

AperTO - Archivio Istituzionale Open Access dell'Università di Torino

Combined phylogenetic analysis of two new Afrotropical genera of Onthophagini (Coleoptera, Scarabaeidae)

This is the author's manuscript

Original Citation:

Availability:

This version is available <http://hdl.handle.net/2318/1596529> since 2018-01-18T11:57:14Z

Published version:

DOI:10.1111/zoj.12498

Terms of use:

Open Access

Anyone can freely access the full text of works made available as "Open Access". Works made available under a Creative Commons license can be used according to the terms and conditions of said license. Use of all other works requires consent of the right holder (author or publisher) if not exempted from copyright protection by the applicable law.

(Article begins on next page)

The logo for IRIS Aperto, featuring the text "IRIS Aperto" in white on a red rectangular background.

UNIVERSITÀ
DEGLI STUDI
DI TORINO

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21

This is the author's final version of the contribution published as:

Angela Roggero, Michael Dierkens, Enrico Barbero, Claudia Palestrini, Combined phylogenetic analysis of two new Afrotropical genera of Onthophagini (Coleoptera, Scarabaeidae) *Zoological Journal of the Linnean Society*, 180 (2), 2017, pagg. 298-320, DOI: 10.1111/zoj.12498

The publisher's version is available at:

<https://academic.oup.com/zoolinnea/issue/180/2>

When citing, please refer to the published version.

This full text was downloaded from iris-Aperto: <https://iris.unito.it/>

22 **Combined phylogenetic analysis of two new Afrotropical genera of Onthophagini**
23 **(Coleoptera, Scarabaeidae)**

24

25 ANGELA ROGGERO^{1*}, MICHAEL DIERKENS², ENRICO BARBERO¹, CLAUDIA
26 PALESTRINI¹

27 ¹Department of Life Sciences and Systems Biology, Via Accademia Albertina 13 – I-
28 10123 Torino, ITALY

29 ²rue du Garet 21, F – 69001, Lyon, FRANCE

30

31 *Corresponding author e-mail: angela.roggero@unito.it

32

33 Running title: Phylogeny of Afrotropical Onthophagini

34

35

36 **ABSTRACT**

37 To reveal the relationships of the Afrotropical *Onthophagus* 32nd group, a combined
38 phylogenetic analysis was employed on a matrix of both discrete and continuous
39 morphological characters. The species of the 32nd group do not constitute a
40 homogeneous group, but two distinct and well-isolated clades of generic rank:
41 *Hamonthophagus* **gen. nov.** with five species and *Morettius* **gen. nov.** with two species,
42 one of which was identified as a new taxon and is described here (i.e., *Morettius utete*
43 **sp. nov.**). The *Hamonthophagus* species were characterized by a wide distribution
44 covering the entire geographic range of Afrotropical grasslands, while the *Morettius*
45 species were restricted to two distinct areas in central Africa and east Africa.
46 Geographical data were integrated with the phylogenetic results and processed by
47 dispersal-vicariance analysis, which confirmed for both genera an evolutionary and
48 biogeographic history in which the ancestral range was located in the central eastern
49 African region.

50

51 **ADDITIONAL KEYWORDS:** Biogeography, dung beetles, *Hamonthophagus*,
52 *Morettius*, new species, geometric morphometrics, Scarabaeoidea, systematics

53

54 INTRODUCTION

55 In 1913, d'Orbigny proposed a full synopsis of the Afrotropical Onthophagini on the
56 basis of external features, providing a useful identification tool for this megadiverse
57 tribe. Although his work still remains a milestone in the study of Onthophagini, the
58 French author classification has now been challenged by new methods of systematic
59 and phylogenetic investigation. In the meantime many new species have been described
60 from the Afrotropical region. In particular, the most speciose *Onthophagus* genus has
61 been found to exceed 1,000 species (Tarasov & Solodovnikov, 2011), to which the
62 former subdivision into 32 groups by d'Orbigny (1913) does not always apply. The
63 d'Orbigny's classification has been substantially confirmed in a few instances (such as
64 for the genus *Phalops* Erichson, 1848, Barbero *et al.*, 2003), but in other cases it has
65 been profoundly modified (Moretto, 2009; Tagliaferri *et al.*, 2012), highlighting how
66 the majority of *Onthophagus* species groups may, indeed, not be phylogenetically
67 homogeneous. In this regard, it was recently showed that the present taxonomic position
68 of some species (including part of the 32nd group species) is doubtful, being they less
69 close to the other *Onthophagus* than it is usually considered (Roggero *et al.*, 2016).
70 The 32nd species group was thus here examined to evaluate if the hypothesized
71 separation from *Onthophagus* should be confirmed. The group includes only six
72 Afrotropical coprophagous and often nocturnal species, generally characterized by a
73 wide distribution in open environments such as savannah, grasslands and pastures:
74 *Onthophagus acutus* d'Orbigny, 1908, *O. bituberculatus* (Olivier, 1789), *O. depressus*
75 Harold, 1871, *O. fallax* d'Orbigny, 1913, *O. laceratus* Gerstaecker, 1871, and *O.*
76 *pallens* d'Orbigny, 1908. Some of these species were accidentally introduced at the
77 beginning of the 20th century into North America and Australia (*O. depressus*), and the
78 Antilles (*O. bituberculatus*), where ostensibly they have adapted quite well, without
79 causing problems to the native fauna.
80 Recently, Wirta *et al.* (2008) included *O. depressus* in their phylogenetic analysis of the
81 Malagasy dung beetle fauna, since this species has also been introduced into
82 Madagascar, hypothesizing a close relationship with the endemic *Mimonthophagus*
83 *hinnulus* (Klug, 1832), which is nevertheless markedly different in external and internal
84 morphology, and thus might not be so closely related to *O. depressus*.

85 The species of the 32nd group lack any complex morphological structures on the head or
86 pronotum, unlike the majority of *Onthophagini*, in which evident exoskeletal structures
87 are relatively common (Emlen *et al.*, 2005, 2006; Moczek, 2006). All of the 32nd group
88 species are ostensibly characterized by low sexual dimorphism, which mainly affects
89 the fore tibiae and the pygidium. Also, these species share similar patterns of
90 intraspecific colour variation, ranging from an evenly black to yellow background with
91 more or less extensive black spots.

92 Despite their wide distribution, and a certain degree of individual variability, the
93 taxonomic history of these species is less problematic than that of other *Onthophagus*
94 groups. Only few synonymies are recognized and employed, and even fewer subspecies
95 or varieties have been defined (see the Taxonomic Account below for further details).

96 The aim of our research was to study the relationships among the species of 32nd
97 *Onthophagus* group applying the combined phylogenetic approach to a dataset of
98 discrete and continuous morphological characters. Once the phylogenetic relationships
99 within the group were clarified, the evolutionary and biogeographic patterns of these
100 species were examined to define which speciation processes led to the current
101 biogeographical ranges, and how. Finally, the taxonomic status of the 32nd group was
102 thoroughly reassessed according to the former phylogenetic results to formalize any
103 reclassification at the generic and specific level.

104

105

106

107 **MATERIAL AND METHODS**

108 To explore the relationships among the *Onthophagus* species of the d'Orbigny 32nd
109 group, a combined phylogenetic approach was applied on morphological data (discrete
110 and continuous characters, see below) based on the external and internal features. This
111 method was selected being it extremely versatile. Formerly, the quantitative data could
112 not be employed “just as they were” in phylogenetic analysis, but were discretized
113 during the analysis (Goloboff *et al.*, 2006; Gold *et al.*, 2014). Thus, the recent
114 formalization of the combined approach (Goloboff & Catalano, 2010; Catalano *et al.*,
115 2010) has opened up huge opportunities for the use of extremely diverse characters that
116 were hitherto inapplicable.

117 The assembled dataset included seven ingroup taxa (i.e., the six already-known species,
 118 plus a new species herein described), and one outgroup taxon, *Digitonthophagus*
 119 *bonasus* (Fabricius, 1775).

120

121 **Material examined**

122 We examined more than fifteen hundred specimens that were lent to us by the following
 123 Institutions:

- 124 - BMNH Natural History Museum, London, UK
- 125 - IRSNB Institut Royal des Sciences Naturelles de Belgique, Bruxelles, Belgium
- 126 - LACM Natural History Museum of Los Angeles County, Los Angeles, USA
- 127 - MCST Museo Civico di Storia Naturale, Trieste, Italy
- 128 - MHNL Musée des Confluences, Lyon, France
- 129 - MNCN Museo Nacional de Ciencias Naturales, Madrid, Spain
- 130 - MNHN Muséum National d'Histoire Naturelle, Paris, France
- 131 - NHMW Naturhistorisches Museum, Wien, Austria
- 132 - NMPC Narodni Muzeum v Praze, Prague, Czech Republic
- 133 - TMSA Ditsong National Museum of Natural History, Pretoria, South Africa
- 134 - ZMHB Museum für Naturkunde der Humboldt-Universität, Berlin, Germany
- 135 - ZSM Zoologische Staatssammlung, München, Germany

136 and by the following private collectors: E. Barbero (EBCT - Torino, Italy), I. Bonato
 137 (IBCT - Torino, Italy), T. Branco (TBCP - Porto, Portugal), I. De Dinechin (IDCL -
 138 Lyon, France), M. Dierkens (MDCL - Lyon, France), O. Montreuil (OMCF - Fleury-
 139 les-Aubrais, France), P. Moretto (PMCT - Toulon, France), and P. Walter (PWCM -
 140 Montségur, France).

141

142 **Taxa coding**

143 The species, named from now on according to the taxonomic rearrangement proposed
 144 below (see the Taxonomic Account), were coded as follows: *Hamonthophagus acutus*
 145 as AC, or red colour, *H. bituberculatus* as BI, or blue colour, *H. depressus* as DE, or
 146 orange colour, *H. fallax* as FA, or purple colour, *H. laceratus* as LA, or green colour,
 147 *Morettius pallens* as PA, or teal colour, and *M. utete* as UT, or burgundy colour.

148

149 **Morphological analysis**

150 Various anatomical parts (i.e., head, mouthparts, pronotum, fore legs, elytra, male and
151 female genitalia) were selected to assess inter and intraspecific morphological
152 differences (Barbero *et al.*, 2009, 2011; Roggero *et al.*, 2015). The mouthparts and
153 genitalia of both sexes were dissected and treated following the methods usually
154 employed to prepare slides in Scarabaeoidea (Barbero *et al.*, 2003). Then, images of
155 internal and external structures were captured using a Leica® DFC320 digital camera
156 connected to a stereoscopic dissecting microscope (Leica® Z16Apo).

157 The nomenclature of the anatomical traits adopted in this study follows that used in
158 Palestini (1992), Tarasov & Solodovnikov (2011), and Roggero *et al.* (2015, 2016).
159 The anatomical traits were examined, and a set of distinctive characters (N = 26) was
160 identified and used to build a discrete data matrix. Usually, a large number of features
161 concur in characterizing taxa. Some could be quantified (see below for the novel
162 approach employed here), but others cannot, and must necessarily be treated using a
163 qualitative approach.

164 The geometric morphometric (GM) approach was here employed to evaluate
165 phylogenetic patterns of diversification according to Gold *et al.* (2014). To test inter-
166 and intraspecific shape variation within the 32nd group species, both landmark and
167 semilandmark methods were applied (Fig. S1 - Appendix 1), choosing the best
168 configuration to capture the overall shape variation of the head (19 points), the
169 epipharynx (17 points), the mentum (22 points), the pronotum (11 points), and the right
170 elytron (14 points). Each landmark configuration was sampled as implemented in
171 tpsDig2 v2.27 (Rohlf, 2016a) and tps Util v1.69 (Rohlf, 2016b). The sampled datasets
172 were then separately analyzed by tpsSmall v1.33 (Rohlf, 2016c) and tpsRelw v1.65
173 (Rohlf, 2016d) to evaluate the reciprocal relationships among the species, retaining for
174 further analyses the Procrustes distances matrices (PD), the relative warp values (RWs),
175 and the aligned configurations (AL). For each structure, the scatterplots of the RWs and
176 the minimum spanning trees (MST) were built using NTSYS v2.21 (Rohlf, 2012).

177

178 **Phylogenetic analysis**

179 To clarify the phylogenetic relationships among the *Onthophagus* 32nd group species, a
180 combined data matrix (Table S1 - Appendix 2) was built, merging together discrete and

181 continuous characters (N = 192). The aligned configurations of each anatomical trait
182 were chosen to avoid the use of the principal components (PCs) of shape (i.e., the RWs,
183 see above) as characters of phylogenetic analysis, as stated by Adams *et al.* (2011). The
184 arbitrary value of 1 was added to the quantitative data employed for the phylogenetic
185 analysis, since TNT (Goloboff *et al.*, 2003, 2006, 2008) cannot analyze negative
186 numbers (Smith & Hendricks 2013; Gold *et al.* 2014). The outgroup method was chosen
187 to root the trees, with *Digitonthophagus* as the root following T. Branco (*pers. comm.*)
188 who has hypothesized that the 32nd group is phylogenetically well-separated from the
189 *Onthophagus* groups, and probably closer to other Onthophagini genera. Also in
190 Roggero *et al.* (2016) the species of 32nd group were closer to *Digitonthophagus* and
191 allied taxa than to the other *Onthophagus* species.

192 To estimate the relationships among the species, a phylogenetic analysis was conducted
193 using the combined approach in TNT (Goloboff *et al.*, 2003, 2008), where each
194 morphometric character was used as a continuous numerical variable, and the
195 quantitative and qualitative characters were treated as separate blocks in the linear
196 parsimony analysis (de Bivort *et al.*, 2010, 2012; Clouse *et al.*, 2010). Implicit
197 enumeration, traditional search and new technology search were run as implemented in
198 TNT following Sharkey *et al.* (2012). The TNT script “stats.run” was then used to
199 evaluate the tree statistics. Relative nodal support values were determined using
200 bootstrap, jackknife and symmetrical resampling with 1,000 iterations, as implemented
201 in TNT (Sharkey *et al.*, 2012). The resulting trees were then drawn by FigTree v1.4.2
202 (Rambaut, 2014).

203

204 **Biogeographical analysis**

205 Specific ranges were identified by employing only material herein examined. Each
206 locality was georeferenced, and coordinates were used to build the digital maps of the
207 distribution for each species (see Appendix 3 for the list of the localities) in the GIS
208 environment through QGIS v2.16 (QGIS Development Team, 2016). Collection
209 localities were then grouped together in eight macroareas (Fig. 1) according to the
210 terrestrial ecoregions proposed by Olson *et al.* (2001), and to the phytogeographical
211 areas proposed by White and Leonard (1991). The distribution data of the species were
212 then summarized in a presence/absence matrix that was employed for the dispersal-

213 vicariance analysis as implemented in RASP (Statistical Dispersal-Vicariance Analysis
214 method, Yu *et al.*, 2010a, 2010b), with the maximum number of ancestral areas set
215 equal to 2.

216 The Vicariance Inference Program (VIP, see Arias *et al.*, 2011) was employed to
217 perform the Spatial Analysis of Vicariance, a method focused on the identification of
218 disjoint (i.e., vicariant, or allopatric) distributions related to the formation of new
219 barriers among sister groups instead of finding the ancestral areas, as in the traditional
220 phylogenetic biogeography. In the analysis, sympatric speciations can also be
221 highlighted since they occur whenever the species distributions overlap. In the VIP
222 approach, the node removal is connected with dispersal, identifying any speciation that
223 cannot be explained by the current reconstruction, and no process can be associated with
224 the “geography of the distribution” of these species. In this framework, the
225 georeferenced distribution data were used as spatial information, while the phylogenetic
226 tree from TNT analysis furnished the phylogenetic information required by VIP.

227 According to Ferretti *et al.* (2012), the analysis was performed using a grid of 1.5x1.5,
228 selecting the Von Neumann neighbourhood and a maximum fill of 1. The default
229 parameters of VIP were employed for the heuristic search, with 100,000 iterations, and
230 the Bremer support was then calculated for each vicariant node. The hypothetical
231 (heuristic) barriers among clades were represented on the maps by Voronoi lines (Arias
232 *et al.*, 2011).

233 The results from VIP were then compared to the former RASP results, to test the
234 hypothesized biogeographic history.

235

236

237

238 **RESULTS**

239 **Morphological analysis**

240 The detailed examination of the mentum, genitalia of both sexes, legs, head, pronotum,
241 and elytra led to the identification of 26 qualitative characters (see the Characters List
242 below), but some features could not be properly defined by a descriptive delineation of
243 the characters. The complexity of anatomical shape was often better appraised by a
244 quantitative approach (such as that provided by GM) than by a qualitative one, so the

245 mentum, head, pronotum, and elytra were also examined from a quantitative point of
246 view. The epipharynx (Fig. 2) was instead examined only by the quantitative approach,
247 which can better highlight even the most subtle shape variations.

248 The discrete and continuous data (characters 1-26 and 27-192, respectively, but see the
249 Characters List below) were then used to build the combined matrix for the
250 phylogenetic analysis (Table S1 - Appendix 2).

251

252 For each dataset, the relationships among the species were examined to test the
253 morphological pattern of diversification applying GM methods. For this, the overall
254 shape variation of each structure was studied separately and the amount of specific
255 difference was quantified and employed in the subsequent phylogenetic analysis (linear
256 parsimony, see below). The congruences/divergences of the identified patterns of
257 morphological variation were also explored at the specific and generic levels.

258 Only the plots of the specimens actually employed to build the phylogenetic matrix of
259 the aligned data were shown for each structure (Figs. 3-4). In each dataset, the typical
260 material (if available) was included, but when types could not be found, topotypical
261 material was selected. On each plot, the MST (built from the Procrustes distances
262 matrix) was added to provide a more thorough insight into the differences among the
263 species.

264

265 In the plot of the two first RWs of the head, well-defined groups were identified, and
266 the species showed clearly differentiated patterns. The variance explained by the first
267 two RWs was 84.74% for the head. The outgroup taxon is more similar to the group
268 UT-PA than to any other species, and the five species included in *Hamonthophagus*
269 share similar patterns of shape variation, with LA more closely related to BI than to the
270 group UT-PA. Besides, according to our results, the head is a structure characterized by
271 two distinctive patterns, allowing us to easily separate the genera, but also the species
272 within each genus can be identified. Examining the deformation grids of RW_1 (Fig. 3),
273 the head was clearly more rounded and the notch on the fore margin was absent (or
274 greatly reduced) in OUT and in UT-PA, while the head was more elongated and deeply
275 notched in the *Hamonthophagus* species. The deformation grids of the RW_2 (Fig. 3)
276 showed a similar pattern, in which the head was broader and shorter in AC-DE and

277 (partially) in FA, elongating gradually in BI and LA on the one hand, and in UT-PA and
278 OUT on the other.

279

280 The shape variation of the pronotum mainly accounted for distinct patterns at the
281 generic level (Fig. 3) in which OUT, the *Hamonthophagus* species, and the *Morettius*
282 species formed three well-separated groups. However, it is noteworthy that in the plot
283 (Fig. 3) LA is separated from the other *Hamonthophagus* species, while OUT is nearer
284 to *Hamonthophagus* than to UT-PA. The deformation grids of the RW_1 showed two
285 distinct patterns, characterized by marked differences in the development of the fore
286 angles, and in the more or less marked posterior expansion of the pronotum. The
287 deformation grids of RW_2 showed instead differences in the lateral expansion of the
288 pronotum, which is slightly rounded in *Hamonthophagus*, broadly more expanded in
289 *Morettius*, and clearly extending outward in the central part in OUT. Here, the variance
290 explained by the first two RWs was 79.96%.

291

292 Noteworthy differences were found mostly at the generic level in the elytron, and again
293 three distinct groups were evident on the plot (Fig. 3), with 89.71% of the variance
294 explained by the first two RWs. The deformation grids of RW_1 highlighted marked
295 differences in the elytron shape, with a more slender and narrow elytron in UT-PA, and
296 a broader one in OUT, with *Hamonthophagus* well-separated and placed in an
297 intermediate position. Also, the deformation grids of RW_2 demonstrated two distinct
298 patterns, in which UT-PA and OUT showed an elytron more elongated than
299 *Hamonthophagus*. As before, LA is the species nearer to OUT, although the most
300 secluded species appears to be FA. On the other hand, UT and PA seem to be more
301 closely related to *Hamonthophagus* than to OUT.

302

303 For the mentum, the variance explained by the first two RWs was 84.37%, showing
304 marked differences at the generic and specific levels in the plot (Fig. 3). The groups
305 were clearly differentiated, with OUT well-characterized and isolated, while UT-PA
306 showed a more marked similarity with *Hamonthophagus*. UT and PA had an ostensibly
307 different mentum, although they remained more closely related to each other than to any
308 *Hamonthophagus* species. Again, LA was the most secluded *Hamonthophagus* species,

309 and partly replicated the situation already evident in the other structures. The
310 deformation grids of RW_1 (Fig. 3) showed conspicuous variations at fore and hind
311 margins, that were more or less deeply notched in OUT and *Hamonthophagus*
312 respectively. Also, the deformation grids of RW_2 (Fig. 3) showed distinct patterns of
313 shape variation with the mentum more squared on the sides in *Hamonthophagus*, and
314 far more rounded and expanded in OUT, with UT-PA in an intermediate position. On
315 the whole, the mentum proved a rather interesting structure, characterized by obvious
316 and marked differences at the specific and generic levels.

317

318 In the plot of the epipharynx (Fig. 4, with 72.04% of the variance explained by the first
319 two RWs), some particularly interesting results were found. Roggero *et al.* (2015) have
320 already pointed out that this structure is a very useful tool for taxa discrimination at the
321 specific and generic levels in Scarabaeidae. Distinct groups are here plainly manifest,
322 with OUT well-separated from the other species, UT-PA closely related, and
323 *Hamonthophagus* forming a third group in which the majority of the species were
324 sorted. LA is clearly distinct from the other *Hamonthophagus* species, but nevertheless
325 remains more closely related to them than to UT-PA. The deformation grids of RW_1
326 (Fig. 4) showed well-defined patterns of variation, particularly on the fore margin (more
327 notched in OUT), the tormae of the haptomerum area (larger in OUT), and the
328 proplegmatium (more downwardly arched in *Hamonthophagus*). The deformation grids
329 of RW_2 accounted mainly for variations of the supporting sclerotized structures (i.e.
330 the tormae), such as the crepis (shorter and larger in *Hamonthophagus* than in UT-PA)
331 and the tormae of the haptomerum (higher in UT-PA, and LA).

332

333 All the structures examined by GM methods provided useful information about the
334 patterns of variation among these species, and contributed in elucidating their
335 relationships based on morphological differences. Thus, the aligned configurations of
336 head, pronotum, right elytron, mentum and epipharynx were employed to build the
337 matrix for the phylogenetic analysis without converting them into linear values.

338

339

340

341 **Characters List**

342 (Figs 2-12)

343 **1. Head:** (0) uniform punctuation in clypeal and frontal parts; (1) punctuation of clypeal
344 part strongly differing from the frontal one.345 **2. Frontal carina:** (0) elongate; (1) intermediate; (2) short.346 **3. Pronotum length:** (0) greater than 2.5mm; (1) smaller than 2.5mm.347 **4. Pronotum width:** (0) greater than 4.5mm; (1) smaller than 4.5mm.348 **5. Pronotum, punctuation:** (0) absent; (1) present.349 **6. Elytral interstria with punctuation:** (0) almost inapparent, with small granules; (1)
350 thick and rasping, with medium-sized granules; (2) more or less large, but always
351 strong, with small and medium granules.352 **7. Elytral stria** (Fig. 5): (0) larger than the points; (1) as large as the points; (2)
353 narrower than the points.354 **8. Pygidium** (Fig. 6) **in males M, width/height ratio:** (0) less than 1.60; (1) more than
355 1.60.356 **9. Pygidium, punctuation constituted by:** (0) few, small and shallow points; (1) large
357 and strong, but scattered points; (2) large, strong and thick points.358 **10. Fore tibia in males, between the first and the second tooth a secondary**
359 **serration:** (0) inapparent; (1) simple; (2) double.360 **11. Fore tibia in males, between the second and the third tooth a secondary**
361 **serration:** (0) inapparent; (1) with two small denticles; (2) with one small denticle.362 **12. Fore tibia in males, after the third tooth a secondary serration:** (0) inapparent;
363 (1) with one small denticle.; (2) with two small denticles.364 **13. Fore tibia in females, between the first and the second tooth a secondary**
365 **serration:** (0) inapparent; (1)) with one small denticle; (2) with two small denticles.366 **14. Fore tibia in females, between second and third tooth a secondary serration:** (0)
367 inapparent; (1 with one small denticle; (2) with two small denticles.368 **15. Fore tibia in females, after the third tooth a secondary serration:** (0) inapparent;
369 (1) with one small denticle.370 **16. Phalloteca, apices of paramers** (Fig. 7): (0) greatly reduced; (1) well-developed.371 **17. Paramers, finger-shaped ventral expansion** (Fig. 7): (0) developed; (1) reduced,
372 almost inapparent.

373 **18. Paramers, the finger-shaped expansion inserted (with respect to the paramers**
 374 **base):** (0) high; (1) low.

375 **19. Endophallus** (Fig. 8) **constituted by:** (0) 2 sclerites; (1) more than 2 sclerites.

376 **20. Endophallus, primary sclerite with a longitudinal development** (Fig. 8): (0)
 377 squat and short; (1) elongate and narrow.

378 **21. Endophallus, primary sclerite carrying at base** (Fig. 8): (0) a convoluted
 379 expansion; (1) an evident hook.

380 **22. Endophallus, primary sclerite apical part** (Fig. 8): (0) little elongate, linear; (1)
 381 very elongated and sinuate.

382 **23. Receptaculum seminis, at base** (Fig. 9): (0) large; (1) narrow.

383 **24. Vagina, sclerotization** (Figs. 10-11): (0) present; (1) absent.

384 **25. Vagina, sclerotization** (Figs. 9-10): (0) groove-shaped, with part 1 inapparent; (1)
 385 funnel-shaped, with part 1 large and deep; (2) funnel-shaped with part 1 deep and
 386 narrow.

387 **26. Mentum, second palpus** (Fig. 12): (0) narrow, sub-cylindrical; (1) expanded, and
 388 rounded.

389 **27-192. Aligned configurations** (quantitative data) of epipharynx (27-60), mentum (61-
 390 104), head (105-142), pronotum (143-164), and right elytron (165-192) (Figs. 3-4).

391

392

393 **Phylogenetic analysis**

394 The linear parsimony analysis on the combined data matrix always gave the same single
 395 tree (Fig. 13), in which two distinct clades are present, one including UT and PA, and
 396 the other including all the other species. The results thus confirmed that the 32nd group
 397 is not a homogenous taxon. The *Morettius* clade is supported by resampling values of
 398 85/87/94, while the *Hamonthophagus* clade is supported by resampling values of
 399 61/83/78 for the Standard Bootstrap, Symmetrical Resampling, and Jackknife
 400 respectively. It is also noteworthy that *H. laceratus* is the most separated species in the
 401 *Hamonthophagus* clade, with high support values, endorsing the observations from the
 402 geometric morphometrics analysis. The following node also shows high support values,
 403 while the last node has much lower support values (Fig. 13), since the species of the

404 clade AC, FA and BI are strictly interrelated, although BI and FA are closer to each
405 other than to AC.

406

407

408 **Biogeographical analysis**

409 The *Hamonthophagus* species were characterized by a wide distribution (Fig. 14)
410 covering at least two macroareas, while *Moretius* species were characterized by more
411 reduced distributions. The georeferenced localities were mapped onto the terrestrial
412 ecoregions, giving clear differences in the species distributions. While AC is present in
413 the more xeric areas, extending only patchily in the savannah ecoregion, BI can be
414 considered a typical savannah species, with a greatly-extended distribution over the
415 entire central area of the Afrotropical region. Also DE and FA are essentially savannah
416 species, with a more southerly distribution than BI, and a much reduced presence in
417 desert areas (Fig. 14). The distribution of LA, covering the whole NE Afrotropical
418 region is also characterized by a prevalent savannah distribution, never reaching the
419 alpine steppe or the rain forest ecoregions.

420

421 The results of RASP (Fig. 15) gave a reconstruction of ancestral areas characterized by
422 three vicariant (green ring) and six dispersal (blue ring) events. At each node, a unique
423 optimal distribution was identified, except for the nodes 10 and 13 in which two
424 equiprobable alternatives were recognized, leading on the whole to four different
425 reconstructions: 1) node 10: DE, and node 13: AD; 2) node 10: DE, and node 13: CD;
426 3) node 10: DF, and node 13: AD; 4) node 10: DF, and node 13: CD. A vicariant event
427 was identified at node 15, in which the Oriental and Afrotropical clades were separated.
428 In the following nodes (Fig. 15), only dispersal events were allowed, with D as an
429 ancestral area. For the clade UT-PA, a vicariant event was evident, with PA being
430 present in AC, and UT in D. For the clade AC-BI-FA, a vicariant event was obtained at
431 Don the ancestral areas (BI-FA). On the basis of these results, it can be therefore
432 hypothesized that these species diversified in D, then extended eastwards (BI, and LA)
433 and southwards (FA, DE, and AC).

434

435 The VIP analysis produced a single possible reconstruction, identifying 3 disjunct sister
 436 pair (vicariant) nodes and 3 node removals (dispersal). The 1st vicariant node (node 8,
 437 Fig. 16) corresponded to the split of the *Morettius* species from the *Hamonthophagus*
 438 species, and the heuristic barrier (shown in red, Fig. 16a) separated A from B-G areas.
 439 The 2nd vicariant event (node 9, Fig. 16) resulted in the separation of PA from UT with
 440 a hypothetical barrier (shown in green, Fig. 16b) running along the Rift Valley and
 441 reaching westward to the Namibian Coast, thus splitting the A-C areas from the D area.
 442 The 3rd vicariant event (node 12, Fig. 16) took place between AC and the clade FA/BI,
 443 with a possible vicariant barrier through the A-D and E-G areas respectively (shown in
 444 blue, Fig. 16e). In the reconstruction, three sympatric speciations were evident in the
 445 nodes 10, 11, and 13 (Fig. 16) where for each there was a high species overlap for the
 446 *Hamonthophagus* species (Fig. 16c,d and f).

447

448

449 **Taxonomic account**

450 **Genus *Hamonthophagus* gen. nov.**

451 **Type species.** *Onthophagus bituberculatus* Olivier, 1789.

452 **Included species.** At present five Afrotropical medium size species (Figs. S2-3 -
 453 Appendix 4) can be included in the genus *Hamonthophagus* **gen. nov.**, namely: *H.*
 454 *acutus* (d'Orbigny, 1908), *H. bituberculatus* (Olivier, 1789), *H. depressus* (Harold,
 455 1871), *H. fallax* (d'Orbigny, 1913), and *H. laceratus* (Gerstaecker, 1871).

456 **Diagnosis.** The species included in the genus *Hamonthophagus* are strictly allied, and
 457 share a combination of characters that distinguishes them from the other Onthophagini:
 458 the anterior margin of the clypeus mostly sinuate and bidentate, and both head carinae
 459 poorly developed (i.e., slightly marked frontal carina and simple vertex carina). The
 460 granulo-punctuate pronotum is not very convex, and has evident anterior angles with
 461 divergent apices, that are inferiorly prolonged to strong prosternal carina. Head and
 462 pronotum are usually black or dark brown. The elytra are flat, narrowed backwards,
 463 with marked striae and interstriae, extremely variable in coloration, ranging from evenly
 464 black to yellow with a discal dark spot.

465 Sexual dimorphism was shown in the protibia (carrying a tooth on the inner margin only
466 in males), and pygidium (far more developed in males than females), as is common in
467 Onthophagini.

468 **Epipharynx.** The epipharynx (Fig. 2) is characterized by a rounded anterior margin
469 gently notched in the middle, abundant and widespread acropariae, and well-developed
470 corypha. The pubescence of the haptomerum is thick, the chaetopariae are almost
471 rectilinear, constituted by short and dense setae. The anterior epitorma is longitudinal
472 and narrow, the proplegmatium well-sclerotized and arched, the apotormae are present,
473 and the crepis is small, sharp and left-turned. The dextiotorma and laeotorma are slightly
474 asymmetrical.

475 **Male genitalia.** The male is characterized by a medium size phallosome (or aedeagus,
476 Fig. 7) with symmetrical parameres, and well-developed apices, carrying ventrally a
477 symmetrical expansion. The membranous internal sac (or endophallus, Fig. 8) carries a
478 hook-shaped and well-sclerotized primary sclerite, and some small accessory sclerites.

479 **Female genitalia.** In females, an asymmetrical, well-sclerotized funnel-shaped area is
480 evident in the vagina (Figs. 9-10), and is perhaps the most obvious character of the
481 genus. The membranaceous and plurisinate infundibulum is barely visible, being
482 basally located at a very low position, just on the oviductus. The receptaculum seminis
483 (Figs. 9-10) is curved in distal third, more enlarged at base, and tapering to apex, with a
484 large desclerotized area medially.

485 **Specific diagnosis.** The *Hamonthophagus* species can be distinguished on the basis of
486 some external features, as the body pubescence, the elytral striae and the punctuation of
487 pronotum and pygidium. Clear differences in shape were underlined by the geometric
488 morphometrics analysis of head, pronotum, elytron and mentum (see above). Marked
489 differences can also be highlighted by the analysis of the epipharynx (always, according
490 to the geometric morphometrics approach) and genitalia.

491 The pubescence covering the body is constituted by thick, ochreous, and truncated setae
492 that are short in *H. depressus*, *H. acutus* and *H. fallax*, and longer in *H. bituberculatus*,
493 while in *H. laceratus* the setae are very elongate, thinner and not truncated.

494 The pronotum has a characteristic punctuation with varying sized, closely spaced,
495 double points often carrying a hook-shaped granule never covering the point. While the
496 points are usually dense (but excluding *H. laceratus*), in *H. bituberculatus* only the

497 larger points bear the minute and flat granules, while in *H. acutus*, *H. fallax* and *H.*
498 *depressus*, the majority of the evident points carry well developed and thick granules.
499 *Hamonthophagus laceratus* is characterized instead by sparse and superficial points,
500 with very minute and scattered granules.

501 The elytral striae are constituted by a very narrow line with larger points, except *H.*
502 *laceratus*, where there are instead large striae with small points. Rasping, dense small
503 setigerous points are present on the interstriae, and in *H. acutus* and *H. bituberculatus*
504 the granules are small, while in *H. depressus* and *H. fallax* they are broader and evident.
505 Again, in *H. laceratus* the points are rade, and almost inapparent, with few, very small
506 granules. *Hamonthophagus fallax* usually carries an evident testaceous dot on the
507 proximal sides of the elytra, that are distally narrower than those in *H. depressus*.

508 In *H. acutus*, the pygidium is covered by superficial points and evident, roundish and
509 small granules, while in *H. depressus* and *H. fallax* the dense, large setigerous points are
510 without granules on the disc, carrying sometimes rough points only on the sides.

511 Besides, the latter species both have an evident and cerebroid microsculpture on the
512 surface, that in *H. acutus* is less marked. The pygidium of *H. bituberculatus* has an
513 opaque, smooth surface with few, scattered, shallow points (sometimes with minute
514 granules), but an evident, very thick microsculpture. Also in *H. laceratus*, the pygidium
515 is almost smooth, with an evident microsculpture, with only few and sparse points
516 lacking granules.

517 The fore margin of the epipharynx (Fig. 2) is only weakly notched in the middle in *H.*
518 *acutus*, while in *H. bituberculatus*, *H. depressus* and *H. fallax* the notch is V-shaped,
519 more marked and large. In *H. laceratus* the fore margin is slightly more squared that in
520 the other species. The apotormae are less developed in *H. bituberculatus* and *H.*
521 *laceratus* than in the other species. The crepis is more reduced in *H. acutus* and *H.*
522 *fallax*. The medial triangular sclerotized area of the proplegmatium is far shorter in *H.*
523 *fallax* than in the other species. In *H. laceratus* the rear sclerotized part between the
524 proplegmatium and crepis is much longer than in any other species.

525 In males, the paramers apices are elongate, large and only slightly hooked in *H. acutus*
526 *H. bituberculatus* and *H. fallax*, and more slender in *H. depressus*. In *H. laceratus*, the
527 parameres of the aedeagus are narrower than in the other *Hamonthophagus*, rounded at

528 apex and slightly downcurved. The small, rounded ventral expansion is well-developed
529 mainly in *H. laceratus* (see Fig. 7 for the comparison among the species).

530 The primary lamella of the endophallus is elongate with a large hook at the base in *H.*
531 *acutus*, *H. bituberculatus*, *H. depressus* and *H. fallax*, with small differences in the
532 longitudinal development among these species. In *H. laceratus* the primary lamella is
533 more peculiar, being tougher and half as long as in the other species, but always hook-
534 shaped (see Fig. 8 for the comparison among the species).

535 These species can also be easily identified by the shape of the peculiar asymmetrical,
536 funnel-shaped sclerotization of the vagina that shows a characteristic and differentiated
537 development in the five species (see Fig. 10 for the comparison among the species).

538 **General remarks.** No preimaginal stages have been described so far.

539 **Distribution.** The genus *Hamonthophagus* is distributed in arid and savannah
540 Afrotropical Regions (Fig. 14).

541 **Etymology.** The new genus was named after the Latin word *hamo*, = hook with
542 reference to the characteristic shape of the primary lamella of the internal sac.

543

544 ***Hamonthophagus acutus* (d'Orbigny, 1908: 171)**

545 (Figs. 2, 7, 8, 10)

546 **Type material.** NAMIBIA [Sud-Ouest africain allemand]: Okahandja [MNHN].
547 Paralectotypes: BOTSWANA: lake Ngami [MNHN]. DEMOCRATIC REPUBLIC OF
548 CONGO: [Tanganyika,] région de Mpala [MNHN]. MALAWI: Malawi Lake [=
549 Nyassa] [not located]. NAMIBIA: Salem [not located]. SOUTH AFRICA: Eastern Cape
550 province [= Cafreterie] [MNHN]. For the morphological account, please refer to the
551 original description.

552 **Geographic distribution** (Fig. 14). The species distribution surely comprises Namibia,
553 SW Botswana, and NW South Africa (see Appendix 3 for a detailed list of the
554 localities). Besides, in the type series d'Orbigny (1908) included also material from the
555 Tanganika area (Democratic Republic of Congo), and Nyassa (i.e., Malawi). The former
556 specimen was reported in the Collection Oberthur, MNHN, and the latter was reported
557 in "coll. du British Museum" (now BMNH) where, however, it has not been traced (M.
558 Barclay, *pers. comm.*). Neither specimen could be examined by us. Since no other
559 collection data from these areas were found within the studied material, these records

560 were here regarded as uncertain until further confirmation. Also, a specimen from the
 561 MNHN labelled as “Sénègal provenance tres douteuse” was not included in the present
 562 analysis.

563

564 ***Hamonthophagus bituberculatus* (Olivier, 1789: 131)**

565 (Figs. 2, 7, 8, 10)

566 **Synonymy.**

567 *Onthophagus discoideus* (Olivier, 1789:171) *teste* Harold 1880

568 **Type material.** At present, the typical material of *H. bituberculatus* could not be found.

569 Although various materials of the Olivier collection were traced in several museum
 570 collections over the years (Bragg, 1996; Staines & Whittington, 2003; Gültekin &
 571 Korotyaev, 2011), most specimens are still missing. The type material of this species
 572 was collected from “Senegal” by Geoffroy de Villeneuve, as well as its synonym *O.*
 573 *discoideus* (that was recorded also from Gorée Island). Since the type material of this
 574 species could not be located at present, no lectotype could be designed here. For the
 575 morphological account, please refer to the original description.

576 **Geographic distribution** (Fig. 14). The species is widely distributed in the whole sub-
 577 Saharan area (Benin, Burkina Faso, Eritrea, Ethiopia, Gambia, Ghana, Guinea, Guinea
 578 Bissau, Ivory Coast, Mauritania, Niger, Nigeria, Senegal (the type locality), Sudan, and
 579 Togo), and in Central and Eastern Africa (Central African Republic, Chad, Democratic
 580 Republic of Congo, Gabon, Kenya, Republic of Congo, and Uganda) extending
 581 eastwards and southwards toward Tanzania and Malawi (see Appendix 3 for more
 582 details). The species is also recorded from Cairo (Egypt, Schatzmayr, 1946; Baraud,
 583 1985) and Arabia (Paulian, 1980), but the data need to be confirmed. Accidental
 584 introduction is reported in Antilles (Martinique), where an anthropic cause was
 585 hypothesized to explain these findings (Matthews, 1966; Chalumeau, 1983).

586

587 ***Hamonthophagus depressus* (Harold, 1871: 116)**

588 (Figs. 2, 7, 8, 10)

589 **Synonymy.**

590 *Onthophagus laceratus* Peringuey 1901 *nec* Harold

591 *Onthophagus carteri* Blackburn, 1904: 147 *teste* Cartwright 1938

592 *depressus* var. *marmoreus* d'Orbigny 1904: 309

593 **Type material.** Lectotype here designated: (male) SOUTH AFRICA: [Caffraria =]
594 Eastern Cape province [ZMHB]. For the morphological account, please refer to the
595 original description.

596 **Geographic distribution** (Fig. 14). The species was originally described from South
597 Africa, Caffraria (now, Eastern Cape Province), but shows a wide distribution (the full
598 list of the localities can be found in Appendix 3) extending in a large part of the
599 Afrotropical region (Angola, Botswana, Burundi, Democratic Republic of Congo,
600 Kenya, Malawi, Mozambique, Namibia, South Africa, Tanzania, Zambia, and
601 Zimbabwe). Accidental introduction in Madagascar, Mauritius, USA (Florida, Georgia
602 and South Carolina), and Australia (New South Wales and Queensland). Howden &
603 Cartwright (1963) reported that specimens were collected at light in Georgia by Fattig.
604 In the USA, *H. depressus* has been recorded in Georgia, SW South Carolina and Florida
605 (Hunter & Fincher, 1996; Hoebeke & Beucke, 1997; Evans, 2014), with a scattered
606 distribution, since 1937 (Cartwright, 1938). The species was unintentionally introduced
607 in Australia surely before 1900 (Matthews, 1972; Woodruff, 1973), when Blackburn
608 (1904) described *H. depressus* specimens as a new species naming it *O. carteri*. This
609 species was later properly identified as *O. depressus* by Arrow (see Cartwright, 1938).
610 It is likely that the first introduction in Australia could be localized to the area near
611 Sydney, from where it began to expand its range starting from 1941 (Matthews, 1972;
612 Woodruff, 1973).

613

614 ***Hamonthophagus fallax* (d'Orbigny, 1913: 471)**

615 (Figs. 2, 7, 8, 10)

616 **Type material.** Lectotype here designated: (male) MALAWI: [Nyassa Zomba haut
617 Chiré=] Zomba, Shire river upper course, Malawi Lake [MNHN]. Paralectotype:
618 (female) TANZANIA [=Afrique Or Alem]: Dar-es-Salaam [MNHN]. For the
619 morphological account, please refer to the original description.

620 **Geographic distribution** (Fig. 14). The species has been described from Malawi and
621 Tanzania, and at present is recorded from Botswana, Burundi, Democratic Republic of
622 Congo, Kenya, Malawi, Namibia, Tanzania, and Zambia. The record from Graaf-Reinet

623 (Eastern Cape province, South Africa) is surely very interesting, but needs to be
624 confirmed by further records. In Appendix 3, a detailed list of the localities is given.

625

626 ***Hamonthophagus laceratus* (Gerstaecker, 1871: 50)**

627 (Figs. 2, 7, 8, 10)

628 **Synonymy.**

629 *Onthophagus laceratus* subsp. *benadirensis* Müller 1942: 82

630 **Type material.** Lectotype here designated: (male) TANZANIA: Zanzibar [ZMHB].

631 Paralectotype: (female) same locality [ZMHB]. The subspecies *benadirensis* from
632 Mogadishu area (Somalia) was examined and no marked differences were evident from
633 the nominal species. For the morphological account, please refer to the original
634 descriptions.

635 **Geographic distribution** (Fig. 14). The species was described from Zanzibar, and
636 shows a wide distribution extending in Burundi, Democratic Republic of Congo,
637 Ethiopia, Kenya, Somalia, Sudan, and Tanzania (see Appendix 3 for the full list of the
638 localities).

639

640

641 **Genus *Morettius* gen. nov.**

642 **Type species.** *Onthophagus pallens* d'Orbigny, 1908

643 **Included species.** *M. pallens* (d'Orbigny, 1908), and *M. utete* **sp. nov.**

644 **Diagnosis.** The species of the genus *Morettius* (Fig. S3 - Appendix 4) are characterized
645 by the mostly rounded and only slightly notched anterior margin of the clypeus, and the
646 pronotum covered by granules, sometimes mixed to points. The pygidium is always
647 smooth, with some rade points. The species show a moderate sexual dimorphism in the
648 fore tibiae, and the pygidium is larger in males than in females.

649 **Epipharynx.** The epipharynx (Fig. 2) fore margin is arched, with a largely V-shaped
650 notch in the middle. The corypha is reduced, the chaetopariae are arched with short,
651 thick, and almost equal length setae. The pubescence of the haptomerum is dense. The
652 proplegmatium is subequal on the whole length, with the posterior triangular
653 sclerotization reaching at least as much or more than half of the length of the anterior
654 epitorma, that is rectilinear, well-sclerotized and thin. The base of the triangular

655 sclerotization reaches the small, thick and upward-turned apophyses. Laeotorma and
656 dextiotorma are symmetrical, short and stout. Pternotormae are well-sclerotized, the
657 crepis is short but evident, with a sharp apex. The plegmatic area is visible.

658 **Male genitalia.** Only the male genitalia of *M. pallens* could be examined. The
659 phallobase (Fig. 7) is short, only slightly arched, slender, of equal size along the whole
660 length. The parameres are symmetrical, and squared, with a small tip at the apex, and a
661 small, rounded protrusion ventrally. The internal sac (Fig. 8) is membranous, with
662 various well-sclerotized parts greatly differing from the *Hamonthophagus* species.

663 **Female genitalia.** The female genitalia are very peculiar (Fig. 11), since the vagina of
664 both species is entirely membranous, and no sclerotization is present at all. Furthermore,
665 the infundibular tube is lowered as in *Hamonthophagus*, but here an expanded portion is
666 identifiable in the central part of vagina, which is differently shaped in the two species.
667 The receptaculum seminis is sickle-shaped, namely slim, arched, and apically sharp
668 (Fig. 9).

669 **Specific diagnosis.** These species can be easily distinguished on the basis of external
670 morphology, epipharynx and female genitalia.

671 The pronotum in *M. pallens* is covered by distinct rasping points mixed with smaller,
672 yellow granules; the granules of the rasping points instead are large, darker than the
673 background surface, and carry long, thick, and light yellow setae. The pronotum of *M.*
674 *utete* is covered by only few rasping setigerous points with thin, yellow setae and many
675 small granules, which are very thick, evenly coloured as the base, and without points.
676 Elytral striae of *M. pallens* are as large as the points, being instead larger than the points
677 in *M. utete*.

678 The smooth pygidium carries in *M. pallens* few, small, rade and deep setigerous points
679 that are not granulated, and in *M. utete* only some large and superficial vanishing points
680 without setae.

681 The epipharynx (see Fig. 2 for the comparison among the species) has the characteristic
682 shape of the *Morettius* species, but can be distinguished from *M. utete* by the more
683 developed apotormae, and the more slender laeotorma and dextiotorma.

684 In both species, the vagina is wholly desclerotized, but carries two globose symmetrical
685 expansions that encircle the desclerotized and lowered infundibulum in *M. pallens*,

686 while in *M. utete* there is a single, large expansion (see Fig. 11 for the comparison
687 among the species)

688 Since the male of *M. utete* is unknown, no comparison can be made between species.

689 **General remarks.** No preimaginal stages have been described so far.

690 **Distribution.** The genus *Morettius* is characterized by a disjoint distribution, being
691 found in central west Africa, and southeastern Africa (Tanzania).

692 **Etymology.** The genus is named after our colleague, the French entomologist Philippe
693 Moretto, who works extensively on African Scarabaeoidea.

694

695 ***Morettius pallens* (d'Orbigny, 1908: 172)**

696 (Figs. 2, 7, 8, 11)

697 **Type material.** Lectotype here designated: (male) CHAD: Kiao-Kata, Moyen-Chari,
698 south to Chad lake [=moyen Chari rives, Kiao-Kata] [MNHN]. Paralectotype: (female)
699 same locality [MNHN]. For the morphological account, please refer to the original
700 description.

701 **Geographic distribution** (Fig. 14). The species was reported from Cameroon, Chad,
702 Nigeria (southern border) Republic of Congo, and Sudan (the collection localities were
703 listed in Appendix 3).

704

705 ***Morettius utete* sp. nov.**

706 (Figs. 2, 11, 17)

707 **Etymology.** The species was named after the collection locality.

708 **Type material.** Holotype: female, TANZANIA: Utete-Rufijikindwjiwi [MHNL].

709 Paratypes: 2 females, same locality [MHNL] [EBCT].

710 **Description**

711 **Male.** Unknown

712 **Female.** Length: 6.1-6.6 mm. Head bronze, transverse (length/width ratio: 0,72), with
713 maximum width just anteriorly to the eyes. Clypeus sinuate, with clypeal edge reddish.
714 Clypeo-genal junction not sinuate. Frontal carina fine, weakly curved, placed at the
715 mid-length of the head, short and low, occupying half of the interocular space. Surface
716 markedly reticulate. Clypeus covered by flat and transverse granules, more or less
717 merged. Genal granules large and round. Vertex unarmed, weakly concave, with round

718 and fine granules. Antennal scape normally shaped, not dentate or serrulate. Antennal
719 club yellow.

720 Pronotum bronze, with hind angles bearing a bronze callus surrounded by a wide
721 yellowish area covering more than the half of the pronotal length and prolonged
722 narrowly on the sides to reach the anterior angles. Pronotal pubescence black and very
723 short, only evident on the lateral edges. Pronotum unarmed, very weakly transverse
724 (length/width ratio: 0.55). Base evenly curved, markedly bordered. Posterior angles not
725 sinuate. Anterior angles strongly sinuate, sharply projected outwards. Surface reticulate.
726 Area surrounding the callus not granulate; remaining pronotal surface entirely covered
727 by small granules.

728 Elytra yellow with black symmetrical spots, one basal on the fifth interstriae, one on the
729 first third of sixth and seventh interstriae, another four connected on the middle of the
730 second to fifth interstriae forming a zig-zig pattern. Juxtasutural interstriae yellow-
731 orange, darkened anteriorly. Basal carina of interstriae bronze. Pubescence yellow, very
732 short and scattered, only evident posteriorly. Elytral ground reticulate. Interstriae
733 bearing small yellow granules, arranged on the juxtasutural interstriae in a regular row.
734 Interstriae weakly convex, basally carinate. Striae narrow, well-marked, yellow.
735 Punctures of striae never wider than the striae. The seventh stria sinuate basally.

736 Pygidium yellow almost smooth, finely microreticulate, with small, rare, hardly
737 perceptible punctures. Base carinate.

738 Epipleura yellow. Sternal thoracic surface bronze, except for the base of propleurae
739 bronze. Abdominal sternites yellow. Metasternal pubescence scattered, short and
740 yellow.

741 Coxae yellow. Trochanters bronze. Femura yellow, apically bronze. Tibiae bronze,
742 meso- and metatibiae apically yellowish. Tarsomeres weakly bronze. Pubescence
743 yellow. Fore tibiae three-toothed. Tibial spur elongate, bent inward, apically rounded.
744 Tarsi normally shaped.

745 **Individual variation.** Paratype: the wide posterior spot of the right elytra is extended
746 on the sixth interstria. Spot of the sccond interstria obviously longer than in the
747 hotolypus. Seventh elytral stria only weakly sinuate.

748 **Epipharynx** (Fig. 2). See the above generic diagnosis.

749 **Male genitalia.** unknown

750 **Female genitalia** (Fig. 11). See the above generic diagnosis.

751 **Geographic distribution** (Fig. 14). The species is known only from the type locality.

752

753

754 **Identification Keys**

755 (Figs. S2-3 - Appendix 4)

756 1. Clypeal anterior edge mostly rounded and only slightly notched; pygidium smooth,
757 sometimes with some fine and scattered points. In males (*M. pallens*), primary sclerite
758 constituted by a well-sclerotized, almost spoon-shaped part and two less sclerotized

759 laminar parts. In the female, vagina entirely membranous *Morettius* (2)

760 1'. Clypeal anterior edge obviously bidentate; points of the pygidium fine or wide but

761 never densely distributed. Hook-shaped and well-sclerotized primary sclerite of

762 endophallus in male. Asymmetrical, well-sclerotized funnel-shaped area of the vagina in

763 female. *Hamonthophagus* (3)

764

765 ***Morettius* gen. nov**

766 2. Pronotum covered by distinct points carrying large granules darker than the surface
767 covering the points mixed with smaller, yellow granules; the rasping points carry long,

768 thick, light yellow setae *pallens* (d'Orbigny)

769 2'. Pronotum covered by many small, very thick evenly coloured granules; only few

770 rasping setigerous points with thin, yellow setae *utete* **sp. nov.**

771

772 ***Hamonthophagus* gen. nov.**

773 3. Pronotum black or dark brown with yellowish spots at the hind angles, and covered

774 by sparse points, with very minute and rade granules *laceratus* (Gerstaecker)

775 3'. Pronotum entirely black or dark brown, and covered by varyingly sized, closely

776 spaced, dense double points often carrying hook-shaped granules 4

777 4. Pronotum with simple, small points and larger points with minute and flat granules.

778 Setae of the dorsal surface obviously longer than wide *bituberculatus* (Olivier)

779 4'. Pronotum covered by granulate points. Setae of the dorsal surface largely as long as

780 wide 5

- 781 5. Pronotum evenly covered by granulate points. Pygidium covered by scattered but
 782 evident granules *acutus* (d'Orbigny)
 783 5'. Pronotum covered by large granulate points and few, very smaller simple points.
 784 Pygidium covered by rather large, more or less dense ocellate points 6
 785 6. In males, apices of paramers elongate, large and only slightly hooked. In females, the
 786 sclerotized area asymmetrically developed with the apex on the right. Elytra black, with
 787 one or several testaceous, symmetrical, small patches *fallax* (d'Orbigny)
 788 6'. In males, apices of paramers more slender and pointed. In females, the sclerotized
 789 area well-developed and triangular-shaped. Elytra usually entirely black or, sometimes,
 790 dark brown *depressus* (Harold)

791

792

793

794 **DISCUSSION**

795 The study of the d'Orbigny 32nd species group has been addressed by employing an
 796 innovative and very powerful approach. The combined phylogenetic method allowed us
 797 to handle together different morphological datasets of discrete and continuous
 798 characters, summarizing the modularized traits. Our first research goal was focused on
 799 testing how several trait configurations could be processed to gain quantitative data, and
 800 then utilized after being combined in a single matrix with the qualitative characters.
 801 The use of the quantitative approach often furnished more detailed information about
 802 various anatomical traits than the qualitative approach, also evidencing at its best the
 803 intra and interspecific differences of shape variation. Therefore, quantifying information
 804 can provide a more accurate dataset and allow more effective analysis of morphological
 805 characters.

806

807 On the whole, the results of the analyses concurred in highlighting a lack of
 808 phylogenetic homogeneity in the d'Orbigny 32nd species group, whereby the recognition
 809 of a generic-level divergence at the basal dichotomy of the tree was well-founded (Fig.
 810 13).

811 The first genus was designated as *Hamonthophagus* and included the majority of the
 812 species (see the Taxonomic Account above), while the second one, namely *Morettius*,

813 included only two species, one of which was identified and described here (see the
814 Taxonomic Account above). The two genera are clearly diversified on the basis of the
815 shape of the epipharynx and genitalia of both sexes. These structure are characterized by
816 marked complexity, evidencing the generic trends, since they usually constitute the
817 synapomorphies founding the basal generic status. Additionally, while for the usefulness
818 of the genitals in taxonomic research is well founded and undisputed (Eberhard, 2010a,
819 2010b, 2011), the epipharynx is still little used, although it has been extremely effective
820 to define and separate even very challenging groups (Roggero *et al.*, 2015).

821

822 The age estimates for the African coprophagous radiations, as evaluated in recent
823 analyses using four nuclear and mitochondrial DNA markers (Ahrens *et al.*, 2014),
824 could also be applied to *Hamonthophagus* and *Morettius*. The scarab divergences were
825 demonstrated in the calibrated Time-Tree showing scenarios closely related to a
826 diversified pattern of herbivores (i.e., dung-producing mammal lineages). Such an
827 evolutionary-ecological context could be allocated to the Miocene when the lineages
828 should have radiated. This period was characterized by climatic changes that caused the
829 spread of the savannah and the dominance of the dung-producing Artiodactyla (Wirta *et*
830 *al.*, 2008; Sole & Scholtz, 2010). In this period, the ancestral generic lineages of most
831 extant Scarabaeini/Onthophagini arose, and it is likely that *Hamonthophagus* and
832 *Morettius* might be involved in these speciation processes, originating in the Eastern
833 Central African area (area D, Fig. 1). Subsequently, these typically tunneling dung
834 beetles split thanks to sequential migrations of herbivorous mammals (Monaghan *et al.*,
835 2007; Philips, 2011) across the entire continent towards the south and north, resulting in
836 various and diversified dispersal events hypothesized in Fig. 18 for *Hamonthophagus*,
837 with part of the group spreading south, and part extending northwards.

838

839

840

841 **ACKNOWLEDGMENTS**

842 This research was made possible thanks to grants from the Fondazione CRT, Research
843 and Education section (Torino, Italy), and was also partly supported by the Italian
844 Ministero dell'Istruzione, dell'Università e della Ricerca (MIUR). We are greatly

845 indebted to the museum curators and private collectors who kindly lent us their material.
846 We are also very grateful to M. Barclay (BMNH, London) for the useful information on
847 32nd group type material. We want also to thank two anonymous reviewers whose
848 comments and suggestions greatly contributed to improving our paper. We are very
849 grateful to our colleague Dan Chamberlain, who kindly corrected the English text.

850

851

852

853 REFERENCES

- 854 Ahrens D, Schwarzer J, Vogler AP. 2014. The evolution of scarab beetles tracks the
855 sequential rise of angiosperms and mammals. *Proceedings of the Royal Society B*
856 281: 20141470. DOI: 10.1098/rspb.2014.1470.
- 857 Adams DC, Cardini A, Monteiro LR, O'Higgins P, Rohlf FJ. 2011. Morphometrics and
858 phylogenetics: Principal components of shape from cranial modules are neither
859 appropriate nor effective cladistic characters. *Journal of Human Evolution* 60:
860 240-243.
- 861 Arias JS, Szumik CA, Goloboff PA. 2011. Spatial analysis of vicariance: a method for
862 using direct geographical information in historical biogeography. *Cladistics* 27:
863 617–628.
- 864 Baraud J. 1985. Coléoptères Scarabaeoidea, Faune du nord de l'Afrique, du Maroc au
865 Sinaï. *Encyclopédie Entomologique* 46: 1-652.
- 866 Barbero E, Palestrini C, Roggero A. 2003. *Revision of the genus Phalops Erichson,*
867 *1848 (Coleoptera: Scarabaeidae: Onthophagini)*. Torino: Museo Regionale di
868 Scienze Naturali.
- 869 Barbero E, Palestrini C, Roggero A. 2009. Systematics and phylogeny of *Eodrepanus*, a
870 new Drepanocerine genus, with comments on biogeographical data (Coleoptera:
871 Scarabaeidae: Oniticellini). *Journal of Natural History* 43: 1835–1878.
- 872 Barbero E, Palestrini C, Roggero A. 2011. *Tibiodrepanus tagliaferrii* - a new
873 Afrotropical Drepanocerina species (Coleoptera: Scarabaeidae: Oniticellini), with
874 notes on phylogeny and distribution of the genus. *Zootaxa* 2923: 27–47.

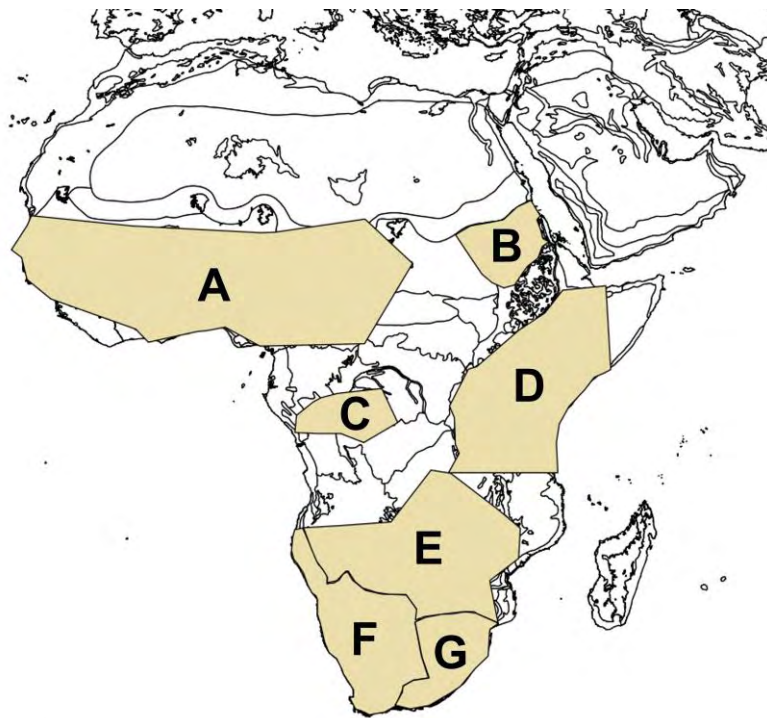
- 875 de Bivort BL, Clouse RM, Giribet G. 2010. A morphometrics-based phylogeny of the
876 temperate Gondwanan mite harvestmen (Opiliones, Cyphophthalmi, Pettalidae).
877 *Journal of Zoological Systematics and Evolutionary Research* 48: 294-309.
- 878 de Bivort BL, Clouse RM, Giribet G. 2012. A cladistic reconstruction of the ancestral
879 mite harvestman (Arachnida, Opiliones, Cyphophthalmi): portrait of a Paleozoic
880 detritivore. *Cladistics* 28, 582–597.
- 881 Blackburn T. 1904. Revision of the Australian Aphodiides, and descriptions of three
882 new species allied to them. *Proceedings of the Royal Society of Victoria* 17: 145-
883 181.
- 884 Bragg PE. 1996. Type specimens of Phasmida in the Nationaal Natuurhistorisch
885 Museum, Leiden (Insecta: Phasmida). *Zoologische Mededelingen* 70: 105-115.
- 886 Catalano SA, Goloboff PA, Giannini P. 2010. Phylogenetic morphometrics (I): the use
887 of landmark data in a phylogenetic framework. *Cladistics* 26: 539–549.
- 888 Chalumeau F. 1983. *Les Coléoptères Scarabaeides des Petites Antilles (Guadeloupe à*
889 *Martinique)*. *Taxonomie-Ethologie-Biogéographie*. Paris: Editions Lechevalier.
- 890 Clouse RM, de Bivort BL, Giribet G. 2010. Phylogenetic signal in morphometric data.
891 *Cladistics* 27: 1-4.
- 892 Cartwright OL. 1938. A South African *Onthophagus* found in the United States.
893 *Entomological News* 49: 114-115.
- 894 d’Orbigny H. 1908. Descriptions d’espèces nouvelles d’Onthophagides africains et notes
895 synonymiques. *Annales de la Société Entomologique de France* 77: 65-208.
- 896 d’Orbigny H. 1913. Synopsis des Onthophagides d’Afrique. *Annales de la Société*
897 *Entomologique de France* 82: 1-742.
- 898 Eberhard WG. 2010a. Evolution of genitalia – theories, evidence, and new directions.
899 *Genetica* 138: 5-18. doi: 10.1007/s10709-009-9358-y.
- 900 Eberhard WG. 2010b. Rapid divergent evolution of genitalia. In: Leonard J., Cordoba A.
901 eds. *The Evolution of Primary Sexual Characters in Animals*. Oxford: Oxford
902 University Press, 40-78.
- 903 Eberhard WG. 2011. Experiments with genitalia: a commentary. *Trends in Ecology and*
904 *Evolution* 26: 17-21. doi:10.1016/j.tree.2010.10.009.

- 905 Emlen DJ, Marangelo J, Ball B, Cunningham CW. 2005. Diversity in the weapons of
906 sexual selection: horn evolution in the beetle genus *Onthophagus* (Coleoptera:
907 Scarabaeidae). *Evolution* 59:1060-1084.
- 908 Emlen DJ, Szafran Q, Corley LS, Dworkin J. 2006. Insulin signaling and limb-
909 patterning: candidate pathways for the origin and evolutionary diversification of
910 beetle 'horns'. *Heredity* 97: 179-191.
- 911 Evans AV. 2014. *Beetles of Eastern North America*. Princeton (NJ): Princeton
912 University Press.
- 913 Ferretti N, González A, Pérez-Miles F. 2012. Historical biogeography of the genus
914 *Cyriocosmus* (Araneae: Theraphosidae) in the Neotropics according to an event-
915 based method and spatial analysis of vicariance. *Zoological Studies* 51: 526-535.
- 916 Gold MEL, Brochu CA, Norell MA. 2014. An Expanded Combined Evidence Approach
917 to the *Gavialis* Problem Using Geometric Morphometric Data from Crocodylian
918 Braincases and Eustachian Systems. *PLoS ONE* 9: e105793.
919 doi:10.1371/journal.pone.0105793.
- 920 Goloboff PA, Catalano SA. 2010. Phylogenetic morphometrics (II): algorithms for
921 landmark optimization. *Cladistics* 27:42-51. doi: 10.1111/j.1096-
922 0031.2010.00318.x.
- 923 Goloboff PA, Farris JS, Nixon KC. 2003. TNT: Tree Analysis Using New Technology.
924 Available at: <http://www.zmuc.dk/public/phylogeny/TNT/> [through the Hennig
925 Society].
- 926 Goloboff PA, Farris JS, Nixon KC. 2008. TNT, a free program for phylogenetic
927 analysis. *Cladistics* 24: 774–786.
- 928 Goloboff, PA, Mattoni CI, Quinteros AS. 2006. Continuous characters analyzed as
929 such. *Cladistics* 22: 589-601. doi:10.1111/j.10960031.2006.00122.x.
- 930 Gültekin L, Korotyaev BA. 2011. *Lixus petiolicola* n. sp. from Northeastern Turkey and
931 *Lixus furcatus* Olivier: Comparative systematic and ecological study (Coleoptera:
932 Curculionidae: Lixinae). *Annales de la Société Entomologique de France* 47: 101-
933 111.
- 934 Hoebeke ER, Beucke K. 1997. Adventive *Onthophagus* (Coleoptera: Scarabaeidae) in
935 North America: geographic ranges, diagnoses, and new distributional records.
936 *Entomological News* 108: 345–362.

- 937 Howden F, Cartwright OL. 1963. *Scarab beetles of the genus Onthophagus North of*
938 *Mexico*. Washington: Smithsonian Institution.
- 939 Hunter JS III, Fincher GT. 1996. Distribution of *Onthophagus depressus*: an introduced
940 species of dung beetle. *Journal of Agricultural Entomology* 13: 319-322.
- 941 Matthews EG. 1966. A taxonomic and zoogeographic survey of the Scarabaeinae of the
942 Antilles. *Memoirs of the American Entomological Society* 21: 1-133.
- 943 Matthews EG. 1972. A revision of the Scarabaeinae dung beetles of Australia. I. Tribe
944 Onthophagini. *Australian Journal of Zoology* 9 (suppl.):1-330.
- 945 Moczek AP. 2006. Integrating micro- and macroevolution of development through the
946 study of horned beetles. *Heredity* 97: 168-178.
- 947 Monaghan MT, Inward DG, Hunt T, Vogler AP. 2007 A molecular phylogenetic
948 analysis of the Scarabaeinae (dung beetles). *Molecular Phylogenetics and*
949 *Evolution* 45: 674-692. doi:10.1016/j.ympev.2007.06.009.
- 950 Moretto P. 2009. Essai de classification des *Onthophagus* Latreille, 1802 africains des
951 5ème et 6ème groupes de d'Orbigny (Coleoptera, Scarabaeidae). *Nouvelle Revue*
952 *d'Entomologie* 25: 145-178.
- 953 Olson DM, Dinerstein E, Wikramanayake ED, Burgess ND, Powell GVN, Underwood
954 EC, D'Amico JA, Itoua I, Strand HE, Morrison JC, Loucks CJ, Allnutt TF,
955 Ricketts TH, Kura Y, Lamoreux JF, Wettengel WW, Hedao P, Kassem KR. 2001.
956 Terrestrial Ecoregions of the World: A New Map of Life on Earth. *BioScience* 51:
957 933-938.
- 958 Palestini C. 1992. Sistematica e zoogeografia del genere *Onthophagus*, sottogenere
959 *Proagoderus* Lansberge. *Memorie della Società entomologica italiana* 71: 1-358.
- 960 Paulian R. 1980. Insects of Saudi Arabia. Coleoptera: Scarabaeoidea (1ère contribution).
961 *Fauna of Saudi Arabia* 2: 141-154.
- 962 Philips TK. 2011. The evolutionary history and diversification of dung beetles. In:
963 Simmons LW, Ridsdill-Smith TJ eds. *Ecology and evolution of dung beetles*.
964 Blackwell Publishing: Oxford. 21-46.
- 965 QGIS Development Team (2016). QGIS v2.16. Geographic Information System User
966 Guide. Open Source Geospatial Foundation Project. – Electronic document,
967 Available at: <http://download.osgeo.org/qgis/doc/manual/<DOCUMENT>>.
- 968 Rambaut A. 2014. FigTree v1.4.2. Available at: <http://tree.bio.ed.ac.uk/software/>.

- 969 Roggero A, Barbero E, Palestini C. 2015. Phylogenetic and biogeographical review of
970 the Drepanocerina (Coleoptera, Scarabaeidae, Oniticellini). *Arthropod Systematic*
971 *and Phylogeny*, 73: 153-174.
- 972 Roggero A, Barbero E, Palestini C. 2016. Revised classification and phylogeny of an
973 Afrotropical species group based on molecular and morphological data, with the
974 description of a new genus (Coleoptera: Scarabaeidae: Onthophagini). *Organisms*
975 *Diversity & Evolution*. DOI: 10.1007/s13127-016-0297-z [online first version,
976 30.7.2016].
- 977 Rohlf FJ. 2012. NTSYSpc: numerical taxonomy system. ver. 2.21r. Setauket, New
978 York: Exeter Software.
- 979 Rohlf FJ. 2016a. tpsDig v2.27. Available at: <http://life.bio.sunysb.edu/morph/>.
- 980 Rohlf FJ. 2016b. tpsUtil v1.69. Available at: <http://life.bio.sunysb.edu/morph/>.
- 981 Rohlf FJ. 2016c. tpsSmall v1.33. Available at: <http://life.bio.sunysb.edu/morph/>.
- 982 Rohlf FJ. 2016d. tpsRelw v1.65. Available at: <http://life.bio.sunysb.edu/morph/>.
- 983 Schatzmayr A. 1946. Gli scarabaeidi coprofagi della Libia e dell'Egitto. *Atti della*
984 *Società italiana di Scienze Naturali* 85: 40-84.
- 985 Sharkey MJ, Carpenter JM, Vilhelmsen L, Heraty J, Liljeblad J, Dowling APG,
986 Schulmeister S, Murray D, Deans AR, Ronquist F, Krogmann L, Wheeler WC.
987 2012. Phylogenetic relationships among superfamilies of Hymenoptera. *Cladistics*
988 28: 80-112. doi: 10.1111/j.1096-0031.2011.00366.x.
- 989 Smith UE, Hendricks JR. 2013. Geometric morphometric character suites as
990 phylogenetic data: extracting phylogenetic signal from gastropod shells.
991 *Systematic Biology* 62: 366–385.
- 992 Sole CL, Scholtz CH. 2010. Did dung beetles arise in Africa? A phylogenetic hypothesis
993 based on five gene regions. *Molecular Phylogenetics and Evolution* 56: 631-641.
994 doi:10.1016/j.ympev.2010.04.023.
- 995 Staines CL, Whittington AE. 2003. Chrysomelidae (Coleoptera) types in the Royal
996 Museum of Scotland Collection. *Zootaxa* 192: 1-8.
- 997 Tagliaferri F, Moretto P, Tarasov SI. 2012. Essai sur la systématique et la phylogénie
998 des *Onthophagus* Latreille, 1802, d'Afrique tropicale appartenant au septième
999 groupe de d'Orbigny. Description d'un sous-genre nouveau et de trois espèces
1000 nouvelles (Coleoptera, Scarabaeoidea, Onthophagini). *Catharsius La Revue* 6:1-

- 1001 31.
- 1002 Tarasov SI, Solodovnikov AY. 2011. Phylogenetic analyses reveal reliable
1003 morphological markers to classify mega-diversity in Onthophagini dung beetles
1004 (Coleoptera: Scarabaeidae: Scarabaeinae). *Cladistics* 27: 1-39.
- 1005 White F, Leonard J. 1991. Phytogeographical links between Africa and Southwest Asia.
1006 In: Engel T, Frey W, Kürschner H. eds. *Contributiones Selectae ad Floram et*
1007 *Vegetationem Orientis*. Flora et Vegetatio Mundi. Band IX. Berlin: Cramer, 229-
1008 246.
- 1009 Wirta H, Orsini L, Hanski I. 2008. An old adaptive radiation of forest dung beetles in
1010 Madagascar. *Molecular Phylogenetics and Evolution* 47: 1076-1089.
1011 doi:10.1016/j.ympev.2008.03.010.
- 1012 Woodruff RE. 1973. *Arthropods of Florida. Vol 8. The scarab beetles of Florida*
1013 *(Coleoptera: Scarabaeidae)*. Gainesville: Florida Dpt. of Agriculture.
- 1014 Yu Y, Harris AJ, He X. 2010a. A Rough Guide to S-DIVA v1.9. Available at:
1015 <http://mnh.scu.edu.cn/S-DIVA/blog/SDIVA/index.html>.
- 1016 Yu Y, Harris AJ, He X. 2010b. S-DIVA (Statistical Dispersal-Vicariance Analysis): A
1017 tool for inferring biogeographic histories. *Molecular Phylogenetics and Evolution*
1018 56: 848–850.
- 1019

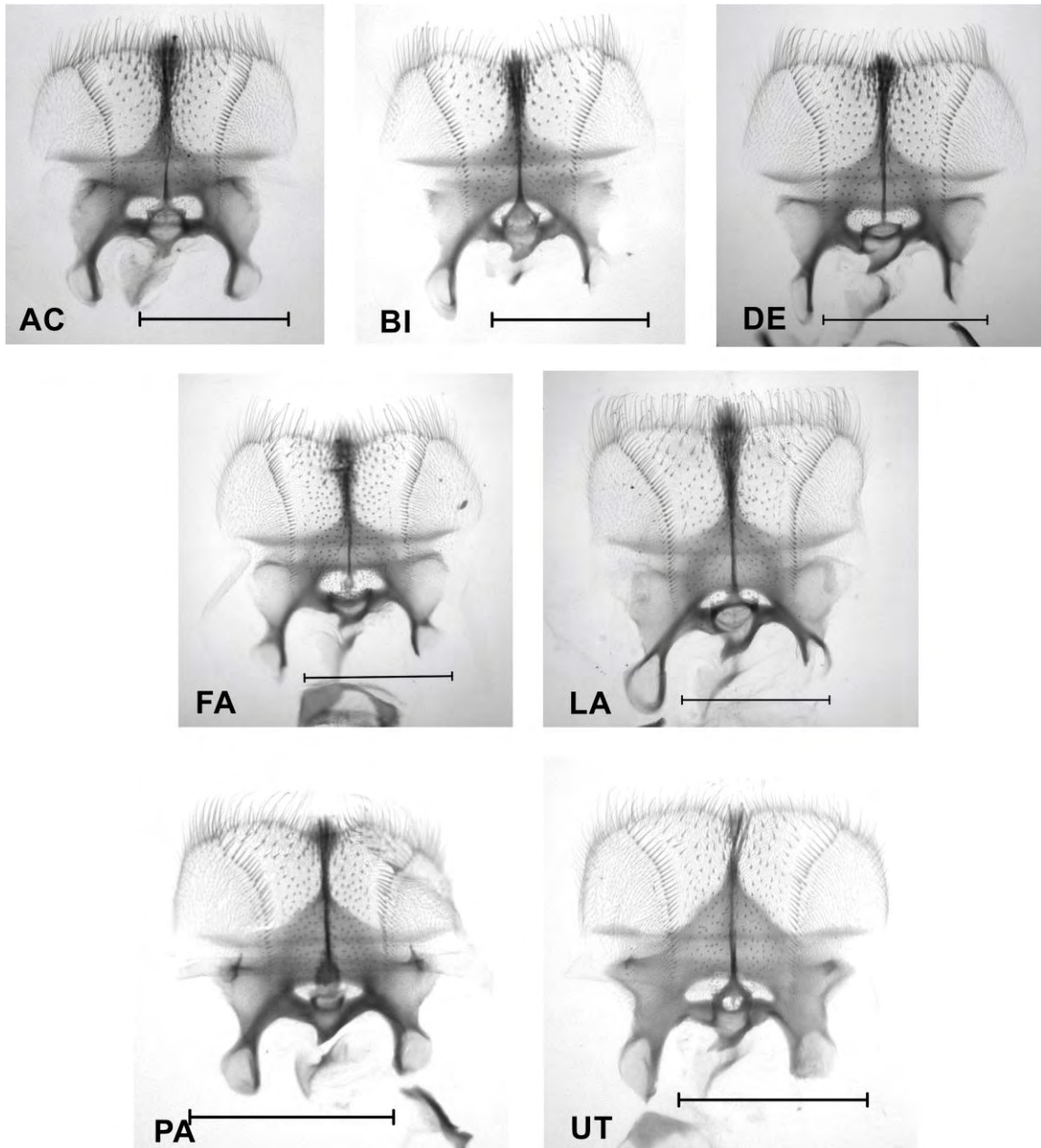
1020 **FIGURES**

1021

1022 **Figure 1.** Macroareas identified for the biogeographical analysis, where A = Guinea-
1023 Congolian area (GCA), B= Eastern Sudanian area (ESA), C= Central Congolian area
1024 (CCA), D = Somalo-Masai area (SMA), E = Zambesian area (ZAA), F = Namib-
1025 Kalahari area (NKA), G = Highveld area (HIA), while the outgroup distribution (H =
1026 Oriental Region, ORA) is not shown on the map.

1027

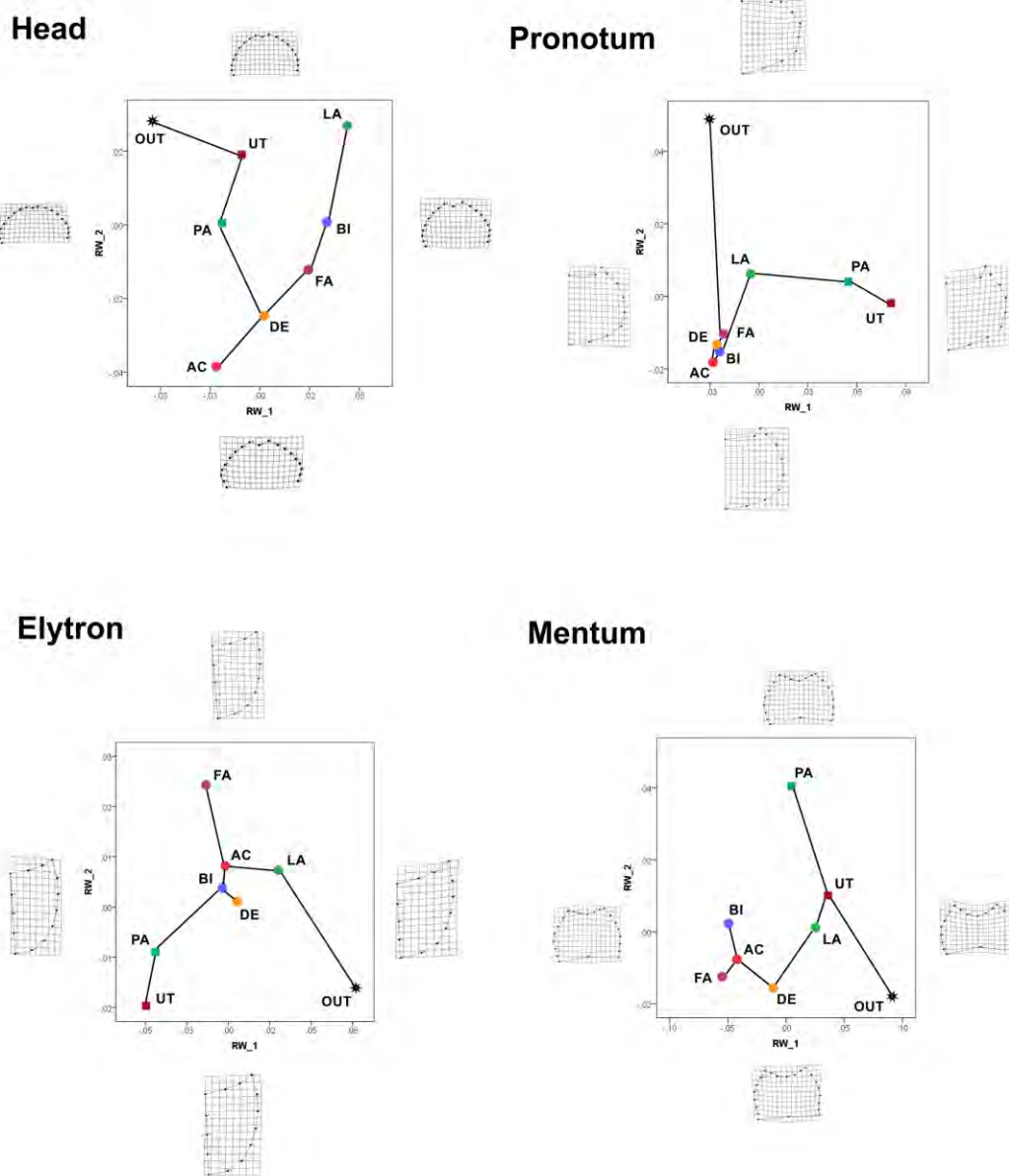
1028



1029

1030 **Figure 2.** Epipharynx of the species of the genera *Hamonthophagus* (AC = *acutus*, BI =
 1031 *bituberculatus*, DE = *depressus*, FA = *fallax*, and LA = *laceratus*) and *Morettius* (PA =
 1032 *pallens*, and UT = *utete*). Scalebar = 0.5 mm.

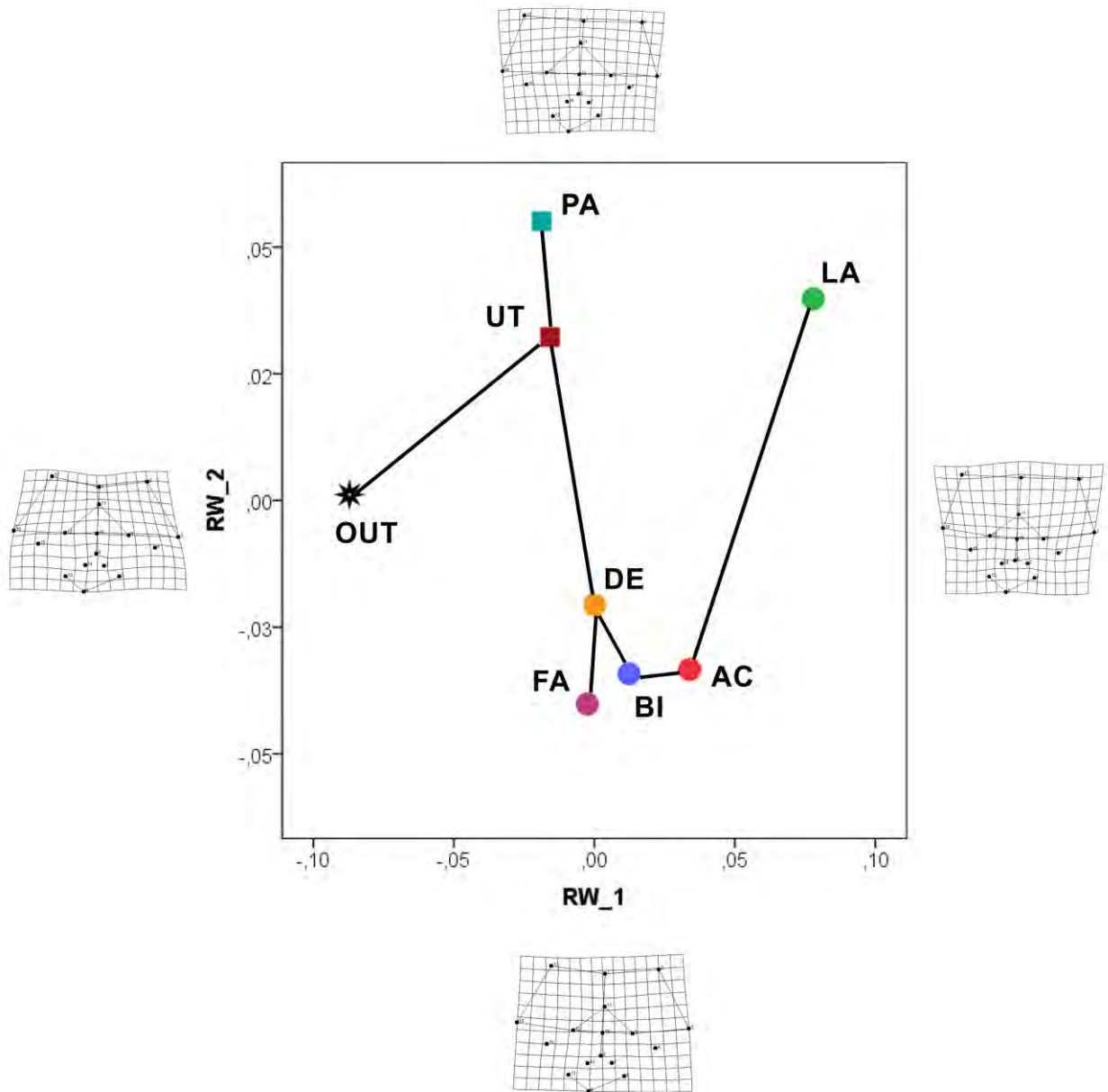
1033



1034

1035 **Figure 3.** Scatterplots of the RW 1 and 2 of head, pronotum, right elytron and mentum
 1036 (semilandmarks method). Only the specimens employed to build the matrix are shown
 1037 here. The deformation grids corresponding to the minimum and maximum values of the
 1038 axes are shown for each anatomical trait. See text for the codes.

1039

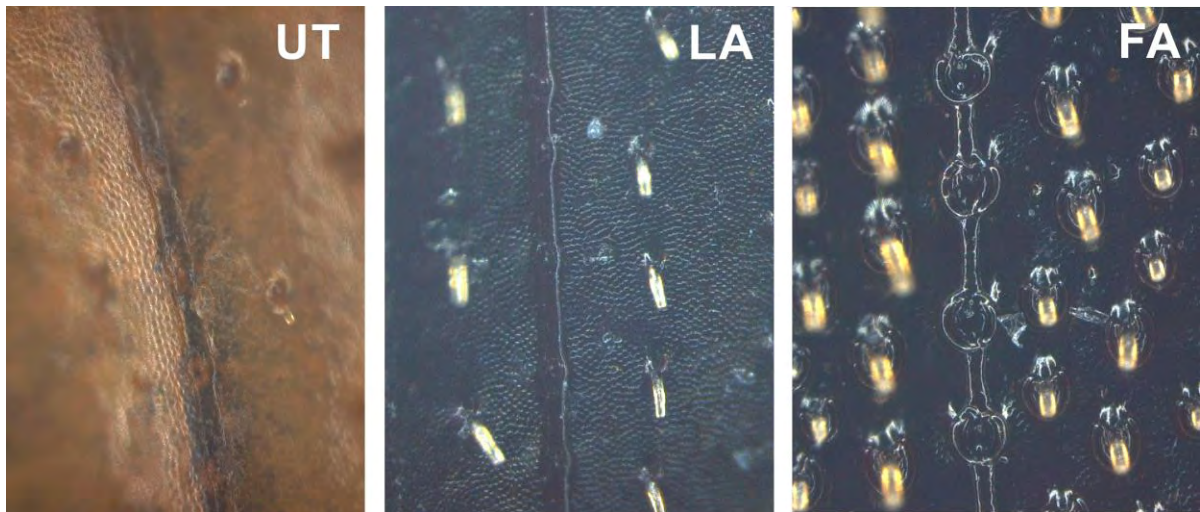


1040

1041 **Figure 4.** Scatterplots of the RW 1 and 2 of epipharynx (landmarks method). Only the
 1042 specimens employed to build the matrix are shown. The deformation grids of the
 1043 minimum and maximum values of the axes are shown. See text for the codes.

1044

1045

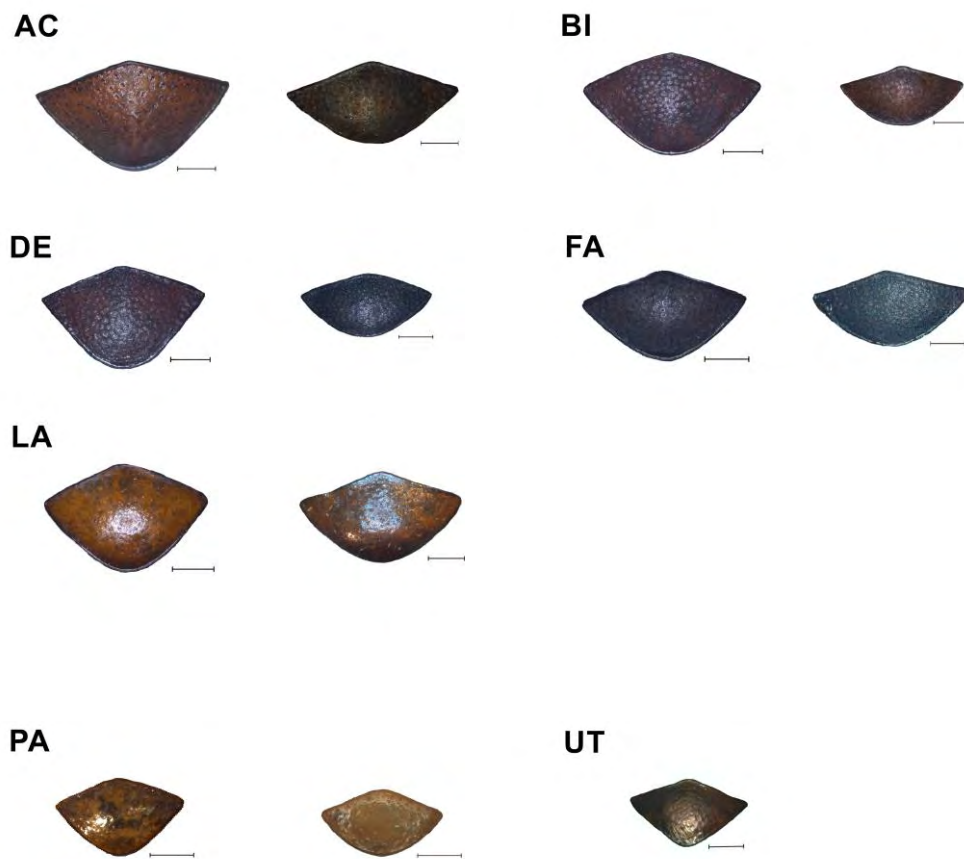


1046

1047 **Figure 5.** Elytral stria, character 7: from left to right state 0 (*Morettius utete* = UT), state
1048 1 (*Hamonthophagus laceratus* = LA), and state 2 (*H. fallax* = FA).

1049

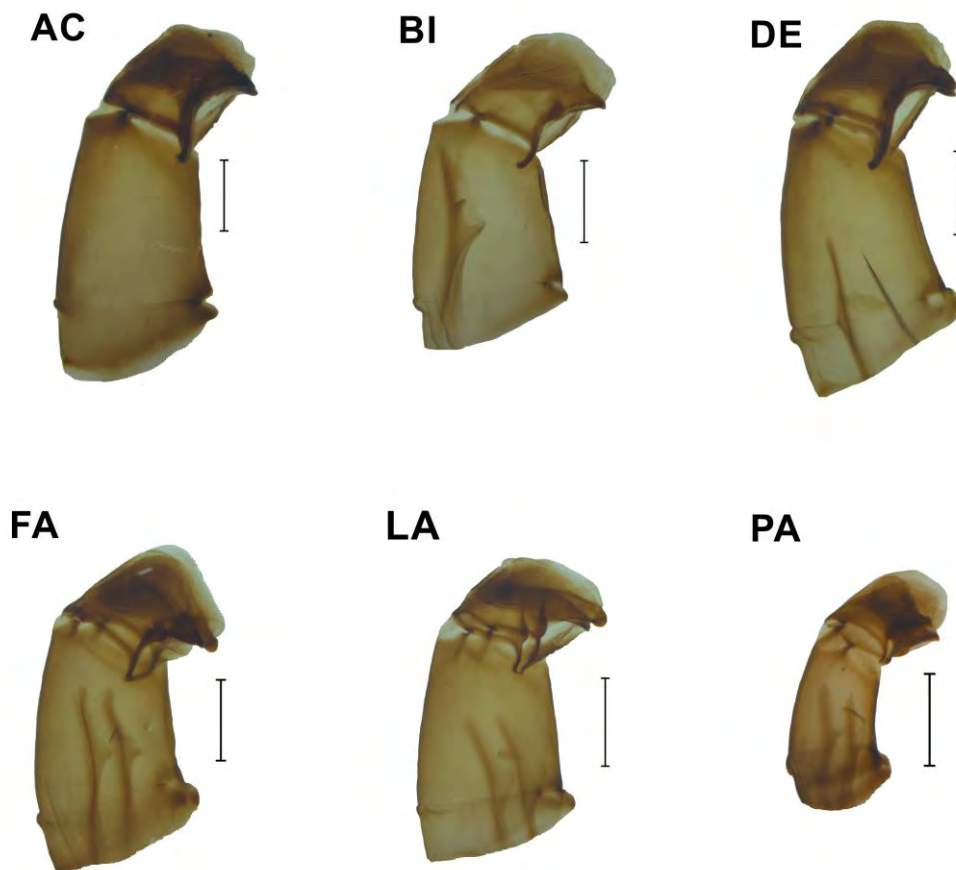
1050



1051

1052 **Figure 6.** Pygidium, male on left and female on right, except *Morettius utete* (UT) in
 1053 which only the female is known, see text for the codes. Scalebar = 0.5 mm.

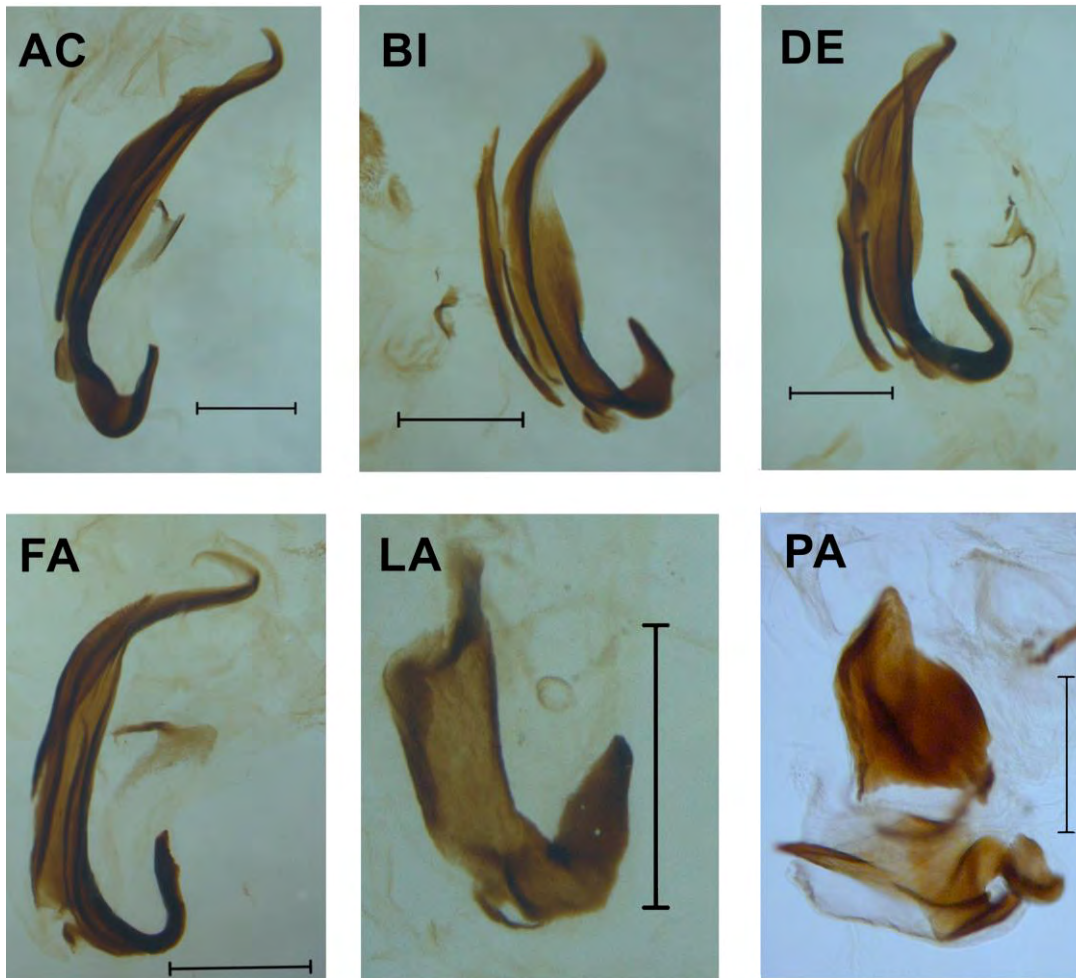
1054



1055

1056 **Figure 7.** Aedeagus of the species of the genera *Hamonthophagus* (AC, BI, DE, FA and
1057 LA) and *Morettius* (PA), see text for the codes. Scalebar = 0.5 mm.

1058



1059

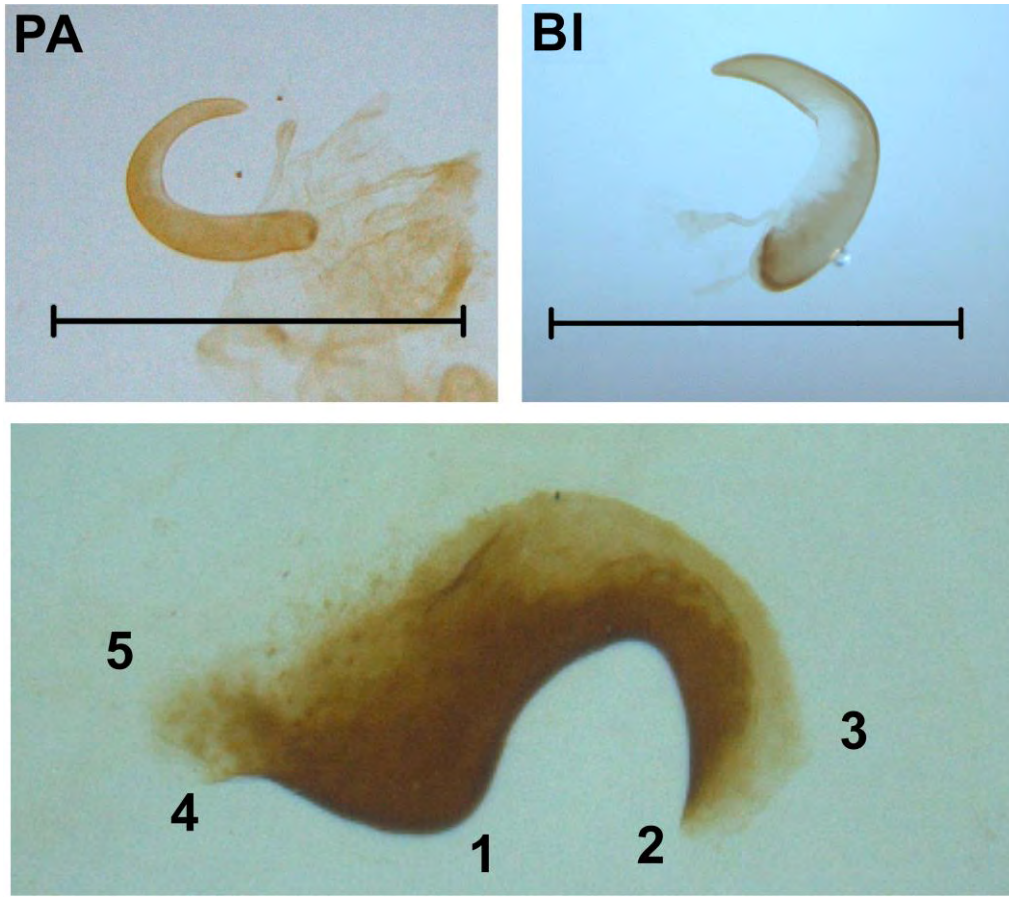
1060 **Figure 8.** Primary lamella of the species of the genera *Hamonthophagus* (AC, BI, DE,

1061

FA and LA) and *Morettius* (PA). Scalebar = 0.5 mm.

1062

1063



1064

1065

1066

1067

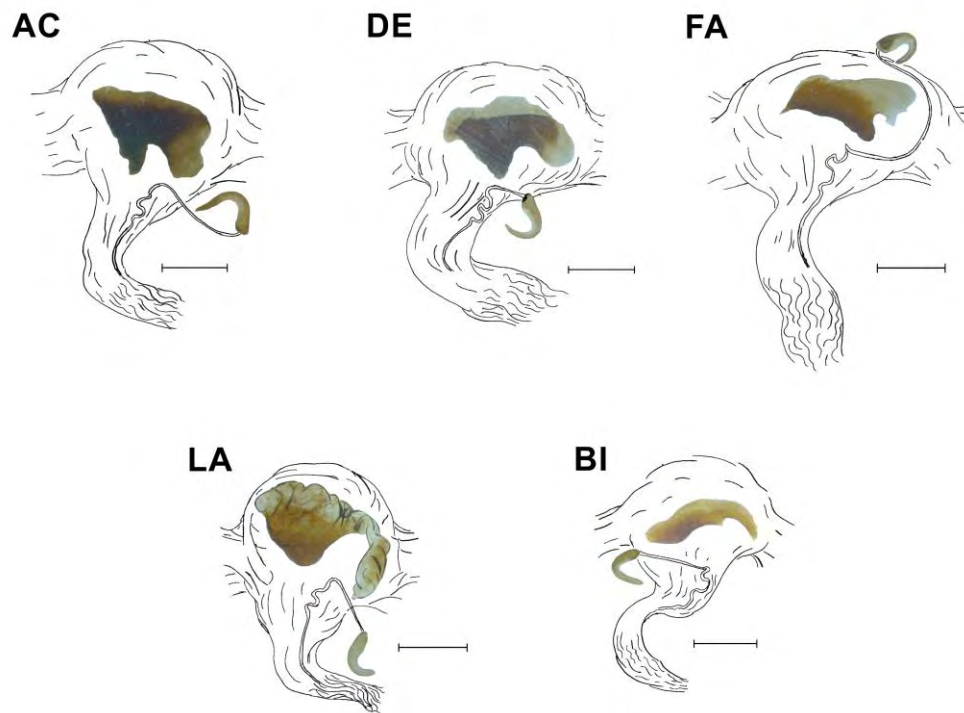
1068

1069

1070

1071

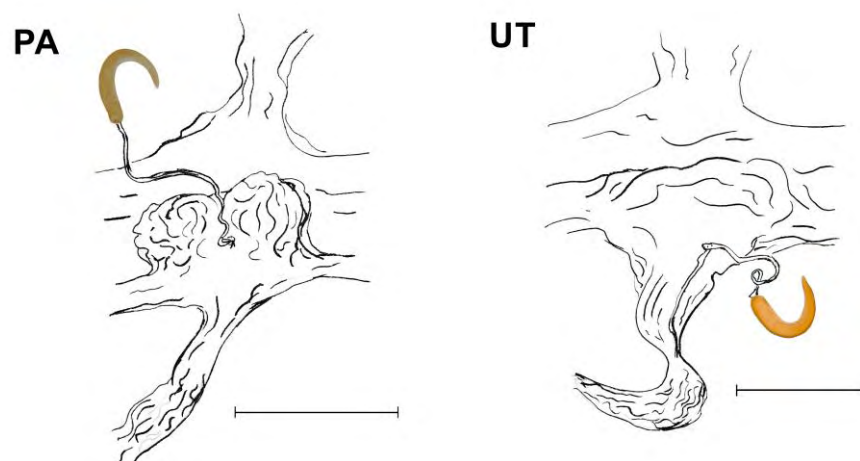
Figure 9. Above, the receptaculum seminis, the features that characterize the two models are clearly represented by *H. bituberculatus* (BI) and *M. pallens* (PA). Scalebar = 0.2 mm. Below, a generic example of the vagina sclerotization (*H. bituberculatus*), after being cleared from membranes and cut off. The various parts were numbered (see text for further details).



1072

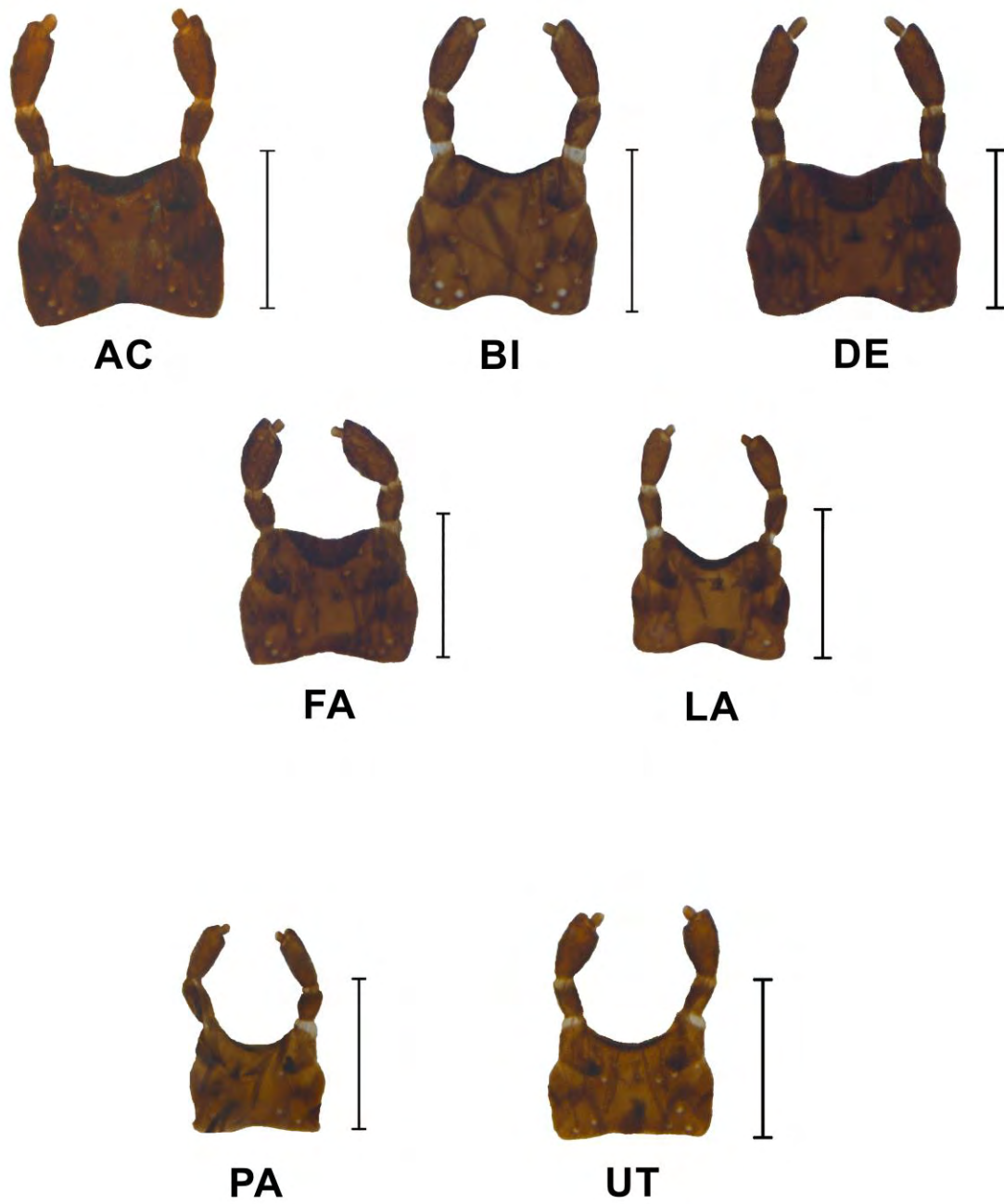
1073 **Figure 10.** Vagina and receptaculum seminis of the species of the genus
 1074 *Hamonthophagus* (AC, BI, DE, FA and LA). Scalebar = 0.5 mm.

1075



1076

1077 **Figure 11.** Vagina and receptaculum seminis of the species of the genus *Moretius* (PA
 1078 and UT). Scalebar = 0.5 mm.

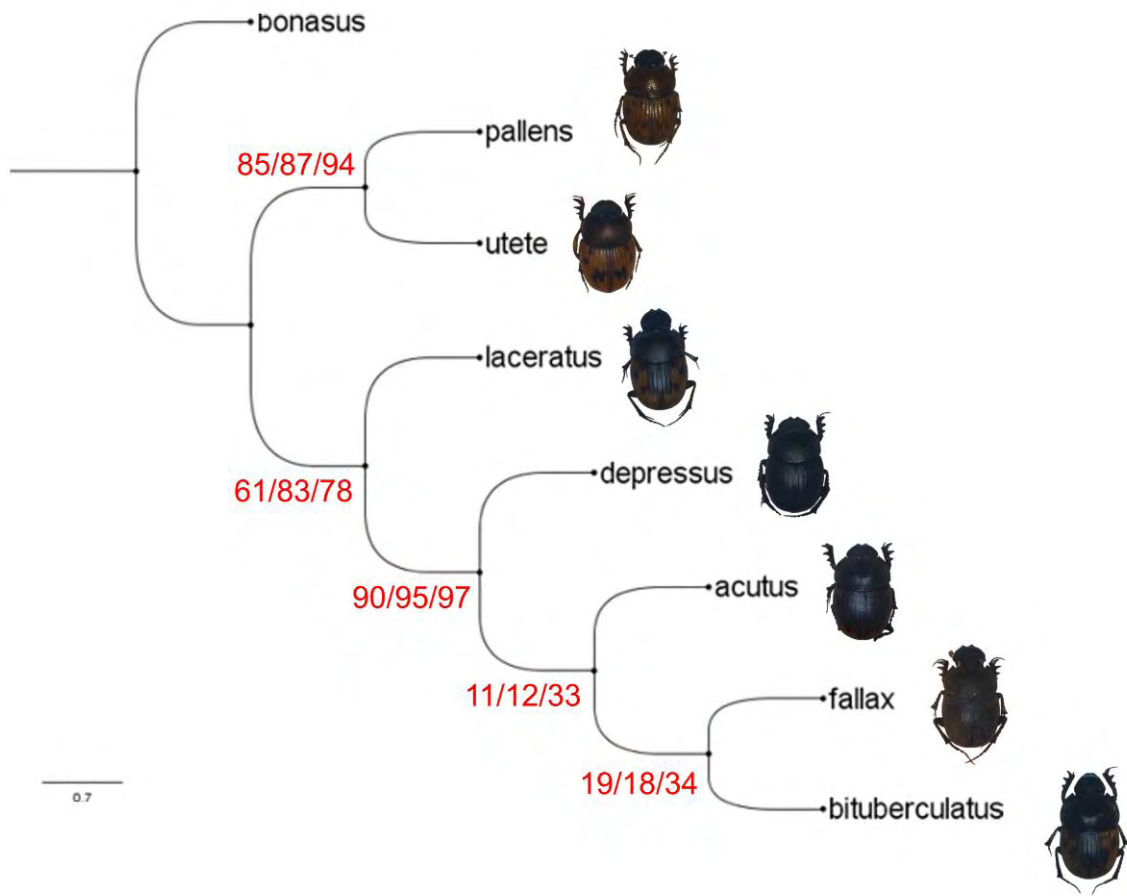


1079

1080 **Figure 12.** Mentum of the species of the genera *Hamonthophagus* (AC, BI, DE, FA and
1081 LA) and *Morettiis* (PA and UT). Scalebar = 0.5 mm.

1082

1083

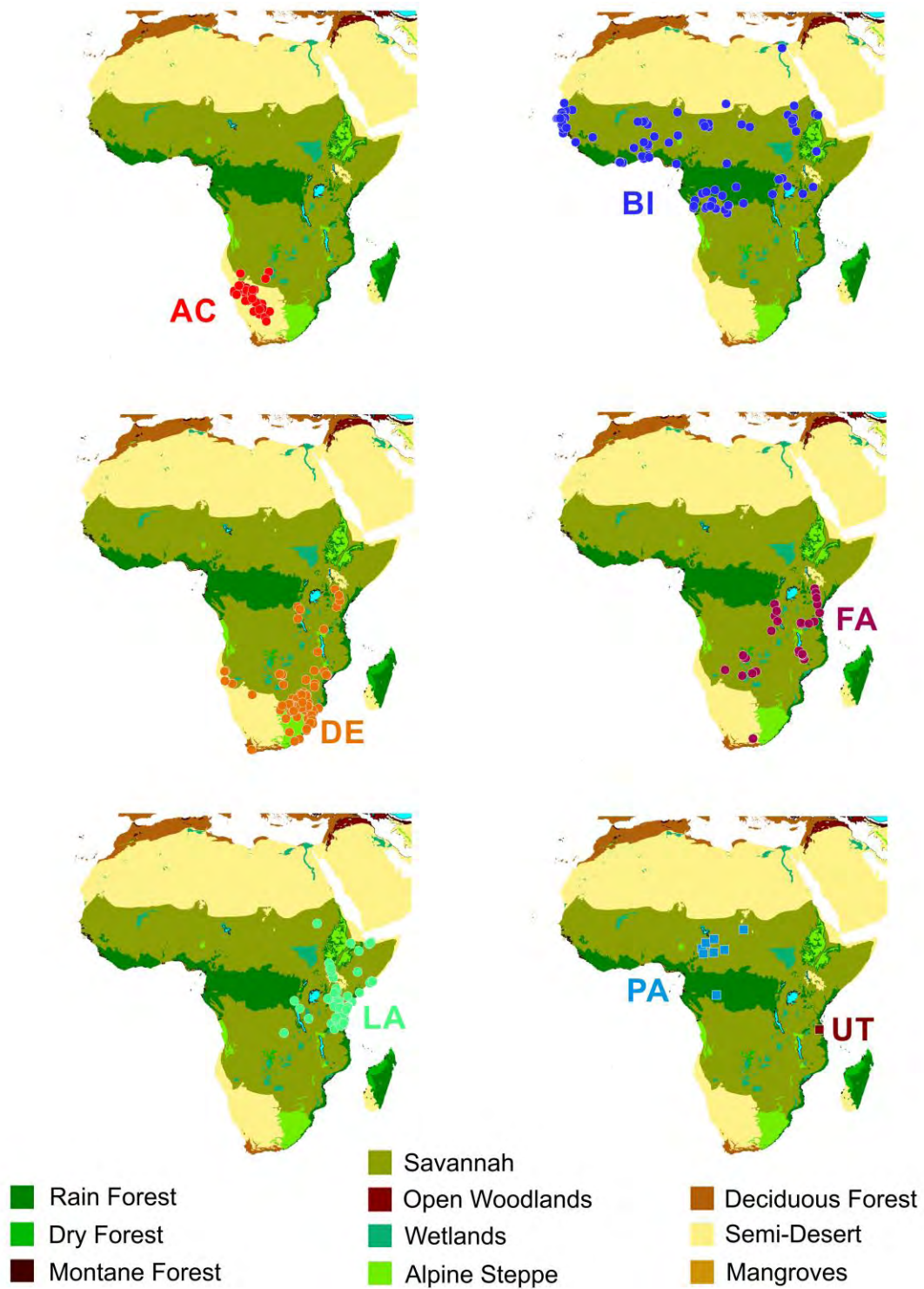


1084

1085 **Figure 13.** Tree from combined analysis, CI = 0.718 and RI = 0.625. Resampling values
 1086 are shown on the branches (Standard Bootstrap, Symmetrical Resampling, and
 1087 Jackknife).

1088

1089

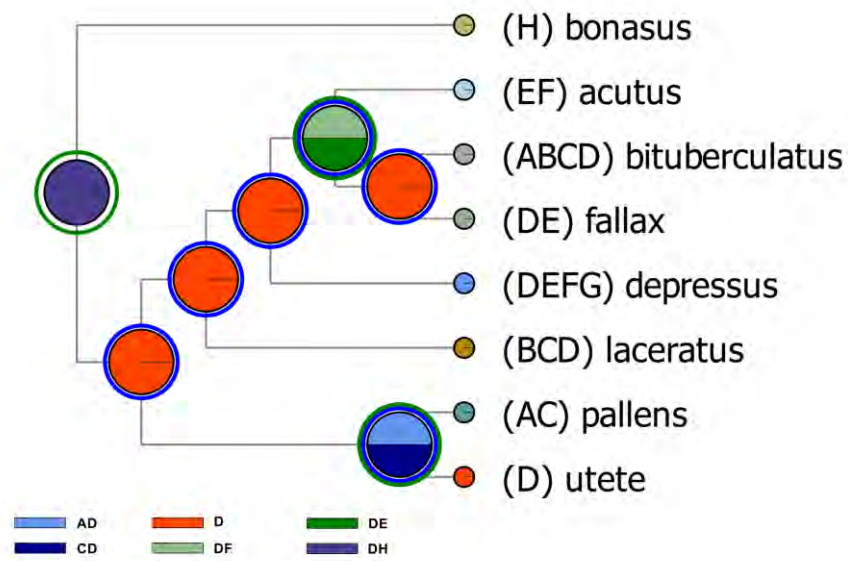


1090

1091 **Figure 14.** Distribution of the species (see Appendix 2 for the list of the localities) with
 1092 the Olson et al. (2000) terrestrial biomes classification.

1093

1094

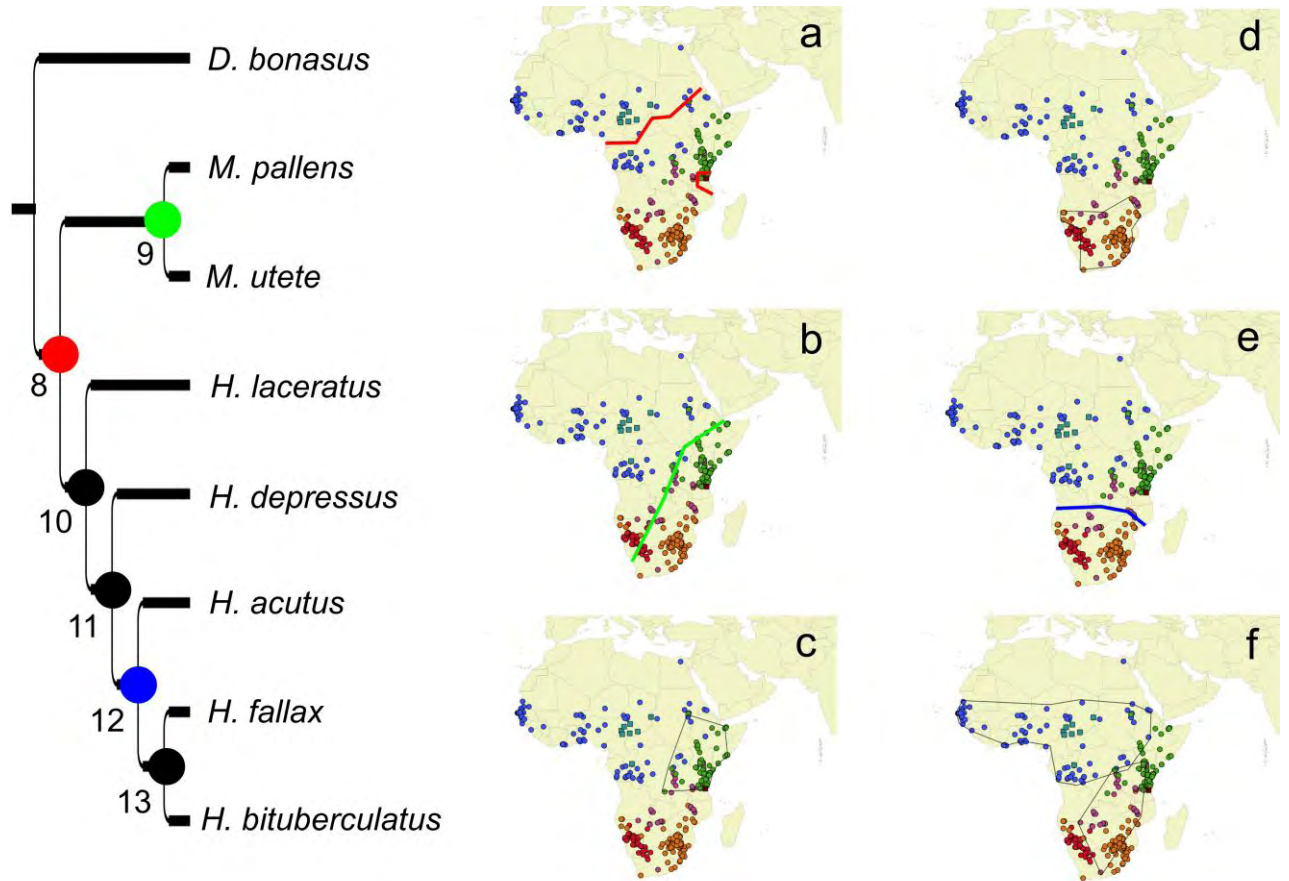


1095

1096 **Figure 15.** Dispersal-Vicariance analysis, with the legend of the ancestral areas. On the
 1097 nodes, the dispersal events are marked by a blue ring, and vicariant events by a green
 1098 ring.

1099

1100



1101

1102 **Figure 16.** VIP analysis, with the vicariant (red, green and blue dots, respectively) and
 1103 dispersal (black dots) events marked on the nodes of the tree. Each node is numbered on
 1104 the tree. The vicariant barriers are shown on the general distribution map (a, b and e),
 1105 while the species distribution is indicated for dispersal events (c, d and f).

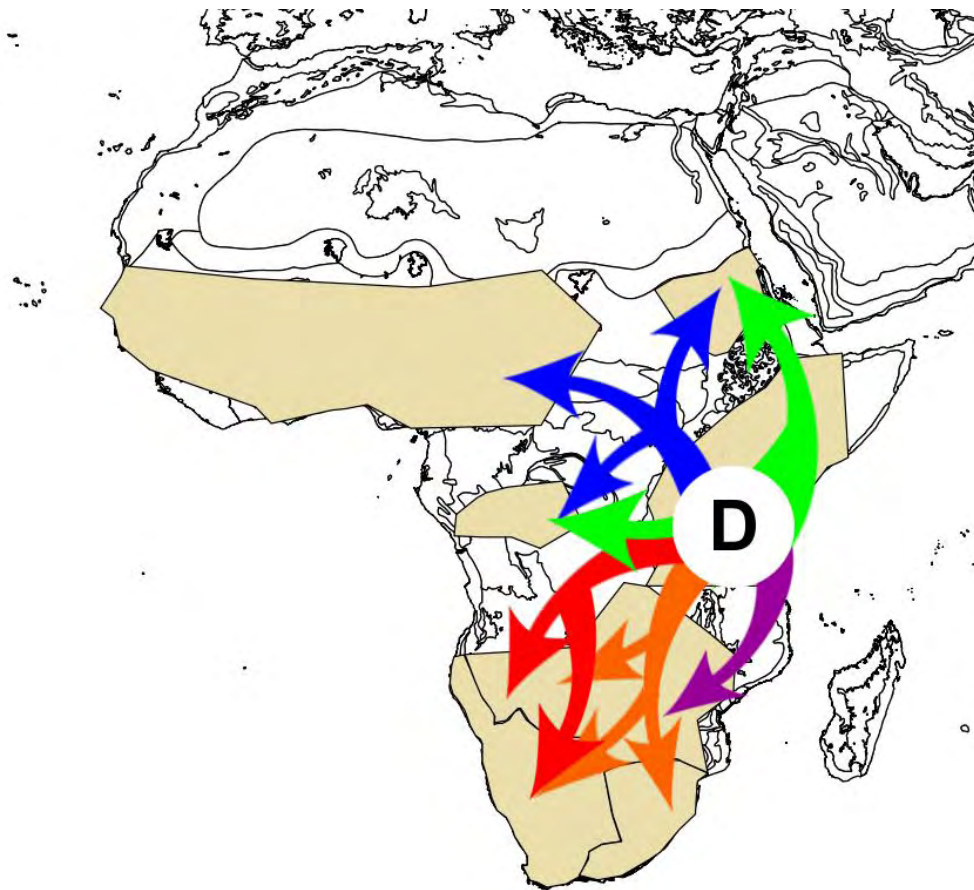
1106



1107

1108 **Figure 17.** *Morettius utete* sp. nov., paratype female facies. Scalebar = 1 mm.

1109



1110

1111 **Figure 18.** Map showing the *Hamonthophagus* dispersal events that have led to the
 1112 current distribution. Blue arrow = *H. bituberculatus*, green arrow = *H. laceratus*, red
 1113 arrow = *H. acutus*, orange arrow = *H. depressus*, and purple arrow = *H. fallax*.

1114