## Grey Thomas Gustafson

## Biology <br> Department

This dissertation is approved, and it is acceptable in quality and form for publication:
Approved by the Dissertation Committee:
Dr. Kelly Miller
Dr. Christopher Witt

## Dr. Joseph Cook

## Dr. Johannes Bergsten

# PHYLOGENETICS AND DIVERSIFICATION OF WHIRLIGIG BEETLES (COLEOPTERA, GYRINIDAE) 

by<br>GREY T. GUSTAFSON

B.S. Biology, The University of Kansas, 2010

## DISSERTATION

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# PHYLOGENETICS AND DIVERSIFICATION OF WHIRLIGIG BEETLES (COLEOPTERA, GYRINIDAE) 

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#### Abstract

Whirligig beetles are a family of aquatic beetles with an estimated 1,000 species distributed globally. Phylogenetic analysis incorporating molecular data has only recently been conducted on the family and many genera have never received modern taxonomic revision. The first chapter of the dissertation provides a taxonomic revision of the Southeast Asian whirligig beetle genus Porrorhynchus, finding the genus to contain 5 species, and no support for the numerous proposed subspecies. For the second chapter a phylogenetic analysis utilizing the most novel phylogenetic reconstructions methods incorporating the new Fossilized Birth-Death macroevolutionary model implemented in the program MrBayes 3.2 .6 was conducted on the whirligig beetle tribe Dineutini. The analysis finds strong support for the monophyly of the tribe, and 4 out of 5 of the proposed genera. The numerous subgenera proposed for the genera Dineutus and Macrogyrus are unsupported, with most synonymized, and the remaining subgenera provided with new diagnoses and constituent species. The second chapter also includes a biogeographic analysis reconstructing the historical biogeography of the Dineutini. The third chapter utilizes the same phylogenetic methods as the second chapter, however, the analysis was performed at the family level, combining the dataset from a previous study as well taxa sampled in chapter two. The analysis reveals the whirligig beetles to be an ancient beetle


group with Triassic origins, and the two currently monotypic subfamilies as the sole survivors of dominant clades from the Mesozoic.

## Introduction

Phylogenetic study of the family Gyrinidae utilizing molecular data has only just begun. A 2012 study analyzed the family in an attempt to primarily elucidate the subfamilial, tribal, and generic relationships. This study proposed a new classification scheme and found strong support for the monophyly of most genera. However, within the tribe Dineutini, three out of the five recognized subgenera had their monophyly in question. Because of this, the tribe Dineutini is the focus of the first and second chapters of the dissertation. Chapter one, Revision of the Southeast Asian whirligig beetle genus Porrorhynchus Laporte, 1835 (Gyrinidae: Gyrininae: Dineutini) investigated the species level diversity within this dineutine genus that has never before received a modern taxonomic revision. This also allows assessment of species-level taxon sampling for the phylogenetic analysis in chapter two. Chapter two, Systematics and evolution of the whirligig beetle tribe Dineutini (Coleoptera, Gyrininae, Gyrinidae) inferred the phylogenetic relationships of the dineutine whirligig beetles from both morphological and molecular datasets using Bayesian and maximum likelihood analyses. This chapter provides a reconstruction of the historical biogeography of the dineutines, during their diversification, to assess the proposed Gondwanan relationships among the genera, as well as an explanation for the genus Dineutus' interesting biogeography. Chapter three, Relicts of early Mesozoic whirligig beetle clades persist in biodiversity hotspots provides a timeframe for the origins and diversification of the family Gyrinidae using a total-evidence fossil calibration approach and the most recent version of the phylogenetic software MrBayes 3.2.6 to provide a time calibrated phylogeny of the whirligig beetles.

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# Revision of the Southeast Asian whirligig beetle genus Porrorhynchus Laporte, 1835 (Gyrinidae: Gyrininae: Dineutini) 

Grey T. Gustafson ${ }^{1}$ and Kelly B. Miller ${ }^{2}$<br>Department of Biology and Museum of Southwestern Biology, University of New Mexico, Albuquerque, NM 87131, USA.

Email: gtgustafson@gmail.com ${ }^{1}$ kbmiller@unm.edu ${ }^{2}$


#### Abstract

The Southeast Asian whirligig beetle genus Porrorhynchus Laporte, 1835 is revised. The genus is found to be composed of five species, P. depressus Régimbart, 1892, P. indicans (Walker, 1858), P. landaisi Régimbart, 1892, P. marginatus Laporte, 1835, and P. misoolensis (Ochs, 1955) new status, and two subgenera the Porrorhynchus s. s.tr. subgenus and Rhomborhynchus Ochs, 1926. The characters uniting the species within the two subgenera are discussed, as is their relation to the genus Dineutus Macleay, 1825. Many new synonyms are established: Ceylorhynchus Brinck, 1955, new synonymy, is synonymized with the s. str. subgenus, $P$. barthelemyi Régimbart, 1907 new synonymy and $P$. landaisi latilimbus Ochs, 1926 new synonymy are synonymized with $P$. landaisi, $P$. marginatus mjoebergi Ochs, new synonymy is synonymized with $P$. marginatus, and $D$. (R.) depressus jamurensis Ochs, 1955 new synonymy, $D$. (R.) depressus versteegi Ochs, 1955: new synonymy; $D$. (R.) depressus moszkowskii Ochs, 1955 new synonymy are synonymized with $P$. (R.) depressus. Porrorhynchus misoolensis (Ochs, 1955) is elevated to full species status. Lectotypes are designated for P. brevirostris Régimbart, 1877, P. tenuirostris Régimbart, 1877, P. landaisi Régimbart, 1892, P. barthelemyi Régimbart, 1907, $P$. marginatus mjoebergi Ochs, 1926, and $P$. landaisi latilimbus Ochs, 1926. Each species has the dorsal habitus imaged, male and female genitalia illustrated, important morphology illustrated, a complete catalog of associated names and their use, distribution mapped, common name proposed, and discussion section. A key to the species is included, as is a checklist of the species including synonyms and common names.


Key words: aquatic beetles, identification keys, morphology, taxonomy

## Introduction

The genus Porrorhynchus Laporte, 1835 contains the largest species of whirligig beetle known ( $P$. landaisi Régimbart, 1892) and some of the most charismatic members within the family Gyrinidae. The genus is widely distributed throughout much of Southeast Asia and New Guinea, where they inhabit streams. Most of the species have been implicated as being of potential use as water quality indicators, appearing restricted to fast-flowing, clean, forested streams (Brinck 1980; Ochs 1927b; Polhemus 2011), increasing the need for their accurate identification. Several water beetle workers have also noticed the need for a revision of the genus and its highly variable species (Balke et al. 2004; Miller and Bergsten 2012). Despite this, the genus has never received a modern taxonomic revision.

Porrorhynchus was original erected by Laporte (1835) for a single characteristic species from Java. Subsequently several other species were added by Régimbart (1892a; b; 1907), as well as another lesser known species originally described by Walker (1858) in the genus Dineutus Macleay, 1825. Since these early descriptions, numerous subspecies have been added
by the works of Georg Ochs (Ochs 1926; 1955). The genus was also carved up into three subgenera by Brinck (1955), and its relationship with the genus Dineutus has been questioned several times, and remains unclear (Brinck 1955; Miller and Bergsten 2012; Ochs 1926).

The purpose of this paper is to revise the species originally proposed to be in the genus and supported by the majority of gyrinid workers (Brinck 1955; Guignot 1950; Miller and Bergsten 2012; Régimbart 1902). Some of the morphological characters supporting the different species' grouping into subgenera and their relationship to the genus Dineutus will be discussed. However, the goal of this paper is to determine how many species are present within this group, while the final consensus on its status as a genus proper in relation to Dineutus and its constituents will be resolved in a forthcoming phylogenetic analysis of the tribe Dineutini.

The common name of "snout whirligig" has been proposed for the genus Porrorhynchus (Jäch and Ji 1998). We here continue to use this common name for the genus, but emend it to "snouted whirligig", as snouted is a more appropriate adjective for the descriptive common name. We propose common names for all the species of Porrorhynchus in order aid future attempts at protection status for these whirligig species of potential conservation concern.

The complete life history has never been described for any of the species of Porrorhynchus and only the imago stage is known. Larvae and pupae remain to be discovered and described. It is likely the larvae are similar in habit to those of the closely related Dineutus, being found in stream bottoms (Hatch 1927), and that pupation takes place on land in a pupal chamber under nearby cover such as rocks or logs, as is done in Dineutus (Wilson 1923).

## Materials and methods

A total of 1362 specimens were examined in this study.
Measurements were taken using a Cen-Tech 4 inch Digital Caliper (ITEM 47256). Total body lengths were measured from the anterolateral margin of the clypeus to the apex of the elytral apices. These areas were chosen for the boundaries of lengths since they are more fixed than other possible boundaries. For example, the labrum may be depressed thereby making it a poor choice as an anterior boundary, and the abdomen may be more or less protruding making it an unsuitable posterior boundary. Width was taken from the widest point of the body, typically just posteriad to the mid-length of the elytra. For each taxon an attempt was made to measure the largest and smallest specimens available for each sex.

Specimens for dissections and imaging were relaxed by placing them in lightly boiling water. The aedeagus was then dissected from relaxed males and placed in warm $10 \% \mathrm{KOH}$ for about 5 minutes. Following removal from KOH the aedeagus was placed in vinegar to neutralize the base and washed in water. Female dissection follows Miller (2001). Genitalia were drawn while in water and kept in glycerin, allowing the genitalia to dry causes them to become distorted and brittle. After dissection and/or illustration, aedeagi, female reproductive tracts, and abdomens were placed in microvials attached to the pin with the original specimens.

Illustrations were first drawn via camera lucida attached to a Zeiss Discovery V8 stereo microscope, then scanned and traced in Adobe Illustrator CS5.

Dorsal and ventral habitus were taken using a Visionary Digital BK+ light imaging system as well as a Passport imaging system (www.visionarydigital.com, R. Larimer).

Habitus images were then edited using Adobe Photoshop CS5 to add scale bars and improve clarity and color.

Distribution maps were created using Arc GIS.
Handwriting on type labels was identified using Horn et al. (1990).
Terminology of dineutine structures follows Gustafson and Miller (2015).
For each species a taxonomic history is provided. The family-group name classification follows Gustafson and Miller (2013).

Depository abbreviations:

| BMNH | The Natural History Museum, London, United Kingdom (C. Taylor) |
| :--- | :--- |
| BPBM | Bernice P. Bishop Museum, Honolulu, Hawaii, USA (S. Myers) |
| CNC | Canadian National Collection of Insects, Ottawa, Ontario, Canada (P. Bouchard) |
| DAPC | Dan A. Polhemus Collection |
| ICRI | Research Institute of Entomology, Sun Yat-Sen University, Zhonghan, Guangzhou, <br> Guangdong, China (F. Jia) |
| IRSB | Institut Royal des Sciences Naturelles de Belgique, Brussels, Belgium (W. Dekoninck) <br> MCZ |
|  | Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts, |
|  | USA (P. Perkins) |

MNHN Musée National d'Histoire Naturelle, Paris, France (A. Mantilleri)
MSBA Museum of Southwestern Biology Arthropod Division, The University of New Mexico, Albuquerque, New Mexico, USA (K. Miller)
MSNG Museo Civico di Storia Naturale, Genoa, Italy (M. Tavano)
MVMA Museum Victoria, Abbortsford, Victoria, Australia (S. Hinkley)
MZLU Museum of Zoology, Lund University, Lund, Sweden (R. Danielson)
NHMW Naturhistorisches Museum Wien, Vienna, Austria (M. Jäch)
NMPC National Museum, Prague, Czech Republic (J. Hájek)
ROME Royal Ontario Museum, Toronto, Ontario, Canada (D. Currie)
SMF Forschungsinstitut und Naturmuseum Senckenberg, Frankfurt-am-Main, Germany (D. Kovac)
UMMZ Museum of Zoology, University of Michigan, Ann Arbor, Michigan, USA (P
UMRM Enns Entomology Museum, University of Missouri, Columbia, Missouri, USA (R. Sites)
ZMUC Zoological Museum, University of Copenhagen, Denmark (A. Solodovnikov)

## Species concept utilized

The evolutionary species concept sensu Wiley (1978) is the preferred definition for a species utilized in this study, defining a species as "... a single lineage of ancestral descendant populations of organisms which maintains its identity form other such lineages and which has it
own evolutionary tendencies and historical fate." Our requirement for species level taxa is evidence from morphology, that the individuals studied exhibit features warranting their membership within a lineage distinct from other such populations, and maintenance of that lineage's unique identity, as inferred through exclusive combinations of morphological characters.

## Structures of taxonomic importance

Antennal flagellomere number: The number of antennomeres in the flagellum of the antenna for Porrorhynchus has been reported previously several times (Brinck 1980; Hatch 1925; 1926; Miller and Bergsten 2012), however, it was found that the number of antennomeres differs between most of the species and some reports were in error. Hatch $(1925 ; 1926)$ reported $P$. landaisi as having 8 flagellomeres, however, we found that $P$. (P.) landaisi had two different antennomere counts between the posterior and anterior faces of the flagellum, 8 posterior, 7 normally on the anterior (Fig. 2A). The anterior face of the flagellum of $P$. (P.) landaisi often varied in degree of the appearance of the 8th, with some specimens having it strongly effaced, while in others it was fairly distinguishable. Porrorhynchus ( $P$.) indicans was reported by Brinck (1980) as having 8 antennomeres in the flagellum, however, it actually has 7 on both faces (Fig. 2B). Miller and Bergsten (2012) reported Porrorhynchus as having 6 antennomeres in the flagellum, having only examined $P$. marginatus. We concur that $P$. (P.) marginatus has 6 flagellomeres (Fig. 2C) but as stated above this is not true of the whole genus. Porrorhynchus (R.) depressus similarly has 6 flagellomeres (Fig. 2D).

Labrum shape: The labra of species of Porrorhynchus are relatively elongate (being at least ca. $1 / 2$ as long as wide), especially in comparison to most members of Dineutus, and vary in shape among the species. Of the species, $P$. (P.) indicans and $P$. (P.) landaisi have relatively short, subtriangular labra, while those of $P$. (P.) marginatus and $P$. (R.) depressus are more triangular in shape. In $P$. ( $P$.) marginatus the labrum is in the form of an isosceles triangle and is the longest among the species of Porrorhynchus. The labrum of $P$. (R.) depressus is in the form of a near equilateral triangle and relatively short in relation to body size.

Labral setation: The setation of the ventral surface of the labrum differs between the two subgenera of Porrorhynchus. Member of the sensu stricto subgenus have the labral setation similar to that of Dineutus with two transverse rows of setae, while Rhomborhynchus has an additional row of setae situated paramedially running longitudinally down the labrum. This character appears to be unique to Rhomborhynchus.

Interorbital ridge coloration: The lateral canthus of the dorsal and ventral eyes, dubbed the interorbital ridge by Brinck (1955), has a character that distinguishes $P$. (P.) marginatus from all other members of Porrorhynchus. In P. (P.) marginatus the yellow margin exhibited on the elytra and pronotum is extended onto the lateral margin of the interorbital ridge. In all other species of Porrorhynchus the interorbital ridge is similarly colored as the frons and vertex.

Labial and maxillary palpi: Within Porrorhynchus the ultimate maxillary palpomere is truncate apically and quadrate in shape. The ultimate labial palpomere on the other hand is different among the species. In $P$. (P.) indicans and $P$. (P.) landaisi the ultimate labial palpomere differs in shape from that of the maxillary by being more elongate in form and apically
subtruncate to weakly rounded (Figs. 6C,G,N). In $P$. (P.) marginatus and $P$. (R.) depressus the ultimate labial palpomere is similar in form to that of the maxillary palps (Fig. 8I,N,S,X,BB). In $P$. (R.) depressus the palps are hatchet-form being short and quadrate with a relatively large highly truncate apical surface (Fig. 11C,G,K,O).

Pronotal transverse impressed line: The pronotum has a transverse impressed line, just removed from the anterior margin and is variously developed in different groups of whirligig beetles (Hatch 1926; Oygur and Wolfe 1991). Within Porrorhynchus this character divides the two subgenera, being absent in the $s$. str. group of species but present in Rhomborhynchus.

Yellow lateral margin of elytra: The lateral margin of the elytra is yellow in color and variable in its extent, width, and whether it is interrupted in its basal $1 / 3$, among the species. In $P$. (P.) indicans (Fig. 1C) the yellow margin typically only extends up to the basal half of the elytra and is reduced to variable degrees in width and extension among individuals. In $P$. (P.) landaisi (Fig. 1B) the yellow margin is well developed, extending most of the elytral length, ending just before the apex, being replaced by a blue reflective spot, and is interrupted at its basal $1 / 3$ by a constriction of its width. This constriction is associated with a swelling resulting from a cavity formed for pro-leg reception. In $P$. ( $P$.) marginatus (Fig. 1A), $P$. (R.) depressus (Fig. 1D), and $P$. (R.) misoolensis (Fig. 1E), the yellow lateral margin of the elytra is complete, reaching the apicolateral margin of the elytral apex. Porrorhynchus (P.) marginatus differs in having the yellow margin interrupted at its basal $1 / 3$ by a dark spot of coloration as well as a darkly colored apicolateral margin of variable length. The dark spot is associated with the swelling created by the cavity for pro-leg reception and is highly variable in size among individuals, as is the apicolateral marginal dark spot.

Dorsal punctation: The dorsal punctation differs greatly between the two subgenera. In the $s$. str. subgenus the dorsal punctation is well evident and composed of fairly large well-impressed punctures. In Rhomborhynchus punctation is present, but the punctures are finer, mostly obscured by the dense microreticulation covering the dorsum. Between the different species of the $s$. str. subgenus the distance between punctures varies considerably, with $P$. (P.) landaisi having the largest and densest dorsal punctation.

Elytral apices: The elytral apices differ among each species and are critical for identification. Porrorhynchus (P.) indicans differs from all other species having non-spinose elytral apices, instead having two large parasutural points, forming a large triangle between the sutural and epipleural angles (Fig. 1C). Both $P$. (P.) landaisi and $P$. ( $P$.) marginatus have saw-tooth-like serration, consisting of large triangular spinose cuticle extensions, running the apicolateral margin of the elytra. The saw-tooth serration becomes larger distally and the final extension is greatly expanded into a spine at the epipleural angle. In additional to this, $P$. ( $P$.) landaisi has the sutural angle produced and two parasutural spines (Fig. 1B). Porrorhynchus ( $P$.) marginatus similarly has the sutural angle produced but only has a single large parasutural spine, as well as having a larger epipleural spine (Fig.1A). Porrorhynchus (P.) depressus has the elytral apices very similar to $P$. (P.) marginatus, but lacks any saw-tooth-serration (Fig. 1D).

Setose furrow of ultimate female protarsomere: The ultimate protarsomere of females of most species have a setose furrow located on its posterolateral face running for most its length (Brinck
1980). This character divides the subgenera, being present in the $s$. str. subgenus, but completely absent in Rhomborhynchus.

Protrochanteric setose patch: The protrochanter of males of the $s$. str. subgenus possess a small patch of setae (Fig.3B) that appear to be recessed in a pit on the ventral face of the protrochanter (Fig. 3A). This sexually dimorphic character has previously been overlooked in the past and appears to constitute a synapomorphy for the $s$. str. subgenus. The position of this setose patch varies among the species. In $P$. ( $P$.) marginatus and $P$. ( $P$.) indicans it is located in the apicolateral corner, while being situated paramedially in $P$. (P.) landaisi. The Protrochanteric setose patch is not found in members of the subgenus Rhomborhynchus.

Profemoral setation pattern: The profemoral setation is another character dividing the two subgenera. In the sensu stricto subgenus the profemur possesses two rows of setae clustered together into large tufts along the anterior and posterior margins of the ventral surface, becoming more closely clustered and larger as they proceed distally (Fig. 4). The subgenus
Rhomborhynchus has only three to four small, narrow clusters of setae situated on the posterior margin of the ventral surface and $4-5$ setigerous punctures on the anterior face, similar to species of Dineutus. These two characters are discussed more in the broader conclusions section.

Protarsus: The shape of the protarsus varies between the different species, being much broader in $P$. (P.) landaisi and $P$. (P.) indicans (Fig. 6A,E,I,L), and more narrow in $P$. ( $P$.) marginatus and $P$. (R.) depressus (Fig. 8A,F,K,P,U,Y, Fig. $11 \mathrm{~A}, \mathrm{E}, \mathrm{I}$ ). There are also differences in the relative proportions of the protarsomeres between some of the species.

Metacoxal process: The form of the metacoxal process varies among most the species. In general the species have a slight production of the distolateral corner of the apex of the metacoxal process. Which is weakly produced in all the species (Fig. 6B,F,J,M), except for $P$. $(P$.$) marginatus in which the process is clearly produced and spinose in most populations (Fig.$ 8B,G,L,Q,V,Z).

Aedeagus: The aedeagus also shows a major division between the subgenera especially in the form of the parameres. In Porrorhynchus s. str. the parameres all have the setae restricted to their apical $1 / 4$, while in Rhomborhynchus the setae is located in the apical $1 / 3$. The parameres also articulate broadly with the median lobe in the $s$. str. subgenus, while in Rhomborhynchus they are much more narrowly articulated. The form of the aedeagus differs between all the species and as such, the aedeagus is a good indicator of species boundary.

Female reproductive tract: The gonocoxae of the female RT can distinguish most the species. All species have the female RT (Fig. 13) with a long tubiform spermatheca and relatively long, narrow laterotergites, and the vaginal shield (Brinck 1980) bordered by a strongly developed sclerotized region anteriorly.

Key to the adults of the known species of Porrorhynchus Laporte, 1835

1 Elytral apices spinose, distinct parasutural spine(s) present. Yellow margin of elytra complete or nearly so, normally extending well beyond half the elytral length.

- Elytral apices not spinose, instead with large parasutural point forming broad triangle between the sutural and epipleural angles. Yellow margin of elytra incomplete and normally strongly reduced, only reaching basal $1 / 2$ to $1 / 3$ of the elytral. Known only from Sri Lanka.

Porrorhynchus (Porrorhynchus) indicans
2 Elytra apicolaterally with saw-tooth-like serration. Yellow margin of elytra normally interrupted in basal $1 / 3$ by constriction of margin's width or darkly pigmented spot. Pronotal transverse impressed lines absent. Distributed in Southeast Asia, west of Wallace's line

- Elytra apicolaterally without saw-tooth-like serration of any kind. Yellow margin of elytra never interrupted in basal $1 / 3$. Pronotal transverse impressed line present. Known only from east of Wallace's line in West Papua, and New Guinea 4

3 Yellow lateral margin continued onto interorbital ridge. One large parasutural spine present on elytral apex. Elytral yellow margin normally interrupted by darkly pigmented spot in basal $1 / 3$. Labrum highly elongate, in the form of an isosceles triangle. Widely distributed in Southeast Asia, west of Wallace's line. More commonly encountered species

Porrorhynchus (Porrorhynchus) marginatus

- Yellow lateral margin not continued onto interorbital ridge. Two parasutural spines of similar size present on elytral apex. Elytral yellow margin interrupted by constriction of its width in basal $1 / 3$. Labrum shorter, sub-triangular in shape. Known from mainland Southeast Asia, primarily in the northeast, and southern China.
$\qquad$
4 Length $9.5-11.6 \mathrm{~mm}$; body form of most populations broader, especially broad at midlength. Metacoxal process with lateral margins less sinuate, less strongly constricted apically. Aedeagus with median lobe as- or nearly as long as parameres, apex not laterally expanded, parallel sided then narrowed in apical $1 / 6$. Labrum larger. Widespread in New Guinea.

Porrorhynchus (Rhomborhynchus) depressus

- Length $9.4-10.1 \mathrm{~mm}$; body form more narrow, broadest just anteriad to mid-length. Metacoxal process with lateral margins more strongly sinuate, more constricted apically. Aedeagus with median lobe shorter than parameres, apex briefly laterally expanded, with lateral margins broadly rounded towards apex. Labrum smaller. Only known from Misool Island. Porrorhynchus (Rhomborhynchus) misoolensis n. stat.


## Taxonomy

## Genus Porrorhynchus Laporte, 1835

Type species: Porrorhynchus marginatus Laporte, 1835: 108 by monotypy.
Trigonocheilus Dejean, 1833: 59 [manuscript name and nomen nudum, synonymy by Aubé, 1838a: 406]; Trigonochilus rostratus: Agassiz 1846: 377 [unjustified emendation of Trigonocheilus rostratus, nomen nudum]; Porrorhynchus Laporte, 1835: 108 [original description]; Porrhorhynchus: Régimbart 1877: 105 [misspelling]; Porrhorrhynchus: Régimbart 1886: 250 [misspelling]; Dineutus (Porrorhynchus): Ochs 1926: 64 [new status];
Porrorhynchus: Guignot 1950: 124 [change in status]; Dineutus (Porrorhynchus): Ochs 1955: 130 [change in status]; Porrorhynchus: Brinck 1955: 103 [change in status].

Diagnosis: Medium to very large whirligig beetles size: $9-26 \mathrm{~mm}$. Body form elongate oval to teardrop-shaped. Antennal flagellum with 6-7 antennomeres, ultimate flagellomere trapezoidal in shape and ca. at least 2 x as long as every other flagellomere. Labrum large, elongate, and more or less triangular in shape. Pronotum and elytra with yellow-margins, often for entirety of their length, but at least in the basal half of the elytra. Protibia with distolateral margin spinose. Venter lightly colored.

## Description:

Head. Vertex and frons with punctation present, reticulation composed of round to ovoid sculpticells. Dorsal eye smaller than ventral eye, anterior margin of ventral eye situated posteriad of anterior margin of dorsal eye; orbital furrow of dorsal eye complete, becoming narrowest anteromedially; exoculata suture well defined. Antenna with cup-like scape; pedicel broad, laterally expanded, dorsoventrally flattened, trapezoidal in form, lateral face with fringe of long fine setae; flagellum with $6-8$ flagellomeres, flagellomere I stalked, triangular in form, ultimate flagellomere trapezoidal in form, at least 2 x longer than flagellomeres II - V or VI, flagellomeres II - V or VI similar in size and shape. Frontal ridge length at least $1.5-2 \mathrm{x}$ width of clypeus at midlength, frontoclypeal suture well developed, posterior margin flat, lateral margins meeting posterior margin at ca. $100-110^{\circ}$ angle. Clypeus weakly to fairly strongly emarginate medially, with reticulation compose of round sculpticells, punctation present, clypealium with long fine setae ventrolaterally. Labrum large, elongate, more or less triangular in shape, densely punctate, with reticulation composed of round sculpticells, ventral margins fringed with long fine setae. Maxilla without galea, maxillary palp 4 segmented, ultimate palpomere ca. as long as proximal three combined, apically truncate. Labial palp 3 segmented with ultimate palpomere as long as all proximal palpomeres combined, apically truncate to subtruncate. Gula well developed, Tshaped with tentorial pits evident, series of long fine setae present anteriad to lateral arms of gular suture.
Thorax. Pronotum with punctation present, reticulation consisting of round sculpticells, lateral margins yellow with fairly broad marginal depressed area, posterolateral corners with several setae. Prosternum with well differentiated prosternal process, prosternal process narrow and paralleled sided for entirety. Prolegs with procoxal process round to lobiform; protrochanter fusiform, posterior face with short field of curved stout setae; profemur nearly parallel sided for most its length, only weakly tapered apically, weakly expanded basally; protibia club-shaped with distolateral marginal apex produced to a more-or-less acute spine, anterior face with
paramedial linear to arcuate series of setigerous punctures in distal half, proximal lateral margin with long grove running distal half of protibiae to distal apex with golden hair-like setae, distomedial apex encircled with short, stout, pointed setae, continuing to posterior face, ventral face of protibia with setose groove running near entire length, setae become larger, more brushlike towards distal apex, setae at distal apex projecting beyond apex's medial process, posterior face of protibia with brush of golden setae in distal $1 / 5$; protarsus 5 segmented, protarsomeres I IV ventral face with long projecting setose patch located in the proximomedial corner, protarsomere V of female with short series of brush-like setae running near entire length of ventral surface, posterior surface of protarsus highly sexually dimorphic, protarsal claws similar between the sexes. Elytra without sutural border, punctation present, reticulation consisting of round sculpticells, lateral margins yellow at least basally. Mesoventrite with well developed cavity for proleg reception; mesoventrite bordered anteriorly, border complete, thinly so at anteromedial projection between procoxae, projection with long setae basolaterally, large shallow punctures present anterolaterally on mesoventral body, mesoventral discrimen well developed, running ca. $1 / 2$ mesoventrite's length; mesepimeron narrow, strap-like. Mesolegs with mesocoxa possessing, shallow setose pit posteromedially; mesotrochanter lobiform; mesofemur broad, stout in distal half, distal apex very flatly rounded, proximal apex strongly attenuate towards trochanter, mesotibia triangular in form, dorsal surface with long natatory setae, distolateral angle of dorsal surface with series of short stout setae, distomedial surface with similar short stout setae, adorned with two spines, anterior shorter, posterior longer, ventral face of mesotibia also with short stout spines for its entire length; mesotarsus 5 segmented, mesotarsomere I in the form of equilateral triangle, 5 times length of mesotarsomere II, mesotarsomere III similar in size and form of II, all with ventral face with short stout setae running their entire length, mesotarsomere IV elongate and narrow, V ovoid with ventrodistal margin produced to spine before claws, both with long natatory setae; mesotarsal claws larger than metatarsal claws, sexually dimorphic in shape. Metaventrite with well developed cavity for proleg reception; metanepisternum lobiform; metacoxal process with posteromedial shallow setose pit, metacoxal process with circular pit anterolaterally, metatrochanter trapezoidal in shape, remainder of leg similar in form to mesoleg except metatarsal claws of both sexes smaller than mesotarsal claw, similar in form.
Abdomen. Abdominal tergites VI-VIII strongly pubescent with long fine setae covering most their surface, medially darkly pigmented, lateral margins lighter yellow in color, reticulation present, composed of round sculpticells, tergite VIII with two types of punctation distinctly present, smaller well impressed punctation covering most of darkly pigmented area, second type consisting of very large, shallowly impressed crater-like punctation situated basomedially extending half its length; abdominal sternite II + III ca. 2 x length of sternite IV, sternite IV ca. 2 x length of sternite V, sternites V,VI,VII, similar in size, sternite VIII triangular in form, nearly 2 x length of sternite VII, emarginate apicomedially, faintly impressed reticulation present over abdominal sternites, composed of ovoid sculpticells. Female reproductive tract with tubiform spermatheca; fertilization duct weakly differentiated; vaginal shield with anterior margin bounded by highly sclerotized bridge.

Sexual dimorphism: Protarsus of male laterally expanded with dense covering of uniform suction cup setae occupying near entirety of posterior surface, missing only from proximomedial corner of protarsomere I. Protarsus of female not laterally expanded, without suction cup setae on posterior surface, instead posterior surface of at least protarsomeres I - IV with smaller patches
of setae located at distal margins. Mesotarsal claws of male more strongly curved ventrally. Female mesotarsal claws less strongly curved, more similar in form to metatarsal claws.

## Subgenus Porrorhynchus Laporte, 1835

Type species: same as for genus.
Trigonocheilus Dejean, 1833: 59 [manuscript name and nomen nudum, synonymy by Aubé, 1838a: 406 Trigonochilus Agassiz, 1846 (nomen nudum); Porrhorhynchus: Régimbart 1877: 105 [misspelling]; Porrhorrhynchus: Régimbart 1886: 250 [misspelling]; Dineutus
(Porrorhynchus): Ochs 1926: 64 [description]; Porrorhynchus (Ceylorhynchus) Brinck, 1955: 103 new synonymy.

Diagnosis: Large to very large whirligig beetles, size: $12-26 \mathrm{~mm}$. Most species with 7 antennal flagellomeres. Gular suture incomplete. Pronotum without transverse impressed line. Elytral lateral margin with swelling at mid-length associated with proleg reception. Males with protrochanteric setose patch. Profemora with unique setation pattern consisting of large clusters of setae arranged along the anterior and posterior margins of the ventral femoral face, becoming larger and more closely clustered apically.

## Description:

Head. Antenna of most species with 7 antennal flagellomeres; pedicel narrow, trapezoidal in form, mildly to noticeably narrowed apically. Labrum ventrally with two transverse linear setose rows in basal $1 / 2$. Gular suture incomplete, lateral arms of gular suture effaced before meeting anterolateral margin of ventral epicranium, posteriad to submentum.
Thorax. Pronotum without transverse impressed line. Elytral lateral margin with swelling at midlength associated with depressed pro-leg reception cavity in meso- and metaventrite. Proleg with protrochanter of male possessing setose patch located paramedially or apicolaterally on posterior face, absent in female; profemur ventrally with two linear rows of setose clusters running entirety of length, clusters consists of long fine golden setae, clusters fairly evenly spaced along most series, becoming longer, larger, more clustered in apical $1 / 4$ of profemur, longest setose cluster ca. $1 / 2$ width of profemur, setose clusters smaller in basal $1 / 4$, anterior face of profemur with sinuate ventral margin, ventral margin of anterior face similar but to lesser degree, posterior face covered with short stout setae in recessed pits, dorsal face similar but to lesser degree, absent on posterior/ventral faces; protarsomere V of female protarsus with setose furrow running near entirety of posterior face. Metasternal wing in form of near equilateral triangle. Metacoxal wing laterally transverse, most species with metacoxal wing ending at apex of metepisternum.
Abdomen. Abdominal sternite VIII deeply emarginate medially. Male genitalia with median lobe of aedeagus broadly articulating basomedially with parameres, lateral proximal longitudinal lists meeting median list, forming trident-shape. Female reproductive tract with medial apodeme extended anteriorly forming continuous, strongly sclerotized bridge forming anterior to lateral boundaries of vaginal shield.

Sexual dimorphism: Males noticeably larger in size in many populations, much more broad in body form being laterally expanded at elytral mid-length. Protibia of very large males often with weak to very strong sinuation. Females with similar but much smaller and sparser profemoral setation. Female elytra without swelling associated with proleg reception.

## Porrorhynchus (Porrorhynchus) indicans (Walker, 1858)

Dineutes indicans Walker, 1858: 205 [original description]; Porrhorhynchus brevirostris Régimbart, 1877: 113, Pl. 6 fig. 5 [original description, dorsal habitus, synonymy by Régimbart, 1886: 250]; Porrhorhynchus brevirostris: Régimbart 1882: 429, pl. 11 fig. 50 [redescription, elytron]; Porrhorhynchus brevirostris: Régimbart 1884: 471 [checklist]; Porrhorrhynchus brevirostris: Régimbart 1886: 250 [misspelling]; Porrhorrhynchus indicans: Régimbart 1886: 250 [new status]; Porrhorrhynchus indicans: Régimbart 1892a: 740 [checklist]; Porrhorrhynchus indicans: Régimbart 1902: 5 [distribution]; Dineutus (Porrorhynchus) indicans: Ochs 1926:139 [checklist]; Dineutus (Porrorhynchus) indicans: Ochs 1929a: 248 [distribution]; Porrorhynchus (Ceylorhynchus) indicans: Brinck 1955: 103 [new status]; Porrorhynchus indicans: Vazirani 1969: 403 [distribution]; Porrorhynchus (Ceylorhynchus) indicans: Brinck 1980: 106, Fig. 2-6 [description, morphology]

Type material examined: Dineutes indicans Walker, 1858: Holotype by monotypy (1 ${ }^{\lambda}$ pinned, Fig. 16B) "Type [beige disc, typed black ink, red circle around font]// Ceylon [blue disc, handwritten in black ink, handwriting unknown]// indicans Walker/ Ann. Nat Hist (Type) [blue label, handwritten in black ink, handwriting unknown]// indicans [beige label, handwritten in black ink, handwriting unknown]// No 464/ examined by/ Prof. Thaxer for/ Laboulbeniaceae. [beige label, typed black ink, except for 464 handwritten in black ink]//" (BMNH).

Porrorhynchus brevirostris Régimbart, 1877 lectotype here designated (1 ठ pinned, Fig. 16A) "Ceylan [beige label, handwritten in black ink, handwriting unknown]// MUSEUM PARIS/ COLL MAURICE REGIMBART/ 1908 [beige label with thin black border, typed in black ink]// TYPE [red label, typed black ink]// LECTOTYPE [red label, typed black ink]" (1 ex. MNHN); paralectotype here designated ( 1 Q pinned, missing right proleg after femur) "Java [beige label, handwritten in black ink, handwriting unknown]// MUSEUM PARIS/ COLL MAURICE REGIMBART/ 1908 [beige label with thin black border, typed in black ink]// TYPE [red label, typed black ink]// PARALECTOTYPE [red label, typed black ink]" (1 ex. MNHN).

Type designation: Régimbart's (1877) original description of $P$. brevirostris indicates that there were 4 specimens used in the description, 2 from "Ceylan" which were stated to be in the Fairmaire and Deyrolle collections, while the other 2 were stated as being from Java (in error) in the Jekel and Régimbart collection. Only two specimens could be located in the MNHN, one with the locality label of Java and the other with Ceylan, both indicated as being from Régimbart's collection, now in the general collection of the MNHN. After searching for the other 2 specimens in the MNHN, they are now presumed lost. Therefore, these two specimens are considered to be part of the syntype series, and the male specimen with the label "Ceylan" is here designated as the lectotype, as P. indicans is truly found in Sri Lanka. Given that Régimbart
himself synonymized his own name, this designation seems quite sound. The specimen from Java is here designated as a paralectotype.

## Additional material examined:

SRI LANKA ("Ceylonia"): "Mus: Brents" (1 ex. ZMUC); "Ceylon": (2 ex. BMNH), Sharp Collection 1905-313 (1 ex. BMNH), "Mus./ Hauschild/ 12-9-1914" (2 ex. ZMUC), "Mus./ Westerm" (1 ex. ZMUC); "Niemer" (1 ex. BMNH), "Niemer" "Fry Collection/ 1905-100" (1 ex. BMNH); leg. P. Brinck, ZML.2010/ 332 ( 1 ex. MZLU), leg. Lewis, Sharp Collection 1905-313 (1 ex. BMNH), leg. G. Lewis, ZML.2010/ 331 (1 ex. MZLU), leg. G. Lewis, Sharp Collection 1905-313 (2 ex. BMNH); 1910, leg. G. Lewis, -320, (4 ex. BMNH); [illegible addition], "Coll.Mus./ Vindob." (3 ex. NHMW); [illegible addition], l"M. Doh", "Coll.Mus./ Vindob." (1 ex. NHMW); Colombo: Sharp Collection 1905-313 (1 ex. BMNH).
No locality information: Sharp Collection 1905-313 (3 ex. BMNH); "Jekyl" (1 ex. BMNH); "6756" (1 ex. BMNH).

Diagnosis: Labrum ovoid and shorter relative to other Porrorhynchus species. Antenna with 7 flagellomeres. Yellow lateral margins incomplete on elytra, extending normally only to the basal $1 / 3$. Elytral apices with blunt parasutural point, without spines.
 mm .
Habitus. Medium sized member of genus; body form elongate oval, attenuated anteriorly in male, female nearly parallel sided in appearance; in lateral view weakly convex, only slightly humped in scutellar region, in general dorsoventrally depressed relative to other species.
Coloration. Dorsally head, pronotum, elytra olive green, pronotum, basal $1 / 3$ of elytra with yellow lateral margins; venter yellowish to reddish orange, ultimate maxillary palpomere black, except for apex; prolegs often slightly darker in coloration, tibia black in proximal $1 / 2$.
Head. Dorsally vertex with sparse weakly impressed punctures; interorbital ridge without yellow margin, similarly colored as vertex; frons with weakly impressed punctures separated by 2 x to 3 x diameter of one puncture, fronto-lateral margins lightly wrinkled, frontoclypeal suture with posterior margin nearly straight, lateral margins nearly straight, meeting posterior margin at ca. $110^{\circ}$ angle; clypeus with punctation most evident at anterior margin, punctures separated by ca. 2 x to 3 x diameter of one puncture; antennal flagellum with 7 flagellomeres (Fig. 2B); labrum ovoid in shape, punctation nearly absent basomedially, strongly present apically, punctation well impressed, dense, separated by 1.5 x to 2 x diameter of one puncture; maxillary/labial palpi somewhat dissimilar in shape, maxillary palps broader with asymmetrical dorsal/ventral margins, ventral margin more strongly curved than dorsal margin, labial palps with anterior/posterior margins more similar, anterior margin nearly straight, posterior margin weakly curved Thorax. Pronotum with shallow weakly impressed wrinkles medially on disc, shallow transverse depression also often present medially, reticulation effaced on medial disc, weakly impressed sparse punctation present, laterally reticulation well impressed, punctation weakly impressed, nearly imperceptible, widely spaced, distance between nearest punctures up to 4 x to 5 x diameter of single puncture, lateral marginal depression present; Protrochanteric setose patch situated apicolaterally; protibial spine projecting forward; male protarsi wide, dorsally convex, shape as in Fig. 6L, male ultimate protarsomere ca. $>2 \mathrm{x}$ as long as wide; ultimate protarsomere of female ca. $2.5 x$ length of penultimate; elytra with reticulation effaced in scutellar/sutural regions,
reticulation present apically/laterally, most strongly impressed marginally, elytral disc with even covering of weakly impressed, fine punctation, distance between nearest punctures ca. 2 x to 3 x diameter of single puncture; lateral marginal depression narrow, of similar length throughout, yellow lateral margin incomplete, typically ending in basal $1 / 3$, when longer never reaching epipleural angle, apicolateral margins of elytra without triangular saw-tooth-like spines, elytral apices (Fig. 1C) not spinose, with parasutural points in form of broad triangle, created by lateral obliquely truncate margin, with rounded inner margin, sutural angle often produced; mesosternal apex not noticeably acuminate; mesocoxae/metacoxae dissimilar, mesocoxae without posteriorly projecting process, mesocoxal process broadly rounded; male mesotarsal claws as in Fig. 60 with ventral margin broadly rounded, anterior claw apically narrowed; metacoxal process as in Fig. 6 M with distinct apicolateral corners, weakly sinuate lateral margins.

Genitalia. Aedeagus (Fig. 5) with median lobe nearly as long as parameres, parallel sided for most its length, strongly acuminate in apical $1 / 5$, apex broadly rounded, in lateral view (Fig. 5B) apex briefly weakly curved dorsally; parameres in dorsal view laterally expanded in apical $1 / 3$, rounded apically, medially reflexed basally; in lateral view ventral margin of parameres evenly curved anteriorly to posteriorly. Female reproductive tract (Fig.13B) with large broad tubiform spermatheca; gonocoxae with lateral sinuation, apically acuminate.

Sexual dimorphism. Females are smaller in size and much more parallel sided in body form.
Differential diagnosis: Porrorhynchus indicans can be distinguished from all other species of Porrorhynchus by the incomplete yellow lateral margins of the elytra, as well as the elytral apices possessing large, triangular, parasutural points, rather than spines.

Distribution: Fig. 14. This species is only known from Sri Lanka. Within the country Brinck (1980) found the species to be primarily known from the southern central region of the island. For this study almost none of the material had specific locality information, aside from the island of Sri Lanka. One specimen indicated it was collected from Colombo, see Brinck (1980) for more precise locality and distribution data.

Biology: Brinck (1980) demonstrated that $P$. indicans is restricted to intact (what he calls "primeval") montane forest. After searching previously known localities, Brinck (1980) was only able to recollect the species from two streams in sheltered ravines within such forests, with water temperatures below $20^{\circ} \mathrm{C}$, and at elevations above $1,200 \mathrm{~m}$. Within the streams the species was mainly found in regions sheltered from the main current (Brinck 1980).

Discussion: Given the few known exact localities from Brinck (1980), the difficulty in recollecting the species in 1980, and imperiled with future potential habitats loss, P. indicans is certainly warranting formal protection status. Future investigation into the distribution and status of this species on the island is highly desirable.

We here propose the common name of the Sri Lankan Snouted Whirligig for $P$. indicans.

Porrhorrhynchus landaisi Régimbart, 1892a: 667, 740 [original description, checklist]; Porrhorrhynchus barthelemyi Régimbart, 1902: 5 [nomen nudum]; Porrhorrhynchus landaisi: Régimbart 1902: 5, fig. 12 [distribution, partial dorsal habitus]; Porrhorrhynchus barthelemyi Régimbart, 1907: 153 [original description] new synonymy; Porrhorrhynchus landaisi: Régimbart 1907: 152 [description of habitat, collection information] ; Porrhorrhynchus landaisi: Peschet, 1923: 123 [review]; Porrhorrhynchus landaisi: Hatch 1925: 437, 450 [size, minor description]; Porrhorrhynchus landaisi: Hatch 1926: 311, Pl. XX 3, 9, 16, 17, 21, Pl. XXI 29, 41, Pl. XXII 47, 60, 64, Pl. XXIII 80, Pl. XXIV 90, 98 [morphology]; Dineutus (Porrorhynchus) barthelemyi: Ochs 1926: 139 [cheklist]; Dineutus (Porrorhynchus) landaisi: Ochs 1926: 139 [checklist]; Dineutus (Porrorhynchus) landaisi latilimbus: Ochs 1926: 139 [nomen nudum in checklist]; Dineutus (Porrorhynchus) landaisi latilimbus Ochs, 1926: 193 [original description] new synonymy; Dineutus (Porrorhynchus) landaisi: Ochs 1929b: 719 [distribution]; Dineutus (Porrorhynchus) landaisi latilimbus: Ochs 1929b: 719 [distribution]; Dineutus (Porrorhynchus) barthelemyi: Ochs 1930: 15 [catalog]; Dineutus (Porrorhynchus) landaisi: Ochs 1930: 16 [catalog]; Dineutus (Porrorhynchus) landaisi latilimbus: Ochs 1930: 16 [catalog]; Dineutus (Porrorhynchus) landaisi: Wu 1931: 71 [distribution]; Dineutus (Porrorhynchus) landaisi latilimbus: Wu 1931: 71 [distribution]; Dineutus (Porrorhynchus) landaisi: Cheo 1934: 222, Pl. 1 Fig. 12, 13 [redescription, elytra]; Dineutus (Porrorhynchus) landaisi latilimbus: Cheo 1934: 223 [diagnosis] Dineutus landaisi latilimbus: Kamiya 1936: 14, fig. 19 [description, dorsal habitus]; Dineutus (Porrorhynchus) landaisi latilimbus: Ochs 1942: 206 [holdings]; Porrorhynchus landaisi landaisi: Mazzoldi 1995: 162 [distribution]; Porrorhynchus landaisi latilimbus: Mazzoldi 1995: 162 [distribution]. Snout Whirligig, Porrorhynchus landaisi latilimbus: Jäch \& Li 1998: foreword [notes on distribution]. Porrorhynchus sp.: Jäch et al. 2012: 66 [distribution].

## Type material examined:

Porrorhynchus landaisi Régimbart, 1892 lectotype here designated (1 $q$ pinned, Fig. 16C) "Environs de/ Cao-Bang./ Tonkin Landais [white label, handwritten in black ink, handwriting appears to be Régimbart's]/ MUSEUM PARIS COLL MAURICE REGIMBART/ 1908 [white label with thin black border, type black ink]// LECTOTYPE [red label, typed black ink]//" (1 ex. MNHN). Paralectotype ( 1 § pinned [prothorax clearly glue back on, head glue back on): "Tonkin, Hanoi/ Landais [white label, handwritten in black ink, handwriting appears to be Régimbart's]// MUSEUM PARIS/ COLL MAURICE REGIMBART/ 1908 [white label with thin black border, typed black ink]// TYPE [red label, typed black ink]// PARALECTOTYPE [red label, typed black ink]//" (1 ex. MNHN); paralectotype (1 + pinned) " Hanoi/ Landais [white label, handwritten in black ink, handwriting appears to be Régimbart's]// MUSEUM PARIS COLL MAURICE REGIMBART/ 1908 [white label with thin black border, type black ink]// PARALECTOTYPE [red label, typed black ink]//" (1 ex. MNHN).

Porrorhynchus barthelemyi Régimbart, 1907 lectotype here designated ( $1 \bigcirc$ pinned, aedeagus dissected on point, Fig. 16E) " MUSEUM PARIS/ ANNAM/ DÉCION DE QUANG NAM/ AUNHA TRANG/ C ${ }^{\text {te }}$ DE BARTHÉLEMY 1899 [brown label, typed black in]// Dans le Aroyos/ des Mois/ à 17000 m . d'altitude [brown label, handwritten in black ink, unknown handwriting]// MUSEUM PARIS/ Annam/ Ct Barthelemy/ 1899 [brown label, MUSEUM PARIS printed in black ink, rest handwritten, unknown handwriting]// MUSEUM PARIS/ COLL MAURICE REGIMBART/ 1908 [white label with thin black border, typed black ink]//

LECTOTYPE [red label, typed black ink]//" (1 ex. MNHN); paralectotype (1 $q$ pinned) same labels as previous except without the final two MUSEUM PARIS labels and with "TYPE [red label, typed black ink]// PARALECTOTYPE [red label, typed black ink]//" (1 ex. MNHN).

Dineutus (Porrorhynchus) landaisi latilimbus Ochs, 1926 lectotype here designated (1 đ pinned, Fig. 16D): " $\delta^{\lambda}$ [white label, printed black ink]// China/ Insel Hainan/ 10.-25.III.09/ H.Schoede S.G. [beige label, typed black ink, except date which is handwritten in black ink]// Coll./ G.Ochs [white label, typed black ink]// Para-/ typoid [red label with black border, typed black ink]// latilimbus Ochs [beige label with black border, handwritten in ink, handwriting appears to be Ochs']// LECTOTYPE [red label typed black ink]" (1 ex. SMF); paralectotype (1 $q$ pinned) same as previous except with $q$ label and without latilimbus Ochs label and "PARALECTOTYPE [red label, typed black ink] (1 ex. SMF).

Type designation: Régimbart in his original description of $P$. landaisi mentions having examined 3 specimens ( 1 male and 2 females) from Tonkin, collected by M. A. Landais (Régimbart 1892a). The exact locality given by Régimbart (1892a) is Ban-Khau, to the south of Cao-bang. Of the material examined in the MNHN Régimbart collection there are 4 specimens with handwritten labels by Régimbart with Landais listed as the collector. Three are listed as being from Tonkin ( 1 male with a glued head and thorax and 2 intact female specimens), the other specimen has only Tuyen Quan as the locality. Therefore, we consider the 3 specimens with Tonkin on the label as the original syntype series. As the only specific locality provided by Régimbart mentions Cao-Bang, we here designate the female specimen with Cao-Bang as the locality and Landais as collector, the lectotype. This specimen is also completely intact. The male specimen from the syntype series, while carrying a type label, is heavily damaged and has not been formerly designated as a lectotype. For this reason we consider the male and the second female specimen as paralectotypes.

Ochs (1926) in his original description of P. l. latilimbus does not specify how many specimens were examined, but does describe both a male and female, implying more than a single specimen involved. Since no specimen was explicitly designated as the holotype the series must be regarded as syntypes. We here designate the large male specimen as the lectotype for $P$. l. latilimbus and the remaining specimens from the syntype series as paralectotypes. A lectotype was also designated for $P$. barthelemyi in order to stabilize the nomenclature. The male with its aedeagus dissected and available for study was selected as the lectotype.

## Additional material examined:

CHINA: leg. G. Liu (1 ex. MCZ); Hainan Island: leg. J Whitehead (1 ex. BMNH), Kiung-ah an Dist., Mt. ran go, 21-22.v.1935, leg. P. K. To (2 ex. BPBM), 5 km NE Tian Chi, Jianfeng mtns, 800m, 22.i.1996, leg. Jäch (4 ex. NHMW), same as previous except: leg. Ji \& Wang (1 ex. NHMW), Jianfengling, 8.ix.1938, leg. "Protector" (1 ex. ICRI), same locality as previous except: 8.xii.1981, leg. B.R. Li (1 ex. ICRI), same as previous except: 22.ii.1982, leg. R.L. Pan (1 ex. ICRI), same as previous except: leg. H.Q. Chen (2 ex. ICRI), Jianfengling Mts., Tiachi Lake env., BiSHU VILLA, $18^{\circ} 44^{\prime} 40^{\prime}{ }^{\prime N}$, $108^{\circ} 50^{\prime} 41^{\prime \prime} E, 950 \mathrm{~m}, 9-11 . v .2011$, leg. M. Fikáček, V. Kubeček \& L. Li , at light (4 ex. NMPC); Tibet: Zayu co., Xiachayu, vii.2011, leg. Li Jingke (1 ex. NHMW); Yunnan: Kunming ("Yunnanfou"), ZML. 2010/ 336 (1 ex. MZLU). VIETNAM: "Annam": 1895, leg. Barthélemy, coll. C.L. Legros (1 ex. MNHN) "Tonkin": (4 ex. NHMW), "Tonkin": ZML. 2010/ 341 (1 ex. MZLU); Bac Kan ("Backan"): viii.1907, leg. P. Lemée,

Oberthur Coll./ 1909-159. (8 ex. BMNH); same as previous except: Oberthür Coll. (1 ex. MNHN), same as previous except: ZML. 2010/ 337-340 (5 ex. MZLU); Bắc Quang: "BacQuang", "Entre Hagiang et Vinh-Tuy", "Vallées de la Haute Riv. Claire", 1908, leg. J. de Retz (2 ex. MNHN); "Thatkhé", coll. R. Peschet ( 2 ex. MNHN). Kon Tum: ca. 20 km NE Ngoc Linh, 14 m trib. of Ngoc Mi River, , $15^{\circ} 08^{\prime} 23.5^{\prime \prime N} 107^{\circ} 54^{\prime} 40.2^{\prime \prime} \mathrm{E}, 980 \mathrm{~m}, 10 . \mathrm{ix} .1998$, leg. B. Hubley, D.C. Currie, \& M. Tseng, $2^{\circ}$ tropical forest, ROM 982314, (2 ex. ROME); Lào Cai: ("Laokay"), in WNW part, 12-13.viii.1934, leg. Ernest R. Tinkham (1 ex. ICRI), Bao Hà ("Bao-Ha"), 24.x.1923, leg. H. Stevens, Sladen-Goodman/ Trust Exped./ B.M.1924-329. (2 ex. BMNH); Nghệ An: Pu Mat ntl. Prk. Moi River, $18^{\circ} 57.085^{\prime} \mathrm{N} 104^{\circ} 48.746^{\prime} \mathrm{E}, 241 \mathrm{~m}, 13 . v i i .2007$, leg. Sites \& Trung, rocky stream, L-1014, (6 ex. UMRM), W of Con Cuong, Khe Moi Forestry Camp, Keh Moi River, $18^{\circ} 56^{\prime}$ N $104^{\circ} 49^{\prime} \mathrm{E}, 308 \mathrm{~m}, 27 . x .1994$, leg. DC Currie, tropical forest, ROM 946108, ROMEnt Spec. No. 17526-17528 (3 ex. ROME), ca. 25 km SW of Con Cuông, Khe Moi River Forestry Camp, $18^{\circ} 56^{\prime} \mathrm{N} 104^{\circ} 49^{\prime} \mathrm{E}, 308 \mathrm{~m}, 4 . \mathrm{vi} .1995$, leg. B. Hubley, pool in Khe Moi River, ROM 956157, ROMEnt Spec. No. 2145, 4595, 4607, 4619, 4643 (7 ex. 5 ROME, 2 CNC); Quảng Ngãi: "Vie Klong", 97 km NE of Kon Tum ("Kontum"), 1140m, 10.vi.1960, leg. R.E. Leech (1 ex. BPBM) Tuyên Quang "Tuyen quan": "Ruiss. Affluents de la Rivieu Claire", leg. Capc A. Landais (1 ex. MNHN); Uncertain locality within Vietnam: "Haut Tonkin": "Rég de Bac Ken Ha-Giang, Quan-Ba et Yen-Minh": 1918, leg. F. de Broissia (3 ex. MNHN).
Uncertain localities: "Kouy-Tchéo": 1909, leg. P. Cavalerie (2 ex. MNHN), "Kouy-Tchéo": "Rég. de Pin-Fa", 1909, leg. P. Cavalerie ( 23 ex. MNHN), same as previous except: coll. R. Peschet ( 3 ex. MNHN); same as previous except: coll. C.L. Legros (10 ex. MNHN).

Diagnosis: Labrum elongate and subtriangular. Antenna with 7 complete flagellomeres and an $8^{\text {th }}$ typically only complete along its posterior face. Yellow lateral margins nearly complete on elytra, extending to just anteriad of elytral apices, interrupted in basal $1 / 3$ by mediad constriction of yellow margin, associated in males with swelling for proleg reception. Elytral apices spinose, apicolaterally with saw-tooth-like serration, sutural angle produced to a short point, two parasutural spines of similar size, last saw-tooth-like spine at the epipleural angle larger and more projecting than rest.
 11.9 mm .

Habitus. Largest member of genus; most specimens normally elongate oval in body form, attenuated anteriorly in large males; in lateral view strongly convex, greatly humped in scutellar region, depressed posteriorly; in anterior/posterior view very steeply sloped towards lateral margins from strongly humped scutellar region.
Coloration. Dorsally head, pronotum, elytra olive green; base of labrum yellow basomedially; pronotum, elytra with yellow lateral margins; turquoise blue reflections apicolaterally just posteriad to end of yellow lateral margins; venter yellow to yellowish orange; ultimate maxillary palpomere black, except for apex; prolegs with tibia black in proximal $1 / 2$, profemur anteriorly with black ventral border for most its length.
Head. Dorsally vertex with even covering of weakly impressed punctures, separated from nearest puncture by ca. 2 x to 3 x diameter of single puncture; interorbital ridge without yellow margin, similarly colored as vertex; frons similarly punctate as vertex, fronto-lateral margins lightly wrinkled, frontoclypeal suture with posterior margin nearly straight, lateral margins nearly straight, meeting posterior margin at ca $120^{\circ}$ angle; clypeus with punctation most evident at
anterior margin, punctures separated from nearest puncture by ca. 2 x to 3 x diameter of one puncture, becoming more densely spaced anteriorly; antennal flagellum with 7 complete flagellomeres, $8^{\text {th }}$ incomplete suture present on posterior face; labrum sub-triangular, punctation absent basomedially in association with yellow coloration, strongly present apically, punctation well impressed, dense, separated by 1.5 x to 2 x diameter of single puncture; maxillary/labial palpi dissimilar in shape, maxillary palps broader, shorter with asymetrical dorsal/ventral margins, ventral margin more strongly curved than dorsal margin, labial palps narrower, more elongate with anterior/posterior margins more similar, anterior margin nearly straight, posterior margin weakly curved.
Thorax. Pronotum densely punctate, punctation consists of medium sized, well impressed punctures separated from nearest puncture by $<1 \mathrm{x}$ to 2 x diameter of single puncture, reticulation less impressed medially, becoming more well impressed laterally, very shallow transverse depression often present medially, lateral marginal depression present; Protrochanteric setose patch situated paramedially; protibial spine projecting anterolaterally; male protarsi wide, fairly dorsally convex, shape as in Fig. 6A, ultimate protarsomere of male ca. $<2 \mathrm{x}$ as long as wide; ultimate protarsomere of female ca. 2.5 x length of penultimate; elytra with reticulation effaced in scutellar region/along elytral suture, reticulation present apically/laterally, being most strongly impressed marginally, elytral disc with even covering of well impressed, medium sized punctation, distance between nearest punctures ca. $1 \mathrm{x}-2 \mathrm{x}$ diameter of one puncture; lateral marginal depression broad, expanded posteriad to humeral region; yellow lateral margin nearly complete, ending just anteriad to elytral apices, interrupted in basal $1 / 3$ by constriction, in males associated with swelling created by cavity for proleg reception, apicolateral margins of elytra with triangular saw-tooth-like spines, final spine at epipleural angle elongate, elytral apices (Fig. 1B) spinose, with 2 parasutural spines, sutural angle produced; mesosternal apex noticeably acuminate; mesocoxae/metacoxae similar, mesocoxal process broadly rounded, without projecting process; male mesotarsal claws as in Fig. 6D, ventral margin broadly rounded, anterior claw apically narrowed; metacoxal process as in Fig. 6B, without distinct apicolateral corners, without sinuate lateral margins.

Genitalia. Aedeagus (Fig. 7A - C, E-G, I-K) with median lobe shorter than parameres, running ca. $5 / 6$ their length, weakly to moderate laterally expanded in apical $1 / 3$, strongly acuminate in apical $1 / 6$, apex truncate, in lateral view apex briefly strongly curved dorsally; parameres in dorsal view weakly laterally expanded in apical $1 / 2$, narrowly rounded, medially reflexed after apical $1 / 4$, basally strongly constricted; in lateral view ventral margin of parameres strongly curved anteriorly to posteriorly after basal $1 / 3$. Female reproductive tract (Fig. 13A) with large tubiform spermatheca; gonocoxae with lateral margin straightly angled towards apex, apex obliquely truncate.

Sexual dimorphism. Males tend to be larger in size than females, some males exhibit a broader body form, having their outline laterally expanded posteriad to elytral midlength, giving large males a more attenuated feel, whereas females are more evenly narrowed anteriorly and posteriorly. The spines of the elytral apices tended to be longer and more pronounced in females, whereas in very large males they tended to be smaller and more blunt.

Variation. There a considerable amount of size variation in this species in terms of body form (Fig. 7D,H,L). Several populations had very large males, especially those from Vietnam (Fig.

7H), while some like those from Hainan Island, China, had very small and narrow males (Fig. 7D), but also some of the largest. Body form tended to change along with size as noted above. The male aedeagus also showed some variation, which is discussed in the discussion section below.

Differential diagnosis: Porrorhynchus landaisi can be distinguished from all other species of Porrorhynchus being dorsally olive green with nearly complete yellow lateral margins of the elytra, ending just anteriad to elytral apex, being interrupted in basal third by a mediad constriction, and in the form of the spinose elytral apices with two parasutural spines.

Distribution: Fig. 14. Known primarily from southern China and northern Vietnam; found as far west as Zayü Co. Tibet in China (Porrorhynchus sp. in (Jäch et al. 2012), through southern China, Vietnam as far south as the Central Highlands, and east to Hainan Island, China.

Biology: This species is lotic, being known from forested streams based on the label data.
Discussion: This is the largest species of gyrinid known, approaching 30 mm in length! The largest specimen examined during this study was ca. 29.5 mm 's from the MNHN in the Legro's collection, including labrum and abdomen. The next largest species of gyrinid known is Dineutus macrochirus Régimbart, 1899 reaching 22.9 mm (without the abdomen) from New Guinea (Brinck 1984).

Porrorhynchus barthelemyi (Fig. 16E) was described from specimens from the Central Highlands region of Vietnam, inhabited by the Degar indigenous ethnic group, called the "mois" by the French. These specimens are more elongate and narrow in dorsal habitus and the dorsal punctation is larger and denser than other populations of $P$. landaisi. However these are the only characters with which these specimens differ from others of $P$. landaisi, thus the name is here synonymized as it represents variation in a more southern population. Furthermore the variation of a narrow body form is exhibited in other populations of $P$. landaisi. The ventral surface of the "barthelemyi" specimens appears to have become unnaturally discolored in certain sclerites (Fig. 16E). Porrorhynchus landaisi latilimbus was described by Ochs from Hainan island, however there is absolutely no distinguishable morphological differences between the Hainan populations (Fig. 6A,B,C,D and Fig. 7A,B,C) and those from the mainland, thus this subspecies is here synonymized.

A very uniquely broad specimen (Fig. 7L) was recently collected from Tibet, collected at a light trap (Jäch et al. 2012). The specimen unfortunately is damaged, missing its labrum and was the only one of its kind collected (Fig. 7L), preventing further access of the populational variation from this area. Aside from the broader habitus the other morphological features are within the variation (Fig. 6I,J,K), including the aedeagus (Fig. 7I, J,K). This extends the known range of the species much further west than previously known.

This species has a very unique antennomere count for gyrinids, having seven distinct, but an 8 th noticeable along its posterior face, most other genera having either 9 or 6 , with only Enhydrus previously known to possess seven (Miller and Bergsten 2012).

No information is available on this species potential for sensitivity to water quality, but given all other species of Porrorhynchus are, it is also likely sensitive as well. This species has a
more northeastern distribution in Southeast Asia (Fig. 14) and one that is considerably smaller than that of $P$. marginatus.

We propose the common name of the Splendid Snouted Whirligig for $P$. landaisi.

## Porrorhynchus (Porrorhynchus) marginatus Laporte, 1835

Trigonocheilus rostratus: Dejean 1833: 59 [manuscript name and nomen nudum, synonymy by Aubé, 1838a: 406], Porrorhynchus marginatus Laporte, 1835: 108; Porrorhynchus marginatus: Laporte 1835: 108 [redescription]; Porrorhynchus marginatus: Aubé 1838a: 406, Pl. 46 fig. 4 [redescription and habitus image]; Trigonocheilus rostratus: Dejean 1836: 67 [manuscript name and nomen nudum, synonymy by Aubé, 1838a: 406]; Porrorhynchus marginatus: Aubé 1838b: 759 [redescription]; Porrorhynchus marginatus: Hope 1838: 145 [list of type species]; Porrorhynchus marginatus: Laporte 1840: 170 [redescription]; Trigonochilus rostratus: Agassiz 1846: 377 [unjustified emendation of Trigonocheilus rostratus, nomen nudum]; Porrorhynchus marginatus: Newman 1847: 48 [specimen holdings]; Porrorhynchus marginatus: Desmarest 1851: 224, fig. 399 [redescription and dorsal habitus]; Porrorhynchus marginatus: Lacordaire 1854: 440 [minor description]; Porrorhynchus marginatus: Wood 1874: 69, fig. 28 [redescription, illustration]; Porrhorhynchus marginatus: Régimbart 1877: 110, Pl. 6 fig. 3 [revision, elytron]; Porrhorhynchus tenuirostris Régimbart, 1877: 110, fig. 4 [description, dorsal habitus]; Porrhorhynchus marginatus: Régimbart 1882: 427, Pl. 12, fig.49, fig. 49a [redescription, range extension, elytron, foreleg]; Porrhorhynchus marginatus var. tenuirostris: Régimbart 1882: 428 [new status, redescription]; Porrhorhynchus marginatus: Régimbart 1884: 270 [checklist]; Porrhorhynchus marginatus var. tenuirostris: Régimbart 1884: 470 [checklist]; Porrorhynchus marginatus: Hagen 1890: 228 [checklist]; Porrorhynchus marginatus: Duncan 1891: 311 [minor description]; Porrhorrhynchus marginatus: Régimbart 1892a: 740 [checklist]; Porrhorrhynchus var. tenuirostris: Régimbart 1892a: 740 [checklist]; Porrhorrhynchus marginatus: Régimbart 1902: 5 [distribution]; Porrhorrhynchus marginatus: Régimbart 1907: 152 [distribution]; Porrorhynchus marginatus: Laporte 1910: 170 [redescription]; Porrhorrhynchus marginatus: Zimmermann 1917: 139 [locality and holdings information]; Porrhorrhynchus marginatus: Peschet, 1923: 122 [review] Porrorrhynchus marginatus: Hatch 1925: 450 [minor description]; Dineutus (Porrorhynchus) marginatus: Ochs 1926: 139 [new status, checklist]; Dineutus (Porrorhynchus marginatus tenuirostris: Ochs 1926: 139 [new status, checklist]; Dineutus (Porrorhynchus) marginatus mjöbergi: Ochs 1926: 139 [nomen nudum, checklist]; Dineutus (Porrorhynchus) marginatus mjöbergi Ochs, 1926: 193 [original description] new synonymy; Dineutus (Porrorhynchus) marginatus: Ochs 1927b: 242 [distribution, habitat note]; Dineutus (Porrorhynchus) marginatus: Ochs 1927a: 116 [distribution]; Dineutus (Porrorhynchus) marginatus: Ochs 1928: 44 [distribution]; Dineutus (Porrorhynchus) marginatus mjöbergi: Ochs 1928: 44 [redescription]; Dineutus (Porrorhynchus) marginatus: Ochs 1929a: 248 [distribution]; Dineutus (Porrorhynchus) marginatus: Ochs 1930: 16 [catalog]; Dineutus (Porrorhynchus) marginatus mjöbergi: Ochs 1930: 17 [catalog]; Dineutus (Porrorhynchus) marginatus tenuirostris: Ochs 1930: 17 [catalog]; Dineutus (Porrorhynchus) marginatus: Ochs 1931: 472 [distribution]; Dineutus (Porrorhynchus) marginatus: Ochs 1937: 111 [locality data]; Dineutus (Porrorhynchus) marginatus: Ochs 1940b: 33 [distribution, habitat, variation]; Dineutus (Porrorhynchus) marginatus: Ochs 1940a: 5 [locality data]; Dineutus (Porrorhynchus) marginatus: Ochs 1953: 220 [locality data]; Porrorhynchus (Porrorhynchus)
marginatus: Brinck 1955: 103 [new status]; Porrorhynchus marginatus: Balke et al. 2004: 570, Fig. 5A [distribution, dorsal habitus]; Porrorhynchus marginatus: Jäch et al. 2012: 66 [distribution]. Porrorhynchus marginatus: Miller and Bergsten 2012: figs. 9D, 12B, 15D, 16A, 20B [morphology].

## Type material examined: Porrorhynchus marginatus Laporte, 1835: Holotype by monotypy

 (1 $\&$ card mounted, Fig. 17A) "Rostratus/ (Java.) [handwritten black ink on card mount base, handwriting unknown]//"(1 ex. MVMA).Porrorhynchus tenuirostris Régimbart, 1877 lectotype here designated ( $1 \curvearrowright$ pinned, Fig. 17D) "MUSEUM PARIS/ COCHINCHINE/ HARMAND 1876 [white label, printed black ink]// yellow disk [underneath is handwritten 9/668 in black ink]// 7 [handwritten in black ink on small beige square]// MUSEUM PARIS/ MUSEUM PARIS COLL MAURICE REGIMBART/ 1908 [white label with thin black border, type black ink]// TYPE [red label, black ink]// LECTOTYPE [red label, black ink]//" ( 1 ex. MNHN); paralectotypes here designated (3: 1 § pinned, 2 q pinned) same labels as previous except without yellow disk or beige square and with "PARALECTOTYPE [red label, black type]//" ( 3 ex. MNHN).

Dineutus (Porrorhynchus) marginatus mjoebergi Ochs, 1926 lectotype here designated (1 đ pinned, with aedeagus on point, Fig. 17G) " $\widehat{\text { }}$ [white label, typed black ink]// Mt. Dulit/ 3,500 f. [beige label, typed black ink]// Coll./ G.Ochs [white label, typed black ink]// Cotypus [red label with black borders, typed black ink]// P. marginatus/ subsp. mjobergi/ Type! Ochs/ 1924 [beige label, handwritten in ink, handwriting appears to be Ochs']// mjobergi Ochs [beige label with black border, handwritten in ink, handwriting appears to be Och's]// LECTOTYPE [red label, typed black ink]//" (1 ex. SMF); paralectotype here designated (1 $\delta^{\lambda}$ pinned) same as previous except $q$ symbol label and without mjobergi Ochs label and with "PARALECTOTYPE [red label, typed black ink] (1 ex. SMF). Nomenclatural note. According to article 32.5.2.1 of The Code (ICZN 1999), the name mjöbergi must be emended to mjoebergi, so it is here emended.

Type designation. Régimbart (1877) mentions having many examples from Cochinchine from Dr. Harmand, with Phu-Quoc in quotes. In the MNHN collection there are numerous specimens collected by Harmand, however, only 4 specifically state Cochinchine on the label and as being collected in 1876. There are many more specimens in the Régimbart collection collected by Harmand, but these have Lakhon as the locality and the collection date as 1878 . Given that the publication date is 1877 , only those aforementioned 4 specimens are here considered as part of the syntype series. The specimen selected as the lectotype (Fig. 17D) is a male specimen already possessing a label that says "TYPE".

## Additional material examined:

INDONESIA: "Borneo": (3 ex. NMPC), "Borneo": Coll. Mus. Vindob. (1 ex. NHMW), "Borneo/ Sunda Isl." (2 ex. UMRM), "Borneo": 1891, leg. Chaper (4 ex. MNHN), same as previous except: 1926, leg. E. Mjöberg, (5 ex. MCZ), "Bivang Riv.", 1926, leg. E. Mjöberg (67 ex. MCZ), "Riv. Mandar", 1897, leg. Fr. Buffat, Coll. MAURICE REGIMBART 1908 (1 ex. MNHN), same as previous except: 7.xi.1897, Coll. MAURICE REGIMBART 1908 (9 ex. MNHN), "Bohac" (1 ex. NMPC); East Kalimantan: Kac. Pujungan, Kayan-Mentarang Nat. Reserve, $2^{\circ} 52^{\prime} \mathrm{N} 115^{\circ} 49^{\prime} \mathrm{E}, 378 \mathrm{~m}$, 20.ii.-4.iii.1993, leg. D.C. Darling, lowland diptero. Forest/

WWF station, Lalut Birai/ rocky stream (Nggeng), IIS 930003, ROMEnt Spec. No. 1739, 3150 (4 ex. 2 ROME, 2CNC), same as previous except: 6.vi.1993, leg. D.C. Darling \& Rosichon U., lowland diptero. Forest/ WWF station, Lalut Birai/ small stream above Nggeng River, IIS 930330, ROMEnt Spec. No. 3098, 3141 (3 ex. ROME); West Kalimantan ("Borneo"): "Riv. Sambey", near Ngabang ("pres Ngabang"), 1897, leg. J.B. Ledru, ZML. 2010/ 342-344 (3 ex. MZLU); Java: ( 1 ex. NMPC), Java: Coll. MAURICE REGIMBART 1908 (1 ex. MNHN), Java: leg. O. Koechlin (1 ex. NMPC), "Jiansberge", Coll. M. SÉDILLOT (2 ex. MNHN); "Sumatra": Coll. MAURICE REGIMBART 1908 ( 1 ex. MNHN), "Sumatra": "Balighe", x.90-iii.91, leg. E. Modigliani (8 ex. ZMUC), same as previous except: ZML. 2010/ 345,346,347,354 (4 ex. MZLU), same as previous except: Coll. MAURICE REGIMBART 1908 ( 1 ex. MNHN), "Sumatra": "Palembang", coll. R. Peschet (7 ex. MNHN),"Sumatra": "Manna" (likely the Manna River), 1901, leg. M. Knappert, Coll. Mus. Vindob. (1 ex. NHMW), same as previous except 1902, Coll. Mus. Vindob. (1 ex. NHMW), "Sumatra": "Indrapoera", Coll. MAURICE REGIMBART 1908 ( 1 ex. MNHN); North Sumatra: "Ober Langkat", "Sumatra/ Sunda Isl.", "ix-x" (2 ex. UMRM), Bohorok, vi.1975, leg. P. Blum (1 ex. NHMW), Nias, "Hili Madjedja", N. Nias, "4 de trim".1895, leg. I.Z. Kannegieter, ZML.2010/ 348-350 (3 ex. MZLU), Gunung Leuser Ntl. Prk., Bukit Lavang env., 26.iv.2004, leg. D. Trávníček, Jiří Hájek Collection (2 ex. NMPC); West Sumatra: "NSG Lemba Harau", 15km NE Payakumbuh ("Payakumbu"), 11.ii.1991, leg. Jäch, , 12b, (3 ex. NHMW), Siberut Island, Salappa vill. Env., Labuan Bajau, ii.2006, leg. St. Jakl, Jiří Hájek Collection (2 ex. NMPC), same as previous except: xii.2005, leg. St. Jakl, Jiří Hájek Collection (2 ex. NMPC), Siberut Island, "Toteburu-Bakeuluk, 17.ii.1991, leg. Schödl, 22 (13 ex. NHMW), same as previous except: leg. Jäch, 22 (26 ex. NHMW), Siberut Island, "Toteburu", "W Muarasiberut", 16.ii.1991, leg. Jäch, 21 (2 ex. NHMW). LAOS: Champasak: Ban Houei Khong ("Houei Kong"), iii.1965, J.A. Rondon Collection (5 ex. BPBM), Muang Paxong, Ban Thongvay, Bolaven plateau, $15^{\circ} 14.398^{\prime} \mathrm{N} 106^{\circ} 31.806^{\prime} \mathrm{E}$, 10001200m, 7-16.vi.2008, leg. A. Solodovnikov \& J. Pedersen, disturbed primary rainforest (2 ex. ZMUC); Khammouane: Phon Tiou, 11-12.vi.1965, J.A. Rondon Collection (3 ex. BPBM), Ban Khoun Nguen env., $18^{\circ} 07^{\prime} \mathrm{N} 104^{\circ} 29^{\prime} \mathrm{E}, 250 \mathrm{~m}, 4-16,25-30 . x i .2000$, leg. E. Jendek \& P. Pacholátko (227 ex. NMW); Luang Namtha: "Nam Youan", 7 km N of Ban Muangsing, $21^{\circ} 14^{\prime} \mathrm{N}$ $101^{\circ} 8^{\prime}$ E, 1.v.1997, leg. W.J. Rainboth \& S. Virawong ( 2 ex. UMMZ), 10-20km NW Luang Namtha, 600m, 17.vi.1996, leg. Schillhammer, 29, (2 ex. NHMW); Luang Prabang: "A THENG", 1888, leg. A. Pavie ( 12 ex. MNHN), Houay Houp ("Houay Houn"), ca. 16km upstream from mouth at Nam Ou, 3.iii.1998, leg. W.J. Rainboth \& K.P. Bounkhamvongsa (2 ex. UMMZ); Sainyabuli ("Sayaboury"): 18.viii.1966, leg. "Native Collector" (1 ex. BPBM), Sainyabuli ("Sayaboury"), 29.iii.1966, leg. "Native Collector" (5 ex. BPBM), same as previous except: 20.v. 1966 (3 ex. BPBM); Vientiane: "Ban Van Eue", 15.vii.1965, leg. "Native Collector" (2 ex. BPBM), same as previous except: 1-15.ix. 1965 (2 ex. BPBM), same as previous except: 30.ix. 1965 (7 ex. BPBM), same as previous except: 30.xi. 1965 ( 5 ex. BPBM), same as previous except: 30.i.1966 (1 ex. BPBM), same as previous except: 15.ii. 1966 ( 1 ex. BPBM), same as previous except: 16.iii.1966 (2 ex. BPBM), same as previous except: 15.v. 1966 (1 ex. BPBM), same as previous except: 31.v. 1966 ( 22 ex. BPBM), same as previous except: 30.xi. 1966 ( 1 ex. BPBM), same as previous except: $15 . x i i .1966$ ( 2 ex. BPBM), same as previous except: $30 . i i i .1967$ ( 1 ex. BPBM), same as previous except: 30 .viii. 1967 ( 1 ex. BPBM), same as previous except: 1-15.ix. 1967 (1 ex. BPBM), Phou Khao Khouay ("Phou Kou Khouei"), 15.ii.1966, leg. "Native Collector" (8 ex. BPBM), same as previous except: 15.iv.1966, leg. (3 ex. BPBM), Phou Khao Khouay, $18^{\circ} 20.369^{\prime} \mathrm{N} 102^{\circ} 48.523^{\prime} \mathrm{E}, 700-800 \mathrm{~m}, 26-31 . \mathrm{v} .2008$, leg. A.

Solodovnikov \& J. Pedersen, strongly disturbed primary rainforest (1 ex. ZMUC), Phou Khao Khouay Ntl. Pr. env., Tad Luek Waterfall, 200m, 1-8.vi.1996, leg. Schillhammer, 15 (1 ex. NHMW), same as previous except: 300m, 16 (4 ex. NHMW); Xiangkhouang "Xieng Khouang": 21.iii.1915, leg. Vitalis, coll. R. Peschet ( 1 ex. MNHN); uncertain locality within Loas: "Bas-Laos": ii.1900, leg. Barthélemy (1 ex. MNHN); "Lakhon", 1878, leg. Harmand, Coll. MAURICE REGIMBART 1908 (13 ex. MNHN); "Tonkin" 1886, leg. Langue (4 ex. MNHN),"Tonkin", "central", 1911, leg. A. Krempf (1 ex. MNHN), "Tonkin", "region de ChimHua et de Tuyen-Quan", 1901, leg. A. Weiss ( 2 ex. MNHN); "Tonpheng", 29.v.1966, leg. "Native Collector" (1 ex. BPBM), same as previous except: 16.xii. 1966 (1 ex. BPBM). MALAYSIA: Melacca: leg. de Morgan ( 1 ex. MNHN); Pahang: 4 km W Rompin, "Selendang", 29.iv-6.v.1993, leg. I. Jenis (3 ex. NHMW), same as previous except: leg. Strba (5 ex. NHMW), Pahang: Taman Negara Ntl. Prk., $3^{\circ} 36^{\prime} 52^{\prime \prime} \mathrm{N} 102^{\circ} 29^{\prime} 11^{\prime \prime} \mathrm{E}$, $21 . v i i i .2003$, leg. G. Svenson, GJS21080301 (5 ex. MSBA); Perak: Kwala-Kangsar, Coll. MAURICE REGIMBART 1908 (2 ex. MNHN), Kwala-Kangsar, 1902, leg. Grubauer, Coll. Mus. Vindob. (3 ex. NHMW); Sabah: Tawau Hills, Tawau River, 7-10.vi.1998, leg. J. Kodada \& F. Ciampor (20 ex. NHMW); Sarawak: "Baram", 1910, leg. H.W. Smith (1 ex. MCZ), "Tuba", 18.i.1979, leg. Gärdenfors, Hall, Hansson, \& Samuelsson (15 ex. MZLU). Kapit, ca. 40km SE Kapit, iii.1994, leg. J. Kodada (9 ex. NHMW), Kapit, "Rumah Ugap vill.", Slut riv., 3-9.iii.1994, leg. J. Horák (12 ex. NHMW), same as previous except: leg. J. Horák, Jiří Hájek Collection (3 ex. NMPC), Kuching, 80km S Kuching, Kampung Ana Rais, 18.ii.1993, leg. M. Jäch (1 ex. NHMW), Miri, 8 km NE Bario, stream in jungle, 24.vi.2003, leg. D. Trávníček, Jiří Hájek Collection (1 ex. NMPC), Miri, Kelabit Hi., 6km E Bario, Pa Ukat, 1000m, 27.ii.1993, leg. M. Jäch, 15 (1 ex. NHMW), Miri, Gunung Mulu Ntl. Prk. ("Mulu N.P."), Long iman, 4.iii.1993, leg. M. Jäch, 20 (1 ex. NHMW), same as previous except: 3-5.iii.1993, leg. Zettel, (14 e) (1 ex. NHMW), same as previous except: small stream, $4^{\circ} 0.207^{\prime} \mathrm{N} 114^{\circ} 49.267^{\prime} \mathrm{E}, 20 . x .2006$, leg. K.B. Miller, KBM20100601 (5 ex. MSBA), Miri, Sungai Kelimau, "ca. 1.3 km above mouth", $3^{\circ} 24^{\prime} 36^{\prime \prime N} 114^{\circ} 3^{\prime} 00$ "E, 12.ix.1980, leg. M.H. Ang, Acc. 4302-3 (EN-0002) (2 ex. ROME), Sri Aman, Batang Ai Ntl. Prk., Engkari riv., "E Bandar Sri Amman", 19-20.ii.1993, leg. Zettel, 7 (3 ex. NHMW).
MYANMAR: Mandalay: 8km E Pyin Oo Lwin, "Pwe Kauk Wf.", $22^{\circ} 03.523^{\prime} \mathrm{N} 96^{\circ} 31.956^{\prime} \mathrm{E}$, 1070m, 19.x.1998, leg. Schillhammer, 20 (1 ex. NHMW); Tanintharyi: "Tenasserim", Mus. Westerm (1 ex. ZMUC), same as previous except: leg. Helfer (14 ex. NMPC).
THAILAND: "Siam": Coll. MAURICE REGIMBART 1908 ( 1 ex. MNHN); THAILAND: viii. 1984 (3 ex. ZMUC); Chaiyaphum: Phu Khieo Wildlife Sanctuary, "Nam Prom", 4.iv.1994, leg. W. Shephard, WDS A 1053 (7 ex. NHMW); Chanthaburi: 15km E Chanthaburi City, stream at Philiu Waterfall, 29.i.1995, leg. P.P. Chen (1 ex. NHMW); Chiang Mai: "SoppongPai", 1800m, 1-8.v.1993, leg. Pacholatko \& Dembicky (1 ex. NHMW); Chumphon: Khun Mae Yam Oum Wildlife Sanct., Haew Lome Waterfall water surf., $09^{\circ} 43^{\prime} \mathrm{N} 98^{\circ} 40^{\prime} \mathrm{E}, 122 \mathrm{~m}$, 22.v.2005, leg. Sites, Vitheepradit, \& Prommi, L-785b (2 ex. UMRM); Kanchanaburi: Amphur Sangkhla Buri, Heuy Kob, $15^{\circ} 13^{\prime} \mathrm{N} 98^{\circ} 22^{\prime} \mathrm{E}, 289 \mathrm{~m}$, 13.iv.2002, leg. UMC and CMU teams, gravel stream, L-339 (11 ex. UMRM), Amphur Sai Yok, Thong Pha Phum Reforestation Station, Mae Nam Noi, $14^{\circ} 31^{\prime} \mathrm{N} 98^{\circ} 37^{\prime} \mathrm{E}, 204 \mathrm{~m}$, 12.iv.2002, leg. UMC and CMU teams, L-335 (9 ex. UMRM), 30km N Thong Pha Phum, sm. Mount. Riv., sec. veget, neustic, $14^{\circ} 58^{\prime} 11^{\prime \prime} \mathrm{N}$ $98^{\circ} 38^{\prime} 35^{\prime \prime} \mathrm{E}, 160 \mathrm{~m}, 13 . x i i .2010$, leg. Freitag, 22b (1 ex. NHMW); Loei: Dan Sai, "Lomie Mt.", 1.iii.1955, leg. R.E. Elbel, ZML. 2010/ 356 (1 ex. MZLU), Na Haeo, river bank, 15.v.2003, leg. P. Grootaert, J. Constant, \& K. Smets, Light trap (3 ex. IRSB), Phu Luang Wildlife Sanctuary, 700-900m, 8-14.x.1984, leg. Karsholt, Lomholdt, \& Nielsen (8 ex. ZMUC), Phu Rua Nat. Prk.,

Nam Tok Huay Pai, waterfall, 10.vi.1998, leg. Sites, Simpson, \& Vitheepradit, L-175 (1 ex. UMRM); Mae Hong Son: "Ban Si Lang", 19 19'N 97 59'E, 1200m, 23-31.v.1991, leg. Dembicky (1 ex. NHMW), Huai Sua Tao, v.1992, leg. Dembicky (1 ex. NHMW); Mukdahan: Phu Pa Yon Ntl. Prk., Keang Pho Waterfall, $16^{\circ} 45.368^{\prime} \mathrm{N} 104^{\circ} 14.736^{\prime} \mathrm{E}, 314 \mathrm{~m}, 23 . \mathrm{iv} .2004$, leg. A. Vitheepradit, margin, L-634 (9 ex. UMRM); Nakhon Nayok: Khao Yai Ntl. Prk., ca. 700m, 29.ix-6.x.1984, leg. Karsholt, Lomholdt, \& Nielsen (6 ex. ZMUC); Nakhon Ratchasima: Khao Yai Ntl. Prk., $14^{\circ} 20^{\prime}$ N $101^{\circ} 30$ 'E, 1000m, 26.iii.1992, leg. E. Hüttinger (10 ex. NHMW), Khao Yai N. D., 700-800m, 31.iv.1990, leg. E. Fuller (3 ex. NHMW); Nan: Ban Pha Khap, 1520.v.1992, leg. P. Pecholatko ( 2 ex. NHMW), Mae Charim Ntl. Prk., Nam Wa River, 18³6'N $100^{\circ} 59^{\prime}$ E, 335m, 13.iii.2002, leg. CMU Team (2 ex. UMRM); Phang Nga: Amphur Khura Buri, Tumbon Bang Won, $08^{\circ} 59^{\prime} \mathrm{N} 98^{\circ} 26^{\prime} \mathrm{E}, 45 \mathrm{~m}, 27 . i v .2002$, leg. Vitheepradit \& Kirawanich, gravel stream, L-375 (23 ex. UMRM), Khao Lak surr., 10.i.2003, leg. H. Forster (6 ex. NHMW), Khura Buri , "Baan Tumnang", W of Si Phang Nga Ntl. Prk., 29.xi.2006, leg. H. Zettel, 48 (4 ex. NHMW); Phetchabun: 5 km E Sila, "Mae Nam Pa Sak", 2.iii.1994, leg. W. Shephard, WDS A 1025 (1 ex. NHMW), Khao Kor Non-Hunting Area, Sridit Waterfall, $16^{\circ} 37^{\prime}$ N $100^{\circ} 56^{\prime} \mathrm{E}, 702 \mathrm{~m}$, 11.v.2004, leg. Vitheepradit \& Prommi, water surface, L-677 (1 ex. UMRM); Rayong: Khao Chamao-Khao Ntl. Prk. ("Khao Chamao NP"), 12.xii.1990, leg. Jäch, 14 (1 ex. NHMW); Sakon Nakhon: "Keek poo", 9.xi.1954, leg. R.E. Elbel ZML.2010/ 357 (1 ex. MZLU), "11 km NE Kham Poem", "Huai Ya", 5.iii.1994, leg. W. Shephard, WDS A 1027 (4 ex. NHMW); Saraburi: Khao Yai Ntl. Prk., Lamtok Khlong, 2.i.2008, leg. R.W. Sites, L-1025 (9 ex. UMRM);
Songkhla: Ton Nga Chang Ntl. Prk., stream at Buddhist temple, 30.i.1995, leg. B.J. Nichols, L81 (3 ex. UMRM), same as previous except: 6.vii.1997, leg. R.W. Sites, L-127 (24 ex. UMRM), same as previous except: 9.vi.2001, L-236 (54 ex. UMRM), Khoa Nam Khang ("Khao Nam Chang"), SW Na Thawi ("SW Nathawee"), 13.i.1995, leg. R. Sites \& B. Nichols (5 ex. UMRM), 102 km SE Amphur Hat Yai, Amphur Na Tawee, Khao Nam Khang Ntl. Prk., 6³6'N 100³6'E, 100m, 15.vi.2001, leg. R.W. Sites, L-246 (5 ex. UMRM); Surat Thani: Khao Sok Ntl. Prk., 59.vi.1999, leg. D. Šanc, Jiří Hájek Collection (5 ex. NMPC), same as previous except: 15.1.2003, leg. H. Forster (5 ex. NHMW); Yala: Betong, "Gunung Cang dun vill.", 25.iii-22.iv.1993, leg. J. Strnad (1 ex. NHMW); uncertain locality within Thailand: Khao Yai Ntl. Prk., 14.xi.1988, leg. Jäch, 2 (3 ex. NHMW).
VIETNAM: Đắk Lắk: ca. 2km SE Ban Don, Yok Don Ntl. Prk. H.Q., $12^{\circ} 53^{\prime} \mathrm{N} 107^{\circ} 48^{\prime} \mathrm{E}$, 2231.v.1997, leg. D.C. Darling \& D.C.Currie, UV/MV light, ROM 974004 (3 ex. ROME), same as previous except: leg. D.C. Darling, D.C. Currie, \& A. Guidotti, MV light, ROM 974019 (1 ex. ROME), 8km SW Ban Don, Yok Don Ntl. Prk., Dak Ken R., $12^{\circ} 53^{\prime} N 107^{\circ} 48^{\prime}$ E, 16-22.vi.1997, leg. B. Hubley, Dipterocarp forest, ROM 974105 (40 ex. ROME); Gia Lai: 20 km N of Pleiku, 650m, 9.v.1960, leg. L.W. Quate (1 ex. BPBM), 25 km SW of Pleiku, 400m, 12.v.1960, leg. L.W. Quate (1 ex. BPBM), 40 km NW An Khe, Buon Luoi, $14^{\circ} 10^{\prime} \mathrm{N} 108^{\circ} 30^{\prime} \mathrm{E}, 620-750 \mathrm{~m}$, $28 . \mathrm{iii}-$ 12.iv.1995, leg. Pacholatko \& Dembicky (47 ex. NHMW), An Khe Dist., Tram Lap. Azun R., 2 km NW on trail from forestry building, $14^{\circ} 27^{\prime} \mathrm{N} 108^{\circ} 33^{\prime} \mathrm{E}, 18 . \mathrm{vi} .1996$, leg. B. Hubley \& D.C. Currie, $1^{\circ}$ rainforest edge/ coffee plantation, ROM 961059 (8 ex. ROME), An Khe Dist., 5.2 km NE Tram Lap on forest road, Dacklest River, $14^{\circ} 2^{\prime} \mathrm{N} 1^{\circ} 8^{\circ} 33^{\prime} \mathrm{E}, 900 \mathrm{~m}, 22-23 . v i .1996$, leg. B. Hubley \& D.C. Currie, 200m upstream bridge/ stream margin/ deep pools, ROM 961086, ROMEnt Spec. No. 17529-17536 (8 ex. ROME), An Khe Dist., 3km E. Buoenloy, small stream 6 km on logging road past Cha River, 25.vi.1996, leg. N. Orlov, ROM 961093 (1 ex. ROME); Hòa Bình ("Hoa Binh"): 1919, leg. R.P.A. de Cooman, ZML.2010/ 355 (1 ex. MZLU); Lâm Đồng: Fyan, 900-1000m, 11.vii-9.viii.1961, leg. N.R. Spencer (5 ex. BPBM), 15 km SW Bảo

Lộc, $11^{\circ} 27^{\prime} \mathrm{N} 107^{\circ} 43^{\prime} \mathrm{E}, 900 \mathrm{~m}, 22-24 . \mathrm{iv} .1995$, leg. Pacholatko \& Dembicky ( 8 ex. NHMW); M'Đrăk: E of Buon Ma Thuot ("BanMeThuot"), 400-600m, 8-19.xii.1960, leg. C.M. Yoshimoto (5 ex. BPBM); Nghệ An: Phúc Sơn ("Phuc-Son"), xi-xii, leg. H Fruhstorfer, ZML.2010/ 353 (1 ex. MZLU), same as previous except: Coll. MAURICE REGIMBART 1908 (1 ex. MNHN); Ninh Thuận: Phan Rang ("Pha-Rang"), leg. H. Fruhstorfer, (1 ex. NHMW), same as previous except: ZML.2010/ 351-352 (2 ex. MZLU); Quảng Trị: "Cua Tung", H.C. Fall Collection (1 ex. MCZ), Da Krong Nature Reserve, $16^{\circ} 37^{\prime} \mathrm{N} 106^{\circ} 47^{\prime} \mathrm{E}, 5-10 . v i i .2011$, leg. J. Constant \& J. Bresseel, day collecting, I.G.:31.933 (4 ex. IRSB), same as previous except: $16^{\circ} 37^{\prime} \mathrm{N} 106^{\circ} 47^{\prime} \mathrm{E}$, Light trap, I.G.:31.933 (1 ex. IRSB); uncertain locality within Vietnam: "Houtabo", "SeSouk", 1897, leg. J.M. Bel (1 ex. MNHN); "Karyu Danar": , 200m, 13-28.ii.1961, leg. N.R. Spencer (10 ex. BPBM).
Uncertain locality: "Inde, Bellary, ou Ceylan", 1896, leg. De Morgan (3 ex. MNHN); "Indes Or.", "Marc 1196-36" (1 ex. MNHN); "Cochinchine", 1878, leg. Pierre (2 ex. MNHN); "Cochinchine", ii.1900, leg. Barthélemy (1 ex. MNHN); "Muang You", 26.v, coll. R. Peschet (1 ex. MNHN).
No locality information: "Patra Ignota" (1 ex. MNHN); [illegible handwriting in ink] (1 ex. NMPC); [illegible handwriting in ink], coll. R. Peschet (1 ex. MNHN); ZML. 2010/ 359 (1 ex. MZLU); 334, ZML.2010/ 358 (1 ex. MZLU); leg. Plason, Coll. Mus. Vindob. (1 ex. NHMW); 1871, "Fieber/ Mulmeir", Coll. Mus. Vindob. (1 ex. NHMW); Mus. Westerm (2 ex. ZMUC); [illegible handwriting in pencil] (1 ex. ZMUC).

Diagnosis: Interorbital ridge with yellow lateral margins, labrum highly elongate and in the form of an isosceles triangular. Antenna with 6 flagellomeres. Dorsally olive green to bronzy-brown, yellow lateral margins complete on elytra, extending to elytral apices, nearly always interrupted in basal $1 / 3$ by a dark spot, associated in males with swelling for proleg reception. Elytral apices spinose, apicolaterally with saw-tooth-like spines, sutural angle produced to a short point, one large parasutural spine, last saw-tooth-like spine at the epipleural angle larger and more projecting than the rest, often strongly produced and spinose.

Description: Size: $\begin{gathered} \\ \text { L }: ~ 13.7-18.1 ~ m m, ~ W: ~ \\ 7.6-10.8 \mathrm{~mm} ; ~ 千+\mathrm{L}: 12.2-15.1 \mathrm{~mm}, \mathrm{~W} 7.1-8.7\end{gathered}$ mm .
Habitus. Medium to very large members of genus; body form often tear dropped shaped, broadest posteriad to middle, attenuated anteriorly, especially in large males, other populations elongate oval, evenly attenuated anteriorly to posteriorly, especially in females; in lateral view convex, strongly humped in scutellar region, depressed posteriorly/anteriorly; in anterior and posterior view steeply sloped towards lateral margins from strongly humped scutellar region. Coloration. Dorsally head, pronotum, elytra olive green to bronzy brown in color; labrum yellow basomedially; pronotum/elytra with yellow lateral margins; elytral margin apicolaterally darkly colored, turquoise blue reflections apicolaterally mediad to end of yellow lateral margins in some individuals; venter yellow; ultimate maxillary palpomere, not darkened in color; prolegs with tibia black in proximal $1 / 2$, profemora black apically.
Head. Dorsally vertex with even covering of lightly impressed punctures, separated from nearest puncture by ca. 2 x to 3 x diameter of a single puncture; interorbital ridge with yellow margin; frons similarly punctate as vertex, punctures mostly concentrated apicomedially, fronto-lateral margins very lightly wrinkled to non-wrinkled, frontoclypeal suture with posterior margin nearly straight, lateral margins nearly straight, meeting posterior margin at ca $120^{\circ}$ angle; clypeus with
punctation most evident at anterior margin, punctures separated from nearest puncture by ca. 2 x to $3 x$ diameter of one puncture, becoming more densely spaced anteriorly; antennal flagellum with 6 complete flagellomeres, ultimate flagellomere at least $2 x$ longer than penultimate, trapezoidal in shape; labrum in form of isosceles triangle, punctation absent basomedially, in association with yellow coloration, strongly present apically, punctation well impressed and dense, separated from nearest puncture by .5 x to 1.5 x diameter of one puncture; maxillary and labial palpi similar in shape, both hatchet-form with anterior margin of maxillary palp/ventral margin of labial palp weakly curved, posterior margin of maxillary palp/dorsal margin of labial palp more strongly curved, apex of both truncate.
Thorax. Pronotum with even covering of punctation, punctation consists of finer, weakly impressed punctures separated from nearest puncture by 2 x to 3 x diameter of single puncture, reticulation less impressed medially, becoming very well impressed laterally, very shallow transverse depression often present medially, lateral marginal depression present; Protrochanteric setose patch situated apicolaterally; protibial spine projecting forward; male protarsi not laterally broadened, not noticeably dorsally convex, shape as in Fig. 8A, ultimate male protarsomere ca. 2 x as long as wide; ultimate protarsomere of female ca. 1.5x length of penultimate; elytra with reticulation effaced in scutellar and sutural region, reticulation present apically/laterally, being very strongly impressed marginally, elytral discs with even covering of finely impressed, punctation, distance between nearest punctures ca. $2 \mathrm{x}-3 \mathrm{x}$ diameter of single puncture; lateral marginal depression broad, strongly expanded posteriad to humeral region; yellow lateral margin complete, ending apicolaterally at elytral apices, nearly always interrupted in basal $1 / 3$ by darkly colored spot, in males associated with swelling created by cavity for proleg reception, apicolateral margins of elytra with triangular saw-tooth-like spines, final spine at the epipleural angle often strongly elongate, spine-like, elytral apices (Fig. 1A) spinose, with single parasutural spine, sutural angle produced; mesosternal apex not noticeably acuminate, evenly narrowed basally to apically; mesocoxae and metacoxae similar, mesocoxae with posteriorly projecting process, processes strongly acuminate, almost spine-like (Fig. 8C); male mesotarsal claws as in Fig. 8E with ventral margin flatly rounded, anterior claw not significantly narrowed apically.

Genitalia. Aedeagus (Fig. 10) with median lobe shorter than parameres, running ca. 5/6 their length, parallel sided in basal $1 / 2$, some populations with medial constriction, moderately laterally expanded in apical $1 / 2$, acuminate in apical $1 / 4$, apex truncate, in lateral view apex briefly and strongly curved dorsally; parameres in dorsal view with lateral margins briefly expanded in apical $1 / 3$, setose in apical $1 / 5$, apex with apicomedial margin straight to oblique reflexed in apical $1 / 5$, medial margin arcuately reflexed after apical $1 / 5$ until basal $1 / 2$; lateral margins often constricted in basal $1 / 5$; in lateral view ventral margin of parameres often evenly curved anteriorly to posteriorly. Female reproductive tract (Fig. 13C) with narrow, elongate, tubiform spermatheca; gonocoxae short, lateral margin straightly angled towards apex, apex obliquely truncate.

Sexual dimorphism. Males tend to be larger in size than females, some males exhibit a broader body form, having their outline laterally expanded posteriad to elytral midlength, giving large males a more attenuated feel anteriorly. Females tend to be much more elongate in appearance, being evenly attenuated posteriorly and apically.

Variation. This widely spread species is highly variable in terms of body form and size. Specimens examined from more northern mainland latitudes tend to be more elongate and narrowly oval in overall body form, with a much more acute labrum (Fig. 9B,C,D), with some larger specimens reaching some fairly long body lengths (Fig. 9B). Specimens from along the Malay Peninsula tended to be smaller in overall body length, with a more evenly oval body form, with a broader labrum (Fig. 9F,G,H). Borneo had specimens that ran the gamut of body forms but was notable for having a unique race containing very large and broad males (described as a formal subspecies $P$. mjoebergi, Fig. 9O). The largest specimens of $P$. marginatus came from East Kalimantan (Fig. 9O). These specimens also have relatively blunt metacoxal processes relative to other populations (Fig. 8Q).

Differential diagnosis: Porrorhynchus marginatus can be distinguished from all other species of the genus in having the interorbital ridge with yellow margins, and elytral apices with a single parasutural spine and buzz-saw-like serration apicolaterally. The species most similar to $P$. marginatus is $P$. depressus, however these two species drastically differ in size (with $P$. marginatus normally being much larger), the shape of the labrum (with that of $P$. marginatus being much more elongate and acuminate, and in the form of an isosceles triangle), and finally the distribution's of the two species should easily separate the two.

Distribution: Fig. 14. The most widely distributed species in the genus. Found throughout most of Southeast Asia from as far northwest as Zayü county China (Jäch et al. 2012), east to Vietnam and south to Java, including Borneo, but notably absent from the Philippines and east of Wallace's line.

Biology: Label data support the previous observation (Ochs 1927b) that $P$. marginatus is found in smaller forested streams, often above 100 m and up to $1,000 \mathrm{~m}$. This species has been implicated as sensitive to water quality, being found only in streams in the Malay peninsula not contaminated by tailings from tin mines (Ochs 1927b).

Discussion: This species exhibits great variation in body form (Fig. 9), however, similar variation was not exhibited elsewhere in external morphology (Fig. 8). The northern populations exhibiting a more narrow body form, formally described by Régimbart as P. tenuirostris 1877, exhibit no other significant variation. Régimbart (1882) identified his "tenuirostris' as being mere variation and synonymized himself. Only later would the name be re-instated by the Ochs (1926). Given this name was only based on variation in outline, with other southern populations having a narrow outline (i.e. Fig. 9I) this name is formally synonymized here again.

The aedeagus shows considerable variation (Fig. 10). Most populations have a narrow elongate median lobe, that is weakly constricted medially and strongly acuminate apically. Populations from Sumatra (Fig. $10 \mathrm{I}, \mathrm{J}, \mathrm{K}$ ) have a shorter and broader aedeagus, with a thicker median lobe. However, all populations, including the Sumatran, exhibit the same general form of the median lobe, being laterally expanded in their apical $1 / 2$, then acuminate in the apical $1 / 4$ with a truncate apex, and having the same length to paramere proportions, being just shorter than the parameres. Importantly in lateral view all medial lobes have the apex curved dorsally. The greatest degree in variation was exhibited by the apex of the parameres. Specimens from Sumatra have broad and mostly rounded apices to the parameres (Fig. 10J,K), while other populations have strongly truncate apices (Fig. 10D,E,H,I), and many with the apex more-or-less
evenly rounded (Fig. 10A,F,L,M,N). However all the parameres had a consistent obliquely truncate medial margin to the apex, and a lateral expansion in their apical $1 / 3$. Furthermore the difference in parameres morphology could not be attributed to any single region. The broader median lobe of the Sumatran populations, were the only consistent morphological differences exhibited among the aedeagi of the $P$. marginatus specimens examined. However, this does not correspond to any formally described subspecies.

Porrorhynchus marginatus mjoebergi was described from very large specimens collected from Borneo's Mount Dulit. Other specimens from Borneo were similarly robust and among the largest specimens examined. However the broad body size was not consistent among specimens from Borneo. Specimens from East Kalimantan matched P. m. mjoebergi well and did exhibit a minor unique morphology apart from the more robust dorsal habitus of males. The metacoxal apex Fig. 8Q had broader and more rounded apices. The labrum of the specimen was also broader and rounder than those of other populations. The aedeagus however, exhibited no substantial difference from other populations (Fig. 10N). Given the potential isolation of the population to the mountains of central Borneo and some of the unique features exhibited, $P . m$. mjoebergi may be deserving of subspecies status pending a phylogenetic analysis. However, morphologically, there is nothing to merit continuing this distinction as a formally named taxon, until phylogenetic evidence gives credence to its distinction.

While the species has been suggested to be sensitive to water quality it has a very large range, unlike the two other members of the sensu stricto subgenus. Given its large range and common occurrence in museums, it is not likely of conservation concern currently.

We propose the common name of the Margined Snouted Whirligig for $P$. marginatus.

## Subgenus Rhomborhynchus Ochs, 1926

Type species: Porrorhynchus depressus Régimbart, 1902 by original designation of Ochs, 1926: 65.

Dineutus (Rhomborhynchus) Ochs, 1926: 65 [original description]; Porrorhynchus
(Rhomborhynchus): Guignot 1950: 124 [new status]; Dineutus (Rhomborhynchus): Ochs 1955: 130; Porrorhynchus (Rhomborhynchus): Brinck 1955: 103 [change status]; Porrorhynchus (Rhomborhynchus): Polhemus 2011: 52 [minor description, habitat]

Diagnosis: Medium sized whirligig beetles: 9 - 11 mm . Antennal flagellum with 6 flagellomeres. Labrum ventrally with longitudinal line of setae paramedially. Dorsal eye situated posteriorly with posterior margin located in plane with that of ventral eye. Gular suture complete. Pronotum with transverse impressed line. Elytral margin without significant swelling associated with proleg reception. Males without Protrochanteric setose patch. Profemur without two linear series of large setose clusters, only with a single linear series of small setose clusters on posterior margin of ventral face, anterior face of profemur with setigerous punctures. Posterior face of protibia with golden setose brush limited to distal $1 / 10$. Posterior face of ultimate protarsomere in female without setose furrow, completely glabrous.

## Description:

Head. Antenna with 6 flagellomeres; pedicel broad, nearly rectangular in form. Labrum ventrally with two transverse linear setose rows in basal $1 / 2$, and an additional longitudinal row anteriad to two basal transverse rows, running near entire length of labrum, situated paramedially. Gular suture complete, lateral arms of gular suture meeting anterolateral margin of ventral epicranium posteriad to submentum.
Thorax. Pronotum with transverse impressed line, situated close to anterior margin of pronotum, running parallel with it, nearly meeting anteromedially, weakly effaced medially. Elytral lateral margin without significant swelling at midlength associated with depressed cavity in meso- and metaventrite which receives proleg, elytra evenly deflexed throughout. Protrochanter of male without setose patch, posterior face completely glabrous, ventral face of males and females with linear series of short sharp setae in distal third; profemur ventrally with linear series of small setose clusters on anterior margin only, running only basal $1 / 3$ to $1 / 2$ of profemur, setose patches composed of one to few long setae, ventral surface with excavation apically for reception of protibia, anterior and posterior margins of ventral surface with series of knobs, especially apically, posterior face covered with short stout setae in recessed pits, as well as linear series of setigerous punctures ( $4-6$ ), seta of setigerous punctures long and narrow, ventral margin of anterior face basally with series of short setae in basal $1 / 2$, posterior face with warty bumps situated basally and towards ventral margin, mostly glabrous; posterior face of protibiae with setose brush limited to distal ca. 1/10 length of tibia; posterior face of protarsomere V of female protarsus without setose furrow, posterior face entirely glabrous. Elytra with elytral striae very faintly visible. Metasternal wing in the form of isosceles triangle. Metacoxal wing obliquely transverse, evenly arcuate, metacoxal wing ending at apical $1 / 9$ of metepisternum length. Abdomen. Abdominal sternite VIII weakly emarginate medially. Male genitalia with median lobe of aedeagus not broadly articulating basomedially with the parameres, parameres with narrow basal bridge, longitudinal lists of medial lob very narrow, lateral lists not meeting medial list.

Sexual dimorphism: No significant size or habitus dimorphism evident. Profemoral setation sexually dimorphic, males often with 2 to 3 tufts of setae, whereas females tend to have 4 or more.

## Porrorhynchus (Rhomborhynchus) depressus Régimbart, 1892

Porrhorrhynchus depressus Régimbart, 1892b: 996 [original description]; Porrhorrhynchus depressus: Régimbart 1902: 5 [distribution]; Dineutus (Rhomborhynchus) depressus: Ochs 1929c: 200 [holdings]; Dineutus (Rhomborhynchus) depressus: Ochs 1955: 133 [redescription];
Dineutus (Rhomborhynchus) depressus jamurensis Ochs, 1955: 133 [original description] new synonymy; Dineutus (Rhomborhynchus) depressus versteegi Ochs, 1955: 134 [original description] new synonymy; Dineutus (Rhomborhynchus) depressus moszkowskii Ochs, 1955: 134 [original description] new synonymy; Porrorhynchus ( Rhomborhynchus) depressus: Brinck 1955: 103 [new status]; Porrorhynchus (Rhomborhynchus) depressus depressus: Polhemus 2011: 52 [locality and habitat]

Type material examined: Porrorhynchus depressus Régimbart, 1892: Holotype (đ card mounted, with aedeagus glued to card, missing labrum, Fig. 17B) "N. Guinea/ Dilo/ Loria Vi.VII. 90 [beige label, typed black ink with black border]// Typus [beige label, typed red ink
with red border]// "Porrhorrhynch./ depressus/ Reg. n.sp. [beige label, handwritten in black ink, handwriting appears to be Régimbart's ]// depressus/ Rég. [beige card label, handwritten in black ink, unknown handwriting, black border]// Porrhorhynch./ depressus/ typus! Rég. [yellow label, handwritten in black ink, unknown handwriting]// Museo Civico/ di Genova [beige label, typed black ink]// Dineutus/ (Rhomborhynchus)/ depressus Rég./ vid. Ochs 1954 [beige label, typed black ink]// Vidit P. Brinck,/ 1982-2006 [white label, typed black ink]// HOLOTYPUS/ Porrorhynchus/ depressus/ Regimbart, 1832 [red label, HOLOTYPUS typed in black ink, rest handwritten in black ink, handwriting unknown]//" (MSNG).

Dineutus (Rhomborhynchus) depressus jamurensis Ochs, 1955: Paratype ( $\begin{gathered}\text { pinned, with }\end{gathered}$ aedeagus on point, Fig. 17C) " đ [white label, black typed ink]// B Jamoer/ 4 VIII 1903 [white label, handwritten in ink, handwriting unkown]// Coll./ G.Ochs [white label, typed black ink]// Para-/ typoid [red label with black border, typed black ink; underneath handwritten in black ink SMF C 9535]// jamurensis/ Ochs [beige label with black border, handwritten in blue ink, handwriting appears to be Ochs']//" (SMF); paratype ( $q$ pinned) same as previous except with O label and underneath Para-typoid label handwritten in black ink SMF C 9536, without jamurensis Ochs label (SMF).

Dineutus (Rhomborhynchus) depressus moszkowskii Ochs, 1955: Holotype (*ð pinned, with aedeagus on point, Fig. 17E) " $\widehat{\text { § }}$ [white label, typed black ink]// Holl. N.-Guinea/ Zentralgeb. Ende XII.10/ Moszkowski S.G. [gray label, typed in black ink, except Zentralgeb. Ende XII. 10 handwritten, handwriting uknown]// 94915 [beige label, handwritten in black ink, handwriting unknown]// Coll./ G.Ochs [white label, typed black ink]// Typus [red label with black border, typed black ink; underneath handwritten in black ink SMF C 9538]// moszkowskii/ Ochs [beige label with black border, handwritten in blue ink, handwriting appears to be Ochs']//" (SMF); paratype (Allotype) ( $~$ q pinned) same label data as holotype except with $q$ label and "Allo/typus [red label with black border, Allo- hand written in black ink, rest typed black ink; underneath handwritten in black in SMF C 9539]//" (SMF).

Dineutus (Rhomborhynchus) depressus versteegi Ochs, 1955: Holotype ( $\overbrace{\text { 1 }}$ pinned, with aedeagus on point, Fig. 17F) " $\widehat{\text { ' [white label, type black ink]// Z. NieuwGuinea/ Versteeg }}$ 1912.13/ Beaufort. II [white label, typed black ink, except Beaufort. II handwritten in black ink]// [blue label, underneath typed in black ink 1919, plus some illegible scribble in pencil]// Museum/ Dresden/ leg. 1928. [white label with thin black border, typed black ink]// Coll./ G.Ochs [white black, typed black ink]// Typus [red label with thick black border, typed black ink; underneath handwritten in black ink SMF C 9537]// versteegi Ochs [beig label with black border, handwritten in blue ink, handwriting appears to be Ochs']//" (SMF); paratype ( $q$ pinned) same locality label as previous, except with $q$ label, as well as "Para-/ typoid [red label with black border, typed black ink; underneath handwritten in black ink SMF C 11258]// det. K.M.HELLER 1915/ Porrorhynchus/ depressus Rég. [beige label, typed black ink, except 15 handwritten in black ink]//" and without blue label and versteegi Ochs label (SMF).

Type material notes. Upon initial receipt of the type specimens of P. d. moszkowskii from SMF close examination revealed that what had been indicated as the male holotype based on the type labels (SMF C 9538), as well as having the male symbol label and pointed aedeagus, was in fact a female specimen. The specimen with the Allotype label (SMF C 9539) and female symbol
label, was the actual male specimen and whose abdomen was propped open, indicative of dissection. It appears at some point during a past examination all the male holotype labels, including the dissected genitalia, were removed, as were those of the female allotype's, at which point they were subsequently switched upon being returned to the specimens. Ochs (Ochs 1955) was explicit when he designated a male as the holotype. Therefore, since it is clear that the male specimen received from SMF was previously dissected, and carrying allotype labels, being likely switched with the present female holding the holotype labels, the holotype labels were returned to the dissected male specimen and the allotype label's were placed back on the female specimen received.

## Additional material examined:

INDONESIA: West Papua ("Irian Jaya"): Nabire, 30km S. Nabire, Kali Cemara, 150m, 15.viii.1998, leg. M. Balke, CE 1 (29 ex. NHMW), Nabire, rd. Nabire-Ilaga, Km 35 Kali Cemara, 100m, 27.ix.1997, leg. M. Balke, IRS97\#6 (41 ex. NHMW), Nabire, rd. Nabire-Ilaga, Km 63, 8.x.1997, leg. M. Balke, IRS97\#9 (62 ex. NHMW), Danau Yamur ("B Jamer"), 4.viii.1905, Coll. MAURICE REGIMBART 1908 ( 1 ex. MNHN), same as previous except: coll. C.L. Legros (1 ex. MNHN), "Nelle Guinea Limmena", Danau Yamur ("B Jamer"), 4.viii.1905, Coll. MAURICE REGIMBART 1908 ( 2 ex. MNHN).
PAPUA NEW GUINEA: : 1912-1913, leg. Versteeg, ZML.2010/ 334,335 (2 ex. MZLU); Central Province: "Astrolabe Geb.", leg. E. Weiske, ZML.2010/ 333 (1 ex. MZLU); Gulf Province: Omo River, at Omo, $6^{\circ} 58^{\prime} 41$ "S $144^{\circ} 18^{\prime} 15{ }^{\prime \prime} \mathrm{E}, 40 \mathrm{~m}, 28 . \mathrm{ii} .1995$, leg. D.A. Polhemus, CL 7001 (6 ex. DAPC).

Diagnosis: Labrum elongate and in the form of a near equilateral triangular. Antenna with 6 flagellomeres. Dorsally bronzy-brown, with pronotal yellow lateral margins broad, completely reaching lateral boundary of pronotum. Yellow lateral margins complete on elytra, extending to elytral apices, never interrupted in basal $1 / 3$ by a dark spot, elytra without swelling associated with proleg reception in males. Elytral broadest at midlength, apices spinose, apicolaterally without saw-tooth-like spines, sutural angle produced to a short point, one large parasutural spine, epipleural angle with a large spine.

Description: Size: § L: 9.5-11.5 mm, W: 5.5-7.0 mm; ㅇ L: 9.6-11.6mm, W 5.2-7.1 mm. Habitus. Small member of genus; body form often tear dropped shaped, broadest posteriad to middle, attenuated anteriorly, and posteriorly, other populations elongate oval, evenly attenuated anteriorly to posteriorly; in lateral view fairly depressed, weakly humped in scutellar region, depressed posteriorly and anteriorly; in anterior and posterior view weakly sloped towards lateral margins, lateral margins explanate.
Coloration. Dorsally head, pronotum, elytral bronzy brown in color; labrum of uniform color, similar to head; pronotum/elytra with yellow lateral margins; elytral margin of uniform yellow color, only darkened at epipleural spine; venter yellow; ultimate maxillary palpomere, not darkened in color; prolegs with tibia somewhat darker in color in proximal $1 / 2$, profemora shortly darker apically.
Head. Dorsally vertex with fairly even covering of lightly impressed, fine punctures, often obscured by strongly reticulation, most readily visible in lateral view, separated from nearest puncture by ca. 2 x to 3 x diameter of single puncture; interorbital ridge without yellow margin; frons punctation similar to vertex, more sparse, punctures sparsest apicomedially, fronto-lateral
margins lightly wrinkled, frontoclypeal suture with posterior margin nearly straight, lateral margins shallowly arced, meeting posterior margin at ca $130^{\circ}$ angle; clypeus with punctation most evident at anterior margin, sparsest medially, punctures separated from nearest puncture by ca. 3 x to 4 x diameter of single puncture, becoming more densely spaced anteriorly and laterally; antennal flagellum with 6 complete flagellomeres, ultimate flagellomere ca. $3 x$ longer than penultimate, trapezoidal in shape; labrum in form of equilateral triangle, punctation absent basomedially, in association with strong reticulation, strongly present apically, punctation more strongly impressed, separated from nearest puncture by 1 x to 2 x diameter of single puncture; maxillary and labial palpi similar in shape, strongly hatchet-form. anterior margin of maxillary palp/ventral margin of labial palp evenly curved, posterior margin of maxillary palp/dorsal margin of labial palp more strongly curved proximally, nearly straight apically, apex of both truncate.
Thorax. Pronotum with even covering of punctation, punctation consists of fine, weakly impressed punctures, most evident medially, nearly imperceptible laterally, punctures separated from nearest puncture by 1 x to 2 x diameter of single puncture, reticulation less impressed medially, becoming very well impressed laterally, shallow transverse depression often present medially, lateral marginal depression absent; protibial spine projecting anterolaterally; male protarsi narrow, somewhat dorsally convex, shape as in Fig. 11A, ultimate protarsomere of male ca. 2 x as long as wide, penultimate protarsomere slightly larger than previous three; ultimate protarsomere of female ca. $.5 x$ longer than length of penultimate; elytra with uniform reticulation, elytral discs with even covering of very weakly impressed, fine punctation, nearly imperceptible, distance between nearest punctures ca. $4 \mathrm{x}-5 \mathrm{x}$ diameter of a single puncture, punctures more closely spaced at suture; lateral marginal depression absent, elytral margins evenly sloped, weakly explanate; yellow lateral margin complete, ending apicolaterally at epipleural spine, never interrupted in basal $1 / 3$ by darkly colored spot, males without swelling associated with cavity for proleg reception, apicolateral margins of elytra without triangular saw-tooth-like spines, the epipleural angle produced to spine, elytral apices spinose (Fig. 1D), with 1 parasutural spine, sutural angle produced; mesosternal apex shortly acuminate with broad apex; mesocoxae/metacoxae dissimilar, mesocoxae without posteriorly projecting process; metacoxal process with without distinct lobes (Fig. 11B); male mesotarsal claws as in Fig. 11L with ventral margin strongly arched, anterior claw not significantly narrowed apically.

Genitalia. Aedeagus (Fig. $12 \mathrm{~A}-\mathrm{C}, \mathrm{E}-\mathrm{G}, \mathrm{I}-\mathrm{K}$ ) with median lobe, nearly as long as parameres, just shorter in some populations, parallel sided for $4 / 5$ length, narrowed towards apex in apical $1 / 5$ with straight apicolateral margins, carinate in apical $1 / 6$, apex in lateral view subtruncate, median lobe evenly arced dorsally; parameres in dorsal view narrow, setose in apical $1 / 2$, weakly laterally expanded in apical $1 / 2$, shallowly arced towards apex, apex flatly rounded, basally with narrow basal bridge, in lateral view ventral margin of parameres very weakly curved anteriorly to posteriorly. Female reproductive tract (Fig. 13D) with narrow, elongate, tubiform spermatheca; gonocoxae elgonate and narrow, apically narrowly rounded.

Sexual dimorphism. No sexual dimorphism appears evident, only differences are those mentioned in the description of the subgenus.

Variation. This species is variable in the extent of lateral expansion of the elytra. Specimens from more eastern regions exhibit a broader dorsal habitus (Fig. 12D, H) from the greater lateral
expansion of the elytra, while those from the western part of the range are more narrow, with only slightly laterally expanded elytra (Fig. 12L).

Differential diagnosis: Porrorhynchus depressus can be distinguished from all other species of the genus in having spinose elytral apices without apicolateral triangular saw-tooth-like spines and from P. misoolensis in the form of the aedeagus and gonocoxae of the female RT. The aedeagus of $P$. d. depressus is parallel-sided for $4 / 5$ its length, straightly narrowed towards apex in apical $1 / 5$ and nearly as long as the narrow parameres, which are narrow, whereas that of $P$. misoolensis is shortly laterally expanded in apical $3 / 4$, then roundly narrowed towards apex in apical $1 / 4$, and distinctly shorter than the broader parameres. The gonocoxae of $P$. depressus are narrower, and more elongate in appearance, with their apex narrowly rounded, compared to those of $P$. misoolensis which has broader, less elongate gonocoxae. In general $P$. depressus is larger in body size and has a broader dorsal habitus, with elytral that are typically much more laterally expanded. The lateral expansion of the elytra occurs near mid-length, as opposed to just anteriad to it, as in P. misoolensis.

Distribution: Fig. 15. Found infrequently across New Guinea.
Biology: This species is known primarily from rivers, and one locality is described in detail by Polhemus (2011). Older records also include the lake, Danau Yamur, but it is unclear if $P$. depressus was collected on the lake or from streams near or feeding the lake, the latter being more likely.

Discussion: Several subspecies of $P$. depressus were described of by Ochs (1955), all from very few specimens, from disparate localities across New Guinea. The subspecies jamurensis was described from Danau Yamur in northwestern New Guinea and based upon minor variation in the yellow lateral margins of the elytra and parameres of the aedeagus. The subspecies versteegi was described from the "Beaufort River", described by Ochs (1955) as a tributary of the Lorentz River, distinguished as being slightly larger and broader than the nominal species, also associated with minor aedeagal variation. Finally moszkowskii, described from the Van Daalen River, was separated by Ochs (1955), due to its smaller size and narrower body form. This minor variation in body size, outline, and parameres served as the basis for separating these subspecies from the typical form from southeastern New Guinea.

As can be seen from Fig. 12D,H,L there is considerable variation in degree of lateral expansion to the elytra, creating relatively broader and narrow dorsal habitus among populations. Newly examined specimens from Nabire are considerably narrower than the southeastern populations, more so than any other previously described subspecies, but similar significant variation is not exhibited in other morphological features (Fig. $11 \mathrm{~A}-\mathrm{L}$ ). As seen with other Porrorhynchus species, the parameres tend to vary considerably among populations (i.e. P. marginatus), but the median lobes tend to be fairly conserved. The most noticeably variation among the median lobes is exhibited in the degree of acumination in the apex. Specimens from Nabire (Fig. 12I) were more weakly constricted, relative to typical specimens (Fig. 12A), while those from the Lorentz River (corresponding to versteegi,Fig. 12E) were more strongly acuminate. However, all the median lobes of all populations are similarly proportioned, being nearly as long as the parameres, with similar features being mostly parallel-sided for nearly their
entire length, acuminate in the apical $1 / 5$, and with a carinate apex. For this reason all the former subspecies are here synonymized, being based on simple populational variation.

This species has not yet been implemented as being sensitive to water quality. However, it is apparently infrequently encountered, suggesting it may be relatively rare compared to other New Guinean gyrinid fauna (Polhemus 2011).

We here propose the common name of Flat Snouted Whirligig for $P$. depressus.

## Porrorhychus (Rhomborhynchus) misoolensis (Ochs, 1955), new status

Dineutus (Rhomborhynchus) depressus misoolensis Ochs, 1955: 135 [original description]; Porrorhynchus (Rhomborhynchus) depressus misoolensis: Polhemus 2011: 53 [locality and habitat information]

Type material examined: Paratype ( $\widehat{\text { pinned, }}$ with aedeagus pointed, Fig. 17H) " $\widehat{\text { }}$ [white label, typed black ink]// MISOOL Id. (W.)/ $0-75 \mathrm{~m}$. Fakal./ 8.ix - 20.x.1948./ M.A. Lieftinck [beige label, typed black ink, except Fakal handwritten in black ink]// Coll./ G.Ochs [white label, typed black ink]// Para-/ typoid [red label with thick black border, typed black ink; underneath handwritten in black ink SMF C 9540]// misoolensis/ Ochs [beige label with black border, handwritten in blue ink, handwriting appears to be Ochs’]//" (SMF); paratype ( $q$ pinned) same as previous except with $q$ label and underneath Para-typoid label handwriting reads SMF C 9541, and without misoolensis Ochs label (SMF).

## Additional material examined:

INDONESIA: West Papua ("Irian Jaya"): Misool Island: Tama River, SE of old Fakal village site, $1^{\circ} 51^{\prime} 38.1^{\prime \prime} \mathrm{S} 129^{\circ} 55^{\prime} 24.1^{\prime \prime} \mathrm{E}, 60 \mathrm{~m}, 22 . \mathrm{iv} .1999$, leg. D.A. Polhemus, CL 7110 (4 ex. DAPC).

Diagnosis: Labrum elongate and in the form of a near equilateral triangular. Antenna with 6 flagellomeres. Dorsally bronzy-brown, with pronotal yellow lateral margins broad, completely reaching lateral boundary of pronotum. Yellow lateral margins complete on elytra, extending to elytral apices, never interrupted in basal $1 / 3$ by a dark spot, elytra without swelling associated with proleg reception in males. Elytra broadest just anteriad to midlenth, apices spinose, apicolaterally without saw-tooth-like spines, sutural angle produced to a short point, one large parasutural spine, epipleural angle with a large spine.

Description: Size: đ L: 9.4-10.1 mm, W: $5.6-5.8 \mathrm{~mm}$; ㅇ L: $9.9-10.1 \mathrm{~mm}$, W $5.6-5.8 \mathrm{~mm}$. Habitus. Smallest member of genus; body form elongate oval, evenly attenuated anteriorly and posteriorly; in lateral view depressed, weakly humped in scutellar region, depressed posteriorly and anteriorly; in anterior and posterior view weakly sloped towards lateral margins, lateral margins explanate.
Coloration. Dorsally head, pronotum, elytra bronzy brown in color; labrum of uniform color, similar to head; pronotum, elytra with yellow lateral margins; elytral margin of uniform yellow color, only darkened at epipleural spine; venter yellow; ultimate maxillary palpomere, not darkened in color; prolegs with tibia darker in color in proximal $1 / 2$, profemora shortly darker apically.

Head. Dorsally vertex with even covering of lightly impressed, fine punctures, often obscured by the strong reticulation, most readily visible in lateral view, separated from nearest puncture by ca. 1.5 x to 2 x diameter of single puncture; interorbital ridge without yellow margin; frons punctation similar to vertex, more sparse, punctures sparsest apicomedially, separated from nearest puncture by ca. $2 x$ to $3 x$ diameter of single puncture, fronto-lateral margins very lightly wrinkled, frontoclypeal suture with posterior margin nearly straight, lateral margins shallowly arced, meeting posterior margin at ca $130^{\circ}$ angle; clypeus with punctation most evident at anterior margin, sparsest medially, punctures separated from nearest puncture by ca. 1.5 x to 2 x diameter of single puncture, becoming more densely spaced anteriorly/laterally; antennal flagellum with 6 complete flagellomeres, ultimate flagellomere ca. $3 x$ longer than penultimate, trapezoidal in shape; labrum in form of equilateral triangle, punctation absent basomedially, strongly present apically, punctation more strongly impressed than remainder of head, separated from nearest puncture by 1 x to 1.5 x diameter of single puncture; maxillary and labial palpi similar in shape, both strongly hatchet-form with anterior margin of maxillary palp/ventral margin of labial palp evenly curved, posterior margin of maxillary palp/dorsal margin of labial palp more strongly curved proximally, nearly straight apically, apex of both truncate. Thorax. Pronotum with even covering of punctation, punctation consists of fine, weakly impressed punctures, most evident medially, nearly imperceptible laterally, punctures separated from nearest puncture by 1 x to 1.5 x diameter of single puncture, reticulation less impressed medially, very well impressed laterally, shallow transverse depression often present medially, lateral marginal depression absent; protibial spine projecting anterolaterally; male protarsi narrow, somewhat dorsally convex, shape as in Fig. 11M, ultimate protarsomere of male ca. 2x as long as wide, penultimate protarsomere slightly larger than previous three; ultimate protarsomere of female ca. . $5 x$ length of the penultimate; elytra with uniform reticulation, elytral discs with even covering of weakly impressed, fine punctation, nearly imperceptible, distance between nearest punctures on average ca. 5 x to 6 x diameter of single puncture, punctures more closely spaced in suture; lateral marginal depression absent, elytral margins evenly sloped, explanate; yellow lateral margin complete, ending apicolaterally at epipleural spine, never interrupted in basal $1 / 3$ by darkly colored spot, males without swelling associated with cavity for proleg reception, apicolateral margins of elytra without triangular saw-tooth-like spines, epipleural angle produced to spine, elytral apices spinose (Fig. 1E), with 1 parasutural spine, sutural angle produced; mesosternal apex shortly acuminate with broad apex; mesocoxae/ metacoxae dissimilar, mesocoxae without posteriorly projecting process (Fig._); metacoxal process without distinct lobes (Fig. 11N); male mesotarsal claws as in Fig. 11P with ventral margin strongly arched, anterior claw not significantly narrowed apically.

Genitalia. Aedeagus (Fig. 12M - O) with median lobe, shorter than parameres, running ca. 5/6 their length, parallel sided in basal 3/4, then laterally expanded, apically arcuately narrowed towards apex in apical $1 / 4$, apex carinate, narrowly rounded, in lateral view apex subtruncate, median lobe weakly arced dorsally; parameres broad, setose in apical $1 / 2$, in dorsal view, laterally expanded in apical $1 / 2$, apically narrowly rounded; in lateral view ventral margin of parameres very weakly curved anteriorly to posteriorly. Female reproductive tract (Fig. 13E) with narrow, elongate, tubiform spermatheca; gonocoxae broad, lateral margin straightly angled towards apex, apex truncate.

Sexual dimorphism. No sexual dimorphism appears evident, only differences are those mentioned in the description of the subgenus.

Variation. Very little variation observed, but only a small number of individuals were available for study.

Differential diagnosis: Porrorhynchus misoolensis can be distinguished from all other species of the genus in having spinose elytral apices without apicolateral triangular saw-tooth-like spines and from $P$. depressus in the form of the aedeagus and female RT. See differential diagnosis under $P$. depressus for further details. Distribution can also be used to separate $P$. misoolensis as it is only known from Misool Island.

Distribution: Fig. 15. Known only from Misool Island, West Papua.
Biology: This species has only been collected once after its original description from a upland forested stream (Polhemus 2011). The habitat is described in detail by Polhemus (2011) who suggested this species may be influenced by stream water chemistry and substrate characteristics.

Discussion: Porrorhynchus misoolensis was only collected from a single locality during sampling of many streams on Misool island by Polhemus in 1999 (Polhemus 2011). This coupled with the species, limited distribution, and potential sensitivity to water chemistry suggests $P$. misoolensis may be of conservation concern. Further investigations into the species distribution on the island and its sensitivity to water quality and chemistry would be beneficial.

We here propose the common name of the Misool Snouted Whirligig for $P$. misoolensis.

## Conclusions

Few features unite all the species currently recognized within Porrorhynchus, they are: 1) an elongate labrum that is at least $1 / 2$ as long as wide; 2 ) a spinose distolateral angle of the protibia; 3) an elongate tubiform spermatheca associated with a vaginal shield. These, however, could simply be symplesiomorphies as members of Rhombodineutus also have elongate labra and elongate spermathecae, and members of Andogyrus have an expanded distolateral angle to their protibia forming a blunt spine (Brinck 1983; Hatch 1925). Members of Porrorhynchus sensu stricto on the other hand present at least three potential synapomorphies: 1) the presence of a protrochanteric setose patch in males; 2) loss of the pronotal transverse impressed line; 3) setation of the ventral face of the profemora consisting of two longitudinal lines of clusters of large setae, progressively becoming denser apically. These features separate members of Porrorhynchus sensu stricto from all other gyrinid genera and may, in fact, represent excellent apomorphies for the true genus Porrorhynchus. Brinck's Ceylorhynchus was erected for $P$. indicans and diagnosed by small setae present at the apicolateral angle of the protibia, but these setae are present on many gyrinid species, and the separation of $P$. indicans from the members of the sensu stricto group seems unjustified given the aforementioned shared features.

Rhomborhynchus on the other hand shares numerous features with the genus Dineutus, these are: 1) profemora with setigerous punctures, 2) setation of the ventral face of profemora composed of small clusters of setae, 3) narrow protibia that are not strongly laterally expanded,
4) median lobe of aedeagus narrowly articulating with parameres. Given these shared features, it is understandable why Ochs $(1926 ; 1955)$ considered Rhomborhynchus a subgenus of Dineutus. Whether Rhomborhynchus is sister to the rest of Dineutus retaining features from a common ancestor with Porrorhynchus, sister to Porrorhynchus sensu stricto retaining features from a common ancestor with Dineutus, or forming a grade into Dineutus from Porrorhynchus will need to be clarified via phylogenetic analysis.

## Checklist of species

The Sri Lankan Snouted Whirligig
Porrorhynchus (Porrorhynchus) indicans (Walker, 1858)
= Porrorhynchus brevirostris Régimbart, 1877
The Splendid Snouted Whirligig
Porrorhynchus (Porrorhynchus) landaisi Régimbart, 1892
= Porrorhynchus barthelemyi Régimbart, 1907
$=$ Dineutus (Porrorhynchus) landaisi latilimbus Ochs, 1926
The Margined Snouted Whirligig
Porrorhynchus (Porrorhynchus) marginatus Laporte, 1835
= Porrorhynchus tenuirostris Régimbart, 1877
= Dineutus (Porrorhynchus) mjoebergi Ochs, 1926
Borneo, China, Java, Laos, the Malay Peninsula, Myanmar, Sumatra, Thailand, Vietnam

The Flat Snouted Whirligig
Porrorhynchus (Rhomborhynchus) depressus Régimbart, 1907
New Guinea
= Dineutus (Rhomborhynchus) depressus jamurensis Ochs, 1955
= Dineutus (Rhomborhynchus) depressus moszkowskii Ochs, 1955
= Dineutus (Rhomborhynchus) depressus versteegi Ochs, 1955
The Misool Snouted Whirligig
Porrorhynchus (Rhomborhynchus) misoolensis Ochs, 1955 new status Misool Island

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Fig. 1. Dorsal habitus of Porrorhynchus species. A) P. (P.) marginatus, B) $P$. (P.) landaisi, C) $P$. (P.) indicans, D) $P$. (R.) depressus, E) $P$. (R.) misoolensis. Scale bar $=5 \mathrm{~mm}$.

Fig. 2. Right antenna of species of Porrorhynchus. For each, above anterior view, below posterior view, scale bars 0.5 mm . A) $P$. (P.) landaisi, B) $P$.(P.) indicans, C) $P$. (P.) marginatus, D) $P$. (R.) depressus.

Fig. 3. Scanning electron microscope image of male $P$. (P.) marginatus. A) Protrochanter, marked area shows protrochanteric setose patch, scale bar $=400 \mu \mathrm{~m}, \mathrm{~B})$ Protrochanteric setose patch, scale bar $=50 \mu \mathrm{~m}$.

Fig. 4. Scanning electron microscope image of male $P$. ( $P$.) marginatus profemur. Scale bar $=$ 1 mm .

Fig. 5. Aedeagus of $P$. (P.) indicans. A) Dorsal view, B) Lateral view, C) Ventral view. Scale bar $=1 \mathrm{~mm}$.

Fig. 6. Morphology of $P$. (P.) landaisi $[\mathrm{A}-\mathrm{K}]$ and $P$. (P.) indicans $[\mathrm{L}-\mathrm{O}]$. A - D specimen from Hainan island. A) Protarsus, scale bar $=1 \mathrm{~mm}, \mathrm{~B}$ ) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{C}$ ) Maxillary palps above, labial palps below, scale bar $=0.5 \mathrm{~mm}, \mathrm{D})$ Mesotarsal claws, scale bar $=$ 0.5 mm . E - H specimen from Vietnam. E) Protarsus, scale bar = 1 mm, F) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{G}$ ) Maxillary palps above, labial palps below, scale bar $=0.5 \mathrm{~mm}, \mathrm{H}$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm}$. I -K specimen from Tibet, China. I) Protarsus, scale bar $=1 \mathrm{~mm}, \mathrm{~J}$ ) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{~K}$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm} . \mathrm{L}-\mathrm{O}$ $P$. (P.) indicans. L) Protarsus, scale bar $=1 \mathrm{~mm}, \mathrm{M}$ ) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{~N}$ ) Maxillary palps above, labial palps below, scale bar $=0.5 \mathrm{~mm}, \mathrm{O})$ Mesotarsal claws, scale bar $=$ 0.5 mm .

Fig. 7. Aedeagus [A - C, E - G, I - K] and dorsal habitus [D, H, L] of $P$. (P.) landaisi. A - D specimen from Hainan Island. A) Dorsal view, scale bar $=1 \mathrm{~mm}, \mathrm{~B})$ Ventral view, C) Lateral view, D) Dorsal habitus, scale bar $=5 \mathrm{~mm} . \mathrm{E}-\mathrm{H}$ specimen from Vietnam. E) Dorsal view, scale bar $=1 \mathrm{~m}, \mathrm{~F}$ ) Ventral view, G) Lateral view, H) Dorsal habitus, scale bar $=5 \mathrm{~mm} . \mathrm{I}-\mathrm{L}$ specimen from Tibet, China. I) Dorsal view, J) Ventral view, K) Lateral view, L) Dorsal habitus, scale bar $=5 \mathrm{~mm}$.

Fig. 8. Morphology of $P$. (P.) marginatus. A - E specimen from Chumphon, Thailand. A) Protarsus, scale bar $=1 \mathrm{~mm}, \mathrm{~B}$ ) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}$, C) Mesocoxal apex, scale bar 0.5 mm , D) Maxillary palp above, labial palp below, scale bar $=0.5 \mathrm{~mm}$, E) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm} . \mathrm{F}-\mathrm{J}$ specimen from Khammouane, Laos. F) Protarsus, scale bar $=1$ mm , G) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{H}$ ) Mesocoxal apex, scale bar 0.5 mm , I) Maxillary palp above, labial palp below, scale bar $=0.5 \mathrm{~mm}, \mathrm{~J}$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm} . \mathrm{K}-$

O specimen from Dac Lac, Vietnam. K) Protarsus, scale bar = 1 mm , L) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{M})$ Mesocoxal apex, scale bar $0.5 \mathrm{~mm}, \mathrm{~N}$ ) Maxillary palp above, labial palp below, scale bar $=0.5 \mathrm{~mm}, \mathrm{O}$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm} . \mathrm{P}-\mathrm{T}$ specimen from East Kalimantan, Malaysia. P) Protarsus, scale bar $=1 \mathrm{~mm}$, Q) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}$, R) Mesocoxal apex, scale bar 0.5 mm , S) Maxillary palp above, labial palp below, scale bar = $0.5 \mathrm{~mm}, \mathrm{~T}$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm} . \mathrm{U}-\mathrm{X}$ specimen from Manna, Sumatra. U) Protarsus, scale bar $=1 \mathrm{~mm}$, V) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{~W}$ ) Mesocoxal apex, scale bar $0.5 \mathrm{~mm}, \mathrm{X}$ ) Maxillary palp above, labial palp below, scale bar $=0.5 \mathrm{~mm} . \mathrm{Y}-\mathrm{C} 2$ specimen from Java. Y) Protarsus, scale bar $=1 \mathrm{~mm}, \mathrm{Z}$ ) Metacoxal apex, scale bar $=0.5 \mathrm{~mm}, \mathrm{~A} 2$ ) Mesocoxal apex, scale bar 0.5 mm , B2) Maxillary palp above, labial palp below, scale bar $=0.5$ $\mathrm{mm}, \mathrm{C} 2$ ) Mesotarsal claws, scale bar $=0.5 \mathrm{~mm}$.

Fig. 9. Dorsal habitus of $P$. (P.) marginatus, populational variation. Scale bar $=5 \mathrm{~mm}$. A) Mandalay Dist., Myanmar, B) Loei Prov., Thailand, C) Khammouane Prov., Laos, D) Đắk Lắk Prov., Vietnam, E) Saraburi Prov., Thailand, F) Chumphon Prov., Thailand, G) Songkhla Prov., Thailand, H) Pahang, Malaysia, I) Gunung Leuser, Sumatra, J) West Sumatra, Sumatra, K) Siberut Island, Sumatra, L) Manna, Sumatra, M) Java, N) Kapit Distr., Sarawak, Malaysia, O) East Kalimantan, Indonesia.

Fig. 10. Aedeagus of $P$. (P.) marginatus, lateral and dorsal view, populational variation. Scale bar $=1 \mathrm{~mm}$. A) Mandalay Dist., Myanmar, B) Loei Prov., Thailand, C) Khammouane Prov., Laos, D) Đắk Lắk Prov., Vietnam, E) Saraburi Prov., Thailand, F) Chumphon Prov., Thailand, G) Songkhla Prov., Thailand, H) Pahang, Malaysia, I) Gunung Leuser, Sumatra, J) Siberut Island, Sumatra, K) Manna, Sumatra, L) Java, M) Kapit Distr., Sarawak, Malaysia, N) East Kalimantan, Indonesia.

Fig. 11. Morphology of $P$. (R.) depressus [A - L] and $P$. (R.) misoolensis $[\mathrm{M}-\mathrm{P}]$, all scale $=0.5$ mm . A - D specimen from Gulf Prov. Papua New Guinea. A) Protarsus, scale B) Metacoxal apex, C) Maxillary palp above, labial palp below, D) Mesotarsal claw. E - H specimen from the Lorentz River, southwestern Irian Jaya. E) Protarsus, F) Metacoxal apex, G) Maxillary palp above, labial palp below, H) Mesotarsal claw. I - L specimen from Nabire, western Irian Jaya. I) Protarsus, J) Metacoxal apex, K) Mesotarsal claws. L- O P. (R.) misoolensis. L) Protarsus, M) Metacoxal apex, N) Maxillary palp above, labial palp below, O) Mesotarsal claws.

Fig. 12. Aedeagus [ $\mathrm{A}-\mathrm{C}, \mathrm{E}-\mathrm{G}, \mathrm{I}-\mathrm{K}, \mathrm{M}-\mathrm{O}$ ] and dorsal habitus $[\mathrm{D}, \mathrm{H}, \mathrm{L}, \mathrm{P}]$ of $P$. (R.) depressus $[\mathrm{A}-\mathrm{L}]$ and $P$. (R.) misoolensis $[\mathrm{M}-\mathrm{P}]$. A - D specimen from Omo River, Gulf Prov., Papua New Guinea. A) Dorsal view, scale bar $=1 \mathrm{~mm}, \mathrm{~B})$ Ventral view, C) Lateral view, D) Dorsal habitus, scale bar $=5 \mathrm{~mm}$. $\mathrm{E}-\mathrm{H}$ specimen from Lorentz River, southwestern Irian Jaya. E) Dorsal view, scale bar $=1 \mathrm{~m}, ~ F$ ) Ventral view, G) Lateral view, H) Dorsal habitus, scale bar = 5 mm . I - L specimen from Nabire, western Irian Jaya. I) Dorsal view, scale bar = 1mm, J) Ventral view, K ) Lateral view, L ) Dorsal habitus, scale bar $=5 \mathrm{~mm} . \mathrm{M}-\mathrm{P} P .(R$.$) misoolensis.$ M) Dorsal view, scale bar $=1 \mathrm{~mm}, \mathrm{~N}$ ) Ventral view, O) Lateral view, P) Dorsal habitus, scale bar $=5 \mathrm{~mm}$.

Fig. 13. Female reproductive tracts of Porrorhynchus species, scale bars $=1 \mathrm{~mm}$. More heavily sclerotized areas indicated in gray. A) $P$. (P.) landaisi, B) $P$. (P.) indicans, C) P. (P.) marginatus, D) $P$. (R.) depressus, D) $P$. (R.) misoolensis.

Fig. 14. Map showing distribution of Porrorhynchus sensu stricto species. Triangles $=P$. $(P$. landaisi, Circles $=P .(P$.$) marginatus, Square =P .(P$.$) indicans$.

Fig. 15. Map showing distribution of species in the subgenus Rhomborhynchus. Triangles $=P$. (R.) depressus, Circle $=P$. (R.) misoolensis.

Fig. 16. Type specimens. A) Porrorhynchus brevirostris Régimbart, 1877, lectotype, B) Dineutes indicans Walker, 1858, holotype C) Porrorhynchus landaisi Régimbart, 1892, lectotype, D) Dineutus (Porrorhynchus) landaisi latilimbus Ochs, 1926, lectotype E) Porrorhynchus barthelemyi Régimbart, 1907, lectotype.

Fig. 17. Type specimens. A) Porrorhynchus marginatus Laporte, 1835, holotype B) Porrorhynchus depressus Régimbart, 1892, holotype C) Dineutus (Rhomborhynchus) depressus jamurensis Ochs, 1955, paratype, D) Porrorhynchus tenuirostris Régimbart, 1877, lectotype E) Dineutus (Rhomborhynchus) depressus moszkowskii Ochs, 1955, holotype, F) Dineutus (Rhomborhynchus) depressus versteegi Ochs, 1955, holotype, G) Dineutus (Porrorhynchus) marginatus mjoebergi Ochs, 1926, lectotype, H) Dineutus (Rhomborhynchus) depressus misoolensis Ochs, 1955, paratype.



A


B


D









$\geqslant 0$


G $\quad \mathrm{H}$



1


H


M


P





 COLL MAURICE REGIMBAET 1908

## LECTOTYPE


G.Ochs

Paratypoid



C


## E



$1 \times 1$


# Systematics and evolution of the whirligig beetle tribe Dineutini (Coleoptera, Gyrininae, Gyrinidae) 

Grey T. Gustafson ${ }^{1}$ and Kelly B. Miller ${ }^{2}$<br>Department of Biology and Museum of Southwestern Biology, University of New Mexico, Albuquerque, NM 87131, USA.<br>Email: gtgustafson@gmail.com ${ }^{1}$ kbmiller@unm.edu ${ }^{2}$


#### Abstract

The phylogeny and evolutionary history of the whirligig beetle tribe Dineutini are inferred from analysis of 56 morphological characters and DNA sequence data from the mitochondrial genes COI, COII, and 12S, and the nuclear genes H 3 and arginine kinase. Bayesian and maximum likelihood analyses were performed. A Bayesian total-evidence approach was taken to provide a time calibrated phylogenetic tree incorporating fossil taxa. Seventy-one species of extant Gyrinidae were included in the analysis, as well as two fossil taxa, representing all dineutine genera and all proposed, non-monotypic subgenera. The resulting trees strongly support the monophyly of the Dineutini and the genera Dineutus, Macrogyrus, Porrorhynchus, and Enhydrus. The results do not support the distinction of Andogyrus as a separate genus, nor do they support the majority of proposed subgenera. A new classification is presented here requiring the following taxonomic changes: Andogyrus stat. n. is relegated to a subgenus of Macrogyrus; the following subgenera are synonymized with Macrogyrus sensu stricto sensu n.: Australogyrus Ochs, 1949 syn. n., Ballogyrus Ochs, 1949 syn. n., Clarkogyrus Ochs, 1949 syn. n., Megalogyrus Ochs, 1949 syn. n., Orectomimus Ochs, 1930 syn. n., Tribologyrus Ochs, 1949 syn n.; the subgenus Stephanogyrus Ochs, 1955 syn. n. is synonymized with the subgenus Cyclomimus; the genus Dineutus now includes two subgenera Cyclous sensu n. and the Dineutus sensu stricto subgenus sensu n.; the following subgenera are synonymized with the subgenus Cyclous: Callistodineutus Ochs, 1926 syn. n., Cyclinus Kirby, 1837 syn. n., Paracyclous Ochs, 1926 syn. n., Protodineutus Ochs, 1926 syn. n., Spinosodineutes Hatch, 1925 syn. n.; the following subgenera are synonymized with the Dineutus sensu stricto subgenus: Rhombodineutus Ochs, 1926 syn. n., Merodineutus Ochs, 1955 syn n. The subgenus Rhomborhynchus incert. s. is tentatively moved to the genus Dineutus, without phylogenetic placement. The analysis confirms Mesodineutes $\dagger$ is a member of the Dineutini. Each genus and subgenus reviewed in detail with (i) a morphological diagnosis, (ii) its taxonomic circumscription, including the placement of species not included in the analysis, (iii) known distribution, and (iv) relevant discussion. A new identification key to the extant genera and subgenera of the Dineutini is provided. Finally a biogeographic analysis reconstructing ancestral ranges was conducted revealing the historical biogeography of the tribe.


## Introduction

The tribe Dineutini contains the most conspicuous members of the whirligig beetles, (Coleoptera, Gyrinidae), being large in size (commonly $\geq 10 \mathrm{~mm}$ in length) (Brinck, 1984; Gustafson \& Miller, 2015) and with a near global distribution (Miller \& Bergsten, 2012). Most species are lotic (Balfour-Browne \& Brinck, 1961; Brinck, 1977, 1983, 1984; Gustafson \& Miller, 2015), but a few are primarily lentic, or found in a variety of freshwater habitats (Brinck, 1955a; Gustafson \& Miller, 2015). Despite their large size and conspicuous nature, new species
are still being discovered, even in well explored regions such as the U.S.A. (Gustafson \& Sites, 2016), and the vast majority of species lack formal descriptions of immature stages and life history. Furthermore, the tribe itself has never specifically been the focus of a phylogenetic analysis.

Régimbart (1882) was the first to formally diagnosis and describe the tribe Dineutini (see the Classification section for more details) who recognized within it four genera, Macrogyrus Régimbart, 1882, Porrorhynchus Laporte, 1835, Enhydrus Laporte, 1835, and Dineutus Macleay, 1825. The genus Dineutus was first to be split into subgenera by Hatch (1925), then extensively split into many subgenera, along with the genus Macrogyrus, by the work of Ochs (1926, 1949, 1955). Ochs (1924) would also erect a new genus within the tribe, Andogyrus Ochs, 1924. The problematic nature of these subgenera has long been recognized (Brinck, 1955b) as has the distinction of Andogyrus from Macrogyrus (Brinck, 1977). The monophyly of the tribe has also been called into question (Beutel, 1990). The first phylogenetic analysis of the family Gyrinidae using molecular and morphological data, provided support for the monophyly of the tribe (Miller \& Bergsten, 2012), but sampling was not extensive enough to strongly test the monophyly of the genera Enhydrus, Andogyrus and Porrorhynchus, nor the numerous subgenera erected within Dineutus and Macrogyrus.

The interesting distribution of the tribe has resulted in hypotheses about the biogeography and origins of the group. Of particular interest are the genera Macrogyrus, Andogyrus and Dineutus. Andogyrus is distributed widely in South America along the Andes (Brinck, 1977) and appears closely related to Macrogyrus found in Australia, New Guinea, and Wallacea (Ochs, 1949). Gondwanan vicariance origins have been invoked to explain this distribution (Hatch, 1925; Ochs, 1949). Furthermore, classic gyrinid taxonomists have debated whether Macrogyrus is descended from a South American (Hatch, 1925) or Australian common ancestor (Ochs, 1949). Dineutus shows a very peculiar distribution, found in Southeast Asia, the Austral regions, throughout Africa, the North American continent, and eastern Palearctic in Korea (Lee \& Ahn, 2015) and the Ryukyu Islands (Satô, 1962), but is absent from South America (Gustafson \& Miller, 2015). There are two possible explanations for this distribution: 1) local extinction within the continent, 2) Dineutus has yet to disperse to South America.

The purpose of this study is to provide the first phylogenetic analysis of the tribe Dineutini to (i) assess the monophyly of the currently proposed genera and numerous subgenera, to improve and stabilize classification; (ii) construct a time calibrated phylogenetic tree to understand the relationships of dineutine species and the timing of their evolution; and (iii) reconstruct the historical biogeography of the group to test the proposed Gondwanan relationship of Macrogyrus and Andogyrus, and provide an explanation to the absence of Dineutus in South America.

## Materials and Methods

## Data

## Taxon sampling

Seventy-three species of Gyrinidae were included in the main time calibrated analysis (Table S1). Ten outgroup species were selected: Heterogyrus milloti for Heterogyrinae, four species from the tribe Gyrinini, four from Orectochilini, and Gyretes giganteus $\dagger$ for a fossil
outgroup member. Within the Dineutini an attempt was made to include at least two members from all currently recognized subgenera. This was mostly attained with the exception of the following monotypic subgenera not sampled for the analysis: Dineutus (Paracyclous) ritsemai (only known from the type series from Sulawesi); Macrogyrus (Ballogyrus) leopoldi (only known from the holotype specimen from New Guinea); Macrogyrus (Stephanogyrus) caledonicus (known from New Caledonia). The subgenus Rhomborhynchus (two species of contentious placement within Porrorhynchus) was represented by a single specimen only coded for morphological data, no molecular grade specimens were available for analysis. The fossil Mesodineutes amurensis $\dagger$ was utilized as the fossil ingroup member. Non-time calibrated analyses utilized a subset of taxa, excluding those available only for morphology and Heterogyrus milloti (Fig. S1, S2).

Ingroup taxa sampled were identifiable from the genera Dineutus, Andogyrus, Enhydrus, and Porrorhynchus to species and subspecies were applicable. The genus Macrogyrus has never received a comprehensive revision. The species from Australia are readily identifiable, thanks to the work of Watts \& Hamon (2010); however, the species from New Guinea and the Lesser Sunda Islands are a major issue for identification. Numerous subspecies have been described by Ochs (1955) based on only a few specimens, with characters primarily relating to general bodyform, providing no illustrations, and poorly constructed identification keys. Therefore many species sampled from New Guinea and the surrounding area cannot be identified reliably beyond the subgeneric level.

## Morphology

Female genitalia were prepared following the methods outlined in Miller \& Bergsten (2012). The genitalia were illustrated in water using a Camera Lucida attached to a Zeiss Discovery.V8 SteREO microscope. Illustrations were then scanned and traced using Adobe Illustrator CS5. Other morphology illustrated were drawn under the camera lucida, scanned, and traced using the same methods. Scanning electron microscope images were taken at the KU Microscopy and Analytical Imaging Laboratory, University of Kansas, Lawrence, KS, U.S.A.

Dorsal and ventral habitus were taken using a Visionary Digital BK+ light imaging system as well as a Passport imaging system (www.visionarydigital.com, R. Larimer). Habitus images were then edited using Adobe Photoshop CS5 to add scale bars and improve clarity and color.

Sperm has been found to be phylogenetically informative (Jamieson, 1987) and the sperm of Dineutus species was found to exhibit a very unique conjugation form (Breland \& Simmons, 1970). For these reasons sperm was examined from several dineutine species and the conjugation type exhibited included as a morphological character in the analysis. Sperm specimens were harvested from the seminal vesicles of specimens in the field. A portion of seminal vesicle was removed from the specimen while in Phosphate Buffer Solution (PBS) then moved to a slide with an additional drop of PBS. The seminal vesicle was then agitated to free sperm. The slide was then allowed to dry and the original specimen was given a unique identifier (SVSK \#) and kept as a voucher deposited in the Museum of Southwestern Biology, Division of Arthropods (MSBA), at the University of New Mexico. The slide was then DAPI stained and mounted with a slide cover. Sperm slides were visualized using a Zeiss AXIO Imager A2 compound microscope with attached Axiocam 506 mono camera.

Fifty-six morphological characters were coded from all body regions, the female RT, male genitalia, and sperm morphology. Full description of morphological characters (Appendix) and coding of morphological characters from (Table S2) is availalbe. Morphology was coded in MacClade 4.08 (Maddison \& Maddison, 2005). Terminology for dineutine external morphology follows Gustafson \& Miller (2015) and Miller \& Bergsten (2012) for female RT, unless otherwise cited.

DNA
DNA was extracted using a Qiagen DNEasy kit (Valencia, California, USA) and the protocol for animal tissue. Thoracic muscle tissue was extracted from a lateral incision via fine forceps. The remaining specimen was retained and given a unique voucher identifier attached to the specimen via a label. Original DNA extractions are deposited at MSBA, as are the voucher specimens, unless indicated otherwise (Table S1).

Portions of five genes were used in the phylogenetic analysis, and a sixth only for some Dineutus specimens used previously in the analysis by Miller \& Bergsten (2012). The six genes are: cytochrome c oxidase subunit I (COI, 1317 bp aligned), cytochrome c oxidase subunit II (COII, 740 bp aligned), 12 S rRNA (12S, 359 bp aligned), histone III (H3, 328 bp aligned), arginine kinase (AK, 712 bp aligned), and elongation factor 1 alpha (EF $1 \alpha, 348 \mathrm{bp}$ aligned). Standard PCR protocols were used for amplification and sequencing following Miller \& Bergsten (2012). Primers and their sources, used for amplification and sequencing are available in Table S3. Gene coverage for each taxon analyzed is available in Table S1. Sequences were edited using Sequencher 4.8 (Gene Codes, 1999). Sequences were aligned using MUSCLE (Edgar, 2004) via EMBL-EBI's website (EMBL-EBI, 2015). Concatenation and clean up implemented in Mesquite 3.01 (Maddison \& Maddison, 2015).

## Partitioning

The dataset broadly overlaps that used by Miller \& Bergsten (2012), and for this reason we used the same partitioning scheme, with codon-position specific nuclear and mitochondrial partitions. This partitioning scheme was previously tested and found to be preferred by a Bayes Factor test over gene-specific partitions (Miller et al., 2009).

## Analyses

## Phylogenetic, Bayesian

Bayesian analysis was implemented using MPI version of MrBayes 3.2.6 (Ronquist et al., 2011; Zhang et al., 2015). No substitution model was selected a priori, instead the reversiblejump MCMC method with gamma rate variation across sites was used to test the probability of different models a posteriori during analysis (Huelsenbeck et al., 2004; Ronquist et al., 2011; Miller \& Bergsten, 2012). A total-evidence approached was taken for time-calibration (Ronquist et al., 2012). This technique is desirable as it incorporates inherent uncertainty regarding placement of fossils during phylogenetic reconstruction, treating fossil taxa as terminals, as opposed to node-based calibration in which fossils are forced to stem positions (Ronquist et al., 2012).

To infer the substitution rate the methods outlined by Ronquist et al. (2012) were followed with the mean age of the fossil Angarogyrus minimus ( 178 Ma ) used to calculate median rate, and the mean age of Mesogyrus antiquus ( 161 Ma ) for the standard deviation. The Fossilization-Birth-Death macroevolutionary model (Heath et al., 2014) was employed using the methods outlined by Zhang et al. (2015). The sampling strategy was set to diversity, with a sample probability of 0.06 as there are 153 known species of Dineutini, the ingroup for the analysis. Fossils were given a uniform age prior based on the age of the fossil. The tree age was given an offset exponential prior based on the age of Mesogyrus antiquus, a likely heterogyrine fossil, as Heterogyrus milloti was used as the furthest outgroup member. A relaxed clock model was used, with the branch length clock prior set to fossilization to use the FBD model, and the clock rate variance prior set to igr. The analysis was run for 10 million generations, using four chains (three heated, one cold), with swap number set to two, and a temperature of 0.1 for the heated chains. MCMC convergence was monitored using Tracer v.1.6 (Rambaut et al., 2013).

Additional analyses using only mitochondrial and nuclear gene data were performed to check their data sets influence on the final total evidence topology (Figs. S1,S2). Certain problematic species were also removed from analyses to test effects on phylogenetic reconstruction (Fig. S4, S5).

## Phylogenetic, Maximum Likelihood

A maximum likelihood analysis was also performed (Figure S6) as a form of reciprocal illumination (Hennig, 1966). The maximum likelihood analysis was implemented using the Hybrid MPI RAxML ver. 8 (Stamatakis, 2014). Model choice for the different genes was tested a priori using jModelTest (Posada, 2008). The GTR+G model was implemented as it was selected as the primary or secondary model for the majority of codon positions for the majority of genes. Each gene was analyzed individually. The genes were then combined to construct a multi-locus species tree using ASTRAL-II (Mirarab et al., 2014; Mirarab \& Warnow, 2015). One hundred replicates of multi-locus bootstrap support (Seo, 2008) were then performed in ASTRAL-II. Morphology was not included in the maximum likelihood analysis. All phylogenetic analyses were run on the super computer cluster Ulam at the Center for Advanced Research Computing, University of New Mexico.

## Biogeographic, Ancestral Range Estimation

The time-calibrated consensus tree from the Bayesian total evidence analysis was used for the biogeographic analysis, with outgroup and fossil taxa pruned, as well as AyTs832 (Macrogyrus albertisi) to remove a polytomy. The analysis was performed using the program R and the package BioGeoBEARS (Matzke, 2013b, a) to estimate the ancestral range of the Dineutini across their entire distribution. The program offers several models and statistical comparison of model fit. Analyses were run under the DEC (Ree et al., 2005; Ree \& Smith, 2008) and DIVALIKE (Ronquist, 1997) models both with and without the +j found-event speciation parameter (Matzke, 2014). Following completion of analyses model fit was compared statistically within BioGeoBEARS.

For the biogeographic regions in the analysis the following abbreviations were used: A, Australia; M, Melanesia, New Guinea and surrounding islands; W, Wallacea; O, Oriental; P, Palearctic; E, Ethiopean region; N, Nearctic; C, Central America; I, West Indies (Fig. 3). Coding
of taxa for biogeography is available in Table S1. The maximum distribution allowed for a species was 5 .

Four time strata (TS) were established for the time stratification, these were: TS1, 120 90 Ma ; TS2, $90-50 \mathrm{Ma}$; TS3, $50-20 \mathrm{Ma}$; TS4, $20-$ Present. TS1 represents the early stages of the final Gondwanan break-up with the rifting of South American and Africa, and the initial break-up of East Gondwana (Storey, 1995). This point also represents the origins of the Dineutini. For this time slice the following areas were made unavailable based on paleogeographic data: Central America (Iturralde-Vinent, 2006), the West Indies (IturraldeVinent, 2006), Melanesia (Toussaint et al., 2014), and Wallacea (Hall, 2001, 2002, 2013). TS2 represents the final stages of the Gondwanan break-up with drifting of South America, Antarctica, Australia (Storey, 1995), and their subsequent final separation (Livermore et al., 2005; Lawver et al., 2011; Reguero et al., 2014). During TS2 the same areas were unavailable, except Central America was made available (Iturralde-Vinent, 2006). TS3 represents the isolation of South America, Antarctica, and Australia (Lawver \& Gahagan, 2003; Lawver et al., 2011); and the first potential emergence of the Caribbean (Iturralde-Vinent, 2006). New Guinea likely had little available land before 25 Ma (Toussaint et al., 2014), but an orogenic event around 35 Ma (van Ufford \& Cloos, 2005) likely created a small island, which persisted to form the oldest regions of the New Guinea (Baldwin et al., 2012). At this point the West Indies, as well as the Melanesia area, are available, but the latter with low dispersal rate multipliers. TS4 represents the appearance of Wallacea (Hall, 2013) and major formation of the terrestrial New Guinean area (Toussaint et al., 2014) with biotic interchange between the regions. At this point Wallacea areas is allowed. The dispersal rate coding followed that of Toussaint et al. (2016), based on the above paleogeographic evidence reference for each time slice.

## Results

## Phylogenetic

The Bayesian analysis (Figs. 1, 2, S1, S2) strongly supports a monophyletic Dineutini (posterior probability, $\mathrm{pp}=0.99$ ) with a Late Cretaceous origin ( $95 \%$ highest probability density, $\mathrm{hpd}=75.75-113.93 \mathrm{Ma}$, median of $\mathrm{hpd}, \mathrm{m}=94.24 \mathrm{Ma}$ ). Within the Dineutini there are two clades, one comprising Dineutus, Porrorhynchus and the extinct genus Mesodineutes $\dagger$; the other with Macrogyrus and Enhydrus. As Mesodineutes $\dagger$ only had few characters available it introduced uncertainty into the analysis, resulting in lower pp for the clades. Removing Mesodineutes $\dagger$ resulted in significantly higher support (Fig. S4) for the two clades (pp = 1.00 for the Porrorhynchus + Dineutus clade and pp $=0.86$ for Enhydrus + Macrogyrus $)$. Both are similar in age with Late Cretaceous origins ( $\mathrm{hpd}=67.38-101.44 \mathrm{Ma}, \mathrm{m}=83 \mathrm{Ma}$ and $\mathrm{hpd}=$ $67.31-105.32 \mathrm{Ma}, \mathrm{m}=85 \mathrm{Ma}$ respectively). The genera Porrorhynchus and Enhydrus are monophyletic with strong support ( $\mathrm{pp}=1.00$ ); both are long branches, and sister to the much larger genera Dineutus and Macrogyrus respectively. Mesodineutes $\dagger$ originated around 83 Ma , and is placed as sister to the extant genera Porrorhynchus and Dineutus, having gone extinct around 64 Ma . While this placement for Mesodineutes $\dagger$ is weakly supported ( $\mathrm{pp}=0.51$ ), the little morphology available is considerably more suggestive of this placement, than with Enhydrus and Macrogyrus (see Mesodineutes $\dagger$ discussion section under classification).

The clade Macrogyrus + Andogyrus (here after referred to as the genus Macrogyrus sensu n.) is strongly supported as monophyletic ( $\mathrm{pp}=1.00$ ) with Paleocene origins ( $\mathrm{hpd}=45.95$
$-72.73 \mathrm{Ma}, \mathrm{m}=59.22 \mathrm{Ma}$ ). The earliest diverging lineage within Macrogyrus, are Neotropical species representing the subgenus Andogyrus, which is strongly supported as a monophyletic group ( $\mathrm{pp}=1.00$ ), sister to the remaining non-South American species. The next branch is a clade of New Guinean species, representing the subgenus Cyclomimus, which is similarly strongly supported as being monophyletic ( $\mathrm{pp}=1.00$ ), diverging in the Eocene ( $\mathrm{hpd}=40.30-$ $65.81 \mathrm{Ma}, \mathrm{m}=51.97 \mathrm{Ma}$ ). Above this branch are species of Macrogyrus from Australia, grading into New Guinean and Wallacean species. This group represents the subgenus Macrogyrus sensu stricto. Macrogyrus striolatus is recovered as sister to the remaining species of the Macrogyrus sensu stricto, but with weak support ( $\mathrm{pp}=0.50$ ). The Australian species M. oblongus, M. rivularis, and $M$. reichei form a strongly supported clade ( $\mathrm{pp}=1.00$ ), but interestingly $M$. oblongus and $M$. rivularis are not recovered as sisters, instead $M$. rivularis is placed as sister to M. reichei $(\mathrm{pp}=1.00)$ with M. oblongus sister to both ( $\mathrm{pp}=1.00$ ). Macrogyrus howittii is placed in an isolated position as sister to the more derived species found in Australia, as well as those from New Guinea and Indonesia, with strong support ( $\mathrm{pp}=1.00$ ). The widespread Australian species Macrogyrus australis is found to be among the youngest (originating around 7 Ma ) and most derived members of Macrogyrus with strong support ( $\mathrm{pp}=0.96$ ).

The genus Dineutus is strongly supported as monophyletic ( $\mathrm{pp}=1.00$ ) with Eocene origins ( $\mathrm{hpd}=40.23-63.16 \mathrm{Ma}, \mathrm{m}=50.31 \mathrm{Ma}$ ). Within Dineutus there is a major split between primarily New Guinean and Southeast Asian species and those found mostly in Africa and North America. This clade is fairly well supported ( $\mathrm{pp}=0.79$ ) and represents the new sensu stricto subgenus as it includes species related to the type species. The other major clade sports the majority of Dineutus species and has strong support for monophyly ( $\mathrm{pp}=0.99$ ), this is the newly defined subgenus Cyclous sensu n. Within the subgenus Cyclous there are two groups, a strongly supported ( $\mathrm{pp}=0.97$ ) North American clade and a mostly African clade, with slightly less support ( $\mathrm{pp}=0.87$ ). The origin of the two subgenera and the major clades within Cyclous are place within the late Eocene (between $44-38 \mathrm{Ma}$ ).

Within the North American Cyclous clade there are two groups of species, a strongly monophyletic ( $\mathrm{pp}=0.96$ ) Nearctic only clade, and a weakly supported widely distributed ( $\mathrm{pp}=$ 0.68 ) clade consisting of mostly Central American species, the Caribbean species, and some Nearctic species. The Nearctic only clade includes some of the largest and the most widely distributed species within North America (e.g. D. ciliatus, D. discolor, and D. robertsi) (Gustafson \& Miller, 2015). Interestingly despite exceptionally similar morphology, D. ciliatus and $D$. robertsi are not recovered as sister species. Instead $D$. ciliatus is strongly supported ( $\mathrm{pp}=$ 0.96 ) as sister to a clade comprising $D$. serrulatus ( $D$. discolor $+D$. shorti). The newly described D. shorti (Gustafson \& Sites, 2016) is recovered as sister to the more widely distributed $D$. discolor ( $\mathrm{pp}=1.00$ ), having diverged from a common ancestor around 7 Ma . The earliest diverging lineage holds the large Central American species, D. truncatus and D. mexicanus which are strongly supported as sisters ( $\mathrm{p} p=1.00$ ) with the Caribbean D. longimanus strongly supported as sister to both ( $\mathrm{pp}=1.00$ ). The next branch has weakly supported placement ( $\mathrm{pp}=0.59$ ) and consists of two species that are strongly supported as sisters ( $\mathrm{pp}=$ 1.00), D. pagdeni and D. fairmairei, known from the Solomon Islands and Fiji respectively. Sister to these island species is a strongly supported monophyletic group ( $\mathrm{pp}=0.97$ ) with Nearctic and Central America species. Dineutus sublineatus is recovered as sister to the remaining members of this clade ( $\mathrm{pp}=0.97$ ). Interestingly another Central American species, $D$. solitarius is also recovered in a isolated position, as sister to a clade of species with a primarily Nearctic distribution ( $\mathrm{p} p=1.00$ ).

The primarily African clade similarly exhibits a large divide between members of the subgenus Protodineutus and those of species placed in the subgenus Spinosodineutes. Spinosodineutes as currently defined is strongly paraphyletic within the analysis. Dineutus australis the type species of the subgenus Cyclous is strongly supported ( $\mathrm{pp}=1.00$ ) as sister to the African species D. fauveli and D. subspinosus, both members of Spinosodineutes. Interestingly the other member of Spinosodineutes included in the analysis, D. striatus is strongly supported ( $\mathrm{pp}=1.00$ ) as sister to the large widespread Malagasy species, D. proximus. The clade containing the members of Protodineutus (including D. striatus of Spinosodineutes) is strongly supported as monophyletic ( $\mathrm{pp}=1.00$ ). Interestingly the Malagasy species $D$. sinuosipennis is recovered as the earliest diverging lineage ( $\mathrm{m}=34 \mathrm{Ma}$ ) and sister to all the species within group ( $\mathrm{pp}=1.00$ ). The other Malagasy species $D$. proximus is distantly related, nested well within a clade or primarily mainland Africa species.

The maximum likelihood (ML) analysis (Figure S5) generally supported the broader conclusions of the analysis. Importantly the ML analysis strongly supports monophyly of the Dineutini (bootstrap support, $\mathrm{bt}=96$ ). There is similarly strong support for the monophyly of Dineutus ( $\mathrm{bt}=97.4$ ) and Macrogyrus $(\mathrm{bt}=97.6)$. Within Macrogyrus there is strong support for the subgenera Andogyrus $(\mathrm{bt}=97.6)$ and Cyclomimus $(\mathrm{bt}=100)$. Enhydrus and Porrorhynchus are each strongly monophyletic $(b t=99)$ but are sister to one another, within a clade with the Gyrinini outgroup members. However, this is clearly a case of long branch attraction occurring in the analysis, known to effect ML analysis, despite selection of correct substitution model (Kück et al., 2012).

## Biogeographic

For the ancestral state estimation the DEC models fit the data significantly better than both DIVALIKE models (Table S4). The DEC +j model, including founder event speciation (Matzke, 2014), had a similar log-likelihood to the DEC model, but the Akaike weights identify this model as the overall best fit for the data (Table S4). Despite the difference in log-likelihood the DEC and DIVALIKE models recovered nearly identical ancestral state reconstructions (Figs. S7-14). The differences in estimation primarily relate to the ancestral ranges of the common ancestor of all Dineutini and the common ancestor of Dineutus subgenus Cyclous, however, with so many possible states the ancestral range is ambiguous for both (Figs. S7-S14). The models either suggest slightly high possibility for a Nearctic Cyclous common ancestor in the DEC +j model, or an Ethiopian and Nearctic ancestral range in the DEC and DIVALIKE models (Figs. S7 - S14). For the common ancestor of all Dineutini, the DIVALIKE models suggest higher likelihood for a common ancestor distributed in both Southeast Asia and South America (Figs. S11-S14).

The ancestor of both Enhydrus and Macrogyrus is recovered as being distributed in South America (Fig. 3). The ancestral state reconstruction supports an origin for Macrogyrus in the Paleocene of South America with subsequent dispersal to Australia around the early Eocene, coinciding with the Early Eocene Climatic Optimum (Fig. 3). The ancestral reconstruction then reveals numerous subsequent dispersal events out of Australia to the Melanesian area around the late Oligocene and early Miocene (Fig. 3). Dispersal to Less Sunda Islands in Wallacea happened most recently around the mid-Miocene.

In the Porrorhynchus and Dineutus clade, the common ancestor is estimated to have been distributed in the Oriental region during the Late Cretaceous (Fig. 3). The common ancestor of

Dineutus likely originated similarly in the Oriental region in the early Eocene, around the Early Eocene Climatic Optimum. In the Dineutus sensu stricto subgenus the common ancestor likely arose in the Oriental region, with subsequent dispersal to Papua New Guinea around the late Eocene (Fig. 3). There is considerable ambiguity related to the ancestral range of the common ancestor of the Dineutus subgenus Cyclous (Figs. S7-S14) preventing any conclusions about its location. The primarily North American clade within Cyclous is estimated to have had a Nearctic ancestor, and the primarily African clade an Ethiopian ancestor (Fig. 3). Given the isolated positions of African and North American at this time, these likely represent two different dispersal events. Within the North American clade dispersal to Central America occurred around the end Eocene (Fig. 3). Several dispersal events out of Central America are then inferred around the early Miocene and late Miocene (Fig. 3).

## Discussion

## Origins of Macrogyrus and the importance of Antarctica

The final stages of the breakup of Gondwana around 100 Ma , involved South America rifting from Africa moving west, while Antarctica and Australia migrated south remaining in close proximity with New Zealand rifting away from Antarctica around 80 Ma (Storey, 1995; Blakey, 2008). However, during the Late Cretaceous the southern tip of South America remained closely associated with western Antarctica into the early Paleogene (Reguero et al., 2014). Data from the Antarctic Peninsula and the southern tip of South America support a continuous landmass connecting South America to Antarctica during the Early Paleogene (Lawver et al., 2011). This landmass, dubbed the Wedellian Isthmus (Reguero et al., 2014), is proposed to have served as a land bridge, allowing faunal exchange between Patagonia and west Antarctica until around 57 Ma (Reguero et al., 2014). During this time ( $60-50 \mathrm{Ma}$ ) the ancestral state reconstruction (Fig. 3) suggests the common ancestor of the Cyclomimus subgenus dispersed from South America, across a cool-temperate Antarctica (Pross et al., 2012) to Australia. In support of this scenario, Patagonia is currently occupied by a single species M. (Andogyrus) seriatopunctatus (Brinck, 1977), which our analysis shows to be earlier diverging than its more northerly Andean sisters, M. (Andogyrus) zimmermanni in Peru and M. (Andogyrus) colombicus in Venezuela and Colombia (Fig. 2).

At around 50 Ma the Drake's passage opened (Fig. 3, DPO), severing the Wedellian Isthmus, separating South America from western Antarctica (Lawver \& Gahagan, 2003; Livermore et al., 2005) and opening of the Tasmanian Gateway to ocean currents cut Antarctic ties with Australia, leading to thermal isolation of Antarctica and global cooling (Bijl et al., 2013) and representing the final breakup of Gondwana (Blakey, 2008). While the ancestral state reconstruction does not support a Gondwanan vicariant origin for the subgenus Andogyrus, the isolation of this subgenus is associated with the final breakup of Gondwana and the dispersal event to Australia was possible via the last remaining Gondwanan connections of South America, Antarctica, and Australia. Antarctica clearly played a critical role in the evolution of Macrogyrus by facilitating dispersal to Australia. Antarctica likely supported a very unique Macrogyrus fauna given its near tropical warmth during the Eocene, until its glaciation around 30 Ma (Pross et al., 2012).

## Origins of Dineutus and its absence from South America

The common ancestor of Dineutus is reconstructed as arising within the Oriental region during the early Eocene (Fig. 3). Subsequent to this dispersal into the Nearctic and the Ethiopian regions likely occurred during the Mid-Eocene Climatic Optimum (Fig. 3). This period of time is way to recent for dispersal to the western hemisphere have occurred over the transatlantic De Geer Route or Thulean Route (Brikiatis, 2014). The most likely route to the Nearctic was through the Beringia, which during the Eocene was a lush swamp forest occupied by such thermophilic species as primates, tapirs, and alligators (Eberle \& Greenwood, 2011). Following the Eocene during the cooling of the Oligocene, dispersal to Central America occurred (Fig. 3). From here subsequent dispersals to the Caribbean occurred either directly, or through the Nearctic during the Miocene (Fig.3). All this data strongly suggest that the absence of Dineutus in South America is because the genus has yet to disperse there. The Dineutus species of the western hemisphere likely dispersed to the western hemisphere through Beringia into North America and have yet to spread further south than Panama.

Interestingly, the species located in the Solomon Islands and Fiji are reconstructed as having diverged from Central American ancestors around the Oligocene (Fig. 3). A transpacific dispersal even out of Central America seems highly unlikely. More probable is a second dispersal back across Beringia to the Palearctic, into the Oriental region and southward to the Melanesian area. A likely critical taxon for reconstructing this path is D. ritsemai, a species known only from Sulawesi. Dineutus ritsemai appears to be closely related to D. pagdeni and $D$. fairairei in that these three species all share a relatively rare morphological feature, the profemoral sub-apicoventral tooth being located on the anterior margin of the profemur's ventral face.

## Broader conclusions

Future sampling in Southeast Asia and the Sunda Islands for Dineutus species will greatly aid in reconstructing the region occupied by the common ancestor of the subgenus Cyclous. However, the reason for Dineutus species' absence from South America seems clear given the young age of the group and the estimated Oriental ancestral range of the common ancestor of Dineutus. The phylogenetic position of Porrorhynchus indicans may also effect the biogeographic reconstruction for the common ancestor of Porrorhynchus and Dineutus, being located in Sri Lanka. Sri Lanka may have held a central position at the heart of Gondwana along with Madagascar (Dissanayake \& Chandrajith, 1999). Given the age and phylogenetic position of $P$. indicans, its presence in Sri Lanka may be exceptionally important for the biogeographic reconstruction and origins of the common ancestor of the Dineutini. The only taxon with a unique distribution missing from the analysis for Macrogyrus is M. caledonicus from New Caledonia. However, this area is unlikely to alter the recovered biogeographic reconstruction.

## Classification

## Tribe Dineutini Desmarest, 1851

Dineutini Desmarest (1851): 225. Type genus Dineutus Macleay, 1825 by original designation. Synonyms: Enhydrini Régimbart, 1882; Dineutini Ochs, 1926; Prothydrinae Guignot (1954); Enhydrusini ICZN (2012)

Diagnosis. Within the Gyrinidae the Dineutini can be diagnosed by having the following combination of characters: 1) maxilla without galea, 2) elytron possessing nine elytral striae without accompanying sutural border, 3) metaventral wings (Hatch, 1925) in the form of a more-or-less equilateral triangle (Fig. 6) (Régimbart, 1882), 4) lobiform metanepisternum, 5) transverse metacoxae (Fig. 6), 6) female RT with greatly expanded, sac-like spermatheca without a well differentiated fertilization duct (Fig. 11, 12) (Miller \& Bergsten, 2012) 7) primary conjugation of sperm via the spermostyle (Fig. 13). The dineutine diagnosable traits are most similar to traits found in Heterogyrus, which also has nine elytral striae, the lateral wing of the metaventrite in the form of an equilateral triangle, and a lobiform metanepisternum. However, the elytra of Heterogyrus have sutural borders, which are absent in all dineutines, and the metacoxae of Heterogyrus are oblique, not transverse as in the dineutines. In regards to the female reproductive tract, the dineutines are most similar to the orectochiline genera Orectochilus and Orectogyrus. The dineutines however, never have the fertilization duct well differentiated or expanded. In Orectochilus the fertilization duct is well differentiated and somewhat removed from the bursa (Miller \& Bergsten, 2012). Most species of Orectogyrus have the fertilization duct greatly expanded, curled, and sclerotized, often forming a snail-shell shape (Brinck, 1956; Miller \& Bergsten, 2012). The lack of maxillary galea is an additional trait shared with orectochilines. Transverse metacoxae are also found in Spanglerogyrus, however, the metacoxae of Spanglerogyrus are weakly developed, and Spanglerogyrus does not have triangular metaventral wings. Some larger Patrus species have transverse metacoxae as well.

Taxonomy. The first formal description and diagnosis of the tribe was by Régimbart (1882). Régimbart (1882) did an excellent job providing potential morphological synapomorphies for the tribe and its constituent genera. Unfortunately an earlier division of the family Gyrinidae was proposed by Desmarest (1851) including some of the genera which Régimbart united in his seemingly new tribe Enhydrini, rendering it a junior synonym of Desmarest's Dineutini. The rediscovery of Desmarest's early name by Bouchard et al. (2011) was quite welcome, however, given the nomenclatural difficulty associated with Régimbart's proposed name for the tribe (Gustafson \& Miller, 2013). The constituent genera were greatly subdivided by the work of Georg Ochs (1924, 1926, 1949), the vast majority of which are not supported by the results of this analysis. A great testament to the outstanding work of Régimbart, we here return to the classification originally proposed by him in 1882 for the Dineutini, with only minor revision. The valid constituent species of the tribe Dineutini have not changed considerably since Régimbart's (1882) work, only growing in number following the taxonomic works of proceeding gyrinid experts.

Distribution. Members of the Dineutini have a global distribution, missing only from more northern latitudes and the arctic regions (Fig. 14).

Discussion. The sperm of Dineutus (Fig. 13D) was first described by Breland \& Simmons (1970), in which they discovered these species had primary conjugation via spermatodesma (as defined by Higginson \& Pitnick (2011)), they dubbed spermatostyles. Because sperm has been found to be phylogenetic informative (Baccetti, 1987), and sperm conjugation is relatively rare phenomenon (Pitnick et al., 2009) the sperm of the dineutine genera were sampled. The study revealed that all Enhydrus (Fig. 13 A - C ), Porrorhynchus (Fig. 13F), and Macrogyrus (Fig. 13E) all exhibit primary sperm conjugation via spermatostyles.

## Genus Dineutus Macleay, 1825

(Figs 1, 4C, 5E, 6D, 7A - D, 8C, 9E, 9G, 11B - D, 13D)
Dineutus Macleay, 1825: 30, type species Dineutus politus Macleay, 1825.
Synonyms: Necticus Laporte, 1835, Dineutes Régimbart, 1882.
Diagnosis. The genus Dineutus can be diagnosed within the Dineutini by the following combination of characters: 1) Gular suture complete, 2) frons without lateral bead (Fig. 4C), 3) antennal flagellum with 6-7 flagellomeres (Fig. 5E), 4) pronotal transverse impressed line present, 5) scutellar shield invisible with elytra closed, 6) protibia and male protarsi narrow (Fig. 9E), 7) mesotarsal claws sexually dimorphic, 8) metaventrite medially triangular in shape (Fig. 6D) and narrow, 9) female RT with vaginal shield (Fig. 11B, C, D) (Brinck, 1980, 1983, 1984). The genus Dineutus lacks a single distinct autapomorphy among gyrinid genera. A character that comes close are sexually dimorphic mesotarsal claws, but this character is a synapomorphy shared with Porrorhynchus, however, the sexual dimorphism is most pronounced among species of Dineutus. The other synapomorphies with Porrorhynchus include the invisible scutellar shield and most noticeably the female RT possessing a vaginal shield. Dineutus can be readily distinguished from all other dineutine genera by the narrowed protibia, which is likely the sole apomorphy separating this genus from Porrorhynchus. Dineutus can be furthered distinguished from Porrorhynchus in having a complete gular suture and the pronotal transverse impressed line present.

Taxonomy. The genus was monotypic when originally erected by Macleay (1825). Régimbart subsequently treated the genus several times, revising it and adding many species (Régimbart, 1882, 1886, 1892, 1907). Hatch (1925) was the first author to divide the genus into subgenera, based primarily on overall body-shape. Georg Ochs $(1926,1955)$ subsequently erected numerous subgenera, including subsuming Porrorhynchus as one of the subgenera. Since Ochs' work, no new subgenera have been proposed, but the composition of the subgenera has been re-arranged by Guignot (1950), and most recently by Brinck (1955b), who attempted to provide distinct morphological traits identifying each subgenus, unsuccessfully.

There are currently 92 species within the genus Dineutus, making it easily the largest genus within the Dineutini.

Distribution. Dineutus has a near global distribution, missing from Europe, and most notably from South America (Fig. 14D) (Mouchamps, 1949b; Brinck, 1955b; Satô, 1962; Brinck, 1976; Mazzoldi, 1995; Watts \& Hamon, 2010; Hájek, 2011; Gustafson \& Miller, 2015; Lee \& Ahn, 2015). Currently the highest diversity is in the Austral region, primarily in New Guinea, but this likely reflect bias due to recent taxonomic work on species in this region (i.e. Brinck (1976, 1981, 1983, 1984). The second highest diversity is found in tropical Africa.

Discussion. This is the largest and most widely distributed genus within the Dineutini.

## Subgenus Dineutus sensu novo

(Figs. 1, 7C, 9E, 11B)
Type species: Dineutus politus Macleay, 1825.

Synonyms: Rhombodineutus Ochs, 1926 new synonymy, Merodineutus Ochs, 1955 new synonymy

Diagnosis. Within Dineutus the sensu stricto subgenus can be diagnosed by the following characters: 1) head capsule of most species with a frons to clypeus ratio less than or equal to 1.5 , 2) a transverse, rounded labrum, 3) distolateral angle of protibia without spine, 4) protrochanter glabrous (Fig. 7C) -without setae apically on ventral face, 5) mesotarsal claws distinctly sexually dimorphic. The Dineutus sensu stricto subgenus contains the largest members of the genus (e.g. Dineutus macrochirus) (Brinck, 1984). Most species exhibit little to no distinguishable sexual dimorphism in terms of elytral shape. The mesotarsal claws are distinctly sexually dimorphic, but not nearly as well developed as those of the Cyclous subgenus.

Taxonomy. There are now 23 species within the sensu stricto subgenus, containing members of the former subgenera Merodineutus and Rhombodineutus. The species of this group were last treated by Mouchamps (1949b) (the original sensu stricto species), Brinck (1983) (the Rhombodineutus species), and Brinck (1984) (Merodineutus species).

Distribution. Primarily distributed in New Guinea and Southeast Asia. One species, D. insularis extends into the far eastern Palearctic being found on the Ryukyu islands.

Discussion. The distinction of Merodineutus from Dineutus was tenuous, based primarily on elytral sculpture, protarsus, and protibial modifications (Brinck, 1984). Brinck (1984) even predicted the derivation of Merodineutus from Dineutus sensu stricto. The subgenus Rhombodineutus was similarly based on elytral modifications resulting in a rhomboid body outline, and a more elongate labrum than other species of Dineutus (Brinck, 1983). Many Dineutus species show unique modifications to the elytral apices and protibial modifications as exhibited by the diversity of North American Dineutus (Gustafson \& Miller, 2015). The large glabrous protrochanters within Dineutus are unique to this clade. For this reason, the other subgenera are synonymized with the Dineutus sensu stricto subgenus.

The close relation found here between Rhombodineutus and Merodineutus is novel. A phylogenetic analysis of the species of this area, including Rhomborhynchus, would prove quite interesting in elucidating directionality of colonization of New Guinea and validity of the numerous described species and subspecies (Brinck, 1983, 1984).

## Subgenus Cyclous Dejean, 1833 sensu novo

(Figs 1, 4C, 5E, 6D, 7A - B, 7D, 8C, 9G, 11C, 13D)
Type species: Dineutus australis (Fabricius, 1775).
Synonyms: Callistodineutus Ochs, 1926 new synonymy, Cyclinus Kirby, 1837 new synonymy, Gyrinodineutus Ochs, 1926, Paracyclous Ochs, 1926 new synonymy, Protodineutus Ochs, 1926 new synonymy, Spinosodineutes Hatch, 1925 new synonymy.

Diagnosis. Within Dineutus the Cyclous subgenus can be diagnosed by the following characters: 1) Head capsule with a frons to clypeus ratio less than or equal to $1.5,2$ ) a transverse, rounded labrum, 3) distolateral angle of protibia without spine, 4) ventral face of protrochanter apically with series of stout setae (Fig. 8C), 5) mesotarsal claws strongly sexually dimorphic 6)
spermatheca not tubiform, less elongate and more rounded. Many species are strongly sexually
dimorphic in elytral shape. This group exhibits the most strongly sexually dimorphic mesotarsal claws.

Taxonomy. This is the largest subgenus, now with 67 species. The species were treated taxonomically most recently by Mouchamps (1949a) (the Spinosodineutes species), Brinck (1955b) (African species), Brinck (1976) (the Callistodineutus species), and Gustafson \& Miller (2015) (the North American species).

Distribution. Widely distributed, found in North America, Africa, Asia, and Australia.
Discussion. The numerous subgenera of Dineutus have long been a source of conflict among gyrinid workers (Hatch, 1925; Ochs, 1926; Guignot, 1950; Brinck, 1955b; Ochs, 1955). The first division of Dineutus into subgenera was proposed by Hatch (1925), but the majority of subgenera was erected by Ochs (1926) during his pre-cladistic systematic treatment of the species of Dineutus (and Porrorhynchus, see below). The subgenera have nearly all been diagnosed in the past by body form, modification to the elytral apex, and/or elytra reticulation. These characters are highly variable among the numerous Dineutus species, and typically not unique to any one subgenus, causing much of the disagreement between constituent species.

The only authority to attempt to propose discrete morphological characters for the subgenera was Brinck (1955b), but was unsuccessful, resorting to the distinction of African species and American species for the subgenera Protodineutus and Cyclinus respectively. However, our analysis shows Callistodineutus to be nested within the North American species, despite a proposed distinct morphological character, suggesting those utilized by Brinck (1955b) were unsuccessful in identifying large natural groups of species. The distinct character of the ventral face of the protrochanter with a series of short stout setae apically, in combination with the other diagnostic features successfully recognizes a large monophyletic group within Dineutus. While D. ritsemai was not included in the phylogenetic study, the taxon was studied for morphology. Dineutus ritsemai has well-developed sexually dimorphic mesotarsal claws, and resembles closely members of the former subgenus Callistodineutus having a single profemoral sub-apicoventral tooth on the anterior face only. Given the former species are nested within the North American members, including this species and synonymizing Paracylous with Cyclous is justified. For this reason we here synonymous the former subgenera. The oldest available name for this grouping is Cyclous initially proposed by Dejean, 1833 for Dineutus australis, one of the most widespread species of Dineutus (Ochs, 1949).

This subgenus is notable for having numerous sexually dimorphic traits. Many species have sexually dimorphic elytral apices, often with one sex having thorn-like productions. This is exhibited in several North American species (Gustafson \& Miller, 2015). This group also exhibits sexually dimorphic modification to the protrochanter, such as the strange waxy region of male Dineutus proximus (Fig. 7B), and most notably the setose brush of D. australis males (Fig. 7A). The male mesotarsal claws are also strongly sexually dimorphic in this group. The North American species exhibit species-specific sexually dimorphic claws, with the claws of D. nigrior being the most extremely dimorphic known (Gustafson \& Miller, 2015). The median lobe of the aedeagus of members of the subgenus Cyclous also present a wide diversity of forms, not seen elsewhere within Dineutini. No other dineutine group exhibits such a suite of sexually selected traits.

## Subgenus Rhomborhynchus Ochs, 1926 incertae sedis

(Figs 11D, S5)
Rhomborhynchus Ochs, 1926: 65
Type species: Porrorhynchus depressus Régimbart, 1907.
Diagnosis. Within Dineutus the subgenus Rhomborhynchus can be diagnosed by the following characters: 1) head capsule with a frons to clypeus ratio of greater than or equal to $1.5,2$ ) labrum elongate and triangular, 3) labrum with a longitudinal paired row of setae, and one transverse row, 4) spinose distolateral corner of the protibia, 5) ventral face of protrochanter apically with series of stout setae, 6) mesotarsal claws weakly sexually dimorphic, 7) female RT with tubiform spermatheca. These species are most similar to members of the former subgenus
Rhombodineutus having relatively elongate labra and a greatly elongate spermatheca (Fig. 11D). But can be distinguished by the spinose distolateral corner of the protibia, the more strongly triangular labrum, and the presence of setae apically on the ventral face of the protrochanter.

Taxonomy. Two species, $D$. depressus and $D$. misoolensis.
Distribution. Known from New Guinea and the neighboring island of Misool. Widespread within New Guinea.

Discussion. Rhomborhynchus was originally erected as a subgenus of Dineutus, however the type species D. depressus has mostly been considered a member of Porrorhynchus for much of its history (Régimbart, 1907; Guignot, 1950; Brinck, 1955b). Ochs (1926) was the first to recognize the different features of D. depressus relative to the members of Porrorhynchus and provided a discussion of why this taxon and several others proposed by him should be considered members of Dineutus (Ochs, 1955). However, Ochs (1926) did not recognize the unique autapomorphies of the other Porrorhynchus species in relation to Dineutus.

This subgenus exhibits numerous similarities to members of the former subgenus Rhombodineutus, such as 1) elongate labra, 2) a more longitudinal orientation to the labral setation, 3) rhomboid body-outline. "Rhombodineutus" species also have a relatively elongate spermatheca (Fig. 11B) compared to other Dineutus members. However, Rhomborhynchus species have setae situation apically on the ventral face of the protrochanter, suggesting placement outside of the Dineutus sensu stricto subgenus and away from the species of the former subgenus Rhombodineutus. The lack of sexually dimorphic traits and weakly sexually dimorphic mesotarsal claws also suggest Rhomborhynchus is not a member of the subgenus Cyclous. Rhomborhynchus species also lack all the synapomorphic characters of Porrorhynchus sharing only seemingly plesiomorphic features like the elongate labrum and the tubiform spermatheca.

Unfortunately, no molecular-grade specimens of Rhomborhynchus were available for this study and analysis only used morphological characters. The Bayesian analysis placed Rhomborhynchus well within Dineutus (Cyclous) in a polytomy with the Malagasy species Dineutus sinuosipennis (Fig. S5), which seems highly unlikely. As the analysis placed the subgenus well within Dineutus and its lack of synapomorphic characters shared with members of Porrorhynchus, it seems safe to tentatively transfer the species to this genus for the time being, but with an incertae sedis in relation to the other Dineutus subgenera. The final placement of this
subgenus is clearly still in question. Future phylogenetic analyses including molecular grade Rhomborhynchus specimens will be necessary to resolve its phylogenetic position.

## Genus Enhydrus Laporte, 1835

(Figs 2, 4A, 5D, 6A, 9F, 12F - G, 13A - C)
Type species: Enhydrus sulcatus (Wiedemann, 1821).
Synonyms: Epinectus Aubé, 1838, Epinectes Régimbart, 1877, Prothydrus Guignot, 1954
Diagnosis. Within the tribe Dineutini Enhydrus can be diagnosed by the following combination of characters: 1) antenna of most species with 7 flagellomeres (Fig. 5D) -one with 6, 2) fons with lateral bead (Fig. 4A), 3) pronotal transverse impressed line present, 4) elytral striae present as strongly impressed lines, 5) scutellar shield visible with elytra closed, 6) protibia laterally expanded apically (as in Fig. 8A), 7) broad, compact male protarsi (Fig. 9F), protarsi of both sexes often with fused segments and large protarsal claws, 8) metaventrite medially pentagonal in shape (Fig. 6A), 9) suture of abdominal sternite II present, 10) female RT without vagina shield, gonocoxae short and stout (Fig. 12G).

Taxonomy. There are four known species in the genus. The species of Enhydrus were last treated taxonomically by Brinck (1978).

Distribution. Disparately distributed in South American and extreme southeastern Central America (Fig. 14C) (Brinck, 1977).

Discussion. The genus Enhydrus lacks a single autapomorphy, however, retention of a fully developed suture to abdominal sternite II is unique to this genus. Fusion of the protarsomeres is unique to Enhydrus as well, but not all species exhibit protarsomere fusion (E. tibialis does not have fused protarsomeres). Molecular data (Fig. S4) however strongly support Enhydrus is a distinct monophyletic group and in general morphology species strongly resemble one another, despite lacking a distinct synapomorphy.

## Genus Macrogyrus Régimbart, 1882

(Figs 2, 4B, 4D, 4E, 5A - B, 6B, 7F, 8A, 8D, 9A - C, 10, 12A - E, 13E)
Type species: Macrogyrus howittii (Clark, 1866).
Diagnosis. Within the tribe Dineutini, Macrogyrus can be diagnosed by the following combination of characters: 1) antennae with 9 flagellomeres (Fig. 5A - B), 2) frons with lateral bead (Fig. 4B, 4D, 4E), 3) pronotal transverse impressed line present, 4) scutellar shield visible with elytra closed, 5) protibia laterally expanded apically (Fig. 8A), 6) protarsus of male broad, discus present ventrally on protarsomere I (described blow) (Fig. 9A - C), 7) metacoxal process bordered posterolaterally (Fig. 6B), 8) female RT without vaginal shield, gonocoxae elongate (Fig. 13B).

Taxonomy. There are now 54 species of Macrogyrus with the inclusion of the former genus Andogyrus. This genus has never received a comprehensive revision.

Distribution. Found in South America, Australia, New Caledonia, New Guinea and surrounding islands, and Lesser Sunda Islands (Fig. 14D) (Ochs, 1949, 1953, 1955; Brinck, 1976, 1977; Watts \& Hamon, 2010).

Discussion. This genus exhibits an excellent autapomorphy: the male protarsus has protarsomere I with a recessed pit possessing adhesive setae with a different suction cup morphology than the remaining adhesive setae (Fig. $9 \mathrm{~A}-\mathrm{C}$, di). This character was first described by Régimbart (1882: 433) and dubbed the discus. This feature is a synapomorphy uniting all the Macrogyrus species.

## Subgenus Andogyrus Ochs, 1924 new status

(Figs 2, 4D, 5B, 6B, 8A, 8D, 9A, 12A)
Type species: Andogyrus ellipticus (Brullé, 1836).
Synonyms: Proteogyrus Mouchamps, 1951.
Diagnosis. Within the genus Macrogyrus, Andogyrus can be diagnosed by the following combination of characters: 1) clypeus narrow, 2) elytra without canaliculate microsculpture 3) metaventrite medially pentagonal in form (Fig. 6B), 4) metaventral discrimen with elongate transverse sulcus ancestrally (as in Fig. 10A). The elongate transverse sulcus of the metaventral discrimen is lost in many species of the subgenus Andogyrus, but its presence in $M$.
seriatopunctatus suggest the absence to be a secondary loss, given its phylogenetic position (Fig. $2)$.

Taxonomy. This subgenus has twenty known species. The species of this subgenus were last treated by Brinck (1977).

Distribution. Found along the Andes of South America, from Venezuela to Argentina (Brinck, 1977).

Discussion. The separation of Andogyrus from Macrogyrus was based primarily on distribution (Ochs, 1924) and Hatch (1925) would prove quite correct in asserting that the Australian Macrogyrus were derived from a common ancestor similar to Andogyrus. As can be seen from the phylogeny (Fig. 2) Andogyrus is far too similar to Macrogyrus to be regarded as a genus distinct from the latter. Instead Andogyrus should be regarded as an early diverging lineage within Macrogyrus. Especially given the very distinct synapomorphy of the male protarsal discus. Separating these two groups into formal genera would also suggest Cyclous and Dineutus sensu stricto deserve separation into distinct genera, using similar phylogenetic logic.

## Subgenus Cyclomimus Ochs, 1949 sensu novo

(Figs 2, 4F)
Type species: Macrogyrus purpurascens Régimbart, 1882.
Synonyms: Stephanogyrus Ochs, 1955 new synonymy
Diagnosis. Within the genus Macrogyrus, Cyclomimus can be diagnosed by the following combination of characters: 1) clypeus considerably enlarged (Fig. 4F), 2) elytra without canaliculate microsculpture 3) metaventral discrimen without transverse sulcus. Some of the

New Guinean species exhibit unique modification to the adhesive setae of the male protarsus. The discus still retains a relatively normal amount of setae, however outside the discus the adhesive setae are reduced in number, nearly absent from the ventral face of the ultimate protarsomere, and have very large suction cups. The species within this group are smaller in body size than most other members of Macrogyrus, but not all.

Taxonomy. Five known species, and the subgenus is returned to its original sense as initially proposed by Ochs (1949). The species were most recently treated by Ochs (1955) (for the New Guinea species) and by Mazzoldi (2010) (for M. caledonicus).

Distribution. Primarily found in New Guinea (fours species) (Ochs, 1955) where it appears widespread, with one species from Grande Terre, New Caledonia (Mazzoldi, 2010).

Discussion. The subgenus Stephanogyrus was erected for the single species M. caledonicus by Ochs (1954) based only on modifications to the elytra. While this species was not included in the formal phylogenetic analysis, specimens were studied and found to exhibit the diagnostic features uniting the monophyletic group of species from New Guinea. Furthermore, this returns Cyclomimus to its original sense, prior to splitting of a single isolated species from New Caledonia.

## Subgenus Macrogyrus sensu novo

(Figs 2, 4B, 5A, 7F, 9B - C, 10, 12D - E, 13F)
Type species: Macrogyrus howittii (Clark, 1866).
Synonyms: Australogyrus Ochs, 1949 new synonymy, Ballogyrus Ochs, 1949 new synonymy, Clarkogyrus Ochs, 1949 new synonymy, Megalogyrus Ochs, 1949 new synonymy, Orectomimus Ochs, 1930 new synonymy, Tribologyrus Ochs, 1949 new synonymy, Tribolomimus Ochs, 1949.

Diagnosis. Within the genus Macrogyrus, the sensu stricto subgenus, can be diagnosed by the following combination of character: 1) clypeus neither narrow nor considerably enlarged (Fig. 4B), 2) elytra with unique canaliculate microsculpture (Fig. 10B - C), 3) metaventral discrimen of most species with well developed transverse sulcus (Fig. 10A). The unique canaliculate microsculpture (Fig. 10B-C) is an excellent autapomorphy for the sensu stricto subgenus. This character is strongly reduced in one species M. sumbawae (Fig. 2), but is still faintly evident apically on the elytra.

Taxonomy. There are now 29 species within this subgenus, a massive increase from the former classification, in which the subgenus only contained the type species, M. howittii (Ochs, 1949). The Australian species are the most well known (Ochs, 1949, 1956) and were recently treated by Watts \& Hamon (2010), making their identification possible. The New Guinean fauna and those of the surrounding islands are in desperate need of revision following the work of Ochs (1955), in which the few known species were divided into numerous subspecies, from disparate locations in New Guinea, based on few specimens. The work of Ochs (1955), including no illustrations, non-discrete morphological characters, and excessive splitting of species, has made the identification of New Guinean specimens exceptionally difficult. For this reason, most species in the analysis were unable to be identified reliably.

Distribution. Primarily known from Australia and New Guinea, also found in the islands surrounding New Guinea, and the Lesser Sunda Islands (Ochs, 1949, 1955).

Discussion. The new definition of the sensu stricto subgenus is based on the earliest diverging taxon suggesting a common ancestor with canaliculate microsculpture (Fig. 10B - C), which in this analysis is M. striolatus. However, the placement of M. striolatus is weakly supported (Fig. $\mathrm{S} 3, \mathrm{~S} 4)$. It is possible that the subgenus Cyclomimus is nested within the sensu stricto subgenus, as examination of the .t tree files from the Bayesian analysis show the placement of M. striolatus fluctuating between a position above or below the Cyclomimus clade. In the case M. striolatus is truly earlier diverging than the Cyclomimus clade, the putative synapomorphic character of the sensu stricto subgenus still stands, with an inferred subsequent loss of the canaliculate microsculpture in Cyclomimus. Reduction of the canaliculate microsculpture is seen in the more derived members of the sensu stricto subgenus, e.g. M. sumbawae (Fig. 2) and other species found in Wallacea. The species of Cyclomimus show their other highly derived characters (e.g. the reduction in number and expansion in size of adhesive setae of the male protarsus; a largely expanded clypeus, reduction of the transverse sulcus of the metaventral discrimen). Therefore, a convergent derived loss of the canaliculate microsculpture is certainly plausible. Because of the strong support for the monophyly of the Cyclomimus subgenus in the analysis is currently retained as a valid subgenus separate from the sensu stricto, but the definition of the Macrogyrus sensu stricto subject to change in future phylogenetic analyses depending upon the placement of M. striolatus.

## Genus Mesodineutes $\dagger$ Ponomarenko, 1977

(Figs. 6E)
Type species Mesodineutes amurensis Ponomarenko, 1977
Diagnosis. Within the tribe Dineutini Mesodineutes can be diagnosed by the following combination of characters: 1) elytral striae present as punctures, 2) elytral apex rounded, without apicolateral sinuation or other modification, 3) metaventrite medially triangular in shape (Fig. 6 E ) and broad, 4) metacoxal process without border posterolaterally (Fig. 6E).

Taxonomy. This fossil genus is monotypic.
Distribution. Described from the Paleocene of southeastern Russian Federation (Ponomarenko, 1977).

Discussion. While the support for the phylogenetic placement of this species was not strong (Fig. S3), the available morphological information and its distribution strongly support its placement with Porrorhynchus and Dineutus. Similar to Porrorhynchus and Dineutus, Mesodineutes has a triangular shaped medial expanse to the metaventrite (Fig. 6 C - E), while Enhydrus and Macrogyrus have a more pentagonal shape (Fig. 6 A - F). Mesodineutes can further be separated from a close relation with Macrogyrus in that it lacks a border to the posterolateral margin of the metacoxae (Fig. 6E compared to 6B). Importantly this species is found in the Palearctic of the Paleocene, which according to the biogeographic analysis suggests it does not belong in the clade with Enhydrus + Macrogyrus whose ancestors evolved in or near Australia. Importantly it
also supports the biogeographic reconstruction that the ancestor of Porrorhynchus + Dineutus was likely found in or near the Oriental region (Fig. 3).

## Genus Porrorhynchus Laporte, 1835

(Figs. 1, 4E, 5C, 6C, 7E, 8B, 9D, 11A, 13F)
Type species: Porrorhynchus marginatus Laporte, 1835
Synonyms: Ceylorhynchus Brinck, 1955
Diagnosis. Within the tribe Dineutini Porrorhynchus can be diagnosed by the following combination of characters: 1) Labrum elongate and triangular in form (Fig. 4E), 2) gular suture incomplete, 3) frons without lateral bead (Fig. 4E), 4) antennal flagellum with 6-8
flagellomeres (Fig. 5C), 4) pronotal transverse impressed line absent, 5) scutellar shield invisible with elytra closed, 5) male protrochanter with setose patch (Fig. 7E), 6) male protarsi narrow (Fig. 9D), 7) protibia expanded distolaterally (8B), 8) ventral face of profemur with two rows of setae arranged into large clusters, progressively becoming denser apically, 9) mesotarsal claws weakly sexually dimorphic, 8) metaventrite medially triangular in shape (Fig. 6C) and narrow, 9) female RT with vaginal shield (Fig.11A). Diagnostic characters (5) and (8) appear apomorphic among all Gyrinidae.

Taxonomy. There are now three species within the genus, following removal of the former subgenus Rhomborhynchus.

Distribution. Widely distributed in Southeast Asia west of Wallace's line, as far northwest as southeastern Tibet (Jäch et al., 2012) and east through southern China (Fig. 14B). One species, P. indicans, known from Sri Lanka (Brinck, 1980).

Discussion. This genus contains the largest known species of whirligig beetle ( $P$. landaisi) and species apparently very sensitive to water quality (Ochs, 1927; Brinck, 1980). Among the Porrorhynchus species, $P$. indicans is of the most concern in terms of conservation, found to already be uncommonly encountered and limited in distribution in 1980's due to deforestation of preferred habitat montane forests within Sri Lanka (Brinck, 1980). This is especially concerning given the unique information $P$. indicans can potentially provide for future analyses (see Discussion).

## Key to the extant genera of the Dineutini

1. Scutellar shield invisible with elytra closed; sexually dimorphic mesotarsal claws (even if weakly so). Female RT with vaginal shield (Fig. 11C, vs).
-Scutellar shield visible with elytra closed; mesotarsal claws not sexually dimorphic. Female RT without vaginal shield (Fig. 12).
2. Pronotum without transverse impressed line; ventral face of profemur with two rows of setae arranged in large clusters, becoming denser apically; protrochanter of male with setose patch (Fig. 7E); mesotarsal claws weakly sexually dimorphic

- Pronotum with transverse impressed line; setae of ventral face of profemur not arranged into large cluster becoming denser apically; protrochanter of male without setose patch, variously modified or not; mesotarsal claws sexually dimorphic, often strongly so

Dineutus
4. Elytra with striae in the form of well impressed lines; protarsus (male and female) compressed often with fused segments; male protarsus ventrally without discus (Fig. 9F).

Enhydrus

- Elytra with striae in the form of punctures or weakly impressed lines, never as well impressed lines; protarsus without compressed or fused segments; male protarsus ventrally with discus (Fig. $9 \mathrm{~A}-\mathrm{C}$ ).

Macrogyrus

## Key to the subgenera of Dineutus

1. Labrum elongate and strongly triangular in form; distolateral corner of protibia produced into a spine.

Rhomborhynchus

- Labrum most often not elongate, strongly rounded, never triangular in form; distolateral corner of protibia not produced into a spine.

2. Protrochanter of both sexes glabrous (Fig. 7C); species without strongly sexually dimorphic elytra.

Dineutus sensu stricto
— Protrochanter of both sexes with setae situated apically on ventral face (Fig. 7D), males of some species with modification (i.e. brushes); many species sexually dimorphic in elytral modification.

Cyclous

## Key to the subgenera of Macrogyrus

1. Elytra with canaliculate microsculpture presenting themselves as distinct "scratches" (Fig. 10B); if scratches barely present or even absent, then species large (ca. 10 mm ), clypeus neither narrow, nor exceptionally large, and without strongly pentagonal medial expanse of the metaventrite (Fig. 10A).

Macrogyrus sensu stricto

- Elytra without canaliculate microsculpture.

2. Clypeus narrow (Fig. 4D); elytral apices unmodified (i.e. without apicolateral sinuation and/or productions); most species very large (i.e. $\geq 10 \mathrm{~mm}$ ). Only found in South America. Andogyrus

- Clypeus wide and greatly enlarged (Fig. 4F); elytral apices modified, with apicolateral sinuation and truncate apex. Most species with males having the adhesive setae of the ventral face of the protarsus modified into fewer and larger cups. New Guinea and New Caledonia.


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## Figure captions

Figure 1. Phylogeny of the Dineutini based on Bayesian total-evidence analysis Part 1. Labels at node denote median age, blue bars indicate $95 \%$ hpd for age, and asterisks above node indicates pp of $\geq 0.95$.The blue clades indicate members of Dineutus, with lighter blue showing the subgenus Cyclous sensu n. and the dark blue the sensu stricto subgenus. Purple indicates the genus Porrorhynchus. Species are approximately to relative scale.

Figure 2. Phylogeny of the Dineutini based on Bayesian total-evidence analysis Part 2. Labels at node denote median age, blue bars indicate $95 \%$ hpd for age, and asterisks above node indicates pp of $\geq 0.95$.The green clades indicate members of Macrogyrus, with the lightest green showing the sensu stricto subgenus, the next darkest members of the subgenus Cyclomimus, and the darkest green showing the subgenus Andogyrus stat. n.. Orange indicates the genus Enhydrus. Species are approximately to relative scale.

Figure 3. Historical biogeography of the dineutine whirligig beetles. The Bayesian totalevidence chronogram is plotted as used in the biogeographic analysis. Blue bars indicate $95 \%$ hpd for age. The circle at the node shows the preferred ancestral state reconstruction from the BioGeoBears results (Figs. S6 - S9). The following abbreviations are used: DG R, De Geers route; Th R, Thulean route; EECO, Early Eocene Climatic Optimum; DPO, Drake's Passage opening; MECO, Mid-Eocene Climatic Optimum. The map legend just below the tree shows the color key for the ancestral state reconstructed. The paleogeographic maps below © Colorado Plateau Geosystems used with permission, show continental positions in the major time slices (Blakey, 2008).

Figure 4. Head capsules of dineutine species, anterior view, scale bars $=1 \mathrm{~mm}$, abbreviations: $\mathrm{lbr}=$ labrum, cly = clypeus, $\mathrm{frb}=$ frontolateral bead. (A) Enhydrus sulcatus; (B) Macrogyrus (Macrogyrus) australis; (C) Dineutus (Cyclous) australis; (D) M. (Andogyrus) seriatopunctatus; (E) Porrorhynchus landaisi; (F) M. (Cyclomimus) purpurascens.

Figure 5. Antennae of dineutine species, above anterior view, below posterior, scale bar $=0.5$ mm. (A) Macrogyrus (Macrogyrus) australis; (B) M. (Andogyrus) zimmermanni; (C) Porrorhynchus landaisi; (D) Enhydrus tibialis;
(E) Dineutus (Cyclous) australis.

Figure 6. Meso- and meta-ventrites of dineutine species and a gyrinine species, ventral view, scale bars $=1 \mathrm{~mm}$, except (F). (A) Enhydrus sulcatus: (B) Macrogyrus (Andogyrus) colombicus, (C) Porrorhynchus marginatus; (D) Dineutus (Cyclous) carolinus: (E) Mesodineutes amurensis $\dagger$ : (F) Gyrinus maculiventris, scale bar $=0.5 \mathrm{~mm}$.

Figure 7. Prtrochanters of male dineutine species, ventral view, abbreviations: $\mathrm{ts}=$ protrochanteric setae, $\mathrm{wx}=$ waxy spot, $\mathrm{pt}=$ protrochanteric setose patch. $(\mathrm{A})$ Dineutus $($ Cyclous $)$ australis, scale bar $=200 \mu \mathrm{~m}, \mathrm{pb}=$ protrochanteric brush; (B) $D$. (Cyclous) proximus, scale bar $=$ $500 \mu \mathrm{~m}$; (C) D. (Dineutus) n. sp., scale bar $=300 \mu \mathrm{~m}$; (D) D. (Cyclous) serrulatus analis, scale bar $=300 \mu \mathrm{~m}$, (E) Porrorhynchus marginatus, scale bar $=400 \mu \mathrm{~m}$; (F) Macrogyrus (Macrogyrus) albertisii, scale bar $=500 \mu \mathrm{~m}$.

Figure 8. Prolegs of male dineutine species, abbreviations: $\mathrm{sb}=$ setose brush, asr $=$ anterior row of profemoral setae, $\mathrm{psr}=$ posterior row of profemoral setae, $\mathrm{sp}=$ setigerous puncture. (A) Macrogyrus (Andogyrus) zimmermanni protibial apex, posterior view, scale bar $=400 \mu \mathrm{~m}$; (B) Porrorhynchus marginatus protibia, posterior view, scale bar $=500 \mu \mathrm{~m}$; (C) Dineutus (Cyclous) australis protrochanter and profemur, ventral view, scale bar $=500 \mu \mathrm{~m}$; (D) M. (A.) zimmermanni protrochanter and profemur, ventral view, scale bar $=1 \mathrm{~mm}$.

Figure 9. Protarsus of male dineutine species, abbreviations: $\mathrm{di}=$ protarsal discus, $\mathrm{sb}=$ setose brush. (A) Macrogyrus (Andogyrus) zimmermanni, scale bar $=500 \mu \mathrm{~m}$; (B) M. (Macrogyrus) sp., scale bar $=500 \mu \mathrm{~m}$; (C) M. (M.) albertisi, scale bar $=500 \mu \mathrm{~m}$; (D) Porrorhynchus marginatus, scale bar $=1 \mathrm{~mm}$; (E) Dineutus (Dineutus) n sp, scale bar $=1 \mathrm{~mm}$; (E) Enhydrus atratus, scale bar $=2 \mathrm{~mm}$; (G) D. $($ Cyclous $)$ australis, scale bar $=500 \mu \mathrm{~m}$.

Figure 10. Sculpture of Macrogyrus (Macrogyrus) albertisi. (A) metaventrite showing, $\mathrm{md}=$ metaventral discrimen, tvs = transverse sulcus, scale bar $=1 \mathrm{~mm}$; $(B)$ elytra with canliculate microsculpture, scale bar $=300 \mu \mathrm{~m}$; (C) canaliculate microsculpture, scale bar $=40 \mu \mathrm{~m}$.

Figure 11. Female reproductive tracts, ventral view, abbreviations: $\mathrm{sp}=$ spermatheca, $\mathrm{fd}=$ fertilization duct, $\mathrm{ov}=$ common oviduct, $\mathrm{bu}=$ bursa, $\mathrm{lt}=$ laterotergite, $\mathrm{vs}=$ vaginal shield, $\mathrm{mp}=$ medial apodeme of gonocoxa, gc = gonocoxa; scale bars $=1 \mathrm{~mm}$. (A) Porrorhynchus landaisi; (B) Dineutus (Dineutus) tetracanthus; (C) D. (Cyclous) discolor; (D) D. (Rhomborhynchus) depressus.

Figure 12. Female reproductive tracts, abbreviations: $b g=$ bursal gland; scale bars $=1 \mathrm{~mm}$. (A) Macrogyrus (Andogyrus) seriatopunctatus, ventral view; (B) gonocoxa of the same; (C) lateral view of the same; (D) M. (Macrogyrus) gouldii, ventral view; (E) gonocoxa of the same; (F) Enhydrus tibialis, ventral view; (G) gonocoxa of the same.

Figure 13. Sperm of dineutine species, exhibiting primary conjugation via spermatostyles. (A) Enhydrus atratus, scale bar $=300 \mu \mathrm{~m}$; (B) the same, scale bar $=50 \mu \mathrm{~m}$; (C) the same with single sperm, scale bar $=20 \mu \mathrm{~m}$; (D) Dineutus emarginatus, scale bar $=50 \mu \mathrm{~m}$; (E) Macrogyrus (Macrogyrus) rivularis, scale bar $=100 \mu \mathrm{~m}$; (F) Porrorhynchus marginatus, scale bar $=100 \mu \mathrm{~m}$.

Figure 14. General distribution maps of dineutine genera. (A) Dineutus; (B) Porrorhynchus); (C) Enhydrus; (D) Macrogyrus.

## Supporting information

Figure S1 Bayesian analysis of mitochondrial genes only (COI, COII, 12S). Using a codon position specific partitioning scheme. Run using the reverse jump technique described in methods section with an invariant gamma distribution and a non-clock model, 16 chains were run, swap number set to 4 , temperature set to 0.2 . Number at nodes indicates posterior probability.

Figure S2 Bayesian analysis of nuclear genes only (H3, AK). Using same settings as those described in Figure S1. Number at nodes indicates posterior probability.

Figure S3 Bayesian total-evidence calibration analysis results including Mesodineutes amurensis. Settings for analysis described in methods. Number at nodes indicate posterior probability.

Figure S4 Bayesian total-evidence calibration analysis results excluding Mesodineutes amurensis. Number at nodes indicate posterior probability.

Figure S5 Bayesian total-evidence calibration analysis results including P. (Rhomborhynchus) depressus. Settings for analysis described in methods. Numbers at nodes indicate posterior probability.

Figure S6 Maximum likelihood tree. Analysis settings outlined in methods section. Numbers at nodes indicate boot strap support.

Figure S7 Ancestral state reconstruction results using DEC model. Label at node indicates probable state.

Figure S8 Ancestral state reconstruction results using DEC model. Pie chart at node indicates probable state.

Figure S9 Ancestral state reconstruction results using DEC +j model. Label at node indicates probable state.

Figure S10 Ancestral state reconstruction results using DEC +j model. Pie chart at node shows probable ancestral states.

Figure S11 Ancestral state reconstruction results using DIVALIKE model. Label at node indicates probable state.

Figure S12 Ancestral state reconstruction results using DIVALIKE model. Pie chart at node shows probable ancestral states.

Figure S13 Ancestral state reconstruction results using DIVALIKE +j model. Label at node indicates probable state.

Figure S14 Ancestral state reconstruction results using DIVALIKE +j model. Pie chart at node shows probable ancestral states.

Appendix Description of morphological characters and coding.
Table S1 Taxa included in the phylogenetic and biogeographic analyses of the Dineutini. Gene coverage for molecular character dataset is indicated for each taxon as is geographical coding used for the biogeographic analysis.

Table S2. Character coding for morphological dataset.
Table S3. Primers used for amplification and sequencing.
Table S4 Statistical comparison of the DEC, DEC +j , DIVALIKE, and DIVALIKE +j model fit.

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## Appendix

## Morphological characters

## Head

1. Maxillary galea. (0) absent; (1) present, one segmented; (2) present, two segmented. The maxillary galea is completely absent in members of the Dineutini and Orectochilini. The Gyrinini have a maxillary galea with a single segment and the Heterogyrinae have a two segmented maxillary galea. This character is treated as ordered in the analyses.
2. Number of antennomeres in scape. (0) nine segments; (1) eight segments; (2) seven segments; (3) six segments. Nine segments are present in the scape of Heterogyrus, gyrinine species, Orectochilus, and all Macrogyrus species. Eight segments are unique to Porrorhynchus landaisi. Seven segments are present in the Enhydrus species. Six segments are present in Porrorhynchus marginatus, nearly all the Dineutus, and all Patrus and Orectogyrus species. This character is treated as ordered in the analyses.
3. Ratio of the frontolateral margin to the width of the clypeus at mid-length. (0) nearly equal or less than one; (1) frontolateral margin at least 1.5 times the longer than the medial clypeal width. The frontolateral margin appears elongate in Heterogyrus, in many orectochilines, in Porrorhynchus, some Dineutus and most Macrogyrus. A reduction of the frons length
is seen in the gyrinines and Dineutus. Several Macrogyrus species in the subgenus Cyclomimus have a greatly enlarged clypeus (Fig. 4F) but not an apparent reduction in the frontolateral margin.
4. Lateral margin of frons with a well developed bead. (0) absent; (1) present. Within the dineutines the bead is absent in Porrorhynchus and Dineutus. A strong frontal bead is present in Macrogyrus and Enhydrus (Fig. 4A, 4B, 4D, 4F). This character is also present in Heterogyrus and the gyrinines. This character cannot be scored for the orectochilines as the lateral margin of the frons is modified into the pseudofrontal ridge (Hatch, 1925).
5. Labral shape. (0) transverse; (1) elongate. A transverse labrum is very common within the Gyrinidae and in these analyses a labrum is coded as being transverse if it is less than half as long as wide. Most species of Gyrinidae have a transverse labrum. An elongate labrum is defined as being at least half as long as wide. An elongate labrum (Fig. 4E) is present in Orectochilus, Orectogyrus, Porrorhynchus, and the Dineutus subgenus Rhomborhynchus.
6. Labral basoventral setation. (0) composed of two transverse rows of setae; (1) composed of one transverse and one longitudinal paired row. This character helps separate the Dineutus subgenus Rhomborhycnhus from Porrorhynchus and the remaining Dineutus.
7. Gular suture. (0) complete, reaching anterior margin; (1) incomplete, effaced prior to anterior margin. An incomplete gular suture unites the species of Porrorhynchus.
8. Clypealium setation. (0) mostly glabrous, only a few sparse setae present basally; (1) strongly setose. The clypealium of gyrinines is mostly glabrous with only setae present basally. The dineutines and orectochilines have a strongly setose clypealium, often with a row of dense long setae medially. Heterogyrus has a setose clypealium, much more setose than the gyrinines, but not as well developed as that of the dineutines and the orectochilines.

## Prothorax

9. Pronotal transverse impressed line. (0) absent; (1) present. The pronotal transverse impressed line (Oygur and Wolfe, 1991) is absent in members of Porrorhynchus and Orectochilus. It is present in all other gyrinid species studied.
10. Prosternal process. (0) not well differentiated; (1) well differentiated and strongly elevated from the remained of the prosternum. In Macrogyrus and Enhydrus the medium portion of the prosternum is not well differentiated into a prosternal process, the posterior margin remains in nearly the same plane as the rest of the prosternum. In Dineutus and Porrorhynchus the prosternum is medially elevated and well differentiated into a distinct often bullet-shaped prosternal process. A similar prosternal process is found in the gyrinines. Heterogyrus and the orectochilines have a different sternum shape that is more cushion like, not comparable to the well differentiated prosternal process discussed previously.
11. Prosternal anteromedial sulcus. (0) absent; (1) present. There is a anteromedial sulcus present on the prosternum in Enhydrus and some Macrogyrus that is absent in Dineutus and Porrorhynchus.

Foreleg
12. Protrochanteric ventral face setation. (0) absent, completely glabrous; (1) present, a short series of short stout setae present apically. These setae are absent in the gyrinines, Heterogyrus, Orectochilus, Porrorhynchus and the Dineutus s. str. subgenus. These setae are present in Orectogyrus, in most of the Dineutus, and present in Macrogyrus only in M. (Andogyrus) seriatopunctatus. The setae are often most easily seen in females of the species.
13. Protrochanteric setose patch. (0) absent; (1) present. The protrochanteric setose patch (Fig. $7 \mathrm{E}, \mathrm{pt})$ is present in species of Porrorhynchus.
14. Protrochanteric setose brush. (0) absent; (1) present. The protrochanteric setose brush (Fig. 7A, pb) is unique to Dineutus australis.
15. Profemoral sub-apicoventral tooth/teeth. (0) absent; (1) present. These teeth are unique to the males of certain species of Dineutus (Gustafson \& Miller, 2015). They are present subapically on the ventral margin of the profemur. As many as two teeth may be present on both the anterior and posterior margins, but many species have only a single tooth present on either margin. Two teeth is a common state for many of the Dineutus s. str. subgenus and the African species of Dineutus.
16. Profemoral sub-apicoventral tooth on anterior margin. (0) absent; (1) present. This feature unites Dineutus fairmairei and D. pagdeni. It is also present in D. ritsemai suggesting this species may also be closely related to the aforementioned two.
17. Profemoral sub-apicoventral tooth on posterior margin. (0) absent; (1) present. This character is present in most of the North American Dineutus species.
18. Setigerous punctures of the anterior face of the profemur. (0) absent; (1) present. A series of setigerous punctures are present on the anterior face of the profemur medially (Fig. 8C D, sp). These punctures are absent in members of Porrorhynchus and Enhydrus but present in all other species examined.
19. Lines of setae of ventral face of profemur. (0) absent; (1) one present on posterior margin (Fig. 8C-D, psr); (2) two present on both posterior and anterior margin (Fig. $8 \mathrm{C}-\mathrm{D}$, psr , asr). Within the Gyrininae at least one line of setae is present on the posterior margin. Two are present in all Porrorhynchus and most Dineutus.
20. Setation of ventral face of profemur. (0) without setation composed of large clumps of setae becoming denser distally; (1) with setation composed of large clumps of setae become denser distally. Profemoral setation composed of two series of large clumps of setae becoming denser distally are present in species of Porrorhynchus.
21. Setose brush of posterior face of protibia. (0) present, not noticeably reduced (Fig. 8B, sb); (1) present but strongly reduced (Fig. 8A, sb); (2) absent indistinguishable from apical setae. A protibial brush is present and not reduced in Heterogyrus, Orectogyrus, Porrorhynchus, and Dineutus. It is absent in Enhydrus, the gyrinines, Orectochilus, Patrus, and some Macrogyrus. The strongly reduced state (Fig. 8A, sb) is seen most often in Macrogyrus species. This character is most variable in Macrogyrus. This character is treated as ordered in the analyses.
22. Protibia apically. (0) not laterally expanded (Fig. 9G); (1) expanded laterally (Fig. 8A - B). The protibia of Enhydrus, Macrogyrus, Porrorhynchus, Orectogyrus, and Patrus is laterally expanded. The protibia is not laterally expanded in all Dineutus, the gyrinines, Orectochilus, and Heterogyrus.
23. Adhesive setose palette of posterior face of male protarsus. (0) completely covered in adhesive setae; (1) adhesive setae reduced to half palette along outer margin. Nearly all
gyrinids have a complete setose palette, the reduced half palette condition (Fig. 9C) was only observed in three species studied. A half palette is present in D. pagdeni and D. fairmairei uniting this two species. A convergent condition is exhibited in Macrogyrus albertisii.
24. Male protarsomere I with recessed pit possessing differently sized adhesive setae. (0) absent; (1) present. The "discus" of Régimbart (1882) (Fig. 9A - C, di), is present in all species of Macrogyrus, absent in all other gyrinid species (Fig. 9 D-G).
25. Posterior face of female protarsomere $V$ with setae. (0) present in well developed furrow; (1) present but without furrow; (2) absent or reduced to a small patch. A well developed furrow is present in Porrorhynchus, and some Dineutus. The large majority of species studied have setae present without a furrow, or largely reduced to absent. This character is ordered in the analyses.

## Metaventrite I

26. Metanepisternum overall shape. (0) not lobiform; (1) lobiform. The metanepisternum of the dineutines and heterogyrines is lobiform (Fig. $6 \mathrm{~A}-\mathrm{D}$ ). The metanepisternum of the orectochilines is neither lobiform nor strongly triangular.
27. Metanepisternum overall shape. (0) not triangular; (1) triangular. The metanepisternum of the gyrinines is strongly triangular (Fig. 6F).

## Mesoventrite

28. Scutellar shield. (0) not visible when elytra closed; (1) visible when elytra closed. Among the species studied only members of Dineutus and Patrus had the scutellar shield not visible when the elytra rae closed.
29. Elytral setation. (0) absent; (1) present. The elytra has fields of setae in Heterogyrus and the orectochlines. The gyrinines and dineutines completely lack setae on the elytra.
30. Elytral serial striae. (0) none evident; (1) 11 striae evident; (2) 9 striae evident. The orectochilines exhibit no serial striae, while gyrinines have 11, and 9 striae are present in heterogyrines and dineutines.
31. Elytral strial appearance. (0) punctures; (1) well impressed lines; (2) weakly impressed lines. The elytral striae appear as punctures in the gyrinines, as well as in M. (Andogyrus) seriatopunctatus and the fossil Meiodineutes amurensis, suggesting that dineutines ancestrally possessed punctate elytral striae. Strongly impressed lines are evident in Heterogyrus and Enhydrus. Weakly impressed lines are present primarily in Dineutus and Macrogyrus. This character is treated as ordered as several gyrinine species exhibit intermediate stages between punctate to strongly impressed lines, suggesting a trend from punctures to strongly impressed lines, with weakly impressed lines as a step towards loss of impressed lines and elytral striae in general.
32. Elytral sutural border. (0) absent; (1) present. The elytra in many species of whirligigs is bordered (Brinck, 1955b), at least apically. A sutural border to the elytral is present in Heterogyrus, many orectochilines, and gyrinines. It is absent in the dineutines.
33. Elytral apex modification. (0) absent; (1) present. Unmodified elytra are attenuated toward the apex, where the apex is regularly rounded. Most gyrinid species exhibit some sort of elytral modification. Unmodified elytra are mostly found in Dineutus and Gyrinus.
34. Elytral apex with sutural production. (0) absent; (1) present. The sutural angle of the elytra is produced in many North American Dineutus (Gustafson and Miller, 2015), the majority of Macrogyrus species, and all Porrorhynchus.
35. Elytral apex with parasutural production. (0) absent; (1) present. The elytra of many dineutines has a production between the sutural and epipleural angles.
36. Elytral apex with epipleural production. (0) absent; (1) present. The epipleural angle has a production in most Macrogyrus, many orectochilines, and Porrorhynchus. This angle is not producted in Dineutus, Enhydrus, and the gyrinines.
37. Elytral apices truncate. (0) absent; (1) present. The elytral apices are truncate in the orectochilines, some Macrogyrus, and only two Dineutus.
38. Elytral apices with serration and irregularities. (0) absent; (1) present. The elytral apices may have serration and irregularities (Gustafson and Miller, 2015). This character is most commonly present in the North American and African Dineutus. It is absent in most other species studied.
39. Elytral apicolateral margins with strongly developed buzz-saw shaped serration. (0) absent; (1) present. This serration is most evident in members of Porrorhynchus. One Dineutus species also presents this serration, Dineutus micans, but it is variously developed among the different subspecies (Brinck, 1955a).
40. Elytral postscutellar pits. (0) absent; (1) present. A pair of postscutellar pits are present in the males of species of the former subgenus Rhombodineutus (Brinck, 1983).
41. Elytra with canalicuate micorsculpture. (0) absent; (1) present. Canaliculated microsculpture (Fig. 10B-C) is present in the Macrogyrus s. str. subgenus, creating a "scratch-like" appearance on the elytra under the dissecting scope. This microsculpture is present only in the Macrogyrus s. str. subgenus with some Macrogyrus species like M. sumbawae exhibiting very strong reduction.

## Mid-leg

42. Male mesotarsal claw sexual dimorphism. (0) absent; (1) present but weakly developed; (2) present strongly developed. The male mesotarsal claws of Dineutus species are strongly sexually dimorphic (Gustafson and Miller, 2015). The claws of Porrorhynchus are also sexually dimorphic but more weakly so, compared to those of Dineutus. No other gyrinid species studied have sexually dimorphic mesotarsal claws. This character is treated as ordered

## Metaventrite II

43. Lateral wings of metaventrite strap-like. (0) not strap-like in form; (1) strap-like in form. The gyrinines have a narrow and strap-like metaventral wing (Fig. 6F), a similarly formed metaventral wing is exhibited in many orectochilines.
44. Lateral wings of metaventrite triangular in form. (0) not triangular in form; (1) triangular in form. The dineutines exhibit a strongly triangular lateral wing of the metaventrite (Fig. $6 \mathrm{~A}-\mathrm{E}$ ), this character is also shared with the heterogyrines.
45. Discrimen of metaventrite with transverse suture. (0) absent; (1) present. The discrimen of the metaventrite has a transverse suture (Fig. 10A, tvs) in some Gyrinidae (Beutel and Roughley, 1988; Beutel, 1990; Miller and Bergsten, 2012). This character was thought to only be present in Spanglerogyrus and Heterogyrus however it is here also found in some members of Macrogyrus (Fig. 10A).

## Hind-leg

46. Anterior margin of lateral wings of metacoxal plate. (0) running more obliquely; (1) running more transversely. The anterior margin of the metacoxal plate is much more oblique (Hatch, 1925) in the dineutines and some Patrus species. In most other gyrinid species the anterior margin of the metacoxal plate is much more obliquely situated.
47. Posterolateral margin of metacoxal plate. (0) without border; (1) bordered. The posterolateral margin of the metacoxal plate exhibits a thick border (Fig. 6B) in species of orectochilines, gyrinines and Macrogyrus species. This border is absent (Fig. 6A, 6C, 6D - E) in Enhydrus, Dineutus, Porrorhynchus, and heterogyrines.

## Abdomen

48. Suture of abdominal sternite II. (0) absent; (1) present. Abdominal sternite II still exhibits a suture in species of Enhydrus (Hatch, 1926; Brinck, 1978), this suture is effaced in all other species studied for the analysis.
49. Overall shape of abdomen. (0) not-cylindrical, broadly rounded; (1) strongly cylindrical. The abdomen of orectochilines is strongly constricted and cylindrical in shape. All other gyrinid species have an overall rounded appearance to the abdomen.
50. Abdominal sternites VII \& VIII with linear series of setae. (0) absent; (1) present. The orectochiline in addition to the constricted cylindrical shape of the abdominal have a linear series of setae postermedially on abdominal sternites VII \& VIII for a sort of "rudder". These setae are not present in any other species studied.
51. Venter coloration. (0) darkly colored; (1) lightly colored. The venter of many species is darkly colored, dark reddish brown to black. Other species have light red to yellowish white.

## Female reproductive tract

52. Spermathecal form. (0) not elongate and sac-like; (1) greatly elongate and sac-like in form. The spermatheca of dineutines and species of Orectochilus and Orectogyrus are greatly elongate and sac-like in form (Miller and Bergsten, 2012). Those of Patrus, gyrinines, and Heterogyrus are not greatly elongate and sac-like.
53. Bursal accessory gland. (0) absent; (1) present. There is an accessory gland (Fig. 12, bg) associated with the bursa of species of Macrogyrus, Enhydrus, and Orectogyrus species (Miller and Bergsten, 2012). This accessory gland is lacking in Dineutus and Porrorhynchus, as well as in gyrinines.
54. Vaginal shield. (0) absent; (1) present. The vaginal shield (Fig. 11C, vs) was first described by Brinck (1980) in Porrorhynchus indicans, then later described again for several Dineutus species (Brinck, 1983, 1984), it is formed by anterior circular bursal sclerites (again described in Miller and Bergsten, 2012) and a posteromedial cone-like projection enclosed in more strongly sclerotized bursal cuticle. This character is present in Dineutus and Porrorhynchus and absent in all other species studied.

## Aedeagus

55. Paramere articulation with median lobe. (0) broadly; (1) narrowly. The aedeagus of Porrorhynchus species broadly articulates with the median lobe via a broad sclerotized basal region, whereas other species of dineutines the median lobe and parameres articulate via a narrow sclerotized bridge. The orectochilines and gyrinines were not coded for this character.

Sperm
56. Spermatostyle primary conjugation (Fig. 13). (0) absent; (1) present. The sperm of Orectogyrus, Orectochilus, and all dineutines is conjugated via a unique spermatostyle (Breland \& Simmons, 1970).

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Supporting information Table S1. Taxa included in the phylogenetic and biogeographic analyses of the Dineutini. Gene coverage for molecular character dataset is indicated for each taxon as is geographical coding used for the biogeographic analysis.

| Genus | Subgenus | species | voucher \# | collection data | Geo | COI | COII | 12S | H3 | AK | Efa |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Andogyrus |  | colombicus | Adco828 | VENEZUELA: Merida State $8^{\circ} 38.006^{\prime} \mathrm{N}, 71^{\circ} 09.762^{\prime} \mathrm{W}$, 2037 m Monte Zerpa area; 20.vii. 2009 leg. Short, Sites, Gustafson, Camacho; stream margin/pools VZ09-0720-01A/L-1098 ABTC-01476. MSBA | S | X | X | X | X | X | - |
| Andogyrus |  | seriatopunctatus | Adsr886 | Argentina. MSBA | S | X | X | X | X | X | - |
| Andogyrus |  | zimmermanni | Adsp648 | see Miller \& Bergsten, 2012 | S | X | X | X | X | - | - |
| Aulonogyrus | Aulonogyrus | striatus | Aost469 | see Miller \& Bergsten, 2012 | - | X | X | X | X | X | - |
| Dineutus | Callistodineutus | fairmairei | DiCf916 | Fiji03. ZSM | M | - | X | X | X | - | - |
| Dineutus | Callistodineutus | pagdeni | DiCp918 | SOLOMON ISLANDS: Guadal-canal ca. 4.5 km S of Barana vill. Forest nr. "Japanese Camp" \& Moka riv. $9^{\circ} 30.3^{\prime} \mathrm{S} 159^{\circ} 58.9^{\prime} \mathrm{E}, 27 \mathrm{~m} .5-6 . x i i .2013$ leg. Jiří Hájek. MSBA | M | X | X | X | X | X | - |
| Dineutus | Cyclinus | americanus | Diam850 | Cuba: Holguín, river/ near Biological Station of PN La Mensura-/ Piloto, $657 \mathrm{~m}, 11 . \mathrm{v} .2013,20.48640 \mathrm{~N}$ 75.79134W, A. Deler-Hernández. NMPC | I | X | X | X | X | X | - |
| Dineutus | Cyclinus | assimilis | Dias819 | USA: KS: Osage Co. pond off hwy 69 nr milemark 54. $38.031^{\circ}-94.705^{\circ} 254 \mathrm{~m} 17 . v i .2013$ leg. CK Faris \& GT Gustafson GTG06171301. MSBA | N | X | X | - | X | X | - |
| Dineutus | Cyclinus | carolinus | Dica821 | USA: TX: Hardin Co. Big Thicket Pres. $30.260^{\circ}$ $94.525^{\circ} 20 \mathrm{~m} 28 . v i .2013$ colr. CK Faris \& GT Gustafson. Mud bottomed bayou GTG06281301. MSBA | I,N | X | X | X | X | X | - |
| Dineutus | Cyclinus | ciliatus | Dici474 | see Miller \& Bergsten, 2012 | N | X | X | X | X | X | X |
| Dineutus | Cyclinus | discolor | Didi473 | see Miller \& Bergsten, 2012 | N | X | X | X | X | X | X |
| Dineutus | Cyclinus | emarginatus | Diem820 | USA: AL: Conecuh Co. nr. Evergreen. $31.291^{\circ}-86.931^{\circ}$ 82m 25.vi.2013. leg. CK Faris \& GT Gustafson. Slow $\mathrm{mud} /$ sand bottom strm GTG06251302. MSBA | N | X | X | - | X | X | - |
| Dineutus | Cyclinus | longimanus longimanus | Dilo818 | DOMINICAN REP.: Pedernales Prov. W of Pedernales on rd. to border with Haiti; roadside sweeping in dry for. \& sec. veg. 15 May $201018.154^{\circ}-71.7582^{\circ}$ colr. G. J. Svenson. MSBA | I | X | X | X | X | X | - |
| Dineutus | Cyclinus | mexicanus | Dimx822 | MÉXICO: Michoacán: Coalcoma, La Nuez, cañada el colorin. 15.ix.2003. col. R Novelo. MSBA | C | X | X | X | X | X | - |
| Dineutus | Cyclinus | productus | Dipd844 | USA: KS: Elk Co. Elk River S of Longton $37.36960^{\circ}$ 96.07873 265m 22.vi. 2014 leg. C.Maier, C.Faris, S.Baca, G.Gustafson Muddy riv. Nr falls. GTG062214A. MSBA | C,N | X | X | X | - | X | - |
| Dineutus | Cyclinus | robertsi | Diro913 | USA: Georgia: Warwoman Wld Mgmt Area. Tuckaluge Cr. $34.90155^{\circ} \mathrm{N} 83.30015^{\circ} \mathrm{W} .533 \mathrm{~m}$. 11July2012. KB Miller colr. KBM11071201. MSBA | N | X | X | X | X | X | - |
| Dineutus | Cyclinus | serrulatus analis | Disa843 | USA: KS: Bourbon Co. Marmaton River, NE Ft. Scott $37.866^{\circ}$, $-94.675^{\circ} 237 \mathrm{~m} 21$.vi. 2014 leg. C.Maier, <br> C.Faris, S.Baca, G.Gustafson GTG062114A. MSBA | N | X | X | X | - | X | - |
| Dineutus | Cyclinus | shorti | Dish895 | USA: AL: Covington Co. Conecuh Nat. For., trib. Of YellowRiv. $31.0929^{\circ},-86.5183^{\circ} 43 \mathrm{~m} 27 . i v .2015$, leg. G.Gustafson \& S.M.Baca GGSB042715C1 | N | X | - | X | X | - | - |
| Dineutus | Cyclinus | solitarius | Diso605 | see Miller \& Bergsten, 2012 | C | X | X | X | X | X | X |
| Dineutus | Cyclinus | $\begin{aligned} & \text { sublinea- } \\ & \text { tus } \\ & \hline \end{aligned}$ | Disu505 | see Miller \& Bergsten, 2012 | C | X | X | X | X | X | X |
| Dineutus | Cyclinus | truncatus | Ditr726 | Panama: Ngäbe-Buglé Comarca, small stream nr Soloy. $8^{\circ} 36.554^{\prime} \mathrm{N}, 82^{\circ} 07.814^{\prime} \mathrm{W}$. 06July2009. Leg K.B. Miller KBM07060903 | C | X | X | X | X | X | - |
| Dineutus | Cyclous | australis | Diau911 | AUSTRALIA: QLD. $19^{\circ} 41.153^{\prime} \mathrm{S} 145^{\circ} 49.536^{\prime} \mathrm{E}$ <br> 14.Mar.2011. colr. KB Miller KBM14031102. MSBA | $\begin{aligned} & \mathrm{P}, \mathrm{O}, \\ & \mathrm{~W}, \\ & \mathrm{M}, \mathrm{~A} \end{aligned}$ | X | X | X | X | - | - |
| Dineutus | Dineutus | fulgidus | DiDf915 | INDONESIA: Sumatra Barat, Solok, Alahan Panjank Road. $1190 \mathrm{~m} .0^{\circ} 56.345$ 'S $100^{\circ} 46.411^{\prime} \mathrm{E}$. ZSM | O | X | X | X | X | X | - |


| Dineutus | Dineutus | n sp | DiDn865 | BALI: Telaga Forest, BLI07. ZSM | O | X | X | X | X | X | - |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Dineutus | Merodineutus | macrochirus | DiMm919 | PAPUA NEW GUINEA: Central Prov. Kokoda Trek. $9^{\circ} 00.338^{\prime} \mathrm{S} 147^{\circ} 44.252^{\prime} \mathrm{E} .1390 \mathrm{~m} . \mathrm{i} .2008$. leg. Posman (PNG 173). ZSM | M | - | X | X | X | - | - |
| Dineutus | Merodineutus | priscus | DiMp917 | PAPUA NEW GUINEA: S Highlands Prov. Sopulkul, $30-35 \mathrm{~km}$ NE Mendi. $6^{\circ} 2.944^{\prime} \mathrm{S} 143^{\circ} 46.485^{\prime} \mathrm{E} .2679 \mathrm{~m}$. 16.vi.2006. Leg. John ex swamp into stream (PNG 79). ZSM | M | - | X | X | X | X | - |
| Dineutus | Protodineutus | aereus | Disp481 | see Miller \& Bergsten, 2012 | E | X | X | X | X | X | - |
| Dineutus | Protodineutus | grossus abyssinicus | Digr855 | CONGO (DRC): Bas Congo Prov. Nduizi River. $5^{\circ} 44.446^{\prime} \mathrm{S} 13^{\circ} 44.385^{\prime} \mathrm{E} 266 \mathrm{~m} .6 \mathrm{Aug}$ 2012. coll: Sites, Shephard, Pwema L-1439. MSBA | E | X | X | X | X | X | - |
| Dineutus | Protodineutus | grossus angolensis | Diga826 | TANZANIA: Kagera Reg. ca. 5 km E of Bhiaramulo $2^{\circ} 38.159^{\prime} \mathrm{S} 31^{\circ} 20.789^{\prime} \mathrm{E} 1427 \mathrm{~m}$. 26Jul2010. coll: R.W Sites \& A. Mbogho L-1154. MSBA | E | X | X | X | X | X | - |
| Dineutus | Protodineutus | indicus | Diin482 | see Miller \& Bergsten, 2012 | O | X | X | X | X | X | X |
| Dineutus | Protodineutus | micans | Disp577 | see Miller \& Bergsten, 2012 | E | X | X | X | X | X | X |
| Dineutus | Protodineutus | proximus | Dipx515 | see Miller \& Bergsten, 2012 | E | X | X | X | - | X | X |
| Dineutus | Protodineutus | sinuosipe nnis | Disn516 | see Miller \& Bergsten, 2012 | E | X | X | X | X | - | X |
| Dineutus | Protodineutus | staudin- <br> geri | Dist825 | TANZANIA: Arusha Reg. Sanaware River at Arusha $3^{\circ} 21.549^{\prime} \mathrm{S} 36^{\circ} 41.619^{\prime} \mathrm{E} 1442 \mathrm{~m} .11 \mathrm{Aug} 2010$. coll. R.W. Sites \& A. Mbogho L-1239. MSBA | E | X | X | X | X | X | - |
| Dineutus | Rhombodineutus | pectoralis | Disp507 | see Miller \& Bergsten, 2012 | M | X | X | X | X | X | - |
| Dineutus | Rhombodineutus | tetracanthus | DiRt908 | PAPUA NEW GUINEA: Madang Prov. Wannang, $5^{\circ} 15.458^{\prime} \mathrm{S} 145^{\circ} 2.389^{\prime} \mathrm{E} 270 \mathrm{~m} .31 . x .2008$. leg. Posman (PNG187) ZSM. | M | X | X | X | X | X | - |
| Dineutus | Spinosodineutes | fauveli | Difa485 | see Miller \& Bergsten, 2012 | E | X | X | X | X | X | - |
| Dineutus | Spinosodineutes | striatus | Disp576 | see Miller \& Bergsten, 2012 | E | X | X | X | X | X | X |
| Dineutus | Spinosodineutes | $\begin{aligned} & \text { subspino- } \\ & \text { sus } \end{aligned}$ | Disu484 | see Miller \& Bergsten, 2012 | E | X | X | X | X | X | X |
| Enhydrus |  | atratus | Ehas646 | see Miller \& Bergsten, 2012 | C, S | X | X | X | X | - | - |
| Enhydrus |  | sulcatus | Ehsu856 | BRAZIL: Rio de Janeiro: Cachoei Ras de Macacu Regua. 01.vi.2013. Ponto do Ganesh (Parte Alta) col. Equipe Coleoptera. MSBA | S | X | X | X | X | X | - |
| Macrogyrus | Australogyrus | oblongus | AyAo857 | AUSTRALIA: QLD. Broken River. $21^{\circ} 10.077$ 'S $148^{\circ}$ 30.386'E. 713m. 17Mar2011. Coll: KB Miller. KBM1703110D. MSBA | A | X | X | X | X | X | - |
| Macrogyrus | Australogyrus | rivularis | AyAr909 | AUSTRALIA: VIC. Off Old Coast Rd, -37.73667 , $148.95166 .30 \mathrm{~m} .12 . \mathrm{i} .2015$ leg. G.Gustafson, K.Miller. Small strm in forest. GTG01122015A. MSBA | A | X | X | X | - | X | - |
| Macrogyrus | Clarkogyrus | reichei | Ayre912 | AUSTRALIA: VIC. Glenelg River nr. Dergholm, 37.36686, 141.2428. 75 m . 13.i.2015. Leg G.Gustafson. MSBA | A | X | X | X | - | - | - |
| Macrogyrus | Cyclomimus | purpura- <br> scens | AyCs841 | PAPUA NEW GUINEA: Morobe Prov. Pindiu. <br> $6^{\circ} 27.147^{\prime} \mathrm{S} 147^{\circ} 29.574^{\prime} \mathrm{E}, 1470 \mathrm{~m} .12 . \mathrm{x} .2009$ leg. Inaho (PNG206). ZSM | M | X | X | X | X | X | - |
| Macrogyrus | Cyclomimus | sp | AyCs831 | PAPUA NEW GUINEA: Sandaun Prov. Mianmin $4^{\circ} 54.570^{\prime} \mathrm{S} 141^{\circ} 35.490^{\prime} \mathrm{E} 990 \mathrm{~m} .23 . x .2008$. leg. Ibalim (PNG193). ZSM | M | X | X | X | X | - | - |
| Macrogyrus | Cyclomimus | toxopeusi | AyCs829 | INDONESIA: Papua: Poga $3^{\circ} 48.382^{\prime} \mathrm{S}, 138^{\circ} 34.780^{\prime} \mathrm{E}$ 2285-2330 m. ZSM | M | X | X | X | X | X | - |
| Macrogyrus | Macrogyrus | howittii | Ayhw887 | AUSTRALIA: Tasmania. Franklin Beach, Lake St. Clair. 10.i. 2015 CHS Watts. MSBA | A | X | - | X | X | X | - |
| Macrogyrus | Megalogyrus | striolatus | Ayst882 | AUSTRALIA: NSW. Megalong Valley, -36.65629, 150.27377, 861 m. 04.i.2015. leg. G.Gustafson Forested strm GTG01042015A. MSBA | A | X | X | X | X | X | - |
| Macrogyrus | Orectomimus | paradoxus | Aypa817 | AUSTRALIA: QLD. Cardstone. $17^{\circ} 46.650^{\prime} \mathrm{S}$ 145³9.015'E. 19Mar2011 KB Miller colr. KBM19031102. MSBA | A | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | albertisi | AyTs832 | Papua New Guinea: Sanduan. Toricelli Mts. 2h walk fr Sibilanga Stm. $350 \mathrm{~m}, 19.1 \mathrm{iv} .2006,03^{\circ} 39.121^{\prime} \mathrm{S}$ $142^{\circ} 29.991^{\prime}$ E. Balke (PNG 44). ZSM | - | X | - | - | X | X | - |
| Macrogyrus | Tribologyrus | australis | Ayau910 | AUSTRALIA: Frog's Hollow Creek off Princes HWY, 36.74363 , $149.81474 .35 \mathrm{~m} .07 . \mathrm{i} .2015$. leg. G.Gustafson. Small strm in open grassland. GTG01072015B. MSBA | A | X | X | X | X | - | - |
| Macrogyrus | Tribologyrus | gouldi | Aygo501 | see Miller \& Bergsten, 2012 | A | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | sexangu- <br> laris | AyTs835 | PAPUA NEW GUINEA: Herowane. ZSM | M | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | sp | Aysp861 | INDONESIA: Timor: Mt. Mutis, creeks and streams. $1580 \mathrm{~m} .9^{\circ} 38.124^{\prime} \mathrm{S} 124^{\circ} 12.800^{\prime} \mathrm{E}$ (TIM04). ZSM | O | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | sp | Aysp863 | INDONESIA: Lombok: Senaru area. $834 \mathrm{~m} .8^{\circ} 19.326^{\prime} \mathrm{S}$ $116^{\circ} 24.989^{\prime} \mathrm{E}$ (LOM004). ZSM | O | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | sp | AyTs833 | PAPUA NEW GUINEA: Central Prov. Woitape. $08^{\circ} 31.290^{\prime} \mathrm{S} 147^{\circ} 13.684^{\prime} \mathrm{E} 1700 \mathrm{~m} . \mathrm{i} .2008$ leg. Posman (PNG166). ZSM | M | X | X | X | X | X | - |
| Macrogyrus | Tribologyrus | sp nr . <br> blanchar- <br> dii | AyTs830 | PAPUA NEW GUINEA: Bougainville Isl.: Kieta $06^{\circ} 12.955^{\prime} \mathrm{S} 155^{\circ} 29.755^{\prime} \mathrm{E} 520 \mathrm{~m} .12$.vi.2008. leg. Posman (PNG180). ZSM | M | X | X | X | X | - | - |


| Macrogyrus | Tribologyrus | sp nr. <br> blanchar- <br> dii | AyTs834 | PAPUA NEW GUINEA: E Highlands Prov. Onerunka, small creek, redsoil rock $6^{\circ} 20.936^{\prime} \mathrm{S}$ 145 ${ }^{\circ} 46.874^{\prime} \mathrm{E} 1700$ m. 21.v. 2006 leg. John \& Balke (PNG71). ZSM | M | X | X | X | X | X | - |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Macrogyrus | Tribologyrus | sumbawae | Aysp860 | INDONESIA: Sumba: dry forest stream in limestone. $370 \mathrm{~m} 9^{\circ} 49.474^{\prime} \mathrm{S} 120^{\circ} 20.856^{\prime} \mathrm{E}$ (SUA08). ZSM | O | X | X | X | X | X | - |
| Mesodineutes |  | amurensis | Meamure | Arkhara, Amur Oblast, Russia $\left(49.4^{\circ} \mathrm{N}, 130.1^{\circ} \mathrm{E}\right.$ : paleocoordinates $52.1^{\circ} \mathrm{N}, 118.2^{\circ} \mathrm{E}$ ). Darmakan formation, Danian (66.0-51.7 mya) Paleocene. PIN | - | - | - | - | - | - | - |
| Porrorhynchus | Porrorhynchus | landaisi | Prla852 | CHINA: Hainan Isl. Jianfengling Mts. Tiachi Lake rd. from Taichi village to 'Sector $5^{\prime} 18^{\circ} 43.6-44.1^{\prime} \mathrm{N} 820-$ $950 \mathrm{~m}, 108^{\circ} 52.1-52.5^{\prime} \mathrm{E}$ 10.v.2011. M.Fikáček \& Sh. Zhao lgt. Small slow-flowing stony river in the primary forest. MSBA | O | X | X | X | X | X | - |
| Porrorhynchus | Porrorhynchus | marginatu <br> s | Prma779 | THAILAND: Kanchanaburi Prov. Huay Ou Long at Ban Ou Long. N14 ${ }^{\circ} 35.177^{\prime}$ W98우․921' 1 Jun2011. 274m col. R.W. Sites. Gravel bottom stream. MSBA | O | X | X | X | - | X | - |
| Porrorhynchus | Rhomborhynchus | misoolensis | Prdepre | INDONESIA: West Papua: Misool Island: Tama River, SE of old Fakal village site, $1^{\circ} 51^{\prime} 38.1^{\prime \prime} \mathrm{S} 129^{\circ} 55^{\prime} 24.1^{\prime \prime} \mathrm{E}$, $60 \mathrm{~m}, 22 . i v .1999$. leg. D.A. Polhemus, CL7110. DAPC. | - | - | - | - | - | - | - |
| Gyretes |  | giganteus | Gegigan | Auvergne, France $\left(46.1^{\circ} \mathrm{N}, 2.9^{\circ} \mathrm{E}\right.$ : paleocoordinates $40.6^{\circ} \mathrm{N}, 0.6^{\circ} \mathrm{E}$ ). Menat formation, Thanetian (58.7-55.8 mya), Paleocene. MNHN. | - | - | - | - | - | - | - |
| Gyrinus | Gyrinus | sericeo- <br> limbatus | Gysp840 | PAPUA NEW GUINEA: Morobe Prov. Pindiu. $6^{\circ} 27.147^{\prime} \mathrm{S} 147^{\circ} 29.574^{\prime} \mathrm{E}, 1470 \mathrm{~m} .12 \times \mathrm{x} .2009$ leg. Inaho (PNG206). ZSM | - | X | X | X | X | X | - |
| Gyrinus |  | maculiventris | Gysp837 | USA: New Mexico, Cibola Co., Zuni Mountains. 18.v.2013. leg. S. Baca. 180513-A. MSBA | - | X | X | X |  | X | - |
| Gyrinus | Oreogyrinus | dimorph- <br> us | Gysp839 | USA: New Mexico, Cibola Co., Zuni Mtns, $35.40178^{\circ} \mathrm{N}$, $108.44956^{\circ}$ W. 30.v.2013. leg. G.Gustafson \& S.Baca. GGSB053013C. MSBA | - | X | X | X | X | X | - |
| Mesogyrus |  | antiquus | Msantiq | Karatau-Mikhailovka, Chimkent, Kazakhstan ( $42.9^{\circ} \mathrm{N}$, $70.0^{\circ} \mathrm{E}$ : paleocoordinates $41.6^{\circ} \mathrm{N}, 73.7^{\circ} \mathrm{E}$ ), lacustrine, siltstone in the Karabastau Formation, <br> Callovian/Oxfordian (157.3-166.1 Ma), Upper Jurassic. PIN | - | - | - | - | - | - | - |
| Orectochilus |  | villosus | Orvi527 | see Miller \& Bergsten, 2012 | - | X | X | X | X | - | - |
| Orectogyrus | Orectogyrus | madagascariensis | Ogmd601 | see Miller \& Bergsten, 2012 | - | X | X | X | X | X | - |
| Orectogyrus |  | sp | Ogsp564 | see Miller \& Bergsten, 2012 | - | X | X | X | X | X | - |
| Patrus |  | productus | Orpr487 | see Miller \& Bergsten, 2012 | - | X | X | X | X | X | - |

Supporting information Table S2. Character coding for morphological dataset.

|  | 1 | 6 | 11 | 16 | 21 | 26 | 31 | 36 | 41 | 46 | 156 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | \| | \| |  | \| | \| | \| |  | \| |  |  | \| | |
| Hsmi596 | 20110 | 0011- | -0000 | 00110 | 0000? | 10112 | 11100 | 01000 | 00011 | 00000 | 1010-0 |
| Orvi527 | 001-1 | -0101 | 12000 | 00010 | 20002 | 01110 | -1100 | 01000 | 00100 | 01011 | 1110-1 |
| Orpr487 | 031-0 | -0100 | 02000 | 00010 | 21002 | 01010 | -1100 | 11000 | 00100 | 01011 | 0010-? |
| Ogmd601 | 031-1 | 10100 | 01000 | 00120 | 01001 | 01110 | -1100 | 11000 | 00100 | 01011 | 01?0-1 |
| Ogsp564 | 031-1 | 10100 | 01000 | 00120 | 01001 | 01110 | -1100 | 11000 | 00100 | 01011 | 11?0-1 |
| Gegigan | ????0 | ????? | ?? ? ? ? | ????? | ????? | ????? | ? ? 100 | 01000 | ????? | ???1? | ?????? |
| Gysp837 | 10010 | 00011 | 10000 | 00120 | 20002 | 01101 | 01000 | 00000 | 00100 | 01000 | 0000-0 |
| Gysp839 | 10000 | 00011 | 10000 | 00120 | 20002 | 01101 | 01000 | 00000 | 00100 | 01000 | 0000-0 |
| Gysp840 | 11010 | 00011 | 10000 | 00120 | 20002 | 01101 | 01000 | 00000 | 00100 | 01000 | 0000-0 |
| Aost469 | 11000 | 00010 | 00000 | 00120 | 10002 | 01101 | 01000 | 00000 | 00100 | 01000 | 0000-0 |
| Diau911 | 03000 | 00111 | 01010 | 00120 | 00001 | 10002 | 20100 | 10100 | 01010 | 10000 | 010111 |
| Ehas646 | 03100 | 00110 | 10000 | 00010 | 11002 | 10102 | 10000 | 01000 | 00010 | 10100 | 011011 |
| Ehsu856 | 02100 | 00110 | 10000 | 00010 | 11002 | 10102 | 10000 | 00000 | 00010 | 10100 | 011011 |
| Ayhw887 | 00110 | 00110 | 02000 | 00120 | 11012 | 10102 | 20100 | 01000 | 10011 | 11000 | 011011 |
| AyAo857 | 00100 | 00110 | 00000 | 00110 | 01010 | $1 ? 102$ | 20111 | 10000 | 10011 | 11000 | 011011 |
| AyAr909 | 00110 | 00110 | 00000 | 00110 | 01010 | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| Ayre912 | 00100 | 00110 | 10000 | 00010 | 01011 | 10102 | 20100 | 10100 | 10011 | 11000 | 011011 |
| Ayst882 | 00110 | 00110 | 00000 | 00110 | 11012 | 10102 | 20100 | 01000 | 10011 | 11000 | 011011 |
| Adsr886 | 00110 | 00110 | 11000 | 00120 | 21012 | 10102 | 00100 | 01000 | 00011 | 11000 | 011011 |
| Adco 828 | 00110 | 00110 | 10000 | 00010 | 11012 | 10102 | 20000 | 00000 | 00010 | 11000 | 011011 |
| Adsp648 | 00110 | 00110 | 10000 | 00010 | 11012 | 10102 | 20000 | 00000 | 00010 | 11000 | 011011 |
| AyCs 829 | 00010 | 00110 | 00000 | 00120 | 11012 | 10102 | 20111 | 10000 | 00010 | 11000 | 111011 |
| AyCs841 | 00010 | 00110 | 00000 | 00120 | 1101? | 10102 | 20111 | 10000 | 00010 | 11000 | 111011 |
| AyTs835 | 00110 | 00110 | 00000 | 00110 | 1101? | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| AyTs830 | 00110 | 00110 | 00000 | 00110 | 1101? | 10102 | 20100 | 11000 | 10011 | 11000 | 011011 |
| Aysp863 | 00110 | 00110 | 02000 | 00120 | 11012 | 10102 | 20110 | 11000 | 00011 | 11000 | 011011 |
| AyTs832 | 00110 | 0011- | 00000 | 00120 | 11110 | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| AyTs834 | 00110 | 00110 | 00000 | 00120 | 1101? | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |


| Ayau910 | 00110 | 00110 | 00000 | 00120 | 11012 | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| AyTs833 | 00110 | 00110 | 00000 | 00110 | 1101? | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| Aysp861 | 00110 | 00110 | 00000 | 00120 | 1101? | 10102 | 20111 | 10000 | 10011 | 11000 | 011011 |
| Aysp860 | 00110 | 00110 | 00000 | 00110 | 1101? | 10102 | 20111 | 10000 | 00011 | 11000 | 011011 |
| AyCs 831 | 00010 | 00110 | 00000 | 00120 | 11012 | 10102 | 20111 | 10000 | 00010 | 11000 | 111011 |
| Aygo501 | 00110 | 00110 | 00000 | 00110 | 11011 | 10102 | 21111 | 10000 | 10011 | 11000 | 111011 |
| Aypa817 | 00110 | 00110 | 10000 | 00110 | 1101? | 10102 | 20110 | 10000 | 10011 | 11000 | 111011 |
| DiDf915 | 03000 | 00111 | 00001 | 11120 | 00000 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| DiDn865 | 03000 | 00111 | 00001 | 11120 | 00000 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Difa485 | 03000 | 00110 | 01000 | 00120 | 00001 | 10002 | 20111 | 10100 | 01010 | 10000 | 110111 |
| Disp507 | 03100 | 00111 | 00000 | 00110 | 00000 | 10002 | 20100 | 10001 | 01010 | 10000 | 110111 |
| DiRt908 | 02100 | 00111 | 00000 | 00120 | 00001 | 10002 | 20101 | 10001 | 01010 | 10000 | 010111 |
| Disu484 | 03000 | 00111 | 01000 | 00120 | 00001 | 10002 | 20100 | 10100 | 01010 | 10000 | 010111 |
| Dipx515 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Dilo818 | 03000 | 00111 | 01001 | 01110 | 00002 | 10002 | 20111 | 00100 | 01010 | 10000 | 110111 |
| Ditr726 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20100 | 01100 | 01010 | 10000 | 010111 |
| Dimx822 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20100 | 01100 | 01010 | 10000 | 010111 |
| Disa843 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20110 | 00100 | 01010 | 10000 | 110111 |
| Disp576 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20110 | 10100 | 01010 | 10000 | 110111 |
| DiCp918 | 03000 | 00111 | 00001 | 10110 | 00101 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| Disn516 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20111 | 00000 | 01010 | 10000 | 010111 |
| Disp481 | 03000 | 00111 | 01001 | 11120 | 00002 | 10002 | 20000 | 00100 | 01010 | 10000 | 010111 |
| Diam850 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20110 | 00100 | 01010 | 10000 | 010111 |
| Dica821 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20000 | 00100 | 01010 | 10000 | 010111 |
| Disu505 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Diso605 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Dias819 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20110 | 00000 | 01010 | 10000 | 010111 |
| Diem820 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Dipd844 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20110 | 00100 | 01010 | 10000 | 010111 |
| Diin482 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Dici474 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Diro913 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| Didi473 | 03000 | 00111 | 01001 | 01120 | 00002 | 10002 | 20110 | 00000 | 01010 | 10000 | 110111 |
| Dish895 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| Disp577 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20100 | 00110 | 01010 | 10000 | 010111 |
| Digr855 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Diga826 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Dist825 | 03000 | 00111 | 01000 | 00120 | 00002 | 10002 | 20000 | 00000 | 01010 | 10000 | 010111 |
| Prla852 | 01101 | 01101 | 00100 | 00021 | 01000 | 10000 | -0111 | 10110 | 01010 | 10000 | 110101 |
| Prma779 | 03101 | 01101 | 00100 | 00021 | 01000 | 10000 | -0111 | 10110 | 01010 | 10000 | 110101 |
| DiCf916 | 03000 | 00111 | 00000 | 00110 | 00102 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| DiMp917 | 03000 | 00111 | 00000 | 00110 | 00000 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| DiMm919 | 02000 | 00111 | 00000 | 00110 | 00000 | 10002 | 20000 | 00000 | 01010 | 10000 | 110111 |
| Meamure | ? ? ? ? | ?? ? ? | ?? ? ? ? | ?? ? ? | ? ? ? ? ? | 10? 02 | 00000 | 00000 | 0? 01? | 10000 | ?????? |
| Prdepre | 03101 | 10111 | 01000 | 00120 | 10002 | 10002 | 20111 | 10000 | 01010 | 10000 | 110111 |

Supporting information Table S3. Primers used for amplification and sequencing.

| Gene | Forward primer (5'-3') | Reverse primer (5'-3') | Reference |
| :---: | :---: | :---: | :---: |
| COI | Jerry, CAACATTTATTTTGATTTTTTGG | Pat, TCCAATGCACTAATCTGCCATATTA | Simon et al. |
| 1994 |  |  |  |

For difficult taxa internal primers were designed for AK183F and AK939R, these are DinAKF1 and DinAKR1. Nested PCR was performed using these primers to amplify Arginine kinase in difficult specimens.

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Supporting information Table S4. Statistical comparison of the DEC, DEC +j, DIVALIKE, and DIVALIKE +j model fit.

|  | LnL | param \# | d | e | j | AIC | AIC_wt |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| DEC | -99.9 | 2 | 0.012 | 0.16 | 0 | 203.8 | 0.25 |
| DEC+ J | -97.78 | 3 | 0.0056 | 0.065 | 0.052 | 201.6 | 0.75 |
| DIVALIKE | -103.7 | 2 | 0.0029 | 0.0007 | 0.061 | 213.5 | 0.0019 |
| DIVALIKE +J | -103.8 | 3 | 0.003 | 0.0011 | 0.064 | 213.7 | 0.0018 |
















Orpr 487
Oamd601












BioGeoBEARS DIVALIKE+J on Dineutine M0_unconstrained
ancstates: global optim, 5 areas max. $\mathrm{d}=0.003 ; \mathrm{e}=0.0011 ; \mathrm{j}=0.0636 ; \operatorname{LnL}=-103.84$



## Relicts of early Mesozoic whirligig beetle clades persist in biodiversity hotspots

Grey T. Gustafson ${ }^{1}$, Alexander A. Prokin ${ }^{2}$, Rasa Bukontaite, Kelly B. Miller ${ }^{1}$, Johannes Bergsten ${ }^{3}$
${ }^{1}$ Department of Biology and Museum of Southwestern Biology, University of New Mexico, Albuquerque, NM 87131, USA. ${ }^{2}$ Papanin Institute for Biology of Inland Waters, Russian Academy of Sciences, Borok, Nekouzskii District, Yaroslavl Region, 152742, Russia. ${ }^{3}$ Department of Zoology, Swedish Museum of Natural History, Box 50007, SE-104 05 Stockholm, Sweden.

Madagascar's distinctive fauna has long fascinated biologists; its unique geologic history including an extensive period of isolation following the break up Gondwana ${ }^{1}$, has been proposed to account for its unique biodiversity ${ }^{2}$. However, most of Madagascar's extant fauna is found to have originated recently during the Cenozoic ${ }^{\mathbf{2 - 4}}$. Whirligig beetles (Coleoptera, Gyrinidae) are a common aquatic insect group, with a global distribution ${ }^{5}$, and numerous endemic Malagasy taxa ${ }^{6,7}$. Here we present the first time-calibrated phylogenetic analysis of the family using novel dating techniques and show a Late Triassic origin, significantly older than the most recent estimate of Middle Jurassic ${ }^{\mathbf{8}}$. The Malagasy striped whirligig beetle, Heterogyrus milloti, is found to be the sole surviving lineage of the Heterogyrinae, the dominant gyrinid fauna of the Mesozoic era. The origin of the Heterogyrinae is dated to a median age of 196 million years ago making Heterogyrus the oldest endemic animal lineage known from Madagascar. The other currently monotypic subfamily, Spanglerogyrinae, has its origin in the Late Triassic. Spanglerogyrus albiventris is known only from the North American Coastal Plain, another global biodiversity hotspot ${ }^{9}$. Our findings show a once widespread Mesozoic taxon isolated on Madagascar, which served as a refugium for the last surviving heterogyrine species. The monotypic relictual gyrinid lineages dated to the Early Jurassic provide further evidence of the

## persistence of coleopteran lineages and new precedence for invertebrate specific conservation in biodiversity hotspots.

The fauna of Madagascar is among the most unique and threatened biodiversity on the planet ${ }^{10}$. The origin of the island's fauna is of great interest given its central position in Gondwana and extensive isolation following separation from India ${ }^{1,2}$. Few studies, however, have revealed taxa with truly Gondwanan relationships or ancient origins ${ }^{2}$. The most iconic fauna, the lemurs, tenrecs, and unique carnivorans rafted to the island ${ }^{11}$ from Africa within the past sixty million years ${ }^{3,4,12}$. A primarily Cenozoic origin is similarly shared among most of the island's other extant endemic vertebrate fauna ${ }^{3,4}$. Few notable exceptions include oplurine iquanids, podocnemidid turtles, mantellid frogs, and cichlid fish, which show Mesozoic origins ${ }^{3,4}$. Invertebrate groups similarly show mostly Cenozoic origins, but among the dated analyses, also contain some of the oldest endemic lineages currently known (Extended Data Tab. 1).

The whirligig beetle fauna of Madagascar is highly endemic, similar to other aquatic invertebrates ${ }^{13}$, with 25 of the 26 known species found nowhere else ${ }^{6,7}$. Notable among the Malagasy fauna is Heterogyrus milloti, the Malagasy striped whirligig beetle, for its intriguing morphology uniting it with several groups of Gyrinidae ${ }^{7}$. Molecular data have only recently been utilized for phylogenetic analysis of the Gyrinidae ${ }^{5}$. This previous study placed the monotypic Heterogyrus in its own subfamily, as sister to the Gyrininae, containing the vast majority of species, but without strong support ${ }^{5}$. The family has also never received a time-calibrated phylogenetic reconstruction, preventing timing of diversification from being known.

Phylogenetic analyses employing novel Bayesian total-evidence fossil calibration ${ }^{14}$ were performed on nine fossil and 128 extant gyrinid taxa. Total-evidence calibration constructs a
time-calibrated tree, while simultaneously placing fossil taxa within the phylogeny during analysis ${ }^{14}$. This calibration technique is desirable to node-based methods as it incorporates inherent uncertainty regarding fossil taxa's relationship to extant diversity ${ }^{14}$. Previous iterations of these analyses implemented in MrBayes 3.2.5 were found to give suspiciously old ages for clades lacking fossils ${ }^{15}$. For this reason a novel Fossilization-Birth-Death (FBD) model ${ }^{16}$ was introduced in MrBayes 3.2.6 to improve dating accuracy ${ }^{15}$.

Origins of the family Gyrinidae are dated to the Late Triassic (95\% highest probability density (hpd) $203-259 \mathrm{Ma}$, median $95 \%$ hpd (m) 228 Ma ). This age is significantly older than the most recent estimate for the family ${ }^{8}$, and while closer to prior estimates ${ }^{17,18}$ is still older. The earliest diverging lineage, Spanglerogyrinae, is found to have originated in the Early Jurassic (hpd $175-200 \mathrm{Ma}, \mathrm{m} 185 \mathrm{Ma}$ ) and the fossil genus Angarogyrus is now a member of this subfamily (Fig. 1), rendering it no longer monotypic. Reexamination of the Angarogyrus fossils reveals remarkable similarity to the extant genus Spanglerogyrus (Fig. 3). Both have a unique quadrate frons, with the frontolateral margins continued posteriorly over the dorsal eye (Fig. 3a', $c^{\prime}, d$ ), a pronotum with a strong medial lobe projecting anteriorly onto the head capsule (Fig. 3a', c', d), and a very small in body size (Fig 3a,b,c). An exceptionally well-preserved elytron of Angarogyrus minimus shows similar coloration to the extant Spanglerogyrus albiventris and covering of setae (Fig. 3e,f). Spanglerogyrus is clearly an exceptionally relictual taxon, exhibiting little morphological change since the Early Jurassic.

The analyses unequivocally support placement of Heterogyrus as sister to the Gyrininae (Fig. 1). Several morphological characters shared with Spanglerogyrus (Extended Data Fig. 1) strongly support this distinction, including the medially divided labrum (Extended Data Fig. 1a, b), retention of the medial spur of the protibia, (Extended Data Fig. 1c, d), and the pronotum with
a well-developed median lobe (Fig. 3d, 4e). The subfamily Heterogyrinae is no longer monotypic. The extinct genera Baissogyrus (Fig. 4c, c'), Cretotortor (Fig 4a, a'), and Mesogyrus are placed within the Heterogyrinae with strong support (Fig. 1). The origin of the Heterogyrinae is placed in the Mid- to Early Jurassic (hpd $157-188 \mathrm{Ma}, \mathrm{m} 170 \mathrm{Ma}$ ). The fossil record shows the heterogyrine genera were widely distributed and found throughout the Mesozoic (Fig. 2), clearly forming the dominant gyrinid fauna. Morphology of the exceptionally preserved fossil Mesogyrus antiquus (Fig. 4g, g') from the well-known Karatau deposits ${ }^{19}$, compared to that of the extant Malagasy Heterogyus milloti (Fig. 4b, d, e, f), shows morphologically heterogyrines have changed very little throughout their evolutionary history. The Karatau deposits are the remains of a large, stable, freshwater, Jurassic lake ${ }^{20}$, indicating M. antiquus was lentic, compared to the extant $H$. milloti known only from small trickling mountain streams ${ }^{5}$, suggesting increased ecological diversity in the past.

No insect fossils have been described from Madagascar, limiting our inference of Mesozoic Malagasy fauna to the well studied fossil vertebrates ${ }^{21}$. The fossil vertebrate fauna at the end Cretaceous of Madagascar is composed of widespread Gondwanan lineages, such as sauropod, non-avian therapod dinosaurs, crocodyliforms, anurans, and snakes ${ }^{22}$. These widespread lineages likely became isolated on Madagascar following Gondwana's breakup, given many were not good dispersers (i.e. therapod and sauropod dinosaurs) and the unlikelihood of a land bridge ${ }^{22}$. The fossil evidence (Fig. 2) shows the Heterogyrinae were widespread across Laurasia, but given both the age of the estimated most recent common ancestor (Fig. 1) and the only extant member being found in Madagascar, were probably similarly distributed in Gondwana. The current Laurasian-only-distribution is very likely a result of sampling bias as the vast majority of known Jurassic insect deposits are Palearctic, with exceptionally few

Gondwanan Jurassic insect fossils known primarily from deposits in Antarctica ${ }^{19}$. The heterogyrine lineage leading to Heterogyrus most likely arrived similarly to other Mesozoic fauna of the time, having become isolated from relatives in Africa or South America following the break up of Gondwana ${ }^{22}$. The phylogeny (Fig. 1) has Heterogyrus diverging from Cretotortor archarensis between $138 \mathrm{Ma}-90 \mathrm{Ma}(\mathrm{hpd})(\mathrm{m}=109 \mathrm{Ma})$, but this estimate is unreliable given the Laurasian distribution of Cretotortor archarensis, the isolated position of Madagascar at this time, the low posterior probability associated with the node, and likely is an artifact of the limited available sampling.

A distinct transition is seen in the fossil record of the Gyrinidae following the K-T boundary (Fig. 2), where the Heterogyrinae are replaced by the Gyrininae as the dominant whirligig beetle fauna. Madagascar appears to have served as a refugium for the last heterogyrine lineage, having been isolated since the break up of Gondwana, while heterogyrines went extinct elsewhere. In Madagascar, the extant Heterogyrus milloti has a very limited distribution, known only from two localities in the southeast with old geologic origins: the mountains near mount Andringitra ${ }^{6,23}$, whose origins date to the Permian; and the other in the nearby mountainous cloud forests of Ranomafana ${ }^{5}$. Here Heterogyrus is only encountered in small trickling streams, where no other gyrinids are found. A variety of gyrinine species are found on the larger streams and ponds in the surrounding area at lower elevations. Fossils of heterogyrines are known from a variety of habitats such as large lakes (Fig. 2) (i.e. Mesogyrus antiquus known from the Karabastau formation and M. elongates from the Sharteg formation ${ }^{24}$ ) as well as fluvial deposits (Cretotortor zherichini from the Kzyl-Zhar deposits), similar to modern Gyrininae, suggesting the current unique habitat of Heterogyrus milloti represents a final stronghold for this relictual species.

The other Malagasy gyrinid fauna, members of the Gyrininae, exhibit a more typical pattern, diverging from African sister taxa, within the Cenozoic (Fig. 1). The notable exception being Orectogyrus cyanicollis and $O$. hastatus, which show a Late Cretaceous divergence (hpd $66 \mathrm{Ma}-99 \mathrm{Ma}, \mathrm{m} 83 \mathrm{Ma}$ ) from their African sister taxa (Fig. 1). The late arrival of the gyrinines lends further support for an early arrival of the ancestor of H. milloti, as gyrinines appear to have replaced heterogyrines elsewhere. The heterogyrine lineage extant today was able to persist via isolation on Madagascar following the break up of Gondwana, and remaining isolated in a unique habitat in an isolated mountain chain, to become the oldest endemic animal lineage on Madagascar.

The Spanglerogyrinae and Heterogyrinae serve as exemplars of the persistence of coleopteran lineages contributing to the diversity of the group ${ }^{18}$, and the unique nature of the fauna of Madagascar. Our findings of two currently monotypic gyrinid subfamilies representing the sole surviving lineages of the Mesozoic fauna are remarkable. Importantly both species are only known from very few localities, within threatened biodiversity hotspots. Given the exceptionally unique evolutionary story arthropod lineages tell, greater consideration should be given to arthropod inclusive conservation ${ }^{25}$ to protect these last remaining Mesozoic gyrinid lineages, and other unique arthropod lineages.

## Figure legends

Figure 1. Phylogenetic relationships of the Gyrinidae. Tree from Bayesian analysis using the FBD macroevolutionary model ${ }^{15,16}$. Blue bars show $95 \%$ highest posterior density age range (hpd), dates at select nodes represent median (m) age from the hpd. The asterisk above select nodes indicates high support ( $\geq 95 \%$ posterior probability). Red branches denote Malagasy
lineages. Gray triangles show selected fossils' phylogenetic placement as terminals, unique to the total evidence approach ${ }^{14}$. Fossils and dineutine exemplar not to scale.

Figure 2. Table of known gyrinid fossils. Green indicates heterogyrine fossil species, teal gyrinine fossil species, and yellow spanglerogyrine fossil species.

Figure 3. Spanglerogyrine species. Arrows indicate important morphological features, abbreviations pro $=$ pronotum, $\mathrm{frs}=$ frons, $\mathrm{frl}=$ frons lateral margin. $\mathbf{a}$, Angarogyrus mongolicus fossil no. 3149/970 (a') line drawing of specimen. b, Spanglerogyrus albiventris dorsal habitus, scale bar $=1 \mathrm{~mm} . \mathbf{c}$, Angarogyrus minimus fossil no. $1670 / 385$, scale bar $=1 \mathrm{~mm}\left(\mathbf{c}^{\prime}\right)$ line drawing of specimen. d, scanning electron microscope (SEM) of Spanglerogyrus albiventris pronotum and head, scale bar $=500 \mu \mathrm{~m} . \mathbf{e}$, Spanglerogyrus albiventris lateral habitus to scale with b. f, Angarogyrus minimus elytron fossil no. $1670 / 385$ scale bar $=1 \mathrm{~mm}$.

Figure 4. Heterogyrine species. Arrows indicate important morphological features. a, Cretotortor zherichini elytron fossil no. 3149/970 (a') line drawing of specimen. b, Heterogyrus milloti elytron SEM, scale bar = 1 mm . c, Baissogyrus savilovi holotype fossil no. 1668/1787, scale $\mathrm{bar}=1 \mathrm{~mm}\left(\mathbf{c}^{\prime}\right)$ line drawing of specimen. $\mathbf{d}$, the Malagasy striped whirligig beetle, $H$. milloti dorsal habitus, scale bar 2 mm . e, pronotum of $H$. milloti. f, thorax and abdomen of $H$. milloti ventral view, scale bar $=2 \mathrm{~mm} . \mathbf{g}$, Mesogyrus antiquus fossil no. 2997/1846, scale bar $=2$ $\mathrm{mm},\left(\mathbf{g}^{\prime}\right)$ line drawing of specimen.

## Methods

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assement.

## Phylogenetic analysis.

The entire phylogenetic dataset from Miller \& Bergsten, 2012 was utilized, with the exception of the morphological portion. Thirty-three additional ingroup gyrinid taxa and one outgroup taxon, Haliplus lineatocollis, were added to the analysis (Extended Data Tab. 2). Additional taxa included sequences for all genes except EF1 $\alpha$ which was found to have multiple copies in Gyrinidae ${ }^{5}$. Thirteen fossil taxa were included, four outgroup taxa, and five ingroup (Extended Data Tab. 3). The fossil Coptoclava longipoda was originally included in the dataset but was removed for the analysis. In total taxon sampling for the analysis included ten outgroup Hydradephaga species and 128 gyrinid taxa.

The same genes and primers were used as in Miller \& Bergsten, 2012. A novel morphological character set of 120 characters was established (Supplementary Data 1) and taxa were coded as in Supplementary Data Table 1. This brought the total character number up from that of Miller \& Bergsten, 2012 to 3438 . The same partitioning scheme was used in Miller \& Bergsten, 2012.

Bayesian phylogenetic analysis was implemented using MPI version of MrBayes 3.2.6 ${ }^{15,26}$ and were run on the super computer cluster 'Ulam' at the Center for Advanced Research Computing (CARC), University of New Mexico. No substitution model was selected a priori, instead the reversible-jump MCMC method with gamma rate variation across sites was used to test the probability of different models a posteriori during analysis ${ }^{5,26,27}$. To infer the substitution rate the methods outlined by Ronquest et al. 2012 were followed with the mean age
of the fossil Moravocoleus permianus ( 293 Ma ), the oldest known Coleopteran ${ }^{28}$, used to calculate median rate, and the mean age of Triaplus sibiricus ( 253 Ma ), the oldest known aquatic Adephagan ${ }^{28}$, for the standard deviation. The Fossilization-Birth-Death macroevolutionary model ${ }^{16}$ was employed using the methods outlined by Zhange et al. 2015. The sampling strategy was set to diversity, with a sample probability of 0.01 as there are an estimated 1000 species of Gyrinidae ${ }^{29}$. Fossils were given a uniform age prior based on the age of the fossil. The tree root was constrained and with a calibration point given a uniform prior with the minimum age based on the age of T. sibiricus and the maximum on M. permianus. The analysis was run for 10 million generations, using four chains (three heated, one cold), with swap number set to two, and a temperature of 0.1 for the heated chains. MCMC convergence was monitored using Tracer v.1. $6^{30}$.

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## Author contributions

G.T.G and J.B.S designed the study. J.B.S. aided in analysis, contributed data, and aid in the field. G.T.G. performed the analyses, constructed morphological data set, coded characters, constructed figures, and wrote the paper. A.A.P. selected outgroup fossil taxa, provided images of fossils, and character coding suggestions. R.B. performed literature review and constructed dated Malagasy lineage table. K.B.M. provided data, specimens, and aid in field work.

## Extended Data

## Extended Data Table 1. Dated Malagasy lineages.







|  |  | Chamaeleonidae Chamaeleonidae | 54 54 | 34 34 | 77 75 | rmc rmc | nc nc | Crottini et al. 2012 <br> Crottini et al. 2012 | $\begin{aligned} & 10.1073 / \text { pnas. } 111248710 \\ & 9 \\ & 10.1073 / \text { pnas. } 111248710 \\ & 9 \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Gerrhosauridae | 61 | 32 | 96 | rmc | nc | Crottini et al. $2012$ | 10.1073/pnas. 111248710 9 |
|  |  | Gerrhosauridae | 37 | 17 | 63 | rmc | nc | $\begin{aligned} & \text { Crottini et al. } \\ & 2012 \end{aligned}$ | ${ }_{9}^{10.1073 / \text { pnas. } 111248710}$ |
|  | Scincidae | Trachylepis | 1 | 9 | 37 | rmc | nc | Crottini et al. 2012 | ${ }_{9}^{10.1073 / \text { pnas. } 111248710}$ |
|  | Scincidae | Trachylepis | 24 | 10 | 39 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  | Scincidae | Scincinae | 65 | 39 | 96 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  | Scincidae | Scincinae | 47 | 23 | 70 | rmc | nc | Crottini et al. $2012$ | 10.1073/pnas. 111248710 <br> 9 |
|  | Gekkonidae | Phelsuma | 62 | 39 | 91 | rmc | nc | Crottini et al. $2012$ | $10.1073 /$ pnas. 111248710 9 |
|  | Gekkonidae | Phelsuma | 49 | 34 | 65 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 9 |
|  | Gekkonidae | Lygodactylus | 62 | 39 | 91 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 9 |
|  | Gekkonidae | Lygodactylus | 49 | 34 | 65 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 9 |
|  | Gekkonidae | Blaesodactylus | 42 | 23 | 68 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  | Gekkonidae | Blaesodactylus | 27 | 10 | 45 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 9 |
|  | Gekkonida | P | 57 | 34 | 86 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 |
|  | Gekkonida | P | 57 | 34 | 86 | rmc | nc | Crottini et al. | 9 10.1073/pnas. 111248710 |
|  | Gekkonidae | Paroedura | 43 | 27 | 60 | rmc | nc | 2012 |  |
|  | Gekkonidae | Uroplatus | 51 | 29 | 78 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  | Gekkonidae | Uroplatus | 38 | 20 | 58 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 |
|  | Gekkonidae | Hemidactylus | 4 | 1 | 10 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 9 |
|  | Gekkonidae | Hemidactylus | 6 | 1 | 11 | rmc | nc | Crottini et al. $2012$ | 10.1073/pnas. 111248710 <br> 9 |
|  | Tortoise | Geochelone | n/p | 11,5* | 17,5* | seqdivR |  | Palkovacs et <br> al. 2002 | $\frac{\text { doi:10.1016/S } 1055-}{7903(02) 00211-7}$ |
|  | Tortoise | Pyxis | n/p | 9.5* | 14.5* | seqdivR |  | Palkovacs et <br> al. 2003 | $\frac{\text { doi:10.1016/S1055- }}{7903(02) 00211-7}$ |
|  | Tortoise | Geochelone | n/p | 14 | 22 | seqdivR | mt | Caccone et <br> al. 1998 | $\frac{\text { doi:10.1006/mpev. } 1998.0}{\underline{594}}$ |
|  | Tortoise | Pyxis | n/p | 8* | 12* | seqdivR | mt | Caccone et al. 1999 | $\frac{\text { doi:10.1006/mpev. } 1998.0}{\underline{594}}$ |
|  |  | Podocnemidid | 112 | 73 | 159 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  |  | Podocnemid | 112 | 73 | 159 | rmc | nc | Crottini et al. |  |
|  |  | Podocnemididae | 87 | 65 | 111 | rmc | nc |  | 10.1073/pnas. 111248710 |
|  |  | Testudinidae | 79 | 33 | 134 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  |  | Testudinidae | 16 | 6 | 30 | rmc | nc | Crottini et al. 2012 | 10.1073/pnas. 111248710 <br> 9 |
|  |  | Crocodylidae | 5 | 0 | 20 | rmc | nc | Crottini et al. $2012$ | $10.1073 /$ pnas. 111248710 |
|  |  |  |  |  |  |  |  | Crottini et al. |  |
|  |  | Crocodylidae | 1 | 0 | 2 | rmc | nc | 2012 | $\begin{aligned} & 10 \\ & 9 \end{aligned}$ |
| $\frac{\tilde{U}}{\frac{0}{2}}$ | orb weaver spider | Nephila spiders | 2.46 | 0.6 | 5.3 | Brc | mt\&nc |  <br> Agnarsson <br> 2011 | $\begin{aligned} & 10.1186 / 1471-2148-11- \\ & 119 \end{aligned}$ |
|  | spider assasin spiders | Madagascar |  |  |  |  |  | Agnarsson |  |
|  |  | Nephilengys | 1.9 | n/p | 7.4 | Brc | mt\&nc | 2011a | $\frac{\frac{10.10 .10}{1.02 .002}}{\underline{1}}$ |
|  |  | Eriauchenius | 154 | 115 | 191 | Brc | $\mathrm{mt} \& \mathrm{nc}$ | Wood et al. $2015$ | 10.1111/evo. 12578 |


| Snails | snail | Madagasikara/P achychilus | 15.6 | 20.3 | 31.5 | smc | mt | Köhler \& Glaubrecht 2012 | $\begin{aligned} & \text { 10.1111/j.1095- } \\ & 8312.2009 .01390 . \mathrm{x} \\ & \hline \end{aligned}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Flatworms | flatworms | Madapolystoma/ Eupolystoma | 116.2 | 95.6 | 134.6 | mdiv | rRNA | Verneau et <br> al. 2010 | 10.1098/rspb.2008.1530 |

*     - only the crown age provided
\# - the age is not provided.
$a$-approximate
Abbreviations for methods

Brc - Bayesian relaxed clock
clEn - clock enforced
coal - coalescence
mc - molecular clock
mDiv - multidivtime
rate - rate smoothing
rmc - relaxed molecular clock
smc - strict molecular clock
mdRc - multidivtime relaxed clock
clEn - clock enforced
PL - penalized likelihood
seqdivR - sequence divergence rate

Abbreviations for genes

| $\mathrm{mt}-$ mitochondrial | cDNA - complimentary DNA |
| :--- | :--- |
| $\mathrm{nc}-$ nuclear | BACseq - Bacterial artificial chromosome |
| rRNA - ribosomal RNA | sequence |

cDNA - complimentary DNA
BACseq - Bacterial artificial chromosome sequence

Highlighted taxa indicate lineages predating the Cenozoic.

## Extended Data Figure 1. Characters shared between Spanglerogyrus and Heterogyrus. a,

SEM of $S$. albiventris anterior view, arrow indicates setose furrow dividing labrum, scale bar = $200 \mu \mathrm{~m}$. b, the same feature on $H$. milloti, scale bar $=500 \mu \mathrm{~m}$. c, prolegs of S. albiventris, arrow indicates medial protibial spur, scale bar $=200 \mu \mathrm{~m}$. $\mathbf{d}$, the same on $H$. milloti scale bar $=300 \mu \mathrm{~m}$.

Extended Data Figure 2. Phylogenetic tree. Bayesian consensus tree resulting from analysis, from which Figure 1 was constructed. Number at nodes indicate posterior probabilities. Bars at node denote $95 \%$ highest probability density interval for age. Scale access in millions of years.

## Supplementary information

Supplementary Data 1. Description of morphological chracters.

## Morphology

First number indicates the character number in the morphology matrix only. Numbers given in parentheses correspond to character numbers in total-evidence matrix found in the Nexus file.

## Head

1 (3319). Head capsule shape excluding labrum. (0) elongate, longer than wide; (1) broad, wider than long. The head capsule of Haliplus and Hygrobia species are distinctly elongate, while those of the remaining species studied are clearly broad.
2 (3320). Divided eyes. (0) absent; (1) present. The eyes of two aquatic beetle families are clearly divided into a dorsal and ventral pair, the Coptoclavidae ${ }^{1}$ and the Gyrinidae. All other species studied exhibit non-divided eyes.
3. (3321) Eyes. (0) bulging; (1) in contour with head. The eyes of most hydradephagans are in contour with the headcapsule. Bulging eyes not in-line the contour of the headcapsule are present in Haliplus and Hygrobria species.
4 (3322). Eye division. (0) narrowly divided by thin canthus; (1) widely divided with well developed ocular ridge. Spanglerogyrus exhibits narrowly divided eyes separated by a thin canthus ${ }^{2}$, while the remaining gyrinid species have widely divided eyes separated by a well-developed interorbital ridge ${ }^{3,4}$.
5 (3323). Antennal form. (0) scape elongate and flagellum filiform; (1) compact flagellum, with an expanded pedicel and cup-like scape. The second character state describes the antennae unique to the family Gyrinidae.
6 (3324). Number of antennomeres in flagellum. (0) more than nine; (1) nine antennomeres; (2) eight antennomeres; (3) seven antennomeres; (4) six antennomeres.
7 (3325). Antennal flagellum apex with long setae. (0) absent; (1) present. Spanglerogyrus and Heterogyrus have antennal scape apices with long setae. These setae are absent in the remaining Gyrinidae and the other Hydradephaga.
8 (3326). Posterior margin of clypeus. (0) complete; (1) incomplete. Hygrobia and the Gyrinidae have a complete posterior margin of the clypeus. In the other hydradephaga studied the clypeal posterior suture is partially effaced.
9 (3327). Ratio of the frontolateral margin to the width of the clypeus at mid-length. (0) frontolateral margin at least 1.5 times the longer than the medial clypeal width; (1) nearly equal or less than one. The frontolateral margin character is specific to the Gyrinidae studied. The frontolateral margin is elongate in Spanglerogyrus, Heterogyrus, in many orectochilines and dineutines. A reduction of the frons length is seen independently in several gyrinid such as the gyrinines, Dineutus, and Gyretes and some Patrus.
10 (3328). Lateral margin of frons with a well developed bead. (0) absent; (1) present. This bead appears in some Gyrinidae such as Heterogyrus, Enhydrus, Macrogyrus, and Gyrinus.

11 (3329). Frons swollen and quadrate with frontolateral ridge continued dorsally to caudal third of dorsal eye ${ }^{5}$. (0) absent, fronts not swollen in appearance, frontolateral ridge not continued dorsally to caudal third of dorsal eye; (1) present. The distinctly swollen, quadrate frons ${ }^{2}$ with the frontolateral margins continue dorsally to caudal third of dorsal eye ${ }^{5}$ are unique to Spanglerogyrus and Angarogyrus.
12 (3330). Pseudofrontal ridge. (0) absent; (1) present, but narrow and weakly developed; (2) present and well developed, broad and often setose. The frons of orectochiline species has an additional lateral, depressed ridge, the pseudofrontal ridge ${ }^{6}$. This ridge is unique to this tribe of whirligig beetles.
13 (3331). Labral shape. (0) transverse; (1) elongate. A transverse labrum is very common within the Hydradephaga and in these analyses a labrum is coded as being transverse if it is less than half as long as wide. An elongate labrum is defined as being at least half as long as wide. An elongate labrum (Fig._) is present in Orectochilus, Orectogyrus, Porrorhynchus.
14 (3332). Labral form. (0) quadrate; (1) rounded, including triangular; (2) emarginate. The Spanglerogyrus and Angarogyrus possess a strongly quadrate labrum, all other gyrinid species have a rounded labrum, as well as Noterus clavicornis. An emarginate labrum is seen in most of the other dytiscoid species.
15 (3333) Labrum basally. (0) with transverse setose division, ventrad to division lightly colored cuticle present, dorsad to division cuticle darkly colored; (1) entire, no division evident. The labrum of Angarogyrus, Spanglerogyrus, and Heterogyrus exhibit a unique basal transverse division. All other species studied had the labrum entire.
16 (3334) Labrum dorsally with setae. (0) absent (1) present. The labrum of Spanglerogyrus, Heterogyrus, and oretochiline species exhibit dorsal setae. These setae are not present in any of the other species studied.
17 (3335) Maxillary galea. (0) two segment; (1) one segmented; (2) absent. The out-group hydradephagan species all have two segmented maxillary galea. Within the Gyrinidae, Spanglerogyrus and Heterogyrus have two segmented maxillary galea, the gyrinines have maxillary galea with a single segment, and the orectochilines and dineutines have the maxillary galea totally absent ${ }^{5-7}$. This character is treated as ordered in the analysis.
18 (3336) Palpi. (0) narrow and elongate; (1) broadened and shortened. Within the Gyrinidae, Spanglerogyrus and Heterogyrus have narrow and elongate labial and maxillary palpi. The other gyrinines have the palpi broadened and relatively shortened.
19 (3337) Palpigers. (0) free, not fused to mentum; (1) palpigers fused to mentum. The out-group hydradephagan species all have free, un-fused palpigers. Within the Gyrinidae, Spanglerogyrus ${ }^{5}$ and Heterogyrus also have free palpigers, while the remaining gyrinid species have the labial palpigers fused to the mentum.
20 (3338) Mentum. (0) weakly tri-lobed; (1) strongly tri-lobed. Heterogyrus milloti and Cretotortor striatus have a mentum with a well developed medial lobe, giving the mentum a strongly tri-lobed appearance. All other species studied have a weakly tri-lobed mentum.
21 (3339) Mental lateral lobe. (0) not strongly expanded; (1) strongly expanded. The lateral mental lobes are greatly expanded in the Gyrinidae ${ }^{5}$.
22 (3340) Clypealium. (0) mostly glabrous, few sparse setae, especially basally; (1) setose with row of long fine setae. The Orectochilini and Dineutini have a strongly setose clypealium. Character not coded for non-gyrinid taxa.

## Prothorax

23 (3341) Pronotum with largely expanded medial lobe. (0) absent; (1) present. The pronotum of Angarogyrus, Spanglerogyrus, Heterogyrus, and Mesogyus has a strong medial lobe. Other pronotum examined did not exhibit such an expanded medial region to the pronotum.
24 (3342) Expanse of lateral margin of pronotum. (0) not reaching anteriad to medial expanse of pronotum; (1) reaching at least equally anteriad to the medial expanse of pronotum, if not beyond. The lateral margins of the pronotum of Spanglerogyrus and Angarogyrus do not reach the medial lobe of the pronotum. In Heterogyrus and Mesogyrus the pronotal lateral margins extend to at least the length of the medial lobe. In the remaining Gyrinidae for which the pronotum is known, the medial lobe of the pronotum is lost and the lateral margins typically extend further anteriad than the medial expanse of the pronotum. Not coded for non-gyrinid taxa

25 (3343) Pronotal lateral bead. (0) absent; (1) present. Some Hydradephaga have a distinct lateral bead to the pronotum.
26 (3344) Pronotal transverse impressed line. (0) absent; (1) present. Most Gyrinidae have a transverse impressed line following the anterolateral margin of the pronotum ${ }^{8}$. This line is absent in Orectochilus and Porrorhynchus species.
27 (3345) Pronotum dorsally with transverse crease. (0) absent; (1) present. Species of Gyrinus exhibit a transverse crease dorsally on the pronotum ${ }^{7,8}$.
28 (3346) Pronotum with basolateral plicae. (0) absent: (1) present. Species of Haliplus exhibit strongly plicae basolaterally on the pronotum ${ }^{9}$.
29 (3347) Pronotal setation. (0) absent; (1) present. Spanglerogyrus, Heterogyrus, and the orectochiline species exhibit setation on the pronotum.
30 (3348) Prosternal medially expanded ventrad and differentiated. (0) Prosternum weakly expanded ventrally relative to lateral margins, weakly differentiated; (1) Prosternum strongly expanded ventrally relative to lateral expanse, medially clearly differentiated. In Spanglerogyrus the Prosternum is weakly humped, but not strongly ventrally expanded relative to the lateral margins. Other Hydradephagan species have the Prosternum strongly differentiated, either expanded ventrally or modified into a prosternal process.
31 (3349) Prosternal differentiation. (0) not cushion shaped; (1) cushion shaped. In Heterogyrus, Mesogyrus striatus, and the orectochilines, the Prosternum is medially expanded into a cushion shape ${ }^{4}$, whose medial region is variously modified.
32 (3350) Prosternal cushion ${ }^{4}$ medially. (0) with depression; (1) entire; (2) with elevated process. In Heterogyrus and Mesogyrus striatus the prosternal cushion has a medial depression. In many orectochilines the prosternal cushion is entire, without a medial depression or elevated process. In some orectochiline species, such as Orectochilus villosus, the prosternal cushion has a medial elevated process.
33 (3351) Prosternum medially. (0) without distinct process; (1) with well differentiated process. The genera Gyrinus and Dineutus have a well differentiated prosternal process as do the out-group hydradephagans studied. Members of Aulonogyrus and Macrogyrus have neither a well differentiated prosternal process, nor a prosternal cushion. However, their
prosterna medially are strongly expanded ventrally, becoming strongly differentiated from the lateral expanse, different from the situation in Spanglerogyrus.
34 (3352) Prosternal process extent. (0) ending at or prior to posterior margin of procoxae; (1) extending just beyond posterior margin of the procoxae; (2) extending between the mesocoxae. The Gyrinidae to not have the prosternal process extending beyond the posterior margin of the procoxae. Coptoclava longipoda and Liadytes longus have the prosternal process extending just beyond the procoxae ${ }^{1}$. The remainder of the out-group hydradephagan species have the procoxae extending between the mesocoxae.
35 (3353) Prosternal process form. (0) not strongly raised and plat-form-like, without truncate posterior margin; (1) strongly raised and plat-form-like, posterior margin truncate. The Haliplidae have strongly raised, plat-form-like prosternal process with a truncate posterior margin ${ }^{10}$.
36 (3354) Pronotum with lateral explanate margin. (0) absent; (1) present. The pronotum of many gyrinid species have a lateral explanat margin to the pronotum.
37 (3355) Pronotum lateral explanate margin color. (0) lightly colored, normally yellow; (1) darkly colored. The lateral explanate margin of most gyrinid species is lightly colored, in some it is darkly colored, as in Enhydrus species.

## Foreleg

38 (3356) Natatory setae. (0) absent; (1) present. Natatory setae is present on the foreleg of outgroup hydradephaga species, with the exception of Coptoclava longipoda.
39 (3357) Protibial medial spur number. (0) absent; (1) one spur; (2) two spurs. The species of the subfamily Gyrininae lack protibial spurs. Heterogyrus and Spanglerogyrus both have a single spur. All the out-group hydradephagans have two medial protibial spurs, with the exception of Noterus clavicornis.
40 (3358) Protibial medial spur modification. (0) unmodified; (1) modified for digging; (2) modified raptorially. Coptoclava longipoda has the protibial medial spurs modified raptorially ${ }^{1}$, being elongate and sharp. Hygrobia and Noterus have the protibial medial spurs modified for digging ${ }^{11,12}$. All other species studied with protibial medial spurs unmodified.
41 (3359) Fringe of setae along dorsal and anterior protibial margins. (0) absent; (1) present. Hygrobia and Noterus have a fringe of short stout setae along the dorsal and anterior apical margins of their protibia ${ }^{13}$.
42 (3360) Protrochanter ventral face. (0) without series of short stout setae; (1) with series of short stout setae apically; (2) with series of short stout setae extending nearly the entire length of the ventral face. The protrochanters of certain dineutines and orectochilines have a series of short stout setae along their ventral face, either limited apically or extending most the protrochanters length.
43 (3361) Protrochanteric setose patch. (0) absent; (1) present. The protrochanters of Porrorhynchus have a distinct setose patch (Gustafson and Miller, TBA).
44 (3362) Profemoral sub-apicoventral tooth/teeth. (0) absent; (1) present. Certain species of Dineutus have a profemoral sub-apicoventral tooth (Gustafson and Miller, 2015).
45 (3363) Setigerous punctures of anterior face of profemur. (0) absent; (1) present. Most species of gyrinid have at least one or more setigerous punctures present on the anterior face of the profemur ${ }^{14}$.

46 (3364) Ventral face of profemur. (0) without lines of setae on either anterior or posterior margin; (1) with one line of setae present on posterior margin only; (2) with two lines of setae on the posterior and anterior margin. Spanglerogyrus lacks lines of setae of the ventral face of the profemur, Heterogyrus has one line of setae along the posterior margin, and nearly all Gyrininae species have either two lines of setae on or at least one.
47 (3365) Setation of ventral face of profemur. (0) not composed of think tufts of setae; (1) composed of thick tufts of setae becoming distally. Porrorhynchus species have the two lines of setae of the ventral face of the profemur modified into thick tufts of setae that becoming denser distally.
48 (3366) Setose brush of posterior face of protibia. (0) absent; (1) reduced; (2) fully present. The protibia of most gyrinid species has some sort. In many species the setose brush is reduced to a small patch at the protibial apex, sometimes continue posteriorly by a very narrow strip of setae. The fully present state is a large triangular brush of setae beginning apically on the protibia and continue down the protibia. Most Dineutus have a fully present setose brush, as do Aulonogyrus species and some Orectogyrus. This character is treated as ordered.
49 (3367) Distolateral corner of protibia. (0) not expanded laterally; (1) expanded laterally. The protibia of orectochiline species and those of most dineutines, except Dineutus species, have the distolateral corner of the protibia laterally expanded and triangular in form, if not pointed. This character is also exhibited in Coptoclava longipoda.
50 (3368) Male protarsomere I posterior face. (0) without recessed pit; (1) with recessed pit containing differently shaped sucker-disc setae. The males of Macrogyrus species possess a recessed pit containing differently shaped sucker-disc setae ${ }^{15}$.
51 (3369) Female protarsomere V posterior face with setae. (0) absent; (1) present but reduced to small patch; (2) present as a line of numerous setae. This character is absent in gyrinines and Spanglerogyrus. But appears variously developed within dineutine and orectochilines ${ }^{4}$. Line of setae is fully present on the posterior face of female protarsomere V in Heterogyrus.

## Mesoventrite

52 (3370) Modification for proleg reception. (0) absent; (1) present. The mesoventrite of all gyrinids except Spanglerogyrus has recessed areas for receiving the prolegs ${ }^{7,16}$.
53 (3371) Mesoventrite size. (0) smaller than metaventirte; (1) larger than metaventrite. The Gyrinidae have a modified and greatly enlarged metaventrite. The out-group hydradephagan species all have the metaventrite much larger than the mesoventrite.
54 (3372) Mesoventrite with recessed hexagonal area for reception of prosternal process. (0) absent; (1) present. Most of the out-group hydradephagan species have the mesoventrite with a hexagonal recessed area for reception of the prosternal process ${ }^{10-12}$, with the exception of Coptoclava longipoda.
55 (3373) Mesoventrite shape. (0) not triangular; (1) triangular and extensive, often shaped similar to the bow of a ship. All gyrinid species with the exception of Spanglerogyrus have the mesoventrite triangular and extensive, shaped similarly to the bow of a ship.
56 (3374) Mesoventral discrimen. (0) absent; (1) present. A medial discrimen of the mesoventirte is present in all Gyrinidae except for Spanglerogyrus.

57 (3375) Mesoventrite with paramedical ridges. (0) absent; (1) present. The fossil gyrinid Baissogyrus savilovi has distinct paramedical ridges on the mesoventrite.
58 (3376) Mesoventral pit. (0) absent; (1) present. Some Gyrinus species have a the mesoventrite basomedially with a distinct pit.
59 (3377) Scutellar shield. (0) visible with elytra close; (1) invisible with elytra closed. The scutellar shield is variously invisible with the elytra closed of the species studied.
60 (3378) Scutellar shield shape. (0) more evenly triangular; (1) transversely triangular. The scutellar shield of Metagyrinus is transversely triangular ${ }^{17}$ and used as a character to distinguish it form Aulonogyrus.
61 (3379) Elytral length. (0) not covering abdominal apex; (1) covering abdominal apex. The elytra cover the apex of the abdomen in most of the hydradephagan out-group taxa. The abdominal apex is not covered in the Gyrinidae and in some out-group taxa.
62 (3380) Elytral setation. (0) absent; (1) present but with distinct glabrous regions to the elytra; (2) present, elytra nearly entirely pubescent. Within Gyrinidae the elytra are glabrous in the gyrinines and the dineutines. Pubescence is present on the elytra but with distinct glabrous regions in Spanglerogyrus, Heterogyrus, Orectogyrus, Patrus, and most Gyretes. Completely pubescent elytra is found in Orectochilus and some Gyretes like Gyretes sericeus.
63 (3381) Elytral explanate lateral margin. (0) absent; (1) present. Many gyrinid species exhibit a broad explanate lateral margin to the elytra.
64 (3382) Elytral explanate lateral margin color. (0) lightly colored, yellow often; (1) darkly colored. Most gyrinids with an explanate latera margin have the margin lightly colored, normally yellow. Rarely is the lateral margin darkly colored, similar in color to the elytral disc, as in Enhydrus species.
65 (3383) Ten or more primary punctures accompanied by numerous secondary punctures. (0) absent; (1) present. Species of Haliplus have distinctly punctate elytra, with ten or more primary punctures associated with numerous secondary punctures ${ }^{9,10}$.
66 (3384) Serial striae number. (0) none evident; (1) nine visible; (2) eleven visible. Within the Gyrinidae the orectochilines have no visible elytral striae, at least dorsally, similarly with Spanglerogyrus, Angarogyrus, and Porrorhynchus. Heterogyrus, Mesogyrus, Cretotortor and most the dineutines, with the exception of Porrorhynchus, have nine elytral striae visible. The Gyrinini all have eleven elytral striae.
67 (3385) Elytral strial appearance. (0) punctures; (1) well impressed lines; (2) faintly evident lines. The elytral striae appear as punctures in the gyrinines, as well as in M. (Andogyrus) seriatopunctatus and the fossil Meiodineutes amurensis, suggesting that dineutines. Strongly impressed lines are evident in Heterogyrus, Mesogyrus, Cretotortor, Metagyrinus and Enhydrus. Weakly impressed lines are present primarily in Dineutus and Macrogyrus. This character is treated as ordered as several Aulonogyrus and Gyrinus species exhibit intermediate stages between punctate to strongly impressed lines, suggesting a trend from punctures to strongly impressed lines, with weakly impressed lines as a step towards loss of impressed lines and elytral striae in general.
68 (3386) Elytral sutural border. (0) absent; (1) present. The elytra is bordered by an additional, non-serial striae (Brinck, 1955). This border is present in most gyrinid species, being completely lost in dineutines, and lost in some orectochilines.

69 (3387) Elytral lateral plica. (0) absent; (1) present. The two species of Angarogyrus studied have a distinct longitudinal plica laterally on the elytra. This character is unique to Angarogyrus.
70 (3388) Elytral apices. (0) unmodified; (1) modified. Unmodified elytra, those that are regularly rounded and attenuated towards the apex are common in the out-group hydradephagans studied, but relatively rare in the Gyrinidae. Unmodified elytra are primarily found in Dineutus and some Gyrinus.
71 (3389) Elytral apex with sutural production. (0) absent; (1) present. The sutural angle of the elytra has a production ${ }^{18}$ in many species of Gyrinidae. Importantly a sutural production is present in both Spanglerogyrus and Angarogyrus.
72 (3390) Elytral apex with parasutural production. (0) absent; (1) present. The elytral apex may have a production between the sutural and epipleural angles, the parasutural production. This is present in dineutine species in members of Dineutus, Porrorhynchus, and Macrogyrus.
73 (3391) Elytral apex with epipleural angle modified. (0) absent; (1) present as prominence; (2) present, spinose. The epipleural angle is modified as a strong spine in many orectochilines and some dineutines. The epipleural prominence is variously present among gyrinid species.
74 (3392) Elytral apices with straight truncation. (0) absent; (1) present. In many gyrinid species the elytral apex has a straightly truncate margin. This is common in many orectochilines and dineutines.
75 (3393) Elytral apices with oblique truncation. (0) absent; (1) present. An oblique truncation to the apex of the elytra is less common within Gyrinidae. Heterogyrus, Mesogyrus, and Cretotortor exhibit this type of truncation, as do some oretochilines and very few dineutines.
76 (3394) Elytral apices with serrations/irregularities. (0) absent; (1) present. Some Dineutus species exhibit serration and/or irregularities to the elytral apices (Gustafson and Miller, 2015)

77 (3395) Elytral apicolateral margins with buzz-saw shaped serration. (0) absent; (1) present. Most species of Porrorhynchus exhibit this time of elytral modification. It is also present in Dineutus micans.
78 (3396) Elytral with canaliculated microsculpture. (0) absent; (1) present. This microsculpture appears as minute "scratch-like" sculpturing of the elytra. It is present on the elytra of the Macrogyrus s. str. species.

## Mid-legs

79 (3397) Mid-legs. (0) not broadened nor flattened; (1) not broadened but dorsoventrally flattened with expanded mesotibia; (2) broadened and flattened dorsoventrally; (3) broadened, flattened dorsoventrally, but also shortened and paddle-like. Most of the outgroup hydradephagan species have mid-legs that are not broadened or flattened. Spanglerogyrus has mid-legs that are not broadened but are dorsoventrally flattened with an expanded mesotibia. The fossil Angarogyrus mongolicus has the mid-legs visible, these are interpreted as being similar to those of Spanglerogyrus given the majority of other similarities the species share in morphology. The legs appear slightly broader than the totally unmodified legs of the out-group taxa, but certainly not broadened like those
of Coptoclava longipoda nor of the other gyrinid species. State (2) is unique in the analysis to Coptoclava longipoda which exhibits broad and flattened midlegs. State (3) is unique to all Gyrinidae except Spanglerogyrus and Angarogyrus mongolicus.
80 (3398) Mesocoxal shape. (0) rounded; (1) triangular and strongly transverse. Triangular and strongly transverse mesocoxae are unique to the Gyrinidae ${ }^{19}$.
81 (3399) Mesocoxae separation ${ }^{6}$. (0) narrowly separated; (1) broadly separated. The mesocoxae are broadly separated in the dineutines, and most of the orectochiline genera Patrus and Gyretes.
82 (3400) Meso- and metatibial medial spurs. (0) both spurs large, greater than or nearly equal to half the length of the first tarsomere; (1) both spurs small, less than half the length of the first tarsomere; (2) the metatibia with the posterior spur larger than or nearly equal to half the length of the first metatarsomere; (3) both the meso- and metatibia with the posterior spur larger than or nearly equal to half the first tarsomere. In the out-group hydradephagan species as well as in Spanglerogyrus ${ }^{5}$ and Heterogyrus the meso- and metatibial medial spurs are both large. In the majority of the Gyrinidae both spurs are short. In some Gyrinus species and orectochiline species the metatibia has the posterior spur large and the anterior spur small. Only in Macrogyrus howittii and in one Gyretes species was a large posterior spur observed in both the meso- and metatibia.
83 (3401) Mesotarsal claws. (0) not sexually dimorphic; (1) weakly sexually dimorphic; (2) strongly sexually dimorphic. The male mesotarsal claws of Dineutus species are strongly sexually dimorphic ${ }^{18}$. Within Porrorhynchus the mesotarsal claws are weakly sexually dimorphic. In the remaining Gyrinidae the mesotarsal claws are not sexually dimorphic.

## Metaventrite

84 (3402) Mesoventrite with anteromedial process. (0) absent; (1) present, receiving posterior expansion of prosternal process. Most of the out-group hydradephagan species have the mesoventrite with an anteromedial process that receives the posterior expansion of the prosternal process, with the exception of Liadytes longus and Coptoclava longipoda ${ }^{1}$.
85 (3403) Metaventrite. (0) not largely expanded anteriorly; (1) largely expanded anterior. Haliplus species have the metaventrite largely expanded anteriorly ${ }^{10}$.
86 (3404) Metaventrite paramedially. (0) not constricted; (1) weakly constricted; (2) distinctly constricted. In the out-group hydradephaga species the metaventrite is not noticeably constricted paramedially by the mesocoxae with the exception of Coptoclava longipoda, which has a weak constriction. Within the Gyrinidae only Spanglerogyrus has the metaventrite weakly constricted paramedially by the mesocoxae, all the remaining Gyrinidae have the metaventrite distinctly constricted, resulting in the formation of "metaventral wings" ${ }^{6}$.
87 (3405) Metaventral wings. (0) absent; (1) in the form of a near equilateral triangle; (2) narrowed and strap-like. The out-group hydradephagan species and Spanglerogyrus lack metaventral wings like other gyrinid species, as per the above character. Heterogyrus, Mesogyrus, Mesodineutes, and the dineutines have the metaventral wings in the form of a near equilateral triangle. The gyrinines and the orectochilines have the metaventral wings strap-like, strongly narrowed medially then gradually broadened laterally.
88 (3406) Medial expanse of metaventrite. (0) relatively narrower and diamond shaped; (1) very broad and pentagonal shaped. The species of Enhydrus have a very broad medial expanse
of the metaventrite (the area medial to the mesocoxae) that is strongly pentagonal shape with a nearly straight posterior margin. Some Macrogyrus species in the subgenus Andogyrus come close to have a similarly shaped medial expanse, however, the posterior margin is not nearly as straight.
89 (3407) Discrimen of metaventrite with transverse sulcus. (0) present and long; (1) present but short; (2) absent. Some of the out-group hydradephagan taxa retain a long transverse sulcus associated with the metaventral discrimen, as does Spanglerogyrus and interesting many Macrogyrus species. An intermediate stage is present in the Gyrinidae, where a transverse sulcus is present but is greatly shortened ${ }^{7}$. This character is present in Heterogyrus, Mesogyrus antiquus, and Baissogyrus.
90 (3408). Metepisternal ostiole. (0) absent; (1) present. The metepsternum of Gyrinus species has an ostiole that is variously developed ${ }^{8}$.
91 (3409). Metanepisternum shape. (0) largely triangular; (1) lobiform; (2) narrow and trapezoidal in form. The out-group hydradephaga have a largely triangular metanepisternum, as do Spanglerogyrus, the gyrinines, and a few Patrus species. A lobiform Metanepisternum is found in Heterogyrus, Mesogyrus, Mesodineutes, and the dineutines. A strongly narrowed and trapezoidal shaped Metanepisternum is unique to most of the orectochilines, with the exception of a few Patrus.
92 (3410). Metanepisternum reaching coxal cavities. (0) absent, ending prior to coxal cavities; (1) present. This character is found in some of the out-group hydradephagan taxa, but is never present in the Gyrinidae.
93 (3411). Noterid platform. (0) absent; (1) present. The distinct "noterid platform" is found only in Noterus clavicornis.

## Hind legs

94 (3412). Hind legs. (0) narrow; (1) narrow and weakly dorsoventrally flattened with expanded metatibia; (2) broadened for swimming, but not dorsovenrally flattened; (3) broadened and significantly dorsoventrally flattened; (4) broadened, significantly dorsoventrally flattened, and shortened to a paddle-like form. Unmodified narrow hind legs are found in the hydradephagan out-group species of Hygrobia, Haliplus, and Liadytes. Narrow and weakly dorsoventrally flattened hindlegs with expanded metatibia are found in Spanglerogyrus, the hind legs of Angarogyrus mongolicus are treated similarly as justified for the mid-legs. Hind legs that are broadened for swimming, but not dorsoventrally flattened, are found in the Dytiscidae and Noterus species examined. The broadened and significantly dorsoventrally flattened hind-legs, that are not shortened, are found in Coptoclava longipoda. Finally the majority of gyrinid species exhibit the paddle-like leg form, with hid legs that are expanded, dorsoventrally flattened, and shortened.
95 (3413). Anterior margin of metacoxae. (0) more transverse; (1) distinctly oblique to very strongly oblique. Most of the hydradephagan out-group species studied (with the exception of the Dytiscidae), Spanglerogyrus, the dineutines, and a few orectochilines have a more transverse anterior margin to the metacoxae. In Heterogyrus, Mesogyrus, the gyrinines, and most of the orectochilines have a distinctly more oblique anterior margin to the metacoxae.

96 (3414). Posterolateral margin of metacoxae. (0) without border; (1) bordered. Many gyrinid species have the posterolateral margin of the metacoxae bordered. Heterogyrus, Mesogyrus, Aulonogyrus, Orectochilus, Porrorhynchus, and Dineutus do not have a border along the posterolateral margin of the metacoxae.
97 (3415). Metacoxae. (0) not expanded anteriorly; (1) greatly expanded anteriorly. The Dysticidae have the metacoxae greatly expanded anteriorly.
98 (3416). Metacoxal plate (sensu Lawrence et al. 2011). (0) present as large plates concealing the basal portion of leg and part of abdomen; (1) present, not concealing leg and abdomen, but continued laterally along anterior margin of metacoxae, (2) strongly reduced present only medially. As Lawrence et al. 2011 defines metacoxal plates: "the excavation of the metacoxae to form at least weak coxal plates", we consider the medial raised region of the metacoxae to be reduced coxal plates, which in the out-group hydradephagan species studied are present only medially, with the exception of Haliplus and Triaplus species ${ }^{1}$, which have large metacoxal plates covering the basal portion of the hind legs ${ }^{10}$. In the Gyrinidae the metacoxal plates are present medially but also continued laterally along the anterior margin of the metacoxae.
99 (3417). Metacoxal plate secondary reduction in Gyrinidae. (0) absent, metacoxal plate region largely triangular; (1) present, metacoxal plate medially rectangular in form with a narrow anterior bridge or with bridge totally reduced. Within the Gyrinidae the dineutines and Spanglerogyrus have large broadly triangular metacoxal plates. Within the orectochilines and gyrinines the metacoxal plates are consricted to a medial rectangular area, with a narrow anterior bridge continued laterally or with the anterior bridge totally reduced in some taxa. This character is associated with more oblique metacoxae.
100 (3418). Metacoxal process apex. (0) straightly truncate; (1) obliquely truncate; (2) rounded and lobiform. The metacoxal process apex of Gyrinidae is most often obliquely truncate. Many orectochilines have this apex straightly truncate, with many Orectogyrus having a rounded or lobiform metacoxal process apex.

## Abdomen

101 (3419). Suture of abdominal sternite II. (0) totally or mostly obliterated; (1) present. The species of Enhydrus still have the suture of abdominal sternite II present ${ }^{3,21}$.
102 (3420). Overal shape of abdominal apex. (0) not cylindrical, clearly broadly rounded; (1) distinctly cylindrical and strongly narrowed apically. The orectochilines have a strongly cylindrical and stinctly narrowed abdominal apex.
103 (3421). Abdominal sternites VII \& VIII. (0) glabrous; (1) with row of long fine setae. The orectochilines have a row of long fine setae posteromedially on abdominal sternites VII \& VIII forming a 'keel' ${ }^{7}$.
104 (3422). Abdominal sternite VIII. (0) apically longitudinally divided; (1) apically biemarginate; (2) entire. This character is discussed in Miller and Bergsten, 2012 and treated as ordered in the analysis.
105 (3423). Posterior margin of penultimate abdominal tergite. (0) not trilobed; (1) weakly trilobed; (2) strongly trilobed. This character is discussed in Miller and Bergsten, 2012.
106 (3424). Tergite VIII divided medially in half. (0) absent, tergite VIII entire; (1) present. The out-group hydradephagan species have tergite VIII divided medially in half.

107 (3425). Venter coloration. (0) entirely lightly colored; (1) distinctly infuscate; (2) entirely darkly colored. Most gyrinid species have either light (ranging from light red, orange, yellow, and even white) or darkly colored venters (dark reddish brown, brown, to black). Some gyrinine and Orectogyrus species exhibit distinctly infuscate venters.

## Female reproductive tract

108 (3426). Spermatheca. (0) not largely expanded nor sac-like; (1) largely expanded and saclike. The spermatheca of dineutines and Orectochilus and Orectogyrus are largely expanded and sac-like to varying degrees. The spermathecae of other gyrinidea are not nearly as expanded or sac-like.
109 (3427). Spermathecal accessory gland. (0) absent; (1) present. An accessory gland attached to the spermatheca is present in Spanglerogyrus, Heterogyrus, and the gyrinines.
110 (3428). Bursal accessory gland. (0) absent; (1) present. An accessory gland attached to the bursa is present in Heterogyrus, the orectochilines, and Enhydrus and Macrogyrus.
111 (3429). Vaginal shield. (0) absent; (1) present. The vaginal shield ${ }^{22-24}$ is present in Porrorhynchus and Dineutus.
112 (3430). Fertilization duct expansion. (0) absent; (1) weakly expanded; (2) strongly expanded. Most orectochilines have the fertilization duct weakly to strongly expanded. The remaining gyrinid species typically do not have the fertilization duct expanded ${ }^{7}$. This character is treated as ordered in the analysis.
113 (3431). Fertilization duct convolution. (0) absent; (1) randomly convoluted; (2) cork-screw shaped convolution. The fertilizaiton duct of orectochilines is often convoluted with random twists and turns. In certain Gyretes species the convolutions are arranged serially into a cork-screw shape ${ }^{7}$. This character is treated as ordered in the analysis.
114 (3432). Fertilization duct curling. (0) absent: (1) weakly curled; (2) strongly curled. The fertilization duct of Orectochilus and Orectogyrus species are curled. In Orectochilus this curling is weakly, with only a single turn back into itself, similarly with some Orectogyrus species. Many Orectogyrus species of the s. str. subgenus have the fertilization duct strongly curled, with numerous recurves, creating a snail-shell shape. Other gyrinids do not have the fertilization duct curled. This character is treated as ordered in the analysis.
115 (3433). Gonocoxae with medial apodeme. (0) absent; (1) present. This character is described by Miller and Bergsten, 2012.

## Aedeagus

116 (3434). Orientation of aedeagus in repose. (0) no rotated; (1) rotated. The aedeagus of Gyrinidae is not rotated in repose, unlike other hydradephaga.
117 (3435). Basal piece. (0) distinctly present; (1) present but fused to parameres; (2) absent/indistinguishable. Spanglerogyrus has a distinct basal piece ${ }^{5}$ which was reconfirmed in this study. Heterogyrus has a distinct additional segment fused to the parameres, which is interpreted here as the basal piece. The remained of the Gyrinidae and the out-group hydradephaga species no longer have a distinguishable basal piece. This character is treated as ordered in the analysis.

118 (3436). Paramere position. (0) ventral to median lobe; (1) lateral to median lobe. The genus Orectogyrus has parameres that are situated lateral to the median lobe. All other gyrinid psecies have the parameres situated ventral to the medial lobe.
119 (3437). Parameres with ventromedial project. (0) absent; (1) present. The gyrinines have a distinct ventromedial projection extending off the parameres. This projection is in a serially homologous postion to the fused basal piece of Heterogyrus. This character is coded here are a distinct character, but it seems reasonable that this character could be interpreted as a transformation series step, and supports the position of the gyrinines as sister to Dineutini + Orectochilini, where this medially project is reduced, representing the total loss of the basal piece.

## Sperm

120 (3438). Spermostyle type sperm conjugation. (0) absent; (1) present. The dineutines and Orectogyrus and Orectochilus have sperm conjugation utilizing a specialized structure, the spermostyle ${ }^{25}$. This type of sperm conjugation appears associated with female reproductive tracts that have a large sac-like spermatheca.

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Supplementary Data Table 1. Character coding for morphology dataset.

| Character \# | 1 | 5 | 10 | 15 | 20 | 25 | 30 | 35 | 40 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
|  | 1 | $\mid$ | $\mid$ | $\mid$ | $\mid$ | $\mid$ | $\mid$ | $\mid$ | $\mid$ |


| Trlati <br> Triaplus laticoxa | 101- | 0???- | ? 000 ? | ????? | 0?-0- | ?000? | 00-00 | 00-?? | ???? |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Nocl503 <br> Noterus clavicornis | 101- | 0001- | 00001 | 10010 | 00-0- | 10000 | 10-12 | 00-11 | 1100 |
| Haho504 Hygrobia hermanni | 000- | 0000- | 00002 | 10010 | 00-0- | 00000 | 10-02 | 00-12 | 1100 |
| Mabi2 Matus bicarinatus | 101- | 0001- | 00002 | 10000 | 00-0- | 10000 | 10-02 | 00-12 | 0000 |
| Plde130 Platynectes decemaculatus | 101- | 0001- | 00002 | 10010 | 00-0- | 10000 | 10-02 | 00-12 | 0000 |
| Lcla91 Lancetes lanceolatus | 101- | 0001- | 00002 | 10000 | 00-0- | 10000 | 10-02 | 00-12 | 0000 |
| Hali Haliplus lineatocollis | 000- | 0001- | 00002 | 100-0 | 00-0- | 10010 | 10-02 | 10-12 | 0000 |
| Hacr Haliplus cretaceus | ???? | ????? | ????? | 1???? | ??-0? | ? 0000 | 10-12 | 1???? | ???? |
| Colg Coptoclava longipoda | 011? | ????? | 000?? | 1???? | 00-0- | 00000 | 10-01 | 00-02 | 20?? |
| Lilo Liadytes longus | 101- | 010?- | 00002 | 1??00 | 00-0- | 10000 | 10-01 | 00-12 | 00?? |
| Mdrh Mesodytes rhantoides | 101- | 0??1- | 00002 | 10??? | ??-0- | 10000 | 10-12 | 00-?? | ???? |
| Spal472 Spanglerogyrus albiventris | 1110 | 11100 | 01000 | 01000 | 01010 | 11001 | 00-00 | 00-01 | 0000 |
| Hsmi596 Heterogyrus milloti | 1111 | 11100 | 10001 | 01000 | 11011 | 01001 | 110-0 | 01001 | 0000 |
| Aoal525 Aulonogyrus alternatus | 1111 | 11001 | 00001 | 10111 | 01001 | 01000 | 10-00 | 01000 | -000 |
| Aoma523 Aulonogyrus marginatus | ???? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ???? |
| Aocr519 Aulonogyrus cristatus | ???? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ???? |
| Aoca604 Aulonogyrus carinipennis | 1111 | 11001 | 10001 | 10111 | 01001 | 01000 | 10-00 | 01000 | -000 |
| Aosp569 Aulonogyrus sp | 1111 | 11001 | 10001 | 10111 | 01001 | 01000 | 10-00 | 01000 | -000 |
| Aobe493 Aulonogyrus bedeli | 1111 | 11001 | 10001 | 10111 | 01001 | 01000 | 10-00 | 01100 | -000 |
| Aoca540 Aulonogyrus caffer | 1111 | 11001 | 10001 | 10111 | 01001 | 01000 | 10-00 | 00-00 | -000 |
| Aost469 Aulonogyrus striatus | 1111 | 11001 | 00001 | 10111 | 01001 | 01000 | 10-00 | 01000 | -000 |
| Aogo529 Aulonogyrus goudoti | 1111 | 11001 | 00001 | 10111 | 01001 | 01000 | 10-00 | 00-00 | -000 |
| Berg1 Metagyrinus sinensis | 1111 | 1??01 | ?0001 | 10??? | ?1001 | ??000 | ????? | ?1000 | -0?? |
| Gyig598 Gyrinus ignitus | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Aost687 Aulonogyrus strigosus | 1111 | 11001 | 10001 | 10111 | 01001 | 01000 | 10-00 | 01000 | -000 |
| Gymi526 Gyrinus minutus | 1111 | 11101 | 00001 | 10111 | 01001 | 01100 | 10-00 | 00-00 | -000 |
| Gymd597 Gyrinus madagascariensis | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Gyna539 Gyrinus natalensis | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Gypl495 Gyrinus plicifer | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |


| Gygi496 Gyrinus gibber | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Gyel494 Gyrinus elevatus | 1111 | 11001 | 10001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Gysp492 Gyrinus sp | 1111 | 11001 | 00001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Gysp628 Gyrinus amazonicus | 1111 | 11001 | 00001 | 10111 | 01001 | 01100 | 10-10 | 00-00 | -000 |
| Difv672 Dineutus fauveli | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disp507 Dineutus pectoralis | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -000 |
| Disu484 Dineutus subspinosus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Diso605 Dineutus solitarius | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Dipx515 Dineutus proximus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disp576 Dineutus striatus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Dici474 Dineutus ciliatus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Didi473 Dineutus discolor | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disn516 Dineutus sinuosipennis | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disp481 Dineutus aereus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disu505 Dineutus sublineatus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Diin482 Dineutus indicus | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Disp577 Dineutus micans | 1111 | 14001 | 00001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -010 |
| Ayau483 Macrogyrus oblongus | 1111 | 11000 | 10001 | 10211 | 01101 | 01000 | 10-00 | 00-00 | -000 |
| Aygo501 Macrogyrus gouldi | 1111 | 11000 | 10001 | 10211 | 01101 | 01000 | 10-00 | 01000 | -000 |
| Ayan502 Macrogyrus australis | 1111 | 11000 | 10001 | 10211 | 01101 | 01000 | 10-00 | 00-00 | -000 |
| Aysp506 Macrogyrus albertisi | 1111 | 11000 | 10001 | 10211 | 01101 | 01000 | 10-10 | 00-00 | -000 |
| Ehas646 Enhydrus atratus | 1111 | 13000 | 10001 | 10211 | 01101 | 01000 | 10-10 | 01100 | -000 |
| Adsp648 Andogyrus zimmermanni | 1111 | 11000 | 10001 | 10211 | 01101 | 01000 | 10-00 | 00-00 | -000 |
| Prte497 Porrorhynchus marginatus | 1111 | 14000 | 00011 | 10211 | 01101 | 00000 | 10-10 | 01000 | -001 |
| Ogca666 Orectogyrus camerunensis | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Gesp615 Gyretes sericeus | 1111 | 11000 | 10001 | 11211 | 01101 | 01002 | 112-0 | 00-00 | -000 |
| Ogsp566 Orectogyrus noctuabundis | 1111 | 14000 | -0201 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Orvi527 Orectochilus villosus | 1111 | 11000 | 10011 | 11211 | 01101 | 00002 | 112-0 | 00-00 | -000 |
| Berg2 Orectochilus bellieri | 1111 | 11000 | 10011 | 11211 | 01101 | 00002 | 112-0 | 00-00 | -000 |


| Ogcy520 Orectogyrus cyanicollis | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Ogha600 Orectogyrus hastatus | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Orpr487 Patrus productus | 1111 | 14000 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Ordi488 Patrus discifer | 1111 | 14001 | 10001 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Orsp499 Patrus sp | 1111 | 14000 | -0101 | 11211 | 01101 | 00001 | 112-0 | 00-00 | -020 |
| Oran486 Patrus andamanicus | 1111 | 14001 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -010 |
| Orvo489 Patrus volubilis | 1111 | 14001 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -010 |
| Orsp500 Patrus sp | 1111 | 14001 | -0101 | 11211 | 01101 | 00001 | 111-0 | 00-00 | -020 |
| Orsp677 Patrus sp | 1111 | 14000 | -0101 | 11211 | 01101 | 00001 | 111-0 | 00-00 | -020 |
| Ogar669 Orectogyrus argenteovittatus | 1111 | 14000 | -0201 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Ogsj671 Orectogyrus sjostedti | 1111 | 14001 | -0211 | 11211 | 01101 | 01001 | 112-0 | 01000 | -020 |
| Ogpi665 Orectogyrus pictimanus | 1111 | 14000 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogde524 Orectogyrus dedalus | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Ogdi491 Orectogyrus discors | 1111 | 14000 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogos667 Orectogyrus oscari | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogpl662 Orectogyrus prolongatus | 1111 | 14001 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -010 |
| Ogdy664 Orectogyrus demeryi | 1111 | 14000 | -0101 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogsp565 Orectogyrus specularis | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogms670 Orectogyrus masculinus | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogsp663 Orectogyrus sp | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogmd601 Orectogyrus madagascariensis | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -010 |
| Ogdo517 Orectogyrus dorsiger | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogsp490 Orectogyrus posticalis | ???? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ???? |
| Ogsp564 Orectogyrus wittei | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogbd661 Orectogyrus bedeli | 1111 | 14000 | -0201 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogsp567 Orectogyrus specularis | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogsp668 Orectogyrus sp | ???? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ???? |
| Ogob595 Orectogyrus oberthuri | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |
| Ogse521 Orectogyrus sedilloti | 1111 | 14000 | -0201 | 11211 | 01101 | 01001 | 111-0 | 01000 | -000 |
| Ogve522 Orectogyrus | 1111 | 14000 | -0211 | 11211 | 01101 | 01001 | 111-0 | 01000 | -020 |


| vestitus |  | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $112-0$ | 01000 | -010 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Gesp626 Gyretes sp | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $111-0$ | 01000 | -010 |  |
| Gesp619 Gyretes sp | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $112-0$ | 01000 | -010 |  |
| Gesp616 Gyretes <br> quadrispinosus | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $111-0$ | $00-00$ | -010 |  |
| Gesp624 Gyretes sp | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $111-0$ | $00-00$ | -010 |  |
| Geir470 Gyretes iricolor | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $111-0$ | $00-00$ | -010 |  |
| Gysp686 Gyretes <br> boucardi | 1111 | 11001 | -0101 | 11211 | 01101 | 01001 | $112-0$ | $00-00$ | -010 |  |
| Gysp685 Gyretes |  |  |  |  |  |  |  |  |  |  |
| acutangulus |  |  |  |  |  |  |  |  |  |  |



| Hacr Haliplus cretaceus | ?????? | ???00 | ???? 1 | -100- | 1--00 | 00000 | 0000? | 0-??0 | 10 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Colg Coptoclava Iongipoda | 0???? 1 | ??000 | 00000 | 0100- | 00-00 | 00000 | 00002 | 0-000 | 01 |
| Lilo Liadytes longus | 0???? 0 | 0?00? | 0? 0 ? | 0100- | 00-00 | ????? | ????0 | 0-0?0 | 00 |
| Mdrh Mesodytes rhantoides | 0????? | ??001 | 0-000 | 0000- | 00-00 | 10001 | 0000? | 0-??1 | 00 |
| Spal472 Spanglerogyrus albiventris | 000-00 | 00010 | 00000 | 0010- | 00-10 | 11000 | 00001 | 10000 | 01 |
| Hsmi596 Heterogyrus milloti | 011010 | 02110 | 11000 | 00110 | 01110 | 10010 | 10003 | 10000 | 02 |
| Aoal525 Aulonogyrus alternatus | 012020 | $0 ? 110$ | 11000 | 00010 | 02010 | 10001 | 00003 | 10100 | 02 |
| Aoma523 Aulonogyrus marginatus | ?????? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ?? |
| Aocr519 Aulonogyrus cristatus | ?????? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ?? |
| Aoca604 Aulonogyrus carinipennis | 012020 | 00110 | 11000 | 00010 | 02110 | 10001 | 00003 | 10100 | 02 |
| Aosp569 Aulonogyrus sp | 012020 | 0? 110 | 11000 | 00010 | 02010 | 10001 | 00003 | 10100 | 02 |
| Aobe493 Aulonogyrus bedeli | 012010 | 00110 | 11000 | 00011 | 02010 | 10001 | 00003 | 10100 | 02 |
| Aoca540 Aulonogyrus caffer | 012020 | 00110 | 11000 | 0000- | 02010 | 10001 | 00003 | 10100 | 02 |
| Aost469 Aulonogyrus striatus | 012020 | 00110 | 11000 | 00010 | 02010 | 10001 | 00003 | 10100 | 02 |
| Aogo529 Aulonogyrus goudoti | 012020 | 00110 | 11000 | 0000- | 02110 | 10001 | 00003 | 10100 | 02 |
| Berg1 Metagyrinus sinensis | ?????0 | ??110 | 1?0?0 | 10010 | $021 ? 0$ | 10001 | 00003 | $1 ? 10$ ? | ?2 |
| Gyig598 Gyrinus ignitus | 012010 | 00110 | 11010 | 0000- | 02010 | 10001 | 00003 | 10100 | 02 |
| Aost687 Aulonogyrus strigosus | 012020 | 00110 | 11000 | 00010 | 02010 | 10001 | 00003 | 10100 | 02 |
| Gymi526 Gyrinus minutus | 012010 | 00110 | 11000 | 0000- | 02010 | 10001 | 00003 | 10100 | 02 |
| Gymd597 Gyrinus madagascariensis | 012010 | 00110 | 11010 | 0000- | 02010 | 00000 | 00003 | 10100 | 02 |
| Gyna539 Gyrinus natalensis | 012010 | 00110 | 11000 | 0000- | 02010 | 10001 | 00003 | 10100 | 02 |
| Gypl495 Gyrinus plicifer | 012010 | 00110 | 11010 | 0000- | 02010 | 00000 | 00003 | 10100 | 02 |
| Gygi496 Gyrinus gibber | 012010 | 00110 | 11010 | 0000- | 02010 | 00000 | 00003 | 10100 | 02 |
| Gyel494 Gyrinus elevatus | 012010 | 00110 | 11010 | 0000- | 02010 | 00000 | 00003 | 10100 | 02 |
| Gysp492 Gyrinus sp | 012010 | 00110 | 11000 | 0000- | 02010 | 10011 | 00003 | 10200 | 02 |
| Gysp628 Gyrinus amazonicus | 012010 | 00110 | 11000 | 0000- | 02010 | 00000 | 00003 | 10200 | 02 |
| Difv672 Dineutus favareli | 012010 | 02110 | 11001 | -000- | 01200 | 11120 | 01003 | 11120 | 02 |
| Disp507 Dineutus pectoralis | 011020 | 02110 | 11001 | -000- | 01200 | 10010 | 10003 | 11120 | 02 |
| Disu484 Dineutus subspinosus | 012010 | 02110 | 11001 | -000- | 01200 | 10020 | 11003 | 11120 | 02 |
| Diso605 Dineutus | 112020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |


| solitarius |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Dipx515 Dineutus proximus | 012020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Disp576 Dineutus striatus | 012020 | 00110 | 11001 | -000- | 01200 | 11021 | 01003 | 11120 | 02 |
| Dici474 Dineutus ciliatus | 012020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Didi473 Dineutus discolor | 112020 | 00110 | 11001 | -000- | 01200 | 11000 | 00003 | 11120 | 02 |
| Disn516 Dineutus sinuosipennis | 012020 | 00110 | 11001 | -000- | 01200 | 11100 | 00003 | 11120 | 02 |
| Disp481 Dineutus aereus | 112020 | 00110 | 11001 | -000- | 01200 | 11000 | 01003 | 11120 | 02 |
| Disu505 Dineutus sublineatus | 112020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Diin482 Dineutus indicus | 112020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Disp577 Dineutus micans | 012020 | 00110 | 11001 | -000- | 01200 | 10000 | 01103 | 11120 | 02 |
| Ayau483 Macrogyrus oblongus | 011021 | 12110 | 11000 | 00011 | 01200 | 11110 | 00013 | 11100 | 02 |
| Aygo501 Macrogyrus gouldi | 011011 | 12110 | 11000 | 00010 | 01210 | 11120 | 00013 | 11100 | 02 |
| Ayan502 Macrogyrus australis | 011011 | 11110 | 11000 | 0000- | 01200 | 11120 | 00013 | 11100 | 02 |
| Aysp506 Macrogyrus albertisi | 011011 | 12110 | 11000 | 00010 | 01200 | 11120 | 00013 | 11100 | 02 |
| Ehas646 Enhydrus atratus | 001001 | 00110 | 11000 | 00010 | 01100 | 10001 | 00003 | 11100 | 02 |
| Adsp648 Andogyrus zimmermanni | 012001 | 10110 | 11000 | 0000- | 00-00 | 00000 | 00003 | 11100 | 02 |
| Prte497 Porrorhynchus marginatus | 002121 | 02110 | 11001 | -0010 | 01200 | 11110 | 01103 | 11110 | 02 |
| Ogca666 Orectogyrus camerunensis | 011021 | 02110 | 11000 | 00110 | 00-10 | 11021 | 00003 | 10100 | 02 |
| Gesp615 Gyretes sericeus | 011001 | 00110 | 11001 | -020- | 00-00 | 10020 | 10003 | 11100 | 02 |
| Ogsp566 Orectogyrus noctuabundis | 001001 | 00110 | 11000 | 00110 | 00-00 | 10001 | 00003 | 10100 | 02 |
| Orvi527 Orectochilus villosus | 001001 | 00110 | 11000 | 0020- | 00-10 | 00000 | 00003 | 10100 | 02 |
| Berg2 Orectochilus bellieri | 001001 | 00110 | 11000 | 0020- | 00-10 | 00000 | 00003 | 10100 | 02 |
| Ogcy520 Orectogyrus cyanicollis | 011021 | 02110 | 11000 | 00110 | 00-10 | 10010 | 00003 | 10100 | 02 |
| Ogha600 Orectogyrus hastatus | 011021 | 01110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Orpr487 Patrus productus | 011001 | 01110 | 11001 | -0110 | 00-00 | 10021 | 00003 | 11100 | 02 |
| Ordi488 Patrus discifer | 011001 | 00110 | 11000 | 00110 | 00-00 | 10011 | 00003 | 11200 | 02 |
| Orsp499 Patrus sp | 001011 | 02110 | 11000 | 00111 | 00-00 | 10010 | 10003 | 11200 | 02 |
| Oran486 Patrus andamanicus | 011001 | 00110 | 11001 | -0110 | 00-00 | 10021 | 00003 | 11100 | 02 |
| Orvo489 Patrus volubilis | 011001 | 00110 | 11000 | 00110 | 00-00 | 10021 | 00003 | 11100 | 02 |
| Orsp500 Patrus sp | 011001 | 00110 | 11000 | 00111 | 00-00 | 10011 | 00003 | 11200 | 02 |


| Orsp677 Patrus sp | 011011 | 01110 | 11000 | 00110 | 00-00 | 10001 | 00003 | 11200 | 02 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Ogar669 Orectogyrus argenteovittatus | 001011 | 00110 | 11000 | 00110 | 00-00 | 10000 | 10003 | 10100 | 02 |
| Ogsj671 Orectogyrus sjostedti | 011021 | 02110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Ogpi665 Orectogyrus pictimanus | 011011 | 02110 | 11000 | 00110 | 00-00 | 10010 | 10003 | 10100 | 02 |
| Ogde524 Orectogyrus dedalus | 001011 | 00110 | 11000 | 00110 | 00-00 | 111-0 | 00003 | 10100 | 02 |
| Ogdi491 Orectogyrus discors | 001011 | 01110 | 11000 | 00110 | 00-00 | 10010 | 10003 | 10100 | 02 |
| Ogos667 Orectogyrus oscari | 001001 | 01110 | 11000 | 00110 | 00-00 | 10020 | 00003 | 10100 | 02 |
| Ogpl662 Orectogyrus prolongatus | 001011 | 0?110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Ogdy664 Orectogyrus demeryi | 011011 | 01110 | 11000 | 00110 | 00-00 | 00000 | 00003 | 10100 | 02 |
| Ogsp565 Orectogyrus specularis | 011021 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogms670 Orectogyrus masculinus | 011011 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogsp663 Orectogyrus sp | 011021 | 0? 110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogmd601 Orectogyrus madagascariensis | 011021 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogdo517 Orectogyrus dorsiger | 011021 | 02110 | 11000 | 00110 | 00-10 | 10020 | 00003 | 10100 | 02 |
| Ogsp490 Orectogyrus posticalis | ?????? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ?? |
| Ogsp564 Orectogyrus wittei | 011021 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogbd661 Orectogyrus bedeli | 011021 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogsp567 Orectogyrus specularis | 011021 | 02110 | 11000 | 00110 | 00-10 | 10021 | 00003 | 10100 | 02 |
| Ogsp668 Orectogyrus sp | ?????? | ????? | ????? | ????? | ????? | ????? | ????? | ????? | ?? |
| Ogob595 Orectogyrus oberthuri | 001021 | 02110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Ogse521 Orectogyrus sedilloti | 001011 | 01110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Ogve522 Orectogyrus vestitus | 001021 | 0?110 | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Gesp626 Gyretes sp | 011001 | 00110 | 11001 | -0110 | 00-00 | 10011 | 00003 | 11300 | 02 |
| Gesp619 Gyretes sp | 011001 | 0? 110 | 11001 | -0110 | 00-00 | 10011 | 00003 | 11100 | 02 |
| Gesp616 Gyretes quadrispinosus | 011001 | 02110 | 11001 | -0110 | 00-00 | 11020 | 00003 | 11200 | 02 |
| Gesp624 Gyretes sp | 011001 | 00110 | 11001 | -010- | 00-00 | 00000 | 00003 | 11100 | 02 |
| Geir470 Gyretes iricolor | 011001 | 00110 | 11001 | -010- | 00-10 | 00000 | 00003 | 11100 | 02 |
| Gysp686 Gyretes boucardi | 011001 | 00110 | 11001 | -010- | 00-00 | 10001 | 00003 | 11100 | 02 |
| Gysp685 Gyretes acutangulus | 011001 | 00110 | 11001 | -010- | 00-00 | 10001 | 00003 | 11200 | 02 |
| Gesp617 Gyretes sp | 011001 | 00110 | 11001 | -010- | 00-00 | 10001 | 00003 | 11200 | 02 |


| Gesp614 Gyretes sp | 011001 | 00110 | 11001 | -010- | 00-00 | 10001 | 00003 | 11200 | 02 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Ayhw887 Macrogyrus howittii | 012001 | 11110 | 11000 | 0000- | 01200 | 10011 | 00013 | 11300 | 02 |
| Ayre912 Macrogyrus reichei | 001021 | 11110 | 11000 | 00011 | 01200 | 10010 | 01013 | 11100 | 02 |
| Ayst882 Macrogyrus striolatus | 011011 | 11110 | 11000 | 00011 | 01200 | 10011 | 00013 | 11100 | 02 |
| Adsr886 Andogyrus seriatopunctatus | 012001 | 10110 | 11000 | 0000- | 01000 | 10011 | 00003 | 11100 | 02 |
| Adco828 Andogyrus colombicus | 012001 | 10110 | 11000 | 0000- | 01200 | 00000 | 00003 | 11100 | 02 |
| AyCs829 Macrogyrus toxopeusi | 011001 | 10110 | 11000 | 0000- | 01200 | 11110 | 00003 | 11100 | 02 |
| AyCs841 Macrogyrus purpurascens | 012001 | $1 ? 110$ | 11000 | 0000- | 01200 | 11110 | 00003 | 11200 | 02 |
| Aysp863 Macrogyrus sumbawae | 011011 | 11110 | 11000 | 0000- | 01200 | 11011 | 00013 | 11100 | 02 |
| AyTs834 Macrogyrus sp | 011011 | 1 ? 110 | 11000 | 00011 | 01200 | 10110 | 00013 | 11100 | 02 |
| AyTs833 Macrogyrus sp | 011011 | 1 ? 110 | 11000 | 00011 | 01200 | 11110 | 00013 | 11200 | 02 |
| AyCs831 Macrogyrus sp | 012001 | 10110 | 11000 | 0000- | 01200 | 11110 | 00003 | 11200 | 02 |
| DiDf915 Dineutus fulgidus | 111020 | 02110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| DiDn865 Dineutus n sp | 111020 | 02110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| DiRt908 Dineutus tetracanthus | 011020 | 02110 | 11001 | -000- | 01200 | 10120 | 00003 | 11120 | 02 |
| Dilo818 Dineutus longimanus | 111020 | 00110 | 11001 | -000- | 01200 | 11100 | 01003 | 11120 | 02 |
| DiCp918 Dineutus pagdeni | 111020 | 01110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Dica821 Dineutus carolinus | 112020 | 00110 | 11001 | -000- | 01200 | 10000 | 01003 | 11120 | 02 |
| Dias819 Dineutus assimilis | 012020 | 00110 | 11001 | -000- | 01200 | 11000 | 00003 | 11120 | 02 |
| Diro913 Dineutus robertsi | 012020 | 00110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Prla852 Porrorhynchus landaisi | 002121 | 02110 | 11001 | -0010 | 00-00 | 11120 | 00103 | 11110 | 02 |
| Ehsu856 Enhydrus sulcatus | 000011 | 00110 | 11000 | 00011 | 01100 | 00000 | 00003 | 11100 | 02 |
| DiCf916 Dineutus fairmairei | 011020 | 01110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Diau911 Dineutus australis | 012020 | 02110 | 11001 | -000- | 01200 | 10011 | 01003 | 11120 | 02 |
| DiMp917 Dineutus priscus | 011020 | 02110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| DiMm919 Dineutus macrochirus | 011020 | 02110 | 11001 | -000- | 01200 | 00000 | 00003 | 11120 | 02 |
| Ogor901 Orectogyrs ornaticollis | 011011 | $0 ? 110$ | 11000 | 00110 | 00-10 | 10011 | 00003 | 10100 | 02 |
| Oghe900 Orectogyrus heros | 011021 | 02110 | 11000 | 00110 | 00-10 | 10010 | 10003 | 10100 | 02 |
| Pasp897 Patrus sp | 001001 | $0 ? 110$ | 11000 | 00110 | 00-00 | 10001 | 00003 | 10100 | 02 |


| Pasp896 Patrus sp | 001001 | 00110 | 11000 | 00110 | 00-00 | 10001 | 00003 | 10100 | 02 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Pasp898 Patrus sp | 001001 | 0? 110 | 11000 | 00110 | 00-00 | 10021 | 00003 | 10100 | 02 |
| Gysp840 Gyrinus sericeolimbatus | 012010 | 00110 | 11010 | 0000- | 02010 | 10001 | 00003 | 10100 | 02 |
| Gysp839 Gyrinus dimorphus | 012010 | 00110 | 11010 | 0000- | 02010 | 10001 | 00003 | 10200 | 02 |
| Gysp837 Gyrinus maculiventris | 012010 | 00110 | 11010 | 0000- | 02010 | 00000 | 00003 | 10100 | 02 |
| Agmo Angarogyrus mongolicus | ?????? | ????? | ????0 | 00?0- | 00-11 | 11000 | 00001 | ????? | ?? |
| Agmi Angarogyrus minimus | ?????? | ????? | ????? | ??10- | 00-11 | 11000 | 0000? | ????? | ?? |
| Basa Baissogyrus savilovi | ?????? | ??110 | 1110? | ????? | ????? | ????? | ????? | 10??0 | 02 |
| Mgan Mesogyrus antiquus | ?????? | ??110 | 11?00 | 00? 10 | 01110 | 10010 | 1000? | 10??0 | 02 |
| Mgst Mesogyrus striatus | ?????? | ????? | ????? | ????? | ????? | 1???1 | 0???? | 10??0 | 02 |
| Crzh Cretotortor zherichini | ?????? | ????? | ????0 | 0??10 | 01110 | 10000 | 1000? | 10??? | ?? |
| Meam Mesodineutes amurensis | ?????? | ??110 | 11??? | ?0?0- | 01000 | 00000 | 0000? | 11??0 | 02 |
| Gegi Gyretes giganteus | ?????? | ????? | ????0 | 00?10 | 00-10 | 10011 | 0000? | ????? | ?? |
| Miin Miodineutes insignis | ?????? | ????? | ????? | ????? | 00-?0 | 10001 | 0000? | ????? | ?? |
| Cresp Cretotortor sp | ?????? | ????? | ????0 | $00 ? 10$ | 01010 | 10010 | 1000? | ????? | ?? |
|  | 87 | 95 | 100 | 105 | 110 | 115 | 120 |  |  |
|  | 1 | I | \| | 1 | 1 | 1 | - |  |  |
| Trlati Triaplus laticoxa | 0-000100 | 0?00- | -10?2 | 0???? | ????? | ????? | ? |  |  |
| Nocl503 Noterus clavicornis | 0-000112 | 0002- | -0002 | 0?0-- | -- | -12- | - |  |  |
| Haho504 Hygrobia hermanni | 0-000000 | 0002- | -0002 | 011-- | ----- | -12-- | - |  |  |
| Mabi2 Matus bicarinatus | 0-200102 | 1012- | -0002 | 010-- | ----- | -12-- | - |  |  |
| Plde130 Platynectes decemaculatus | 0-200102 | 1012- | -0002 | 012-- | ----- | -12-- | - |  |  |
| Lcla91 Lancetes lanceolatus | 0-200102 | 1012- | -0002 | 010-- | ----- | -12-- | - |  |  |
| Hali Haliplus lineatocollis | 0-000000 | 0000- | -0002 | 010-- | ----- | -12-- | - |  |  |
| Hacr Haliplus cretaceus | 0-0?0000 | 0000- | -?002 | 01??? | ????? | ????? | ? |  |  |
| Colg Coptoclava longipoda | 00200003 | 0002- | -0002 | 01??? | ????? | ????? | ? |  |  |
| Lilo Liadytes longus | 0-000100 | 0012- | ????? | ????? | ????? | ?12-- | ? |  |  |
| Mdrh Mesodytes rhantoides | 0-200100 | 1012? | ??0?2 | 01??? | ????? | ?12-- | ? |  |  |
| Spal472 Spanglerogyrus albiventris | 00000001 | 01010 | 10000 | 00001 | 00000 | 00000 | 0 |  |  |
| Hsmi596 Heterogyrus milloti | 10101004 | 00011 | 10001 | 00001 | 10000 | 00100 | 0 |  |  |
| Aoal525 Aulonogyrus alternatus | 20200004 | 10011 | 10002 | 20101 | 00000 | 00201 | ? |  |  |


| Aoma523 Aulonogyrus marginatus | ???????? | ????? | ????? | ????? | ????? | ????? | ? |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Aocr519 Aulonogyrus cristatus | ???????? | ????? | ????? | ????? | ????? | ????? | ? |
| Aoca604 Aulonogyrus carinipennis | 20200004 | 10011 | 10002 | 20001 | 00000 | 00201 | ? |
| Aosp569 Aulonogyrus sp | 20200004 | 10011 | 10002 | 20001 | 00000 | 00201 | ? |
| Aobe493 Aulonogyrus bedeli | 20200004 | 10011 | 10002 | 20101 | 00000 | 00201 | ? |
| Aoca540 Aulonogyrus caffer | 20200004 | 10011 | 10002 | 20101 | 00000 | 00201 | ? |
| Aost469 Aulonogyrus striatus | 20200004 | 10011 | 10002 | 20201 | 00000 | 00201 | ? |
| Aogo529 Aulonogyrus goudoti | 20200004 | 11011 | 10002 | 20201 | 00000 | 00201 | 0 |
| Berg1 Metagyrinus sinensis | ??????04 | 1?0?? | ??002 | 10?01 | 00000 | 0020? | ? |
| Gyig598 Gyrinus ignitus | 20210004 | 11011 | 00002 | 00201 | 00000 | 00201 | 0 |
| Aost687 Aulonogyrus strigosus | 20200004 | 11011 | 10002 | 20201 | 00000 | 00201 | 0 |
| Gymi526 Gyrinus minutus | 20200004 | 11011 | 10002 | 00101 | 00000 | 00201 | ? |
| Gymd597 Gyrinus madagascariensis | 20210004 | 11011 | 00002 | 00201 | 00000 | 00201 | 0 |
| Gyna539 Gyrinus natalensis | 20210004 | 10011 | 00002 | 00201 | 00000 | 00201 | ? |
| Gypl495 Gyrinus plicifer | 20210004 | 11011 | 10002 | 00201 | 00000 | 00201 | 0 |
| Gygi496 Gyrinus gibber | 20210004 | 11011 | 10002 | 00201 | 00000 | 00201 | 0 |
| Gyel494 Gyrinus elevatus | 20210004 | 11011 | 10002 | 00201 | 00000 | 00201 | ? |
| Gysp492 Gyrinus sp | 20200004 | 11011 | 10002 | 00001 | 00000 | 00201 | ? |
| Gysp628 Gyrinus amazonicus | 20200004 | 11011 | 00002 | 00001 | 00000 | 00201 | ? |
| Difv672 Dineutus favareli | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Disp507 Dineutus pectoralis | 10201004 | 00010 | 10002 | 00110 | 01000 | 10200 | ? |
| Disu484 Dineutus subspinosus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Diso605 Dineutus solitarius | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Dipx515 Dineutus proximus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |
| Disp576 Dineutus striatus | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Dici474 Dineutus ciliatus | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | 1 |
| Didi473 Dineutus discolor | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |
| Disn516 Dineutus sinuosipennis | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |
| Disp481 Dineutus aereus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Disu505 Dineutus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |


| sublineatus |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Diin482 Dineutus indicus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Disp577 Dineutus micans | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Ayau483 Macrogyrus oblongus | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | 1 |
| Aygo501 Macrogyrus gouldi | 10001004 | 01010 | 10002 | 00010 | 10000 | 10200 | ? |
| Ayan502 Macrogyrus australis | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| Aysp506 Macrogyrus albertisi | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| Ehas646 Enhydrus atratus | 11201004 | 00010 | 11002 | 00210 | 10000 | 10200 | 1 |
| Adsp648 Andogyrus zimmermanni | 10201004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| Prte497 Porrorhynchus marginatus | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | 1 |
| Ogca666 Orectogyrus camerunensis | 20202004 | 11011 | 10112 | 00010 | 10202 | 10210 | ? |
| Gesp615 Gyretes sericeus | 20202004 | 11011 | 00112 | 00200 | 10110 | 10200 | ? |
| Ogsp566 Orectogyrus noctuabundis | 20202004 | 11011 | 00112 | 00010 | 10101 | 10210 | ? |
| Orvi527 Orectochilus villosus | 20202004 | 10011 | 00112 | 00010 | 10101 | 10200 | 1 |
| Berg2 Orectochilus bellieri | 20202004 | 10011 | 00112 | 000?? | ????? | ????? | ? |
| Ogcy520 Orectogyrus cyanicollis | 20202004 | 11011 | 00112 | 00210 | 10201 | 10210 | ? |
| Ogha600 Orectogyrus hastatus | 20202004 | 11011 | 00112 | 00110 | 10201 | 10210 | ? |
| Orpr487 Patrus productus | 20202004 | 11011 | 20112 | 00200 | 10010 | 10200 | ? |
| Ordi488 Patrus discifer | 20200004 | 11011 | 00112 | 00100 | 10010 | 10200 | ? |
| Orsp499 Patrus sp | 20200004 | 01011 | 00112 | 00200 | 10210 | 10200 | ? |
| Oran486 Patrus andamanicus | 20202004 | 11011 | 00112 | 00200 | 10210 | 10200 | ? |
| Orvo489 Patrus volubilis | 20202004 | 01011 | 00112 | 00200 | 10210 | 10200 | ? |
| Orsp500 Patrus sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Orsp677 Patrus sp | 20202004 | 01011 | 00112 | 00200 | 10??? | 10200 | ? |
| Ogar669 Orectogyrus argenteovittatus | 20202004 | 11011 | 00112 | 00010 | 10201 | 10210 | ? |
| Ogsj671 Orectogyrus sjostedti | 20202004 | 11011 | 10112 | 00110 | 10202 | 10210 | ? |
| Ogpi665 Orectogyrus pictimanus | 20202004 | 11011 | 00112 | 00010 | 10201 | 10210 | ? |
| Ogde524 Orectogyrus dedalus | 20202004 | 11011 | 00112 | 00010 | 10101 | 10210 | ? |
| Ogdi491 Orectogyrus discors | 20202004 | 11011 | 20112 | 00010 | 10101 | 10210 | ? |
| Ogos667 Orectogyrus | 20202004 | 11011 | 20112 | 00010 | 10202 | 10210 | ? |


| oscari |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Ogpl662 Orectogyrus prolongatus | 20202004 | 11011 | 20112 | 00010 | 10??? | 10210 | ? |
| Ogdy664 Orectogyrus demeryi | 20202004 | 11011 | 20112 | 00010 | 10202 | 10210 | ? |
| Ogsp565 Orectogyrus specularis | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | ? |
| Ogms670 Orectogyrus masculinus | 20202004 | 11011 | 20112 | 00010 | 10202 | 10210 | ? |
| Ogsp663 Orectogyrus sp | 20202004 | 11011 | 20112 | 00010 | 10??? | 10210 | ? |
| Ogmd601 Orectogyrus madagascariensis | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | 1 |
| Ogdo517 Orectogyrus dorsiger | 20202004 | 11011 | 20112 | 00210 | 10202 | 10210 | ? |
| Ogsp490 Orectogyrus posticalis | ???????? | ????? | ????? | ????? | ????? | ????? | ? |
| Ogsp564 Orectogyrus wittei | 20202004 | 11011 | 20112 | 00010 | 10202 | 10210 | ? |
| Ogbd661 Orectogyrus bedeli | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | ? |
| Ogsp567 Orectogyrus specularis | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | ? |
| Ogsp668 Orectogyrus sp | ???????? | ????? | ????? | ????? | ????? | ????? | ? |
| Ogob595 Orectogyrus oberthuri | 20202004 | 11011 | 20112 | 00010 | 10202 | 10210 | ? |
| Ogse521 Orectogyrus sedilloti | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | 1 |
| Ogve522 Orectogyrus vestitus | 20202004 | 11011 | 10112 | 00110 | 10202 | 10210 | ? |
| Gesp626 Gyretes sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Gesp619 Gyretes sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Gesp616 Gyretes quadrispinosus | 20202004 | 11011 | 20112 | 00200 | 10110 | 10200 | ? |
| Gesp624 Gyretes sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Geir470 Gyretes iricolor | 20202004 | 11011 | 00112 | 00200 | 10220 | 10200 | 0 |
| Gysp686 Gyretes boucardi | 20202004 | 11011 | 00112 | 00200 | 10220 | 10200 | ? |
| Gysp685 Gyretes acutangulus | 20202004 | 11011 | 00112 | 00200 | 10220 | 10200 | ? |
| Gesp617 Gyretes sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Gesp614 Gyretes sp | 20202004 | 11011 | 00112 | 00200 | 10??? | 10200 | ? |
| Ayhw887 Macrogyrus howittii | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| Ayre912 Macrogyrus reichei | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | 1 |
| Ayst882 Macrogyrus striolatus | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | 1 |
| Adsr886 Andogyrus seriatopunctatus | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| Adco828 Andogyrus colombicus | 10201004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| AyCs829 Macrogyrus | 10201004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |


| toxopeusi |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| AyCs841 Macrogyrus purpurascens | 10201004 | 01010 | 10002 | 00010 | 10000 | 10200 | ? |
| Aysp863 Macrogyrus sumbawae | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| AyTs834 Macrogyrus sp | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| AyTs833 Macrogyrus sp | 10001004 | 01010 | 10002 | 00210 | 10000 | 10200 | ? |
| AyCs831 Macrogyrus sp | 10201004 | 01010 | 10002 | 00010 | 10000 | 10200 | ? |
| DiDf915 Dineutus fulgidus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| DiDn865 Dineutus n sp | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| DiRt908 Dineutus tetracanthus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Dilo818 Dineutus longimanus | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| DiCp918 Dineutus pagdeni | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Dica821 Dineutus carolinus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |
| Dias819 Dineutus assimilis | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | 1 |
| Diro913 Dineutus robertsi | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Prla852 Porrorhynchus landaisi | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Ehsu856 Enhydrus sulcatus | 11201004 | 00010 | 11002 | 00210 | 10000 | 10200 | ? |
| DiCf916 Dineutus fairmairei | 10201004 | 00010 | 10002 | 00010 | 01000 | 10200 | ? |
| Diau911 Dineutus australis | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| DiMp917 Dineutus priscus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| DiMm919 Dineutus macrochirus | 10201004 | 00010 | 10002 | 00210 | 01000 | 10200 | ? |
| Ogor901 Orectogyrs ornaticollis | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | 1 |
| Oghe900 Orectogyrus heros | 20202004 | 11011 | 20112 | 00110 | 10202 | 10210 | 1 |
| Pasp897 Patrus sp | 20202004 | 11011 | 20112 | 0020? | ????? | ????? | 0 |
| Pasp896 Patrus sp | 20202004 | 10011 | 20112 | 0020? | ????? | ????? | 0 |
| Pasp898 Patrus sp | 20202004 | 11011 | 20112 | 0010? | ????? | ????? | 0 |
| Gysp840 Gyrinus sericeolimbatus | 202?0004 | 11011 | 10002 | 00201 | 00000 | 00201 | ? |
| Gysp839 Gyrinus dimorphus | 20210004 | 11011 | 10002 | 00201 | 00000 | 00201 | ? |
| Gysp837 Gyrinus maculiventris | 20210004 | 11011 | 10002 | 00101 | 00000 | 00201 | ? |
| Agmo Angarogyrus mongolicus | ???????1 | ????? | ??0?? | ?0??? | ????? | ????? | ? |
| Agmi Angarogyrus minimus | ???????? | ????? | ??0?? | ????? | ????? | ????? | ? |


| Basa Baissogyrus savilovi | 101?100? | 00011 | 1000? | ??0?? | ????? | ????? | ? |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Mgan Mesogyrus antiquus | 1010100? | 00011 | 1000? | ??0?? | ????? | ????? | ? |
| Mgst Mesogyrus striatus | 10?? 100 ? | 0?011 | 1?0?? | ????? | ????? | ????? | ? |
| Crzh Cretotortor zherichini | ???????? | ????? | ????? | ????? | ????? | ????? | ? |
| Meam Mesodineutes amurensis | 1000100? | 00010 | 10002 | 00??? | ????? | ????? | ? |
| Gegi Gyretes giganteus | ???????? | ????? | ??1?? | 00??? | ????? | ????? | ? |
| Miin Miodineutes insignis | ???????? | ????? | ??1?? | 00??? | ????? | ????? | ? |
| Cresp Cretotortor sp | ???????? | ????? | ??0?? | ?0??? | ????? | ????? | ? |

Supplementary Data Table 2. Additional taxa added to dataset.

| Genus | species | voucher \# | Collection data | COI | COII | 12S | H3 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Haliplus | lineatocollis | Hali | GenBank | X | - | X | X |
| Macrogyrus | howittii | Ayhw887 | AUSTRALIA: Tasmania. Franklin Beach, Lake St. Clair. 10.i. 2015 CHS Watts. MSBA AUSTRALIA: VIC. Glenelg River nr. Dergholm, - | X | - | X | X |
| Macrogyrus | reichei | Ayre912 | 37.36686, 141.2428. $75 \mathrm{~m} .13 . \mathrm{i} .2015$. Leg <br> G.Gustafson. MSBA <br> AUSTRALIA: NSW. Megalong Valley, -36.65629, <br> $150.27377,861 \mathrm{~m} .04 . i .2015$. leg. G.Gustafson | X | X | X | - |
| Macrogyrus | striolatus seriatopunctat | Ayst882 | Forested strm GTG01042015A. MSBA | X | X | X | X |
| Andogyrus | us | Adsr886 | Argentina. MSBA <br> VENEZUELA: Merida State $8^{\circ} 38.006^{\prime} \mathrm{N}$, $71^{\circ} 09.762^{\prime} \mathrm{W}, 2037 \mathrm{~m}$ Monte Zerpa area; 20.vii. 2009 leg. Short, Sites, Gustafson, Camacho; stream margin/pools VZ09-0720-01A/L-1098 ABTC-01476. | X | X | X | X |
| Andogyrus | colombicus | Adco828 | MSBA | X | X | X | X |
| Macrogyrus | toxopeusi | AyCs829 | INDONESIA: Papua: Poga $3^{\circ} 48.382^{\prime} \mathrm{S}, 138^{\circ} 34.780^{\prime} \mathrm{E}$ 2285-2330 m. ZSM <br> PAPUA NEW GUINEA: Morobe Prov. Pindiu. <br> $6^{\circ} 27.147^{\prime} \mathrm{S} 147^{\circ} 29.574^{\prime} \mathrm{E}, 1470 \mathrm{~m} .12 . x .2009 \mathrm{leg}$. | X | X | X | X |
| Macrogyrus | purpurascens | AyCs841 | Inaho (PNG206). ZSM | X | X | X | X |
| Macrogyrus | sumbawae | Aysp860 | INDONESIA: Sumba: dry forest stream in limestone. $370 \mathrm{~m} 9^{\circ} 49.47 \mathrm{l}^{\prime} \mathrm{S} 120^{\circ} 20.856^{\prime} \mathrm{E}$ (SUA08). ZSM PAPUA NEW GUINEA: E Highlands Prov. | X | X | X | X |
| Macrogyrus | sp nr . <br> blanchardii | AyTs834 | Onerunka, small creek, redsoil rock $6^{\circ} 20.936$ 'S $145^{\circ} 46.874^{\prime} \mathrm{E} 1700 \mathrm{~m} .21 . v .2006$ leg. John \& Balke (PNG71). ZSM | X | X | X | X |
| Macrogyrus | sp | AyTs833 | PAPUA NEW GUINEA: Central Prov. Woitape. $08^{\circ} 31.290^{\prime} \mathrm{S} 147^{\circ} 13.684^{\prime} \mathrm{E} 1700 \mathrm{~m} . \mathrm{i} 2008$ leg. Posman (PNG166). ZSM | X | X | X | X |
| Macrogyrus | sp | AyCs831 | PAPUA NEW GUINEA: Sandaun Prov. Mianmin $4^{\circ} 54.570^{\prime} \mathrm{S} 141^{\circ} 35.490^{\prime} \mathrm{E} 990 \mathrm{~m} .23 . x .2008$. leg. Ibalim (PNG193). ZSM | X | X | X | X |
| Dineutus | fulgidus | DiDf915 | INDONESIA: Sumatra Barat, Solok, Alahan Panjank Road. $1190 \mathrm{~m} .0^{\circ} 56.345^{\prime} \mathrm{S} 100^{\circ} 46.411^{\prime} \mathrm{E}$. ZSM | X | X | X | X |
| Dineutus | n sp | DiDn865 | BALI: Telaga Forest, BLI07. ZSM | X | X | X | X |
| Dineutus | tetracanthus | DiRt908 | PAPUA NEW GUINEA: Madang Prov. Wannang, $5^{\circ} 15.458^{\prime} \mathrm{S} 145^{\circ} 2.389^{\prime} \mathrm{E} 270 \mathrm{~m} .31 . x .2008$. leg. Posman (PNG187) ZSM. <br> DOMINICAN REP.: Pedernales Prov. W of | X | X | X | X |
| Dineutus | longimanus longimanus | Dilo818 | Pedernales on rd. to border with Haiti; roadside sweeping in dry for. \& sec. veg. 15 May $201018.154^{\circ}$ $-71.7582^{\circ}$ colr. G. J. Svenson. MSBA | X | X | X | X |



Depository abbreviations:
MSBA: Museum of Southwestern Biology, University of New Mexico, Albuquerque, NM, USA.
ZSM: Zoologische Staatssammlung München, Munich, Germany

Supplementary Data Table 2. Fossil taxa examined.

| group | Genus | species | Age Ma | Locality of deposit | Depository |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Out- | Triaplus | laticoxa | $221-235$ | Madygen area, Batken Region,Osh Oblast, <br> Kyrgyzstan | PIN |
| Out- | Haliplus | cretaceus | $112-125$ | Bon-Tsan, outcrop 87.8, Bayankhongor <br> Province, Mongolia | PIN |
| Out- | Liadytes | longus | $125-150$ | Daya settlement, Glushkovo Formation, Russian <br> Federation | PIN |
| Out- | Mesodytes | rhantoides | $122-125$ | Yixian Formation, Liutiaogou, China | NIG |
| In- | Angarogyrus | mongolicus | $112-125$ | Gurvan-Eren Formation, Govi-Altai, Mongolia <br> Cheremkhovskaya Formation, Irkutsk, Russian <br> Federation | PIN |

PIN: Borissiak Paleontological Institute, Russian Academy of Sciences, Moscow, Russia
MNHN: Muséum National d'Histoire Naturelle, Paris France
NIG: Nanjing Institute of Geology and Palaeontology, Chinese Academy of Sciences, Nanjing, China
ESSI: Erdwissenschaftliche Sammlung, Swiss Federal Institute of Technology, Zurich, Switzerland







