## KIM RIBEIRO BARÃO

Morfologia comparada, descrição de novos táxons e filogenia de um gênero de Pentatomidae (Hemiptera: Heteroptera)

Tese apresentada ao Programa de Pós-Graduação em Biologia Animal, Instituto de Biociências, Universidade Federal do Rio Grande do Sul, como requisito parcial à obtenção do título de Doutor em Biologia Animal.

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Aprovada em $\qquad$ de $\qquad$ de $\qquad$ .

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## RESUMO

Pentatomidae tem sido extensivamente estudada taxonomicamente, com espécies novas e táxons supragenéricos ainda sendo descritos, porém tanto as suas relações filogenéticas quanto a classificação supra-genérica permanecem em discussão. Este trabalho aborda comentários sobre análises filogenéticas realizadas na família, morfologia comparada como subsídio para análises filogenéticas, descrição de espécies novas e análise filogenética de Dichelops. Ressaltamos a necessidade da descrição e ilustração claras de caracteres, amostragem ampla de grupos externos, codificação de exemplares ao invés do uso de groundplan, da avaliação e consideração de dados faltantes e do emprego claro de métodos filogenéticos, concluindo que análises filogenéticas em Pentatomidae deveriam, preferencialmente, incluir grupos externos mais representativos, a fim de melhor testar as hipóteses de monofilia. Estudando comparativamente a morfologia de algumas estruturas da tíbia das pernas anteriores de Pentatomidae, Scutelleridae e Thyreocoridae, concluimos que a estrutura pré-tarsal em Pentatomidae é pouco variável e que não permite a identificação de grupos taxonômicos, à exceção de Phyllocephalinae que apresenta pulvilos distintos; que algumas características pré-tarsais permitem diferenciar o pré-tarso de Thyrecoridae do prétarso das outras duas famílias; que o número de cerdas que compõem o aparato tibial varia consideravelmente dentro de Pentatomidae, possuindo um padrão por subfamílias, e entre subfamílias. Sugerimos que o número de cerdas do aparato tibial e o número de projeções espatulares do pente tibial sejam usadas e exploradas como caracteres contínuos em análises filogenéticas em níves supra-genéricos. Uma nova espécie de Dichelops (Diceraeus) e o macho de Dichelops (Prodichelops) divisus são descritos. As relações filogenéticas de Dichelops, incluindo 44 espécies de grupo externo, foram estudadas por parcimônia utilizando-se dados morfológicos e discretos. Dichelops foi sempre recuperado parafilético.

Decisões taxonômicas são tomadas e discussão sobre metodologia cladística e tratamento de caracteres contínuos são fornecidos.

## INTRODUÇÃO GERAL

Pentatomoidea reúne aproximadamente 7000 espécies em 15 famílias (sensu Grazia et al. 2008: Acanthosomatidae, Canopidae, Cydnidae, Dinidoridae, Lestoniidae, Megarididae, Pentatomidae, Phloeidae, Plataspididae, Saileriolidae, Scutelleridae, Tessaratomidae, Thaumastellidae, Thyreocoridae, Urostylididae), é composta basicamente por insetos fitófagos, representados em todas as regiões biogeográficas. A sua diversidade morfológica é notável, desde espécies com aparência de casca-de-árvore (Phloeidae) até espécies com pernas especializadas para cavar (Cydnidae). Além da sua diversidade morfológica e hábitos, a interação com plantas cultiváveis têm chamado a atenção de biólogos e agrônomos para este grupo.

Estudo morfológicos comparados na superfamília já exploraram a morfologia das coxas (Lis 2010a), tíbias (Lis \& Schaefer 2005), pré-tarso (Bonatto 1988; Lis 2010b; Lis \& Ziaja 2010), sistema eferente odorífero externo das glândulas metatorácicas (Kment \& Vilímová 2010a,b; Parveen et al. 2014) e genitália de machos (McDonald 1966; DavidováVilímová \& McPherson 1991) e fêmeas (Scudder 1959). A base para a proposição de caracteres filogenéticos tem vindo destes e outros estudos.

Relaçães dentro de Pentatomoidea foram estudadas por diversos autores, alguns empregando métodos filogenéticos (Gapud 1991; Grazia et al. 2008) e outros propondo classificações por comparações morfológicas (Singh-Pruthi 1925; Leston 1958; China \& Miller 1959; Cobben 1968, 1978). Grazia et al. (2008) utilizaram parcimônia para analisar dados moleculares, alinhados por homologia dinâmica, e morfológicos da mais representativa amostra taxonômica do grupo até o momento. Os resultados encontrados por Grazia et al. (2008) concordam com outros (Singh-Pruthi 1925; Leston 1958; China \& Miller 1959; Cobben 1968, 1978; Gapud 1991) somente na posição de Urostylididae como grupo irmão
dos demais pentatomóideos. Demais relações são reduzidas a duas politomias próximas à base do clado, que incluem 12 das 15 famílias. Neste pente, são reconhecidos um grupo formado por Cydnidae s.l. e outro por Scutelleridae + Plataspididae. Esforços amostrais maiores, tanto em grupos taxonômicos chave quanto em caracteres morfológicos e moleculares, são necessários para ajudar a resolver as relações nesse grupo.

Pentatomidae é a família mais diversa de Pentatomoidea, com aproximadamente 870 gêneros e 4.700 espécies. Com exceção de um grupo de espécies predadoras, i.e. Asopinae, todas as demais são fitófagas. Pentatomidae tem sido extensivamente estudada taxonomicamente, com espécies novas e táxons supragenéricos ainda sendo descritos (p.ex., Rider 2000; Rider \& Brailovsky 2014), porém tanto as suas relaçães quanto a classificação supra-genérica permanecem em discussão.

A monofilia de Pentatomidae é suportada em uma série de estudos (Gapud 1991; Hasan \& Kitching 1993; Grazia et al. 2008) e recentemente novas subfamílias foram incluídas nesta (Rider 2000: Stirotarsinae; Grazia et al. 2008: Aphylinae, Cyrtocorinae). Gapud (1991) foi o único que estudou as relações filogenéticas infra-familiares em Pentatomidae, porém a metodologia e os táxons utilizados não são claros, dificultando a interpretação, repetibilidade e aplicabilidade dos resultados encontrados. Ainda assim, Gapud (1991) encontrou Pentatominae e Podopinae polifiléticas e Asopinae, Aphylinae, Discocephalinae, Edessinae e Phyllocephalinae monofiléticas.

A falta de uma filogenia robusta para Pentatomidae fortemente influencia a sua classicação sistemática. Autores discordam quanto ao número de subfamílias e tribos em Pentatomidae, sendo aceitas até nove subfamílias (sensu Grazia et al. 2008: Aphylinae, Asopinae, Cyrtocorinae, Discocephalinae, Edessinae, Pentatominae, Phyllocephalinae, Podopinae, Stirotarsinae). A falta de classificação em Pentatominae é ainda mais confusa, 43 tribos já foram propostas (Rider in litt.) com base em similaridade morfológica e poucas
foram testadas com métodos filogenéticos claros (Shwertner \& Grazia 2012). Hasan \& Kitching (1993) foram os únicos a estimar uma filogenia para Pentatominae, mas assim como em Gapud (1991) para Pentatomidae, diversas questões metodológicas impedem uma clara interpretação dos resultados.

Carpocorini, uma das tribos mais diversas de Pentatominae, com cerca de 450 espécies incluídas em 100 gêneros, não possui uma diagnose formal. Historicamente, autores reconhecem grupos de gêneros tratados empiricamente como próximos e que são incluídos na tribo. Foi originalmente descrita por Mulsant \& Rey (1866) e reconhecida, com diferentes composições de gêneros e espécies por autores subsequentes (Stål 1876; Distant 1902; Putshkov 1965; Ahmad et al. 1974; Gross 1975). Rider (in litt.) agrupou 100 gêneros em Carpocorini, levando em consideração informações de literatura que hipotetizaram a sua relação baseando-se em similaridade morfológica.

O relacionamento entre tribos de Pentatomidae foi estudado por Hasan \& Kitching (1993). Neste, Menidini, Carpocorini e Catacanthini formam um grupo relacionado pela presença de mesoesterno sulcado. O agrupamento de Carpocorini e Catacanthini é suportado por duas características: (i) búcula mais longa que o primeiro segmento do lábio e (ii) mesoesterno liso. Entretanto, estas sinapormorfias são evidências fracas da monofilia do grupo. De fato, neste trabalho não foram incluídas subfamílias e tribos exclusivamente neotropicais, cuja inclusão pode alterar as hipóteses de relacionamentos (Pollock et al. 2002).

Outros estudos sobre Carpocorini têm caráter regional, explorando as relações filogenéticas do grupo de espécies presentes, catálogos ou listas regionais (ex., Cassis \& Gross 2002; Rider 2006), chaves dicotômicas para gêneros (ex., Ribes \& Pagola-Carte 2009), entre outros. Zaidi (1987) propõe uma hipótese filogenética para Carpocorini, entretanto utiliza somente espécies da fauna do "subcontinente Indo-Paquistanês". Desta forma,

Carpocorini ainda carece de estudos que confirmem sua monofilia, que estabeleçam as relações filogenéticas intergenéricas e sua relação com outras tribos.

Dentre a fauna Neotropical de Carpocorini, Dichelops é um dos gêneros melhor estudados. Entretanto, ainda carece de atenção. Dichelops é composto por 16 espécies incluídas em três subgêneros: Dichelops Spinola, 1837 (10), Diceraeus Dallas, 1851 (5) e Prodichelops Grazia, 1978 (1). Grazia (1978) revisou extensivamente o gênero descrevendo novas espécies e produzindo a primeira proposta de relaçães entre os subgêneros. Klein et al. (2012) descreveram uma nova espécie e propuseram chave de identificação para as espécies do subgênero Dichelops. Segundo Grazia (1978), espécies de Dichelops podem ser facilmente identificadas por possuírem as placas mandibulares mais longas que o clípeo. Cada subgênero é suportado por uma série de características de morfologia geral e de genitália interna e externa de ambos os sexos.

Segundo Grazia (1978), Dichelops é morfologicamente próximo a Euschistus Dallas, 1851, Berecynthus Stål, 1862 e Padaeus Stål, 1862: entre outras características, todos apresentam mesosterno carenado. Por outro lado, Dichelops diferencia-se destes por possuir as placas mandibulares mais longas que o clípeo. Por similaridade em cor, tamanho e aparência geral, Rolston (1974) associa indiretamente Dichelops a nove gêneros que compõem o grupo Euschistus: Proxys Spinola, 1837, Galedanta Amyot \& Serville, 1843, Hymenarcys Amyot \& Serville, 1843, Agroecus Dallas, 1851, Tibraca Stål, 1860, Menecles Stål, 1867 e Sibaria Stål, 1872, além daqueles listados por Grazia (1978). Além desses gêneros, podem ser incluídos nesta lista gêneros subsequentemente relacionados taxonomicamente a Euschistus, como Acledra Signoret, 1864, Caonabo Rolston, 1974, Coenus Dallas, 1851, Ladeaschistus Rolston, 1973, Mcphersonarcys Thomas, 2012, Padaeus Stål, 1862, Oenopiella Bergroth, 1891 e Spinalanx Rolston \& Rider, 1988. De acordo com

Rider (1995), a presença de um par de processos dorsais da phalloteca é característica diagnóstica do grupo Euschistus.

Nenhuma das propostas de agrupamento supracitadas, em nível de tribo ou inferior, foi testada filogeneticamente. A resolução das relações em Pentatomidae é necessidade primária na classificação de Pentatomoidea (Schuh \& Slater 1995; Grazia et al. 2008). Análises filogenéticas com grupos de gêneros em Carpocorini possibilitarão a verificação do relacionamento entre gêneros historicamente tratados como próximos e a proposição de diagnoses mais precisas para os grupos analisados e classificação da tribo com base nas relações de parentesco.

## Estrutura da tese

Esta tese está estruturada em formato de artigos, que compõem quatro capítulos. No Capítulo I escrevemos uma resposta ao trabalho de Memon et al. (2011), que versa sobre uma análise filogenética de Halyini ocorrentes no sub-continente Indo-Paquistanês. Na Carta ao Editor escrita por nós, fizemos um apanhado geral do avanço da sistemática filogenética nos últimos 50 anos e enumeramos boas práticas que deveriam ser seguidas em qualquer análise filogenética. Dentre elas, descrição e ilustração claras de caracteres, amostragem ampla de grupos externos, codificação de exemplares ao invés de plano básico, da avaliação e consideração de dados faltantes e do emprego de métodos filogenéticos.

No Capítulo II, estudamos comparativamente a morfologia de algumas estruturas da tíbia das pernas anteriores de Pentatomidae, tendo sido amostradas 81 espécies ( 55 gêneros) representantes de sete subfamílias. A titulo de comparação estudamos Scutelleridae e Thyreocoridae, representadas por três e 10 espécies, respectivamente.

No Capítulo III, descrevemos uma nova espécie de Dichelops (Diceraeus) Dallas, 1851 e o macho de Dichelops (Prodichelops) divisus (Walker, 1867). No Capítulo IV
testamos filogeneticamente a monofilia de Dichelops e do grupo Euschistus, utilizando caracteres morfológicos discretos e contínuos em uma série de análises com pesos iguais e implicados e uma ampla amostragem de grupos externos.

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## CAPÍTULO I ${ }^{1}$

# Phylogeny of the South Asian Halyini? Comments on Memon et al. (2011) towards a better practice in Pentatomidae phylogenetic analysis 

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Recently, Memon et al. (2011) published a paper entitled "Phylogeny of the South Asian Halyine stink bugs (Hemiptera: Pentatomidae: Halyini) based on morphological characters" (Ann. Entomol. Soc. Am. 104(6):1149-1169). The main goal of the authors was to estimate the phylogenetic relationships of the Halyini South Asian genera based morphological characters. In our understanding, the authors ignore some basic assumptions of a phylogenetic analysis, including the interpretation of a priori hypothesis of relationship, ingroup and outgroup choice, character and character state definitions, and interpretations of the results. Hereafter, we do not intend to determine the best practices in phylogenetic analysis but to discuss some points of their analysis that are under-developed and cannot be disregarded.

Ingroup and Outgroup sampling - According to Memon et al., Halyini is monophyletic, despite the fact that there is no published phylogeny, and its taxonomy and systematic position need revision. The taxon sampling of Memon et al. is restricted to South

[^0]and Central Asian species; they did not include species from outside the studied area (the tribe is known to include species from North America, Africa, and Australia) or taxa that have been allied to Halyini. Therefore, the absence of outgroups and the restricted ingroup sampling do not allow the authors to investigate their main question or, much less, Halyini monophyly. A group monophyly is tested by the optimal placement of the outgroup terminals outside of the ingroup (Farris 1972). Even without changing ingroup monophyly, outgroup taxa can change their relationships. Furthermore, at least two outgroup terminals are required to test ingroup monophyly. These include one terminal that serves as the root and another that is free to potentially fall inside the ingroup. Memon et al. claim that all analyzed genera belong to Halyini due to basic halyine tribal characters (for such characters see p. 1150), which are used by all the researchers to identify tribe members. In the most comprehensive, although unpublished, phylogeny of Halyini, Wall (2004) described these characters as homoplasious and the tribe as paraphyletic, thereby making it impossible to define the tribe with an exclusive sinapormorphy.

Terminal taxa - Memon et al. adopted the ground plan character coding approach in their analysis. The authors' data matrix has supraspecific taxa as terminals (p. 1151, for a species list see Appendix), but they do not mention the criteria used to manage intra- and interspecific variations nor how they coded the terminals. The ground plan character coding approach consists of a variety of methods that are frequently not specified by the authors and considered to be intuitive (some methodological explanations can be found in Prendini 2001). The main methodological objection to the use of supraspecific taxa as terminals is the assumption of its monophyly, which is particularly dangerous in very diverse taxa such as the Halyini. The exemplar coding approach seems to be superior in this regard because species as terminal taxa are defined on diagnosability criteria rather than monophyly. Therefore, the exemplar approach is preferable for interpreting character polarity because it uses observable
and verifiable data, rather than hypothetical states and character combinations (Prendini 2001).

Characters and character states - The methods of Memon et al. do not adhere to any logical procedure for coding morphological characters. In the same character, is possible to observe information regarding quantitative traits and form as well as the presence or absence of particular characters, such as the character "Shape of Lateral Margins of Pronotum" (p. 1155). Brazeau (2011) analyzed the effects of different coding practices in morphological phylogenetic analysis and suggested that certain practices are undesirable and should be avoided. Particularly, multistate character information may impose congruence artificially by linking more than one character variable to a particular state. Additionally, their character explanations are biased and result in the a priori establishment of the typical character states of a particular taxon, e.g.: "A dentate lateral margin to the pronotum is a halyine character differentiating genera of the tribe from most others (apart from some Australian genera)". Within a cladistics framework, such a conclusion should be resultant of an analysis and is dependent on the taxa included.

Missing data - Memon et al. justified the exclusion of some taxa from the analysis on the basis that taxa with large proportions of missing characters decrease the accuracy of phylogenetic inference. However, the level of data completeness alone should not guide the exclusion of taxa, and most studies suggest that it is generally possible to accurately place incomplete taxa in phylogenies if enough informative characters are sampled (Wiens \& Morrill 2011). In addition, their matrices do not have large amounts of missing data, and the authors did not perform tests to ascertain whether such taxa would decrease the accuracy of phylogenetic inference to justify their exclusion.

Phylogenetic analysis - Although not explicitly stated, the phylogenetic analysis completed by Memon et al. supposedly followed the procedures of successive weighting
using the rescaled consistency index with the sole purpose of reducing the number of equally parsimonious trees found. According to Goloboff (1993), some authors still incorrectly advocate weighting as a method only for selecting a tree among the shortest trees under equal weights. In the study by Memon et al., this argument does not apply because the tree determined under successive weighting was none of the equally parsimonious trees found in an analysis with equal weights. Under an equal weights parsimony analysis, the authors found 5825 equally parsimonious trees for the complete matrix ( 31 terminal genera) and 52 trees for the reduced matrix ( 22 terminal genera). This occurred because no collapsing rule was applied during the heuristic search, which was not mentioned. If one uses a collapsing rule, such as "collapsing branches if supported ambiguously" ( min . length $=0$ : PAUP rule 1 ), 419 and 11 equally parsimonious trees would be determined for the same respective data sets (swapping algorithm $=\mathrm{TBR}, 3000$ RAS, saving 150 trees per replication).

Final comments - As challenging as it may be to elect an outgroup taxon for cladistics analysis in Pentatomidae, such difficulty does not justify the exclusion or noninclusion of any taxa. There are some cues to guide such choices, e.g., Hasan and Kitching (1993) suggest a monophyletic clade comprising Halyini, Megarrhamphini, Tetrodini and Phyllocephalini, and Wall (2005) states that Halyini putative genera do not form a monophyletic assemblage in any of his analyses. Moreover, as a first attempt, one can select outgroup taxa based on shared taxonomic history or by the morphological diversity of analyzed characters.

In the last three decades, cladistic analysis has improved with the development of several new procedures. There are current debates concerning relevant practices adopted in cladistic analysis, such as how to describe and encode characters; whether to weight all characters equally or based on some optimization criteria other than the number of steps (e.g., homoplasy); how to choose outgroup taxa and how many are needed; and which measure of
branch support is the more appropriate to evaluate the results. Most of these questions are still dependent on the researcher's methodological background and theoretical beliefs, but we advocate that whatever method is chosen should be explained and justified.

Aiming a better practice of phylogenetic systematics and an improvement on the development and interpretations on the Pentatomidae phylogeny, we suggest the following: (i) the use of broader outgroup samples, allowing more reliable tests of monophyly; (ii) the use of more than one species per genus and its individual entry in the data matrix, avoiding the ground plan approach for character coding; (iii) encoding of the characters clearly and objectively, making clear the procedure coding type; (iv) illustration of the main characters and (v) clearly explaining all the methodological procedures, allowing for analytic repeatability.

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## CAPÍTULO II $^{\mathbf{2}}$

# Comparative morphology of selected characters of the Pentatomidae foreleg (Hemiptera, Heteroptera) 

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#### Abstract

Heteropteran legs are very diverse within and among taxa, and such variation is frequently correlated with life habits. Structural modifications are commonly present in the legs of the Pentatomoidea but are poorly studied. Using scanning electron microscopy, the tibia and pretarsal microstructure of 82 species of Pentatomidae (Heteroptera), three species of Scutelleridae, and ten species of Thyreocoridae were described, focusing on the pretarsal structure, the foretibial apparatus, and the foretibial comb. The Pentatomidae, the Scutelleridae, and the Thyreocoridae have uniform pretarsal structures. Variation can be found in the length of the parempodial setae and in the shape of the parempodial projections. The foretibial combs of the Pentatomidae, the Thyreocoridae, and the Scutelleridae are described for the first time, and we have demonstrated that there is low structural variation in the foretibial comb complex of the studied species. The setae organization and distribution on


[^1]the foretibial apparatus is uniform in the families studied. However, the Asopinae (Pentatomidae) bear a foretibial apparatus that is uniquely organized. The taxonomic and phylogenetic relevance of the pretarsal traits, the foretibial apparatus, and the foretibial comb are discussed.

## Keywords

Asopinae; foretibial apparatus; pretarsus; Scutelleridae; Thyreocoridae; tibial comb

## 1. Introduction

Heteropteran legs and their structures are very diverse within and among taxa, and such variation is frequently correlated with life habits (Lis and Schaefer, 2005). Structural modifications are commonly present on the posterior legs of the Pentatomoidea, whereas the anterior and median legs are less variable. Variations in the coxae (e.g., Lis, 2010a), the tibiae (e.g., Lis and Schaefer, 2005; Grazia et al., 2008), the tarsi (e.g., Grazia et al., 2008), and the pretarsi (e.g., Bonatto, 1988; Lis, 2010b; Lis and Ziaja, 2010) have been described in the Pentatomoidea. However, except for the work of Hasan (1990), and the unpublished thesis of Bonatto (1988), little attention has been given to the Pentatomidae, the most diverse group in the superfamily. Because the Pentatomidae have lower macrostructural variation compared to other groups of the Heteroptera (e.g., Schuh, 1975; Weirauch et al., 2011), the anterior legs have been neglected as taxonomic characters.

Although understudied, the forelegs bear some structures of potential phylogenetic interest, particularly the tibia and pretarsus. The foretibial apparatus, a group of modified setae found on the inner surface of the foretibia, located approximately one-third the distance from the apex, is a unique feature of the Pentatomoidea (Grazia et al., 2008). Previously described by McAtee and Malloch (1928), the foretibial apparatus has been explored in four
studies (McDonald and Edwards, 1978; Dolling, 1981; Bonatto, 1988; Grazia et al., 2008), and this is the only information available about its morphology and possible phylogenetic relevance. A grooming comb that is located distally on the foretibia, called a tibial comb, is usually associated with cleaning the head appendages (Schuh and Slater, 1995). Its presence is widespread in the Heteroptera but it varies in the number and rows of setae (Lis and Schaefer, 2005). Within the Pentatomoidea, tibial combs have been described only in Thaumastella species (Thaumastellidae; Jacobs, 1989) and in the Cydnidae (Lis and Schaefer, 2005).

By contrast, the pretarsus has been widely investigated (e.g., Dashman, 1953a,b; Goel and Schaefer, 1970; Goel, 1972; Vasarhelyi, 1986; Hasan, 1990, 1995) and has proven to be phylogenetically relevant in the Heteroptera at higher group levels (e.g., Wheeler et al., 1993; Grazia et al., 2008) and in subordinate groups (e.g., Schuh, 1976; Weirauch, 2008). However, its structural variability and phylogenetic significance are not yet well known in the Pentatomoidea. The study with the broadest pretarsal sampling in the Pentatomoidea is the unpublished thesis of Bonatto (1988), which found large structural variation in the claws and pulvilli. Hasan and Nasreen (1994) studied eight pentatomoid families and found structural variations that were of significant taxonomic value, reflecting the systematics of the superfamily. More recently, Lis et al. (2002), Lis (2010b), and Lis and Ziaja (2010) presented new morphological comparisons on the pretarsus of the pentatomoid families Dinidoridae, Parastrachiidae, and Cydnidae.

We aim to explore the morphological diversity of the tibial and the pretarsal structures of the Pentatomidae foreleg and to draw attention to characteristics of possible phylogenetic interest using scanning electron microscopy. We intend to provide a basis for future systematic interpretations of the foreleg traits, which along with a wider set of characters,
should be subjected to congruence testing in future phylogenetic analyses of higher level groups in the Pentatomidae.

## 2. Material and methods

### 2.1. Taxon sampling

Although the Pentatomidae is a group that has been extensively studied taxonomically, both the phylogeny and systematics remain confusing. The monophyly of the Pentatomidae has been supported by a series of studies (Gapud, 1991; Hasan and Kitching, 1993; Grazia et al., 2008). The sole study concerned with the internal relationships of the Pentatomidae is that of Gapud (1991), but the phylogeny was biased, i.e., the characters were polarized by a priori concepts of ancestral-derived sequences of evolution (see Nixon and Carpenter, 1993), the character optimization was based on characters that were considered a priori to be synapomorphies to structure the phylogenetic tree, and characters were eliminated to avoid homoplasy (see Farris, 1983). Such procedures hamper an unbiased interpretation of character evolution and taxa relationships.

Thus, the lack of a robust phylogeny of the Pentatomidae also influences its systematics. There is disagreement among heteropterists on the subfamilial and tribal groupings within the Pentatomidae (e.g., Schuh and Slater, 1995; Grazia et al., 2008). More confusing is the Pentatominae tribal composition; without phylogenies to settle the tribal arrangements, the tribal classification varies from eight (Schuh and Slater, 1995) to 42 tribes (Rider, 2012).

Such problems make it difficult to reliably sample the Pentatomidae phylogenetic diversity. In an attempt to minimize such problems, we sampled all the Pentatomidae subfamilies, except for the monotypic Stirotarsinae and the diverse Podopinae, and some of the most common and diverse tribes that are recognized by Rider (2012).

A total of 81 species ( 55 genera) representing seven pentatomid subfamilies and seven tribes of Pentatominae were examined. Additionally, 13 species that were assigned to the Scutelleridae (3) and the Thyreocoridae (10) were examined as outgroup representatives. The Pentatomidae classification follows Grazia et al. (2008) and Rider (2012) for subfamilies and tribes, respectively. The species examined and specimen collection data are listed in Tables S1 and S2, respectively. Both sexes were sampled for each species, but sexual differences were not observed. Images of all the studied species are available as Supplementary Figures S1-S27. The specimens used in this study were borrowed from the following institutions: AMNH, American Museum of Natural History, New York, United States; DARC, David Rider Collection, North Dakota, United States; DZUP, Museu de Entomologia Pe. Jesus Santiago Moure, Universidade Federal do Paraná, Paraná, Brazil; FIOC, Fundação Instituto Oswaldo Cruz, Rio de Janeiro, Brazil; FSCA, Florida State Collection of Arthropods, Florida, United States; HDEC, Dodge Engleman Collection, Canal Zone, Panama; IBSP, Instituto Butantan, São Paulo, Brazil; IFML, Instituto Fundación Miguel Lillo, Tucumán, Argentina; INBio, Instituto Nacional de Biodiversidad, Costa Rica; J. E. Eger, Joe E. Eger Collection, Florida, United States; MACN, Museo Argentino de Ciencias Naturales "Bernardino Rivadavia", Buenos Aires, Argentina; MCNZ, Museu de Ciências Naturais da Fundação Zoobotânica do Rio Grande do Sul, Rio Grande do Sul, Brazil; MGAP, Museu Anchieta de Ciências Naturais, Rio Grande do Sul, Brazil; MIZA, Museo Del Instituto de Zoología Agrícola, Venezuela; MNRJ, Museu Nacional, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil; MZSP, Museu de Zoologia da Universidade de São Paulo, São Paulo, Brazil; UFRG, Departamento de Zoologia, Universidade Federal do Rio Grande do Sul, Rio Grande do Sul, Brazil; NMNH, National Museum of Natural History, Washington D.C., United States; SAM, South Australian Museum, Adelaide, Australia.

### 2.2. Foreleg preparation

The tegumentary ultrastructure was studied using scanning electron microscopy (SEM) at the Centro de Microscopia Eletrônica of UFRGS. The prothoracic legs of pinned specimens were removed, cleaned manually, kept submersed in contact lens solution for 24 hours, and then agitated in a sonicator with water and detergent solution for three minutes. After, the legs were dehydrated by an alcohol series, stove dried at $40^{\circ} \mathrm{C}$ for 24 hours, coated with gold and observed by SEM. The number of setae on the tibial comb and on the foretibial apparatus was counted on the SEM photographs. The values for each species are available in Table S3.

The tarsal and pretarsal terminology that we use follows Weirauch (2005). The tibial comb complex and the foretibial apparatus terminology follow Lis and Schaefer (2005) and Grazia et al. (2008), respectively.

## 3. Results

### 3.1. General morphology of the pretarsus in the Pentatomidae

Although the pretarsus comprises structures that are distal to the last tarsomere, the structures that are functionally correlated with the pretarsus are also present in the last tarsomere and are described here. The sclerotized structures of the pretarsus comprise paired claws, pulvilli, and parempodial sclerites with paired parempodial setae, a median sclerite, and the unguitractor plate, all of which are connected to one another by membranous areas (Figs. 1B,D).

The last tarsomere opens ventrodistally and has reinforced edges. There are two cylindrical paired setae along its margin. The ventralmost seta (vs, Figs. 1B,D) is directed outwards from the tarsomere margin, whereas the lateral seta (ls, Figs. 1B,D) is directed posteriorly. One asymmetric seta (as, Fig. 1D) is located at the internal ventrolateral angle of the tarsal rim.

The unguitractor plate is uniform among the Pentatomidae; it is ellipsoidal and tripartite (vsup, lsup, Figs. 1B,D, 3). The proximal portion of the unguitractor plate is beset with denticles, and the distal portion is divided into a median ventral row and paired lateral rows of transverse ridges (Figs. 1B,D). Distally, the unguitractor plate is apparently fused to the parempodial sclerites.

The parempodial sclerites are separated from one another by a deep and broad, median area (psc, Figs. 1B-F). Distally, the parempodial sclerites bear the paired parempodial setae, and dorsally, they have paired projections (pp, ps, Figs. 1D-F). The parempodial setae are slender, circular in cross-section, and equipped with a longitudinal ridge along the first third of their lengths (Figs. 1E, 4). A triangular median sclerite (ms, Fig. 1C) is located dorsal to the parempodial sclerites. The claws are flattened, with large bases, and taper from the base to the apex. Large and robust pulvilli are connected ventrally to the claws and divided into basi- and distipulvilli.

### 3.2. Comparative morphology of the pretarsal structures in the Pentatomidae

Differences in the pretarsal structures that are congruent with any taxonomic rank within the Pentatomidae could not be found. However, the variation that was found is described and discussed below for each particular case.

The tarsomere rim bears two setae in all examined Pentatomidae; the lateral seta is always longer than the ventral seta. The ventral seta varies in length, being a minute projection of one-sixth the length of the lateral seta, such as in Dichelops melacanthus (Fig. $3 F$ ), or one-third the length of the lateral seta (Fig. 3E). The asymmetric seta is always present in the Pentatomidae at the internal ventrolateral angle of the tarsomere rim and is the same length as the lateral seta (Fig. 3E), or is three times shorter than the lateral seta (Figs. 3D,F,I), which is the most common state.

The ventral surface of the unguitractor plate is one or two times wider than the lateral surfaces in most Pentatomidae. The number of ridges is variable depending on the overall size of the unguitractor plate, but the lateral surfaces have approximately twice the number of ridges as the ventral surface, except in Lopadusa augur (Fig. 3C), in which the number of ridges is similar. The ventral surface protrudes distally beyond the lateral surface in most pentatomids, but both can be equal, such as in Alitocoris parvus (Fig. 3B) and Antestia degenera (Fig. 3D). The ventral and lateral surfaces may bear microtrichia or small denticles, as observed in Alcaeorrhynchus grandis (Fig. 3A) and Rhyncholepta meinanderi (Fig. 3H), or they may be smooth, such as in Cuspicona simplex (Fig. 3E) and Dichelops melacanthus (Fig. 3F).

The parempodial sclerites are rather uniform throughout the Pentatomidae. The parempodial projection arises dorsally of the parempodial setae socket (Figs. 1E,F) and can be elongate with a rounded (Fig. 4A) or an acute apex (Fig. 4B), elongate with a median constriction (Fig. 4C), or star-shaped (Fig. 4D), or with a straight and smooth apex (Fig. 4E) or with different developmental degrees of fingerlike projections (Figs. 4F-H). The parempodial setae vary in length and are more commonly half the length of the claws or less commonly, approximately the same size as the claws (Fig. 2) but are diminutive in the single examined representative of the Phyllocephalinae (Fig. 2M).

The claws in most of the Pentatomidae are flattened, with large bases, and they taper from the base to the apex (Fig. 2), but the claws are slender and elongate in Chloropepla vigens (Fig. 2J), Neotibilis spp. (Fig. 2K), Rhyncholepta spp., and Janeirona stali, and they have very robust bases and sharply taper toward the apex in Macrina juvenca (Fig. 2M). The claw orientation is variable because of the membranous nature of the claw insertion area, but in dried-preserved specimens, the claws are commonly held obliquely to the axis of the
tarsus. The claws lack projections and may be smooth (Fig. 2A) or bear reticulations near the base (Fig. 2C) or have a series of longitudinal ridges (Fig. 2G).

The typical pentatomoid pulvillus was the most common within the Pentatomidae, except for the long and narrow pulvillus of the single examined species of Phyllocephalinae (Fig. 2M). Because of its membranous nature, the distipulvilli shape is highly variable and therefore difficult to characterize by SEM.

### 3.3. Pretarsal structures in the outgroup representatives

The overall morphology of the Scutelleridae (Fig. 2N) and the Thyreocoridae (Fig. 2O) pretarsus is similar to the Pentatomidae pretarsus. The lateral and ventral setae and campaniform sensilla are present at the tarsomere rim, following the proportions that have been described for the Pentatomidae. The lateral and ventral surfaces of the unguitractor plate of the Thyreocoridae are the same width. The parempodial sclerite shape is conical in the Scutelleridae and the Thyreocoridae; the parempodial projections are variable in the Scutelleridae and homogeneous in the Thyreocoridae, elongate with an acute apex on the latter. The parempodial setae of the Scutelleridae follow the same pattern as in the Pentatomidae (Fig. 2N), whereas in the Thyreocoridae, the setae are less than half the length of the claws (Fig. 2O). The claws are flattened with large bases, and taper from the base to the apex in the Scutelleridae. In the Thyreocoridae, the claws are slender, smooth, and bear a longitudinal ventrolateral groove. Pulvilli similar to those found in the Pentatomidae were found in the Scutelleridae and the Thyreocoridae.

### 3.4. Foretibial comb

The tibial comb of the Pentatomidae and the Scutelleridae show little variation and consist of a single row of stout setae located almost apically on the inner surface of the tibiae (Figs. 1A,
5). The tibial fossula is covered by setae and consists of a broad depression that begins at the tibial comb and narrows and becomes shallower toward the distal sixth of the tibia.

The Thyreocoridae differ from the pentatomids and scutellerids because of a series of components that are associated with the tibial comb called the tibial comb complex (Figs. 50,P). The tibial comb is bound by two outer setae, which are apically sharpened and longer than the remaining regular setae. Additionally, two apical tibial longitudinally ridged spines border the tibial comb. Apart from these differences, the tibial combs of the Pentatomidae, the Scutelleridae, and the Thyreocoridae differ only in the number of setae (Table 1).

### 3.5. Foretibial apparatus

The foretibial apparatus, which is always located on the ventral surface of the foreleg, approximately one-third or less from the apex, is characterized by a set of stout setae that are surrounded by a region lacking setae (Figs. 1A, 6). This region may be either delimited by elevations (Fig. 6M) or by depressions (Fig. 60) in the integument, or have no indication of tegument differentiation (Fig. 6D).

The foretibial apparatus setae are arranged in two longitudinal rows that extend distally, forming a medial groove (Fig. 6). Each seta has a broad base and sinuously narrows toward an acute apex.

The foretibial apparatus of the Pentatomidae, the Scutelleridae and the Thyreocoridae differ only in the number of setae (Table 1), ranging from three to 23 . The number of setae does not overlap in the Scutelleridae and the Thyreocoridae (Table 1). In the Pentatomidae, the number of setae on the foretibial apparatus overlaps with the other two families (Table 1).

Within the Pentatomidae, the Asopinae foretibial apparatus has the largest amount of setae and the largest variation in the number of setae (Table 1), whereas the other pentatomid subfamilies are closer to the family average. The Pentatominae subfamily is second after the

Asopinae regarding the number of setae and the variation of the foretibial apparatus setae, and the tribes Catacanthini, Rhynchocorini and Antestiini have the largest amounts of setae (Table 1).

## 4. Discussion

### 4.1. The pretarsal morphology

To the extent that one can generalize from the few species that we studied, the Pentatomidae, the Scutelleridae, and the Thyreocoridae are uniform regarding the pretarsal structure, which is consistent with studies of other pentatomoid families (e.g., Lis et al., 2002; Lis, 2010b; Lis and Ziaja, 2010). Indeed, studying the pretarsus in the Pentatomomorpha, including the Aradidae, the Alydidae, the Coreidae, the Largidae, the Lygaeidae, the Pyrrhocoridae, and almost all the families of Pentatomoidea, Bonatto (1988) concluded that the pretarsus is essentially similar in all the groups. However, Bonatto (1988), Hasan and Nasreen (1994), and Hasan (1995) found structural variation in the claws and pulvilli that were of significant taxonomic value and that reflected the so far phylogeny of the Pentatomoidea.

Flattened claws with large bases that taper from the base to the apex are found in all the Pentatomoidea, whereas cylindrical claws that have nearly equal diameters of the base and the apex are found in many pentatomomorphan families (Grazia et al., 2008). As in the Pentatomidae, the Scutelleridae, and the Thyreocoridae, the claws of most pentatomoid families are usually smooth and may bear small reticulations near the base or may have longitudinal grooves. However, the Urostylididae, the Saileriolidae, and the Acanthosomatidae bear articulated bristles on the claws (Bonatto, 1988). The bristles are also found on the claws of some coreoid Coreidae (Bonatto, 1988).

In contrast to the results of Bonatto (1988), Hasan and Nasreen (1994), and Hasan (1990), we found little variation in the structure of the pulvilli of the Pentatomidae, the

Scutelleridae, and the Thyreocoridae. Indeed, the most common pulvilli structure in the Pentatomoidea is divided into a round, lamellate, membranous distipulvillus that is supported by a narrow and stout basipulvillus (Bonatto, 1988; Lis et al., 2002; Grazia et al., 2008). Thus far, pulvilli that differ from the standard are scattered across pentatomoid families. Pulvilli that are long and narrow, usually flattened, and undivided into basi- and distipulvilli are found on some Cydnini and Geotomini (Cydnidae, Cydninae; Lis and Ziaja, 2010), on the Phloeidae (Bonatto, 1988), and on the single species of the Phyllocephalinae (Pentatomidae) that we studied. Very large, robust, and stout pulvilli were documented as a unique feature of the Parastrachiidae (Lis, 2010b), but we consider the structure essentially the same as the pulvilli of Pentatomoidea. The absence of pulvillar structures has been documented only for Scaptocoris minor (Cydnidae, Cephalocteinae, Scaptocorini; Grazia et al., 2008; Lis and Ziaja, 2010).

The unguitractor plate, the parempodia, and the parempodial setae are largely similar in the terrestrial heteropterans (Goel, 1972). Our findings on the unguitractor plate of the Pentatomidae, the Thyreocoridae, and the Scutelleridae are concordant with those of Goel (1972): ventral and lateral surfaces of the unguitractor plate are formed by sclerotized scales, which may bear microtrichia. We also found variation in the shape of the parempodial projections and the unguitractor plate, variation in the length of parempodial setae, and variation in the number of scale rows on the unguitractor plate of the Pentatomidae, the Thyreocoridae, and the Scutelleridae. Dashman (1953b) noted variation in the parempodia and suggested that the parempodia could be used to help family identification. Our observations are the first detailed compilation of these structures, and the lack of a phylogeny for the Pentatomidae, the Scutelleridae, or the Thyreocoridae hamper any interpretation of their evolution or their phylogenetic value.

### 4.2. The foretibial comb

In this study, the foretibial comb of the Pentatomidae, the Thyreocoridae, and the Scutelleridae is described for the first time, and we have demonstrated low structural variation of the foretibial comb complex within the studied tribes, subfamilies, and families.

The Thyreocoridae presented the greatest morphological variation in the elements of the foretibial comb complex. A foretibial comb complex composed of a tibial comb, outer setae, basal spines, and tibial fossula is characteristic of the Cydnidae (Lis and Schaefer, 2005). On the Pentatomidae and the Scutelleridae, the foretibial comb complex is rather uniform and composed of the tibial comb and fossula. Although the tibial comb remains understudied in the Pentatomomorpha, in groups that are supported by several morphological characteristics (e.g., Asopinae, Edessinae), the variation in setae number on the tibial comb is less than in groups of uncertain composition (e.g., Pentatominae).

### 4.3. The foretibial apparatus

Despite that the morphology of the foretibial apparatus is conserved in the Pentatomoidea, the arrangement and number of setae can be useful in taxonomic and phylogenetic frameworks. Within the Pentatomomorpha, the foretibial apparatus is a unique feature of the Pentatomoidea, absent only in the Megarididae, the Urostylididae, and the Saileriolidae and according to Grazia et al. (2008), its absence in the Megarididae is most likely because of secondary loss. Thus, the foretibial apparatus may be an important diagnostic feature for the Pentatomoidea.

According to Bonatto (1988), the setae organization and distribution on the foretibial apparatus is uniform among the pentatomoid families. His survey on the Acanthosomatidae, the Canopidae, the Dinidoridae, the Pentatomidae, the Phloeidae, the Plataspididae, the Scutelleridae, and the Tessaratomidae species, found that the number of setae varies between
four and eight. Grazia et al. (2008) found a different number in the Acanthosomatidae, at least fifteen setae in Bebaeus punctipes. Additionally, we found 9-12 setae that comprise the foretibial apparatus of the Scutelleridae, in contrast to the 4-8 of Bonatto (1988). The species that were sampled by Bonatto (1988) and those that were sampled by us are different, so there may be more variation in the number of setae in species that are not yet sampled.

Within the Pentatomidae, five to six setae is the most common condition in the foretibial apparatus found both by Bonatto (1988) and by us. However, the Asopinae is an exception, bearing at least sixteen setae that are distributed in two longitudinally arranged rows. Members of the Asopinae are distinguished from the other pentatomid subfamilies by their essentially predaceous feeding habits (De Clercq, 2000), by possessing a crassate rostrum, and by the presence of genital plates with a thecal shield in the males (Thomas, 1992). The Asopinae have been considered to be a natural group (Gapud, 1991), but only further phylogenetic analysis can establish the group's monophyly and determine whether the foretibial characteristics, including a heavy foretibial apparatus, are synapomorphies.

### 4.4. Foreleg traits and the Pentatomoidea phylogenetic hypothesis

Of the foreleg traits that we studied, some were coded as characters in previous phylogenetic analyses of the Pentatomomorpha and the Pentatomidae (Gapud, 1991; Hasan and Kitching, 1993; Grazia et al., 2008). Both the foretibial apparatus and the tibial comb were only coded as characters in the phylogenetic analysis of the Pentatomoidea by Grazia et al. (2008).

Foretibial apparatus presence is considered a synapomorphy of the Pentatomoidea, except of Urostylididae and Saileriolidae (Grazia et al., 2008), but no distinction in its organization has been considered. We and Bonatto (1988) found that the foretibial apparatus varies in the number of setae within the pentatomoids. Although incipient, our results are
consistent with those of Bonatto (1988), mainly regarding the number of the Asopinae foretibial apparatus setae, as commented in the Discussion section 4.3.

A tibial comb consisting of a complex of different setal types is characteristic of the Cydnidae s.l. (Lis and Schaefer, 2005). Although not coded in a phylogenetic analysis, the tibial comb complex may be a diagnostic characteristic of the families Cydnidae, Parastrachiidae, Thaumastellidae, and Thyreocoridae, once according to Grazia et al. (2008), these families are closely related. Thus, these families may share a common evolutionary history of the foretibial comb complex, but it remains necessary to explore this characteristic in the Parastrachiidae and the Thaumastellidae and to code the foretibial comb complex in a phylogenetic analysis.

Despite recent efforts, the foretibial apparatus and tibial comb remain understudied in the Pentatomoidea, thus their phylogenetic signal cannot be evaluated. However, the variation in the setae number of such structures could be coded as a continuous character, as proposed by Goloboff et al. (2006).

By contrast, the pretarsus was used in three studies regarding the Pentatomomorpha, the Pentatomoidea or the Pentatomidae (Gapud, 1991; Hasan and Kitching, 1993; Grazia et al. 2008). Large amounts of morphological variation in the pretarsi appears to be phylogenetically informative in higher group levels (Bonatto, 1988; Gapud, 1991; Hasan and Kitching, 1993; Grazia et al. 2008), e.g., the variation in the number of tarsal segments, the presence or absence of the claw cuticular appendages, and the overall morphology of the pretarsi.

In the light of the current phylogeny of the Pentatomoidea (Grazia et al., 2008), the claws and the pulvilli structure variation found in the superfamily most likely evolved independently in each lineage.Likewise, minor characteristics of the pretarsi that were coded by Hasan and Kitching (1993), such as unguitractor plate sculpturing, are found within the

Pentatomoidea and Pentatomidae (Bonatto, 1988). Hasan and Kitching (1993) used the ground plan approach to code their characters, and they do not list the analyzed species; this, compounded by the fact that the group's systematics have been continuously modified during the last two decades, make it difficult to evaluate the composition of the groups that were analyzed by them. We were able to find variation in the unguitractor plate and in other structures in the species that we studied, but we still cannot evaluate their phylogenetic value.

In summary, the lack of either a phylogenetic analysis of the Pentatomidae, or of the availability of information about the structures that we evaluated in most groups within Pentatomoidea, hamper further interpretations of the evolution of the pretarsal and the tibial structures. Our results show that both the pretarsal and foretibial structures in the Pentatomidae, the Scutelleridae, and the Thyreocoridae are rather uniform and the morphological differences between the taxa are of low taxonomic value. By contrast, the variation in the foretibial apparatus and the foretibial combs may be of interest in phylogenetic analysis for use as continuous characters. Further studies of other pentatomoid families are required to shed light on the taxonomic and phylogenetic value of the understudied structures, particularly on the foretibial apparatus and the foretibial comb complex.

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Table 1. Mean, standard error, and minimum and maximum values of setae in the tibial comb and the foretibial apparatus. [mean $\pm$ standard error (range)]

| Family | Subfamily | Tribe | Number of setae |  |
| :---: | :---: | :---: | :---: | :---: |
|  |  |  | Tibial comb | Foretibial apparatus |
| Pentatomidae |  |  | $30.1 \pm 0.56$ (16-46) | $6.0 \pm 0.29$ (3-23) |
|  | Aphylinae |  | 20 | 4 |
|  | Asopinae |  | $30.7 \pm 1.70$ (26-37) | $16.5 \pm 1.00$ (13-23) |
|  | Cyrtocorinae |  | $21.6 \pm 3.17$ (18-28) | $5.3 \pm 0.33$ (5-6) |
|  | Discocephalinae |  | $27.7 \pm 1.86$ (24-30) | $5.3 \pm 0.18$ (5-6) |
|  |  | Discocephalin <br> i | 24 | 5 |
|  |  | Ochlerini | $32.4 \pm 1.44$ (29-37) | $5.5 \pm 0.29$ (5-6) |
|  | Edessinae |  | $34.3 \pm 0.63$ (33-36) | $5.1 \pm 0.14$ (5-6) |
|  | Pentatominae |  | $30.3 \pm 0.60$ (16-46) | $5.0 \pm 0.10$ (3-10) |
|  |  | Antestiini | $24.5 \pm 0.29$ (24-25) | $5.6 \pm 0.57$ (5-9) |
|  |  | Carpocorini | $27.5 \pm 0.64$ (21-33) | $4.3 \pm 0.09$ (3-5) |
|  |  | Catacanthini | $34.6 \pm 0.60$ (33-36) | $8.0 \pm 0.62$ (5-10) |
|  |  | Chlorocorini | $35.4 \pm 1.03$ (30-42) | $4.7 \pm 0.14$ (4-6) |
|  |  | Pentatomini | $31.1 \pm 1.60$ (16-43) | $5.0 \pm 0.14$ (4-7) |
|  |  | Procleticini | $26.0 \pm 1.45$ (21-30) | $4.8 \pm 0.20$ (4-5) |
|  |  | Rhynchocorin |  |  |
|  |  | i | $32.5 \pm 6.5$ (26-39) | $6.3 \pm 0.67$ (5-9) |
| Scutelleridae Thyreocorida e | Phyllocephalina e |  |  |  |
|  |  |  | $36.0 \pm 1.00$ (35-37) | 5 |
|  |  |  | $27.6 \pm 1.20$ (26-30) | $10.7 \pm 0.88$ (9-12) |
|  |  |  |  |  |
|  |  |  | $17.0 \pm 0.53$ (14-21) | $4.2 \pm 0.11$ (4-5) |
|  | Corimelaeninae |  | $17.3 \pm 0.56$ (14-21) | $4.2 \pm 0.12$ (4-5) |
|  | Thyreocorinae |  | $15.0 \pm 1.00$ (14-16) | 4 |



Fig. 1. Foreleg and pretarsus of the Pentatomidae. A, lateral view of the tibial and the tarsal segments of the pentatomid leg, the foretibial apparatus is indicated by a red arrow, whereas the tibial comb, by a dashed red square; B, Ventrodistal view of the foreleg pretarsus (Dichelops furcatus); C, distal view of the foreleg pretarsus (Mormidea v-luteum); D, structures on the rim of the distal tarsomere, unguitractorial plate and parempodia (Dichelops furcatus); E-F, parempodia (Agroecus griseus and Rhyssocephala infuscata, respectively). as, asymmetric seta; bp, basipulvillus; c, claw; dp, distipulvillus; ls, lateral seta; lsup, lateral surface of unguitractor plate; mmic, membrane with microtrichia; ms, median sclerite; pp, parempodial projection; ps, parempodial seta; psc, parempodial sclerite; tr, rim of distal tarsomere; vs, ventral seta; vsup, ventral surface of unguitractor plate. Scales: B-C, $50 \mu \mathrm{~m}$; EF, $10 \mu \mathrm{~m}$.


Fig. 2. Pretarsus structures in the Pentatomidae, the Scutelleridae and the Thyreocoridae. AM, Pentatomidae; A, Aphylinae sp.; B, Oplomus salamandra (Asopinae); C, Cyrtocoris trigonus (Cyrtocorinae); D, Edessa quadridens (Edessinae); E-L, Pentatominae; E, Plautia stali (Antestini); F, Cosmopepla decorata (Carpocorini); G, Dolycoris baccarum (Carpocorini); H, Rhyssocephala infuscata (Catacanthini); I, Chlorocoris (Chlorocoris) complanatus (Chlorocorini); J, Chloropepla vigens (Chlorocorini); K, Neotibilis compascens (Pentatomini); L, Phalaeucus pustulatus (Pentatomini); N, Augocoris gomesi (Scutelleridae); O, Galgupha (Galgupha) cruralis (Thyreocoridae). Scales $=100 \mu \mathrm{~m}$.


Fig. 3. Unguitractorial plate and related structures in the Pentatomidae. Arrows indicate the asymmetric sensilla. A, Alcaeorrhynchus grandis (Asopinae); B, Alitocoris parvus (Discocephalinae); C, Lopadusa augur (Edessinae); D, Antestiade genera (Antestini); E, Cuspicona simplex (Rhynchocorini); F, Dichelops (Diceraeus) melacanthus (Carpocorini); G, Chlorocoris (Monochrocerus) subrugosus (Chlorocorini); H, Rhyncholepta meinanderi (Chlorocoris); I, Banasa patagiata (Pentatomini); J, Pellaea stictica (Pentatomini); K, Lobepomis peltifera (Procleticini). Scale: $25 \mu \mathrm{~m}$.


Fig. 4. Variation of the parempodial projections in the Pentatomidae. A, Agroecus griseus (Carpocorini); B, Taurocerus achilles (Pentatomini); C, Cosmopepla decorata(Carpocorini); D, Rhyssocephala infuscata (Catacanthini); E, Oplomus salamandra (Asopinae); F, Mormidea v-luteum (Carpocorini); G, Banasa zeteki (Pentatomini); H, Thoreyella trinotata (Procleticini). Scales: $10 \mu \mathrm{~m}$.


Fig. 5. Tibial combs in the Pentatomidae, the Scutelleridae and the Thyreocoridae. A-L, Pentatomidae; A, Antestiopsisanchora (Pentatominae, Antestini), arrows indicate the location of the tibial comb at the apex of tibia; B, Oplomus salamandra (Asopinae); C, Podisus nigrispinus (Asopinae); D, Lincus securiger (Discocephalinae); E, Edessa quadridens (Edessinae); F-L, Pentatominae; F, Plautia stali (Antestini); G, Dichelops (Dichelops) punctatus(Carpocorini); H, Arocera placens (Catacanthini); I, Chlorocoris (Monochrocerus) flaviviridis (Chlorocorini); J, Neotibilis compascens (Pentatomini); K, Thoreyella trinotata (Procleticini); L, Biprorulus bibax (Rhynchocorini); M, Augocoris gomesi (Scutelleridae); N, Orsilochides leucoptera (Scutelleridae); O, Galgupha (Euryscytus) difficilis (Thyreocoridae); P, Thyreocoris scarabaeoides (Thyreocoridae). Scales: $100 \mu \mathrm{~m}$.


Fig. 6. Foretibial apparatus in the Pentatomidae, the Scutelleridae and the Thyreocoridae. A, Augocoris gomesi (Scutelleridae); B, Galgupha (Galgupha) cruralis (Thyreocoridae); C, Cydnoides (Cydnoides) renormatus (Thyreocoridae); D-O, Pentatomidae; D, Aphylinae sp.; E, Oplomus cruentus (Asopinae); F, Cyrtocoris trigonus (Cyrtocorinae); G, Lincus securiges (Discocephalinae); H, Brachystethus geniculata (Edessinae); I-O, Pentatominae; I, Antestia ellenriederi (Antestini); J, Dichelops (Dichelops) melacanthus(Carpocorini); K, Arocera apta (Catacanthini); L, Loxa deducta (Chlorocorini); M, Pentatoma rufipes (Pentatomini); N, Thoreyella trinotata (Procleticini); O, Biprorulus bibax (Rhynchocorini); P, Macrina juvenca (Phyllocephalinae). Scales: $50 \mu \mathrm{~m}$.

## CAPÍTULO III ${ }^{3}$

# Contributions to the knowledge of Dichelops Spinola: description of a new species of Dichelops (Diceraeus) and of the male of Dichelops (Prodichelops) divisus (Heteroptera: Pentatomidae: Pentatominae: Carpocorini) 

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#### Abstract

A new species of Dichelops (Diceraeus) Dallas and the so-far unknown male of Dichelops (Prodichelops) divisus (Walker, 1867) are described, based upon morphological characters. D.(Dice.) caatinguensis sp. nov. is compared to the other four species of Diceraeus. Revised keys to separate the subgenera of Dichelops and the species of Diceraeus are also presented. $D$. (P.) divisus male is described and compared to the type species. Comparative illustrations of external and internal genitalia of the species are provided.


Key words: genitalia, Hemiptera, Heteroptera, Neotropical, taxonomy.

[^2]
## Introduction

The Neotropical genus Dichelops Spinola, 1837 was revised by Grazia (1978); it comprises three subgenera, Dichelops Spinola, 1837, Diceraeus Dallas, 1851 and Prodichelops Grazia, 1978. Subgenus Dichelops is the most diverse, bearing ten species (Grazia, 1978; Klein et al., 2012). The genus was recently revised by Klein et al. (2012): a new species was described, the genitalia terminology was updated, and an adapted key to the subgenus was provided.

The subgenera Diceraeus is, so far, represented by four species: D. (Dice.) furcatus (Fabricius, 1775), D. (Dice.) lobatus Grazia, 1978, D. (Dice.) melacanthus (Dallas, 1851) and D. (Dice.) phoenix Grazia, 1978. The study of material collected in northeast Brazil revealed a species, whose morphological characteristics resemble Diceraeus, but do not match with any of the species already described. Herein we describe Dichelops caatinguensis sp. nov. which is included in Diceraeus by serrated anterolateral and sinuated posterolateral margins of pronotum, presence of ventral process of conjunctiva, pars intermedialis twisted, and capsula seminalis lacking processes. Morphological comparisons of male and female genitalia among these species are presented. Additionally, a new key to the species of Diceraeus, adapted from Grazia (1978), is provided.

The subgenus Prodichelops is monotypic and was described from one female specimen, the holotype of $D$. (P.) divisus (Walker, 1867). It is characterized by humeral angles developed; anterolateral margins of pronotum crenulated, posterolateral margins sinuated, lacking projections, and capsula seminalis lacking processes. Recently, three specimens (one male and two females) were collected in the Brazilian and Colombian Amazonia. Here we describe the male of $D$. (P.) divisus, aiming to recognize the subgenus characteristics and to do comparisons with other subgenera. A revised key to separate the subgenera of Dichelops is also provided.

## Material and methods

The study of Dichelops (Diceareus) caatinguensis sp. nov. was based on 40 specimens from the following collections: Fundação Oswaldo Cruz, Rio de Janeiro, Brazil (FIOC); Museu de Entomologia Pe. Jesus Santiago Moure, Universidade Federal do Paraná, Brazil (DZUP); Museu Nacional, Universidade Federal do Rio de Janeiro, Brazil (MNRJ); Museu de Zoologia, Universidade de São Paulo, Brazil (MZSP). The Dichelops (P.) divisus male description was based on the single specimen available, belonging to the Instituto Nacional de Pesquisas da Amazonia, Manaus, Brazil (INPA) collection.

The genital terminology follows Dupuis (1970), Schaefer (1977), Grazia (1978), and Klein et al. (2012). The structures of the pygophore named "processos do diafragma" by Grazia (1978) are here treated as "superior process of dorsal rim" (Grazia \& Schwertner, 2008; Klein et al., 2012). In the female the "vesicula" present on pars intermedialis is considered here a twist of the duct in this structure.

For the study of internal genitalia, specimens were boiled in hot water before dissection and then prepared with a hot saturated potassium hydroxide solution. Measurements (average $\pm$ standard deviation, minimum and maximum values) are given in millimeters.

## Results

## Key to the subgenera of Dichelops (modified from Grazia, 1978)

1. Anterolateral margins of pronotum toothed or crenulated; black spiracles; abdominal punctures darker than the abdominal surface; conjuntiva lacking processes; pars intermedialis not twisted; capsula seminalis with or without processes

1'. Anterolateral margins of pronotum serrated; posterolateral margins sinuated or crenulated (Fig. 1); spiracles and abdominal punctures concolorous with abdominal surface; process of conjunctiva present (Fig. 10); pars intermedialis twisted (Fig. 15); capsula seminalis lacking processes (Fig. 15) ... Diceraeus
2. Anterolateral margins of pronotum toothed; posterolateral margins emarginated, with rhomboid projection between humeral angles and hemelytra base; process of conjunctiva absent; capsula seminalis with processes ... Dichelops

2'. Anterolateral margins of pronotum crenulated; posterolateral margins sinuated, lacking projections; pygophore not dissected; capsula seminalis lacking processes ... Prodichelops

## Dichelops (Diceraeus) Dallas, 1851

Diceraeus Dallas, 1851: 193, 208.
Dichelops (Neodichelops) Grazia, 1978: 14, 63-65.
Dichelops (Diceraeus) Rider, 1998: 505.

Type species: Cimex furcatus Fabricius, 1775 for Neodichelops Grazia, 1978 by original designation; Diceraeus melacanthus Dallas, 1851 for Diceraeus by monotypy.

Diagnosis. Anterolateral margins of pronotum serrated, posterolateral margins sinuated or crenulated; spiracles, and abdominal punctures concolorous with abdominal surface. Apex of scutellum without punctures, bearing a rounded or striped yellowish callus. Male: posterolateral angles of pygophore lacking bristle tufts; superior process of dorsal rim conspicuous, in a tubercle or forming a flap projected towards paramere base; parameres elongated, robust, and flattened; paramere head as long as base; ventral process of conjunctiva present. Female: pars intermedialis twisted; capsula seminalis lacking processes.

Grazia (1978) described the presence of the vesica and the processus vesicae in the phallus of Diceraeus. Actually, the processus vesicae formerly described by Grazia (1978) corresponds to the process of the conjunctiva (Fig. 10), and the former vesica corresponds to the processus vesicae (Fig. 12). The processus vesicae is the same structure observed in the subgenus Dichelops by Klein et al.(2012).

The vesicle of pars intermedialis (Grazia, 1978), present in all species of Diceraeus, is here indicated by a twist in this structure. We chose this designation, because the word "vesicle" denotes a secretory function, which cannot be ascertained by now. The extension of the twisted area is variable, occupying two-thirds of the pars intermedialis length in $D$. furcatus, D. melacanthus, and D. phoenix, and the entire pars intermedialis's length in $D$. lobatus. In all these species the duct is tightly-twisted; however in D. caatinguensis sp. nov. the duct is loosely-twisted and occupies the basal half of the pars intermedialis.

## Key to the species of subgenus Diceraeus (modified from Grazia, 1978)

1. Connexivum lacking spots or with diffuse castaneous spots; scuttelum apex with yellowish stripe ... 2

1'. Connexivum with dark spots at anterior and posterior margins; scuttelum apex with yellowish callus ... D. (Dice.) caatinguensis sp. nov.
2. Jugae acute; humeri varying from underdeveloped to forming long spines; pars intermedialis tightly-twisted along two-thirds of its length ... 3

2'. Jugae obtuse; humeri with rhomboid projections laterally expanded; pars intermedialis tightly-twisted all its length ... D. (Dice) lobatus Grazia
3. Gonocoxites 8 inflated, with the apical quarter folded dorsally, forming a conspicuous arch in lateral view; ventral rim of the pygophore straight or moderately sinuated at the middle; superior process of dorsal rim of the pygophore variable ... 4

3＇．Gonocoxites 8 less inflated，almost flattened；ventral rim of the pygophore medially bisinuated；superior process of dorsal rim in a scalloped flap structure ．．．D．（Dice．） melacanthus Dallas

4．Superior process of dorsal rim of the pygophore in an entire flap structure；outer face of paramere with a hook projection；black punctures in the anterior half of the pronotal anterolateral margins，around the pronotal cicatrices and，forming three parallel stripes along the costa，in exochorion，often distinct ．．．D．（Dice．）phoenix Grazia

4＇．Superior process of dorsal rim of the pygophore in a tubercle placed inside the genital cup；paramere lacking projections；black punctures in the pronotum anterolateral margins and in exochorion，seldom distinct ．．．D．（Dice．）furcatus Fabricius

## Dichelops（Diceraeus）caatinguensis Grazia \＆Poock－da－Silva sp．nov．

（Figs．1，3－15）

Etymology．The name is related to its geographic distribution（Caatinga biome，northeast Brazil，Rio Grande do Norte，Paraíba，Pernambuco，and Bahia states）

Type material．Holotype $\widehat{\lambda}$ ，labeled：BRAZIL，Paraíba，Soledade，Juazeirinho， 08．VII．1956，A．G．A．Silva col．，Coleção Campos SEABRA（MNRJ）．Paratypes：1才̂，1q， BRAZIL，Rio Grande do Norte，Baixa Verde，VII．1950，B．J．Souza col．，Coleção Campos SEABRA（UFRG）；1q，Paraíba，Soledade，Juazeirinho，22．III．1956，A．G．A．Silva col．， Coleção Campos SEABRA（MNRJ）；1才，Paraíba，Soledade，Juazeirinho，08．VII．1956， A．G．A．Silva col．，Coleção Campos SEABRA（MNRJ）；1才，10ㅇ，Pernambuco，Casa Nova， V．1974，J．C．M．Carvalho col．，Caatinga，（ $1 \delta^{\lambda}, 2 q$ UFRG， $8 q$ MNRJ）； $6 q$ ，Pernambuco， Petrolina，V．1974，J．C．M．Carvalho col．，Caatinga（2 $q$ MCNZ， $4 q$ MNRJ）；1q， Pernambuco，Petrolina，V．1969，M．Alvarenga col．（MNRJ）；1ㅇ，Pernambuco，Petrolina，

Estrada Picos, V.1974, J.C.M. Carvalho col., caatinga (MNRJ); 1 , divisa entre Pernambuco e sul do Piauí, Estrada Picos Km3, V.1974, J.C.M. Carvalho col., caatinga (MNRJ); 1ठ, Bahia, Anajê, 16.V.1975, C. \& P. Elias col. (MCNZ); $1 \delta^{\lambda}, 1$ ㅇ, Bahia, Brumado, 9.V.1975, C. \& P. Elias col. (DZUP 212768, 212766); 1§, 4 4 , Bahia, Juazeiro, V.1974, J.C.M. Carvalho col., Caatinga ( $\uparrow$ MNRJ); 1 Q, Bahia, Juremal, Estrada do Juazeiro, V.1974, J.C.M. Carvalho col., Caatinga (MNRJ); 1q, Bahia, G. Bondar col.; $1 \widehat{ }^{\lambda}, 2$ Q , nordeste, 1933, Ihering col. ( $1 \delta^{\star}$ UFRG; $2 q$ FIOC); $2 q$ (MZSP 72407, 72408).

Description.Ovoid, medium-sized $(8-10 \mathrm{~mm})$ species. Dorsal color castaneous (Fig. 1), lighter castaneous ventrally.

Head slightly longer than wide (Fig. 1). Dorsal surface with coarse punctures regularly distributed, except in a circular area adjacent to eyes; ventral surface with sparsely distributed punctures. Juga clearly surpassing clypeus, length before clypeus equals one-third of first antennal segment; lateral margins straight, often convergent, and outlined by black, apex obtuse. Proportion of antennal segments: I $<$ II $<$ III $<$ IV $>V$. Rostrum slightly surpassing metacoxae. First rostral segment entirely contained between buccula. Proportion of rostral segments: I<II>III>IV.

Pronotum. Anterolateral margins serrated. Posterolateral margins sinuated. Humeri varying from rhomboid to acutely projected, black at apex. Cicatrices sometimes delimited by dark punctures. Darker punctures concentrated at middle pronotum, forming a rounded spot.

Scutellum. Surface uniformly punctured, forming rounded areas lacking punctures. Two pairs of black spots formed by concentrated punctures, one basally and other medially. Distal region without punctures, bearing a yellow callus. Small black fovea at basal angles.

Hemelytra. Rusted-red punctures uniformly distributed. Reddish radial vein. Black spots near distal apex of radial vein.

Ventral surface light castaneous, punctures uniformly distributed and thinner than those on dorsal surface. Spiracles concolorous to ventral surface. Connexivum light castaneous, with dark spots at anterior and posterior margins. Legs light castaneous, with reddish spots in all segments.

Male genitalia. Pygophore subquadrangular (Figs. 5-8); genital cup well exposed dorsally (Figs. 5-6). Dorsal rim medially sinuated (Figs. 5-6). Superior process of dorsal rim conspicuous, as a sinuated flap projected into genital cup towards paramere base (Fig. 6). Posterolateral angles slightly projected, without bristle tufts. Ventral rim bisinuated, in a 'V' excavation at middle, with bristle tufts (Figs. 7-8). Parameres elongated, robust and flattened, directed dorsally. Paramere head as long as base, with a digitiform projection on inner surface (Fig. 9). Phallus almost pyriform in lateral view (Fig. 12); processus phallothecae not surpassing conjunctiva (Figs. 10, 12); ventral process of conjunctiva projecting towards phallotheca ventral wall (Figs. 10, 12); processus vesicae embracing ductus seminis distalis (Fig. 11).

Male measurements ( $\mathrm{n}=6$ ). Body length $8.81 \pm 0.22$ (8.48-9.12); abdominal width $5.09 \pm 0.28$ (4.8-5.44); head length $2.13 \pm 0.096$ (2.08-2.32), width $2.07 \pm 0.12$ (1.92-2.24); interocular distance $1.24 \pm 0.067$ (1.2-1.36); length of antennal segments: I $-0.55 \pm 0.06$ ( $0.48-$ 0.64 ), II $-0.83 \pm 0.082$ ( $0.8-0.96$ ), III $-1.13 \pm 0.15$ ( $0.96-1.36$ ), IV $-1.16 \pm 0.62$ (1.12-1.84), V $-1.16 \pm 0.61$ ( $0.96-1.36$ ); pronotum length $2.04 \pm 0.13$ (1.84-2.24), width $4.92 \pm 0.34$ (4.45.36), length to the height of the spine $5.89 \pm 0.4$ (5.52-6.64); scutellum length $3.0 \pm 0.13$ (2.83.12), width $3.11 \pm 0.12$ (2.88-3.20).

Female genitalia.Gonocoxites 8 almost triangular, not covering gonocoxites 9; sutural margins sinuated, overlapping at base, sutural angles acute, tumid areas occupying $1 / 2$ of each plate (Figs. 13-14). Laterotergites 8 acute at apex, but not forming a spine. Laterotergites 9 obtuse at apex, not surpassing laterotergites 8 (Figs. 13-14).Posterior margin of gonocoxites

9 biconvex.Chitinellipsen rounded (Fig. 15). Ductus receptaculi before vesicular area almost three times longer than ductus after vesicular area (Fig. 15). Internal wall of vesicular area conical at base; median wall dilated at basal quarter of vesicular area (Fig. 15). Basal half of pars intermedialis loosely-twisted; anterior annular flange flat, posterior annular flange convergent. Capsula seminalis rounded, lacking processes.

Female measurements ( $\mathrm{n}=31$ ). Body length $9.56 \pm 0.37$ (9.2-10.24); abdominal width $5.57 \pm 0.22$ ( $5.28-5.92$ ); head length $2.13 \pm 0.2$ (1.92-2.48), width $2.22 \pm 0.07$ (2.16-2.32); interocular distance $1.34 \pm 0.09$ (1.28-1.52); length of antennal segments: I $-0.56 \pm 0.07$ ( $0.48-$ $0.64)$, II $-0.85 \pm 0.065$ ( $0.8-0.96$ ), III - $1.2 \pm 0.06$ (1.12-1.28), IV $-1.50 \pm 0.14$ (1.36-1.68), V - $1.4 \pm 0.1$ (1.28-1.52); pronotum length $2.28 \pm 0.16$ (2.08-2.48), width $5.47 \pm 0.3$ (5.12-5.92), length to the height of the spine $6.47 \pm 0.5$ (5.92-7.2); scutellum length $3.64 \pm 0.13$ (3.44-3.76), width $3.61 \pm 0.21$ (3.28-3.84).

Comments. Dichelops caatinguensis sp. nov. is similar to Dichelops lobatus. From D. lobatus it can be distinguished by being smaller in length (male: $8.81 \pm 0.22 \mathrm{~mm}$; female: $9.56 \pm 0.36 \mathrm{~mm})$; by the presence of a yellow spot at the scutellum apex, which in $D$. lobatus is a stripe; connexivum with $1+1$ dark spots at the anterior and posterior margins, while immaculate in $D$. lobatus; jugae convergent or juxtaposed, while parallel in $D$. lobatus. In the male genitalia both species have the paramere and the superior process of the dorsal rim similar in shape; Dichelops caatinguensis sp. nov. can be distinguished from D. lobatus by the ventral rim's being bisinuated, excavated in a ' V ' at the middle, and with bristle tufts. Dichelops caatinguensis sp. nov. females can be distinguished from all Diceraeus species by the posterior margin of the gonocoxites 9 biconvex and by the pars intermedialis being loosely twisted along the basal half of its length.

Distribution. Northeast Brazil, corresponding to Caatinga, Cerrado and Para biogeographic provinces (sensu Morrone, 2006).

## Dichelops (Prodichelops) Grazia, 1978

Dichelops (Prodichelops) Grazia, 1978: 14, 87-89.

Type species: Diceraeus divisus Walker, 1867 by original designation.

Diagnosis. Head ventrally with narrow black stripe, sub-adjacent to juga outer margin, extending from antennal tubercle to bucculae apex. Pronotum with a transversal orange or yellowish stripe, without punctures; anterolateral margins crenulated; posterolateral margins sinuated, without projections (Fig. 2). Humeral angles developed. Male: pygophore dorso posteriorly opened, genital cup widely exposed; ventral rim of pygophore ventrally projected in a beaklike structure; parameres scythe-like, long, and slender. Female: pars intermedialis not twisted; capsula seminalis without processes.

## Dichelops (Prodichelops) divisus (Walker, 1867)

(Figs. 2, 16-21)
Diceraeus divisus Walker, 1867: 250.
Dichelops divisus; Lethierry \& Severin, 1893: 129; Kirkaldy, 1909: 68.
Dichelops (Prodichelops) divisus; Grazia, 1978: 90-91.

Type: BMNH, HEM. 1056, female, Bates col., deposited on "British Museum (Natural History)", examined.

Type locality. "Amazon Region"; according to label information, the specimen was collected in "St. Paulo" [São Paulo de Olivença, Rio Solimões, Amazonas, Brazil].

Head. Male proportion of antennal segments: $\mathrm{I}<\mathrm{II}=\mathrm{III}<\mathrm{IV}>\mathrm{V}$; female proportion of antennal segments: $\mathrm{I}<\mathrm{II}<\mathrm{III}<\mathrm{IV}<\mathrm{V}$.

Male genitalia. Pygophore quadrangular (Fig. 21); genital cup widely open dorso posteriorly (Figs. 16-19). Dorsal rim in an open 'U', sinuate at base, slightly projected over segment X (Figs.18-19). Superior processes of dorsal rim spine-like placed before middle of lateral margins of dorsal rim, clearly distant from paramere apices. Posterolateral angles almost truncate not projected posteriorly, with dense and convergent bristle tufts (Figs. 1619). Ventral rim bisinuated, at middle in a shallow 'U' (Figs. 16-17, 20-21); inferior layer of ventral rim projecting ventrad in a beaklike structure, well-observed in profile and/or in posterior view of pygophore (Figs. 16-17). Bristles along ventral rim, especially over folded area inside genital cup, forming $1+1$ tuft adjacent to posterolateral angles, $1+1$ tuft laterad to 'U' concavity, and one tuft at middle of 'U' (Figs. 17, 21). Parameres long and slender, scyte like, laterad directed. Segment X almost pyriform, with $1+1$ minute tubercles at middle (Fig. 19).

Male measurements $(\mathrm{n}=1)$. Body length 8.96; abdominal width 5.44; head length 1.92, width 2.0 ; interocular distance 0.96 ; length of antennal segments: $\mathrm{I}-0.72$, II $-0.8, \mathrm{III}-0.8$, IV $-1.12, \mathrm{~V}-1.6$; pronotum length 2.16 , width 5.6 , length to the height of the spine 6.4 ; scutellum length 3.28 , width 3.2.

Female measurements ( $\mathrm{n}=2$ ). Body length $9.32 \pm 0.06$ (9.28-9.36); abdominal width $5.68 \pm 0.11$ (5.6-5.76); head length $1.8 \pm 0.06$ (1.76-1.84), width $2.2 \pm 0.06$ (2.16-2.24); interocular distance $1.12 \pm 0.23$ (0.96-1.28); length of antennal segments: I $-0.52 \pm 0.06(0.48-$ 0.56 ), II $-0.76 \pm 0.06(0.72-0.8)$, III $-0.92 \pm 0.06$ ( $0.88-0.96$ ), IV $-1.2, \mathrm{~V}-1.6$; pronotum length $1.96 \pm 0.06$ (1.92-2.0), width $6.2 \pm 0.06$ (6.16-6.24), length to the height of the spine $7.56 \pm 0.51$ (7.2-7.92); scutellum length $3.8 \pm 0.06$ (3.76-3.84), width $3.76 \pm 0.34$ (3.52-4.0).

Material examined. BRAZIL, Amazonas, Estirão da Preta, Rio Liberdade, 1115.V.2011, D. Takyia col., sweep net (INPA), $1 \widehat{O}^{\lambda}, 1$ q. COLOMBIA, Amazonas, Prefectura de Tabatinga [Leticia], 12.IV.1992, A. Saenz col. (UFRG), 1 ㅇ.

Comments. The male genitalia of Dichelops (Prodichelops) divisus are similar to the species of Dichelops (Dichelops) by the superior process of the dorsal rim as a spine-like structure, and by the parameres' shape, wide at the base, tapering toward the apex. Dichelops (Prodichelops) divisus can be distinguished from the other subgenera by the dorsal rim of the pygophore being medially bisinuated, and by the ventral rim of the pygophore being ventrally projected, forming a beaklike structure.

Distribution. Western Amazonia, in Brazil and Colombia, which corresponds to Emeri, Varzea and Madeira biogeographic provinces (sensuMorrone, 2006).

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FIGURES 1-2. Dorsal view: 1, Dichelops (Diceraeus) caatinguensis sp. nov.; 2, Dichelops divisus $($ Scales $=1 \mathrm{~mm})$.


FIGURES 3-12. Dichelops (Diceraeus) caatinguensis sp. nov., male genitalia. 3-4, pygophore, posterior view; 5-6, pygophore, dorsal view; 7-8, pygophore, ventral view; 9, left paramere, dorsal view; 10-12, phallus, ventral, dorsal and lateral views, respectively. (cj, conjunctiva; dr, dorsal rim; dsd, ductus seminis distalis; gs, secondary gonopore; pa, paramere; pcj, process of conjunctiva; ph, phallotheca; prph, processus phallothecae; prv, processus vesicae; spdr, superior process of dorsal rim; vr, ventral rim; X, proctiger) (Scales: figs. 3-8 $=0.5 \mathrm{~mm} ; 9-12=0.15 \mathrm{~mm}$ ).



FIGURES 16-21. Dichelops (Prodichelops) divisus, pygophore. 16-17, posterior view; 1819, dorsal view; 20-21, ventral view. (dr, dorsal rim; pa, paramere; spdr, superior process of dorsal rim; vr, ventral rim; $X$, tenth segment $)($ Scales $=0.5 \mathrm{~mm})$.

## CAPÍTULO IV ${ }^{4}$

# Phylogeny of Dichelops (Heteroptera, Pentatomidae) 

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#### Abstract

Dichelops comprises 16 species, included in three subgenera, with exclusive Neotropical distribution. The genus was revised recently and each subgenus have a set of distinct morphological characters, which allow their precise identification, and different distributional patterns, thus raising questions about the genus monophyly and the subgenera evolutionary relationships. The genus monophyly was tested under parsimony using discrete and continuous characters and two character weighting methods, equal weights and implied weights. The genus was paraphyletic in every analysis and the characters formerly proposed as diagnostic for the genus and subgenera were found to be shared with other taxa. Discussion about continuous characters treatment and use of weighting methods are provided.


Keywords: continuous characters, implied weighting, Neotropics, systematics.

[^3]
## Introduction

The cosmopolitan Pentatominae (Hemiptera: Pentatomidae) comprises about $60 \%$ of Pentatomidae diversity. Tribal classification of pentatomines has been changing for the last four decades and is still under dispute, ranging from eight (Schuh \& Slater, 1995) to 43 (D.A. Rider in litt.) recognized tribes. In the latter tribal arrangement, the Neotropical Pentatomini sensu Rolston \& McDonald (1979; 1980a,b; 1984) was divided into seven tribes: Antestiini Distant, 1902, Carpocorini Mulsant \& Rey, 1866, Menidini Atkinson, 1888, Nezarini Atkinson, 1888, Pentatomini Leach, 1815, Piezodorini Atkinson, 1888, and Strachiini Mulsant \& Rey, 1866.

Carpocorini, the most diverse pentatomine tribe (ca. 450 spp . and 100 genera), occurs in all continents except Antarctica. The highest tribal diversity occurs in the New World (ca. 290 spp. and 55 genera) and the most diverse genera are Euschistus Dallas, 1851 (74 spp.), Mormidea Amyot \& Serville, 1843 (34 spp.), and Dichelops Spinola, 1837 (16 spp.). Some groups of genera within Carpocorini have been proposed based on morphological similarities and taxonomic history, but have not been tested with up-to-date cladistic methods.

Euschistus-group is one of those. Rolston (1974) grouped Agroecus Dallas, 1851, Dichelops, Euschistus, Galedanta Amyot \& Serville, 1843, Hymenarcys Amyot \& Serville, 1843, Menecles Stål, 1867, Proxys Spinola, 1837, Sibaria Stål, 1872, and Tibraca Stål, 1860, by similarities in coloration, size, and general morphology. One could add to this group, genera that have been erected from the above genera, i.e. Ladeaschistus Rolston, 1973 and Mcphersonarcys Thomas, 2012; or lately related taxonomically to them, i.e. Acledra Signoret, 1864, Berecynthus Stål, 1862, Caonabo Rolston, 1974, Coenus Dallas, 1851, Padaeus Stål, 1862, Oenopiella Bergroth, 1891, and Spinalanx Rolston \& Rider, 1988. According to Rider (1995), a diagnostic feature of the group is the presence of a pair of dorsal processes on the phallotheca (sic dorsal thecal appendages). Recently, Tibraca has been
referred as belonging to the Mecocephala group, which also has dorsal processes on the phalloteca, but is distinct in having very reduced or absent parameres (Frey-da-Silva et al., 2002; Schwertner et al. 2002).

The Euschistus-group includes some species associated to crops of legumes and which are important crop pests in the Neotropics. However, the Euschistus group has never been cladistically tested before, neither was the Carpocorini. In an effort to best understand the relationships and biogeographic history of Neotropical pentatomids, the Carpocorini and some genera of the Euschistus-group are under formal phylogenetic investigations (e.g. Agroecus, Euschistus, Proxys).

Here we study the phylogenetic relationships of Dichelops. The genus comprises 16 species, collectively recognized by having the mandibular plates longer than the clypeus, included in three subgenera: Dichelops Spinola, 1837 (10 spp.), Diceraeus Dallas, 1851 (5 spp.), and Prodichelops Grazia, 1978 (1 sp.). The genus was revised by Grazia (1978) and few species have been described since then (Klein et al., 2012; Poock-da-Silva et al., 2013). Each subgenus has a set of distinct morphological characters, which allow their precise identification, and different distributional patterns, thus raising questions about the genus monophyly and the sub-genera evolutionary relationships.

In this work two issues are addressed: (1) the monophyly of Dichelops and (2) the hypothesized generic affinities of the Euschistus-group.

## Material and methods

## Ingroup, outgroup and root

In the absence of a phylogenetic hypothesis to the Euschistus-group and to the Carpocorini, outgroup selection was based on taxonomic history and morphological diversity of the characters here studied (Barão et al., 2012). Rolston (1974) grouped on the same
dichotomous key genera he considered similar to Euschistus, being followed by subsequent authors in considering those genera as belonging to the Eushistus-group.

The ingroup taxa (Table S1) comprises the 16 species currently assigned to Dichelops subgenera: 10 of Dichelops, five of Diceraeus, and one of Prodichelops. Out of the remaining 15 genera considered part of Euschistus-group, 27 species belonging to 13 genera were included (Table S1; Mcphersonarcys and Sibaria were not included). Other 17 species (belonging to 11 genera) were included, representing taxa not related taxonomically to Euschistus-group (Table S1) but included in Carpocorini sensu Rider (in litt.). Trees were rooted on Carpocoris, the Carpocorini type genus. Appendix S1 is a list of examined specimens and its depositories.

## Terminology

Terminology derives from Tsai et al. (2011) for general morphology; Baker (1931), Dupuis (1970) and Schaefer (1977) for genital terminology; and Kment \& Vilímová (2010) for external scent efferent system (ESES) of the metasternal gland.

## Character matrices

Of the 149 characters included, 10 are continuous and 139 are discrete characters. Discrete characters represent general morphology (73), female (24) and male genitalia (42). Terminals were coded for characters directly from series of available specimens (Appendix S1) and coded from literature when specimens were not available (Table S1).

Matrix of discrete characters was constructed on Mesquite 3.03 (Maddison \& Madisson, 2015) and matrix of continuous characters was constructed using a spreadsheet. Continuous characters comprise the range of one standard error around the mean (i.e. [mean -
$\mathrm{SE}]-[$ mean +SE$]$ ). The two data matrices were combined into one matrix and formatted to run on TNT using a text editor. Matrix is available on Appendix S2.

Character statements were elaborated following the logical basis proposed by Sereno (2007), especially regarding the use of reductive coding (Forey \& Kitching, 2000; Brazeau, 2011). Inapplicable data were coded as "-" and missing data as "?" - except for continuous characters where inapplicable data were coded as "?", because of TNT continuous characters handling options.

The following notations for characters are used in Results and Discussion sections: " $\mathrm{X}(\mathrm{Y})$ ", in which X represents the character and Y represents the state; " s ", to the number of steps; "CI", to consistency index (Kluge \& Farris, 1969); and "RI", to retention index (Farris, 1989).

## Morphological methods

Internal male and female genitalia were dissected, macerated in a supersaturated KOH solution, stained with Congo Red and/or Chlorazol Black, and observed with a stereo microscope. Images were taken in multiple focal planes using a DS-Fi2 camera coupled to a Nikon AZ100M scope and stacked with the NIS Elements AR software available at the Departament of Zoology, UFRGS.

Structures analyzed using scanning electron microscopy (SEM) were manually cleaned, subsequently kept submersed in contact lens solution (Renu®) for 24 h to degrade proteins, and then sonicated at 5400 kHz during three minutes in a solution of water and detergent. Followed by dehydration by alcoholic series and stove dried at $40^{\circ} \mathrm{C}$ for 24 h , sputter-coated with gold, and observed in a SEM at the Centro de Microcopia Eletrônica (CME) of UFRGS.

Measurements were taken with a micrometer scale attached to the scope's ocular lens. Morphometric parameters (Table S2) were measured for all terminals, in an effort of 20 specimens per species ( 10 spec . per sex), following the recommendations of Marie-Stephane (2013) for Acari. Because of material availability, the number of measured specimens was lower for some species (Table S1). Morphometric parameters and measurements (mean, standard error, maximum and minimum values) are available in Table S2.

## Phylogenetic analyses

Phylogenetic analyses were performed by parsimony using software TNT v.1.1 (Goloboff et al., 2008b); discrete characters were treated as nonadditive and continuous characters as additive (Goloboff et al., 2006). Continuous characters were always rescaled to unit in order to reduce dominance of large characters influence on matrix (Mongiardino et al., 2014). Three matrices were used, with the same set of terminals but varying datasets: discrete characters only, continuous characters only, and continuous + discrete characters.

Matrices were analysed under equal weights (EW) and implied weights (IW). Implied weighting (Goloboff, 1993) downweights characters according to their degree of homoplasy: the higher their homoplasy, the lower their weight. Goloboff et al. (2008a) demonstrated that weighting against homoplasy on morphological datases improves jackknife-frequencies and produces more stable trees. Intending to explore the effect of datasets (continuous and discrete characters) and different weighting methods (EW and IW), five analyses (A-E) were performed with different combinations of datasets and weighting methods (Fig. 1). Matrices of discrete characters and continuous characters were analysed under EW and IW, whereas matrix of continuous character only was analysed solely under IW (Fig. 1). Strict consensus was calculated for each analysis as well as jackknife frequencies. The analysis of continuous data only was exploratory and not intended to discuss phylogenetic relationships.

Since Dichelops was recovered paraphyletic, we conducted other five analyses (AcEc; Fig. 1) constraining the genus monophyly, to reflect the alternative a priori taxonomic hypothesis. We understand constraining the monophyly of Dichelops as a way to evaluate how better the results of not constrained analyses are, measured as tree length and/or fit. Constrained analysis followed the same analytical procedures as detailed to analyses A-E. No phylogenetic nor taxonomic conclusions were drawn from it.

Independent of the weighting method employed, heuristic searches were always performed under 700 random addition sequences, TBR branch swapping algorithm retaining 150 trees per replication, random seed 0 , and holding up to 100,000 trees in memory (rseed 0 ; mult $700=$ tbr hold 150). For IW analyses, the concavity value ( $k$-value $=14$ ) was determined as described below and employed in all.

Topology stability was calculated to each analyses by jackknife with symmetric resampling, recording absolute-group frequencies and GC (Group present/Contradicted) frequency differences (Goloboff et al., 2003), using 1000 pseudo-replicates and removal probability of $33 \%$. GC values are informative for the amount of contradictory information in the dataset: if absolute frequencies and GC values are equal, probably no contradictory groups are supported by the data. Compatibility of strict consensuses resultant of the 10 analyses (A-E, Ac-Ec) were compared by Similitud index and Robinson-Foulds distances.

Defining k-value. Approaches to choosing the $k$-value have included exploring an arbitrary range of $k$-values and subsequently evaluating results through consensus methods, and exploring a range of $k$-values adjusted to the fit $(F)$ of an average character (Mirande, 2009).

The protocol proposed by Mirande (2009) was employed to define the $k$-value used in all IW analyses performed in this study. Mirande (2009) proposed using a measure of the "average homoplastic character", based on the number of observed steps of the shortest trees
found under equal weights. Implied weighting formula was thus rewritten as $[k=(F * S) /(1$ - $F$ )], in order to obtain $k$-values, where $S$ is a measure of the average homoplasy per character, calculated as $[S=(($ number of observed steps $)-($ minimum number of steps $)) /$ (number of characters)]. We determined $k$-value based on the discrete dataset only.

Mirande's protocol was set to 24 distortion groups, that assign to an "average" character fits from 50 to $90 \%$, finding the best tree five times, and ratchet and drift set to 10 iterations. Remaining parameters were set as default. The 24 runs resulted in 40 trees; strict consensus of each run was calculated and consensuses were compared through Similitud index (tcomp). Similitud index calculates RI resulting from mapping the Matrix Representation Parsimony of each tree to each other: the higher the index, more similar the trees. Tree topologies resultant from $k$-values ranging from 9.3 to 20.3 were the most similar, and the $k$-value used by us is its average ( $k$-value $=14$ ). Mirande's protocol output and tree comparisons are available on Appendix S3.

## Results

## Character statements

## Continuous characters.

0. Head length
1. Head width between eyes
2. Eyes width
3. Length of mandibular plates ahead of clypeus
4. Interocellar distance
5. Pronotum length
6. Pronotum width at humeral angles
7. Pronotum width at distal margin
8. Scutellum width at basal margin

## 9. Scutellum length

## Discrete characters.

## Head

10. Mandibular plates, length related to apex of clypeus: (0) shorter (Fig. 5); (1) equal (Fig. 4); (2) longer (Fig. 2). Modified from Grazia (1997); Kocorek \& Lis (2000); Weiler (2011).
11. Mandibular plates, form apically: (0) obtuse; (1) pointed.
12. Mandibular plates, apex position in relation to clypeal apex, in lateral view: (0) inferior; (1) leveled; (2) superior. Modified from Weiler (2011).
13. Mandibular plates, inner margins ahead of clypeus: (0) contiguous (Fig. 3); (1) not contiguous (Fig. 2).
14. Mandibular plates, outline color laterally related to background color of head: (0) concolour (Fig. 3); (1) not-concolour (Fig. 2).
15. Mandibular plates, orientation of lateral margins: (0) flat; (1) explanate (Fig. 3); (2) declivous (Fig. 5).
16. Mandibular plates, medial margins, unpunctuated area: (0) absent; (1) present (Fig. 4). Modified from Weiler (2011).
17. Clypeus, proximal limit of clypeal suture related to an imaginary line across anterior margin of eyes: (0) anterior; (1) posterior. Modified from Weiler (2011).
18. Clypeus, form of apex: (0) obtuse; (1) acute. Modified from Weiler (2011).
19. Clypeus, height related to mandibular plates, longitudinally: (0) leveled; (1) higher (Fig. 5).
20. Bucculae, anterior margin, form: (0) truncate; (1) lobate; (2) tapering toward apex of head.
21. Bucculae, posterior margin, form: (0) tapering toward base of head; (1) truncate; (2) lobate. Modified from Bernardes et al. (2009).
22. Labium, length of first segment related to bucculae: (0) contained; (1) surpassing.
23. Labium, placement of apex: (0) between pro- and meso-coxae; (1) between meso- and meta-coxae; (2) beyond metacoxae.
24. Antenniferous tubercles, lateral process: (0) absent; (1) present. Modified from Ferrari (2009).
25. Antenniferous tubercles, in dorsal view of head: (0) not visible; (1) visible. Modified from Grazia et al. (2008).
26. Antenniferous tubercles, ventral process: (0) absent; (1) present. Modified from Ferrari (2009).
27. Anteocular processes: (0) absent; (1) present.
28. First antennal segments, length related to anterior margin of head: (0) shorter; (1) subequal; (2) longer. Modified from Kocorek \& Lis (2000).

## Thorax

29. Pronotum, anterior angles, process: (0) absent; (1) present.
30. Pronotum, anterior angles, shape of process apically: (0) obtuse; (1) acute.
31. Pronotum, anterolateral margins, outline: (0) flat; (1) explanate (Fig. 7).
32. Pronotum, anterolateral margins, form: (0) straight (Fig. 8); (1) concave (Fig. 6); (2) convex (Fig. 7). Modified from Fernandes (1993).
33. Pronotum, anterolateral margins, ornamentation: (0) smooth (Fig. 7); (1) serrated (Fig. 6); (2) crenulated (Fig. 8). Modified from Grazia (1997).
34. Pronotum, humeral angles: (0) developed, (1) obsolete.
35. Pronotum, humeral angles, form: (0) obtuse; (1) pointed (Fig. 6); (2) quadrate (Fig. 9). Modified from Gapud (1991); Grazia (1997); Fortes \& Grazia (2005); Bernardes et al. (2009); Grazia et al. (2008); Weiler (2011).
36. Pronotum, posterolateral margins, ornamentation, form: (0) smooth; (1) crenulated (FIG). Modified from Grazia (1997).
37. Pronotum, posterolateral margins, median tubercle: (0) absent; (1) present (Fig. 6).
38. Pronotum, anterior lobe related to posterior lobe: (0) leveled (Fig. 10); (1) sloping (Fig. 11).
39. Pronotum, transhumeral band: (0) absent; (1) present.
40. Pronotum, transhumeral band, sculpturing: (0) smooth; (1) punctuated.
41. Scutellum, apex, callus(es): (0) absent; (1) present. Modified from Weiler (2011).
42. Scutellum, apex, calloused area: (0) continuous; (1) divided in $1+1$. Modified from Weiler (2011).
43. Scutellum, apex, callus shape: (0) rounded; (1) thin band.
44. Hemelytrum, corium, radial vein apex: (0) punctuated (Fig. 12); (1) calloused (Fig. 13); (2) smooth (Fig. 14). Modified from Barcellos \& Grazia (2003); Weiler (2011).
45. Hemelytrum, membrane, venation: (0) parallel; (1) reticulate. Modified from Gapud (1991); Henry (1997); Grazia (1997); Grazia et al. (2008); Weiler (2011).
46. Hemelytrum, length related to abdominal apex: (0) surpassing; (1) not surpassing.
47. Mesosternum, carina: (0) absent; (1) present.
48. ESES, Peritreme, shape: (0) ruga (Fig. 19); (1) spout (Fig. 20); (2) reniform (Fig. 21). Modified from Kment \& Vilímová (2010).
49. ESES, Peritreme, tegument surface finish: (0) smooth (Fig. 23); (1) circumvolutions (Fig. 24); (2) scales (Fig. 25); (3) cylindrical projections (Fig. 26).
50. Metapleura, range of evaporatorium related to metapleura width: (0) less than half (Fig. 15); (1) more than half (Fig. 16).
51. Metapleura, form of outer margin of evaporatorium: (0) convex (Fig. 15); (1) concave (Fig. 16); (2) straight (Fig. 17).
52. Metapleura, form of antero-lateral margin of evaporatorium: (0) rounded (Fig. 16); (1) acute (Fig. 15); (2) evanescent.
53. Mesopleura, range of evaporatorium related to mesopleura width: (0) less than half (Fig. 17); (1) more than half (Fig. 18).
54. Mesopleura, evaporatorium along the outer margin, in ventral view: (0) absent; (1) present.
55. ESES, Evaporatorium, punctures: (0) absent; (1) present (Fig. 22).
56. ESES, Evaporatorium, sensillum: (0) absent; (1) present (Fig. 22).
57. ESES, Evaporatorium, setae: (0) absent; (1) present (Fig. 22).
58. Metathoracic spiracle, form: (0) narrow (Fig. 27); (1) wide (Fig. 28). Modified from Kment \& Vilímová (2010).
59. Metathoracic spiracle, filter system, surface ornamentation: (0) smooth (Figs. 29, 31); (1) spiny (Fig. 30).
60. Metathoracic spiracle, filter, lateral connections: (0) absent (Fig. 29); (1) present (Figs. $30,31)$.
61. Legs, rounded pigmentation at bristles base: (0) absent; (1) present. Modified from Weiler (2011).
62. Legs, width of pigmentation at bristles base, related to bristles diameter: (0) less than twice ; (1) more than twice. Modified from Weiler (2011).
63. Legs, fore femorae, area around base of setae: (0) smooth; (1) elevated.
64. Legs, anterior tibiae, setae, appearance: (0) thin; (1) stout.
65. Pronotum, cicatrices, callus at postero-median margin: (0) absent; (1) present.
66. Pronotum, humeral angle, orientation: (0) laterad; (1) anteriad.
67. Scutellum, fovea, at latero-anterior angles: (0) absent; (1) present.
68. Scutellum, fovea, width related to ocelli width: (0) smaller; (1) wider.
69. Mesopleura, evaporatorium, diagonal extension towards antero-lateral angle of mesopleura: (0) absent; (1) present (Fig. 18).

Abdomen (pre-genital segments)
70. Sternites, longitudinal stripe medially: (0) absent; (1) present (Fig. 32).
71. Sternites, longitudinal stripes sub-laterally: (0) absent; (1) present (Fig. 32).
72. Sternites, postero-lateal angles protruding from sternite edge: (0) absent; (1) present
73. Sternites, postero-lateral angles, form of apex: (0) obtuse (Fig. 36); (1) acute (Fig. 37).
74. Sternites, coloration pattern of antero-lateral angles related to discal coloration: (0) concolour (Fig. 34); (1) not-concolour (Fig. 35).
75. Sternites, coloration pattern of postero-lateral angles related to discal coloration: (0) concolour (Fig. 34); (1) not-concolour (Fig. 35).
76. Sternites, calluses along posterior margin: (0) absent; (1) present (Fig. 33).
77. Sternite 7, development of postero-lateral angles related to postero-lateral angles of remaining abdominal sternites: (0) equal (Fig. 36); (1) longer (Fig. 38).
78. Sternite 7, length medially related to length at lateral margin in females: (0) equal; (1) longer.
79. Spiracles, color in relation to abdominal color: (0) concolour; (1) not-concolour.
80. Connexivum, in dorsal view, related to mesosternal wing development: (0) exposed; (1) concealed.
81. Connexivum, coloration pattern of antero-lateral angles related to discal coloration: (0) concolour; (1) not-concolour.
82. Connexivum, coloration pattern of postero-lateral angles related to discal coloration: (0) concolour; (1) not-concolour.

Female genitalia
83. Genital plates, gonocoxites 8 , basal portion of sutural margins related to each other: (0) juxtaposed (Fig. 41); (1) overlapping (Fig. 40). Modified from Weiler (2011).
84. Genital plates, gonocoxites 8, posterior margin, form: (0) sinuous (Fig. 41); (1) straight (Fig. 40); (2) convex (Fig. 39).
85. Genital plates, gonocoxites 8, excavations in the inner folding: (0) absent (Fig. 42); (1) present (Fig. 43, 44).
86. Genital plates, gonocoxites 8 , number of excavations in the inner folding: ( 0 ) one (Fig. 43); (1) two (Fig. 44).
87. Genital plates, gonocoxites 8 , development degree over gonocoxite 9: (0) partially covering (Fig. 39); (1) completely covering.
88. Genital plates, gonocoxite 9 , posterior margin, shape: (0) concave; (1) straight.
89. Genital plates, gonocoxite 9 position in relation to segment X : (0) levelled; (1) oblique, in an obtuse angle. Modified from Weiler (2011).
90. Genital plates, gonocoxites 9, surface: (0) flat; (1) swollen; (2) depressed.
91. Genital plates, gonocoxites 9, esclerotized lateral arms: (0) absent; (1) present (Fig. 45).
92. Genital plates, laterotergites 8, posterior margin, form: (0) straight (Fig. 39); (1) acutely projected (Fig. 40); (2) obtusely projected (Fig. 41). Modified from Weiler (2011).
93. Genital plates, laterotergites 8 , spiracle: (0) absent; (1) present.
94. Genital plates, laterotergites 9, base, surface: (0) flat; (1) depressed; (2) swollen.
95. Genital plates, laterotergites 9, apex in relation to tergite 8: (0) not surpassing (Fig. 39); (1) surpassing (Fig. 41). Modified from Fortes \& Grazia (2005); Weiler (2011).
96. Genital plates, gonapophyses 8 , visible in ventral view: (0) absent; (1) present.
97. Genital plates, gonapophyses 9 , sclerotized areas: (0) absent; (1) present.
98. Ectodermal ducts, thickening of vaginal intima, length of arcuate posterior portion related to round anterior portion: (0) shorter (Fig. 46); (1) longer (Fig. 45).
99. Ectodermal ducts, proximal ductus receptaculi, length in relation to vesicular area: (0) shorter; (1) longer. Modified from Hasan \& Kitching (1993); Weiler (2011).
100. Ectodermal ducts, vesicular area, median wall, shape sub-proximally: (0) cylindrical (Fig. 48); (1) enlarged (Fig. 47).
101. Ectodermal ducts, distal ductus receptaculi: (0) straight (Fig. 48); (1) convolute (Fig. 47).
102. Ectodermal ducts, distal ductus receptaculi, before anterior annular flange, form: (0) tubular; (1) dilated. Modified from Gapud (1991); Weiler (2011).
103. Ectodermal ducts, pars intermedialis, form: (0) rectilinear (Fig. 49); (1) twisted (Fig. 50); (2) enlarged (Fig. 51). Modified from Fortes \& Grazia (2005); Campos \& Grazia (2006); Weiler (2011).
104. Ectodermal ducts, posterior annular flange, width related to capsula seminalis width: (0) thinner; (1) wider.
105. Ectodermal ducts, capsula seminalis, form: (0) globose (Fig. 50); (1) ovate (Fig. 51); (2) elongate (Fig. 49). Modified from Hasan \& Kitching (1993); Grazia (1997); Fortes \& Grazia (2005), Campos \& Grazia (2006); Weiler (2011).
106. Ectodermal ducts, capsula seminalis, process(es): (0) absent; (1) present (Fig. 51).
107. Pygophore, dorsal rim, marginal process: (0) absent; (1) present (Fig. 53).
108. Pygophore, dorsal rim, superior process: (0) absent (Fig. 54); (1) present.
109. Pygophore, dorsal rim, superior process, form: (0) flap-like (Fig. 56); (1) spine-like (Fig. 57); (2) tumescence (Fig. 58). Modified from Barcellos \& Grazia (2003); Fortes \& Grazia (2005); Ferrari et al. (2010); Weiler (2011).
110. Pygophore, postero-lateral angle, shape: (0) rounded (Fig. 53); (1) quadrate (Fig. 55); (2) acute (Fig. 52).
111. Pygophore, postero-lateral angles, bristles tufts: (0) absent; (1) present (Fig. 55). Modified from Fortes \& Grazia (2005); Weiler (2011).
112. Pygophore, ventral rim, bristle tufts medially: (0) absent; (1) present (Fig. 55).
113. Pygophore, ventral rim, superior layer projected toward genital cup: (0) absent; (1) present (Fig. 54).
114. Pygophore, longitudinal ridges: (0) absent; (1) present (Fig. 55).
115. Pygophore, superior ridge, form: (0) concave; (1) convex.
116. Pygophore, superior ridge, processes: (0) absent; (1) present (Fig. 53).
117. Pygophore, genital cup, transversal striations: (0) absent; (1) present.
118. Pygophore, paramere: (0) well-developed; (1) reduced (Fig. 54).
119. Pygophore, paramere, length of head related to length of base: (0) shorter; (1) as long as; (2) longer.
120. Pygophore, segment X, processes: (0) absent; (1) present. Modified from Weiler (2011).
121. Pygophore, segment $X$, processes shape: (0) tumescence; (1) carina.
122. Pygophore, superior ridge, projecting over segment X : (0) absent; (1) present.
123. Pygophore, segment X, processes, placement of insertion: (0) basally; (1) medially.
124. Paramere, head, vestiture, setae: (0) absent; (1) present.
125. Paramere, head, vestiture, microtrichia: (0) absent; (1) present.
126. Phallus, phallotheca, length medially related to width apically: (0) shorter; (1) longer.
127. Phallus, phallotheca, ductus seminis proximalis insertion on ejaculatory reservoir, position related to phallotheca: (0) near base of; (1) beyond apical half of.
128. Phallus, phallotheca, ejaculatory reservoir, aspect: (0) smooth (Fig. 60); (1) striate (Fig. $61)$.
129. Phallus, phallotheca, postero-dorsal margin, projection(s), medially: (0) absent; (1) present (Figs. 62, 63).
130. Phallus, phallotheca, postero-dorsal margin, number of projection(s), medially: (0) one (Fig. 63); (1) two (Fig. 62).
131. Phallus, postero-lateral margins, rounded projections: (0) absent; (1) present (Fig. 63).
132. Phallus, phallotheca, thecal shield: (0) absent; (1) present (Fig. 64).
133. Phallus, phallotheca, processes of phallotheca: (0) absent; (1) present (Figs. 59-61).
134. Phallus, phallotheca, processes of phallotheca, width basally related to width medially: (0) narrower; (1) wider; (2) as wide as.
135. Phallus, phallotheca, processes, ventro-basally: (0) absent; (1) present.
136. Phallus, conjunctive, dorsal lobe: (0) absent; (1) present (Fig. 60).
137. Phallus, conjunctive, lateral lobes: (0) absent; (1) present (Fig. 63).
138. Phallus, conjunctive, median lobes: (0) absent; (1) present (Fig. 63).
139. Phallus, conjunctive, median lobes, aspect: (0) entirely membranous (Fig. 63); (1) sclerotized apically (Fig. 64); (2) entirely sclerotized (Fig. 59).
140. Phallus, conjunctive, median lobes, entirely sclerotized, shape apically: (0) rounded; (1) acute.
141. Phallus, conjunctive, ventral lobes: (0) absent; (1) present.
142. Phallus, processes of vesica: (0) absent; (1) present.
143. Phallus, processes of vesica, shape: (0) cup-like (Fig. 65); (1) sac-like (Fig. 66); (2) flap-like (Fig. 67); (3) keel-like; (4) geniculate (Fig. 68); (5) tubular.
144. Phallus, cup-like processes of vesica, form ventrally related to each other: (0) connected; (1) independent.
145. Phallus, vesica, length related to phallothecal lenght: (0) shorter; (1) longer (Fig. 67). Modified from Hasan \& Kitching (1993); Barcellos \& Grazia (2003); Fortes \& Grazia (2005); Campos \& Grazia (2006); Bernardes et al. (2009); Weiler (2011).
146. Phallus, vesica, aspect: (0) coiled (Fig. 67); (1) loose. Modified from Hasan \& Kitching (1993); Barcellos \& Grazia (2003); Fortes \& Grazia (2005); Campos \& Grazia (2006); Bernardes et al. (2009); Weiler (2011).
147. Phallus, vesica, sclerotized cap basally: (0) absent; (1) present (Fig. 67).
148. Phallus, vesical aperture, shape: (0) circular; (1) spatulate; (2) bifid.

## Phylogenetic analysis

Missing data represent about $7 \%$ of the matrix and is largely due to inapplicable (-) characters resulted from reductive coding; only data from male genitalia of Spinalanx monstrabilis and male internal genitalia of $D$. (P.) divisus correspond to missing data (?) due to unavailability of specimens.

Analyses A resulted in six most parsimonious trees (MPT) of length of 878 steps. Only one MPT was found on each of the remaining analyses (Table 1). Consensus of analyses A is highly resolved, with two polytomies: one polytomy is basal and the other within Dichelops (Dichelops).

Topological comparisons by Similitud Index and Robinson-Foulds distances indicate that analyses A-D are more similar to each other than to E (Table S3). The highest tree topologies congruence, according to both tree comparison metrics, was A \& B followed by C
\& D. The remaining ranking of tree comparisons changed according to tree comparison metrics (Table S3).

Relationships among taxa were in broad agreement irrespective of datasets (discrete and discrete + continuous) on analyses A-D, with relatively minor topology differences (Figs 69-72), which are discussed subsequently. Analyses E, of continuous data only, resulted in a topology in great disagreement with remaining analyses (Fig. S1).

Monophyly of Dichelops was not recovered, independent of datasets or weighting method employed (Figs 69-72): subgenera Dichelops and Prodichelops were grouped and subgenus Diceraeus was grouped elsewhere. Sister group relationships to Dichelops subgenera was consistent on analyses A-D (Figs 69-72). Most genera which we have included more than one species were recovered monophyletic, to the exception of Euschistus. Stability of subgenera of Dichelops was high, with jackknife absolute frequencies higher than $50 \%$ in analyses A-D (Figs 69-72).

When constraining monophyly of Dichelops, six MPT of 884 steps were found on analysis Ac. Consensus of analysis Ac was highly resolved (Fig. S2), with one basal polytomy and other polytomy within Dichelops (Dichelops). Only Euschistus was not recovered monophyletic. Overall jackknife absolute and GC frequencies were higher for constrained analysis than unconstrained (Table 1; Figs. S3-6).

## Discussion

## Datasets and analyses

One of the main issues in using continuous characters in phylogenetics is the differential influence of characters depending on the scale in which they are measured. Two strategies have been independently used to circumvent the scaling issue: range rescaling and implied weighting. Mongiardino et al. (2014) demonstrated that even though IW reduces continuous characters influence in determining a topology by 10 times, it does so only
partially and there is still a major influence of characters scale. Therefore, characters are more accurately weighted according to their homoplasy after they have been rescaled (Mongiardino et al., 2014). Accordingly, continuous characters were more influent in our analyses when we did not use IW, even though most of its asymmetric influence was already circumvented by the rescaling procedure. IW decreases influence of homoplastic characters, thus reducing continuous characters influence in analysis (Goloboff et al., 2006).

Continuous features can be the result of heritable variation, but phenotypic variation in continuous characters might be so large as to confound the heritable variation. After rescaling continuous characters and analyzing it under IW, an estimate of how much phylogenetic information is contained on continuous characters can be obtained (Goloboff et al., 2006). According to Goloboff et al. (2006), the extent of the conflict or concordance between continuous and discrete datasets can be examined by the increase or decrease of groups supported by discrete characters alone when continuous characters are added to the dataset. In general in our datasets, irrespective of weighting method employed, the addition of continuous characters have not changed tree topology nor added support to basal nodes, however have increased support of clades close to branch tips (Figs 69-72).

Cladogram topologies resultant of analyses A-D were highly congruent, as demonstrated by both tree comparison metrics (Table S3). The results of analyses A and B are the most similar, followed by results of analyses C and D . Thus, comparing the topologies by those metrics, indicate that results of analyses were first grouped by weighting method disregarded datasets used, i.e. in our data, weighting methods had major influence in results than the different datasets.

Analysing continuous characters along with discrete characters improved clade support, but did not change topology. Thus, continuous characters had not highly influenced
tree search on our datasets. Hence, using EW or IW had a major impact on analyses by weighting differentially homoplasy (Goloboff et al., 2008).

Moreover, because of the disproportional number of discrete characters related to continuous characters, the former play a more important role on tree topology. Characters with a strong influence in determining the optimal tree of the entire partition will, when isolated, support a similar tree (Mongiardino et al., 2014), what is evidenced by the few topological differences between the analyses of discrete dataset only and discrete + continuous datasets, on both weighting methods. Jackknife values were slightly higher when IW was employed, compared to EW analyses (C to A and D to B).

For the matter of discussing phylogenetic relationships and taxonomic decisions, we will follow the reasoning of Kluge (1989) concerning the philosophy of total evidence in phylogenetic analyses and the reasoning of Goloboff et al. (2008) for weighting character against homoplasy. Topology of analysis D (discrete + continuous characters, under IW) is the one that fits such criteria.

Constraining monophyly of Dichelops. As expected, tree length and adjusted homoplasy increased when forcing ingroup monophyly (Table 1). Topologies with constrained monophyly represent a priori hypotheses of relationships and, in the case of our analyses, suboptimal phylogenetic hypotheses. Topologies resultant of EW analyses were six steps shorter than the same analyses under constrained groups.

## Taxa relationships

As argued above, phylogenetic relationships and taxonomic decisions were made based on topology resultant of analysis D. Subsequently, we comment the relationships of some clades, as numbered on Figs. 72-73.

Clade I. This clade was not supported by jackknife resampling; however was consistently found on every analysis. Genera in this clade comprise those attributed to the Euschistusgroup, plus Agroecus and Prionosoma. Features sustaining this clade are the antero-lateral margins of pronotum serrated [33(1)], postero-lateral margins crenulated [36(1)] and membrane of hemelytrum not surpassing abdominal apex [46(1)], but several reversions occurs throughout the clade. A pair of dorsal processes on phallotheca, a characteristic diagnostic for the group suggested by Rider (1995), was a synapomorphy for the clade with reversions in Galedanta and Caonabo. Even though most basal clades were not supported by jackknife, relationships can be considered stable because were consistent in all analyses performed.

Clade II. Dichelops was found as sister group to Proxys + Padaeus, with high support values. The three genera share with each other, truncate posterior margin of bucculae [21(1)], outer margin of evaporatorium concave [51(1)] (but in some species it is straight), antero-lateral margin of evaporatorium acute [52(0)] (but in some species it is evanescent), absence of secondary thickenings of gonapophyses 9 [97(0)], and capsula seminalis oval [105(1)]. Grazia (1978) had already related Dichelops to Padaeus because of the shape of the bucculae.

Clade III. This clade comprises in part what Grazia (1978) considered Dichelops. Grazia (1978) produced extensive work concerning the taxonomy of Dichelops and was considered by some as the first phylogenetic analysis produced in Brazil: a list of eight characters was provided and analysed by congruence, resulting on Diceraeus as sister group to $D$. $($ Dichelops $)+D$. (Prodichelops). As characteristics shared by D. (Dichelops) and D.
(Prodichelops), Grazia (1978) enumerates calluses on posterior margin of pronotal cicatrices, scutellum foveate, laterotergites 8 with spiracles and pars intermedialis cylindrical.

All diagnosing characteristics proposed by Grazia are shared with other genera; an outcome of wide outgroup sampling, allowing better understanding of character evolution as well as taxa. Based on character optimization Dichelops can be diagnosed by a combination of characters: mandibular plates longer than clypeus [10(2)], presence of a black stripe on the outline of mandibular plates [14(1)], posterior margin of gonocoxites 8 concave [88(0)], presence of bristle tufts on postero-lateral angles (except on D. nigrum) [111(1)]. The subgenera Dichelops and Prodichelops can be recognized by a series of characteristics (Fig. 73); we highlight the presence of an unpunctuated transhumeral band [39(1), 40(0)] and postero-lateral angles of sternites [75(1)] and lateral angles of connexivum with black spots [81(1), 82(1)] in Prodichelops and in Dichelops a median-tubercle on the postero-lateral margins of pronotum [37(1)], gonocoxites 8 excavated in internal view [85(1)] and processes on capsula seminalis [106(1)].

Clade IV. This clade comprises other group previously included in Dichelops by Grazia (1978). Diceraeus stat. nov. is supported as a monophyletic genus, independent of Dichelops. Apex of scutellum calloused [41(1)], pars intermedialis twisted [103(1)], and process of vesical opened ventrally [134(1)] are some of the characteristics of Diceraeus, but also shared with other taxa. All diagnostic characters of Diceraeus proposed by Grazia (1978) are shared with other genera. Also, some of those diagnosing features were reinterpreted here: calloused areas on the posterior margin of pronotal cicatrices were found on Diceraeus, as well as very reduced, but present, fovea on the scutellum.

## Proposed classification of Dichelops

Based on the results of Analyses A-D, which consistently recovered Dichelops paraphyly, we propose the new taxonomic arrangements below:

## Dichelops Spinola, 1837

Type species: Dichelops punctatus Spinola, 1837
Dichelops (Dichelops) Spinola, 1837
Dichelops (D.) australis Grazia \& Klein, 2012

Dichelops (D.) avilapiresi Grazia, 1978
Dichelops (D.) bicolor Distant, 1890
Dichelops (D.) leucostigmus (Dallas, 1851)
Dichelops (D.) miriamae Grazia, 1978
Dichelops (D.) nigrum Bergroth, 1914
Dichelops (D.) peruanus Grazia, 1978
Dichelops (D.) pradoi Grazia, 1978
Dichelops (D.) punctatus Spinola, 1837
Dichelops (D.) saltensis Grazia, 1978
Dichelops (Prodichelops) Grazia, 1978
Type species: Diceraeus divisus Walker, 1867
Dichelops (Prodichelops) divisus (Walker, 1867)

## Diceraeus Dallas, 1851 stat. nov.

Type species: Diceraeus melacanthus Dallas, 1851
Diceraeus caatinguensis (Grazia \& Poock-da-Silva, 2013) nov. comb.
Diceraeus furcatus (Fabricius, 1775) nov. comb.
Diceraeus lobatus (Grazia, 1978) nov. comb.
Diceraeus melacanthus Dallas, 1851 stat. nov.

Diceraeus phoenix (Grazia, 1978) nov. comb.

## Conclusions

Rescaling continuous characters, combining datasets, and downweighting homoplasy has great influence on tree topology. Our results showed that the phylogentic component of the continuous characters employed are low, but still enough for adding support to nodes closer to tips. The factors most influencing analyses were firstly, weighting method and, secondarily, datasets according to the different tree comparison metrics.

Dichelops was found paraphyletic in all not-constrained analyses. Results support Dichelops as comprising the subgenera Dichelops and Prodichelops, and the new generic status of Diceraeus. Conclusions about sister group relationships are still clouded by low values of branch support. However, to the exception of Euschistus, every genus with more than one species included in the analyses were recovered monophyletic with high values ( $>50 \%$ ) of jackknife frequencies.

This is the first phylogenetic analysis of a pentatomid genus, under up-to-date cladistics methods, concerned in sampling a broad outgroup within a tribe. As such, can shed light on morphological diversity under phylogenetic analysis and on the necessity of broad taxonomic sampling for future taxonomic and systematic decisions on the group.

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Table 1. Summary of analyses results.

| Analyses | \#MPT | Length | Adjusted <br> Homoplasy | CI | RI | Jackknife |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  | Absolute Frequency | GC |
| A | 6 | 878 | - | 0.194 | 0.624 | 43.9 | 48.4 |
| B | 1 | 917.842 |  | 0.195 | 0.618 | 48.8 | 50 |
| C | 1 | - | 33.73161 | 0.192 | 0.621 | 49.4 | 49.4 |
| D | 1 | - | 33.86172 | 0.194 | 0.615 | 48.6 | 50 |
| E | 1 | - | 0.68842 | 0.455 | 0.786 | 17.3 | 18.3 |
| AC | 4 | 884 | - | 0.192 | 0.621 | 47.4 | 52.3 |
| BC | 1 | 924.603 | - | 0.194 | 0.615 | 52.7 | 54.5 |
| CC | 1 | - | 33.93576 | 0.191 | 0.617 | 55.5 | 53.8 |
| DC | 1 | - | 34.07655 | 0.193 | 0.612 | 54.4 | 54.6 |
| EC | 1 | - | 0.78547 | 0.42 | 0.753 | 20.4 | 20.1 |

\#MPT, number of most parsimonious trees; Length, number of steps of tree; CI, Consistency Index; RI, Retention Index.


Fig. 1. Scheme of 10 analyses performed with different matrices and weighting algorithms. Three matrices (discrete data only, continuous data only and discrete + continuous data) were analyzed under the same searching parameters, however, with varying weighting criteria (EW, equal weights, and IW, implied weights), corresponding to analyses A-E. The consensus of the resultant most parsimonious trees were calculated for each analysis, as well as stability of nodes by Jackniffe with symmetric sampling recording absolute-group and GC frequencies. The same set of analyses were performed forcing Dichelops monophyly (analyses Ac-Ec). The resultant 10 consensus were then compared by Coefficient Distances and Robinson-Foulds distances.


Figs. 2-14. Illustration of thoracic characters. (2-5) head in dorsal view; (2) Dichelops (Dichelops) leucostigmus; (3) Acledra albocostata; (4) Euschistus (Lycipta) sharpi; (5) Tibraca limbativentris; (6-9) pronotum in dorsal view; (6) Dichelops (Dichelops) leucostigmus; (7) Menecles insertus; (8) Acledra albocostata; (9) Eushistus (Mitripus) grandis; (10-11) head and prothorax in lateral view; (10) Hypatropis sternalis; (11) Dichelops (Dichelops) leucostigmus; (12-14) apex of radial vein on hemelytra; (12) Carpocoris purpureipennis; (13) Dichelops (Dichelops) nigrum; (14) Dichelops (Diceraeus) melacanthus. Scale bars: 2-11 = $1 \mathrm{~mm} ; 12-14=10 \mu \mathrm{~m}$


Figs. 15-31. Illustration of characters of external scent efferent system of mestasternal glands. (15-18) meso and meta pleura in ventral view; (15) Glyphepomis setigera; (16) Dichelops (Dichelops) pradoi; (17) Coenus delia; (18) Dolycoris baccarum; (19-21) peritreme; (19) Carpocoris purpureipennis; (20) Proxys victor; (21) Glyphepomis setigera; (22) evaporatorium surface, Galedanta bituberculata; (23-26) peritremal surface; (23) Caribo fasciatus; (24) Acledra fraterna; (25) Agroecus scabricornis; (26) Spinalanx rolstoni; (27-28) metathoracic spiracle; (27) Hypatropis inermis; (28) Dolycoris baccarum; (29-31) filtering system of metathoracic spiracle; (29) Carpocoris purpureipennis; (30) Dichelops (Dichelops) peruanus; (31) Proxys victor. Scale bars: 15-18, 27-28 $=500 \mu \mathrm{~m} ; 19-21=100 \mu \mathrm{~m} ; 22-26=$ $5 \mu \mathrm{~m} ; 29-31=10 \mu \mathrm{~m}$.


Figs. 32-38. Illustration of abdominal characters. (32-33) abdomen in ventral view; (32) Padaeus trivittatus; (33) Dichelops (Diceraeus) melacanthus; (34-35) lateral margin of $4^{\text {th }}$ and $5^{\text {th }}$ abdominal sternites; (34) Dichelops (Dichelops) bicolor; (35) Dichelops (Prodichelops) divisus; (36-38) lateral margin of $5-7^{\text {th }}$ abdominal segments in dorsal view; (36) Agroecus scabricornis; (37) Dichelops (Dichelops) punctatus; (38) Euschistus (Euschistus) variolarius. Scale bars: 1 mm .


Figs. 39-51. Illustration of female genital characters. (39-41) external genitalia; (39) Caribo fasciatus; (40) Dichelops (Diceraeus) melacanthus; (41) Hymenarcis nervosa; (42-44) gonocoxite 8; (42) Dichelops (Diceraeus) furcatus; (43) Dichelops (Dichelops) peruanus; (44) Dichelops (Dichelops) bicolor; (45-46) dissected genital plates; (45) Dichelops (Diceraeus) furcatus; (46) Dolycoris baccarum; (47-51) spermatheca; (47) Hymenarcis nervosa; (48) Dichelops (Diceraeus) melacanthus; (49) Cosmopepla decorata; (50) Dichelops (Diceraeus) furcatus; (51) Dichelops (Dichelops) peruanus. Scale bars: 39-41 = 1 $\mathrm{mm} ; 42-48=0.3 \mathrm{~mm} ; 49-51=0.1 \mathrm{~mm}$.


Figs. 52-68. Illustration of male genital characters. (52-55) pygophore in dorsal view; (52) Dolycoris baccarum; (53) Mormidea maculata; (54) Tibraca limbativentris; (55) Dichelops (Dichelops) avilapiresi; (56-58) pygophore in latero-dorsal view; (56) Dichelops (Diceraeus) melacanthus; (57) Dichelops (Dichelops) avilapiresi; (58) Euschistus (Mitripus) grandis; (59) apex of phallus in lateral view, Dichelops (Dichelops) leucostigmus; (60-63) phallus in dorsal view; (60) Dichelops (Diceraeus) caatinguensis; (61) Dichelops (Dichelops) saltensis; (62) Tibraca limbativentris; (63) Berecynthus hastator; (64-68) phallus in lateral view; (64) Cosmopepla decorata; (65) Dichelops (Diceraeus) melacanthus; (66) Tibraca limbativentris; (67) Hymenarcis nervosa; (68) Berecynthus hastator. Scales: 39-45 $=0.5 \mathrm{~mm} ; 47-50=0.3$ $\mathrm{mm} ; 46,51-55=0.2 \mathrm{~mm}$.


Fig. 69. Strict consensus of six equally parsimonious cladograms resultant of Analysis A (discrete characters only, equal weights). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. S: 878; CI: 0.19; RI: 0.624.


Fig. 70. Cladogram resultant of Analysis B (discrete + continuous characters, equal weights).
Values above and below branches represent jackknife absolute frequencies and GC values, respectively. S: 917.842; CI: 0.195; RI: 0.618 .


Fig. 71. Cladogram resultant of Analysis C (discrete characters only, implied weights).
Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 33.73161; CI: 0.192; RI: 0.621.


Fig. 72. Cladogram resultant of Analysis D (discrete + continuous characters, implied weights). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 33.86172 ; CI: $0.194 ;$ RI: 0.615 .


Figure 73a. Optimization of discrete characters on tree resultant of analysis D.


Figure 73b. (Continuation) Optimization of discrete characters on tree resultant of analysis D.

## CONCLUSÕES

No Capítulo I escrevemos uma resposta ao trabalho de Memon et al. (2010), que versava sobre uma análise filogenética de Halyini ocorrentes no sub-continente IndoPaquistanês. Concluimos que análises filogenéticas em Pentatomidae deveriam, preferencialmente, incluir grupos externos mais representativos, a fim de melhor testar as hipóteses de monofilia, e utilizar procedimentos metodológicos claros em todas as etapas das análises filogenéticas.

No Capítulo II, estudamos comparativamente a morfologia de algumas estruturas da tíbia das pernas anteriores de Pentatomidae, tendo sido amostradas 81 espécies ( 55 gêneros) representantes de sete subfamílias. A titulo de comparação estudamos Scutelleridae e Thyreocoridae, representadas por três e 10 espécies, respectivamente. Concluimos que a estrutura pré-tarsal em Pentatomidae é pouco variável e que não permite a identificação de grupos taxonômicos, à exceção de Phyllocephalinae que apresenta pulvilos distintos. Por outro lado, algumas características pré-tarsais permitem diferenciar o pré-tarso de Thyrecoridae do pré-tarso das outras duas famílias: em Thyreocoridae foi observado um vinco na face interna das garras, característica não observada nas demais famílias estudadas. As outras estruturas estudadas foram o aparato tibial e o pente tibial; ambas estruturas são formadas por projeções cuticulares, a primeira por cerdas espessas organizadas lado a lado e a segunda por projeções espatulares do tegumento organizadas em linha. Não observamos diferenças no aparato tibial tanto entre quanto dentre famílias. Por outro lado, o número de cerdas que compõem o aparato tibial varia consideravelmente dentro de Pentatomidae, possuindo um padrão por subfamílias, e entre subfamílias. Sugerimos que o número de cerdas do aparato tibial e o número de projeções espatulares do pente tibial sejam usadas e exploradas como caracteres contínuos em análises filogenéticas em níves supra-genéricos.

No Capítulo III, descrevemos uma nova espécie de Dichelops (Diceraeus) Dallas, 1851 e o macho de Dichelops (Prodichelops) divisus (Walker, 1867). A descrição da genitália masculina de $D$. (P.) divisus permitiu reconhecer que este é mais similar à genitália de Dichelops (Dichelops) Spinola, 1837, pela presença de processo superior do bordo dorsal triangular, parâmeros falciformes e ângulos póstero-laterais com tufos de cerdas. Dichelops (D.) caatinguensis Grazia \& Poock-da-Silva, 2013 foi descrita com base em espécimes provenientes da Caatinga. A descrição dessa espécie ampliou a distribuição do gênero para essa eco-região e também ampliou a diversidade morfológica do subgênero: D. caatinguensis é única no subgênero por possuir calo arrendondado no ápice do escutelo e manchas escuras nas margens anteriores e posteriores do conexivo, nos machos, processo ventral da conjuntiva no pênis e, nas fêmeas, gonocoxitos 9 côncavos posteriormente. A descrição desta espécie para uma região onde até então o gênero não ocorria traz a atenção para a importância de coletas e do estudo da fauna de regiões pouco exploradas, como a Caatinga.

No Capítulo IV, investigamos as relações filogenéticas de Dichelops. O gênero foi sempre recuperado parafilético. Os resultados suportam Dichelops como formado pelos subgêneros Dichelops e Prodichelops e o novo status genérico de Diceraeus. Conclusões sobre as relações de parentesco ainda são obscurecidas por baixo valores de suporte para os clados. Entretanto, excetuando-se Euschistus, a monofilia de todos os gêneros amostrados por mais de uma espécie foi recuperada com altos valores de suporte por frequências de Jackknife (>50\%). Também, ao explorar as diferentes análises, concluiu-se que reescalonar caracteres contínuos, combinar os conjuntos de dados contínuos e discretos e pesar os caracteres contra homoplasia tiveram grande influência nas análises. Os resultados demonstraram que o componente filogenético nos caracteres contínuos é pequeno, mas que ainda assim é suficiente para adicionar suporte aos nós próximos aos terminais. Conforme as métricas de comparações de árvores, os fatores que mais influenciaram as análises foram, primeiramente,
os métodos de pesagem de caracteres e, secundariamente, o conjunto de dados (somente caracteres discretos, somente caracteres contínuos, ou matrizes combinadas).

## APÊNDICE I

## Material suplementar ao Capítulo II

## Comparative morphology of selected characters in the Pentatomidae foreleg (Hemiptera: Heteroptera)

Table S1. List of examined species. The Pentatomidae clasification follows Grazia et al. (2008) and Rider (2012) for subfamilies and tribes, respectively.

| Family | Subfamily | Tribe | Species |
| :---: | :---: | :---: | :---: |
| Pentatomidae | Aphylinae |  | Aphylinae sp. |
|  | Asopinae |  | Alcaeorrhynchus grandis (Dallas, 1851) |
|  |  |  | Oplomus cruentus (Burmeister, 1835) |
|  |  |  | Oplomus salamandra (Burmeister, 1835) |
|  |  |  | Podisus connexivus Bergroth, 1891 |
|  | Cyrtocorinae |  | Cyrtocoris egeris Packauskas \& Schaefer, 1998 |
|  |  |  | Cyrtocoris trigonus (Germar, 1839) |
|  | Discocephalinae | Discocephalini | Antiteuchus mixtus (Fabricius, 1787) |
|  |  |  | Eurystethus microlobatus Ruckes, 1966 |
|  |  | Ochlerini | Alitocoris parvus (Distant, 1880) |
|  |  |  | Catulona pensa Rolston, 1992 |
|  |  |  | Lincus securiger Breddin, 1904 |
|  | Edessinae |  | Brachystethus geniculata (Fabricius, 1787) |
|  |  |  | Brachystethus vexillum Breddin, 1903 |
|  |  |  | Edessa quadridens Fabricius, 1803 |
|  |  |  | Edessa rufomarginata (De Geer, 1773) |
|  |  |  | Lopadusa (Lopadusa) augur Stål, 1860 |
|  |  |  | Olbia elegans (Stål, 1862) |
|  | Pentatominae | Antestiini | Antestia ellenriederi Breddin, 1900 |
|  |  |  | Antestiopsis anchora (Thunberg, 1783) |
|  |  |  | Plautia stali Scott, 1874 |
|  |  |  | Starioides degenera (Walker, 1867) |
|  |  | Carpocorini | Acledra albocostata (Spinola, 1852) |
|  |  |  | Agonoscelis nubilis (Fabricius, 1775) |
|  |  |  | Agroecus griseus Dallas, 1851 |
|  |  |  | Berecynthus hastator (Fabricius, 1794) |
|  |  |  | Coenus delius (Say, 1831) |
|  |  |  | Cosmopepla decorata (Hahn, 1834) |
|  |  |  | Dichelops (Diceraeus) furcatus (Fabricius, 1775) |
|  |  |  | Dichelops (Diceraeus) melacanthus (Dallas, 1851) |
|  |  |  | Dichelops (Dichelops) punctatus Spinola, 1837 |
|  |  |  | Dichelops (Dichelops) saltensins Grazia, 1978 |

Dichelops (Prodichelops) divisus (Walker, 1867)
Diploxys sp.
Dolycoris baccarum (Linnaeus, 1758)
Hypatropis inermis (Stål, 1872)
Mormidea quinqueluteum (Lichtenstein, 1796)
Mormidea ypsilon (Linnaeus, 1758)
Oebalus poecilus (Dallas, 1851)
Tibraca limbativentris Stål,1860
Catacanthini Arocera apta (Walker, 1867)
Arocera contalineata Pirán, 1955
Rhyssocephala infuscata (Rider, 1992)
Rhyssocephala rufolimbata (Stål, 1872)
Runibia decorata (Dallas, 1851)
Runibia perspicua (Fabricius, 1798)
Chlorocorini Arvelius albopunctatus (De Geer, 1773)
Chlorocoris (Chlorocoris) complanatus Guérin-Méneville, 1831
Chlorocoris (Chlorocoris) distinctus Signoret, 1851
Chlorocoris (Monochricerus) flaviviridis Barber, 1914
Chlorocoris (Monochricerus) subrugosus Stål, 1872
Chloropepla rolstoni Grazia-Vieira, 1973
Chloropepla vigens (Stål, 1860)
Loxa deducta Walker, 1867
Loxa flavicollis (Drury, 1773)
Rhyncholepta grandicallosa Bergroth, 1911
Rhyncholepta meinanderi Becker \& Grazia-Vieira, 1971
Pentatomini Banasa dolabrata Thomas, 1988
Banasa patagiata (Berg 1879)
Banasa zeteki Sailer, 1959
Janeirona stali (Kormilev, 1956)
Lelia octopunctata (Dallas, 1849)
Myota aerea (Herrich-Schäffer, 1842)
Neotibilis (Laeviscutis) compascens
Neotibilis (Neotibilis) parva (Distant, 1893)
Pallantia macula (Dallas, 1851)
Pallantia macunaima Grazia, 1980
Pellaea stictica (Dallas, 1851)
Pentatoma japonica (Distant, 1882)
Pentatoma metallifera (Motschulsky, 1859)
Pentatoma rufipes (Linnaeus, 1758)
Phalaecus pustulatus de Geer, 1773
Taurocerus achilles (Stål, 1862)
Procleticini Dendrocoris arizonensis Barber, 1911
Dendrocoris pini Montandon, 1893
Lobepomis peltifera Berg, 1891
Thoreyella cornuta Berg, 1883
Thoreyella trinotata Berg, 1878

|  | Rhynchocorini Biprorulus bibax Breddin, 1900 |  |
| :---: | :---: | :---: |
|  |  | Cuspicona simplex Walker, 1867 |
|  | Phyllocephalinae | Macrina juvenca (Burmeister, 1835) |
| Scutelleridae |  | Augocoris gomesi Burmeister, 1835 |
|  |  | Orsilochides leucoptera (Germar, 1839) |
|  |  | Pachycoris torridus Scopoli, 1772 |
| Thyreocoridae | Corimelaeninae | Corimelaena (Corimelaena) tibialis (Fabricius, 1803) |
|  |  | Corimelaena (Parapora) extensa Uhler, 1867 |
|  |  | Cydnoides (Cydnoides) ciliatus Uhler, 1863 |
|  |  | Cydnoides (Cydnoides) renormatus (Uhler, 1895) |
|  |  | Galgupha (Euryscytus) difficilis (Breddin, 1914) |
|  |  | Galgupha (Galgupha) atra Amyot\&Serville, 1843 |
|  |  | Galgupha (Gyrocnemis) cruralis (Stål, 1862) |
|  |  | Galgupha (Microcompsus) vinculata (Germar, 1839) |
|  |  | Galgupha (Nothocoris) terminalis (Walker, 1867) |
|  | Thyreocorinae | Thyreocoris scarabaeoides (Linnaeus, 1758) |

Table S2. List of examined specimens.

| Classification |  |  |  |  |  | Sex | Data collection |  |  |  |  | Collection |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Family | Subfamily | Tribe | Genus | Subgenus | Species |  | Country | State/Province | Locality | Data | Colector |  |
| Pentatomidae | Aphylinae |  |  |  |  |  |  |  |  | V. 1967 | R. Edwards | S. A. Museum |
| Pentatomidae | Asopinae |  | Alcaeorrynchus |  | grandis | F | Venezuela | Zulia | Snza. Rosa Dzo. Colon | 19.II. 1961 | N. Ange \& R. Veja | MIZA |
| Pentatomidae | Asopinae |  | Alcaeorrynchus |  | grandis | M | Venezuela | Aragua | Ramizito | 1.X. 1945 | FFY \& FDR | MIZA |
| Pentatomidae | Asopinae |  | Oplomus |  | cruentus | F | Brazil | Rio Grande do Sul | Estação Ecológia do Taim | 18.III. 1982 | J. Grazia | UFRG |
| Pentatomidae | Asopinae |  | Oplomus |  | cruentus | M | Brazil | Rio Grande do Sul | Rio Grande | 04.II. 1981 | J. Grazia | UFRG |
| Pentatomidae | Asopinae |  | Oplomus |  | cruentus | M | Brazil | Rio Grande do Sul | Rio Grande | 05.II. 1981 | J. Grazia | UFRG |
| Pentatomidae | Asopinae |  | Oplomus |  | salamandra | F | Venezuela | Aragua | Rancho Grande | 3.VIII. 1953 | Ferd. Kern | MIZA |
| Pentatomidae | Asopinae |  | Oplomus |  | salamandra | M | Brazil | Pará | Bujaru | 24.III. 1978 | M. Carvalho | UFRG |
| Pentatomidae | Asopinae |  | Oplomus |  | salamandra | M | Brazil | Pará | Serra Norte | 18.X. 1984 | T. Pimentel | UFRG |
| Pentatomidae | Asopinae |  | Podisus |  | connexivus | M | Brazil | Rio Grande do Sul | Derrubadas | 16.IV. 1983 | S. Bonatto | UFRG |
| Pentatomidae | Asopinae |  | Podisus |  | connexivus | M | Brazil | Rio Grande do Sul | Derrubadas | 22.XII. 1982 | S. Bonatto | UFRG |
| Pentatomidae | Asopinae |  | Podisus |  | connexivus | M | Brazil | Rio Grande do Sul | Derrubadas | 22.XII. 1982 | S. Bonatto | UFRG |
| Pentatomidae | Asopinae |  | Podisus |  | mellipes | F | Venezuela | Zulia | Kasmera | 19.IX. 1961 | C.J. RosaUFRG \& F. Fernandez | MIZA |
| Pentatomidae | Asopinae |  | Podisus |  | mellipes | M | Brazil | Rio Grande do Sul | Porto Alegre | 19.V. 1954 |  | MGAP |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | egeris |  | Brazil | Rio Grande do Sul | Maquiné, Garapiá | 21.XII. 2006 | L.M. Weiler \& R.C. Bins-Neto | UFRG |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | egeris |  | Brazil | Rio Grande do Sul | Maquiné, Trilha 4 | 02.IV. 2006 | J.L.C. Bernardes | UFRG |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | egeris |  | Brazil | Rio Grande do Sul | Maquiné, Trilha 4 | 02.IV. 2006 | J.L.C. Bernardes | UFRG |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | trigonus | M | Brazil | Espírito Santo | Linhares | 8.VII. 1995 | A.J. Cardoso | UFRG |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | trigonus | M | Brazil | Espírito Santo | Linhares | 8.VII. 1995 | A.J. Cardoso | UFRG |
| Pentatomidae | Cyrtocorinae |  | Cyrtocoris |  | trigonus | M | Brazil | Espírito Santo | Linhares | 8.VII. 1995 | A.J. Cardoso | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Antiteuchus |  | mixtus | F | Brazil | Pará | Belém, EMBRAPA-CPATU | 15.I. 1997 | J.A.M. Fernandes | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Antiteuchus |  | mixtus | F | Brazil | São Paulo | Campinas, Taquaral | 18.V. 1977 | I. Sagima | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Antiteuchus |  | mixtus | M | Brazil | Espírito Santo | São Mateus | 5.VI. 1992 | A.D. Araújo | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Antiteuchus |  | mixtus | M | Brazil | São Paulo | Campinas | 26.III. 1976 |  | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Eurystethus |  | microlobates | F | Brazil | Minas Gerais | Santana do Riacho | X. 2007 | T.J.A. Guerra | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Eurystethus |  | microlobates | F | Brazil | Minas Gerais | Santana do Riacho | X. 2007 | T.J.A. Guerra | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Eurystethus |  | microlobates | M | Brazil | Minas Gerais | Santana do Riacho | X. 2007 | T.J.A. Guerra | UFRG |
| Pentatomidae | Discocephalinae | Discocephalini | Eurystethus |  | microlobates | M | Brazil | Minas Gerais | Santana do Riacho | X. 2007 | T.J.A. Guerra | UFRG |


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\text { M } & \text { United States } \\
\text { M } & \text { United States }
\end{array}
$$

| Arizona | Huachaca | VII． 1905 |  |
| :---: | :---: | :---: | :---: |
| Arizona | Huachaca | 23．VII． 1932 | D．K．Duncan |
| Arizona | Huachaca | 31．X． 1937 |  |
|  |  | 12－14．XII． 2009 | L．Pöllumaa \＆O．Maasikas |
|  |  | 15．II． 2010 | J．E．Eger |
| Rio Grande do Sul | Palmares do Sul | 14．XI． 2003 | Equipe Probio |
| Rio Grande do Sul | Palmares do Sul | 14．XI． 2003 | Equipe Probio |
| Rio Grande do Sul | Porto Alegre | 26．XII． 1984 | J．Grazia |
| Rio Grande do Norte | Natal | 11．VII． 1981 | J．Grazia |
| Pará | Tucuru | I． 1979 | M．Alvarenga |
| Pará | Tucuru | I． 1979 | M．Alvarenga |
| Amazonas | Manaus | 18－21．II． 2004 | J．A．Rafael |
| Amazonas | Manaus | 19－22．III． 2004 | J．A．Rafael |
| Amazonas | Manaus | 10－13．XII． 2004 | C．S．Motta |
| Amazonas | Manaus | 21－24．I． 2004 | C．S．Motta |
| El Oro | Zaruma | 10．IV． 1965 | L．E．Pena |
| Zamora | Santiago | 27．III． 1965 | L．E．Pena |
| Bahia | Encruzilhada | XI． 1972 | M．Alverenga |
| Bahia | Encruzilhada | XI． 1973 | M．Alverenga |
| Alta Verapaz | Pancajché | 14．VII． 1947 | C．\＆P．Vaurie |
| Quintana Roo | Xcan Nuevo | 6－7．VI． 1959 | P．\＆C．Vaurie |
| São Paulo | Salesópolis | 25．III． 1961 | Reichardt |
| São Paulo | Salesópolis | 25．III． 1961 | Reichardt |
| Rio de Janeiro | Rio de Janeiro | 1940 | M．Alvarenga |
| São Paulo | São Paulo | 10．XII． 1993 | G．Ckuk |
| Pará | Serra Nova | 22．I． 1985 |  |
|  | Pipeline Road | 12．VII． 1976 |  |
|  | Pipeline Road | 17．VII． 1976 |  |
| Rio de Janeiro | Alto da Boa Vista | 27．X． 1989 | R．Kernshak |












| Parque Estadual do Nonoai | III. 1993 | J.A.M. Fernandes | UFRG |
| :---: | :---: | :---: | :---: |
| Londrina | II. 1995 | L.M. Vivan | UFRG |
| Londrina | II. 1996 | L.M. Vivan | UFRG |
| Limatambo | 10.III. 1967 | F. Carrasco | UFRG |
| Maracay | 2.X. 1982 | F. Fernandez | MIZA |
| Mount Tasiro | 20.VIII. 1971 | Y. Kurosanja | NMNH |
| Dunhua | 28.VI-2VII. 1982 | P. Schaefer | NMNH |
| Radde, Amur |  | H.L. Parker | NMNH |
| Chvojnice |  | J.L. Stehlik | AMNH |
|  |  |  | AMNH |
| Goiânia | 26.III. 1985 | M.J. Ferreira | UFRG |
|  | X. 1943 | A. Parko | UFRG |
| Itatiaia | 21.XI. 1948 | Wygodzinsky | MNRJ |
| Itatiaia | II. 1943 | Wygodzinsky | MNRJ |
| Lemon Cave | III. 1934 | Univ Berkeley Calif Insect Survey | UFRG |
| Lemon Cave | III. 1934 | Univ Berkeley Calif Insect Survey | UFRG |
| Lemon Cave | III. 1934 | Univ Berkeley Calif Insect Survey | UFRG |
| Clayton | 27.V. 1961 | Univ Berkeley Calif Insect Survey | R. L. Langston |
| Monterey | 9.V. 1959 | Univ Berkeley Calif Insect Survey | NZ |
| Rodeo | 25.I. 1958 | R. Golbach | IFML |
| Lago Muyoj | 29.III. 1957 | R. Golbach | IFML |
| Porto Alegre, Morro do Coco | 19.II. 1962 |  | UFRG |
|  |  |  | MACN |
|  | 15.X. 1896 | S. Venturi | MACN |
|  | 19.X. 1896 | S. Venturi | MACN |
| Buta |  | F.R. Wulsin | AMNH |
| Coari, Rio Urucu | 27.X. 2008 | J.A.M. Fernandes e equipe col. | MCNZ |
| Coari, Rio Urucu | 28.X. 2008 | J.A.M. Fernandes e equipe col. | MCNZ |
| Coari, Rio Urucu | 30.X. 2008 | J.A.M. Fernandes e equipe col. | MCNZ |
| Criciuma | 2008 | F.M. Bianchi | MCNZ |
| Criciuma | 2008 | F.M. Bianchi | MCNZ |
| João Pessoa |  |  | UFRG |



| macula | F | Brazil |
| :---: | :---: | :---: |
| macunaima | M | Brazil |
| macunaima | M | Brazil |
| stictica | F | Peru |
| stictica | F | Venezuela |
| japonica | F |  |
| metallifera | F | China |
| metallifera | M | China |
| rufipes | F |  |
| rufipes | F |  |
| pustulatus | F | Brazil |
| achilles | F | Brazil |
| achilles | M | Brazil |
| achilles | M | Brazil |
| arizonensis | F | United States |
| arizonensis | M | United States |
| arizonensis | M | United States |
| pini | F | United States |
| pini | M | United States |
| peltifera | F | Argentina |
| peltifera | M | Argentina |
| cornuta | F | Brazil |
| cornuta | M | Argentina |
| trinotata | F | Argentina |
| trinotata | M | Argentina |
| juvencus | M | Congo |
| gomesii |  | Brazil |
| gomesii |  | Brazil |
| gomesii |  | Brazil |
| leucoptera |  | Brazil |
| leucoptera |  | Brazil |
| torridus |  | Brazil |


| Pentatomidae | Pentatominae | Pentatomini | Pallantia |
| :---: | :---: | :---: | :---: |
| Pentatomidae | Pentatominae | Pentatomini | Pallantia |
| Pentatomidae | Pentatominae | Pentatomini | Pallantia |
| Pentatomidae | Pentatominae | Pentatomini | Pellaea |
| Pentatomidae | Pentatominae | Pentatomini | Pellaea |
| Pentatomidae | Pentatominae | Pentatomini | Pentatoma |
| Pentatomidae | Pentatominae | Pentatomini | Pentatoma |
| Pentatomidae | Pentatominae | Pentatomini | Pentatoma |
| Pentatomidae | Pentatominae | Pentatomini | Pentatoma |
| Pentatomidae | Pentatominae | Pentatomini | Pentatoma |
| Pentatomidae | Pentatominae | Pentatomini | Phalaeucus |
| Pentatomidae | Pentatominae | Pentatomini | Taurocerus |
| Pentatomidae | Pentatominae | Pentatomini | Taurocerus |
| Pentatomidae | Pentatominae | Pentatomini | Taurocerus |
| Pentatomidae | Pentatominae | Procleticini | Dendrocoris |
| Pentatomidae | Pentatominae | Procleticini | Dendrocoris |
| Pentatomidae | Pentatominae | Procleticini | Dendrocoris |
| Pentatomidae | Pentatominae | Procleticini | Dendrocoris |
| Pentatomidae | Pentatominae | Procleticini | Dendrocoris |
| Pentatomidae | Pentatominae | Procleticini | Lobepomis |
| Pentatomidae | Pentatominae | Procleticini | Lobepomis |
| Pentatomidae | Pentatominae | Procleticini | Thoreyella |
| Pentatomidae | Pentatominae | Procleticini | Thoreyella |
| Pentatomidae | Pentatominae | Procleticini | Thoreyella |
| Pentatomidae | Pentatominae | Procleticini | Thoreyella |
| Pentatomidae | Phyllocephalinae | Macrina |  |
| Scutelleridae |  |  | Augocoris |
| Scutelleridae |  |  | Augocoris |
| Scutelleridae |  |  | Augocoris |
| Scutelleridae |  |  | Orsilochides |
| Scutelleridae |  |  | Orsilochides |
| Scutelleridae |  |  | Pachycoris |



| Pachycoris |  | torridus |  | Brazil | Paraíba | João Pessoa |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Pachycoris |  | torridus |  | Brazil | Paraía | João Pessoa |  |  |
| Corimelaena | Corimelaena | tibialis | F | Brazil | Rio Grande do Sul | Palmares do Sul | 20.V. 2004 | Eq. Probio |
| Corimelaena | Corimelaena | tibialis | F | Trinidad and Tobago | Saint George | Valencia | 29.VIII. 1981 | R.M. Baranowski |
| Corimelaena | Corimelaena | tibialis | M | Brazil | Rio Grande do Sul | Palmares do Sul | 20.V. 2004 | Eq. Probio |
| Corimelaena | Corimelaena | tibialis | M | Trinidad and Tobago | Saint George | Valencia | 29.VIII. 1981 | R.M. Baranowski |
| Corimelaena | Parapora | extensa | F | United States | Idaho | Idaho | 5.VIII. 1990 | G.M. Stephens |
| Corimelaena | Parapora | extensa | M | United States | Idaho | Idaho | 15.VII. 1990 | G.M. Stephens |
| Cydnoides | Cydnoides | ciliatus | F | United States | Florida | Miami | 18.XII. 2005 | T. Smith |
| Cydnoides | Cydnoides | ciliatus | F | United States | Wyoming | Sweetwater Co. | 13.VI. 2008 | P. \& P. Skelley, R. Gordon |
| Cydnoides | Cydnoides | ciliatus | M | United States | Arizona | Navajo Co. | 7.IX. 2003 | C.W. \& L. O'Brien |
| Cydnoides | Cydnoides | ciliatus | M | United States | Florida | Miami | 18.XII. 2005 | T. Smith |
| Cydnoides | Cydnoides | renormatus | F | United States | Arizona |  | 2.IX. 2005 | C.W. \& L.O'Brien |
| Cydnoides | Cydnoides | renormatus | M | United States | Arizona |  | 30.VII. 2000 | W.B. Warner |
| Galgupha | Euryscytus | difficilis | F | Brazil | Rio Grande do Sul | Maquiné | 1.IV. 2006 | J.L.C. Bernardes |
| Galgupha | Euryscytus | difficilis | F | Brazil | Rio Grande do Sul | Maquiné, Garapiá | 10.I. 2007 | C.F. Schwertner |
| Galgupha | Euryscytus | difficilis | M | Brazil | Rio Grande do Sul | Esmeralda | 18-19.III. 1981 | J. Grazia |
| Galgupha | Euryscytus | difficilis | M | Brazil | Rio Grande do Sul | Maquiné | 1.IV. 2006 | V.C. Matesco |
| Galgupha | Galgupha | atra | F | United States | Florida | Osceola Co. Deer Park | 27.IV. 1989 | J.E. Eger |
| Galgupha | Galgupha | atra | F | United States | Florida | Osceola Co. Deer Park | 27.IV. 1989 | J.E. Eger |
| Galgupha | Galgupha | atra | M | United States | Florida | Osceola Co. Deer Park | 27.IV. 1989 | J.E. Eger |
| Galgupha | Gyrocnemis | cruralis | F |  |  |  |  |  |
| Galgupha | Gyrocnemis | cruralis | F |  |  |  |  |  |
| Galgupha | Gyrocnemis | cruralis | M | Brazil | Paraná | Bandeirantes | III. 2008 | J. Raudo |
| Galgupha | Gyrocnemis | cruralis | M | Brazil | Paraná | Bandeirantes | III. 2008 | J. Raudo |
| Galgupha | Microcompsus | vinculata | F | Brazil | Amazonas | Coari, Rio Urucu | 26.X. 2008 | J.A.M. Fernandes |
| Galgupha | Microcompsus | vinculata | F | Brazil | Amazonas | Coari, Rio Urucu | 31.X. 2008 | J.A.M. Fernandes |
| Galgupha | Microcompsus | vinculata | M | Brazil | Amazonas | Coari, Rio Urucu | 26.X. 2008 | J.A.M. Fernandes |
| Galgupha | Microcompsus | vinculata | M |  |  |  |  |  |
| Galgupha | Nothocoris | terminalis | F | Brazil | Rio Grande do Sul | Porto Alegre, Morro Santana | 09.XII. 2010 |  |
| Galgupha | Nothocoris | terminalis | F | Brazil | Rio Grande do Sul | Porto Alegre, Morro Santana |  | UFRG |
| Galgupha | Nothocoris | terminalis | M | Brazil | Rio Grande do Sul | Maquiné, Garapiá | 21.XII. 2006 | R.C. Bins-Neto \& L.M. Weiler |


| oridae | Galgupha | Nothocoris | terminalis | M | Brazil | Rio Grande do Sul | Porto Alegre, Morro Santana | UFRG |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| oridae | Thyreocoris |  | scarabaeoides | F | Suécia |  |  |  | AMNH |
| oridae | Thyreocoris |  | scarabaeoides | F | Suécia |  |  |  | AMNH |
| oridae | Thyreocoris |  | scarabaeoides | M | Suécia |  |  |  | AMNH |
| oridae | Thyreocoris |  | scarabaeoides | M |  |  | Box Hill 2.X. 1949 | D. UFRGton | AMNH |
| F, Fe <br> DZU <br> Brazi <br> Brazi <br> Unite <br> Zoob <br> Muse <br> Unive | m Acrony gia Pe. Jes Collection ación Mig eo Argent do Sul, Br de Federal Grande | ': AMN <br> Santiag <br> of Arthro <br> Lillo, <br> o de Cie <br> zil; MGA <br> do Rio de <br> Sul, Bra | American <br> Moure, Un ds, United umán, Arg iasNaturale Museu An neiro, Bra NMNH, |  | um o idade s; HD ; INB ernard de C ZZSP, nal M | History, Unite do Paraná, Cur ge Engleman uto Nacional d davia", Buenos Naturais, Brazil Zoologia da Natural Histo | States; DARC, David Rider ba, Brazil; FIOC, Fundação ollection, Canal Zone, Panam Biodiversidad, Costa Rica; J. Aires, Argentina; MCNZ, M MIZA, Museo del Instituto de niversidade de São Paulo, Br Washington D.C., United | ction, Nor uto Oswal SP, Instit ger, Joe E de Ciênci logía Agr UFRG, D SAM, S |  | Adelaide, Australia.

Table S3. Mean, standard error, and minimum and maximum values of setae in the tibial comb and the foretibial apparatus, and number of
examined specimens ( N ), for each of the examined species of Pentatomidae, Thyreocoridae, and Scutelleridae . [mean $\pm$ standard error (range)]

| Family | Subfamily | Tribe | Especies | Tibial comb | N | Foretibial apparatus | N |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Pentatomidae | Aphylinae |  | Aphylinae sp. | 20 | 1 | 4 | 1 |
|  | Asopinae |  | Alcaeorrhynchus grandis (Dallas, 1851) |  |  | $20.5 \pm 2.5$ (18-23) | 2 |
|  |  |  | Oplomus cruentus (Burmeister, 1835) | 37 | 1 | $14.5 \pm 0.5$ (14-15) | 2 |
|  |  |  | Oplomus salamandra (Burmeister, 1835) | $29.6 \pm 2.03$ (26-33) | 3 | $13.6 \pm 0.67$ (13-15) | 3 |
|  |  |  | Podisus nigrispinus (Dallas, 1851) | $17.7 \pm 1.38$ (15-21) | 4 | $29 \pm 3$ (26-32) | 4 |
|  | Cyrtocorinae |  | Cyrtocoris egeris Packauskas \& Schaefer, 1998 | $18.5 \pm 0.5$ (18-19) | 2 | 5.0 | 2 |
|  |  |  | Cyrtocoris trigonus (Germar, 1839) | 28.0 | 1 | 6.0 | 1 |
|  | Discocephalinae | Discocephalini | Antiteuchus mixtus (Fabricius, 1787) | 24.0 | 1 | 5.0 | 1 |
|  |  |  | Eurystethus microlobatus Ruckes, 1966 |  |  | 5.0 | 2 |
|  |  | Ochlerini | Alitocoris parvus (Distant, 1880) | $29.5 \pm 0.5$ (29-30) | 2 | 5.0 | 1 |
|  |  |  | Catulona pensa Rolston, 1992 | 36.0 | 1 | 6.0 | 1 |
|  |  |  | Lincus securiger Breddin, 1904 | $35.5 \pm 1.5$ (34-37) | 2 | $5.5 \pm 0.5$ (5-6) | 2 |
|  | Edessinae |  | Brachystethus geniculatus (Fabricius, 1787) | 34.0 | 1 | 5.0 | 1 |
|  |  |  | Brachystethus vexillum Breddin, 1903 | 34.0 | 1 | 5.0 | 1 |
|  |  |  | Edessa quadridens Fabricius, 1803 | 33.0 | 1 | $5.5 \pm 0.5$ (6-5) | 2 |
|  |  |  | Edessa rufomarginata (De Geer, 1773) | 36.0 | 1 |  |  |
|  |  |  | Olbia elegans (Stål, 1862) |  |  | 5.0 | 2 |
|  | Pentatominae | Antestiini | Antestia ellenriederi Breddin, 1900 | 24.0 | 1 | $7 \pm 2$ (5-9) | 2 |
|  |  |  | Antestiopsis anchora (Thunberg, 1783) | 25.0 | 1 | 5.0 | 1 |
|  |  |  | Plautia stali Scott, 1874 | 25.0 | 1 | 5.0 | 2 |
|  |  |  | Starioides degenera (Walker, 1867) | 24.0 | 1 | 5.0 | 2 |
|  |  | Carpocorini | Acledra albocostata (Spinola, 1852) | $25 \pm 1$ (24-26) | 2 | 4.0 | 2 |
|  |  |  | Agonoscelis nubilis (Fabricius, 1775) | 31.0 | 1 | 4.0 | 2 |
|  |  |  | Agroecus griseus Dallas, 1851 | $28 \pm 0.58$ (27-29) | 3 | 3.0 | 1 |
|  |  |  | Berecynthus hastator (Fabricius, 1794) | $28.5 \pm 0.5$ (28-29) | 2 | 4.0 | 1 |




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Procleticini
Rhynchocorini

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\begin{aligned}
& \text { Shyllocephalinae } \\
& \text { Scutelleridae } \\
& \text { Thyreocoridae Corimelaeninae }
\end{aligned}
$$

$$
\begin{aligned}
& \text { Rhyncholepta meinanderi Becker \& Grazia-Vieira, } 1971 \\
& \text { Banasa dolabrata Thomas, 1988 } \\
& \text { Banasa patagiata (Berg 1879) } \\
& \text { Banasa zeteki Sailer, 1959 } \\
& \text { Janeirona stali (Kormilev, 1956) } \\
& \text { Lelia octopunctata (Dallas, 1849) } \\
& \text { Myota aerea (Herrich-Schäffer, 1842) } \\
& \text { Neotibilis (Laeviscutis) compascens (Bergroth, 1914) } \\
& \text { Neotibilis (Neotibilis) parva (Distant, 1893) } \\
& \text { Pallantia macula (Dallas, 1851) } \\
& \text { Pallantia macunaima Grazia, 1980 } \\
& \text { Pellaea stictica (Dallas, 1851) } \\
& \text { Pentatoma japonica (Distant, 1882) } \\
& \text { Pentatoma metallifera (Motschulsky, 1859) } \\
& \text { Pentatoma rufipes (Linnaeus, 1758) } \\
& \text { Phalaecus pustulatus de Geer, 1773 } \\
& \text { Taurocerus achilles (Stå, 1862) } \\
& \text { Dendrocoris arizonensis Barber, 1911 } \\
& \text { Dendrocoris pini Montandon, 1893 } \\
& \text { Thoreyella cornuta Berg, 1883 } \\
& \text { Thoreyella trinotata Berg, 1878 } \\
& \text { Biprorulus bibax } \text { Breddin, 1900 } \\
& \text { Cuspicona simplex Walker, 1867 } \\
& \text { Macrina juvenca (Burmeister, 1835) } \\
& \text { Augocoris gomesi Burmeister, 1835 } \\
& \text { Orsilochides leucoptera (Germar, 1839) } \\
& \text { Corimelaena (Corimelaena) tibialis (Fabricius, 1803) } \\
& \text { Corimelaena (Parapora) extensa Uhler, 1867 } \\
& \text { Cydnoides (Cydnoides) ciliatus Uhler, 1863 } \\
& \text { Cydnoides (Cydnoides) renormatus (Uhler, 1895) }
\end{aligned}
$$

| Galgupha (Euryscytus) difficilis (Breddin, 1914) | 18.0 | 1 | 4.0 | 1 |
| :--- | :---: | :--- | :--- | :--- |
| Galgupha (Galgupha) atra Amyot \& Serville, 1843) | $17.5 \pm 0.5(17-18)$ | 2 | 4.0 | 2 |
| Galgupha (Gyrocnemis) cruralis (Stål, 1862) | $20 \pm 1(19-21)$ | 2 | 4.0 | 1 |
| Galgupha (Microcompsus) vinculata (Germar, 1839) | 17.0 | 1 | 4.0 | 2 |
| Galgupha (Nothocoris) terminalis (Walker, 1867) | $18.5 \pm 0.5(18-19)$ | 2 | 4.0 | 2 |
| Thyreocoris scarabaeoides (Linnaeus, 1758) | $15 \pm 1(14-16)$ | 2 | 4.0 | 1 |

Tibial comb and foretibial apparatus of the following species were not observed: PENTATOMIDAE: EDESSINAE: Lopadusa (L.) augur Stål, 1860; PENTATOMINAE: Procleticini: Lobepomis peltifera Berg, 1891; SCUTELLERIDAE: Pachycoris torridus Scopoli, 1772.


Figure S1. Pretarsus, tibial comb, and tibial apparatus of Aphylinae and Phyllocephalinae (Pentatomidae). A-C, Aphylinae sp.; D-F, Macrina juvencus (Phyllocephalinae). Scales: 100 $\mu \mathrm{m}$.


Figure S2. Pretarsus, tibial comb, and tibial apparatus of Asopinae (Pentatomidae). A-B, Alcaeorrynchus grandis; C-E, Oplomus cruentus; F-H, Oplomus salamandra; I-K, Podisus connexivus; L, Podisus mellipes. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S3. Pretarsus, tibial comb, and tibial apparatus of Cyrtocorinae (Pentatomidae). A-C, Cyrtocoris egeris; D-G, Cyrtocoris trigonus. Scales: $100 \mu \mathrm{~m}$.


Figure S4. Pretarsus, tibial comb, and tibial apparatus of Discocephalini (A-G) and Ochlerini (H-L) (Discocephalinae, Pentatomidae). A-C, Alitocoris parvus; D, Catulona pensa; E-G, Lincus securiges; H-I, Eurystethus microlobates; J-L, Antiteuchus mixtus. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S5. Pretarsus, tibial comb, and tibial apparatus of Edessinae (Pentatomidae). A-C, Brachystethus geniculata; D-F, Brachystethus vexillum; G-I, Edessa quadridens; J-K,

Edessa rufomarginata; L, Lopadusa algur. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S6. Pretarsus, tibial comb, and tibial apparatus of Edessinae (Pentatomidae). A-C, Olbia elegans. Scales: $100 \mu \mathrm{~m}$.


Figure S7. Pretarsus, tibial comb, and tibial apparatus of Antestini (Pentatominae, Pentatomidae). A-C, Starioides degenera; D-F, Antestia elleriederi; G-I, Antestiopsis anchora; J-L, Plautia stalli. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S8. Pretarsus, tibial comb, and tibial apparatus of Carpocorini (Pentatominae, Pentatomidae). A-C, Acledra albocostata; D-F, Agonoscelis nubilis; G-I, Agroecus griseus; J-L, Berecynthus hastator. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S9. Pretarsus, tibial comb, and tibial apparatus of Carpocorini (Pentatominae, Pentatomidae). A-C, Coenus delius; D-F, Cosmopepla decorata; G-I, Dichelops (Dichelops) punctatus; J-L, Dichelops (Dichelops) saltensis. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S10. Pretarsus, tibial comb, and tibial apparatus of Carpocorini (Pentatominae, Pentatomidae). A-C, Dichelops (Diceraeus) furcatus; D-F, Dichelops (Diceraeus) melacanthus; G-I, Dichelops (Prodichelops) divisus; J-L, Diploxys sp. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S11. Pretarsus, tibial comb, and tibial apparatus of Carpocorini (Pentatominae, Pentatomidae). A-C, Dolycoris baccarum; D-F, Hypatropis inermis; G-I, Mormidea quinqueluteum; J-L, Mormidea ypsilon. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S12. Pretarsus, tibial comb, and tibial apparatus of Carpocorini (Pentatominae, Pentatomidae). A-C, Oebalus poecilus; D-F, Tibraca limbativentris. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S13. Pretarsus, tibial comb, and tibial apparatus of Catacanthini (Pentatominae, Pentatomidae). A-C, Arocera apta; D-F, Arocera contralineata; G-H, Rhyssocephala infuscata; I-K, Rhyssocephala rufolimbata. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S14. Pretarsus, tibial comb, and tibial apparatus of Catacanthini (Pentatominae, Pentatomidae). A-C, Runibia decorata; D-E, Runibia perspicua. Scales: $100 \mu \mathrm{~m}$; insets, 20 $\mu \mathrm{m}$.


Figure S15. Pretarsus, tibial comb, and tibial apparatus of Chlorocorini (Pentatominae, Pentatomidae). A-C, Arvelius albopunctatus; D-F, Chlorocoris (Chlorocoris) complanatus; G-I, Chlorocoris (Chlorocoris) distinctus; J-L, Chlorocoris (Monochricerus) flaviviridis. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S16. Pretarsus, tibial comb, and tibial apparatus of Chlorocorini (Pentatominae, Pentatomidae). A-C, Chlorocoris (Monochricerus) subrugosus; D-F, Chloropepla rolstoni; G-I, Chloropepla vigens; J-L, Loxa deducta. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S17. Pretarsus, tibial comb, and tibial apparatus of Chlorocorini (Pentatominae, Pentatomidae). A-C, Loxa flavicolis; D-F, Rhyncholepta grandicallosa; G-I, Rhyncholepta meinderi. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S18. Pretarsus, tibial comb, and tibial apparatus of Pentatomini (Pentatominae, Pentatomidae). A-C, Banasa dolabrata; D-F, Banasa patagiata; G-I, Banasa zeteki; J-L, Janeirona stali. Scales: $50 \mu \mathrm{~m}$; insets, $10 \mu \mathrm{~m}$.


Figure S19. Pretarsus, tibial comb, and tibial apparatus of Pentatomini (Pentatominae, Pentatomidae). A-B, Lelia octopunctata; C-E, Myota aerea; F-H, Neotibilis compascens; IK, Neotibilis parva. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S20. Pretarsus, tibial comb, and tibial apparatus of Pentatomini (Pentatominae, Pentatomidae). A-C, Pallantia macula; D-F, Pallantia macunaima; G-I, Pellaea stictica; JL, Pentatoma japonica. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S21. Pretarsus, tibial comb, and tibial apparatus of Pentatomini (Pentatominae, Pentatomidae). A-C, Pentatoma metallifera; D-F, Pentatoma rufipes; G-H, Phalaeucus pustulatus; I-K, Taurocerus achilles. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S22. Pretarsus, tibial comb, and tibial apparatus of Procleticini (Pentatominae, Pentatomidae). A-C, Dendrocoris arizonensis; D-F, Dendrocoris pini; G, Lobepomis peltifera; H-J, Thoreyella cornuta; K-L, Thoreyella trinotata. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S23. Pretarsus, tibial comb, and tibial apparatus of Rhynchocorini (Pentatominae, Pentatomidae). A-C, Biprorulus bibax; D-F, Cuspicona simplex. Scales: $100 \mu \mathrm{~m}$; insets, 20 $\mu \mathrm{m}$.


Figure S24. Pretarsus, tibial comb, and tibial apparatus of Scutelleridae. A-C, Augocoris gomesi; D-F, Orsilochides leucoptera. Scales: $100 \mu \mathrm{~m}$; insets, $20 \mu \mathrm{~m}$.


Figure S25. Pretarsus, tibial comb, and tibial apparatus of Thyreocoridae. A-B, Corimelaena (Corimelaena) tibialis; C-E, Corimelaena (Parapora) extensa; F-H, Cydnoides (Cydnoides) ciliatus; I-K, Cydnoides (Cydnoides) renormatus. Scales: $50 \mu \mathrm{~m}$; insets, $10 \mu \mathrm{~m}$.


Figure S26. Pretarsus, tibial comb, and tibial apparatus of Thyreocoridae. A-C, Galgupha (Euryscytus) difficilis; D-E, Galgupha (Galgupha) atra; F-H, Galgupha (Galgupha) cruralis; I-K, Galgupha (Microcompsus) vinculata. Scales: $50 \mu \mathrm{~m}$; insets, $10 \mu \mathrm{~m}$.


Figure S27. Pretarsus, tibial comb, and tibial apparatus of Thyreocoridae. A-C, Galgupha (Nothocoris) terminalis; D-E, Thyreocoris scarabaeoides. Scales: $50 \mu \mathrm{~m}$; insets, $10 \mu \mathrm{~m}$.

## APÊNDICE II

# Material suplementar ao Capítulo IV <br> Phylogeny of Dichelops (Hemiptera: Pentatomidae) 

## Appendix S1: List of examined material

## Depositories (curators within parentheses)

AMNH American Museum of Natural History, New York, United States (R.T. Schuh).
DARC David Rider Collection, North Dakota, United States (D.A. Rider).
DZUP Museu de Entomologia Pe. Jesus Santiago Moure, Universidade Federal do Paraná, Paraná, Brazil (R.R. Cavichioli).

FIOC Fundação Instituto Oswaldo Cruz, Rio de Janeiro, Brazil (J.Costa).
IFML Instituto Fundación Miguel Lillo, Tucumán, Argentina (L. Claps).
MACN Museo Argentino de Ciencias Naturales "Bernardino Rivadavia", Buenos Aires, Argentina (A. Bachmann).
MCNZ Museu de Ciências Naturais da Fundação Zoobotânica do Rio Grande do Sul, Rio Grande do Sul, Brazil (A. Barcellos).

MIZA Museo del Instituto de Zoología Agrícola , Maracay, Venezuela (E. Osuna).
MNNC Museo Nacional de Historia Natural, Santiago, Chile (M. Elgueta).
MNRJ Museu Nacional, Universidade Federal do Rio de Janeiro, Rio de Janeiro, Brazil (G. Mejdalani).

MPUJ Pontificia Universidad Javeriana, Museo Javeriano de Historia Natural, Laboaratorio de Entomologia, Bogotá, Colombia (D. Forero).

MZSP Museu de Zoologia da Universidade de São Paulo, São Paulo, Brazil (M. Duarte).

RMNH National Museum of Natural History (Naturalis), Leiden, The Netherlands (Y. van Nierop).

UEMA Universidade Estadual do Maranhão, São Luiz, Maranhão, Brazil (F. Oliveira).
UFRG Departamento de Zoologia, Universidade Federal do Rio Grande do Sul, Rio Grande do Sul, Brazil (J. Grazia).

USNM National Museum of Natural History, (United States National Museum), Washington D.C., Estados Unidos (T. Henry).
ZUEC Museu de Zoologia "Adão José Cardoso", Universidade Estadual de Campinas, Campinas, São Paulo, Brazil (A. Freitas).

## Dichelops (Dichelops) australis Grazia \& Klein, 2012

Holotype, ${ }^{\text {T, }}$, BRAZIL: Rio Grande do Sul, Eldorado do Sul (J. Klein) (MCNZ). Paratypes, ARGENTINA: 4 đ̃, 3 ㅇ, Santa Fe, El Sombrerito, 11.ix. 1964 (A.A. Pirán) (MACN); BRAZIL: 1 §, São Paulo, Campinas (UFRG); 2 §, Rio Grande do Sul, Cachoeirinha, 17.vii. 1985 (Albuquerque) (UFRG).

## Dichelops (Dichelops) avilapiresi Grazia, 1978

Paratype, 1 §, BRAZIL: Santa Catarina: Pinhal, xii. 1948 (A. Maller) (DARC). Additional material examined. BRAZIL: São Paulo: 1 §̃, Americana, 11.xi. 1976 (W.A. Lacerda) (UFRG); 1 ㅇ, Santo André, 20.ii. 1962 (L. Stowbunenko) (UFRG); Paraná: Pinhão, 1 ㅇ, VII. 1992 (G. Skuk) (UFRG), 1 đ, 26.x. 1991 (R.P. da Rocha) (UFRG); Santa Catarina: Nova Teutônia, 300-500m, $27^{\circ} 11^{\prime} \mathrm{S} 52^{\circ} 23^{\prime} \mathrm{E}$ (F. Plaumann); 2 §, ix. 1939 (DARC); 1 §, ii. 1969 (DARC); 2 ㅇ, iii. 1971 (DARC); 1 đ, xi. 1973 (DARC); 1 §, 1 ㅇ, xii. 1973 (DARC); 1 ㅇ,
 Derrubadas, 16.iv. 1983 (S.L. Bonatto) (UFRG); 1 §, 4 \& , no data.

## Dichelops (Dichelops) bicolor Distant, 1890

2 §, 1 q, BRAZIL: Mato Grosso, Juína, 01.xi. 1978 (K.S. Brown Jr.) (ZUEC); 1才, 1q, COLOMBIA: Valle del Cauca, Tuluá, 24-31.viii. 1996 (M. Reyes) (MPUJ); 1 §, ECUADOR: Napo, Santa Cecilia, 240m, 8.vi-1.viii. 1968 (W.G. Saul) (DARC); PERU: 1 §, Junin, Valle Chanchamayo, 800m, 13.viii. 1956 (Weirauch) (DARC); 1 §, Satipo, ii. 1944 (P. Paprzyck) (UFRG).

## Dichelops (Dichelops) leucostigmus (Dallas, 1851)

BRAZIL: 1 \& , Acre, Rio Branco, 06-10.v. 1981 (J. Arias) (UFRG); Amazonas, 1 ภ, 3 q, Alexo, 06.vii. 1941 (Parko) (FIOC); 1 §, 2 个, Manaus, 06.vii. 1941 (Parko) (UFRG); 1 §, Manaus, 25.v. 1982 (L.R. Loterre) (UFRG); 1 §, Manaus, 24.iii. 1986 (R. Sampaio) (UFRG); 1 ¢, Serra dos Porcos, viii. 1977 (M. Franklin) (UFRG); 1 , Distrito Federal, Planaltina, 23.iii. 1977 (V.O. Becker) (CPAC); 1 §', Goiás: Goiânia, 13.i. 1975 (E. Ferreira) (DARC); Maranhão: 1 ठ̉, Bom Jardim, Reserva Biológica Gurupi, 17-27.i. 2010 (F. Limeira-deOliveira, R.O. Souza \& M.B. Aguiar Neto) (UEMA); 6 §, 9 ㅇ, Carolina, Fazenda Cincorá, 17-22.x. 2009 (F. Limeira-de-Oliveira, R.O. Souza \& M.B. Aguiar Neto) (UEMA); Mato

Grosso, 1 q, Jaçanã, P.N. Xingu, xi. 1960 (W. Alvarenga) (DZUP); 2 §, 1 q, Aripuana, $10^{\circ} 25^{\prime}$ S $59^{\circ} 28^{\prime}$ E, 300m, 17-22.iii. 1977 (D. Engleman) (DARC); Rondônia: 1 q, Porto Velho, 15.iii. 1979 (J. Campbell) (UFRG); 1 \&, 62km SW Ariquemes, 30.iii-10.iv. 1992 (J.E. Eger) (DARC); $1 \delta^{\lambda}, 62 \mathrm{~km}$ SW Ariquemes, 25.x. 1993 (U. Schmitz) (DARC); $1 \jmath^{\lambda}, 1$ \&, 62km SW Ariquemes, 08-20.xi. 1994 (J.E. Eger \& L.B. \& C.W. O’Brien) (DARC); COLOMBIA: 1 ${ }^{\text {², }}$, Meta, Puerto Lopez, ix. 1991 (Martinez Guerra) (MPUJ).

## Dichelops (Dichelops) miriamae Grazia, 1978

Paratypes, BRAZIL: Rio Grande do Norte: Natal, 1 §, i. 1950 (M. Alvarenga) (UFRG); 1 §, 1 ㅇ, ii. 1950 (M. Alvarenga) (FIOC, MNRJ); 1 §', xii. 1952 (M. Alvarenga) (MNRJ); 1 , iii. 1952 (M. Alvarenga) (MNRJ). Additional analyzed specimens. BRAZIL: $1{ }^{\lambda}, 1$, Bahia, Encruzilhada, xi. 1972 (M. Alvarenga) (DARC).

## Dichelops (Dichelops) nigrum Bergroth, 1914

ARGENTINA: 1 q, Misiones, Puerto Iguazu, 30.xii. 1991 (DARC); BRAZIL: Amazonas: 1 ठె, Manaus, 06.vii. 1941 (Parko) (UFRG); 1 \&, Manaus, 06.iii. 1988 (J.C. Furtado) (UFRG); 1 §, Espírito Santo, Santa Tereza, 07.xii. 1964 (C. Elias) (DZUP); 1 §, Maranhão, Bom Jardim, R.B. Gurupi, 05-15.vi. 2010 (F. Limeira-de-Oliveira, R.O. Souza \& M.B. Aguiar Neto) (UEMA); Rio de Janeiro: 1 §, Rio de Janeiro, 25.xi. 1952 (Zajciv) (UFRG); 1 \&, Rio de Janeiro, vi. 1949 (Carvalho) (UFRG); 1 \&, Rio Grande do Sul, Derrubadas, P.E. Turvo, 17.xii. 1982 (S.L. Bonatto) (UFRG); Santa Catarina: Nova Teutonia, 300-500m, 27¹1'S $52^{\circ} 23^{\prime} \mathrm{E}$ (F. Plaumann) 1 q iv. 1973 (DARC), $1 \delta^{\star}$ xii. 1975 (DARC); COLOMBIA: $1 才, 1$ \& , Meta, Restrepo (G. Gómez) (UFRG); VENEZUELA: 1 §̧, 1 q, Amazonas, San Juan de Manapiare, 03.iv. 1958 (UFRG); $1 \AA^{\lambda}$, Aragua, Cata, $15 . \mathrm{ix}$.1965 (J. \& B. Bechyné) (UFRG).

## Dichelops (Dichelops) peruanus Grazia, 1978

Paratypes, PERU: Junin: Satipo, 2 ㅇ, i. 1937 (P. Paprzyck) (FIOC, MNRJ); 12 \&, xii. 1937 (P. Paprzyck) (FIOC); 1 \&, xii. 1943 (P. Paprzyck) (FIOC); 1 §, 1 ㅇ, iii. 1944 (P. Paprzyck) (FIOC). Additional analyzed specimens. ECUADOR: Napo: Limón Cocha, 1 \&, 1965 (C.R. Patrick) (DARC); 1 §, 1 q, 17.ix. 1964 (G. Dozier) (DARC); 1 §, 17.viii. 1964 (G. Dozier) (DARC); 1 ㅇ, 19.vi. 1965 (C.R. Patrick) (DARC); 2 ठ, 1ㅇ, 25.vi. 1965 (C.R. Patrick) (DARC); PERU: Huanuco: $1 \delta^{\lambda}$, Tingo Maria, Monzon Valley, 02.xi. 1954 (E.I. Schlinger \& E.S. Ross) (DARC); 1 入, Tingo Maria, 650m, $5 . i v .1963$ (T.H. Hubbell \& L.E. Pena) (DARC).

## Dichelops (Dichelops) pradoi Grazia, 1978

Paratypes, 1 ㅇ, BRAZIL: Goiás, Aragarças, xi. 1965 (Alvarenga) (MZSP); 1 ㅇ, BRAZIL: Mato Grosso do Sul, Corumbá, Serra do Urucum, 29.xi. 1940 (K. Lenko) (MZSP). Additional
analyzed specimens．BRAZIL：Maranhão： $1 \delta^{\lambda}$ ，Caxias，Morro do Alecrim，01－10．i． 2010 （F． Limeira－de－Oliveira）（UEMA）；São Paulo： 1 §，Araçatuba，Rio Jacarecatinga，x． 1961 （Lane \＆Rabello）（MZSP）．

## Dichelops（Dichelops）punctatus Spinola， 1837

BRAZIL： $1 \jmath^{\lambda}$ ，Minas Gerais：Viçosa，iii． 1963 （F．Werner，U．Martins \＆L．Silva）（UFRG）； Rio de Janeiro： 1 ¢，Petrópolis，05－07．iii． 1962 （J．Bechyné）（UFRG）； 1 \＆，Rio de Janeiro， x． 1944 （Carvalho）（MNRJ）；São Paulo： 1 §，São Paulo，ii． 1938 （E．Schur）（UFRG）； 1 q，São Paulo，17．ii． 1934 （MZSP）．

## Dichelops（Dichelops）saltensis Grazia， 1978

ARGENTINA： 1 ¢，Salta，S．Martin Pocitos，i． 1962 （A．Martinez）（MZSP）；Tucumán： 1 q， Aconquija，xi． 1946 （R．Golbach）（DARC）； 7 §， 23 \＆，Tucumán，iii． 1959 （Golbach）（UFRG）； 1 đ̂， 1 \＆，Tucumán，25．i． 1962 （W．Weirauch）（IFML）．

## Dichelops（Diceraeus）caatinguensis Grazia \＆Poock－da－Silva， 2013

Holotype ठ̉，BRAZIL：Paraíba，Soledade，Juazeirinho， $08 . v i i .1956$（A．G．A．Silva）（MNRJ）． Paratypes： 1 §， 1 q，BRAZIL：Rio Grande do Norte，Baixa Verde，vii． 1950 （B．J．Souza） （UFRG）； 1 q，BRAZIL：Paraíba，Soledade，Juazeirinho， $22.1 i i .1956$（A．G．A．Silva）（MNRJ）； 1 §，BRAZIL；Paraíba，Soledade，Juazeirinho，08．vii． 1956 （A．G．A．Silva）（MNRJ）； 1 §， 10 ＋，BRAZIL：Pernambuco，Casa Nova，v．1974，Caatinga（J．C．M．Carvalho）（UFRG，MNRJ）； 6 \＆，BRAZIL：Pernambuco，Petrolina，v．1974，Caatinga（J．C．M．Carvalho）（MCNZ，MNRJ）； 1 ¢，BRAZIL：Pernambuco，Petrolina，v． 1969 （M．Alvarenga）（MNRJ）； 1 q，BRAZIL： Pernambuco，Petrolina，Estrada Picos，v．1974，Caatinga（J．C．M．Carvalho）（MNRJ）； 1 q， BRAZIL：divisa entre Pernambuco e sul do Piauí，Estrada Picos Km3，v．1974，Caatinga （J．C．M．Carvalho）（MNRJ）； 1 入̄，BRAZIL：Bahia，Anajê，16．v． 1975 （C．\＆P．Elias）
 BRAZIL：Bahia，Juazeiro，v．1974，Caatinga（J．C．M．Carvalho）（MNRJ）； 1 \＆，BRAZIL： Bahia，Juremal，Estrada do Juazeiro，v．1974，Caatinga（J．C．M．Carvalho）（MNRJ）； 1 q， BRAZIL：Bahia（G．Bondar）； 1 §， 2 个，BRAZIL：nordeste， 1933 （Ihering）（UFRG，FIOC）．

## Dichelops（Diceraeus）furcatus（Fabricius，1775）

ARGENTINA： 2 万，Buenos Aires，Pergamino（P．Rebagliati）（UFRG）； 1 q，Chaco，Pozo La Gringa，i． 1995 （DARC）； $1 \widehat{J}^{\lambda}$ ，Entre Rios，Concordia，Parque San Carlo（P．Rebagliati） （UFRG）； 1 q，Mendoza，Junin， $20 . i i i .1945$（A．Silva）（UFRG）； $1 ઊ^{\lambda}$ ，Santa Fé，Huges，iii． 2000 （P．Rebagliati）（UFRG）；BOLIVIA： 1 §，Cochabamba，Cristal Mayu，600m，x． 1962 （DARC）； 1 §＇，Santa Cruz，Buena Vista，xi． 1961 （DARC）；BRASIL： 1 q，Minas Gerais， Viçosa，01．xii． 1990 （J．E．M．Leite）（UFRG）；Paraná： 2 \＆，Cascavel，10．viii． 1993 （J．B．Nessa）
（UFRG）； 2 §，Guarapuava， $13.1 i .1974$（A．R．Panizzi）（UFRG）；Rio de Janeiro： 1 §，Campos do Jordão，Morundo，viii． 1978 （M．Alvarenga）（UFRG）； 1 §，Itatiaia，v． 1949 （W．Zikán） （UFRG）；Rio Grande do Sul： 13 ठ̉，$^{\text {T，}} 27$ ，Bagé，27－30．iii． 2006 （Cohen \＆Schwertner） （UFRG）； 4 §， 3 中，Derrubadas，P．E．Turvo，26－30．ix． 1983 （S．L．Bonatto）（UFRG）；Santa Catarina： 2 §̉， 3 ค，Água Doce，11．iii． 2012 （E．Foelkel）（UFRG）； 3 §̉，Chapecó，18．iii． 1983 （J．M．Milani）（UFRG）；1才，1q，São Paulo，Piracicaba，12．iv． 1938 （A．Silva）（UFRG）； PARAGUAY：Central： 1 đ̋，Asunción，15．i． 1983 （E．G．Riley）（DARC）； 1 §， 1 q，Nemby， 11．i． 1983 （E．G．Riley）（DARC）； 1 §，San Lorenzo，16．i． 1983 （E．G．Riley）（DARC）； 1 §， Cordillera，Caacupe，17－20．i． 1983 （E．G．Riley）（DARC）； 3 §， 1 q，Presidente Hayes， 42 km NW Benjamin Aceval，31．i． 1983 （E．G．Riley）（DARC）；URUGUAY： 1 q，Colonia，La Estanzuela，11．v． 1945 （A．Silva）（UFRG）； 1 ¢，Maldonado，Piriapolis，Punta Maldonado， 17．ii． 1997 （P．Araújo）（UFRG）．

## Dichelops（Diceraeus）lobatus Grazia， 1978

ARGENTINA： 1 Q，Catamarca，El Rodeo，20－28．ii． 1958 （Golbach）（IFML）； 1 §，Cordoba， La Caleta，ii． 1956 （A．Willink）（IFML）；Salta： 1 \＆，Cafayate，12．xi． 1948 （Monrós－Willink） （IFML）； 1 § ${ }^{\text {® }}$ ，Cafayate， $10 . \mathrm{iii} 1951$（K．Hayward）（IFML）．

## Dichelops（Diceraeus）melacanthus（Dallas，1851）

ARGENTINA： 1 §，Catamarca，Andalgala， $7 . x i i .1971$（D．J．Brothers）（DARC）；Chaco： 4 q， Pozo La Gringa，i． 1995 （DARC）； 1 §，Charata，xi．1994－i． 1995 （DARC）； 1 §§，Cordoba：Agua de Oro，3．iii． 1976 （A．Willink）（DARC）；Corrientes： 1 §， 3 \＆，7km S Bella Vista，16．i．1989 （C．W．\＆L．O’Brien \＆G．Wibmer）（DARC）； 1 \＆，3km W Valencia，23．i． 1989 （C．W．\＆L． O＇Brien \＆G．Wibmer）（DARC）； 2 §， 7 \＆，24km W Ituzaingo，29．i． 1989 （C．W．\＆L．O＇Brien \＆G．Wibmer）（DARC）； 3 §， 4 ¢，Entre Rios，Colon，iii． 1990 （DARC）； 1 §， 1 \＆，Formosa， 5km N．Pirane，27．i． 1989 （C．W．\＆L．O’Brien \＆G．Wibmer）（DARC）； 2 §， 2 ㅇ，Santa Fe， Santa Fe，31．i． 1989 （C．W．\＆L．O’Brien \＆G．Wibmer）（DARC）；BOLIVIA：Santa Cruz： 1 q， 8mi S Santa Cruz de la Sierra，18．iv． 1978 （O’Brien \＆Marshall）（DARC）； 1 ठ， 5 km N Warnes，14．v． 1980 （D．Foster）（DARC）；BRAZIL： 1 q，Amapá，Porto Platon，ii． 1964 （J．C．M． Carvalho）（MNRJ）；Ceará： 1 đ̉，Aracati，iii． 1952 （F．S．Silva）（MNRJ）； 1 q，Fortaleza， 02．xi． 1967 （F．Costa）（UFRG）； $1 \delta^{\lambda, 1} 1$ \＆，Goiás，Minaçu，Serra da Mesa，19－30．xi． 1996 （L． Moura）（UFRG）； 3 万， 6 ㅇ，Maranhão，Caxias，Campus UEMA，Morro do Alecrim，01－ 08．iii． 2010 （D．W．A．Marques）（UEMA）； 2 §̉， 1 ㅇ，Mato Grosso，Barranco Branco， 12．xii． 1935 （Souto Maior）（UFRG）； 1 §， 1 \＆，Mato Grosso do Sul，Dourados，xi． 1993 （A．R． Panizzi）（UFRG）； 1 §̉， 1 ¢，Paraná，Londrina，25．xi． 1998 （V．R．Chocorosqui）（UFRG）；Rio Grande do Sul： 3 §̂，Santa Maria，xii． 1981 （D．Link）（UFRG）； 1 §ె， 2 ㅇ，São Sepé， 1981 （Costa \＆Link）（UFRG）；COLOMBIA：Cundinamarca： 3 万，Tolemaida，Quebrada La Naranjola，18．v． 1968 （E．Ramirez Z．）（MHNM）； 1 \＆，Villeta，28．iv． 1968 （G．Aguiar） （MHNM）； 1 \＆，Girardo，v． 1969 （Arias）（MHNM）； 1 §，Meta，Puerto Lopez，ix． 1991 （H． Rodriguez V．）（MPUJ）； 1 Q，Tolima，Mariquita，ix． 1992 （MPUJ）；FRENCH GUIANA： $1 \delta^{\lambda}$ ， La Guyane，Highway N1 to Saint Laurent，9km SE Iracoubo，7．vi． 1986 （E．G．Riley \＆D．A．

Rider）（DARC）；GUYANA： 1 §，Demerara－Mahaica，Georgetown，Ogle airport，vi． 1965 （M．Medina）（DARC）；PARAGUAY：Central： 1 \＆，Asunción，5．ii． 1983 （E．G．Riley） （DARC）； 1 q，Capiatá，21．ii． 1994 （G．Arviagada）（DARC）； 1 q，San Lorenzo，20．i． 1983 （E．G．Riley）（DARC）； 2 \＆，Presidente Hayes，42km NW Benjamin Aceval，31．i． 1983 （E．G． Riley）（DARC）；PERU： 1 §，Cusco，Limatambo，30．iv． 1968 （UFRG）；VENEZUELA： 1 ふ， 1 \＆，Zulia，Maracaibo，viii． 1971 （J．Maldonado）（DARC）．

## Dichelops（Diceraeus）phoenix Grazia， 1978

Paratypes， 1 中，BRAZIL：Rio de Janeiro，Ramos，i． 1953 （E．Lebato）（FIOC）； 1 §，BRAZIL： São Paulo，Pirassununga，iv． 1948 （FIOC）．Additional analyzed specimens．BOLIVIA： 1 §， 1早，Santa Cruz，Buena Vista，xi． 1961 （DARC）；BRAZIL：Distrito Federal：Brasília，Campus
 Brasília，Fazenda UNB， $15 . \mathrm{ii} 1969$（I．C．B．）（UFRG）； 1 §， 1 q，Mato Grosso do Sul，Itaum， Dourados，iii． 1974 （M．Alvarenga）（DARC）； 1 \＆Paraná，Laranjeiras， $25^{\circ} 24^{\prime} \mathrm{S} 52^{\circ} 23^{\prime} \mathrm{E}$ ， 900m，iii． 1963 （F．Plauman）（DARC）； 1 ठ，Santa Catarina，Nova Teutônia， $27^{\circ} 11^{\prime}$＇S 52ํ2＇́E，300－500m，7．i． 1964 （F．Plaumann）（DARC）．

## Dichelops（Prodichelops）divisus（Walker，1867）

BRAZIL： 1 §， 1 ㅇ，Amazonas，Estirão da Preta，Rio Liberdade，11－15．V．2011，sweep net（ $D$ ． Takyia）（INPA）；COLOMBIA： 1 ¢，Amazonas，Leticia，12．iv． 1992 （A．Saenz）（UFRG）．

## Acledra albocostata（Spinola，1852）

ARGENTINA： 1 §， 5 ㅇ，Chubut，El Maiten，08．ii． 1965 （UFRG）； 1 §，Río Negro，El Bolson， 20．xii． 1963 （UFRG）；CHILE： 7 ठ $^{\lambda}, 5$ \＆，Coquimbo，Condoriaco，10xi． 1964 （L．E．Pena） （UFRG）； 1 §，Coquimbo，Condoriaco（L．E．Pena）（UFRG）．

## Acledra fraterna（Stål，1859）

ARGENTINA： 2 q，Distrito Federal，Buenos Aires（MACN）； 1 §，San Luis，San Luis， vi． 1953 （UFRG）；CHILE：Santiago：Santiago， 4 §， 4 ¢，27．vii． 1964 （UFRG）； 3 〕， 4 q， Santiago，xi． 1947 （T．Ramirez）（UFGR）．

## Agroecus griseus Dallas， 1851

BRAZIL：Amazonas： 1 §̉， 1 q，Uaupes，07－12．vii． 1956 （M．Alvarenga）； 1 §，Uaupes， vi． 1949 （J．C．M．Carvalho）； 1 ，Mato Grosso，Utiariti，viii． 1961 （K．Lenko）（MNRJ）； Paraná： 1 q，Arapongas，ii． 1952 （A．Mallep）（UFRG）； 2 §， 2 q，Londrina，vi． 2010 （M．M．

Rodrigues) (UFRG); 2 中, Rio de Janeiro, Rio de Janeiro, Tijuca, 1954 (Zajciv) (UFRG); $1 \jmath^{\lambda}$, Rio Grande do Sul, Porto Alegre, 26.xi. 1991 (J.A.M. Fernandes) (UFRG); 1 ㅇ, São Paulo, Barueri, 22.i. 1961 (K. Lenko) (UFRG); PERU: Junin: 1 §, 1 q, Satipo, i. 1937 (P. Paprzyck) (UFRG); 1 §, Satipo, i. 1937 (UFRG). 1 §, VENEZUELA: Aragua, Tiara, 07.viii. 64 (F. Fernandes \& C.J Rosales) (UFRG).No data, 2 ふ̉, 08.iv. 1938 (W. Zikán) (UFRG), 1 q, 25.xi. 1952 (Zajciv) (UFRG).

## Agroecus scabricornis (Herrich-Schäffer, 1844)

BRAZIL: Rio Grande do Sul: 1 q, Catiporã, 22.ix. 1986 (P. Marson) (UFRG); 1 §, Torres, 03.i. 1989 (R. Hildebrand) (UFRG); 2 , Santa Catarina, Florianópolis, 28.i. 1986 (B.P. Mohe); São Paulo: 1 ㅇ, Caraguatatuba, 02.iv. 1962 (M. Reichardt \& Silva) (UFRG); 1 §', Caraguatatuba, 22.v-1.vi. 1962 (UFRG); 2 , Ilha dos Búzios, 16.x-04.xi. 1963 (UFRG); 1 , , Itanhaem, 01-05.v. 1961 (U. Martins) (UFRG); 1 §', Salesópolis, 12.ii. 1963 (L. Silva \& H. Reichardt) (UFRG); 1 §̉, Santo André, 20.ii. 1962 (L. Stowbunenko) (UFRG); 1 §, Santos, 17.x. 2000 (UFRG); 1 \&, São Sebastião, ii. 1992 (F. Silveira).

## Berecynthus hastator (Fabricius, 1798)

BRAZIL: 3 §, 4 ㅇ, Mato Grosso, 17-22.iii. 1977 (D. Engleman) (UFRG); COLOMBIA: $2 \delta^{\lambda}$, 1 \&, Amazonas, Leticia, 02-07.iv. 1975 (D. Engleman) (UFRG); HONDURAS: 1 q, Olancho, Catacamas, 13.vi. 1974 (C.W \& L.O'Brien \& Marshall) (UFRG); PANAMA: Panama: 1 §', Coco, 25.iv. 1972 (D. Engleman) (UFRG); 1 ¢, Fort Kobbe, 20.vi. 1976 (UFRG); 1 §̉, Las Cumbres, 02.vi. 1975 (Henk Wolda) (UFRG); 1 §', Veraguas, Santiago, 06.x. 1973 (D. Engleman) (UFRG); SURINAME: Paramaribo, 1 §, 16-18.vii. 1975 (D. Engleman) (UFRG); 2 ㅇ, 22-30.vii. 1975 (D. Engleman) (UFRG); VENEZUELA: Trujillo: 1 ㅇ, Agua Viva, 19.iv. 1952 (L. Rey \& J.Araujo) (UFRG); 1 §', Escuque, 07.xii. 1969 (R. Casares \& J.B. Teran) (UFRG);

## Caonabo pseudoscilax (Bergroth, 1891)

BRAZIL: Rio Grande do Sul: Itapuã, 2 §, 1 ¢, 16.iii. 2002 (E. Silva \& A. Barcellos) (UFRG); 41 ㅇ, 25.iii. 2002 (E. Silva \& C. Schwertner) (UFRG); 1 §, São Francisco de Paula, $01 . \mathrm{ii} 2007$ (B.B. Furstenau \& L.M. Weiler) (UFRG); 1 §, 1 \&,São Francisco de Paula, 1719.xii. 2005 (P.R.S. Bunde) (UFRG); 2 §̉, 2 O, São Francisco de Paula, $18 . \mathrm{iii} 2007$ (C.F. Schwertner) (UFRG); 1 §, 2 \&, São Francisco de Paula, 18.iii. 2007 (J.L.C. Bernardes) (UFRG); 1 §, São Paulo, Bocaina, 24.iv (UFRG); $1 \widehat{J}^{\lambda}$, no data (UFRG).

UNITED STATES: Puerto Rico: 3 §̉, Guanica, Bosque Estatal de Guánica, Ballena Trail, 26.vii.2004, $17^{\circ} 58^{\prime} 49^{\prime \prime N} 66^{\circ} 51^{\prime} 74 " \mathrm{~W}$, Blacklighting (S.W. Ligafelter) (USNM); $1 \delta^{\lambda}$, Lajas, Parguera, Island Magueyes, 19.xii. 1962 (Paul \& Phyllis Spangler) (USNM). DOMINICAN REPUBLIC: 1 § ${ }^{\lambda}, 1$ ㅇ, La Veja, RD-149 Loma La Golondrina, Reserve Ebano Verde, 11.vii.2003, $1^{\circ} 03.498^{\prime} \mathrm{N} 70^{\circ} 32.670^{\prime} \mathrm{W}$ (D. Perez, R. Bastardo, B. Hierro) (USNM); 1 +, Barahona, 5km NE of Barahona, 29-30.iv. 1978 (Agr. Exp. Sta.) (USNM). BRITISH VIRGIN ISLANDS: 1 §̃, Guana Island, 5-23.vii. 1985 ( S.E. \& P.M. Miller) (USNM); 1 q, Prickley Pear Island, Vixen Point (J.F.G. Clarke) (USNM).

## Carpocoris purpureipennis (DeGeer, 1783)

1 ¢, AUSTRIA: Niederosterreich, 26.vi. 1988 (Stella \& Tatro) (DARC); 1 §, 1 ¢, INDIA (M.M. Carleton) (AMNH); 1 §', KAZAKHSTAN: 15.v. 1992 (DARC); 1 ㅇ, POLAND: Warsaw, 20-30.v. 1988 (M.A. Ivie) (DARC); 1 ठ', UKRAINE: Donets'ka Oblast', Novotroitskoye, 14.iii. 1995 (DARC).

## Coenus delius (Say, 1832)

UNITED STATES: 1 ค, Minnesota, Homer, 10.ix. 1927 (F.M. Uhler) (USNM); 1 §, Missouri, Clayton, 01.x. 1919 (USNM); 1 §, Montana, Musselshell, 16.viii. 1917 (USNM); New York: 1 \& Ithaca, 10.v. 1911 (MNRJ); 1 \&, McLean, 29.v. 1915 (MNRJ); 1 §, Olean, 05.ix. 1913 (MNRJ); 1 Q , no data (USNM).

## Cosmopepla decorata (Hahn, 1834)

MEXICO: 1 §, 1 q, Hidalgo, Tlanchinol, 31.vii. 1982 (L. O'Brien \& G. Wibmer) (AMNH); 1〕̄, Michoacan de Ocampo, San Jose Purua, 05.v. 1962 (Harffer \& Pereira) (MZSP); 1 q, Puebla, Teziutlan, 18.vii. 1982 (L. O'Brien \& G. Wibmer) (AMNH)

## Cosmopepla lintneriana Kirkaldy, 1909

UNITED STATES: $1 \delta^{\lambda}$, Illinois, Chicago, 13.vi. 1956 (J.B. Rayes) (AMNH); 1 ㅇ, Kentucky, 13.vii. 1959 (J.M. Campbell) (AMNH); 1 \&, Michigan, East Lansing, 1965 (E.D. Evans) (AMNH); New York: 1 đ̃, Ithaca, 19.v. 1946 (F. Fernandez) (MIZA); 1 §̂, 1 , Ithaca (UFRG); 2 \&, Oregon, Gardiner, 27.v. 1964 (J.D.Vertrees) (AMNH)

1 ㅇ, CYPRUS: Yermasoyia River, 01.iii. 1950 (G. Mavromoustakis) (AMNH); Unknown country: 3 §̂, 1 中, Médio Rio Amur, vi. 1968 (A. Maslov) (MNRJ); 1 đ̉, 13.v. 1950 (D. Leston) (AMNH).

## Euschistus (Euschistus) emoorei Rolston, 1972

VENEZUELA: Aragua: 2 ㅇ, Cagua, 28.v. 1958 (A. Fernandez) (MIZA); 1 §, 1 ¢, El Limón, 01.vi. 1965 (E. Osuma) (MIZA); 1 §̂, 1 ㅇ, El Limón, 03.vi. 1965 (F. Fernandez) (MIZA); 1〕, El Limón, 12.V. 1963 (E. Osuma) (MIZA); 2 §, 2 ¢, Maracay, 02.v. 1948 (F. Fernandez) (MIZA); 1 §, Rancho Grande, 20.xi. 1965 (F. Romero) (MIZA); Cojedes: 2 §, El Tinaco, 24.v. 1948 (F. Fernandez) (MIZA); 2 §, 1 O, San Carlos, 24.v. 1948 (F. Fernandez \& L. Salas) (MIZA); 1 ㅇ, Lara, Sarare, 27.vii. 1949 (F. Aponte) (MIZA).

## Euschistus (Euschistus) heros (Fabricius, 1794)

BRAZIL: Rio Grande do Sul: 2 §, 2 q, Cruz Alta, 24.v. 2006 (M.T.B. da Silva) (UFRG); 1 q, Derrubadas, 13.iv. 1983 (S.L. Bonatto) (UFRG); 2 §, 1 \& , Derrubadas, 26-30.ix. 1983 (S.L. Bonatto) (UFRG); 1 §, 1 ㅇ, Santa Maria, x. 1980 (D. Link) (UFRG); 1 §, Rondônia, Porto Velho, 30.xii. 1964 (R.T. Lima) (MZSP); Santa Catarina: 1 §̉, Guatambú, 04.xii. 2004 (UFRG); 1 Q, Guatambú, 06.xi. 2004 (UFRG); 1 §§, 1 Q, Guatambú, 27.xi. 2004 (UFRG); São Paulo: 1 đ̉, Campinas, 04.ii (UFRG); 1 \&, Iporanga, xii. 1987 (G. Accacio) (UFRG); 1 q, São Paulo, 02.ii. 1975 (C.M. De Caro) (UFRG); 1 ¢, São Sebastião, 07-16.i. 1992 (J.A.M. Fernandes) (UFRG); 1 §, Teodoro Sampaio, xi. 1977 (M. Alvarenga) (UFRG).

## Euschistus (Euschistus) variolarius (Palisot de Beauvois, 1817)

UNITED STATES: 1 ㅇ, New York, Ithaca, 07.ix. 1977 (UFRG); North Dakota: 1 ㅇ, Oakville Prairie, 22.vi. 2012 (D.A. Rider) (DARC); 1 §̂, 1 \&, Sheldon, 27.vi. 2012 (D.A. Rider) (DARC); 1 §, Sheyenne National Grasslands, $27 . v i .2012$ (D.A. Rider) (DARC); $1 \delta^{\lambda}, 1$ q, Wild Rice River at Red River, 01. viii. 2012 (D.A. Rider) (DARC).

## Euschistus (Lycipta) sharpi (Bergroth, 1891)

BRAZIL: Rio Grande do Sul: 1 ̉, Estação Ecológica do Taim, 23.iii-23.iv. 1981 (J. Grazia) (UFRG); 1 Q, Pelotas (UFRG); São Paulo: 1 đ̃, 1 Y, Santo Amaro, x. 1960 (J. Lane) (MZSP); $1 \delta^{\lambda}, 1$ \&, Santo Amaro, xi. 1960 (J. Lane) (MZSP); 1 \&, Santo Amaro, xii. 1969 (J. Lane) (MZSP); $4 \widehat{\delta}, 6$ q, no data (MZSP).

BRAZIL： 1 §，Distrito Federal（Carvalho）（MNRJ）；Rio Grande do Sul： 1 ㅇ，Maquiné， $01 . \mathrm{iv} .2006$（V．C．Matesco）（UFRG）； 1 §，Maquiné，12．x． 2005 （M．O．Marchiori）（UFRG）； 1 §̉，São Francisco de Paula，18．iii． 2007 （C．F．Schwertner）（UFRG）；Santa Catarina： 1 q， Corupá，xi． 1953 （Maller）（MNRJ）； 1 §，Maracajá，19．iv． 2006 （T．Bertolin）（UFRG）；São Paulo： 1 \＆，Ilhabela，ii． 1953 （E．Rabello）（MZSP）； 1 §，Ilhabela，13．iv． 1963 （H．M．Canter） （MZSP）； 1 §，Ilhabela，03－05．v． 1963 （H．Urban）（MZSP）； 1 ठ，São Sebastião，03．vii． 1961 （K．Lenko）（MZSP）； 1 \＆，no data，07．ii． 2008 （UFRG）； $1 \delta^{\imath}$ ，no data．

## Euschistus（Mitripus）acutus Dallas， 1851

BOLIVIA： 1 \＆，Beni，Rurrenabaque，x． 1956 （Dirings）（MZSP）； 1 q，Santa Cruz，Roboré，06－ 08．ii． 1958 （Monrós）（IFML）；BRAZIL：Amazonas： 2 §， 1 \＆，Benjamin Constant，xi． 1962 （A． Silva）（MZSP）； 1 \＆，Itacoatiara，vi． 1960 （Dirings）（MZSP）； 1 ¢，Rio Purus，Lago de Beruri， 06－08．iv． 1967 （Exp．Perm．Amaz．）（MZSP）； 1 §，Goiás，Jataí，i． 1955 （M．Canera）（MZSP）； 1 ㅇ，Maranhão，Igarapé Gurupi－Una，Aldeia Araçu，ii． 1966 （Malkin）（MZSP）； $1 \delta^{\lambda}, 1$ 中，Mato Grosso，Barra do Tapiragé，14．xii． 1964 （R．Malkin）（MZSP）； 1 §，Pará，Maraba，05．v． 1959 （Alvarenga）（UFRG）； 1 ¢，Rio de Janeiro，Itatiaia，xii． 1950 （L．Iravamos）（UFRG）； 1 ô， Roraima，Boa Vista，17－18．ix． 1964 （A．Mones）（UFRG）；PERU： 1 đ，Cuzco，Carrasco， 20．iv． 1967 （N．Bellavista）（UFRG）；Junin： 1 \＆，Satipo，iii． 1944 （P．Paprzyck）（FIOC）； 1 ㅇ，Satipo，v． 1937 （P．Paprzyck）（FIOC）； 2 §，Valle Chanchamayo，viii． 1944 （Weyrauch） （IFML）； 1 ふ̊，VENEZUELA：Barinas，Ticoporo，22－28．v． 1968 （M．Galbez \＆J．Salcedo） （MIZA）．

## Euschistus（Mitripus）grandis Rolston， 1978

BRAZIL： 1 §，Paraná，Morretes，21．ix． 1974 （Pe．Moure）（DZUP）；Rio de Janeiro： 1 q， Itatiaia，15．xii． 1966 （H．Reichardt）（DZUP）； 1 Q，Rio de Janeiro， 1954 （Zajciv）（MNRJ）； 2 §， Serra dos Órgãos，29．viii． 1957 （J．Bécker）（MNRJ）； 1 đ̃，Teresópolis， 1954 （Zajciv）（MNRJ）； Rio Grande do Sul： 1 §̉，São Francisco de Paula，20－21．xii． 2006 （L．Moura）（UFRG）； 1 中， São Francisco de Paula， 13.1 .2007 （C．F．Schwertner）（UFRG）； 1 §̃，São Francisco de Paula， 18．iii． 2007 （L．M．Weiler）（UFRG）； 3 §̉， 2 \＆，São Francisco de Paula，11．x． 2007 （B．Marks \＆ L．M．Weiler）（UFRG）； 1 §， 1 \＆，São Francisco de Paula，16．iii． 2008 （B．Marks \＆L．M． Weiler）（UFRG）；São Paulo： 1 ¢，Campos do Jordão，29．iv． 1962 （J．Halik）（MZSP）； 1 §， Capão Bonito，12．x． 1991 （J．A．M．Fernandes）（UFRG）； 1 \＆，Paranapiacaba，25．ii． 1962 （L． Stowbunenko）（MZSP）； 1 q，São José do Barreiro，04．xi． 1965 （F．M．Oliveira）（DZUP）．

## Galedanta bituberculata Amyot \＆Serville， 1843

1 ô，ARGENTINA：Misiones，Eldorado，18．viii，1967（DARC）；BRAZIL： 1 q，Rio de Janeiro，Rio de Janeiro（Zajciv）（MNRJ）；Rio Grande do Sul： 1 Y，Canela，04．xi． 1998 （L． Moura）（MCNZ）； 1 §̉，Maquiné，11－13．x． 2005 （R．Ott）（MCNZ）； 1 ㅇ，Porto Alegre，

25．ix． 1957 （MCNZ）； 1 \＆，Torres，13．i． 2005 （L．Kaminski）（MCNZ）； 1 \＆，Triunfo，18．ix． 2006 （R．Ott \＆A．Barcellos）（MCNZ）； 3 \＆，Triunfo，19．xi． 2002 （A．Barcellos）（MCNZ）； 1 中， Triunfo，19．xi． 2002 （M．P．de Barros）（MCNZ）； 1 \＆，Triunfo，30．xi． 2004 （R．Ott \＆A． Barcellos）（MCNZ）； 2 ふ̉，10．xi． 1966 （F．M．Oliveira）（MNRJ，ZUEC）；São Paulo： 1 q， Guanabara，x． 1975 （MNRJ）； 1 §，São Sebastião，05－08．x． 1979 （F．M．P．Balestieri）（UFRG）．

## Glyphepomis adroguensis Berg， 1891

1 ，ARGENTINA：Corrientes，San Tome，ii． 1927 （ UFRG）；BRAZIL：Rio Grande do Sul： 1 Q，Cachoeirinha，17．vii． 1985 （Albuquerque）（UFRG）； 1 ㅇ，Capão do Leão，ii． 1988 （L．C．Belarmino）（UFRG）； 1 §̃，Estação Ecológica do Taim，23．iii－4．iv． 1981 （J．Grazia） （UFRG）； 1 ㅇ，Gramado，09．xii． 1990 （J．Grazia）（UFRG）； 1 ㅇ，Guaíba，04．vii． 1985 （Albuquerque）（UFRG）； 1 个，Guaíba，21．ix． 1982 （G．R．P．Moreira）（UFRG）； 1 §，Pelotas， 01．v． 1963 （UFRG）； 1 个，Porto Alegre，02．i． 1994 （L．A．Campos）（UFRG）； 1 §，Porto Alegre， 19．vi． 1992 （J．A．M．Fernandes）（UFRG）； 2 §， 1 \＆P Porto Alegre，31．I． 1969 （Soffel）（UFRG）； 1 §， 1 ¢，Santa Maria，08．iii． 1986 （D．Link）（UFRG）； 1 ¢，Santo Antônio da Patrulha， 24．vii． 1985 （Becker）（UFRG）； 1 §，Taquara，03．ix． 1989 （UFRG）； 1 §，Tocantins，Gurupi， 01．vi． 1988 （K．Kishing）（UFRG）．

## Glyphepomis setigera Kormilev \＆Pirán， 1952

BRAZIL：Rio Grande do Sul： 3 §， 11 q，Capão do Leão，ii． 1988 （L．C．Belarmino）（UFRG）； 2 đ，Santa Maria，i． 1990 （D．Link）（UFRG）；No data： 1 ठ，07．iii． 1972 （UFRG）； 1 ภ， 11．iii． 1974 （UFRG）．

## Hymenarcis nervosa（Say，1831）

1 ㅇ，MEXICO：Tamaulipas，Nuevo Laredo，02．vii． 1970 （USNM）；UNITED SATES： 2 § Maryland，Odenton，13．i． 1985 （W．E．Steiner \＆J．E．Lowry）（USNM）； 1 \＆，Texas，Punkin Cave，04．ix． 1965 （I．Reddell）（USNM）；Virginia： 1 O＇，Vienna，16．vii． 1922 （H．G．Barber）$^{\text {（H．}}$ （USNM）； 1 §，Vienna，08．vii． 1925 （USNM）．

## Hypatropis inermis（Jensen－Haarup，1928）

BRAZIL：Pará： 1 ¢，Belém，14．iii． 1929 （Bonifácio）（UFRG）； 1 ¢，Belém，29．i． 1964 （A． Souza）（UFRG）； 2 ，，Belém，i． 1997 （UFRG）；Rio Grande do Sul： 2 ，Cachoeirinha，
 ठ̉，Pelotas，10．x． 1967 （UFRG）； 1 ㅇ，Pelotas，08．v． 1978 （Elói）（UFRG）； 1 q，Triunfo， 25．v． 1990 （L．Moura）（MCNZ）； 1 q，Santa Catarina，Morro das Pedras，19．i． 1957 （UFRG）；

São Paulo: 1 O, Paulínia, 21.v. 1982 (C. Paiva) (UFRG); 1 §, Ubatuba, 16-27.iii. 1964 (UFRG); $1 \AA^{\lambda}$, URUGUAY: Montevideo, Santiago Vazquez, 21.x. 1960 (UFRG).

## Hypatropis sternalis (Stål, 1869)

1 §, ARGENTINA: Buenos Aires, Rosas (J.B. Daguerre) (MACN); BRAZIL: Rio Grande do Sul: 1 \& , Imbé, ii. 1961 (E. \& L. Buckup) (UFRG); 1 \&, Pelotas, ii. 1950 (UFRG); 1 §̂, Pelotas, iv. 1975 (J. Claudino) (UFRG); 1 §̉, Santa Catarina, Morro das Pedras, 22.i. 1957 (Pe. Buck) (UFRG).

## Ladeaschistus armipes (Stål, 1872)

BRAZIL: 2 §, Bahia, Rio de Contas, 17.ii. 2002 (M.J.S. Lopes) (UFRG); 1 §, Distrito Federal, Planaltina, 14.iii. 1979 (Kishino) (UFRG); Mato Grosso: 1 q, Chapada dos Guimarães, xi. 1963 (M. Alvarenga) (UFRG); 1 \&, Chapada dos Guimarães, 01-04.ii. 1965 (S. Laroca) (UFRG); 1 ¢, Cuiabá, i. 1963 (M. Alvarenga ); São Paulo: 2 ㅇ, Itirapina, 01.x. 2005 (L. Kaminski) (UFRG); 1 đ̉, 1 ㅇ, Jundiaí, 14.x. 1961 (Werner) (MZSP); 2 ㅇ, Pirassununga, 06-13.i. 1983 (Exc. DZ USP) (MZSP); 1 ㅇ, Pirassununga, 12.x. 1991 (F. Silveira) (UFRG).

## Ladeaschistus bilobus (Stål, 1872)

ARGENTINA: Misiones: 1 q, Panambi, 24.xi. 1951 (Willink-Monrós) (IFML); 1 q, San Ignacio, 21.xi. 1951 (F. Monrós) (IFML); 1 ㅇ, San Javier, 20.xi. 1973 (Willink-Tomsic) (IFML); BRAZIL: 1 +, Mato Grosso, Riacho do Herv, Rio Paraná (Dirings) (MZSP); Rio Grande do Sul: 1 ㅇ, Catuípe, 07.xi. 2004 (F.L. Santos) (UFRG); 2 §’, Catuípe, 27.xi. 2004
 Derrubadas (UFRG); Santa Catarina: 1 入ె, Guatambú, 19.ix. 2004 (UFRG); 1 Q, Guatambú, 02.x. 2004 (UFRG); 1 ㅇ, São Paulo, Porto Cabral, iii-iv. 1944 (L. Travassos) (FIOC); 1 q, PERU: Cusco, Quilla Bamba, 31.i. 1952 (F. Monrós) (IFML); No data, 4 ठ (UFRG).

## Mecocephala acuminata Dallas, 1851

ARGENTINA: Buenos Aires: 1 \&, General Alvarado, 18.xi. 1946 (Mirama \& Bachmann) (MACN); $1 \AA^{\lambda}$, Mar del Plata, 22.xii. 1919 (A. Frers) (UFRG); no data, $1 \jmath^{\lambda}$ (UFRG).

UNITED STATES: Arizona: 1 \&, Portal, 20-23.viii. 1922 (W.J Gertsch) (AMNH); 1 §, Portal, 20.viii. 1966 (AMNH); 1 \&, Maryland, Plummers Island, 14.xii. 1913 (W.L. McAtee) (USNM); 1 §, 1 ㅇ, New Jersey, Brigantine, 02.viii. 1931 (J.C. Lutz) (USNM); Pennsylvania: 1 đ, Gulph Mills, 22.v. 1938 (Ed. Endy) (USNM); 1 §, Philadelphia, 15.vii. 1946 (J.C. Lutz) (USNM).

## Mormidea cornicollis (Stål, 1860)

BRAZIL: Minas Gerais: 1 §, 2 \&, Belo Horizonte, 22.i. 1963 (N. Papavero) (MNRJ, UFRG); 1 §, Caxambu, ii. 1971 (J. Jurberg \& A.L. Perachi) (FIOC); 1 §, Pouso Alegre, xii. 1953 (P. Pereira) (MNRJ); Rio de Janeiro: 1 §, Itatiaia, i. 1978 (Carvalho \& Schaffner) (MNRJ); 1 q, Petrópolis, 03.vii. 1941 (Parko) (MNRJ); 1 \&, Teresópolis, ii. 1969 (J. Jurberg) (FIOC); Rio Grande do Sul: 1 Q, Esmeralda, 18-19.iii. 1981 (J. Grazia) (UFRG); 1 §, Pelotas, 11.iii. 1982 (J. Grazia) (UFRG); 1 Q, São Francisco de Paula, 20.xii. 1959 (Pereira, Ditadi, Petersen, Meyrer \& Volkmer) (MNRJ); 2 §, 1 q, São Francisco de Paula, $28.1 i i .1959$ (Pereira, Ditadi, Petersen, Meyrer \& Volkmer) (MNRJ); São Paulo: 1 §, 1 个, Barueri, xi. 1965 (K. Lenko) (MNRJ); 1 中, Mairiporã, 04-13.i. 1967 (C. Costa) (MNRJ); 1 §, Salesópolis, 15.iii. 1963 (E.X. Rabello) (MNRJ); 1 ㅇ, São Paulo, 13.iv. 1962 (Lenko \& Reichardt) (MNRJ); 1 §, São Paulo, 18.i. 1962 (J. Halik) (MNRJ).

## Mormidea maculata Dallas, 1851

BRAZIL: Amazonas: 1 ¢, Manaus, 19.vii. 1917 (J. Grazia) (UFRG); 1 §, Manaus, 12.v. 1976 (UFRG); 1 đ, 1 q, Nova Olinda, 15.ii. 1963 (E. Cerqueira) (UFRG); 2 §, 2 q, Pará, Belém, 25.i. 1959 (I.C. Leite) (UFRG); 2 §, 2 \&, São Paulo, Ilha Seca, 19-26.ii. 1940 (FIOC); PERU: Junin: 1 §, 1 \&, Satipo, i. 1937 (P. Paprzyck) (FIOC); 1 \&, Satipo, xii. 1937 (P. Paprzyck) (FIOC); 1 đ, Satipo, i. 1944 (P. Paprzyck) (FIOC); 1 đ̉, Satipo, iii. 1944 (P. Paprzyck) (FIOC); VENEZUELA: 1 ㅇ, Aragua, Tocorón, 06.xi. 1947 (F. Fernandez) (MIZA); Territorio Federal: 1 \&, Maracay, 15.iv. 1965 (F. Fernandez) (MIZA); 1 §, Maracay, 15.iv. 1966 (F. Fernandez) (MIZA).

## Mormidea ypsilon (Linnaeus, 1758)

BRAZIL: Pará: 1 §, 05-08.vii. 1978 (UFRG); 1 q, 23.iii. 1978 (M. Andrade) (UFRG); 1 q, 25.iii. 1978 (UFRG); Rio Grande do Sul: $1 \AA^{\imath}$, Derrubadas, 13.iv. 1983 (S.L. Bonatto) (UFRG); 1 §, Derrubadas, 15.iv. 1983 (S.L. Bonatto) (UFRG); 1 §, 1 \&, Derrubadas, 16.iv. 1983 (S.L. Bonatto) (UFRG); 1 ㅇ, Derrubadas, 17.xii. 1982 (S.L. Bonatto) (UFRG); São Paulo: $1 \delta^{\lambda}$, Picinguaba, 18.xi. 1991 (A.C. Marques) (UFRG); 2 ㅇ, Porto Cabral, iii-iv. 1944 (L.Travassos) (UFRG); 1 Q, São Paulo, 18-26.ii. 1940 (UFRG); PERU: Junin: 1 đ̃, Satipo, v. 1937 (P. Paprzyck) (UFRG); 1 §, Satipo, x. 1943 (P. Paprzyck) (UFRG); 2 \& , Satipo, xii. 1963 ( $P$. Paprzyck) (UFRG); VENEZUELA: Aragua: 1 §̉, El Limón, 30.v. 1965 (F. Fernandez)
（UFRG）； 1 ㅇ，Rancho Grande，13．ii． 1965 （F．Romero）（UFRG）； 1 §，Bolivar，Santa Elena， 13．xi． 1966 （J \＆B Bechyne，E．Osuna）（UFRG）； 1 §，no data，15．iv． 1965 （F．Fernandez） （UFRG）．

## Oenopiella punctaria（Spinola，1859）

ARGENTINA： 5 §ె， 5 ㅇ，Buenos Aires，Tandil，Buenos Aires，ii． 1960 （F．H．Walz）（RMNH）； BRAZIL：Rio Grande do Sul： 1 q，Pelotas，iv． 1991 （Piero）（UFRG）； 1 §，Pelotas（UFRG）； CHILE： 1 §，Santiago，Rangue，13．xii． 1983 （G．Arriagada）（MNNC）； 1 §， 2 ㅇ，Aisen del General Carlos Ibanez del Campo，Aysen，12．ii．1960－61（L．Pena）（AMNH）；URUGUAY： 1 ठ， 2 个，Montevideo，Montevideo，1．i． 1953 （C．Biezanko）（AMNH）； 1 §，Colonia，La Estanzuela，16．i． 1953 （C．Biezanko）（AMNH）．

## Oenopiella unidentata（Stål，1859）

CHILE： 1 §，Atacama，Copiapó，22．vi． 1955 （L．E．Pena）（AMNH）；Valparaiso： 1 〕，Concon， 13．iii． 1964 （AMNH）； 6 万̉， 1 q，El Convento，San Antonio，4．xii． 1964 （L．E．Pena）（AMNH）； Santiago： 2 万，Apoquindo，vi． 1953 （L．E．Pena）（AMNH）； 3 \＆，Apoquindo，vii． 1954 （L．E． Pena）（AMNH）； 1 ¢，La Rinconada，19．i． 1956 （R．H．González）（UFRG）； 1 ㅇ，Maipu， 2．v． 1961 （AMNH）；Biobio： 4 \＆，Río Chirihuillin，9．ii． 1965 （T．Cekalovic）（MZSP）．

## Padaeus trivitatus Stål， 1872

MEXICO： 1 \＆，Distrito Federal，Pedregal de San Ángel，13．viii． 1969 （Diaz Batrez） （MHNM）；Mexico： 1 ठ，Coatlinchan，29．vi． 1968 （M．E．Díaz B．）（MHNM）； 1 ठ，Villa de Allende，06．vii． 1911 （M．A．Morón）（MHNM）； 1 §̂，Michoacan de Ocampo，Patzcuaro， 18．vi． 1968 （G．Halffier）（UFRG）； 1 ठ＇，Morelos，Tepoztlán，13．xi． 1971 （A．Espinoza） （MHNM）．

## Padaeus viduus（Vollenhoven，1868）

$1 \delta^{\lambda}$ ，COSTA RICA：San Rafael，10．ii．1935（ MIZA）； 1 §， 1 ㅇ，MEXICO：Mexico， Tejupilco，09．xii． 1973 （A．Barrera）（MHNM）．

## Pedinonotus catarinensis Fernandes \＆Grazia， 2002

Holotype §＇，BRAZIL：Santa Catarina，Nova Teutônia，xii． 1939 （F．Plaumann）（AMNH）． Paratypes： 1 ठ， 1 ㅇ，BRAZIL：Santa Catarina，Nova Teutônia，ix． 1939 （F．Plaumann） （AMNH，UFRG）； 1 ㅇ，BRAZIL：Santa Catarina，Corupá，x． 1953 （A．Maller）（UFRG）．

Additional analyzed material. BRAZIL: Santa Catarina: 2 §, 1 q, Nova Teutônia, ix. 1939 (F. Plaumann) (AMNH); 1 \&, Nova Teutônia, xii. 1939 (F. Plaumann) (AMNH); 1 ¢, Nova Teutônia, 29.iv. 1966 (F. Plaumann) (AMNH); 1 ô, 1 \& , Nova Teutônia, v. 1967 (F. Plaumann) (AMNH); 1 ㅇ, Nova Teutônia, x. 1969 (F. Plaumann) (AMNH); 1 đ, Nova Teutônia, x. 1974 (F. Plaumann) (AMNH); 1 §, Nova Teutônia, xi. 1975 (F. Plaumann) (AMNH).

## Prionosoma podopioides Uhler, 1863

UNITED STATES: 1 \&, Arizona, Douglas, 21.viii. 1932 (USNM); 1 ㅇ, California, Hammil, 19.vii. 1952 (W.J. Gertsch \& R. Schrammel) (UFRG); 1 q, Colorado (USNM); 1 入, Iowa, Iowa, 30.x. 1945 (USNM); $1 \delta^{\lambda,}$ Oklahoma, Fort Sill, $29.1 i i .1954$ (J.C. Schaffner) (DARC); 1 đ, Texas, vii. 1917 (USNM).

## Proxys albopunctulatus (Palisot, 1811)

BRAZIL: 1 đ̋, Amazonas, Itacoatiara, 19.vi. 1952 (O.M. Rego) (MNRJ); Espírito Santo: 1 q, Linhares, 02-07.viii. 1974 (C. Elias) (DZUP); 1 \&, Linhares, 09-15.i. 1975 (C. Elias) (DZUP); 2 §̃, Linhares (D. Zajclw) (MNRJ); 1 ¢, Minas Gerais, Araxá, 22.iv. 1965 (C. Elias) (DZUP); $1 \jmath^{\lambda, 1} 1$ Q, Paraná, Maringá, $07 . x i i .1965$ (V. Graf \& L. Azevedo) (DZUP); 2 § , Rio Grande do Norte, Natal, v. 1950 (M. Alvarenga) (MNRJ); Rio Grande do Sul: 1 q, Derrubadas, 11.x. 1981 (S.L. Bonatto) (UFRG); 1 ¢, Derrubadas, 22.xii. 1982 (S.L. Bonatto) (UFRG); 1 \&, Derrubadas, 24-25.v. 1984 (S.L. Bonatto) (UFRG); 1 §, Triunfo, 15.ix. 1977 (M.H. Galileo) (MCNZ); COLOMBIA: Bolivar: $1 \jmath^{\lambda}$, Monterrey (F. Fernandez \& G. Ulloa) (UFRG); 1 q, Totumito, 09.xii. 1993 (F. Fernandez) (UFRG); UNITED STATES: 1 q, Florida, Silver Springs, 04.vi. 1969 (J. Slater, T.Schuh \& J. Harrington) (AMNH); VENEZUELA: Aragua: 1 O, Cagua, 28.v. 1958 (A. Fernandez) (UFRG); 1 ठ, 1 O, El Limón, 31.v. 1957 (F. Fernandes \& L. Salas) (UFRG).

## Proxys victor (Fabricius, 1775)

1 §, COLOMBIA, Valle del Cauca, Bajo Calima, 22.iii. 1995 (UFRG); 1 đ, ECUADOR, S. Domingos, ix. 1962 (J.C.M. Carvalho) (UFRG); 1 §, PUERTO RICO, 1967 (N. Virkki); VENEZUELA: Aragua: 2 q, Cagua, 28.v. 1958 (A. Fernandez) (UFRG); 1 §, El Limón, 12.v. 1963 (UFRG); 1 \&, Maracay, 24.v. 1950 (F. Fernandez) (UFRG); 1 q, Barinas, Barinas, 4.ix. 1949 (F.A. Ponte) (UFRG); 1 §, 4 \&, Cojedes, San Carlos, $24 . v .1948$ (F. Fernandes \& L. Salas) (UFRG); 2 §, Distrito Federal, El Valle, 14.VII. 1942 (UFRG); 1 q, Monagas, Jusepin, 07.x. 1965 (F. Fernandes \& C.J Rosales) (UFRG); 1 §, no data, 13.iv. 1960 (UFRG).

## Spinalanx monstrabilis Rolston \& Rider, 1988

BRAZIL: Pará: 1 Q, Belém, 07-30.iv. 1966 (Malkin) (UFRG); 1 §, Itaituba, ii. 1965 (Dirings) (UFRG).

## Spinalanx rolstoni Thomas, 1995

BRAZIL: Rio Grande do Sul: 1 ㅇ, São Francisco de Paula, 17-19.xii. 2005 (P.R.S. Bunde) (UFRG); 1 §, São Francisco de Paula, 16.iii. 2008 (L.M. Weiler) (UFRG); 1 §, São Francisco de Paula, 18.xii. 2010 (F.M. Bianchi) (UFRG)

## Tibraca limbativentris Stål, 1860

1 §, BRAZIL: Espírito Santo, Santa Tereza, 19.x. 1964 (C. Elias) (DZUP); 1 §, Maranhão, São Luís, 04.11974 (E. Chagas) (UFRG); 1 q, Mato Grosso do Sul, Rio Brilhante, ii-iii. 1995 (UFRG); Rio Grande do Sul: 1 入, Capão do Leão, ix. 1995 (L. Rodrigues) (UFRG); 1 q, Guaíba, 04.vii. 1985 (E. Albuquerque) (UFRG); 1 §, Pelotas, xii. 1993 (J. F. Fonseca) (UFRG); 1 +,Santa Maria, 29.iv. 1988 (Equipe DFS) (UFRG); 1 ㅇ, Santa Catarina, Lages, viii. 1988 (H. Kalvelage) (UFRG); Tocantins: 1 \&, Gurupi, 01.vi. 1988 (K. Kishino) (UFRG); 1 §, Gurupi, 24.v. 1989 (K. Kishino) (UFRG); 1 \&, COLOMBIA: Valle del Cauca, Palmira, 10.x. 1989 (A. Baena) (DARC); 1 \& COSTA RICA: Heredia, Sarapiqui, 24.vi. 1963 (T.R. Everett) (DARC); 1 §, 1 ㅇ, DOMINICAN REPUBLIC: 13.xi. 1997 (De Benitez) (UFRG); PERU: 1 §, 1 \&, Cajamarca, Bagua, 10-15.i. 1954 (W. Ebeling) (DARC); 1 §, Junin, Satipo, ii. 1938 (P. Paprzyck) (FIOC); 1 §, Loreto, Yurimaguas, 27.x. 1983 (M. Calderon) (DARC); VENEZUELA: 1 §̉, Portuguesa, Payara, 14.viii. 1957 (M. Angeles) (MIZA); 1 ㅇ, v. 1948 (MIZA).

## Tibraca simillima Barber, 1941

Paratypes: 1 ठ, 1 ㅇ, ECUADOR (AMNH). Additional analyzed specimens. ECUADOR: 6 §, 4 ? , Azuay, Cuenca, 10.iii. 1963 (J. Davis) (AMNH); 2 §, 5 q, Portoviejo (F. Campos) (FIOC); 1 §, no data (UFRG).

## Appendix S2: Complete dataset: 10 continous characters and 139 discrete characters.



DDpunctatus DDleucostigmus DDbicolor DDnigrum DDavilapiresi DDmiriamae DDperuanus DDpradoi DDsaltensis DDaustralis Ddfurcatus Ddmelacanthus Ddphoenix Ddlobatus Ddcaatinguensis DPdivisus Aalbocostata Afraterna Agriseus Ascabricornis Bhastator Cpseudoscylax Cfasciatus Cpurpureipennis Cdelia
Cdecorata
Clintneriana
Dbaccarum EMacutus
EEemoorei EMgrandis
EEheros
ELsharpi
ELtriangulator EEvariolarius Gbituberculata Gadroguensis
Gsetigera
Hnervosa
Hinermis
Hsternalis
Larmipes
Lbilobus

## Macuminata

Minsertus
Mcornicolis
Mmaculata
Mypsilon
Opunctaria
Ounidentata
Ptrivittatus
Pviduus
Pcatarinensis
Ppodopioides
Palbopunctulatus
Pvictor
Smonstrabilis
Srolstoni
Tlimbativentris
Tsimilima
$\begin{array}{llllll}1 & 2 & 3 & 4 & 5 & 6\end{array}$

| 0 | 0 | 0 | 0 | 0 | 0 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |

$211110010001021100011011011110-0--101111112111101110-0011100$ $211110010001021100011011011110-0--10111111111111011110-0011100$
 $212110010001021100011011011110-0-1011111112111101111110010100$ $211110010001021101011011111110-0-201111111111110111110010100$ $212110010001021101011011011110-0-2001111112111101111110010100$ $212110010001021100011011011110-0-2011111111121101110-0010100$ $211110010001021100011011111110-0--100111111111110111110010100$ $211110110001021101011011111110-0-20111111111110111110010100$ $211110010001021101011011111110-0--2011111211111101111110000100$ 211110110001021110010011011011010120111112211010111100010100 211110110001021100010011011011010120111112211010111100010100 21111011000102111001001101101101012011111221 ? 0101111100010100 $201110110001011100010011001010-101201111122110010111100010100$ $201110000000021100010011001010-1002011111122110101111100010100$ $20111010000102110001101200101100-10011$ ? 1 ? ? 111 ? 01 ? ? 111010100 $201001010001001100010102100010-0--201111122110101110-00000-0$ 101-01000001011110010102100010-101001111122110101110-00000-0 111-10010000121101011001101010-1101011121102101101110-1000110 111-100100001211010110110110110110101112102101101110-1011110
 $102-01010001011100010010011010-0--201111110211010111110011100$ $101-0001000012010001001010001110--000000101110010100110000101$ 101-10010011021010010100100010-0-0001001?110010100100000101 $100-10000112011110010000100000-0--011122121000100111111000110$ $101-0001000212110000-00010001101002001111$ ? $11110100010-00000-0$ $101-0001000212100000-00010001101002001111$ ? $11011000010-00000-0$ $201110000001021010010120100010-1000001121$ ? 1110011110100000101 $201100010000011100111012010010-0--1001101221$ ? 0110 ? ? 1110010100 101-10110000011100010001010010-101200111122100101111001000-0 $201100000000121100011011021010-0--200111110211010111110010110$ $001-10110000021100010011010010-100000111122110110111100101100$ $101-00110000021110010001101010-0-0001111102110101110-0000110$ $001-10011000011110011011011010-10011111111021$ ? 1110111110011100 101-10110000121100010011010010-0--20011211211010111110100100 $200001010000110100010011021010-0--2111131102101111011110000110$ $000-12010120011100010010010010-0--20012200210100101100010110$ $000-12010120011100011010010010-0--200122002101001011100000110$ 101-10110102011110010020100010-10121111112210110011100100110 $101-00010000001111110002100100-0--10101211211110001110010110$ $201001010000001111110001100100-0--101011111111110001100010110$ $101-00110000121100110002100010-0--100110122100101110-1000100$ $101-00010000121100111011010010-0--10011012210010011111010100$ $000-02000120020000010100000000-0-1011111121110101110$ $101-00110000020000010120100000-0--211111121111100111110100100$ $001-00010000111110010010110010-0--200111000210110001110010100$ $001-10010000111110010010110010-100100111122101100011100100-0$ $001-10010000121110010010110010-100100111122100100011100100-0$ 101-02010101011110010120100000-0--20102112201100111100001100 101-02010101011110010120100000-0--20102112201110111100001100 101-00110001020100011010011010-0--10111111110 ? 010101110011100 101-00110001020100011010011010-0--101111111110110101110011100 $200111010100011101110102100000-0--0011101221111010101110000110$ $211102010102011100110010020010-0--2010220020$ ? $00101000-0000110$ 010-02011001021100011010010010-100101111111111110001110011100 $010-02011001021100011010010010-100101111121111110001110011100$ $000-00000100121100010011011010-0-101122002111100111111010110$ $011-00010100121111010011011010-0--10112300211110201110010110$ $000-02000120021000010010010010-0--101111121111010101100010111$ $000-02000120021000010010100010-0--10111012111010101100000111$



## Appendix S3: Defining $\boldsymbol{k}$-value

Results of explored parameters under Implied Weighting using Mirande (2009) protocol. Parameter used were default, except for distortion groups (24), hits of best trees (5), and drift and ratchet iterations (10).

| Runs | distref | kref | length | trees | fit |
| :--- | :---: | ---: | ---: | ---: | ---: |
| k0 | 50.000 | 4.165 | 898 | 3 | 65.745 |
| k1 | 51.739 | 4.465 | 898 | 3 | 63.826 |
| k2 | 53.478 | 4.787 | 898 | 3 | 61.897 |
| k3 | 55.217 | 5.135 | 898 | 3 | 59.957 |
| k4 | 56.957 | 5.511 | 898 | 3 | 58.003 |
| k5 | 58.696 | 5.918 | 893 | 3 | 56.028 |
| k6 | 60.435 | 6.361 | 893 | 3 | 54.032 |
| k7 | 62.174 | 6.845 | 890 | 3 | 52.017 |
| k8 | 63.913 | 7.376 | 885 | 1 | 49.978 |
| k9 | 65.652 | 7.96 | 885 | 1 | 47.911 |
| k10 | 67.391 | 8.607 | 885 | 1 | 45.823 |
| k11 | 69.130 | 9.327 | 884 | 1 | 43.71 |
| k12 | 70.870 | 10.132 | 884 | 1 | 41.573 |
| k13 | 72.609 | 11.04 | 884 | 1 | 39.411 |
| k14 | 74.348 | 12.071 | 884 | 1 | 37.222 |
| k15 | 76.087 | 13.251 | 884 | 1 | 35.003 |
| k16 | 77.826 | 14.617 | 884 | 1 | 32.753 |
| k17 | 79.565 | 16.216 | 884 | 1 | 30.468 |
| k18 | 81.304 | 18.112 | 884 | 1 | 28.147 |
| k19 | 83.043 | 20.396 | 884 | 1 | 25.787 |
| k20 | 84.783 | 23.203 | 878 | 1 | 23.372 |
| k21 | 86.522 | 26.735 | 878 | 1 | 20.911 |
| k22 | 88.261 | 31.312 | 878 | 1 | 18.404 |
| k23 | 90.000 | 37.482 | 878 | 1 | 15.848 |

Comparison matrix of tree consensuses topologies found for each run of Mirande's script. Values are Coefficient Distances: the higher the index, more similar the trees. Range of $k 11-19$ was found to be the $k$ 's best adjusted to the matrix. $k$-value defined to perform Implied

Table S1. List of terminals included on the analysis, with information on the genitalia studied per species and sex (species marked with "*" only data on external genitalia was available; species marked with "L", data on internal genitalia was retrieved from literature), the availability of SEM of external scent efferent system of the metasternal glands (ESES), number of specimens measured per species and sex, and literature used for species determination when available.

| Species | Genitalia |  | ESES SEM | \# spec measured |  |  | Literature for determination |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | § | ¢ |  | Total | ठ | + |  |
| Acledra albocostata (Spinola, 1852) | X | X | X | 20 | 10 | 10 | Faúndez \& Verdejo (2009) |
| Acledra fraterna (Stål, 1859) | X | X | X | 18 | 8 | 10 | Faúndez \& Verdejo (2009) |
| Agroecus griseus Dallas, 1851 | X | X | X | 20 | 10 | 10 | Rider \& Rolston (1987) |
| Agroecus scabricornis (Herrich-Schäffer, 1844) | X | X | X | 13 | 5 | 8 | Rider \& Rolston (1987) |
| Berecynthus hastator (Fabricius, 1798) | X | X | X | 20 | 10 | 10 | Grazia \& Hildebrand (1982) |
| Caonabo pseudoscylax (Bergroth, 1891) | X | X | X | 19 | 9 | 10 | Rolston (1974) |
| Caribo fasciatus Rolston, 1984 | X | X | X | 12 | 9 | 3 | Rolston \& McDonald (1984) |
| Carpocoris purpureipennis (DeGeer, 1783) | X | X | X | 6 | 3 | 3 |  |
| Coenus delius (Say, 1832) | X | X | X | 7 | 3 | 4 | Rider (1995) |
| Cosmopepla decorata (Hahn, 1834) | X | X | X | 4 | 2 | 2 | McDonald (1986) |
| Cosmopepla lintneriana Kirkaldy, 1909 | X | X | X | 8 | 3 | 5 | McDonald (1986) |
| Dichelops (Diceareus) caatiguensis Grazia \& Poock-da-Silva, 2013 | X | X | X | 16 | 6 | 10 | Poock-da-Silva et al. (2013) |
| Dichelops (Diceareus) furcatus (Fabricius, 1775) | X | X | X | 20 | 10 | 10 | Grazia (1978), Poock-da-Silva et al. (2013) |
| Dichelops (Diceareus) lobatus Grazia, 1978 | X | X | X | 4 | 2 | 2 | Grazia (1978), Poock-da-Silva et al. (2013) |
| Dichelops (Diceareus) melacanthus (Dallas, 1851) | X | X | X | 20 | 10 | 10 | Grazia (1978), Poock-da-Silva et al. (2013) |
| Dichelops (Diceareus) phoenix Grazia, 1978 | X | X | X | 9 | 6 | 3 | Grazia (1978), Poock-da-Silva et al. (2013) |
| Dichelops (Dichelops) australis Grazia \& Klein, 2012 | X | X | X | 11 | 8 | 3 | Klein et al. (2012) |
| Dichelops (Dichelops) avilapiresi Grazia, 1978 | X | X | X | 9 | 3 | 6 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) bicolor Distant, 1890 | X | X | X | 6 | 4 | 2 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) leucostigmus (Dallas, 1851) | X | X | X | 20 | 10 | 10 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) miriamae Grazia, 1978 | X | X | X | 8 | 5 | 3 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) nigrum Bergroth, 1914 | X | X | X | 16 | 7 | 9 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) peruanus Grazia, 1978 | $\mathrm{X}^{\text {L }}$ | X | X | 11 | 2 | 9 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) pradoi Grazia, 1978 | $\mathrm{X}^{\mathrm{L}}$ | X | X | 4 | 2 | 2 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) punctatus (Spinola, 1837) | $\mathrm{X}^{\text {L }}$ | X | X | 5 | 2 | 3 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Dichelops) saltensis Grazia, 1978 | X | X | X | 17 | 7 | 10 | Grazia (1978), Klein et al. (2012) |
| Dichelops (Prodichelops) divisus (Walker, 1867) | X* | X* | - | 3 | 1 | 2 | Grazia (1978), Poock-da-Silva et al. (2013) |
| Dolycoris baccarum (Linnaeus, 1758) | X | X | X | 6 | 4 | 2 |  |
| Euschistus (Euschistus) emoorei Rolston, 1972 | X | X | X | 18 | 10 | 8 | Rolston (1974) |
| Euschistus (Euschistus) heros (Fabricius, 1794) | X | X | X | 20 | 10 | 10 | Rolston (1974) |


| Euschistus (Euschistus) variolarius (Palisot de Beauvois, 1817) | X | X | X | 7 | 3 | 4 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Euschistus (Lycipta) sharpi (Bergroth, 1891) | X | X | X | 17 | 7 | 10 | Rolston (1982) |
| Euschistus (Lycipta) triangulator (Herrich-Schäffer, 1842) | X | X | X | 12 | 8 | 4 | Rolston (1982) |
| Euschistus (Mitripus) acutus Dallas, 1851 | X | X | X | 20 | 10 | 10 | Rolston (1978a) |
| Euschistus (Mitripus) grandis Rolston, 1978 | X | X | X | 20 | 10 | 10 | Rolston (1978a) |
| Galedanta bituberculata Amyot \& Serville, 1843 | X | X | X | 15 | 5 | 10 | Grazia (1967), Grazia (1981) |
| Glyphepomis adroguensis Berg, 1891 | X | X | X | 18 | 8 | 10 | Campos \& Grazia (1998) |
| Glyphepomis setigera Kormilev \& Pirán, 1952 | X | X | X | 17 | 7 | 10 | Campos \& Grazia (1998) |
| Hymenarcis nervosa (Say, 1831) | X | X | X | 6 | 4 | 2 | Rolston (1973b) |
| Hypatropis inermis (Jensen-Haarup, 1928) | X | X | X | 14 | 4 | 10 | Fernandes \& Grazia (1996) |
| Hypatropis sternalis (Stål, 1869) | $\mathrm{X}^{\text {L }}$ | X | X | 5 | 3 | 2 | Fernandes \& Grazia (1996) |
| Ladeaschistus armipes (Stål, 1872) | X | X | X | 13 | 4 | 9 | Rolston (1973a) |
| Ladeaschistus bilobus (Stål, 1872) | X | X | X | 19 | 9 | 10 | Rolston (1973a) |
| Mecocephala acuminata Dallas, 1851 | $\mathrm{X}^{\text {L }}$ | X | X | 3 | 2 | 1 | Schwertner et al. (2002) |
| Menecles insertus (Say, 1832) | X | X | X | 7 | 4 | 3 | Rolston (1972) |
| Mormidea cornicollis (Stål, 1860) | X | X | X | 20 | 10 | 10 | Rolston (1978b) |
| Mormidea maculata Dallas, 1851 | X | X | X | 20 | 10 | 10 | Rolston (1978b) |
| Mormidea ypsilon (Linnaeus, 1758) | X | X | X | 20 | 10 | 10 | Rolston (1978b) |
| Oenopiella punctaria (Spinola, 1859) | X | X | X | 20 | 10 | 10 | Fernández-Aldea et al. (2014) |
| Oenopiella unidentata (Stål, 1859) | X | X | X | 20 | 10 | 10 | Fernández-Aldea et al. (2014) |
| Padaeus trivittatus Stål, 1872 | X | X | X | 5 | 4 | 1 |  |
| Padaeus viduus (Vollenhoven, 1868) | X | X | X | 3 | 2 | 1 |  |
| Pedinonotus catarinensis Fernandes \& Grazia, 2002 | $\mathrm{X}^{\text {L }}$ | X | X | 14 | 7 | 7 | Fernandes \& Grazia (2002) |
| Prionosoma podopioides Uhler, 1863 | X | X | X | 6 | 2 | 4 |  |
| Proxys albopunctulatus (Palisot, 1811) | X | X | X | 20 | 10 | 10 |  |
| Proxys victor (Fabricius, 1775) | X | X | X | 17 | 8 | 9 |  |
| Spinalanx monstrabilis Rolston \& Rider, 1988 | - | $\mathrm{X}^{\text {L }}$ | X | 2 | 1 | 1 | Rolston \& Rider (1988), Thomas (1995) |
| Spinalanx rolstoni Thomas, 1995 | X | X | X | 3 | 2 | 1 | Rolston \& Rider (1988), Thomas (1995) |
| Tibraca limbativentris Stål, 1860 | X | X | X | 20 | 10 | 10 | Fernandes \& Grazia (1998) |
| Tibraca similima Barber, 1941 | X | X | X | 20 | 10 | 10 | Fernandes \& Grazia (1998) |

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error (minimum - maximum)].

| Species | Measurements |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | HD | IOD | EW | LMP | IOC | PL | PWH | PWM | SWB | SL |
| Acledra albocostata | $\begin{array}{r} \hline 2.12 \pm 0.02 \\ (1.96-2.27) \end{array}$ | $\begin{array}{r} 1.53 \pm 0.02 \\ (1.42-1.65) \end{array}$ | $\begin{array}{r} \hline 0.74 \pm 0.01 \\ (0.65-0.85) \end{array}$ | $\begin{array}{r} 0.07 \\ (0.04-0.08) \end{array}$ | $\begin{array}{r} \hline 0.98 \pm 0.01 \\ (0.85-1.04) \end{array}$ | $\begin{array}{r} \hline 2.33 \pm 0.03 \\ (2.12-2.58) \end{array}$ | $\begin{array}{r} 6.03 \pm 0.05 \\ (5.69-6.38) \end{array}$ | $\begin{array}{r} 5.64 \pm 0.06 \\ (5.08-6.15) \end{array}$ | $\begin{gathered} 3.51 \pm 0.04 \\ (3.23-3.92) \end{gathered}$ | $\begin{array}{r} 3.95 \pm 0.05 \\ (3.58-4.35) \end{array}$ |
| Acledra fraterna | $\begin{aligned} & 1.52 \pm 0.02 \\ & (1.27-1.69) \end{aligned}$ | $\begin{aligned} & 1.28 \pm 0.01 \\ & (1.15-1.38) \end{aligned}$ | $\begin{aligned} & 0.67 \pm 0.01 \\ & (0.61-0.81) \end{aligned}$ | $\begin{aligned} & 0.35 \pm 0.02 \\ & (0.23-0.46) \end{aligned}$ | $\begin{aligned} & 0.83 \pm 0.02 \\ & (0.69-0.92) \end{aligned}$ | $\begin{gathered} 2.15 \pm 0.03 \\ (1.96-2.42) \end{gathered}$ | $\begin{aligned} & 5.87 \pm 0.07 \\ & (5.38-6.46) \end{aligned}$ | $\begin{aligned} & 5.42 \pm 0.07 \\ & (4.88-5.85) \end{aligned}$ | $\begin{aligned} & 3.38 \pm 0.04 \\ & (3.12-3.73) \end{aligned}$ | $\begin{aligned} & 3.55 \pm 0.05 \\ & (3.19-3.92) \end{aligned}$ |
| Agroecus griseus | $\begin{aligned} & 1.93 \pm 0.03 \\ & (1.73-2.15) \end{aligned}$ | $\begin{aligned} & 1.27 \pm 0.02 \\ & (1.19-1.42) \end{aligned}$ | $\begin{aligned} & 0.97 \pm 0.02 \\ & (0.85-1.15) \end{aligned}$ |  | $\begin{aligned} & 0.88 \pm 0.02 \\ & (0.77-1.04) \end{aligned}$ | $\begin{aligned} & 2.38 \pm 0.05 \\ & (2.04-2.73) \end{aligned}$ | $\begin{aligned} & 6.13 \pm 0.11 \\ & (5.42-6.92) \end{aligned}$ | $\begin{aligned} & 5.85 \pm 0.10 \\ & (5.08-6.62) \end{aligned}$ | $\begin{aligned} & 3.86 \pm 0.07 \\ & (3.42-4.42) \end{aligned}$ | $\begin{aligned} & 3.70 \pm 0.08 \\ & (3.08-4.31) \end{aligned}$ |
| Agroecus scabricornis | $\begin{aligned} & 2.00 \pm 0.03 \\ & (1.88-2.31) \end{aligned}$ | $\begin{aligned} & 1.28 \pm 0.02 \\ & (1.15-1.38) \end{aligned}$ | $\begin{aligned} & 0.88 \pm 0.01 \\ & (0.80-0.97) \end{aligned}$ |  | $\begin{aligned} & 0.79 \pm 0.01 \\ & (0.69-0.85) \end{aligned}$ | $\begin{aligned} & 2.37 \pm 0.05 \\ & (1.92-2.62) \end{aligned}$ | $\begin{aligned} & 7.35 \pm 0.15 \\ & (6.54-8.27) \end{aligned}$ | $\begin{aligned} & 5.87 \pm 0.11 \\ & (5.00-6.35) \end{aligned}$ | $\begin{aligned} & 3.92 \pm 0.07 \\ & (3.50-4.35) \end{aligned}$ | $\begin{aligned} & 3.66 \pm 0.09 \\ & (3.19-4.04) \end{aligned}$ |
| Berecynthus hastator | $\begin{aligned} & 2.15 \pm 0.02 \\ & (2.04-2.31) \end{aligned}$ | $\begin{aligned} & 1.28 \pm 0.01 \\ & (1.19-1.38) \end{aligned}$ | $\begin{aligned} & 0.80 \pm 0.02 \\ & (0.42-0.88) \end{aligned}$ |  | $\begin{aligned} & 0.77 \pm 0.01 \\ & (0.69-0.81) \end{aligned}$ | $\begin{aligned} & 1.90 \pm 0.03 \\ & (1.77-2.23) \end{aligned}$ | $\begin{aligned} & 5.69 \pm 0.13 \\ & (5.00-7.12) \end{aligned}$ | $\begin{aligned} & 4.51 \pm 0.05 \\ & (4.12-4.92) \end{aligned}$ | $\begin{aligned} & 2.92 \pm 0.03 \\ & (2.73-3.19) \end{aligned}$ | $\begin{aligned} & 3.04 \pm 0.04 \\ & (2.81-3.38) \end{aligned}$ |
| Caonabo pseudocylax | $\begin{aligned} & 1.80 \pm 0.02 \\ & (1.65-2.04) \end{aligned}$ | $\begin{aligned} & 1.13 \pm 0.01 \\ & (1.04-1.19) \end{aligned}$ | $\begin{aligned} & 0.62 \pm 0.01 \\ & (0.54-0.70) \end{aligned}$ | - | $\begin{aligned} & 0.61 \pm 0.01 \\ & (0.54-0.69) \end{aligned}$ | $\begin{aligned} & 1.90 \pm 0.03 \\ & (1.69-2.15) \end{aligned}$ | $\begin{aligned} & 5.65 \pm 0.09 \\ & (5.00-6.23) \end{aligned}$ | $\begin{aligned} & 4.62 \pm 0.07 \\ & (4.04-5.12) \end{aligned}$ | $\begin{aligned} & 2.95 \pm 0.06 \\ & (2.46-3.31) \end{aligned}$ | $\begin{aligned} & 3.05 \pm 0.06 \\ & (2.58-3.42) \end{aligned}$ |
| Caribo fasciatus | $\begin{aligned} & 1.62 \pm 0.02 \\ & (1.52-1.72) \end{aligned}$ | $\begin{aligned} & 1.06 \pm 0.02 \\ & (0.96-1.16) \end{aligned}$ | $\begin{aligned} & 0.81 \pm 0.04 \\ & (0.64-1.04) \end{aligned}$ | - | $\begin{aligned} & 0.79 \pm 0.02 \\ & (0.72-0.88) \end{aligned}$ | $\begin{aligned} & 1.49 \pm 0.03 \\ & (1.36-1.60) \end{aligned}$ | $\begin{aligned} & 4.85 \pm 0.09 \\ & (4.56-5.36) \end{aligned}$ | $\begin{aligned} & 4.60 \pm 0.08 \\ & (4.24-5.00) \end{aligned}$ | $\begin{aligned} & 2.99 \pm 0.05 \\ & (2.80-3.28) \end{aligned}$ | $\begin{aligned} & 2.80 \pm 0.07 \\ & (2.56-3.16) \end{aligned}$ |
| Carpocoris purpureipennis | $\begin{aligned} & 2.20 \pm 0.02 \\ & (2.14-2.24) \end{aligned}$ | $\begin{aligned} & 1.41 \pm 0.03 \\ & (1.33-1.48) \end{aligned}$ | $\begin{aligned} & 0.93 \pm 0.04 \\ & (0.81-1.00) \end{aligned}$ | - | $\begin{aligned} & 0.87 \pm 0.03 \\ & (0.76-0.95) \end{aligned}$ | $\begin{aligned} & 2.38 \pm 0.06 \\ & (2.19-2.52) \end{aligned}$ | $\begin{aligned} & 6.91 \pm 0.15 \\ & (6.38-7.24) \end{aligned}$ | $\begin{aligned} & 6.33 \pm 0.12 \\ & (5.90-6.62) \end{aligned}$ | $\begin{aligned} & 4.07 \pm 0.08 \\ & (3.76-4.29) \end{aligned}$ | $\begin{aligned} & 4.32 \pm 0.14 \\ & (3.86-4.86) \end{aligned}$ |
| Coenus delia | $\begin{aligned} & 2.02 \pm 0.03 \\ & (1.92-2.15) \end{aligned}$ | $\begin{aligned} & 1.56 \pm 0.03 \\ & (1.46-1.65) \end{aligned}$ | $\begin{aligned} & 0.69 \pm 0.01 \\ & (0.65-0.73) \end{aligned}$ | - | $\begin{aligned} & 0.95 \pm 0.01 \\ & (0.92-1.00) \end{aligned}$ | $\begin{aligned} & 2.08 \pm 0.03 \\ & (2.00-2.19) \end{aligned}$ | $\begin{aligned} & 4.62 \pm 0.09 \\ & (4.31-5.00) \end{aligned}$ | $\begin{aligned} & 4.56 \pm 0.09 \\ & (4.23-5.00) \end{aligned}$ | $\begin{aligned} & 3.13 \pm 0.07 \\ & (2.88-3.42) \end{aligned}$ | $\begin{aligned} & 3.43 \pm 0.10 \\ & (3.08-3.85) \end{aligned}$ |
| Cosmopepla decorata | $\begin{aligned} & 1.30 \pm 0.06 \\ & (1.15-1.42) \end{aligned}$ | $\begin{aligned} & 1.07 \pm 0.03 \\ & (1.00-1.12) \end{aligned}$ | $\begin{aligned} & 0.60 \pm 0.01 \\ & (0.58-0.61) \end{aligned}$ | - | $\begin{aligned} & 0.67 \pm 0.01 \\ & (0.65-0.69) \end{aligned}$ | $\begin{aligned} & 1.42 \pm 0.08 \\ & (1.27-1.65) \end{aligned}$ | $\begin{aligned} & 4.16 \pm 0.21 \\ & (3.69-4.54) \end{aligned}$ | $\begin{aligned} & 4.07 \pm 0.20 \\ & (3.62-4.42) \end{aligned}$ | $\begin{aligned} & 2.61 \pm 0.16 \\ & (2.23-2.92) \end{aligned}$ | $\begin{aligned} & 2.72 \pm 0.08 \\ & (2.54-2.85) \end{aligned}$ |
| Cosmopepla lintneriana | $\begin{aligned} & 1.24 \pm 0.03 \\ & (1.15-1.38) \end{aligned}$ | $\begin{aligned} & 1.02 \pm 0.02 \\ & (0.94-1.09) \end{aligned}$ | $\begin{aligned} & 0.47 \pm 0.02 \\ & (0.41-0.56) \end{aligned}$ | - ${ }^{-}$ | $\begin{aligned} & 0.61 \pm 0.01 \\ & (0.56-0.68) \end{aligned}$ | $\begin{aligned} & 1.41 \pm 0.03 \\ & (1.26-1.50) \end{aligned}$ | $\begin{aligned} & 3.65 \pm 0.09 \\ & (3.32-4.06) \end{aligned}$ | $\begin{aligned} & 3.61 \pm 0.09 \\ & (3.29-4.00) \end{aligned}$ | $\begin{aligned} & 2.35 \pm 0.07 \\ & (2.18-2.71) \end{aligned}$ | $\begin{aligned} & 2.21 \pm 0.06 \\ & (2.00-2.44) \end{aligned}$ |
| Dichelops (Dichelops) australis | $\begin{aligned} & 1.99 \pm 0.02 \\ & (1.88-2.08) \end{aligned}$ | $\begin{aligned} & 1.35 \pm 0.01 \\ & (1.31-1.38) \end{aligned}$ | $\begin{aligned} & 0.85 \pm 0.01 \\ & (0.77-0.92) \end{aligned}$ | $\begin{aligned} & 0.18 \pm 0.01 \\ & (0.12-0.23) \end{aligned}$ | $\begin{aligned} & 0.86 \pm 0.01 \\ & (0.81-0.92) \end{aligned}$ | $\begin{aligned} & 2.38 \pm 0.05 \\ & (2.19-2.62) \end{aligned}$ | $\begin{aligned} & 6.04 \pm 0.09 \\ & (5.69-6.62) \end{aligned}$ | $\begin{aligned} & 5.86 \pm 0.07 \\ & (5.54-6.15) \end{aligned}$ | $\begin{aligned} & 3.76 \pm 0.05 \\ & (3.54-4.04) \end{aligned}$ | $\begin{aligned} & 3.92 \pm 0.04 \\ & (3.77-4.23) \end{aligned}$ |
| Dichelops (Dichelops) avilapiresi | $\begin{aligned} & 1.75 \pm 0.09 \\ & (1.08-1.88) \end{aligned}$ | $\begin{aligned} & 1.16 \pm 0.01 \\ & (1.12-1.19) \end{aligned}$ | $\begin{aligned} & 0.77 \pm 0.01 \\ & (0.70-0.81) \end{aligned}$ | $\begin{aligned} & 0.09 \pm 0.01 \\ & (0.04-0.15) \end{aligned}$ | $\begin{aligned} & 0.76 \pm 0.01 \\ & (0.73-0.85) \end{aligned}$ | $\begin{aligned} & 2.20 \pm 0.06 \\ & (2.00-2.62) \end{aligned}$ | $\begin{aligned} & 5.49 \pm 0.06 \\ & (5.23-5.85) \end{aligned}$ | $\begin{aligned} & 5.09 \pm 0.05 \\ & (4.92-5.38) \end{aligned}$ | $\begin{aligned} & 3.26 \pm 0.05 \\ & (3.08-3.46) \end{aligned}$ | $\begin{aligned} & 3.37 \pm 0.06 \\ & (3.12-3.73) \end{aligned}$ |
| Dichelops (Dichelops) bicolor | $\begin{aligned} & 2.09 \pm 0.03 \\ & (2.00-2.21) \end{aligned}$ | $\begin{array}{r} 1.3 \pm 0.03 \\ (1.23-1.42) \end{array}$ | $\begin{aligned} & 0.83 \pm 0.02 \\ & (0.77-0.89) \end{aligned}$ | $\begin{aligned} & 0.20 \pm 0.01 \\ & (0.15-0.23) \end{aligned}$ | $\begin{aligned} & 0.83 \pm 0.02 \\ & (0.77-0.92) \end{aligned}$ | $\begin{aligned} & 2.28 \pm 0.03 \\ & (2.19-2.38) \end{aligned}$ | $\begin{aligned} & 6.82 \pm 0.13 \\ & (6.23-7.15) \end{aligned}$ | $\begin{aligned} & 5.78 \pm 0.06 \\ & (5.58-5.92) \end{aligned}$ | $\begin{aligned} & 3.69 \pm 0.05 \\ & (3.58-3.85) \end{aligned}$ | $\begin{aligned} & 4.06 \pm 0.10 \\ & (3.77-4.42) \end{aligned}$ |
| Dichelops (Diceraeus) caatinguensis | $\begin{aligned} & 1.71 \pm 0.04 \\ & (1.42-2.00) \end{aligned}$ | $\begin{aligned} & 1.08 \pm 0.02 \\ & (1,00-1.20) \end{aligned}$ | $\begin{aligned} & 0.71 \pm 0.01 \\ & (0.65-0.84) \end{aligned}$ | $\begin{array}{r} 0.15 \\ (0.12-0.16) \end{array}$ | $\begin{aligned} & 0.69 \pm 0.01 \\ & (0.64-0.76) \end{aligned}$ | $\begin{aligned} & 1.83 \pm 0.04 \\ & (1.58-2.08) \end{aligned}$ | $\begin{aligned} & 5.14 \pm 0.12 \\ & (4.44-5.92) \end{aligned}$ | $\begin{aligned} & 4.55 \pm 0.08 \\ & (3.92-5.12) \end{aligned}$ | $\begin{aligned} & 2.92 \pm 0.05 \\ & (2.60-3.28) \end{aligned}$ | $\begin{aligned} & 2.77 \pm 0.06 \\ & (2.44-3.16) \end{aligned}$ |
| Dichelops (Prodichelops) divisus | $\begin{aligned} & 2.09 \pm 0.06 \\ & (1.96-2.15) \end{aligned}$ | $\begin{aligned} & 1.20 \pm 0.01 \\ & (1.19-1.23) \end{aligned}$ | $\begin{aligned} & 0.96 \pm 0.06 \\ & (0.85-1.04) \end{aligned}$ | $\begin{aligned} & 0.09 \pm 0.01 \\ & (0.08-0.12) \end{aligned}$ | $\begin{aligned} & 0.76 \pm 0.05 \\ & (0.65-0.81) \end{aligned}$ | $\begin{aligned} & 2.42 \pm 0.07 \\ & (2.31-2.54) \end{aligned}$ | $\begin{aligned} & 7.27 \pm 0.20 \\ & (6.88-7.54) \end{aligned}$ | $\begin{aligned} & 5.90 \pm 0.20 \\ & (5.54-6.23) \end{aligned}$ | $\begin{aligned} & 3.57 \pm 0.08 \\ & (3.42-3.65) \end{aligned}$ | $\begin{aligned} & 3.53 \pm 0.07 \\ & (3.38-3.62) \end{aligned}$ |
| Dichelops (Diceraeus) furcatus | $\begin{aligned} & 2.00 \pm 0.03 \\ & (1.80-2.32) \end{aligned}$ | $\begin{aligned} & 1.35 \pm 0.02 \\ & (1.12-1.48) \end{aligned}$ | $\begin{aligned} & 0.77 \pm 0.01 \\ & (0.68-0.88) \end{aligned}$ | $\begin{aligned} & 0.47 \pm 0.01 \\ & (0.40-0.60) \end{aligned}$ | $\begin{aligned} & 0.86 \pm 0.01 \\ & (0.76-0.96) \end{aligned}$ | $\begin{aligned} & 2.12 \pm 0.04 \\ & (1.76-2.36) \end{aligned}$ | $\begin{aligned} & 6.90 \pm 0.14 \\ & (6.00-8.16) \end{aligned}$ | $\begin{aligned} & 5.48 \pm 0.10 \\ & (4.80-6.08) \end{aligned}$ | $\begin{aligned} & 3.46 \pm 0.06 \\ & (2.96-3.92) \end{aligned}$ | $\begin{aligned} & 3.35 \pm 0.08 \\ & (2.36-3.84) \end{aligned}$ |
| Dichelops (Dichelops) leucostigmus | $\begin{aligned} & 1.78 \pm 0.03 \\ & (1.58-2.19) \end{aligned}$ | $\begin{aligned} & 1.15 \pm 0.01 \\ & (1.08-1.23) \end{aligned}$ | $\begin{aligned} & 0.79 \pm 0.01 \\ & (0.73-0.89) \end{aligned}$ | $\begin{aligned} & 0.34 \pm 0.01 \\ & (0.23-0.46) \end{aligned}$ | $\begin{aligned} & 0.70 \pm 0.01 \\ & (0.65-0.77) \end{aligned}$ | $\begin{aligned} & 1.98 \pm 0.04 \\ & (1.77-2.46) \end{aligned}$ | $\begin{aligned} & 6.11 \pm 0.11 \\ & (5.38-6.92) \end{aligned}$ | $\begin{aligned} & 4.94 \pm 0.05 \\ & (4.65-5.31) \end{aligned}$ | $\begin{aligned} & 3.15 \pm 0.03 \\ & (2.88-3.46) \end{aligned}$ | $\begin{aligned} & 3.14 \pm 0.04 \\ & (2.81-3.54) \end{aligned}$ |
| Dichelops (Diceraeus) lobatus | $\begin{aligned} & 1.94 \pm 0.05 \\ & (1.85-2.08) \end{aligned}$ | $\begin{aligned} & 1.35 \pm 0.03 \\ & (1.27-1.42) \end{aligned}$ | $\begin{aligned} & 0.71 \pm 0.01 \\ & (0.69-0.73) \end{aligned}$ | $\begin{aligned} & 0.26 \pm 0.02 \\ & (0.23-0.31) \end{aligned}$ | $\begin{aligned} & 0.85 \pm 0.02 \\ & (0.81-0.88) \end{aligned}$ | $\begin{aligned} & 2.04 \pm 0.04 \\ & (1.92-2.12) \end{aligned}$ | $\begin{aligned} & 5.66 \pm 0.20 \\ & (5.08-6.00) \end{aligned}$ | $\begin{aligned} & 5.22 \pm 0.13 \\ & (4.85-5.42) \end{aligned}$ | $\begin{aligned} & 3.30 \pm 0.08 \\ & (3.12-3.46) \end{aligned}$ | $\begin{aligned} & 3.14 \pm 0.08 \\ & (2.92-3.27) \end{aligned}$ |
| Dichelops (Diceraeus) melacanthus | $\begin{aligned} & 1.88 \pm 0.02 \\ & (1.68-2.08) \end{aligned}$ | $\begin{aligned} & 1.27 \pm 0.01 \\ & (1.16-1.36) \end{aligned}$ | $\begin{aligned} & 0.65 \pm 0.01 \\ & (0.60-0.72) \end{aligned}$ | $\begin{aligned} & 0.43 \pm 0.01 \\ & (0.32-0.52) \end{aligned}$ | $\begin{aligned} & 0.83 \pm 0.01 \\ & (0.72-0.92) \end{aligned}$ | $\begin{aligned} & 1.94 \pm 0.03 \\ & (1.64-2.15) \end{aligned}$ | $\begin{aligned} & 6.63 \pm 0.12 \\ & (5.77-7.60) \end{aligned}$ | $\begin{aligned} & 5.05 \pm 0.07 \\ & (4.60-5.44) \end{aligned}$ | $\begin{aligned} & 3.25 \pm 0.05 \\ & (2.84-3.50) \end{aligned}$ | $\begin{aligned} & 2.97 \pm 0.05 \\ & (2.52-3.35) \end{aligned}$ |
| Dichelops (Dichelops) miriamae | $\begin{aligned} & 1.81 \pm 0.02 \\ & (1.73-1.92) \end{aligned}$ | $\begin{aligned} & 1.20 \pm 0.02 \\ & (1.15-1.27) \end{aligned}$ | $\begin{aligned} & 0.81 \pm 0.02 \\ & (0.73-0.89) \end{aligned}$ | $\begin{gathered} 0.37 \pm 0.02 \\ (0.27-0.46) \end{gathered}$ | $\begin{aligned} & 0.79 \pm 0.02 \\ & (0.69-0.88) \end{aligned}$ | $\begin{aligned} & 2.08 \pm 0.05 \\ & (1.92-2.31) \end{aligned}$ | $\begin{aligned} & 6.03 \pm 0.11 \\ & (5.69-6.54) \end{aligned}$ | $\begin{aligned} & 4.84 \pm 0.07 \\ & (4.42-5.08) \end{aligned}$ | $\begin{aligned} & 3.18 \pm 0.05 \\ & (2.96-3.38) \end{aligned}$ | $\begin{aligned} & 3.41 \pm 0.06 \\ & (3.23-3.65) \end{aligned}$ |






| $0.77 \pm 0.01$ |
| :--- |
| $(0.69-0.85)$ |
| $0.72 \pm 0.01$ |
| $(0.65-0.77)$ |
| $0.82 \pm 0.02$ |
| $(0.73-0.88)$ |
| $0.80 \pm 0.05$ |
| $(0.67-0.92)$ |
| $0.72 \pm 0.02$ |
| $(0.65-0.77)$ |
| $0.75 \pm 0.01$ |
| $(0.69-0.84)$ |
| $0.99 \pm 0.02$ |
| $(0.92-1.08)$ |
| $0.54 \pm 0.02$ |
| $(0.48-0.81)$ |
| $0.92 \pm 0.01$ |
| $(0.83-1.00)$ |
| $0.78 \pm 0.01$ |
| $(0.65-0.88)$ |
| $0.89 \pm 0.01$ |
| $(0.75-1.00)$ |
| $0.83 \pm 0.01$ |
| $(0.75-0.92)$ |
| $0.69 \pm 0.01$ |
| $(0.65-0.76)$ |
| $1.01 \pm 0.01$ |
| $(0.96-1.08)$ |
| $1.63 \pm 0.03$ |
| $(1.46-1.96)$ |
| $0.56 \pm 0.01$ |
| $(0.50-0.62)$ |
| $0.69 \pm 0.16$ |
| $(0.47-3.18)$ |
| $0.59 \pm 0.02$ |
| $(0.54-0.65)$ |
| $0.69 \pm 0.04$ |
| $(0.50-0.92)$ |
| $0.60 \pm 0.01$ |
| $(0.58-0.62)$ |
| $0.83 \pm 0.02$ |
| $(0.71-1.06)$ |
| $0.65 \pm 0.01$ |
| $(0.57-0.76)$ |
| $0.82 \pm 0.03$ |
| $(0.76-0.88)$ |
| $0.97 \pm 0.04$ |
| $(0.82-1.12)$ |






| Dichelops (Dichelops) nigrum |
| :--- |
| Dichelops (Dichelops) peruanus |
| Dichelops (Diceraeus) phoenix |
| Dichelops (Dichelops) pradoi |
| Dichelops (Dichelops) punctatus |
| Dichelops (Dichelops) saltensis |
| Dolycoris baccarum |
| Euschistus (Mitripus) acutus |
| Euschistus (Euschistus) emoorei |
| Euschistus (Mitripus) grandis |
| Euschistus (Euschistus) heros |
| Euschistus (Lycipta) sharpi |
| Euschistus (Lycipta) triangulator |
| Euschistus (Euschistus) variolarius |
| Galedanta bituberculata |
| Glyphepomis adroguensis |
| Glyphepomis setigera |
| Hymenarcis nervosa |
| Hypatropis inermis |
| Hypatropis sternalis |
| Ladeaschistus armipes |
| Ladeaschistus bilobus |
| Mecocephala acuminata |
| Menecles insertus |


| Mormidea cornicollis | $1.85 \pm 0.02$ | $1.22 \pm 0.02$ | $0.79 \pm 0.01$ |  | $0.56 \pm 0.01$ | $1.71 \pm 0.03$ | $6.00 \pm 0.14$ | $4.65 \pm 0.06$ | $3.01 \pm 0.04$ | $3.09 \pm 0.05$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | (1.72-1.96) | (1.08-1.36) | (0.68-0.88) |  | (0.44-0.65) | (1.52-1.96) | (5.04-6.80) | (4.08-5.12) | (2.72-3.36) | (2.76-3.40) |
| Mormidea maculata | $1.54 \pm 0.01$ | $0.97 \pm 0.01$ | $0.75 \pm 0.01$ | - | $0.45 \pm 0.01$ | $1.46 \pm 0.02$ | $4.65 \pm 0.10$ | $3.83 \pm 0.05$ | $2.42 \pm 0.03$ | $2.69 \pm 0.04$ |
|  | (1.40-1.64) | (0.88-1.00) | (0.68-0.84) |  | (0.40-0.48) | (1.20-1.60) | (3.60-5.20) | (3.28-4.36) | (2.12-2.60) | (2.28-3.00) |
| Mormidea ypsilon | $1.59 \pm 0.02$ | $1.06 \pm 0.04$ | $0.63 \pm 0.13$ |  | $0.44 \pm 0.01$ | $1.44 \pm 0.03$ | $4.74 \pm 0.08$ | $3.88 \pm 0.06$ | $2.52 \pm 0.05$ | $2.85 \pm 0.05$ |
|  | (1.40-1.76) | (0.92-1.76) | (1.76-0.84) |  | (0.40-0.48) | (1.20-1.68) | (4.04-5.52) | (3.20-4.40) | (2.08-2.96) | (2.40-3.28) |
| Oenopiella punctaria | $1.61 \pm 0.03$ | $1.32 \pm 0.02$ | $0.73 \pm 0.01$ |  |  | $1.84 \pm 0.01$ | $4.74 \pm 0.04$ | $4.56 \pm 0.04$ | $2.94 \pm 0.03$ | $3.17 \pm 0.03$ |
|  | (1.38-1.82) | (1.19-1.42) | (0.59-0.81) |  |  | (1.70-1.94) | (4.43-5.14) | (4.24-4.86) | (2.71-3.12) | (2.94-3.38) |
| Oenopiella unidentata | $1.67 \pm 0.02$ | $1.30 \pm 0.01$ | $0.65 \pm 0.01$ |  |  | $1.75 \pm 0.03$ | $4.31 \pm 0.06$ | $4.17 \pm 0.07$ | $2.71 \pm 0.04$ | $2.93 \pm 0.05$ |
|  | (1.55-1.89) | (1.21-1.38) | (0.52-0.75) | - |  | (1.56-1.97) | (3.90-4.86) | (3.71-4.62) | (2.41-3.03) | (2.65-3.38) |
| Padaeus trivitatus | $2.17 \pm 0.03$ | $1.19 \pm 0.02$ | $0.87 \pm 0.02$ |  | $0.83 \pm 0.01$ | $2.11 \pm 0.05$ | $6.69 \pm 0.09$ | $5.96 \pm 0.14$ | $3.63 \pm 0.06$ | $4.11 \pm 0.10$ |
|  | (2.08-2.23) | (1.15-1.23) | (0.85-0.93) |  | (0.81-0.85) | (2.00-2.23) | (6.38-6.92) | (5.58-6.35) | (3.46-3.85) | (3.88-4.46) |
| Padaeus viduus | $2.19 \pm 0.03$ | $1.19 \pm 0.03$ | $0.83 \pm 0.08$ |  | $0.83 \pm 0.02$ | $2.28 \pm 0.11$ | $6.76 \pm 0.20$ | $5.68 \pm 0.24$ | $3.53 \pm 0.17$ | $3.96 \pm 0.12$ |
|  | (2.15-2.23) | (1.15-1.23) | (0.73-0.93) |  | (0.81-0.85) | (2.08-2.46) | (6.42-7.12) | (5.38-6.15) | (3.27-3.85) | (3.81-4.19) |
| Pedinonotus catarinensis | $1.71 \pm 0.02$ | $1.11 \pm 0.01$ | $0.74 \pm 0.01$ | $0.11 \pm 0.01$ | $0.58 \pm 0.01$ | $1.90 \pm 0.03$ | $4.47 \pm 0.05$ | $4.29 \pm 0.05$ | $2.72 \pm 0.04$ | $3.14 \pm 0.04$ |
|  | (1.57-1.86) | (1.00-1.14) | (0.66-0.81) | (0.05-0.19) | (0.48-0.67) | (1.76-2.05) | (4.14-4.71) | (3.95-4.57) | (2.43-2.86) | (2.81-3.48) |
| Prionosoma podopioides | $1.91 \pm 0.14$ | $1.39 \pm 0.04$ | $0.58 \pm 0.02$ | $0.10 \pm 0.02$ | $0.95 \pm 0.01$ | $2.01 \pm 0.04$ | $5.69 \pm 0.13$ | $5.04 \pm 0.10$ | $3.21 \pm 0.08$ | $3.86 \pm 0.14$ |
|  | (1.23-2.15) | (1.27-1.50) | (0.54-0.65) | (0.04-0.15) | (0.92-1.00) | (1.85-2.15) | (5.31-6.15) | (4.73-5.42) | (2.96-3.42) | (3.54-4.50) |
| Proxys albopunctulatus | $2.47 \pm 0.02$ | $0.92 \pm 0.01$ | $0.82 \pm 0.01$ |  | $0.61 \pm 0.01$ | $2.16 \pm 0.04$ | $7.76 \pm 0.14$ | $4.98 \pm 0.07$ | $3.10 \pm 0.05$ | $3.41 \pm 0.06$ |
|  | (2.31-2.58) | (0.81-0.96) | (0.77-0.89) |  | (0.50-0.73) | (1.69-2.38) | (6.73-9.23) | (4.19-5.46) | (2.54-3.35) | (2.46-3.73) |
| Proxys victor | $2.50 \pm 0.03$ | $0.87 \pm 0.01$ | $0.85 \pm 0.02$ |  | $0.58 \pm 0.01$ | $2.15 \pm 0.04$ | $7.96 \pm 0.13$ | $4.87 \pm 0.06$ | $3.03 \pm 0.05$ | $3.44 \pm 0.04$ |
|  | (2.27-2.69) | (0.81-0.92) | (0.74-0.96) |  | (0.54-0.62) | (1.77-2.46) | (7.15-8.85) | (4.54-5.38) | (2.65-3.46) | (3.15-3.77) |
| Spinalanx monstrabilis | $2.29 \pm 0.04$ | 1.33 | $0.89 \pm 0.03$ | - | $0.91 \pm 0.05$ | 1.9 | $5.79 \pm 0.02$ | $5.26 \pm 0.07$ | $3.43 \pm 0.10$ | $3.29 \pm 0.15$ |
|  | (2.24-2.33) |  | (0.86-0.91) |  | (0.86-0.95) |  | (5.76-5.81) | (5.19-5.33) | (3.33-3.52) | (3.14-3.43) |
| Spinalanx rolstoni | $2.47 \pm 0.05$ | $1.51 \pm 0.01$ | $1.02 \pm 0.02$ | - | $0.97 \pm 0.02$ | $1.97 \pm 0.03$ | $6.21 \pm 0.08$ | $5.41 \pm 0.07$ | $3.62 \pm 0.08$ | $3.63 \pm 0.07$ |
|  | (2.38-2.52) | (1.48-1.52) | (1.00-1.05) |  | (0.95-1.00) | (1.90-2.00) | (6.05-6.33) | (5.29-5.52) | (3.48-3.76) | (3.52-3.76) |
| Tibraca limbativentris | $2.89 \pm 0.03$ | $1.51 \pm 0.02$ | $0.94 \pm 0.01$ | - | $0.86 \pm 0.02$ | $2.67 \pm 0.05$ | $7.21 \pm 0.10$ | $6.64 \pm 0.09$ | $4.40 \pm 0.06$ | $4.88 \pm 0.07$ |
|  | (2.59-3.12) | (1.35-1.71) | (0.88-1.00) | - | (0.76-1.00) | (2.29-3.12) | (6.35-8.12) | (5.82-7.35) | (3.88-4.88) | (4.29-5.35) |
| Tibraca simillima | $2.81 \pm 0.02$ | $1.54 \pm 0.01$ | $0.95 \pm 0.01$ |  | $0.93 \pm 0.01$ | $2.66 \pm 0.03$ | $6.77 \pm 0.04$ | $6.39 \pm 0.05$ | $4.19 \pm 0.04$ | $4.54 \pm 0.06$ |
|  | (2.59-2.94) | (1.47-1.65) | (0.88-1.17) | - | (0.88-1.00) | (2.47-2.88) | (6.35-7.00) | (6.00-6.76) | (3.82-4.41) | (4.12-4.88) |

HD, head length; IOD, interocular distance; EW, eyes width; LMP, length of mandibular plates ahead of clypeus; IOC, interocellar distance; PL, pronotum lenght; PWH, pronotum width at humeral angles; PWM, pronotum width at distal margin; SWB, scutellum width at basal margin; SL, scutellum length.
Table S3. Comparison of analyses consensuses. Values above diagonal correspond to Similitud Index and below diagonal, to Robinson-
Foulds distances.

|  | A | B | C | D | E | Ac | Bc | Cc | Dc | Ec |
| :--- | ---: | ---: | :--- | ---: | ---: | ---: | ---: | ---: | ---: | ---: |
| A | - | 0.994485 | 0.963768 | 0.955908 | 0.401934 | 0.967985 | 0.965328 | 0.949821 | 0.949821 | 0.473438 |
| B | 0.004784 | - | 0.963415 | 0.955857 | 0.395442 | 0.96745 | 0.97193 | 0.953448 | 0.953448 | 0.468278 |
| C | 0.10179 | 0.099116 | - | 0.98325 | 0.385942 | 0.957219 | 0.958478 | 0.982993 | 0.982993 | 0.447761 |
| D | 0.139401 | 0.135754 | 0.050483 | - | 0.39922 | 0.949653 | 0.951096 | 0.968491 | 0.968491 | 0.456934 |
| E | 0.842857 | 0.846957 | 0.846957 | 0.846957 | - | 0.379263 | 0.376 | 0.377632 | 0.377632 | 0.872922 |
| Ac | 0.080593 | 0.080708 | 0.117547 | 0.15434 | 0.844248 | - | 0.996409 | 0.975309 | 0.975309 | 0.466872 |
| Bc | 0.082686 | 0.075887 | 0.114246 | 0.150405 | 0.846957 | 0.003489 | - | 0.976027 | 0.976027 | 0.463964 |
| Cc | 0.123763 | 0.118057 | 0.046454 | 0.087598 | 0.846957 | 0.070277 | 0.067792 | - |  | 1 |
| Dc | 0.123763 | 0.118057 | 0.046454 | 0.087598 | 0.846957 | 0.070277 | 0.067792 | 0 | - | 0.464497 |
| Ec | 0.825054 | 0.829657 | 0.829657 | 0.829657 | 0.267491 | 0.80233 | 0.805798 | 0.805798 | 0.805798 | - |



Fig. S1. Cladogram resultant of Analysis E (continuous characters only, implied weights). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 0.68842 ; CI: 0.455 ; RI: 0.786 .


Fig. S2. Strict consensus of four equally parsimonious cladograms resultant of Analysis Ac (discrete characters only, equal weights, forcing Dichelops monophyly). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. S: 884; CI: 0.192; RI: 0.621.


Fig. S3. Cladogram resultant of Analysis Bc (continuous characters only, equal weights, forcing Dichelops monophyly). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. S: 924.603; CI: 0.194; RI: 0.615.


Fig. S4. Cladogram resultant of Analysis Cc (continuous + discrete characters, implied weights, forcing Dichelops monophyly). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 33.93576; CI: 0.193; RI: 0.615 .


Fig. S5. Cladogram resultant of Analysis Dc (continuous + discrete characters, implied weights, forcing Dichelops monophyly). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 34.07655; CI: 0.191; RI: 0.617 .


Fig. S6. Cladogram resultant of Analysis Ec (continuous characters only, implied weights, forcing Dichelops monophyly). Values above and below branches represent jackknife absolute frequencies and GC values, respectively. F: 0.78547 ; CI: $0.42 ; \mathrm{RI}: 0.753$.

## ANEXO I <br> NORMAS PARA PUBLICAÇÃO:

## ANNALS OF THE ENTOMOLOGICAL SOCIETY OF AMERICA

Disponível em: http://www.entsoc.org/pubs/publish/style

## ESA Style Guide

Find everything you need to know about writing and formatting your manuscript for Annals of the ESA, Environmental Entomology, Journal of Economic Entomology, Journal of Insect Science, Journal of Medical Entomology, and American Entomologist.

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## Order of Elements

Order of elements are as follows: title page; Abstract and key words; introduction (no heading); Materials and Methods; Results; and Discussion (or Results and Discussion); Acknowledgments; References Cited; footnotes; tables; figure legends; and figures.
The introduction should clearly state the basis of your study along with the background of the problem and a statement of purpose. The Materials and Methods section should include a clear and concise description of the study design, experimental execution, materials, and method of statistical analysis. Results should be clearly differentiated from the interpretation of your findings in the Results section or within the Results and Discussion. Cite tables and figures in numerical order as they should appear in the text. Include suggestions for direction of future studies, if appropriate.

## Title Page

The title page should include the name, complete address, phone number, fax number, and email address of corresponding author.
Include a running head of $<65$ characters, including author names. Example: Smith and Jones: Biological Control of C. capitata (no period). For more than two authors, use the senior author's name followed by et al. Example: Smith et al.: Biological Control of C. capitata (no period).
Include the section of the journal.
The title should be concise and informative. Include either the ESA approved common name or the scientific name, but not both of the subject. Common names used in the title must be listed in the ESA Common Names of Insects \& Related Organisms. Do not include authors of scientific names in the title. Do not capitalize the following words in the title or subheadings: a, an, and, as, at, be, by, for, in, of, on, per, to, the. Insert (Order: Family) immediately after the name of the organism.

Affiliation line includes a complete address. If appropriate, designate current addresses for all authors by numbered footnotes (superscripted numbers) placed at the bottom of the title page. Example:

1Department of Entomology, University of Colorado, 345 East 7th Street, Denver, CO 78095.

Include all authors' names below the title. Footnote numbers are placed outside commas in multi-authored articles.

## Abstract and Key Words

Abstract. On a separate page, provide an abstract of fewer than 250 words. Give scientific name and authority at first mention of the subject organism. Do not cite references, figures, tables, probability levels, or results. Refer to results only in the general sense.
Keywords. Place three to five key words, separated by commas, on a line below the abstract. Use only singular words/noun. Spell out scientific names (e.g., spell out Aedes albopictus instead of Ae. albopictus). Do not combine different subjects as one key word (e.g., "pesticides and grass," should be two separate keywords, "pesticide, grass." Do not use scientific names and common name at the same time as one key word [e.g., use "coffee, Coffea Arabica" (as 2 key words) instead of coffee (Coffea Arabica).
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## Methods and Results

Heading Levels
First-level headings are centered and boldfaced on their own line. Initial capital letters. Used to divide the manuscript into major sections (e.g., Materials and Methods, Results).
Second-level headings are flush left, boldface, and are also on their own line with initial capital letters. Second-level headings are rarely used except in taxonomic articles where multilevels of headings may be necessary.)
Third-level headings are boldfaced, paragraph indented, have initial capital letters, and are followed by a period. Third-level headings are used to divide first-level sections into smaller sections.

Fourth-level headings are italicized (but not boldfaced), paragraph indented, have initial capital letters, follow immediately after a third-level heading or start a new paragraph, and are followed by a period. Fourth-level headings are used to divide third-level sections into smaller sections.

## In-text Citation

## Single Author

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Two Authors
(Smith and Jones 1993)

## Multiple Citations

(Smith 1996, Smith et al. 1997, Jones 1998)
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Personal Communications
(Jones 1988; L. J. Smith, personal communication). Obtain and forward (at submission) a letter of permission to use citations to personal communications (from those other than authors).

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In Press
(Smith 1997) for in press, cite projected year of publication.

## Software

(PROC GLM, SAS Institute 1999) for software user's manual.

## Manufacturers

In parentheses, provide manufacturer's name and location (city, state) and model number of relevant materials and equipment. Example: (Model 3000, LI-COR, Lincoln, NE). Use generic names when possible (e.g., self-sealing plastic bags).

Reporting Requirements for Statistical Tests
All data reported (except for descriptive biology) must be subjected to statistical analysis. Descriptive biology should include information such as sample sizes and number of replications. Authors are responsible for the statistical method selected and for the accuracy of their data. Authors should be able to justify the use of a particular statistical test when requested by an editor. Results of statistical tests may be presented in the text, in tables, and in figures. Statistical methods should be described in Materials and Methods with appropriate references. Experimental designs should also be described fully in Materials and Methods. Descriptions should include information such as sample sizes and number of replications. See specific section in this style guide for suggestions on formatting statistical results. Only t-tests and analyses of variance require no citation. Cite the computer program user's manual in the References Cited.

## Probit/logit

When presenting results of probit/logit analysis, these columns should be included in tables (in this order, left to right); n, slope + SE, LD (or LC) ( $95 \%$ CL), and chi-square. When a ratio of one LD versus another is given, it should be given with its 95\% CI.

Statistical tests to show what model best fits data intended to estimate the $99.9986 \%$ level of effectiveness should be presented to justify use of any model, including the probit model. Thus, we do not recommend use of the Probit 9 without tests to show that the probit model fits the data.

## Analysis of Variance or t-test

When presenting the results of analysis of variance or a t-test, specify F (or t ) values, degrees of freedom, and P values. This information may be placed in parentheses in the text. Example: $(\mathrm{F}=9.26 ; \mathrm{df}=4,26 ; \mathrm{P}<0.001)$. If readability of the text is affected by the presence of repeated parenthetical statistical statements, place them in a table.

## Regression

In regressions, specify the model, define all variables, and provide estimates of variances for parameters and the residual mean-square error. Italicize variables in equations and text.

Variance and sample size
Include an estimate of the variance and sample size for each mean regardless of the method chosen for unplanned multiple comparisons. The use of Duncan's Multiple Range Test (DMRT) is not acceptable as a mean separation test as it is no longer commonly accepted as a method for post hoc mean separation anlysis.

## Model Analysis

At the beginning of the manuscript, authors should state clearly the goals of their model construction and analysis. Evaluation by reviewers depends upon these goals and the type of model. Authors should attempt to describe the main conclusions, limitations, and sensitivity of results to assumptions. For stochastic models, describe the variability in the results.

## Modeling Guidelines

The following guidelines pertain to any mathematical model calculated for purposes other than statistical analysis. Authors must adequately describe both model structure and model analysis. Authors must explain and justify original equations and computer programs or justify the selection of a published software package used in the computation of models. Model structure and steps in the analysis must be described in the Materials and Methods section. Without presenting extensive computer code, the text must permit an understanding of the model that would allow most mathematically inclined scientists to duplicate the work. Present all equations that represent the biology of the system being modeled. Unless their derivation is self-evident, show how the equations were derived and mention the underlying assumptions. Express how the equations are solved over time and space. Provide references for standard techniques (e.g., matrix manipulation, integration). Define all variables and parameters in each equation and describe their units (e.g., time, space, and mass). In the Materials and Methods or Results section, present the range of parameter values included in the model, and describe the uncertainty in or range of validity of these values.

## Equations

Consult Mathematics into Type for correct formatting of equations and mathematical variables. Italicize all mathematical variables. Center more complex equations on a separate line.

$$
\begin{equation*}
\mathrm{R}=\mathrm{A} \text { barrtype }+\mathrm{Blog} 10(\mathrm{f}) \tag{2}
\end{equation*}
$$

Validation or the Testing of Model Results
Authors must state why the model did not require testing (e.g., theoretical study), why it cannot be tested (e.g., lack of data), or how it was tested. Data used for testing must be independent of data used to build or calibrate the model. Describe the data and procedures in Materials and Methods. Authors should be aware that the testing of models is an important step that should be a part of most studies.

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For models solved or simulated by computers, mention the programming language and computer used. Describe the important numerical methods used in calculating the model (e.g., integration and random number generation). Mention how the program's logic and algorithms were tested and verified. When published software is computed, provide a reference and state which procedures were used. Discuss in any section of the manuscript the limitations of the published software. Original computer programs should be made available at the request of reviewers and readers.

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Inclusion of a GenBank/EMBL accession number for primary nucleotide and amino acid sequence data is a criterion for the acceptance of a manuscript for publication. Sequences from new species and new genes must indicate the proportion of the gene sequenced and should include data from both strands. The accession number may be included in the original manuscript or the sequence may be provided for review and an accession number provided when the manuscript is revised. A manuscript will not be accepted for publication until the accession number is provided.
GenBank may be contacted at their website at http://www.ncbi.nlm.nih.gov/Genbank/submit.html. The EMBL Data Library may be contacted at their website at http://www.ebi.ac.uk/embl/Submission/index.html.

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Follow the International Code of Zoological Nomenclature, 4th ed., for taxonomic style. Center the heading that indicates the name of the taxon in bold type. Center figure numbers in parentheses under the main heading; do not use bold type. Start all synonomies at the left margin with runovers indented. Include authors and date. References must appear in References Cited section. Use telegraphic style throughout descriptions.
For Journal of Medical Entomology Authors only: Please refer to the journal's Policy on Names of Aedine Mosquito Genera and Subgenera if writing about these insects.
Taxonomy Headings
Use only acceptable 3rd-level subheadings such as:
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Female
Material Examined
Type Material
Distribution
Etymology
Biology
Discussion
Avoid using Description as a subheading.
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Use Roman numerals I through XII to designate month of collection. Use arabic numerals 00 through 99 to designate collection years in the 20th century. Do not abbreviate other years, including the 21st century. Express data in this format: day-month (use a Roman numeral)year. Example: 2-V-97.
Locality Other than Principal Types
Start with the largest area followed by successively smaller areas separated by colons. Capitalize countries. Arrange data for each locality in the following order: count of specimens and sex or stage (as applicable), city or vicinity, date, collector, and depository. Example: MEXICO: Tamaulipas: 1 male, 1 female, Ciudad Mante, 15-III-97, K. Haack; 5 females, Ciudad Victoria, 3-VII-99, C. Hughes, MCZ. Arrange localities alphabetically. Use
a semicolon to separate data for different localities. Define depositories in the Materials and Methods.

## Type Material

Start description with the principal type in capital letters. Follow this immediately with count and sex of specimens (use male and female symbols if possible), then place additional data in the order of locality, date, additional data, and collector. Separate these items with commas. Example: HOLOTYPE: 1 male, Locust Grove, VA, 22-X-98, on Cercis canadensis, R. H. Foote. PARATYPES: 2 males, same data.
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Voucher specimens of arthropods serve as future reference for published names used in scientific publications. Although the deposition of voucher specimens is not required as a condition for publication, authors are encouraged to deposit specimens in an established, permanent collection and to note in the published article that the expected deposition has been made and its location. Authors should contact the curator of a voucher repository before deposition concerning the procedures required for curation to ensure that the collection will accept the voucher materials. The designation and proper labeling of voucher specimens is the author's responsibility. When available, at least three specimens should be deposited. Each specimen should have the following information provided at the time of deposition:
Standard label data that are required for the specimens collection (i.e., locality, date of collection, collector, host, ecological data, whether the specimen is from a laboratory collection, etc.).
An identification label that includes the identifier and date of identification.
A label that designates the specimen as "voucher."
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Pathogens. Reference should be made to Biological Use Authorization approved by an institutional Environmental Health and Safety committee or similar body.
Sample notification: The collection and infection of wild birds with encephalitis viruses was done under Protocol 11184 approved by the Institutional Animal Care and Use Committee of the University of California, Davis, California Resident Scientific Collection Permit 80104902 by the State of California Department of Fish and Game, and Federal Fish and Wildlife

Permit No. MB082812-0. Use of arboviruses was approved under Biological Use Authorization \#0554 by Environmental Health and Safety of the University of California, Davis, and USDA Permit \#47901.

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## References Cited

Cite only those articles published or formally accepted for publication (in press). Include all references mentioned in text. Include enough information to allow reader to obtain cited material (e.g., book and proceedings citations must include name and location [city and state or country] of publisher).
Abbreviate journal titles according to the most recent issue of BIOSIS Serial Sources. For non-English titled journals that are cited in the references, the title of the journal should be spelled out, and not abbreviated. Systematics-related articles may specify that all serial titles be spelled out for final publication. Citations and References should not be numbered.

References Cited: Alphabetical order (chronological for one author or more than two authors, and alphabetical order [by surname of second author] for two authors)

## Journal Articles

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Tyler, A. 2001. Western corn rootworm, vol. 2. Publisher, city, state abbreviation or country.

## Article/Chapter in Book

Tyler, A. 2001. Article or chapter title, pp. 000-000. In T.A.J. Royer and R. B. Burns (eds.), Book title. Publisher, city, state abbreviation or country.
Tyler, A., R.S.T. Smith, and H. Brown. 2001. Onion thrips control, pp. 178-195. In R. S. Green and P. W. White (eds.), Book title, vol. 13. Entomological Society of America, Lanham, MD.

## No Author Given

(USDA) U.S. Department of Agriculture. 2001. Title. USDA, Beltsville, MD.
(IRRI) International Rice Research Institute. 2001. Title. IRRI, City, State or Country.
Patents
Harred, J. F., A. R. Knight, and J. S. McIntyre, inventors; Dow Chemical Company, assignee. 1972 Apr 4. Epoxidation process. U.S. patent 3,654,317.
Proceedings
Martin, P. D., J. Kuhlman, and S. Moore. 2001. Yield effects of European corn borer (Lepidoptera: Pyralidae) feeding, pp. 345-356.In Proceedings, 19th Illinois Cooperative Extension Service Spray School, 24-27 June 1985, Chicago, IL. Publisher, City, State.

Rossignol, P. A. 2001. Parasite modification of mosquito probing behavior, pp. 25-28. In T. W. Scott and J. Grumstrup-Scott (eds.), Proceedings, Symposium: the Role of Vector-Host Interactions in Disease Transmission. National Conference of the Entomological Society of America, 10 December 1985, Hollywood, FL. Miscellaneous Publication 68. Entomological Society of America, Lanham, MD.
Theses/Dissertations
James, H. 2001. Thesis or dissertation title. M.S. thesis or Ph.D. dissertation, University of Pennsylvania, Philadelphia.
Software
SAS Institute. 2001. PROC user's manual, version 6th ed. SAS Institute, Cary, NC.
Online Citations
Reisen, W. 2001. Title. Complete URL (protocol://host.name/path/file.name) and/or DOI (Digital Object Identifier)
[return to top]

## Tables

Place tables after the References Cited section. Double-space and number all tables. Boldface table title. Do not repeat data already presented in text. If a table continues on more than one page, repeat column headings on subsequent page(s).
Title
Title should be short and descriptive. Boldface table number and title only. Include "means + SEM" in title if applicable. Do not footnote title; use the unlettered first footnote to include general information necessary to understand the table (e.g., define terms, abbreviations, and statistical tests).

## Lines

Use horizontal lines to separate title from column headings, column headings from data field, and data field from footnotes. Do not use vertical lines to separate columns. All columns must have headings.

## Abbreviations

Use approved abbreviations. Use abbreviations already defined in the text and define others in the general footnote. Use the following abbreviations in the body or column headings of tables only: amt (amount), avg (average), concn (concentration), diam (diameter), exp (experiment), ht (height), max (maximum), min. (minimum), no. (number), prepn (preparation), temp (temperature), vs (versus), vol (volume), wt (weight). Use the following abbreviations for months: Jan., Feb., Mar., April, May, June, July, Aug., Sept., Oct., Nov., and Dec.

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Repeat operational signs throughout data field. Insert a space on either side of sign (1.42 $\pm$ 1.36).

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Leave no space between lowercase letters and their preceding values (e.g., 731.2ab).

## Footnotes to Tables

Use footnotes to define or clarify column headings or specific datum within the data field. Do not footnote the title; use the unlettered first footnote to include general information necessary to understand the table (e.g., define terms, abbreviations, and statistical tests). The use of asterisks is reserved for statistical significance only.

## Example:

Means within a column followed by the same letter are not significantly different ( $\mathrm{P}<0.05$; Student t-test [Abbott 1925]). *, P < 0.05; **, P < 0.01; ***, P < 0.001; NS, not significant).

Use lowercase italicized superscripted letters to indicate footnotes. Footnote letters should appear in the table in consecutive order, from left to right across the table then down the page.

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Although figures of any size can be submitted, figures that fit exactly the width of 1 column ( 72 mm ) or 2 columns ( 148 mm ) expedite the publication process. Figures should be no longer than 195 mm from top to bottom. Separate parts of the same figure must be grouped together and arranged to use space efficiently. Wherever possible, it is best to avoid using a full page for a set of illustrations. That is, authors should attempt to have each figure appear separately from the others and should consider numbering illustrations as separate figures rather than as multiple parts of the same figure.
When choosing a font size, remember that it should be large enough so that reduction to fit the journal page will not make lettering difficult to read. Final lettering size should be 8 or 9 point using the fonts Arial or Helvetica or Times New Roman or Times Roman. Letter locants on figures composed of more than one element should match those in the text (either upper- or lowercase). Use a scale bar in lieu of magnification, and define scale in the figure caption. Figures will not be relettered nor will flaws be corrected.
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See the Digital Art Preparation section on how to submit photographs. Remember to scan black and white photos as grayscale and not color. For color photos, use the CMYK color mode, not RGB. Save photos in the TIFF format.

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Type all captions double-spaced on a separate page. All captions should be in paragraph form as shown by the example below.
Fig. 1. Relationship between percentage of defoliation of oak trees and gypsy moth population density. (A) Defoliation and egg mass density. (B) Defoliation of egg density.
Letter locants on figures composed of more than one element should match those in the text (either upper- or lowercase). Do not use equal signs to define abbreviations; use commas (e.g., Ap, barometric pressure).

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Supplemental Material may be submitted in the form of one or more ( 8 maximum) files to accompany the online version of an article. Such material often consists of large tables, data sets, or videos which normally are not possible or convenient to present in print media. Supplemental Material represents substantive information to be posted on the ESA journal website that enhances and enriches the information presented in the main body of a paper. However, the paper must stand on its own without the need for the reader to access the
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Scientific names and authorities must be spelled out (except for Fabricius and Linnaeus, which are abbreviated as F. and L., respectively) the first time a species is mentioned in the abstract and again in the main body of text.

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Use only those common names cited in the current ESA Common Names of Insects \& Related Organisms online database, or those names approved by the ESA Common Names Committee. Do not use any other common name. Do not abbreviate common names (e.g., CPB for Colorado potato beetle).
Give scientific name and authority at first mention of each organism (including plants) in the abstract and again in the text.
Use of "Stadium," "Stage," and "Instar"
Manuscripts received for publication in ESA periodicals refer to arthropods and the periods of time in their development in various ways. These designations should be used consistently.
Stadium (Plural: Stadia): The period of time between two successive molts.
Stage: One of the successive principal divisions in the life cycle of an arthropod (e.g., egg, nymph, larva, prepupa, pupa, subimago, and adult).
Instar: The arthropod itself between two successive molts. For the purposes of the definition, hatching is considered a molt.

## Examples of Usage:

Nymphs feed on the underside of leaves during the first stadium.
Larvae of some dermestids go through an indefinite number of stadia (or have an indefinite number of instars).
The nymphs were reared through the fifth stadium. Immature stages (e.g., eggs, larvae, and pupae; eggs and nymphs) are illustrated.

First instar of cerambycids make galleries in wood.
Some 200 first-instar spiderlings were collected. The predators fed readily on early instars of the face fly.
Notes on Formatting

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Do not capitalize the following words in titles or subheadings: a, an, and, as, at, be, by, for, in, of, on, per, to, the.

## Abbreviations

Use standard abbreviations as listed in the Council of Science Editors' Scientific Style and Format or those listed in this guide. Avoid nonstandard abbreviations.

## Abbreviations for Time

Use the following abbreviations for time: h (hour), min (minute), s (second), yr (year), mo (month), wk (week), d (day). Do not add "s" to create plurals (e.g., wks).
Fig./Figs.
Use "Fig." if singular and "Figs." if plural (e.g., Fig. 1; Figs. 2 and 3).
Dates
When citing dates in the text (not in tables or taxonomic reports), do not abbreviate month, and use this format: 26 January 1997.

## Metric Units

Use metric units. English units may follow within parentheses only if they are of direct practical purpose.

## Liter

Do not abbreviate "liter" by itself or when accompanied by a numeral.
\% versus percentage
Use "\%" only with numerals and in tables and figures. Close up space to numerals (e.g., $50 \%$ ). Otherwise, use the word percentage (e.g., percentage of defoliation).

Per versus slash
Use "per" rather than a slash unless reporting measurements in unit to unit (e.g., insects per branch, not insects/branch; but $\mathrm{g} / \mathrm{cm} 2$, not g per cm 2 .

## Numbers

Spell out numbers at the beginning of a sentence. Spell out the numbers one through nine (10 and up are always used as numerals), unless they are used as units of measure (e.g., eight children, three dogs, $8 \mathrm{~g}, 3 \mathrm{ft}, 0600$ hours; NOT 8 children, 3 dogs, eight grams, three feet, or six o'clock am). This includes spelling out the ordinals first through ninth, along with twofold, one-way ANOVA, and one-half. Ordinals from 10 and higher are numerals, such as 10 th or 51 st. In some cases, such as where there is a long list of items (e.g., 8 flies, 6 mosquitoes, 4 butterflies, and 10 bees), exceptions can be made if the editor concurs. The editorial staff will have flexibility in interpreting the rule.
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All numbers $<1$ must be preceded by a zero (e.g., $\mathrm{P}<0.05$ ).

## Commas

When a number is $>1,000$, use a comma to separate hundreds from thousands.

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Use a semicolon to separate different types of citations (Fig. 4; Table 2).
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It is not necessary to repeat symbols or units of measure in a series (e.g., 30,40 , and $60 \%$, respectively).
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Avoid footnotes in the text. Use unnumbered footnotes only for disclaimers and animal use information. Place all footnotes on a separate page after References Cited. Examples of footnotes are:

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In conducting the research described in this report, the investigators adhered to the "Guide for the Care and Use of Laboratory Animals," as promulgated by the Committee on Care and Use of Laboratory Animals of the Institute of Laboratory Animal Resources, National Research Council. The facilities are fully accredited by the American Association of Laboratory Animal Care.

## ANEXO II

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INTRODUCTION
Types of paper

## Types of Contributions

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State the objectives of the work and provide an adequate background, avoiding a detailed literature survey or a summary of the results.
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Results
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## Appendices

If there is more than one appendix, they should be identified as A, B, etc. Formulae and equations in appendices should be given separate numbering: Eq. (A.1), Eq. (A.2), etc.; in a subsequent appendix, Eq. (B.1) and so on. Similarly for tables and figures: Table A.1; Fig. A.1, etc.

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Ensure that each illustration has a caption. Supply captions separately, not attached to the figure. A caption should comprise a brief title (not on the figure itself) and a description of the illustration. Keep text in the illustrations themselves to a minimum but explain all symbols and abbreviations used.

## Tables

Number tables consecutively in accordance with their appearance in the text. Place footnotes to tables below the table body and indicate them with superscript lowercase letters. Avoid vertical rules. Be sparing in the use of tables and ensure that the data presented in tables do not duplicate results described elsewhere in the article.

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Hallberg, E., Johansson, K.U.I. and Wallé n, R.,1997. Olfactory sensilla in crustaceans: morphology, sexual dimorphism, and distribution patterns. Int J Insect Morphol Embryol 26, 173-180.

Book Chapter:
Locke, M., 1998. Epidermis. In: Harrison, F.E. and Locke, M. (Eds), Microscopic Anatomy of Invertebrates, vol. 11A. Wiley-Liss, New York, pp. 75-138.

## Book:

Chapman, R.F., 1998. The Insects: Structure and Function. 4th edition. Cambridge Univ. Press, Cambridge.
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All necessary files have been uploaded, and contain:

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Further considerations

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- All references mentioned in the Reference list are cited in the text, and vice versa
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## ANEXO III

# NORMAS PARA PUBLICAÇÃO: 

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## Information for authors

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obituary in memory of deceased systematic zoologists (e.g. Zootaxa 545: 67-68)
taxonomic/nomenclatural notes of importance
book reviews meant to introduce readers to new or rare taxonomic monographs (interested authors/publishers must write to subject editors before submitting books for review; editors then prepare the book review or invite colleagues to prepare the review; unsolicited reviews are not published)
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4) The abstract should be concise and informative. Any new names or new combinations proposed in the paper should be mentioned. Abstracts in other languages may also be included in addition to English abstract. The abstract should be followed by a list of key words that are not present in the title. Abstract and key words are not needed in short correspondence.
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Smith, A. \& Smith, B. (2000) Title of the Chapter. In: Smith, A, Smith, B. \& Smith, C. (Eds), Title of Book. Publisher name and location, pp. $x-y$.
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Smith, A., Smith, B. \& Smith, C. (2001) Title of Book. Publisher name and location, xyz pp.
D) Internet resources

Author (2002) Title of website, database or other resources, Publisher name and location (if indicated), number of pages (if known). Available from: http://xxx.xxx.xxx/ (Date of access).
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## ANEXO IV

## NORMAS PARA PUBLICAÇÃO:

## SYSTEMATIC ENTOMOLOGY

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3113/homepage/ForAuthors.html

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Papers may be of any length up to about 50 printed pages but should be in clear, concise English. The editors reserve the right to make minor textual corrections and changes that do not alter the author's meaning. Detailed lists of examined material are welcome at any length, but will be included as freely available electronic supplements on Synergy. Concise lists and all details of the type material will be printed.

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The title of the paper should be informative but concise. A short title (for page headlines) should be given. The abstract should be a concise summary of the significant content and salient conclusions of the work. Tables should be used sparingly and should be self-
explanatory. Tables should be at the end of the file and must be formatted according to the journal's style (consult a current issue).

Figures should be of publishable quality and about twice their published size, labelled large enough to allow reduction and numbered consecutively in the sequence referred to in the text. Drawings and photographs that make up separate parts of the same figure must be mounted together and labelled consecutively (preferably left to right, top to bottom) with capital letters. Reference in the text should be: Fig. 1, Figs 2, 3 and Figs 1; 2A,C. Figure legends should be grouped together on a separate sheet. The format for figure legends is: Fig. 1. Genus species. A, Habitus;B,C, male genitalia; D-F, head (D), thoras (E) and abdomen (F). In the full-text online edition of the journal, figure legends may be truncated in abbreviated links to the full screen version. Therefore the first 100 characters of any legend should inform the reader of key aspects of the figure.

Colour illustrations are welcome. An appropriate editor should be contacted to negotiate the cost of publication.

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