks．－No material additional to the type ma－ terial has been examined．
H．gaiseri is similar（and probably closely related） to $H$ ．helleri（see below）．It differs from this species in the distally more pointed subgenital plates and in the position of the lateral spinose aedeagal processes which are located subapically in H．gaiseri and in midlength of shaft in $H$ ．belleri．
Distribution．－Although this species was described from Ivory Coast it is likely that a similar male spec－ imen from Sierra Leone is conspecific（Webb \＆ Heller 1990：4）．

## 5．Hengchunia murtus（Webb \＆Heller）comb．n． （figs．11－19）

Pseupalus murtus Webb \＆Heller，1990：6．Holotype ô， Philippines（вмNн）［examined］．

Material examined（additional to the type material of Webb \＆Heller 1990）．－Indonesia：Sulawesi： $10^{\top}$ ，Sulawesi Utara，Dumoga Bone National Park，rainforest near base－ camp Toraut，19．x．1985；10 ，ibid．，banks of Toraut river；
 id．，trail to Tumpah river， 23 ．xi． 1985 ； $10^{\star}$ ，ibid．，trail to Tumpah river uphill to ca． $400 \mathrm{~m}, 20 . \mathrm{x} .1985$ ； 1 ㅇ，ibid． banks of Tumpah river；10 ，Molosso I．，opposite Lolok （N．－coast），11．xi．1985；all：on Imperata grass，Project Wallace Expedition，M．Asche \＆H．Hoch（Ah，вmnh）．

Remarks．－H．murtus closely resembles H．digitus in the shape of the male genitalia（see under $H$ ．digi－ tus）．
Distribution．－This species was originally descri－ bed from Luzon I．in the Philippines（Webb \＆Heller 1990）．Its occurrence in Sulawesi extends its range to the south．

## 6．Hengchunia helleri sp．n． （figs．20－26）

Type material．－Holotype ©゙：Zaire，Lubumbasha （＝Elisabethville），30．iii．1939，at light，H．－J．Brédo（ISNB）．－ Paratypes：Zaire：19才，43 9,2 ？（sex unknown），same data as holotype（various dates）（ISNB，MRAC，BMNH）； $30^{\top}, 3$ 우， Libenge，Savane Liki－Bembe，23－28．ii． 1948 （one without date）（R．Cremer \＆M．Neuman，（IsNb）；1오，Kivu， Kayimvira（Uvira），ii－iii．1955，G．Marlier（MRAC）； 20. Kinchassa，Waelbroeck， $24 . \mathrm{ii}$ and 3．iv． 1899 （ISNB）； $10^{\hat{\lambda}}, 1$ 우， Maka，Lualaba，25．i．1939，H．－J．Bredo（IsNB）．－Cameroun： $10^{\text {® }}$ ，Bertoua，14．ix．1960，F．J．Breteler（AUWN）．

Description．－Habitus，bodily proportions and colouration as in generic description（see Webb \＆ Heller 1990）．
Male genitalia：Subgenital plate（figs．20－23）distal－ ly broadly rounded；toothed area rather long，extend－ ing from apex to about half of the length of the sub－ genital plate mediobasad，apical tooth and basal teeth


Figs. 20-26. Hongchunia helleri sp.n., paratype of from Cameroun. - 20, male genitalia, ventral view; 21, same, left lateral view; 22. left subgenital plate, dorsal view; 23 , same, dorsolateral view; 24 , left style, dorsal view; 25 , aedeagus and connective, ventrocaudal view; 26, aedeagus, left lateral view. Scales: 0.1 mm .


Figs. 27-35. Hengchunia indica sp.n., holotype ot from India. - 27, male pygophore, left lateral view; 28, left subgenital plate, dorsal view; 29, same, dorsolateral view; 30, subgenital plates, ventral view; 31, left style, dorsal view; 32, aedeagus and connective, ventrocaudal view; 33, aedeagus, left lateral view; 34, same, dorsal view; 35, apex of aedeagus, left dorsocaudal view. Scales: 0.1 mm .
strongest. Style (fig. 24) with short inner and long outer apophysis, the latter continuously tapering to apex, distal margin with about 7 tooth-like projections. Aedeagus (figs. 25, 26) with shaft short, evenly curved dorsad, on both sides at midlength a spinose process which is directed basad; phallotreme apically, exposed to the ventrocaudal side; processes of preatrium long and slender, straight.
Length: ${ }^{\text {on }}: 3.5-3.6 \mathrm{~mm}$, $\frac{7}{}: 3.8 \mathrm{~mm}$.

Remarks. - H. belleri is similar (and probably closely related) to $H$. gaiseri; it can be distinguished mainly by the shape of the subgenital plates which are apically less pointed than in $H$. gaiseri and which bear a more extended toothed area, as well as by the position of the aedeagal processes which arise at midlength of the shaft (not subapically as in H. gaiseri).
Distribution. - The occurrence of $H$. belleri in Zaire and Cameroun may indicate a wider distribution, at least in Central- and West Africa.


Figs. 36-39. Hengchunia pakistanica sp.n., holotype ơ from NE Pakistan. - 36, head and thorax, dorsal view; 37, head, frontal view: 38 , left tegmen; 39, left hind wing. Scales: 0.5 mm .
7. Hengchunia indica sp. n.
(figs. 27-35)
Type material. - Holotype $\delta:$ India, Bihar, Pusa, 14.:1931, T. B. Fletcher (вммnh).

Description. - Habitus, bodily proportions, and colouration as in generic description (see Webb \& Heller 1990).

Male genitalia: Subgenital plate (figs. 27-30) rather stout with distal margin very little convex, almost truncate; toothed area with a triangular portion at margin and an arm-like extension directed mediobasad. Style (fig. 31) comparatively short and stout, devoid of a mediodistal apophysis, subapically with a short lohe-like projection. Aedeagus (figs, 32-35) with shaft evenly curved dorsad; ventrobasad of the apical phallotreme with a gently curved horn-shaped process which arises on a broad base; medially on dorsal side of shaft a longer, apically bifurcate process, and basad of it a shorter single process; processes of
preatrium slender, slightly curved ventrad.
Length: © : 3.0 mm .
Remarks. - Within the genus, H. indica is most similar to H. pakistanica (see below) with which it shares the short and distally truncate subgenital plates and a single horn-shaped aedeagal process basad of the phallotreme on the ventral side. However, it differs considerably from this species in the shape of the toothed area of the subgenital plates and in details of the aedeagus, especially by the presence of paired processes of the preatrium which are entirely missing (probably reduced) in H. pakistanica.

Distribution. - Only known from the BiharRegion in North India.

## 8. Hengchunia pakistanica sp. n. <br> (figs. 36-50)

Type material. - Holotype ot: Pakistan, Gujrat (= Guja-


Figs. 40-50. Hengchunia pakistanica sp.n., holotype ơ from NE Pakistan; Figs. 45, 48: paratype ô from SE Pakistan. - 40, male genitalia, ventrocaudal view; 41 , same, left lateral view; 42 , left subgenital plate, dorsal view; 43 , same, dorsolateral view; 44, 45, left style, dorsal view; 46, aedeagus and connective, ventrocaudal view; 47, 48, aedeagus, left lateral view; 49, same, dorsal view; 50 , apex of aedeagus, left dorsocaudal view. Scales: 0.1 mm .


Figs. 51-54. Hengchunia javana sp.n., holotype of from Java. - 51, head and thorax, dorsal view; 52, head, frontal view; 53, left tegmen; 54, left hind wing. Scales: 0.5 mm .
rat) Nausari, 16.i.1981, C.A. Viraktamath (bMnH). Pararypes: 49 , same data as holotype (UASB, BMNH); $1 \delta^{\star}$, 18. Pakistan, Hyderabad, Patanduru, vii-ix.1980, Bernays \& Woodhead (BMNH).

Description. - Habitus, bodily proportions, and colouration as in generic description (see Webb \& Heller 1990).

Male genitalia: Subgenital plate (figs. 40-43) short, distally truncate, toothed area separated in an apical and a median field of teeth which are connected by a ridge. Style (figs. 44,45 ) distally with a short median edge and a long outer apophysis with a short subapical lobe. Acdeagus (figs. 46-50) with shaft short, slightly curved dorsad; a single long, hornshaped, ventrobasad directed process arising basad of phallotreme on the ventral side; shaft submedially with a pair of latcral processes; preatrium devoid of elongate paired processes.

Length: ठ: $3.0 \mathrm{~mm}, 9: 3.1-3.3 \mathrm{~mm}$.

Remarks. - H. pakistanica can readily be distinguished from all other congeners by the lack of processes of the aedeagal preatrium. The aedeagal shaft with its single horn-shaped process on the ventral side is similar to that of $H$. indica but it differs in other genital characters, as noted above. Slight differences could by found in the genital structures (especially aedeagal spines and styles) between the males from North and South Pakistan but they seem too small to warrant species or even subspecific status.

Distribution. - Only known from the GujratRegion in the North-East and from the Sind-Region in the South-East of Pakistan.

## 9. Hengchunia javana sp. n.

 (figs. 51-62)Type material. - Holotype ô: Indonesia: Java, Bogor, on Imperata, 24.xi.1989, M.R. Wilson (BMNH). - Paratype: Iq, same data as holotype (BMNH).


Figs. 55-63. Hengchunia species. - 55-62. Hengchunia javana sp.n., holotype of from Java; 55, male genitalia, ventrocaudal view; 56, same, left lateral view; 57, left subgenital plate, dorsal view; 58 , same, dorsolateral view; 59 , left style, dorsal view; 60, aedeagus and connective, ventrocaudal view; 61, aedeagus, left lateral view; 62, apex of aedeagus, ventrocaudal view. Scales: 0.1 mm .63 ; H. koshunensis (Matsumura), aedeagus, redrawn and modified from Vilbaste, 1969, Fig. 6 E.

Descriprion. - Habitus, bodily proportions, and colouration as in generic description (see Webb \& Haller 199()).

Male genitalia: Subgenital plate (figs. 55-58) distalfy convex, roothed area devided in an apical and a median portion, both portions with prominent teeth. Srile (fig. 59) with a short inner edge and an elongate outer apophysis with undulate margin. Aedeagus (figs. 60-62) with shaft short, in middle almost rectangularly bent dorsad; phallotreme apically, dorsally on each side flanked by a pair of short, fork-like spinose processes directed dorsocaudad; paired processes of the preatrium conspicuously surpassing the aedeagal shaft, almost straight.

Length: $0: 3.4 \mathrm{~mm} ; ~ ㅇ: 3.6 \mathrm{~mm}$.
Remarks. - Within the genus, H. javana is rather isolated. It can readily be separated from the other congeners by the configuration of the aedeagal shaft with its fork-like, dorsocaudad directed (not reflected) spinose processes at apex.

Distribution. - Only known from Bogor in West Java.

## 10. Hengchunia sp.

We have examined one female from Australia, Queensland, Cairns (ridge at Edge Hill, rainforest), collected on Imperata grass, on 10.vi. 1987 by M. Asche \& H. Hoch (AH). Externally, this specimen resembles $H$. mutus, but its specific identity can only be clarified after a corresponding male becomes available: however, this finding extends the geographic range of the genus Hengchunia to NE Australia.

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## References

Dlabola, J., 1961. Die Zikaden von Zentralasien, Dagestan und Transkaukasien (Homopt. Auchenorrhyncha). Acta Entomologica Musei Nationalis Pragae 34: 241358.

Emeljanov, A. F., 1962. New tribes of leafhoppers of the subfamily Euscelinae (Auchenorrhyncha, Cicadellidae). Entomological Review 41: 236-240.
Emeljanov, A. F., 1972. New Palearctic leafhoppers of the subfamily Deltocephalinae (Homoptera, Cicadellidae). Entomological Review 51: 63-68.
Emeljanov, A. F. \& V. I. Kirilova, 1989. Trends and modes of karyotype evolution in the Cicadina (Homoptera) I. (Cicadelloidea). - Entomologicheskoe Obozrênie 68: 587-603.
Hamilton, K. G. A., 1975. Review of the tribal classification of the leafhopper subfamily Aphrodinae (Deltocephalinae of authors) of the Holarctic region (Rhynchota: Homoptera: Cicadellidae). - Canadian Entomologist 107: 477-498.
Ossiannilsson, F., 1983. The Auchenorrhyncha (Homoptera) of Fennoscandia and Denmark. Part 3: The family Cicadellidae: Deltocephalinae. - Fauna Entomologica Scandinavica 7: 594-979.
Matsumura, S., 1914. Die Jassinen und einige neue Acocephalinen Japans. - Journal of the College of Agriculture, Tohoku Imperial University 5: 165-240.
Rao, R., 1989. Descriptions of some new leafhoppers (Homoptera: Cicadellidae) with notes on some synonymies and imperfectly known species from India. - Hexapoda 1: 59-84.
Remane, R. \& M. Asche, 1980. Neue Zikaden-Taxa aus dem Tribus Paralimnini Distant, 1908, aus dem Mittelmeergebiet (Homoptera Cicadina Cicadelloidea, Cicadellidae). - Marburger Entomologische Publikationen 1 (4): 67-166.
Webb, M. D. \& F. R. Heiler, 1990. The leafhopper genus Pseupalus in the Old World Tropics, with a check-list of the Afrotropical and Oriental Paralimnini (Homoptera: Cicadellidae: Deltocephalinae). - Stuttgarter Beiträge zur Naturkunde, Serie A (Biologie), 452: 1-10.
Vilbaste, J., 1969. On some East-Asiatic leafhoppers described by Professor S. Matsumura (Homoptera: Cicadinea: Iassidae). - Insecta Matsumurana, Supplement 6. 12 pp .

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# A NEW GENUS OF NOMADINE BEES FROM NORTH AFRICA (HYMENOPTERA: APOIDEA, ANTHOPHORIDAE) 


#### Abstract

Baker, D. B., 1994. A new genus of nomadine bees from North Africa (Hymenoptera: Apoidea, Anthoporidae). - Tijdschrift voor Entomologie 137: 155-159, figs. 1-4. [ISSN 00407496]. Published 15 December 1994. Aethammobates prionogaster gen. et sp. n. Anthophoridae: Nomadinae) is described from a unique male collected in Egypt. The new genus is referred to the Holcopasitini and its relationships with other holcopasitine bees are discussed. D.B. Baker, Hope Entomological Collections, University Museum, Oxford ox 1 3pw, U.K. Keywords. - Hymenoptera; Anthophoridae: Holcopasitini; Aethammobates prionogaster, new genus; new species; North Africa.


Among a number of epeoline bees received many years ago from the late Prof. Dr. H. Priesner was a single male of a strange nomadine bee that could not be referred to any described genus. A description and figures were prepared, and the specimen was then set aside in the expectation that further specimens, including possibly the female, would sooner or later come to light. After more than 40 years, this expectation has not been fulfilled, although the locality where the bee was obtained (Gebel el Asfar, near Cairo) was one much visited by both pre-War and more recent local collectors. Since the locality is now being devastated by construction works (C.G. Roche, in litt., 6 September, 1992), making the recovery of further material less likely, and since also the systematics of the Nomadinae have recently come under review by several authors, notably Alexander (1990) and Roig-Alsina (1991), it seems desirable to place the new genus and species on record.

## Aethammobates gen. nov.

Description. - Head in frontal aspect transverse, the eyes rather small, separated by 1.2 times their length, their inner margins convex, subparallel; clypeus short, carinate laterally along epistomal suture, paraocular area adjacent to carina slightly concave and narrowly impunctate; labrum elongate (length to breadth ration $1.44: 1$ ), apically entire, lacking discal tubercles; antennal sockets at mid-level of eyes; interantennal carina divided, forming a $V$-shaped protuberance, the open end of the $V$ upward; facial foveae absent; vertex longer than inter-ocellar distance; preoccipital ridge carinate; malar area rudimentary.

Antennae 13 -segmented, scape long (length to breadth ratio 3 : 1); pedicel free; first flagellar segment long (length to breadth ratio $3: 1$ ), equal to segments $2+3$; segments $2-10$ passing from transverse to quadrate. Mandibles simple, with weak basal protuberance opposed to lateral termination of clypeal carina; only posterior articulation in contact with eye. Stipital comb absent; maxillary palpi 5 -segmented; blade of galea moderately, uniformly sclerotized, acuminate, extreme apex very narrowly rounded.

Mesosoma broad, dorso-ventrally compressed. Pronotum with well developed dorsal surface limited anteriorly by an uninterrupted carina; anterior surface broad, flat, rounded abruptly into collum. Mesoscutum little convex; parapsidal lines well marked, about as long as tegula; axillae not protuberant; scutellum extending laterad slightly beyond axillae, with distinct dorsal and posterior vertical surfaces separated by a weak, irregular (interrupted by punctation) carina, slightly overhanging subvertical, coplanar surfaces of metanotum and propodeum; omaulus continuous across mesepisternum, forming prominent angles on either side of a median, semicircular inflexion; mesepisternum anterior to intermediate coxae strongly transverse, transversely concave. Metanotum with prominent lateral lobes. Wings (fig. 4) short, forewing about 0.6 body length. Forewing bare basally, becoming pubescent apically, especially beyond cells; marginal cell broad, rather broadly rounded at apex, apex separated from wing-tip by less than cell length; distal abscissa of Rs absent; two submarginal cells, the second much shorter than the first, the first receiving 1 m -cu near its apex, the second 2 m -cu beyond the middle. Hindwing with jugal lobe
very short: sceond abscissa of $\mathrm{M}+\mathrm{Cu}$ abour 1,5 times longer than cu-v, not quite half as long as M . Legs of normal proportions. Anterior coxae proximate, transverse, the trochanters widely separated; tarsus sub-bipectinate, basitarsus and segments two to four apicalIy each with a few strong setae on either side; basitarsus flattened and slightly expanded apicad. Intermediate coxae widely separated, short, much shorter than distance from their summits to posterior wing bases; femur broad; tibia without anterior setose area, coarsely spiculate. Posterior coxae large, broad, inwardly weakly, externally strongly carinate; tibiae coarsely spiculate. Arolia present, normally developed.

Mctasoma broad, depressed, greatest width at apex of segment 2. Tergum 1 with vertical anterior and horizontal dorsal surfaces separated by a deeply arcuate carina, in dorsal aspect with prominent lateral angles; marginal areas of terga abruptly and progressivcly more strongly depressed caudad, the margins of the discal areas passing from subserrate on tergum 1 to strongly denticulate on terga 4-6; pygidial plate strong, rostriform, rounded apically, not constricted basally, considerably exceeding apical margin of segment; sternum 6 laterally dentate.

Vestiture generally squamiform, on the terga forming basal and marginal fasciae, not maculae; simple, erect hairs largely confined to mandibles and disc of labrum; anterior angles of malar areas not penicillate; eyes bare; sterna without subapical fimbriae of modified hairs, but sterna 5 and 6 mesially with fine, dense, erect pubescence, on 6 , towards the lateral teeth, becoming longer and directed mesad.

Integument black with extensive pale areas, the appendages and metasoma predominantly pale; moderately strongly, moderately densely to subreticulately punctate, microsculpture not evident at 37,53 .

Distribution. - N.E. Africa.
Host. - Not known.
Type species. - Aethammobates prionogaster, sp. n.
Derivatio nominis. - Gr. $\alpha \eta \theta \eta s$, strange, + Ammobates, nom. propr. Hymenoptera

Aerhammobates is separated from other nomadines by a varicty of autapomorphies, but for purposes of recognition its general habitus and the modifications of the pronotum and first metasomal tergum are immediately diagnostic (see Discussion).

[^12]
## Description

Male. - Structural characters: See generic descrip-
tion. $\mathrm{POL}=$ OOL. Hamuli 8 . Length 7.5 mm , forewing 4.75 mm .

Vestiture. The face, dorsal surface of pronotum, peripheries of mesoscutum and scutellum, mesepisterna, and outer ventral surfaces of posterior coxae, all more or less densely clothed with white squamiform or subsquamiform pubescence, that on the pronotum apparently particularly dense (matted in type); elsewhere on head and mesosoma sparser, the individual hairs becoming longer and less squamiform. Terga with broader, white, medially emarginate, laterally expanded, basal fasciae and narrow, entire, apical fasciae, filling the depressed marginal areas, of squamiform pubescence; tergum 5 nearly completely covered. Disc of pygidial plate with similar pubescence. Sterna 2-4 with weak submarginal fasciae, sterna 5 and 6 medially with fine, dense, erect pubescence (as noted in generic description - possibly homologous with pre-genital brush in Nomada).

Integument of head and mesosoma black with extensive pale areas. Clypeus, anterior extremities of paraocular areas, malar areas, labrum, small maculae adjacent to summits of eyes, larger maculae on genal areas adjacent to mandibles, extending narrowly upwards almost to summits of eyes, labrum and all head appendages (except the mandibles apically) castaneous, the flagellum darker. Lateral lobes of pronotum, axillae, tegulae, sclerites of wing bases, wing veins (the costa darker), and legs (the coxae basally darker) castaneous. Metasoma castaneous, terga 2-5 discally progressively, irregularly (possibly the result of post-mortem decomposition) darker; sterna 2 and 3 discally largely, 4 and 5 almost wholly, dark. Labrum smooth, moderately strongly, irrergularly punctate; anterior face of pronotum smooth, impunctate; mesoscutum and scutellum smooth, moderately densely, simply punctate; terga smooth, medially densely, laterally reticulately, punctate, the punctation coarsest on tergum 2, on tergum 1 about equal to that of mesoscutum, on terga 3-5 progressively finer. Margins of pygidial plate reflexed.

Derivatio nominis. - Gr. $\pi \rho \imath \sigma \nu-\omega \prime \delta \eta s / \omega \tau o ́ s$, serrated, + gaster .

## Discussion

Both Alexander (1990) and Roig-Alsina (1991) have recently published cladistic analyses of nomadine bees. Roig-Alsina's analysis was based primarily (15 out of 22 characters) on characters exclusive to the female, and is consequently of limited use in the present context. Alexander's analyses were based on (1) larval characters only, (2) on adult characters of those genera where larvae were known, (3) on both larval and adult characters of these genera, and (4) on the adult characters of all genera known to him. It is this last analysis, (4), that is relevant to the placement

 and 5

Characters

|  | 1 | 2 | 4 | 5 | a | 6 | 7 | 8 | 9 | 10 | 11 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Holionusites | 1 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 |
| Silmiedeknechria | 1 | 0 | 0 | 0 | , | 0 | 0 | 0 | 0 | 0 | 0 |
| Acthammobates | 0 | 0 | 1 | 0 | 0 | 0 | $-{ }^{2}$ | 0 | 1 | 0 | 2 |
|  | 12 | 13 | 14 | 15 | 16 | 17 | b | c | 18 | d | 19 |
| Holocopasies | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 14 | 1 | 0 |
| Schmiedelnechriar | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | $1{ }^{4}$ | 1 | 0 |
| Aeshammobares | 1 | 1 | 0 | 0 | 0 | $0^{3}$ | 0 | 1 | 1 | 1 | 1 |
|  | 20 | 21 | e | f | $\mathrm{g}^{5}$ | 22 | h | 23 | 24 | 29 | 32 |
| Holocopasites | 1 | 0 | 1 | 0 | 0 | 0 | 2 | 1 | 0 | 1 | 0 |
| Sthmiedeknechria | 1 | 0 | 1 | 0 | 0 | 0 | 2 | 1 | 0 | 1 | 0 |
| Aeshammobates | 1 | 0 | 1 | ? | 0 | 0 | 2 | $1{ }^{6}$ | 0 | 0 | 07 |

Notes on matrix

1) character a: Antennal sockets below mid-level of eyes, but not far below; no code. - (2) character 7: 'Length/width of first flagellomere' = 3: no code. - (3) character 17: No 'patches of appressed squamiform setae': Alexander presumably refers to diactue metanomal maculations of Holocopasites and Schmiedeknechtia. - (4) character 18: Incorrectly coded '0' by Alexander: : c:n int r-m of forewing is absent (two submarginal cells). - (5) character g: 'arising from' in Alexander's Appendix 5 presumwh: mean "riving from (i.c.. in lateral aspects bends upward from ventral plane of mesosoma immediately before intermediate coxae). - (6) character 23: a weak, short, dorso-basal carina present on posterior coxa. - (7) character 32: Coded ' 0 ' since apex narrowly rounded, subacuminate, but galea is more than weakly sclerotized.
of Aethammobates. Alexander's data matrix (his Appendix 6) employed 46 characters, of which six applied solely to the female, leaving 40 in which Aethammobates could potentially be compared. In practice, since the holotype and only known specimen of Aethammobates has not been dissected (cf. Roig-Alsina's comment, 1991: 25, on the pre-episrernal sulcus in Rhopalolemma), comparison has been limited to 33 characters. These have been covered in the description given above, and are now given in coded form, following Alexander's Appendices 3 and 5, as an extension of his data matrix, in Table 1. This table reproduces Alexander's codings, but substitutes observed values for six characters marked with a quae$r e$ in his matrix.

It should be noted that Alexander's codings, here and elsewhere, cannot always be accepted because he did not see certain described and undescribed taxa that would have necessitated different coding or the adoprion of additional codes; however, a complete revision of his data matrix, which might well result in significant changes in his cladogram, is beyond the scope of the present paper. Important characters that should be taken into account in future work on the phylogeny and classification of the Nomadinae include the presence of facial foveae in some Nomada (a group of, chicfly, Panurgus parasites); the development of an anal truncation, with associated modifications of the apical sterna, among the pasitines (Bischoff 1923: 585. Analstutz); and modifications of
the antenna in such genera as Morgania and Pasitomachthes, where the scape and pedicel may be coalescent, forming a single functional unit (Baker 1971: 7, footnote).
Inspection of the extended matrix confirms a priori impressions that Aethammobates shares a preponderance of characters with Holcopasites and Schmiedeknechtia and must be placed in the Holcopasitini.

While obviously most nearly related to Holcopasites [Nearctic, revised by Linsley (1943, as Neopasites) and by Hurd and Linsley (1972)] and Schmiedeknechtia [western palaearctic, revised by Popov (1933)], and sharing such derived characters as the lateral processes of the metanotum and certain venational details, Aethammobates is not close to either. It differs from both in, among other characters:

1. The dorso-ventrally compressed form of mesosoma and metasoma.
2. The mid-dorsally exposed pronotum with sharply differentiated dorsal, densely pubescent and anterior, glabrous surfaces. (In lateral aspect, the pronotum resembles that of Pae (Sphecoidea) as illustrated by Bohart \& Menke (1976: 15, fig. 3A),] In Holcopasites and Schmiedeknechtia the pronotum is, as in most other bees, recessed beneath the mesoscutum with only its lateral lobes and, when the head is deflexed, the collum conspicuous in dorsal aspect.
3. The $V$-shaped interantennal carina.
4. The strongly expanded intermediate femora.
5. The pronounced basal truncation of the mesosoma.
6. The form of the first metasomal tergum, which is broadly, deeply emarginate, with prominent ante-ro-lateral angles (see fig. 3) and the anterior and dorsal surfaces sharply, rectangularly separated.
7. The presence of basal and marginal tergal fasciae and the absence of of discrete spot- or bar-like markings.
Of these, characters 1-6 appear as autapomorphies. Aethammobates differs further, from male Holcopasites, in the 13-segmented antennae, and from Schmiedeknechtia in the form of the pygidial plate, which is not narrowed basad; in not having the inner orbits convergent below; in not having the anterior mandibular articulations in contact with the eyes; and in the venation: in Schmiedeknechtia either both 1 m cu and $2 \mathrm{~m}-\mathrm{cu}$ are received by second Rs, or (Cyrtopasites, a single species with convex mesoscutum and fine and sparse punctation) 1 m -cu is interstitial with or slightly precedes 1 r -m.

## Biology

It is interesting to speculate on the possible host of Aethammobates. Known hosts of Holcopasites and Schmiedeknechtia are panurgines. Camptopoeum and Meliturgula are possible candidates, but the former. which is normally abundant where it does occur, and which appears not to have been recorded from the Cairo area (it is not represented in recent, extensive Egyptian collections), has species of Parammobatodes as its regular parasites. The somewhat flattened shape of Aethammobates suggests some correlation with
habits or host. Both sexes of Meliturgula have a conspicuously broad or flattened metasoma: regional species are few in number and rare in collections; nothing is known of their biology.

## References

Alexander, B., 1990. A cladistic analysis of the nomadine bees (Hymenoptera: Apoidea). - Systematic Entomology 15: 121-152.
Baker, D. B., 1971. A new Pasitomachthes from Rhodesia (Hymenoptera, Apoidea). - Novos Taxa entomológicos, no. 98: 1-8.
Bischoff, H., 1923. Zur Kenntnis afrikanischer Schmarotzerbienen. - Deutsche entomologische Zeitschrift 1923: 585-603.
Bohart, R. M. \& A. S. Menke, 1976. Sphecid wasps of the world: a generic revision: ix +695 pp. - University of California Press, Berkeley.
Hurd, P. D. \& E. G. Linsley, 1972. Parasitic bees of the genus Holcopasites Ashmead (Hymenoptera: Apoidea). Smithsonian Contributions to Zoology, no. 114: 1-41.
Linsley, E. G., 1943. A revision of the genus Neopasites (Hymenoptera: Nomadidae). - Transactions of the American Entomological Society 69:119-140.
Popov, V. B., 1933. Notes on the parasitic bees allied to the genus Biastes Panz. (Hymenoptera, Nomadidae). Trudy zoologicheskogo Instituta Akademy Nauk SSSR 2: 51-75.
Roig-Alsina, A., 1991. Cladistic analysis of the Nomadinae s.str. with description of a new genus (Hymenoptera: Anthophoridae). - Journal of the Kansas Entomological Society 64: 23-37.

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# FOUR SPECIES ADDED TO THE BAETURIA NASUTA GROUP, WITH NOTES ON TAXONOMY AND BIOGEOGRAPHY (HOMOPTERA, TIBICINIDAE) 


#### Abstract

Boer, A. J. de, 1994. Four species added to the Baeturia nasuta group, with notes on taxonomy and biogeography (Homoptera, Tibicinidae). - Tijdschrift voor Entomologie 137: 161-172, figs. 1-39. [ISSN 0040-7496]. Published 15 December 1994. A new concept is proposed for the nasuta group of the cicada genus Baeturia Stål, 1866. Four species are added to this group, bringing the total number of its species to twelve. One of these species (B. guttulipennis Blöte, 1960) is redescribed and three (B. gibberosa, B. splendida, and B. retracta) are described as new. Implications for the biogeographic pattern of the group are, that the nasuta group can no longer be seen as a typical central New Guinea group; most of the species can apparently be found in northern New Guinea. A.J. de Boer, Institute for Systematics and Population Biology (Zoölogisch Museum), University of Amsterdam, P.O. Box 94766,1090 GT Amsterdam, The Netherlands Key words. - Baeturia, nasuta group, taxonomy, biogeography, New Guinea.


The nasuta group was recognized as a monophyletic group consisting of eight New Guinean species of the genus Baeturia Stål (de Boer 1982). The monophyly of that group is based on two presumed apomorphies: a narrow and angularly swollen postclypeus and four sclerotized ridges on the tymbals. $B$. guttulipennis Blöte was not included in the nasuta group, since it does not share these apomorphies. However, now that most of the remaining species of Baeturia have been revised and placed in monophyletic species groups, it appears that B. guttulipennis is more closely related to the nasuta group than to any other species group. It is proposed here to change the concept of the nasuta group and to include B. guttulipennis in that group. Furthermore, three new species can be added to the nasuta group, these species do share the original apomorphies of that group.

Species of the nasuta group, as previously defined, were mainly distributed in the central mountain ranges of New Guinea. Its distribution pattern has been compared to that of the cicada genus Cosmopsaltria Stål, which also centres in the montane parts of central New Guinea (de Boer 1982; Duffels 1986; Duffels \& de Boer 1990) and it was suggested that the similarities between these patterns might find a common cause in the paleogeology of New Guinea. The present additions to the nasuta group greatly alter its general distribution pattern and necessitate a reconsideration of its paleogeographic history.

## Material and methods

The material examined for this study is deposited in the following collections:
amnH: American Museum of Natural History, New York; Ams: Australian Museum, Sydney; Bmnh: Natural History Museum (formerly: British Museum (Natural History)), London; BPBM: Bernice P. Bishop Museum, Honolulu; Moul: Personal collection Mr M.S. Moulds, Sydney; RmnH: Nationaal Natuur historisch Museum (formerly: Rijksmuseum van Natuurlijke Historie), Leiden; SMN: Staatliches Museum für Naturkunde, Stuttgart; ZMA: Institute for Systematics and Population Biology (Zoölogisch Museum), Amsterdam.

The following geographical sources have been used: Atlas van tropisch Nederland (1938), The Times Atlas of the World (1968) and the 'List of New Guinea localities' published by the Bishop Museum (1966).

After overnight softening, male genitalia were examined by pulling out the pygofer with a sharp needie inserted between pygofer and 8th abdominal segment. The aedeagus was pulled out at the same time, by inserting the needle between the claspers. Measurements are based on all available specimens.

## Phylogeny

Baeturia belongs to a larger group of genera, the

Bucruria and related genera complex', as defined earlier (de Boer 1990). A phylogeneric analysis of this genus complex, with the oriental Prasiini as defined by De Jong (1985) as ourgroup, is in preparation. The ensuing phylogeneric discussions are based on the preliminary results of that analysis.

## The monophyly of the nasuta group

The monophyly of the nasuta group is based on three presumed apomorphies. Males of the nasuta group are easily identified by the almost transparent lateral sides of the abdomen, usually restricted to segments 3-7. Only B. marmorata does not show this character. A similar transparency of the male abdomen was found in a monophyletic group of three species of the genus Gymnotympana (unpublished), but not in other related taxa. A laterally transparent abdomen is regarded as apomorphous for the nasuta group ( 1 in fig. 1b) and the similar transparency in three Gymnotympana species must be explained by parallelism.
A strongly curved aedeagus, with lateral lobes at its basal curve, is regarded apomorphous for the genus Bacturia. The aedeagus of the species of the nasuta


[^13]group differs from that of other species by a longer and more slender apical part, distally of the lateral lobes (fig. 11), which is regarded apomorphous (2 in fig. 1b). A very similarly elongate, but stouter, aedeagus was found in B. loriae and B. pigrami of the loriae group (de Boer 1994a). The elongation of the aedeagus in these two groups cannot be regarded as homologous in the most parsimonious solution.

The male pygofer is characterized by a strongly bent and dorsally flattened caudodorsal beak. This shape of the caudodorsal beak is shared by all species and regarded apomorphous for the nasuta group ( 3 in fig. 1b). In other species, which have a similarly strongly bent beak, the beak is convexly curved dorsally (comp. the viridis group, de Boer 1992).

## The phylogenetic position of the nasuta group

Baeturia is a monophyletic genus, for which a frontally rounded and protruding postclypeus, a triangleshaped medial thorn on the fore femur, and lobate lateral crests on the aedeagus are the presumed apomorphies (1, 2, 3 in fig. 1a). Baeturia can be subdivided into seven monophyletic species groups: the bloetei group (de Boer 1989), the conviva group (de Boer 1986), the exhausta group (de Boer 1994b) the guttulinervis group (de Boer 1994c), the loriae group (de Boer 1994a), the viridis group (de Boer 1992), and the nasuta group (de Boer 1982).

The nasuta group is presumed to form a monophyletic group with the conviva, guttulinervis, and viridis groups. The large, often conically-shaped, distinctly posteriorly projecting protuberance on the lateral lobe of pygofer is regarded synapomorphous for these four species groups together ( 4 in fig. 1a). Furthermore, these four groups share a very stout and strongly bent caudodorsal beak. The occurrence of a similarly bent caudodorsal beak in several Melanesian species of the bloetei group must be explained by parallel development.

The nasuta and viridis groups are sister groups, sharing an angularly bent caudodorsal beak as supposed synapomorphy ( 5 in fig. 1a). In the species of these two groups the dorsal margin of the pygofer is not continuously rounded with the dorsal margin of the caudodorsal beak. Furthermore, these species share a generally truncate, but sometimes bluntly rounded or bicuspidate, caudodorsal beak. Species of other groups of Baeturia generally have a more narrowly rounded or pointed beak, while a truncate beak only sporadically occurs. The conviva and guttulinervis groups are sister groups based on a shared apical swelling of the clasper (this swelling is almost globular in the guttulinervis group) and a dorsally strongly rounded pygofer ( 6 in fig. 1a), but the relative position of the remaining species groups of Baeturia is not clear.

## Ingroup phylogeny

The species of the nasuta group are very similar in general appearance. The species differ mainly in the shape of the claspers and some clasper characters indicate phylogenetic relationships. Most other differences observed are either species specific characters, or characters that also occur outside the nasuta group. It is not possible to construct a fully resolved and unambiguous cladogram for the group, based on the limited number of characters now available. Nevertheless, some remarks on ingroup phylogeny can be made (see fig. 1b).
B. guttulipennis presumably forms the sister taxon of all other species of the nasuta group; these other species share four tymbal ridges (three in B. parva) and a more angularly swollen postclypeus, which are the original apomorphies for the nasuta group ( 4,5 in fig. 1b). B. guttulipennis and B. nasuta however, share similar brown markings on the tegmina and an apically rounded male caudodorsal beak, and could be sister species. $B$. nasuta and $B$. intermedia share an almost identical clasper and are presumably either sister species or part of a paraphyletic subgroup.
B. arabuensis, B. bipunctata, B. laminifer, B. mamillata, B. retracta and B. gibberosa share a distinct dorsal protrusion on the clasper as presumed synapomorphy ( 6 in fig. 1b). That B. nasuta and B. intermedia should also be included in this group, as was suggested before (de Boer 1982) seems unlikely, since the weakly developed dorsal crest of the claspers of $B$. nasuta and B. intermedia is very similar to that found in many other Baeturia species. The remaining species of the nasuta group share a dorsally smoothly rounded clasper.
B. arabuensis and B. mamillata share a posteriorly directed and almost finger-shaped protrusion, and are possibly sister species, B. bipunctata, B. gibberosa, B. laminifer and B. retracta have a much broader dorsal protrusion on the clasper, which is regarded as synapomorphous (8 in fig. 1b). B. laminifer and B. retracta share a laminiform and square-shaped dorsal protrusion on the clasper as presumed synapomorphy ( 9 in fig. 1b). B. bipunctata and B. gibberosa have a more triangular, hump-shaped, protrusion; these are either sister species or part of a paraphyletic subgroup.

Females of most of the species that have a dorsal protrusion on the clasper, as far as they could be identified, generally have a truncate caudodorsal beak, which character is unique for these species and presumably apomorphous at that level ( 7 in fig. 1b). Only the females that presumably belong to B. arabuensis and the females of $B$. gibberosa have a pointed beak as those in B. guttulipennis, B. nasuta, and $B$. parva. The female of $B$. splendida has a bluntly rounded, almost truncate, caudodorsal beak.

## BIOGEOGRAPHY

The nasuta group is distributed in New Guinea and some adjacent islands, some species reach to the Bismarck Archipelago or the Admiralty Islands.

Study of the New Guinean cicadas indicates that the various genera or monophyletic species groups show different patterns of distribution. The genera seem concentrated with most, and often endemic, species in different parts of the island. These parts of New Guinea, thus recognized as areas of endemism (Duffels \& de Boer 1990), largely coincide with frag-


Fig. 2. Distribution of Baeturia gibberosa, B. guttulipennis, B. retracta, and B. splendida.
ments of a historic island arc, that collided at different times and in various places with the northern craton of the Australian continent, to form present day New Guinea (for geological reviews see Daly et al. 1991; Rangin et al. 1990a, 1990b; Pigram \& Davies 1987). It is supposed that the various genera of New Guinean cicadas evolved on isolated island arc fragments previous to their collision with the Australian continent (de Boer 1994c).

The nasuta group was regarded as a typical central New Guinea group (de Boer 1982) and, as such, has been compared to the genus Cosmopsaltria Stål (Duffels 1986; Duffels \& de Boer 1990). Cosmopsaltria and the B. nasuta group both appeared to be concentrated in the central mountain ranges of New Guinea. The distributions of B. arabuensis, B. mamillata and B. nasuta, from the Wissel Lakes to well into the Papuan Peninsula, are indeed very similar to that of many species of Cosmopsaltria (Duffels 1983). A similar historical explanation for the distribution patterns found in these two groups seemed required. Such an explanation can be found in supposing that both Cosmopsaltria and the nasuta group evolved on the Sepic Arc terrane as defined by Pigram \& Davies (1987), which is the first of the island arc fragments that collided with the Australian continent, and which developed into the central mountain ranges of New Guinea. However, the phylogenetic level of the groups (Cosmopsaltria as sister group of Diceropyga, Rhadinopyga, Aceropyga, and Moana (Duffels 1986, 1993), and the nasuta group as a subgroup of Baeturia) possibly indicates a considerable age difference, which contradicts to a contemporal evolution on the Sepic Arc fragment.

With the four species here added to the nasuta group, this group as a whole no longer has a typical central New Guinea distribution. In fact, the number of species that occurs in northern New Guinea about equals that of central New Guinea. Only four species (B. arabuensis, B. intermedia, B. mamillata, and B. nasuta) can be said to have a central mountain range distribution, though two (B. mamillata, and B. nasuta) also occur in the Papuan Peninsula. The distributions of $B$. laminifer and B. retracta are very similar; they are recorded from southern New Guinea (unpublished data for laminifer), the most eastern part of northern New Guinea, and the Papuan Peninsula. B. laminifer is recently recorded from New Britain (unpublished) and B. retracta is recorded from the Admiralty Islands. These two species are absent from the western parts of the central mountain ranges. $B$. gibberosa is possibly restricted to the Papuan Peninsula and Normanby Island. B. splendida and, possibly, B. guttulipennis are endemic to northern New Guin (it is not certain that the female allotted to the latte1 pecies really belongs to B. guttulipennis).

Also B. parva is distributed in northern New Guinea, but its distribution extends to the Vogelkop Peninsula, Misool, Biak, the Bismarck Archipelago, and Admiralty Islands. The two remaining species ( $B$. bipunctata and B. marmorata), both from Araucaria camp and Rattan camp on the northern slopes of the central mountains of western New Guinea, should presumably be regarded more as northern, than as central New Guinean: several other species groups with a predominantly northern New Guinea distribution (e.g. the viridis group and the guttulinervis group) also have species in that area.
The distribution pattern of the nasuta group in its present concept is more in agreement with that of other groups of Baeturia, most of which seem to concentrate in northern New Guinea, while similarities with Cosmopsaltria have become less obvious and a paleobiogeographical comparison with that genus seems no longer relevant.

## Taxonomy

## Description of the nasuta group

Species of the nasuta group are all very small sized. The specimens are shorter than 2.5 cm and generally do not exceed 2.0 cm . Males are easily recognized by their laterally transparent abdomen (this character is absent in B. marmorata). Body generally yellowish brown or reddish brown coloured and covered with many irregularly shaped brown speckles. Females generally darker coloured and more densely speckled (B. splendida unspeckled with greenish head and thorax).

Head (fig. 3a): Narrow, 2.3-2.8 x as wide as long. Vertex narrower than in most other Baeturia species, ocelli close together. Distance between lateral ocelli less than 2 x as wide as, and sometimes narrower than, frontal ocellus, and generally about as long (0.7-1.2 x ) as distance between lateral ocellus and eye. Vertex $1.6-2.5 \mathrm{x}$ as wide as postclypeus. Postclypeus in dorsal view narrow (1.1-2.2 x as wide as long) and triangularly protruding beyond vertex lobes, though with rounded anterior margin. Postclypeus distinctly swollen ventrally, its anterior margin (lateral view) often angularly bent (fig. 3b).
Legs: Fore femur (Fig. 6) with row of three erect and sharply pointed spines, diminishing in length towards tibia. The most distal spine often bicuspidate, forming a very small 4th spine. Most proximal spine slightly shorter than distance to middle spine. Middle spine triangular, very broad at its base.
Tegmina and wings: Hyaline, though tegmina sometimes with darkly suffused spots along veins. Tegmina with 8 , wings with 6 apical areas. Hyaline border along hind margin of tegmen distinct, though narrow. This border slightly broader in wing. Veins
ochraceous, sometimes tinged with red, and set with many long setae.

Tymbals: With 3-5, generally 4, parallel sclerotized ridges from dorsal to ventral tymbal margin. Often traces of red between dorsal parts of tymbal ridges.

Opercula: Male operculum fairly small, its distal part not covering tymbal cavity in ventral view and nearly always shorter than meracanthus, generally not reaching to anterior margin of 2 nd abdominal segment. Operculum in ventral view reaching to under, or medially of, meracanthus. Female operculum smaller than that of male, sickle-shaped and erect.

Abdomen: Male abdomen conspicuous by almost transparent sides, its dorsal and distal parts darkened and brown speckled, ventral parts unspeckled. Lateroventral row of dark spots generally very distinct. Hind margins of abdominal segments often reddish. First and second sternites adjacent. First tergite very short and often partly hidden under metanotum (in B. arabuensis distinctly longer than in other species). Second tergite medially with almost straight anterior margin. Auditory capules well developed, globularly swollen. Female abdomen more robust than that of male, darker coloured, and more densely speckled. Ovipositor sheaths sometimes reaching beyond apex of caudodorsal beak. Female caudodorsal beak often truncate at apex.

Male genitalia: Caudodorsal beak strongly bent, making an almost right angle with dorsal margin of pygofer. Dorsal margin of pygofer not continuously rounded into dorsal margin of beak, and weakly concave near apex of beak. Caudodorsal beak very broad and straight, dorsally flattened, and with bluntly rounded or truncate, often weakly concave, apex. Lateral lobe of pygofer with strongly swollen, often conically shaped, protuberance, projecting posteriorly beyond margin of pygofer. Claspers fairly long, parallel, and generally directed posteriad. Apical part of clasper hardly bent down, with large and sharply edged ventral hollow. Aedeagus S-curved with lateral lobes at basal curve. Apical part of aedeagus extremely long and slender (compared to other species of Baeturia). Aedeagal pore narrow and oval.

## Baeturia guttulipennis Blöte

(figs. 2-13)
Baeturia guttulipennis Blöte, 1960: 69, fig. 15; de Boer, 1994c.

Material examined: Bewani R. territ., $1200 \mathrm{~m}, 1939$, W. Stüber, ơ holotype, RMNH; Bivak 39A, Star Range, 1150 m, 6.vii.1959, 1 \& Baeturia guttulipennis? det. H.C. Blöte, RMNH.
B. guttulipennis is easily separated from other spe-


Figs. 3-6. Baeturia guttulipennis Blöte, 1960. - 3a, head in dorsal view, holotype; 3b, head in lateral view, holotype; 4, female caudodorsal beak in dorsal view; 5, female genital segment in lateral view; 6, fore femur, holotype.
cies of the nasuta group by the dark spots along the veins of tegmina and by its five tymbal ridges. Similar dark spots in tegmina were found in several other species of Baeturia, e.g. B. guttulinervis Blöte (de Boer 1994c), B. pigrami De Boer and B. silveri De Boer (de Boer 1994a), and, though less distinct, in B. nasuta Blöte (de Boer 1982).

## Description

Body of male light ochraceous brown and weakly speckled with brown. Female darker, red-brown and more densely speckled. Female head and thorax larger than in male. Tegmina of males 1.2 x as long as body length, of females 1.5 x . Male abdomen 1.3 x as long as head and thorax, of females 1.1 x .

Head (fig. 3a): Ochraceous, brown-stained mainly on vertex lobes. Postclypeus bluntly rounded anteriorly, weakly protruding in male, more strongly in female. Postclypeus 1.4-2.0 x as wide as long. Postclypeus in lateral view (fig. 3b) distinctly swollen, anterior margin forming an almost right angle at $1 / 2$ its length, and concave to anteclypeus.

Thorax: Pronotum ochraceous with dark speckling predominantly in medial band and on medial parts of


Blote. 19(x): - pegofer in lateral view, holotype; 8, clasper, holotype; 9, male operculum, holotype: 10. make caudodorsal beak in dorsal view, holotype; 11, aedeagus from behind, holotype; 12, aedeagus in lateral : ( mate operculum: 18, head in lateral view; 19, female operculum; 20, female caudodorsal $\therefore \because$.u : in lateral vew: 22. aedeagus from behind. Lettering: $\mathrm{bp}=\mathrm{basal}$ part of operculum; $\mathrm{cb}=\mathrm{cau}-$ and operulum: $\mathrm{di}==$ distal margin of pygofer; $\mathrm{dmc}=$ distomedial corner of operculum; do $=$ dorsal margin of pygnfer: $\mathrm{dp}=$ distal part of operculum; he = clasper heel; ho = clasper hollow; $1=$ lateral margin of operculum: $m=$ medial margin of operculum; $p=$ protuberance on lateral lobe of pygofer; ve $=$ ventral margin of pygofer.
pronotal lobes, unspeckled in and along its fissures. Mesonotum greenish brown, weakly brown speckled in male, but densely speckled in female. Male with two brown spots in front of cruciform elevation.

Legs: Ochraceous, often reddish towards tarsi, and brown speckled. Fore femur (Fig. 6) with row of three or four sharply pointed and erect spines.

Tegmina and wings: Tegmina with dark brown markings concentrated along veins. Wings hyaline. Veins ochraceous brown, densely set with setae.

Tymbal organ: Five darkly sclerotized parallel ridges spanning the tymbal from dorsal to ventral margin. Four intercalary ridges form a lateral band across tymbal.

Opercula: Male operculum (fig. 9) very small and not covering tymbal cavity. Distal part of operculum erect and reaching to about $2 / 3$ the length of meracanthus, but not reaching to abdomen. Lateral margin of distal part short, concavely bent into crest around distolateral corner of basal part, and convexly bent into long and almost straight distal margin. Distomedial corner rounded, medial margin straight. Female operculum (fig. 13) almost as large as that of male, broadly sickle-shaped and curved to abdomen, with almost continuously rounded distal margin.

Abdomen: Male abdomen weakly inflated, yellow brown and brown speckled, though ventrally unspeckled. Tergites 3-6 almost transparent laterally. A distinct lateroventral row of brown spots on tergites 3-7. First tergite very short and medially partly hidden under metanotum. Female abdomen more robust than that of male, dark reddish brown, and densely speckled. Lateroventral spots on tergites 3-7 almost lost in speckling. Ovipositor sheaths not reaching to apex of caudodorsal beak (fig. 5). Female caudodorsal beak in dorsal view (fig. 4) sharply pointed at apex.

Male genitalia: Pygofer in lateral view as in fig. 7. Dorsal margin weakly convex, strongly bent into almost straight caudodorsal beak. Distal margin concave, gradually bent into straight margin of beak. Protuberance on lateral lobe of pygofer rounded and conically projecting posteriad. Pygofer lobe forming an angular corner just above this protuberance. Ventral margin almost straight, but angularly bent just under lateral protuberance. Ventral margins converge to sharp angle at base of pygofer opening. Caudodorsal beak in dorsal view (fig. 7) rounded at apex. Clasper (fig. 8) almost straight and directed posteriad, with long and slender, dorsally smoothly rounded, apical part. Basal part of clasper with weakly developed clasper heel. Clasper hollow fairly long. Aedeagus (fig. 12) with very long and slender apical part and well developed lateral lobes. Aedeagus from behind (fig. 11) very slender, aedeagal pore narrow, oval shaped and truncate at apex.

Measurements: Body length $\delta: 18.6 \mathrm{~mm}$, $.7: 18.2$ mm ; tegmen length $\mathbf{\delta}^{2}: 22.9 \mathrm{~mm}, \quad:: 28.0 \mathrm{~mm}$; head length $\delta: 1.4 \mathrm{~mm}, \quad,: 1.8 \mathrm{~mm}$; pronotum length ot: $2.4 \mathrm{~mm}, ~ ㅇ: 2.8 \mathrm{~mm}$; mesonotum length ${ }^{6}: 3.8 \mathrm{~mm}$, 우: 4.8 mm ; head width $\delta$ ㅇ: $3.7 \mathrm{~mm}, \quad$ ㅇ: $: 4.3 \mathrm{~mm}$; width of pronotal collar $\delta^{6}: 5.3 \mathrm{~mm}, \quad:=6.5 \mathrm{~mm}$.

Distribution (fig. 2). - The male holotype of $B$. guttulipennis comes from Bewani in northeastern Irian Jaya; a female that possibly belongs to the species comes from the Star Range in central eastern Irian Jaya.

## Baeturia splendida sp. n.

(figs. 2, 14-22)
Holotype male: 'NEW GUiNEA NE / Torricelli Mts. / Mokai Vill. $750 \mathrm{~m} /$ xii-16-31-1958' (print); 'W.W. Brandt / Collector / Bishop' [print], BPBM. - Paratypes: same data as holotype 5 우, BPBM ; same data but 8 -15.xii. 1958, 3 여; 123.i.1959, 1 ㅇ, all BPBM.

Only one male and several females are known of this species. B. splendida is easily recognized by its greenish colour, with a pattern of brown markings. Due to this colour pattern the females too are easily identified. The colour pattern is unique within the nasuta group.

## Description

Head and thorax olive green, abdomen yellowish brown. Tegmina of male 1.2 x as long as body length, of females 1.3-1.4 x . Male abdomen 1.2 x as long as head and thorax, of females 0.9-1.1 x .

Head: Olive-green. Vertex lobe with dark brown spot between eye and lateral ocellus. Postclypeus distinctly protruding, bluntly rounded anteriorly, and 1.1-1.8 x as wide as long. Postclypeus in lateral view (fig. 18) distinctly and angularly swollen, anteriorly almost globularly inflated.

Thorax: Pronotum greenish and unmarked. Mesonotum green with two large semi-oval paramedian brown spots at pronotal margin, reaching to about half-length of mesonotum. No dark spots in front of cruciform elevation.

Legs: Greenish, yellow-brown towards tarsi. Fore femur with row of four sharply pointed and erect spines.

Tegmina and wings: Hyaline. Veins greenish, and set with setae.

Tymbal organ: Four weakly sclerotized parallel ridges spanning the tymbal from dorsal to ventral margin. Three intercalary ridges form a lateral band across tymbal.

Opercula: Male operculum (fig. 17) very small and not covering tymbal cavity. Distal part of operculum erect and reaching to about half-length of meracan-

thus, not reaching to abdomen. Lateral margin of distal part short, concavely bent into crest around distolateral corner of basal part, and convexly bent into long and almost straight distal margin. Distomedial corner rounded, medial margin straight. Female operculum (fig. 19) almost as large as that of male, broadly sickle-shaped and erect, with almost continuously rounded distal margin.

Abdomen: Male abdomen hardly inflated, yellow brown, darker brown in distal part. Third and 4th tergites almost transparent laterally. A distinct lateroventral brown spot on 3rd tergite and a slightly smaller lateral spot on 8th tergite. First tergite very short and medially partly hidden under metanotum. Female abdomen more robust than that of male, light brown, with a dark brown lateroventral spot on 3rd tergite. Ovipositor sheaths not reaching to apex of caudodorsal beak. Female caudodorsal beak in dorsal view (fig. 20) weakly convex, almost truncate, at apex.

Male genitalia: Pygofer in lateral view as in fig. 14. Dorsal margin weakly convex, angularly bent into almost straight caudodorsal beak. Distal margin concave, almost continuously bent with margin of beak. Protuberance on lateral lobe of pygofer bluntly rounded and weakly projecting posteriad. Pygofer lobe forming an angular corner just above this protuberance. Ventral margin angularly bent at about $1 / 3$ its length and weakly concave towards base. Ventral margins converge to sharp angle at base of pygofer opening. Caudodorsal beak in dorsal view (fig. 15) almost truncate at apex, but ending in very small and sharply pointed medial thorn. Clasper (fig. 17) curving upwards with long and slender, dorsally smoothly rounded and recurving, apical part. Basal part of claspér without clasper heel. Clasper hollow fairly long. Aedeagus (fig. 21) with very long and slender apical part. Basal part of aedeagus with small and angular dorsolateral protuberances. Lateral lobes small, hardly projecting in lateral view. Aedeagus from behind (fig. 22) very slender. Aedeagal pore narrow, oval-shaped and truncate at apex.

Measurements: Body length $\delta: 16.5 \mathrm{~mm}$, 오: $16.0-19.1 \mathrm{~mm}(\overline{\mathrm{x}} 18.0 \mathrm{~mm} \pm 0.9)$; tegmen length $\delta^{*}:$ $20.6 \mathrm{~mm}, \quad,: 22.0-25.5 \mathrm{~mm}(\overline{\mathrm{x}} 24.6 \mathrm{~mm} \pm 1.0)$; head length $\delta^{\prime}: 1.5 \mathrm{~mm}$, ㅇ: : $1.6-1.9 \mathrm{~mm}(\overline{\mathrm{x}} 1.8 \mathrm{~mm})$; pronotum length $\delta: 2.3 \mathrm{~mm}$, , $:=2.8-3.2 \mathrm{~mm}(\overline{\mathrm{x}} 2.9$ mm ); mesonotum length $\delta^{\hat{1}}: 4.0 \mathrm{~mm}, ~ ㅇ: 7.4 .0-5.2 \mathrm{~mm}$ ( $\overline{\mathrm{x}} 4.8 \mathrm{~mm}$ ); head width $\delta: 3.3 \mathrm{~mm}, ~ ㅇ: 4.0-4.8 \mathrm{~mm}$ ( $\overline{\mathrm{x}} 4.6 \mathrm{~mm}$ ); width of pronotal collar $\delta \mathbf{~}: 5.1 \mathrm{~mm}$, $\circ$ :
$5.7-7.0 \mathrm{~mm}(\overline{\mathrm{x}} 6.6 \mathrm{~mm})$.
Distribution (fig. 2). - B. splendida is only known from the Torricelli mountains in northern Papua New Guinea.

Etymology. - Splendidus (Latin) refers to the beautiful colour pattern of this species.

## Baeturia gibberosa sp. n.

(figs. 2, 23-31)
Holotype: 'New Guinea SE / Popondetta / 25 m vi. 66 ' (print); 'Shanahan- Lippert / Light Trap / Bishop Museum' (print), © ©, BPBM. - Paratypes: NEW GUINEA: Popondetta, 60 m, 1-4.ix.1963, J. Sedlacek, 2ठ, BPBM; Popondetta, Inbora Plant'n, 23.ix. 1963, P. Shanahan, 20ै, bpbm; Normanby: Wakaiuna, Sewa Bay, 5-9.xi.1956, W.W. Brandt, if, вРвм; same data but $1-10$ xii. 1956,1 ; ; 11-20.xii. 1956 , 1 ㅇ, both BPBM; Wamula, $400-500 \mathrm{~m}, 31$.xii. 1988 , R. de Keyzer, $3{ }^{\circ}$, Moul.
B. gibberosa is easily recognized by the rectangular dorsal corner of its clasper.

## Description

Body ochraceous brown, sometimes tinged with red, and densely speckled with brown. Tegmina of males $1.1-1.2 \mathrm{x}$ as long as body length, of females 1.3 x . Male abdomen 1.3-1.5 x as long as head and thorax, of females 1.0 x .

Head: Ochraceous, brown speckled. Postclypeus weakly protruding and bluntly rounded anteriorly, $1.7-2.2 \mathrm{x}$ as wide as long. Postclypeus in lateral view distinctly swollen with straight anterior margin, concave towards anteclypeus.

Thorax: Pronotum ochraceous, brown speckled. Mesonotum greenish brown, slightly brown speckled. Two dark spots in front of cruciform elevation.

Legs: Ochraceous and weakly speckled. Fore femur with row of four sharply pointed and erect spines.

Tegmina and wings: Hyaline. Veins ochraceous, moderately set with setae.

Tymbal organ: Four weakly sclerotized parallel ridges spanning the tymbal from dorsal to ventral margin. Three intercalary ridges form a lateral band across tymbal.

Opercula: Male operculum (fig. 30) small and only partly covering tymbal cavity. Distal part of operculum oval shaped, slightly erect and reaching to about $2 / 3$ of length of meracanthus, in some specimens reaching to anterior matgin of 2 nd abdominal

Figs. 23-31. Baeturia gibberosa sp. n. -23 , pygofer in lateral view; 24, pygofer from aslant; 25 , clasper; 26 , female caudodorsal beak in dorsal view; 27 , aedeagus from behind; 28 , aedeagus in lateral view; 29 , male caudodorsal beak in dorsal view; 30 , male operculum; 31, female operculum
Figs. 32-39. Baeturia retracta sp. n. - 32, female caudodorsal beak in dorsal view; 33, pygofer in lateral view; 34, aedeagus from behind; 35 , aedeagus in lateral view; 36 , pygofer from aslant; 37 , male caudodorsal beak in dorsal view; 38 , female operculum; 39, male operculum.
segment. Lateral margin of distal part fairly long and directed mesiad, concavely bent into crest around distolateral corner of basal part, and forming an obtuse angle with straight distal margin. Medial part of operculum angularly rounded, extending medially of meracanthus. Female operculum (fig. 31) short, sickleshaped and erect.
Abdomen: Male abdomen not inflated, ochraceous brown and brown speckled, though ventrally unspeckled. Segmental hind margins reddish. Tergites 3-6 almost transparent laterally. A distinct lateroventral row of brown spots on tergites 3-8. First tergite very short and medially partly hidden under metanotum. Female abdomen red-brown and densely speckled. Female caudodorsal beak (fig. 26) sharply pointed at apex.

Male genitalia: Pygofer in lateral view as in fig. 23. Dorsal margin weakly concave, but convexly bent into almost straight caudodorsal beak and concave near apex of beak. Distal margin concave, and continuously bent into margin of beak. Protuberance on lateral lobe of pygofer bluntly rounded and weakly projecting posteriad. Pygofer lobe forming a small angular corner just above this protuberance. Ventral margin almost straight, but convexly bent just under lateral protuberance. Ventral margins converge to sharp angle at base of pygofer opening (fig. 24). Caudodorsal beak in dorsal view (fig. 29) truncate at apex. Clasper (fig. 25) almost straight and directed posteriad. Apical part of clasper forming a distinct and angularly protruding dorsal crest, giving the clasper a humped appearance. Clasper hollow fairly short. Basal part of clasper gradually widening, not forming an angular clasper heel. Aedeagus (fig. 27) with fairly long and slender apical part, but much shorter than in most other species of this group, and well developed lateral lobes. Aedeagus from behind (fig. 28) very slender, aedeagal pore narrow, oval-shaped and truncate at apex.

Measurements: Body length ${ }^{\text {ox}}$ : 14.7-17.8 mm ( $\overline{\mathrm{x}}$ $15.9 \mathrm{~mm} \pm 1.0), \quad \circ: 16.5-17.7 \mathrm{~mm}(\overline{\mathrm{x}} 17.2 \mathrm{~mm} \pm$ 0.5 ); tegmen length ô: $17.1-19.1 \mathrm{~mm}(\overline{\mathrm{x}} 18.1 \mathrm{~mm} \pm$ $0.8), \quad \circ: 21.9-22.3 \mathrm{~mm}(\bar{x} 22.1 \mathrm{~mm} \pm 0.9)$; head length $\bar{\delta}: 1.2-1.5 \mathrm{~mm}(\overline{\mathrm{x}} 1.3 \mathrm{~mm})$, 우: $1.5-1.9 \mathrm{~mm}$ ( $\overline{\mathrm{x}}$ 1.7 mm ); pronotum length $\delta$ : $1.9-2.3 \mathrm{~mm}(\overline{\mathrm{x}} 2.1$ $\mathrm{mm}), \circ: 2.6-2.9 \mathrm{~mm}(\overline{\mathrm{x}} 2.8 \mathrm{~mm})$; mesonotum length $3: 2.9-3.5 \mathrm{~mm}(\overline{\mathrm{x}} 3.2 \mathrm{~mm}), \quad \%: 4.1-4.6 \mathrm{~mm}(\overline{\mathrm{x}} 4.4$ $\mathrm{mm})$; head width $\delta: 2.5-3.7 \mathrm{~mm}(\overline{\mathrm{x}} 3.3 \mathrm{~mm})$, $: 9: 4.0-$ $4.3 \mathrm{~mm}(\bar{x} 4.2 \mathrm{~mm})$; width of pronotal collar ${ }^{\text {ot }}: 4.3-$ $5.2 \mathrm{~mm}(\bar{x} 4.6 \mathrm{~mm}), \mp: 5.9-6.4 \mathrm{~mm}(\bar{x} 6.2 \mathrm{~mm})$.

Distribution (fig. 2). - B. gibberosa is recorded from Popondetta on the Papuan Peninsula and from Normanby Island.
Etymology. - Gibberosus (Latin) humpbacked refers to the hump-shaped dorsal protrusion of the clasper.

## Baeturia retracta sp. n.

(figs. 2, 32-39)
Holorype: 'papua new guinea / Tabubil, Western Province / $5^{\circ} 15^{\prime} \mathrm{S} 140^{\circ} 13^{\prime}$ E / 13 oct. 1992 / R.B. Lachlan' (print, day and month written); 'On loan from / M.S. Moulds / Sydney' [print, violet label], ठै, AMS). - Paratypes: same data as holotype but 17.x.1992, 1 §े, Moul; same data but $29 . x i .1992$, $10^{\circ}$, Moul; Garaina, 11-14.vii.1969, J.L. Gressitt, $10^{\circ}$, BPBM; Gurakor, Wampit R. Valley, 45 mi from Lae, $670 \mathrm{~m}, ~ 8 . v .1959$, L.J. Brass, Sixth Archbold Exped. to Papua New Guinea, 1 'f, AMnH; Gurakor, 7.vii. 1965, H. Pyka, 10 , SMN; Iongai, $1700-1900 \mathrm{~m}$, 9.xi. 1965, J. \& M. Sedlacek, 1 ㅇ, BPBM; Iongai, 10 km E of Mt. Albert Edward, 1450-1800 m, 8-10xi. 1965, J. Sedlacek, 10゙, вРвм; Kiunga, 18.vii. 1970, O.K. McCaw, $10^{\circ}$, AMS; Kiunga, $5^{\circ} 15^{\prime}$ S $141^{\circ} 05^{\prime}$ E, 2280 m , 25.v. 1970 , O.K. McCaw, 1 if Moul; Kiunga, Fly River, 35 m, viii.1969, J. Sedlacek, 1f, вРвм; Koitaki, 1500 ft, xxi. 1928, Pemberton, 1 太, BPBM; Wau, Morobe Distr., $1050-$ 1100 m, 15.xii.1961, J., J.H. \& M. Sedlacek, $1 \delta^{\top}$, врвм; Yagaum, Madang, 3-5.vi. 1965, H. Pyka, $1 \delta^{\circ}$, SMN; MANUS: Manus isl., 24.i. 1966, 10 ${ }^{\text {or }}$ SMN.
B. retracta can be recognized by its erect and retracted claspers, which lie for the greater part concealed within the pygofer. The species is closely related to $B$. laminifer Blöte, sharing a very similar clasper with a similarly square, laminiform, dorsal protrusion. This protrusion, however, is completely concealed within the pygofer, and therefore hardly visible in B. retracta. B. retracta is regarded as a separate species, since the retraction of the clasper appears to be a constant character. Furthermore, B. retracta tends to have a more strongly swollen postclypeus, while males have a slightly shorter operculum and a shorter, more rounded, protuberance on the lateral lobe of the pygofer.

## Description

Body of males light yellow-brown or greyish brown and distinctly speckled with brown. Females more red-brown and more densely speckled. Female head and thorax larger than in male. Tegmina of males 1.11.3 x as long as body length, of females 1.2-1.4 x . Male abdomen 1.2-1.5 x as long as head and thorax, of females $1.0-1.3 \mathrm{x}$.

Head: Ochraceous, densely brown speckled. Postclypeus weakly protruding, almost oblongshaped and bluntly rounded anteriorly. Postclypeus $1.4-2.0 \mathrm{x}$ as wide as long. Postclypeus in lateral view distinctly swollen with strongly convex anterior margin.

Thorax: Pronotum ochraceous or greyish brown, with brown speckling concentrated in, often somewhat reddish, medial band and on medial parts of pronotal lobes. Mesonotum greenish brown, densely brown speckled. Two brown spots in front of cruciform elevation.

Legs: Ochraceous, often reddish towards tarsi, and weakly speckled. Fore femur with row of three or four sharply pointed and erect spines.

Tegmina and wings: Hyaline. Veins ochraceous, and densely set with setae.
Tymbal organ: Four weakly sclerotized parallel ridges spanning the tymbal from dorsal to ventral margin. Three intercalary ridges form a lateral band across tymbal.

Opercula: Male operculum (fig. 39) very small and not covering tymbal cavity. Distal part of operculum erect and reaching to about half-length of meracanthus, but not reaching to abdomen. Lateral margin of distal part short, concavely bent into crest around distolateral corner of basal part, and angularly bent into long and concave distal margin. Distomedial corner rounded, medial margin straight. Male from Iongai with larger, more oval-shaped, operculum. Female operculum (fig. 38) almost as large as that of male, broadly sickle-shaped and erect, with almost continuously rounded distal margin.

Abdomen: Male abdomen not inflated, ochraceous brown and brown speckled, though ventrally unspeckled. Segmental hind margins sometimes reddish. Tergites 3-6 almost transparent laterally. A distinct lateroventral row of brown spots on tergites 3-8. First tergite very short and medially almost completely hidden under metanotum. Female abdomen more robust than that of male, dark reddish brown, and densely speckled. Lateroventral spots on tergites 3-8 very distinct. Ovipositor sheaths not reaching to apex of caudodorsal beak. Female caudodorsal beak in dorsal view (fig. 32) truncate at apex (pointed in Iongai specimen).

Male genitalia: Pygofer in lateral view as in fig. 33. Dorsal margin concave, but convexly bent into almost straight caudodorsal beak and concave near apex of beak. Distal margin almost straight, but concavely bent into straight margin of beak. Protuberance on lateral lobe of pygofer bluntly rounded and weakly projecting posteriad. Pygofer lobe sometimes forming a small rounded corner just above this protuberance. Ventral margin almost straight, but convexly bent just under lateral protuberance and weakly concave near base. Ventral margins converge to sharp angle at base of pygofer opening (fig. 36). Caudodorsal beak in dorsal view (fig. 37) weakly bicuspidate at apex. Clasper strongly upcurved, reaching into basal part of caudodorsal beak. Dorsal part of clasper with laminiform and square-shaped protrusion. Clasper hollow fairly short. Aedeagus (fig. 35) with long apical part, but shorter and broader than in most other species of this group, and with well developed lateral lobes. Basal part of aedeagus very stout. Aedeagus from behind (fig. 34) long and slender, though fairly broad between its lateral lobes. Aedeagal pore narrow, oval-
shaped and truncate at apex.
Measurements: Body length ô: $14.9-19.6 \mathrm{~mm}(\bar{x}$ $17.2 \mathrm{~mm} \pm 1.4$ ), $\circ: 16.3-18.1 \mathrm{~mm}(\bar{x} 17.5 \mathrm{~mm} \pm$ 0.7 ); tegmen length ${ }^{\hat{\prime}}: 18.2-21.9 \mathrm{~mm}(\overline{\mathrm{x}} 20.6 \mathrm{~mm} \pm$ 1.1), ㅇ: $21.4-25.9 \mathrm{~mm}(\overline{\mathrm{x}} 23.0 \mathrm{~mm} \pm 1.7$ ); head length of: 1.3-1.5 mm ( $\bar{x} 1.4 \mathrm{~mm}$ ), 오: 1.6-1.8 mm ( $\bar{x}$ 1.7 mm ); pronotum length $\overline{0}: 2.1-2.5 \mathrm{~mm}$ ( $\overline{\mathrm{x}} 2.2$ $\mathrm{mm}), ~ i: 2.4-2.7 \mathrm{~mm}(\overline{\mathrm{x}} 2.6 \mathrm{~mm})$; mesonotum length ठ': $3.2-4.1 \mathrm{~mm}(\overline{\mathrm{x}} 3.7 \mathrm{~mm}), \quad+: 3.9-4.8 \mathrm{~mm}(\overline{\mathrm{x}} 4.2$ $\mathrm{mm})$; head width $\delta$ : $3.4-3.8 \mathrm{~mm}(\overline{\mathrm{x}} 3.7 \mathrm{~mm})$, $8: 3.9-$ 4.2 mm ( $\bar{x} 4.1 \mathrm{~mm}$ ); width of pronotal collar ${ }^{\text {or }}: 4.7$ $5.6 \mathrm{~mm}(\overline{\mathrm{x}} 5.1 \mathrm{~mm}), \quad \circ: 5.7-6.3 \mathrm{~mm}(\overline{\mathrm{x}} 5.9 \mathrm{~mm})$.

Distribution (fig. 2). - B. retracta is widely distributed in Papua New Guinea, and also recorded from Manus Island of the Admiralty Islands.
Etymology. - Retracta refers to the retracted claspers of this species.

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## References

Anonymous, 1938. Atlas van tropisch Nederland: [i-ix], maps 1-31b, legenda 1-31b, 1-17 (Koninklijk Nederlands Aardrijkskundig Genootschap \& Topografische Dienst in Nederlandsch-Indië, Amsterdam/Batavia).
Anonymous, 1966. List of New Guinea localities: 1-19 (Bernice P. Bishop Museum, Honolulu).
Anonymous, 1968. The Times atlas of the world: i-xliii, 1272, pls. 1-123. (Times Newspapers Ltd., London).
Blöte, H. C., 1960. The genus Baeturia Stål as represented in New Guinea (Homoptera, Cicadidae). - Zoologische Mededelingen, Leiden 37: 61-80.
Boer, A. J. de, 1982. The taxonomy and biogeography of the nasuta group of the genus Baeturia Stål, 1866 (Homoptera, Tibicinidae). - Beaufortia 32 (4): 57-78.
Boer, A. J. de, 1986. The taxonomy and biogeography of the conviva group of the genus Baeturia Sràl, 1866 (Homoptera, Tibicinidae). - Beaufortia 36 (7): 167-182.
Boer, A. J. de, 1989. The taxonomy and biogeography of the bloetei group of the genus Baeturia Stal, 1866 (Homoptera, Tibicinidae). - Beaufortia 39 (1): 1-43.
Boer, A. J. de, 1990. Aedeastria, a new cicada genus from New Guinea, its phylogeny and biogeography

Homoprera. lïbicinidace, preceded by a discussion on the tanomoms of the New Guinean Tibicinidac. Beautorth - (1) (3): 63- $\mathbf{3}^{-2}$
Buer. A. I. de. 14) 2. The taxonomy and biogeography of the : artai, group of the genus Bacturia Srâl, 1866 Homoprera. Tibbicinidae). - Bijdragen tot de Dierkunde 61 (3): 16,3-183.
Beer, A. I. de, 190 ta. The taxonomy and biogeography of the lorize group of the genus Baterria Stål, 1866 Homoptera. Tihicinidac). - Tijdschrift voor Entomologie 13": 1-26.
Boer. A. I. de. 19) ft. The taxonomy and biogeography of the ahtausaz group of the genus Baeturia Stål, 1866 1Homopeera, Tibicinidace) - Beaufortia (in press).
Beer. A. J. de, 1994c. The taxonomy and biogeography of the gursulinervis group of the genus Baeturia Stål, 1866 (Homoptera. Tibicinidac) - Bijdragen tot de Dierkunde (2.4: $8^{-}-100$

Daly: M. C.. M. A. Cooper, I. Wilson, D. G. Smith, \& B. G. [). Hooper. 1991. Cenozoic plate tectonics and basin evolution in Indonesia. - Marine and Petroleum Geology . 1 PGD) $8(1): 1-21$.
1)uffels. I. P.. 1983. Taxonomy, phylogeny and biogeography of the genus Cosmopsaltria, with remarks on the historic biogeography of the subtribe Cosmopsaltiaria (Homoptera: Cicadidae). - Pacific Insects Monographs 34): 1-12 $2^{-}$.

Duffels. J. P., 1986. Biogeography of Indopacific Cicadoidea, a tentative recognition of areas of endemism. - Cladistics 2: 318-336.

Duffels. J. P., 1993. The systematic position of Moana expansad (Homoptera: Cicadidae), with reference to sound organs and the higher classification of the superfamily

Cicadoidea. - Journal of Natural History 27: 1223-1237.
Duffels, J. P. \& A. J. de Boer, 1990. Areas of endemism and composite areas in East Malesia. In: P. Baas, C. Kalkman \& R. Geesink (eds.), The plant diversity of East Malesia: Proccedings of the Flora Malesiana Symposium commemorating Professor Dr. C.G.G.J. van Steenis, Leiden, August 1989: 249-272 (Kluwer Academic Publishers, Dordrecht).
Duffets, J. P. \& P. A. van der Laan, 1985. Catalogue of the Cicadoidea (Homoptera, Auchenorhyncha) 1956-1980. Series Entomologica 33: i-xvi, 1-414.
Jong, M. R. de, 1985. Taxonomy and biogeography of oriental Prasiini 1: The genus Prasia Stål, 1863 (Homoptera, Tibicinidae). - Tijdschrift voor Entomologie 128: $165-$ 191.

Pigram, C. J., \& H. L. Davies, 1987. Terranes and the accretion history of the New Guinea orogen. B. M. R. Journal of Australian Geology \& Geophysics 10: 193-212.
Rangin, G., L. Joliver, \& M. Pubellier, 1990a. A simple model for the tectonic evolution of the southeast Asia and Indonesia region for the past 43 my . - Bulletin de la Société géologique de France (8) 6:889-905.
Rangin, G., M. Pubellier, J. Azema, A. Briais, P. Chotin, H. Fontaine, P. Huchon, L. Jolivet, R. Maury, C. Muller, J. P. Rampnoux, J.-F. Stephan, J. Tournon, N. Cottereau, J. Dercourt, \& L. E. Ricou, 1990b. The quest for Tethys in the western Pacific. 8 paleogeodynamic maps for Cenozoic time. - Bulletin de la Société géologique de France (8) 6 (6): 907-913

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## A REVISION OF THE NEW WORLD GENUS

ERPETOGOMPHUS HAGEN IN SELYS
(ODONATA: GOMPHIDAE)


#### Abstract

Garrison, R. W., 1994. A revision of the New World genus Erpetogomphus Hagen in Selys (Odonata: Gomphidae). - Tijdschrift voor Entomologie 137: 173-269, figs. 1-230, tabs. 1-4 [issn 0040-7496]. Published 15 December 1994. This revision of adult Erpetogomphus includes a phylogenetic assessment of all 21 species using outgroup comparison and parsimony algorithm, descriptive biogeography, keys to both sexes, synonymies, descriptions, type designations, and illustrations, including distribution maps of al! species. Six new species are described: E. agkistrodon, E. leptophis, E. elaphe, E. liopeltis, E. bothrops, E. heterodon. Erpetogomphus coluber is considered a junior synonym of $E$. compositus, $E$. natrix is considered a subspecies of $E$. lampropeltis, and a neotype is designated for $E$. cophias. Phylogenetic assessment of 41 mostly somatic characters shows Erpetogomphus to be partitioned into three monophyletic groups: 1) six dark green species (E. constrictor, E. sabaleticus, E. tristani, E. agkistrodon, E. schausi, E. ophibolus) with mostly allopatric or parapatric distributions along the eastern coast of Mexico south into northern Colombia and Venezuela, 2) two species (E. leptophis, E. eutainia) with distributions from southern Texas south through Mexico and up the west coast to Michoacan states, and 3) 13 remaining species (E. elaphe, E. elaps, E. Liopeltis, E. botbrops, E. viperinus, E. designatus, E. sipedon, E. lampropeltis, E. crotalinus, E. heterodon, E. compositus, E. boa, E. cophias) with distributions in the central United States south through Mexico to Costa Rica. Derived characters states were gleaned mostly from primary and secondary genitalic characters (head structure, penis, hamules, caudal appendages, vulvar lamina); but satisfactory resolution of terminal clades, especially of the third group, is difficult due to apparent character reversals. Dr. R. W. Garrison, Los Angeles County Agricultural Commissioner's Office, 3400 La Madera Ave., El Monte, CA 91732, USA


Key words. - Odonata; Gomphidae; Erpetogomphus, systematics; keys; cladistics; phylogeny; biogeography.

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Introductory part

## Introduction

No other group of Middle American Gomphidae has been in need of revision as much as the genus Erpetogomphos Hagen in Selys. The last species were described by Williamson and Williamson (1930), and although several odonatists have since received species which are possibly undescribed, there has been a hesitancy to describe them because so many species are rare or poorly known. Several years ago, I collecsed a few specimens of one species I thought new, but $\|$ realized that an examination of most of the types would be necessary before I could resolve the problem.

This paper is the result of those investigations. My purpose here is to: 1) describe and figure all species currently going under the generic name Erpetogomphus, 2) select lectotypes and establish type localities, if possible, and 3) construct a cladogram of all the known species so that phylogenetic relationships and distribution patterns can be adduced for the genus. I have examined types for all taxa except for two species, E. cophias (type lost), and E. ophibolus. The last species is distinct, and no confusion exists about its identity or type deposition. A neotype is selecred for $E$. cophias. I also provide keys for all species.

Twenty-three names (table 1) have been included in Erpetogomphus, but three, E. severus Selys, Herpetogomphus rupinsulensis Walsh, and H. pictus Needham (= Ophiogomphus rupinsulensis (Walsh)) have been transferred to Ophiogomphus. Another, E. menetriesii (Selys) I consider to be a nomen dubium, for reasons detailed below; and six new species are described. One species thought lost, E. boa Selys, has been rediscovered, and $E$. viperinus auctorum is really an undescribed species. I suspect that several new species will be found in Mexico and Central America, and I hope this paper will make it easier for others to describe those new species.

The difficulties I have encountered during this study have mostly been due to the paucity of specimens of various species which have been considered rare. I have successfully associated both sexes of all current species except the male of E. agkistrodon, which is unknown. My task of recording aspects of intraspecific variation has been made considerably casier by various museum authorities and individuals listed under acknowledgements.

## History

The genus Erpetogomphus was described (as a subgenus) by Selys (1858) to receive two previously described species, Gomphus menetriesii Selys, 1850, and Ophiogomphus crotalinus Hagen in Selys (the latter originally described from a pair in 1854), as well as
two new species, Erpetogomphus elaps Selys, and $E$. cophias Selys. In the 'Appendice' of the same paper, Hagen described $E$. compositus and $E$. designatus. Selys described E. boa in 1859 and E. viperinus in 1868. Some of these species were briefly redescribed or discussed by Selys and Hagen in 1859, 1869, 1873, and 1878. In 1879, Selys redefined the subgenus Herpetogomphus (an unjustified emendation of Erpetogomphus, see Cowley, 1934), under which were included eight species: $E$. compositus, $E$. designatus, $E$. viperinus, E. menetriesii, E. elaps, E. boa, E. cophias, and E. crotalinus. Calvert (1899) redescribed E. viperinus (now E. bothrops sp. n.) based on material collected in Tepic, and included a diagnosis of the females of E. viperinus and E. elaps. Kirby (1890), in a synonymic catalogue of the world Odonata, designated Herpetogomphus crotalinus as the type species of the genus. Calvert (1905) included a synopsis for all Erpetogomphus from Middle America and described E. eutainia, E. ophibolus, E. sipedon, and E. diadophis. Calvert (1912a) added E. tristani. Ris (1917) described $E$. constrictor and redescribed what he thought to be the second known male of E. boa. In 1918 Williamson described E. sabaleticus from a pair from Colombia, and Kennedy (1918) described E. lampropeltis from California. Calvert (1919) described E. schausi from a male from Guatemala. Finally, Williamson and Williamson (1930) described two new species from Baja California, E. coluber and $E$. natrix. In the same paper, they presented a synopsis of what was known for all species. Their summary makes clear that only a few common species were represented in collections, and that many others were poorly known and/or described from insufficient material.

## The status of the name Erpetogomphus menetriesii (Selys)

This name was first applied by Selys to an incomplete male supposedly from Brazil. Selys (1850) briefly compared it to Paragomphus genei, as follows: 'Its colouration yellow, almost without spots, prevents confusion of this species with any other in Europe, but it resembles in this respect G. pallidus (Ramb.) of Southern America whose stature is stronger, and Gomphus menetriesii (new) from Brazil, but the feet of the last are mostly black, the ot abdomen is bordered by black on the two sides, etc...'. The same specimen was redescribed it in 1854 as follows: 'Head yellow, except for space between ocelli, that between the eyes and ciliated margin of the occiput black. Front of thorax yellow, with 2 median, contiguous stripes, almost obliterated, pale red, antehumeral and humeral stripe equidistant.
ot Appendages lacking. From: Brazil?' The type is said to be in the 'Musée de St- Pétersburg'.

Table 1. Species-group names used in the genus Erpetogomphus

| Name | Original <br> Genus | Original <br> Reference | Type locality | Location of type | Type starus | Reference for first placement in Erpetogomphus | Present placement |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| agkistrodon | Erpetogomphus | This paper | Jalapa, Mexico | UNAM ( ${ }^{\circ}$ ) | Holotype | This paper | E. agkistrodon |
| boa | Erpetogomphus | Selys 1859 | Vera Cruz, Mexico | IRSN ( ${ }^{\circ}$ ) | Lectotype | Selys 1859 | E. boa |
| bothrops | Erpetogomphus | This paper | Rio Otapa, Veracruz State, Mexico | USNM ( ${ }^{\text {® }}$ ) | Holorype | This paper | E. bothrops |
| coluber | Erpetogomphus |  <br> Williamson 1930 | San José de Comandu, Baja Calif., Mexico | UMMZ ( ${ }^{\text {* }}$ ) | Holotype |  <br> Williamson 1930 | E. compositus |
| compositus | Erpetogomphus | Hagen in <br> Selys 1858 | Pecos River, Texas; corrected to vicinity of Roswell, NM | MCZC (\%) | Holotype | $\begin{aligned} & \text { Hagen in Selys } \\ & 1858 \end{aligned}$ | E. compositus |
| constrictor | Erpetogomphus | Ris 1917 | Misantla, Veracruz, Mexico | SMF (\%) | Lectotype | Ris 1917 | E. constrictor |
| cophias | Erpetogomphus | Selys 1858 | Mexico | MNHP (\%) | Neorype | Selys 1858 | E. cophias |
| crotalinus | Ophiogomphus | Hagen in <br> Selys 1858 | Mexico | MCZC ( ${ }^{\text {( ) }}$ | Lecrotype | $\begin{aligned} & \text { Hagen in Selys } \\ & 1858 \end{aligned}$ | E. crotalinus |
| designatus | Erpetogomphus | Hagen in Selys 1854 | Pecos River, Texas; corrected to vicinity of Roswell, NM | MCZC ( ${ }^{\text {a }}$ ) | Lectotype | $\begin{aligned} & \text { Hagen in Selys } \\ & 1858 \end{aligned}$ | E. designatus |
| diadophis | Erpetogomphus | Calvert 1905 | Texas | BMNH ( ${ }^{\text {\% }}$ ) | Holotype | Calvert 1905 | eutainia |
| elaphe | Erpetogomphus | This paper | Costa Rica | FSCA ( $\delta^{\circ}$ ) | Holotype | This paper | E. elaphe |
| elaps | Erpetogomphus | Selys 1858 | Mexico | MNHP (\%) | Holotype | Selys 1858 | E. elaps |
| eutainia | Erpetogomphus | Calvert 1905 | Guerrero, Mexico | BMNH ( ${ }^{\text {a }}$ ) | Holotype | Calvert 1905 | E. eutainia |
| beterodon | Erpetogomphus | This paper | Aragon, New Mexico | USNM ( ${ }^{\text {( })}$ | Holotype | This paper | E. heterodon |
| lampropeltis | Erpetogomphus | Kennedy 1918 | Fillmore, California | USNM ( ${ }^{\circ}$ ) | Holotype | Kennedy 1918 | E. l. lampropeltis |
| leptophis | Erpetogomphus | This paper | Blue Creek, Belize | FSCA (\%) | Holotype | This paper | E. leptophis |
| liopeltis | Erpetogomphus | This paper | La Estanzuela, Nuevo Leon, Mexico | FSCA ( $\left.{ }^{( }\right)$ | Holotype | This paper | E. liopeltis |
| menetriesii | Ophiogomphus | Selys 1854 | Brazil? | Lost |  | Selys 1858 | Nomen dubium |
| montanus | Herpetogomphus | Selys 1878 | Yellow Town, Montana | IRSN ${ }^{\text {a ( }}$ ( ${ }^{\text {a }}$ ) | Holotype | Selys 1879 | Ophiogomphus severus montanus |
| natrix | Erpetogomphus | Williamson \& Williamson 1930 | San Jose de Comandu, Baja Calif., Mexico | UMMZ ( ${ }^{\text {® }}$ ) | Holotype | Williamson \& Williamson 1930 | E. lampropeltis natrix |
| ophibotus | Erpetogomphus | Calvert 1905 | Atoyac, Veracruz, Mexico | BMNH ( ${ }^{\text {( ) }}$ | Holotype | Calvert 1905 | E. ophibolus |
| pictus | Herpetogomphus | Needham 1897 | Ithaca, New York | CUIC ( ${ }^{\text {( ) }}$ | Holotype | Needham 1897 | Ophiogomphus rupinsulensis ${ }^{2}$ |
| rupinsulensis | Herpetogomphus | Walsh 1862 | Rock Island, Illinois | Lost |  | Walsh 1862 | Ophiogomphus rupinsulensis |
| sabaleticus | Erpetogomphus | Williamson 1918 | Cristalina, Colombia | UMMZ ( ${ }^{\text {a }}$ ) | Holotype | Williamson 1918 | E. sabaleticus |
| schausi | Erpetogomphus | Calvert 1919 | Purulta, Guatemala | ANSP ( $0^{\circ}$ ) | Holotype | Calvert 1919 | E. schausi |
| severus | Ophiogomphus | Hagen 1874 | Colorado | Lostơ ( $\begin{aligned} & \text { ( }\end{aligned}$ |  | Selys $1878^{\circ}$ | Ophiogomphus severus |
| sipedon | Erpetogomphus | Calvert 1905 | Guadalajara, Mexico | BMNH (\%) | Holotype | Calvert 1905 | E. sipedon |
| tristani | Erpetogomphus | Calvert 1912 | Oricuajo, Costa Rica | ANSP (0) | Holorype | Calvert 1912 | E. tristani |
| viperinus | Erpetogomphus | Selys 1868 | Orizaba, Veracruz, Mexico | IRSN ( ${ }^{\text {O }}$ ) | Lectotype | Selys 1868 | E. viperinus |

[^14]In 1858. Selys discussed E. menetriesii under E. croralinus (in translation):
'N.B. In the Synopsis des Gomphines, I described under the name of Ophiogomphus? Menetriesii an incomplete male individual, which was communicated to me a long time ago by the knowledgeable M. Menerries. Now, I am inclined to believe that this is not a species distinct from crotalinus.

One reason that made me suppose the species different. was its occurrence in Brazih but it was still necessary that this fact be duly confirmed and that the missing anal appendages be known.

This $\delta$ is a little smaller than our specimens [the presently described males and female of E. crotalinus], the space around the ocelli is distinctly black, the appearance of the reddish antehumeral stripes more closely approaches the median [stripes], the humeral is more distinct and more elongated, the base of the abdomen is less swollen, the dorsal yellow stripe of 3 rd, 4th, and 5th segments seems modified: it consists of spots of three lobes [each], whose posterior isn't visibly narrowed into a little round head. (This note is based on diagnosis and a description made a long time ago, so that I have not been able to compare the specimen since I have had crotalinus at hand.)

A female, same indication of origin [Brazil] from the Mus. St. Petersburg, communicated to M. Hagen by M. Menetries differs very slightly from crotalinus by the following: 6 cells under the pterostigma.; costa yellower, tarsi black, the first article [tarsomere] of the posterior [leg] yellow, black tibiae bilineated with yellow outside; femora yellow with a double black external line shorter and finer at the ends. Abdomen thicker, marked with black; 9th segment black above with a dorsal round spot, touching the posterior margin. [Abdominal segment] 10 and appendages light red, the end of these last [structures] pale not marked with black [italics are Selys']. Vertex more black. Dimensions a little more robust.
Not being certain of its identity, I fear to cause confusion later by putting the name $E$. menetriesii in true crotalinus from Mexico, even though I have already published this name [menetriesi] in speaking of another species (page 102 Revue des Odonates, 1850; and Synopsis des Gomphines, 1854).

If its identity is confirmed, it would be necessary to replace the name of $E$. crotalinus with that of $E$. Menetriesii.

Finally, Selys (1878) described Herpetogomphus menetriesii based on 'un mâle in complet [in MNHP]' and female [in IRSN] from Guatemala. He states: 'I Colicee it idential with the example described in Synopsis No. 20 [1854] and reported with doubt as rotalinus in the Monographie [1858].' Calvert 1905 pointed out the inconsistencies of the three $\therefore \quad \therefore$ riptions and left open the question of what $E$.
menetriesii really is.
According to the original description (Selys, 1854), E. crotalinus cannot be $E$. menetriesii. A supplementary male of E. crotalinus described in 1858 (in zмнв, examined) is unusual in having faint antehumeral stripes, but no $E$. crotalinus I have examined fits the description of E. menetriesii of 1854 (with middorsal, antehumeral and humeral stripes).

The 1854 description is too brief to indicate what E. menetriesii is, and I cannot associate it with any known species. According to Hagen (1861), it was destroyed. I have been unable to trace the 1858 female described by Selys, but I was able to examine the 'pair' described in 1878. The two specimens, which have determination labels by Selys, are really two females of $E$. eutainia Calvert. However, I cannot be certain that these females are conspecific with Selys' original description of $E$. menetriesii. Selys and others apparently had difficulty in associating specimens of E. eutainia, because a male from Texas in the IRSN is given the manuscript name of Erpetogomphus berus, another female is identified as Cyanogomphus ?mexicanus (by R. Martin), and, of course, the two females, one in the IRSN, the other in the MNHP, are labelled as E. menetriesii.

Thus, the original description of $E$. menetriesii may refer to $E$. eutainia, but because no one can ever be sure of this, I propose that $E$. menetriesii should be considered a nomen dubium.

## Biology

Little is known of the biology of the genus, though their habits as recorded in the literature mirror the behaviour of other Gomphidae. Adults are most commonly found near shores of streams and rivers, but they may also be found in agricultural stubble or on tree branches near streams. Many species are seldom encountered and only a few species appear to be common in collections.

Available notes taken from the literature, personal correspondence, and personal observations cited under various species accounts indicate that female Erpetogomphus oviposit by swiftly tapping the water's surface or by hovering motionless over moderately swift water systems. Details of copulation and copulation time are largely unknown, although the bizarre male and especially female morphological adaptations for achieving the tandem position in E. tristani have been detailed by Calvert (1912a).

Larvae of $E$. designatus, $E$. compositus, and $E$. lampropeltis were keyed by Needham and Westfall (1955), and illustrations and full descriptions of $E$. crotalinus and E. lampropeltis natrix were given by Novelo and González (1991). The larva of $E$. sabaleticus was recently described and illustrated by Belle (1992).

Table 2. Characters used in cladistic analysis of Erpetogomphus adults. oे (male) or $\%$ (female) in parentheses indicates sex to which character pertains. An asterisk (*) indicates an ordered multistate character. Numbers in parentheses following character states refer to figure numbers.

| Number/character | Primitive state | Derived state(s) |
| :---: | :---: | :---: |
| 1. Postocciput widh (\%) | tumid, convex | linear |
| 2. Postocciput (\$) | not visible from above | visible from above |
| 3. Occiput (\%) | semicircular | transverse |
| 4. Vertex (\%) | with long median trough | with no trough |
| 5. Postocellar ridge (f) | incomplete | complete |
| 6. Vertex: | sexually dimorphic | not sexually dimorphic |
| 7. Frontoclypeal suture | with black | with no black |
| 8. Hind margin of metepimeron | with black | with no black |
| 9. Hind wing 3-celled anal loop | present | absent |
| 10. Anterior hamule* ( ${ }^{\text {a }}$ ) | states 1-5 (fig. 208) | state 0 (fig. 208) |
| 11. Posterior hamule* ( ${ }^{\text {² }}$ ) | states 1-4 (fig. 209) | state 0 (fig. 209) |
| 12. Tip of posterior hamule* ( ${ }^{\text {( })}$ | states 1-5 (fig. 210) | state 0 (fig. 210) |
| 13. Posterior lobes of penis segm. 1 ( $\delta^{*}$ ) | small | well developed, divided lobes |
| 14. Penis: segment 3 dorsally ( ${ }^{\text {a }}$ ) | with pair of tubercles | lacking tubercles |
| 15. Penis: segment 4 ( $0^{*}$ ) | about twice as long as wide (fig. 211) | about as long as wide (fig. 211) |
| 16. Prepuce of penis* ( ${ }^{\text {a }}$ ) | states 1-3 (fig. 212) | state 0 (fig. 212) |
| 17. Lateral lobe of penis-shape* ( ${ }^{\text {a }}$ ) | states 1-5 (fig. 213) | state 0 (fig. 213) |
| 18. Membranous hood of penis ( $\mathbf{\delta}^{\text {) }}$ | not overlapping distally | overlapping |
| 19. Shape of cornua of penis* ( ${ }^{(8)}$ | states 1-6 (Fig. 214) | state 0 (fig. 214) |
| 20. Development of cornua of penis* ( ${ }^{\text {² }}$ ) | reduced (1), absent (2) | present (fig. 214) |
| 21. Length of vulvar lamellar plates ( $(\%)$ | about 0.25 length of sternite | $\geq 0.50$ length of sternite |
| 22. Medial margins of vulvar lamina (\%) | folded under | not folded under |
| 23. Postlamellar ridge ( $(\%)$ | Y-shaped | semicircular |
| 24. Postlamellar ridge with juncture ( $\ddagger$ ) | posterior to plates (e.g. figs. 184, 195) | at or before margin of plates (fig. 192) |
| 25. Ventral base of cercus ( ${ }^{\text {a }}$ ) | with large blunt tooth | with no tooth (fig. 94). |
| 26. Ventral base of cercus ( $\delta$ ) | with no carina | with a carina |
| 27. Distal part of cercus ventrally ( ${ }^{\text {® }}$ ) | with no carina | wish a carina |
| 28. Dorsal surface of cercus ( ${ }^{\text {d }}$ ) | with a tubercle | without a rubercle |
| 29. Ventral surface of cercus ( ${ }^{\circ}$ ) | linear or concave | convex |
| 30. Curvature of epiproct with tips (\%) | about parallel to base | at about $90^{\circ}$ to base |
| 31. Tips of epiproct ( ${ }^{\text {d }}$ ) | spatulate or bidentate | pointed |
| 32. Penis segment 3 ( ${ }^{\text {d }}$ ) | short and stocky | longer than wide |
| 33. Penis guard ( $\delta$ ) | quadrate in cros--section | circular in cross-section |
| 34. Epiprocts (8) | approximate and curved at distal 0.50 | not approximatebarely curved |
| 35. Spermatheca ( $¢$ ) | states 1-2 (fig. 215) | state 0 (fig. 215) |
| 36. Dorsal surface of epiproct ( ${ }^{\text {² }}$ ) | with dorsal spine (fig. 95) | unarmed |
| 37. Ventral base of cercus ( $\delta$ ) | with pebble-like carina | otherwise (fig. 97) |
| 38. Membranous hood of penis ( ${ }^{\circ}$ ) | partially (1) or wholly (2) exposing subcuticular membrane | completely covering subcuticular membrane |
| 39. Ventral base of cercus ( $\delta$ ) | with carina ending in tooth (fig. 116) | with carina (if present) not ending in tooth |
| 40. Postocciput (\%) | with lateral depressions (fig. 165) | with no lateral depressions |
| 41. Occiput (\%) | with a posteriorly directed medial spine fig. 151 | without a spine |

## Phylogenics and biogeography

## Phylogenetic analysis

I have attempted to construct a cladogram based on an algorithm which infers phylogenies based on the presence of shared derived characters. Using the outgroup comparison method of Watrous and Wheeler (1981), I consider Ophiogomphus to be the outgroup to Erpetogomphus, following Carle (pers. comm.) in his assessment of gomphid dragonfly classification. In this analysis, I have used the most austral species, $O$. severus Hagen.

The genus Erpetogomphus has never been easily defined (Calvert 1905, Needham and Westfall 1955), these authors using a series of characters to characterize the genus. A synapomorphy apparently unique to Erpetogomphus is the condition of the epiprocts of the males: all have gently (at least $110^{\circ}$ ) to strongly dor-
sally curved, non-divergent epiprocts.
Carle (1986), in his treatment of the higher classification of the Gomphidae, included Erpetogomphus in the tribe Onychogomphini of the subfamily Onychogomphinae. In his key to the eight subfamilies of Gomphidae, he characterized the Onychogomphinae as follows: 1) hind femur shorter than head, 2) anal triangle typically 4 -celled with small rectangular cell along inner margin, 3) anterior hamuli with shoulder and end hook forming lateral u-shaped notch, and, 4) female sternum 9 with large U - or V-shaped basal membranous area. Within this subfamily, the tribe Onychogomphini (including Erpetogomphus) is distinguished from the tribe Crenigomphini by the following characters (antagonistic characters for Crenigomphini in parentheses): postgenal suture present (absent), male tibial laminae
present (absent), penile segment one (= peduncle) without cuplike thin-walled hood (with bilobed cuplike thin-walled hood), female sternum 9 with domeshaped membranous area not extending to distal half of sternum (this structure extending to distal half of sternum). None of these characters is stated by Carle (1986) to be uniquely derived for any of the above categories. Carle (pers. comm.) and I agree that Ophiogomplius is the most likely outgroup to Erpetogomphus, and we agree that both should be placed in the Onychogomphini. Three characters seem unique to the New World Onychogomphini, which include only Ophiogomphus and Erpetogomphus: 1) rami of male epiprocts contiguous along all or most of their length, 2) female sternum 8 with a well-defined costate postlamellar ridge separating the soft membranous area from the posteriorly sclerotized area (this ridge reduced in Ophiogomphus), 3) presence of a lateral lobe on the ventrolateral margin of the fourth penile segment (fig. 213) (reduced in Ophiogomphus, and not specialized in the E. ophibolus group). This last character seems to be a unique synapomorphy for Ophiogomphus and Erpetogomphus.
Carle (1992) further separated Erpetogomphus into three subgenera, two of which were new: Calogomphus (type species E. eutainia) and Erpetocyclops (type species E. ophibolus). Both Carle (pers. comm.) and I consider Erpetogomphus to be composed of three monophyletic groups as discussed below. While several earlier versions of my cladogram using series of characters and different states generally support a conservative grouping of the genera into three monophyletic groups, I do not think it necessary to provide subgeneric rank to those groups.
A suite of 41 characters, 7 of which are multistate (table 2), was analyzed using the Hennig 86 computer program (Farris, 1988), with the following options: The character set was first run using the ie* (guaranteed to find the most parsimonious trees) option where all characters were ordered and each was cqually weighted (default options). All autapomorphies were included from this analysis. The final set of 41 characters was finally chosen from larger sets, each with different coding sequences, because this set seemed to be the most heuristic. The resultant analysis yielded six equally parsimonious trees. A Nelsen (consensus) tree (length 106, consistency index 0.62) was then generated from the six trees. The resultant characters, tree, and matrix are shown in figs. 208216 , and tables 2-3. As this manuscript was in review, Jan van Tol kindly ran the same data set on paup 3.1.1 for the Macintosh (Swofford 1993) using unordered as well as ordered sequence of multistate characters. His results, which he kindly communicated to me, resulted in 61 semistrict consensus trees each with a length of 94 for unordered multicharacter
states. His resulting cladogram is similar to that in fig. 216 except that the entire $E$. crotalinus group (except for three sister groups: $E$. elaphe and $E$. elaps, $E$. sipedon and E. lampropeltis, and E. boa and E. cophias) collapses to a polytomious assemblage. (I recorded a similar cladogram when multistate characters were listed as unordered for hennig86.) When all multistate characters were run as ordered on paup 3.1.1, the resulting cladogram was the same as shown for fig. 216.

I have not been totally successful in resolving several of the terminal clades, due to high degree of apparent homoplasies. The best characters are those of the secondary genitalia of the male (hamules, penis); but, despite a thorough examination of these structures, I am still uncertain of the correct interpretation of character polarity for several of these complex, multistate character sets (figs. 208-210, 213-214). Little difficulty was encountered in assigning states to members of the E. ophibolus and E. eutainia groups, and their generalized tracks also independently supplement in understanding their phylogeny; but the same characters in males of the $E$. crotalinus group show, if my cladogram is correct, a bewildering complex of character reversals for some characters. The generalized tracks within the E. crotalinus group (fig. 226) also provide few clues to possible speciation events, except for that between the E. elaps and E. crotalinus (s.s.) groups.

Individual clades are numbered as shown in fig. 216, each of which is discussed below.

Clade 1.1. - E. ophibolus group (six species: $E$. constrictor, E. sabaleticus, E. tristani, E. ophibolus, E. agkistrodon, E. schausi) $=$ subgenus Erpetocyclops Carle

Character 3. - female occiput semicircular (parallel development in Clade 1.22112, E. liopeltis)

Character 5. - female postocellar ridge incomplete (parallel development in Clade 1.211, E. leptophis)

Character 6. - vertex sexually dimorphic (state not known for male $E$. agkistrodon, but probably follows for other members of this clade)

Character 11, state 1. - male with posterior hamule triangular (fig. 209)

Character 12, state 1. - male with tip of posterior hamule acuminate (fig. 210)

Character 16, state 1. - male with prepuce well-developed (fig. 212)

Character 17, state 1. - male with lateral lobe vestigial, broadly angulate with no serrations (fig. 213)

Clades 1.11 ( $E$. constrictor, $E$. sabaleticus, $E$. tristani, E. ophibolus), 1.12 (E. agkistrodon), 1.13 (E. schausi). Clade 1.11 is defined by the following synapomorphies:

Character 20, state 2 - male with loss of cornuae of penis (fig. 214). Most Gomphidae (including the outgroup, Ophiogomphus severus), have a pair of long, strap-like cornuae. I have hypothesized the character transformation for this structure throughout the genus (fig. 214).

Character 25. - ventral base of cercus of male with large blunt tooth (fig. 94)

Character 28. - dorsal surface of cercus of male with a tubercle (fig. 94)

I have found no synapomorphy for uniting 1.12 (E. agkistrodon) and 1.13 (E. schausi). The former is known only from the holotype female, the latter by two males and a teneral female.

Clades 1.111 (E. constrictor), 1.112 (E. sabaleticus), 1.113 (E. tristani)

Character 29. - curvature of epiproct of male with tips completely recurved so as to lie parallel to base

Clade 1.21. - E. eutainia group (two species: $E$. eutainia, E. leptophis) $=$ subgenus Calogomphus Carle

Character 10, state 1. - anterior hamule with equally divided arms (fig. 52-53, 208) at distal 0.25

Character 11, state 2. - posterior hamule digitshaped (fig. 209)

Character 14. - dorsal surface of third segment of penis of male with two tubercles

Character 16, state 2. - prepuce of male reduced, largely hidden by frill-like lateral lobe (fig. 212)

Character 17, state 2. - lateral lobe of penis of male frill-like (fig. 213)

Character 35, state 1. - spermatheca of female a single tube (fig. 215). I was only able to study this structure in E. eutainia, as only the allotype of E. leptophis is known. Due to other synapomorphies listed above, I hypothesize that $E$. leptophis will possess single, rather than paired, spermatheca.

Clade 1.22. - E. crotalinus group (13 species: $E$. elaphe, E. elaps, E. liopeltis, E. bothrops, $E$. viperinus, E. designatus, E. sipedon, E. lampropeltis, E. crotalinus, E. heterodon, E. compositus, E. boa, E. cophias) $=$ subgenus Erpetogomphus Hagen in Selys

Character 16, state 3. - prepuce of penis of male lacking

Character 35, state 2. - spermatheca of female paired, each forming a long, narrow sac (fig. 215)

Clade 1.221. - E. elaps subgroup (five species: $E$. elaphe, $E$. elaps, $E$. liopeltis, $E$. bothrops, $E$. viperinus)

Character 17, state 3. - unique shape of lateral lobe (fig. 213)

## Clade 1.2211 (E. elaphe, E. elaps, E. Liopeltis, E. bothrops)

No satisfactory synapomorphy has been found for this clade. One character, the spatulate tip of the epiproct (fig. 129) separates E. viperinus from its sister species.

Clade 1.22111 (E. elaphe, E. elaps, E. liopeltis)
No synapomorphy has been found, but the transverse condition of the female postocciput in $E$. bothrops and in E. viperinus (fig. 166) split E. bothrops from the other three sister species.

## Clade 1.221111 (E. elaphe, E. elaps)

Character 22. - vulvar lamina of female with margin folded under ventrally (figs. 180-181)

Clade 1.222 ( $E$. designatus, E. sipedon, $E$. lampropeltis, E. crotalinus, E. heterodon, E. compositus, E. boa, E. cophias)
No satisfactory synapomorphy has been found to unite the E. crotalinus group (sensu stricto [s.s.]). The one aberrant species in the group is $E$. designatus. It alone has the following characters which separate it from the other 7 species: 1) tip of posterior hamule lacking a tooth (recurrence of this state from the $E$. eutainia group) and 2) unique shape and position of the lateral lobe (fig. 82). The shape of the anterior hamule of $E$. designatus (fig. 60) is, however, like that of E. sipedon, E. lampropeltis, E. crotalinus, and E. compositus, which thus unites these species into a clade.

The widely distributed $E$. designatus has been difficult to place cladistically in this study. It combines characters of the E. elaps group (for example, linear condition of lateral lobe and acuminate condition of the cornua) and even apomorphic characters common to the E. crotalinus (s.s.) group as noted above.

Clade 1.2222 ( E. sipedon, E. lampropeltis, $E$. crotalinus, $E$. heterodon, E. compositus, E. boa, E. cophias)

Character 12, state 5. - tip of posterior hamule of male with anteapical tooth

Character 17, state 5. - unique shape of lateral lobe of fourth segment of penis of male (fig. 213). The small, semicircular condition of the lateral lobe present in Clade 1.2222 is similarly developed in Clade 1.12 ( $E$. schausi).

Clades 1.22221 ( $E$. sipedon), 1.22222 ( $E$.
lampropeltis), and 1.22223 (E. crotalinus, E.
beterodon, E. compositus, E. boa, E. cophias)
No satisfactory apomorphy has been found to dichotomize these clades. The distinctive shape of the cornuae of $E$. sipedon and $E$. lampropeltis (figs. 83-85) has separated these two species from clade 1.22223 .

Table 3. Character matrix used in producing Nelsen (consensus) tree shown in Fig. 216.

| Character No. 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 | 16 | 17 | 18 |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Ancestor | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| Oph. severus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| consrricror | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 |
| sabalericus | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 |
| ristani | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 |
| ophibolus | 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 |
| agkistrodon' | 0 | 0 | 1 | 0 | 1 | $1 ?$ | 1 | 1 | 0 | $0 ?$ | $1 ?$ | 1 ? | 1 | 0 | 0 | 1 ? | $?$ | 1 ? |
| schausi | 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 5 | 1 |
| eutainia | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 1 | 0 | 1 | 2 | 2 | 1 | 1 | 0 | 2 | 2 | 1 |
| leprophis | 1 | 1 | 0 | 0 | 1 | 0 | 1 | 1 | 0 | 1 | 2 | 2 | 1 | 1 | 0 | 2 | 2 | 0 |
| elaphe | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 3 | 2 | 1 | 0 | 1 | 3 | 3 | 1 |
| elaps | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 3 | 5 | 1 | 0 | 1 | 3 | 3 | 1 |
| liopelris | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 3 | 4 | 1 | 0 | 1 | 3 | 3 | 1 |
| bothrops | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 3 | 5 | 1 | 0 | 1 | 3 | 3 | 1 |
| viperinus | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 | 3 | 3 | 1 | 0 | 1 | 3 | 3 | 1 |
| designarus | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 2 | 0 | 0 | 1 | 3 | 4 | 0 |
| sipedon | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| Lampropelris | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| croralinus | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| heterodon | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| compositus | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 | 3 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| boa | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 4 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |
| cophias | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 4 | 5 | 1 | 0 | 1 | 3 | 5 | 1 |

Only the holotype + is known for this species. Male characters were scored as? (unknown), on 0? or 1?, based on affinity with rest of the E. ophibolus group. ${ }^{2}$ Character prevalent in this species. ${ }^{3}$ Basal carina weak in E. boa.
This structure not examined for this species. Plesiomorphy/apomorphy hypothesized by examination of spermatheca in closely related species, and placement of these taxa in cladogram due to other synapomorphies.

Clades 1.222231 ( $E$. crotalinus) and 1.222232 ( $E$. heterodon, E. compositus, E. boa, E. cophias)

No satisfactory apomorphy has been found for clade 1.222232 . Erpetogomphus crotalinus was separated because of the unique shape of its cornuae (fig. 86).

Clades 1.2222321 ( $E$. beterodon), 1.2222322 ( $E$. compositus), 1.2222323 (E. boa, E. cophias)

The unique shape of the cornua (figs. 87-89) has segregated $E$. beterodon and $E$. compositus from clade 1.2222323. Apomorphies for the last clade, representing the two closely related species, $E . b o a$ and $E$. cophias, are:

Character 10, state 5. - unique shape of the anterior hamule of male (fig. 208)

Character 11, state 4. - unique shape of the posterior hamule of male (fig. 209)

Character 19, state 6. - unique shape of the cornua of male (fig. 214)

The clades mentioned above have problems primarily within the E. crotalinus group (sensu lato, [s.l.]), and most dichotomies are based on one or two highly complex character states of the penis. I realize that my interpretation of the morphoclines of this character is open to reevaluation, which could result in a somewhat different shuffling of some species
within the larger group. Only two species in this group, E. boa and E. cophias, are easily characterized as a monophyletic group.

A further analysis to resolve some of the difficulties described above will have to await further specimens and a proper analysis of their larvae. As stated earlier, I have not assigned formal names to subordinate clades, because further investigation may involve transfer of some monophyletic or paraphyletic groups to other clades, nor can I see any purpose in a proliferation of infrageneric names.

## Cladospecies

Of the 21 species described here, seven ( $33 \%$ ) can be defined by autapomorphies and can be termed cladospecies (Ackery and Vane-Wright, 1984). These cladospecies with their uniquely derived character states are shown below.

Clade 1.112. E. ophibolus. Character 36, dorsal surface of epiproct of male with a dorsal spine (fig. 95)

Clade 1.211. E. leptophis. Character 41, female occiput with a posteriorly directed medial spine (fig. 151)

Clade 1.212. E. eutainia. Character 37, ventral base of cercus of male with a pebble-like carina.

Clade 1.2211112. E. elaphe. Character 10, state 3,

2 | 23 | 24 | 25 | 26 | 27 | 28 | 29 | 30 | 31 | 32 | 33 | 34 | 35 | 36 | 37 | 38 | 39 | 40 | 41 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 1 | 0 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 1 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 1 | $0^{4}$ | 1 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | $0 ?$ | 0 | $1 ?$ | $0 ?$ | 1 | 0 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 1 | 0 | 1 | 0 | 1 | 0 | $1 ?$ | 1 | 1 | 1 | $0 ?$ | 0 | 0 | 0 | 0 | 0 | 0 |
| 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | $0^{4}$ | 0 | 1 | 0 | 0 | 0 | 0 |
| 0 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 1 |
| 1 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | $2^{4}$ | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | $2^{4}$ | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 0 | 1 | 1 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 2 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 2 | 0 | 0 | 1 | 0 | 0 | 0 |
| 1 | 1 | 0 | $1^{3}$ | 1 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 2 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | $2^{4}$ | 0 | 0 | 0 | 1 | 1 | 0 |

unique shape of anterior hamule of male (fig. 208)
Clade 1.22221. E. sipedon. Character 38, state 2, membranous hood of penis of male wholly exposing subcuticular membrane.

Clade 1.2222322. E. compositus. Character 38, state 1 , membranous hood of penis of male partially exposing subcuticular membrane.

Clade 1.22223232. E. cophias. Character 39, ventral base of cercus in male with carina ending in tooth (fig. 116), and Character 40, postocciput of female with lateral depression (fig. 165). These two characters are probably correlated: I hypothesize that the postoccipital depressions of the female receive the inferior carinal tooth of the cerci of the male during copulation.

I have been unsuccessful in finding autapomorphies for the 14 other species, and they can be termed paraspecies (Ackery and Vane-Wright, 1984). A further, more detailed analysis of these species may yield autapomorphies. The cladist may question the status of species recognized by single or combinations of plesiomorphic characters, but I agree with Ackery and Vane-Wright (1984: 10), who cogently argue: 'the traditional groupings often (but by no means always) turn out to be good. The cladist must, therefore, always retain respect for the 'Gestalt' approach (espe-
cially when practised by field workers), unless armed with concrete evidence for some contrary arrangement.'.

## Distribution patterns

Table 4 details the distributional patterns of all New World Gomphidae according to five political regions: Canada/Alaska, U. S. A., Mexico, Central America, and South America. All speciose (i.e., $\mathrm{N} \geq 3$ species) gomphid genera in the New World have proportionally more species in the United States and in South America than Erpetogomphus. Erpetogomphus has the most species in Mexico (76\%). Central America follows with $48 \%$, the United States with $24 \%$. No other genus is proportionally as speciose in Mexico. The Gomphus complex of genera (Arigomphus, Gomphus and subgenera, Stylurus), and Ophiogomphus are dominantly North American, and several genera (Agriogomphus, Aphylla, Archaeogomphus, Cyanogomphus, Gomphoides, Neogomphus, Phyllocycla, Phyllogomphoides, Progomphus, and Zonophora) are South American. Epigomphus is most speciose in Central America, where $46 \%$ of all described species occur, closely followed by South America.

Figure 217 shows the density and approximate overall distribution of species of Erpetogomphus as
shown by 150 km squares. Of the five species found in the United States, most are found in the southwestern states. The most northerly distribution for the genus is Grant County, Washington ( $46^{\circ} 50^{\circ} \mathrm{N}$ ), for E. compositus (Paulson and Garrison, 1977). The greatest concentration in species is in south-central Mexico between approximately $18^{\circ}-20^{\circ} \mathrm{N}$ with eight species $(31 \%)$ of all the total number of species in the genus found in southern Veracruz alone.

## Descriptive biogeography

In this section I describe the current distribution range ( $=$ tracks) of various monophyletic groups based on my cladogram. Unfortunately, the numerous problems associated with the cladistic analysis concomitant with the poorly known or spotty distributions (most of which may be due to inadequate collecting in parts of Mexico) preclude a discussion of vicariance biogeography. However, I do discuss possible speciation sequences within the $E$. ophibolus and $E$. eurainiag groups.
The cladogram (fig. 216) for Erpetogomphus divides the 21 known species into three monophyletic groups. The generalized track for the genus is shown in fig. 218. The first two groups, the $E$. ophibolus and E. eutainia groups, have generalized tracks encompassing souch Texas, eastern Mexico south into northern Colombia and Venezuela (figs. 219-220). The E. ophibolus group comprises six closely related species united by seven derived character states. Figure 221 shows that the distribution of all six species except $E$. schausi are largely allo- or parapatric. Little can be said of $E$. agkistrodon or $E$. schausi due to paucity of records. Erpetogomphus ophibolus occupies an area between two disjunct populations of $E$. constrictor. Further collecting may show E. constrictor to be symparric or parapatric with E. ophibolus. Aside from E. agkistrodon and E. schausi, the track of $E$. ophibolus was probably the first to separate from the E. ophibolus group, a hypothesis consistent with the results of my cladogram (fig. 216). The three remaining species, E. constrictor, E. tristani, and E. sabaleticus, are closely related sister taxa whose tracks are largely parapatric. Presumably their differentiation by allopatric speciation?) took place from a common ancestor which occupied part of the present range of all three species.

The tracks of the second monophyletic group comprises just two species, the relatively widespread $E$. :ancourand tha perpheral /. leprophis (fig. 222). The latter, known only from the holotype and allotype from Belize, is sympatric with E. ophibolus, but its genealogical affinities are unquestionably with $E$. eutai$\because$ fig. 216). Erpetogomphus leptophis probably arose :s a peripheral isolate from E. eutainia, or from a common ancestor, since it is a more apomorphic spe-
cies than E. eutainia.
The remaining 13 species comprise two monophyletic subgroups, one predominantly austral, the other more boreal. Their generalized tracks are shown in figs. 223-226. The E. elaps subgroup contains five species, whose somewhat limited distributions occupy the western and eastern lowlands of Mexico south along the mountainous areas into Costa Rica. Erpetogomphus elaps and E. elaphe are closely related sister species which are paraparric in Guatemala. A slight but consistent southern variant of $E$. elaps occupies the area between more northerly typical populations of $E$. elaps and $E$. elaphe (see discussion under $E$. elaps): it may be an incipient species. These two (or three?) species, like those for $E$. constrictor, E. tristani, and $E$. sabaleticus, probably arose from an $E$. elaps-like ancestor or from one another. Interpretations of the tracks of the remaining three species is problematical due to the uncertain position of $E$. liopeltis in my cladogram. It possesses characters common to $E$. bothrops and $E$. viperinus. No convincing apomorphy unites $E$. liopeltis with the sister taxa $E$. elaps and $E$. elaphe, so its genealogical relationship must remain in doubt. The two species with the greatest ranges ( $E$. elaps and $E$. bothrops) are broadly sympatric; E. liopeltis and $E$. viperinus are allopatric.

The E. crotalinus (s.s.) subgroup comprises the boreal group of eight species, with at least one representative occurring throughout most of the southern half of the United States. Their complex, mostly overlapping distributions (fig. 226) and questionable phylogeneric sequence in the cladogram preclude any meaningful discussion on their biogeography. Erpetogomphus boa and E. cophias stand out because of three synapomorphies. Their limited distribution and close proximity indicate speciation, possibly by allopatric means. Erpetogomphus sipedon and $E$. heterodon make up the only other allopatric species pair in this complex subgroup. Although distribution of $E$. beterodon and E. compositus overlap, I have seen no evidence of these two species collected together.

It is obvious that a more thorough knowledge of the phylogeny of the genus will be necessary before interpretation of biogeographical events is possible. Of particular interest will be a methodological interpretation of the biogeography of the genus, for then we may be able to determine which species tracks are the result of dispersal and gene flow or vicariant events.

## Characters

The best diagnostic characters for males are found in the caudal appendages, accessory genitalia, and overall body maculation. Most species are easily identified by the caudal appendages. The cerci (superior appendages) assume a wide variety of shapes (figs. 92-

Table 4. Distribution of species of New World Gomphidac. Number under each region = area of region in square miles; $\% / \%$ directly under area $=\%$ of total land area; $N=$ total number of species; number under each region = total number of species; number in parentheses () = percent of total number of species in genus. Phylogenetic classification is after Carle (1986). Genera are arranged alphabetically under each tribe.

| Genus | N | Canada/Alaska 4, 438,221 $\mathrm{mi}^{2}$ 29.5\% | $\begin{aligned} & \text { U.S.A. }{ }^{1} \\ & 3,022,261 \mathrm{mi}^{2} \\ & 20.2 \% \end{aligned}$ | $\begin{aligned} & \text { Mexico } \\ & 761,530 \mathrm{mi}^{2} \\ & 5.2 \% \end{aligned}$ | $\begin{aligned} & \text { Cent. Amer. }{ }^{2} \\ & 208,800 \mathrm{mi}^{2} \\ & 1.1 \% \end{aligned}$ | South Amer. $6,597,386 \mathrm{mi}$ 43.9\% |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Hageniinae |  |  |  |  |  |  |
| Hagenini | 1 | 100) | (100) |  |  |  |
| Octogomphinae |  |  |  |  |  |  |
|  |  |  |  |  |  |  |
| Hemigomphini | 3 |  |  |  |  | 3(100) |
| Octogomphini |  |  |  |  |  |  |
| Lanthus | 2 | 1 (50) | 2 (100) |  |  |  |
| Octogomphus | 1 | 1 (100) | 1 (100) | 1 (100) |  |  |
| Stylogomphus | 1 | 1 (100) | 1 (100) |  |  |  |
| Gomphinae |  |  |  |  |  |  |
| Gomphini |  |  |  |  |  |  |
| Arigomphus | 7 | 3 (43) | 7 (100) |  |  |  |
| Dromogomphus | 3 | 1 (33) | 3 (100) |  |  |  |
| Gomphus ${ }^{3}$ | 38 | 13 (34) | 38 (100) | 1 (3) |  |  |
| Stylurus | 13 |  | 12 (92) | 4 (31) |  |  |
| Epigomphinae |  |  |  |  |  |  |
| Epigomphini |  |  |  |  |  |  |
| Epigomphus | 26 | 6 (23) | 12 (46) | 10 (38) |  |  |
| Austrogomphinae |  |  |  |  |  |  |
| Archaeogomphini |  |  |  |  |  |  |
| Archaeogomphus | 5 |  |  | 1 (20) | 1 (20) | 5 (100) |
| Cyanogomphini |  |  |  |  |  |  |
| Agriogomphus | 4 |  |  | 1 (25) | 1(25) | 3 (75) |
| Cyanogomphus | 6 |  |  |  |  | $6(100)$ |
| Tibiagomphus | 2 |  |  |  |  | 2(100) |
| Onychogomphinae |  |  |  |  |  |  |
| Onychogomphini |  |  |  |  |  |  |
| Erpetogomphus | 21 |  | 5 (24) | 16 (76) | 10 (48) | 1 (5) |
| Ophiogomphus | 18 | 8 (44) | 18 (100) |  |  |  |
| Lindeniinae |  |  |  |  |  |  |
| Gomphoidini |  |  |  |  |  |  |
| Aphylla | $19^{4}$ |  | 3 (16) | 2 (11) | 3 (16) | 18 (95) |
| Gomphoides | 3 |  |  |  |  | 3 (100) |
| Idiogomphoides | 2 |  |  |  |  | 2 (100) |
| Peruviogomphus | 2 |  |  |  |  | 2 (100) |
| Phyllocycla | 31 |  |  | 3 (10) | 4 (13) | 27 (87) |
| Phyllogomphoides | 43 |  | 2 (5) | 12 (28) | 7 (17) | 27 (63) |
| Lindeniini |  |  |  |  |  |  |
| Cacoides | 1 |  |  |  |  | 1 (100) |
| Melanocacus | 2 |  |  |  |  | 2 (100) |
| Mitragomphus | 1 |  |  |  |  | 1 (100) |
| Progomphini |  |  |  |  |  |  |
| Progomphus | $59^{5}$ |  | 4 (7) | 9 (15) | 7 (12) | 52 (77) |
| Zonophorini |  |  |  |  |  |  |
| Desmogomphus | 2 |  |  |  | 1 (50) | 2 (100) |
| Diaphlebia | 2 |  |  |  |  | 2 (100) |
| Perigomphus | 1 |  |  |  | 1 (100) | 1 (100) |
| Zonophora | 10 |  |  |  |  | 10 (100) |
| TOTAL | 329 | 35 (10.6) | 97 (29.4) | 50 (15.2) | 47 (14.3) | 180 (54.7) |

${ }^{1}$ Excluding Hawaii; ${ }^{2}$ Belize, Guatemala, El Salvador, Honduras, Nicaragua, Costa Rica, Panama; ${ }^{3}$ Includes subgenera Gomphurus, Gomphus, Phanogomphus, Stenogomphurus; " One species, A. caraiba, is endemic to Hispaniola and Cuba, not included in N; ${ }^{5}$ Three species, $P$. integer, $P$. serenus, $P$. zephyrus, are endemic to the Greater Antilles, not included in N.
116), but they are never longer than segments 9 and 10 combined. All males of the E. ophibolus group, except for $E$. schausi (and possibly the unknown male of E. agkistrodon) have a dorsal appendage on the distal third of the cercus. The dorsal surface of the cercus may be straight as in $E$. elaps, strongly curved as in $E$. schausi and $E$. viperinus, angulate as in most species in the $E$. crotalinus group, or concave as in the $E$. boa and $E$. cophias group. The inferior third of the cercus forms a strong carina in E. cophias (weakly so in $E$. boa), in the E. crotalinus group, and in E. eutainia. In the lighter coloured species of the E. crotalinus group, the carina does not contrast with the rest of the appendage. A unique series of pebble-like structures characterizes the basal carina in E. eutainia. The inferior carina was used by Calvert (1905), although he wrongly placed E. eutainia among individuals lacking such a carina. Instead of a basal carina, the $E$. ophibolus group, $E$. bothrops and $E$. viperinus have an inferior carina on the distal third of the appendage. This structure is weak in E. elaps.

The epiprocts (inferior appendages) are not as varied as the cerci. In most of the $E$. ophibolus group, they are strongly curved so that their distal third is nearly parallel to the basal 0.30 . The epiproct of $E$. ophibolus is unique in having an anteriorly directed projection on its dorsal surface near the middle. The epiprocts are remarkably similar throughout the remaining groups: they form a gentle curve with the tips at approximately $90^{\circ}$ to the basal third of the appendage.

The penes of most species possess a semi-hyaline pair of flagella or cornuae best seen in dorsal view. The cornuae are considerably reduced in the $E$. ophibolus group (except for $E$. schausi). Their shape provides a useful character to differentiate among $E$. designatus, E. compositus, and E. Lampropeltis. The tip of each cornua is bluntly rounded in the $E$. crotalinus group (except for $E$. crotalinus). In all others, the tip forms a sharp point.

The spiny lateral lobes of the penis are well developed and frill-like in E. eutainia and E. leptophis, primarily knife-like or pointed in E. viperinus, E. liopeltis, E. bothrops, E. elaps, and E. elaphe, and a protruding semicircle in most members of the E. croralinus and E. cophias groups. When the penis is viewed laterally, the lateral lobe is on edge, because this structure rotates posteriorly with its flat portion placed mediodistally. The lateral lobes are poorly developed in E. constrictor, E. tristani, and E. sabaleticus (possibly also in E. agkistrodon), but those species and E. schausi have a unique ventral, arcuate prepuce.

The anterior hamules are always divided, but the posterior arm in the E. tristani group forms more of a shoulder than a branch. The bifurcation occupies the distal 0.25 of the hamule in the E. ophibolus, E. elaps,
and $E$. eutainia groups, the distal 0.50 in the $E$. crotalinus group, and the entire length of the appendage in E. boa and E. cophias.

The posterior hamules are always more prominent than the anterior pair. They are broadly triangular in the E. ophibolus group, but they possess an anterior shoulder in the E. crotalinus group. In E. eutainia and E. leptophis, the posterior hamule forms a small fin-ger-like appendage. All known species except $E$. designatus, E. elaphe, E. leptophis, and E. eutainia possess a black apical tooth. This distal tooth is rotated laterally in E. viperinus and in some $E$. bothrops.

The best structural characters for separating the females are those associated with the vertex, occiput, postocciput (note: my use of the term, 'postocciput' is not equivalent to the structure immediately surrounding the occipital foramen, but, instead refers to the medioposterior side of the head behind and below the occipital crest), and vulvar lamina. Calvert (1905) relied on the curvature of the posterior border of the occiput, but it is variable within species. I do not use it here. Better characters are the shapes of structures on the vertex. In $E$. constrictor, $E$. tristani, and $E$. sabaleticus, the median ocellus is placed within a deep longitudinal trough which accommodates the strongly curved epiproct of the male. In all other species may be found a transverse trough (figs. 145-146) or a pair of anterolateral pits (fig. 137) anterior to the median ocellus. These pits correlate with the shape of the tip of the male epiproct, and these depressions are also present on the vertex of males. Their shape is useful in distinguishing between the closely related pairs, $E$. viperinus and $E$. bothrops, and between $E$. compositus and E. lampropeltis. A complete postocellar ridge is present only in all members of the $E$. tristani group and E. leptophis. In all others, the middle part of the ridge is lacking, leaving only postocellar tubercles. Females of a few species have autapomorphic characters, such as the raised median area of the occiput of $E$. designatus (figs. 156-157), the posteriorly directed pointed protuberance on the occiput of $E$. leptophis (fig. 151) or the postoccipital depressions of E. cophias (fig. 165).

The vulvar laminae vary in the shape of the plates. The simple horizontal plates are bisected by a small V-shaped cleft in the E. ophibolus group. The plates are corrugated and more membranous in $E . b o a$ and E. cophias, often succumbing to much post mortem distortion. In the E. crotalinus group, each plate is a large, rectangular, planar structure which meets the other only at the anterior angle. The resulting median cleft is only a third or less the width of each plate; but the cleft is wider and U-shaped in E. designatus. A further widening of the cleft is evident in the E. elaps group. The plates form two small, widely separated, digit-like structures in E. eutainia and E. leptophis.

A Y-shaped postlamellar ridge is found posterior to the vulvar lamina. The posterior stem of the Y is absent in E. eutainia and E. leptophis, so that the ridge forms a semicircle (figs. 178-179).
I have found no structural differences between the females of $E$. heterodon and $E$. sipedon. Body pattern must be used to differentiate females of those species. Erpetogomphus tristani and E. sabaleticus, and E. elaphe and E. elaps are also closely related pairs, and their females are distinguishable only by locality.
In well-preserved specimens, body maculation provides useful characters. Dark thoracic maculation predominates in the E. tristani group, in E. leptophis, and in southernmost specimens of E. eutainia. All of those have a well-defined dark stripe bordering the posterior margin of the metepimeron. Erpetogomphus cophias, E. crotalinus, and most $E$. elaps have nearly immaculate body patterns. Thoracic maculation is usually stable, but there is variability in E. lampropeltis, $E$. compositus, $E$. designatus, $E$. elaps, and $E$. eutainia. Two formerly named species, E. natrix and $E$. coluber, I consider subspecies of $E$. lampropeltis and $E$. compositus, respectively: I have found no morphological characters which separate those forms. Thoracic patterns in E. elaps are often nearly absent (fig. 13), and more extensive dark markings characterize populations of $E$. eutainia in the north, less extensive in the south (figs. 7-9).
Abdominal patterns are characterized by a combination of light and dark banding patterns separated dorsally by pale longitudinal stripes. The least amount of black occurs in E. cophias and E. crotalinus, the most in E. agkistrodon.
A sharp, prominent, middorsal posterior spine on abdominal segments 8 and 9 is found in many males of $E$. ophibolus, E. bothrops, E. liopeltis, and E. viperinus. The spine is variable and cannot be used reliably to distinguish among species. For example, many $E$. bothrops have no indication of a middorsal spine, but it is well developed in others. The spines are also found on abdominal segment 9 in some females of the $E$. ophibolus group.

## Erpetogomphus species groups

I divide the 21 species into five groups. I believe the unknown male of one species ( $E$. agkistrodon) will follow those characters listed for that sex discussed below.

1. E. ophibolus group (= subgenus Erpetocyclops Carle, 1992). Males (probably so for E. agkistrodon, when discovered): With a distinct curved prepuce, posterior hamule forming a triangle culminating in a sharp black tooth. Females: Vulvar lamina simple, small, relatively undifferentiated postocellar ridge complete, though curved in E. tristani, E. sabaleticus, and $E$. constrictor. Six species: E. tristani, E. sabaleti-
cus, E. constrictor, E. ophibolus, E. agkistrodon, E. schausi.
2. E. eutainia group (= subgenus Calogomphus Carle, 1992). Males: Prepuce small, obscured by lateral lobes of penis; lateral lobes of penis strongly developed, forming a spiny frill; posterior hamule fin-ger-like, lacking an apical tooth. Reduced prepuce present. Females: Vulvar lamina small, finger-like, widely separated; post-laminar ridge semicircular, not Y- shaped. Two species: E. leptophis, E. eutainia.
3. E. elaps group. Males: Lateral lobe of penis knifelike. Females: Vulvar lamina connected anteriorly, but separated by a cleft 1.0 or more the width of each plate. Five species: E. bothrops, E. liopeltis, E. viperinus, E. elaphe, E. elaps.
4. E. crotalinus group. Males: Superior surface of cercus angulate (except for E. compositus); anterior hamule divided at distal half. Females: Vulvar lamina large, planar, rectangular, meeting at anterior margins only, resultant cleft one third the width of each plate (except for $E$. designatus). Six species: $E$. crotalinus, $E$. lampropeltis, E. compositus, E. heterodon, E. sipedon, E. designatus.
5. E. cophias group. Males: Anterior hamule talonlike, divided along its full length; posterior hamule broadly spatulate, surmounted by a small, black, curved anteapical tooth. Females: Vulvar lamina diagonally corrugated; distal and lateral margins with a strongly raised semicircular ridge. Two species: $E$. boa, E. cophias.

## Systematic part

## Introduction

Under each species, I have endeavoured to include a complete synonymy (some minor references may be lacking for $E$. designatus, a common species in the eastern United States), a full description, type data, diagnosis, remarks, biology, distribution, and material examined. For new species, I have designated allotypes when the opposite sex was available, but I have not established allotypes for syntypic material. Under comments, I have quoted older authors (Ris, Selys, Hagen), when this information contributes to the history of the species. For E. agkistrodon, I include a discussion of what the male should be like, based on my cladogram. Unless otherwise stated, measurements are taken from 20 males and 20 females, when available, and from as many localities as possible. Abdominal measurements include caudal appendages. Full locality data are given for all species, except for the following species where abundant material was available: $E$. eutainia, $E$. elaps, $E$. viperinus, $E$. designatus, E. l. lampropeltis, E. l. natrix, E. crotalinus, and $E$. compositus. For these species, I have listed states, counties, brief data, and repository. Full locality
data for these specimens are available from me. Maps represent distribution records from collections. For some common and well known species (such as $E$. borhrops, $E$. compositus, $E$. crotalinus, $E$. designatus), I include records from published sources I consider reliable.
Abbreviations for figure legends and synonymies are: abd. = illustration(s) of abdomen; app. = illustrations(s) of caudal appendages; cat. $=$ listed in catalogue, descr. = description of male, female, larva, or all, may include keys; distr. = documents or mentions distribution of taxon; ept. $=$ illustration(s) of epiproct; sep. = [page number of] separate (see Cowley (1937) for pagination of reprint of Selysian monographs and synopses of Odonata); thx. = illustration(s) of thorax; vl. = illustration(s) of vulvar laminae.
Abbreviations in descriptions are as follows: anx = antenodal crossveins, pnx = postnodal crossveins, cs = crossveins.
New names proposed in this paper follow the ophidian example set by Selys, Hagen, Calvert, and Williamson. All should be considered as nouns in apposition.
Most illustrations are from type material, which I believe will aid further researchers when describing new species. All illustrations were executed with the aid of a camera lucida with a Wild M-8 stereoscopic microscope.

## Erpetogomphus Hagen in Selys, 1858

Erpetogomphus Selys, 1858 (Type species Ophiogomphus crotalinus [Hagen in Selys], 1854; designated by Kirby 1890: 61). - Selys 1858: 329 ( 69 sep.) (characters of genus); Selys 1859: 535 ( 9 sep.) (addition of generic characters); Hagen 1861: 98 (brief descr. of genus); Walsh 1863: 253 (comparison with other American gomphid genera); Brauer 1868: 372 (in key to genera of Gomphidae); Karsch 1890: 371 (mentioned in Gomphidae classification); Calvert 1905: 147 (key to Middle American Gomphidae); Calvert 1909; 468 (distr.); Calvert 1912a: 289 (distr. of genus); Kennedy 1917a: 544 (comments on larvae); Tillyard 1917: 296 (distr); Ris 1921: 343 (comparison with Mesogomphus [= Paragomphus]); Garman 1927: 125 (note on genus); Needham and Heywood 1929: 78 (descr.); Byers 1930: 52 (descr.); Cowley 1934: 241 (nomenclature of Erpetogomphus); Tinkham 1934: 218 (comments on distr. in Tex.); Fraser 1940: 544 (possible affinities with Onychogomphus based on penis morphology); Ncedham 1941: 240 (comments on larva); Needham 1944: 172 (verification table, larva); Wright and Peterson 1944: 152 (in key to larvae of nearctic Gomphidae); Needham \& Westfall 1955: 139 (treatment of genus); Pritchard \& Smith 1956: 114 (key to larvae of nearctic Gomphidae, key to genera of Gomphidae); Ferguson-Beatty 1956: 369 (proventriculus of larva); Musser 1962: 13 (diagnosis of larvae); Gloyd 1963: 147 (molar structure of adult); Westfall 1984: 158 (in key to larvae of nearctic Gomphidae); Chao 1984:79
(uncertainty of Erpetogomphus in subfamily Onychogomphinae); Carle 1986: 313 (characters of genus within Onychogomphini); Belle 1988: 99 (in key to neotropical genera of Gomphidae); Bridges 1991: II. 17 (cat.); Novelo-G. \& Peña-O. 1991: 129 (origin of Erpetogomphus in Mexico); Gonzalez-S. \& Novelo-G. 1991: 97 (adults in Mexico); Novelo-G. \& Gonzalez-S. 1991: 150 (larvae in Mexico); Belle \& Quintero 1992: 93 (in key to adults and larvae of Panamanian Gomphidae); Carle 1992: 148 (key, Erpetocyclops subg. n. [type $=E$. ophibolus Calvert], Calogomphus subg. n. [type $=$ E. eutainia Calvert]).
Herpetogomphus Walsh, 1862: 388 (unjustified emendation of Erpetogomphus). - Hagen 1875a: 42 (cat.); Selys 1879: 63 (2 sep.) (modification of generic characters); Kirby 1890: 60 (cat.); Carpenter 1897: 452 (distr.); Needham 1897: 166 (in key to nearctic Gomphidae); Needham 1899: 234 (venation); Calvert 1899: 386 (comments on Selys' classification); Needham \& Hart 1901: 53 (in key to larvae of North American Gomphidae); Förster 1914: 73 (comparison with Ammogomphus); Seemann 1927: 19 (key to larvae, adults of western U. S. Gomphidae); Needham 1940: 389 (key, verification table).

Medium sized to small neotropical Gomphidae of slender build.

Head. - Eyes widely separated, characters of the mouth parts, clypeus, frons typical of the Gomphidae. Vertex variable, often interspecifically and, in the E. ophibolus group, sexually dimorphic. Anterior margin of vertex with a transverse trough, or a pair of transverse pits anterior or anterolateral to median ocellus, median ocellus anterior to lateral ocelli, or in some members of the E. ophibolus group, this structure at posteriormost part of deep, longitudinal trough, thus causing median ocellus to be posterior to lateral ocelli (figs. 145-146); median surface of vertex at vicinity of lateral ocelli and postoccipital tubercle mostly planar, but members of $E$. ophibolus group with a complete postocellar ridge, which may be bilobed (E. agkistrodon, fig. 148) or, in remainder of group, vestigial; occiput highly variable according to species and sex, E. ophibolus group with occiput small, semicircular, in other groups roughly trapezoidal, with posterior part broader than anterior part, its dorsal surface roughly planar to nearly vertically inclined, some species with slight to strongly tumid area medially, one species (female of $E$. leptophis) with a posteriorly projected medial spine; rest of occiput variable, semicircular, linear, weakly to strongly notched medially, or sinuate; condition of crest often variable within a few species; postocciput variable, most in the E. ophibolus group convexly arcuate, in others slightly convex to linear, postocciput mostly not visible from above, but conspicuously so in others (e.g., female of E. compositus); lateral margins of one species (females of $E$. cophias) with transverse depressions.

Thorax. - Prothorax and synthorax typical of the

Gomphidae, but with great differences in maculation and colouration due to species and age differences. Pale colour pale green, apple green to vivid blue green, usually lighter ventrally and often with a slight dusting of white pruinosity ventrally and around coxae. Synthorax immaculate to heavily patterned with full complement of thoracic stripes. Legs moderately short, with femora slightly swollen and usually pale basally, with dark brown to black distally; tibiae in almost all species black, prothoracic tibiae of males with small keel occupying distal 0.15 of mesal ventral margin.
Wings (figs. 227-230) variable inter- and intraspecifically. Fore wing moderately narrow, with 11-21 antenodal crossveins, $5-16$ postnodal crossveins, no subcostal crossvein; pterostigma prominent, about 4 times as long as wide, widest medially with its posterior margin slightly convex, surmounting 4-8 crossveins, the proximal a brace vein; anterior margin of costa in many species pale, becoming dark at pterostigma; arculus at vicinity of second antenodal, the sectors not stalked, separated at base by distance greater than width of each sector; supratriangular, triangular, and subtriangular crossveins absent; paranal cells variable in number, often numbering 5-7, smaller species usually with zero or few supplementary marginal cells behind paranal cells, other species with irregular row of up to 6-7 supplementary cells; no apical or medial planates, discoidal field of two rows. Hind wing with 7-14 antenodal crossveins, 7-14 postnodal crossveins; anal triangle of 2-4 cells, often with one of them in the latter condition small; anal loop absent (some specimens with a vestigial anal loop of 3 cells, but this condition rare); one paranal cell at Y interspace, two paranal cells at X interspace. Wing colouration entirely hyaline or flavescent at bases in some species. Hind wing length $21-36 \mathrm{~mm}$.

Abdomen cylindrical, narrowest medially in males, transverse carinae on segments 3-7; auricles well developed in male, inner margin denticulate; auricle vestigial in female; male posterior margin of tergites 8 and especially 9 in a few species prolonged into a medial point, but this condition variable intraspecifically; segments 8-9 moderately clubbed, with moderate foliate extensions; female with apical abdominal segments hardly widened, foliar expansions small. Colour pattern and maculation variable between species and within sexes; all species with a pale green to pale orange middorsal stripe, sometimes extending full length of segment, but progressively less so on more posterior segments; dark brown to black dorsolateral stripe present on first 7 segments, usually constricted or separated medially, giving most of abdomen a serial dark and pale appearance; posterior part of segment 7 , all of segments $8-10$ tan to dark red brown, especially middorsally and ventrolaterally.

Length of abdomen including appendages: ठ: 29-40 mm , ㅇ: : $29-41 \mathrm{~mm}$.

Abdominal appendages of male with cerci of great diversity according to species, but these structures always as long as segment 10 , cercus usually gently decumbent at apical 0.30 ; rarely with a dorsal appendage. Ventral carina present basally, or distally, or absent, depending on species, epiprocts divided at base but contiguous along most of their length, gently to strongly curved at distal 0.50 to 0.30 , depending on species; this structure lacking any accessory spines, or protuberances (except for E. ophibolus, fig. 95); tips of epiprocts assuming a wide variety of shapes and forms, and often variable intraspecifically (e.g., E. bothrops, figs. 127-128).

Vulvar lamina of female variable interspecifically, but never more than 0.50 length of segment 9 , vulvar plates small, digit-like to forming large, contiguous, triangular or quadrangular plates, a moderate to large V- or U-shaped notch medially; postlamellar ridge always present, semicircular in only two species ( $E$. eutainia, E. leptophis), in all others forming an inverted Y ; stem of Y originating before or often at level of vulvar plates, some species with a well defined semicircular depression on each side of posterior stem.

Male genitalia variable among species: anterior lamina not prominent; anterior hamule smaller than posterior hamule, usually with the distal end hidden from view by posterior lamina; anterior lamina dark, divided at various heights according to species, but with posterior branch usually less conspicuous, smaller, and anterior branch more prominent; posterior lamina usually pale, linear in two species (E. eutainia, E. leptophis); in all others with a moderately wider base, its tip usually adorned with an apical or anteapical hook directed cephalad or cephalolaterad. Penis with basal (first) segment prominent, with a semicircular rim, especially so laterally, posteriorly with a pair of prominent lobes, or these lobes vestigial; second segment moderately long, third short, with a pair of small, nipple-like sclerotizations dorsally in two species (E. eutainia, E. leptophis); fourth segment highly variable specifically, with or without a prepuce, most species with a well defined lateral lobe; in most of the E. ophibolus group, lateral lobe small and pointed; membranous hood short to long and overlapping; a pair of flattened membranous cornuae of various lengths and shapes present on apical part of segment; in most of the E. ophibolus group, cornuae absent or vestigial; penis guard short, quadrate in cross-section.

A more complete discussion of the male and female genitalia is given under discussions of the species group accounts and in the cladistics section.

Keys
The keys should work for most specimens, but the worker may have difficulty when trying to determine poorly preserved material. I have used morphological characters for both sexes wherever possible, but some species pairs (e.g., females of $E$. beserodon and E. sipedon, or E. tristani and E. sabalericus) show little or no morphological differentiation. For them, I have had to rely on colour pattern. Abdomens of some females, particularly those laterally compressed, may need to be relaxed so that the vulvar lamina can be seen.

Key to Erpetogomphus males

1. Cercus with a prominent superior tooth at about 0.75 of appendage length (figs. 92-94)

2
Cercus with dorsal surface convexly angulate (fig. 112), smoothly curved (fig. 103), straight (fig. 100), or with a concavity (fig. 115)
.. 5
2(1). Epiproct smoothly curved, with tip pointing dorsally or posterodorsally, a small, sharp, anteriorly directed superior tooth at middle of epiproct (fig. 95). Southern Mexico, Belize .ophibolus Epiproct strongly curved so that distal 0.30 is parallel to basal 0.30 , tip of epiproct pointing anteriorly, superior surface of epiproct with no tooth .. 3
3(2). Ventral margin of cercus smoothly concave (fig. 92). Northeastern Mexico to Costa Rica constrictor

- Ventral margin with a large ventral tubercle at 0.50-0.75 length of cercus (figs. 93-94). Costa Rica south to Colombia and Venezuela
4.3). In lateral view, superior tooth of cercus as long as rest of appendage, so that appendage appears to end in two equal branches (fig. 93). Panama, Colombia, Venezuela ..... sabaleticus In lateral view, superior tooth of cercus less than 0.50 as long as remainder of appendage (fig. 94). Costa Rica, northern Panama tristani
$=1$. Metathoracic tibiae either entirely yellow externally (fig. 169), or yellow with median longitudinal black line. Western Mexico, southeastern Arizona $\qquad$ crotalinus Metathoracic tibiae entirely brown or black externally; never with any yellow (e.g, figs. 168.170)

6 ~. Basal 0.25 to 0.30 of lower margin of cercus with a distinct longitudinal carina which
may end in a small ventral tooth (e.g., E. cophias, fig. 116)

7
Basal part of cercus with no ventral carina, though a small tubercle may be present at basal $0.15-0.20$ of appendage (e.g., figs. 99, 102)

14
7 (6). With a dark stripe bordering posterior margin of metepimeron (figs. 8-9). Southern Mexico, Belize through Costa Rica eutainia (in part) Without a dark stripe on posterior margin of metepimeron

8
8 (7). In lateral view, ventral margin of cercus with carina terminating in a distinct ventral tooth at basal 0.30 of appendage (fig. 116); thorax almost immaculate, at most barely an indication of a dark antehumeral stripe (fig. 27). Michoacan and Morelos states of western Mexico
cophias

- Ventral margin of cercus with longitudinal carina not terminating in a ventral tooth; thorax with well-defined dark antehumeral stripe, though it may be an elongated, isolated spot (e.g., fig. 23)

9
9 (8). Dorsal surface of cercus distinctly angulate in lateral view (figs. 110, 112)

10
Dorsal surface of cercus smoothly curved (fig. 112)

13
10 (9). Sides of thorax almost entirely green, with only a small, ill-defined dark humeral stripe; second lateral stripe incomplete, dark only below metathoracic spiracle, or absent; and third lateral stripe present only on upper 0.30 of suture (fig. 23), or absent (fig. 19)...
.11

- Sides of thorax green with usual complement of dark thoracic stripes; second lateral stripe may be incomplete, but humeral and third lateral stripes always present (figs. 17, 20-21)

12
11 (10). Dark antehumeral stripe incomplete, not touching humeral stripe (fig. 23); tip of cercus acute, but not gradually narrowing posteriorly (figs. 110-111); cornuae of penis without mesal lobes (fig. 87). New Mexico, western Texas, northern Mexico

## heterodon

Dark antehumeral stripe complete, usually connected to humeral stripe (fig. 19); tip of cercus drawn out into a narrow cylindrical point (figs. 105-106); cornuae with mesal lobes (fig. 83). Durango, Jalisco, Morelos, and Puebla states of central Mexico
sipedon
12 (10). Tip of cercus strongly acuminate (fig. 104), dark antehumeral stripe not connected to
collar (fig. 17) (except in some Mexican individuals), base of wings flavescent (except in some Mexican individuals, fig. 18), median area of occiput with a strongly raised tubercle (fig. 156). Eastern U.S. to Arizona, northern Mexico
designatus

- Tip of cercus blunt, not acuminate (figs. 107-108); dark antehumeral stripe always connected to collar (ssp. natrix, fig. 21), or largely connected to humeral stripe (ssp. lampropeltis, fig. 20); base of wings hyaline, median area of occiput only slightly raised (fig. 160). Southern California, Arizona, New Mexico, western Texas, Baja California, Durango state, Mexico lampropeltis
13 (9). Ventral carina at base of cercus black, contrasting with remainder of appendage; this carina usually denticulate; thorax blue-green in life; sides of posterior hamules parallel, linear (fig. 52). Southern Texas to southern Mexico eutainia (in part) Ventral carina at base of cercus pale, the same colour as rest of appendage, this carina smooth; thorax pale green in life; sides of posterior hamules converging toward tip, not linear (fig. 66). Southwestern United States, Baja California $\qquad$ compositus
14 (6). Distal 0.5 of cercus concave dorsally, this area covered with thick, long bristles (fig. 115). Southern Veracruz state, Mexico ....... boa
- Distal 0.5 of cercus straight or convex (e.g., figs. 99, 103)15

15 (14). Ventral margin of cercus straight (figs. 99, 100 ) or forming a gentle concave curve (fig. 98) so that appendage appears linear ..... 16

- Apical 0.30 of ventral margin of cercus curved downward .................................. 18
16 (15). Thorax entirely yellow-green (fig. 13), or with only an antehumeral and faint indication of a dark middorsal and antehumeral stripe (figs. 11-12); face entirely pale. Mexico south to Costa Rica. 17
- Thorax blue-green with full complement of dark thoracic stripes (fig. 10); face bluegreen with extensive dark brown frontoclypeal stripe. Belize
leptophis
17 (16). Distal branch of anterior hamule thick, as high as gap separating the two branches (fig. 54); posterior hamules with no distal tooth (fig. 54). Guatemala south to Costa Rica .... ........................................................... elaphe Distal branch of anterior hamule narrow, not as high as gap separating the two branches (figs. 55-56); posterior hamules
with a distal tooth (fig. 55). Mexico to Guatemala
elaps
18(15). Thorax blue-green, with full complement of dark stripes, including second lateral stripe and stripe on posterior margin of metepimeron (fig. 6); face (fig. 142) predominantly dark, with a well defined blue green spot on frons. Guatemala, Costa Rica schausi
- Thorax blue-green or yellow-green; sides lacking complete dark second lateral stripe (except in some viperinus, incomplete in some bothrops) and without dark stripe on posterior margin of metepimeron; face predominantly pale, with dark areas limited to sutures

19
19(18). Tip of epiproct in posterior view terminating in a blunt point (fig. 129); southern Veracruz state, Mexico. .viperinus

- Tip of epiproct in posterior view spatulate (fig. 126), broadly truncate (fig. 127), or bidentate (fig. 128)

20
20 (19). Larger species (hind wing 30-31 mm); epiprocts in lateral view curved, extending 0.50 or less the length of cerci (fig. 101), epiprocts in posterior view truncate or slightly bidentate; dark antehumeral and humeral stripes combined or nearly so (fig. 14); posterior hamular tooth prominent, directed laterocephalad or cephalad. Nuevo Leon, Michoacán, San Luis Potosí states, Mexico. (fig. 199) liopeltis Smaller species (hind wing 21-28 mm); epiprocts in lateral view curved, extending 0.75-0.90 the length of cerci (fig. 102), epiprocts in posterior view bidentate or truncate (fig. 128); dark antehumeral and humeral stripes separate (fig. 15); posterior hamular tooth not as prominent, more strongly directed cephalad. Tamaulipas, Mexico, south to El Salvador (fig. 199)
bothrops

## Key to Erpetogomphus females

1. Medial area of occiput with a posteriorly pointed protuberance (fig. 151) ... Leptophis

- Medial area of occiput variously shaped, but never with a posteriorly directed protuberance2

2 (1). Occiput with a shallow oval depression on each side posterior to elevated rim (fig. 165). Michoacán and Morelos, Mexico .... cophias

- Occipital area posterior to elevated rim entire, with no shallow depressions 3
3 (2). Outer surfaces of metathoracic tibiae entirely yellow, or yellow with a median longitudinal black line (fig. 169). Western Mexico,
southeastern Arizona ..... crotalinus
Outer surfaces of metathoracic tibiae entire ly black (e.g., figs. 168, 170) ..... 4
4.3). Median ocellus posterior to lateral ocelli, theformer lying within a deep longitudinaltrough (figs. 145-146)5
Median ocellus at level of or anterior to lat-eral ocelli, no longitudinal trough6
5 (4). Dorsal surface of vertex with sides of troughroughly parallel, or only slightly converginganteriorly (fig. 145). Northeastern Mexico,south to Costa Rica
$\qquad$ constrictor
Dorsal surface of vertex with sides of troughstrongly converging anteriorly ......(fig. 146)Costa Rica, western Panama
$\qquad$ tristani
Central Panama, Colombia, Venezuelasabaleticus
6 (4). Vertex with transverse ocellar ridge bilobedbehind median ocellus (figs. 148-149) ..... 7
Vertex with transverse ocellar ridge entire(fig. 147), low and almost vestigial (fig.150 ), or absent, its lateral ends forming ovaltubercles posterior to lateral ocelli (e.g., fig.150)8
- (6). Occiput transversally narrow, forming ashallow semicircle (fig. 148). Jalapa, Mexicoagkistrodon
Occiput wide, forming a full semicircle (fig.149). Guatemala, Costa Rica .......... schausi
8 (6). Vertex with transverse ocellar ridge entire, prominent, and concave posteriorly (fig. 147). Southern Veracruz, Mexico, Belize .... ophibolus
- Vertex with median area bare, or at most only slightly raised (e.g., fig. 150) 9
9 (8). Vulvar lamina followed on segment 9 by distinct and prominent semicircular ridge, never with a posteriorly directed arm (fig. 178). Southern Texas, eastern Mexico, Belize to Guatemala and Costa Rica $\qquad$ eutainia Vulvar lamina followed on segment 9 by a Y-shaped ridge (e.g., fig. 195)10
10 (9). Vulvar lamina with each plate strongly cor-rugated diagonally, distal and lateral mar-gins of each plate forming a strongly raisedsemicircular ridge (figs. 195-196); a mediantumid swelling on postocciput immediatelybehind occipital ridge (fig. 167) [includes li-opeltis, but properly goes to $10^{\prime}$ ]. SouthernVeracruz, Méexico ................................. boa
Vulvar lamina with each plate entire (fig.
185) or bisected diagonally by only one
groove (fig. 186); distal and lateral margins
of each plate planar, not forming a strongly
raised semicircular ridge; postocciput not
differentiated, evenly curved (fig. 166) [tu-
mid in liopeltis] 11
11 (10). Median surface of occiput with a strongly raised tubercle (fig. 156). Eastern United States west to Arizona, northern Mexico ..... designatus
- Median surface of occiput planar or at most slightly raised (fig. 152) 12
12 (11). Lobes of vulvar lamina separated by an almost U-shaped interval 3 to 4 times as wide as either lobe (figs. 180-181)
Mexico south to Guatemala
elaps
Guatemala to Costa Rica .................. elaphe
- Lobes of vulvar lamina separated by a triangular or semicircular interval 0.5 to 1.0 the width of each lobe (e.g., figs. 182, 187) ... 13
13 (12). With a second complete dark lateral stripe on thorax (e.g., figs. 20-21, 24) 14
- Second dark lateral stripe completely lacking or vestigial, at most extending from base to just above metathoracic spiracle (e.g., fig. 36)

15
14 (13). Occiput in dorsal view narrow, its width less than width between median ocellus and occiput; postoccipital area easily visible (fig. 163); base of wings with flavescent infusion between Sc and R at least up to first anx and cubitoanal area; top of abdomen with light areas ivory or with a tinge of orange, usually not different from white on sides. Southwestern United States, Baja California compositus Occiput in dorsal view wide, its width almost equal to width between median ocellus and occiput; postoccipital area not visible (fig. 160); base of wings hyaline; top of abdomen with light areas much darker than white on sides. Southwestern United States, Baja California, Durango state, Mexico
lampropeltis
15 (13). Occiput narrow, vestigial, with only a small medially arched area (fig. 153); post-occiput with a median tumid area (as in fig. 167). Nuevo Leon, Michoacan, San Luis Potosí states, Mexico
liopeltis

- Occiput not especially narrow, extending full width (e.g., figs. 154-155, 159, 162); post-occiput planar or, at most, only slightly convex medially (as in fig. 166)

16
16 (15). Stem of postlamellar ridge with a well defined circular or oval depression on either side; junction of Y of postlamellar ridge usually posterior to hind margin of vulvar laminar plates (fig. 183) 17 Stem of postlamellar ridge with only a slight depression or, more often, without any circular or oval depression on either side; junc-
tion of Y of postlamellar ridge at or anterior to hind margin of vulvar laminar plates (figs. 187-188, 193) 18
17 (16). A deep pit at anterior margin of frons anterolateral to median ocellus (fig. 155); cleft between vulvar laminar plates wide, forming an obtuse arc greater than $100^{\circ}$; dark humeral and antehumeral stripes combined or nearly so (fig. 16); larger species (hind wing $31-34 \mathrm{~mm}$ ). Southern Veracruz, Mexico .... ..viperinus

- Anterior margin immediately anterior to median ocellus forming a narrow V-shaped trough with base of postfrons (fig. 154); cleft between vulvar laminar plates narrow, forming a V - shaped notch of not more than $90^{\circ}$ (fig. 183); dark humeral and antehumeral stripes separate (fig. 15); smaller species (hind wing $17-31 \mathrm{~mm}$ ). Tamaulipas, Mexico, south through El Salvador
bothrops
18 (16). Dark antehumeral stripe complete, usually connected at its upper end to narrow humeral stripe (fig. 19). Jalisco, Morelos, and Puebla, Mexico sipedon Dark antehumeral stripe vestigial, forming an isolated elongate spot, not connected to vestigial humeral stripe (fig. 23). Southwestern United States, northern Mexico ...................................... heterodon


## Key to Erpetogomphus males based primarily on characters of abdominal segments 2 and 3

I include this key for male specimens lacking the posterior abdominal segments. The key should allow a high probability of identification, though some species (E. ophibolus, E. sabaleticus, E. tristani, E. constrictor, for example) appear inseparable by penis structure and are distinguished only by the caudal appendages. For those species, I have added other somatic characters and/or geographic data with which to separate them. The key does not include $E$. agkistrodon, the male of which is unknown.

1. Ental surface of penis with a prepuce (e.g., fig. 76), this structure reduced and partially hidden by frill-like lateral lobe in leptophis (fig. 76) and eutainia (fig. 75); thoracic pale colouration in life deep blue green, dark thoracic stripes always present on sides (e.g., fig. 10)

## .2

- Ental surface of penis lacking a prepuce (e.g., fig. 90); thoracic pale colour in life yellow green (except for viperinus and probably liopeltis); dark lateral thoracic stripes often (but not always - figs. 16,18 ) reduced or absent

2 (1). Posterior hamule digit-shaped (figs. 52-53), its tip broadly rounded; anterior hamule divided at upper 0.30 to 0.25 , its posterior branch well defined, its tip almost meeting larger anterior branch (figs. 52-53); lateral lobe of penis composed of large spinulose frill (figs. 52-53); cornuae well developed, their apices pointed $\qquad$

- Posterior hamule triangular, its tip armed with a spine (fig. 47); anterior hamule with only a posterior shoulder (e.g., fig. 47); lateral lobe of penis reduced to a small, semicircular, spinulose frill (fig. 74) or vestigial (fig. 70); cornuae vestigial (fig. 74) or present (fig. 70)
3 (2). Membranous hood of penis about 3 times as long as wide, apices overlapping (fig. 76) [Note: this character is based on the only known specimen, the holotype; other specimens may show variability similar to eutainid.]; dark second and third lateral stripes connected at upper ends, forming an isolated pale spot below subalar carina; dark lateral stripe along posterior margin of metepimeron well defined (fig. 10); caudal appendages as in fig. 98; Belize ..... leptophis Membranous hood of penis reduced, less than 2 times as long as wide, though apices may overlap (fig. 75); dark second and third lateral thoracic stripes not connecting (fig. 7-9); dark lateral stripe along posterior margin of metepimeron absent (fig. 7) or present primarily in populations of central Mexico (fig. 8) south through Costa Rica (fig. 9); caudal appendages as in fig. 97 ......

4 (2). Cornuae of penis present; their tips divergent (fig. 74); lateral lobe a small, circular, spinulose frill (fig. 74); Guatemala and Costa Rica ..................................... schausi Cornuae vestigial (fig. 71); lateral lobe vestigial, forming a small, acute point (fig. 71) ..

5 (4). Appendages as in fig. 93; Central Panama south to Colombia and Venezuela sabaleticus Mexico south through Costa Rica and northern Panama 6
6 (5). Epiproct smoothly curved, with tip pointing dorsally or posterodorsally, a small, sharp, anteriorly directed superior tooth at middle of epiproct (fig. 95). Southern Mexico, Belize ............................................ ophibolus Epiproct strongly curved so that distal 0.30 is parallel to basal 0.30 , tip of epiproct
pointing anteriorly, superior surface of epiproct with no tooth 7
-(6). Appendages as in fig. 92; Mexico south through Costa Rica
.................... constrictor Appendages as in fig. 94; northern Costa Rica through northern Panama ..... tristani
8(1). Anterior hamule divided at base; its branches resembling a claw; posterior hamule broadly spatulate (figs. 68-69) 9 Anterior hamule divided at lower 0.30 to upper 0.25 posterior arm short and truncate or acute; posterior hamule broadly triangular 10
9(8). Dark antehumeral and humeral stripes united, forming a broad, dark stripe over first lateral suture (fig. 26); southern Veracruz, Mexico
boa

- Thorax almost entirely pale, with only a vestigial antehumeral stripe (fig. 27); Michoacán and Morelos states of Mexico ... cophias
10 (8). Lateral lobe of penis linear, at least 3 times longer than wide; its posterior margin spinulose (figs. 77-78, 82, etc.) 11 Lateral lobe of penis round, about as long as wide, its margin spinulose (fig. 86) ........ 16
11 (10). Lateral lobe of penis almost recumbent against ental margin of fourth segment (fig. 82); anterior hamule divided at distal 0.5 of segment; posterior hamule lacking a distal tooth (fig. 60); posterior margin of first segment (peduncle) of penis strongly cleft, each lobe on either side of cleft protruding well beyond posterior margin of segment; southern United States west to Arizona, northern Mexico ...................................... designatus Lateral lobe of penis prominent, directed almost perpendicularly to ental margin of fourth segment; anterior hamule divided at distal 0.30 of segment, posterior hamule with a distal tooth (except for elaphe from Guatemala south through Costa Rica); posterior margin of first segment of penis weakly bilobed, lobes on either side of emarginarion only slightly extending beyond posterior margin of segment; northern Mexico south through Costa Rica ......... 12
12 111). Distal branch of anterior hamule thick, as tall (high) as gap separating the two branches (fig. 54); posterior hamule with no distal tooth (fig. 54); Guatemala south to Costa Rica elaphe Distal branch of anterior hamule narrow, not as tall (high) as gap separating the two branches (figs. 55-56); posterior hamule with a distal tooth 13

13 (12). Distal tooth of posterior hamule bent anteriorly so that it lies in same direction as the planar surface of hamule (figs. 55-56); Mexico south to Guatemala $\qquad$ elaps

- Distal tooth of posterior hamule bent laterally or anterolaterally so that it points distally and perpendicularly from planar surface of hamule (fig. 59) [Note: This condition appears universally only in $E$. viperinus, but variation in this condition occurs in $E$. bothrops and E. liopeltis; some specimens of these last two species may not be separable from E. elaps by this character]; Mexico south through Guatemala

14
14 (13). Base of postfrons medially with a pair of deep pits, each anterolateral to median ocellus (fig. 137). Southern Veracruz, Mexico .. viperinus

- Base of postfrons medially with a transverse arcuate trough immediately anterior to median ocellus (fig. 136)

15
15 (14). Smaller species (hind wing 21-28 mm); dark antehumeral and humeral stripes separate (fig. 15); posterior hamule with apical tooth shorter, less prominent, more strongly curved laterally over tip of hamule. Tamaulipas, Mexico, south through El Salvador (fig. 199)
bothrops

- Larger species (hind wing 30-31 mm); dark antehumeral and humeral stripes connected (fig. 14); posterior hamule with apical tooth longer, more linear, not as strongly curved laterally over tip of hamule. Nuevo Leon, Michoacan, San Luis Potosí, Mexico
liopeltis
16 (10). Cornuae of penis with tips long, narrow, and attenuate; medial lobe present (fig. 86); extensor surfaces of tibiae yellow, or yellow with a median longitudinal black line; south-eastern Arizona south through central Mexico . crotalinus Cornuae of penis with tips bluntly rounded (figs. 87-88); medial lobe present (figs. 8384) or absent (figs. 87-88); extensor surfaces of tibiae black, never with yellow 17
17 (16). Tips of cornuae bluntly rounded; medial lobe absent (figs. 87-88) 18
$-\quad$ Tips of cornuae with a well defined medial lobe (figs. 83-84)

19
18 (17). Median part of base of postfrons with a transverse arcuate trough immediately anterior to median ocellus (as in fig. 138); western United States $\qquad$ compositus Base of postfrons medially with a pair of deep pits, each anterolateral to median ocellus (as in fig. 138); western Texas, western

New Mexico, Chihuahua, Mexico
heterodon
19 (17). Penis in dorsal view with membranous hood short, exposing a subcutaneous membranous hood which forms the posterior margin of segment [be sure to examine closely: rotation of penis to dorsolateral view will reveal a raised hood anterior to subcutaneous membrane], (fig. 83); Jalisco, Morelos, and Puebla, Mexico $\qquad$ sipedon Penis in dorsal view with membranous hood long, completely covering subcutaneous membrane, thus forming posterior margin of segment (figs. 84-85); western Texas, southern and central New Mexico and Arizona, southwestern California, Baja California, Durango state, Mexico
lampropeltis

## Erpetogomphus constrictor Ris

(figs. 1-thx, 28-abd, 97-hamules, 70-penis, 92-app, 145-오 vertex, 171-vl, 198-distr)
Erpetogomphus constrictor Ris, 1917: 154 ('Nach 30', 1 아 von Misantla, Vera Cruz, Mexiko, VI 1911, die ich dem schweizerischen Sammler W. Gugelmann verdanke.'). Williamson and Williamson 1930: 12 (summary of status); Montgomery 1973: 239 (derivation of name); Paulson 1982: 255 (Mex.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat.); Dunkle 1988: 46 (Honduras); Maes et al. 1988: 36 (Nicaragua); Bridges 1991: VII. 51 (cat.); Tsuda 1991: 95 (cat.).

## Description

Male. - Labium grey, becoming dark grey medially, labrum entirely brown to grey green with dark brown margin and large inverted medial triangular spot; base of mandibles grey green, anteclypeus grey green; postclypeus brown except for small triangular green spots above lateral lobes; ventral margin of frons brown joining brown of postclypeus, thus forming a large brown frontoclypeal stripe; remainder of frons green; extreme base of frons, vertex, and occiput dark brown; area immediately anterior to median ocellus with a shallow longitudinal trough, postocellar tumid areas prominent, a lower incomplete transverse ridge posterior to median ocellus but not continuous with each postocellar tumid area laterally; occiput planar, its crest costate, linear, fringed with dark brown hairs; postocciput brown, linear to gently concave medially; rear of head brown.
Prothorax brown, paler dorsally, anterior and posterior lobes green, some specimens with two small green midlateral spots (almost touching) on median lobes; synthorax (fig. 1) with prominent, well defined dark stripes on green background as follows: middorsal stripe narrowing anteriorly at collar, costal mar-
gins of antealar sinus and area along this structure connecting with humeral stripe; antehumeral stripe connecting with humeral basally and dorsally, but its upper end often not touching brown below antealar crest, or, if so, then isolating small area of green; second and third lateral stripes connecting at upper 0.25 , isolating small green spot, second lateral stripe swollen in vicinity of metaspiracle and, in some specimens, enlarged and touching third lateral stripe; posterior margin of metepimeron, metasternum, and venter of thorax grey green to tawny. Venter of profemora grey green, remainder of femora dark brown, becoming black distally; tibiae, tarsi, armature black.

Wings hyaline, venation and pterostigma black.
Venational statistics. Fifth (rarely fourth or sixth) antenodal thickened; number of marginal cells behind fore wing paranal cells: $0-1 / 0-1$; anx: fore wing 13-17/14-17, hind wing 10-12/10-13; pnx: fore wing 10-14/9-14, hind wing 9-12/9-13; cs under pterostigma.: fore wing 5-7/4-7, hind wing 4-7/5-7; anal triangular cells: $3-5 / 3-4$. Hind wing $25-30 \mathrm{~mm}$.

Abdomen with segment 1 brown, green laterally and with a green middorsal stripe; segment 2 brown except for green auricles, ventral margin, and narrow middorsal stripe, annulus black; segment 3 primarily black with green middorsal stripe extending from black annulus of segment 2 and ending at distal annulus, a pale grey green anterolateral spot becoming narrower along ventral margin and disappearing at 0.25 to 0.75 of segment length; segments $4-6$ similar to segment 3 but with pale middorsal stripe ending at posterior 0.10 to 0.25 of segment, segment 6 in some specimens with anterolateral white connecting dorsally with pale green middorsal stripe; segment 7 grey green on anterior 0.50 to 0.60 , except for dark transverse carina, posterior part of segment brown to black; segments $8-10$ red brown, becoming darker dorsally and ventrally, especially along foliate margins, posterior medial margin of segment 9 bluntly pointed or slightly so. Abdomen $31-35 \mathrm{~mm}$.

Cercus (fig. 92) grey green, becoming dark brown posteriorly, arcuate with a prominent laterally compressed dorsal tubercle on distal 0.75 of cercus, extreme base with a black ventral tooth (mostly hidden by posterior margin of abdominal tergite 10), a ventral carina on distal 0.25 of cercus, tip of cercus with a black tooth; epiprocts dark brown, strongly curved at distal 0.50 so that distal 0.5 of appendage is parallel, distal 0.30 sulcate medially on exterior surface. Accessory genitalia. Hamules (fig. 47) small; anterior hamule brown, divided at distal 0.30, posterior branch a well developed shoulder; posterior hamule roughly triangular, its tip with a spine; penis (fig. 70) with cornuae slightly bilobate or transverse at base, a well developed prepuce, lateral lobe a small tooth.

Female. - Head markings similar to male but with
dark median spor on labrum larger, often connecting with ventral brown, thus separating green into two lateral spots; some specimens with an entirely brown labrum; brown on frontoclypeal suture more extensive laterally, so that entire postclypeus may be brown: brown at base of antefrons with a median extension which, in some specimens, connects with brown frontoclypeal suture, thus dividing frontal green into two spots. Morphology of vertex and occiput differs from male as follows: a large longitudinal trough on vertex with median ocellus recessed posterior to lateral ocelli (fig. 145); bottom of median furrow with longitudinal convex area; lateral margins of trough formed by median ocellar protuberances connecting posteriorly forming a semicircular ridge; occiput small, forming a narrow semicircle, crest covered with long brown hairs; postocciput brown, convex.

Pro- and synthorax as in male, but with dark areas more extensive.

Venational statistics ( $\mathrm{n}=15$ ). Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: 0-5/0-3; anx: fore wing 14-17/14-17, hind wing 11-12/11-12; pnx: fore wing 11-14/11-14, hind wing 10-14/11-13; cs under pterostigma.: fore wing 5-7/5-7, hind wing 5-7/5-7. Hind wing 28-31 mm.

Abdomen (fig. 28) with segment 1 brown, a green spot posterolaterally and with a complete green middorsal stripe; segment 2 brown except for following green areas: wide middorsal stripe narrowing posteriorly, longirudinal lateral stripe connecting just anterior to auricle and narrowing slightly to posterior margin of segment, a thin pale margin at venter of segment; segment 3 similar to segment 2 but lateral pale stripe ending posteriorly at transverse carina, middorsal green narrowly acuminate posteriorly, forming a mere line on posterior 0.25 of segment; segments 4-6 similar to segment 3 but with middorsal green progressively shorter posteriorly, lateral pale stripe white and forming a small quadrangular spot at anteroventral margin of segment; segment 7 with anterior 0.50 pale green, interrupted by black transverse carina, posterior 0.50 becoming red brown; segments 8-10 all red brown, often darker dorsoposteriorly; cerci pale grey green. Vulvar lamina (fig. 171) with plates broadly connected at basal 0.75 , relatively unspecialized, cleft obtusely V- or U-shaped, about as wide as each lobe; postlamellar ridge with $Y$-juncture posterior to plates, base of stem well defined and with oval depression laterally. Abdomen $30-35 \mathrm{~mm}$.

## Diagnosis

Males of E. constrictor are easily separated from its closest allies, E. tristani and E. sabaleticus, by the sha$\because$ of the cercus. This structure is concave ventrally in E. constrictor, not convex as in the other two species.

It superficially resembles $E$. schausi in body colouration and is diagnosed under that species.

Females seem to be separable from the same two species by the width of the median trough of the vertex. This structure is slightly divergent anteriorly in $E$. constrictor (figs. 145), but is largely parallel in the other two species (fig. 146).

## Remarks

Variation. - Venational details of lectotype male of E. constrictor. No marginal cells behind fore wing paranal cells; anx: fore wing $15 / 14$, hind wing $11 / 11$; pnx: fore wing $11 / 12$, hind wing $12 / 13$; cs under pterostigma: fore wing $7 / 7$, hind wing $7 / 6$; number of anal triangular cells: $4 / 3$. Hind wing 29 mm .

Material I examined indicates that specimens from more southerly parts of its range (Guatemala, Honduras, Nicaragua) are more melanic than those from eastern Mexico. For example, black markings on the labrum, postclypeus, and thoracic striping are generally more extensive, further reducing or isolating green areas.

Two females, one from San Luis Potosí (El Salto Falls) and the paralectotype are unusual in having $5 / 3$ and $4 / 3$ marginal cells behind fore wing paranal cells in right and left fore wings, respectively. All other female specimens ranged from 0-1 marginal cells.

Biology. - E. González (in litt.) found this species to be abundant at the Rio Huichihuayan. Males perched on leaves, vines, and snags at the edge of the river. Judging from specimens in collections, E. constrictor is the northern counterpart of E. tristani, and appears to be common in certain areas.

Erpetogomphus constrictor appears to be a lowland species of wet tropical forest habitats along the east coast of Mexico south through Costa Rica (Guanacaste Prov.: Miravalles, 26 April 1991 (C. Esquivel), $3 \sigma^{\star}$, A. Ramirez, in litt.). Altitudes range from near sea level ( 70 m , San Luis Potosí, km 410) to about 550 m (Nuevo Leon, entrance to Horsetail Falls). Collection dates range from 26 April to 4 October. One specimen was collected in an ultraviolet light trap.

Distribution (fig. 198). - This species occurs along the eastern Mexican states of Nuevo Leon, Tamaulipas, San Luis Potosí, Veracruz, south through Guatemala, Honduras, and Nicaragua, to Guanacaste Prov., Costa Rica. It is apparently parapatric or allopatric with E. tristani in Costa Rica.

## Material

Type data. - Lectotype $\delta$ by present designation with the following data in Ris' hand: '15098' in green ink in an unknown hand, Erpetogomphus / n. sp. ő / constrictor Ris 1917/ Misantla, Vera Cruz / Mexico VI 1914 / W. Gugelmann. Two paralectotype males
and one female with same data. One male is labelled ' 15092 ' in green ink, another male in poor condition (head partially fragmented, abdomen broken in several places) is labelled 'Type' probably by Ris. The left pair of wings is missing and probably served for figure 6 in the original description. The female is labelled ' 15096 ' in green ink. In SMF.

Other material ( 53 ox, 15 , including lectorype of and paralectorypes). - Mexico: Nuevo Leon: Cola Caballo, SW Monterrey, 6 July 1960 (W. B. Cutts), 19 (ummz); Rio Ramos, 3 km S of Allende, $500 \mathrm{~m}, 7$ Sept. 1963 (T. W. Donnelly), $1 \delta^{\text {( }} \mathrm{RwG}$ ); south of Monterrey, Rio Elizondo, 19-20 June 1965 (O. S. Flint, Jr.), $6{ }^{\text {ot }}$ (UsNm, RWG); stream at entrance to Horsetail Falls, 550 m, 25 June 1965 (D. R. Paulson), 1 (Drp); Tamaulipas: Gomez Farias, Río Frio, 4 Oct. 1985 (F. Arias, L. Cervantes, M. Garcia), 'colectado en trampa U.V. en la noche', 1 ( (RwG); San Luis Potosí: El Salto, 4 June 1967 (O. S. Flint, Jr.), 1 ( (UsNm); Palitla, N of Tamazunchale, 25 June 1965 (O. S. Flint, Jr.), 90 (USNM, RwG); Huichihuayan, rte 85, km 399, 7 Aug. 1966 (O. S. Flint, Jr.), $5{ }^{\text {® }}$ (UsNm, Cc, TwD); stream crossing Hwy $85\left(1000^{\circ}\right) ; 6 \mathrm{mi} \mathrm{N}$ of Tamazunchale, 4 Sept. 1957 (G. H. Beatty, III), 10 (fsca); km 410, 'Sam Brown' Hacienda, 250 ft, 27 Sept. 1938 (L. J. Lipovsky), $1 \begin{gathered}\text { § (UMmz); Route }\end{gathered}$ 85, km 399, Huehuetlán, 26 June 1965 (O. S. Flint, Jr.), $40^{\circ}, 1$ 오 (USNM); Rio Huichihuayan, Mpio de Huchuetlan, 100 m, 26 May 1987 (E. González-S, 20 (RwG); upstream Cascadas Micos just $S$ of town $N$ of Ciudad Valles off Hwy 85,27 June 1990 (J. S. Daigle), 1 of (rwg); Guatemala: Suchitepequez Dept., Cuyotenango, Finca San Rafael Olimpo, 1700 ft, 10-20 June 1966 (O. S. Flint, Jr., M. A. Ortiz-B.), $80^{\circ}$ (usnm, rwg); honduras: Comayagua Dept., Rancho Chiquito, 11.8 mi SE Villa de San Antonio, 2-3 Aug. 1967 (O. S. Flint, Jr., M. A. Ortiz-B.), $3 \delta^{\star}, 7$ 우 (usnm, cc, Rwg); Francisco Morazan Dept., 30 km E of Tegucigalpa, 13 June 1982 (Fernandez, Garcia), 1 ㅇ (swd); [no date], (Carlos Jaramillo), 1 it (swd); El Salvador: Abuachapan Dept., Loma de Paya, Bosque El Impossible, San Francisco Menendez, 24 July 1987 (A. Campus), 10 (vн); 20 Aug. 1986 (V. Hellebuyck), 1 ( (vн) 4 Aug. 1987 (V. Hellebuyck), 1 ( (vh); nicaragua: Chobales Dept., La Flor, rte 7, km 159, 4 mi W of Acoyapajit, 29 July 1967 (O. S. Flint, J., M. A. Ortiz- B.), $40^{\circ}$ (UsNM); Managua Dept., stream along Pan American Hwy, 7 km S of Nandaime, 21 June 1962 (T. W. Donnelly), $1 \delta^{\hbar}$ (Twd).

## Erpetogomphus sabaleticus Williamson

(figs. 2-thx, 48-hamules, 71 -penis, 93 -app, 172-vl, 198-distr)
Erpetogomphus sabaleticus Williamson, 1918: 1 (descr. holotype ${ }^{\text {on }}$, allotype ¢). - Williamson and Williamson 1930: 12 (summary of status); Paulson 1982: 255 (Panama, South America); Davies and Tobin 1985: 28 (cat.); Tsuda 1986: 87 (cat.); De Marmels 1990: 338 (Venez.); Bridges 1991: VII. 184 (cat.); T suda 1991: 95 (cat.); Belle 1992: 32 (descr. larva); Donnelly 1992: 85 (Panama); Belle and Quintero 1992: 99 (key, Panama).

## Description

A thorough comparative description of the holotype male and allotype female is given by Williamson
(1918). I illustrate the thorax (fig. 2), caudal appendages (fig. 93) and details of the accessory genitalia (fig. 48) of the holotype, and the vulvar lamina of the allotype (fig. 172).

Male ( $\mathrm{n}=6$, including holotype). - Venational statistics. Fifth (rarely sixth) antenodal thickened; number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing 13-18/15-17, hind wing 10-12/912; pnx: fore wing 11-13/15-16, hind wing 11-14/11-13; cs under pterostigma: fore wing 5-6/6-7, hind wing 5-7/6-7; anal triangular cells 3-4/3-4. Hind wing $26-27 \mathrm{~mm}$. Abdomen $30-33 \mathrm{~mm}$.

Female ( $\mathrm{n}=4$, including allotype). - Venational statistics. Fifth or sixth (rarely fourth) antenodal thickened; number of marginal cells behind fore wing paranal cells: $0-1 / 0-1$; anx: fore wing $15-17 / 15-16$, hind wing 11-12/9-12; pnx: fore wing 11-15/13, hind wing 10-12/11-13; cs under pterostigma: fore wing 6-7/6-7, hind wing 6-7/6-7. Hind wing 28-29 mm . Abdomen $31-32 \mathrm{~mm}$.

## Diagnosis

Erpetogomphus sabaleticus is diagnosed under $E$. tristani.

## Remarks

Variation. - Venational details of holotype male: number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing $15 / 16$, hind wing $11 / 11$; pnx: fore wing $13 / 13$, hind wing $12 / 11$; cs under pterostigma: fore wing $6 / 6$, hind wing $6 / 6$; anal triangular cells: $3 / 3$. Hind wing 26 mm .

Venational details of allotype female: number of marginal cells behind fore wing paranal cells: $1 / 0$; anx: fore wing $16 / 15$, hind wing $12 / 11$; pnx: fore wing $12 / 13$, hind wing $10 / 11$; cs under pterostigma: fore wing $6 / 6$, hind wing $6 / 7$. Hind wing 29 mm .

The living colours of the holotype male were recorded by Williamson (1918).

Biology. - Williamson (1918) collected the allotype female and holotype male along the upper San Juan near Maraquita in Colombia. The male was taken along the Quebrada Sabaleticus as it rested on a flat leaf about five feet above the water. Only one other specimen of this species was seen (but not collected) during their stay in Colombia. A description of the type locality is given by Williamson (1918).

Michael May collected one male and three female larvae of the species at the Quebrada Juan Grande, along Pipeline Road in the Panama Canal Zone on 28 January 1975 and 23 January 1977. These were brought back to Gainesville, Florida, where the adults emerged about three months later.

This species, like E. tristani, is apparently rare. Other odonatists and I have collected along Pipeline Road in Panama, but have failed to find the species.

Distribution (fig. 198), - Erperogomphus sabaleticus is the most austral of the genus and is currently known from one locality in northwestern Venezuela, one locality in northern Colombia, and Panama. It is apparently the only member of the genus found in the first two countries. Its northern limit is unknown, but it is probably parapatric with the more northerly E. tristani. Collection dates range from 3 February (Maraquita, Colombia) through 24 June (Panama Canal Zone).

## Material

Type data. - Holotype male: Colombia: Dept. Antioquia: Cristalina, 28 km on railroad above Puerto Berrio, 19 Feb. 1917 (J. H. and E. B. Williamson); allorype female: Colombia: Dept. Tolima: Maraquita, 3 Feb. 1917 (J. H. and E. B. Williamson). Both specimens in ummz.

Other marerial ( $60^{*}, 4$, including holotype ${ }^{\text {o }}$ and allorype 9 ). - Panama: Canal Zone, Quebrada Juan Grande, Pipeline Road, 28 Jan. 1975 (M. L. May), 1 I larva, emerged 19 June 1975 ( FSCA ); same data but 23 Jan. 1977 (FSCA), 10 , 29 larvae, male emerged 16 May 1977, females emerged 20, 26 May 1977; Pipeline Road, 1.7-4.8 mi NW of Gamboa, 2nd bridge, 24 June 1970 (E. S. Morton), 10 (TWD); Venezuela: Tachira Dept.: Tachira, 11 April 1920 (J. H. and E. B. Williamson, W. H. Ditzler), $3 \delta^{\star}$ ( $2 \delta^{\star}$ in UMmZ, $1 \delta^{\circ}$ in FSCA).

## Erpetogomphus tristani Calvert

(figs. 3-thx, 49-hamules, 72-penis, 94 -app, 120-ept, 146-vertex, 173,174-vl, 198-distr)
Erpetogomphus tristani Calvert, 1912a: 290 (descr. of © , ㅇ). - Calvert 1912b: 384 (mentions mating adaptations); Williamson 1918: 1 (comparison with E. sabaleticus); Ris 1917: 154 (Panama); Ris, 1918: 154 (Panama); Calvert 1920b: 339 (mentions mating adaptations); Williamson and Williamson 1930: 12 (summary of status); Paulson 1982: 256 (Costa Rica, Panama); Davies and Tobin 1985: 28 (car.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 212 (cat.); Tsuda 1991: 95 (cat.); Belle and Quintero 1992: 100 (key, Panama).

## Description

Male. - A thorough description of the holotype male and allorype female is given by Calvert (1912). I illustrate the caudal appendages of the holotype (fig. 94) and vulvar lamina of the allorype (fig. 173). In addition to Calvert's description, I add the following: A strong ventral tooth at base of cercus (plainly seen in lateral view, fig. 94), which in most specimens is partially hidden by lateroposterior margin of tergite 10; tips of epiprocts narrow in posterior view (fig. 120); anterior hamule black, divided at apical 0.25 , lower branch a rounded shoulder; anterior (upper)
branch well developed; posterior hamule pale, acutely triangular, with a prominent apical tooth; penis with well developed prepuce, which, when viewed laterally, is hidden by ventrolateral margin of segment 4; lateral lobe small, rudimentary, barely visible, and with spinulose margin; cornua rudimentary, forming an ental membranous plate with a quadrate margin (fig. 72). Abdomen $31-33 \mathrm{~mm}$.

Venational statistics. - ( $\mathrm{n}=5$, including holotype). Fifth antenodal thickened; number of marginal cells behind fore wing paranal cells: $0-1 / 0$; anx: fore wing 13-15/14-15, hind wing 10-12/10-11; pnx: fore wing 11-13/11-13, hind wing 11-13/10-12; cs under pterostigma: fore wing 5-6/5-6, hind wing 5-6/5-6; anal triangular cells: $3 / 3$. Hind wing $25-27 \mathrm{~mm}$.

Female. - Vulvar lamina (fig. 174, allotype; 173) broadly connected, relatively unspecialized, and with a gently U-shaped notch; basal plate poorly developed; juncture of Y-shaped postlamellar ridge posterior to posterior margin of lamina; well defined arcuate depression on either side of central stem. Abdomen $31-33.5 \mathrm{~mm}$.

Venational statistics. - ( $\mathrm{n}=4$, including allotype). Fifth antenodal thickened; number of marginal cells behind fore wing paranal cells: $0-1 / 0$; anx: fore wing 15-16/15-17, hind wing 11-12/10- 12; pnx: fore wing $12-14 / 11-12$, hind wing $12-13 / 12-13$; cs under pterostigma: fore wing 5-7/6-7, hind wing 6-7/5-6. Hind wing 29 mm .

## Diagnosis

Erpetogomphus tristani, E. sabaleticus, and E. constrictor form a compact group which is easily separable from all other congeners by the unique shape of the epiprocts of the male (figs. 92-94, 120) and by the morphology of the vertex of the female (figs. 145146). Males of E. tristani differ from the other two species only in the shape of the cerci. The ventral margin is distinctly convex at the apical $0.60-0.80$ in E. tristani (fig. 94) and E. sabaleticus (fig. 93), not smoothly concave as in E. constrictor (fig. 92). All three species have a prominent dorsal tooth near the distal end, but the tooth in E. constrictor is broader in lateral view and more vertical than in $E$. tristani and E. sabaleticus.

Males of E. tristani and E. sabaleticus approach one another closely. Williamson (1918: 4-5) found males of both species inseparable except by the male caudal appendages. Williamson sent a pair of $E$. sabalecticus to Calvert, who also supported the statement of differences between the males of both species. Calvert was unable to separate the females of these two species. I have found the females of these three species (E. constrictor, E. tristani and E. sabaleticus) to be extremely similar, more so than their males. The sides
of the longitudinal trough of the vertex of $E$. constrictor are more parallel (fig. 145), than those of the other two species. Like Calvert and Williamson, I have not found any differences between females of the small series of $E$. tristani and $E$. sabaleticus.

## Remarks

Variation. - Little variation is expressed among the small series (males, $\mathrm{n}=5$; females, $\mathrm{n}=4$, including holotype and allotype).

Venational details of holotype male: number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing $14 / 14$, hind wing $11 / 11$; pnx: fore wing $12 / 12$, hind wing $13 / 12$; cs under pterostigma: fore wing $5 / 5$, hind wing $5 / 5$. Hind wing 27 mm .

Venational details of allotype female: number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing $15 / 15$, hind wing $11 / 11$; pnx: fore wing $14 / 12$, hind wing $13 / 12$; cs under pterostigma: fore wing $7 / 7$, hind wing $7 / 6$. Hind wing 29 mm .

Biology. - The species is relatively rare in collections, despite intensive collecting in Costa Rica in recent years. Collection dates are February, June, and July. Nothing else is known of its biology.

Distribution. (fig. 198). - Erpetogomphus tristani is thus far known from only two northwestern provinces of Costa Rica and Panama (Lino, Chiriqui Prov. at about $1600 \mathrm{~m}, 8^{\circ} 48^{\circ} \mathrm{N}, 82^{\circ} 26^{\circ} \mathrm{W}, 1 \mathrm{\sigma}^{\circ}, 1$ ㅇ, Ris 1917). It is replaced to the north by $E$. constrictor (fig. 198) and to the south by E. sabaleticus. Erpetogomphus tristani will probably be found in southern Nicaragua, but its southernmost distribution is unknown.

## Material

Type data. - Holotype male with following label data (all handwritten by P. P. Calvert unless otherwise noted): 'Oricuajo,/ Costa Rica./ July, 1911/ J. F. Tristan', 'Erpetogomphusl tristanil $\delta /$ Calvert/ Type.', 'Needham/ fig' [in Needham's hand?], red label: '9244 тyPe [printed] 944/ Erpetogomphusl tristani ơ/ Calvert.', '2nd ham-/ ules/ sheath of/ penis.' Allotype female with same label as holotype except for following: 'Labium \&/ Maxillae.',' 'Erpetogomphus/ trista$n i$ i / Calvert'. Both specimens were originally pinned, but each has been placed in a clear envelope. Both specimens in ansp.

Other material ( $5 \delta, 4$ ㅇ, including holotype $\delta$ and allotype f). - Costa Rica: Guanacaste Prov:: Nicoga, Feb. 1912 (J. F. Tristan), 10 (Ummz); Quebrada Azul, 2.5 mi W of Tilaran, 24 July 1967 (O. S. Flint, Jr., and Ortiz B.), 1 § , 1 ㅇ (usnm); Rio Santa Rosa, 3.7 mi E of Las Canas, 25 July 1967 (O. S. Flint, Jr., and Ortiz B.), 1 if (usnm); Puntarenas Prov.: 8 mi WNW of Esparta, 26 June 1967 (O. S. Flint, Jr., and Ortiz B.), $10^{\text {® }}$ (UsNM); stream 8.1 mi WNW of

Esparta, 26 June 1967 (O. S. Flint, Jr., and Ortiz B.), 10 (Drp); San Jose Prov:: Res. Biol. El Rodeo, 7 km W of Villa Colon, $9^{\circ} 54^{\prime} \mathrm{N} 84^{\circ} 16^{\prime} \mathrm{W}, 800 \mathrm{~m}, 10-13$ July 1990 (T. W. Donnelly), 1 ¢ (TwD).

## Erpetogomphus ophibolus Calvert

(figs. 4 -thx, 29 -abd, 50 -hamules, 73 -penis, 95 -app, 121 -ept, 147 -vertex, 175 -vl, 198-distr)
Erpetogomphus ophibolus Calvert, 1905: 163 (Mexico: Atoyac in Veracruz, H. H. Smith, $2 \mathbf{0}^{\circ}$ ). - Calvert 1909: 489 (distr., Mex.); Muttkowski 1910: 87 (cat.); Calvert 1912a: 294 (comparison with E. tristanz); Ris, 1917: 155 (comparison with E. constrictor); Williamson and Williamson 1930: 489 (summary of status); Kimmins 1969: 296 (type in bмnн); Montgomery 1973: 239 (derivation of name); Paulson 1982: 255 (Mex.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 154 (cat.); Tsuda 1991: 95 (cat.).
Erpetogomphus (Erpetocyclops) ophibolus. - Carle 1992: 148 (key to subgenera, type species of Erpetocyclops subg. n.).

## Description

Male. - Labium, labrum, mandibles, anteclypeus brown; postclypeus entirely brown, or with small lateral green triangular spot; frons green except for brown along frontoclypeal suture, and at base of antefrons; vertex dark brown; area anterior to median ocellus undifferentiated, postocellar tubercles well developed, connected medially by a low transverse ridge, slightly tuberculate medially; occiput planar, trapezoidal, hind margin linear or slightly concave medially, crest covered with long brown hairs; postocciput brown, transversely concave; rear of head brown.

Prothorax brown, anterior lobe green, posterior lobe green brown; synthorax (fig. 4) with prominent well defined dark stripes on green background as follows: middorsal stripe slightly narrowing dorsally, antealar sinus and area along this structure connecting to humeral stripe, antehumeral stripe of same width as humeral connecting ventrally and (but not always) dorsally, and in some specimens broadly so, antehumeral not or just touching antealar sinus; second and third lateral stripes connecting at upper 0.75 , isolating round green spot below subalar carina; posterior margin of metepimeron, metasternum and venter of thorax grey brown; venter of profemora grey green, remainder of femora dark brown, becoming black distally; tibiae, tarsi, armature black. Wings hyaline, venation and pterostigma black.

Venational statistics. Fifth (rarely fourth or sixth) antenodal thickened; number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing 13-15/13-16, hind wing 10-12/10-12; pnx: fore wing $10-13 / 8-12$, hind wing $9-12 / 9-12$; cs under pterostigma: fore wing 4-7/5-8, hind wing 5-6/4-7; anal tri-
angular cells: $3 / 3$. Hind wing $25-27 \mathrm{~mm}$.
Abdomen (fig. 29) with segment 1 brown, green posterolaterally and with a green middorsal stripe; segment 2 brown except for green auricles, ventral margin, and narrow middorsal stripe; annulus dark brown; segment 3 primarily dark brown, becoming black distally with green middorsal stripe extending from annulus of segment 2 and narrowing posteriorly to end, a pale grey anterolateral spor becoming narrower along ventral margin and disappearing at 0.25 to 0.50 of segment length; segments $4-6$ similar to segment 3 but with pale middorsal stripe ending at posterior 0.10 to 0.75 of segment and anterolateral pale spot ending at transverse carina; segment 7 grey green on anterior 0.50 to 0.60 , except for dark transverse carina, posterior part of segment brown to black; segments $8-10$ red brown, becoming darker dorsally and ventrally, especially along foliate margins; posterior medial margin of segment 9 bluntly pointed or slightly so. Abdomen $30-33 \mathrm{~mm}$.

Cercus (fig. 95) ivory brown, becoming brown distally, arcuate with a prominent laterally compressed dorsal tubercle on distal 0.75 , extreme base with a black ventral tooth (mostly hidden by posterior margin of abdominal segment 10), though not as prominent as in $E$. constrictor, a ventral carina on distal 0.30 of cercus, tip of cercus with a black tooth; epiprocts brown, gently curved at distal 0.50 , tips slightly diverging, each forming a blunt point (fig. 121), a well defined anterodorsally projecting tooth at middle of medial margin of epiproct.
Accessory genitalia. Hamules (fig. 50) small; anterior hamule brown, divided at distal 0.30 , posterior branch a well developed shoulder; posterior hamule roughly triangular, its tip with a spine; penis (fig. 73) with cornuae slightly bilobate or transverse at base; a well developed prepuce, lateral lobe a small tooth.
Female. - Head markings similar to male but with some green on mediolateral areas of labrum, brown at base of antefrons with a median extension connecting with frontoclypeal suture, thus dividing frontal green into two spots. Morphology of vertex and occiput similar to male with following differences: transverse postocellar ridge (fig. 147) complete, slightly arcuate medially, with a slight shallow notch laterally before resuming at postocellar tubercle; occiput a narrow planar semicircle, crest covered with long brown hairs; postocciput tumid, brown; rear of head brown.

Pro- and synthorax as in male, with dark markings in some specimens less extensive.

Venational statistics ( $\mathrm{n}=11$ ). Fifth (rarely fourth or sixth) antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $0-1 / 0-2$; anx: fore wing $14-16 / 14-17$, hind wing $10-12 / 10-13$; pnx: fore wing 10-13/9-13, hind wing 10-13/10-14; - 5 under pterostigma: fore wing 5-8/5-8, hind wing

5-7/4-7. Hind wing 26-29 mm.
Abdomen with segment 1 brown, a green spot posterolaterally and a complete green middorsal stripe; segment 2 brown with following areas green: wide middorsal stripe narrowing posteriorly, incomplete lateral stripe extending from just anterior to auricle to black posterior annulus, a thin pale margin at venter of segment; segment 3 similar to segment 2 but brown becoming darker; middorsal stripe narrowing posteriorly and forming a mere line on posterior 0.25 of segment; lateral pale stripe ending posteriorly at transverse carina; segments 4-6 similar to segment 3 but middorsal green becoming increasingly smaller on successive segments, so that only a small middorsal spot occupies basal 0.10-0.50 of segment 6 ; segment 7 with anterior 0.50 pale green white, interrupted by brown transverse carina, posterior 0.50 becoming red brown to almost black; segment 8-10 all red brown to dark brown, darker dorsoposteriorly; cerci pale grey green. Vulvar lamina (fig. 175) with plates broadly connected at basal 0.75 ; relatively unspecialized, cleft obtusely V- or U-shaped, about as wide as each lobe; postlamellar ridge with Y - juncture posterior to plates, base of stem well defined and with oval depression laterally. Abdomen $30-33 \mathrm{~mm}$.

## Diagnosis

Males of $E$. ophibolus are unique in possessing an anterodorsal spine on the mediodorsal surface of the epiproct (fig. 95). The cercus of this species is most similar to that of $E$. constrictor (fig. 92). Morphology of the vertex and occiput serve to distinguish female $E$. ophibolus from similarly marked species. The median ocellus is in line with the lateral ocelli (fig. 147), separating this species from $E$. sabaleticus, E. tristani (fig. 146), and E. constrictor (fig. 145). Female E. ophibolus more closely resemble females of E. schausi, E. agkistrodon, and E. eutainia. However, E. schausi and E. agkistrodon have a transverse postocellar ridge which is medially bilobed (figs. 148-149), while E. eutainia lacks any postocellar ridge and has postocellar tubercles only. The postocellar ridge in E. ophibolus is arcuate and entire medially (fig. 147). Erpetogomphus opbibolus and E. eutainia are often sympatric, and the females superficially resemble one another. The second and third thoracic stripes in E. eutainia are always separate (figs. 7-9), but are connected at their upper 0.30 in E. ophibolus (fig. 4). Finally, females of E. eutainia lack the basal stem of the postlamellar ridge (fig. 178) present in E. ophibolus (fig. 175).

## Remarks

Variation. - Most specimens of E. ophibolus that I examined are from southern Veracruz, and they show little variability in maculation. The single male from Chiapas has slightly more extensive black thoracic
markings：the pale area between the dark humeral and antehumeral stripes is a narrow line．The two females from Belize have the apical dark brown on abdominal segment 7 laterally extending anteriorly separating the lateral pale area into one anterodorsal and one an－ teroventral spot．The acuminate condition of the pos－ terior medial margin of abdominal segment 9 varies intraspecifically．

Biology．－This small green species often occurs with E．eutainia at small rivulets along the lowland Gulf of Mexico drainage systems of eastern Mexico． In 1976 I collected many by flushing them from agri－ cultural stubble bordering trees and shrubs along the Rio Otapa in central Veracruz．Adults，when flushed， did not fly far and were consequently easy to collect． Six other gomphids collected at the same site under the same circumstances included Phyllocycla brevi－ phylla Belle，P．volsella（Calvert），Phyllogomphoides suasus（Selys），P．duodentatus Donnelly，Progomphus clendoni Calvert，Erpetogomphus eutainia，and E． bothrops．
Raúl López（in litt．），from information given to him by Enrique González，writes that E．ophibolus is the most common gomphid in Veracruz，found year－ round in the vicinity of the Los Tuxtlas Biological Station near Catemaco．Altitudinal range is from sea level to 615 m （Chiapas）．Collection dates range from 8 June（Belize）to 13 September（Veracruz）．

Distribution（fig．198）．－This species is thus far known from southern Veracruz（southeast of $19^{\circ} \mathrm{N}$ ， $\left.97^{\circ} \mathrm{W}\right)$ ，central Chiapas，Guatemala，and Belize． Tineke Boomsma（in litt．）collected a pair of this spe－ cies in the Mountain Pine Ridge area of Belize（Cayo Distr．：Privassion Creek， 24 July 1993）．

## Material

Type data．Holotype male：［Mexico］Vera Cruz， Atoyac，May（H．H．S［mith］）／Erpetogomphus ophi－ bolus Calv．type ठ̃．P．P．Calvert det．1905．B．C．A． Neur．，p．164．Original of pl．7，figs．30－32， 46 （Kimmins 1969）．In bмnн．

Other material（ 43 すे， 12 ））．－Mexico：Veracruz： Rio Otapa， 8 km S of La Tinaja， $90 \mathrm{~m}, 13$ Aug．1976， （R．W．and J．A．Garrison）， $18 \delta^{\star}, 3$ 오，（RWG）； 20 Aug． 1976，12才， 1 if（RWG）；Playa Escondida，20－24 June 1981，（E．González）， 1 ㅇ，（Unam）；Arroyo cerca de Playa Escondida，17－24 July 1981，（E．González）， 1 우， （UNAM）；arroyo nr．Playa Escondida，about 30 km NE of Catemaco， $18^{\circ} 34-36^{\prime} \mathrm{N}, 95^{\circ} 04-09^{\prime} \mathrm{W}, 23$ July－ 6 Aug．，9－13 Aug．1982，（R．W．Garrison），50＇，（RWG）； Los Tuxtlas，arroyo despues de Laguna Escondida， 25 May 1980，（E．González）， 1 ®ै $^{\text {，（swd）；}} 11$ June 1980， （R．Novelo）， 10 太 ，（UNAM）；Los Tuxtlas，stream nr． Jicacal Beach， 10 July 1979，（Gerado Jimenez）， 1 오， （cc）；Colonia Apachital， $16 \mathrm{~km} \mathrm{~S}, 10 \mathrm{~km}$ E of Tierra Blanca， 10 Sept．1965，（T．W．Donnelly），2ठ̃， 1 ㅇ，
（Twd，cc，unam）；Rio Hondo，on rd．to＇Colonia la Apachital＇ $50 \mathrm{ft}[15 \mathrm{~m}], 6 \mathrm{mi}$ E of Hwy from Tierra Blanca to Cd．Aleman，25－26 Aug．1957，（G．H． Beatty III）， $1 \mathbf{\delta}^{\circ}$ ，（ FSCA ）； 3 km N of Santiago Tuxtla， 13 Sept．1965，（T．W．Donneily）， 1 if，（cc）；Chiapas： stream 20.1 mi N Ocozocoautla， 2000 ft ［ 615 m ］， 25 Aug．1967，（D．R．and M．L．Paulson）， $10^{\circ}$ ，（ FSCA ）； Belize：Toledo Distr：Blue Creek Village，earth－ watch Belize Expedition，1981， 8 June 1981，（D．H． Messersmith，W．E．Steiner，et al．）， 19 ，（UsNM）；Cayo Distr：：Mountain Pine Ridge，Rio Frio at Augustine， $16^{\circ} 58^{\prime} \mathrm{N}, 88^{\circ} 59^{\prime} \mathrm{W}, 500 \mathrm{~m}, 22-25$ July 1983，（T．W． Donnelly）， 1 甲，（Twd）；Guatemala：El Progreso Dept．： 6.5 km N of Est．de la Virgén， 29 Aug．1965，（T．W． Donnelly）， 10 ， 2 ㅇ，（TWD）．

## Erpetogomphus agkistrodon sp．n．

（figs．5－thx， 30 －abd， 148 －vertex， 176 －vl， 198 －distr）
Type material．－Holotype 9, Mexico：Veracruz State，Parque Javier Clavijero，Jalapa， 23 Aug．1982， 1300 m （R．López）in unam．

## Description

Male．－Unknown．
Holotype female．－Labium pale grey，becoming dark grey medially；labrum，clypeus，base of mandible dark brown washed with dark olive green along me－ dian area of labrum，ante－clypeus，and above lateral lobes；pale green spot along lateral margin of labium； broad green stripe above frontoclypeal suture contin－ uing to near base of vertex；antennae，vertex，occiput dark brown；transverse postocellar ridge（fig．148） complete，emarginate medially；rear of occiput broad－ ly semicircular；rear of head dark brown．
Prothorax entirely brown except for green anterior lobe；synthorax（fig．5）with dark brown thoracic stripe well developed on green background；triangular middorsal stripe interrupting inverted green＇ 7 ＇at base，upper end of middorsal stripe connecting with antehumeral and humeral stripes，these two united leaving only pale spot below antealar sinus，and small green streak below；second lateral stripe connecting with humeral and third lateral stripes above and be－ low，this stripe also connecting with third lateral be－ low antealar sinus，thus isolating a green spot；meta－ spiracle black；posterior margin of metepisternum with a dark brown stripe；metasternum dark brown－ grey．Femora red－brown，becoming black distally；re－ mainder of legs and armature black．

Wings hyaline，venation and pterostigma black．
Venational details．Sixth antenodal thickened in all wings；number of marginal cells behind fore wing par－ anal cells： $3 / 2$ ；anx：fore wing $15 / 17$ ，hind wing $12 / 11$ ； pnx：fore wing 14／16，hind wing $15 / 15$ ；cs under pt： fore wing $6 / 5$ ，hind wing $5 / 6$ ．Hind wing 31 mm ．

Abdomen (fig. 30) predominandy black with vesriges of a narrow green middorsal stripe on segments 2-6; anterior 0.25 of segment 7 green, becoming white laterally; broad brown midlateral stripe on segment 2 becoming darker on segment 3 and more extensive on succeeding segments, thus encircling small white anterolateral spots; auricle on segment 2 green; scgments 8-10 entirely black, cercus white. Vulvar lamina as shown in fig. 176. Abdomen 31 mm .

## Diagnosis

Erpetogomphus agkistrodon belongs to the E. trista$n$ igroup because of the simple condition of the vulvar lamina and the complete postocellar ridge. This species is distinguished from females of all other species by the emarginate condition of the postocellar ridge. That of $E$. ophibolus is entire. The female of $E$. agkistrodon is similar to the only known female of $E$. schausi and is diagnosed under the latter species.

## Remarks

I suspect the male of $E$. agkistrodon will have a penis similar to that of E. ophibolus or E. schausi; it should have a prepuce and the lateral lobes should be small, semi-circular and with or without spinules.
The thickened sixth antenodal in this species is an unusual condition: the fifth antenodal is usually thickened throughout the genus. However, this may be an anomaly of the holotype.

No conspicuous depressions or pits are present anterior to the median ocellus. This area, instead, forms a shallow transverse V . Based on female morphology, I suspect the male epiprocts will be relatively thick, gently curved at posterior 0.5 , forming a $90^{\circ}$ angle, tip broadly spatulate.

Biology. - The female was collected along a shaded creek in company with Cordulegaster diadema godmani McLachlan. Erpetogomphus boa were also taken at the locale, though they were flushed from low vegetation about 50 m from the stream.
Distribution (fig. 198). - Known only from the type locality.

## Erpetogomphus schausi Calvert

(figs. 6-thx, 31-abd, 51-hamules, 74-penis, 96-app, 142-face, 149-vertex, 177 -vl, 198-distr)
Erpetogomphus schausi Calvert, 1919:33 (desc. holotype ©). Calvert 1920a: 113 (note on type); Williamson and Williamson 1930: 13 (summary of status); Paulson 1982: 255 (Guat.); Davies and Tobin 1985: 28 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 187 (cat.); Tsuda 1991: 95 (cat.).

## Description

Male. - A thorough description of the holotype male accompanies the original description. I illustrate
the thorax (fig. 6), abdomen (fig. 31), accessory genitalia (fig. 51), penis (fig. 74), head (fig. 142), and caudal appendages (fig. 96) of the holotype. The specimen has been skewered with a bristle. The cercus of the holotype (fig. 96) is strongly curved at distal 0.30 , the ventral margin has a remnant of an inferior carina at its extreme base, but is well developed again on the distal 0.50 .

Venational statistics (based on holotype $\begin{gathered} \\ \text { and one }\end{gathered}$ t from Costa Rica): number of marginal cells behind fore wing paranal cells: $3 / 2-3$; anx: fore wing $16-$ 18/16-20, hind wing 12-14/12-13; pnx: fore wing $15 / 14-15$, hind wing $13-14 / 13-15$; cs under pterostigma: fore wing 5-6/5-6, hind wing 6/5-6; anal triangular cells: $4 / 4$. Hind wing $29-30 \mathrm{~mm}$. Abdomen 3233 mm .

Female. - Labium grey; labrum brown with indication of green mediolaterally, ante- and postclypeus brown with possible green medially on postclypeus; frons green with brown along frontoclypeal suture and base of antefrons, especially medially, with offshoot almost touching frontoclypeal suture, thus almost separating green; base of antefrons with a slight longitudinal raised area with a slight concavity on each side; antennae, vertex brown; transverse postocellar ridge (fig. 149) complete, emarginate medially; occiput brown, semicircular crest covered with long brown hairs; postocciput brown, convex, rear of head brown.

Prothorax entirely brown, paler on anterior and posterior lobes; synthorax as in holotype male (fig. 6); femora grey brown, tibiae and tarsi darker, armature black.

Wings hyaline, venation and pterostigma black.
Venational statistics ( $n=1$ ). Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $2 / 0$; anx: fore wing $16 / 16$, hind wing $12 / 11$; pnx: fore wing $15 / 15$, hind wing 14/14; cs under pterostigma: fore wing $6 / 5$, hind wing $6 / 6$. Hind wing 31 mm .

Abdomen compressed laterally with markings mostly unrecognizable, but probably similar to E. agkistrodon (fig. 30). Vulvar lamina simple (fig. 177), similar to that of $E$. agkistrodon. Abdomen about 32 mm .

## Diagnosis

Erpetogomphus schausi is unique in possessing a combination of characters of the penis (with well developed prepuce) and caudal appendages (with gently decumbent cercus (fig. 96)). Although the morphology of its penis allies this species with $E$. sabaleticus, $E$. tristani, and E. constrictor, the penes of the other three species lack well developed cornuae and a spinulate lateral lobe, typical of E. schausi. The only known female differs from all other known species except $E$.
agkistrodon in possessing a complete transverse postocellar ridge which is notched medially. The only morphological difference that I have noted between these two species is the broader, more semicircular occiput (figs. 149) of E. schausi, compared with the narrower, more broadly semicircular condition found in the holotype of E. agkistrodon (fig. 148). The sixth antenodal crossvein is thickened in all four wings of E. agkistrodon, while the fifth is thickened in this female of $E$. schausi; but, as stated under $E$. agkistrodon, I suspect the condition in the holotype is atypical.

## Remarks

Variation. - The male sex of this species is known from only two specimens, the holotype male described in detail by Calvert (1919), and another male from Costa Rica collected by J. Belle and loaned to me for inclusion in this paper. The Costa Rican male is slightly smaller $(H W=29 \mathrm{~mm})$ than the holotype ( $\mathrm{HW}=30 \mathrm{~mm}$ ) and differs as follows (condition for holotype in parentheses): small green triangle just lateral to medial area of anteclypeus (all dark brown); round green spot below antealar sinus of mesepisternum smaller (bigger); anterior pale abdominal spots extending well beyond transverse carina on segments 3-7 (these spots smaller and confined to anterior 0.25 of segments 4-6, to 0.5 of segment 7); sides of abdominal segments $8-10$ light brown (dark brown); lateral lobe of penis more angulate (more rounded); second thickened antenodal in fore wing 5-5 (6-7) and in hind wing 5-6 (6-6).

The holotype appears to be a fully mature male: traces of pruinosity are present on the venter of its thorax. The Costa Rican male seems to be less mature: it lacks pruinosity, and the dark brown areas of the holotype are pale in this specimen. The green areas of the thorax of the Costa Rican male are slightly more restricted than for the holotype.

The teneral female described above seems to represent the female of this species. The combination of characters allying it with the E. tristani group and the occurrence of a male from Puntarenas Province, lead me to suspect that I have associated this specimen correctly.

Before the discovery of the two E. schausi from Costa Rica, I considered the possibility that E. agkistrodon might be the female of E. schausi: morphological considerations in light of the characters for other members of the $E$. tristani group (e.g., vulvar lamina, shape of occiput) supported this view. However, the geographic distance (about 1700 km ), coupled with the local occurrence of both forms, tend to discount this. It is possible that the morphological differences I detect in the occiput of female $E$. agkistrodon and $E$. schausi, though major, may be due to geographic variation, which would render E. agkistrodon a junior
synonym of $E$. schausi.
Biology. - Nothing is recorded of the circumstances of capture of the holotype. Jean Belle records the following for the Costa Rican male: 'This gomphid was secured on the grass-covered bank of an almost impassable mountain rivulet where the insect alighted after a clash with a Hetaerina specimen. The colors when alive are as follows: compound eyes dark brown; head and thorax dark brown with light green; middorsal pale spots of abdominal segments 3 to 6 and pale spot of abdominal 7 yellow.'

Altitudinal data range from 1220 m to 1500 m ; the female was collected 8-9 April, the Costa Rican male on 30 August.

## Material

Type data. - Holotype male: Guatemala: Dept. Purulhá, Baja Vera Paz, 1220 m , forest stream (W. Schaus, J. Barns), 7 July [no date], in ansp.

Other material ( 2 추, 1 우, including holotype $\delta^{\top}$ ). Costa Rica: Puntarenas Prov.: Monte Verde, 1500 m, rivulet, 30 Aug. 1986 (J. Belie), 1 ( (RNHL); Rio Bellavista, ca. 1.5 km NW of Las Alturas (8.951 N, 82.846 W), elev. $1400 \mathrm{~m}, ~ 8-9$ April 1987 (Holzenthal, Hamilton, Heyn), 1 it (usnm).

Distribution (fig. 198). - Known only from the type locality in Guatemala and Puntarenas Province, Costa Rica. It probably occurs in intervening Central American republics.

## Erpetogomphus eutainia Calvert

(figs. 7, 8, 9-thx, 32, 33-abd, 52-hamules, 75 -penis, 97 -app, 122, 123, 124, 125-ept, 143, 144- face,
150 -vertex, 178 -vl, 199-distr)
Herpetogomphus menetriesii. - Selys, 1878: 429 (24 sep.) ('Guatemala. (Un mâle [sic, female] incomplet. Musée de Paris, une femelle. - Coll. Selys.')).
Erpetogomphus eutainia Calvert, 1905: 162 (10*, Rio Papagayo in Guerrero, Mexico). - Calvert 1919: 35 (comparison with $E$. schaust); Kimmins 1969: 293 (type in BMNH); Paulson 1982: 255 (U.S., Mex., Guat.); Davies and Tobin 1985: 27 (cat.); Tsuda, 1986: 87 (cat.); Bridges 1991: VII. 72 (cat.); Tsuda 1991: 95 (cat.).
Erpetogomphus diadophis Calvert, 1905: 167 (20, Texas). Muttkowski 1910: 87 (cat.); Needham and Heywood 1929: 79 (descr.); Needham and Westall 1955: 147 (descr.); Borror 1963: 104 (common name); Montgomery 1968: 133 (distr.); Kimmins 1969: 293 (type in bMnH); Paulson, 1982: 266 (synonymy of $E$. diadophis and E, eutainia); Bridges 1991: VII. 60 (cat.).
Erpetogomphus? diadophis. - Calvert 1919: 36 (Guat., possible conspecificity with $E$. diadophis)
Herpetogomphus diadophis. - Byers 1928: 5 (larva unknown).
Erpetogomphus (Calogomphus) eutainia - Carle 1992 (key to subgenera, type species of Calogomphus subg. n.).

## Description

Male. - Labium grey to dark grey, labrum all green with a small medial spot on anterior margin (fig. 144) to entirely brown with large central green spot (fig. 143), base of mandibles green, tips becoming black, anteclypeus all green to largely brown; postclypeus green with brown medially along frontoclypeal suture and descending laterally along lateral lobes, thereby isolating medial green spor; in more southerly specimens, brown expanding so that green confined to lateral margins; frons green with brown along frontoclypeal suture and at extreme base of antefrons; vertex brown with medial transverse area at postocellar tubercles green, to entirely brown; occiput brown, transverse triangular pits anterolateral to medial ocellus, postocellar tubercles low, incomplete medially; occiput trapezoidal, largely planar except for medial swelling; crest costate, linear to slightly concave, rimmed with long brown hairs; postocciput green medially to all brown, concave; rear of head brown.
Prochorax brown along middle lobe, remainder green or becoming entirely brown except for pale anterior lobe; synthorax (figs. 7-9) with prominent, well-defined dark brown stripes on green background as follows: middorsal stripe expanding toward collar, divided by pale middorsal thoracic carina, not touching collar, medial 0.5 or entire antealar crest, linear antehumeral, not touching upper margin; humeral, second and third lateral stripes, medial area of mesinfraepisternum; more southerly specimens with these stripes becoming thicker (figs. 8-9) so that antehumeral and humeral connect at lower and upper ends; a vestigial stripe along posterior margin of metepimeron; venter of thorax and metasternum pale grey green to grey. Coxae, trochanters pale green, venter of profemora green, remainder of profemora black; metafemora dark brown to black except for pale area ventrally; basal 0.40 to 0.60 of metafemora pale green to dark grey, becoming black apically; tibiae, tarsi and armature black.

Wings hyaline, venation and pterostigma black; more northerly specimens with pale, narrow line along costa disappearing proximal to pterostigma.

Venational statistics. Fifth, rarely fourth, antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $0-1 / 0-1$; anx: fore wing 11-15/11-15, hind wing 8-11/8-10; pnx: fore wing 8-11/8-12, hind wing 8-11/8-11; cs under pterostigma: fore wing 4-6/4-6, hind wing 4-6/4-6; anal triangular cells: 3-4/3-4. Hind wing length $23-25 \mathrm{~mm}$.

Abdomen (fig. 32) with segment 1 green and a pale brown dorsolateral stripe on each side; segment 2 brown except for green auricles, ventrolateral margin, and middorsal stripe; segment 3 with lateral brown *pe imerruptad medially just posterior to transverse carina, and with an irregular pale spot within
anterior brown spot, posterior brown more prominent posteriorly and meeting dorsally on posterior 0.10 of segment; more southerly specimens with lateral brown not interrupted medially and with pale anterolateral spot reduced or almost absent; incomplete middorsal stripe blue green, pale areas becoming white laterally; segments $4-6$ similar to segment 3 but with dark brown more prominent posteriorly and with anterolateral white spot confluent with ventral margin; segment 7 pale blue green on anterior 0.50 0.60 , except orange brown transverse carina, posterior part of segment dark red brown; segments 8-10 red brown, becoming darker dorsally and ventrally, especially along foliate margins; posterior medial margin of segment 9 smoothly carinate, or with a moderately blunt point. Abdomen 29-32 mm.

Cercus (fig. 97) ivory, becoming brown distally, gently arcuate at distal 0.30 , with a small black tooth, a ventral costate carina on basal $0.35-0.40$ of cercus, often ending in a small, isolated, pebble-like tooth; in other specimens, the posterior part of this carina may be pebble- like; epiprocts (figs. 122-123) brown, moderately curved along posterior 0.50 , tip of epiproct in lateral view thick, bluntly pointed, with lateral costate margin; tips in posterior view (figs. 124125) planar, roundly divergent.

Accessory genitalia. Hamules (fig. 52) smaill, anterior hamule brown, distal 0.25 divided, the superior branch meeting the inferior branch, thus enclosing an oval space; posterior hamule small, digit-shaped, no apical hook; penis (fig. 75) with long pointed cornuae; dorsal membranous hood prominent, its ends parallel; lateral lobes large, frill-like, their margins with spinules; dorsal area of membranous third segment with a pair of mediolateral blunt chitinized tubercles.

Female. - Head as in male with following differences: brown at base of antefrons projecting anteriorly , partially isolating dorsal green; vertex morphology similar to male, postoccipital tubercles small, isolated medially, occiput trapezoidal as in male, planar with faintly convex arcuate crest; postocciput brown, planar.

Thorax as in male.
Venational statistics. Fifth, rarely sixth, antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $0-2 / 0-3$; anx: fore wing 13-16/13-16, hind wing 9-11/8-11; pnx: fore wing 8-12/8-11, hind wing 8-12/8-13; cs under pterostigma: fore wing 4-7/4-7, hind wing 4-7/4-7. Hind wing 2528 mm .

Abdomen (fig. 33) with segment 1 green with a brown dorsolateral stripe; segment 2 similar to segment 1 , but with wash of brown along ventral margin of tergite; segment 3 with dorsolateral stripe darker, expanding posteriorly and connecting above at posterior 0.10 to 0.20 or separated by thin hairline of
green, ventral margin with brown interrupted by white below brown dorsolateral stripe; in more northerly specimens, connecting above with middorsal green stripe; posterior part of brown connecting with dorsolateral stripe posteriorly, transverse carina dark brown; segments $4-6$ similar to segment 3 but brown darker, more definite, connecting posterodorsally on $0.25-0.50$ on progressive segments, lateral white confined to spots, one at anteroventral margin, another ventromedially, these two spots sometimes connected by narrow line of white ventrally; more northerly specimens with medioventral white connecting with middorsal green stripe; segment 7 with anterior 0.50 white except for darkened transverse carina, posterior 0.50 dark brown; segments $8-10$ dark brown; cerci white, their tips black. Vulvar lamina (fig. 178) with plates small, almost digit-shaped; U-shaped cleft about as wide as or wider than each plate; postlamellar ridge with a costate semicircular ridge. Abdomen $29-33 \mathrm{~mm}$.

## Diagnosis

The characteristically shaped posterior hamules (fig. 52) and penile structures (short prepuce, wide spinulate lateral lobe [fig. 75]) separate males of this species from all others except E. leptophis. Erpetogomphus eutainia possesses a straight or pebblelike ventral carina on the basal 0.25 of the cercus (fig. 97); no such carina is present in E. leptophis (figs. 98).

Females of $E$. eutainia and $E$. leptophis are unique in possessing a semicircular postlamellar ridge; in all other congeners, this structure is a Y-shaped ridge. The posteriorly pointed occipital protuberance in $E$. leptophis (fig. 151) is distinct from the relatively unmodified condition present in E. eutainia (fig. 150).

## Remarks

Variation. - Venational details of holotype male of E. eutainia. No marginal cells behind fore wing paranal cells; anx: fore wing $14 / 14$, hind wing $10 / 10$; pnx: fore wing $10 / 9$, hind wing $10 / 10$; cs under pterostigma: fore wing $5 / 5$, hind wing $5 / 6$; number of anal triangular cells: $4 / 4$. Hind wing 24.5 mm .

Venational details of holotype female of $E$. diadophis number of marginal cells behind fore wing paranal cells: $0 / 1$; anx: fore wing $13 / 14$, hind wing $10 / 10$; pnx: fore wing $10 / 9$, hind wing $10 / 11$; cs under pterostigma: fore wing $6 / 7$, hind wing $6 / 5$. Hind wing 28 mm .

This small species has a widespread distribution and, consequently, exhibits some degree of clinal variation north to south. Specimens from southeastern Texas and northern Mexico are the palest: the dark transverse markings of the face are narrower, and green is often present medially on the vertex and occiput. The labrum is mostly green (fig. 144), the dark
thoracic stripes are narrow, and none is present on the posterior margin of the metepimeron (fig. 7). Medial bands of white encircle abdominal segments 3-6 in both sexes. The dark somatic markings are darker, broader, and more definite in specimens from southern Veracruz. Vestiges of a stripe on the posterior margin of the metepimeron can be seen (fig. 8), and the pale white medial abdominal bands are obscured, entirely or almost entirely interrupted by the dorsolateral brown stripe. Specimens from Costa Rica are the most melanic (figs. 9, 143); no green is present on the vertex or occiput, and the middorsal green stripe of abdominal segments 3-6 is always separate from the medioventral white.

The ventral carina of the cercus is also variable. Some specimens have a continuous costate carina with a small, isolated, pebble-like tooth distally; but others may have several antepenultimate pebble-like teeth in addition to the longer ultimate one. The tips of the epiproct may also be variable. In most specimens, these structures are posteriorly flattened (fig. 122 ) with the outer margin slightly to strongly concave (fig. 123). The male from El Progreso, Guatemala, has the epiprocts similar to the condition described above, but they are narrower (fig. 125).

The clinal differences described above probably led Calvert (1905) to interpret the female of $E$. diadophis as a separate species from E. éutainia. Calvert (1919) later received another female from Guatemala, which he postulated may be the same as the paratype of $E$. diadophis, he made no mention of E. eutainia.

The large series of males and females examined from various parts of its range convince me that Paulson (1982) was correct in synonymizing the two names. The status of the two females (not male and female) described by Selys (1878) as Herpetogomphus menetriesii is discussed under the status of $E$. menetriesii. I also examined a female labelled 'Cyanogomphus? / mexicanus/ n. sp. / coll. R. Martin / of', from Honduras, in the mNHP. $_{\text {I }}$.

## Biology

I collected this small, colourful species in southeastern Texas over several years. They were flushed from stubble bordering agricultural fields next to the Gonzales River. They were sympatric with the larger E. designatus, and were about as common. However, the former species behaved more like a damselfly, for they never flew far, rested on tips of stubble or barbs of barbed-wire fences, and were always easy to take with a net. I collected one male which was the prey of a robber fly (Asilidae); another, I rescued from an orb-weaver spider web. I was able to photograph a pair in copula in the late afternoon.

In southern Veracruz at the Rio Otapa, E. eutainia had similar habits, as described under E. ophibolus.

The female from El Salvador (Rio Palio) has the following by V. Hellebuyck: 'on bushes along a fast running river, clear water, rocky bottom \& shady banks'. Altitudinal ranges are from near sea level ( 20 m , Medina Bank, Belize) to 615 m (Oaxaca). Collection dates range from 19 May (Costa Rica) through October (Veracruz, La Gloria; Guerrero, Rio Papagaio).

Distribution (fig. 199). - This species occurs sourheast of the Edwards Plateau in southern Texas, south along the lowlands and foothills of the Sierra Madre Oriental in northern and central Mexico. The farthest western locality is El Sabino in Michoacan state. Specimens have been collected sporadically in Belize, Guatemala, Honduras, El Salvador, and northern Costa Rica.

## Material

Type data. - Holotype male of E. eutainia [labels all printed unless otherwise stated]: 'R. Papagayo, / Guerrero, 1200 ft. / Oct. H. H. Smith.', 'Brit. Mus. / 1911-339.', 'erpetogomphus/ eutainia Calvert [written]/ P. P. Calvert, det. 1905 [written]/ B. C. A. Neur., p. 162 [written]/ Original of Pl. 7, f. 24-27, 39 [all written]', small round label with red border: 'Holo-/ type', in вмnн. Holotype female (not male, as erroneously stated by Kimmins, 1969) of $E$. diadophis: [labels all printed unless otherwise stated]: 'Texas [written by an unknown hand]', pale violet label: 'McLachlan Coll./ B. M. 1938-674.', 'Erpetogomphus $q$ [written]/ diadophis Calv. TYPE [written]/ P. P. Calvert, det. 1905 [written]/ B. C. A. Neur., p. 167 [written]/ original of Pl. VII, ff. 35, 47 [all written]', in BMNH.

Material examined ( $748^{*}, 42$ 早, including holotypes of $E$. eutainia and $E$. diadophis). - U. S. A.: Texas: Caldwell Co.: San Marcos R. at Luling, ( $\mathrm{FSCA}, \mathrm{CC}$ ); Gonzales Co.: Guadalupe River, 4 mi S of Gonzales, ( CC, rwg); near Otting, Palmetto State Park, (FSCA); Palmetto State Park, 6 mi S of Luling, (Rwg); Mexico: Michoacan: El Sabino, 20 mi SSE of Uruapan (Ummz, Rwg); Oaxacaz La Escondida, route 190, km 727, 2000', (UsNM); San Luis Potosi: Tamazunchale, (usnm); Tamaulipas: 3 mi S of Ciudad Victoria, (UsNm); Veracruz Cardel, La Gloria, (UsNm); pond 2.7 mi of La Tinaja, 300 ft . (DRP); Rio La Palma, 25 km N of Catemaco, (rwg); Rio Otapa, 8 km S of La Tinaja, 90 m , (rwg); Sontecomapan, Coscoapan, (unam); Belize: Toledo Dist.: Medina Bank, 20 m , (tb); Guatemala: El Progreso Dept.: San Agustin Ac., (fsCA); Zacapa Dept: Zacapa, (FSCA); unknown locality and date ('Rodig.' [illegible]) [labelled as Herperogomphus menerriesil], (IRSN); unknown locality, date, and collector [labelled 'E. menetriesi? Selys'], (MNHP); Honduras: Unknown locality, date, and collector [labelled 'Cyanogomphus?') mexicanus/ n. sp./ (Coll. R. Martin/ ${ }^{\prime}$ '], (mnhp); El Salvador: La Libertad Deptr: Rio Palio, San Juan Opico, 400 m , (vh); Costa Ruca: Alajuela Prov:: Quebrada Mina 0.2 mi S Hwy, 11 on Hwy, (FSCA); Guanacaste Prov:: Hda. Taboga, 100 ft., (DRP); Rio Santa Rosa, 3.5 mi N of Cañas, 300 ft (DRP).

Erpetogomphus leptophis sp. n.
(figs. 10 -thx, 34, 35 -abd, 53 -hamules, 76 -penis, 98 app, 151-vertex, 179-vl, 200-distr)

Type material. - Holotype male. Belize: Toledo District, I. Z. E. field station, Blue Creek, 25 June 1983 (M. L. May) (fSCA). Allotype female. Belize: Toledo District, Blue Creek Village (Earthwatch) Belize Expedition 1981, 8 June 1981 (D. H. Messersmith, W. H. Steiner, et al.) (USNM).

## Description

Holotype male. - Labium light grey, becoming dark grey medially; labrum green with dark brown margin and medial line almost separating green into two spots; base of mandibles grey-green, anteclypeus green, postclypeus brown except for small green spots above lateral lobes; ventral margin of frons brown, joining brown of postclypeus, thus forming a large frontoclypeal stripe; remainder of frons green; a large transverse furrow at base of antefrons, this area with a pair of deeper pits, each anterolateral to median ocellus; postocellar ridge complete, but with medial part lower and at level of lateral ocelli; extreme base of frons, entire vertex, and occiput dark brown, occiput tumid medially, hind margin mostly straight but peaked medially so that hind margin seems slightly angularly convex; rear of head dark brown.

Prothorax entirely brown except for following green areas: anterior lobe, and two small midlateral spots (almost touching) on median lobe; synthorax (fig. 10) with dark brown thoracic stripes well developed on green background; dark middorsal stripe triangular, its wide base touching collar but not connecting with antehumeral; antehumeral and humeral connected along upper 0.25 , thus isolating small green spot; second lateral as shown in fig. 10, a posterior branch connecting with third lateral along upper 0.30 , isolating a green spot; posterior margin of metepimeron brown, all thoracic stripes connected by brown along antealar sinus; metasternum greybrown. Venter of profemora grey-green, remainder of legs dark brown; tibiae, tarsi, armature black.

Wings hyaline, venation and pterostigma black.
Venational details. Fifth antenodal thickened in all wings; no supplementary marginal cells behind fore wing paranal cells; anx: fore wing 14/13, hind wing 10/10; pnx: fore wing $10 / 11$, hind wing $10 / 10$; cs under pterostigma: fore wing $6 / 5$, hind wing $5 / 5$; anal triangular cells: 4 . Hind wing 29 mm .

Abdomen as in fig. 34. Segment 1 brown, green laterally and with a green middorsal spot; segment 2 brown except for green auricles, ventral margin, and narrow middorsal spot with a cordate expansion in the middle; segment 3 black with green middorsal stripe ending at basal 0.30 of segment and grey ante-
rolateral spot; segment 4 black with anterior 0.2 white laterally, becoming green dorsally; segments 56 like segment 4 ; segment 7 with anterior 0.5 white laterally, becoming green dorsally, posterior 0.5 of segment dark red-brown; segments 8 -10 red-brown, posterior medial margins of segments 8-9 slightly pointed. Abdomen 31 mm .

Cercus (fig. 98) light green, simple, with no ventral carinae, slightly arcuate, tip black; epiprocts brown, gently curved at distal 0.5 , tips viewed posteriorly with parallel sides, forming bluntly rounded tips.

Accessory genitalia. Hamules (fig. 53) small; anterior hamule brown, distal 0.25 divided, the superior branch meeting the inferior branch, thus enclosing an oval space; posterior hamule small, finger-like, no apical hook; penis (fig. 76) with long, pointed cornuae, dorsal membranous hood long, its ends crossing (fig. 76); lateral lobes large, frill-like, their margins with spinules.

Allotype female. - Head as in male except on labrum, brown medial line separating green into two spots; base of antefrons as in male (fig. 151) with a raised bilobed ridge between lateral ocelli; posterior margin of occiput with a prominent, posteriorly directed, pointed protuberance, its base tumid so that entire raised area assumes shape of a plumbob.

Thorax as in male.
Wings hyaline, similar to male.
Venational details. No supplementary marginal cells behind fore wing paranal cells; anx: fore wing $14 / 16$, hind wing $9 / 11$; pnx: fore wing $11 / 13$, hind wing $10 / 11$; cs under pterostigma: fore wing $5 / 4$, hind wing 5/6. Hind wing 25 mm .
Abdomen (fig. 35) with segment 1 brown, becoming dark green-brown laterally; segment 2 brown with a narrow green middorsal stripe and green lateral stripe encircling auricle and extending entire length of segment; segment 3 like segment 2, but brown becoming darker and green lateral stripe abbreviated, forming an elongate spot along basal 0.5 of segment; segments 4-6 all black with white basal band, extending ventroposteriorly along ventral margin of tergites; segment 7 with anterior 0.30 white, remainder dark red-brown; segments 8-10 dark red-brown, cercus white. Vulvar lamina (fig. 179) with plates connected anteriorly, plates wider than in E. eutainia; postlamellar ridge semicircular as in E. eutainia. Abdomen 30 mm .

## Diagnosis

The structure of the hamules and penis ally $E$. leptophis with E. eutainia, but the superior appendages easily distinguish the two species (figs. 97-98). Erpetogomphus eutainia has a pebble-like ventral carina on the basal 0.25 of the cercus; but no such structure is present in E. leptophis. The posteriorly directed
pointed occipital protuberance easily identifies the female of $E$. leptophis. The female shares with $E$. eutainia the semicircular postlamellar ridge, but $E$. leptopbis possesses a bilobed ridge between the lateral ocelli. This surface is planar in $E$. eutainia.

## Remarks

Although the types were taken by different collectors two years apart, the female characters indicate it is the female of $E$. leptophis.
Biology. - Michael May collected the male along a creek in the forest.

Distribution (fig. 200). - Known only from the type locality.

## Erpetogomphus elaphe sp. n.

(figs. 11-thx, 54-hamules, 77 -penis, 99 -app, 180-vl, 200-distr)
Erpetogomphus elaps. - Calvert 1907: 398 (Costa Rica, figures anterior hamules).
Erpetogomphus sp. n. near elaps. - Dunkle 1988: 46 (Honduras).

## Description

Holotype male. - Labrum pale grey green; anteclypeus, labrum, base of mandibles pale green, tips of mandibles dark brown; postclypeus and frons pale green, a slight wash of brown on lateral margins of frontoclypeal suture, antefrons pale green, slightly darker at base, vertex dark brown; a deep transverse trough in front of median ocellus; pedicel, scape, and flagellum brown; occiput trapezoidal, pale green, mostly planar, slightly tumid medially, crest slightly prominent, slightly emarginate medially and covered with long brown hairs; postocciput green, transverse when viewed dorsoposteriorly; rear of head yellow brown.
Prothorax green, becoming brown on middle lobe laterally and anteromedially. Synthorax (fig. 11) apple green with following dark areas: poorly defined middorsal stripe not extending to collar and interrupted medially by middorsal thoracic carina and antealar crest, well defined antehumeral stripe narrowing dorsally but touching antealar crest and extending ventrally to mesinfraepisternum, a small posterior offshoot from antehumeral stripe at upper 0.10 connecting with vestigial first lateral stripe which is abbreviated to upper 0.50 ; venter of synthorax green, coxae and trochanters grey green, femora pale green becoming dark brown on distal extensor surfaces, these dark markings occupying distal 0.80 of profemora, about distal 0.60 of mesofemora, and distal 0.30 of metafemora; tibiae and tarsi black; armature black.
Wings hyaline, anterior margin of costa yellow to pterostigma, thereafter black; yellow at base of costa darkening to brown at costal triangle, pterostigma
brown, veins bordering it black.
Venational details. Fifth antenodal thickened in all wings: number of marginal cells behind fore wing paranal cells: $1 / 1$; anx: fore wing $13 / 13$, hind wing 10/10; pnx: fore wing 9/9, hind wing 9/10; cs under pt: fore wing $6 / 5$, hind wing $4 / 6$; anal triangular cells: $4 / 3$. Hind wing 26 mm .

Abdomen with segment 1 yellow green with a wash of brown dorsolaterally; segment 2 yellow green dorsally and ventrally, with a brown midlateral stripe expanding ventrally behind auricle, anterior 0.5 of this stripe poorly defined, annulus dark brown; segment 3 light yellow green dorsally interrupted laterally by a longitudinal brown stripe well marked at transverse carina, expanding again at posterior 0.25 of segment and connecting dorsally at black annulus; inferior margin of tergite probably ivory (post mortem preservation has partially obscured this area); segments 4-6 similar to segment 3 , with anterior 0.10 of dark midlateral interrupted and with gradual dorsolateral expansion of stripe so that inverted dorsal pale green wedge-shaped middorsal stripe disappears at distal 0.40 of segment 6 ; segment 7 pale green brown becoming tawny on posterior 0.10 of segment; segments 8-10 red brown becoming darker dorsolaterally on segment 8; foliate expansion and denticles black. Abdomen 33 mm .
Cercus (fig. 99) pale ocher becoming dark brown at tip; linear, gradually narrowing distally, tips roundly pointed, a weakly defined ventral carina on distal 0.50 of cercus; epiprocts brown, gently curved at $90^{\circ}$ angle; tips spatulate when viewed posteriorly.
Accessory genitalia (fig. 54). Anterior hamule green basally, becoming dark brown distally, branched at distal 0.50 ; superior (upper) branch greatly enlarged, as long as stem of hamule with resulting interval between branches small, semicircular; posterior hamule (fig. 54) almost digit-shaped, wider at base, with a poorly developed anterior shoulder, tip broadly rounded, without an apical tooth; penis with a prominent knife-like serrated lateral lobe; membranous hood well developed but lobes not overlapping, cornua pointed but with median shoulder as in fig. 77.
Allotype female. - Similar to male in markings and colouration. Head with transverse trough anterior to median ocellus as in male, but occipital region differing as follows: occiput small, erect, barely visible in dorsal view, convex medially; postocciput with median tumid area. Thoracic, leg, and abdominal patterns as in male, but dark areas more restricted; middorsal and antehumeral stripes faint, first lateral stripe vestigial, confined to upper 0.20 of thorax.

Venational details. Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $1 / 2$; anx: fore wing $14 / 15$, hind wing 10/10; pnx: fore wing $10 / 11$, hind wing 11/12; cs
under pterostigma: fore wing $6 / 5$, hind wing $5 / 6$. Hind wing 29 mm .

Vulvar lamina (fig. 180) with basal plate prominent, lobes narrow with medial margins distinctly concave, their bases not touching; distal part of lobe with a ventral flap bent under; its costate rim visible externally and disappearing medially near base of lobe (fig. 180); with resultant cleft forming a wide, Ushaped interval; Y-shaped juncture of postlamellar ridge not extending beyond posterior margin of plate, stem with diagonally ovaloid depression on each side. Abdomen 34 mm .

## Diagnosis

Erpetogomphus elaphe males differ from males of the more northerly $E$. elaps by only two morphological characters (contrasting characters for E. elaps in parentheses): 1) The superior branch of the anterior hamule is robust and is as high as the basal 0.5 of the hamule so that the resulting gap between upper and lower branches is small (fig. 54), (superior branch less robust, thinner; resultant gap between hamular branches wider [figs. 55-56]), and 2) The tip of the posterior hamule is rounded, (tip of posterior hamule with a cephalad directed tooth). The posterior hamule of E. elaphe appears to be more digit-shaped and is reminiscent of those of $E$. eutainia and $E$. leptophis. The same structure appears more triangular in E. elaps. However, this last character is subtle and is best detected when specimens of both species are in hand: I do not consider it a good diagnostic character. I have found no significant differences in the penis or caudal appendages between the closely related and allopatric E. elaphe and E. elaps. I have been unable to find any differences between females of the two species.

Differences between E. elaphe and other similar species parallel those of E. elaps and are discussed under that species.

## Remarks

Variation. - Erpetogomphus elaphe does not show as great variation in thoracic markings as does $E$. elaps. The holotype male is the most boldly marked of the series. All paratype males ( $\mathrm{n}=15$ ), two of which are teneral, have an antehumeral stripe, although it is obscure in one male from Costa Rica. The humeral stripe in all but one of the pararypes is restricted to the upper 0.20 of the humeral suture, and the middorsal stripe is vestigial or absent.

Female paratypes of $E$. elaphe are similar in maculation to the allotype, although one female from Costa Rica has a poorly defined antehumeral stripe. The small humeral stripe is reduced to the upper 0.20 and barely visible.

Venational statistics for paratype males ( $\mathrm{n}=18$ ); number of marginal cells behind fore wing paranal
cells: $0-2 / 0-3$; anx: fore wing $12-15 / 12-15$, hind wing 9-11/9-11; pnx: fore wing 8-11/9-11, hind wing 9-11/9-11; cs under pterostigma: fore wing 3-6/4-7, hind wing 4-6/4-7; anal triangular cells: $3-5$. Hind wing $25-28 \mathrm{~mm}$. Abdomen $30-34 \mathrm{~mm}$.

Venational statistics for paratype females ( $n=6$ ); number of marginal cells behind fore wing paranal cells: $0-4 / 0-3$; anx: fore wing 11-17/14-16, hind wing 10-12/10-12; pnx: fore wing 9-12/10-12; hind wing $9-12 / 9-13$; cs under pterostigma: fore wing 4-6/4-5, hind wing 5-6/4-6. Hind wing $27-30 \mathrm{~mm}$. Abdomen $33-34 \mathrm{~mm}$.

Biology. - Little is known of the biology of this species, though its habits probably mirror those of $E$. elaps. The female from Cartago Province was collected in a 'marshy area by grassy hillside'. Altitudinal gradients for $E$. elaphe range from about 900 m (nr. Camotán, Guatemala) to about 1700 m (SE of Cartago, Costa Rica). Collection dates range from 2 June (Agua Caliente, Guatemala) to 5 August (San Jose, Costa Rica).

Distribution (fig. 200). - Erpetogomphus elaphe is known only from Guatemala, Honduras, and Costa Rica, but it probably exists in El Salvador and Nicaragua. Its distribution is allopatric (or possibly parapatric) with the form of $E$. elaps with thin anterior hamules.

## Material

Type data. - Holotype male: Guatemala, Guatemala Dept., El Fiscal, 6 June 1909 (E. B. Williamson). Allotype female: same data, but 4 June 1909. Both in ummz.

Paratypes examined (188, 7우): Guatemala: Chiquimula Dept.: Alda dos Quebradas nr. Camotán, 900 m (T. W. Donnelly), 1 if (тwD); Guatemala Dept.: Agua Caliente, 2 June 1909 (E. B. Williamson), 1 영 El Fiscal, 4 June 1909 (E. B. Williamson), $1 \delta$, 1 여 5 June 1909 (E. B. Williamson), $1 \delta^{\star} ; 6$ June 1909, $2 \delta^{\star}$ (all ummz); Finca El Rosario, 36.5 km SE of Guatemala City, 15 June 1975 (J. E. Hafernik, Jr.), 2ठ (rwg); honduras: Francisco Morazan Dept.: 30 km ESE Tegucigalpa, 29 June 1985 (Pinto and Ranch), 1 if (swd); 24 July 1983 (L. Cordoba), 1 す' $^{\text {(swD }}$ ); COSTA rica: [no locality data], 1920 (Paul Serre), 30 (мnнp); Cartago Prov., 5 mi SE Cartago, $5500 \mathrm{ft}, 14$ June 1963 (F. G. Thompson), 1 if (Drp); San José Prov:: (all collected by H. Schmidt), San José, 18 June [no year stated], 10 (H. Kahl Coll'n, Acc. 12676); 24 June, 10 ; 4
 21 July, 1 ठे; 5 August, $1 \delta^{\text {A }}$ (FSCA).

## Erpetogomphus elaps Selys

(figs. 12, 13-thx, 55, 56-hamules, 78 -penis, 100 app, 152-vertex, 181 -vl, 200-distr)

Erpetogomphus elaps Selys, 1858: 330 (70 sep.) ( 3 descr. 'Le Mexique, d'après un exemplaire unique du Museum de Paris, rapporté par M. Sal[1]e.'). - Selys 1859: 538 (12 sep.) (descr. ठ̃); Selys 1869: 175 (12 sep.) (descr. ठ, \&); Selys 1873b: 519 ( 75 sep.) (list); Calvert 1905: 163 (localities in Mex.); Calvert 1907: 399 (comparison with E. boa); Calvert 1909: 481 (distr. in Mex.); Muttkowski 1910: 87 (cat.); Ris 1917: 153 (mentions Calvert's [1907] comments on comparison with E. boa); Williamson and Williamson 1930: 13 (summary of status); Montgomery 1973: 239 (derivation of name); Paulson 1982: 255 (Mex., Guat., Costa Rica); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII, 68 (cat.); Tsuda 1991: 95 (cat.).
Gomphus elaps. - Hagen, 1861: 100 (descr. ©)
Herpetogomphus elaps. - Hagen 1875a: 42 (cat.); Selys 1879: 64 (note on classification); Kirby 1890: 60 (cat.); Needham 1897: 182 (name mentioned); Calvert 1899: 386 (descr. $\delta$, comparison with E. viperinus auct.), 415 (gizzard); Higgins 1901: 128 (mentions Calvert, 1899).

## Description

Male. - Labrum grey white with wash of brown around margin of median lobe; anteclypeus, labrum, base of mandibles pale grey green, tips of mandibles dark brown; postclypeus and frons light green, some specimens with brown at base; vertex dark brown; a deep transverse trough in front of median ocellus, a small tubercle behind each lateral ocellus; pedicel, scape and flagellum brown; occiput trapezoidal, light green, mostly planar; crest slightly prominent, straight, slightly notched medially, and covered with long brown hairs; postocciput green, transverse when viewed dorsoposteriorly; rear of head yellow brown, paler laterally.

Prothorax and median lobe red brown becoming pale apple green medially, especially on anterior and posterior lobes. Synthorax ranging from entirely apple green (fig. 13) with slight hint of dark antehumeral stripe to having following brown areas (fig. 12): poorly defined middorsal stripe divided by pale middorsal thoracic carina, well-defined antehumeral stripe just touching antealar crest dorsally and not touching mesinfraepisternum ventrally; a vestigial first lateral stripe confined to upper $0.30-0.40$ of suture and sometimes with a smail anterior offshoot touching antehumeral; venter of synthorax, coxae, trochanters, grey green; femora pale green, becoming dark brown on distal extensor surfaces; these dark markings occupying distal 0.80 of profemora, distal 0.50 of mesofemora, and distal 0.20 to 0.40 of metafemora; tibiae and tarsi dark brown to black; armature black.

Wings hyaline, anterior margin of costa yellow to pterostigma, thereafter black; yellow at base of costa darkening to brown costal triangle; pterostigma dark brown, veins bordering it black.

Venational statistics. Fifth antenodal (occasionally fourth or sixth) thickened in all wings; number of marginal cells behind fore wing paranal cells: $0-2 / 0-3$;
anx: fore wing 11-15/12-15, hind wing 9-11/8-11; $p n x$ : fore wing 8-11/8-11, hind wing 8-11/8-11; cs under pr: fore wing 4-6/4-7, hind wing 4-6/4-7; anal triangular cells: 3-5. Hind wing 24-29 mm.

Abdomen with segment 1 yellow green with a wash of brown dorsolaterally; segment 2 yellow green dorsally with a large midlareral brown spot dorsoposteriorly to auricle, or, in well- marked specimens, a distinct dark brown midlateral stripe expanding ventrally behind auricle, annulus dark brown; segment 3 broadly light yellow green dorsally interrupted lateratly by a longitudinal brown stripe wellmarked at transverse carina and flaring again at posterior 0.25 of segment to annulus; in palely marked specimens, brown midlateral stripe interrupted behind transverse carina; in well-marked specimens, midlateral stripe broad and meeting dorsally on posterior 0.10 of segment; inferior margin of tergite ivory, in heavily marked specimens limited to a triangular spot an anterior 0.10 of segment; segments 4-6 similar to segment 3 , but some specimens with dark midlateral stripe more prominent and not interrupted behind transverse carina; in heavily marked specimens, midlateral stripe almost black and covering entire segment, except for basal pale annulus at basal 0.10 of segment and dorsal inverted pale green wedge- shaped middorsal stripe gradually narrowing posteriorly and disappearing at distal 0.20 of segment 4; these dorsal pale areas lacking on segment 5 and 6; segment 7 pale green, becoming tawny on posterior 0.10 of segment, or with posterior 0.20 dark brown, becoming black dorsally, transverse carina brown; segments 8-10 generally red brown becoming darker dorsally; in darkly marked specimens becoming almost black dorsally; foliate expansion and denticles black. Abdomen 29-38 mm.

Cercus (fig. 100) pale ocher to light brown, linear, gradually narrowing distally, the tips roundly pointed, a weakly defined ventral carina on distal 0.50 of cercus; epiprocts brown, gently curved at a $90^{\circ}$ angle, tips when viewed posteriorly slightly spatulate, broadly rounded, or obliquely truncate.
.Accessory genitalia (fig. 55). Anterior hamule dark brown to black, branched at distal 0.30 , superior branch larger than inferior branch; superior branch moderately thickened in specimens from northern Mexico sourh to Chiapas, Mexico; specimens south of Chiapas (fig. 56) with superior arm of hamule more slender (see remarks); posterior hamule (figs. 55-56) pale, roughly triangular, with an anterior shoulder, tip with a tooth pointed cephalad; penis with prominent, knife-like serrated lateral lobe; membranous hood well developed but lobes not overlapping, cornua pointed but with median shoulder as shown in fig. 78.

Female. - Similar to male in markings and colour-
ation; head with transverse trough anterior to median ocellus as in male, but occipital region differing as follows: occiput small, planar, encompassed posteriorly by arcuate (convex) crest (fig. 152); postocciput with median tumid area. Thoracic, leg, and abdominal patterns as in male, but dark areas more restricted, especially on femora and abdomen; cerci pale. Abdomen $30-36 \mathrm{~mm}$.
Venational statistics. Fifth (rarely sixth) antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $0-3 / 0-3$; anx: fore wing 12-16/13-16, hind wing $9-12 / 9-12$; pnx: fore wing 8 $11 / 9-12$, hind wing $8-12 / 9-12$; cs under pt: fore wing $5-6 / 5-7$, hind wing 5-7/5-7. Hind wing $26-30 \mathrm{~mm}$.

Vulvar lamina (fig. 181) with basal plate prominent, lobes narrow with medial margin distinctly concave, their bases not touching, with resultant cleft forming a wide, U-shaped interval; apical 0.30 of plate with mesal margin folded ventrally, beneath main plate, the costate rim sometimes visible along lateral margin of plate (fig. 181); Y-shaped juncture of postlamellar ridge not extending beyond posterior margin of plate, stem with diagonally ovaloid depression on each side.

## Diagnosis

Erpetogomphus elaps and E. elaphe are closely related species which are diagnosed under $E$. elaphe. In males, the linear cercus of $E$. elaps (fig. 100) easily distinguishes it from the decumbent cercus (fig. 102) of E. bothrops. Body colouration is similar in E. elaps and E. bothrops, but all $E$. bothrops have well-marked antehumeral and humeral stripes (fig. 15). Many specimens of E. elaps have these markings poorly developed or nearly absent (fig. 13).

With the exception of E. elaphe, whose females appear indistinguishable, females of $E$. elaps are most similar to $E$. bothrops. The differences in the vulvar lamina as discussed by Calvert (1899) are an easy way to distinguish them. Secondly, the occiput of E. elaps is small and convexly arcuate (fig. 152); in E. bothrops, this structure is wider (fig. 154) and not nearly as arcuate.

## Remarks

Variation. - The description is based on 49 males and 21 females. The holotype is in fair condition, although most of dorsum of the prothorax and part of the mesepisternum have been eaten away by dermestids. It has a completely pale thorax and the black midlateral stripes on abdominal segments 3-6 are interrupted only on the basal 0.10 of each segment. The hamules (fig. 55) and cerci (fig. 100) show that this specimen is the apple-green species ranging from northern Mexico south through Guerrero and into southern Veracruz.

Venational details of holotype male: number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing $12 / 12$, hind wing $9 / 10 ;$ pnx: fore wing $9 / 8$, hind wing $8 / 9$; cs under pt: fore wing $4 / 4$, hind wing 5/5; anal triangular cells: $3 / 4$. Hind wing 24 mm . Abdomen 31 mm .

I consider $E$. elaps to be the most variable of any species in the genus. Calvert (1907) figured three forms of the anterior hamules of this species when he examined a male from San Jose, Costa Rica $(=E$. elaphe). His fig. 31 (Guerrero, Rincon) and fig. 32 (Morelos, Cuernavaca) correspond to the holotype of E. elaps (fig. 55), which is characterized by possessing a moderately thickened anterior branch whose tip almost overlaps the posterior branch when viewed laterally. Calvert's fig. 30 (Guerrero, Rincon) is typical of specimens I have seen from Chiapas, Mexico, and northern Guatemala (fig. 56). The anterior branch forms a slender hook which rarely overlaps the posterior branch when viewed posteriorly. Figures 33 and 34 of Calvert (1907) represent E. elaphe and are discussed under that species. Calvert (1907) stated that he was unable to find any other characters which varied correlatively, and I have found no other characters to separate E. elaps and the narrow hamular form. I have not found any intermediate conditions in the shape of the anterior hamule. Their distributions appear to be largely allopatric, although Calvert (1907) figures both forms from Rincon in Guerrero.

Interestingly, the distribution of the form with the thin anterior hamular branch lies between typical $E$. elaps and $E$. elaphe. Because I have been unable to find any other diagnostic characters, I am inclined to treat the Rincon male and others from Chiapas and northern Guatemala as a variant of $E$. elaps. The divergent morphology of southernmost $E$. elaps may represent character displacement between $E$. elaps and $E$. elaphe.

Both males and females show remarkable range in size and maculation patterns throughout its range. The smallest specimens examined are from Morelos (HW: males 23 mm , females 26 mm ), the largest from the Rio Metlac region of Veracruz (HW: males 29 mm , females 31 mm ).

Specimens collected at 1220 m in Sinaloa State are melanic, characterized by pale areas of abdominal segments $4-6$ reduced to basal 0.10 laterally; pale areas on dorsum of segments 3-6 reduced laterally, this pale area a thin hairline on segment 5 , a basal ring on segment 6 .

I find no correlation of presence or absence of the dark thoracic stripe to locality. A series of specimens from Palapita, Nayarit, ranges from an entirely pale thorax to possessing antehumeral and humeral stripes. All specimens from Veracruz that I have examined lack dark thoracic stripes, but all southern variants with the thin anterior hamular branch from

Chiapas and northern Guatemala possess them.
Biology. - This small species has habits similar to those described for $E$. bothrops. The following field notes accompany specimens collected by E. B. Williamson: Nov. 6 (Tepic): ' $q$ Gomphine - from brushy vegetation between road and river at rapids above mill'; Nov. 7: 'Gomphine o - Have seen these only in bushes along bank of Rio de Tepic - usually not over foot above ground - resting or flying. One $\&$ was ovipositing in rather still water below the rapids'; Nov. 14 (Jalisco: San Diego Rancho): 'Very hard to catch in net. Sits tight on rock and lets it [net] pass over and then flies. Easily approached and I finally began catching them by dropping the net over their rock \& letting them rise into the bag.'; and Nov. 16 (Jalisco: San Diego Rancho): ' [ 0 ,,$\uparrow]$ caught in grass in banana patch early in AM, sluggish'.
López (in litt.) records it as one of the most common members of the genus in Mexico, having a wide tolerance for diverse stream habitats. I have seen no records of this species taken near sea level, unlike $E$. bothrops. Altitudinal records range from 670 m (Nayarit, Palapita) to about 1500 m (Cuernavaca), dates of capture from 9 June (Veracruz, Teocelo) to 23 November (Jalisco, San Diego Rancho).

Distribution (fig. 200). - Both this species and E. bothrops appear to be the most widespread members of this genus in Mexico. Erpetogomphus elaps occurs in mesic areas on either side of the Sierra Madre Oriental and Occidental. It has been collected as far north as Nuevo León in the east, and southern Sinaloa in the west, southward through Jalisco, Morelos, Guerrero, and Veracruz.

## Material

Type data. - Holotype male with following data: small round white label with handwritten ' $2 / 44$ '; rectangular manila label with handwritten (in R . Martin's hand) 'Ophiog. elaps/ of De Selys/ Mexique? a renvoyer.'; large green label with black border with handwritten 'E. elaps, Selys/ Mexique'; small white label with printed 'museum paris'; narrow red label with printed 'ноLotype'; white printed label with 'E. elaps holotype/ Det. J. Belle, $1977 /$ No. 2, 1844. Insects [sic] de/ [reverse side] differents Ordres envoyes/de Mexique par/ Mr. Ghresbreght!' and in lower right hand corner 'ne pas/ Sal[1]e!' In MNHP. The caudal appendages and accessory genitalia are shown in figs. 55, 78, 100.

Other material ( 49 ô, 21 ㅇ). - Mexico: Chiapas: Pacific slope, $800-1000 \mathrm{~m}$ ), (USNM); stream 20.1 mi NE Tapanatepec, Oaxaca, $2700 \mathrm{ft}$. ( FSCA ); San Geronimo, Volcan Tacana, 450 m (UMmz); Distrito Soconusco, Finca Juarez, (Ummz); Guerrero: Chilpancingo, $4600^{\prime}$, (FSCA, BMNH); Tepetlapa, 3000 ft., (bмnн); Jalisco: San Diego Rancho nr. Cocula,
(UMMZ); 15 mi on road to Tequila, (UMMZ); Michoacan: El Sabino, 20 mi SSE of Uruapan, (ummz); Morelos: Alpuyeca, S of Cuernavaca on Rt. 95, (FSCA); Cerro de Higuerón, Jojutla, (UNAM); Cuautla. (ummz); Cuernavaca, ca. $1200-1500 \mathrm{~m}$, (UMMZ, uSNM, RWG); 5 mi S of Cuernavaca, (UMmz); 16 mi S of Cuernavaca, (UMmz); Itzamatitlan, 5 km SW of Yautepec, (rwg); Nayarit: Acaponeta, (bmnh); Jumatán, (unam); Tepic, (ummz, usnm); Jalisco: Palapita, 670 m (UNAM); Nuevo León: Municipio de Monterrey, Cerro El Mirador, (FSCA); Sinaloa: stream 27.9 mi NE of Concordia, 4000 ft ., (DRP, FSCA, CC, RWG); Veracruz: Barranca de Cayoapa, Teocelo, (UNAM); Jalapa, (UsNm); 4.9 km N of Coscomatepec by Mex. Hwy 156 (rwG); Rio Merlac, 2 km WNW of Fortin, 900 m , (RWG); Guatemala: Suchitepequez Dept.: Finca Moca Grande, Rio Bravo, (FSCA).

## Erpetogomphus liopeltis sp. n.

(figs. 14-thx, 36-abd, 57-hamules, 79-penis, 101app, 117 -cercus, 126 -ept, 140,141 -occiput, 153 vertex, 182-vl, 201-distr, 227-wings)
Erpetogomphus elaps. - Novelo \& Peña, 1991: 130 (misidentification).

Type material. - Holotype male: Mexico: Nuevo Leon State, Municipio de Monterrey, La Estanzuela, 2 July 1987 (M. J. Westfall, Jr., H. Quiróz, A. Contreras). In fsca. Allotype female: Nuevo Leon State, Monterrey, El Diente, 5 mi SE, 2 July 1960 (W.B. Cutts). In ummz.

Paratypes examined ( $150^{\circ}, 3$ 여, including holotype 3 and allorype ) ). - Mexico: Hidalgo: Pemuxtitla, Rio Zacuala, 1000 m, 22 April 1984 (R. Novelo), 10 © (RWG); Michoacán: 12 km W Apatzingán, 369 m (1200 f.). , 4th Hoogstraal Mexican Biological Expedition, 12 Aug. 1941 (Harry Hoogstraal), 1 아 (Ummz); Nuevo Leon: Chipingue, 12 July 1960 (W.B. Cutts), 1 ㅇ (RwG); Municipio de Monterrey, Cerro El Mirador, 9 Aug. 1984 (A. Contreras), $2 \mathbf{\delta}^{\circ}$ (rori); same data as holotype, 30 ( FSCA ); Municipio de Santiago, El Cercado, Arroyo Dolores, 17 Aug. 1984 (A. Contreras), $2 \delta^{\hat{\sigma}}$ (rorr); Municipio de Santiago, Rancho Los Pinos, 3 July 1987 (M. J. Westfall, Jr., Jr., G. Luna, A. Contreras), 50 (FSCA, RWG); San Luis Potosí́ La Conchita (Camino a Xilitla), 7 May 1950 (no collector), $10^{\text {º }}$ (cc).

## Description

Holotype male. - Entire head pale green, labrum dull grey green, base of mandible, anteclypeus dull grcen, tip of mandible black; basal 0.25 and all of vertex brown, a transverse trough at anterior margin of vertex at juncture of base with antefrons, this trough
with two slightly deeper pits, one each anterolateral to median ocellus; occiput with slightly medial tumid area, occipital crest distinctly emarginate medially (figs. 140-141), dark brown, rimmed with long dark brown hairs; postocciput green, becoming black laterally, concave medially; rear of head brown, becoming yellow brown laterally; antennae dark brown.

Prothorax brown, anterior and posterior lobes and small dorsolateral area of median lobe pale green. Thorax (fig. 14) entirely green (probably blue green in life) with following dark brown: obscure middorsal stripe ending before collar, becoming darkest near middorsal thoracic carina, lateral margins of this stripe gradually fading to green on mesepisternum; posterolateral rim of antealar crest; lateral 0.40 of mesepisternum except for dorsal emargination, with no indication of pale green separating coalesced humeral and antehumeral stripes; this stripe narrowing slightly ventrally but connecting with lateral margin of collar and anterior 0.50 of mesinfraepisternum; rim below subalar carina at obsolete second lateral suture; narrow partial third lateral stripe extending ventrally to 0.50 of suture before merging with green. Metasternum pale grey brown, tawny anteriorly. Coxae, trochanters grey; base of femora brown, becoming black distally; tibiae and armature black.

Wings hyaline, anterior margin of costa dark brown at base, becoming yellow distally to proximal level of pterostigma, remainder of venation and pterostigma black.
Venational details. Fifth antenodal thickened in all wings; no supplementary marginal cells behind fore wing paranal cells; anx: fore wing $14 / 15$, hind wing $10 / 10$; pnx: fore wing $8 / 9$, hind wing $9 / 9$; cs under pterostigma: fore wing $7 / 6$, hind wing $6 / 5$; anal triangular cells: $4 / 3$. Hind wing 31 mm .

Abdomen (fig. 36) with segment 1 green dorsally, paler laterally with a midlateral stripe of brown expanding anteriorly at base and darkening posteriorly at articulation point; segment 2 similar to segment 1 but with a ventral stem of brown behind auricle from dark midlateral stripe, midlateral brown becoming wider and darker along posterior 0.50 of segment, annulus black; segment 3 with pale green middorsal stripe acuminate and disappearing at posterior 0.10 of segment, black midlateral stripe expanding slightly at transverse carina and again widening at posterior 0.10 of segment, this area with a ventral offshoot terminating anteriorly at 0.20 of segment; segments 4-6 similar to segment 3 , but with middorsal pale green stripe shorter and narrower on each succeeding segment, thus black midlateral stripe connecting dorsally at posterior 0.40 of segment 6 , these midlateral stripes interrupted at anterior 0.10 of each segment with small longitudinal wedge-shaped white area along ventral margin of tergite behind transverse cari-
na; segment 7 with basal 0.60 pale grey green, except for irregular diffuse black spot at transverse carina, remainder of segment red brown, becoming black dorsally; segments $8-10$ red brown, becoming darker dorsally; carinae and lateral foliations black, posterior medial margin of segment 9 pointed.

Cercus (fig. 101) pale ocher, decumbent at posterior 0.30 , largely parallel in lateral view except for distal 0.20 , tip with a shiny black tooth; black inferior carina on distal 0.25 ; epiproct brown, about 0.45 as long as cercus, U-shaped in lateral view, tip of epiproct when viewed posteriorly (fig. 126) broadly truncate. Abdomen 40 mm .

Accessory genitalia (fig. 57). Anterior hamule dark brown, branched at distal 0.30 , superior branch larger than inferior branch; posterior hamule pale, roughly triangular with a well developed anterior shoulder, distal 0.75 curved posteromedially with a well developed black tooth twisted laterally from broad axis of hamule; penis with cornuae acuminate as shown in fig. 79, lateral lobe prominent, knife-like, serrated posteriorly.

Allotype female. - Similar to male with following differences: labrum slightly darker (possibly due to post mortem effects); transverse trough at juncture of antefrons and vertex well defined with slightly larger and deeper pit anterolateral to median ocellus; occiput strongly reduced to a small, gently convex, largely perpendicular medial arch; postocciput tumid medially, dark middorsal stripe vestigial, with only a wash of brown on either side of lower arms of antealar sinus, a narrow poorly defined incomplete green stripe separating dark brown antehumeral and humeral stripes up to dorsal 0.75 . Abdominal segment 3 with acuminate black spot posterolateral to black dorsolateral stripe projecting anteriorly to black transverse carina; dorsolateral black stripes on segments 47 more extensive; pale middorsal stripe reduced to basal 0.75 (segments 4-5) to basal 0.50 (segment 7), and anteriorly touching annulus on each segment; segments 8-9 black dorsally, red brown laterally. Abdomen 38 mm .

Vulvar lamina (fig. 182) small, cleft larger than each plate, each plate connected basally, each with a raised shelf occupying anterior 0.5 , postlamellar ridge with Y-shaped juncture posterior to hind margin of lamina, central stem with a well defined circular depression on either side.

Venational details. Fifth antenodal thickened in left fore wing; the sixth in remaining wings; Number of marginal cells behind fore wing paranal cells: 2/3; anx: fore wing $16 / 17$, hind wing 13/11; pnx: fore wing $10 / 12$, hind wing $12 / 11$; cs under pterostigma: fore wing $6 / 7$, hind wing $6 / 7$. Hind wing 34 mm .

## Diagnosis

This remarkable species approaches $E$. viperinus in size, colour, and maculation, but is most closely related to $E$. bothrops in morphology. Males of $E$. liopeltis and $E$. bothrops are similar in possessing epiprocts with broadly truncate tips (figs. 126-128); but those of $E$. bothrops are usually, but not always, bidentate (fig. 128). In lateral view, the epiprocts of $E$. liopeltis are more strongly curved and extend about 0.5 or less the length of the cercus; the same structures in $E$. bothrops extend 0.75 or more the length of the cercus (fig. 102). The apical tooth of the posterior hamule of $E$. liopeltis is longer and not as tightly recurved as in $E$. bothrops (figs. 57-58). Abdominal segments 1-3 in E. liopeltis (fig. 36) are more heavily marked than in E. bothrops (fig. 37); and the dark thoracic markings are also more extensive (fig. 14) than in E. bothrops (fig. 15).

In size and overall colouration, E. liopeltis seems indistinguishable from $E$. viperinus, but differences in the tips of the epiproct (figs. 126, 129) and structure of the area anterior to the vertex in both sexes (figs. 136 [as for E. bothrops], 137) easily distinguish the two species. Subtle differences between these two species exist in the morphology of the distal 0.5 of the cercus. In E. liopeltis, these structures are thicker and more cylindrical when viewed dorsally (fig. 101) and mediodorsally (fig. 117). The cercus of E. viperinus is more laterally compressed (fig. 119). The tip of the cercus in E. liopeltis terminates in a sharp black tooth; this tooth is lacking in E. viperinus.

Females of $E$. liopeltis and $E$. bothrops can be separated by overall size ( $E$. liopeltis hind wing $32-34 \mathrm{~mm}$; E. bothrops hind wing $28-31 \mathrm{~mm}$ ) and thoracic maculation and colouration, as stated for males. Structural differences of the occiput and postocciput easily distinguish these species. The occiput of E. liopeltis is rudimentary, with only a gently rounded vertical medial crest. The medially tumid postocciput is easily visible in dorsal view (fig. 153). In E. bothrops, the occiput is narrow, but the anterior and posterior margins are roughly parallel except for the angulate lateral margins (fig. 154). In E. bothrops, the crest is vertical and the exposed postocciput is easily visible; but its posterior margin is planar or only slightly tumid. Receptacles for the tips of the male cerci differ slightly between the females. The transverse trough located anterior to the median ocellus narrows laterally in E. bothrops (fig. 136); but the lateral area of the greater U-shaped trough tends to curve lateroposteriorly around the anterior margin of the median ocellus in E. Liopeltis (as in fig. 139).

Like males, females of E. liopeltis and E. viperinus can show striking similarities, but greater morphological differences manifest themselves in structures of the head and vulvar laminae. The trough anterior to the median ocellus of $E$. liopeltis is unlike the antero-
lateral pits of $E$. viperinus (fig. 137). The occiput of $E$. viperinus (fig. 155 ) is much broader, and the anterior margin of the occiput and curvilinear occipital crest are roughly parallel. The median postoccipital swelling of $E$. liopeltis is absent in E. viperinus. The vulvar laminae of the two species are similar, but the Yshaped postlamellar ridge generally arises at or just before the posterolateral margins of the lamellar plates in E. liopeltis, the postlamellar ridge usually surpasses the vulvar lamina in E. viperinus. The median cleft between the vulvar lamellae is more obtuse in $E$. viperinus (fig. 184) than in E. liopeltis (fig. 182).

Erpetogomphus liopeltis and E. viperinus are allopatric and $E$. viperinus generally has more dense venation, despite wing lengths (E. liopeltis hind wing 29$31 \mathrm{~mm}[\mathrm{n}=13$ males, 3 females], E. viperinus hind wing 29-31 mm [ $\mathrm{n}=20$ males, 20 females $]$ ). The number of marginal cells behind fore wing paranal cells in male E. liopeltis ranges from 0-2 (only one wing has 3), compared with 1 (in 4 wings only) to 4 in $E$, viperinus.

Both sexes of $E$. liopeltis and $E$. elaps are easily diagnosed in the key by differences in body colouration, shape, and position of the posterior hamular tooth and shape of the cercus (males) and shape of the vulvar laminae (females). The occiput of females of $E$. claps, E. elaphe, and $E$. liopeltis are similar in being reduced to a perpendicularly raised medial area (figs. 152-153) and a medial tumid area on the postocciput.

## Remarks

Variation. - I found little variation among the 12 paratype males and 2 paratype females. The hind margin of the occiput (crest) is medially concave in the holotype (fig. 140), but ranges to almost linear (fig. 141) in other specimens. A poorly defined narrow green stripe is present between the coalesced dark antehumeral and humeral thoracic stripes in 7 males, but this condition varies from a short, narrow mark to one occupying the medial 0.50 of the lateral margin of the mesepisternum.

Wing variation among paratypes: Males: anx: fore wing 13-15/13-16, hind wing 9-12/10-12; pnx: fore wing 8-10/8-10, hind wing 9-13/9-12; cs under pterostigma: fore wing 5-7/4-6, hind wing 4-6/5-7; anal triangular cells: $3-4 / 3-4$. Hind wing $29-31 \mathrm{~mm}$. tbdomen $37-40 \mathrm{~mm}$.

Females: Anx fore wing 16/14-15, hind wing 10/10; pnx fore wing 8-10/9-10, hind wing 11/9-11; cs under pterostigma fore wing $6-7 / 6$, hind wing 6 --o.
Other aspects of wing venation variability are charicterized under the diagnosis for $E$. liopeltis and $E$. vi:"rmus.
some ( $\mathrm{n}=5$ ) males have no marginal cells behind
fore wing paranal cells, while two others (fig. 227) have one marginal cell in only one wing. All females have at least two such cells, except for the left fore wing of the Chipingue female. The fifth antenodal is almost always thickened; but the sixth is thickened in two males from La Estanzuela and in the right fore wing of the Apatzingan female.

One male from La Estanzuela has aberrant hind wing venation and was not included in the measured samples. Most specimens, including the holotype, had been preserved in acetone; and their resultant preservation leads me to believe that the pale colouration is similar to the blue green of $E$. viperinus.

Biology. - Novelo (pers. comm.) misidentified this species as E. elaps in Novelo and Peña (1991). They collected E. liopeltis in Hidalgo state from 22 April to 25 July (Rio Zacuala, Pemuxtitla) and 27 July (Calnali). They were taken as they perched on rocks along narrow, shaded, shallow, rocky streams. Erpetogomphus liopeltis was collected at Cerro El Mirador with E. elaps. Data for the female collected in Michoacan indicate that it was collected on 'side of stream'. Collection dates range from 22 April (Rio Zacuala) to 17 August (Arroyo Dolores).

Distribution (fig. 201). - Erpetogomphus liopeltis is only known from the states of Hidalgo, Michoacan, Nuevo Leon, and San Luis Potosi in northeastern Mexico.

## Erpetogomphus bothrops sp. n.

(figs. 15 -thx, 37 -abd, 58 -hamules, 80 -penis, 102app, 118 -cercus, 127,128 -ept, 136 -base of
postfrons, 154 -vertex, 183 -vl, 201-distr, 228-wings)
Erpetogomphus viperinus, - Calvert 1899: 385 (described as E. viperinus Selys); Calvert 1905: 163 (described as E. viperinus Selys); Calvert 1909: 35 (misidentified as $E$. viperinus, compared with E. schausi); Muttkowski 1910:87 (in part, cat.); Williamson \& Williamson 1930: 11 (in part, refers to true E. viperinus but specimens in E. B. Williamson Coll. misidentified); Calvert 1947: 608 (status of Tepic specimens); Paulson 1982: 256 (in part, as $E$. viperinus from Mex., Guat.); Davies and Tobin 1985: 28 (in part, cat.); Tsuda, 1986: 87 (in part, cat.); Maes et al. 1988: 36 (as E. viperinus from Nicaragua); Tsuda 1991: 95 (in part, cat.).

Type material. - Holotype male: Mexico: Veracruz: Rio Otapa, 8 km S of La Tinaja, elev. 90 m , 13 Aug. 1976 (R. W. Garrison). Allotype female: same data, but 20 Aug. 1976. In usnm.
Pararypes examined ( 61 ô, 28 ) ). - Mexico: Guerrero: Dos Arroyos, 1000 ft., Sept. 1888 (H. H. Smith), 1 ㅇ (вмnн); Jalisco: San Diego Rancho near Cocula, 14 Nov. 1923 (J. H. Williamson), 10 त (bmnh); Michoacan: El Sabino, 20 mi SSE of Uruapan, 27 July, 1 Aug. 1936 (H. Devlin Thomas), 8 ô, 3 ㅇ (ummz, fSCA); Morelos: Puente de Ixtla, 3

July 1900 (C.C. Deam), 1 ㅇ (ummz); Cerro del Higuerón-Jojutla, July 1983 (C. Deloya), 1 ठ (unam); Nayarit: Acaponeta, 1-2 Nov. 1923 (J. H. Williamson), 9 §̂, 1 ㅇ (ummz); Tepic, 7 Nov. 1923 (J. H. Williamson), $1 \delta^{+}$(ummz); Jumatán, 10 Sept. 1980 (G. Jiménez), 2 ơ (Unam); San Luis Potosi, $^{\text {a }}$ Cascadas Micos nr. aqueduct, 27 June 1990 (K. J. Tennessen), 1 (KJT); El Salto, $400 \mathrm{~m}, 6$ Sept. 1963 (T. W. Donnelly), $1 \delta$, 1 if ( Twd ); Huichihuayan, km 410, S of Valles, 'Sam Brown' Hacienda, 25 Sept. 1938 (L. J. Lipovsky), 1̊̂, 2 우 (Ummz); Tamaulipas. Rio Corona, 20 mi N of Ciudad Victoria, nr. Mex. Hwy 101, 1000 ft., 26 July 1968 (R. W. Garrison), $10^{\text {th }} 29$ (RWG); Veracruz: Atoyac, 400 m , (Schumann), 1 ㅇ (вмNн); 16.5 miS of Catemaco, by Hwy 180, 25 June 1985, (I. S. Askevold), 20 (CC); Cordoba 12-25 July 1964 (E. Fisher, D. Verity), 1 ㅇ (Lacm); Isla, 17 July 1969 (R. Wind), 10 (cc); K375-390 Cordoba Rd., 11 Aug. 1961 (R. and K. Dreisbach), $1 \delta^{\star}$ (USNM); 4.9 km N of Coscomatepec, by Mex. Hwy 156, 11 Aug. 1976 (R. W. and J. A. Garrison), 1 ( RWG ); Rio Otapa, 8 km S of La Tinaja, 90 m, 13, 20 Aug. 1976 (R. W. and J. A. Garrison), $10 \delta^{\hat{*}}, 2$ ( PWG ); Rio Hondo, on road to 'Colonia la Apachital' ( 50 '), Soteapan, 500 m , JulyAug. 1990, (collector unknown), 6ठ, 49 (PM, RWG); 6 mi E of Hwy from Tierra Blanca to Ciudad Aleman, 25-26 Aug. 1957 (G. H. Beatty, III), 10 đ (FSCA); Salto Eyipantla, 8 km S of San Andres Tuxtla and Mex. Hwy 180, 15 Aug. 1976 (R. W. and J. A. Garrison), $2 \delta^{\star}$ (Rwg); Tierra Colorada, nr. Veracruz, 17 July 1932 (H. M. Smith), 1 if (ummz); Oaxaca: Candelaria Loxicha, $500 \mathrm{~m}, 7$ Sept. 1973, (E. C. Welling-M.), $1 \delta^{\star}$ (cc); 5 July 1974, (E. C. WellingM.), $10^{\text {º }}$ (swd); Chiapas. river 26.1 mi NE Tapanatepec, 2100 ft., 2 Aug. 1965 (D. R. Paulson), $10^{\circ}$ (DRP); stream 15.4 mi NE Arriaga on Mex. 195, 2300 ft., 24 July 1965 (D. R. Paulson), $2 \delta^{\star}$ (DRP); El Aguacero, nr. Ocozocoautla, 26 Oct. 1986 (E. Fisher), $1 \begin{aligned} & \text { (r) (rwg); Guatemala: Baja Vera Paz Dept.: }\end{aligned}$ San Geronimo, 1879-80 (G. C. Champion), 10 (вмлн); Chiquimula Dept.: streams vic. Tierra Colorado, 800 m, 20 July 1962, (T. W. Donnelly), $1 \delta^{\top}, 2$ (TwD); small stream vic. Veguitas, $600 \mathrm{~m}, 10$ July 1962 (T. W. Donnelly), 1才, 1 와 (тwD); Dept. Zacapa: Gulan, 16-17 June 1909 (E. B. Williamson), $2 \sigma^{\circ}$ (ummz); La Union, $850 \mathrm{~m}, 31$ Aug. 1972 (E. C. Welling M.), 1 it (Ummz); El Salvador: Anuachopan Dept.: (all collected by V. Hellebuyck): Bosque El Imposible, San Francisco Mendez, 16 Aug. 1987, 3 ơ ( $\mathrm{vH}, \mathrm{RWG}$ ); May 1987, 1 ot, 1 ㅇ (vH); 12
 Aug. 1987, $1 \delta^{\circ}(\mathrm{vH})$; 3 Sept. 1987, 3 ठे, 1 ㅇ ( 1 pair in copula) (vH); El Coyolar, El Imposible, 2 Aug. 1987, $2 \delta^{\circ}$ (vн); Valle de la Puerta, Bosque El Imposible, 3 Sept. 1987, $30^{\circ}$ (vf).

## Description

Holotype male. - Labium grey-white with wash of brown around margin of median lobe; entire face including vertex and occiput yellow green, a transverse trough in front of median ocellus, lateral ends slightly deeper than medial area; a small tubercle behind each ocellus; pedicel, scape, flagellum brown; occiput yellow green, mostly planar, slightly tumid medially, crest green, slightly prominent, straight, slightly notched medially and covered with long brown hairs; postocciput green, transverse when viewed dorsoposteriorly: rear of head yellow brown, paler laterally; lateral margins of labrum and base of mandible ocher; rear of head yellow brown.

Prothorax yellow green except for brown on anterior margin of median lobe and posterior area of anterior lobe. Synthorax (fig. 15) entirely yellow green except for following brown areas: slight vague stripe lateral to middorsal carina; antehumeral stripe; spot on upper end of humeral suture connected anteriorly to antehumeral, and gradually disappearing basally toward mesinfraepisternum; line bordering antealar crest; metasternum pale yellow green. Coxae, femora pale yellow green, becoming dark brown dorsodistally; tibiae, tarsi and armature black.

Wings hyaline, anterior margin of costa yellow, remainder of venation black, pterostigma brown.

Venational details. Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $1 / 3$; anx: fore wing $15 / 15$, hind wing 11/11; pnx: fore wing $11 / 13$, hind wing 13/13; cs under pterostigma: fore wing 6/6; hind wing 7/6; anal triangular cells: 4 . Hind wing 27 mm .

Abdomen (fig. 37) with segment 1 yellow green with a wash of brown dorsally; segment 2 yellow green with diffuse brown spot posterodorsally to auricle; segment 3 broadly light green dorsally, interrupted laterally by a longitudinal brown stripe flaring at transverse carina, inferior margin of tergite ivory; segments 4-6 similar to segment 3, but lateral brown stripes progressively more extensive and each meeting its neighbor dorsally at posterior end of segment; segment 7 with anterior 0.66 pale green white except for black transverse carina, posterior 0.30 of segment red brown; segments $8-10$ red brown, darker dorsally, carinae and lateral foliations black; posterior medial margin of 9 pointed. Abdomen 35 mm .

Cercus (fig. 102) pale ocher, gently decumbent at posterior 0.30 , gradually narrowing toward end, tip with a shiny black tooth; epiproct brown, gendy curved as is characteristic for the genus, tip of epiproct when viewed posteriorly (fig. 128) bidentate.

Accessory genitalia. Anterior hamule dark brown, branched at distal 0.30 , superior branch larger than inferior branch; posterior hamule (fig. 58) pale, roughly triangular with an anterior shoulder, tip with
a black rooth twisted laterally from broad axis of hamule: penis with cornuae pointed but each with median shoulder as shown in fig. 80; lateral lobe prominent, knife- like, serrated posteriorly.
Allotype female. - Similar to male with following differences: synthorax with definite dark triangular middorsal stripe, its base not touching collar; broad, diffuse brown dorsolateral stripe on abdominal segment 2 , brown lateral stripes on segments $3-6$ with their anterior ends ending at basal 0.20 of each segment so that lateral white connects with dorsal yel-low-white middorsal stripe; cercus pale.

Vulvar lamina (figs. 183) small, cleft about as large as each plate; each plate connected basally, each with a raised shelf occupying anterior 0.5 ; postlamellar ridge with Y-shaped juncture posterior to hind margin of lamina; central stem with a well defined circular or oval depression on each side. Abdomen 32 mm .
Venational details. Fifth antenodal thickened in all wings; no marginal cells behind fore wing paranal cells; anx: fore wing $12 / 13$, hind wing $9 / 9$; pnx: fore wing $8 / 9$, hind wing $10 / 10$; cs under pterostigma: fore wing $4 / 5$, hind wing $5 / 5$. Hind wing 27 mm .

## Diagnosis

Selys' (1868) description of Erpetogomphus viperinus was too brief and inexact for Calvert (1899) to detect a specific difference between his specimens of $E$. viperinus ( $=$ E. bothrops) and the true E. viperinus. Examination of the lectotype male and female of $E$. viperinus shows that the Erpetogomphus viperinus of Calvert and all subsequent authors is referable to the new species $E$. bothrops.
E. bothrops is most similar to E. liopeltis and is diagnosed under that species. Erpetogomphus bothrops also resembles $E$. viperinus, but in life, the pale colouration of $E$. bothrops is yellow green (deep green in $E$. viperinus). Erpetogomphus viperinus is generally larger (hind wing male 29-31 mm) and darker. In males, the cercus of $E$. bothrops is pale with a distinct black rooth; while in E. viperinus this structure is dark red brown with no apical tooth. In E. bothrops, the distal 0.5 of the cercus gradually narrows, so the distal 0.5 of the cercus in lateral view is not as thick or robust as the base (fig. 102). In E. viperinus, the distal 0.5 of the appendage is robust and is as thick as its base (fig. 103). In dorsoposterior view, the posterior 0.30 of the cercus of $E$. bothrops gradually narrows (fig. 118), and is not as laterally compressed as in E. viperinus (fig. 119). The inferior distal margin of the cercus of E. viperinus is strongly carinate, not so in $E$. bothrops. The tip of each epiproct of $E$. bothrops in posterior view is bidentate (fig. 128) or truncate (fig. 127), not bluntly pointed as in E. viperinus (fig. 129). Other characters are given in the key.

Females of these species are easily distinguished by
overall body colouration in life, and thoracic maculation. Two structural characters separate E. bothrops from $E$. viperinus the anterior margin of the vertex of E. viperinus contains two well defined pits, each dorsolateral to the median ocellus. Each pit is the receptacle for the bluntly pointed tip of the male epiproct when in copulation. In $E$. bothrops, these pits are replaced by a continuous groove deepest medially, which accommodates the explanate tips of the male epiprocts. Secondly, the V-shaped notch of the vulvar lamina in E. bothrops is more acute (fig. 183) than the more widely divergent notch of E. viperinus (fig. 184). The shape of the vulvar lamina of $E$. bothrops (fig. 183) distinguishes it from similarly marked females of E. sipedon (figs. 187, 188) and E. elaps (fig. 181).

## Remarks

Variation. - Venational variation among the paratype series of males (fig. 228): number of marginal cells behind fore wing paranal cells: $0-3$; anx: fore wing 14-16/13-16, hind wing 9-12/10-12; pnx: fore wing $9-11 / 8-13$, hind wing $10-13 / 10-13$; cs under pterostigma: fore wing 5-7/5-7, hind wing 5-7/5-7; number anal triangular cells: 3-4. Hind wing 21-28 mm.

Variation exists in the direction of the apical tooth of the posterior hamule. It is present in all males and, in most, is twisted so that its tip is directed laterally (i.e., distally) to the direction of the planar surface of the hamule. In a few males, the tooth, though rotated, points in a more lateroanterior direction.

The tip of the epiproct, when viewed posteriorly, is planar and bidentate (e.g., fig. 128); but in some, the distal and mesal teeth are reduced. In rare cases, the tip is almost straight (fig. 127); variations link these extremes. I have found no correlation of the posterior hamular condition with the epiproct condition; nor do these conditions seem to vary according to altitude or locality. For example, of 5 males from Nayarit, Jamatan, only one has hamular teeth pointing predominantly cephalad. This same specimen has a bidentate condition of tips of the epiprocts. Another male has a bidentate condition, another has an intermediate condition, and two others have truncate tips. Of another series of 9 males (including holotype) from the Veracruz, Rio Otapa, two have the hamular tooth pointing predominantly cephalad. These two specimens have bidentate epiprocts, although the right epiproct of one male is intermediate. The remaining 7 males have bidentate epiprocts, but one has an intermediate condition of its right epiproct.

The membranous hood of the penis is variable among and within populations. Most specimens, including the holotype, possess a short hood, exposing a subcutaneous membrane (fig. 80); but a male from

San Luis Potosi, Cascadas Micos, has a long, acute, non-overlapping hood. Another two males, one from Guatemala (Dept. Chiquimula, small stream, vic. Veguitas), and all specimens I examined from El Salvador (Dept. Anuachopan, Bosque El Imposible, San Francisco Mendez), have long, overlapping hood membranes similar to that illustrated for the holotype male of $E$. leptophis (fig. 76). With the exception of the San Luis Potosí male, specimens with long, acuminate hoods seem to typify most southerly specimens. I can find no other characters differentiating these specimens from others.

Venational variation among the paratype series of females ( $n=18$ ): number of marginal cells behind fore wing paranal cells: 1-3; anx: fore wing 12-16/12-16, hind wing 9-11/8-11; pnx fore wing 8-12/9-12, hind wing 9-13/9-13; cs under pterostigma: fore wing 4 $7 / 5-7$, hind wing 5-8/5-7. Hind wing $27-31 \mathrm{~mm}$.

Biology. - I have often seen this species along margins of cut agricultural fields bordering streams and canals. Its apple green colouration renders it difficult to detect among vegetation, but when disturbed, it does not fly far. The type locality is a wide, shallow stream bordered by trees in extensively cultivated farmland. I collected E. bothrops next to cut fields along with six other gomphids: Erpetogomphus eutainia, E. ophibolus, Phyllocycla breviphylla Belle, P. volsella (Calvert), Phyllogomphoides duodentatus Donnelly, and Progomphus clendoni Calvert. Twentyfive other species of Odonata were collected there.
R. Novelo (pers. comm.) observed a pair in copula in Morelos state (Cerro del Higuerón) at 1250 m at 18.20 hr on 14 Sept. 1988.

Williamson records the following on two Acaponeta males: 'Eas[il]y caught while sitting in brushy weed patch on bank of river, sun behind cloud,' and 'Quite common flying over water just above the real rapids and lighting on willow-like stems and twigs along the bank. Saw no $i$ gomphine today.' López (in litt.) writes that they perch along sides of streams or on exposed rocks in the middle of streams. He states that they do not remain long at a spot, moving constantly along the length of the stream.

Elevation gradients range from near sea level ( 50 m, Veracruz, Agua Caliente) to 1250 m (Morelos, Cerro del Higuerón). Collection dates range from May (El Salvador) to November (Mexico, Jalisco).

Distribution (fig. 201). - This species and E. elaps are the most widely distributed species in Mexico and Guatemala. Erpetogomphus bothrops ranges from Tamaulipas in eastern Mexico $\left(24^{\circ} \mathrm{N}\right)$ south to El Salvador $\left(14^{\circ} \mathrm{N}\right)$. It also occurs in mesic areas in western Mexico from Nayarit $\left(22^{\circ} \mathrm{N}\right)$ south. Its distribution indicates an avoidance of the xeric areas in northwestern Mexico and the high mountain plateau in northern Mexico.

## Erpetogomphus viperinus Selys

(figs. 16-thx, 38 -abd, 59 -hamules, 81-penis, 103app, 119-cercus, 129 -pet, 137 -base of postfrons, 155 -vertex, 166 -postocciput, 184 -vl, 201-distr, 229wings)
Erpetogomphus viperinus Selys, 1868: 68 (3 sep.) (descr. of o', 우, 'D'Orizaba'). - Selys 1869: 176 (13 sep.) (redescription of $\delta$ and $q$ from Orizaba); Selys 1873b: 519 (75 sep.) (list); Muttkowski 1910: 87 (in part, cat.); Williamson \& Williamson 1930: 14 (summary of status); Paulson 1982: 256 (Mex.); Davies and Tobin 1985: 28 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 220 (cat.); Tsuda 1991: 95 (cat.).
Herpetogomphus viperinus. - Hagen 1875a: 42 (cat.); Selys 1879: 64 (2 sep.) (status of classification); Kirby 1890: 60 (car.).

## Description

Male. - Entire head pale green (blue green in life), slightly darker on anterior of ante- and postclypeus and labrum, lateral margins of labrum ocher, base of mandible and labium dull grey green; basal 0.25 of postfrons and all of vertex brown, a pair of pits at anterior margin of vertex at juncture of base of antefrons (fig. 137), one each anterolateral to median ocellus; antennae dark brown; occiput with well developed medial tumid area, green, occipital crest slightly emarginate medially, dark brown to black, rimmed with long dark brown hairs; postocciput brown, green medially, transverse, with a slight vertical depression medially; rear of head yellow brown.

Prothorax brown, anterior and posterior lobes green. Synthorax (fig. 16) entirely green (blue green in life) with following dark brown: obscure middorsal stripe ending before collar, becoming darkest at base of antealar crest and extending as narrow wash of brown below rim of antealar crest and joining wide stripe along lateral 0.30 of mesepisternum; this stripe coalescing with normal humeral stripe; often with a narrow isolated stripe of green separating these two stripes; combination antehumeral and humeral stripe reaching collar; all of mesinfraepisternum and ventral part of thorax; rim below subalar carina, often with small extension on obsolete second lateral suture, which in some specimens may form an indistinct second lateral stripe; narrow but well defined third lateral stripe. Metasternum pale grey green. Coxae, trochanters grey, becoming brown exteriorly; base of femora brown, becoming black distally; tibiae and armature black.

Wings (fig. 229) hyaline, anterior margin of costa dark brown at base, becoming yellow distally to proximal level of pterostigma, remainder of venation and pterostigma black.

Venational statistics. Fifth (occasionally sixth) antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: 1-4/1-4; anx: fore
wing 13-18/13-18, hind wing 9-12/10-13; pnx: fore wing $8-13 / 10-12$, hind wing $10-14 / 10-13$; cs under pterostigma: fore wing 5-7/4-7, hind wing 5-7/5-7; anal triangular cells: 4 (rarely 5). Hind wing 29-31 mm .
Abdomen (fig. 38) with segment 1 green with brown dorsolaterally except for green on posterior dorsal 0.50 of segment; segment 2 with middorsal green stripe; brown dorsolaterally surrounding green auricle; ventrolateral area pale green, annulus black; segments 3-6 primarily black with following pale areas: basal 0.10 of each segment, dorsally forming an incomplete blue green middorsal stripe extending to posterior 0.20 of each segment; ventrolateral wedge shaped white spot at middle of each segment, connecting in some specimens to basal pale ring; segment 7 with dorsal 0.50 light green, posterior 0.50 red brown, black along transverse carina; segments 8-10 predominantly red brown, darker dorsally, denticulated posterior margin of these segments black; posterior margin of segment 9 often forming a posteriorly directed point. Abdomen $36-40 \mathrm{~mm}$.

Cercus (fig. 103) pale ocher, slightly darker ventrally, robust and strongly curved at posterior 0.30 , inner margin of posterior 0.30 of cercus planar (fig. 119); same structure in posterodorsal view narrow, inferior carina along posterior 0.30 of cercus; epiproct brown, becoming black posteriorly, gently curved as is characteristic for the genus; tip of epiproct, when viewed posteriorly (fig. 129) forming a narrow, blunt point.

Accessory genitalia. Anterior hamule (fig. 59) black, divided at distal 0.30 ; smaller posterior branch almost as large as anterior branch, its pointed tip almost meeting tip of anterior (larger) branch; posterior hamule (fig. 59) pale, roughly triangular, swollen at distal 0.25 to 0.50 of appendage, tip with a black tooth rwisted laterad from broad axis of hamule; penis with cornuae pointed but each with well developed median shoulder (or median lobe in some specimens), as shown in fig. 59; lateral lobe prominent, knife-like, arcuate laterally, its outer margin strongly serrated.

Eemale. - Similar to male, with following differences: Vertex (fig. 155) without postocellar ridge, a well defined pit anterolaterad to median ocellus as in male; occiput green, narrow, mostly planar, or with only a slight tumid area medially, hind margin slightly sinuate, barely emarginate at middle; postocciput green, transverse; synthorax with no vestige of second lateral stripe; abdomen with dorsolateral dark stripe on segment 1 reduced or vestigial, middorsal dark stripe on segments 2-6 narrower, anteriorly touching black annulus, widened at transverse carina, conitricted near center and widening at distal 0.25 of each segment, ventral white narrow but expanded near center of segment, sometimes a longitudinal
wedge-shaped spot isolated by narrow posterior and anterior lateral offshoots of black dorsolateral stripe, white lateral wedge- shaped spots largest on segments 2 and 3, dividing or partially dividing dorsolateral brown stripe; segment 7 similar to segment 6 , but dorsal pale orange green connecting anteriorly with lateral white at basal 0.10 of segment; segments 8-10 dark brown, becoming black dorsally and posterolaterally to all black with ill-defined white lateral spot on each segment; cercus, paraproct brown. Abdomen $37-41 \mathrm{~mm}$.
Vulvar lamina (fig. 184) small, cleft as large as each plate; each plate broadly connected basally, each with a well developed raised shelf occupying anterior 0.50 ; medial margin of each plate slightly concave, its tip falcate; postlamellar ridge with Y -suture posterior to hind margin of lamina; central stem short with a welldefined circular or oval depression on each side.
Venational statistics. Anx: fore wing 15-20/15-21, hind wing 10-14/11-14; pnx: fore wing $10-13 / 10-15$, hind wing $10-15 / 9-14$; cs under pterostigma: fore wing 5-8/5-8, hind wing 5-8/6-7. Hind wing 31-34 mm.

## Diagnosis

Erpetogomphus viperinus is most closely related to E. bothrops, and is diagnosed under that species. It is also superficially similar to E. liopeltis and is diagnosed under that species. The thoracic pattern of $E$. viperinus is superficially similar to that of $E$. sipedon, but is easily separated by characters listed under that species and in the key.

## Remarks

Variation. - The description is based on 49 males and 38 females from Veracruz State, Mexico. The lectotype and paralectotype are in reasonably good condition, but post mortem preservation is poor and would yield an inadequate description. Venation characters for this pair are as follows:

Lectotype male: number of marginal cells behind fore wing paranal cells: $3 / 1$; anx: fore wing $16 / 17$, hind wing 12/12; pnx: fore wing $13 / 12$, hind wing 11/13; cs under pterostigma: fore wing $6 / 6$, hind wing $6 / 5$; anal triangular cells: 4/4.

Paralectotype female: number of marginal cells behind fore wing paranal cells: $4 / 3$; anx: fore wing $15 / 18$, hind wing $12 / 11$; pnx: fore wing $10 / 12$, hind wing $12 / 13$; cs under pterostigma: fore wing $7 / 8$; hind wing 7/6.

Examination of 20 males shows variability is some venational characters. All specimens have marginal cells behind the fore wing paranal cells, ranging from 1-4 cells. The fifth antenodal is most often thickened, but rarely the fourth (one wing) or sixth ( 7 wings). The sixth antenodal is thickened on both wings of the lectotype.

Little variation exists in body colouration. The second lateral thoracic stripe is incomplete or vestigial in most males. It is complete on only one male from Rio Metlac. Females show a greater tendency toward a dense venation than do males. Marginal cells behind the fore wing paranal cells were present in all 20 females I examined, ranging from 1 (one wing) to 8 (two wings). The fifth antenodal is generally thickened, but it is the fourth on the left wing of the paralectotype, sixth in 17 wings (including right wing of paralectotype) and seventh in four wings.

Biology. - Although of restricted distribution, $E$. viperinus can be common during certain years. I collected a few specimens of both sexes in the mostly shaded understory at the Rio Metlac. The locality consists of a fast running river in a steep canyon. The narrow valley contained many coffee plants. I flushed various specimens, and they alit on low vegetation, where their deep blue green colour made them difficult to detect unless the eye followed closely where each specimen landed. Once approached they were easy to take with a net. Other Odonata taken at the same site were Hetaerina cruentata Rambur, Argia sp. nr. fissa Selys, Argia extranea Hagen in Selys, Aeshna psilus Calvert, Brechmorhoga pertinax (Hagen), B. vivax Calvert, Cannaphila vibex Hagen, Erythrodiplax fusca (Rambur), and E. umbrata (Linnaeus).

The species has been collected more frequently farther north at Teocelo under similar circumstances. The Odonata assemblage there is more varied and consists of Palaemnema sp. n., Paraphlebia zoe Selys, Argia extranea, Argia sp. n., Brechmorhoga pertinax, B. rapax Calvert, B. tepeaca Calvert, Cannaphila vibex, Libellula herculea Karsch.

This species has been taken at elevations of 8001300 m (Teocelo and vicinity). Collection dates range from 6 June through 18 September (Teocelo and vicinity).

I have abstracted the following biological notes from López (in litt.): Tenerals and adults travel far from the stream, hiding in vegetation in open areas, where they feed by making short sallies from perches. They capture primarily small flies and butterflies (Lycaenidae) passing by. Copulation takes place in open fields away from streams, as males grab females which fly by. The copulating pair then retires to the dense forest. Females oviposit while flying above water, dropping their eggs from about 30 cm . At dusk, males return to feeding sites, often in considerable numbers, where threat display patterns among males and females have been observed when two or more individuals try to occupy the same perch. Specimens are often preyed upon by spiders (Araneidae).

Distribution (fig. 201). - Erpetogomphus viperinus is apparently restricted to central Veracruz. As stated under the species account for $E$. bothrops, all pub-
lished records of this species since Calvert's redescription of $E$. viperinus refer to the widespread $E$. bothrops. I examined one male from the Paris Museum with a handwritten label, 'N. Carolina', but I have seen no specimens of $E$. viperinus from the United States. I believe that locality is in error.

## Material

Type data. - Lectotype male by present designation with following data: small green handwritten label 'Mex/B[ouchard]'; white handwritten label 'Herp. / viperinus /S.[elys]/ ठ"; two manila coloured labels each with ' 26 ' handwritten in pencil in an unknown hand; rectangular red label with printed 'iectotype'/and handwritten: 'Erpetogompbus/ viperinus Selys $1869 /$ ơ/des. [printed] R. W. Garrison 1984'. The caudal appendages are shown in fig. 103. Paralectotype female: small green handwritten label 'Mex./B[ouchard]'; white handwritten label 'Herp./ viperinus/ S[elys]/ P', two manila coloured labels, each with ' 28 ' handwritten in pencil in an unknown hand; rectangular white label with printed 'iectotype'/ and handwritten 'Erpetogomphus/viperinus Selys 1869/ ㅇ/ des. [printed] R. W. Garrison 1984'. Both specimens in IRSN.
 and paralectotype of). - Mexico: Veracruz: nr. Municipio Teocelo at Puente Teocelo, ca. 1150 m , (unam, Rwg); Barranca de Cayoapa, Teocelo, 750890 m, (UNAM, JB, CE, PSM, DALD); Barranca de Cayoapa, Tejeria-Teocelo, 800 m , (UNAM); road to Monte Blanco, Teocelo, (unam); El Trapiche, Teocelo, 1100 m (Unam); road to Santa Rosa, Teocelo, (unam); Santa Rosa, Teocelo, (unam); Cascada de Xico, Teocelo, 1100 m , (unam); Cascada Texolo, (unam); Parque Javier Clavijero, Jalapa, 1300 m (UNAM); Fortín de las Flores, 1010 m (RWG); Rio Metlac, ca. 3.5 km WNW of Fortín de las Flores, 900 m (RWG); km 327 W of Cordoba, (FSCA); Cordoba, (UsNM).

## Erpetogomphus designatus Hagen in Selys

(figs. 17, 18 -thx, 60 -hamules, 82 -penis, 104 -app, 130, 131-ept, 156, 157-vertex, 185, 186-vl, 202distr)

Erpetogomphus designatus Hagen in Selys, 1858: 661 (401 sep.) (descr. o', ㅇ, 'Pecos River, Texas'). - Hagen in Selys 1859: 536 (10 sep.) (descr. ơ, 아); Selys 1873b: 519 (75 sep.) (list); Calvert 1899: 386 (mentioned); Calvert 1905: 166 (Ohio, Ind., Mexico); Muttkowski 1910: 86 (cat.); Calvert 1912a: 289 (Mex., distr.); Hine 1913: 96 (Ohio); Williamson 1914b: 447 (Tex.); Kennedy 1917a: 544 (larva, notes); Williamson 1917: 8 (Ind.): Kennedy 1917b: 137 (Kans.); Kennedy 1918: 298 (notes); Williamson 1923: 8 (Kentucky); Montgomery 1925: 386 (Ind., habits); Montgomery 1927: 289 (Ind.);

Kennedy 1928:373 (seasonal distribution); Montgomery 1929: 340 (Ind.); Needham \& Heywood 1929: 80 (key, descr.); Byers 1930: 53 (Fla. Key, descr., habits); Williamson \& Williamson 1930: 12 (summary of status); Bvers 1931: 51 (Tenn.); Bird, 1932: 51 (Okla.); Williamson 1932: 23 (Mo., habits); Tinkham 1934: 216 (Tex.); Montgomery 1935: 234 (Ind.); Borror 1935: 453 (Ohio): Borror 1937: 186 (Ohio); Montgomery 1937: 207 (Ind., habits); Ahrens 1938: 11 (Mo., predation); Wright 1938: 27 (Tenn.); Ferguson 1940: 5 (Tex.); Montgomery 1940: 289 (S. C.); La Rivers 1940b: 63 (Nev.); Montgomery 1941: 230, 238 (Ind.); Ferguson 1942: 146 (Tex.); Montgomery 1947: 165 (distr.); Bick 1951: 179 (Okla.); Montgomery 1951: 207 (Ind.); Needham and Westfall 1955: 146 (key, descr.); Cross 1955: 11 (S. C.); Montgomery 1955: 133 (Ind.); Cross 1956:4 (Fla.); Bick and Bick 1957: 2 (Okla.); Kormondy 1957: 108 (Tenn.); Bick 1957: 80 (La.); Gloyd 1958: 8 (Tex.); Bick and Bick 1958: 240 (Okla.); Bick 1959: 131 (Ark.); Donnelly 1961:7 (Md. Va., D. C.); Alrutz 1961: 23 (Ohio); Borror 1963: 104 (common name); Macklin and Cook 1967: 120 (Ky.); Montgomery 1967: 127 (distr.); Roback \& Westfall 1967: 114 (water quality data); Montgomery 1968: 134 (distr.); Pinhey 1969: 189 (tandem linkage); Resener 1970: 37 (Ky.); Huggins et al. 1976: 16 (Kans.); Harp \& Rickett 1977: 50 (Ark.); Huggins 1978: 2 (Kans.); Young \& Bayer 1979: 90 (larva, key, Tex.); Carle 1979: 322 (status in Virginia); White et al. 1980: 26 (S. C.); Carle 1982: 339 (W.Va.); Dunkle \& Westfall 1982: 32 (status in Fla.); Paulson 1982: 255 (U.S., Mex.); Huggins \& Brigham 1982: 4: 39 (N. C., S. C.); Harp \& Rickett 1985: 132 (Ark.); Davies \& Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat., attributes authorship to Selys); Bridges 1991: VII. 60 (cat.); Tsuda 1991: 95 (cat.); Dunkle 1992: 39 (Fla).
Gomphus designatus. - Hagen 1861: 99 (descr. ठै, 우).
Gomphus sp.. - Cabor, 1872: 4 (descr. larva as Gomphus sp. No. 6, Poles Creek, Tex.).
Herpetogomphus designatus. - Walsh 1862: 389 (compared with Ophiogomphus rupinsulensis); Hagen 1875a: 42 (cat.); Selys 1879: 64 (2 sep.) (characters of genus); Hagen 1885: 255 (larva of Cabot identified); Kirby 1890: 60 (cat., attributes authorship to Selys); Banks 1892: 179 (Kans.); Calvert 1899:386 (characters of genus); Adams 1900: 622 (Ark.); Williamson 1903: 226 (Tenn.); Van der Weele 1906: 177 (mentioned); Tucker 1907: 79 (Kans.); Tucker 1908: 99 (Tex.); La Rivers 1938: 76 (Nev., habits); La Rivers 1940a: 112 (Nev.).

## Description

Male. - Entire face pale green, more vivid on postclypeus and frons, vertex with wash of brown around ocelli or encirely dark brown, scape dark brown, pedicel light brown, flagellum dark brown; anterior margin of vertex forming a trough at junction of antefrons, this trough with a pair of deeper pits anterolateral to median ocellus; occiput green, wide, tumid medially, crest slightly sinuate, barely emarginate medially or straight, its hind margin covered with long brown hairs; postocciput not visible dorsally, green; rear of head brown, darker toward occipital foramen.

Prothorax predominantly brown, becoming green
dorsally; anterior and posterior lobes green; synthorax (fig. 17) green with following dark brown: well defined middorsal stripe widening to collar, its upper end extending along antealar sinus connecting with narrow humeral stripe; an isolated antehumeral stripe not touching dorsal or ventral margin (except in specimens from Durango and Chihuahua - see remarks); lower part of humeral stripe extending posteriorly to form a fragmented second lateral stripe, though in some specimens a vestige of a complete stripe exists; narrow third lateral stripe. Coxae, trochanters pale green, femora green becoming brown posterolaterally at distal 0.20 ; distal 0.05 of metafemora green with narrow brown line extending distally from subapical brown area; tibiae dark brown with yellow along lateral carinae of basal 0.5 of mesotibiae, a vestige of yellow or none on metatibiae; tarsi dark brown, armature black.

Wings hyaline with wash of yellow basally, venation dark brown, basal wing venation light brown, especially anteriorly, anterior margin of costa yellow, except for anterior margin of pterostigma; pterostigma dark brown, veins bordering it black.

Venational statistics. Fifth (rarely fourth) antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: $2-5 / 1-4$; anx: fore wing 11-14/11-14, hind wing 8-11/8-11; pnx: fore wing $7-10 / 8-11$, hind wing $7-11 / 8-12$; cs under pterostigma: fore wing 5-7/5-7, hind wing 4-7/4-7; anal triangular cells: 3-5/3-4. Hind wing 28-31 mm.

Abdomen. Segment 1 green with dorsolateral brown stripe occupying basal 0.75 of segment; segment 2 similar to segment 1, but dark brown expanding ventrolaterally behind auricle, contracting again and connecting with dark brown annulus; segment 3 similar to segment 2 but dark brown dorsolateral stripe interrupted at anterior 0.25 and again at distal 0.50 to 0.75 of segment, dark brown prominent at transverse carina, posterior 0.25 to 0.30 of segment with dorsolateral brown expanding laterally to encircle segment at posterior 0.10 of segment; segments 46 similar to segment 3 but dorsolateral brown interrupted only at anterior 0.25 to 0.30 of segment and expanding abruptly at transverse carina; segment 7 with narrow dark brown on transverse carina and poorly defined brown on lateral 0.50 of segment; segments $8-10$ yellow brown with dorsolateral red brown occupying basal 0.40 to 0.75 of segments; pale areas of segments $2-6$ orange green dorsally, becoming white below. Abdomen $35-37 \mathrm{~mm}$.

Cercus (fig. 104) yellow brown, strongly angulate near middle; tip of cercus beyond angulation narrow, drawn out into a fine point; ventral margin with inferior carina at basal 0.45 . Epiprocts yellow brown, gently curved dorsally at $90^{\circ}$, tips obliquely truncate (fig. 130) or spatulate (fig. 131).

Accessory genitalia (fig. 60). Anterior hamule black, divided at upper 0.50 ; lower branch forming a small, posteriorly directed truncate appendage; posterior hamule pale green, triangular, with anterior basal shoulder, tip evenly round, black; penis with long, decumbent lanceolate lateral lobe, its posterior end serrated; cornuae separated, tips narrowly rounded, with no medial lobe; membranous hood moderately long but usually with ends not overlapping.

Female. - Head as in male, but median swelling of occiput more prominent; prothorax and synthorax as in male; wings with yellow at base more extensive, often with markings at base of arculus, to cubitoanal crossvein; abdomen with definite pale orange dorsally, white to orange white laterally, these areas separated by brown dorsolateral stripes as in male; these stripes on each segment may be narrower, especially so between transverse carina and posterior 0.20 to 0.30 of segment; segments $8-9$ mostly dark brown dorsally, sides tawny, annulus pale; segment 10 yellow brown, cercus ocher. Abdomen $33-36 \mathrm{~mm}$.

Vulvar lamina (figs. 185-186) with prominent basal plates; quadrangular laminar plates acuminate posteriorly, meeting at medioanterior margin; cleft between plates broadly U-shaped; juncture of Yshaped postlamellar ridge at or before level of posterior margin of lamina. Abdomen $33-36 \mathrm{~mm}$.

Venational statistics: number of marginal cells behind fore wing paranal cells: $2-5 / 1-5$; anx: fore wing 12-15/11-15, hind wing 8-12/8-11; pnx: fore wing $7-$ 12/7-12, hind wing $8-12 / 7-12$; cs under pterostigma: fore wing 5-7/5-7, hind wing 4-8/5-7. Hind wing 2832 mm .

## Diagnosis

Little difficulty should be encountered in identifying either sex of this species. The strongly angulate cercus superficially allies it with $E$. sipedon, $E$. heterodon, and E. lampropeltis, but the structure of the occiput and the penile characters easily separate males of these species. The broadly defined occiput, with the large medial tumid area characteristic of $E$. designatus, is not present in the other three species. Male E. lampropeltis have a broad occiput, as do male $E$. designatus, but the medial area of the occiput is only slightly tumid.

The long, recumbent lateral lobe of the penis is unique to this species (fig. 82). In E. lampropeltis and E. sipedon, this structure forms a spinulose semicircle (figs. 83-85).

As in males, females of $E$. designatus are distinguished from all other similar species by the tumid median area of the occiput (fig. 156). The vulvar lamina in E. designatus (figs. 185-186) more closely approaches that of $E$. bothrops (fig. 183) than of $E$. sipedon (figs. 187-188), E. beterodon (fig. 193), or E.
lampropeltis (figs. 189-190). Each lameilar plate of $E$. designatus is smaller and the notch in the medial margin is more concave than in the other species.

## Remarks

Variation. - Calvert (1907) described a specimen from San Pedro, Coahuila, Mexico, which differed from typical $E$. designatus in the following characters (typical condition in parentheses): 1) dark antehumeral stripe reaching down to mesinfraepisternum (lower part of stripe isolated), and 2) darker, more pronounced markings on abdominal segments as follows: segment 7 with a black spot on each side of posterior 0.50 of segment (like area red brown, blending anteriorly with ocher pale area); segment 8 with a black stripe on each side for almost its entire length (this area red brown); segment 9-10 with black at basal dorsolateral 0.50 to 0.75 (this area red brown with some black on dorsal 0.50 of segment). Calvert (1907) considered the San Pedro male to represent $E$. designatus. This specimen is indicated as in the ANSP, but it could not be found (Azuma in litt., 1984). However, I have seen three males (Durango and Chihuahua) and a female (Durango) whose markings agree with Calvert's San Pedro male. I also find the following other differences: 1) no wash of yellow present on wing bases, 2) dark thoracic stripes thicker and more pronounced (fig. 18), 3) a black spot anteroventrally to auricle of abdominal segment 2 , and 4) tip of epiproct, viewed posteriorly, is spatulate (fig. 131), not obliquely truncate (fig. 130) as is typical of other specimens. The single female has a more arcuate occiput (fig. 157) than typical E. designatus (fig. 156), but the dark anteroventral spot on segment 2 is small and inconspicuous.

Specimens from Durango, Chihuahua, and Coahuila could be a well-defined subspecies characterized by overall darker colouration and slight morphological modification of the female occiput (if typical for this form) and male epiproct. I detect no morphological differences in the penes, hamules, or cerci and, like Calvert, I am inclined to treat these specimens as $E$. designatus. Further evidence to support their conspecificity comes from a male from Nuevo Leon (in foothills of the northernmost range of mountains) in which the antehumeral stripe extends to the mesinfraepisternum but is in all other characters typical of $E$. designatus.

One male from McLennan Co., Texas, has the tips of the posterior hamules with a small, anteriorly recurved hook. This male and another from the same locality have vestiges of an anteroventral dark spot on abdominal segment 2 .

I have not seen sufficient material of this species throughout eastern parts of its range to render any conclusions about geographic variability. Numerous
specimens, primarily from Gonzales and Williamson counties. Texas, show liede variability.

Although the type locality for E. designatus and many other species collected by Capt. John Pope of the Smithsonian Institution and described by Hagen, is said to be the Pecos River in western Texas $\left(32^{\circ} \mathrm{N}\right.$, $\left.104^{\circ} \mathrm{W}\right)$, Needham and Cockerell (1903), citing Scudder and Cockerell (1902), argue that the type locality was probably at or near Roswell, Chaves Co., New Mexico, as follows (Needham and Cockerell 1903: 138-139): 'In Proc. Davenport Acad. Sci. vol. 9 (1902), p. 51, it is inferred that Pope collected his material at very different localities above the river, since he obtained both Melanoplus bivittatus and $M$. differentialis, which inhabit different life-zones. However. in 1902 it was observed that these two grasshoppers do actually occur together at Roswell. As it is not very likely that their ranges overlap very much, it thus seems likely that Pope's collections, if all from one place, were from the vicinity of the present town of Roswell, rather than from Lat. $32^{\circ}$ (the present boundary between New Mexico and Texas), as Hagen indicates. In all probability, however, the collections were made in several places.

No year is indicated on the data of any of the type material, nor have I found any reference to the year of Capt. Pope's expedition. However, a public display on the recent human history of the caverns at the visitor center at Carlsbad Caverns National Park, Eddy Co., New Mexico, states that Pope's expedition occurred in 1854-1855.

Venational details of lectotype male: number of marginal cells behind fore wing paranal cells: $1 / 2$; ; anx: fore wing $12 / 13$, hind wing $10 / 10$; pnx: fore wing $8 / 9$, hind wing $9 / 9$; cs under pterostigma: fore wing $6 / 6$, hind wing 7/6; anal triangular cells: $3 / 4$.

Biology. - The synonymy indicates that more is known about this species than any other congener. Erpetogomphus designatus seems to be a species of mesic deciduous forests where there are clear streams and rivers. Montgomery (1925) writes of its habits in Indiana: "This species was common in a well-cultivated orchard near Vincennes on August 7, 1924, resting on the ground with wings spread out against the soil but taking flight so readily that only three specimens were taken in two hours although I saw 20 or more.' In Hamilton Co., Indiana, Montgomery 1937) records it as '...rather numerous at ripples in the river, flying just above the surface of the water, and alighting on rocks. However, it was exceedingly difficult to caprure...'. Williamson (1932) observed - his species flying back and forth near the surface over ripples in Missouri. Others alighted on vegetation or $\because$. . 1 maitem hads. La Rivers (1938) described he flight of! in ont: at Hiko Springs in the l'ahranagar 'Valley, Nevada, as 'low, swift, and change
of position rapid...' Ahrens (1938) describes the capture of a male along with a male Hagenius brevistylus Selys near Lebanon, Missouri: 'The H. brevistylus had struck and seized the designatus with such force that both were carried into the water. Both were dipped from the river an instant after they struck the surface.'

Dunkle and Westfall (1982) assign E. designatus as 'threatened' in Florida, because it is thus far known only from Liberty and Gadsden counties in the panhandle. In 1975 and 1977, I found this species abundant along cultivated stubble bordering trees next to the Guadalupe River in Gonzales County, Texas. It was more abundant than E. eutainia; and both species were easily flushed from low or cut vegetation. Members of both sexes did not fly far, and I could easily photograph and capture specimens.

Collection dates range from 6 May in Texas (Williamson, 1914b) to 5 October in Tennessee (Wright, 1938).

The larva has been described by Cabot (1872), Needham and Heywood (1929) and Needham and Westfall (1955)

Distribution (fig. 202). - Erpetogomphus designatus and E. compositus are the most boreal of the genus, with $E$. designatus generally replacing the western $E$. compositus in the eastern United States. Erpetogomphus designatus occurs as far north as Maryland and Washington, D. C., west through West Virginia, Ohio, Indiana, Missouri, Kansas, southeastern Colorado, eastern New Mexico, into western Texas. Sidney Dunkle (in litt., 1991) mentions one female from South Dakota (Fall River Co., 9.7 mi SW Hot Springs, 29 July 1976, G. and J. Bick, L. Hornuff), erroneously published as Stylurus intricatus (Selys) (Bick et al., 1977: 151), which is the northernmost record for this species. It occurs generally throughout the southeastern United States, but does not penetrate peninsular Florida (Dunkle and Westfall, 1982). Farthest western records include Hiko Springs, Lincoln Co., Nevada (La Rivers, 1938) and Arizona (Maricopa Co., Salt River at Corn Bluff Campground, 15 mi NE of Mesa). Erpetogomphus designatus is found as far south as Nuevo Leon (24$26^{\circ} \mathrm{N}, 100^{\circ} \mathrm{W}$ ) and west to Coahuila (Calvert, 1907), Durango, and Chihuahua, Mexico.

It is partially sympatric with $E$. compositus in Arizona, Nevada, New Mexico, and western Texas. I collected it with E. eutainia in Gonzales Co., Texas.

## Material

Type data. - Lectotype male by present designation with following data: 'July/ $16^{\prime}$ [handwritten], 'designatus' [handwritten], 'Hagen' [printed], small white label with printed 'Type' with red label glued to lower edge with handwritten '1864'; red rectangular label with printed 'Lectotype'/ and written
'Erpetogomphus designatusl Hagen ơ/ des. [printed] R. W. Garrison 1984'. Another ơ, a paralectotype, has the handwritten labels, 'Erp. designatus/ $\delta /$ Pecos River' [in Hagen's hand] and 'Pecos River/ Uhler 1860.' in an unknown hand. A female paralectotype, collected 'July/ 15' has the additional label: [all printed unless otherwise stated] 'erpetogomphus/ designatus hagen [stamped] / P. P. Calvert, det [written] $1905 /$ B. C. A. Neur., p. [written] 166.' Type locality herein restricted to vicinity of Roswell, Chaves Co., New Mexico (see remarks).
 and ô of paralectotypes). - U. S. A.: Florida: Gadsden Co., Apalachicola River, Aspalaga Landing, Hwy I-10 (jd, rwg); Georgia: Houston Co., Ocmulgee River at 96, (rwg); Virginia: Louisa Co., South Anna River, Rte 657, (csuc); North Anna River, Rte 601, (csuc); Kansas: Chatauqua Co., Big Caney River, $11 / 2 \mathrm{mi}$ W of Elgin, (csuc); Texas: Caldwell Co., Plum Creek at US Hwy 90 (rwg); Goliad Co., 3 mi N of Goliad, (rwg); Gonzales Co., Guadalupe River, 4 mi S of Gonzales (RWG); Palmetto State Park, 6 mi S of Luling, nr. US Hwy 183, (rwg); Hidalgo Co., Bentsen, Rio Grande State Park, (csuc); McLennan Co., Hog Creek, (rwg); Real Co., Nueces River at Tex Hwy 55, S of Barksdale, (RWG); Reeves Co., roadside irrigation ditch W of Balmorhea (RWG); Williamson Co., 3 mi SW of Taylor, (rwg); Mustang Creek by Carlos G. Parker Blvd (= Loop 427), Taylor, (Rwg); Colorado: Las Animas Co., Purgatory River, Bent Cyn nr. Delphi (csuc, rwg); New Mexico: Guadalupe Co., Santa Rosa, 4600 ft. (ummz); Arizona: Maricopa Co., Salt River at Corn Bluff Campground, 15 mi NE Mesa, (fSCA); Mexico: Nuevo Leon: in foothills of the northernmost range of mountains, (Ummz); Apodaca, just NE of Monterrey, near course on Instituto farm, (ummz, rwg); Chihuahua: La Cruz, (rwg); Naica (ummz); Durango: 3.7 mi NW of Gomez Palacio on Hwy 49, 3700 ft ., (Ummz); 8.2 mi N of Gomez Palacio on Hwy 49, (ummz); Rio Nazas at Mex. Hwy $49,3.4 \mathrm{mi} \mathrm{S}$ Ciudad Leon Guzman, 4200 ft . (DRP).

## Erpetogomphus sipedon Calvert

(figs. 19-thx, 39, 40 -abd, 61-hamules, 83 -penis, 105,106 -app, 158, 159 -vertex, 168 -leg, 187, 188vl, 203-distr)
Erpetogomphus sipedon Calvert, 1905: 165 (descr. \& Jalisco: Guadalajara). - Calvert 1907: 399 (号 'Las Bocas in Durango...'); Calvert 1908c: $\operatorname{xxx}$ (Erpetogomphus sipedon, type ㅇ, Guadalajara, found by Schumann); Calvert 1909: 481 (Cuernavaca, Mexico); Williamson and Williamson 1930: 13 (summary of status); Kimmins 1969: 297 (type in bмnн); Paulson 1982: 256 (Mex.); Davies and Tobin 1985: 28 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 195 (cat.); Tsuda 1991: 95 (cat.).

## Description

Male. - Entire face light green; this colour more vivid on frons, post- and anteclypeus; base of mandible, labrum light green, vertex with brown around and on ocellar prominences, remainder light green; antennal pedicel dark brown dorsally, light green ventrally; second segment and flagellum dark brown; occiput light green, crest slightly emarginate medially, its hind margin covered with long brown hairs; rear of head brown, darker toward occipital foramen, becoming light green laterally.

Prothorax predominantly dark brown, becoming light green dorsally; anterior and posterior lobes light green. Synthorax entirely light green (probably a vivid apple green in life) with following dark brown: antehumeral stripe ending dorsally before antealar sinus, narrow stripe along humeral suture, this stripe widening at dorsal 0.30 of suture and with anterior margin just touching upper part of antehumeral stripe, anterior 0.50 of mesinfraepisternum. Coxae, trochanters ivory, femora light green with black along external surfaces, more so on mesofemora; most of lateral surface of mesofemora dark brown, these markings more restricted and limited to apical 0.30 of metafemora; tibiae black with small lateral ivory stripes along basal 0.30 ; tarsi and armature black.

Wings hyaline, venation dark brown, basal wing venation brown, anterior margin of costa yellow; pterostigma brown, black along margins.
Venational statistics ( $\mathrm{n}=13$ ). Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: 1 ; anx: fore wing 12 -13/12-15, hind wing 9-10/9-10; pnx: fore wing $7-$ 10/7-9, hind wing 9-11/8-10; cs under pterostigma: fore wing 5-6/5-6, hind wing 5-6/6; anal triangular cells: 4 . Hind wing $28-31 \mathrm{~mm}$.
Abdomen (fig. 39). Segment 1 light green with dorsolateral brown stripe; segment 2 similar to segment 1 but dark brown expanding posteriorly behind auricle, contracting again and interrupted just before brown annulus; segment 3 similar to segment 2 but with dark brown at transverse carina, anterior end of dorsolateral not on or just touching segment 2, its posterior end broadly connecting with dark brown annulus; segments $4-6$ similar to segment 3 with posterior 0.40 wider than anterior 0.60 ; segment 7 largely pale except for brown along transverse carina, becoming red brown posteriorly, especially so laterally; segment 8 largely red brown with ill-defined light yellow brown middorsal and ventrolateral stripes; segment 9 similar to segment 8 ; segment 10 red brown basally, becoming pale yellow brown distally. Abdomen 33-37 mm.

Cercus (figs. 105-106) ivory, strongly angulate near middle; tip of cercus beyond angulation narrow, drawn out to a fine point; ventral margin with inferi-
or carina at basal 0.30 . Epiprocts yellow brown, typical of genus.

Accessory genitalia (fig. 61). Anterior hamule black, divided at upper 0.30 , lower branch forming a small, truncate appendage; posterior hamule white, triangular, with anterior basal shoulder, tip with small, anteriorly recurved tooth; penis with small serrated lateral lobes, membranous hood short, exposing a subcutaneous membrane which forms posterior margin of segment (fig. 83); cornuae (fig. 83) moderately long, widely separated, medial lobe present, usually 0.5 to 1.0 length of outer lobe.

Holotype female. - Face entirely pale except for small black area at base of mandible, occiput (fig. 159) dark brown around ocelli, remainder pale. Prorhorax mostly pale, darker along pleura. Synthorax (fig. 19) mostly pale except for brown margin around antealar sinus and incomplete antehumeral stripe, its upper end joined with narrow humeral stripe. Coxae, trochanters, and femora pale with apical 0.30 black, especially mesofemora, metathoracic legs with black less extensive (fig. 168), forming black streaks externally; tibiae entirely black except for pale line laterally, tarsi and armature black.
Abdomen (fig. 40) similar to male but with dark brown dorsolateral stripe of uniform width on segment 2 , segments $8-9$ mostly black dorsally, sides tawny, annulus pale, cerci lacking. Abdomen 35 mm .
Venational details: number of marginal cells behind fore wing paranal cells: $2 / 2$; anx: fore wing $13 / 13$, hind wing $9 / 11$; pnx: fore wing $9 / 9$, hind wing $10 / 12$; cs under pterostigma: fore wing $6 / 7$, hind wing $7 / 6$. Hind wing 31 mm .
Vulvar lamina (fig. 187) with broadly rectangular flaps meering at anterior margin; juncture of Yshaped postlamellar ridge at level of posterior margin of lamina.

## Diagnosis

Males of Erpetogomphus sipedon are most similar to E. heterodon but are easily distinguished by the structure of the penis and cercus. The cornua in E. sipedon has a medial lobe (fig. 83), which is lacking in $E$. heterodon (fig. 87), and the tip of the cercus in E. sipedon are slender and finely attenuate (figs. 105-106), not robust and short as in E. heterodon (figs. 110111). Erpetogomphus heterodon and E. sipedon are the only two species of the $E$. designatus group which share a similar thoracic design. However, E. sipedon lacks any dark thoracic stripes on the second and third lateral sutures; a vestigial incomplete second lateral stripe and complete narrow third lateral stripe exist on $E$. heterodon. The thoracic pattern of $E$. boa is also simi'ar to E. sipedon, but E. boa is well differentiated by genital characters given in the keys. Although the difference in somatic patterns between E. lampropeltis
and $E$. sipedon is great, these two species are separable by the structure of the penis. In E. sipedon, the hood is abbreviated, exposing a subcutaneous membrane, which latter forms the posterior margin of the segment. In E. lampropeltis, the hood is of normal length and it forms the posterior margin of the segment.

Although females of $E$. sipedon are easily differentiated from all other similar species by differences in colour and pattern, their separation from the same species using only structural characters is more difficult. Species morphologically similar to $E$. sipedon include E. crotalinus, E. compositus, E. viperinus, $E$. bothrops, and $E$. beterodon.

The occiput of E. sipedon is straight or, at most, weakly notched, (notch in E. crotalinus not clearly seen in the paralectotype female, [fig. 161]). The longitudinal distance of the occiput (hereafter called width) is narrower in E. sipedon than in E. crotalinus, and the posterolateral ocellar tubercle is less pronounced in E. sipedon than in E. crotalinus.

The occipital crest of $E$. compositus is sinuous (mostly linear in E. sipedon) and the postocciput is easily visible in dorsal view (fig. 163); the same structure is barely visible in E. sipedon (fig. 159).

The structure of the postlamellar ridge provides the best means of separation between $E$. sipedon and sister taxa E. bothrops and E. viperinus. In E. sipedon, this Yshaped structure is short: the juncture of the lateral arms meets at or near the hind margin of each vulvar plate. The longitudinal stem also does not possess any circular or oval depressions on either side (figs. 187188). In E. bothrops and E. viperinus, the juncture of the Y-stem is well posterior to the hind margin of each vulvar plate, and a well-defined oval or circular depression exists on either side of the longitudinal ridge (figs. 183-184).

I have been unable to find any morphological differences between females of $E$. sipedon and $E$. heterodon. Body pattern (figs. 19, 23) and distribution (figs. 201-204) will serve to separate females of these closely related species.

## Remarks

Variation. - The male description is based on 11 males from Jalisco, Morelos, and Durango states, Mexico. The four from Durango are slightly larger (hind wing $30-31 \mathrm{~mm}$ ); and their thoracic pattern is similar to that of the holotype female (fig. 19). The other seven males are smaller (hind wing 28-29 mm), and the lower 0.60 of the dark humeral stripe is narrower. The Durango males, though larger, have a sparser venation. For example, three have no marginal cells behind fore wing paranal cells, one has one marginal cell in both wings; and all other specimens have one marginal cell, except for the left fore wing of one Acatlipa male, which has no marginal cells. Two
of the Durango males have a 3-celled anal triangle in both hind wings, while all others possess 4 cells. The cerci of the Durango males are less angulate and the tips more robust (fig. 106) than the other males, but other characters of the body and penis leave no doubt that they represent $E$. sipedon.

Variability for females is less pronounced than for males, and I can find no overall differences in venational patterns among the 12 females examined.

Female venational statistics ( $\mathrm{n}=12$, including holotype): number of marginal cells behind fore wing paranal cells: $1-3 / 0-3$; anx: fore wing 12-14/12-15, hind wing 9-12/9-11; pnx: fore wing 8-10/8-10, hind wing $9-11 / 8-12$; cs under pterostigma: fore wing $5-$ 7/5-7, hind wing 5-7/4-6. Hind wing $30-34 \mathrm{~mm}$. Abdomen $36-37 \mathrm{~mm}$.

When Calvert (1905) described this species, he had only 6 females available; and he noted differences in the condition of the occiput between the holotype and a paratype from Matamoros, Puebla (figs. $158-$ 159). Both forms were represented in his key. I cannot consider the differences noted by Calvert as useful, as much variation seems to exist in the shape of the occipital ridge in females of this and other species.

At the time of the original description, Calvert (1905) postulated that the female of E. sipedon could be $E$. boa, which was then known only from one incomplete teneral male. The discovery of more specimens of both species, and especially the capture of both males and females of E. sipedon in Durango, leaves no doubt that both are valid species.

Biology, - González (in litt.) collected this species at Palo Bolero, where adults perched on herbaceous vegetation on the banks of the Rio Sabinos. Its known distribution suggests that it is more widely distributed, but specimens are rare in collections. Specific ecological requirements may restrict its occurrence to certain sites, but a more likely answer to its apparent rarity is a lack of thorough collecting within its range. Collection dates range from 21 May (Cuernavaca) to 13 August (Nombre de Dios).

Distribution (fig. 203). - Erpetogomphus sipedon is known from northern Durango (Las Bocas [Calvert, 1907] settlement on the Rio Florida in extreme northern Durango near Villa Ocampo, approx. $26^{\circ} 25^{\prime} \mathrm{N} 105^{\circ} 25^{\prime} \mathrm{W}$, [Selander and Vaurie 1962]) south through Morelos and Puebla (approx. $\left.18^{\circ} 50^{\prime} \mathrm{N}, 99^{\circ} \mathrm{W}\right)$ and seems restricted to the highland area of central and western Mexico from altitudes of 1000 m (Palo Bolero) to about 2073 m (Las Bocas). González (in litt.) collected this species in Mexico state (Tonatico) during July, 1992.

## Material

Type data. - Holotype female: white printed label: '[Mexico] Guadalajara,/Jalisco,/July. Schumann:',
white, partially printed label: ‘Erpetogomphus 9 /sipedon Calv. Type /P. P. Calvert, det. 1905/B. C. A. Neur., p. xxx, 166/orig. of Pl. VIII, ff. 34, $40^{\circ}$ with '(form/a)' handwritten on left side; printed label 'Brit. Mus./1911-339.'; small round label with red margin: 'Holo--type'.

Other material (110 , 12 우, including holotype ¢). - Mexico: Durango: Nombre de Dios, just SE of Durango on Mex. Hwy 45 ( $23^{\circ} 51^{\prime} \mathrm{N}, 104^{\circ} 14^{\circ} \mathrm{W}$ ), $5900 \mathrm{ft} .(1800 \mathrm{~m}), 13$ Aug. 1947 (W. Gertsch, M. Cazier), 4 §§, 7 오 (AMNH, RWG); Morelos: Cuernavaca, 21 May 1898 (no collector), 1 (cuic); Cuernavaca, 7 July 1900 (C.C. Deam), 2 \& (Ummz); Palo Bolero, km. 18.3, route $95,5 \mathrm{~km}$. S. Acatlipa, 1000 m (approx. $18^{\circ} 45^{\prime} \mathrm{N}, 99^{\circ} 15^{\circ} \mathrm{W}$ ), 26 June 1985 (E. González), 2 ờ (UNAM); 18 June 1986 (E. González), 2 (UNAM), 12 July 1986 (E. González, V. Garcia), $30^{\text {on (rwg) }}$ ); Morelos: Pueblo Cocyotla, Hwy 421, about 30 km SW of Cuernavaca, 23 July 1992 (J. Daigle), $1 \delta^{\text {(JD }}$ (J); Puebla: Matamoros, no date, (Otis W. Barrett), 1 i (ANSP).

## Erpetogomphus lampropeltis Kennedy

## Description

Male. - Face entirely pale with dark markings as follows: along basal margin of labium, along frontoclypeal suture (less developed in E. l. natrix), base of antefrons; vertex all dark brown; antennae dark brown; anterior margin of vertex with an arcuate pit, one each anterolaterad to median ocellus; area between these pits tumid (fig. 138), occiput white green, wide, slightly tumid medially, crest entirely black, slightly raised, linear or slightly convex posteriorly, some with a small median notch, its hind margin covered with pale brown hairs; postocciput slightly concave, pale; rear of head dark brown.

Prothorax largely brown with pale green or grey green areas on anterior and posterior lobes, and with two small median spots on median lobe; contrasting dark/pale areas more prominent in nominotypic subspecies; pale colours of synthorax grey or grey green (E. l. lampropeltis) to green (E.l. natrix) with following dark brown areas: middorsal stripe, widening basally to collar, its upper end covering antealar sinus, extending posteriorly and connecting with broad, well defined antehumeral and humeral stripes; well defined sinuate second lateral stripe; third lateral stripe. Coxae, trochanters pale grey green with slight wash of brown posteriorly; tibiae pale grey green with defined superior surfaces brown, becoming dark brown distally; tibiae black with narrow strip of pale grey green along basal 0.30 to 0.50 of lateral margin; base and armature black.

Wings hyaline, basal wing venation brown; anterior margin of costa yellow up to pterostigma; pteros-
rigma black; fifth antenodal thickened in all wings (with rare exceptions).

Abdomen. Abdominal segment 1 pale grey green to green with basal 0.25 brown and with dorsolateral brown stripe; segment 2 with dorsolateral stripe sending a ventral stem posterior to auricle and (often) an isolated spot on anteroventral margin of segment, dorsolateral stripe often connecting with black annulus or ending just before; segment 3 with isolated black dorsolateral spor surrounding lateral carina; its anterior end tapering and somerimes touching anterior margin of segment, distal 0.25 of segment with dorsolateral black increasing in width posteriorly and touching dorsally near black annulus; segment 4 similar to segment 3, but two black spots broadly connected forming a continuous dorsolateral stripe occupying distal 0.80 of segment, this stripe often constricted in middle; segments 5 and 6 similar to segment 4 , but with middorsal pale areas becoming darker; segment 7 with anterior 0.50 pale, except for black transverse carina, becoming entirely tawny to black posteriorly; segments $8-10$ yellow brown, becoming black dorsally in most specimens; foliar extensions dark brown to black.

Cercus (figs. 107-108) ocher, in lateral view slightly concave dorsally near base, posterior 0.25 moderately angulate, the short tips slightly recumbent; cercus enlarged ventrally at basal 0.40 to 0.50 with inferior carina along same area, remainder slightly concave with tip forming a blunt point. Epiprocts yellow brown, gendly curved dorsally at $90^{\circ}$, tips (fig. 132) dorsoventrally flattened and divergently bluntly pointed.

Accessory genitalia (figs. 62-63). Anterior hamule dark brown, divided at basal 0.50 , lower branch forming a small, posteriorly directed truncate appendage; posterior hamule grey white, triangular, with anterior basal shoulder, tip bluntly rounded and armed with a small anteriorly directed tooth on posterior border; penis with lateral lobe small, roughly semicircular, serrate; cornuae (figs. 84-85) moderately widely separated, tips broadly rounded with well developed median lobes.

Female. - Head as in male, but with crest of occiput slightly more erect and convexly arcuate; posterior margin of postocciput slightly concave; prothorax and thorax as in male, but with pale areas generally more extensive; wings hyaline or with saffron infusion at base in some E.l. natrix, abdomen in well preserved specimens with pale colours primarily white with dull orange dorsally; black dorsolateral stripes and spots more reduced than in male, these stripes sometimes separated or greatly constricted on segments 4 and 5; cercus ocher.

Vulvar lamina (figs. 189-190) with prominent ba$\therefore$ plates, quadrangular laminar plates acuminate
posteriorly, meeting at anteromedial margin; cleft between plates forming a $90^{\circ}$ angle; V juncture of Y shaped postlamellar ridge at level of posterior margin of lamina.

## Diagnosis

This species is most similar to its congeneric relatives, $E$. sipedon and $E$. compositus, and is diagnosed under those species.

## Remarks

The original description of $E$. lampropeltis is brief and is based on an unspecified number of males and females from Sespe Creek, Fillmore, Ventura County, California. Williamson and Williamson (1930) originally described $E$. natrix in detail from 21 males and two females from Baja California, Mexico. An abstract of their diagnosis of males $E$. natrix from E. lampropeltis is as follows:
'The head and thorax of lampropeltis are duller and paler than in natrix.... the rear of the head is brown in lampropeltis and black in natrix; the thoracic brown is paler in lampropeltis and the pale areas are grayer, giving much less contrast in the thoracic pattern; the two lateral dark stripes are always joined, often very broadly in lampropeltis, and are not joined in.....natrix, .... the femora are brown in lampropeltis.....black in natrix, and the inferior dilated edge of abdominal segments 8 and 9 is black in lampropeltis and, in natrix, is colored light vivid brown like the sides of the segment adjacent to the edge......

The apparent differences in morphology described above fall well within the latitude of individual and geographic variation and do not, in my opinion, constitute specific differences. Further, several specimens of $E$. natrix from various parts of its range have maculation patterns more closely approaching those of $E$. lampropeltis. For example, the antehumeral and humeral stripes are briefly connected, the dorsum of abdominal segments 7-10 have black as in E. lampropeltis, and the inferior dilated margin of abdominal segments 8 and 9 is black. I can also detect no differences in the shape of the vulvar lamina between females of $E$. lampropeltis and $E$. natrix.

The forms are allopatric, separated by the Colorado Desert. The large series of E. lampropeltis (mostly topotypes) that I examined does show consistent differences in pale thoracic colouration and minor differences in thoracic maculation. I consider the forms to represent subspecies separable as follows:

1. Pale thoracic colouration grey green, antehumeral and humeral stripes broadly connected on dorsal 0.25 , after isolating a small pale spot below antealar sinus; second lateral stripe about twice as wide on dorsal 0.5 and often connecting with narrow third lateral stripe; dark thoracic stripes,
especially second and third, often not well defined (fig. 20); Ventura, San Diego, Los Angeles, and western San Bernardino counties, California

## E. lampropeltis lampropeltis

- Pale thoracic colouration vivid green, antehumeral and humeral stripes not connected, or only narrowly so; second lateral stripe narrower than in E. l. lampropeltis, not connecting with third lateral stripe (fig. 21); central Baja California, Arizona, New Mexico, western Texas, south through Durango, Mexico
E. lampropeltis natrix

Erpetogomphus lampropeltis lampropeltis Kennedy (figs. 20 -thx, 62 -hamules, 84 -penis, 107 -app, 132 ept, 138 -base of postfrons, 160 -vertex, 189 -vl, 204distr)

Erpetogomphus lampropeltis Kennedy, 1918: 297 (descr. $\delta$ Sespe Creek, Fillmore, Ventura Co., California). Needham and Heywood 1929: 80 (descr. ठิ, $\uparrow$ ); Williamson and Williamson 1930: 13 (summary of status); Needham and Westfall 1955: 147 (descr.); Pritchard and Smith 1956: 116 (key); Musser 1962: 14 (notes on larva); Borror 1963: 104 (common name); Montgomery 1968: 133 (distr.); Paulson and Garrison 1977: 157 (Calif.); Paulson 1982: 255 (U.S.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 115 (cat.).

Herpetogomphus lampropeltis. - Byers 1928: 5 (larva unknown).
Erpetogomphus lampropeltis lampropeltis. - Tsuda 1991: 95 (cat.).

## Description

Male. - Colouration and maculation as given under species account, key and in fig. 20.

Venational statistics: number of marginal cells behind fore wing paranal cells: 0-3/1-3. - anx: fore wing 9-14/10-14, hind wing 8-10/8-10; pnx: fore wing $7-$ $9 / 5-10$, hind wing $7-12 / 7-12$; cs under pterostigma: fore wing 4-5/4-5, hind wing 3-5/3-6; number of anal triangular cells: 3-4/3-5. Hind wing 28-31 mm. Abdomen $34-37 \mathrm{~mm}$.

Female. - Venational statistics ( $n=6$ ): number of marginal cells behind fore wing paranal cells: $0-4 / 1-3$; anx: fore wing 13-15/12-16, hind wing 9-11/9-11; pnx: fore wing 9-11/9-10, hind wing 9-12/9-12; cs under pterostigma: fore wing $5-6 / 5-6$, hind wing $5-$ $6 / 5-6$. Hind wing 32-36 mm. Abdomen 33-36 mm.

## Diagnosis

See account under $E$. compositus and key to subspecies of E. lampropeltis.

## Remarks

Variation. - Venational details of holotype male: number of marginal cells behind fore wing paranal
cells: $1 / 2$; anx: fore wing $12 / 12$, hind wing $9 / 9 ;$ pnx: fore wing $8 / 9$, hind wing $9 / 9$; cs under pterostigma: fore wing $5 / 5$, hind wing $5 / 4$; number of anal triangular cells: $4 / 5$. Hind wing 28 mm .

The large series of specimens primarily from the type locality shows little somatic colour and/or pattern variation, due, possibly, to its limited distribution. The fifth antenodal in the fore wings is generally thickened, but 7 of 40 wings of 20 males examined have the fourth antenodal thickened; and one wing has the sixth antenodal thickened. A 3-celled anal triangle is common in this subspecies, as almost half (19/40) of the wings possess this condition.

The female from Matillija Hot Springs has flavescent wing bases.

Biology. - At the type locality, this species was abundant during early September, 1982. I collected 54 males within about two hours. All preferred to sit on exposed rocks and were easily collected with a net. Hundreds of cast skins were attached to the exposed rocks. I did not find the species in smaller rivulets emptying into Sespe Creek. The only female I saw and collected was found dead in the water.

I returned to the locality in September, 1984, but the entire area had been fenced off. Five males were collected in the general vicinity, below Matillija Hot Springs, but they were rare and more wary.

This is a late summer species: the specimens from San Bernardino Co. collected in June are all teneral. Flight dates for specimens from Ventura Co. range from 10 July to 8 September.

Distribution (fig. 204). - The nominotypic subspecies has been collected only at three localities in Ventura County one in Sar Bernardinc County, one in Los Angeles County, and one in San Diego County, California. It has the most restricted distribution of any Odonata form in southern California.

## Material

Type data. - Holotype male: calif: Ventura Co., Sespe Creek, Fillmore, 7 Aug. 1915 (C. H. Kennedy); in usnm. I have illustrated the thorax (fig. 20), caudal appendages (fig. 107), and penis (fig. 84) of the holotype.

Other material ( $70 \delta^{\star}, 6$ 우, including holotype $\delta^{\pi}$ ). - U. S. A.: California: Los Angeles Co., San Gabriel Mtns., jct of N and W Fork of San Gabriel River, E of Hwy 39, 11 Aug. 1986 (D. Swinney), 10 (rwg), Ventura Co., Sespe Creek, 4.0 mi N of Fillmore, 600 ft., 8 Sept. 1971 (D. Paulson), $2 \delta^{\circ}$ (cc, RWG); 5 Sept. 1982 (R. W. and J. A. Garrison), $54{ }^{\circ}, 1$ if (RwG); Matillija Hot Springs, 14 Aug. 1927 (T. Craig), 10 , 1 ㅇ ( FSCA ); just below Matillija Hot Springs, 1400 ft ., 2 Sept. 1984 (R. W. and J. A. Garrison), 50 (RWG); Wheeler Hot Springs, 10 July 1927 (T. Craig), 40 , (fsca); San Bernardino Co., Deep Creek Public

Camp, 15 June 1957 (A. Menke, L. Strange), $1 \delta^{\top}$, 4 :
(Lacm); San Diego Co., La Jolla, La Jolla Indian Res. Campground, San Luis Rey River, 25 Aug. 1991 (W. F. Mauffray), $10^{\text {( }}$ (RwG).

Erpetogomplous Lampropeltis natrix Williamson and Williamson stat. n.
(figs. 21 -thx, 63 -hamules, 85 -penis, 108 -app, 1990vl. 204-distr)
Erpetogomphus narrix-Williamson \& Williamson, 1930: 19 (descr. of, if Baja Calif. Sur, Purissima, 12 October 1923). - Byers 1939: 50 (quotes Williamson \& Williamson, 1930); Fraser 1940: Pl. 6 (penis); Needham \& Westfall 1955: 148 (descr. §); Borror 1963: 104 (common name); Montgomery 1968: 133 (distr.); Paulson 1982: 255 (U.S., Mex.); Davies and Tobin 1985: 28 (cat.); Tsuda 1986: 87 (cat.).
Erpetogomphus lampropeltis. - Tinkham 1934: 215 (Tex.), Needham \& Westfall 1955: 147 (Tex.); Gloyd 1958: 8 (Tex.); Kormondy 1960: 122 (Ariz.); González \& Novelo 1991: 97 (Mex.); Novelo \& González 1991: 154 (descr. larva, Mex.).
Erpetogomphus lampropeltis natrix. - Bridges 1991: VII. 144 (cat.); Tsuda 1991: 95 (cat.). [Demotion of $E$. natrix to subspecies status was advised by me in litt. to Bridges and Tsuda.]

## Description

Male. - Colouration and maculation as given under species account, key, and fig. 21.

Venational statistics: number of marginal cells behind fore wing paranal cells: $0-3 / 0-2$; anx: fore wing 11-15/11-14, hind wing 9-11/9-11; pnx: fore wing 6-11/7-10, hind wing 8-14/7-12; cs under pterostigma: fore wing 4-6/4-6, hind wing 4-6/4-6; number of anal triangular cells: $3-4 / 2-4$. Hind wing $24-33 \mathrm{~mm}$. Abdomen $30-37 \mathrm{~mm}$.
Female. - Venational statistics ( $\mathrm{n}=17$, including allotype): number of marginal cells behind fore wing paranal cells: 0-6/0-5; anx: fore wing 12-15/12-15, hind wing 9-11/9-11; pnx: fore wing 7-11/7-12, hind wing $9-12 / 9-12$; cs under pterostigma: fore wing 4 $6 / 4-7$, hind wing 4-7/5-7. Hind wing $30-35 \mathrm{~mm}$. Abdomen $36-41 \mathrm{~mm}$.

## Diagnosis

See comments under $E$. compositus and key to subspecies of $E$. lampropeltis.

## Remarks

Venational details of holotype male: number of marginal cells behind fore wing paranal cells: $2 / 1$; anx: fore wing $13 / 13$, hind wing $9 / 9$; pnx: fore wing 9/9, hind wing 10/9; cs under pterostigma: fore wing 4/4, hind wing 5/4; number of anal triangular cells: 4/3. Hind wing 30 mm .

Venational details of allotype female: number of marginal cells behind fore wing paranal cells: $2 / 1$; anx: fore wing 13/13, hind wing $9 / 10$; pnx: fore wing $7 / 8$, hind wing $10 / 9$; cs under pterostigma: fore wing $6 / 5$, hind wing $5 / 5$. Hind wing 30 mm .

Variation. - Intrapopulational variation of body maculation appears to be as great as interpopulational variation. The antehumeral and humeral stripes may be connected or separate in any small series from a given locality: 1 of 3 specimens from Cane Springs, Mojave Co., 3 of 12 specimens from Cave Creek, Maricopa Co. (4 have these stripes barely connected), and 2 of 11 specimens from 15 mi E of Douglas, Cochise Co., Arizona, all have connected antehumeral and humeral stripes. Similar degrees of variation exist for the dorsal and dorsolateral darkening of abdominal segments $7-10$. Some specimens have little black on these segments (as noted by Williamson and Williamson, 1930, for their type series of $E$. natrix), but others have various degrees of darkening, and some specimens are as dark as I have observed for nominotypic E. lampropeltis.

A male I collected at Las Parras, Baja California, has the posterior 0.5 of the dorsum of abdominal segments 7 and all of segments $8-9$ with black. The male from 'Corralitos' is unusual in having narrow second and third lateral thoracic stripes. The upper 0.5 of the second lateral stripe is also lacking, but in all other aspects, this specimen is E. lampropeltis natrix by morphology and overall body colouration and maculation.

Biology. - Williamson and Williamson (1930) described the type locality. Gonzalez (in litt.) collected this subspecies with E. crotalinus in Durango state (La Michiliá). I collected one male of the species at Las Parras in September, 1985 (Garrison, 1986) as it sat upon an exposed rock in the middle of the small stream. It was the only individual I saw.

This subspecies can be common at certain sites in Arizona and New Mexico. Jo A. Garrison and I collected E. lampropeltis natrix on the East Fork of the Gila River at Grapevine Campground, where it was equally as common as $E$. heterodon. Adults landed on exposed rocks, exposed bleached logs in the center of the stream, or on the gravel shores. I have also taken it on sandy-bottomed desert streams, often in company with E. compositus. Novelo (pers. comm.) found this subspecies with the more common E. crotalinus in Durango state (La Michiliá).

I observed oviposition on 1 August 1992 at the Agua Fria River, Yavapai County, Arizona. Females appeared over the stream surface in the late afternoon (about 6: $15 \mathrm{p} . \mathrm{m}$. ), and flew rapidly over the water, dispersing their eggs by dipping the abdomen briefly into the water. This behaviour continued until sundown (about 7: 30 p.m.). A few males were still
present, sitting on emergent rocks near oviposition sites. No copulations were observed.

Altitudinal gradients for the species range from near 923 m (Tangle Creek, Yavapai Co., Ariz.) to 1754 m in New Mexico and southeastern Arizona (W of Portal). Like E. l. lampropeltis, E. l. natrix is primarily a late summer to autumn species. Flight dates range from 27 June to 10 October.

Distribution (fig. 204). - Though originally described from Baja California, E. lampropeltis natrix is widely distributed throughout the southwestern United States. It has been collected as far east as western Texas (Tinkham, 1934, Gloyd, 1958) and Durango and Chihuahua, Mexico. The most northerly records are from northcentral Arizona. I have seen no specimens from central or eastern New Mexico, although it probably occurs in those areas. The species apparently does not penetrate north of the Mogollon Rim in Arizona northwest into the Colorado Desert.

## Material

Type data. - Holorype male: Mexico: Baja California Sur: Purissima, 12 Oct. 1923 (J. H. Williamson); allotype female, same data but 6 Oct. 1923 (both examined), in ummz. I have illustrated the thorax (fig. 21), hamules (fig. 63), caudal appendages (fig. 108), and penis (fig. 85) of the holotype, and vulvar lamina (fig. 190) of the allotype.

Other material ( $101 \delta^{*}, 229$, including holotype of and allotype f). - U. S. A.: Arrzona: Cochise Co.: Bear Creek, Rte 61, 6 mi SE Parker Lake, Coronado National Forest (csuc); mouth Miller Canyon, Huachuca Mtns., (fsca, ummz), pond N of Slaughter Ranch, 15 mi E of Douglas, 3800 ft , (RwG); springs just E of Slaughter Ranch, 17 mi E of Douglas, (Rwg); Bear Creek, Rte 61, 6 mi SE Parker Lake, Coronado Nat'l Forest, (csuc), San Pedro River at Hwy 90 (FsCA); San Pedro River, jct. Charleston Rd., (byuc); San Pedro River, 9 mi SW of Tombstone (byuc); San Pedro River at Hereford Rd., (rwg); Southwestern Research Station, 5 mi W of Portal at John Hands Picnic Grounds, 5400 ft , (fsCA); Gila Co., Tonto Natural Bridge (Rwg); Webber Creek at Camp Geronimo, 8 mi ENE of Payson, (Rwg); Maricopa Co., Cave Creek by Ocotillo Rd, Cave Creek, (Rwg); Camp Creek by Cave Creek Rd., 12 mi NE of Scottsdale Rd., 2700 ft ., Tonto Nat'l. Forest, (rwg); Sycamore Cyn., 3.3 mi WNW of Hwy 87 (rwg); Mojave Co., irrigation ditches E of Cave Springs, (DRP); Pima Co., Sabino Cyn, N of Tucson (fsCA); Lower Sabino Cyn, (rwg); Upper Sabino Cyn, ca. 3 mi N of visitor center, N of Tucson (rwg); Santa Cruz Co., Sonoita Creek Ranch (on way to Salero Cyn), along Santa Cruz River, along Ariz. Hwy 82, 2 mi SW of Patagonia, (RWG); White Rock Campground, Peña Blanca Lake, 4200 ft , (RWg, JD); Yavapai Co., Agua Fria River at Black Canyon City, (rwg); Tangle Creek, nr Tangle Creek cabin, ca 33 mi N of Carefree, T9N-R5E, sec. 1, 2800 ff (RWG); on limestone rim of Montezuma's (sic) Well (National Monument) (Rwg); Sycamore Canyon, Atasco Mtns (lacm); New Mexico: Catron Co., Gila Cliff Dwellings, (rwg); West Fork Gila River at Gila Cliff Dwellings Nat'l Mon., 5700 ft , (Rwg); Grant Co., Gila River at Cliff (FSCA); Mangus Springs Creek,

Hwy 180, Mangus Springs, (rwc); Grapevine Campground, East Fork of Gila River at N. Mex. Hwy 15, 5600 ft , (Rwg); Texas: Jeff Davis Co., Fr. Davis, (fsca); Limpia Creek, 2-3 mi N of Ft. Davis, (ummz); Musquiz Creek, $61 / 2 \mathrm{mi}$ SE Ft. Davis (fsCa); Mexico: Chihuahua: Cascada de Basaseachic National Park, (csuc, byuc); San Diego Canyon, (byuc); Durango: Arroyo El Temazcal (camino a El Aleman), Reserva de la Biosfera La Michilíá, (unam); km 21 Camino Suchil-Reserva de la Biosfera La Michiliá, 'Corralitos', (unam); Sonora: Maycoba River, (BYUC); Maycoba River, W of Maycoba, (ByUC); 8 mi W of Maycoba River, (csuc); Maycoba River, Hwy 16, 13 mi W of Maycoba at San Francisco Bridge, (csuc); Rio Magdalena, Hwy 15 S of Immuris, (Byuc, csuc); small river W of Vecora, Rte 15, (csuc, byuc); Baja California Sur: Purissima (fSCA); Las Parras, about 18 km W of Hwy 1 on route to San Javier, SW of Loreto, (RWG).

## Erpetogomphus crotalinus (Hagen in Selys)

(figs. 22-thx, 41 -abd, 64 -hamules, 86 -penis, 109 -
app, 133-ept, 161 -vertex, $169-$ leg, 191, 192-vl, $205-$ distr)

Ophiogomphus crotalinus Hagen in Selys, 1854: 40 (21 sep.) ( 0 , $\frac{9}{}$ descr. from Mexico).
Erpetogomphus crotalinus. - Hagen in Selys 1858: 332 (72 sep.) (descr. ©ै, f, relationship with E. menetriesii); Walsh 1863: 253 (mentioned); Selys 1869: 174 (11 sep.) (Mex., attributes authorship to Selys); Selys 1873b: 519 ( 75 sep.) (list, attributes authorship to Selys); Kirby 1890: 61 (cat., attributes authorship to Selys); Calvert 1899: 38 (Tepic, Mex.); Calvert 1905: 165 (Mex.); Calvert 1907: 399 (Mex.), Calvert 1909: 481 (distr. in Mexico); Muttkowski 1910: 86 (cat.); Ris 1917: 153 (comparison with $E$. boa); Kennedy 1918: 298 (comparison with $E$. lampropeltis); Byers 1939: 50 (quotes Calvert 1907); Williamson and Williamson 1930: 12 (summary of status); Montgomery 1968: 133 (distr.); Paulson 1982: 255 (cat.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat., attributes authorship to Selys); Bridges 1991: VII. 54 (cat.); Tsuda 1991: 95 (cat.); Cannings and Garrison 1991: 478 (Mex.); González and Novelo 1991: 97 (Mex.); Novelo and González 1991: 150 (descr. larva, Mex.).
Herpetogomphus crotalinus. - Hagen 1875a: 43 (cat.); Selys 1879: 64 ( 2 sep.) (characters of genus); Kirby 1890: 61 (cat.).

## Description

Male. - Labium grey white, base of mandibles pale green becoming red brown distally, lips black, remainder of face pale green except for narrow wash of brown at base of antefrons, vertex mostly dark brown with green medially behind postoccipital tubercles; antennae dark brown; anterior margin of vertex with two lanceolate pits, each anterolateral to median ocellus; occiput bright green, broad, gently tumid medially, crest black, linear to emarginate medially, its rim covered with row of long brown hairs; postocciput green, smoothly concave or somewhat angulate medially, rear of head light green with brown near occipital foramen.

Prothorax predominantly green, most of middle lobe and posterior of frontal lobe marked with dark brown, a small middorsal green spot longitudinally divided by narrow dark brown line medially on middle lobe. Synthorax (fig. 22) entirely green with brown along margins of antealar crest in a few specimens with a hint of a dark brown antehumeral stripe. Coxae, trochanters pale green, femora green with streak of dark brown becoming wider distally, these streaks restricted to lateral surface of femora (fig. 169) and brown reduced to apical 0.50 of metafemora; a supplementary narrow, linear streak of dark brown more posteriorly placed on femora and, in some specimens, touching lateral streaks of brown distally; tibiae ranging from all yellow green except dark brown laterally. to all brown with yellow on lateral carinae; tarsi and armature black or with some yellow on exterior surfaces of second and third tarsomeres.

Wings hyaline, venation dark brown, anterior margin of costa yellow to base of pterostigma, this struccure brown, veins bordering it black.

Venational statistics. Fifth, rarely fourth or sixth, antenodal thickened in all wings; marginal cells behind fore wing paranal cells: $1-3 / 1-3$; anx: fore wing 11-13/11-13, hind wing 8-9/8-10; pnx: fore wing $7-$ 10/7-10, hind wing 8-10/8-10; cs under pterostigma: fore wing 4-6/4-7, hind wing 4-7/4-6; anal triangular cells: 3-4/3-5. Hind wing $26-31 \mathrm{~mm}$.

Abdomen (fig. 22). Segment 1 green, slightly fuscous basally, with small line of black at posterior articulation point; segment 2 all green with narrow dorsolateral streak of black above auricle and with poorly defined dorsolateral brown spot on posterior 0.25 of segment, dorsal 0.75 to 0.50 of annulus black; segment 3 pale green, becoming white ventrally with black along transverse carina and elongate black dorsolateral spot on posterior 0.25 of segment connecting to black annulus; segments 4-7 similar to segment 3 , but with narrow anterior offshoot of black laterally on transverse carina and with posterior midlateral black occupying posterior 0.50 of segments, middorsal pale area of these segments becoming progressively more tawny posteriorly, especially so on posterior 0.50 of segment 7 ; segments 8-9 ocher yellow with well defined dark brown dorsolateral stripe; segment 10 ocher becoming red brown at basal 0.50 . . 1 bdomen $31-36 \mathrm{~mm}$.
Cercus (fig. 109) yellow, strongly to moderately angulate dorsally at distal $0.50-0.30$; tip of cercus beyond angulation attenuate, drawn out to a blunt point, ventral margin with inferior carina at basal 0.40 to 0.50 . Epiprocts yellow brown, gently curved dorsally at $90^{\circ}$, extending to distal 0.25 or almost to level of tip of cercus, tips forming divergent points, dorsoventrally flattened (fig. 133).

Accessory genitalia (fig. 64). Anterior hamule
black, divided at upper 0.50 , lower branch forming small posteriorly directed truncate appendage; posterior hamule pale green, triangular, with anterior basal shoulder, tip rounded with a small anteriorly directed tooth on posterior margin; penis with semicircular spinose lateral lobe, cornuae separated; with medial lobe, and a long narrow flap at outer distal margin, membranous hood present but not overlapping. Abdomen 31-37 mm.
Female. - Head as in male but with crest of occiput (fig. 161) more emarginate medially, in some specimens with lateral margin arcuate; prothorax and synthorax as in male; legs with black areas more reduced than in male; wings with slight wash of yellow at base (more so in tenerals); abdomen as in male but with black areas more reduced on abdominal segments 1 3 ; dorsolateral brown stripes almost complete on segments 4-7 except for basal 0.05 of each segment, these stripes complete on segments $8-9$; cercus ocher. Abdomen $30-36 \mathrm{~mm}$.
Vulvar lamina (figs. 191-192) with broadly rectangular flaps meeting at anterior margin, juncture of $Y$ shaped postlamellar ridge at level of posterior margin of lamina.

Venational statistics: number of marginal cells behind fore wing paranal cells: $1-4 / 1-3$; anx: fore wing 11-13/11-13, hind wing 8-10/8-10; pnx: fore wing $7-$ 10/7-9, hind wing 8-10/8-10; cs under pt: fore wing 4-7/4-6, hind wing 4-6/4-6. Hind wing $26-32 \mathrm{~mm}$.

## Diagnosis

Erpetogomphus crotalinus is unique in having the extensor surfaces of the tibiae yellow (fig. 169). Some specimens have largely brown tibiae, but the lateral costate ventral margins are always yellow; all other congeners have dark brown to black tibiae. The almost immaculate synthorax is shared by only two other species: E. cophias (fig. 27) and some E. elaps (fig. 13), but the male cerci of these two species (figs. 100, 116) are never angulate as in E. crotalinus (fig. 109). The cornua of the penis (fig. 186) has a well developed median lobe and a long, acuminate tip, a condition unique to this species. This species is superficially similar to $E$. heterodon and is diagnosed under that species.

Females of $E$. crotalinus differ from other species by three characters of the head, the presence of anterolateral pits at the base of the frons, the relatively broad occiput, and the medial notch of the crest. The vulvar lamina (figs. 191-192) provides no discriminatory characters, and is similar to those of E. lampropeltis (figs. 189-190), E. compositus (fig. 194), E. heterodon (fig. 193), and E. sipedon (figs. 187-188).

[^15]les. The lectotype male is reasonably well preserved, although it lacks the accessory genitalia. I illustrate the cercus (fig. 109) and right hind leg (fig. 169) of the lectotype, and the vertex, occiput (fig. 161), and vulvar lamina (fig. 191) of the paralectotype.

Venational details for lectotype male: number of marginal cells behind fore wing paranal cells: 2/3; anx: fore wing $13 / 13$, hind wing $9 / 8 ;$ pnx: fore wing $10 / 10$, hind wing $10 / 10$; cs under pt: fore wing 6/6, hind wing 6/6; anal triangular cells: $4 / 5$. Hind wing 31 mm . Abdomen 36 mm .

Venational details for paralectotype female: number of marginal cells behind fore wing paranal cells: $1 / 1$; anx: fore wing $11 / 11$, hind wing $9 / 9$; pnx: fore wing $9 / 9$; hind wing $9 / 9$; cs under pt: fore wing $5 / 4$, hind wing $4 / 5$. Hind wing 29 mm . Abdomen 33 mm .

Variation. - Two teneral males of E. crotalinus reared in Arizona are superficially similar to $E$. heterodon in the blunt shape of the cercus, the vestige of an isolated antehumeral stripe in one specimen, and the fuscous condition of the extensor surfaces of the metafemora. However, the penis structure of both specimens is unmistakably that of E. crotalinus. Because the two Arizona specimens are teneral, I cannot determine if dark thoracic stripes similar to those of $E$. heterodon would have developed, or whether the maculation would have been the same as observed when mature.

The male described by Hagen in Selys (1858) has the darkest and most distinct antehumeral stripes that I have seen for any specimen of this species. The tibiae of a majority of specimens I examined are entirely dark brown with the lateral costae yellow; others, including the lectotype, have the entire ventral margin yellow.

Biology. - Specimens are apparently common at favored localities. Boris Kondratieff (in litt., 1990) states that adults of this species were 'on every [exposed] rock [in, Rio Macoba] - hundreds and hundreds of them.' During a two-day stay at Ajijic near Laguna Chapala, Oliver S. Flint, Jr., and M. A. Ortiz collected 28 males and 23 females. Calvert (1907) records the following concerning its habits: '[Progomphus borealis] and Erpetogomphus were found on sand- or mud-banks, hardly above water-level, along sides of an irrigating ditch into which the water from the baths of Santa Rosalia empties. On alighting on these banks, $P$. obscurus borealis held its abdomen slanting upward to form an angle of $45^{\circ}-60^{\circ}$ with the bank, while E. crotalinus held it nearly horizontal.' The male collected at Lago de Patacuaro in Michoacan is noted as taken on 'stream flowing into lake', the two males and one female from the pyramids at S. Juan Teotihuacan as 'on high grassy plains far from water'. González and Novelo (1991) record
the following about E. crotalinus (in translation): 'This is the most abundant gomphid in the area [the Biosphere Reserve of La Michiliá, Durango]. Like other members of the family, the adults of E. crotalinus have a seasonal flight pattern that, although it has not been exactly established, apparently is restricted to the months of May to October. The individuals of this species are characteristic inhabitants of both seasonal and permanent streams, easily recognized in the field by the bright green colour of the thorax. Males have the habit of resting on emergent rocks, a position they defend aggressively from other conspecific males.'

Erpetogomphus crotalinus has been collected with $E$. designatus, E. lampropeltis, and E. beterodon. Collection dates range from May through October (Calvert 1905).

Distribution (fig. 205). - Erpetogomphus crotalinus is restricted to the higher xeric areas of central and western Mexico. The species was reared from larvae collected by Minter Westfall, Jr., from the Slaughter Ranch, 15 mi W of Douglas, Cochise Co., Arizona; but no other collectors have encountered it there. I have also seen a pair from New Mexico (Eddy Co., Sitting Bull Falls), which may be strays from Mexico; its occurrence in the southwestern United States appears sporadic, and other odonatists have failed to find the species during their work in Arizona and New Mexico. Altitudinal data show a range of 1225 m (Mexico: Chihuahua: Santa Rosalia Springs) to 2290 m (Mexico: Mexico: Pyramids, S. Juan Teotihuacan). In the Paris Museum is a male labelled, 'Veracruz', which is probably in error.

## Material

Type data. - Lectotype male by present designation with following data: white handwritten label by Hagen, 'G. lineatus / Mexico'; white handwritten label, 'crotalinus' in an unknown hand; white, blackbordered printed label: 'Mus. Berol.'; two small printed labels, 'Hagen'; red label with 'Type [printed]/12334' (written by N. Banks); rectangular label [all printed unless otherwise stated]: 'erpetogomphus / crotalinus hagen [stamped]/ P. P. Calvert, det. 1905 [written]/ B. C. A. Neur. p. 165 [written]'. In mczc. The penis and hamules are missing and probably served as the basis for the freehand illustrations of these structures for Hagen in Selys (1858). Paralectotype female with following data: green printed label: 'Mexico/ Deppe.'; red printed label 'Typus'; yellow printed label, 'Zool. Mus./ Berlin', on reverse side written in ink, 'cat nr./ 2314; white label: 'syntypus $\frac{q}{}$ von: / Ophiogomphus / crotalinus Hagen' [written in an unknown hand]; yellow label (handwritten except as noted), 'Paralectotype [printed] 우 / Ophiogomphus / crotalinus Hagen/ des.
（printed）R．W．Garrison 1984＇．In ZMHB．
There is a male in the ZMHB with the same data as the paralectotype female，except as follows：green la－ bel，＇Mexico Deppe＇；green label，＇crotalinus／Hag．＇ ［both handwritten by Selys］；printed label，＇2314＇； and＇Syntypus of von／Ophiogomphus／crotalinus Hagen＇．Although the specimen is labelled as a type，I believe it is not a syntype and，in fact，that it repre－ sents the second male described by Hagen in Selys， 1858．I decided that the male in the MCzC and the fe－ male in the $Z M H B$ represent the original syntypes for the following reasons：The original（1854）descrip－ tion of Ophiogomphus crotalinus is brief and was based on a male and female．Hagen gave measurements for both sexes（abdomen：male 36 mm ，female 33 mm ； hind wing male 31 mm ，female 29 mm ）．These meas－ urements and the description of the male（＇Front of thorax yellow，with indication of two median stripes and a pale red humeral stripe．．．＇）correspond to the mCzC male and the zMHB female．Hagen in Selys （1858）described this species in greater detail and had access to a second，smaller male（abdomen 33 mm ， hind wing 29 mm ）．He stated，＇In the other specimen ［i．e．，the second male］，the stripes are brown and straight，quite visible．．．＇The original male is again characterized as follows，＂Thorax yellow green with－ out spots and．．．one can scarcely see the appearance of rwo median stripes and a thick antehumeral stripe．．．， which tallies with the earlier description of 1854 ．The male from the ZMHB corresponds with the smaller of the two measurements given by Hagen in 1858；and it also has a pair of easily visible antehumeral stripes， an apparently rare feature for this species．The zMHB male is unique in this respect．Even though the origi－ nal description by Hagen gives＇Musée de Berlin＇as the repository of the types，I strongly suspect that Hagen took the male with him when he was invited to join the faculty at the Museum of Comparative Zoology in 1867.

Other material（ $97 \delta^{*}, 44$ ㅇ，including lectorype $\delta$ and paralectotype 9）．－U．S．A．：Arizona：Cochise Co．，San Bernardino Ranch， 15 mi E of Douglas（FSCA）；NEW Mexico：Eddy Co．，Sitting Bull Falls，Guadalupe Mens．， riwg．WM）：Mexico：Chihuahua：Chihuahua Creek， 4 mi SW of Col．Juarez，（csuc）；Rio Pacheco，（csuc）；Santa Rosalia Springs， 1219 m ，（ANSP）；Durango：Durango City， ：Asp）；Rio Dalita and marshes， 6 mi E of Durango， 1890

Jalisco：Jopopau， 1667 m ，（ANSP）；Guadalajara， MCzC）：Laguna de Chapala（nr．Mex．Hwy 15）， 75 mi SE of Guadalajara，（rwg）；Ajijic nr．Laguna Chapala（USNM，

Ajiijic，（fSCA）；Rio La Sanguijuela，Degollado， －．．．（．）：rours 15，km 206，San Lorenzo， 9 mi W ．，．．．．．C）；Jalisco［no other locality］，（BMNH）； ．．．．I W of La Piedad，（UNAM）；Lago de ro． 2.6 mi W of Quiroga， 2042 m ，（FSCA）；Route 15， ．．Jan Lorenzo． 8 mi W of Tuxpan，（USNM）；Roure Carapan，（USNM）；Route 15，km 291，near Nacional Insurgente Morelos，（UsNm）；
stream 0.5 mi S of Tuxpan， 1798 m ，（DRP）；Hidalgo：ditch－ es around Tasquillo， 1700 m ，（DRP）；Rio Tula at Puente Tasquillo，（USNM）；Tecozautla，Rio San Juan， 1710 m ， （RWG）；Tecozautla，Rio Tecozautla， 1750 m （RWG）； Tecozautla，arroyo San José del Desierto， 1730 m（RWG）； Mexico：Route 15，km 125，El Salitre，（USNM）；Pyramids，S． Juan Teotihuacan， 2290 m ，（FSCA，cUIC）；Morelos： Cuernavaca（BMNH，ANSP，CUIC）；Puebla：ditch just NW of Tehuacan， 1700 m ，（DRP，TWD）；Veracruz：Veracruz［prob－ ably erroneous locality］，（MNHP）；Mexico［no other local－ ity］，（ZMHB）．

## Erpetogomphus heterodon sp．n．

（figs．28－thx，42，43－abd，65－hamules，87－penis， 110，111－app，162－vertex，170－leg，193－vl，206－ distr，230－wings）

Type data．－Holotype male．U．S．A．：New Mexico：Catron Co．，Tularosa River just E of Aragon on N．Mex．Hwy 12， 28 July 1984 （R．W．and J．A． Garrison）．Allotype female．Same data as male．Both in USNM．

Paratypes（ $360^{\star}, 2$ ） ）．－New Mexico：Catron Co．， same data as holotype and allotype，27－28 July 1984， 22 むै， 1 ㅇ（RWG）；same data， 23 August 1964 （Clifford Johnson）， 3 むt， 1 오（DRP，FSCA）；Gila Cliff Dwellings， 30 July 1983 （John E．Hafernik，Jr．）， 4 © （rwg）；Grant Co．，Grapevine Campground，East Fork of Gila River at N．Mex．Hwy 15， 5600 ft．， 27 July 1984 （R．W．and J．A．Garrison）， 11 （RWG）； Texas：Jeff Davis Co．，Limpia Canyon in Davis Mtns． State Park， 23 June 1958 （M．J．Westfall，Jr．）， 10 º （fsCA）；Limpia Creek at Tex．Hwy 17， 3.4 mi NE of Ft．Davis， 4500 ft．， 24 August 1977 （Dennis Paulson， Susan Hills）， 10 （DRP）；Reeves Co．，Balmorhea State Park， 13 Sept． 1983 （S．W．Dunkle）， 1 ô（swd）； Mexico：Chihuahua： 6 mi NNE of Boquilla， 16 July 1960 （Scheibner）， $1 \delta^{\top}$（RWG）；Chihuahua Creek， 4 mi S of Col．Juárez， 25 Aug． 1986 （B．C．Kondratieff）， 10 （csuc）；＇Bosuchel［probably Basuchi］＇， 18 Aug． 1950 （R．F．Smith）， 2 o $^{\circ}$（AMNH，RWG）．

## Description

Holotype male．－Entire face pale light green， slightly darker on postclypeus and anterior of frons； labrum，base of mandibles，labium ivory；vertex and antennae black；base of antefrons with transverse fur－ row，tumid medially，laterally with a pair of pits；po－ stocellar tubercles small，no postocellar ridge；occiput pale green，slightly tumid medially，hind margin frin－ ged with long hairs，crest with black along lateral margin，slightly emarginate medially；postocciput gr－ een，transverse；rear of head dark brown，becoming pale green laterally．

Prothorax light green with dark brown on middle lobe extending laterally to pleura，two adjacent later－ odorsal small green spots on medial lobe．Synthorax
(fig. 23) light green with following dark brown: inverted wedge-shaped middorsal stripe, its ventral end not touching collar, its dorsal end forming a narrow stripe below antealar carina and meeting narrow humeral stripe; small isolated antehumeral stripe; vestigial second lateral stripe extending from metacoxa to metaspiracle; upper end of third lateral stripe. Coxae, trochanters ivory, femora ivory with black external surfaces spreading to lateral areas at distal 0.25 (fig. 170), tibiae black with small lateral ivory stripes along basal 0.5 ; tarsi and armature black.

Wings (fig. 230, paratype) hyaline, venation dark brown, basal wing venation brown; anterior margin of costa yellow; pterostigma brown, darker around margin.

Venational details. Fifth antenodal thickened in all wings; number of marginal cells behind fore wing paranal cells: 2 ; anx: fore wing $14 / 12$, hind wing $9 / 9$; pnx: fore wing $9 / 8$, hind wing $10 / 10$; cs under pterostigma: fore wing $4 / 6$, hind wing $6 / 4$; anal triangular cells: 4 . Hind wing 33 mm .

Abdomen (fig. 42). Segment 1 pale orange-yellow dorsally with narrow black lateral stripe, white laterally; segment 2 similar to segment 1 but with black along anterior margin of segment laterally; segment 3 similar to segment 2 but with black along transverse carina and posterior annulus; segments 4-7 similar to segment 3 but black dorsolateral stripe incomplete anteriorly so that dorsal orange-yellow merges with lateral white and with black along transverse carina, expanding ventrally, and with a similar mark posteriorly just before black annulus; segments 8-9 predominantly yellow-orange with black denticles dorsally, incomplete black dorsolateral stripe extending from anterior margin of segment but ending just before posterior margin; segment 10 yellow-orange posteriorly, orange-brown anteriorly. Abdomen 40 mm .

Cercus (fig. 110) yellow brown, strongly angulate near middle, ventral margin with inferior carina at basal 0.30 . Epiprocts curved at distal 0.75 , tips darkened, bluntly pointed, slightly divergent.

Accessory genitalia (fig. 65, paratype). Anterior hamule black, divided at upper 0.30 , lower branch forming a small, truncate appendage; posterior hamule white, triangular, with anterior basal shoulder, tip with small, anteriorly recurved tooth; penis with small serrated lateral lobes, cornuae (fig. 87, paratype) moderately long, widely separated tips broadly rounded.

Allotype female. - Overall colouration similar to male but with more extensive pale areas on abdomen and legs; furrow anterior to medial ocellus as in male; vertex (fig. 162) with postocellar tubercles not connected but with small remnant of ridge extending medially; occiput simple, posterior margin raised with row of hairs; postocciput green, transverse.

Venational details. Fifth antenodal thickened in all
wings; number of marginal cells behind fore wing paranal cells: $3 / 2$; anx: fore wing $13 / 12$, hind wing 10/10; pnx: fore wing $7 / 8$, hind wing $9 / 10$; cs under pterostigma: fore wing $6 / 6$, hind wing $6 / 6$. Hind wing 36 mm .

Abdomen (fig. 43) as in male, but with black dorsolateral stripe reduced, with black extending anteriorly before transverse carina. Abdomen 37 mm .
Vulvar lamina (fig. 193) with broadly planar rectangular flaps meeting at anterior margin; juncture of Y-shaped postlamellar ridge at level of posterior margin of lamina.

## Diagnosis

This species is related to E. crotalinus and E. sipedon. Erpetogomphus crotalinus lacks the dark thoracic stripes present in $E$. beterodon (fig. 22); the extensor surfaces of the tibiae of $E$. beterodon are black (fig. 170) (yellow in E. crotalinus); and the cornuae of the penis in $E$. heterodon are broadly rounded (fig. 87) (long and acutely pointed in E. crotalinus [fig. 86]).
Males of $E$. heterodon are most similar to $E$. sipedon, but they differ in thoracic maculation shape of the epiprocts, and structure of the penis. In E. sipedon, the antehumeral stripe is long and joined with the humeral stripe (fig. 19); in E. heterodon, this stripe is vestigial and disjunct. In E. sipedon, the tips of the superior appendages are slender and acuminate (figs. 105-106), not obtusely acuminate as in $E$. heterodon (figs. 110-111). The cornua of the penis in $E$. sipedon (fig. 83) possess a mesal lobe absent in $E$. heterodon (fig. 87).
Structurally, the females of E. crotalinus, E. heterodon, and E. sipedon approach one another closely. The latter two species have a relatively narrow occiput with the width across the crest about 4 times as long as its greatest length. The occiput in E. crotalinus is wider, about 3 times the width. I have been unable to detect morphological differences between females of E. heterodon and E. sipedon. Body maculation must be used to separate females.

## Remarks

Variation. - Slight variation occurs in males within the type series. The dark dorsolateral stripe on abdominal segment 3 may be broken in the middle. In the male from Mexico ( 6 mi NNE Boquilla), this stripe is reduced to an isolated spot on the basal 0.30 of the segment. Similarly, the extent of black markings on abdominal segments 8-10 may be reduced.

Venational statistics for type series. Males: Number of marginal cells behind fore wing paranal cells: 0-3; anx: fore wing 11-14/11-15, hind wing 8-11/8-11; pnx: fore wing $7-11 / 7-11$, hind wing $8-12 / 8-11$; cs under pterostigma: fore wing 4-7/4-6, hind wing $4-$ 7/4-7; anal triangular cells: 3-4. Hind wing 32-34
mm . Abdomen $36-40 \mathrm{~mm}$.
Females: Number of marginal cells behind fore wing paranal cells: 1-4; anx: fore wing 13/12-13, hind wing $8-10 / 9-10$; pnx: fore wing 7-9/8-9, hind wing 8-10/8-10; cs under pterostigma: fore wing 5-6/5-6, hind wing 5-6/5-6. Hind wing $34-36 \mathrm{~mm}$. Abdomen $36-37 \mathrm{~mm}$.

Biology. - Adult males were taken along sandy margins of the Tularosa River in New Mexico. They often sat on the bank facing the river and were easy to approach. I collecred one female as it hovered over the river ovipositing. Another teneral female was flushed from nearby vegetation. At the East Fork of the Gila River, males sat on emergent rocks in the stream or by the river's edge. They occasionally made swift sorties, usually to chase another male. At this locale, E. heterodon was sympatric with E. lampropeltis natrix, Altitudinal data show $E$. heterodon to occupy high, mostly xeric areas of the southwestern United States and northern Mexico. Collection dates are from 23 June to 13 September at elevations of 1370 m (Limpia Creek, Texas) to 1700 m (Grapevine Campground, New Mexico).

Distribution (fig. 206). - Erpetogomphus heterodon is known only from western Texas, New Mexico, and northern Mexico. It has been collected with $E$. crotalinus in at least one locality in Mexico (Chihuahua state, Rio Pacheco, 27 Aug. 1986, B. Kondratieff, S. Dunkle, pers. comm.), and is apparently allopatric with E. sipedon.

## Erpetogomphus compositus Hagen in Selys

(figs. 24, 25-thx, 66, 67-hamules, 88, 89-penis, 112, 113 -app, 145 -ept, 139 -base of postfrons, 163 -vertex, 194-vl, 206-distr)

Erpetogomphus compositus Hagen in Selys, 1858: 660 (400 sep.) ( $q$, 'Rivière Peros [sic. Pecos] (Texas occidental)'). Hagen in Selys 1859: 536 ( 10 sep.) (descr. ㅇ) ; Hagen in Selys 1873a: 740 ( 12 sep.) (descr. ò, ㅇ, 'Le nord de la Californie"); Selys 1873b: 519 (75 sep.) (list); Calvert 1905: 166 (Ariz.); Calvert 1908a: 45 (Ariz.); Muttkowski 1910: 86 (cat.); Calvert 1912a: 289 (mentioned); Williamson 1914a: 226 (Ariz.); Kennedy 1917a: 544 (Calif.); Seemann 1927: 22 (Calif.); Byers 1928: 51 (larva unknown); Needham and Heywood 1929: 80 (descr.); Williamson and Williamson 1930: 12 (summary of status); Tinkham 1934: 215 (W. Tex.); Ahrens 1938: 11 (Ariz., Utah); Fraser 1940: pl. 5 (penis); Ferguson 1940: 5 (Tex.); Needham and Westfall 1955: 144 (descr.); Pritchard and Smith 1956: 116 (key); Gloyd 1958: 8 (W. Tex.); Musser 1961: 54 (larva, Utah); Musser 1962: 14 :larva, Utah); Borror 1963: 104 (common name); Cruden 1964: 81 (Calif.); Montgomery 1968: 133 (distr.); Paulson and Garrison 1977: 151 (Washington); Molnar and Lavigne 1979: 130 (Wyo.); Paulson 1982: 255 (distr.); Paulson 1983: 67 (Wash.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat., attributes authorship to Selys); Bick 1990: 3 (Idaho); Bridges 1991:

VII: 49 (cat.); Tsuda 1991: 5 (cat.).
Gomphus compositus. - Hagen 1861: 99 (desc. ㅇ, Tex.).
Gomphus (Herpetogomphus)? viperinus. - Hagen, 1873 (misidentification, Yellowstone).
Herpetogomphus compositus. - Hagen 1874: 597 (Yellowstone, Tex.); Hagen 1875a: 42 (Tex.); Hagen 1875b: 918 (N. Mex.); Selys 1879: 64 (2 sep.) (characters of genus); Kirby 1890: 60 (cat., attributes authorship to Selys); Calvert 1899: 386 (characters of genus); Currie 1903: 303 (Ariz.); Osburn 1905: 186 (mentioned); Byers 1928: 5 (larva unknown); La Rivers 1938: 85 (Nev.); La Rivers 1940a: 112 (Nev.); La Rivers 1941: 177 (Nev.).
Erpetogomphus coluber Williamson and Williamson, 1930: 17 ( ${ }^{\top}$, San José de Comandu, Baja Calif. Sur). Needham and Westfall 1955: 143 (desc.); Borror 1963: 104 (common name); Montgomery 1968: 133 (distr.); Montgomery 1973: 239 (comment on name); Paulson 1982: 255 (distr.), 266 (synonymy with E. compositus); Bridges 1991: VII: 49 (cat.).

## Description

Male. - Entire face pale grey green with dark markings as follows: along basal margin of labium and median extension ending before distal margin (almost absent in some specimens), frontoclypeal suture, base of antefrons; vertex all dark brown or with pale median area extending posteriorly from median ocellus to anterior margin of occiput; antennae dark brown; anterior margin of vertex forming a trough at junction of postfrons, this trough with a pair of deeper pits anterolateral to median ocellus (fig. 139); occiput white green, wide, planar with slight tumid median area, crest barely convex, slightly to broadly emarginate medially, its hind margin covered with pale brown hairs; postocciput white green, transverse, not visible dorsally, rear of head dark brown.

Prothorax pale green with brown dorsolaterally on middle lobe; synthorax (fig. 24) predominantly pale green with following dark brown: well defined middorsal stripe widening to collar, its upper end extending along antealar sinus connecting with well defined humeral stripe; antehumeral stripe widened dorsally and connected basally at mesinfraepisternum; lower part of humeral stripe extending posteroventrally but not connecting with well defined sinuate second lateral stripe; lower 0.50 of this stripe encompassing metaspiracle, constricted along posterior margin at 0.50 and abruptly widening dorsally before turning anterodorsally toward subalar carina; complete third lateral stripe ending behind posterior margin of metacoxa. Pale colour of thorax light green with lighter tone almost becoming white in areas between antehumeral and humeral stripes, and between second and third lateral stripes. Coxae, trochanters white grey with slight wash of brown, tibiae pale grey green with defined superior surfaces black, basal 0.40 of metafemora with streaks of grey green or with basal 0.60 of this area largely pale, tibiae black with narrow stripe of pale grey green along basal 0.30 of lateral margin in
some specimens; tarsi and armature black.
Wings hyaline with slight wash of yellow at extreme base (more so on teneral specimens), basal wing venation brown, especially anteriorly, anterior margin of costa yellow white except for anterior margin of pterostigma, costa beyond pterostigma white or darkened, but contrasting with black of pterostigma; pterostigma black.

Venational statistics. Fifth antenodal thickened in all wings; marginal cells behind fore wing paranal cells: $0-4$; ; anx: fore wing $10-15 / 10-15$, hind wing $7-$ 11/7-11; pnx: fore wing 6-11/6-10, hind wing 7 $11 / 7-11$; cs under pterostigma: fore wing 3-6/3-6, hind wing 3-6/3-6; anal triangular cells: 2-5. Hind wing $25-32 \mathrm{~mm}$.

Abdomen. Segment 1 pale grey green with broad basal semicircle of brown laterally, its posterior end touching lateral articulated area; segment 2 white with dorsolateral brown stripe sending a ventral stem posterior to auricle, and connecting with black annulus, a small brown spot at anteroventral margin of segment; segment 3 with isolated black dorsolateral spot surrounding lateral carina, distal 0.25 of segment with dorsolateral black increasing in width posteriorly and touching dorsally near black annulus; segment 4 similar to segment 3 , but isolated black of transverse carina longer and touching or connecting with distal black, thus forming an incomplete midlateral stripe occupying distal 0.75 of segment and isolating ventral longitudinal wedge of white; segments 5 and 6 like segment 4 , but with middorsal white becoming fuscous; segment 7 with anterior 0.5 white, except for tawny transverse carina, becoming entirely tawny posteriorly, some specimens with varying degrees of lateral black on posterior 0.25 of segment or in specimens from Baja California ( $E$. coluber) with tawny areas becoming black; segments $8-10$ yellow brown with darker red brown dorsally, in Baja California specimens this dark red brown replaced with well defined black on segments 8 and 9 , and dark brown on segment 10 . Abdomen 32-39 mm.

Cercus (figs. 112-113) ocher, in lateral view slightly concave dorsally near base and with tips slightly recumbent, cercus enlarged ventrally at basal 0.45 with inferior carina along this same area, remainder slightly concave with tip forming a blunt point. Epiprocts yellow brown, gently curved dorsally at $90^{\circ}$, tips (fig. 134) dorsoventrally flattened and truncate or obliquely truncate.

Accessory genitalia (figs. 66-67). Anterior hamule dark brown, divided at basal 0.5 ; lower branch forming a small, posteriorly directed truncate appendage; posterior hamule grey white, triangular; with anterior basal shoulder, tip bluntly rounded and armed with a small anteriorly directed tooth on posterior border; penis with lateral lobe small, roughly semicircular,
serrate; cornuae (figs. 88-89) moderately long, widely separated, tips broadly rounded.
Female. - Head as in male but pale areas more extensive, especially on vertex, where dark brown may be reduced to areas around ocelli and postocellar protuberances; occiput narrower than in male, with erect curvilinear occipital crest separating easily visible postocciput (fig. 163), posterior margin of postocciput slightly concave; prothorax and thorax as in male, but with pale areas more extensive; wings with yellow at base more extensive, often with markings within second series of antenodals to level of arculus and cubitoanal area in both wings.
Abdomen in well preserved specimens with pale colours primarily white with pale orange dorsally, black dorsolateral stripes reduced and separated in their middle on segments $3-6$; segment 7 similar to male but with posterior 0.25-0.30 darker laterodorsally; segments $8-10$ tawny, with varying amounts of dark brown, especially dorsally, on segments 8-9; cerci ocher. Abdomen 31-37 mm.
Vulvar lamina (fig. 194) with prominent basal plates, quadrangular laminar plates acuminate posteriorly, meeting at anteromedial margin; cleft between plates forming a $90^{\circ}$ angle; stem of Y-shaped postlamellar ridge at level of posterior margin of lamina.

Venational statistics: number of marginal cells behind fore wing paranal cells: $1-6 / 0-5$; anx: fore wing 12-15/12-14, hind wing 8-11/9-11; pnx: fore wing $7-$ 11/8-10, hind wing 9-11/8-11; cs under pterostigma: fore wing $3-5 / 4-6$, hind wing. Hind wing $30-32 \mathrm{~mm}$.

## Diagnosis

This common, distinctive species is easily separated from its nearest ally, E. lampropeltis, by several structural and maculation characters. Males of E. compositus differ from E. lampropeltis in lacking the characteristic middorsal angulation of the cercus. The tip of the epiproct in $E$. compositus is truncate (fig. 134), but this structure in E. lampropeltis is obliquely and bluntly pointed (fig. 132). Depressions in male and female anterolateral to median ocellus will also separate the species: in E. compositus, an arcuate trough has more pronounced pits anterolateral to the median ocellus (fig. 139). In E. lampropeltis, the anterolateral pits are deeper and the intervening area directly anterior to the median ocellus is not as depressed (fig. 138). This structure in E. lampropeltis more closely approaches the condition found in E. viperinus (fig. 137).

Expetogomphus compositus, though a boldly marked species, is lighter than E. lampropeltis. The occipital crest in $E$. compositus is largely pale, though there may be some darkening along the lateral 0.30 of margins in specimens from Baja California. In E. Lampropeltis, the occipital crest is entirely dark brown and provides
a striking contrast to the pale occiput. The thoracic patterns of the two species are different (figs. 20-21, 24-25), and the dorsolateral black stripes of abdominal segments 4-6 of E. lampropeltis are more extensive.

In the field, pale thoracic colouration will easily separate both species. Erpetogomphus compositus is unique in having a pale green thorax with intervening grey white areas between the antehumeral and humeral stripes and second and third lateral stripes. In E. lampropeltis, the thoracic colouration is grey green (E. Lampropeltis lampropeltis) or dark green (E. lampropeltis natrix).

Females of E. compositus differ from E. lampropeltis in the shape of the occiput. In E. compositus, the occiput is narrow and the postocciput is clearly visible in dorsal view (fig. 163). In E. lampropeltis, the occiput is broad and the postocciput is not visible dorsally (fig. 160). Secondly, the area in front of the median ocellus mirrors the condition stated for males above. The vulvar laminae of the two species appear indistinguishable.

## Remarks

Williamson and Williamson (1930) described in detail Erpetogomphus coluber, comprising 31 males from San José de Comandu, Baja California. In comparing $E$. coluber with other species, they stated that E. coluberwould, in Calvert's (1905) key to the genus '...run to AA, page 160 , and if the individual be one with the facial dark markings reduced, it will run to H. compositus. Drs. Calvert and Kennedy regard coluber as distinct from any described species.' The Williamsons sent two males to Calvert, who commented, 'No special reason for thinking it diadophis $[=$ E. eutainia]; like a small compositus, differs in size; has broader metepisternal dark stripe; brown on [abdominal segments] 3-7 more extended; brown on 810 much darker; darker lines or stripes on the frontoclypeal and clypeo-labral sutures'. Finally, the Williamsons commented on the affinities of $E$. coluber to E. compositus as follows: 'Compositus is certainly its closest relative and the derivation of coluber from compositus, through geographical isolation in Baja California, is almost certain and is a case exactly parallel, so far as speciation goes, to that of certain species, of several genera, endemic in Florida. Coluber is separated from compositus by its darker color, especially of the last four abdominal segments and by venational characters, especially the two-celled anal triangle and the single row of cells posterior to A in the front wing.
I have seen the holotype and 15 paratypes of E. coluber and believe, as did Paulson (1982), that they represent diminutive, melanic examples of $E$. composit:s. The specific differences ascribed to E. coluber are
those of venation only, and I believe these to be correlated to their smaller size. I have found no differences in body morphology, including the accessory genitalia.

Variation. - Two specimens from San Diego County appear intermediate to $E$. coluber and $E$. compositus s.s. These males have more pronounced dark thoracic stripes, but the dorsum of abdominal segments 8-10 lacks dark brown markings typical of $E$. coluber. However, other E. compositus have varying amounts of dark brown on these segments. The hind wing anal triangle in 3 of 16 specimens of $E$. coluber has 3 anal triangular cells instead of 2 , and a 2 -celled anal triangle, though rare, does exist in $E$. compositus (s.s.). For example, of 21 males I measured from California, Arizona, and another from northern Baja California, one of the San Diego males and another from Riverside, California, had one wing each with two cells. These two specimens had shorter hind wings ( 26 mm and 28 mm , respectively) than is typical for $E$. compositus (s.s.). Statistics for the number of anal triangular cells show it to be variable within $E$. compositus (s.s.). The number of wings with range of stated conditions is (number of wings follows in parentheses): 3 (19), 4 (19), 5 (2). Hind wing lengths from the San Diego males are $25-26 \mathrm{~mm}$, well within the range for $E$. coluber. The next smallest specimen is a male from Riverside ( 28 mm ), the same male with two anal triangular cells in one hind wing.

Most wings of $E$. coluber which I have examined have no marginal cells behind the fore wing paranal cells, but 6 of 32 (19\%) did. Eight of 42 (19\%) wings of $E$. compositus (s.s.) had no marginal cells.

Williamson and Williamson (1930) warned of fallibility of some of their characters among the paratypes they examined: 'The face markings are very pronounced in some and scarcely discernible in others; there is considerable variation in the extent of the dark thoracic markings [fig. 25] and this variation occurs independently on the mesepisternum and metepisternum, so an individual may have the dark humeral and antehumeral relatively extensive and the two dark lateral stripes relatively reduced and vice versa.

With the exception of specimens from Baja California and the two specimens from San Diego, body colouration appears remarkably constant throughout its range. The male from Baja California Norte (Guadalupe Hot Springs) is more like typical E. compositus, but it has no marginal cells behind the fore wing paranal cells. The hind wing length (29 mm ) and other venational characters indicate a closer relationship to more northerly populations.

The female holotype was collected with the type series of $E$. designatus and was probably collected in the vicinity of Roswell, New Mexico in 1854-1855. See
remarks under $E$. designatus for further details.
Venational details. Holotype female of E. compositus. number of marginal cells behind fore wing paranal cells: 2/4; anx: fore wing 13/13, hind wing 9/10; pnx: fore wing $8 / 9$, hind wing $9 / 8$; cs under pterostigma: fore wing $5 / 6$, hind wing $5 / 6$. Holotype male of E. coluber: number of marginal cells behind fore wing paranal cells: $0 / 0$; anx: fore wing $12 / 12$, hind wing $9 / 9$; pnx: fore wing $7 / 7$, hind wing $9 / 9$; cs under pterostigma: fore wing $2 / 2$, hind wing $4 / 4$.

Biology. - This species and Progomphus borealis McLachlan in Selys are the two most conspicuous gomphid elements along most desert streams and irrigation ditches in the southwestern United States. In Arizona, this species occurs commonly in the Lower Sonoran Zone, and in ecotonal fingers into the Upper Sonoran Zone. Kennedy (1917a), who collected E. compositus at certain sites in the northern Central Valley of California, writes: 'One female, a teneral, was taken on the irrigating ditch across the river from Oroville.' Collecting along the lower Truckee River south of Pyramid Lake, Nevada, Kennedy (1917a) mentions that 'This widely spread species occurred sparingly on the riffles of the lower Truckee. The males appeared to be more nervous and more touchy, flying farther for conflict with passing males than the males of Ophiogomphus morrisoni nevadensis.'

The two males collected at the Boyce Thompson Southwestern Arboretum in Pinal County, Arizona, had the following annotation by H. K. Gloyd: '[in] Creek, arboretum. Canyon, after rain and run-off almost over. between $5 \& 7: 00$ p.m., water low, few dragonflies seen'. Williamson and Williamson (1930) provided a lengthy account of the type locality and assemblage of Odonata present with Erpetogomphus coluber. In August, 1972, I collected several E. compositus with Stylurus plagiatus (Selys) at Riverside Park, Yuma, Arizona. Adults were taken in the late afternoon on grassy areas shaded by trees. Adults of this species and Progomphus borealis were abundant along Big Chico Creek in Chico, Butte County, California in June and July of 1974, 1976, and 1978. They sat on exposed sand bars near the edge of the creek, and both were difficult to approach. Progomphus borealis was the more wary. In the late afternoon, I have collected E. compositus away from water, resting on dry desert scrub. Collection dates range from 24 May (Nevada: Clark Co.) to 13 Dec. (Calif.: San Bernardino Co.).

Distribution (fig. 206). - Erpetogomphus composit$u s$ is a desert species found commonly in the southwestern United States. It has been taken as far east as Dallas (Hagen, 1875a), but most records from Texas are from the western portion of the state. It occurs in eastern New Mexico and is common in low desert regions of Arizona, especially southwest of the

Mogollon Rim. It is common in the arid regions of southern California and penetrates north through the Central Valley to Chico. Paulson and Garrison (1977) list it from south central Washington. The species was listed from Oregon by Hagen (1875a, 1875b) with no specific locality); but specimens were recently collected from the John Day River, Deschutes Co., Oregon (Valley 1993). Erpetogomphus compositus occupies river systems in the Great Basin of Nevada, but its most easterly distribution is unknown. Bick (1990) lists a specimen from Owyhee Co., Idaho. Hagen (1874) records it from the Yellowstone, but it was probably taken in a more arid environment around what is present-day Yellowstone National Park. Molnar and Lavigne (1979) also cite it without specific locality from Wyoming, based on previous records given by Needham and Heywood (1929), Needham and Westfall (1955), and Pritchard and Smith (1956). All of these records probably originated from the original Hagen (1874) citation. In Utah, the species apparently penetrates only the southwesternmost part of the state. The most southerly distribution for $E$. compositus is also unknown. Except for E. coluber, the only records I have observed from mainland Mexico are two males collected by J. H. Williamson in Hermosillo, Sonora. The distribution of E. composit$u s$ in Texas, New Mexico, and Arizona indicates that it must be further widespread in northern Mexico.

## Material

Type data. - Of E. compositus: Holotype female with following label data: 'Aug/ 16' [handwritten], 'Hagen' printed [two of these labels attached], 'compositus' written in an unknown hand; red label 'Holotype [printed] / Erpetogomphus compositus / Hagen 1857 [sic, should be 1858] 'O?' handwritten by RWG. In mczc. Of $E$. coluber. Holotype male dry in envelope with following data: ' $E$. coluber' written in pencil by E. B. Williamson, 'Mexico/ State of Baja California/ San Jose de Comandu/ J. H. Williamson/ Oct 10 1923' [all stamped]/ o' 214.' handwritten by EBW, 'Type' handwritten in pencil by EBW on side of label. In ummz.

Type locality of E. compositus herein restricted to vicinity of Roswell, Chaves Co., New Mexico (see remarks under $E$. designatus).
Other material ( $151 \mathrm{o}^{\pi}, 46$ 오, including holorype 우 $E$. compositus and holotype of E. coluber). - U. S. A.: Arizona: Cochise Co., Miller Cyn., Huachuca Mins., (cc); pond ar Slaughter Ranch, San Bernardino Valley, 15 mi E of Douglas, 1169 m , (RWG); 5 mi E of Hereford (cc); San Pedro River, 9 mi SW of Tombstone, (Byuc); San Pedro River, at Hereford Rd., (Rwg); Coconino Co., Havasu Canyon, 3 mi N of Supai, 923 m , (rwg); Supai, (cdea); Grabam Co., 3 mi SE of Bylas, valley of the Gila, 2000 ft ., (UMmz); Roper Lake, 5 mi S of Safford (rwg); Maricopa

Co., Granite Reef Dam, (ummz); Mesa, (ummz); Phoenix, (RwG); slough ponds by Verde River, by Ariz. Hwy 87, Ft. McDowell Indian Reservation (rWG); Tempe (UMMZ); Mohave Co., spring. Hwy 15 bridge, Littlefield, (bXUC, RWG); Virgin River, Big Bend, (bXUC); Pima Co., Organ Pipe Cacrus Nat'l. Mon., Quitobaquito, (Long Beach State Univ.); Quitobaquito Springs, 15 mi S of Lukeville, Organ Pipe Cactus Nat'l. Mon., (rwg); Pinal Co., Boyce Thompson Southwestern Arboretum, $41 / 2 \mathrm{mi}$ SW of Superior, (Ummz); 8 mi NW of Florence, (ummz); Santa Cruz Co., Santa Cruz River, Sonoita Creek Ranch (on way ro Salero Cyn.), ca. 2 mi W of Patagonia, by Ariz. Hwy 82, (RWG); Yavapai Co., Oak Creek at Cornville, 1077 m , (csuc); Yuma Co. Ave 3 E at Co. 14 St., SE of Yuma, (RWG); Riverside Park, nr. Colorado River, Yuma, (rwg); N. R. Adair Park, McPhaul Bridge, by Gila River, ca. 15 mi ENE of Yuma, (rwg); California: Butte Co., Bidwell City Park, by Big Chico Creek, Chico, 61 m , (RWg); Oroville, (UMMz); Fresno Co., Friant, San Joaquin River, (ummz); Imperial Co. Calexico, (byuc); canal 8 mi E of Holtville, sea level, (DRP); ditch 4.7 mi E of Bond's Corner, (Dre); Hot Mineral, (LACM); irrigation canal 7.3 mi E of Holtville, (DRP); irrigation canal at Winterhaven, (Drp); Inyo Co., Hunter Cyn., Salino Valley, (LACM); Laws, Owens River, (Ummz); Lone Pine, Owens River, (ummz); Los Angeles Co., Tanbark Flat, (CDFA); Mono Co., 5 mi N of Benton Station, 5300 ft ., (Ummz, Lacm); Riverside Co. Blythe, (cc); Coachella Valley Preserve, NE of Thousand Palms, (J. Cole); Palm Springs, (lacm); Whitewater River nr. Salton Sea, (drp); San Bernardino Co., Colorado River at Moabi Rd., S of Needles, (rwg); Lost Palm Cyn., Joshua Tree N. Mon., (Long Beach State Univ.); Parker Dam, (rwg); San Diego Co.: T14S, R5E, (CASC); San Felipe Creek, 13.8 mi E of Julian, 615 m , (DRP); San Felipe Creek at Scissors Crossing, 11.9 mi E of Julian, 738 m, (DRP, RWG); Sentenac Cyn., (LACM); Vallecito, (Lacm); Yolo Co., Cache Creek at William H. 'Bill' Davis Memorial Picnic Area, by Calif. Hwy 16, 6 mi N of Rumsey, 132 m, (RWG); Nevada: Clark Co., Logandale, (byuc); Elko Co., Carlin, Humboldt River, (ummz); 8.5 mi NW of Currie, (Ummz); Humboldt Co., Can Spring, 3 mi S of Pahuhe Meadows, (UMMZ); Lincoln Co., 6 mi S of Alamo, (ummz); Nye Co., Amargosa R., 2 mi below Beatty, (ummz); Pershing Co., Lovelock, Humboldt River, (ummz); Washoe Co., Pyramid Lake, Truckee River, (ummz); White Pine Co., 2 mi NW of Preston, (Ummz); Texas: Brewster Co., Big Bend Nat'l Park, 1 mi N of Rio Grande Village, (RWG); Boquillas Cyn., Big Bend Nat'l. Park, (csuc); Rio Grande Village, Big Bend Nar'l. Park, (csuc); Maverick Co., Quemado, (Lacm); Utah: Washington Co., St. George, (cc); Beaver Dam Wash, Terry Ranch, (Byuc); Beaver Dam Wash, Lytle Ranch, (byUc); Gunlock, W of Veyo, (BXUC); Virgin River, Virgin, (byuc); Mexico: Baja Calif. Norte. Guadalupe Hot Spgs., Guadalupe Cyn., Sierra Juarez, (csuc); Baja Calif. Sur. San José de Comandu, 10 Oct. 1923 (J. H. Williamson), $150^{\circ}$ (all pararypes of E. coluber) (UMMZ, RWG, FSCA, USNM); Sonora: Hermosillo, (UMmZ).

## Erpetogomphus boa Selys

(figs. 26-thx, 44, 45-abd, 68-hamules, 90 -penis, 114, 115 -app, 135-ept, 164-vertex, 167 postocciput, 195, 196-vl, 207-distr)
Erperogomphus boa Selys, 1859: 37 (11, sep.) (descr. ơ' 'Vera Cruz, Mexique. Par M. Salle. (Collect. Selys.)'). - Walsh 1863:253 (mentioned); Selys 1873b: 519 (75, sep.) (list);

Calvert 1905: 165 (notes); Calvert 1907: 399 (possible identity with E. elaps); Muttkowski 1910: 86 (cat.); Ris 1917: 153 (notes and descr. of 2nd known $\delta^{\top}$ ); Williamson and Williamson 1930: 11 (summary of status); Paulson 1982: 255 (Mex.); Davies and Tobin 1985: 27 (cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 31 (cat.); Tsuda 1991: 95 (cat.).
Erpetogomphus crotalinus, nec Hagen in Selys, 1854. - Selys 1859: 537 (11, sep.) ( $\ddagger$ descr. 'Vera Cruz, Mexique. Par M. Salle. (Collect. Selys.)' = E. crotalinus Hagen in Selys, 1858.

Gomphus boa. - Hagen 1861: 100 (descr. from Selys).
Herpetogomphus boa. - Walsh 1862: 389 (mentioned); Hagen 1875a: 42 (cat.); Selys 1879: 64 (2, sep.); Calvert 1899: 386 (list, English translation of Selys, 1879); Calvert 1908b: 693 (mentioned).

## Description

Male. - Labrum, clypeus, and frons light blue green, paler along sides of labrum and lateral lobes of postclypeus; base of mandibles pale green, tips black; base of antefrons anterior to median ocellus with narrow, arcuate, V-shaped sulcus, shallower medially; vertex dark brown with usual postocellar tubercles, occiput tumid medially, pale green brown, its posterior margin ciliated, transverse to slightly concave, slightly emarginate in the middle, or more rarely smoothly curved, postocciput green, tumid; rear of head red brown.

Prothorax red brown, synthorax (fig. 26) predominantly green, area around middorsal carina washed with brown, often appearing as a faded middorsal stripe; antehumeral and humeral stripes dark brown, united into one large stripe; most of this stripe occupying posterior 0.30 of mesepisternum, metastigma black, a wash of brown between metepisternum and metepimeron often forming an obscure, narrow thoracic stripe, subalar carina darkened with brown, mesinfraepisternum and metinfraepisternum brown; metasternum green-brown. Coxae pale grey- brown with some green on exterior surfaces, trochanters grey brown, femora tawny at base, becoming dark brown distally, tibiae and tarsi black.

Venational statistics ( $n=13$, including lectotype $\delta^{\top}$ ). Fifth antenodal (occasionally fourth, sixth, or seventh) thickened in all wings; marginal cells behind fore wing paranal cells: $0-4 / 0-4$; anx: fore wing $13-$ 15/13-16, hind wing 9-11/10-12; pnx: fore wing 9-12/9-11, hind wing $10-13 / 10-13$; cs under pterostigma: fore wing 5-7/5-7, hind wing 5-7/5-7; anal triangular cells: 4-5. Hind wing $33-34 \mathrm{~mm}$.

Abdomen (fig. 44). Segment 1 brown dorsally, becoming green laterally; segment 2 with blue green middorsal stripe, with dorsolateral stripe of brown surrounding green auricle, lateral margin neat hamules pale greenish white; segment 3 with green middorsal stripe bounded on sides by dorsolateral stripes of brown, lateral areas of tergites pale green becoming white posteriorly, anterior transverse carina edged
with black expanding laterally interrupting white lateral areas of segment, thus forming two pale spots, posterior transverse carina also black, expanding laterally so that posterior 0.2 of segment is black; segments 4-6 similar to segment 3 , but with middorsal brown stripe narrowed anteriorly so that whitish green side almost touches middorsal green; segment 7 with anterior 0.5 dull white, conspicuously traversed by black anterior transverse carina, posterior 0.5 of segment red brown; segments 8-10 red brown with ventral and posterior borders of each segment black. Appendages red brown with apices becoming black. Abdomen $37-$ 40 mm .

Cercus (fig. 115) nearly straight, swollen anteriorly, apical 0.5 abruptly concave dorsally and covered with thick series of strong black bristles. Epiprocts about 0.5 the length of superiors, strongly curved dorsally; tips slightly divergent, each a thick, blunt point (fig. 135).

Accessory genitalia (fig. 68). Anterior hamule black, glabrous, deeply forked with posterior branch slightly longer than anterior, the whole structure resembling a talon; anterior branch of hamule with a small semi-oval planar surface on outer side, this area covered with small hairs. Posterior hamule pale green or white, becoming brown near tip; in lateral view, tip obtusely pointed with posterior margin gently convex. Penis (fig. 90) with lateral lobes poorly developed posteriorly, almost circular, its margin serrated.

Female. - Overall colouration as in male, but with pale areas on abdomen and legs more extensive. On abdomen (fig. 45), brown on terminal segments extending anteriorly so that green middorsal and dorsolateral brown stripes become ill-defined. Lateral margins of abdominal segments mostly white and similar to male except that white extends posteriorly to segment 8 . abdomen $37-40 \mathrm{~mm}$.

Vertex with slightly tumid, oval areas posterior to lateral ocelli; posterior margin of occiput (fig. 164) elevated and nearly straight; postocciput green, tumid medially (fig. 167).

Venational statistics. Fifth antenodal (occasionally fourth, sixth, or seventh) thickened in all wings; anx: fore wing 13-16/13-16, hind wing 10-12/10-11; pnx: fore wing $9-13 / 9-12$, hind wing $10-13 / 10-14$; cs under pt: fore wing 5-7/5-8, hind wing 5-8/5-8. Hind wing $34-37 \mathrm{~mm}$.

Vulvar lamina (fig. 195) membranous, diagonally corrugated, outer margin of plate with strong costate ridge, area immediately mesal to ridge furrow-like, with a smaller tumid area occupying the center, medial area tumid, posterior margin of each plate a thin, membranous ridge, postlamellar ridge long, greatly surpassing hind margin of lamina, its $Y$-shaped juncture at distal 0.30 of sternum. Abdomen $37-40 \mathrm{~mm}$.

## Diagnosis

Erpetogomphus boa is most similar to E. cophias, but is easily distinguished by the thoracic pattern and the male caudal appendages. A well-defined combined antehumeral and humeral stripe is present in E. boa, but no such stripe is present in E. cophias. The superior appendages of $E$. cophias possess a large ventral ridge which ends in a prominent tooth 0.30 the length of the appendage (fig. 116). This structure is lacking in $E$. boa (fig. 115). Dorsally, the apical 0.30 of the superior appendages of $E$. cophias contains a few strong bristles, not as many or as stout as in $E$. boa.

The females of these two species are similar morphologically, but may be distinguished by the thoracic pattern. The postocciput of $E$. boa (fig. 164) does not have the posterior depressions on each side found in E. cophias (fig. 165).

Females of $E$. boa superficially resemble females of E. viperinus, but the latter have a pair of circular depressions on abdominal sternite 9 posterior to the juncture of the postlamellar ridge which are absent in $E$. boa. The vulvar lamina of $E$. viperinus (fig. 184) is totally unlike that of $E$. boa (figs. 195-196).

## Remarks

This species has been known from only two specimens: Selys' type and an incomplete male described by Ris (1917). The lectotype is a teneral male with the apical 0.5 of the cerci missing (fig. 114), but it was apparently complete at the time of the original description. The original description of the male is as follows (translated from the French): 'Pterostigma light yellow brown. Head and thorax yellow, femora yellow, with an external brown stripe, short, the anterior four tarsi brownish black.
'[male]. Superior appendages swollen at their base, with an obtuse tooth above at the end of the swelling; their points rounded, slightly bent inwards, hairy. Inferior appendage divided, branching a little distance, attaining $1 / 2$ length of the superior. Occiput nearly straight. Tibiae brown.'

Selys' (1859) description of the female led Calvert (1905) to believe that it was really a specimen of $E$. crotalinus, and my examination of this female confirms Calvert's assertion. Erpetogomphus crotalinus appears to be restricted to the Pacific side of Mexico, and I doubt that the locality ('Vera Cruz') is correct for this specimen. In the supplement to the Biologia (Calvert, 1907), Calvert received drawings of the incomplete abdominal appendages of the type which led him to remark that $E$. boa seemed hardly different from E. elaps. Ris (1917) provided a history of the type male of $E$. boa and remarked (translated from the German): 'I examined a long time ago in the Hamburg Museum an old male that was different
from elaps and corresponds with the description of boa，that its identity［with boa］seemed likely．The lo－ cality is，unfortunately，uncertain，the label reads ＇Agua Caliente＇withour further information．Its pres－ ervation is fair，the tip of the inferior appendages are missing about the distal fourth as shown in the fig－ ure．．．

Examination of the hamules of the lectotype veri－ fies the identity of a series of $E$ ．boa all collected in southern Veracruz；and the Agua Caliente specimen of Ris is also $E$ ．boa．

Calvert（1905）postulated that $E$ ．sipedon might be the female of $E$ ．boa．Although I have not seen a pair of $E$ ．boa in copula or in tandem，I am confident in as－ cribing these females to $E$ ．boa．The similarity of the vulvar laminae in E．boa and E．cophias（whose males are closely related）further strengthens my conviction that $E$ ．sipedon does not represent the female of $E$ ． boa．López（in litt．）has observed tandem pairs，con－ firming the female sex of this restricted species．

The male epiprocts，which are robust and form a sharp point，are apparently responsible for structural damage to the vertex of many females during mating． Of the 26 females I examined， 14 （54\％）had two holes medially to the postoccipital tubercles．Some of these specimens had encrusted haemolymph sur－ rounding the wounds．Dunkle $(1984,1991)$ reviews cases of traumatic mating among other anisopteran Odonata，but their negative effect，if any，is un－ known．Of the remaining 12 females，which were un－ injured， 5 were teneral and had not yet mated．

Venational details of lectotype male：fifth anteno－ dal thickened in fore wings，fourth in hind wings； marginal cells behind fore wing paranal cells： 1 ？／2； anx：fore wing $14 / 14$ ，hind wing $10 / 10$ ；pnx：fore wing $10 / 10$ ，hind wing $10 / 11$ ；cs under pterostigma： fore wing 5／6，hind wing 5／5；anal triangular cells： 4. Hind wing 34 mm ．

Biology．－Most specimens were obtained at a city park near Jalapa．Individuals were found by flushing them from low vegetation about 50 m from a small stream．Adults spent most of their time resting and were difficult to detect among the green foliage．Its habirs are similar to those reported for E．viperinus． Adults prefer parts of small，low volume sandy－bot－ tomed streams where banks are often covered by trees．Reproductive maturation continues through July．Copulation apparently occurs primarily in open areas near streams．Ensuing pairs fly some distance from these sites．Some males settle at the edge of the stream，but passing females are unreceptive there． Females，like other members of this genus，oviposit unattended，flying over the water，stopping briefly to drop eggs from a height of about 40 cm ．López has observed E．boa captured by spiders（Araneidae）and robber flies（Asilidae）．

Collection dates are July and August．Altitudes of capture range from 1150 m （Orizaba）to 1300 m （Jalapa）．
Distribution（fig．207）．－Erpetogomphus boa has been taken only in central Veracruz between $18^{\circ} 50^{\prime} \mathrm{N}$ and $19^{\circ} 30^{\prime} \mathrm{N}$ along the eastern foothills of the Sierra Madre Oriental．

## Material

Type data．－Lectotype male by present designa－ tion with following data：small green label＇Vera Cruz／Salle＇，yellow rectangular label：＇Erpet．boa／ठ＇ ［this last label probably added by Selys during or after 1879］，all in Selys＇hand；two pencilled labels on red tags：＇No． 100 ＇；and two pencilled white labels：＇ 28 ＇， all in an unknown hand．The caudal appendages and accessory genitalia are shown in fig．114．The female specimen originally described as the female of $E$ ．boa is actually $E$ ．crotalinus and possesses the following la－ bels：small green label：＇Vera Cruz／Salle＇；and white label：＇Herp．boa S．／ 9 ＇，both in Selys＇hand．Both specimens in IRSN．

Other material（ $130^{\star}$ ，including lectotype ó； 26 ）．－Mexico：Veracruz：Parque Javier Clavijero， Jalapa， $1300 \mathrm{~m}, 21$ June 1980 （Raul López）， 1 § $^{*}$ （RWG）；（same data）， 11 Aug． 1980 （A．Garcés）， 10 ڤ （UNAM）；（same data）， 13 Aug． 1980 （R．López）， 10 º （UnAM）；（same data）， 16 Aug． 1980 （A．Garcés）， 10 ڤ （UnAM）；（same data）， 20 Aug．1980， 1 if（unam）； （same data）， 5 Aug．1981，（R．López）， 2 む̊， 6 ㅇ（RWG）； （same data）， 6 Aug．1981， 10 （UnAM）；（same data）， 1 Aug．1982， 3 む̀， 1 ㅇ（RWG）；（same data）， 9 Aug．1982， 1 if（UNAM）；（same data）， 19 Aug．1982， 3 여（UNAM）； （same data），Bosque Mesofilo de Montaña， 12 June 1981 （R．López）， 1 우（Unam）；Barranca de Cayoapa， Teocelo，（no collector）， 1 \＆（ RWG ）； 4.8 mi N of Coscomatopec，（M．A．Ortiz，O．S．Flint）， 10 ， 2 아 usnm）； 2.5 mi S of Huatusco， 23 July 1966 （M．A． Ortiz，O．S．Flint）， 1 if（USNM）； 4.7 mi N of Huatusco，Puente Ruiz Cortines， 31 July 1966 （M． A．Ortiz，O．S．Flint）， 3 ㅇ（UsNM）；Orizaba，Ojo de Agua， 3800 ft ．（D．R．Paulson）， 10 ， 5 ㅇ（DRP，FSCA）．

## Erpetogomphus cophias Selys

（figs．27－thx，46－abd，69－hamules，91－penis，116－
app，165－vertex，197－vl，207－distr）
Erpetogomphus cophias Selys，1858： 332 （72 sep．）（descr．$\delta^{\top}$ ＇Le Mexique，d＇après un mâle du Museum de Paris＇）．－ Selys 1859： 537 （ 11 sep．）（descr．© ）；Selys 1869： 175 （12 sep．）（descr．우）；Selys 1873b： 519 （ 75 sep．）（list）；Calvert 1899： 386 （mentioned）；Calvert 1905： 164 （descr．む， ㅇ）；Calvert 1907： 398 （mention of new figure）；Calvert 1909： 481 （seasonal distr．）；Muttkowski 1910： 86 （cat．）； Williamson and Williamson 1930： 12 （summary of stat－ us）；Montgomery 1973： 239 （comment on name）； Paulson 1982： 255 （Mex．）；Davies and Tobin 1985： 27
(cat.); Tsuda 1986: 87 (cat.); Bridges 1991: VII. 52 (cat.); Tsuda 1991: 95 (cat.).
Gomphus cophias. - Hagen 1861: 100 (descr. ઠ) .
Herpetogomphus cophias. - Walsh 1862: 389 (mentioned); Hagen 1875a: 42 (cat.); Selys 1879: 64 (2 sep.) (characters of genus); Kirby 1890: 60 (cat).

## Description

Neotype male. - Entire face pale green; base of mandibles pale green, tips black; antefrons pale green with wash of brown at base; vertex brown, with some green on tubercle behind each lateral ocellus; two prominent cone-shaped pits anterolateral to median ocellus; scape and pedicel dark brown, flagella missing; occiput broad, its dorsal surface evenly convex, especially medially, its posterior margin slightly arcuate; crest with long brown hairs; postocciput light green, evenly concave, rear of head red brown.

Prothorax primarily brown, with light green medially on anterior, median, and posterior lobes; synthorax entirely pale green with wash of brown ventrally above coxae. Coxae and trochanters pale grey green, femora pale grey green becoming tawny distally, a dark brown streak on lateroextensor surfaces of femora occupying almost all of profemora, about 0.60 of mesofemora, and distal 0.30 of metafemora; tibiae, tarsi, and armature black. Wings hyaline, venation dark brown basally, black distally; anterior margin of costa pale yellow to proximal end of pterostigma; pterostigma brown, veins bordering it black.

Venational details. Fifth antenodal thickened in all wings; no marginal cells behind fore wing paranal cells; anx: fore wing $13 / 13$, hind wing $9 / 9$; pnx: fore wing $7 / 7$, hind wing $7 / 9$; cs under pterostigma: fore wing $5 / 5$, hind wing $5 / 4$; anal triangular cells: 4 Hind wing 29 mm .

Abdomen predominantly pale with following dark brown markings: vestige of midlateral stripe on segment 1 ; this stripe more defined on segment 2 , especially behind auricle; annulus; midlateral stripe on segment 3, transverse carina and darker annulus; segments 4-6 similar to segment 3 but stripes darker, each beginning at a little beyond beginning of segment, enlarging at transverse carina, narrowing and then widening, especially posteriorly, but not touching dorsally; segment 7 with transverse carina and posterior 0.40 of segment laterally; segment 8 pale yellow brown with ill-defined dorsolateral dark stripe; segment 9 similar to segment 8 , but with a longitudinal lateral middorsal spot on posterior 0.75 of segment; segment 10 pale ocher with dark red brown along anterior 0.20 of segment. Pale areas of segments 1-7 primarily pale olive dorsally and ventrally; pale areas of segments $8-10$ red brown dorsally and ventrally. Abdomen 35 mm .

Cercus yellow brown, linear, slightly concave dorsally beyond basal 0.30 , tip smoothly rounded, ventral
margin with inferior carina at basal 0.30 , terminating in a distinct ventral tooth; dorsal concave area of cerci with scattered thick bristles. Epiprocts yellow brown, distal 0.50 curved dorsally at $90^{\circ}$, as is typical of genus, tips slightly divergent, each with a thick, blunt point.

Accessory genitalia. Anterior hamule black, glabrous, deeply forked, with posterior branch slightly longer than anterior, the whole structure resembling a talon; distal 0.20 of anterior branch with a longitudinal, obtuse V-shaped area. Posterior hamule spatulate, pale green, becoming brown near tip; in lateral view, tip obtusely pointed with a small, blunt cephalad directed tooth on rear margin. Peduncle of penis with prominent rounded foliate erect lateral lobes, its posterior margin not prominent, forming a gentle bilobed area; penis with lateral lobes poorly developed posteriorly, forming a serrated semicircle; membranous hood not overlapping; cornuae well developed, parallel sided, their tips evenly rounded.

Female. - Overall colouration as in male, head with prominent conical pits anterolateral to median ocellus as in male; vertex with tumid oval area posterior to lateral ocelli; occiput narrow, crest prominent, straight to slightly sinuate along medial 0.30 , lateral arms bent posteriorly; prominent transverse postoccipital pit (fig. 165) immediately behind lateral arms of crest, medial area of postocciput tumid.

Pro- and synthorax as in male, femora with brown areas reduced with little brown on extensor surface of metafemora.

Venational statistics ( $\mathrm{n}=2$ ): number of marginal cells behind fore wing paranal cells: 2-4/1-3; anx: fore wing $13 / 13$, hind wing $9-11 / 9-10$; pnx: fore wing $9 / 8-$ 9 , hind wing 9-10/10; cs under pterostigma: fore wing $5 / 4-5$, hind wing 4-6/4-5. Hind wing 32-34 mm.

Abdomen with brown dorsolateral stripe more reduced than male, more prominent and expanded around lateral carinae; segments 8-9 tawny, cercus pale ocher. Vulvar lamina (fig. 197) membranous, diagonally corrugated; posterior margin of each plate gently curved, a prominent ridge along its border, a prominent long diagonal depression immediately anterior to posterior margin; plates meeting medially at tumid area; cleft between plates a small notch; juncture of Y-shaped postlamellar ridge well posterior to vulvar laminar plates; area on each side of central stem of postlamellar ridge darkened, with a slight depression. Abdomen $35-37 \mathrm{~mm}$.

## Diagnosis

Erpetogomphus cophias is most similar to E. boa and is diagnosed under that species. The characteristic ventral tooth of the cercus in the male and the postoccipital depressions in the female are autapomorphic characters.

## Remarks

This is apparently a rare species. Williamson and Williamson (1930) stated that there were only 10 known specimens. Only one other specimen from Michoacan, collected in 1941, has come to my attention. I have examined only 3 males and 4 females, of which one male and one female are teneral and in poor condition. I can see no noticeable differences among the few specimens examined.

Variation. - Venational details of one additional male: number of marginal cells behind fore wing paranal cells: $3 / 2$; anx: fore wing $11 / 12$, hind wing $9 / 8$; pnx: fore wing $9 / 8$, hind wing $9 / 10$; cs under pterostigma: fore wing $5 / 5$, hind wing $5 / 5$; number of anal triangular cells $3 / 3$. Hind wing 30 mm .

Biology. - Nothing is known of the biology of this species.

Distribution (fig. 207). - Like E. boa, E. cophias has a restricted distribution and apparently replaces E. boa in the highlands of west-central Mexico south of $20^{\circ} \mathrm{N}$ and west of $100^{\circ} \mathrm{W}$. Records indicate an elevational gradient of 1525 m (Cuernavaca) to 2438 m (Omilteme) and flight during June and July.

## Material

Type data. According to Dr. J. Legrand (in litt. 11 May 1984), the holotype male from 'Le Mexique, d'après un mâle du Museum de Paris', is missing, there remaining only a large rectangular green label with [handwritten] 'E. cophias, Selys' which was originally attached to the specimen. Dr. Legrand was kind enough to send a male of which he states, '...we have from ex Martin's collection a specimen called $O$. cophias Selys (determined by Selys himself, according to Martin...),' which he suggested I designate as neotype. This I now do. The neotype male contains the following data: white label in R. Martin's hand: 'Ophiogomphus / cophias Selys/ Det. De Selys/ Mexique', small green label printed: 'museum paris / Coll. R. martin 1920', and red rectangular label handwritten: 'Erpetogomphus / cophias Selys ō/ neoTYPE / des./ R. W. Garrison 1986'. The abdomen is detached and is in a triangular envelope pinned beneath the specimen.

Other material ( $30^{*}, 4 \%$, including neotype $\delta^{*}$ ). Mexico: Guerrero: Omilteme, 8000 ft ., July 1888 (H. H. Smith), 1 ᄋ, (bmnh); Michoacan: Tancítaro, $6000 \mathrm{ft} . \quad(1846 \mathrm{~m})$, 4th Hoogstraal Mexican Biological Expedition, 28 July 1941 (H. Hoogstraal), $10^{\circ}$ (UMmz); Morelos: Cuernavaca, June 1897 (O. W. Barrett), 10, 1 ㅇ (ANSP); June 1888 (H. H. Smith), $1 \delta^{\circ}$ (вмnh); 8 July 1900 (C.C. Deam), 2 ㅇ (Ummz).

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## References

Ackery, P. R. \& R. I. Vane-Wright, 1984. Milkweed butterflies their cladistics and biology being an account of the natural history of the Danainae, a subfamily of the Lepidoptera, Nymphalidae. - Cornell University Press, Ithaca: ix +425 .
Adams, C. C., 1900. Odonata from Arkansas. Entomological News 11: 621-622.
Ahrens, C., 1938. A list of dragonflies taken during the summer of 1936 in western United States (Odonata). Entomological News 49: 9-16.
Alrutz, R. W., 1961. Notes and records of Ohio dragonflies and damselflies (Odonata). - Ohio Journal of Science 61 (1): 13-24.

Ander, K., 1929. Über die Nymphe von Mesogomphus Hageni Selys (Odonata). - Konowia 8 (2): 159-162.
Banks, N., 1892. A synopsis, catalogue, and bibliography of the neuropteroid insects of temperate North America. Transactions of the American Entomological Society 19: 327-373.
Belle, J., 1972. An unknown gomphid larva from Surinam, possibly Progomphus geijskesi Needham, 1944 (Odonata). - Odonatologica 1 (2): 113-116.

Belle, J., 1988. A synopsis of the species of Phyllocycla Calvert, with descriptions of four new taxa and a key to the genera of neotropical Gomphidae (Odonata, Gomphidae). - Tijdschrift voor Entomologie 131: 73102.

Belle, J., 1992. Studies on ultimate instar larvae of neotropical Gomphidae, with the description of Tibiagomphus gen. nov. (Anisoptera). - Odonatologica 21 (1): 1-24.
Belle, J. \& D. Quintero, 1992. Chapter 6. Clubtail dragonflies of Panama (Odonata: Anisoptera: Gomphidae), pp. 91-101. - In D. Quintero \& A. Aiello, ed. Insects of

Panama and Mesoamerica: selected studies. Oxford University Press: xxii +692 pp.
Bick, G. H., 1951. Notes on Oklahoma dragonflies. Journal of the Tennessee Academy of Science 26: 178180.

Bick, G. H., 1957. The Odonata of Louisiana. - Tulane Studies in Zoology 5 (5): 71-135.
Bick, G. H., 1959. Additional dragonflies (Odonata) from Arkansas. - The Southwestern Naturalist 4 (3): 131-133.
Bick, G. H., 1990. Unpublished records in Florida Stare Collection of Arthropods (fSCA). - Argia, the News Journal of the Dragonfly Society of America 2 (1-4): 3-4.
Bick, G. H. \& J. C. Bick, 1957. The Odonata of Oklahoma. - The Southwestern Naturalist 2 (1): 1-18.

Bick, G. H. \& J. C. Bick, 1958. The ecology of the Odonata at a small creek in southern Oklahoma. - Journal of the Tennessee Academy of Science 33 (3): 240-251.
Bick, G. H., J. C. Bick and L. E. Hornuff, 1977. An annotated list of the Odonata of the Dakotas. - Florida Entomologist 60 (3): 149-166.
Bird, R. D., 1932. Dragonflies of Oklahoma. - Publications of the University of Oklahoma Biological Survey 4 (1-2): 50-57.
Borror, D. J., 1935. New records of Ohio dragonflies (Odonata). - Ohio Journal of Science 35 (6): 451-456.
Borror, D. J., 1937. An annotated list of the dragonflies (Odonata) of Ohio. - Ohio Journal of Science 37 (3): 185-196.
Borror, D. J., 1963. Common names for Odonata. Proceedings North Central Branch, Entomological Society of America 18: 104-107.
Brauer, F., 1868. Verzeichniss der bis jetzt bekannten Neuropteren im Sinne Linne's. - Verhandlungen der zoologisch-botanischen Gesellschaft in Wien 18: 359416, 711-742.
Bridges, C. A., 1991. Catalogue of the family-group, genusgroup and species-group names of the Odonata of the world. - Available from the author, 502 W. Main St., Urbana, Illinois, U. S. A., 61801 : xiv + 704 pp.
Byers, C. F., 1928. The unknown nymphs of North American Odonata. - Canadian Entomologist 60: 4-6.
Byers, C. F., 1930. A contribution to the knowledge of Florida Odonata. - University of Florida Publication. Biological Science Series 1 (1): 1-137.
Byers, C. F., 1931. Dixie dragonflies collected during the summer of 1930 (Odonata). - Entomological News 42: 113-119.
Byers, C. F., 1939. A study of the dragonflies of the genus Progomphus (Gomphoides) with a description of a new species. - Proceedings of the Florida Academy of Sciences 4: 19-85.
Cabot, L., 1872. The immature state of the Odonata. Part I. - Subfamily Gomphina. - Illustrated Catalog of the Museum of Comparative Zoology, at Harvard College 5: 1-17.
Calvert, A. S. \& P. P. Calvert, 1917. A year of Costa Rican natural history. - Macmillan Co., New York: xix + 577 pp.
Calvert, P. P., 1895. The Odonata of Baja California, Mexico. - Proceedings of the California Academy of Sciences (2) 4: 463-558.
Calvert, P. P., 1899. Odonata from Tepic, Mexico, with supplementary notes on those of Baja, California. Proceedings of the California Academy of Sciences (3) 1: 371-418.
Calvert, P. P., 1905. Odonata, in Biologia Centrali-

Americana: Insecta Neuroptera. - R. H. Porter and Dulau Co., London: pp. 145-212.
Calvert, P. P., 1907. Odonata, in Biologia Centrali Americana: Insecta Neuroptera, - R. H. Porter and Dulau Co., London: pp. 309-404.
Calvert, P. P., 1908a. List of Odonata taken by Dr. Henry Skinner in Carr Canyon, Huachuca Mountains, Arizona. - Entomological News 19: 45.

Calvert, P. P., 1908b. The present state of our knowledge of the Odonata of Mexico and Central America. - Science 28 (724): 692-695.
Calvert. P. P., 1908c. Introduction to the Odonata, in Biologia Centrali-Americana: Insecta Neuroptera. - R. H. Porter and Dulau Co., London: pp. i-xxx.

Calvert, P. P., 1909. The composition and ecological relations of the odonate fauna of Mexico and Central America. - Proceedings of the Academy of Natural Sciences of Philadelphia 60: 460-491.
Calvert, P. P., 1912a. Studies on Costa Rican Odonata. IV. Erpetogomphus in Costa Rica, with descriptions of a new species having complex structural mating adaptations. Entomological News 23 (7): 289-295.
Calvert, P. P., 1912b. [Note on Erpetogomphus tristani.]. Entomological News 23: 384.
Calvert, P. P., 1919. Odonata Anisoptera from Guatemala. - Entomological News 30: 72-78.

Calvert, P. P., 1920a. [Note on Guatemalan Odonata]. Entomological News 31: 113.
Calvert, P. P., 1920b. The Costa Rican species of Epigomphus and their mutual adaptations (Odonata). Transactions of the American Entomological Society 46: 323-354.
Calvert, P. P., 1942. Increase in knowledge of the odonate fauna of Mexico, Central America, and the West Indies since 1908. - Proceedings: Eighth American Scientific Congress, Biological Sciences: Zoology 3: 323-331.
Calvert, P. P., 1947. The Odonate collections of the California Academy of Sciences from Baja California and Tepic, Mexico, of 1889-1894. - Proceedings of the California Academy of Sciences (4) 23: 603-609.
Cannings, R. \& R. W. Garrison, 1991. Sympetrum signifer$u m$, a new species of dragonfly (Odonata: Libellulidae) from western Mexico and Arizona. - Annals of the Entomological Society of America 84 (5): 474-479.
Carle, F. L., 1979. Environmental monitoring potential of the Odonata, with a list of rare and endangered Anisoptera of Virginia, United States. - Odonatologica 8 (4): 319-323.

Carle, F. L., 1982. Ophiogomphus incurvatus. a new name for Ophiogomphus carolinus Hagen (Odonata: Gomphidae). - Annals of the Entomological Society of America 75 (3): 335-339.
( Carle, F. L., 1986. The classification, phylogeny and biogeography of the Gomphidae (Anisoptera). I. Classification. - Odonatologica 15 (3): 275-326.
Carle, F. L., 1992. Ophiogomphus (Ophionurus) australis spec. nov. from the gulf coast of Louisiana, with larval and adult keys to American Ophiogomphus (Anisoptera: Gomphidae). - Odonatologica 21 (2): 141-152.
Carle, F. L. \& C. Cook, 1984. A new Neogomphus from South America, with extended comments on the phylogeny and biogeography of the Octogomphini trib. nov. (Anisoptera: Gomphidae). - Odonatologica 13 (1): 5570.

Carpenter, G. H., 1897. The geographical distribution of
dragonflies. - Scientific Proceedings of the Royal Dublin Society 8 (N. S.) (5) 55: 439-468.
Chao, H-f., 1984. Reclassification of Chinese gomphid dragonflies, with the establishment of a new subfamily and the descriptions of a new genus and species (Anisoptera: Gomphidae). - Odonatologica 13 (1): 7180.

Cowley, J. C., 1934. Notes on some generic names of Odonata. - Entomologists Monthly Magazine 70: 240247.

Cowley, J. C., 1937. The pagination of the reprints of the Selysian monographs and synopses of Odonata. - Journal of the Society for the Bibliography of Natural History 1 (3): 73-81.

Cross, W. H., 1955. Anisopteran Odonata of the Savannah River Plant, South Carolina. - Journal of the Elisha Mitchell Scientific Society 71 (1): 9-17.
Cross, W. H., 1956. Dragonflies in the Tallahassee region. Florida Entomologist 39 (1): 9-16.
Cruden, R. W., 1964. Notes on Brechmorhoga mendax (Hagen): Odonata. - Entomological News 75 (3): 79-82.
Currie, R. P., 1903. The Odonata collected by Messrs. Schwarz and Barber in Arizona and New Mexico. Proceedings of the Entomological Society of Washington 5 (4): 298-303.
Davies, D. A. L. \& P. Tobin, 1985. The dragonflies of the world: a systematic list of the extant species of Odonata. Vol. 2 Anisoptera. - Societas Internationalis Odonatologica Rapid Communications (Suppl.) 5: xi +151 pp .
De Marmels, J., 1990. An updated checklist of the Odonata of Venezuela. - Odonatologica 19 (4): 333-345.
Donnelly, T. W., 1961. The Odonata of Washington, D. C., and vicinity. - Proceedings of the Entomological Sociery of Washington 63 (1): 1-13.
Donnelly, T. W., 1992. Chapter 5. The Odonata of Central Panama and their position in the neotropical odonate fauna, with a checklist, and descriptions of new species, pp. 52-90. In D. Quintero and A. Aiello, ed. Insects of Panama and Mesoamerica: selected studies. - Oxford University Press: xxii + 692 pp.
Dunkle, S. W., 1984. Head damage due to mating in Ophiogomphus dragonflies (Anisoptera: Gomphidae). Notulae Odonatologicae 2 (4): 63-64.
Dunkle, S. W., 1988. A list of the Odonata of Honduras. Ceiba 29 (1): 41-49.
Dunkle, S. W., 1991. Head damage from mating attempts in dragonflies (Odonata: Anisoptera). - Entomological News 102 (1): 37-41.
Dunkle, S. W., 1992. Distribution of dragonflies and damselflies (Odonata) in Florida. - Bulletin of American Odonatology 1 (2): 29-50.
Dunkle, S. W. \& J. J. Belwood, 1982. Bat predation on Odonata. - Odonatologica 11 (3): 225-229.
Dunkle, S. W. \& M. J. Westfall, Jr., 1982. Order Odonata, pp. 32-45. In R. Franz, ed. Invertebrates, vol. 6. Rare and endangered biota of Florida. - University Presses of Florida: xx +131 pp .
Farris, J. S., 1988. Hennig86 version 1.5 [Phylogenetic program for PC's]. - Port Jefferson, NY.
Ferguson, A., 1940. A preliminary list of the Odonata of Dallas County Texas. - Field and Laboratory. Journal of the Graduate Research Center, Southern Methodist University 8 (1): 1-10.
Ferguson, A., 1942. Scattered records of Texas and Louisiana Odonata with additional notes on the Odonata
of Dallas County. - Field and Laboratory. Journal of the Graduate Research Center, Southern Methodist University 10 (2): 145-149.
Ferguson-Beatty, A., 1956. An inquiry into the significance of the larval proventriculus in the taxonomy of Odonata. - Proceedings Tenth International Congress of Entomology 1: 367-372.
Förster, F., 1914. Beiträge zu den Gattungen und Arten der Libellen. III. - Archiv für Naturgeschichte (A) 80 (2): 5983.

Fraser, F. C., 1940. A comparative study of the penes of the family Gomphidae (Order Odonata). - Transactions of the Royal Entomological Society of London 90 (20): 541-550.
Garman, P., 1927. Guide to the insects of Connecticut. Part V. The Odonata or dragonflies of Connecticut. Bulletin. Connecticut State Geological and Natural History Survey 39: 1-331.
Garrison, R. W., 1986. In the wilds of Baja. - Selysia 15 (1): 15-17.
Gloyd, L. K., 1958. The dragonfly fauna of the Big Bend region of trans-Pecos, Texas. - Occasional Papers of the Museum of Zoology, University of Michigan 593: 1-23.
Gloyd, L. K., 1963. A movable molar in the Odonata Proceedings, North Central Branch Entomological Society of America 18: 147-149.
González-S., E. \& R. Novelo-G., 1990. Dos nuevas especies de Phyllogomphoides Belle 1970 (Odonata: Gomphidae) del estado de Morelos, Mexico. - Folia Entomologica Mexicana 79: 33-43.
González-S., E. \& R. Novelo-G., 1991. Odonata de la Reserva de la Biosfera Michilia, Durango, Mexico. Parte I. Imagos. - Folia Entomologica Mexicana 81: 67-105.

Hagen, H. A.., 1861. A synopsis of the Neuroptera of North America. Smithsonian Miscellaneous Collections, Washington, D. C.: xx +347 pp.
Hagen, H. A., 1873. Odonata from the Yellowstone. In F. V. Hayden. Sixth annual report of the United States Geological Survey of the Territories, embracing portions of Montana, Idaho, Wyoming, and Utah; being a report of progress of the explorations for the year 1872. - Report of the United States Geological Survey of the Territories 6: 727-729.
Hagen, H. A., 1874. Report on the Pseudo-neuroptera collected by Lieut. W. L. Carpenter in 1873 in Colorado. In F. V. Hayden. Annual report of the United States Geological and Geographical Survey of the Territories, embracing Colorado, being a report of progress of the exploration for the year 1873. - Report of the United States Geological Survey of the Territories 7: 571-606.
Hagen, H. A., 1875a. Synopsis of the Odonata of America. - Proceedings of the Boston Society of Natural History 18: 20-96.
Hagen, H. A., 1875b. Chapter 14. Report upon the collections of Neuroptera and Pseudo-neuroptera, made in portions of Colorado, New Mexico, and Arizona during the years 1872, 1873, and 1874. In Volume 5. Zoology. Report upon geographical and geological explorations and surveys west of the one hundredth meridian, in charge of First Lieut. Geo. M. Wheeler, corps of engineers, U.S. Army, under the direction of Brig. Gen. A. A. Humphreys, chief of engineers, U.S. Army, pp. 909-922.
Hagen, H. A., 1885. Monograph of the earlier stages of the Odonata. Sub-families Gomphina and Cordulegastrina. - Transactions of the American Entomological Society

12: 249-291.
Harp, G. L. \& J. D. Rickett, 1977. The dragonflies (Anisoptera) of Arkansas. - Arkansas Academy of Science Proceedings 31: 50-54.
Harp, G. L. \& J. D. Rickett, 1985. Further distributional records for Arkansas Anisoptera. - Arkansas Academy of Science Proceedings 39: 131-135.
Higgins, H. T., 1901. The development and comparative structure of the gizzard in the Odonata Zygoptera. Proceedings of the Academy of Natural Sciences of Philadelphia 126-141.
Hine, J. S., 1913. Additions and corrections to the Odonata of Ohio. - Ohio Naturalist 13 (5): 94-96.
Huggins, D. G., 1978. Additional records of Kansas Odonata. - Technical Publications of the State Biological Survey of Kansas 6: 1-35.
Huggins, D. G. \& W. U. Brigham, 1982. Chap. 4, Odonata, pp. 4.1-4.100. In A. R. Brigham, W. U. Brigham \& A. Gnilka, eds. Aquatic insects and oligochaetes of North and South Carolina. Midwest Aquatic Enterprises, Mahomet, IL: 837 pp.
Huggins, D. G., P. M. Liechti and D. W. Roubik, 1976. Species accounts for certain aquatic macroinvertebrates from Kansas (Odonata, Hemiptera, Coleoptera and Sphaeriidae). In J. Caldwell, ed. New records of the fauna and flora of Kansas for 1975, Technical Publications of the State Biological Survey of Kansas 1: 13-77.
Karsch, F., 1890. Ueber Gomphiden. - Entomologische Nachrichten 16: 370-382.
Kennedy, C. H., 1917a. Notes on the life history and ecology of the dragonflies (Odonata) of central California and Nevada.. - Proceedings of the United States National Museum 52: 483-635.
Kennedy, C. H., 1917b. The dragonflies of Kansas. The Odonata of Kansas with reference to their distribution. Bulletin of the University of Kansas Biological Series 11: 127-143.
Kennedy, C. H., 1918. New species of Odonata from the southwestern United States. Part II. - Canadian Entomologist 50 (9): 297-299.
Kennedy, C. H., 1928. Evolutionary level in relation to geographic, seasonal and diurnal distribution of insects. Ecology 9 (4): 367-379.
Kimmins, D. E., 1969. A list of the type-specimens of Odonata in the British Museum (Natural History) Part II. - Bulletin of the British Museum Natural History 23 (7): 287-314.

Kirby, W. F., 1890. A synonymic catalogue of Neuroptera Odonata, or dragonflies, with an appendix of fossil species. - Gurney and Jackson, London: ix + 202 pp.
Kormondy, E. J., 1957. New knowledge of the Odonata of Tennessee. - Journal of the Tennessee Academy of Science 32 (2): 106-115.
Kormondy, E. J., 1960. New North American records of anisopterous Odonata. - Entomological News 71 (5): 121130.

La Rivers, I., 1938. An annotated list of the Libelluloidea (Odonata) of southern Nevada. - Pomona College Journal of Entomology and Zoology 30: 73-85.
La Rivers, I., 1940a. A preliminary synopsis of the dragonflies of Nevada. - Pan-Pacific Entomologist 16 (3): 111123.

La Rivers, I., 1940b. Some dragonfly notes from northern Nevada. Pomona College Journal of Entomology and Zoology 32: 61-68.

La Rivers, I., 1941. Additions to the list of Nevada dragonflies (Odonata). - Entomological News 52: 126-130, 155-157.
Larsen, W. P., 1952. The dragonflies (Anisoptera) of Utah. - Unpublished Master's Thesis, Deparment of Zoology, University of Utah: 95 pp.
Macklin, J. A. \& C. Cook, 1967. New records of Kentucky Odonata. - Proceedings North Central Branch, Entomological Society of America 22: 120-121.
Maes, J-M., J-M. Desmedt \& V. Hellebuyck, 1988. Catalogo de los Odonata de Nicaragua. Revista nicaraguense entomologica 4: 29-43.
Molnar, D. R. \& R. J. Lavigne, 1979. The Odonata of Wyoming (dragonflies and damselflies). - University of Wyoming Agricultural Experiment Station Scientific Monograph 37: 1-142.
Montgomery, B. E., 1925. Records of Indiana dragon-flies-1. - Proceedings of the Indiana Academy of Sciences 34: 383-389.
Montgomery, B. E., 1927. Records of Indiana dragon-flies-II. - Proceedings of the Indiana Academy of Sciences 36: 287-291.
Montgomery, B. E., 1929. Records of Indiana dragonflies, III. - 1927-1928. Proceedings of the Indiana Academy of Sciences 1929: 335-343.
Montgomery, B. E., 1935. Records of Indiana dragonflies, VIII. 1934. - Proceedings of the Indiana Academy of Sciences 44: 231-235.
Montgomery, B. E., 1937. Records of Indiana dragonflies, IX. 1935-1936. - Proceedings of the Indiana Academy of Sciences 46: 203-210.
Montgomery, B. E., 1940. The Odonata of South Carolina. - Journal of the Elisha Mitchell Scientific Society 56 (2): 283-301.
Montgomery, B. E., 1941. Records of Indiana dragonflies, X. 1937-1940. - Proceedings of the Indiana Academy of Sciences 50: 229-241.
Montgomery, B. E., 1947. The distribution and relative seasonal abundance of Indiana species of five families of dragonflies (Odonata: Calopterygidae, Petaluridae, Cordulegasteridae, Gomphidae and Aeshnidae). Proceedings of the Indiana Academy of Sciences 56: 163169.

Montgomery, B. E., 1951. Notes and records of Indiana Odonata, 1941-1950. - Proceedings of the Indiana Academy of Sciences 60: 205-210.
Montgomery, B. E., 1955. Notes and records of Indiana Odonata, 1953-54. - Proceedings of the Indiana Academy of Sciences 64: 131-135.
Montgomery, B. E., 1967. Geographical distribution of the North Central States. - Proceedings North Central Branch, Entomological Society of America 22: 121-129.
Montgomery, B. E., 1968. The distribution of western Odonata. - Proceedings North Central Branch, Entomological Society of America 23 (2): 126-136.
Montgomery, B. E., 1973. Why snakefeeder? Why dragonfly? Some random observations on etymological entomology. - Proceedings of the Indiana Acaderny of Sciences 82: 235-241.
Musser, R. J., 1961. Some noteworthy dragonfly records from Utah (Odonata: Anisoptera). - Entomological News 72 (2): 53.
Musser, R. J., 1962. Dragonfly nymphs of Utah (Odonata: Anisoptera). - University of Utah Biological Series 12 (6): vii + 74 pp.

Muttkowski, R. A., 1910. Catalogue of the Odonata of North America. - Bulletin of the Public Museum of the City of Milwaukee 1 (1): 1-207.
Needham, J. G., 1897. Preliminary studies of N. American Gomphinae. - Canadian Entomologist 29: 164- 168, 181-186.
Needham, J. G., 1899. Ophiogomphus. - Canadian Entomologist 31 (9): 233-238.
Needham, J. G., 1911. Notes on a few nymphs of Agrioninae (Order Odonata) of the Hagen collection. Entomological News 22: 342-345.
Needham, J. G., 1940. Studies on neorropical Gomphine dragonflies. (Odonata). - Transactions of the American Entomological Society 65: 363-394.
Needham, J. G., 1941. Life history studies on Progomphus and its nearest allies (Odonata: Aeschnidae). Transactions of the American Entomological Society 67: 221-245.
Needham, J. G., 1943. Notes on some Gomphine dragonflies from Venezuela and Guatemala. - Boletin de Entomología venezolana 2 (4): 197-206.
Needham, J. G., 1944. Further studies on neotropical Gomphinae (Odonata). - Transactions of the American Entomological Society 69: 171-224.
Needham, J. G. \& M. H. Anthony, 1903. The skewness of the thorax in Odonata. - Journal of the New York Entomological Society 11: 117-124.
Needham, J. G. \& T. D. A. Cockerell, 1903. Some hitherto unknown nymphs of Odonata from New Mexico. Psyche 10: 134-139.
Needham, J. G. \& C. A. Hart, 1901. The dragonflies (Odonata) of Illinois. Part I. Petaluridae, Aeschnidae, and Gomphidae. - Bulletin of the Illinois State Laboratory of Natural History 6: 1-94.
Needham, J. G. \& H. B. Heywood, 1929. A handbook of the dragonflies of North America. - Charles C. Thomas, Springfield: viii + 372 pp.
Needham, J. G. \& M. J. Westfall, Jr., 1955. A manual of the dragonflies of North America (Anisoptera). - Univ. of Calif. Press, Berkeley: xii +615 pp.
Novelo-G., R. \& E. González-S., 1991. Odonata de la Reserva de la Biosfera La Michiliá, Durango, Mexico. Parte II. Náyades. - Folia Entomologica Mexicana 81: 107-164.
Novelo-G., R. \& J. Peña-O., 1991. Odonata from the northern mountain range of Hidalgo State, Mexico. Notulae Odonatologicae 3 (8): 129-131.
Osburn, R. C., 1905. The Odonata of British Columbia. Entomological News 16: 184-196.
Paulson, D. R., 1982. Odonata, pp. 249-277. In S. H. Hurlbert \& A. Villalobos-Figueroa, eds. Aquatic biota of Mexico, Central America and the West Indies. San Diego State University, San Diego, CA: xv + 529 pp.
Paulson, D. R., 1983. A new species of dragonfly, Gomphus (Gomphurus) lynnae spec. nov., from the Yakima River, Washington, with notes on the pruinosity in Gomphidae (Anisoptera). - Odonatologica 12 (1): 59-70.
Paulson, D. R. \& R. W. Garrison, 1977. A list and new distributional records of Pacific Coast Odonata. - PanPacific Entomologist 53 (2): 147-160.
Pinhey, E., 1969. Tandem linkage in dichoptic and other Anisoptera (Odonata). - Occasional Papers of the National Museums of Rhodesia 4 (28B): 137-207.
Pritchard, A. E. \& R. F. Smith, 1956. Odonata, pp. 106153. In R. L. Usinger, ed. Aquatic insects of California. -

University of California Press, Berkeley: ix + 508 pp.
Resener, P. L., 1970. An annotated check list of the dragonflies and damselflies (Odonata) of Kentucky. Transactions of the Kentucky Academy of Science 31 (12): 32-44.

Ris, F., 1908. In L. Schultze, Forschungsreise im westlichen und zentralen Südafrika, ausgeführt in den Jahren 19031905. Denkschriften der medizinisch-naturwissenschaftlichen Gesellschaft 13: 303-346.
Ris, F., 1909. Abessinische Libellen, gesammelt von Dr. Eduard Rüppell. - Bericht über die Senckenbergische naturforschende Gesellschaft, Frankfurt am Main 40: 2127.

Ris, F., 1917. über drei Arten Erpetogomphus (Odonata). Archiv für Naturgeschichte (A) 82 (3): 152-158.
Ris, F., 1918. Libellen (Odonata) aus der Region der amerikanischen Kordilleren von Costarica bis Catamarca. Archiv für Naturgeschichte (A) 82 (9): 1-197.
Ris, F., 1921. The Odonata or dragonflies of South Africa. - Annals of the South African Museum 18 (3): 245-452.

Roback, S. S. \& M. J. Westfall, Jr., 1967. New records of Odonata nymphs from the United States and Canada with water quality data. - Transactions of the American Entomological Society 93: 101-124.
Scudder, S. \& T. Cockerell, 1902. A first list of the Orthoptera of New Mexico. - Proceedings of the Davenport Academy of Sciences 9: 1-60.
Seemann, T. M., 1927. Dragonflies, mayflies and stoneflies of Southern California. - Journal of Entomology and Zoology (Pomona College) 19 (1): 1-69.
Selander, R. B. \& P. Vaurie, 1962. A gazetteer to accompany the 'Insecta' volumes of the 'Biologia CentraliAmericana'. - American Museum Novitates 2099: 1-70.
Selys-Longchamps, Edm. de, 1850. Revue des Odonates ou Libellules d'Europe. - Mémoirs de la Société royale des Sciences Liège 6: xxii +408 pp .
Selys-Longchamps, Edm. de, 1854. Synopsis des Gomphines. - Bulletin de lAcadémie royale de Belgique 21 (2): 23-112 (3-93 separate).
Selys-Longchamps, Edm. de, 1858. Monographie des Gomphines. - Mémoirs de la Société royale des Sciences Liège 11: 257-720 (1-460 separate).
Selys-Longchamps, Edm. de, 1859. Additions au synopsis des Gomphines. - Bulletin de 1Académie royale de Belgique (2) 7: 530-552 (1-26 separate).
Selys-Longchamps, Edm. de, 1868. Note sur quelques Odonates nouveaux du Mexique. - Comptes Rendus de la Société entomologique de Belgique 11: lxvi-lxxi (1-6 separate).
Selys-Longchamps, Edm. de, 1869. Secondes additions au synopsis des Gomphines. - Bulletin de l'Académie royale de Belgique (2) 28: 168-208 (1-45 separate).
Selys-Longchamps, Edm. de, 1873a. Troisièmes additions au synopsis des Gomphines. - Bulletin de l'Académie royale de Belgique (2) 35: 732-774 (1-46 separate).
Selys-Longchamps, Edm. de, 1873b. Appendices aux troisièmes additions et liste des Gomphines, descrites dans le synopsis et ses trois additions. - Bulletin de l'Académie royale de Belgique (2) 36: 492-531 (47-87 separate).
Selys-Longchamps, Edm. de, 1878. Quatrièmes additions au synopsis des Gomphines. - Bulletin de l'Académie royale de Belgique (2) 46: 408-698 (1-106 separate).
Selys-Longchamps, Edm. de, 1879. Revision des Ophiogomphus et descriptions de quatre nouvelles

Gomphines Americaines. - Comptes Rendus de la Société Entomologique de Belgique 22: Ixii-lxx (1-8 separate).
Swofford, D. L. 1993. PAUP: Phylogenetic analysis using parsimony, version 3.1. - Computer program distributed by the Illinois Natural History Survey, Champaign, Illinois.
Tennessen, K. J. \& J. A. Louton, 1984. The true nymph of Gomphus (Gomphurus) crassus Hagen (Odonata: Gomphidae), with notes on adults. - Proceedings of the Entomological Society of Washington 86 (1): 223-227.
Tillyard, R., 1917. The biology of dragonflies (Odonata or Paraneuroptera). - Cambridge University Press: xii + 396 pp.
Tinkham, E. R., 1934. The dragonfly fauna of Presidio and Jeff Davis counties of the Big Bend region of trans-Pecos, Texas. - Canadian Entomologist 66: 213-218.
Tsuda, S., 1986. A distributional list of world Odonata. Preliminary edition. - Privately published, Osaka: viii + 246 pp.
Tsuda, S., 1991. A distributional list of world Odonata. Privately published, Osaka: 362 pp.
Tucker, E. S., 1907. Some results of desultory collecting of insects in Kansas and Colorado. - Kansas University Science Bulletin 4 (2): 51-112.
Tucker, E. S., 1908. Incidental captures of Neuropterous insects at Plano, Texas: - Psyche 15: 97-100.
Valley, S., 1993. D[ragonfly ]S[ociety of $]$ A[merica] meeting in Bend, Oregon. - Argia 5 (2): 3-6.
Van der Weele, H. W., 1906. Morphologie und Entwicklung der Gonapophysen der Odonaten. Tijdschrift voor Entomologie 49: 99-198.
Walsh, B., 1862. List of the pseudoneuroptera of Illinois contained in the cabinet of the writer, with descriptions of over forty new species, and notes on their structural affinities. - Proceedings of the Academy of Natural Sciences of Philadelphia 14: 361-402.
Walsh, B., 1863. Notes [on Pseudoneuroptera], pp. 182272. In Observations on certain N. A. Neuroptera, by H. Hagen, M. D., of Koenigsberg, Prussia; translated from the original French MS., and published by permission of the author, with notes and descriptions of about twenty new N. A. species of Pseudoneuroptera. - Proceedings of the Entomological Society of Philadelphia 2 (1): 167-272.
Watrous, L. E. \& Q. D. Wheeler., 1981. The out-group comparison method of character analysis. - Systematic Zoology 30(1): 1-11.
Westfall, M. J., Jr., 1984. Odonata, pp. 126-176. In R. W. Merritt and K. W. Cummins, eds. An introduction to the aquatic insects of North America (2nd ed.). Kendall/Hunt, Dubuque: xiii + 722 pp.
Westfall, M. J., Jr., 1987. Order Odonata, pp. 95-117. In F. W. Stehr, ed. Immature insects. Kendall/Hunt, Dubuque: xiv + 754 pp .
White, T. R., K. J. Tennessen, R. C. Fox, P. H. Carlson, 1980. The aquatic insects of South Carolina. Part I: Anisoptera (Odonata). - Station Bulletin, South Carolina Agricultural Experiment Station, Clemson University, Clemson: 632: 1-153.
Williamson, E. B., 1902. Additions to the Indiana list of dragonflies, with a few notes. No. II. - Proceedings of the Indiana Academy of Sciences 119-127.
Williamson, E. B., 1903. The dragonflies (Odonata) of Tennessee, with a few records for Virginia and Alabama. - Entomological News 14: 221-229.

Williamson, E. B., 1914a. September dragonflies about

Mesa, Arizona (Odon.). Entomological News 25: 225$22(0$.
Williamson, E. B., 1914b. Dragonflies (Odonata) collected in Texas and Oklahoma. - Entomological News 25: 411 -$415,444-454$.
Williamson, E. B., 1917. An annotated list of the Odonata of Indiana. - Miscellaneous Publications of the Museum of Zoology. University of Michigan 2: 1-13.
Williamson, E. B., 1918. Results of the University of Michigan-Williamson expedition to Colombia, 1916-17. 1. Two interesting new Colombian Gomphines (Odonara). - Occasional Papers of the Museum of Zoology, University of Michigan 52: 1-14.
Williamson, E. B., 1923. Odonatological results of an auto trip across Indiana, Kentucky and Tennessee. Entomological News 34: 6-9, 37-40.
Williamson, E. B., 1932. Dragonflies collected in Missouri. - Occasional Papers of the Museum of Zoology, University of Michigan 240: 1-40.
Williamson, E. B. \& J. H. Williamson, 1930. Five new Mexican dragonflies (Odonata). - Occasional Papers of the Museum of Zoology, University of Michigan 216: 134.

Wright, M., 1938. A review of the literature on the Odonata of Tennessee. - Tennessee Academy of Science 13 (1): 26-33
Wright, M. \& A. Peterson, 1944. A key to the genera of anisopterous dragonfly nymphs of the United States and Canada (Odonata, Suborder Anisoptera). - Ohio Journal of Science 44 (4): 151-166.
Young, W. C. \& C. W. Bayer, 1979. The dragonfly nymphs (Odonata: Anisoptera) of the Guadalupe River Basin, Texas. - Texas Journal of Science 31 (1): 85-98.

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Figs. 1-14. Synthorax of Erpetogomphus species, lateral view (males, except given ocherwise). - 1, constrictor (Honduras); 2, sabaleticus (holotype); 3, tristani (Costa Rica); 4, ophibolus (Mexico: Veracruz State); 5, agkistrodon (female holotrpe); 6, schaut si (holotype); 7, eutainia (Texas: Gonzales Co.); 8, eutainia (Mexico: Veracruz State); 9, eutainia (Costa Rica); 10, Leptophis (holotype); 11, elaphe (holotype); 12, elaps (Mexico: Sinaloa State); 13, elaps (Mexico: Morelos State); 14, liopeltis (holortpe). Scale line: 3 mm .


[^16]28. constrictor 9

29. ophibolus ${ }^{\circ}$

30. agkistrodon


31. schausi $\begin{gathered} \\ \end{gathered}$

32. eutainia $\delta^{\star}$

33. eutainia +

34. leptophis 아

35. liopeltis $\delta$


Figs. 28-36. Abdomen, lateral view. - 28, constrictor ㅇ (Honduras); 29, ophibolus ơ (Mexico: Veracruz State); 30, agkistrodon $\circ$ (holotype); 31, schausi ${ }^{\star}$ (holotype); 32, eutainia đै (Texas: Gonzales Co.); 33, eutainia 9 (idem); 34, leptophis ${ }^{\circ}$ (holotype); 35, leptophis 우 (allotype); 36, liopeltis ơ (holotype). Scale line 10 mm .
37. bothrops ${ }^{\text {o }}$

38. viperinus $\delta$

39. sipedon $\delta$

40. sipedon 9
41. crotalinus $\sigma^{\top}$

42. heterodon ${ }^{\circ}$

43. heterodon ${ }_{9}$

44. boa ठ

45. boa 9

46. cophias ठ


 $\therefore$.-...inu: : allorype); 44, boa ơ (Mexico: Veracruz State); 45, boa 오 (idem); 46, cophias oै (Mexico: Morelos State). Scale :ime $i 11 \mathrm{~mm}$.


48. sabaleticus

50. ophibolus

52. eutainia
53. leptophis

54. elaphe

55. elaps

56. elaps

57. liopeltis

58. bothrops

Figs. 47-58. Anterior and posterior hamules. First view is anterolateral view showing profile of anterior hamule; second view (when shown) is lateral view showing profile of posterior hamule. - 47, constrictor (Mexico: San Luis Potosi State); 48, sabaleticus (holotype); 49, tristani (Costa Rica); 50, ophibolus (Mexico: Veracruz State); 51, schausi (holotype); 52, eutainia (Texas: Gonzales Co.); 53, leptophis (holotype); 54, elaphe (Costa Rica); 55, elaps (holotype); 56, elaps (variant) (Mexico: Chiapas State); 57, liopeltis (paratype); 58, bothrops (paratype). Scale line 2 mm .


[^17]
70. constrictor

72. tristani

74. schausi

76. leptophis

78. elaps


73. ophibolus


75. eutainia

77. elaphe

79. liopeltis

## 80. bothrops

Figs. 70-80. Penis segment 4, left: dorsal view, right: lateral view. - 70, constrictor (Mexico: San Luis Potosi State); 71, sabaleticus (holotype); 72, tristani (Costa Rica); 73, ophibolus (Mexico: Veracruz State); 74, schausi (holotype); 75, eutainia (Texas: Gonzales Co.); 76, leptophis (holotype); 77, elaphe (Costa Rica); 78, elaps (holotype); 79, liopeltis (paratype); 80, bothrops (Mexico: Tamaulipas State). Scale line 1 mm .

83. sipedon

85. lampropeltis natrix

87. heterodon
89. compositus


## 91. cophias

Figs. 81-21. Penis segment 4, left: dorsal view, right: lateral view. - 81, viperinus (Mexico: Veracruz State); 82, designatus Texas: Gonzales Co.); 83, sipedon (Mexico: Morelos State); 84, l. Lampropeltis (California: Ventura Co.); 85, l. natrix Arizona: Cochise Co.); 86, crotalinus (Mexico: Michoacan State); 87, beterodon (New Mexico: Catron Co.); 88, compositus CA: Yolo Co.); 89, compositus (paratype of coluber); 90, boa (lectotype); 91, cophias (Mexico: Morelos State). Scale line 1 mon.


Figs. 92-103. Caudal appendages of male, above $=$ lateral view, below $=$ dorsal view of cerci. -92 , constrictor (Mexico: Tamaulipas State); 93, sabaleticus (holotype); 94, tristani (holotype); 95, ophibolus (Mexico: Veracruz State); 96, schausi (holotype); 97, eutainia (Texas: Gonzales Co.); 98, leptophis (holotype); 99, elaphe (Costa Rica); 100, elaps (holotype); 101, Iiopeltis (holotype); 102, bothrops (holotype); 103, viperinus (lectotype). Scale line 5 mm .

104. designatus

108. lampropeltis natrix

112. compositus

117. liopeltis 118 . bothrops 119. viperinus

106. sipedon


## 110. heterodon


115. boa

107. lampropeltis lampropeltis

111. heterodon

116. cophias
$\qquad$

Fig. 104-116. Caudal appendages of male, above = lateral view, below $=$ dorsal view of cerci. -104 , designatus (lectotype); 105, sipedon (Mexico: Morelos State); 106, sipedon (only lateral view) (Mexico: Durango State); 107, l. lampropeltis California: Ventura Co.); 108, L. natrix (holotype); 109, crotalinus (lectotype); 110, heterodon (holotype); 111, beterodon Mexico: Durango State); 112, compositus (California: Yolo Co.); 113, compositus (paratype of coluber); 114, boa (only lateral v) (lectotype); 115, boa (Mexico: Veracruz State); 116, cophias (Mexico: Morelos State). Figs. 117-119. Right cercus, melatcral view. - 117 , liopeltis (Mexico: Nuevo Leon State); 118, bothrops (Mexico: Nayarit State); 119, viperinus (Mexico: State). Scale line 5 mm .

120. tristani
121. ophibolus

122. eutainia

124. eutainia

127. bothrops

130. designatus

125. eutainia

128. bothrops

131. designatus

134. compositus

123. eutainia

126. liopeltis

129. viperinus

132. lampropeltis lampropeltis

135. boa

Figs. 120-135. Erpetogomphus. - 120, tip of epiproct of male, posterior view, tristani (Costa Rica); 121, idem, ophibolus (Mexico: Veracruz State); 122, epiproct, lateral view, eutainia (Texas: Gonzales Co.); 123, idem, dorsolateral view; 124-135, tip of epiprocts of male, posterior view, 124, eutainia (Texas: Gonzales Co.); 125, eutainia (Guatemala); 126, liopeltis (Mexico: Nuevo Leon State); 127, bothrops (Mexico: Nayarit State); 128, bothrops (holotype); 129, viperinus (Mexico: Veracruz State); 130, designatus (Texas: Gonzales Co.); 131, designatus (Mexico: Durango State); 132, l. lampropeltis (California: Ventura Co.); 133, crotalinus (Mexico: Jalisco State); 134, compositus (Arizona: Coconino Co.); 135, bod (Mexico: Veracruz State). Scale line 2 mm (figs. 122-123), 1 mm (figs. 120-121, 124-135).


Figs. 136-153. Erpetogomphus. - 136-139. Anterior part of frons, dorsal view. 136, bothrops $\circ$ (Mexico: San Luis Potosi State); 137, viperinus ㅇ (Mexico: Veracruz State); 138, l. Lampropeltis ơ (California: Ventura Co.); 139, compositus ơ (AZ: Maricopa Co.). - Figs. 140-141. Crest outline of occiput, dorsal view. 140, liopeltis (holotype), 141, liopeltis (Mexico: Nuevo Leon State). - Figs. 142-144. Head, anterior view. 142, schausi (holotype); 143, eutainia (Costa Rica); 144, eutainia (Texas: Gonzales Co.). - Figs. 145-153.Vertex, occiput of female, dorsal view. 145, constrictor (Honduras); 146, tristani (Costa Rica); 147, ophibolus (Mexico: Veracruz State); 148, agkistrodon (holotype); 149, schausi (Costa Rica); 150, eutainia (Texas: Gonzales Co.); 151, leptophis (allotype); 152, elaps (Mexico: Morelos State); 153, liopeltis (allotype). Scale line 4 mm (figs. $142-144) ; 1 \mathrm{~mm}(136-141,145-153)$.

154. bothrops

157. designatus

159. sipedon

162. liopeltis

160. lampropeltis lampropeltis

163. compositus

166. viperinus

156. designatus

161. crotalinus

164. boa

167. boa

Figs. 154-165. Vertex, occiput of female, dorsal view. 154, bothrops (allotype); 155, viperinus (Mexico: Veracruz State); 156. designatus (paralectotype); 157, designatus (Mexico: Durango State); 158, sipedon (form 'b') (Mexico: Puebla State); 159, sipedon (holotype); 160, l. lampropeltis (California: Ventura Co.); 161, crotalinus (paralectotype); 162, heterodon (allotype); 163, compositus (holotype); 164, boa (Mexico: Veracruz State); 165, cophias (Mexico: Guerrero State). - Figs. 166-16 ${ }^{-}$. Crest and part of rear of head (postocciput) of female, dorsal view. 166, viperinus (Mexico: Veracruz State); 167, boa (Mexico: Veracruz State). Scale line 1 mm .


Fig. 168-170. Mctathoracic leg. lateral view. -168 , sipedon of (holotype); 169 , crotalinus of (lectotype); 170, heterodon of tholntyper. - Figs. $1^{-1} 1$ 182. Vulvar lamina, dorsal view. - 171, constrictor (Honduras); 172, sabaleticus (allotype); 173, triatmi (allotype): $1^{-4}$, tristani (Costa Rica); 175, ophibolus (Mexico: Veracruz State); 176, agkistrodon (holotype); 177, schausi '( onta Rical Idistorted on right side): 178, eutainia (Mexico: Veracruz State); 179, leptophis (allotype); 180, elaphe (Costa Riva": 181. chaps $\backslash$ Mexico: Míorelos State); 182, Liopeltis (Mexico: Nuevo Leon State). Scale line 4 mm (figs. 168-170), 2 mm fig. 171-182).

183. bothrops

184. viperinus


185. designatus

186. designatus

189. lampropeltis 190. lampropeltis lampropeltis
 natrix

188. sipedon

192. crotalinus

193. heterodon

194. compositus

191. crotalinus
187. sipedon

195. boa

196. boa


Figs. 183-197. Vulvar lamina, dorsal view. - 183, bothrops (allotype); 184, viperinus (Mexico: Veracruz State); 185, designatus (paralectotype); 186, designatus (Texas: Gonzales Co.); 187, sipedon (holorype); 188, sipedon (paratype); 189, L. kzmpropeltis (California: Ventura Co.); 190, l. natrix (allotype); 191, crotalinus (paralectotype); 192, crotalinus (Mexico: Jalisco State); 193, heterodon (allotype); 194, compositus (Arizona: Maricopa Co.); 195, boa (Mexico: Veracruz State); 196, boa (Veracruz State); 197, cophias (Mexico: Guerrero State). Scale line 2 mm .


Figs. 198-202. Distribution patterns. - 198, E. ophibolus group; 199, E. eutainia group; 200, E. elaps group (in part); 201, E. elaps group (in part); 202, E. designatus.


Figs. 203-207. Distribution patterns. 203, E. sipedon; 204, E. lampropeltis, 205, E. crotalinus, 206, E. beterodon and E. compositus, 207, E. boa and E. cophias.

O. severus
constr.
eut. elaps
cophias

O. severus
sabalet.
eut.
viper.
liop.
lampro.



211-a
Ophiogomphus
severus


211-b tristani


211-c
viperinus


211-d compositus


Figs. 208-211. - 208, morphocline of anterior hamule, character states 0-5; 209, morphocline of posterior hamule, character states 0-4;210, morphocline of tip of posterior hamule, character stares $0-5 ; 211$, dorsal view of fourth segment of penis (S4); figs. 211 a and 211 b have 4 th segment about as long as wide, figs. $211 \mathrm{c}-\mathrm{e}$ about twice as long as wide. Scale line 2 mm (figs. 208-209): 1 mm (figs. 210-212).


| Ophiogomphus |
| :---: |
| severus |


| ophibolus |
| :---: |
| group | | eutainia |
| :---: |
| group |$\quad$| elaps |
| :---: |
| group |$\quad$ designatus | crotalinus <br> subgroup <br> (except <br> designatus) |
| :---: |



Figs. 212-215. - 212, morphocline of prepuce of penis, character states 0-3; 213, morphocline of shape of lateral lobe, character states $0-5 ; 214$, morphocline of cornua: character states $0-6$ for character 19 and character states $0-2$ for character 20 ; 215 , morphocline of spermatheca in dorsal (internal) view character states $0-2$ (figures show dorsal (internal) view of sternum 9 (including postlamellar ridge). Spematheca is dorsad of vulvar lamina). Scale line 1 mm (figs. 212, 215); 0.5 mm (figs. 213214).


Fig. 216. cladogram


Fig. 217. geographic distribution of Erpetogomphus collected in the U. S. A., Mexico, and Central America. Each square is approximately 150 kilometers square.


Figs. 218-224. - 218, Gincralized track for genus Erpetogomphus, 219, generalized tracks for Erpetogomphus ophibolus / eutuinia and E. claps / crotalinus groups; 220, generalized tracks for Erpetogomphus ophibolus and E. eutainia groups; 221, tracks (ff species within the Erpetogomphus ophibolus group; 222, tracks of species within the Erpetogomphus eutainia group; 223, generalized tracks for Erpetogomphus crotalinus and E. elaps groups; 224, tracks of species within the Erpetogomphus elaps group (in part).


227. liopeltis

229. viperinus

230. heterodon

Figs. 225-230. - 225, tracks of species within the Erpetogomphus elaps group (in part); 226, tracks of species within the Erpetogomphus crotalinus group; 227, pair of wings of liopeltis oै $^{7}$ (Mexico: Nuevo Leon State); 228, idem, bothrops ठ $^{\top}$ (Mexico: Nayarit State); 229, idem, viperinus ơ (Mexico: Veracruz State); 230, idem, heterodon oै (N. Mexico: Grant Co.)

# THE PHYLOGENY OF THE SUBGENUS TIPULA <br> (SAVTSHENKIA) (DIPTERA: TIPULIDAE), WITH 

SPECIAL REFERENCE TO THE WESTERN<br>MEDITERRANEAN FAUNA

In memory of E. N. Savchenko (12.VII. 1909 - 8.I. 1994 )


#### Abstract

Jong, H. de, 1994. The phylogeny of the subgenus Tipula (Savtshenkia) (Diptera: Tipulidae), with special reference to the western Mediterranean fauna. - Tijdschrift voor Entomologie 137: 271-323, figs. 1-202, tabs. 1-5, appendices A-C. [ISSN 0040-7496]. Published 15 December 1994.

The phylogeny of the currently recognized 87 species and subspecies of the subgenus Tipula (Savtshenkia Alexander) (Diptera, Tipulidae) is discussed in reference to 87 morphological characters of the adults. The distribution of the species and subspecies of Savtshenkia is briefly outlined. Taxonomic novelties include the description of three new western Palaearctic species, viz. alpha, omega, and trinacria; the establishment of the synonymy of broweri Alexander under fragilina Alexander, insignifica Alexander under invenusta Riedel, nebulipennis Alexander under alpium Bergroth, packardi Alexander under invenusta Riedel, and perparvula Alexander under ignobilis Loew; the elevation of the subspecies subsignata baennii Dufour to species rank; and the removal of the species convexifrons Holmgren from Savtshenkia. H. de Jong, Department of Entomology, Institute for Systematics and Population Biology (Zoological Museum), Plantage Middenlaan 64, 1018 DH Amsterdam, The Netherlands. Key words. - Diptera, Tipulidae, Savtshenkia, phylogeny, western Mediterranean.


As part of a research project on the historical biogeography of the western Mediterranean region, the phylogeny of the subgenus Tipula (Savtshenkia Alexander) is discussed. Savtshenkia as defined here, contains 87 described species and subspecies, of which 22 have a restricted distribution in the western Mediterranean. Special emphasis is laid on the establishment of the phylogenetic position of these western Mediterranean endemics, as they offer potential clues to a better understanding of the historical biogeography of the area.

The species constituting the subgenus Savtshenkia were formerly grouped in the marmorata species group of the genus Tipula Linnaeus (Alexander 1919a, 1919b et seqq.). The same assemblage of species has also been known as the fragilis group (Alexander 1934 et seqq.) and obsoleta group (Lackschewitz 1936). Usually, these species groups were referred to the subgenus Oreomyza Pokorny (Alexander 1934, Edwards 1931). Savchenko (1961), misinterpreting Riedel (1913) regarding the type species of Pterelachisus Rondani, synonymized Oreomyza under the latter taxon. Savchenko's lapsus resulted in the improper use of the name Pterelachisus for the species at present placed in Savtshenkia. Although

Savchenko's reasoning was fallacious, Mannheims (1962a) showed that the synonymy of Oreomyza and Pterelachisus sensu Rondani in itself was valid and introduced the name Savtshenkia to replace Pterelachisus sensu Savchenko. As Mannheims omitted to denote the type species of this new taxon, Savtshenkia remained a nomen nudum until Alexander (1965a) designated rufina Meigen as the type species. (In the literature, a paper by Alexander published in the 1965 volume of the Philippine Journal of Science is usually cited as the relevant reference for the authorship of the name Savtshenkia. As the pertaining publication was actually issued August 9, 1966, Alexander's paper that appeared in Pacific Insects of June 20, 1965, should be accepted as the correct reference for the type designation.)

Savtshenkia has an essentially Holarctic distribution (fig. 1), with 65 species and subspecies occurring in the Palaearctic Region and eight in the Nearctic Region. Both regions have two species in common. In addition to the Holarctic members of the subgenus, five Oriental and 12 Afrotropical species and subspecies are known. The Oriental fauna of Savtshenkia has a single species in common with the Palaearctic fauna, the Afrotropical fauna shows no species overlap


Fig. 1. Global distribution of the subgenus Tipula (Savtshenkia).
with that of any other biogeographical region. At the end of this paper a brief account of the distribution of the species of Savtshenkia is provided. I will present more detailed information on the distribution of the western Mediterranean species and their closest relatives in a forthcoming paper on the historical biogeography of the western Mediterranean. In Appendix A, three western Palaearctic species of Savtshenkia are described as new, five synonymies are established, while one subspecies is raised to species rank and one species is removed from the subgenus. The resulting list of species and subspecies of Savtshenkia as recognized in this paper is given in table 1 .

Savchenko (1961, under Pterelachisus) and Theowald (1957, under Oreomyza, 1967, 1973, 1978) arranged the (western) Palaearctic members of Savtshenkia in a number of species groups. Theowald's 1957 and 1967 arrangements were based on characters of the preadult stages of western Palaearctic species and contained four species groups (table 2). Savchenko (1961) recognized five species groups plus a miscellaneous group within the fauna of the former USSR (table 3). Theowald's 1973/1978 classification of the adults of Savtshenkia in 14 species groups basically is a refinement of Savchenko's 1961 concept (table 3). The arrangements of both authors will be compared with the results of the present study in the 'Discussion of adopted phylogeny', below.

## Material, methods and terminology

During the course of this investigation material representing 78 species and subspecies of Savtshenkia was examined (table 1). Most specimens used for the present study originated from the Institute for Systematics and Population Biology (Zoological Museum), Amsterdam. Supplementary material was studied during a short stay at the United States National Museum of Natural History, Smithsonian Institution, Washington D.C. In addition to this, material was borrowed from the following institutions: Academy of Natural Sciences, Philadelphia; Musée d'Histoire Naturelle, Neuchâtel; Natural History Museum, London; Staatliches Museum für Naturkunde, Stuttgart; United States National Museum of Natural History, Smithsonian Institution, Washington D.C.; Zoologiska Institutionen, Systematiska avdelningen, Lund.

The majority of specimens consisted of pinned material; in addition to this, material preserved in alcohol was studied from the collections in Amsterdam, Lund, and Neuchâtel.

Preparations of the male and female terminalia were made by removing these parts and clearing them in a nearly boiling $10 \% \mathrm{KOH}$ solution for about five minutes. After rinsing with water and $70 \%$ alcohol, the terminalia were transferred to glycerol. Examination of the specimens and their terminalia

Table 1. List of species and subspecies of Tipula (Savtshenkia) as recognized in this paper. Abbreviations: +: examined; - : not examined; u: unknown; Afro: Afrotropical Region; Nea: Nearctic Region; Or: Oriental Region; Pal: Palacarctic Region; Ih., Duf. \& Oost.: Theowald, Dufour \& Oosterbroek.

|  | ठ | 아 | Region |  | ${ }^{3}$ | \% | Region |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| aberdareca Alexander, 1956 | + | u | Afro | letifera Alexander, 1951 | - | $u$ | Pal |
| a. ulugurica Alexander, 1962 | u |  | Afro | limbata Zetterstedt, 1838 | + | + | Pa |
| akeleyi Alexander, 1956 | + | $u$ | Afro | lundbladi Mannheims, 1962 | + | + | Pal |
| alpha sp. n. | + | + | Pa | macaronesica Savchenko, 1961 | + | $u$ | Pal |
| alpium Bergroth, 1888 | + | + | Nea, Pal | mannheimsi Theowald, 1973 | + | $u$ | Pal |
| asbolodes Speiser, 1909 | + | - | Afro | minuscula Savchenko, 1971 |  | - | Pal |
| aspromontensis Theowald, 1973 | + | + | Pal | mohriana Alexander, 1954 | + | + | Pal |
| aster Theischinger, 1983 | + | u | Pal | multipicta Becker, 1908 | + | + | Pal |
| atlas Pierre, 1924 | + | + | Pa | nephrotomoides Alexander, 1924 | + | + | Afro |
| baltistanica Alexander, 1936 | + | + | Or, Pal | n. invariegata Alexander, 1956 | $u$ | + | Afro |
| benesignata Mannheims, 1954 | + | + | Pal | nielseni Mannheims, \& Theowald, 1959 |  | + | Pal |
| boreosignata Tjeder, 1969 | + | u | Pal | nivalis Savchenko, 1961 |  | - | Pal |
| breviantennata Lackschewitz, 1933 | + | + | Pal | obsoleta Meigen, 1818 | + | + | Pal |
| caligo Alexander, 1956 | + | + | Afro | odontsstyla Savchenko, 1961 | + | - | Pal |
| cheethami Edwards, 1924 | + | + | Pal | omega sp. n. | + | u | Pal |
| chrysocephala Mannheims, 1958 | + | u | Afro | ornata Theowald \& Oosterbroek, 1987 |  | + | Pal |
| confusa Van der Wulp, 1887 | + | + | Pal | pagana Meigen, 1818 | + | + | Pal |
| corsosignata Th., Duf. \& Oost. 1982 | + | + | Pal | pechlaneri Mannheims \& |  |  |  |
| cyrnosardensis Th., Duf. \& Oost. 1982 | + | + | Pal | Theowald, 1959 | + | + | Pal |
| draconis Alexander, 1964 |  | $u$ | Afro | persignata Alexander, 1945 | + | $u$ | Pal |
| eleonorae Theischinger, 1978 | + | $u$ | Pal | p. tofina Alexander, 1945 | + | u | Pal |
| elgonensis Alexander, 1956 | + | $u$ | Afro | phoroctenia Alexander, 1919 | + | + | Nea |
| eugeni Theowald, 1973 | + | + | Pal | postposita Riedel, 1919 | + | - | Pal |
| fragilina Alexander, 1919 | + | + | Nea | productella Alexander, 1928 |  | $u$ | Nea |
| fragilis Loew, 1863 | + | + | Nea | rufina Meigen, 1818 | + | + | Pal |
| gimmerthali Lackschewitz, 1925 | + | + | Pal | r. maderensis Lackschewitz, 1936 | + | + | Pal |
| g. mattheyi Theowald \& Dufour, 1983 |  | $u$ | Pal | sardosignata Mannheims \& |  |  |  |
| g. pteromaculata Th., Duf. \& Oost. 1982 |  | u | Pal | Theowald, 1959 | + | + | Pa |
| glaucocinerea Lundström, 1915 | + | + | Pal | sciadoptera Alexander, 1964 | + | u | Or |
| goriziensis Strobl, 1893 | + | + | Pal | serrulifera Alexander, 1942 | + | + | Pa |
| graciae Alexander, 1947 | + | $u$ | Nea | signata Staeger, 1840 | + | + | Pal |
| grisescens Zetterstedt, 1851 | + | + | Pal | simulans Savchenko, 1966 | + | + | Pal |
| baennii Dufour, 1991 stat.n. | + | - | Pal | sordidipes Alexander, 1961 | + | + | Or |
| bancocki Alexander, 1956 | + | + | Afro | staegeri Nielsen, 1922 | + | + | Pal |
| bartigiana Th., Duf. \& Oost. 1982 | + | + | Pal | subalpium Savchenko, 1961 | - | u | Pa |
| boloptera Edwards, 1939 | + | + | Pal | subnodicornis Zetterstedt, 1838 |  | + | Pal |
| ignobilis Loew, 1863 | + | + | Nea | subsignata Lackschewitz, 1933 | + | + | Pal |
| imperfecta Riedel, 1914 | - | - | Afro | s. cazorla Dufour, 1991 | + | $u$ | Pal |
| interserta Riedel, 1913 | + | + | Pal | subvafra Lackschewitz, 1936 | + | + | Pal |
| invenusta Riedel, 1919 | + | - | Nea, Pal | tetragramma Edwards, 1928 | - | $u$ | Or |
| i. microinvenusta Dufour, 1990 |  | + | Pal | trinacria sp. n. | + | + | Pal |
| i. subinvenusta Slípka, 1950 | + | + | Pal | tulipa Dufour, 1983 | + | $u$ | Pal |
| jeekeli Mannheims \& Theowald, 1959 | + | + | Pal | venerabilis Alexander, 1936 | + | $u$ | Or |
| kiushiuensis Alexander, 1925 |  | + | Pal | villenenvii Strobl, 1909 | + | + | Pal |
| koreana Alexander, 1934 | + | + | Pal |  |  |  |  |

was carried out using a Wild stereomicroscope with a magnification of up to 100 times. Drawings were made with the aid of a drawing tube attached to the microscope. Illustrations of the genital structures were made from macerated specimens. For permanent storage, the terminalia were transferred to a microvial containing a drop of glycerol. The microvial was pinned with the relevant specimen.

The parsimony program Hennig86, version 1.5 (Farris 1988) was used to analyse the phylogeny.

More information on the calculation procedures followed is given in 'Discussion of adopted phylogeny'.

The terms for the structures of Savtshenkia as used in the character discussion are in general in accordance with the terms employed by McAlpine (1981), with a few additions for particular features of the Tipulidae of the study group. The terms are explained in figs. 3-6, 12, and 58. It should be noted that I consider the structure lateral of the gonapophysis in Savtshenkia homologous with the fragmentum


Fig. 2. Strict consensus tree of 14 equally parsimonious trees derived after successive weighting for the species and subspecies of Tipula (Savtshenkia). -: synapomorphy; $\square$ : homoplasy; $X$ : reversal.
in the sense of Rees \& Ferris (1939) and Frommer (1963) and name it accordingly (figs. 3, 4). Mannheims (1951) denoted the same sclerite A9s (appendage of sternite 9). Suture x is employed in the sense of Neumann (1958). A recent overview of the morphology of the terminalia of Tipulidae was given by Tangelder (1985) to which paper the reader is referred to for further details.

Acronyms used throughout the text are the following:
bmnh: Natural History Museum, London, England; bsnh: Boston Society of Natural History, Boston, Massachusetts, U.S.A.; CNC: Canadian National Collection, Biosystematics Research Centre, Ottawa, Ontario, Canada; mczc: Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts, U.S.A.; mnhn: Muséum National d'Histoire Naturelle, Paris, France; mrac: Koninklijk Museum voor Midden Afrika, Tervuren, Belgium; usnm: United States National Museum, Smithsonian Institution, Washington D.C., U.S.A.; zlas: Zoological Institute, Russian Academy of Sciences, St. Petersburg, Russia; zman: Institute for Systematics and Population Biology, Department of

Entomology, Zoological Museum, Amsterdam, the Netherlands.

## Biology

In general, the members of Savtshenkia are adapted to either mountainous, high latitudinal, or autumnal conditions. Within these limits, adults of the respective species of Savtshenkia show a wide variety of preferred biotopes. They can be found in alpine Rhododendron assocations, the heath zone at about 3300 m , mountainous wet or dry Pinus forests and mixed woods, low woods, low moorlands, marshes, bogs, and gardens. The species may occur in the vegetation near the sources and on the banks of streams, on boulders in streams, on dripping rocks, and in the splash zone of cascades (Alexander 1961, Audcent 1932, Brunhes \& Dufour 1992, Bryce 1956, Coulson 1959, 1962, Dufour 1983, 1986, 1990, 1991, Mannheims \& Pechlaner 1963, Stubbs 1992, Theowald 1973, Theowald, Dufour \& Oosterbroek 1982).

The larvae of the species of Savtshenkia are associated with mosses and liverworts. They are recorded

from mosses on stones and $\log s$ in and along cold streams, from wet moorland mosses and mosses of cold bogs of tundra and high mountains, as well as from relatively dry mosses on walls, stones and trees (Gelhaus 1986).
As in most Tipulidae, the majority of species of Savtshenkia have a one-year life cycle. A few are, or appear to be, bivoltine (viz. alpium, breviantennata, rufina rufina, and rufina maderensis, Dufour 1986, Mannheims 1951, Theowald 1973). On the other hand, Lantzov (1982) suggested a long term larval development for the arctic species glaucocinerea of up to six years. [Multi-annual life cycles are known of other species of Tipulidae with a northerly distribution; see Pritchard (1983) for an overview of life cycles in Tipulidae.]

Details on the copulation of a few species of Savtshenkia were given by Hemmingsen (1954, 1962; viz. limbata, signata, staegeri, and subsignata). Hemmingsen (1952) provided a short account on oviposition of the species confusa (as marmorata) and rufina.

## Systematic position

The species and subspecies of Savtshenkia can be distinguished from other Tipulidae by the following
derived character states: (1) Adult, male terminalia: sp2 medially blackish sclerotized (character 53, below); (2) Adult, female terminalia: dorsal margin of hypogynial valve blackish sclerotized (character 83, below); (3) Larva: eight anal papillae arranged in four pairs. The number of anal papillae in other groups of Tipulidae is either six, four, two, or zero (Gelhaus 1986, Theowald 1957, 1967). As both adult characters occur also in other genera and subgenera of Tipulidae, the larval character seems to offer the best criterion to distinguish Savtshenkia from the remainder of Tipulidae.

The systematic position of Savtshenkia within the Tipulidae is not yet fully established. Savchenko $(1966,1979,1983)$ placed Savtshenkia next to the subgenus Tipula (Mediotipula Pierre), a relationship substantiated by the presence in the female pupa of a laterally placed small tubercle at the apex of the sheath of the hypogynial valve (see Theowald 1957, 1967). In species of the genus Dolichopeza Curtis, the sheath of the hypogynial valve is also provided with an extension, but here the extension is much bigger than in Mediotipula and Savtshenkia, and is placed near midlength of the sheath (Byers 1961, Theowald 1957, 1967, Wood 1952). Dolichopeza probably is not closely related to Mediotipula and Savtshenkin, and the extension of the sheath of the hypogynial


Figs. 3-6. Tipula (Savtshenkia) mifina rufina. - 3, male terminalia, lateral view; 4, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 5 , sperm pump, lateral view; 6 , female terminalia, lateral view.
Abbreviations: aed: aedeagus; aed gd: aedeagal guide; ant apod: anterior apodeme; cerc: cercus; comp apod: compressor apodeme: endoph: endophallus; fragm: fragmentum; gen fk: genital fork; gon: gonapophysis; goncx: gonocoxite; hyp vlv: hypogynial valve: i gonst: inner gonostylus; midv area: midventral area: o gonst: outer gonostylus; post apod: posterior apodeme; prct: proctiger; sp2: lateral part of genital bridge; spm pmp: sperm pump; spmth: spermatheca; st7 etc.: sternite 7 etc.; sut $x$ : surure $x$; $\operatorname{tg} 7$ etc.: tergite 7 etc.
valve in Dolichopeza supposedly is not homologous to that of Mediotipula and Savtshenkia. Theowald (1978) accepted Savchenko's views regarding a sister group relationship of Mediotipula and Savtshenkia. Another character state which could be indicative of a close phylogenetic relationship of Mediotipula and Savtshenkia is the shape of wing cell ml . In both subgenera this cell is 'bell-shaped', narrowing towards the wing margin, whereas it usually widens towards the wing margin in other Tipulidae (character 4, below). The genital fork in both Mediotipula and Savtshenkia is broadest at its anterior part, a presumed apomorphy within Tipulidae (character 82, below).

Mediotipula at present contains 12 recognized species with a primarily Mediterranean distribution (Oosterbroek \& Theowald 1992).

Conflicting with a presumed sister group relationship of Mediotipula and Savtshenkia is the presence of a separate sclerite lateral of the gonapophysis in Savtshenkia. As stated above (section 'Material, methods and terminology'), I consider this sclerite homologous with the so-called fragmentum as found in other Tipulidae. The presence of a fragmentum is a derived character state within the Tipulidae and could substantiate a phylogenetic relationship of Savtshenkia with the complex of Lunatipula and allies
(here after referred to as the Lunatipula group). The fragmentum is absent in Mediotipula. The Lunatipula group contains about 1200 species of Tipulidae, that is circa one fourth of the total number of species of the family. Besides the subgenus Lunatipula Edwards, the Lunatipula group includes the following subgenera of Tipula: Beringotipula Savchenko, Eremotipula Alexander, Eumicrotipula Alexander, Hesperotipula Alexander, Labiotipula Alexander, Lindnerina Mannheims, Odonatisca Savchenko, Pectinotipula Alexander, Pterelachisus Rondani, Ramatipula Alexander, Serratipula Alexander, Setitipula Alexander, Triplicitipula Alexander, and Vestiplex Bezzi. The phylogenetic relationships of the species of the Lunatipula complex are still largely unresolved, but it is nevertheless clear that not all of the subgenera in their present delimitation represent monophyletic taxa (e.g., Lunatipula and Pterelachisus).

In the discussion of the characters, I will refer to the species of the Lunatipula group and Mediotipula as outgroups of Savtshenkia.

## Character discussion

In this section I will discuss the characters used in the phylogenetic analysis of Savtshenkia. Preceding a more detailed discussion of each character, a couplet shortly defines the alternative character states and gives their codes. The character state matrix is presented in table 4. Although I did not examine the male of gimmerthali mattheyi, this subspecies is, according to its authors, very similar to the nominotypical form, differing only in minute discriminating characters. Therefore, I feel safe to copy the score of the male characters of gimmerthali gimmerthali for this subspecies. In the discussion of the characters below, the resulting strict consensus tree (fig. 2) will often be mentioned to point out the species which show a particular character state. I have done this in order to save space and for convenience, rather than presenting a list of species for each of the distinguished states. The results of the phylogenetic analysis are discussed in the next section. Species and subspecies of Savtshenkia which for various reasons are not included in the character discussion below, are dealt with in Appendix B. A list of autapomorphies of the species and subspecies of Savtshenkia is presented in Appendix C.

## Antenna

1.     - Flagellum: (0) well developed in male; (1) short, about the length of head or shorter.

The male antenna of most species of Savtshenkia is well developed and relatively long, with the flagellomeres being about 2.5 times as long as their diameter at the greatest width, or longer (figs. 7, 11, 171).

A similar state represents the presumed groundplan condition of the Lunatipula group and Mediotipula. A few members of Savtshenkia are characterized by a conspicuously short flagellum in the male. The flagellomeres of the pertaining species are about two times as long as their diameter or shorter (fig. 8). The species with exceptionally short male antennae are atlas, breviantennata, cyrnosardensis, and villeneuvii.
2. - Flagellum: (0) male flagellomeres well developed; (1) male flagellomeres longer than standard length.
As noticed under the previous character, the flagellomeres of the males of Savtshenkia are usually about 2.5 times as long as their diameter at greatest width or slightly longer. Flagellomeres distinctly longer than this are found in alpha, grisescens, holoptera, pagana, and most Afrotropical species (the clade elgonensis to chrysocephala; figs. 9, 10, 152). In the pertaining species, the male flagellomeres are about 5 to 6.5 times as long as wide. The only Afrotropical species of Savtshenkia of which the male antennae are not conspicuously long are hancocki and nephrotomoides (and the unexamined imperfecta which doubtfully belongs to Savtshenkia; see Appendix B).
3. - Flagellum: (0) male flagellomeres with slightly enlarged base, remainder cylindrical; (1) flagellomeres more or less reniform.

In the genus Tipula sensu lato, the flagellum is usually composed of eleven segments of which the intermediate ones have a slightly bulbous base and a cylindrical apical part (figs. 7-10, 152, 171). This character state probably represents the groundplan condition of the Lunatipula group and Mediotipula. The same state is also found in most species of Savtshenkia. The species lundbladi and multipicta differ from the other species of Savtshenkia by having the intermediate flagellomeres more or less reniform (fig. 11). In the Lunatipula group, a similar form of the flagellomeres is present in most species of the subgenus Vestiplex. These species appear, however, to represent a more derived group within Vestiplex.

## Wing

4.     - Cell m1 shape: (0) widening towards wing margin; (1) constricted before wing margin.

In the Tipulidae veins M1 and M2, enclosing cell m1, usually diverge towards the wing margin (fig. 12). This situation probably represents the groundplan condition of the Lunatipula group, which contains only a few species that have cell ml constricted near the margin of the wing. In most species of Mediotipula, cell ml is bell-shaped or paunchy ('bauchig', Mannheims \& Pechlaner 1963; Theowald 1973) with veins M1 and M2 approaching towards


Figs. -15.-7-11, male antenna; 7, Tipula (Savtshenkia) hartigiana; 8, T. (S.) atlas, 9, T. (S.) grisescens, 10, T. (S.) elgonensis, 11, T. (S.) lundbladi; 12-15, tip of wing; 12, T. (S.) grisescens, 13, T. (S.) goriziensis, 14, T. (S.) cheethami; 15, T. (S.) akeleyi. Abbreviations: M1 etc.: medial vein 1 etc.; R4 etc.: radial vein 4 etc.; r-m: radial-medial crossvein.
the margin of the wing, a state which probably represents the groundplan condition of Mediotipula. The majority of species of Savtshenkia also show this latter character state (figs. 13-15). Only a few species of Savtshenkia have veins M1 and M2 diverging towards the wing tip, viz. alpha, glaucocinerea, grisescens, and postposita. The species subnodicornis usually has cell ml narrowing, but specimens of this species are known which show a widening cell ml.
5. - Cells of wing tip: (0) membrane covered with microtrichia only; (1) membrane partly covered with macrotrichia.

Most species of Tipulidae have the membrane of the wing tip covered with microtrichia only, appearing bare under low magnification (figs. 12, 13, 15). This situation is found in the species of the Lunatipula group, Mediotipula, and in most species of Savtshenkia. The species cheethami and simulans are


Figs. 16-31. - 16, Tipula (Savtshenkia) grisescens, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 17, T. (S.) hartigiana, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 18, T. (S.) atlas, aedeagal guide with right gonapophysis, posteromedial view; 19, T. (S.) draconis, aedeagal guide with gonapophyses, ventral view after microscopic slide holotype; $20, T$. (S.) aberdareica, aedeagal guide with right gonapophysis, posteromedial view; 21 , T. (S.) obsoleta, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 22, T. (S.) fragilis, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 23, T. (S.) limbata, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 24, T. (S.) pechlaneri, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 25-27, T. (S.) baennii, 25, aedeagal guide, tip, posterior view; 26, right gonapophysis, medial view; 27. right fragmentum, posteromedial view; $28-30, T$. (S.) corsosignata, 28 , aedeagal guide, tip, posterior view; 29 , right gonapophysis. medial view; 30 , right fragmentum, posteromedial view; 31, T. (S.) subsignata subsignatt, right gonapophysis, medial view.
unique within Savtshenkia by having macrotrichia on the membrane of the wing tip. The macrotrichia cover the wing to a larger extent in simulans than in cheethami. The density of the macrotrichia is also higher in the former species. Macrotrichia are usually restricted to cells $\mathrm{r} 4, \mathrm{r} 5$, and mI in cheethami (fig. 14), while in simulans cell m 2 and the tip of cell m 3 are also provided with macrotrichia.
6. - Crossvein r-m: (0) apical section directed towards wingtip, or at most transverse to
longitudinal axis of wing; (1) apical section directed towards base of wing.

In Tipulidae crossvein r-m usually lies in an oblique position relative to the longitudinal axis of the wing, with its apical part directed towards the wing tip. This situation is found in the species of the Lunatipula group, Mediotipula, and in the majority of species of Savtshenkia (figs. 12-14). Three species of Savtshenkia have the apical end of $\mathrm{I}-\mathrm{m}$ directed towards the base of the wing, viz. akeleyit caligo, and chrysocephala (fig. 15).

Male terminalia
7. - Aedeagal guide, apex: (0) tapering towards tip; (1) at tip with flaring rim.

As is usual in the Tipulidae, the apex of the aedeagal guide is the narrowest part of this structure in the species of the Lunatipula group, Mediotipula, and the majority of species of Savtshenkia (figs. 4, 16, 17, 19, $20,22-25,28,162,182,194)$. The species atlas, breviantennata, and obsoleta have the apex of the aedeagal guide bent outwards as a flaring rim (figs. 18, 21).
8. - Aedeagal guide, apex: (0) gradually tapering towards tip; (1) subapically abruptly narrowed, terminating in a slender point.

The aedeagal guide in Tipulidae normally gradually narrows towards its apex, as it does in almost all species of Savtshenkia (caudal view, fig. 25). The apex of the aedeagal guide of corsosignata, subsignata subsignata, and subsignata cazorla differs from that of the other species of Savtshenkia by the abruptly constricted apex which terminates in a slender tip (fig. 28).
9. - Aedeagal guide, gonapophysis: (0) well developed; (1) very short or almost completely absent.

Gonapophyses are usually well developed in the Tipulidae, a state which probably represents the groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also have well developed gonapophyses which occur in a wide range of sizes and shapes (figs. 4, 17-24, 26, $29,31,182,194)$. The gonapophyses are very short or almost completely absent in the species alpha, alpium, boreosignata, grisescens, interserta, and macaronesica (figs. 16, arrow, 162).
10. - Aedeagal guide, gonapophysis: (0) dorsal margin evenly fused with base of aedeagal guide; (1) anterodorsally slightly extended, thus with free-lying dorsal tip.

In general, the dorsal margin of the gonapophysis in the Tipulidae gradually merges with the base of the aedeagal guide. This situation probably represents the groundplan condition of the Lunatipula group and Mediotipula. The gonapophyses of most species of Savtshenkia also show this character state (figs. 16, 21-24, 162, 182, 194). The species of the clade hartigiana to simulans are characterized by their typically shaped gonapophyses, which differ from the gonapophyses of the other species of Savtshenkia by the presence of a free anterodorsal extension (figs. 17, arrow, 18).
11. - Aedeagal guide, gonapophysis: (0) pubescent or short haired; (1) long haired.

The covering of the gonapophyses in the Tipulidae usually consists of pubescence or short hairs, as is the presumed groundplan state of the Lunatipula group and Mediotipula. A similar condition is found in the majority of species of Savtshenkia (figs. 4, 17-19, 2124). A number of Afrotropical species of Savtshenkia are distinguished from the remainder of the subgenus by the dense and elongate hairing of the gonapophyses, viz. aberdareica, akeleyi, asbolodes, caligo, chrysocephala, and hancocki (fig. 20).
12. - Aedeagal guide, gonapophysis: (0) present, variously shaped; (1) a large and elongate structure.

The gonapophyses of the Tipulidae show a wide range of shapes. When present, they are usually well developed and rather slender. This is the state that occurs in most of the species of Savtshenkia and is the presumed groundplan condition of its outgroups (figs. 4, 17-21, 23, 24, 182, 194). The Nearctic species fragilis and phoroctenia differ from the majority of species of Savtshenkia by their extremely large and elongate gonapophyses (fig. 22). The only other species of Savtshenkia with similarly enlarged gonapophyses is the western Mediterranean cymosardensis.
13. - Aedeagal guide, gonapophysis: (0) a relatively slender structure; (1) a large blade.

As noted under the previous character, the plesiomorphous condition of the gonapophyses in Savtshenkia is probably represented by a well developed, rather slender structure (figs. 4, 17-21, 23, 24, 182, 194). Large mediolaterally flattened gonapophyses are found in corsosignata, haennii, subsignata subsignata, and subsignata cazorla (figs. 26, 29, 31; see also next character).
14. - Aedeagal guide, gonapophysis: (0) at apex at most moderately enlarged; (1) dorsally and ventrally extended and thus widening towards broad apex.

The two subspecies subsignata subsignata and subsignata cazorla are characterized by their expanded gonapophyses (fig. 31). This character state corresponds with character 6 of Dufour (1991).
15. - Aedeagus: (0) short, directly curving from sperm pump to aedeagal guide, anteriorly reaching as far as abdominal segment 8 ; (1) elongate, anteriorly running to segment 7 or beyond.

A relatively short aedeagus which loops in the median sagittal plane from the sperm pump through abdominal segment 8 and from there to the aedeagal guide, is the presumed groundplan state of this feature in the Lunatipula group. The species of Mediotipula also show this character state, as do a number of species of Savtshenkia (figs. 3, 155). Other species of Savtshenkia have a longer aedeagus. The ae-
deagus in these species reaches anteriorly at least to segment 7 and often lies coiled inside the abdomen (figs. 173, 185). The species involved constitute the clades confusa to aster and fragilis to omega. Extremely long aedeagi were found in corsosignata, ornata, and staegeri, species in which the aedeagus reaches abdominal segment 1 . The species pair boreosignata and eleonorae is characterized by a peculiarly modified aedeagal complex (character 52). Although belonging to the clade fragilis to omega, they have a short aedeagus.
16. - Fragmentum: (0) either absent, or posterior margin convex or slightly concave; (1) posterior margin subapically with sharp emargination.

The fragmenta of the species of the Lunatipula group and Savtshenkia occur in a diversity of forms. Usually, the posterior margin near the dorsal tip of the structure is convex. This is the presumed groundplan condition of the Lunatipula group and the common character state in Savtshenkia (figs. 4, 16, 17, 22$24,27,30,162,182,194)$. (The fragmentum is absent in the species of Mediotipula.) The fragmenta of the species holoptera, obsoleta, and pagana differ from the standard situation of Savtshenkia by the presence of a distinct emargination just below the dorsal apex (fig. 21, arrow).
17. - Fragmentum: (0) either absent, or anterior part sclerotized as the remainder of the structure; (1) anterior part dorsally blackish sclerotized.

In general, the fragmentum is yellowish-brown in colour and uniformly sclerotized in the species of the Lunatipula group and Savtshenkia (figs. 4, 16, 17, 21, $22,24,162,182,194)$. A unique state is found in the species fragilina, limbata, and subvafra, where the dorsal margin of the anterior part of the fragmentum is blackish sclerotized (fig. 23).
18. - Fragmentum: (0) either absent, or anterior part a rounded lobe; (1) anterior part a separate anterocaudally flattened extension with slender tip.

The fragmentum consists of a single undivided sclerite in the species of the Lunatipula group and most species of Savtshenkia (figs. 4, 16, 17, 21-24, 162, 182, 194). In corsosignata, baennii, subsignata subsignata, and subsignata cazorla, the anterior part of the fragmentum is characteristically modified in a slender tipped structure which stands apart from the posterior part (figs. 27, 30). The few hairs of the brush on top of the anterior part of the fragmentum are very short in these four taxa.
19. - Gonocoxite, midventral area: (0) variously shaped, usually tumescent; (1) with cone-shaped membranous extension.

The midventral area in between the gonocoxites is
represented by a tumescent structure in the presumed groundplan state of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also have a tumescent midventral area (fig. 32), but a number of species show a modification of this part of the terminalia. The species atlas, breviantennata, cheethami, bartigiana and simulans have the midventral area provided with a ventrally directed cone-shaped membranous projection. The projection is relatively low in atlas and breviantennata (fig. 34), in the other species listed it is longer and more slender (figs. 33, 35). The structure is placed near the posteroventral margin of the gonocoxites in cheethami (fig. 35), in the four other species it has a more anterior position (figs. 33, 34).
Other species of Savtshenkia with a projection to the area in between the gonocoxites are alpium, aspromontensis, macaronesica, ornata, phoroctenia, and a few Afrotropical species. As the actual shapes of the projections differ notably among these species, a number of different character states are distinguished. In the species alpium and macaronesica, the appendage is membranous, dorsoventrally flattened and midventrally placed (character 21; fig. 39), in ornata it is swollen, sclerotized and posteriorly projecting (fig. 40), in phoroctenia it is slender, sclerotized and posteriorly projecting (fig. 41). The modifications of the midventral area in ornata and phoroctenia are characteristic of the respective species and represent autapomorphies (see Appendix C). Males of the species aspromontensis originating from Greece (Peloponnisos, Crete) have the midventral area posteriorly terminating in a small and low extension. A similar extension is absent in specimens of aspromontensis from the type locality in southern Italy. I could not find other distinguishing characters between male specimens of the populations from Greece and from Italy and consider them conspecific. The Afrotropical species of Savtshenkia with a modified midventral area include akeleyi caligo, chrysocephala, and nephrotomoides. These species are characterized by a relatively large and bulbous extension to the midventral area (next character).
20. - Gonocoxite, midventral area: (0) posterior part not bulbous; (1) bulbous.
As noted under the previous character, the midventral area in Savtshenkia usually is tumescent. The Afrotropical species caligo, chrysocephalas nephrotomoides, and, to a lesser extent, akeleyí are distinguished by a posteroventrally projecting lobe near the posterior margin of the midventral plate. The lobe is large in the first three species listed (fig. 36), in akeleyi it is relatively low (fig. 38).
21. - Gonocoxite, midventral area: (0) without


Figs. 32-45. - 32-41, male sternite 9, left gonocoxite and midventral area, lateral view; 32, Tipula (Savtshenkia) grisescens, 33, T. (S.) hartigiana; 34, T. (S.) atlas, 35, T. (S.) cheethami; 36, T. (S.) nephrotomoides, 37, T. (S.) aberdareica; 38, T. (S.) akeleyí 39, T. (S.) alpium; 40, T. (S.) ornata; 41, T. (S.) phoroctenia; 42-45, male sternite 9, gonocoxites and midventral area, ventral view; 42, T. (S.) baltistanica; 43, T. (S.) lundbladi; 44, T. (S.) nielseni; 45, T. (S.) baennii.
Abbreviations: midv area: midventral area; sut x : suture x .
flattened appendage; (1) at about midlength with dorsoventrally flattened appendage.

As stated above, the midventral area in between the gonocoxites is tumescent in most species of Savtshenkia. The species alpium and macaronesica are distinguished from the other species of Savtshenkia by the presence of a dorsoventrally flattened membra-
nous appendage at about midlength of the midventral area. The appendage is long in alpium (fig. 39), in macaronesica it is considerably shorter. (The unexamined subalpium has a small tongue-shaped appendage to the midventral area and is probably closely related to alpium and macaronesica; see Appendix B).


Figs. 46-51. Male tergite 9-10 and dorsal part of left gonocoxite plus appendages, lateral view. - 46, Tipula (Savtshenkia) hartigiana; 47, T. (S.) tulipa; 48, T. (S.) baennii; 49, T. (S.) boreosignata; 50, T. (S.) benesignata; 51, T. (S.) persignata tofina.
22. - Gonocoxite, midventral area: (0) broad and usually separated from gonocoxites by membranous zone; (1) narrow and fused with gonocoxites.

The midventral area is probably broad in the groundplan states of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also have a broad midventral area in between the gonocoxites (figs. 42-44). It either is membranous, or partly or entirely sclerotized. In most Savtshenkia species with a sclerotized midventral area, a membranous zone (partly) separates the plate from the gonocoxites.

The species pechlaneri and the members of the clade baennii to omega have a slender parallel sided and sclerotized midventral area which is largely fused with the ventromedial margins of the gonocoxites (fig. 45).
23. - Gonocoxite, posterodorsal margin: (0) with scattered short setae; (1) with concentration of long and strong setae.

The gonocoxites of most species of Tipulidae are covered with scattered setae, a situation which probably represents the groundplan condition of this char-


Figs. 52-63. Left inner gonostylus. - 52, Tipula (Savtshenkia) grisescens, dorsal view; 53, T. (S.) villeneuvii, dorsal view; 54, $T$. (S.) goriziensis, dorsal view; 55, T. (S.) rufina rufina, lateral view; 56, T. (S.) aberdareica, lateral view; 57, T. (S.) akeleyi, lateral view; $58, T$. (S.) confusa, lateral view; $59, T$. (S.) ornata, lateral view; $60, T$. (S.) serrulifera, lateral view; $61, T$. (S.) aspromontensis, lateral view; 62, T. (S.) fragilis, medial view; 63, T. (S.) odontostyla, medial view.
Abbreviations: ant pt: anterior part; lat rdg: lateral ridge; post pt: posterior part.
acter for the Lunatipula group and Mediotipula. It is also found in most species of Savtshenkia (figs. 3, 46, 48, 49, 155). Within Savtshenkia, the species of the clades tulipa to simulans and benesignata to omega are distinguished by a concentration of strong setae along the posterodorsal margin of the gonocoxites (figs. 47 , arrow, $50,51,173,185)$. The setae are placed in two or three irregular rows. They extend dorsally towards the membranous suture separating the gonocoxite from tergite 9-10 in the species of the clade tulipa to simulans and in benesignata (figs. 47, 50). In the species of the clade persignata to omega, the setae are confined to a more ventral position (fig. 51, 173, 185).

## 24. - Gonocoxite, posterodorsal margin: (0)

posterodorsal corner rounded off; (1) with posterior projection.

The posterodorsal corner of the gonocoxites is presumably rounded off in the groundplan condition of the Lunatipula group, as it is in the majority of species of Savtshenkia. In the species of Mediotipula and a number of species of Savtshenkia, the posterodorsal margin of the gonocoxite is extended into a more or less well developed projection (fig. 48, arrow). The species of Savtshenkia which show this character state
are those of the range fragilina to subsignata cazorla.
25. - Gonocoxite, suture x: (0) angle between suture $x$ and the median line about $45^{\circ}$ or less; (1) angle between suture and median line much wider, up to $90^{\circ}$.

Suture x usually stands oblique to the median line of the body under an angle of about $45^{\circ}$ or less in the Tipulidae. This probably is the groundplan condition of the character for the Lunatipula group and Mediotipula, and is the general state in Savtshenkia (figs. $42,44,45$ ). In the species lundbladi, multipicta, rufina, and the Afrotropical members of the subgenus, the angle is much wider, often reaching a magnitude of about $90^{\circ}$ (fig. 43). (The direction of suture $x$ could not be adequately studied in the celluloid preparation of the holotype of elgonensis, the only known specimen of this species.)
26. - Gonocoxite, suture x: (0) narrow; (1) broad.

Suture x usually is rather narrow in the Tipulidae, a condition considered the groundplan state of the Lunatipula group and Mediotipula. A narrow suture x is also found in the majority of species of Savtshenkia (figs. 3, 32-39, 41-44). The species and subspecies
corsosignata, baennii, subsignata subsignata, and subsignata cazorla differ from the other members of Savtshenkia by their broad suture x (fig. 45, arrow). The suture is straight in these four taxa. Other species of Savtshenkia with a broad suture x are fragilis and ornatd. In fragilis, suture x is sigma-shaped, in ornata it is, as many other genital features in this species, very different from that of the remainder of species of Savtshenkia. In ornata, suture $x$ encompasses an isolated sclerite (fig. 40, arrow; see Appendix C).
27. - Inner gonostylus, anterior part: (0) ventral surface smooth, or at most rugged; (1) serrate.

The ventral surface of the anterior part of the inner gonostylus usually is smooth in Tipulidae. It is considered the groundplan state of the Lunatipula group and Mediotipula, and is found in the majority of species of Savtshenkia (all figures that depict the inner gonostylus in side view, except 57). Three Afrotropical members of Savtshenkia are characterized by a partly serrate lower margin of the anterior part of the inner gonostylus, viz. akeleyi, caligo, and chrysocephala (fig. 57, arrow).
28. - Inner gonostylus, anterior part: (0) crest absent on posterodorsal margin of anterior part; (1) crest present.

The anterior part of the inner gonostylus probably lacks a crest in the groundplan state of the Lunatipula group. All species of Mediotipula are devoid of a crest on this part, as are most species of Savtshenkia (the majority of figures depicting the inner gonostylus). Three species of Savtshenkia have a well developed membranous crest on the posterodorsal margin of the anterior part of the inner gonostylus, viz. aspromontensis, aster, and serrulifera. The crest is dorsally rounded in aster and serrulifera (fig. 60), but it is grotesquely widening towards its slightly concave dorsal margin in aspromontensis (fig. 61).
29. - Inner gonostylus, anterior part: (0) anteriorly terminating in various ways; (1) anteriorly terminating in a prolonged nipple.

The anterior part of the inner gonostylus shows a wide range of forms in the Lunatipula group, Mediotipula, and Savtshenkia. In Savtshenkia it is often short and terminates anteroventrally in a more or less acute tip (the majority of figures showing the inner gonostylus). The species aspromontensis, aster, and serrulifera have a uniquely shaped anterior part which gradually tapers towards an anteriorly extended slender nipple-like projection (figs. 60, 61). The extension is longest in aspromontensis (fig. 61) and aster.
30. - Inner gonostylus, anterior part: (0) tip single; (1) tip bifid.

The anterior part of the inner gonostylus terminates in a single tip in the majority of Tipulidae, a situation representing the presumed groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also have the tip of the anterior part of the inner gonostylus undivided (figs. 55-63, 68, 70-75, 161, 181, 193). The species corsosignata, nielseni, and the representatives of the clade fragilina to gimmerthali differ from the other species of Savtshenkia in having a bifid anterior tip of the inner gonostylus. The tip in most of these species shows a relatively broad dorsal and finer lateroventral point (figs. 64-67). In corsosignata (fig. 69), the smaller tip lies in a more lateral position compared with that of the other species with a bifid tip.
31. - Inner gonostylus, anterior part: (0) narrowing towards pointed tip; (1) abruptly bent, almost truncate at tip.
The majority of species of Savtshenkia are characterized by a gradually curved and pointed tip of the anterior part of the inner gonostylus (lateral view; see figs. 55-72). The same situation probably is the groundplan condition of the Lunatipula group and Mediotipula. Within Savtshenkia, the species of the clade persignata to omega are distinguished from the remainder of the subgenus by the broad, almost par-allel-sided anterior part of the inner gonostylus which terminates in a rather steep front end (figs. 73-75, 181, 193).

## 32. - Inner gonostylus, base: (0) posteriorly

rounded; (1) posteromedially terminating in pointed extension.

Usually, the base of the inner gonostylus is rounded at its posterior margin in the Tipulidae, a situation representing the presumed groundplan condition of the Lunatipula group and Mediotipula. It is also found in most species of Savtshenkia (dorsal view, fig. 52, arrow). A number of species of Savtshenkia are distinguished by the presence of an acute point at the posteromedial corner of the base of the inner gonostylus (figs. 53, arrow, 54). The species concerned constitute the clade bartigiana to simulans.
33. - Inner gonostylus, lateral ridge, ventral part: (0) without extensions; (1) with one or two acute blackish pointed extensions.
The ventral part of the lateral ridge of the inner gonostylus gradually merges with the main body of the inner gonostylus in most species of Savtshenkia. The three species cheethami, goriziensis, and simulans differ from the remainder of species of Savtshenkia by the presence of one or two acute blackish points at the end of the lower part of the lateral ridge of the inner gonostylus. The points are located near midheight of


Figs. 64-75. Left inner gonostylus, lateral view. - 64, Tipula (Savtshenkia) limbata; 65, T. (S.) pechlaneri; 66, T. (S.) gimmerthali gimmerthali; 67, T. (S.) gimmerthali pteromaculata; 68, T. (S.) haennii; 69, T. (S.) corsosignata; 70, T. (S.) subsignata subsignata; $71, T$. (S.) boreosignata; 72, T. (S.) benesignata; 73, T. (S.) persignata tofina; 74, T. (S.) signata; 75, T. (S.) mannheimsi.
the body of the inner gonostylus. In goriziensis, a single point is present (fig. 54, arrow), while cheethami and simulans have two points, the posterior of which is bifid in simulans.
34. - Inner gonostylus, lateral ridge: (0) ventrally directed and terminating at about midlength of base of inner gonostylus; (1) in ventral part angularly
bent and anteriorly extended, terminating in front of base of inner gonostylus.

The lateral ridge of the inner gonostylus is ventrally directed in the majority of species of Savtshenkia. The lateral ridge of subsignata subsignata and subsignata cazorla is present as a sharp edge directed towards the anterior margin of the inner gonostylus. The ridge makes a distinct angle in its posteroventral
section near the base of the inner gonostylus (fig. 70).
Dufour (1991, character 3) recognized the presence of a ridge, on what he referred to as the median part of the inner gonostylus, as a synapomorphy of the subspecies subsignata subsignata and the then newly described subsignata cazorla and subsignata baennii. Furthermore, Dufour (1991, character 4) considered the continuity of the ventral and posterior parts of the ridge, as found in subsignata subsignata and subsignata cazorla, a synapomorphy of these two subspecies. Regarding the shape of the ridge in other species of Savtshenkia, it appears, however, that a continuous ridge should be judged a plesiomorphy in subsignata subsignata and subsignata cazorla. Consequently and contrary to Dufour, I regard the separation of the posterior and ventral parts of the ridge in haennii an autapomorphy of that species (fig. 68; see Appendix C).
35. - Inner gonostylus, posterior part: (0) variously shaped; (1) posterolaterally extended, cup-shaped, laterodorsal margin blackened and serrate.

The groundplan condition of the posterior part of the inner gonostylus in the Lunatipula group and Mediotipula is probably represented by a relatively simple and single structure. The posterior part is rounded and little developed in most species of Savtshenkia. The species confusa, eugeni, and jeekeli differ from the other species of Savtshenkia by the configuration of the posterior part. In these three species, it projects posterolaterally and carries acute processes along its blackish sclerotized laterodorsal margin (fig. 58). The species confusa has a serrate and blackened rim running medially of and parallel to the posterolateral margin (fig. 58; see Appendix C).
36. - Inner gonostylus, posterior part: (0) variously shaped; (1) a posteriorly projecting, often rather acute, structure.

As stated above, the posterior part of the inner gonostylus is moderately developed and rounded off in most species of Savtshenkia. The members of the clade odontostyla to omega are characterized by a more caudal development of the posterior part. In these species, the posterior part either is a rather slender and often somewhat acute structure (fig. 63), or a more enlarged and posteriorly produced extension (figs. 68-75, 181, 193; see also next character). A caudally produced posterior part of the inner gonostylus is also present in the species ornata (fig. 59).
37. - Inner gonostylus, posterior part: (0) a relatively small projection; (1) posteriorly extended, large.

As mentioned under the previous character, the posterior part of the inner gonostylus is usually little developed in the species of Savtshenkia. The species
benesignata, mannheimsi, omega, persignata, signata, and trinacria are distinguished among the species of the subgenus by a strikingly large posterior part which is of about the same length as the anterior part (figs. $72-75,181,193)$. The posterior part in these species is vertically orientated and lies more or less in line with the anterior part.
38. - Inner gonostylus, posterior part: (0) dorsal surface even; (1) dorsal surface corrugated.

The dorsal surface of the posterior part of the inner gonostylus is even in the majority of species of Savtshenkia, as is the probable groundplan condition of the Lunatipula group and Mediotipula. Within Savtshenkia the dorsal surface of this part of the inner gonostylus is corrugated in both persignata persignata and persignata tofina (fig. 73).
39. - Inner gonostylus, posterior part: (0) variously shaped; (1) anterior section anterodorsally extended towards anterior part of inner gonostylus.

As noted under character 37, a number of species of Savtshenkia have a caudally produced and large posterior part of the inner gonostylus. The extended posterior part of the inner gonostylus is low in its anterior half in benesignata, persignata persignata, and persignata tofina (figs. 72,73 ), as it is in the other species of Savtshenkia. In the species mannheimsi, omega, signata, and trinacria it is anterodorsally protruding (figs. 74, 75, 181, 193).
40. - Inner gonostylus: (0) variously shaped; (1) similarly shaped, with high anterior part and blackish sclerotized triangular projection posterolaterally.

The inner gonostylus shows a wide variety of forms throughout Savtshenkia. The two subspecies rufina rufina and rufina maderensis display a remarkable similarity in the structure of the inner gonostylus which differs most notably from that of other species of Savtshenkia by the shape of the anterior part and the location of the posterolateral projection (fig. 55).
41. - Inner gonostylus: (0) variously shaped; (1) similarly shaped, anterior part anterodorsally straightened, gradually narrowing towards bifid tip.

Throughout Savtshenkia a wide variety of forms of the inner gonostylus can be found. The three recognized subspecies of gimmerthali can be separated from the other species of the subgenus by the unique shape the inner gonostylus, of which the anterior part gradually narrows towards a bifid tip (figs. 66, 67).
42. - Inner gonostylus: (0) variously shaped; (1) similarly shaped, with long sloping caudodorsal margin.


Figs. 76-99. - 76-88, left outer gonostylus, lateral view; 76, Tipula (Savtshenkia) grisescens, 77, T. (S.) hartigiana; 78, T. (S.) lundbladi; 79, T. (S.) multipicta; 80, T. (S.) jeekeli; 81, T. (S.) aspromontensis, 82, T. (S.) fragilis, 83, T. (S.) gimmerthali gimmerthal; 84, T. (S.) gimmerthali pteromaculata; $85, T$. (S.) nielseni; 86, T. (S.) sardosignata; 87, T. (S.) signata; $88, T$. (S.) mannheimsi; 89-99, sperm pump and appendages; 89, T. (S.) subnodicornis, lateral view; 90, T. (S.) baltistanica, lateral view; 9193, T. (S.) alpium, 91, lateral view; 92, compressor apodeme, posterior view; 93, anterior view; 94-96, T. (S.) jeekeli, 94, lateral view; 95, compressor apodeme, posterior view; 96, anterior view; 97, T. (S.) confusa, lateral view; 98, T. (S.) graciae, lateral view; 99, T. (S.) boreosigata, lateral view.

The species boreosignata and eleonorae are distinguished among the species of Savtshenkia by their uniquely shaped and very similar inner gonostyli. The overall structure is rather long, with a well developed anterior part and a long caudodorsal margin which gradually slopes towards the low posterior part (fig. 71).
43. - Outer gonostylus: (0) anterior margin convex
or slightly and evenly concave; (1) concave, with distinct bend at about two-fifth of length from base.

The anterior margin of the outer gonostylus is either entirely convex or evenly concave in its lower half in the presumed groundplan condition of the Lunatipula group and Mediotipula, and in the majority of species of Savtshenkia (figs. 76, 78-88, 160, 180, 192). The three species atlas, breviantennata, and hartigiana have the anterior margin of the outer
gonostylus angularly concave near two-fifth of its height (fig. 77, arrow).
44. - Outer gonostylus: (0) upright structure; (1) inclined forward.

The outer gonostylus in Tipulidae usually is an upright structure, as is the presumed groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia are also distinguished by the presence of a vertically orientated outer gonostylus (figs. 76, 77, 80, 82-88, 160, 180, 192). The outer gonostylus of the species aspromontensis, lundbladi, and multipicta leans over anteriorly (figs. $78,79,81)$. The actual shapes of the outer gonostyli of the three pertaining species differ notably. The base of the outer gonostylus is low and elongate in lundbladi (fig. 78) and multipicta (fig. 79), whereas it is rather short in aspromontensis (fig. 81).
45. - Outer gonostylus: (0) anterior part rather broad; (1) slender and elongate.

In general, the outer gonostylus is present as a single oblong and rather broad structure in the Tipulidae, a situation which probably represents the groundplan condition of this character for the Lunatipula group and Mediotipula. A broad outer gonostylus is found in about half of the species of Savtshenkia (figs. 76-81, 160). The species of the clade fragilis to omega are distinguished by the presence of a rather slender and elongate anterior substructure of the outer gonostylus (figs. 82, 83, 85-88, 180, 192). A similar outer gonostylus with slender anterior section is found furthermore in the species eugeni and ornata. The subspecies gimmerthali pteromaculata, which is included in the clade fragilis to omege, has a relatively wide anterior part of the outer gonostylus (fig. 84).
46. - Outer gonostylus: (0) base variously shaped, usually relatively narrow; (1) broad, laterally concave and blackish sclerotized.

As noticed under the previous character, the outer gonostylus generally is an oblong structure in the Tipulidae. The base of the outer gonostylus is usually relatively narrow compared with the length of the structure, a situation present in the majority of species of Savtshenkia (figs. 76, 77, 80, 82-84, 87, 88, $160,180,192)$. The species nielseni (fig. 85) and sardosignata (fig. 86) differ from the remainder of Savtshenkia by the presence of a large base of the outer gonostylus. The base is laterally concave, partly blackish sclerotized and carries a strong posterior projection.
47. - Sperm pump, endophallic lumen: (0) bulbous, ventrally expanded; (1) small.

Throughout the Tipulidae, the sperm pump has a rather large and bulbous endophallic lumen, a situation which probably represents the groundplan condition of the Lunatipula group and Mediotipula. Within Savtshenkia a similar condition is found in the species alpha, grisescens, ignobilis, subnodicornis, and the species of the clade hartigiana to simulans (figs. 89, 163). The majority of species of Savtshenkia, however, have an endophallic lumen that is hardly ventrally expanded (figs. 5, 90, 91, 94, 97, 98, 183, 195). The species concerned include the members of the clade baltistanica to omega.
48. - Sperm pump, compressor apodeme: (0) a flattened blade; (1) posteriorly provided with U shaped or V-shaped brace.

The groundplan condition of the compressor apodeme of the Lunatipula group and Mediotipula probably consists of a flattened blade with a more or less distinct dorsomedial emargination. Within Savtshenkia a similar compressor apodeme is found in the species that range from alpha to macaronesica (figs. 92, 93, 164). The other species of Savtshenkia are distinguished by the presence of a U-shaped or Vshaped support on the posterior side of the compressor apodeme (fig. 95, arrow, 96, 184, 196). The species involved constitute the clade jeekeli to omega.
49. - Sperm pump: (0) ventral wall convex; (1) ventral wall flattened or concave.
In Tipulidae, the body of the sperm pump usually is a bulbous structure which has a convex ventral wall. This situation probably reflects the groundplan condition of the Lunatipula group and Mediotipula and is found in about two-fifth of the species of Savtshenkia (the range alpha to macaronesica; frontal view, figs. 93, 164). Within Savtshenkia the bulb of the sperm pump has a flat or concave ventral wall in most species of the clade jeekeli to omega (figs. 96, 184, 196). There are, however, a few species within this latter clade which have a convex ventral wall of the sperm pump, viz. aspromontensis, eugeni, and interserta.
50. - Sperm pump, posterior apodemes: ( 0 ) well developed and posteriorly or posterodorsally directed; (1) elongate and anteriorly extended.
In Tipulidae, the posterior apodemes of the sperm pump are generally well developed, rather broad and posteriorly or posterodorsally directed. This character state probably represents the groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also display this character state (figs. 5, 89-91, 94, 163, 183, 195). In contrast, the members of the clade confusa to aster are distinguished by the presence of elongate and anteriorly extended posterior apodemes. The tips of the posterior


Figs. 100-112. - 100, 101, sp2 and part of tergite 9-10, posteromedial view; 100, Tipula (Savtshenkia) odontostyla; 101, T. (S.) limbata; 102-112, male sternite 8 , posterior margin; 102, T. (S.) grisescens, ventral view; 103, T. (S.) lundbladi, ventral view; 104, T. (S.) rufina rufina, posterior view; 105, T. (S.) alpium, ventral view; 106, T. (S.) obsoleta, ventral view; 107, T. (S.) boloptera, ventral view; 108, T. (S.) confusa, ventral view; 109, T. (S.) aster, ventral view; 110, T. (S.) limbata, ventral view; 111, T. (S.) pechlaneri, ventral view; 112, T. (S.) invenusta invenusta, ventral view.
Abbreviations: sp2: lateral part of genital bridge; tg9-10: tergite 9-10.
apodemes in these species lie alongside the compressor apodeme (figs. 97, 98).
51. - Sperm pump, posterior apodemes: (0) well developed, broad; (1) slender.
As noticed under the previous character, the posterior apodemes of the majority of species of Savtshenkia are well developed and broad (figs. 5, 89-
$91,94,98,163,183,195)$. Three species are distinguished from the remainder of the subgenus by their long and slender posterior apodemes, viz. confusa, interserta, and ornata (fig. 97). The posterior apodemes of these three species are anteriorly extended (see previous character).
52. - Sperm pump: (0) general shape of the standard
tipulid type; (1) general shape different, partly heavily sclerotized.

The sperm pump in Tipulidae usually consists of a bulbous main body, which is surrounded in the horizontal plane by a pair of anterior and a pair of posterior apodemes, and perpendicular and dorsal to these a single compressor apodeme. The anterior and posterior apodemes are attached to the main body of the sperm pump at about midheight and the aedeagus leaves the sperm pump in between the anterior apodemes. This configuration doubtlessly represents the groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also show a sperm pump of this form (figs. 5, 89-98, $163,164,183,184,195,196)$. However, the species boreosignata and eleonorae possess a peculiarly built sperm pump, which has the heavily sclerotized main body lying just dorsal of (boreosignata, fig. 99) or on (eleonorae) a membranous disc-like structure which contains the remnants of the anterior and posterior apodemes. The heavily sclerotized compressor apodeme seems to be fused with the main body of the pump. In boreosignata, the short aedeagus appears to be attached to the ventral side of the main body, after which it makes a loop at the level of the disc-like structure containing the anterior and posterior apodemes. Compared with the sperm pump of the other species of Savtshenkia, the whole structure is small in both boreosignata and eleonorae.
53. - Sp2: (0) uniformly and moderately sclerotized; (1) medially blackish sclerotized.

Throughout the Tipulidae, sp2 generally is uniformly sclerotized. This situation is found in the Lunatipula group and in Mediotipula and represents the groundplan state of this character for both taxa. All species of Savtshenkia as recognized in this paper have the medial margin of sp 2 blackish sclerotized (figs. 100, 101, 159, 179, 191). The only other Tipulidae in which a similar situation was studied are species of the genus Nephrotoma Meigen and of the subgenus Tipula (Schummelia Edwards). In Nephrotoma, a partly blackish sclerotized sp 2 can be found in for instance the Palaearctic aurantiocincta Alexander, biarmigera Alexander, medioproducta Alexander and the Oriental citricolor Alexander, integra Alexander, progne Alexander, subumbonis Alexander, and umbonis Alexander. In the pertaining species of Nephrotoma, the general shape of sp2 differs from that of the species of Savtshenkia in that sp2 is short and broad, anteriorly notched, and rather deeply concave. All species of Schummelia studied (viz. abrensi Savchenko, butzi Edwards, variicornis Schummel, yerburi Edwards, zernyi Mannheims, and zonaria Goetghebuer) have the anteromedial margin of sp2 more or less darker sclerotized than the re-
mainder of the structure. The general shape of sp2 in the species of Schummelia is similar to the elongate form as found in the majority of species of Savtshenkia. The phylogenetic positions of the pertaining species of Nephrotoma and Schummelia within the respective taxa are still unestablished. Except for the similar modification of sp 2 , there are at the moment, however, no indications to suggest that either Nephrotoma or Schummelia, or both combined, represent(s) the closest relative of Savtshenkia (see also next character).
54. - Sp2: (0) dorsally separate from tergite 9-10; (1) dorsally fused with ventromedial part of tergite 9-10.

Sp2 and tergite 9-10 are separate from each other in the greater majority of Tipulidae, a situation considered the groundplan condition of the Lunatipula group and Mediotipula. More than half of the species of Savtshenkia also show this character state (fig. 100, arrow). In the species of the clade fragilina to omega, sp 2 is fused with the ventromedial sclerotization of tergite 9-10 (fig. 101). Both structures are firmly connected over their entire width. Outside Savtshenkia, I found a similar character state in species of the subgenus Tipula (Schummelia) only. In the examined species of Schummelia (see previous character), sp2 and the ventral sclerotization of tergite 9-10 are approximated. Some of these species show a partial fusion of sp2 and tergite 9-10 (viz. ahrensi, variicomis, and yerburi). In zernyi, both structures are more firmly connected. The studied species of Schummelia all have a western Palaearctic distribution and represent only a minor fraction of the subgenus which contains about 75 species distributed further in the Afrotropical, Nearctic and especially Oriental Regions. Fusion of tergite 9-10 with sp2 probably evolved independently in Schummelia. As stated under the preceding character, I do not consider Schummelia a potential sister group of Savtshenkia.
55. - Sternite 8, posterior margin: (0) unmodified; (1) caudally extended.

An unmodified sternite 8 in Tipulidae consists of a relatively short structure of which the posterior margin is evenly convex or slightly concave and lacks any trace of spinosity. This situation probably represents the groundplan condition of this character for the Lunatipula group and Mediotipula, In a number of species of Savtshenkia, a modification of the posterior margin of sternite 8 is found, which generally consists of a posterior elongation of the structure (figs. 103, $105-110,112-117,173,177,178,185,189,190)$. Elongation of sternite 8 is usually combined with the presence of spines or strong setae along the posterior margin (character 57). A caudally produced posterior


Figs. 113-128. - 113-117, male sternite 8, posterior margin; 113, T. (S.) cyrnosardensis, posterior view; 114, 115, T. (S.) haenniל, 114, posterior view; 115, ventral view; 116, T. (S.) signata, posterior view; 117, T. (S.) mannheimsi, posterior view; 118120, male sternite 9, anteroventral appendage, dorsal view; 118, T. (S.) sordidipes, 119, T. (S.) eugeni; 120, T. (S.) limbata; 121-128, male tergite $9-10 ; 121,122, T$. (S.) grisescens, 121 , dorsal view; 122 , posterior view; $123, T$. (S.) hartigiana, dorsal view; 124, T. (S.) atlas, dorsal view; 125, T. (S.) goriziensis, dorsal view; 126, T. (S.) cheethami, dorsal view; 127, T. (S.) ignobilis, dorsal view; 128, T. (S.) subnodicornis, dorsal view.
margin of sternite 8 is found in the species lundbladi, multipicta, rufina, and those of the clade alpium to omega. Within the latter clade a short sternite 8 is present in aspromontensis, gimmerthali, interserta, pechlaneri, sordidipes, and venerabilis (see fig. 111).
56. - Sternite 8, posterior margin: (0) variously
shaped; (1) with V-shaped ventral carina.
Sternite 8 is present as an evenly curved sclerite in the presumed groundplan condition of the Lunatipula group and Mediotipula. Within Savtshenkia a variety of forms of the posterior margin of sternite 8 is found. The two subspecies rufina rufi$n a$ and rufina maderensis are distinguished from the
remainder of the subgenus by the presence of a midventral $V$-shaped carina in the posterior part of sternite 8 (posterior view, fig. 104).
57. - Sternite 8, posterior margin: (0) unarmed; (1) provided with strong setae or black spines.

In most Tipulidae, sternite 8 is unarmed, a situation considered to represent the groundplan state of this character for the Lunatipula group and Mediotipula. About half of the species of Savtshenkia have the posterior margin of sternite 8 provided with strong setae or short spines (figs. 105-109, 113-117, $177,178,189,190$ ). The number of spines or setae and their location varies among the different species of Savtshenkia. The species with an armed posterior margin of sternite 8 include baltistanica, multipicta, and most species of the clade alpium to omega. Within the latter clade a number of species show an unarmed sternite 8, viz. interserta, macaronesica, orna$t a$, and the species of the clade fragilina to invenusta (see figs. 110-112).
58. - Sternite 8, posterior margin: (0) variously shaped; (1) with medial tongue-like extension, posteriorly concave.

As stated above, a number of species of Savtshenkia have a caudally produced posterior margin of sternite 8. The species holoptera, obsoleta, and pagana are distinguished among these by the peculiarly shaped posterior extension of sternite 8, which appears as a tongue-like blade which is caudally more or less emarginate (figs. 106, 107). In most specimens of pagana, the caudal margin of the posterior extension is nearly straight, in the species holoptera and obsoleta it is distinctly concave.
59. - Sternite 8, posterior margin: (0) unarmed or armed with more than 10 spinous setae; (1) number of spinous setae less, ranging from seven to one.

The number of strong setae or spines on the posterior margin of sternite 8 in the species of Savtshenkia which have this structure armed usually is (considerably) more than 10 (figs. 105, 113-117, 177, 178, 189, 190). In the species obsoleta, there are about 13 to 15 spines on the posterior margin of sternite 8 (fig. 106). The species holoptera and pagana have less than 10 spines along the posterior margin of sternite 8 . Sternite 8 of pagana usually has about four to eight spinous setae, in holoptera the number varies between one and three (fig. 108; see Eiroa, 1987).
60. - Sternite 8, posterior margin: (0) medial membranous area absent, small or, if large, without rim of longish setae; (1) membranous area broadly V-shaped, anteriorly extending, its margins set with long slender setae.

As noticed above, the groundplan condition of sternite 8 for the Lunatipula group and Mediotipula probably consists of an unmodified sclerite. The posterior margin of the sternite in this situation is uniformly sclerotized. In Savtshenkia, sternite 8 generally has no membranous area at its posterior margin or the membranous area is restricted to a narrow zone (figs. 102, $103,105-107,111,112,115,158,177,189)$. The three species fragilina, limbata, and subvafra are distinguished from the remainder of Savtshenkia by a large and V-shaped membranous area which is bordered by a fringe of thickly set longish slender setae (fig. 110). The species aster and confusa are the only other species of Savtshenkia that also have a relatively large membranous area on the posterior margin of sternite 8 . In as ter, a V-shaped membranous area is present in between the caudolateral extensions, the extensions themselves are provided with a few black spines (fig. 109; see Appendix C). The species confusa has a broad and rounded membranous zone (fig. 108; see Appendix C). In both species, the margin of the membranous area is surrounded by scattered setae only.
61. - Sternite 8, posterior margin: (0) ventrally directed medial extension absent; (1) unarmed medial extension present rostrally of lateral extensions, or at this position with sclerotized edge.

In the majority of species of Savtshenkia, the posterior margin of sternite 8 medially either is unmodified or produced into a rather broad lobe provided with spines or strong setae (figs. 102, 103, 105-111, 158). An unarmed medial extension rostrally of a pair of lateral extensions on the posterior margin of sternite 8 can be found in the three subspecies of invenusta and in the species of the clade nielseni to omega. The size, shape and sclerotization of the medial extension varies among the species involved. In the subspecies of invenusta, and in mannheimsi, omega, and trinacria the medial extension is relatively small and moderately sclerotized (figs. 112, 117, 177, 178, 189, 190), it is large and partly membranous in nielseni, whereas it is more pronounced and heavily sclerotized in most other species involved (figs. 114, 116). It should be noted, however, that the species benesignata, boreosignata, and eleonorae lack a protruding medial extension. In these three species, a sclerotized rim marks the position of the medial extension.
62. - Sternite 8, posterior margin: (0) extension(s) absent or dorsoventrally flat; (1) lateral extensions appearing as bulbous structures, densely set with spines.

The extensions of sternite 8 are relatively flat and follow the curve of sternite 8 when seen in posterior view in about half of the species of Savtshenkia with a modified sternite 8. Most members of the clade cyr-


Figs. 129-141. Male tergite 9-10.- 129, T. (S.) baltistanica, dorsal view; 130, T. (S.) elgonensis, posterior margin, dorsal view, after celluloid preparation holotype; 131, 132, T. (S.) aberdareica, 131, posterior view; 132, dorsal view; 133, T. (S.) sordidipes, dorsal view; 134, T. (S.) serrulifera, posterior view; 135, T. (S.) aspromontensis, posterior view; 136, T. (S.) invenusta invenusta, dorsal view; 137, T. (S.) nielseni, dorsal view; 138, T. (S.) boreosignata, dorsal view; 139, T. (S.) persignata tofina, posterior view; 140, T. (S.) signata, posterior view; 141, T. (S.) mannheimsi, posterior view.
nosardensis to omega are characterized by the presence of bulbously swollen lateral extensions of sternite 8 . In the majority of these species, the lateral extensions are thickly set with black spines (figs. 113-116), but in the species boreosignata only a few spines on the lateral lobes are found. Moreover, the three species mannheimsi, omega, and trinacria have a relatively flat pair of lateral extensions (figs. 117, 178, 190; character 65).
63. - Sternite 8, posterior margin: (0) lateral extensions absent, or, if present, dorsally not connected; (1) dorsally connected by a sclerotized bridge.

In most species of Savtshenkia which carry a pair of lateral extensions on the posterior margin of sternite 8 , the extensions are dorsally separated by a membranous area (figs. 113, 116, 117, 178, 190). The species and subspecies corsosignata, haennii, subsignata subsig-
nata, and subsignata cazorla are distinguished by the presence of a sclerotized dorsal bridge that connects the lateral extensions of sternite 8 (fig. 114).
64. - Sternite 8, posterior margin: (0) variously shaped; (1) in between medial and lateral extensions a pair of medially directed projections.

The ventral margin of sternite 8 in between the medial and lateral extensions is straight or evenly curved in most species of Savtshenkia with a threelobed sternite 8 . As an extra feature, corsosignata, baennii, subsignata subsignata, and subsignata cazorla have an acute and dorsomedially directed sclerotized projection on each side of the medial extension (fig. 115, arrow).
65. - Sternite 8, posterior margin: (0) variously shaped; (1) lateral extensions dorsoventrally flattened, posterior margin straight, spines confined to posterior margin.

The bulbously swollen lateral extensions of sternite 8 in most species of the clade cyrnosardensis to omega are more or less circular in cross section and covered with spines on their posterior half, dorsally as well as ventrally (figs. 113-116). The species mannheimsi, omega, and trinacria have the lateral extensions of sternite 8 dorsoventrally flatter and straight along their posterior margins (figs. 117, 177, 178, 189, 190). The spines in these three species are concentrated along the posterior margin of the sternite.
66. - Sternite 9, midventral anterior appendage: (0) apically rounded; (1) apically bifid.

Sternite 9 often carries a more or less developed anterior midventral appendage in the Tipulidae. When present, this appendage has a rounded anterior margin, as it has in the majority of species of Savtshenkia (figs. 118, 120). In the species aspromontensis, aster, eugeni, and serrulifera the anterior appendage has a bifid apex (fig. 119).
67. - Sternite 9, midventral anterior appendage: (0) a plain structure; (1) dorsally provided with medial carina.

In the presumed groundplan condition of the Lunatipula group and Mediotipula, the ventromedial part of sternite 9 , including the anterior appendage, is a dorsoventrally flattened structure. The majority of species of Savtshenkia show an anterior midventral appendage of sternite 9 which looks like a simple tongue-like extension without further modifications (fig. 118). A different form of this structure is present in the species of the clade fragilina to omega, where the appendage is provided with a medial carina on its dorsal side (fig. 120). Although the height of the carina varies among the pertaining species, it is usually
well developed and clearly visible. A similar anterior appendage was also found in the species hartigiana. Within the clade fragilina to omega only pechlaneri, sardosignata, and subsignata cazorla lack a distinct medial carina.
68. - Tergite 9-10, posterior margin: (0) extensions variously shaped; (1) narrowing toward caudally produced tip.

In Tipulidae, tergite 9-10 often has a pair of low and rounded extensions at its posterior margin, a situation that probably represents the groundplan condition of the Lunatipula group and Mediotipula. Within Savtshenkia, only a few species show similar extensions (figs. 121, 122, 125, 126, 127, 156, 157). The species concerned lie in the range alpha to ignobilis, with the exception of the species atlas, breviantennata, and bartigiana. In the three latter species, the extensions are caudally produced and constricted before their tips. The medial margins of the extensions in these species are slightly concave in dorsal view (figs. 123, 124).

## 69. - Tergite 9-10, posterior margin: (0)

posteroventral sclerotization uninterrupted along posterior margin; (1) separated into two parts by medial incision.

Tergite 9-10 in Tipulidae often has a sclerotized rim running along the ventral side of the posterior margin. An uninterrupted rim probably represents the groundplan state of this character for the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also have a complete and broad posteroventral sclerotized rim along the posterior margin of tergite 9-10 (figs. 121, 123, 125-129, etc., after macerated specimens). Seen in dorsal view, this rim seems to be intersected by the narrow and deep medial incision of tergite $9-10$ in the species atlas and breviantennata (fig. 124).
70. - Tergite 9-10, posterior margin: (0) tips of extensions broadly separate; (1) close together, separated by a narrow gap.

A broad emargination or medial notch usually separates the extensions on the posterior margin of tergite 9-10 in Tipulidae, a situation reflecting the presumed groundplan condition of the Lunatipula group and Mediotipula. In the majority of species of Savtshenkia, the posterior extensions of tergite 9-10 are separated by a broad V-shaped or U-shaped emargination (figs. 121-125, 127, 129, etc.). In cheethami and simulans, the extensions approach each other medially and are separated by a rather deep and narrow fissure (fig. 126).
71. - Tergite 9-10, posterior margin: (0) extensions


Figs. 142-151. - 142, 143, genital fork, dorsal view; 142, Tipula (Savtshenkia) hartigiana; 143, T. (S.) atlas, 144, 145, ovipositor, lateral view; 144, T. (S.) grisescens, $145, T$. (S.) bartigiana; 146,147 , spermatheca; $146, T$. (S.) bartigiana; 147, T. (S.) villeneuvi; 148, 149, female sternite 8 and hypogynial valves, ventral view; 148, T. (S.) grisescens, 149, T. (S.) ignobilis, 150, 151, female sternite 9, medial part, ventral view; 150, T. (S.) jeekeli; 151, T. (S.) obsoleta.
posteriorly directed and rounded at tip; (1) laterally curved and pointed at tip.
As noted above, the groundplan condition of tergite 9-10 for the Lunatipula group and Mediotipula probably shows a pair of broad and rounded extensions on the posterior margin of the tergite. The majority of species of Savtshenkia, however, have the tips of the extensions of tergite $9-10$ pointed and outwardly curved when seen in dorsal view (figs. 128-$130,137,139-141,174,175,186,187)$.
72. - Tergite 9-10, posterior margin: (0) provided with black spines; (1) spines absent.

The section of the family Tipulidae to which Savtshenkia and its outgroups belong, is, among other things, characterized by the presence of black spines on the posterior margin of tergite 9-10. Although such spines are absent in the majority of species of the Lunatipula group, there are several species within the group which have the posterior margin of tergite 9-10 armed with spines. For this reason I, putatively, consider this latter state the groundplan
condition of the Lunatipula group. All species of Mediotipula and the majority of species of Savtshenkia also show a spinose posterior margin of tergite 9-10 (figs. 121-128, 133-136, 139, 140, 187). Within Savtshenkia, total absence of spines on the posterior margin of tergite $9-10$ is found in the species alpha (figs. 156, 157) and the species of the clade baltistanica to chrysocephala (figs. 129-132). The species glaucocinerea has a few knob-like protuberances on the lower margin of the extensions of tergite 9-10, but lacks distinct spines. Spines along the ventral margin of the extensions of tergite 9-10 are absent in the species mannheimsi and omega, but in both species a few small spines are present in the dorsal part of the medial cleft (figs. 141, 175).
73. - Tergite 9-10, posterior margin: (0) extensions variously shaped, usually long and laterally directed; (1) extensions present as short spineless points.

The laterally directed extensions of the species of the clade subnodicornis to omega usually terminate in slender and elongate points (figs. 128-130, 137-141,

174, 175, 186, 187). A number of Afrotropical species of Savtshenkia are characterized by the presence of short and acute extensions, viz. the species aberdareica, akeleyi, asbolodes, chrysocephala, draconis, and hancocki (figs. 131, 132).
74. - Tergite 9-10, posterior margin: (0) spines confined to the extensions; (1) spines present above medial notch.

As is usual in Tipulidae which have tergite 9-10 provided with spines, the spines are concentrated on the posterior margin of the tergite in the majority of species of Savtshenkia (figs. 121-128, 139, 140, 175, 187). Spines are present also in the area above the medial notch of tergite 9-10 in the species aspromontensis, aster, eugeni, odontostyla, serrulifera, sordidipes, and venerabilis (figs. 133-135). In the latter five species, there are about 10 spines in this area (figs. 133, 134), while aspromontensis and aster carry 20 to 30 spines above the notch (character 78).
75. - Tergite 9-10, posterior margin: (0) variously shaped; (1) broadly $V$-shaped emarginate with rounded corners.

The Kashmirian species sordidipes and venerabilis differ from all other species of Savtshenkia by the shape of the posterior margin of tergite 9-10. Tergite 9-10 in both species has a broad V-shaped emargination with a small median notch, while the posterior corners are broadly rounded off (fig. 133). The species sordidipes and venerabilis resemble each other very much, including in details of the male terminalia. Differences are found in the colour of the legs and abdomen and in the height of the anterior part of the inner gonostylus.
76. - Tergite 9-10, posterior margin: (0) single pair of extensions present; (1) two pairs present.

As in most other Tipulidae, the standard number of extensions on the posterior margin of tergite 9-10 in Savtshenkia is two (figs. 121-127, 129-133, 136141, 156, 157, 174, 175, 186, 187). The species serrulifera, aspromontensis, and aster differ from this by having a medial and a sublateral pair of extensions. Both pairs of extensions are provided with black spines in aster and serrulifera (fig. 134), in aspromontensis the sublateral pair is unarmed (fig. 135).
77. - Tergite 9-10, posterior margin: (0) dorsal surface of extensions rounded; (1) flattened.

In general, the extensions on the posterior margin of tergite 9-10 are rounded on their dorsal surface in Tipulidae, a situation found also in the majority of species of Savtshenkia. As a special trait, the species aspromontensis and aster have the medial pair of extensions of the posterior margin of tergite 10 flattened.

The dorsal surface of the medial extensions of both species is glossy black.
78. - Tergite 9-10, posterior margin: (0) dorsal surface in between extensions unarmed or armed with up to 10 spines; (1) dorsal surface in between extensions armed with 20 to 30 spines.

As discussed under character 74 , a few species of Savtshenkia have the area above the medial notch of tergite 9-10 armed with up to 10 spines (fig. 133, 134). The number of spines is higher in the species aspromontensis and aster. In aspromontensis, there are about 30 spines above the medial notch (fig. 135), in aster there are about 20.
79. - Tergite 9-10, posterior margin: (0) extensions present; (1) margin straightened, with small medial notch.

As discussed above, the majority of species of Savtshenkia have a pair of lateral extensions on the posterior margin of tergite 9-10, either low and rounded or elongate, diverging and pointed (figs. 128-130, 134, 135, 137-141, 174, 175, 186, 187). The species interserta, the three subspecies of invenus$t a$, and the species subvafra are characterized by a straightened posterior margin of tergite 10 (fig. 136). In subvafra, a pair of small posteroventral projections flank the narrow medial notch; such projections are absent in interserta and invenusta.
80. - Tergite 9-10: (0) broader than long, or about as long as broad; (1) elongate, distinctly longer than broad.

In Tipulidae, tergite 9-10 usually is a relatively short and broad structure, a situation which probably reflects the groundplan condition of the Lunatipula group and Mediotipula. The majority of species of Savtshenkia also show a similar character state (figs. 121, 123-129, 132, 133, 136, 156). In the pertaining species, tergite $9-10$ is broader than long and in most of the species shows a broad medial membranous zone. The species and subspecies of the clade nielseni to omega differ from this in having an elongate tergite 9-10 which is markedly longer than broad (fig. 137, $138,174,186$ ). The medial membranous area of tergite 9-10 in these taxa is restricted to a narrow zone, with the exception of the species boreosignata which has a relatively wide membranous area (fig. 138).
81. - Tergite 9-10, posterior margin: (0) extensions variously shaped; (1) rather broad, ventral margin straightened, terminating in an acute lateral tip.

As noted above (character 71), most species of Savtshenkia have the extensions on the posterior margin of tergite 9-10 laterally directed and pointed at the tip. In the majority of these species, the ventrome-
dial margin of the extensions is distinctly concave when seen in posterior view (fig. 139). The species corsosignata, mannheimsi, omega, signata, and trinacria are characterized by rather broad extensions of tergite 9-10 of which the ventral margin appears to be almost straight in posterior view (figs. 140, 141, 175, 187).

## Female terminalia

82.     - Genital fork: (0) broadest width in posterior half; (1) broadest width in anterior half.

In general, the genital fork in Tipulidae is a rather slender structure that widens in its apical posterior part. This situation probably represents the groundplan condition of the Lunatipula group. Within Mediotipula, the genital fork is slender throughout, with the widest portion lying in the anterior part. The general structure of the genital fork is rather uniform in the species of Savtshenkia, being broad with its greatest width lying in the anterior half (figs. 142, 169, 201; lower side in illustrations). The species atlas and breviantennata are the only studied representatives of Savtshenkia which have the broadest part of the genital fork situated in the posterior section (fig.

143; upper side in illustration).
83. - Hypogynial valve, dorsal margin: (0) not blackish sclerotized; (1) blackish sclerotized.
The hypogynial valves are usually uniformly sclerotized in Tipulidae and this condition probably represents the groundplan state of this character for the Lunatipula group and Mediotipula. All species of Savtshenkia of which the female was examined have the dorsal margin of the hypogynial valve blackish sclerotized.
84. - Ovipositor: (0) well developed, elongate; (1) short.

The ovipositor in Tipulidae generally is well developed and has elongate cerci and hypogynial valves, a situation which represents the groundplan condition of the Lunatipula group and Mediotipula. The same state occurs in the majority of species of Savtshenkia (figs. 145, 165, 197). Cerci and hypogynial valves are short in the females of grisescens, holoptera, obsoleta, pagana, and subnodicornis (fig. 144).
85. - Spermathecae: (0) globular; (1) elongate.

Table 2
Theowald's (1957, 1967) arrangement of species groups of Tipula (Savtshenkia) based on larvae and pupae (1957), and pupae (1967).

|  | 1957 | 1967 |
| :---: | :---: | :---: |
| rufina group | cheethami <br> rufina | cheethami <br> rufina |
| subnodicornis group | goriziensis <br> (not mentioned) <br> subnodicornis | (not filed) <br> grisescens <br> subnodicornis |
| subvafra group | atlas (as breviantennata) | atlas (as breviantennata) |
| signata group | alpium <br> confusa <br> nielseni <br> obsoleta <br> (not mentioned) <br> (not mentioned) <br> signata <br> staegeri | alpium <br> confusa <br> (not mentioned) <br> obsoleta <br> odontostyla <br> pagana <br> signata <br> staegeri |

## Table 3

Division of the Palaearctic species and subspecies of Tipula (Savtshenkia) into species groups according to Savchenko (1961) and Theowald (1973, 1978). Theowald's group numbering is given in parentheses, a '?' denotes a species placed by Theowald under restriction in the pertaining group.
Notes: 1: placed in miscellaneous group by Savchenko, under restriction in convexifrons group by Theowald; 2: referred to by Theowald only: 3: placed in rufina group by Savchenko, in pagana group by Theowald; 4: placed in miscellaneous group by Savchenko. under restriction in pagana group by Theowald; 5: placed in cheethami group by Savchenko, in alpium group by Theowald: 6: placed in rufina group by Savchenko, under restriction in obsoleta group by Theowald; 7: referred to by Savchenko only,

| Savchenko 1961 | Theowald 1973, 1978 | Species and subspecies |
| :---: | :---: | :---: |
| subnodicornis group | subnodicornis group (1) | postposita <br> subnodicornis |
|  | convexifrons group (2) | convexifrons interserta ? venerabilis |
| limbata group | grisescens group (3) | goriziensis ${ }^{2}$ <br> grisescens (as macrocera in Savchenko 1961) <br> villeneuvii ${ }^{2}$ (as goriziensis villenewvii) |
|  | invenusta group (4) | invenusta invenusta invenusta subinvenusta subvafra |
|  | limbata group (5) | limbata |
|  | pagana group (6) | baltistanica ${ }^{3}$ <br> glaucocinerea <br> holoptera (as pagana holoptera) <br> ? kiushiuensis ${ }^{2}$ <br> ? koreana ${ }^{4}$ <br> ? minuscula ${ }^{2}$ (as nana) <br> nivalis <br> pagana (as pagana pagana) |
|  | gimmerthali group (7) | gimmerthali pechlaneri ${ }^{2}$ |
| rufina group | rufina group (8) | rufina rufina <br> rufina maderensis |
|  | alpium group (11) | alpium <br> subalpium macaronesicas baltistanica ${ }^{3}$ odontostyla ${ }^{6}$ |
| cheethamigroup | atlas group (9) | atlas ${ }^{2}$ <br> breviantennata cheethami macaronesicas |
| marmorata group | obsoleta group (10) | aspromontensis ${ }^{2}$ <br> lundbladi ${ }^{2}$ <br> ? mohriana <br> multipicta ${ }^{2}$ <br> obsoleta <br> ? odontostyla ${ }^{6}$ <br> serrulifera |
|  | marmorata group (12) | confusa (as marmorata) eugeni ${ }^{2}$ jeekeli |
|  | staegeri group (13) | boreosignata ${ }^{2}$ <br> nielseni ${ }^{2}$ <br> sardosignata <br> staegeri <br> subsignata |
|  | signata group (14) | benesignata <br> mannheimsi ${ }^{3}$ <br> persignata persignata <br> persignata tofina <br> signata |
| miscellaneous group |  | koreana * <br> letifera <br> venerabilis ${ }^{1}$ |

The shape of the (three) spermathecae in Tipulidae is usually spherical to somewhat ovoid. The groundplan condition of this character for the Lunatipula group and Mediotipula is probably represented by this state. The majority of females of Savtshenkia examined also show this spermathecal form (figs. 146, 170, 202). Only the species cheethami, goriziensis, simulans, and villeneuvii are characterized by the presence of elongate spermathecae (fig. 147). It should be noted that the female of tulipa is not yet known.
86. - Sternite 8: (0) ventromedial membranous area near base of hypogynial valves $V$-shaped, encompassing base of medial margins of hypogynial valves; (1) posteriorly terminating in between hypogynial valves.

The ventromedial membranous area of sternite 8 broadens posteriorly and encompasses the bases of the hypogynial valves medially in the species of the Lunatipula group and Mediotipula, thus probably representing the groundplan condition of this character for the pertaining taxa. The same state is found in the species of Savtshenkia which lie in the range alpha to simulans (fig. 148). The membranous area lies in between the bases of the hypogynial valves in the species of the clade ignobilis to omega (fig. 149). In these species, the valves are usually broadly separate at the posterior end of the membrane.
87. - Sternite 9: (0) with long and slender medial projection; (1) with broad and short medial projection.

The medial projection of sternite 9 usually takes the form of an elongate slender and pointed structure in Tipulidae. This situation, which probably reflects the groundplan condition of the Lunatipula group and Mediotipula, is present in the majority of species of Savtshenkia (figs. 150, 168, 200). The species benesignate, eugeni, boloptera, obsoleta, and pagana have a differently built medial projection, which consists of a rather broad structure that is narrowed near midlength (fig. 151). In eugeni, the cavities next to the medial projection are restricted to the anterior part of sternite 9 , whereas they run along the whole length of the medial projection in the four other species.

## DISCUSSION OF ADOPTED PHYLOGENY

The character state matrix as given in table 4 was used to establish the phylogeny of the species and subspecies of Savtshenkia. The character states are binary coded, polymorphies are coded '-", absent data '?'. Initially all characters employed had the same weight 1 . The character state matrix of table 4 was run under the command ' $\mathrm{mh}^{*}$ '; $\mathrm{bb}^{\text {'; ' ' of the parsimo- }}$ ny program Hennig86. The first part of the com-
mand ( $\mathrm{mh}^{*}$;) constructs several trees to which it applies branch-swapping, retaining one tree for each initial tree, while the second part of the command ( $\mathrm{bb}^{* ;}$ ) applies extended branch-swapping to each of the input trees, retaining all most parsimonious trees. Although it does not guarantee to find all trees of minimal length, applying ' $\mathrm{mh}^{*}$; $\mathrm{bb}^{*}$;' to extensive data sets offers a good alternative for the time-consuming implicit enumeration options of Hennig86 that definitely result in finding all trees of minimal length (Farris 1988). The procedure followed resulted in 28 equally most parsimonious trees with length 157 , consistency index 55 , and retention index 88.

Subsequently successive weighting was carried out under the command ' $\mathrm{mh}^{*}$; $\mathrm{bb}^{*}$; xs w; cc;'. This procedure uses the fit of the characters to the input trees to calculate the weights as the product of the character consistency and character retention indices, which are then scaled in the range $0-10$ (Farris 1988). Successive weighting resulted in 14 equally most parsimonious trees with length 723, consistency index 84, and retention index 96 . This result was obtained after a single weighting run and did not improve upon further repeats. The calculated weights for the characters are given in table 5, the strict consensus tree of the 14 equally most parsimonious trees is shown in fig. 2. The strict consensus tree of the 14 trees resulting of successive weighting (fig. 2) differs from the strict consensus tree of the 28 trees without successive weighting in the presence of the clade battistanica to chrysocephala in the former. The internode leading to this clade is not recognized in the consensus tree of the 28 trees without weighting, lowering the grouping baltistanica to chrysocephala one level relative to its position in fig. 2. For the remainder the topologies of both consensus trees are identical. The results of the phylogenetic analysis will be discussed with reference to fig. 2.

At the base of the cladogram a trichotomy of the clades alpha and grisescens, hartigiana to simulans, and ignobilis to omega is found. The species pair alpha and grisescens probably are sister species as exemplified by two homoplasies and one reversal [characters 2 (length of male antenna), 9 (size of gonapophysis), and 4 (shape of cell ml ), respectively].

The second clade seems to be better substantiated by the characteristic shape of the gonapophysis (character 10) and the acute posterior corner of the base of the inner gonostylus (character 32). The first lineage within this clade leads to a triplet containing the species hartigiana as the sister species of the pair atlas and breviantennata. In its original description, bartigiana was referred to the atlas group sensu Theowald (table 3) (Theowald, Dufour \& Oosterbroek 1982). Besides the species atlas, breviantennata, and hartigiana, the atlas group sensu Theowald also included the species
cheethami. As the present analysis shows, the latter species seems to be more closely related to the species of the second group within the clade hartigiana to simulans. This group, the clade tulipa to simulans, consists of the species tulipa, villeneuvit, goriziensis, and the pair cheethami and simulans. Dufour (1983), on describing the species tulipa, placed it in the grisescens group sensu Theowald which by then contained the species goriziensis, grisescens, and villeneuvii (table 3). Dufour (1983) suggested that grisescens stands apart from the other three species, including villeneuvii, thus countering Theowald \& Oosterbroek's (1981) contention that villeneuvii should be regarded a subspecies of grisescens. The results of the present analysis endorse Dufour's views.

The third basal clade in fig. 2 contains the remainder of Savtshenkia. It opens with the successive branching off of the species ignobilis and subnodicornis, followed by the remainder of species of Savtshenkia which constitute the monophyletic group baltistanica to omega. The species subnodicornis was placed either near goriziensis, grisescens, or postposita by Savchenko (1961) and Theowald (1957, 1967, 1973, 1978) (tables 2, 3). The species listed belong to the phylogenetically more basal species of Savtshenkia, but none of them appears to be the actual sister species of subnodicornis.

The clade baltistanica to chrysocephala contains the majority of species of Savtshenkia with a non-spinous posterior margin of tergite $9-10$ (character 72). The only other species of Savtshenkia in which the spines on the extensions of tergite 9-10 are absent, are the phylogenetically basal species alpha and the derived species mannheimsi and omega. The basal trichotomy of the clade baltistanica to chrysocephala is constituted of the species baltistanica, the species pair glaucocinerea and postposita, and the clade lundbladi to chrysocephala. The clade lundbladi to chrysocephala contains almost all strictly Atlantic (viz. lundbladi, multipicta, and rufina maderensis), and all Afrotropical species of Savtshenkia (viz. nephrotomoides to chrysocephala). (The only remaining Atlantic species of Savtshenkia is macaronesica from the Azores, a species closely related to alpium, see below.) As table 3 shows, the Palaearctic species of the clade baltistanica to chrysocephala were placed in various groupings by Savchenko (1961) and Theowald (1973). (The phylogeny of the Afrotropical species presented here should be regarded as preliminary only. Most species are known of just a few specimens, thus giving the strong impression that the Afrotropical fauna is poorly known. Furthermore, the species are in general very similar to each other and homoplasy in this group seems to be rather common. As the present study is primarily concerned with the western Mediterranean species of Savtshenkia, this issue is not further ex-
plored here.)
Modifications of the posterior margin of the male sternite 8 (characters 55,57 ) suggest that the remainder of species of Savtshenkia, the clade alpium to ome$g a$, constitute a monophyletic group. The contours of this group were recognized by Theowald (1957, 1967) in his signata group (table 2), by Savchenko (1961) in his marmorata group, and by Theowald (1973) in his obsoleta, alpium, marmorata, staegeri, and signata groups (table 3). Within the clade alpium to omega, the basal lineage is composed of the species pair alpium and macaronesica. Theowald (1973) correctly grouped both species in his alpium group, but Savchenko (1961) placed macaronesica in his cheethami group (table 3). The species subalpium is probably closely related to alpium and macaronesica (see Appendix B). The sister group of alpium and macaronesica is substantiated by apomorphies of the sperm pump (characters 48, 49). The basal phylogeny of this clade (jeekeli to omega) could not be fully resolved. Four lineages constitute a polytomy, with the first containing the single species jeekeli, the second the clade obsoleta to pagana, the third the clade confu$s a$ to aster, and the fourth the clade fragilis to omega. The species of the clade obsoleta to pagana are most readily distinguished from the other species of Savtshenkia by the peculiar form of sternite 8 (character 58). Notwithstanding this, Savchenko (1961) as well as Theowald (1973) placed obsoleta in a species group different from holoptera and pagana (table 3). II here retain holoptera as a taxon separate from pagana. Both taxa are usually distinguished by the length of the wing and build of the legs in the female (macropterous with slender legs in holoptera, brachypterous with stoutish legs in pagana), and the shape of the posterior margin of male sternite 8 and the number of setae it carries (more deeply emarginate with one seta on tubercle in holoptera, less emarginate to truncate with about five to seven setae on tubercles in pagana) (Coe 1950; Edwards 1939). However, numbers of setae on the posterior margin of sternite 8 intermediate to those of typical holoptera and pagana can be found in certain specimens (Eiroa 1987; see also Hutson \& Vane-Wright 1969). The occurrence of intermediate stages of fully winged and brachypterous forms has been recorded within Savtshenkia for the female of gimmerthali (Dufour \& Brunhes 1984). As long as a similar variability of wing length is not recorded for females of the pair holoptera and pagana, I prefer to keep them apart. This the more so as the studied male specimens of holoptera show a slight difference with those of pagana in the presence of a small acute and blackish sclerotized extension at the posterior end of the dorsal edge of the inner gonostylus.]

The clade confusa to aster, which is substantiated by the elongate aedeagus (character 15 ) and the presence

Table 4. Character state matrix of Tipula (Lunatipula), T. (Mediotipula), and the species and subspecies of T. (Savtshenkia).

| Character | 1 | 2 | 3 |
| :--- | :--- | :--- | :--- |
|  | 0 | 0 | 0 |


| Lunatipula | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mediotipula | 00010 | 00000 | 00000 | 00000 | 00010 | 00000 | 00000 |
| aberdareica | 01010 | 00000 | 10000 | 00000 | 00001 | 00000 | 00000 |
| akelcyi | 01010 | 10000 | 10000 | 00001 | 00001 | 01000 | 00000 |
| alpha | 01000 | 00010 | 00000 | 00000 | 00000 | 00000 | 00000 |
| alpium | 00010 | 00010 | 00000 | 00000 | 10000 | 00000 | 00000 |
| asbolodes | 01010 | 00000 | 10000 | 00000 | 00001 | 00000 | 00000 |
| aspromontensis | 00010 | 00000 | 00001 | 000-0 | 00000 | 00110 | 00000 |
| aster | 00010 | 00000 | 00001 | 00000 | 00000 | 00110 | 00000 |
| atlas | 10010 | 01001 | 00000 | 00010 | 00000 | 00000 | 01000 |
| baltistanica | 00010 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| benesignata | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 00000 |
| boreosignata | 00010 | 00010 | 00000 | 00000 | 01000 | 00000 | 00000 |
| breviantennata | 10010 | 01001 | 00000 | 00010 | 00000 | 00000 | 01000 |
| caligo | 01010 | 10000 | 10000 | 00001 | 00001 | 01000 | 00000 |
| cheethami | 00011 | 00001 | 00000 | 00010 | 00100 | 00000 | 01100 |
| chrysocephala | 01010 | 10000 | 10000 | 00001 | 00001 | 01000 | 00000 |
| confusa | 00010 | 00000 | 00001 | 00000 | 00000 | 00000 | 00001 |
| corsosignata | 00010 | 00100 | 00101 | 00100 | 01010 | 10001 | 00000 |
| cyrnosardensis | 10010 | 00000 | 01001 | 00000 | 00010 | 00000 | 00000 |
| draconis | 01010 | 00000 | 00000 | 00000 | 00001 | 00000 | 00000 |
| eleonorae | 00010 | 00000 | 00000 | 00000 | 01000 | 00000 | 00000 |
| elgonensis | 01010 | 00000 | 00000 | 00000 | 0000? | 00000 | 00000 |
| eugeni | 00010 | 00000 | 00001 | 00000 | 00000 | 00000 | 00001 |
| fragilina | 00010 | 00000 | 00001 | 01000 | 00010 | 00001 | 00000 |
| fragilis | 00010 | 00000 | 01001 | 00000 | 00000 | 10000 | 00000 |
| gimmerthali | 00010 | 00000 | 00001 | 00000 | 00010 | 00001 | 00000 |
| g. mattheyi | 00010 | 00000 | 00001 | 00000 | 00010 | 00001 | 00000 |
| g. pteromaculata | 00010 | 00000 | 00001 | 00000 | 00010 | 00001 | 00000 |
| glaucocinerea | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| goriziensis | 00010 | 00001 | 00000 | 00000 | 00100 | 00000 | 01100 |
| graciae | 00010 | 00000 | 00001 | 00000 | 00000 | 00000 | 00000 |
| grisescens | 01000 | 00010 | 00000 | 00000 | 00000 | 00000 | 00000 |
| haennii | 00010 | 00000 | 00101 | 00100 | 01010 | 10000 | 00000 |
| bancocki | 00010 | 00000 | 10000 | 00000 | 00001 | 00000 | 00000 |
| hartigiana | 00010 | 00001 | 00000 | 00010 | 00000 | 00000 | 01000 |
| holoptera | 01010 | 00000 | 00000 | 10000 | 00000 | 00000 | 00000 |
| ignobilis | 00010 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| interserta | 00010 | 00010 | 00001 | 00000 | 00000 | 00000 | 00000 |
| invenusta | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| i. microinvenusta | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| i. subinvenusta | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| jeekeli | 00010 | 00000 | 00000 | 00000 | 00000 | 00000 | 00001 |
| limbata | 00010 | 00000 | 00001 | 01000 | 00000 | 00001 | 00000 |
| lundbladi | 00110 | 00000 | 00001 | 00000 | 00001 | 00000 | 00000 |
| macaronesica | 00010 | 00010 | 00000 | 00000 | 10000 | 00000 | 00000 |
| mannheimsi | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |


| $\begin{aligned} & 5 \\ & 0 \end{aligned}$ |  | 60 |  | 7 |  | 8 |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00 |
| 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 01000 | 00 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? | $\because:$ |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 00000 | 01000 | 00000 | 01100 | 00 |
| 01000 | 00101 | 01000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? | ? ? |
| 01101 | 00100 | 01000 | 00000 | 10000 | 00010 | 11100 | 01100 | 10 |
| 01111 | 00101 | 01000 | 00000 | 10000 | 00010 | 11100 | 0? ? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 00110 | 00000 | 00000 | 00100 | 00 |
| 01000 | 00100 | 01000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11000 | 01000 | 10000 | 00001 | 01100 | 11 |
| 01010 | 01111 | 01000 | 11000 | 01000 | 00000 | 00001 | 0? ? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 00110 | 00000 | 00000 | 00100 | 00 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 00000 | 00100 | 00000 | 00000 | 00001 | 00000 | 00000 | 01101 | 00 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? ? | ?? |
| 01111 | 10101 | 01000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11110 | 01000 | 10000 | 00001 | 11100 | 10 |
| 01110 | 00111 | 01000 | 01000 | 01000 | 10000 | 00000 | 01100 | 10 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? ? | ?? |
| 01010 | 01111 | 01000 | 11000 | 01000 | 10000 | 00001 | 0? ? ? ? | ? ${ }^{\text {l }}$ |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11000 | 00000 | 0? ? ? ? | ?? |
| 01101 | 00101 | 01000 | 00000 | 10000 | 10010 | 00000 | 01100 | 11 |
| 01110 | 00111 | 00001 | 00000 | 01000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00101 | 01000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00110 | 00000 | 00000 | 01000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00110 | 00000 | 00000 | 01000 | 10000 | 00000 | 0? ? ? ? | ! |
| 01110 | 00110 | 00000 | 00000 | 01000 | 10000 | 00000 | 0? ? ? ? | ?? |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 00000 | 00100 | 00000 | 00000 | 00000 | 00000 | 00000 | 01101 | 00 |
| 01111 | 00101 | 01000 | 00000 | 00000 | 00000 | 00000 | 0? ? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 00000 | 00000 | 00000 | 01110 | 00 |
| 01110 | 00111 | 01000 | 11110 | 01000 | 10000 | 00001 | 0? ? ? ? | ?? |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11100 | 00000 | 0? ? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 01100 | 00000 | 00000 | 01100 | 00 |
| 01110 | 00101 | 01110 | 00000 | 00000 | 10000 | 00000 | 01110 | 11 |
| 00000 | 00100 | 00000 | 00000 | 00000 | 00000 | 00000 | 01100 | 10 |
| 01101 | 10100 | 00000 | 00000 | 00000 | 00000 | 00010 | 01100 | 10 |
| 01110 | 00111 | 00000 | 10000 | 01000 | 00000 | 00010 | 0? ? ? ? | ?? |
| 01110 | 00111 | 00000 | 10000 | 01000 | 00000 | 00010 | 01100 | 10 |
| 01110 | 00111 | 00000 | 10000 | 01000 | 10000 | 00010 | 01100 | 10 |
| 01110 | 00101 | 01000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00111 | 00001 | 00000 | 01000 | 10000 | 00000 | 01100 | 10 |
| 01000 | 00101 | 00000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 01000 | 00101 | 00000 | 00000 | 00000 | 10000 | 00000 | 0? ? ? ? | ?? |
| 01110 | 00111 | 01000 | 11001 | 01000 | 11000 | 00001 | 1???? | ? ${ }^{\text {a }}$ |

Table 4. (continued) Character state matrix of Tipula (Lunatipula), T. (Mediotipula), and the species and subspecies of T. (Sav

| Character |  | $\begin{aligned} & 1 \\ & 0 \end{aligned}$ |  | $\begin{aligned} & 2 \\ & 0 \end{aligned}$ |  | 30 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| multipicta | 00110 | 00000 | 00001 | 00000 | 00001 | 00000 | 00000 |
| nephrotomoides | 00010 | 00000 | 00000 | 00001 | 00001 | 00000 | 00000 |
| nielseni | 00010 | 00000 | 00001 | 00000 | 00010 | 00001 | 00000 |
| obsoleta | 00010 | 01000 | 00001 | 10000 | 00000 | 00000 | 00000 |
| odontostyla | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| omega | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |
| ornata | 00010 | 00000 | 00001 | 00000 | 00000 | 10000 | 00000 |
| pagana | 01010 | 00000 | 00000 | 10000 | 00000 | 00000 | 00000 |
| pechlaneri | 00010 | 00000 | 00001 | 00000 | 01010 | 00001 | 00000 |
| persignata | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |
| p. tofina | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |
| phoroctenia | 00010 | 00000 | 01001 | 00000 | 00000 | 00000 | 00000 |
| postposita | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| rufina | 00010 | 00000 | 00000 | 00000 | 00001 | 00000 | 00000 |
| r. maderensis | 00010 | 00000 | 00001 | 00000 | 00001 | 00000 | 00000 |
| sardosignata | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| serrulifera | 00010 | 00000 | 00001 | 00000 | 00000 | 00110 | 00000 |
| signata | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |
| simulans | 00011 | 00001 | 00000 | 00010 | 00100 | 00000 | 01100 |
| sordidipes | 00010 | 00000 | 00001 | 00000 | 00000 | 00000 | 00000 |
| staegeri | 00010 | 00000 | 00001 | 00000 | 00010 | 00000 | 00000 |
| subnodicornis | 000-0 | 00000 | 00000 | 00000 | 00000 | 00000 | 00000 |
| subsignata | 00010 | 00100 | 00111 | 00100 | 01010 | 10000 | 00010 |
| s. cazorla | 00010 | 00100 | 00111 | 00100 | 01010 | 10000 | 00010 |
| subvafra | 00010 | 00000 | 00001 | 01000 | 00010 | 00001 | 00000 |
| trinacria | 00010 | 00000 | 00001 | 00000 | 01100 | 00000 | 10000 |
| tulipa | 00010 | 00001 | 00000 | 00000 | 00100 | 00000 | 01000 |
| venerabilis | 00010 | 00000 | 00001 | 00000 | 00000 | 00000 | 00000 |
| villeneuvii | 10010 | 00001 | 00000 | 00000 | 00100 | 00000 | 01000 |

of elongate and anteriorly directed posterior apodemes of the sperm pump (character 50), contains as its first lineage the triplet confusa, interserta, and orna$t$. Although differing in most genital features, these three species share the presence of the quite remarkable slender elongate and anteriorly extended posterior apodemes of the sperm pump (character 51). In fig. 2, the remainder of the clade confusa to aster is depicted as a monophyletic group which is substantiated by a single reversal only. The reversal concerns the redirection and reduction of the extensions of the posterior margin of tergite 9-10 (character 71). The extensions are rather differently shaped in the species of this grouping and its actual monophyly is open to question. The species eugeni, which is included in this grouping, has the extensions of tergite 9-10 well developed and laterally extended.

Theischinger (1983) correctly considered his species aster to be the closest relative of aspromontensis.

Theowald (1973) placed the species confusa (as
marmorata), eugeni, and jeekeli in his marmorata group on account of the peculiar shape of their inner gonostyli (character 35). When other characters are taken into consideration, the three species appear not to be closely related, forcing the conclusion that the characteristic shape of the inner gonostylus must be due to homoplasy.

Next in fig. 2 is the clade fragilis to omega, in which the Nearctic species pair fragilis and phoroctenia branch off first, followed by the species odontostyla.

The remainder of species of Savtshenkia constitute the monophyletic group fragilina to omega of which the first lineage leads to the clade fragilina to invenusta microinvenusta. This latter clade contains species which are characterized by the presumably secondary absence of spines on the posterior margin of male sternite 8 (character 57). It is composed of species which Theowald (1973) arrayed in his gimmerthali, invenusta, and limbata groups (table 3), together with the Nearctic species fragilina (the species mohriana

| $\begin{aligned} & 5 \\ & 0 \end{aligned}$ | $0$ |  |  | $\begin{aligned} & 7 \\ & 0 \end{aligned}$ |  | $\begin{aligned} & 8 \\ & 0 \end{aligned}$ |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 01000 | 00101 | 01000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 11110 | 00111 | 01000 | 11000 | 01000 | 10000 | 00001 | 01100 | 10 |
| 01110 | 00101 | 01100 | 00000 | 00000 | 10000 | 00000 | 01110 | 11 |
| 01110 | 00101 | 01000 | 00000 | 00000 | 10010 | 00000 | 0? ? ? | ? ? |
| 01110 | 00111 | 01000 | 11001 | 01000 | 11000 | 00001 | 1?? ? | ?? |
| 01111 | 10101 | 00000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00101 | 01110 | 00000 | 00000 | 10000 | 00000 | 01110 | 11 |
| 01110 | 00110 | 00000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11000 | 01000 | 10000 | 00001 | 0? ? ? | ? ? |
| 01110 | 00111 | 01000 | 11000 | 01000 | 10000 | 00001 | 0? ? ? | ?? |
| 01110 | 00101 | 01000 | 00000 | 00000 | 10000 | 00000 | 01100 | 10 |
| 01000 | 00100 | 00000 | 00000 | 00000 | 11000 | 00000 | 0? ? ? | ?? |
| 01000 | 00101 | 10000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 01000 | 00101 | 10000 | 00000 | 00000 | 11000 | 00000 | 01100 | 10 |
| 11110 | 00111 | 01000 | 11000 | 00000 | 10000 | 00001 | 01100 | 10 |
| 01111 | 00101 | 01000 | 00000 | 10000 | 00010 | 10000 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11000 | 01000 | 10000 | 00001 | 11100 | 10 |
| 00000 | 00100 | 00000 | 00000 | 00001 | 00000 | 00000 | 01101 | 00 |
| 01111 | 00100 | 01000 | 00000 | 00000 | 00011 | 00000 | 01100 | 10 |
| 01110 | 00111 | 01000 | 01000 | 01000 | 10000 | 00000 | 01100 | 10 |
| 00000 | 00100 | 00000 | 00000 | 00000 | 10000 | 00000 | 01110 | 10 |
| 01110 | 00111 | 01000 | 11110 | 01000 | 10000 | 00001 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11110 | 00000 | 10000 | 00001 | 0? ? ? | ? ? |
| 01110 | 00111 | 00001 | 00000 | 01000 | 10000 | 00010 | 01100 | 10 |
| 01110 | 00111 | 01000 | 11001 | 01000 | 10000 | 00001 | 11100 | 10 |
| 00000 | 00100 | 00000 | 00000 | 00000 | 00000 | 00000 | 0? ? ? ? | ? ? |
| 01111 | 00100 | 01000 | 00000 | 00000 | 00011 | 00000 | 0? ? ? | ?? |
| 00000 | 00100 | 00000 | 00000 | 00000 | 00000 | 00000 | 01101 | 00 |

and productella probably also belong here; see Appendix B).
The clade fragilina to invenusta microinvenusta contains two trichotomies. The exact relationships of the species fragilina, limbata and subvafra as well as those between the three subspecies of gimmerthali are hard to establish. The subspecies gimmerthali pteromaculata was described as differing from the nominotypical subspecies by its darker colour, spotted wings, and relatively short second flagellomere (Theowald, Dufour \& Oosterbroek 1982). Examination of material of the nominotypical form of gimmerthali, however, revealed that the length of the second flagellomere varies considerably within this subspecies. The relative length of the second flagellomere thus offers no arguments to distinguish gimmerthali pteromaculata from the nominotypical subspecies. [The length of the second male flagellomere has also been used to distinguish gimmerthali from pechlaneri (Mannheims \& Pechlaner 1963). Considering the
variability of this character within gimmerthali it has proved to be non-applicable here. The species gimmerthali and pechlaneri are more easily distinguished on account of the structure of the inner gonostylus and the shape of male tergite 9-10 (see also Appendix C). The monophyly of the three subspecies of gimmerthali could be demonstrated by the shape of the inner gonostylus only (character 41)]. Useful differences between gimmerthali gimmerthali and gimmerthali pteromaculata are found in the autapomorphous larger outer gonostylus and relatively plump anterior tip of the inner gonostylus of gimmerthali pteromaculata (cf. figs. 66 and 67, and 83 and 84). The subspecies gimmerthali mattheyi, known of the male holotype only (not examined), apparently differs from the other subspecies by the autapomorphous absence of a posterior projection on the outer gonostylus and the reduced armature of the posterior margin of tergite 910 (Theowald \& Dufour 1983). [The females of both gimmerthali mattheyi and gimmerthali pteromaculata

Table 5. Weights of characters after single weighting run using the command ' $\mathrm{mh}^{*}$; bb'; xs w; cc; of Hennig86.

| Character | Weight | Character | Weight | Character | Weight |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 1 | 31 | 10 | 61 | 4 |
| 2 | 1 | 32 | 10 | 62 | 10 |
| 3 | 10 | 33 | 10 | 63 | 10 |
| 4 | 1 | 34 | 10 | 64 | 10 |
| 5 | 10 | 35 | 0 | 65 | 10 |
| 6 | 10 | 36 | 4 | 66 | 10 |
| 7 | 2 | 37 | 10 | 67 | 1 |
| 8 | 10 | 38 | 10 | 68 | 10 |
| 9 | 1 | 39 | 10 | 69 | 10 |
| 10 | 10 | 40 | 10 | 70 | 10 |
| 11 | 10 | 41 | 10 | 71 | 1 |
| 12 | 2 | 42 | 10 | 72 | 2 |
| 13 | 10 | 43 | 10 | 73 | 4 |
| 14 | 10 | 44 | 2 | 74 | 4 |
| 15 | 1 | 45 | 2 | 75 | 10 |
| 16 | 10 | 46 | 10 | 76 | 10 |
| 17 | 10 | 47 | 10 | 77 | 10 |
| 18 | 10 | 48 | 4 | 78 | 10 |
| 19 | 3 | 49 | 2 | 79 | 1 |
| 20 | 3 | 50 | 10 | 80 | 10 |
| 21 | 10 | 51 | 10 | 81 | 3 |
| 22 | 4 | 52 | 10 | 82 | 2 |
| 23 | 4 | 53 | 10 | 83 | 10 |
| 24 | 2 | 54 | 10 | 84 | 1 |
| 25 | 10 | 55 | 1 | 85 | 10 |
| 26 | 2 | 56 | 10 | 86 | 10 |
| 27 | 10 | 57 | 1 | 87 | 1 |
| 28 | 10 | 58 | 10 |  |  |
| 29 | 10 | 59 | 10 |  |  |
| 30 | 2 | 60 | 10 |  |  |

remain unknown. Those of gimmerthali gimmerthali are usually brachypterous, but populations of the nominotypical form of gimmerthali are known from Switzerland which have female wing lengths ranging from the brachypterous to the fully winged condition (Dufour \& Brunhes 1984). Dufour (1990) furthermore recorded hemipterous females of gimmerthali from the eastern side of the Sierra Nevada, Spain.]

The species invenusta presently contains three subspecies. Differences between the subspecies can be found in details of the male terminalia and in the ventral distance between the eyes. The examined specimens of invenusta invenusta conform to Riedel's (1919) description of the species as having rounded posterolateral corners on tergite 9-10 ['Lam. term. sup. (...) mit abgerundeten Ecken']. However, illustrations of the terminalia of 'invenusta invenusta' by Savchenko (1961) and Theowald (1973) depict tergite 9-10 with acute posterior extensions. Examination of material present in ZMAN showed that Theowald actually figured a specimen that can only be identified as invenusta subinvenusta. However, considering Savchenko's illustration there still is a possibility that the structure of the posterior margin of tergite 9-10 varies within invenusta invenusta. [All
three described subspecies bear spines on the posteroventral margin of tergite 9-10. Having examined several of its paratypes, I can not corroborate Dufour's (1990) observation that spines are absent from the posterior margin of tergite 9-10 in his invenusta microinvenusta.] Another complicating factor in maintaining the three recognized subspecies is the observation that Tipulidae occurring at high geographical latitudes or high altitudes usually have smaller eyes than their relatives living under more moderate ecological conditions. Therefore, the size of the eyes in specimens of invenusta might well be an ecotypic differentiation in response to local environmental conditions and thus would be of little value in separating the subspecies. The complicated nature of the taxonomy of invenusta is furthermore illustrated by the fact that Alexander described invenusta invenusta under three different names, viz. docilis (correctly synonymized with invenusta by Savchenko 1961), and insignifica and packardi (synonymized in this paper; see Appendix A, synonymy). Hancock (1988) suggested that yet another subspecies might be introduced to accommodate a morphologically slightly deviating population from Andorra (Pyrenees). I here would like to stress the need to study the variability of this
taxon over its immense Holarctic range before even considering to introduce new formal names for single isolated populations. [The most recently published map showing the Palaearctic distribution of invenusta is the one given by Dufour (1992).]

The clade cyrnosardensis to omega contains the species of Theowald's (1973) staegeri and signata species groups (table 3). Most species of this clade are easily recognized by the presence of the bulbous and spinous extensions on the posterior margin of sternite 8 (character 62). The clade opens with a trichotomy consisting of the species cyrnosardensis and staegeri in combination with a clade which includes the remainder of the species. Both cyrnosardensis and staegeri are species distinguished by a number of autapomorphies but apparently lack any further character states that could enable the resolution of their exact phylogenetic position.

The species nielseni and sardosignata are depicted here as sister species as is suggested by their broad based outer gonostyli, although the overall shape of this structure differs remarkably in both species (character 46). The two species are, moreover, distinguished from each other by the autapomorphous form of the inner gonostylus and the shape of sternite 8 . Their tentative sister species relationship could not be substantiated by other apomorphies.

Although on quite different grounds, the clade baennii to subsignata cazorla contains the four taxa which were recognized as a monophyletic group by Dufour (1991). The characters employed in the present analysis result in a sequence of relationships between these taxa which differs from the solution provided by Dufour. Compared with Dufour's cladogram, corsosignata and baennii have changed places in the present phylogeny, necessitating the elevation in rank of baennii from subspecies of subsignata to species (see Appendix A). Dufour substantiated his concept of a monophyletic group containing corsosignata, baennii, subsignata subsignata, and subsignata cazorla by the degree of sclerotization and shape of the ventromedial appendage of male sternite 8 and the degree of sclerotization of female sternite 9 (Dufour 1991, characters 1 and 2). I can see no essential differences between the sclerotization and shape of the ventromedial extension of male sternite 8 in this clade of four taxa and that of other derived members of Savtshenkia such as persignata, sardosignata, and signata, and prefer to consider it a plesiomorphy at this level. The presence of an unsclerotized female sternite 9 as a synapomorphy for the members of the clade haennii to subsignata cazorla has not been employed in this paper because it is not always clear whether sternite 9 should be valued as sclerotized or not in species of Savtshenkia. Moreover, the female of subsignata cazorla remains unknown. Dufour considered
haennii the sister species of the pair subsignata subsignata and subsignata cazorla because of the presence of a ridge on what he denoted as the median part of the inner gonostylus (Dufour 1991, character 3). As discussed under character 34, I consider the presence of the anteriorly extended part of the lateral ridge of the inner gonostylus an apomorphy of the pair subsignata subsignata and subsignata cazorla. Of the three apomorphies of Dufour supporting the monophyly of subsignata subsignata and subsignata cazorla, two appear to be invalid (Dufour 1991, characters 4 and 5). The continuity of the posterior and ventral parts of the lateral ridge of the inner gonostylus in these two subspecies (Dufour 1991, character 4) probably is a plesiomorphy (see discussion under character 34). The relatively broad base of the outer gonostylus in subsignata subsignata and subsignata cazorla (Dufour 1991, character 5) probably also represents a plesiomorphy considering the situation in most related species. The width of the base of the outer gonostylus furthermore varies within the species and subspecies and differs only slightly from that in haennit. Dufour's third argument for the monophyly of the pair subsignata subsignata and subsignata cazorla (Dufour 1991, character 6, shape of the gonapophysis) is employed in this paper as well (character 14).

Following on his description of eleonorae, Theischinger (1978) suggested that this species is most closely related to benesignata. I am, however, convinced that the sister species of eleonorae is boreosignata, which is substantiated by the almost identical shape of the inner gonostylus (character 42) and the bizarre modification of the aedeagal complex (character 52). Both boreosignata and eleonorae are known of their holotypes only, with the type localities lying about 2500 kilometers apart, that of boreosignata in northern Sweden and that of eleonorae in western Rumania!

The clade benesignata to omega contains the species which Theowald (1973) included in his signata group, with the addition of the species omega and trinacria described herein as new.

## Distribution

In this section I will give a brief account on the distribution of the species and subspecies of Savtshenkia. As noted above, more detailed information on the distribution of the western Mediterranean species and subspecies and their closest relatives will be given in a forthcoming paper on the historical biogeography of the western Mediterranean area. A survey of the distribution of the species and subspecies of Savtshenkia is given for the Afrotropical Region by Hutson (1980, under Pterelachisus), the Nearctic Region by Alexander (1965b), the Oriental Region by Alexander \&

Alexander (1973), and the Palaearctic Region by Oosterbroek \& Theowald (1992). The sequence of taxa in the summary below follows that of fig. 2.

The two species of the pair alpha and grisescens differ considerably in their ranges. The first species is known of its type locality in the Rodna Mountains of northern Rumania only, while grisescens is distributed over the central belt of Europe eastwards to the Altai Mountains in central Asia.

The species of the clade hartigiana to simulans are confined to the western Palaearctic, with most of them having a restricted range. The species hartigiana is known from Corsica and Sardinia only, atlas is recorded from Algeria and Morocco, and breviantennata from Spain, southern Switzerland, Corsica, Italy, including Sardinia, and northern Algeria. The species tulipa was described from two localities in southern Switzerland, villeneuvii is known from central Spain, while goriziensis, cheethami and simulans are more widely distributed over the mountainous areas of central Europe. The species ignobilis has an eastern Nearctic distribution between about $35^{\circ}$ and $50^{\circ} \mathrm{N}$ latitude. The species subnodicornis has a wide range extending from the Faeroes and Ireland in the west over central Europe towards Lake Baykal in the east.

The clade baltistanica to chrysocephala contains species which cover a wide geographic area. The species baltistanica occurs in the Caucasus, Tadzhikistan and Kashmir, glaucocinerea has an arctic Siberian distribution, as does its presumed sister species postposita, which is furthermore known from northern Finland, the Altai Mountains and Mongolia. The species lundbladi is an endemic of Madeira, the species multipicta is presently recorded from the Canary Islands only. The nominotypical form of rufina is widespread, ranging from Iceland and north-west Africa towards Tadzhikistan in the east, while the subspecies rufina maderensis is endemic to Madeira. The majority of species of the grouping nephrotomoides to chrysocephala is distributed in the mountainous areas surrounding Lake Victoria in eastern Africa. The only member of this group that does not occur in this region is the species draconis from South Africa (Natal).

The species alpium has a wide distribution which includes most of Europe. It also occurs along the eastcoast of the Nearctic Region between about $42^{\circ}$ and $55^{\circ} \mathrm{N}$ latitude (see Appendix A, synonymy, under nebulipennis). The species macaronesica is endemic to the Azores.

As is presently known, the phylogenerically rather isolated species jeekeli has a disjunct distribution throughout the Mediterranean, ranging from Galicia in north-west Spain to Cyprus in the east.

The species obsoleta is distributed over most of Europe, extending its range towards Georgia in the south-east, while the pair holoptera and pagana is con-
fined to the central belt of Europe, with holoptera having a disjunct distribution in south-west England, north-west Spain, and Czechia.

Of the species of the clade confusa to aster, confusa occurs over most of Europe, including the Faeroes and Ireland in the north-west. The species interserta has a distribution ranging from the central European mountains in the west towards the Tuva in the east. The beautiful ornata was described on the basis of a single male from northern Israel, but a series of this species, including females, was collected recently on Cyprus by Pjotr Oosterbroek and Cita Hartveld. It seems that the species graciae still is known from its type locality in southern California only. Both sordidipes and venerabilis are restricted to Kashmir. The species eugeni is presently known from Corsica, Sicily, and southern Italy. The species serrulifera has a more extensive distribution, occurring in mountainous areas from the Pyrenees in the west to the Caucasus in the east. Theowald (1973) described aspromontensis from southern Italy (Aspromonte), but more recently it was collected also in Greece (Peloponnisos, Crete). The species aster still is known of its holotype only, originating from the Greek island Thasos in the northern Aegean Sea.

The presumed sister species fragilis and phoroctenia have a Nearctic distribution which extends across the continent in the region of the 50th degree of latitude. The species odontostyla is known from the north-eastern coastal areas of the Black Sea and the Greek island Thasos.

The species fragilina has a Nearctic distribution, being recorded from the north-western part of the continent, ranging from Alaska to Colorado, as well as from Maine on the east coast (under its synonym brower). The species limbata is widespread in the mountainous areas of Eurasia, subvafra seems to be confined to continental western Europe. The members of the clade pechlaneri to gimmerthali pteromaculata are distributed in the mountainous areas of the western Palaearctic Region, with pechlaneri having a disjunct distribution from Andorra in the Pyrenees to northern Iran, gimmerthali gimmerthali ranging from the Sierra Nevada of Spain in the south towards northern Scandinavia and eastwards to the Caucasus, gimmerthali mattheyi being restricted to southern France, and gimmerthali pteromaculata to Corsica.

The three subspecies of invenusta differ considerably in the extent of their distribution area. The subspecies invenusta microinvenusta is presently known only from southern Spain (Sierra Nevada), the subspecies invenusta subinvenusta has a wider range in the mountains of central Europe, while the nominotypical subspecies disjunctly ranges across Eurasia and also occurs in eastern North America (see Appendix A, synonymy, under insignifica and packardi).

The species cyrnosardensis is endemic to Corsica and Sardinia, staegeri occurs in Europe from the Faeroes to northern Spain in the west towards the coastal areas of the Baltic Sea in the east. Both species of the pair nielseni and sardosignata have a restricted range, with nielseni being confined to the Alps, and sardosignata occurring on Corsica and Sardinia, as well as in southern France.

The four members of the clade haennii to subsignata cazorla are also restricted to Europe, with haennii being recorded from the French and Spanish Pyrenees, corsosignata occurring on Corsica, subsignata cazorla in northern Spain, and subsignata subsigna$t a$ being more widely distributed in the mountains of central Europe. As mentioned above, both boreosignata and eleonorae are known of their holotypes only, with the type localities lying in northern Sweden and western Rumania respectively. The distribution area of the species benesignata consists of widely separated populations ranging from the Alps towards northern Scandinavia and eastwards to the Tien Mountains. The original descriptions of the two subspecies persignata persignata and persignata tofina were based on material originating from North Korea, to which area the distribution range of the nominotypical form seems to be restricted. Later records of persignata tofina, however, showed that this subspecies ranges across all of northern Eurasia.

The species signata is widespread in Europe, covering most of the area. The triplet trinacria to omega is restricted to Italy, with trinacria occurring on Sicily, mannheimsi in the north of Italy, and omega on Sardinia.

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## References

Alexander, C. P., 1919a. New Nearctic cranc-flies (Rhyphidae and Tipulidae) part VII. - Canadian Entomologist 51: 162-172.
Alexander, C. P., 1919b.The crane-flies of New York. Part I. Distribution and taxonomy of the adult flies. - Cornell University Agricultural Experiment Station, Memoir 25: 76-993.
Alexander, C. P., 1924. New or little known crane-flies from New England. -- Occasional Papers of the Boston Society of Natural History 5: 115-118.
Alexander, C. P., 1925. Undescribed species of Japanese crane-flies. Part V. - Annals of the Entomological Society of America 17: 431-448.
Alexander, C. P., 1926. Undescribed species of crane-flies from the United States and Canada, part II. - Insecutor Inscitiae Menstruus 14: 114-122.
Alexander, C. P., 1928. New or little-known species of the genus Tipula from Labrador (Tipulidae, Dipt.). Canadian Entomologist 60: 95-101.
Alexander, C. P., 1934. New or little-known Tipulidae from Eastern Asia (Diptera), XXI. - Philippine Journal of Science 55: 19-60, pls. 1-3.
Alexander, C. P., 1940. Undescribed species of crane-flies from the eastern United States and Canada (Dipt:: Tipulidae). Part VI. - Entomological News 51: 83-85, 99-103.
Alexander, C. P., 1942. See Alexander 1966b (reprint).
Alexander, C. P., 1951. New or little-known Tipulidae (Diptera). - XCI. Oriental-Australasian species. - Annals and Magazine of Natural History (12) 6: 1072-1102.
Alexander, C. P., 1953. The Oriental Tipulidae in the collection of the Indian Museum. Part III. - Records of the Indian Museum 50; 321-357.
Alexander, C. P., 1954. The Tipulidae of Mount Ishizuchi and the Omogo Valley, Iyo, Shikoku, Japan. - Transactions of the Shikoku Entomological Society 4: 23-31.
Alexander, C. P., 1955. Records and descriptions of Japanese Tipulidae (Diptera), part IV. The crane-flies of Shikoku, IV. - Philippine Journal of Science 83: 263306, pls. 1-4.
Alexander, C. P., 1956. Tipulidae. - Ruwenzori Expedition 1934-35 1: 129-380, pls. XXVII-XXVIII.
Alexander, C. P., 1961. New or little-known Tipulidae from Eastern Asia (Diptera), L. - Philippine Journal of Science 90: 397-445, pls 1-3.
Alexander, C. P., 1962. Mission zoologique de PI.R.S.A.C. en Afrique orientale. (P. Basilewsky et N. Leleup, 1957) LXXX. - Diptera Tipulidae. - Annales Musée Royal de l'Afrique Centrale, Série 8vo (Zool.) 110: 343-362.
Alexander, C. P., 1964. New or little-known Tipulidae from Eastern Asia (Diptera), LII. - Philippine Journal of Science 93: 77-130, pls. 1-6.
Alexander, C. P., 1965a. New subgenera and species of crane-flies from California (Diptera: Tipulidae). - Pacific Insects 7:333-386.
Alexander, C. P., 1965b. Family Tipulidae. - In: Stone, A. et al. (eds.) A catalog of the Diptera of America north of Mexico. Agriculture Handbook, Agriculture Research Service 276: 16-90.
Alexander, C. P., 1966a. New or little-known Tipulidae from Eastern Asia (Diptera), LVI. - Philippine Journal of Science 94: 235-286, pls. 1-5.
Alexander, C. P., 1966b. Family Tipulidae. - In: Crampton, G. C. et al. (eds.) Guide to the insects of Connecticut.

Part VI. The Diptera or true flies of Connecticut. First fascicle. Bulletin of the Connecticut State Geological and Natural History Survey 64: 196-486b. (Reprint of Alexander 1942).
Alexander, C. P. \& M. A. Alexander, 1973. Tipulidae. - In: Delfinado, M. D. \& D. E. Hardy (eds.) A catalog of the Diptera of the Oriental Region. Volume I. Suborder Nematocera: 10-224. Honolulu.
Audcent, H., 1932. British Tipulinae (Diptera, Tipulidae). - Transactions of the Entomological Society of the South of England 8: 1-34, pls. I-III.
Bergroth, E. 1888. Oesterreichische Tipuliden, gesammelt von Professor J. A. Palmén im Jahre 1870. Verhandlungen der Zoologisch-Botanischen Gesellschaft in Wien 38: 645-656.
Brindle, A., 1958. A field key for the identification of Tipula larvae (Dipt.: Tipulidae). - Entomologist's Gazette 9: 165-182.
Brindle, A., 1959. Notes on the larvae of the British Tipulinae (Dipt., Tipulidae) part 7. - The larvae of the Tipula marmorata group. - Entomologist's Monthly Magazine 95: 204-205.
Brindle, A., 1960. The larvae and pupae of the British Tipulinae (Diptera: Tipulidae). - Transactions of the Society for British Entomology 14: 63-114.
Brunhes, J. \& C. Dufour, 1992. Érudes structurales et dynamiques sur les écosystèmes de Tourbières acides. III - le peuplement des Tipulides (Diptera, Tipulidae). Bullerin d'Écologie 23: 17-26.
Bryce, D., 1956. Notes on the life-history of Tipula cheetha$m i$ Edw. (Dipt., Tipulidae). - Entomologist's Monthly Magazine 92: 104-106.
Byers, G. W., 1961. The crane fly genus Dolichopeza in North America. - University of Kansas Science Bulletin 42: 665-924.
Coe, R. L., 1950. Family Tipulidae. - Handbooks for the Identification of British Insects IX 2: 1-66.
Coulson, J. C., 1959. Observations on the Tipulidae (Diptera) of the Moor House Nature Reserve, Westmoreland. - Transactions of the Royal Entomological Society of London 111: 157-174, pls. I, II.

Coulson, J. C., 1962. The biology of Tipula subnodicornis Zetterstedt, with comparative observations on Tipula paludosa Meigen. - Journal of Animal Ecology 31: 1-21.
Dufour, C., 1983. Tipula (Savtshenkia) tulipa sp.n. from xenothermic valleys of the Swiss Alps (Diptera: Tipulidae). - Mitteilungen der Schweizerischen Entomologischen Gesellschaft 56: 275-281.
Dufour, C., 1986. Les Tipulidae de Suisse (Diptera, Nematocera). - Documenta Faunistica Helveriae 2: 1-$187,+1-149$.
Dufour, C. 1990. Tipula (Acutipula) nevada sp.n. and Tipula (Savtshenkia) invenusta microinvenusta ssp.n. from the heights of Sierra Nevada in Spain (Diptera: Tipulidae). - Mitteilungen der Schweizerischen Entomologischen Gesellschaft 63: 227-232.
Dufour, C., 1991. Tipula (Savtshenkia) subsignata haennii ssp.n. and Tipula (S.) subsignata cazorla ssp.n. from the Iberian Peninsula (Diptera, Tipulidae). - Mitteilungen der Schweizerischen Entomologischen Gesellschaft 64: 243-249.
Dufour, C., 1992. High altitude Tipulidae in Switzerland (Diptera, Nematocera). - Acta Zoologica Cracoviensia 35: 113-134.

Dufour, C. \& J. Brunhes, 1984. Les Tipulidae brachyptères de la région paléarctique occidentale avec les descriptions des $ㅇ+$ Y holoptères de Tipula (Savtshenkia) gimmerthali Lackschewitz et de Tipula (Platytipula) luteipennis agilis ssp.n. (Diptera, Tipulidae). - Mitteilungen der Schweizerischen Entomologischen Gesellschaft 57: 133-151.
Edwards, F. W., 1928. Some Nematocerous Diptera from Yunnan and Tibet. - Annals and Magazine of Natural History (10) 1: 681-703, pl. XIX.
Edwards, F. W., 1931. Some suggestions on the classification of the genus Tipula (Diptera, Tipulidae). - Annals and Magazine of Natural History (10) 8: 73-82.
Edwards, F. W., 1939. Additions to the list of British craneflies. - Entomologist's Monthly Magazine 75: 241-249.
Eiroa Alvarez, M. E。, 1987. Estudio de los Tipúlidos (O. Dípteros) en Galicia. - Thesis, Facultad de Biologia, Universidad de Santiago de Compostela: 1-422 + unnumbered pages.
Farris, J. S. 1988. Hennig86, version 1.5 + reference. - Port Jefferson Station, New York.
Frommer, S. I., 1963. Gross morphological studies of the reproductive system in representative North American crane flies (Diptera: Tipulidae). - Kansas University Science Bulletin 44: 535-626, pls. I-XX.
Gelhaus, J. K., 1986. Larvae of the crane fly genus Tipula in North America (Diptera: Tipulidae), - Kansas University Science Bulletin 53: 121-182.
Hancock, E. G., 1988. A cranefly, Tipula (Savtshenkia) invenusta Riedel (Diptera: Tipulidae), new to the British Isles. - Entomologist 107: 96-98.
Hemmingsen, A. M., 1952. The oviposition of some cranefly species (Tipulidae) from different types of localities. Videnskabelige Meddelelser fra Dansk Naturhistorisk Forening i Kjobenhavn 114: 365-430.
Hemmingsen, A. M., 1954. The function of the peculiar processes of the 8th sternite in the males of Tipula (Oreomyza) staegeri Peder Nielsen and Tipula (Oreomyza) signata Staeger. - Videnskabelige Meddelelser fra Dansk Naturhistorisk Forening i Kjobenhavn 116: 411-418.
Hemmingsen, A. M., 1962. Copulatory adaptations of male hypopygium to female tergal ovipository valves (cerci) in certain crane-flies (Tipulidae). - Videnskabelige Meddelelser fra Dansk Naturhistorisk Forening i Kjobenhavn 124: 135-163, figs. 1-18.
Hutson, A. M., 1980. Tipulidae. - In: Crosskey, R. W. (ed.) Catalog of the Diptera of the Afrotropical Region: 47-91. London.
Hutson, A. M. \& R. I. Vane-Wright, 1969. Corrections and additions to the list of British Nematocera (Diptera) since Kloet and Hincks' 'A check list of British insects' (1945), part 1. - Entomologist's Gazette 20: 231-256.
Kandybina, M. N., V. I. Lantzov \& E. N. Savchenko, 1987. A catalog of the type-specimens in the collection of the Zoological Institute, Academy of Sciences of the USSR, Insecta, Diptera, N3. Families Tanyderidae, Trichoceridae, Limoniidae, Tipulidae: 1-68 + inside covers. - Nauka, Leningrad. (In Russian).
Lackschewitz, P., 1936. A new species of Tipula from the Azores (Diptera). - Proceedings of the Royal Entomological Society of London (B) 5: 118.
Lantzov, V. I., 1982. Ecology and preimaginal developmental stages of Tipula gleucocinerea (Diptera, Tipulidae). Zoologicheskii Zhurnal 61: 1913-1916.
Loew, H., 1863. Diptera Americae Septentrionalis indigena.

Centuria quarta, - Berliner Entomologische Zeitschrift 7: 275-326.
Mannheims, B., 1951. Tipulidae. - Fliegen der Palaearktischen Region 15, Lieferung 167: 1-64, pls. I-VII.
Mannheims, B., 1962a. Zur Synonymie der europäischen Tipuliden (Dipt.) VII. - Bonner Zoologische Beiträge 13: 193-195.
Mannheims, B., 1962b. Die Tipuliden Madeiras (Dipt. Tipulidae). - Notulae Entomologicae 42: 130-136.
Mannheims, B. \& E. Pechlaner, 1963. Die Tipuliden Nordtirols (Dipt.). - Stuttgarter Beiträge zur Naturkunde 102: 1-29.
McAlpine, J. F., 1981. Morphology and terminology adults. - In: McAlpine, J. F. et al. (eds.) Manual of Nearctic Diptera. Volume 1. Research Branch, Agriculture Canada, Monograph 27: 9-63.
Neumann, H., 1958. Der Bau und Funktion der männlichen Genitalapparate von Trichocera annulata Meig. und Tipula paludosa Meig. (Dipt. Nematocera). Deutsche Entomologische Zeitschrift, Neue Folge 5: 235-298.
Oosterbroek, P. \& B. Theowald, 1992. Family Tipulidae. In: Soós, A., L. Papp \& P. Oosterbroek (eds.) Catalogue of Palearctic Diptera 1: 56-178. Budapest.
Pritchard, G., 1983. Biology of Tipulidae. - Annual Review of Entomology 28: 1-22.
Rees, B. E. \& G. F. Ferris, 1939. The morphology of Tipula reesi Alexander (Diptera: Tipulidae). - Microentomology 4: 143-178.
Riedel, M. P., 1913. Die paläarktischen Arten der Dipteren(Nematocera polyneura-) Gattung Tipula L. (Dipt.). Abhandlungen des Vereins für Naturwissenschaftliche Erforschung der Niederrheins- (Bezirksgruppe des deutschen Lehrervereins für Naturkunde.) I. Band 1913: 1122, pls. I-III.
Riedel, M. P., 1914. Nematocera polyneura. - In: Voyage de Ch. Alluaud et R Jeannel en Afrique orientale (19111912). Résultats scientifiques. Diptera, III: 69-100. Schulz, Paris.
Riedel, M. P., 1919. Résultats scientifiques de l'expédition des frères Kuznecov (Kouznetzov) à l'Oural arctique en 1909, sous la direction de H. Backlund. 8. Nematocera polyneura. - Zapiski Rossiiskoi Akademii Nauk 28 (8): 1-10.
Savchenko, E. N., 1961. Crane flies (Fam. Tipulidae), subfam. Tipulinae: genus Tipula L. (part 1). - Fauna SSSR (N.S. 79) Two-winged insects II (3): 1-488. (In Russian).

Savchenko, E. N., 1966. Crane flies. - Fauna Ukraini 14(1): 1-552. (In Ukrainian).
Savchenko, E. N., 1968a. On a new species of the crane-fly (Diptera, Tipulidae) from Kamchatka. - Zoologicheskii Zhurnal 47: 1567-1570. (In Russian, English summary).
Savchenko, E. N., 1968b. Crane-flies (Diptera, Tipulidae) new for the fauna of the USSR. Dopovidi Akademii Nauk Ukrainskoi RSR 5: 469-472. (In Ukrainian, Russian and English summaries).
Savchenko, E. N., 1969. On some little-known species of crane flies (Diptera, Tipulidae) of Transcaucasica. Zbirnyk Prats Zoolohichnoho Muzeyu 33: 51-60. (In Ukrainian, English summary).
Savchenko, E. N., 1979. Phylogenie und Systematik der Tipulidae. - Tijdschrift voor Entomologie 122: 91-126. (German translation of Savchenko 1966: 63-88).
Savchenko, E. N., 1983. Crane flies of the family Tipulidae. General part and beginning of systematic part. Subfam.

Dolichopezinae; subfam. Tipulinae (part), - Fauna SSSR (N.S. 127) Two-winged insects II (1-2): 1-585. (In Russian).
Stubbs, A. E., 1992. Provisional atlas of the long-palped craneflies (Diptera: Tipulinae) of Britain and Ireland: 1134. - Biological Records Centre, NERC Institute of Terrestrial Ecology, Monks Wood, Huntingdon, UK.
Tangelder, I. R. M., 1985. Phylogeny of the Nephrotoma dorsalis species-group (Diptera, Tipulidae), mainly based on genital characters. -Beaufortia 35: 135-174.
Theischinger, G., 1978. Tipula (Savtshenkia) eleonorae sp.nov. aus Rumänien (Diptera, Tipulidae). Entomologische Berichten 38: 45-46.
Theischinger, G., 1983. Über eine Aufsammlung von Savtshenkia Alexander, 1965, aus dem östlichen Mittelmeerraum (Diptera: Tipulidae). - Entomologische Berichten 43: 26-28.
Theowald, B. 1957. Die Entwicklungsstadien der Tipuliden (Diptera, Nematocera), insbesondere der west-palaearktischen Arten. - Tijdschrift voor Entomologie 100: 195308.

Theowald, B. 1967. Familie Tipulidae. (Diptera, Nematocera). Larven und Puppen. - Bestimmungsbücher zur Bodenfauna Europas 7: 1-100.
Theowald, B., 1973. Tipulidae. - Fliegen der Palaearktischen Region 15, Lieferung 300: 321-404.
Theowald, B., 1978. Tipulidae. - Fliegen der Palaearktischen Region 15, Lieferung 318: 405-436.
Theowald, B. \& C. Dufour, 1983. Tipula (Savtshenkia) gimmerthali mattheyi ssp.n. from the French Pyrenees. Mitteilungen der Schweizerischen Entomologischen Gesellschaft 56: 283-284.
Theowald, B., C. Dufour \& P. Oosterbroek, 1982. The zoogeography of the western Palaearctic Tipulidae (Diptera). Part IV: the Tipulidae of Corsica and Sardinia with a note on Dolichopeza fuscipes Bergroth. Mitteilungen der Schweizerischen Entomologischen Gesellschaft 55: 317-332.
Theowald, B. \& P. Oosterbroek, 1981. Zur Zoogeographie der westpalaearktischen Tipuliden II. Die Tipuliden der iberischen Halbinsel (Diptera, Tipulidae). - Beaufortia 30: 179-192.
Wood, H. G., 1952. The crane-flies of the South-West Cape (Diptera, Tipuloidea). - Annals of the South African Museum 39: 1-327.

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## Appendix A

In this appendix taxonomic changes pertaining to Savtshenkia are dealt with. The changes include the description of three new western Palaearctic species of the subgenus, the establishment of five synonymies, the elevation of one subspecies to species rank, and the removal of one species from Savtshenkia.

## Descriptions of new species

Tipula (Savtshenkia) alphasp.n.
(figs. 152-170)
Type material. - Holotype $\delta$ : Rumania, Mara-


Figs. 152-164, Tipula (Savtshenkia) alpha sp. n. - 152, male head and appendages, lateral view; 153, male head, dorsal view, antennac and mouthparts omitted; 154 , female antenna, basal segments; 155 , male terminalia, lateral view; 156 , male tergite 9-10, dorsal view; 157, male tergite 9-10, posterior view; 158 , male sternite 8 , posterior margin, ventral view; 159 , right sp2, dorsal view; 160, left outer gonostylus, lateral view; 161, left inner gonostylus, lateral view; 162, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 163, sperm pump and appendages, lateral view; 164 , sperm pump and appendages, anterior view.
mures, Muntii Rodnei, Borsa, subalpine lake 'Stiolul', main sources of river Bistrita Aurie, 1700$1800 \mathrm{~m}, 31 . \mathrm{V} .1974$, L. Botosaneanu (zman). Paratypes $50^{\circ}, 1$ ? same data as holotype (zMAN). Material preserved in alcohol.

## Description

Body length $10-12 \mathrm{~mm}$ (\$), 14 mm (f); wing length $12-13.5 \mathrm{~mm}$ ( $\delta$ ), 13.5 mm ( (\%); antennal length $5-5.5 \mathrm{~mm}(\delta)$ ) ? mm ( $\circ$; antenna broken, fig. 154).

Colour. - Ground colour greyish brown. Head and thorax dark greyish. Antenna with scapus dark greyish, remainder of segments dark brown. Palpus dark brown. Prescutum with single broad medial and pair of lateral dark stripes. Wing veins brown; pterostigma brown, distinct; membrane unpatterned. Haltere yellowish brown, including knob. Trochanters of same dark greyish colour as coxae, femora yellowish brown in basal section, from about half length darkening towards tip, remainder of legs dark brown. Abdomen dark brown, terminalia slightly darker. (Colour de-


Figs. 165-170. Tipula (Savtshenkia) alpha sp. n., female. - 165, ovipositor, lateral view; 166, sternite 8 and hypogynial valves, ventral view; 167, sternite 8 and hypogynial valves, dorsal view; 168 , sternite 9 , medial part, ventral view; 169 , genital fork. dorsal view; 170 , spermathecae.
duced by approximation, material preserved in alcohol).

Head (figs. 152-154). - Rostrum shorter than remainder of head, nasus well developed; eyes small, dorsally and ventrally separated by about four times diameter of scapus; occiput perpendicular to lateral sides of head (dorsal aspect, fig. 153); male antenna about four times as long as head, longest verticils about one and a half times as long as diameter of flagellomeres at base; female antenna shorter (flagella of both antennae broken beyond basal flagellomeres in female paratype, fig. 154).

Thorax. - Wing fully developed in both sexes, squama devoid of macrotrichia; cell m 1 gradually widening towards wing margin. Tarsal claws toothless in both sexes.

Male terminalia (figs. 155-164). - Compactly built, tergite $9-10$ and gonocoxites separated by membranous zone (fig. 155). Tergite 9-10 (figs. 156, 157) broad and short, with medial membranous area occupying about one third of total width; posterior margin widely V-shaped, with shallow emargination near median line; margin ventrally set with scattered hairs, without spines; lateral corners broadly rounded. Posterior margin of sternite 8 unmodified (fig. 158). Gonocoxites ventrally separated by membranous area, membrane with pair of weakly sclerotized bands extending anteriorly from base of aedeagal guide. Suture x narrow, about one fourth length of gonocoxite (fig. 155). Sp2 (fig. 159) long, medial blackish sclerotized margin moderately serrate; dorsally separate from tergite 9-10; connexion between sp2 and posterior margin of foramen of gonostyli short (fig. 159, arrow). Outer gonostylus (fig. 160) broad, its tip anteriorly inclined, posterior margin only moderately blackened.

Inner gonostylus (fig. 161) with relatively slender blackish sclerotized anterior part; lateral rim in its ventral part directed towards base of gonostylus; posterolaterally near base with blackened rim (fig. 161, arrow). Aedeagal guide (fig. 162) a short and plump blackish sclerotized structure, gonapophysis reduced to a sclerotized strip (fig. 162, arrow). Fragmentum (fig. 162) a rounded lobe with a few long hairs dorsomedially. Sperm pump (figs. 163, 164) with anterior apodemes short, posterior apodemes well developed and obliquely placed, compressor apodeme dorsally emarginate, blackish; lumen of endophallus well developed; ventral wall convex. Aedeagus short, anteriorly reaching segment 8 (fig. 155), tubular throughout, gradually narrowing towards tip.

Female terminalia (figs. 165-170). - Cercus and hypogynial valve well developed, robust, relatively short (fig. 165). Sternite 8 dorsolaterally near base of hypogynial valve with concavity (fig. 165, arrow); ventral membrane in posterior part of sternite 8 enclosing medial base of hypogynial valves (fig. 166). Dorsomedial margin of hypogynial valve only moderately blackened (fig. 167). Sternite 9 with slender medial projection (fig. 168). Genital fork elongate, anteriorly broader (fig. 169). Three spermathecae, globular to oblong, of about same size, moderately sclerotized (fig. 170).

Etymology. - The name of this species was prompted by its phylogenetic position within Savtshenkia and is the pendant of Tipula (Savtshenkia) omega sp. n. A noun in apposition.

Remarks. - Tipula (Savtshenkia) alpha differs from its presumed sister species $T$. (S.) grisescens by the


Figs. 171-176. Tipula (Savtshenkia) omega sp. n., male. - 171, head and appendages, lateral view; 172, head, dorsal view, anrennae and mouthparts omitted; 173, terminalia, lateral view; 174, tergite 9-10, dorsal view; 175, tergite 9-10, posterior view; 176, left sclerotization proctiger, lateral view.
colour of the coxae (dark greyish in alpha, yellowish brown in grisescens), the less elongate male antenna (see figs. 9 and 152), the separated gonocoxites (fused in grisescens), the broader outer gonostylus (see figs. 76 and 160), the more slender anterior part of the inner gonostylus and the presence of a sclerotized rim posterolaterally at the base of the same structure, the shallowly emarginate compressor apodeme of the sperm pump (deeply emarginate in grisescens), the rounded body of the sperm pump (with conical anteroventral extension in grisescens).

The type locality of alpha lies above the tree line on the subalpine plain at an altitude of $1700-1800 \mathrm{~m}$.

The locality has a characteristic vegetation of Pinus mugo Turra and the grass Nardus stricta Linnaeus. At the end of May thaw has set in, but the surface of the area is still extensively snow- and ice-covered. Under sunny conditions alpha emerges in large numbers as soon as holes begin to appear in the ice-covering. Although fully winged, the insects tend to crawl around on the ice and surrounding grasses. Under cloudy conditions alpha shows no sign of activity, suggesting that the behaviour of the species is largely influenced by the environmental temperature. In these aspects, the biology of alpha resembles that of the trichopteron Chionophylax czarnohoricus Dziedz.,


Figs. 177-184. Tipula (Savtshenkia) omega sp. n., male. - 177, sternite 8, posterior margin, ventral view; 178, sternite 8, posterior margin, posterior view; 179 , right sp2, dorsal view; 180, left outer gonostylus, lateral view; 181, left inner gonostylus, lateral view; 182, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 183, sperm pump and appendages, lateral view; 184, sperm pump and appendages, anterior view.
a species which occurs in the same locality (Dr Botosaneanu, pers. comm.).

## Tipula (Savtshenkia) omega sp. n.

 (figs. 171-184)Type material. - Holotype ô: Italy, Sardinia, Fiume Taloro, $670 \mathrm{~m}, 9^{\circ} 31 / 40^{\circ} 06,21 . \mathrm{X} .1981, \mathrm{H}$. Malicky (zman). - Paratypes: 10 , Italy, Sardinia, Rio Aratu, $970 \mathrm{~m}, 9^{\circ} 15 / 40^{\circ} 02$, 22.X.1981, H. Malicky (ZMAN). Material preserved in alcohol.

## Description

Body length $12-14.5 \mathrm{~mm}$ (す); wing length 14 15.5 mm ( $\delta^{\circ}$ ); antennal length $5-6 \mathrm{~mm}$ ( $\mathrm{\delta}^{\text {o }}$ ).

Colour. - Ground colour yellowish brown. Vertex of head and dorsal and lateral sides of thorax darker, probably slightly greyish pruinose in dry specimens. Antenna yellowish brown, scapus and pedicel somewhat lighter coloured than flagellum. Palpus yellowish brown. Presutum with confluent medial and pair
of lateral dark stripes indicated. Wing veins yellowish brown; pterostigma yellowish brown, distinct; membrane vaguely patterned. Halteres yellowish brown, including knob. Legs yellowish brown, femora and tibiae darkened at extreme tip, tarsi darkening towards tip. Abdomen yellowish at base, darkening towards tip. (Colour deduced by approximation, material preserved in alcohol).

Head (figs. 171, 172). - Rostrum about as long as remainder of head, nasus well developed; eyes well developed, dorsally separated by about two and a half times diameter of scapus, ventrally by about two times diameter of scapus; occiput gradually narrowing towards cervix (dorsal aspect, fig. 172); male antenna just less than two times length of head, longest verticils about as long as flagellomeres (fig. 171).

Thorax. - Wing fully developed, squama devoid of macrotrichia; cell ml elongate bell-shaped, narrowing towards wing margin. Tarsal claws with small medial teeth in male.

Male terminalia (figs. 173-176). - Large, tergite 9-


Figs. 185-191. Tipula (Savtshenkia) trinacria sp. n., male. - 185, terminalia, lateral view; 186, tergite 9-10, dorsal view; 187, tergite 9-10, posterior view; 188, left sclerotization proctiger, lateral view; 189, sternite 8, posterior margin, ventral view; 190, sternite 8, posterior margin, posterior view; 191, right sp2, dorsal view.

10 and gonocoxites separated by membranous zone (fig. 173). Tergite 9-10 (figs. 174, 175) elongate, with narrow medial membranous area; posterior margin with pair of broad laterally diverging extensions; extensions not blackish sclerotized, devoid of spines along ventral margin. Proctiger with rather diffuse and narrow lateral sclerotization (fig. 176). Sternite 8 (figs. 173, 177, 178) posteriorly with diverging broad dorsoventrally flattened lobes, each lobe carrying about $35-40$ black spines. Gonocoxites ventrally fused with narrow medial carina; at level of insertion of inner and outer gonostyli with concentration of
long and strong setae (fig. 173). Suture x narrow, about two-fifth length of gonocoxite (fig. 173). Sp2 (fig. 179) medially protruding, blackish sclerotized medial margin entire; sp2 fused with tergite 9-10; connexion between sp2 and posterior margin of foramen of gonostyli long. Outer gonostylus (fig. 180) with elongate slender anterior part, posterior blackish sclerotized part relatively broad, with short posterodorsal extension, slightly concave. Inner gonostylus (fig. 181) with anterior and posterior parts about as long, posterior part caudally terminating in a densely pubescent lobe. Aedeagal guide (fig. 182) long and


Figs. 192-196. Tïpula (Savtshenkia) trinacria sp. n., male. - 192, left outer gonostylus, lateral view; 193, left inner gonostylus, lateral view; 194, aedeagal guide with right gonapophysis and fragmentum, posteromedial view; 195, sperm pump and appendages, lateral view; 196, sperm pump and appendages, anterior view.
slender, blackish sclerotized, gonapophysis short, dorsally directed. Fragmentum (fig. 182) anteriorly with sclerotized bar which dorsally carries a bundle of long hairs, posteriorly weakly sclerotized. Sperm pump (figs. 183, 184) with large anterior apodemes, posterior apodemes caudally directed, compressor apodeme dorsally deeply emarginate, wings broad; lumen endophallus small, body wall ventrally concave. Aedeagus elongate (fig. 173), anteriorly reaching segment 3, tubular throughout.

Etymology. - The phylogenetic position of this species within Savtshenkia suggested its name. A noun in apposition.

Remarks. - The species omega very much resembles the species mannheimsi and trinacria sp . n. It differs from these species by the number of spines on the posterior lobes of male sternite 8 (ca. 35-40 in omega to ca. 50 in mannheimsi and ca. 65 in trinacria), the absence of spines on the ventral margin of male tergite 10 in omega (spines absent also in mannheimsi, present in trinacria), and in the shape of the inner and outer gonostyli.

The female of omega remains unknown.

Tipula (Savtshenkia) trinacria sp. n.
(figs. 185-202)
Type material. - Holotype ô: Italy, Sicily, Nébrodi, 10 km north of Capizzi, Fiume Troina, 1200-1400 m, 11-14.X.1993, P. Oosterbroek \& C. Hartveld (zman). - Paratypes: 13ó, 3 ㅇ, same data as holotype; $4 \mathbf{\delta}^{\hat{\prime}}$, Italy, Sicily, Nébrodi, 5 km north of Monte Soro, Biviere di Cesaro, 1300 m, 12.X. 1993, P. Oosterbroek \& C. Hartveld; 20, Italy, Sicily, Nébrodi, 6 km east of Serra del Re, Foresta Vécchia, Fiume di Saracena, 1100 m, 13.X.1993, P. Oosterbroek \& C. Hartveld; 40 , 3 우, Italy, Sicily, Madonie, 6 km north of Monte San Salvatore, Torrente Vicaretto, $1300 \mathrm{~m}, ~ 15-21 . X .1993, \mathrm{P}$. Oosterbroek \& C. Hartveld, (zman).

## Description

Body length $13.5-16 \mathrm{~mm}$ (ot), $15.5-17 \mathrm{~mm}$ ( $\%$ ); wing length $16-19 \mathrm{~mm}\left(\delta^{\top}\right), 16.5-18 \mathrm{~mm}$ (o); antennal length 6-7 mm (ð), 5-5.5 mm (¢).

Colour. - Ground colour yellowish brown. Excepting rostrum, head greyish pruinose. Antenna with yellowish scapus and pedicel, flagellum almost uniformly brownish, basal flagellomeres somewhat lighter. Palpus yellowish brown. Dorsal and lateral sides of thorax slightly greyish pruinose. Presutum


Figs. 197-202. Tipula (Savtshenkia) trinacria sp. n., female. - 197, ovipositor, lateral view; 198, sternite 8 and hypogynial valves, ventral view; 199, sternite 8 and hypogynial valves, dorsal view; 200, sternite 9, medial part, ventral view; 201, genital fork, dorsal view; 202, spermathecae.
with pair of approximating slender medial and pair of lateral dark stripes, medial stripes anteriorly lighter coloured. Wing veins brown; pterostigma brown, distinct; membrane vaguely patterned and iridescent. Halteres with yellowish brown pedicel, knob brown. Legs yellowish brown, femora and tibiae darkened at extreme tip, tarsi largely dark brown. Abdomen yellowish at base, darkening towards tip.

Head. - Very much as that of omega sp. n. (see figs. 171, 172; see for description of head under that species, above).

Thorax. - Wing fully developed in both sexes, squama devoid of macrotrichia; cell m 1 elongate bellshaped, narrowing towards wing margin. Tarsal claws with small medial teeth in the male, female claws toothless.

Male terminalia (figs. 185-196). - Large, tergite 910 and gonocoxites separated by membranous zone (fig. 185). Tergite 9-10 (figs. 186, 187) elongate,
with narrow medial membranous area; posterior margin with pair of broad laterally diverging extensions; extensions blackish sclerotized along ventral margin, ventral margin set with small black spines. Proctiger with rather well demarcated slender lateral sclerotization (fig. 188). Sternite 8 (figs. 189, 190) posteriorly with diverging broad flattened lobes, each lobe carrying about 65 slender black spines. Gonocoxites ventrally fused with narrow medial carina. Suture x narrow (fig. 185). Sp2 (fig. 191) medially protruding, blackish sclerotized medial margin entire; sp2 fused with tergite 9-10; connexion between sp 2 and posterior margin of foramen of gonostyli long. Outer gonostylus (fig. 192) with elongate slender anterior part, posterior blackish sclerotized part elongate, upcurved. Inner gonostylus (fig. 193) with anterior and posterior parts about as long, posterior part anteriorly extended, posteriorly terminating in cup-shaped structure. Aedeagal guide (fig. 194) long and slender,
blackish sclerotized, gonapophysis short, dorsally directed. Fragmentum (fig. 194) anteriorly with sclerotized bar which dorsally carries a bundle of long hairs, posteriorly weakly sclerotized. Sperm pump (figs. 195, 196) with large anterior apodemes, posterior apodemes caudally directed, at tip somewhat upturned, compressor apodeme dorsally deeply emarginate, with narrow cleft, wings broad; lumen endophallus small, wall ventrally concave. Aedeagus elongate (fig. 185), anteriorly reaching segment 4 , tubular throughout.

Female terminalia (figs. 197-202). - Cercus and hypogynial valve well developed, cercus approximately straight (fig. 197). Ventral membrane in posterior part of sternite 8 terminating in between bases of hypogynial valves (fig. 198). Dorsomedial margin of hypogynial valve only moderately blackened (fig. 199), ventrally at base with concavity (figs. 197, 198). Sternite 9 with relatively narrow and slender medial projection, concave lateral of this structure (fig. 200). Genital fork short and broad, broadest at anterior side (fig. 201). Three spermathecae, globular, of about same size, blackish sclerotized (fig. 202).

Etymology. - Trinacria is an old name of Greek origin for Sicily. A noun in apposition.

Remarks. - The species trinacria very much resembles mannheimsi and omega sp. n. For distinguishing characters see under omega. The type specimens of trinacria were captured along rivers and a lake in deciduous forests. The specimens collected in the period from 15-21 October were taken from the moss covered river banks of the Torrente Vicaretto. Among these latter specimens was one pair in copula.

## Synonymy

insignifica Alexander, 1924 = invenusta invenusta Riedel, 1919 syn. n.

Alexander (1924) described insignifica as a new species based on a single male originating from Mount Washington, New Hampshire, U.S.A. Alexander (loc. cit.) recorded the collection of bSNH as the depository of the holotype (holotype not examined). I have studied specimens labelled by Alexander as metatypes, i.e. specimens compared with the type specimen(s), originating from Mount Katahdin, Maine and Mount Madison, New Hampshire (USNM). Examination of these specimens revealed only slight differences between the Nearctic insignifica and the Palaearctic invenusta invenusta in the distance of the eyes on the ventral side of the head (see also 'Discussion of adopted phylogeny'). No other structural differences between insignifica and invenusta invenusta were found and I therefore consider insignif-
ca a junior synonym of invenusta invenusta. [Contrary to Alexander's $(1942,1966 \mathrm{~b})$ assertion, the posterior margin of tergite 9-10 is posteroventrally armed with small black spines in insignifica, as it is in invenusta].
broweri Alexander, 1940 = fragilina Alexander, 1919 syn. n.

Alexander (1940) described broweri based on 12 males and two females from Mount Katahdin, Maine, U.S.A. (types in UsNM). Despite Alexander's claim that broweri should be entirely distinct from its relatives, examination of type material of broweri showed that it is morphologically identical with fragilina. I therefore consider broweri a junior synonym of fragilina.
nebulipennis Alexander, 1919 = alpium Bergroth, 1888 syn. n.

Alexander (1919a) introduced nebulipennis as a new species based on two males from Battle Harbour, Labrador, Canada (holotype in Brooklyn Museum, New York, U.S.A., paratype in USNM). Later, nebulipennis was also recorded from Quebec and New Hampshire (Alexander 1965b). Examination of material of nebulipennis in the Alexander collection (USNM) revealed no significant morphological differences between these specimens and the European species at pium. I therefore consider nebulipennis a junior synonym of alpium. The larvae of alpium were recorded from terrestrial mosses (Brindle 1958, 1959, 1960). Theowald $(1957,1967)$ recorded the larvae furthermore from the litter layer beneath oaks. Possibly alpi$u m$ has been introduced into North America in a way similar to the European species Tipula (Tipula) paludosa Meigen (see Alexander 1965a).
packardi Alexander, 1928 = invenusta invenusta
Riedel, 1919 syn. n.
Alexander (1928) described Tipula packardi based on a single male from Hopedale, Labrador, Canada. The holotype of packardi is preserved in CNCI (holotype not examined). An additional male specimen originating from the type locality of packardi is present in USNM. This latter specimen was compared by Alexander with the holotype of packardi and considered conspecific. Examination of the metatype of packardi in USNM showed that it has the eyes on the ventral side of the head slightly more approximate than the examined specimens of invenusta invenusta (see also 'Discussion of adopted phylogeny', and $i n$ significa, this section). No other structural differences were found and I therefore consider packardia junior synonym of invenusta invenusta.
perparvula Alexander, 1926 = ignobilis Loew, 1863 syn.n. (partim?)

Loew (1863) described the species ignobilis based on the female sex from material captured by OstenSacken in the District of Columbia, U.S.A. (holotype in mczc, not examined). According to its labels, Alexander compared a male specimen captured by Osten Sacken in Catskill, New York, with the type specimen of ignobilis and considered the two conspecific. Examination of this inferred male specimen of ignobilis and the paratype male of the species perparvula (both in USNM) revealed that these two specimens belong to the same species. Alexander (1926), describing perparvula based on two males originating from Aweme, Manitoba, Canada (holotype in CNCI, paratype in USNM), explicitly stated that perparvula has a relatively stout nasus. Examination of the paratype of perparvula shows that this structure is entirely absent in the specimen, as it is in the species ignobilis. Either the holotype and paratype of perparvula belong to two different species, with the paratype being conspecific with ignobilis, or both type specimens are conspecific and belong to ignobilis. Examination of the holotype of perparvula should settle this question.

## Subspecies raised to species rank

## baennii Dufour, 1991

This taxon was originally described as a subspecies of subsignata (Dufour 1991). The tip of the aedeagal guide of haennii does not show the abrupt constriction which probably represents a synapomorphy of a monophyletic group containing corsosignata, subsignata subsignata and subsignata cazorla (character 8, above). Because of its phylogenetic position next to this clade, haennii is elevated from subspecific to specific rank.

## Species removed from Savtshenkia

convexifrons Holmgren, 1883
Savchenko (1961) placed the species convexifrons Holmgren in his subnodicornis group of Pterelachisus sensu Savchenko, nec Rondani. Savchenko (1961: 223) based its membership of the subnodicornis group on the shape of the inner gonostylus, and its incorporation in Pterelachisus sensu Savchenko on the proximal position of the fork of vein $\mathrm{M} 3+4$. At the same time, Savchenko underlined the isolated position of convexifrons within this grouping. As noticed in the introduction to the present paper, the name Savtshenkia replaces Pterelachisus sensu Savchenko. Theowald (1973) followed Savchenko in considering convexifrons a species of Savtshenkia, but split the subnodicornis group sensu Savchenko into two species groups (table 3). I remove convexifrons from Savtshenkia because of the absence of the defining apomorphies 53 and 83 of Savtshenkia. Furthermore,
the species convexifrons differs from the species of Savtshenkia by the shape of male tergite $9-10$, the ventral structure of the fused gonocoxites, the shape of the elongate aedeagal guide and its gonopophyses, and, mose noteworthy, by the presence of a complete genital bridge (a plesiomorphy; the medial part of the genital bridge is absent in all species of Savtshenkia). The fork of vein $\mathrm{M} 3+4$ is located proximally in many genera and subgenera of Tipulidae other than Savtshenkia. The resemblance of the inner gonostylus of convexifrons to the inner gonostyli of plesiomorphous species of Savtshenkia is only superficial. At the moment I am unable to establish the actual phylogenetic relationships of convexifrons and can not allocate it to any other subgenus or species group of Tipulidae.

## Appendix B

In this appendix are listed the species and subspecies of Savtshenkia which, because of deficiency of data, were not incorporated in the character state matrix given in table 4. Incompleteness of data resulted of several causes. For some of the pertaining taxa no material was examined, for others the material examined was incomplete as regards the structures of the male terminalia, for others again only the female is known, thus excluding their incorporation in a phylogeny which is largely based on characters of the male terminalia. Where possible, the presumed phylogenetic position of the species and subspecies concerned is indicated. For the deposition of the type material of species described by Savchenko see Kandybina, Lantzov \& Savchenko (1987).

## Unexamined

imperfecta Riedel, 1914. - Description based on single brachypterous male specimen from Kenya (Mount Kenya) and single brachypterous female specimen, possibly belonging to the same species, from Tanzania (Mount Kilimanjaro). Types in mnhn. Species doubtfully belongs to Savtshenkia. Distribution: Kenya, Tanzania.
letifera Alexander, 1951. - Description based on single male from China (Tibet, Shugden Gompa a.k.a. Hsiu-teng). Type in bmnh. Inner and outer gonostyli quite similar to those of Kashmirian sordidipes and venerabilis. Differences with these species in structures of posterior margins of tergite 9-10 and sternite 8 . Shape of sternite 8 suggests relationship with Nearctic graciae and Palaearctic kiushiuensis and koreana. Alexander (1953) illustrated details male terminalia of letifera. Distribution: China (south-east Tibet).
minuscula Savchenko, 1971. - Originally described as Tipula (Savtshenkia) nana by Savchenko (1968a) based on two males and single female from USSR (Kamchatka). Types in zlas. Savchenko, probably
correctly, considered minuscula closely related to Nearctic ignobilis, with which it should differ in colour characteristics. Details male terminalia of mi nuscula illustrated by Savchenko (1968a). Distribution: Russia (Kamchatka).
nivalis Savchenko, 1961. - Description based on three males and four females from USSR (Stavropolskiy kray and Gruzinskaya SSR). Types in zias. Savchenko (1961) compared nivalis with glaucocinerea. Theowald (1973) considered both very closely related and suggested that nivalis might be considered a subspecies of glaucocinerea. Details male terminalia of nivalis illustrated by Savchenko (1961) and Theowald (1973). Distribution: Russia (Stavropolskiy kray), Georgia.
productella Alexander, 1928. - Description based on single male from Canada (Labrador). Type in cnci. According to Alexander (1928) most closely allied to Nearctic fragilina. Structure of sternite 8 of productella of same build as found in clade fragilina to subvafra of present paper; productella probably belongs to this clade; synonymy with one of its species possible. Distribution: Canada (Labrador).
subalpium Savchenko, 1961. - Description based on single male from USSR (Krasnodarskiy kray, Glavniy Mountains). Type in zias. Savchenko (1961) compared subalpium with alpium, with which it shares same general colour and structure of inner and outer gonostyli and to which it is probably closely related. Differs quite remarkably from alpium in structure of posterior margin of sternite 8: whereas posterior margin of sternite 8 is U -shaped emarginate and carries spines along its edge in alpium (fig. 105), emargination is filled up with membrane covered with short setae in subalpium. According to Savchenko, there is a short pubescent extension in between what he denotes sternites 8 and 9. This structure could well be homologous with midventral extension found in alpium and macaronesica, which in dry material is often hardly visible or seems to originate from posterior margin of sternite 8 (see fig. 39). Savchenko (1969) depicted male terminalia of subapium in lateral and posterior view. Distribution: Russia (Krasnodarskiy kray), Georgia.
tetragramma Edwards, 1928. - Description based on single male from China (Yunnan). Type in bmnн. Edwards (1928) was of the opinion that tetragramma must bear a rather strong resemblance to kiushiuensis. Distribution: China (Yunnan).

## Incomplete male terminalia

kiushiuensis Alexander, 1925. - Description based on single male and single female from Japan (Kyushu, Mount Kirishima). Types in usnm. Glass slides with terminalia male holotype and additional male specimen (USNM) do not allow study of all relevant struc-
tures. Related to species in range jeekeli to aster of cladogram fig. 2. Posteriorly produced unarmed medial part of posterior margin male sternite 8 suggests relationship with koreana. Savchenko (1968b) illustrated details male terminalia of kiushiuensis. Distribution: Russia (Primorskiy kray), Japan (Kyushu, Shikoku).
koreana Alexander, 1934. - Description based on single male and four females from North Korea (Mount Kongo). Types in usnm. Glass slide with terminalia male holotype does not allow study of all relevant structures. Related to species in range jeekeli to aster of cladogram fig. 2. Posteriorly produced unarmed medial part of posterior margin male sternite 8 suggests relationship with kiushiuensis. Alexander (1934) illustrated details male terminalia and wing of koreana. Distribution: Russia (Sakhalin), North Korea, Japan (Shikoku).
mohriana Alexander, 1954. - Description based on single male and single female from Japan (Shikoku, Mount Ishizuchi). Types in UsNm. Glass slide with terminalia male holotype does not allow study of all relevant structures. Most probably sister species of clade fragilina to subvafra of present paper as implied by fused sp2 and tergite 9-10 (character 54), slender anterior part outer gonostylus (character 45), bifid tip of anterior part of inner gonostylus (character 30), and widely emarginate bilobed and unarmed posterior margin of male sternite 8 (cf. character 60). Differing most notably from species of clade fragilina to subvafra by presence of spinous gonapophysis (autapomorphy) and absence of blackened dorsal margin of fragmentum (cf. character 17). Details male terminalia and wing of mohriana illustrated by Alexander (1955). Distribution: Japan (Shikoku).
sciadoptera Alexander, 1964. - Described as guttulifera by Alexander (1961) based on two males from India (Himalayas). Types in usnm. Glass slides with terminalia male holotype and paratype do not allow study of all relevant structures. Probably related to species in range baltistanica to chrysocephala of cladogram in fig. 2. Alexander (1961) illustrated details male terminalia of sciadoptera (as guttulifera). Distribution: India (Sikkim).

## Known in female sex only

aberdareica ulugurica Alexander, 1962. Description based on single female from Tanzania (Uluguru Mountains). Type in Mrac, not examined. According to Alexander (1962) closely related to nominotypical form, sharing several colour characteristics with chrysocephala. Distribution: Tanzania.
nephrotomoides invariegata Alexander, 1956. Description based on single female holotype from Uganda (Ruwenzori Range). Type in BMNH. Agreeing in its dark colour with the nominotypical form from Zaire and Uganda. Distribution: Uganda.

## Appendix C

## Autapomorphies of the species and subspecies of

## Savtshenkia

aberdareica. no autapomorphy recognized.
aberdareica ulugurica no autapomorphy recognized (known of female holotype only).
akeleyi: inner gonostylus with dorsal margin anterior part undulating.
alphar posterior rim of inner gonostylus ventrally blackish sclerotized (fig. 161, arrow).
alpium: shape of posterior margin of male tergite 9-
10 ; shape of posterior margin of male sternite 8 (fig.
105); shape of extension of midventral area (long; fig. 39).
asbolodes: inner gonostylus with anterior part rather slender.
aspromontensis: shape of inner gonostylus (crest, fig. 61); shape of male tergite 9-10 (two pairs of pointed extensions, fig. 135).
aster. shape of aedeagal guide; gonapophysis slender, downcurved and pointed; shape of male tergite 9-10; shape of posterior margin of male sternite 8 (fig. 109).
atlas: extensions of male tergite 9-10 mediolaterally compressed.
baltistanica: arrangement of setae on posterior margin of male sternite 8.
benesignata: shape of inner gonostylus (fig. 72).
boreosignata: broad membranous zone of tergite 9-
10 (fig. 138); reduction of spines on lobes on posterior margin male sternite 8 ; fragmentum with low anterior part; shape of aedeagal complex (fig. 99).
breviantennata: anterior part of inner gonostylus widening towards apex, lateral ridge of inner gonostylus short and angled.
caligo: shape of posterior margin of male tergite 910.
cheethami: midventral area posteriorly with long conical extension (fig. 35).
chrysocephala: anterior part of inner gonostylus constricted before tip (lateral view).
confusa: inner gonostylus with two blackish sclerotized dentate ridges on lateral side of posterior part (fig. 58); male tergite 9-10 with truncate extensions close together; shape of posterior margin of male sternite 8.
corsosignata: shape of inner (fig. 69) and outer gonostyli.
cyrnosardensis: inner gonostylus caudodorsally with posteriorly produced pubescent lobe; outer gonostylus with slender extension on posterior margin at about two-thirds from base; apex of elongate fragmentum medially curved.
draconis shape of inner gonostylus; reduced nasus. eleonorae. no autapomorphy recognized.
elgonensis. no autapomorphy recognized
eugeni: extensions of male tergite 9-10 upcurved (lateral view); shape of inner gonostylus; anterior apodemes sperm pump elongate and dorsally directed, compressor apodemes mediolaterally flattened in apical part.
fragilina: inner gonostylus with acute sclerotized extension at ventral tip of posterior ridge.
fragilis: aedeagal guide elongate and slender (fig. 22); suture $x$ broad, sigma-shaped.
gimmerthali: inner gonostylus with anterior part gradually sloping towards bifid apex (fig. 66).
gimmerthali mattheyi: outer gonostylus without posterior sclerotized appendage (present in both other subspecies).
gimmerthali pteromaculata: inner gonostylus with relatively broad anterior part (fig. 67); outer gonostylus broadened above midlength (fig. 84).
glaucocinerea: shape of extensions of tergite 9-10.
goriziensis shape of inner gonostylus (fig. 54).
graciae posterior margin of male sternite 8 medially with rounded extension set with long and strong setae.
grisescens. extensions of male tergite 9-10 dorsoventrally flattened (figs. 121, 122).
baennii: fragmentum 3-lobed (fig. 27); inner gonostylus with ridge separated in posterior and ventral part, ventral part ridge extended over posterior part of inner gonostylus, and dorsal crest anteriorly abruptly terminating (fig. 68).
bancocki: male antenna reduced in length.
hartigiana: shape of inner gonostylus (long anterior part, strongly protruding lateral ridge); extension of midventral area gonocoxites (fig. 33).
holoptera: posterior margin of male sternite 8 with a few strong setae (fig. 107).
ignobilis: shape of outer gonostylus; shape of sp2.
imperfecta: wings reduced in both sexes.
interserta: shape of inner gonostylus; outer gonostylus large, posteriorly inclined.
invenusta: shape of posterior margin of male tergite 9-10 (fig. 136).
invenusta microinvenusta: inner gonostylus anteriorly more curved forward than in both other forms; differing primarily in smaller size and darker colour from both other forms.
invenusta subinvenusta: inner gonostylus relatively slender.
jeekeli: shape of inner gonostylus; shape of outer gonostylus (fig. 80); shape of posterior margin of male sternite 8 ; sperm pump with sclerotized sperm duct; apex of aedeagus with spines.
kiushiuensis: shape of posterior margin of male sternite 8 .
koreana: shape of posterior margin of male sternite 8.
letifera: no autapomorphy recognized (probably very similar to sordidipes and venerabilis).
limbata: anterior part of inner gonostylus produced into long and bifid extension (fig. 64).
lundbladi: shape of inner gonostylus (elongate, with short crest); shape of outer gonostylus (fig. 78); shape of posterior margin of sternite 8 (fig. 103); surface of sp2 anterodorsally with cone-shaped extension.
macaronesica: shape of extension of midventral area (short).
mannheimsi: outer gonostylus with incision ventral of posterior sclerotized spur.
minuscula: no autapomorphy recognized (probably very similar to ignobilis).
mobriana: gonapophysis as slender curved spine.
multipicta: shape of inner gonostylus (high anterior part with long and slender anterior apex); shape of outer gonostylus (oblong).
nephrotomoides: blackish species.
nephrotomoides invariegata: no autapomorphy recognized (known of female holotype only).
nielseni: shape of inner gonostylus; shape of outer gonostylus (fig. 85); shape of medial extension of male sternite 8 .
nivalis: no autapomorphy recognized (probably very similar to glaucocinerea).
obsoleta: shape of inner and outer gonostyli; shape of posterior margin of male sternite 8 (fig. 106); spinous extensions on posterodorsal part fragmentum (fig. 21).
odontostyla: shape of male tergite $9-10$; inner gonostylus dorsally with long and dense pubescence (fig. 63); apical part of fragmentum bent and posteriorly directed; gonapophysis twisted.
omega: inner gonostylus with hump in anterior half of posterior part (fig. 181).
ornata: wing cell a 2 almost symmetrical in distal tip; posterior extensions male tergite 9-10 dorsoventrally flattened, laterally curved, ventrally carrying long thin spinous setae; shape of posterior margin of male sternite 8 ; midventral anterior extension of male sternite 9 elongate, membranous; shape of appendage of midventral area gonocoxites (fig. 40); suture x broad, enclosing isolated sclerite; female tergite 9 medially reduced to strip.
pagana: female brachypterous.
pechlaneri: male tergite 9-10 with pair of low extra extensions lateral of usual extensions on posterior margin; shape of inner gonostylus (fig. 65).
persignata: posterior part of inner gonostylus corrugated (fig. 73).
persignata tofina: differing in details of male terminalia from nominotypical form.
phoroctenia: shape of male tergite 9-10; shape of posterior margin of male sternite 8; lateral ridge of inner gonostylus posterodorsally angular; suture x of
gonocoxite Y -shaped; shape of extension of midventral area gonocoxites (fig. 41).
postposita: shape of outer gonostylus.
productella: no autapomorphy recognized.
rufina: extensions of tergite 9-10 extremely long and slender; shape of inner gonostylus; tip of gonapophysis medially curved; posterior margin of male sternite 8 bulbously curved.
rufina maderensis: larger than nominotypical species.
sardosignata: shape of inner gonostylus; shape of outer gonostylus (fig. 88); shape of posterior margin of male sternite 8 .
sciadoptera: shape of inner gonostylus.
serrulifera: shape of inner gonostylus (fig. 60); shape of outer gonostylus; shape of posterior margin of male sternite 8 .
signata: posterior part of inner gonostylus anterior-
ly rounded (fig. 74); midventral extension on posterior margin male sternite 8 large, with dorsal sclerite (fig. 116).
simulans: inner gonostylus with lateral ridge dorsally produced into pointed extension; shape of extension of midventral area.
sordidipes. legs extensively blackened.
staegeri: extensions of male tergite 9 widely separate, mediolaterally flattened; male sternite 8 midventrally conical; posterior extensions of male sternite 8 elongate; aedeagal guide posteriorly curved; shape and length of gonapophysis; shape of inner gonostylus; female sternite 8 dorsally extended; base of hypogynial valve laterally bulging.
subalpium: shape of posterior margin of male sternite 8 .
subnodicornis: tergite 9-10 with extension in between lateral pair (fig. 128).
subsignata: together with next subspecies distinguished by lateral ridge being acutely bent and anteriorly produced in ventral part (fig. 70); differing from subsignata cazorla in details of inner gonostylus.
subsignata cazorla: differing from nominotypical subspecies by shape of inner gonostylus (anterior part dorsally abruptly narrowed, lateral view).
subvafra: shape of male tergite 9-10; shape of inner gonostylus.
tetragramma: wing pattern.
trinacria: outer gonostylus with extremely long posterior sclerotized appendage (fig. 192); sclerotization of proctiger well demarcated (fig. 188).
tulipa: inner gonostylus with relatively short lateral ridge; midventral area gonocoxites broad and long pubescent.
venerabilis. no autapomorphy recognized (very similar to sordidipes).
villenewvii: small and posteriorly curved outer gonostylus; gonocoxite posterodorsally acutely angled.

# THREE NEW SPECIES OF PSEUDEXECHIA 

TUOMIKOSKI FROM TANZANIA AND THAILAND

## (DIPTERA: MYCETOPHILIDAE)


#### Abstract

Kjxrandsen, J., 1994. Three new species of Pseudexechia Tuomikoski from Tanzania and Thailand (Diptera: Mycetophilidae). - Tijdschrift voor Entomologie 137: 325-330, figs. 1-4. [Issn 0040-7496]. Published 15 December 1994. Pseudexechia longistylus sp. n. and $P$. lanceostylus sp. n. from Tanzania and $P$. inthanonensis sp. n. from Thailand are described, based on adult males. The terminalia are figured. The presence of a pair of medium sized bristle-like dorsocentral setae on scutum in the two African species and an aberrant wing venation in $P$. inthanonensis sp.n. is commented on. J. Kjerandsen, Museum of Zoology, University of Bergen, Muséplass 3, N-5007 Bergen, Norway. Key words. - Diptera; Mycetophilidae; Pseudexechia; Tanzania; Thailand; taxonomy.


Pseudexechia Tuomikoski, 1966 constitutes one of 14 genera in the tribus Exechiini, subfamily Mycetophilinae (Tuomikoski 1966). The genus is characterised by the absence of large discal setae on the mesoscutum, by the ovate clypeus and on characters in the male terminalia such as the bud-like sternal process. Chandler (1978) treated all eight holarctic species known at that time. Three palaearctic species were added later; viz. P. trilobata Ostroverkhova, 1979, P. ussurensis Zaitzev, 1982 and P. altaica Zaitzev, 1988. Matile (1970) revised the Afrotropical Pseudexechia comprising four species. A fifth species, P. tanganyikae (Lindner, 1958) originally described as Exechia, was later referred to this genus as well (Matile 1980). No species of Pseudexechia are hitherto known from the oriental region. Thus, 16 species of the genus Pseudexechia were known: 9 palaearctic, 2 nearctic and 5 afrotropical. Of only one of the five Afrotropical species both sexes were described, the four other were all described on single females. Female holotypes make it difficult to describe new species from the region as certain association of the sexes is not easy to establish without large samples or reared specimens. However, based on differences in external characters as coloration and wing venation the present specimens can not belong to any of the described species.

## Material

The material consists of two males collected in the West Usambara Mountains in Tanzania in 1990, and one male collected in the Doi Inthanon mountain in

Thailand in 1991. The holotypes are deposited in the Museum of Zoology, Bergen (zMBN).

## Methods and terminology

The specimens were cleared and slide mounted in Canada balsam. The general terminology follows McAlpine (1981).

Wing measurements (fig. 1): Total wing length is measured from the extreme base of the distal median plate. $\mathrm{A}=$ distance between the points where $\mathrm{R}_{1}$ and $\mathrm{R}_{4,5}$ reach wing margin. $\mathrm{B}=$ distance between the points where $R_{4+5}$ and $M_{1}$ reach wing margin. $C=$ width of $M$-fork at wing margin. $\mathrm{D}=$ width of Cu fork at wing margin. $\mathrm{E}=$ distance from extreme base of the distal median plate to base of M -fork. $\mathrm{F}=$ distance from extreme base of the distal median plate to base of Cu -fork. Other measurements are self-explanatory and indicated in Fig. 1.

The tibial spur formula is the length of each spur in relation to the apical diameter of tibia in the following order: fore tibial spur; the two mid tibial spurs; the two hind tibial spurs.

## Systematic part

## Pseudexechia longistylus sp. n.

(figs. 1-2)
Type material. - Holotype male, tanzanta: Tanga region, W. Usambara Mis, Mazumbai, 1440 m a.s.l., 22.XI. 1990, G. E. E. Soli, sweep net (zmbn No. 179).

Diagnostic characters. - The male imago is charac-


Fig. 1. Pseudexechia longistylus sp. n., right wing showing measuring points. - Total wing length is measured from the extreme base of the distal median plate to tip of wing. $\mathrm{A}=$ distance between the points where $\mathrm{R}_{1}$ and $\mathrm{R}_{4.5}$ reaches wing margin. B $=$ distance between the points where $\mathrm{R}_{4,5}$ and $\mathrm{M}_{1}$ reaches wing margin. $\mathrm{C}=$ width of M -fork at wing margin. $\mathrm{D}=$ witdh of Cu-fork at wing margin. $\mathrm{E}=$ distance from extreme base of the distal median plate to point of M furcation. $\mathrm{F}=$ distance from extreme base of the distal median plate to point of Cu furcation. $\mathrm{M}-\mathrm{p} .=\mathrm{M}$-petiole.

Fig. 2. Pseudexechia longistylus sp. n., male terminalia. $-A$, ventral view; $B$, internal face of the gonostylus. Scale 0.50 mm .

terised by the enlarged terminalia in which the dorsal lobe of gonostylus ends in a very elongated lobe with a setose cushion basally, and by the shape of the bifurcate internal process. The ventral lobe of the gonostylus also is enlarged compared to other species.

Etymology. - From Latin, longus, long, and stylus, style, referring to the elongated gonostylus. A noun in apposition.

Description of male holotype. - Length of thorax 0.86 mm . Wing length 2.62 mm . Wing length / length of thorax 3.1. Wing length / length of fore femur 3.1.

Coloration. Antenna with scape, pedicel and basal part of flagellum yellow, gradually becoming darker and more greyish dusted towards tip. Head yellow. Maxillary palp greyish yellow. Thorax mainly yellow; scutum with narrow pale yellow or somewhat silvery shining stripe along lateral margin, delimited by darker margins below and above; scutellum darker yellow. Wing unmarked, yellow tinted, paler towards tip and hind margin. Wing veins yellow with dark setae. Halter greyish yellow. Legs mainly yellow; coxae paler yellow, hind coxa posteriorly with darker stripe. Abdominal tergites mainly yellowish brown; tergite 1 and 2 with pale lateral markings; tergite 3 to 5 with pale posterolateral markings, almost reaching middle; tergite 6 almost entirely dark. Terminalia mainly yellow, darker apically on gonocoxite, on sternal process and on parts of dorsal lobe of gonostylus.
Head. Total length of flagellum 1.35 mm . First flagellomere 1.5 times as long as second flagellomere. Second flagellomere 2.2 times as long as wide. Five large orbital setae. Small round median ocellus present. Diameter of lateral ocellus / diameter of median ocellus 1.7. Clypeus bearing 31 setae; width / length 0.8 . Fourth palpomere about 1.5 times as long as third palpomere.
Thorax. Pronotum with vertical row of 3 erect setae. Scutum with pair of medium sized bristle-like dorsocentral setae posteromedially in front of scutellum (prescutellar setae); anterior and lateral margin with large setae; otherwise uniformly clothed with small setae (but larger and fewer than in other Pseudexechia except in P. lanceostylus sp. n.). Scutellum with pair of strong scutellar setae and pair of very short incurved setae basad of them. Proepisternum with 1 strong, 1 medium sized and 5 small setae. Laterotergite with 6-7 large and about 50 small setae.

Wing (fig. 1). Crossvein $h$ with 1 ventral seta. $\mathrm{R}_{1}$ with 11 ventral setae apically. $\mathrm{R}_{4+5}$ with $26-27$ ventral setae apically. Wing length / length of $\mathrm{R}_{1}$ 2.6. Wing length / length of $\mathrm{R}_{4+5}$ 1.9. $\mathrm{R}_{45}$ nearly straight. Length of $\mathrm{r}-\mathrm{m} /$ length of M -petiole 1.7. A / B 1.4. E / length
of $M_{1} 0.7$. E / length of $M_{2} 0.8$. Base of Cu -fork well beyond base of M -fork. M-fork veins distaly divergent. Cu-fork veins divergent. F / E 1.3. F / length of $\mathrm{CuA}_{1}$ 1.7. F / length of $\mathrm{CuA}_{2} 2.5$. C / D 1.4. Vein CuP reaching as far as base of $\mathrm{Cu}-$ fork. Vein $\mathrm{A}_{4}$ weak, shorter.

Legs. Length of fore basitarsus / length of fore tibia 1.3. Tibial spur formula 3.4; $8.8,5.6 ; 4.6,4.6$. Fore tibia with $1 \mathrm{ad}, 36 \mathrm{p}$ and 3 v setae. Mid tibia with 40 a, 5 pd and 11 p setae. Hind tibia with 8 a, 5 pd and 3 p setae. Posterior sensillae placodea on basal part of tibia 3; 3; 3.

Terminalia (fig. 2A, B). Length of gonocoxite 0.60 mm . Sternal process narrow with about 40 tiny setae; length / width 1.9. Dorsal lobe of gonostylus 0.84 mm long with very elongated lobe, basally with strongly setose cushion. Sclerotized internal process bifurcate with both prongs widening apically. Ventral lobe of gonostylus large, bearing 6 fan-tipped setae basally. One lobe of the inner lamellate parts of the gonostylus apically with a slightly curved, spine-like seta thus forming a hook. Cerci thin and slender, reaching to about apical level of gonocoxite, with two short but strong apical setae (not figured).

## Pseudexechia lanceostylus sp. n.

(fig. 3)
Type material. - Holotype male, tanzania: Tanga region, W. Usambara Mts., Mazumbai, 1530 m a.s.l., 02.-03.XI.1990, G. E. E. Soli, Malaise tent (zmbn No. 180).

Diagnostic characters. - The male imago is characterised by the shape of the dorsal lobe of gonostylus which ends in a slender lobe, and by the shape of the bifurcate internal process.

Etymology. - From Latin, lancea, a small light spear, and stylus, style, referring to the slender lance-like dorsal lobe of gonostylus.

Description of male holotype. - Length of thorax 0.98 mm . Wing length 3.02 mm . Wing length / length of thorax 3.1. Wing length / length of fore femur 3.1.

Coloration. Antenna with scape, pedicel and basal part of flagellum yellow, gradually becoming darker and more greyish dusted towards tip. Head and maxillary palp yellow. Thorax mainly yellow; scutum uniformly brownish yellow with pale yellow or somewhat silvery shining stripe along lateral margin, delimited by darker margins below and above; scutellum brownish yellow with faint paler median stripe. Wing unmarked, yellow tinted, slightly paler towards hind margin. Wing veins yellow with dark setae. Halter pale yellow with three darker yellow stripes.


Fig. 3. Pseudexechia lanceostylus sp. n. male terminalia. - A, ventral view; B, internal face of the gonostylus. Scale 0.50 mm .

Legs mainly yellow; coxae paler yellow, fore coxa anteriorly with darker stripe. Abdominal tergites mainly yellowish brown; tergite 1 to 3 with pale lateral markings; tergite 4 and 5 with pale posterolateral markings, almost reaching middle; tergite 6 almost entirely dark. Terminalia mainly yellow, darker on sternal process and on parts of dorsal lobe of gonostylus.

Head. Total length of flagellum 1.53 mm . First flagellomere 1.7 times as long as second flagellomere. Second flagellomere 2.3 times as long as wide. Six large orbital setae. Tiny median ocellus present. Diameter of lateral ocellus / diameter of median ocellus 2.4. Clypeus bearing about 60 setae; width / length 0.8. Fourth palpomere about 1.5 times as long as third palpomere.

Thorax. Pronotum with vertical row of 3 erect setae. Scutum with pair of medium sized bristle like dorsocentral setae present posteromedially in front of scutellum (prescutellar setae); anterior and lateral margins with large setae; otherwise uniformly clothed with small setae (but larger and fewer than in other

Pseudexechia except in P. longistylus sp. n.). Scutellum with pair of strong scutellar setae and 2 pairs of very short posteriorly curved setae basad of them. Proepisternum with 1 strong, 2 medium sized and 8 small setae. Laterotergite with $8-10$ large and about 60 small setae.
Wing. Crossvein h with 1-2 ventral setae. $\mathrm{R}_{1}$ with $5-8$ ventral setae apically. $\mathrm{R}_{4+5}$ with $3-4$ ventral setae basally and 36-39 ventral setae apically. Wing length $/$ length of $\mathrm{R}_{1} 2.6$. Wing length / length of $\mathrm{R}_{4.5} 1.8$. $\mathrm{R}_{4+5}$ nearly straight. Length of $\mathrm{r}-\mathrm{m}$ / length of M-petiole 1.6. A / B 1.5. E / length of $M_{1} 0.7$. E / length of $\mathrm{M}_{2} 0.8$. Base of Cu -fork well beyond base of M -fork. M-fork veins distaly divergent. Cu-fork veins divergent. F / E 1.3. F / length of $\mathrm{CuA}_{1} 1.5$. $\mathrm{F} /$ length of $\mathrm{CuA}_{2}$ 2.3. C/D 1.1. Vein CuP reaching as far as base of Cu -fork. Vein $\mathrm{A}_{1}$ strong, shorter.

Legs. Length of fore basitarsus / length of fore tibia 1.8. Tibial spur formula $3.7 ; 9.8,7.5 ; 5.4,5.3$. Fore tibia with $1 \mathrm{ad}, 39 \mathrm{p}$ and 3 v setae. Mid tibia with 41 a, 5 pd and 7 p setae. Hind tibia with 9 a, 5 pd and 4


Fig. 4. Pseudexechia inthanonensis sp. n. male terminalia. - A, ventral view; B, internal face of the gonostylus. Scale 0.50 mm .
p setae. Posterior sensillae placodea on basal part of tibia $3 ; 4 ; 3$.

Terminalia ( Fig. 3A, B). Length of gonocoxite 0.57 mm . Sternal process rounded with about 55 tiny setae; length / width 1.4. Dorsal lobe of gonostylus 0.51 mm , ending in slender lobe. Sclerotized internal process bifurcate, the the prongs pointing in almost opposite directions. Ventral lobe of gonostylus broad and rounded, bearing 6 fan-tipped setae. Inner lamellate parts of gonostylus with lobe ending in strong seta-like hook. Cerci thin, not reaching apical level of gonocoxite, with two strong apical and one strong subapical setae (not figured).

## Pseudexechia inthanonensis sp. n.

(fig. 4).
Type material. - Holotype male, thalland: Chiang Mai province, Doi Inthanon, about 2200 m a.s.l., 12.IV.1991, J. Kjerandsen, sweep net (zmbN No. 181).

Diagnostic characters. - The male imago is separable from other species by the shape of the dorsal lobe of gonostylus, which has a smooth rounded hook at the tip and a large internal process with 25 teeth, by the shape of the sternal process and by the short stem. of the cubital fork.

Etymology. - From Doi Inthanon, a mountain in northern Thailand. A noun in genitive case.

Description of male holotype. - Length of thorax 1.20 mm . Wing length 3.82 mm . Wing length / length of thorax 3.2. Wing length / length of fore femur 3.3.
Coloration. Antenna with scape, pedicel and first flagellomere for half of its length pale yellow, rest of flagellum greyish yellow. Head mainly brown, frons and sides yellowish. Maxillary palp yellow. Pronotum yellowish brown. Scutum brown with three yellow stripes, median stripe broadening to fore margin, and narrow yellow lateral margins. Scutellum brown with yellowish lateral margins. Pleura yellowish brown. Wing unmarked, greyish yellow tinted, paler on basal half posterior of R-stem. Wing veins yellowish brown with dark setae. Halter pale yellow with three greyish yellow stripes. Legs mainly yellow; coxae paler yellow, each coxa with yellowish brown stripe. Abdominal tergites mainly brown; tergite 2 to 5 with pale posterolateral markings, reaching middle; tergite 6 dark brown. Terminalia mainly yellow, darker on sternal process and on parts of dorsal lobe of gonostylus. Tip of dorsal lobe of gonostylus whitish.
Head. Total length of flagellum 1.84 mm . First flagellomere 1.7 times as long as second flagellomere.

Second flagellomere 2.0 times as long as wide. Four to five large orbital setae. Small median ocellus present. Diameter of lateral ocellus / diameter of median ocellus 2.1. Clypeus bearing about 40 setae; width / length 0.8 . Fourth palpomere about 1.4 times as long as third palpomere.

Thorax. Pronotum with vertical row of 2 strong and 2 medium sized upcurved setae. Anterior and lateral margin of scutum with setae of different sizes; scutum otherwise uniformly cloched with small dark setae. Scutellum with pair of strong scutellar setae and pair of short posteriorly curved setae basad of them. Proepisternum with 1 large, 2 medium sized and 24 small setae. Anepisternum with 6 small setae. Laterotergite with 12-14 large and about 200 tiny setae.

Wing. Crossvein $h$ without ventral setae. $\mathrm{R}_{1}$ with 9 ventral setae apically. $\mathrm{R}_{\text {t. }}$ with $4-5$ ventral setae basally and $58-65$ ventral setae apically. Wing length / length of $\mathrm{R}_{1} 2.3$. Wing length / length of $\mathrm{R}_{4,5}$ 1.7. $\mathrm{R}_{4,5}$ distinctly downcurved. Length of $\mathrm{r}-\mathrm{m} /$ length of M petiole 1.0. A / B 1.9. E/ length of $\mathrm{M}_{1} 0.7$. E / length of $\mathrm{M}_{2} 0.8$. Base of Cu -fork not beyond base of M-fork. M-fork veins only slightly divergent apically. Cu -fork large, veins divergent. F/E 0.98. F / length of $\mathrm{CuA}_{1}$ 1.2. F / length of $\mathrm{CuA}_{2}$ 1.7. C / D 0.9. Vein CuP reaching well beyond base of Cu -fork. Vein $\mathrm{A}_{1}$ strong, shorter.

Legs. Length of fore basitarsus / length of fore tibia 1.2. Tibial spur formula $3.5 ; 6.7,6.4 ; 4.6,4.5$. Fore tibia with $6 \mathrm{ad}, 45 \mathrm{p}$ and 8 v setae. Mid tibia with 49 a, 5 pd and 4 p setae. Hind tibia with 6 a, 5 pd and 8 $p$ setae. Posterior sensillae placodea on basal part of tibia $3 ; 3 ; 3$.

Terminalia ( Fig. 4A, B). Length of gonocoxite 0.67 mm . Sternal process narrow without setae; length / width 1.8. Dorsal lobe of gonostylus 0.67 mm long with smooth rounded hook at tip and relativly large sclerotized internal process with 25 teeth. Ventral lobe of gonostylus bearing 4-5 fan-tipped ventral setae and one strong dorsal setae apically. Inner lamellate parts of gonostylus with 5 small setae. Cerci thin and slender, reaching to about apical level of gonocoxite, without strong setae (not figured).

## Discussion

The presence of a pair of medium sized bristle-like dorsocentral setae posteromedially above scutellum in Pseudexechia longistylus sp. n. and P. lanceostylus sp. n. is not in accordance with the generic description. Tuomikoski (1966) used absence of 'discal bristles' as one of the main diagnostic characters for Pseudexechia. The other setae on the scutal disc in these two species are also relatively strong, erect and bristle-like when compared with other species in the
genus, and they are fewer in numbers. Hence, this character should be used with care. However, other generic characters clearly place these species within Pseudexechia.
P. inthanonensis sp. n. is evidently closely related to the Palaearctic P. trisignata (Edwards, 1913) based on the structure of the male terminalia. However, $P$. inthanonensis sp. n . is rather different in coloration (yellow thoracic stripes on brown ground) and shows an aberrant and interesting wing venation. The short Cu -stem places the base of Cu -fork slightly before base of M-fork ( $\mathrm{F} / \mathrm{E}=0.98$ ). A long Cu -stem (short Cu -fork) has been used to group the species of Exechia s.l. (including Exechiopsis and Pseudexechia) (e.g. Edwards 1925).

These findings strengthen the view of Tuomikoski (1966) that Pseudexechia is more closely related to Allodiopsis and Allodia than to Exechia and Exechiopsis.

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## References

Chandler, P. J., 1978. Notes on the Holarctic species of Pseudexechia Tuomikoski (Diptera: Mycetophilidae), with the description of a new British species. Entomologist's Record and Journal of Variation 90: 44-51.
Edwards, F. W., 1925. British fungus-gnats (Diptera, Mycetophilidae). With a revised generic classification of the family. - Transactions of the Royal Entomological Society of London 1924: 505-670.
Matile, L., 1970. Pseudexechia de la région éthiopienne (Dipt. Mycetophilidae). - Bulletin de la Société entomologique de France 75: 209-214.
Matile, L., 1980. 15. Family Mycetophilidae. - In: Crosskey, R. W., Catalogue of the Diptera of the Afrotropical region. - British Museum (Natural History), London., pp. 216-230.
McAlpine, J. F., 1981. Morphology and terminology Adults. - In: J. F. McAlpine et al. (eds.), Manual of Nearctic Diptera. Volume 1. - Research Branch Agriculture Canada, Monograph no. 27. Ottawa, Ontario. Pp. 9-63.
Tuomikoski, R. 1966. Generic taxonomy of the Exechiini (Dipt., Mycetophilidae). - Annales entomologici fennici 32(2): 159-194.

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# SYNONYMICAL NOTES ON THE NEPA CINEREA LINNAEUS COMPLEX (NEPIDAE: HETEROPTERA) 


#### Abstract

Polhemus, J. T., N. Nieser \& S. L. Keffer, 1994. Synonymical notes on the Nepa cinerea Linnaeus complex (Nepidae: Heteroptera). - Tijdschrift voor Entomologie 137: 331-336. [rssn 0040-7496]. Published 15 December 1994. The Nepa cinerea Linnaeus complex is distributed over Europe, North Africa, the Middle East and northern Asia. Within this complex we recognize three species, $N$. cinerea, $N$. sardiniensis Hungerford and $N$. anopbthalma Décu, Gruia, Keffer \& Sarbu. The following are considered to be junior synonyms of $N$. cinerea Linnaeus, 1758: Nepa rubra Linnaeus, 1758; Nepa scorpioaquaticus De Geer, 1773; Nepa cinerea var. minor Puton, 1886, syn. n.; Nepa seurati Bergevin, 1926, syn. n.; Nepa cinerea var. major Bergevin, 1926, syn. n.; Nepa dollfusi Esaki, 1928, syn. n.; Nepa cinerea var. orientalis Esaki, 1928, syn. n.; Nepa rubra meridionalis Poisson, 1961, syn. n.; Nepa remyi Poisson, 1961, syn. n.; Nepa cinerea poissoni Tamanini, 1973. Correspondence: Dr. John T. Polhemus, University of Colorado Museum, 3115 S. York St., Englewood, Colorado 80110. Key words. - Heteroptera; Nepidae; Nepa cinereat; synonymy.


The Nepa cinerea Linnaeus complex is a tightly knit and very widespread species group in the Old World, distributed throughout Europe, the Mediterranean region, and as far east as northwest China and the Amur region of Siberia. Nepa cinerea was one of the earliest water bugs to be studied (Hoefnagel 1592, Moufet 1634, Frisch 1728, Swammerdam 1737-38), and the morphology has been extensively investigated, with an excellent treatment and summary of earlier studies by Hamilton (1931). Despite of this wealth of knowledge, the species group nomenclature concerning this insect remains unsettled. A definitive synonymical treatment is needed for the Catalogue of Palearctic Heteroptera (B. Aukema \& C. Rieger, editors) now being prepared for publication.

The early nomenclatural dilemma regarding the priority and synonymy of Nepa cinerea versus Nepa rubra Linnaeus, 1758 , first addressed in detail by Esaki (1926), then by Tamanini (1973), was finally resolved by the ICZN (Opinion 1335/1985) in response to a petition by Kerzhner (1981). Tamanini (1973) has declared the type locality to be Sweden, because the first specimens were collected there; he cites inclusion in Linnaeus' Fauna Svecica (1761) as evidence for this. Over the years however, a number of additional taxa have been proposed, as species, subspecies and varietal forms, that are either synonymous or very closely allied with $N$. cinerea. Esaki (1928), Poisson (1961) and Tamanini (1973) addressed this
species-group problem, however, far from resolving the status of these species-group taxa, each of these authors exacerbated the situation by adding at least one additional variety.

The following is a brief summary of the taxa and their supposed differences from Nepa cinerea sensu stricto, with comments on the validity of the characters used for differentiation.

Nepa cinerea var. minor Puton, 1886 and Nepa cinerea var. major Bergevin, 1926 were established solely on the basis of size differences, which is not a differentiating characteristic, as it varies widely within single populations. Jaczewski (1934) examined specimens of $N$. cinerea minor from Algeria, and stated that 'they seem to be conspecific with the typical $N$. cinerea L.' We therefore synonymize both varieties.

Bergevin (1926) established Nepa seurati n. sp. from Tunisia on the basis of a number of somatic differences, which he compared, in a table, with Nepa cinerea var. minor Puton. Esaki (1928) provided a photograph of the type of seurati which does indeed have a slightly different shape than the cinerea specimens he illustrates, with the lateral abdominal margins straight along much of the basal part, widening posteriorly, and abruptly incurved beyond, as noted by Poisson (1961: 631). Also the fore femur basally is wider than the $N$. cinerea specimens illustrated. This taxon would therefore seem to be separable from Nepa cinerea cinerea, however in long series taken from single populations in Morocco (nr. Tangier bor-
der, 8.VIII.1955, D. R. Lauck, Polhemus Coll.) and Portugal (nr. Portinas, Riba la Banho, 15.IV.1976, N. Nieser, Nieser and Polhemus Collns.) all of these character states are present along with intermediates. If considered as a valid subspecies, then many of the populations from north Africa would probably be attributable to this taxon.

Esaki (1928) described Nepa dollfusi from a single female collected in Morocco. Some of the differences from $N$. cinerea he cited may be attributable to alary polymorphism (see Larsén 1949, 1955), as the flightless forms, such as dollfusi, have an altered morphology, e. g. straighter hind margin of the pronotum, and reduced hemelytral membrane. Esaki provided a photograph of the type of dollfusi which appears to represent a malformed specimen, quite possibly resulting from damage during emergence from the last nymphal stage. The specimen also has reduced antennae, discussed below, that may have also resulted from a traumatic ecdysis. The red abdominal dorsum noted for European specimens of $N$. cinerea in contrast to dollfusi is unreliable, as it varies from grey brown to yellowish to pink or red in individual series from Europe and the Mediterranean region. Theiry (1981) noted the great variation in size and coloration (of both ground colour and abdominal tergites) in the $100+$ specimens of Nepa collected over a wide range of elevation in Morocco, all of which he assigned to $N$. cinerea, noting that the parameres and antennae of all forms were of the 'type cinerea.' Lindberg (1929) considered $N$. dollfusi to be only an aberrant specimen of $N$. cinerea.

Nepa cinerea var. orientalis Esaki, 1928 was founded on the basis of size and slight differences in antennal and head morphology, all unreliable. We have not been able to study a long series from the Far East to determine if there is any discernable difference in the male genitalia from the western European stock, but for the present the available evidence leads us to synonymize this form with $N$. cinerea.

Hungerford (1928) described Nepa sardiniensis from the island of that name. He separated it from related taxa by the lack of a prolongation on the second antennal segment and slender fore femur. These alone are questionably enough to justify the status of a separate species for the populations of Sardinia and Corsica, however Mazza $(1971,1978)$ has given additional evidence for separate species status (see below). Nepa sardiniensis is the only geographically isolated taxon in the $N$. cinerea complex, aside from the unique spatially isolated cavernicolous $N$. anophthalma Décu et al (in press). Linnavuori (1960) reported this species from Israel, Transcaspica and Turkestan, but according to Tamanini (1973) these records refer instead to $N$. cinerea cinerea.

For the Palearctic region, Stichel (1955) listed as
species Nepa seurati Bergevin, Nepa dollfusi Esaki, Nepa sardiniensis Hungerford, and N. cinerea (as Nepa rubra), with minor Puton, major Bergevin and orientalis Esaki clearly designated as forms of the latter.

Linnavuori (1960) claimed separation of $N$. cinerea and $N$. sardiniensis on the basis of the short second antennal segment, broader sternite VII, and differently formed male clasper of the latter. He stated that $N$. cinerea does not occur in Palestine. He also stated that the anterior femora of Nepa seurati are much broader basally than in $N$. sardiniensis.

Poisson (1961) gave figures of the antennae, female subgenital plate and male genitalia of $N$. dollfusi Esaki, $N$. cinerea cinerea (as N. rubra rubra), N. cinerea meridionalis (as $N$. rubra meridionalis), and Nepa remyi Poisson, 1961 from Morocco. N. remyi was separated on the basis of slight differences in the shapes of the male subgenital plate ('opercule génital'), male paramere, and the base of the fore femur, all of which have been seen to be somewhat variable within populations. We consider this variety to fall within the variability of $N$. cinerea, thus a synonym. Poisson also gave a key to separate all of the forms and species known in 1961 (including Nepa apiculata Uhler from North America and Nepa hoffmanni Esaki from China), based on differences in the antennae, the respiratory siphon, shape of the anterior femur, overall body length.

Seidenstücker (1963) investigated the status of $N$. dollfusi, $N$. sardiniensis and $N$. seurati, relying mainly on the antennal structure to conclude that Nepa seurati is a good species, and that Nepa sardiniensis Hungerford, 1928 (June) is a junior synonym of Nepa dollfusi Esaki, 1928 (April), and occurs in Turkey as well as Corsica and Sardinia. If Seidenstücker's analysis were accepted, the $N$. cinerea complex would include $N$. anophthalma, $N$. cinerea, N. dollfusi ( $=$ N. sardiniensis), and N. seurati. Later authors have not accepted the synonymy of $N$. sardiniensis (see below).
Mazza (1968) studied the variability of the number of respiratory horns of 5081 eggs of $N$. cinerea from Italy. He noted that the modal number of horns documented by Hinton (1961) for a British population (7) was different than the modal number (6) for the Italian population from La Spezia Province. Later he analyzed many specimens and concluded that the geographical forms were impossible to delimit, and that they form a continuum (Mazza 1974). This evidence suggests clines within $N$. cinerea.

Mazza (1971) noted constant differences in the number of respiratory horns (without giving details) between $N$. sardiniensis collected on Corsica and $N$. cinerea collected in Toscana (Italy), and further stated that these two populations were reproductively isola-
ted, as cross breeding produced sterile F1 hybrids. Later he (Mazza, 1978) examined about 1000 specimens of each species, and using several somatic characters in first order equations he showed consistent separation, therefore $N$. sardiniensis and $N$. cinerea are presently accepted as distinct species.

Tamanini (1973) discussed all of the previously mentioned forms except Nepa remyi Poisson, 1961, and proposed yet another subspecies, Nepa cinerea poissoni Tamanini from the Pyrenees Mountains of France and Spain, based on differences in pronotal morphology and antennae. He compared his new taxon to $N$. sardiniensis and $N$. cinerea cinerea. The differences he cites are not sufficient to separate this taxon from the latter, and it was synonymized with the latter by Nieser \& Montes (1984). Tamanini claims that $N$. sardiniensis is endemic to Corsica and Sardinia, and states that $N$. cinerea dollfusi and $N$. cinerea meridionalis are of 'uncertain systematic rank.' For all of Europe and North Africa, Nieser (1978) listed only Nepa sardiniensis Hungerford and Nepa cinerea, and recognized as subspecies of the latter $N$.c. poissoni Tamanini and N. c. seurati Bergevin. Later Nieser \& Montes (1984) synonymized N. c. poissoni with $N$. cinerea, without comment.

Nepa anophthalma Décu, Gruia, Keffer \& Sarbu (in press) is separable from all other species of the $c i$ nerea complex by five characters; 1. A yellow ground color (vs. dark), 2. Eyes absent (vs. present), 3. Hemelytra brachypterous, posterior margin sinuate (vs. macropterous, posterior margin rounded ), 4. Midlongitudinal groove of mesosternum absent (vs. present), 5. Paraterga hirsute (vs. glabrous except for hairs ringing stigmata). This is clearly the most annectant species of the complex, yet the male genitalia are very similar to the other members of the group. It is endemic to the Movile cave system in Romania.

As discussed above, most of the characters used by various authors to separate species-group taxa in the Nepa cinerea complex are as variable within populations as they are between populations, thus must be considered unreliable for the separation of taxa, especially when dealing with single specimens or small series. The antennae have been extensively cited as a separating characteristic notwithstanding their variability. Tamanini (1973) has shown that the antennal morphology is quite variable, even in a single specimen, thus the antennal morphology is not a completely reliable specific character in this genus, although their form may indicate trends, certainly vary between some populations, and seem to have a norm within certain populations (vide Seidenstücker, 1963; Tamanini, 1973). Esaki (1928) illustrated the antennae of a topotypic specimen of $N$. seurati, the type of $N$. dollfusi, and of $N$. cinered (no provenance given). In dollfusi the prolongation of the second seg-
ment is lacking and segments two and three are fused and short; in this regard it is very similar to N . sardiniensis, in fact so similar that Seidenstücker (1963) proposed the synonymy of the two taxa. In seurati the prolongation of the second segment is moderately long, and segments two and three are separated and moderately long, very similar to cinerea except in the latter the prolongation of the second segment is slightly longer. Tamanini (1973) studied many specimens from various localities, and illustrated the antennal forms most frequently encountered as well as aberrant forms rarely seen. The range of morphology within the species essentially covers the spectrum of differences used to separate the three species group taxa discussed above, although the 'normal' form of each conforms roughly to those illustrated by Esaki (1928). Mancini (1936), who tentatively identified specimens from Libya as $N$. cinerea minor, questioned whether the shape of the antennal segments was a valid character.

One of us (SLK) has studied the male genitalic structure of several populations of the Nepa cinerea complex from Finland, Morocco, Sardinia, Romania, and Iran and can find only one significant genitalic difference in any of the taxa (see below), although the genitalia of the cinerea complex are considerably different than the genitalia of the apiculata group which contains Nepa apiculata and Ňepa hoffmanni (Keffer et al. 1990). However, this general lack of male genitalic difference should not be taken as evidence that species or subspecies designations within the cinerea complex are invalid. Keffer (1991 and personal observation) has found that male genitalia are often of litthe taxonomic value at the species level in the Nepidae. For example, in his revision of the New World waterscorpion genus Curicta, Keffer (1991) found male genitalia to be diagnostic for the genus but largely invariant throughout the range of the genus. Similarly, Keffer (in prep.) has studied a majority of the species across the family Nepidae and found that male genitalic characters often have generic or species group significance but are often of little value for species determination. One genitalic character, paramere shape, has been used extensively by two nepid taxonomists of the recent past, Jose De Carlo and Raymond Poisson, in their numerous new species descriptions. Keffer, however, has found that parameres may vary either little or else randomly across species groups, or even genera, and thus are not generally valid as a species specific character.

Keffer's observations with Curicta and other nepid genera are affirmed in the genus Nepa. Male genitalia do separate the cinerea and apiculata species groups (Keffer et al. 1990). However, within the cinerea complex phallic structures, with one exception, are either invariant, or they vary randomly, across all
taxa. For example, parameres appear to vary randomly and are therefore of no value in making species and subspecies determinations. Taminini (1973, figs. 6178) figured parameres for $N$. cinerea cinerea, $N$. cinerea poissoni, $N$. cinerea seurati, and $N$. sardiniensis. We have observed $N$. cinerea cinerea specimens from Finland with paramere shapes similar to those figured by Taminini for $N$. cinerea poissoni and we have observed $N$. sardiniensis parameres that are similar to those figured by Taminini for $N$. cinerea cinerea. Furcher, Décu et al. (in press) have shown that $N$. anophthalma has parameres similar to those of N . cinerea poissoni.
Only one genitalic character, the shape of the secondary struts, appears to be of value for separating some cinerea group taxa. During copulation, the male's secondary struts and ventral diverticulum lodge in the female bursa copulatrix (Larsen 1938, fig. 26; secondary struts $=$ 'gr, Griffel', bursa $=$ 'vt, Vaginaltasche'; Keffer 1991, p.3) where they may have a holdfast and/or stimulatory function. If, in fact, the struts are stimulating the female during copulation, and thus subject to sexual selection, they could exhibit species specificity. Taminini (1973: 233) noted that, in ventral view, the secondary struts ('processi a spatola della coniunctiva') of $N$. cinerea cinerea specimens from Sicily and continental Italy converge toward the midline distally, whereas $N$. sardiniensis secondary struts are nearly straight. We would add to Taminini's description two of our own observations. First, converging secondary struts, as in cinerea cinerea, are also found in the cave dwelling cinerea group taxon, $N$. anophthalma and in apiculata group taxa. Second, in both $N$. cinerea cinerea and $N$. sardiniensis the secondary struts are dorsoventrally flattened whereas in $N$. anophthalma and in the apiculata group taxa the struts are tubular in shape. In sum, it appears that secondary struts can be used to separate $N$. sardiniensis (secondary struts nearly straight, not converging distally) and $N$. anophthalma (secondary struts tubular, not dorsoventrally flattened) from other ci nerea group taxa.
The status of four species group taxa must be decided; they are $N$. cinerea, $N$. dollfusi, $N$. sardiniensis and $N$. seurati. The first question to be settled is the proposed synonymy of dollfusi and $N$. sardiniensis, by Seidenstücker (1963). This synonymy seems to rest on the strength of the similarity of the antennae of the two taxa. If this synonymy is accepted, and the modal antennal form is accepted as a valid discriminating character, then the range of $N$. dollfusi, the senior synonym, is Morocco, Tunisia, Corsica and Sardinia. Tamanini (1973) rejected this synonymy, however, and also rejected the citation of $N$. sardiniensis for Israel by Linnavuori (1960) and for Turkey, Transcaspica and Turkestan by various authors, as be-
ing narrowly based on antennal form only. In this contention he is supported by Mazza (1971, 1978), thus current opinion supports the recognition of $N$. sardiniensis as a valid species endemic to Corsica and Sardinia, with $N$. dollfusi relegated to synonymy under $N$. cinerea.

Tamanini (1973) contends that $N$. cinerea cinerea occurs only as far south as northern Italy, and that $N$. cinerea seurati is distributed from southern Italy through Sicily to Libyia and Tunisia (we would possibly add Morocco in part). He shows a gradation of characters on a latitudinal cline through Italy, which suggests a single species rather than subspecies. If the subspecies status of seurati were to be accepted, then dolffusi should be considered as a synonym with aberrent antennae. This would leave $N$. cinerea cinerea as a variable subspecies with a range throughout northern Europe and as far south as Morocco nearest Spain, across the Balkans, through the Middle East, and as far east as Siberia.

We here reject the subspecies concept for seurati, however, as based on insufficient evidence and unreliable characters, and synonymize it under cinerea.

Mayr \& Ashlock (1991:43) give the following definition of the subspecies: 'A subspecies is an aggregate of phenotypically similar populations of a species inhabiting a geographic subdivision of the range of that species and differing taxonomically from other populations of that species'. We accept this definition, and all of the species group taxa of the $N$. cinerea complex that fail to conform to this definition are placed in synonymy of $N$. cinerea below. We have seen no convincing evidence that the characters upon which these taxa were founded are anything more than individual variations within and between populations, scattered more or less randomly, and without significant separation as required for subspecies characterization. It is indeed possible that sufficient evidence may be presented in the future to reestablish one or more of the synonymized taxa as valid subspecies, but for the present such clear evidence is lacking. We therefore recognize only three species, without subspecies, in the cinerea complex; Nepa anophthalma (endemic to the Movile Cave system of Romania), $N$. sardiniensis (endemic to Corsica and Sardinia), and the widespread $N$. cinerea. The results of our investigations are summarized as follows:

Nepa anophthalma Décu, Gruia, Keffer \& Sarbu, in press.

Nepa anophthalma Décu, Gruia, Keffer \& Sarbu, in press. Holotype: ${ }^{\circ}$, Movile Cave, Romania. (ERSI) [examined].
Distribution. - Europe, Romania, nr. Mangalia, Movile Cave. Endemic.

## Nepa cinerea Linnaeus, 1758

Nepa cinerea Linnaeus 1758: 440. Syntypes, ${ }^{\text {ox, }}$, Q, Sweden (type locality restricted to Sweden by Tamanini 1973: 226). (uziu) [not examined].

Nepa rubra Linnaeus 1758: 440. Holotype, 9 , locality unknown. (Uziu) [not examined]. (syn. of Nepa cinerea Linnaeus 1758; see Opinion 1335/1985).
Nepa scorpioaquaticus De Geer 1773: 361. Syntypes, 5, sex unknown, type locality unknown. (coll De Geer, NHRS) [not examined]. Unneccessary new name for N . cinerea L. (see Retzius 1783: 90).
Nepa cinerarea P. L. S. Müller 1774: 472. Incorrect subsequent spelling.
Nepa cinerea var. minor Puton 1886: 8. Syntypes, sex unknown, Tunisia, Oued Tessa, 'Sidi-Mohamed-ben-Ali', Oued Bateha, Oued Eddedj, Gafsa, Gabés. (Depository unknown) [not examined] (syn. Jaczewski 1934, suspected). Syn. n..
Nepa seurati Bergevin 1926: 290. Holotype, ס', Tunisia, Kebili. (mNHN) [not examined]. Syn. n.
Nepa cinerea var. major Bergevin 1926: 294. Holotype, ðै, Morocco, Taza. (Depository unknown; coll. Bergevin; mnhn?) [not examined]. Syn. n.
Nepa dollfusi Esaki 1928: 434 (April). Holotype, ㅇ, Morocco, Oued Djenanimès près Khénifra. (mnHn) [not examined]. (syn. Lindberg 1929: 9; restored by Seidenstücker 1963: 315, and named senior synonym of Nepa sardiniensis, synonymy with sardiniensis rejected by inference by Nieser 1978: 283). Syn. n.
Nepa cinerea var. orientalis Esaki 1928: 436. Syntypes, 3 아, Russia, Amur (bMnH) [not examined]. Syn. n.
Nepa rubra meridionalis Poisson 1961: 635. Syntypes, ${ }^{7}$, 9 , France, Tunisia, 'Iberian Peninsula'. (coll. Poisson, USNM) [examined]. Syn. n.
Nepa remyi Poisson 1961: 636. Syntypes, 1ठ, 2 ㅇ, Morocco, region d'Ifrane, Tagbaloute. (coll. Poisson, USNM) [examined]. Syn. n.
Nepa cinerea poissoni Tamanini 1973: 239. Holotype, ơ, France, Pyrenées-Orientales, Collioure. (Ltrc) [not examined]. (syn. Nieser \& Montes 1984: 38).

Distribution. - Europe, North Africa, Middle East, northern Asia.

Note. - Nepa annulipes (non Laporte 1833): Kolenati 1857: 481 is a misidentification referring to a Laccotrephes species, not collected in the Caucasus; see Kiritshenko 1918: 171; see also Esaki 1928: 434.

Nepa sardiniensis Hungerford, 1928
Nepa sardiniensis Hungerford, 1928: 120 (June). Syntypes, 10 , 1 우, Italy, Sardinia. (SEMC) [examined]. (syn. with dollfusi by Seidenstücker 1963: 322; considered as valid species by Mazza 1971: 539, 1978: 85, Tamanini 1973: 241).

Distribution. - Europe (Corsica, Sardinia)

## Depositories

Abbreviations (codens) for depositories follow Arnett, Samuelson \& Nishida (1993), with the addition of: ERSI (Emil Racovitâ Speleological Institute,

Bucharest); ltrc (Livio Tamanini Collection, Rovereto, Italy).

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## References

Arnett, R. H., Jr., G. A. Samuelson \& G. M. Nishida, 1993. The insect and spider collections of the world. - Sandhill Crane Press, Gainesville, vi +308 pp.
Bergevin, E. de, 1926. Description d'une nouvelle espèce de Nepa (Hémiptère Nepidae) de sud Tunisien. - Bulletin de la Société Histoire Naturelle de Afrique Nord 17: 290294, 1 fig.
Décu, V., M. Gruia, S. L. Keffer \& S. M. Sarbu, in press. A stygiobiotic waterscorpion, Nepa anophthalma n. sp. (Heteroptera: Nepidae), from a sulfurous cave in Romania. - Annals of the Entomological Society of America.
De Geer, C., 1773. Memoirs pour servir a l'histoire des Insectes. - Hosselberg, Stockholm, Vol. 3, 2 + 696 pp, 44 pls.
Esaki, T., 1926. Remarks on the Linnaean species of Nepa and Laccotrephes (Heteroptera: ${ }^{\circ}$ Nepidae). - Bulletin of the Brooklyn Entomological Society (N. S.) 21: 177-181.
Esaki, T., 1928. Contribution to the knowledge of the genus Nepa (Hemiptera; Nepidae). - Annals and Magazine of Natural History (10) 1:434-441, 1 pl .
Frisch, J. L., 1728. Beschreibung von allerley Insecten in Teutschland, nebst nützlichen Anmerkungen und nöthigen Abbildungen von diesem kriechenden un fliegenden inländischen Gewürme, zur Bestätigung und Fortsetzung der gründlichen Entdeckung, so einige von der Natur dieser Creaturen herausgegeben, und zur Ergänzung und Verbesserung der andern. - Nicolai, Berlin, Teil 7, pp. 8 +31 , figs. I-XXII.
Hamilton, M. A., 1931. Morphology of the water scorpion Nepa cinerea Linn. (Rhynchota, Heteroptera). Proceedings of the Linnaean Society of London 1931: 1067-1136. 6 pls., 22 figs.
Hinton, H. E., 1961. The structure and function of the eggshell in the Nepidae (Hemiptera). - Journal of Insect Physiology 7: 224-257.
Hoefnagel, G., 1592. Archetypa Studiaque Patris Georgii Hoefnagelii. - Jacobus F. Francoforti ad Moenum, 4 pts., each with 12 pls . and title page, 52 pp .
Hungerford, H.B., 1928. A new Nepa (Hemiptera Nepidae). - Bulletin of the Brooklyn Entomological Society (N. S.) 23: 119-123, 1 pl.
Jaczewski, T., 1934. Notes on some palaearctic aquatic and semiaquatic Heteroptera, chiefly from south-eastern Europe. - Annales Musei Zoologici Polonici 10: 267288.

Keffer, S. L., 1991. Taxonomic revision of the Neotropical genus Curicta Stål (Insecta: Heteroptera: Nepidae). PhD Dissertation, Southern Illinois University, Carbondale, xvii +264 pp .

Keffer, S. L., J. E. McPherson \& J. T. Polhemus, 1990. What is Nepa hoffmanni (Heteroptera: Nepidae)? Male genitalia hold the answer, and delimit species groups. Journal of the New York Entomological Society 98 (2): 154-162.
Kerzhner, I. M., 1981. Nepa cinerea Linnaeus, 1758 (Insecta, Heteroptera, Nepidae): proposed conservation under the plenary powers. Z. N. (S.) 2144. - Bulletin of Zoological Nomenclature 38: 138-141.
Kiritshenko, A. N., 1918. Hemiptera-Heteroptera of the fauna of the Caucasian Region. I. (Hemipt. Het. Faunae Caucasicae). - Mémoires du Musée du Caucase, Tiflis (A) 6: 1-177. (in Russian)

Kolenati, F. A., 1857. Meletemata Entomologica. Fasc. VI. Hemipterorum Heteropterorum Caucasi. - Bulletin de la Société Impériale des Naturalistes de Moscou 29: 419491, 1856.
Larsén, O., 1938. Untersuchungen über den Geschlechrsapparat der Aquatilen Wanzen. - Opuscula Entomologica, Lund, Suppl. 1, 338 pp., 151 figs.
Larsén, O., 1949. Die Ortsbewegungen von Ranatra linearis L. Ein Beitrag zur vergleichende Physiologie der Lokomotionsorgane der Inseckten. - Lunds Universitets Arsskrift N. F. Avd. 2, 45: 1-82.
Larsén, O., 1955. Der Bau des Flugapparats bei Nepa cinerea L. Ein Vergleich zwischen der flugfähigen Imago und der gewöhnlichen Form mit reduzierten Flugorganen. Opuscula Entomologica 20: 170-173.
Lindberg, H., 1929. Inventa entomologica itineris Hispanici et Maroccani, quod a. 1926 fecerunt Harald et Håkan Lindberg. I. Hemiptera Heteroptera Hydrobiotica. - Commentationes Biologicae. Helsingfors 3: 1-12, 1 pl.
Linnaeus, C., 1758. Systema naturae per regna tria naturae, secundum classes, ordines, genera, species, cum characteribus, differentiis, synonymis, locis. Editio decima, reformata. - L. Salvii, Holmiae, v + 824 pp.
Linnaeus, C., 1761. Fauna Svecica sistens animalia sveciae regni: Mammalia, Aves, Amphibia, Pisces, Insecta, Vermes, distributa per classes $\&$ ordines, genera $\&$ species. - Laurentii Salvii, Stockholmiae, frontis. + xlvi + 578 pp., 2 pls.
Linnavuori, R., 1960. Hemiptera of Israel I. - Annales Zoologici Societas Zoologicae-Botanicae Fennicae, Vanamo, Helsinki 22: 1-71.
Mancini, C., 1936. Contributo alla conoscenza degli emitteri della Tripolotania e della Cirenaica. - Annali del Museo Civico di Storia Naturale di Genova 59: 195-201.
Mayr, E. \& P. D. Ashlock, 1991. Principles of sytematic zoology. Second Edition. - McGraw Hill, New York, xx +475 pp .
Mazza, M., 1968. Osservazioni sulla variabilità del numero dei prolungamenti respiratori nelle uova di Nepa rubra (Rincote Eterottero). - Bollettino di Zoologia, Roma 35: 448-449.
Mazza, M., 1971. Ricerche su Nepa sardiniensis. - Bollettino di Zoologia, Roma 38: 359.
Mazza, M., 1974. Variabilità ed anomalie negli scorpioni d'acqua euromediterranei (Heteroptera: Nepidae). - Atti della Socierà Toscana di Scienze Naturali, Pisa, Memorie (Serie B) 81: 209-247, 1 pl., 10 tables.
Mazza, M., 1978. Caratteristiche discriminanti in scorpioni
d'acqua euromediterranei (Heteroptera: Nepidae). - Atti XI Congrèsso nazionale italiano Entomologica 1976: 8589.

Moufet, T., 1634. Insectorum sive minimorum animalium theatrum. - T. Cotes, London, $9+326+2 \mathrm{pp}$.
Müller, P. L. S., 1774. Des Ritters Carl von Linné Königlich Schwedischen Leibarztes k. k. vollständiges Natursystem nach der zwölften latineischen Ausgabe und nach Anleitung des Holländischen Houttuynischen Werks, mit einer ausführlichen Erklärung. - Gabriel Nicolaus Kaspe, Nürnberg, Vol. 5, part 1, $4+8+758$ pp.
Nieser, N., 1978. Heteroptera. pp. 280-285. In: Illies, J., Ed., Limnofauna Europaea. - G. Fischer Verlag, Stuttgart, New York; Swets \& Zeitlinger B. V., Amsterdam.
Nieser, N. \& C. Montes, 1984. Listas faunistica y bibliografica de los Heteropteros Aquaticos (Nepomorpha \& Gerromorpha) de España y Portugal. - Lista de la Flora y Fauna las aguas continentales de la Peninsula Ibérica, Publ. No. 1, 69 pp.
Opinion 1335/1985: Nepa cinerea Linnaeus, 1758 (Insecta, Heteroptera): Conserved. - Bulletin of Zoological Nomenclature 42: 241-243.
Poisson, R., 1961. A propos d'une nouvelle espece Palearctique du genre Nepa L. 1758 (Heteroptera, Nepidae). - Vie et Milieu 11: 628-640, 1960.
Puton, A., 1886. Enumération des Hémiptères recueillis en Tunisie in 1883 et 1884 . In: MM. Valery Mayet et Maurice Sédillot, Exploration Scientifique de la Tunisie. - Paris, ii + 24 pp.

Retzius, A. J., 1783. Caroli Lib. Bar. De Geer, genera et species insectorum e generosissimi auctoris scriptis extaxit, digessit, latine quoad partem reddidit, et terminologiam insectorum Linneanam addidit. - Siegfried Lebrecht Crusium, Lipsiae, 220 pp.
Seidenstücker, G., 1963. Zur Aufklärung von Nepa dollfusi (Heteroptera). Reichenbachia 1: 315-322.
Stichel, W., 1955-56. Illustrierte Bestimmungstabellen der Wanzen. II. Europa. (Hemiptera - Heteroptera Europae). - Selbstverlag, Berlin-Hermsdorf, Vol. 1, 168 pp. (pp. 132: 14.iii.1955; 33-64: 25.vi.1955; 65-96: 1.viii.1955; 97-128: 1.xi.1955; 129-160: 15.xii.1955; 161-168: 1.ii.1956.)

Swammerdam, J., 1737-1738. Biblia naturae; sive Historia Insectorum, in classes certas redacta, nec non exemplis, et anatomico variorum animaculorum examine, aeneisque tabulis illustrata. - Severin etc., Leydae, 2 vols, $910+56$ $+36+124$ pp., 53 pls.
Tamanini, L., 1973. Priorità e sinonimia di Nepa cinerea Linneo e Nepa rubra Linneo. Regione tipica e valore delle razze europee di Nepa cinerea Linneo, 1758 (Hemiptera Heteroptera, Nepidae). - Studi Trentini di Scienze Naturali (B) 50: 222-259.
Thiery, A., 1981. Contribution a la connaissance des Hétéroptères du Maroc: Les Hétéroptères aquatiques du Haut Atlas Occidental. - Bulletin de l'Institut Scientifique, Rabat 1981: 13-34.

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Synonymical notes on the Nepa cinerea Linnaeus complex (Nepidae: Heteroptera).


[^0]:    Abb. 1-21. - 1,2,3,5,6, Horisme vitalbata; 4, Synopsia sociaria; 7, Archiearis notha; 8, Aethalura punctulata; 9, 10, Siona lineata; 11, Chesias legatella; 12, Schistostege treitschkei; 13, 14, Aplocera praeformata; 15, Carsia sororiata; 16, 17, Operophthera brumata; 18, 19, Alsophila quadripunctaria; 20, 21, Theria rupicapraria.
    1, Habitusbild der Puppe, Ventral; 2, Dorsalsicht; 3, 4, Frons und Umgebung; 5, Abdominalende in Lateralsicht; 6, idem, in Dorsalsicht; 7, Kremaster in Dorsalsicht; 8, Kopf in Ventralsicht; 9, Frontalhälfte der Puppe in Ventralsicht; 10, 14, 15. 17. 18, 20, Abdominalende in Dorsalsicht; 11, Abdominalhälfte der Puppe in Dorsalsicht; 12, 13, idem in Ventralsicht; 16, Kopf und Beine in Ventralsicht; 19, 21, Augen und Umgebung.
    $\mathrm{A}_{1}-\mathrm{A}_{10}=1 .-10$. Abdominalsegment, $\mathrm{Af}=$ Analfeld, $\mathrm{An}=$ Analnaht, $\mathrm{Ant}=$ Antennen, $\mathrm{As}=$ abdominale $\mathrm{Spiracula}, \mathrm{C} 1=$ Clypeus, $\mathrm{D}_{1}, \mathrm{D}_{2}=$ Dorsalborsten $1,2, \mathrm{Dr}=$ Dorsalrinne, Es = Epicranialnaht, $\mathrm{Fa}=$ Frontalausläufer der Dorsalrinne, $\mathrm{Fr}=$ Frons, $\mathrm{G}=\mathrm{Genae}, \mathrm{Gn}=$ Genitalnaht, $\mathrm{Hb}=$ Hinterbeine, $\mathrm{Hfl}=$ Hinterflügel, $\mathrm{Ka}=$ Kaudalausläufer der Dorsalrinne, $\mathrm{Kr}=$ Kremaster, $\mathrm{L}_{1}=$ Lateralborste $1, \mathrm{Lb}=$ Labium, $\mathrm{Lbr}=$ Labrum, $\mathrm{Le}=$ Lateraleinschnitt, $\mathrm{Mb}=$ Mittelbeine, $\mathrm{Msn}=$ Mesonotum, $\mathrm{Mtn}=$ Metanotum, $\mathrm{O}=$ Auge, $\mathrm{Pdl}=$ Postclypeus, $\mathrm{Pn}=$ Pronotum, $\mathrm{Pr}=$ Proboscis, $\mathrm{Sd} \mathrm{S}_{1}=$ Subdorsalborste $1, \mathrm{Ts}=$ thorakales Spiraculum, V = Vertex, $\mathrm{Vb}=$ Vorderbeine, Vfl $=$ Vorderflügel, Vs $=$ Vorderschenkel

[^1]:    Abb. 58-87. - 58, 59, Parectropis similaria; 60, 61 Tephrina arenacearia; 62, Deileptenia ribeata; 63, Arichanna melanaria, 64, Stegania dilectaria; 65, Peribatodes rhomboidaria; 66, Semiothisa signaria; 67, 68, Itame brunneata; 69, 71, Diastictis artesiaria; 70, Narraga fasciolaria; 72, Abraxas grossulariata; 73, Ennomos autumnaria; 74, 76, Apeira syringaria; 75, 77, 78, Hemistola biliosata; 79, Jodis putata; 80, 81, Pseudoterpna pruinata; 82-84, Thalera fimbrialis, 85, Chlorissa cloraria; 86, Hemithea aestivaria; 87, Epirrita christyi.
    58, 61-63, 66, 67, 70-73, 80, Abdominalende in Dorsalsicht; 59, 60, 64, 65, 68, 69, 75, 78, 89, Basis des 5. Abdominalsegmentes in Lateralsicht; 75, 79, Abdominalende in Ventralsicht; 76, 82, 86, Habitusbild in Ventralsicht; 77, Kopf und Beine in Ventralsicht; 81, Abdominalende in Lateralsicht; 84, 85, Augen und Umgebung.

[^2]:    Abb. 146-178. - 146, Catarhoe cucullata; 147, C. rubidata; 148, Epirrhoe alternata; 149, 150, Nycterosea obstipata; 151, 152, Camptogramma bilineata; 153, 154, Scopula marginepunctata; 155, Idaea aureolaria; 156, I. rufaria; 157, I. deversaria, 158, 159, Rhodostrophia vibicaria; 160, 161, Baptria tibiale, 162, Glacies alticolaria; 163, Gnophos unicoloraria; 164, Colotois pennaria; 165, 167, Nothacasis sertata; 166, 171-173, Anticollix sparsata; 168, Acasis appensata; 169, Trichopteryx polycommata; 170, T. carpinata; 174-176, Anticles badiata; 177, Discoloxia blomeri; 178, Venusia cambrica.
    146, 148, 150, 152, 154, 157, 159, 161-164, 167, 168, 171, 172, 174, 177, 178, Abdominalende in Dorsalsicht; 147, Dorsalrinne; 149, 151, 153, Pronotum, 1. Abdominalsegment; 158, Erhebung kaudal von dem thorakalen Spiraculum; 160, Metanotum, 1. Abdominalsegment; 165, Kopf und Beine in Ventralsicht; 166, Vorderschenkel; 169, 170, 175, Abdominalende in Lateralsicht; 176, Punktgrübchen an der Basis des 2. Abdominalsegmentes.

[^3]:    Characters. - The combination of characteristics of the female genitalia as exhibited in the Dipseudopsidac (having synapomorphy 2, lateral papillae of segment X present, but lacking synapomorphy 6, abdominal sternite VIII not subdivided into lateral lobes) does not represent an apomorphic state, but only a unique combination of characteristics which happens to define the family very well, but only in a taxonomic sense. Synapomorphy 7 , inferior appendage one-segmented is a homoplasy in Polycentropodidae and Eenomidac. Three synapomorphies 8 10 support monophyly of Dipseudopsidac, and the lirst of these, antennal bases close together, is not unique, but within the Hydropsychoidea is also present in Hydropsychinae and Macronematinae Hydropsychidac). However, since this characteristic ahsent for the most part in Arctopsychinae and Diplectroninace. it is not part of the groundplan of Hydropsychidace and thus is a homoplasy. A more accurate description of this condition as a homologue is

[^4]:    India. Nepal

[^5]:    Specimens examined．－INDONESIA：Sumatra：Pematang Siantar：58゙，1．V11．84－13．IV．1985；30゙，1．VI－12．XII．1985； 32．22．IV－11．VII．1986；58，1．IX．88－15．I．1989；10̊，9．III－ 10．X．1989．D）ichl（citrm）．

[^6]:    . 11 matc semulin: A. latcral: B, proanal appendages, segments IX and X, dorsal; C, inferior ... ...e.al: L.. phatlus, ventral: l. segments IX, X, dorsal; G, ventral; H, segment IX, dorsal. I-J, : ... drint. ventral. Fig. 22, I). contorta Banks. Fig. 23, D. diehli Malicky and Weaver. Fig. 24, .. ....ehleri Ulmer

[^7]:    $\therefore . . .2 . . . .$. UcLachlan, 1866: 269, pl. 17: fig. 6, DONESA: Sulawesi: Ujung i'andane. ${ }^{\circ}$ Celches Macassar, Wallace coll. (BMNH). The - . . mounted on rwo separate slides by Mosely

[^8]:    ————ma Ulmer. 1905: 92-93, 96, figs. 68b,
    

[^9]:    Dipseudopsis martynovisp．n．
    he．37）
    「ype material．－Holotype $\mathbf{\delta}$ ：PHILIPPINES：Sulu： larawakan NE，12．II． 1957 Yoshio Kondo（BPPM）． $\therefore \quad . \quad$ ：Tawitawi，Tarawakan，north of Batu Batu， $\therefore$ Dan Exp．（ZMmi＇C）：10才，21．X．1961；4才， 20．X．1961：28，13．XI．1961；28：14．XI．1961．

[^10]:    1才, Rangoon, McL. coll 1:63 49. Phnom Penh (UsNM, ZMHB). West Malaysia: Kedah: 18, Kuala Ketil (BMNH). bong Lima. light trap (tssim). Thalland: $2 \mathbf{o}^{\text {a }} 19$,
     I = (M N N ) : 53 1f. Chiang Mai, Ban-tin-doi, $310 \mathrm{~m}(-$ ${ }^{2}$ Rang Sit. at light (bminh, Nhmw). $10^{\circ}$, Bung Borapet (USNM). 43 . det. Banks nin. Siam", Patalung, I.V.1924, I. H. XX. Evans (MčC. BM:H). 10. Patalung, Paknam

[^11]:    Ect: 14. left subgenital plate dorsal view: 15 , same, dorsolateral view; 16, left style, dorsal view; 17 , aedeagus and connectie. ventrocaudal view: 18 . aedeagus. dorsal view; 19 , same, left lateral view. Scales: 0.1 mm .

[^12]:    Aethammobates prionogaster sp. n. 11es. 1-4

    「ype material. - Holotype ठ : 'Coll. A. Mochi / 27.V. 37 (ebb. Astar / Egitto (in coll. Baker)

[^13]:    Cladogram of Bacturia, numbers .. . : issed in the section on the phylogenasuta group; b. Cladogram of the - uagroup, numbers refer to characters discussed ..... .. he monophyly and ingroup phylogeny of

[^14]:    ${ }^{2}$ See remarks under $E$. designatus;: ${ }^{2}$ Transferred to Ophiogomphus by Needham (1899); ${ }^{3}$ Transferred to Ophiogomphus by Hagen (1874) As Herpetogomphus; ${ }^{5}$ Transferred to Ophiogomphus by Selys (1879)

[^15]:    Remarks
    The description is based on 91 males and 43 fema-

[^16]:    1 19. 15-27. Synthorax of Erpetogomphus species, lateral view (males, except given otherwise). - 15, bothrops (holotype); 16, itherm: Mcxico, \'eracruz State); $1^{-}$, designatus (lectotype); 18, designatus (Mexico, Durango State); 19, sipedon (female hoIontye: 21. I. Lampropethis (holotype); 21, lampropeltis natrix (holorype); 22, crotalinus (lectotype); 23, beterodon (holotype); 2. cmpomms "California, Yolo Co.); 25, compositus (paratype of coluber); 26, boa (Mexico: Veracruz State); 27, cophias Mexico: Morelos State). Scale line 3 mm .

[^17]:    :Sa. $\sqrt{ } 4-(5)$. Anterner and posterior hamules. First view is anterolateral view showing profile of anterior hamule; second view :Hen hown is lateral view showing profile of posterior hamule. - 59, viperinus (Mexico: Veracruz Stare); 60, designatus (lec-- Wp. ais sedon (Mexico: Morelos State); 62, l. Lampropeltis (California: Ventura Co.); 63, l. natrix (holorype); 64, crot.anz. . vico: Michoacan State); 65, heterodon (New Mexico: Catron Co.); 66, compositus (Arizona: Pima Co.); 67, compo:nt. or of coluber); 68, boa (lectotype); 69, cophias (Mexico: Morelos State). Scale line 2 mm .

