

Taxonomy of the hyper-diverse ant genus *Tetramorium* Mayr in the Malagasy region (Hymenoptera, Formicidae, Myrmicinae) – first record of the *T. setigerum* species group and additions to the Malagasy species groups with an updated illustrated identification key

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Abstract

In this study we provide an update to the taxonomy of the ant genus *Tetramorium* Mayr in Madagascar. We report the first record of the *T. setigerum* species group in Madagascar and describe the only Malagasy representative as *T. cavernicola* sp. n., which is known only from a cave in Ankarana. In addition, we provide an overview of the 19 proposed Malagasy species groups, and discuss their zoogeography and relationships to other groups and larger lineages within the hyper-diverse genus *Tetramorium*. At present, we recognise a highly unique Malagasy *Tetramorium* fauna with 113 species endemic to the island of Madagascar out of a total of 125 translating into an endemism rate of 93%. We hypothesise that this fauna is based on one or a few colonisation events from the Afrotropical region, with subsequent adaptive radiation in Madagascar. Furthermore, we present an updated and illustrated identification key to the *Tetramorium* species groups in the Malagasy region.

Keywords

Endemism, Madagascar, taxonomy, *Tetramorium*, *T. setigerum* species group, zoogeography

Introduction

The genus *Tetramorium* Mayr, widely distributed throughout all zoogeographical regions, is among the most species-rich ant genera in the world. Currently, we recognise around 600 valid species, but because the authors are aware of a larger number of undescribed species, we expect the total count to be closer to 700 or more species. Most *Tetramorium* species are found in the tropics and subtropics of the Old World, where the genus can be considered hyper-diverse by the definition of Wilson (2003), and are key elements of most local ant communities, especially in the Afrotropical and Malagasy regions. Recent studies in the latter region have revealed an astonishingly diverse and highly endemic *Tetramorium* fauna consisting of 107 valid species plus approximately 17 undescribed species (Hita Garcia and Fisher 2011, 2012a, 2012b, 2014b, unpublished data). Based on these figures, *Tetramorium* is by far the most species-rich ant genus in Madagascar, where it seems to have undergone a radiation that was particularly successful in the forested eastern and northern areas of the island.

On a global scale, Bolton (1976, 1977, 1979, 1980) revised the taxonomy of most regional faunas with the exception of the Palaearctic region. These works provided an excellent foundation for the many later revisions or treatments of species groups/complexes or regions/subregions (e.g. Csösz et al. 2007; Csösz and Schulz 2010; Hita Garcia et al. 2010; Hita Garcia and Fisher 2011, 2012a, 2012b, 2013, 2014a, 2014b; Bharti and Kumar 2012; Sharaf et al. 2012; Vásquez Bolaños et al. 2011). The Malagasy *Tetramorium* fauna was first monographed by Bolton (1979), who treated eight species groups with 36 species (29 of these endemic to Madagascar). The later synonymisation of *Triglyphothrix* Forel (Bolton 1985) under *Tetramorium* added an additional species group with one tramp species; two additional tramp species have been recorded since then (Blard et al. 2003; Roberts and McGlynn 2004). This means that 39 Malagasy *Tetramorium* species were known prior to 2011. In that year, we began a large-scale taxonomic revision of the genus for the Malagasy region based initially on more than 160 morphospecies with more than 40,000 mounted specimens. As a foundation for a series of monographs, we proposed 14 species groups for the Malagasy region and provided a preliminary identification key to these groups (Hita Garcia and Fisher 2011). In addition, we revised the *T. bicarinatum*, *T. obesum*, *T. sericeiventre*, and *T. tosii* species groups. In that study we described one species and sank another to the rank of junior synonym, which did not change the total species richness for the region. Based on that work, we revised the *T. bessonii*, *T. bonibony*, *T. dysalum*, *T. kelleri*, *T. marginatum*, *T. tortuosum*, *T. tsingy*, and *T. weitzackeri* species groups shortly afterwards (Hita Garcia and Fisher 2012a, 2012b). These studies treated 58 species, of which 45 were described as new, and raised the species count for the region to 84. We also proposed additional species groups leading to a total of 18 for the region. In the most recent study we revised four additional groups: *T. naganum*, *T. plesiarum*, *T. schaufussii*, and *T. severini* (Hita Garcia and Fisher 2014b). We treated 31 species, of which 22 were newly described, and raised one junior synonym to species status. This increased the current species count for Malagasy *Tetramorium* to 107 (not 106 as mentioned in the introduction of Hita Garcia and Fisher 2014b).

In this study we report the first record of the presence of the Afrotropical *T. setigerum* species group on Madagascar and describe the single representative in the region as a new species, *T. cavernicola* sp. n.. With the *T. setigerum* group, there are now 17 Malagasy species groups that have undergone a current taxonomic revision. Nevertheless, the last two groups, the *T. ranarum* and the *T. simillimum* groups, have not been revised since Bolton (1979). The revisions of these two groups are currently in preparation. In addition to the revision of the *T. setigerum* group, in this study we also present an updated discussion on the currently recognised Malagasy species groups. We give an overview, discuss their biographical affinities, and try to assess their relationships to other key lineages within this hyper-diverse genus. Also, as a consequence of the recent revisions (Hita Garcia and Fisher 2012a, 2012b, 2014b; this study) that proposed additional species groups and added species not known during the preparation of the species group key published in Hita Garcia and Fisher (2011), we present an updated illustrated key to the 19 proposed species groups.

Abbreviations of depositories

The collection abbreviations follow Evenhuis (2014). The material upon which this study is based is located and/or was examined at the following institutions:

BMNH	The Natural History Museum (British Museum, Natural History), London, U.K.
CASC	California Academy of Sciences, San Francisco, California, U.S.A.
HLMD	Hessisches Landesmuseum Darmstadt, Darmstadt, Germany
MCZ	Museum of Comparative Zoology, Harvard University, Cambridge, Massachusetts, U.S.A.
MHNG	Muséum d'Histoire Naturelle de la Ville de Genève, Geneva, Switzerland
NHMB	Naturhistorisches Museum Basel, Basel, Switzerland

Material and methods

The material examined for this study and the previous Malagasy *Tetramorium* revisions (Hita Garcia and Fisher 2011, 2012a, 2012b, 2014b) was collected during ant inventories carried out in the Malagasy region from 1992 to 2013. These inventories included material from more than 6,000 leaf litter samples, 4,000 pitfall traps, and 9,000 additional hand collecting events (see Fisher 2005 for additional details). All new type material and all imaged specimens can be uniquely identified by specimen-level codes affixed to each pin (e.g. CASENT0247028). Digital colour montage images were created using a JVC KY-F75 digital camera and Syncroscopy Auto-Montage software (version 5.0), or a Leica DFC 425 camera in combination with the Leica Application Suite software (version 3.8). All images used for the colour plates illustrat-

ing the identification key or for the presentation of species are available online and can be seen on AntWeb (<http://www.antweb.org>). We predominantly have used images of valid species, but in a few cases used images of undescribed species. The latter have morphospecies codes (e.g. *Tetramorium fhg-forc*) and can be seen under their respective codes on AntWeb. The distribution map provided below was generated with R (R Core Team 2014). Morphometric measurements were performed with a Leica MZ 12.5 equipped with an orthogonal pair of micrometres at a magnification of 100×. Measurements and indices are presented as minimum and maximum values with arithmetic means in parentheses. In addition, all measurements are expressed in mm to two decimal places. The measurements and indices used in this study follow Hita Garcia and Fisher (2011, 2012a, 2012b, 2013, 2014a, 2014b):

- HL** Head length: maximum distance from the midpoint of the anterior clypeal margin to the midpoint of the posterior margin of head, measured in full-face view. Impressions on the anterior clypeal margin and the posterior head margin reduce head length.
- HW** Head width: width of the head directly behind the eyes measured in full-face view.
- SL** Scape length: maximum scape length excluding basal condyle and neck.
- EL** Eye length: maximum diameter of compound eye measured in oblique lateral view.
- PH** Pronotal height: maximum height of the pronotum measured in lateral view.
- PW** Pronotal width: maximum width of the pronotum measured in dorsal view.
- WL** Weber's length: diagonal length of the mesosoma in lateral view from the posteroventral margin of propodeal lobe to the anteriormost point of pronotal slope, excluding the neck.
- PSL** Propodeal spine length: in dorsofrontal view the tip of the measured spine, its base, and the centre of the propodeal concavity between the spines must all be in focus. Using a dual-axis micrometre the spine length is measured from the tip of the spine to a virtual point at its base where the spine axis meets orthogonally with a line leading to the median point of the concavity.
- PTH** Petiolar node height: maximum height of the petiolar node measured in lateral view from the highest (median) point of the node to the ventral outline. The measuring line is placed at an orthogonal angle to the ventral outline of the node.
- PTL** Petiolar node length: maximum length of the dorsal face of the petiolar node from the anterodorsal to the posterodorsal angle, measured in dorsal view excluding the peduncle.
- PTW** Petiolar node width: maximum width of the dorsal face of the petiolar node measured in dorsal view.
- PPH** Postpetiole height: maximum height of the postpetiole measured in lateral view from the highest (median) point of the node to the ventral outline. The measuring line is placed at an orthogonal angle to the ventral outline of the node.
- PPL** Postpetiole length: maximum length of the postpetiole measured in dorsal view.
- PPW** Postpetiole width: maximum width of the postpetiole measured in dorsal view.
- OI** Ocular index: $EL / HW \times 100$

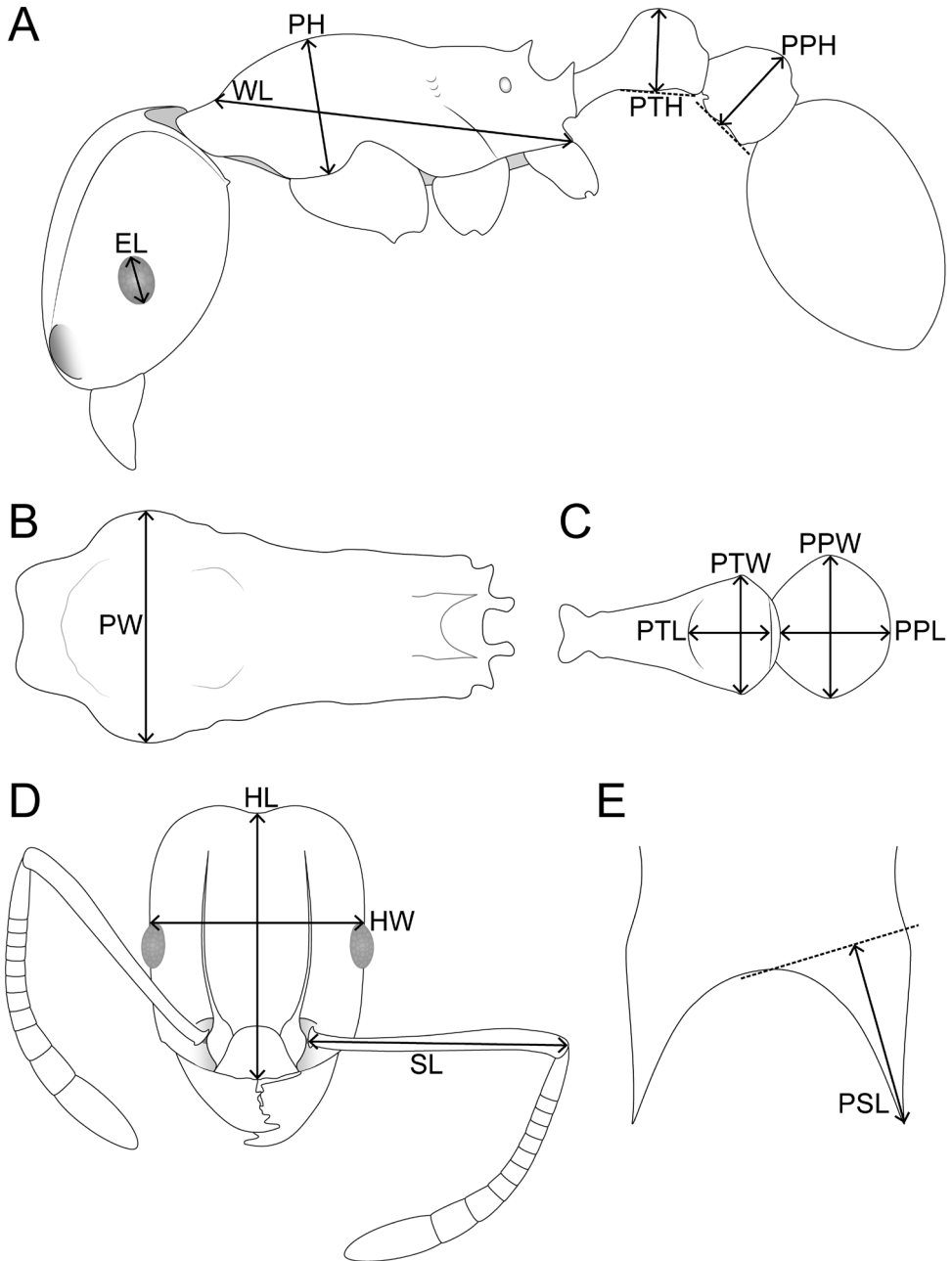


Figure 1. Schematic line drawings of *Tetramorium cavernicola* sp.n. illustrating the used measurements. **A** Body in profile with measuring lines for EL, WL, PH, PTH, and PPH **B** Mesosoma in dorsal view with measuring line for PW **C** Petiole and postpetiole in dorsal view with measuring lines for PTL, PTW, PPL, PPW **D** head in full-face view with measuring lines for HL, HW, and SL **E** Dorsocaudal view of the propodeum with measuring line for PSL.

CI	Cephalic index: $HW / HL \times 100$
SI	Scape index: $SL / HW \times 100$
DMI	Dorsal mesosoma index: $PW / WL \times 100$
LMI	Lateral mesosoma index: $PH / WL \times 100$
PSLI	Propodeal spine index: $PSL / HL \times 100$
PeNI	Petiolar node index: $PTW / PW \times 100$
LPeI	Lateral petiole index: $PTL / PTH \times 100$
DPeI	Dorsal petiole index: $PTW / PTL \times 100$
PpNI	Postpetiolar node index: $PPW / PW \times 100$
LPpI	Lateral postpetiole index: $PPL / PPH \times 100$
DPpI	Dorsal postpetiole index: $PPW / PPL \times 100$
PPI	Postpetiole index: $PPW / PTW \times 100$

Pubescence and pilosity are often of high diagnostic value within the genus *Tetramorium* (e.g. Bolton 1976, 1980, 1985; Hita Garcia et al. 2010, Hita Garcia and Fisher 2012a, 2012b). The varying degrees of inclination of pilosity is particularly important for the diagnosis of groups or species. In this context we use the terms “erect”, “suberect”, “subdecumbent”, “decumbent”, and “appressed” following Wilson (1955). The terminology used for the description of surface sculpturing follows Harris (1979) and Bolton (1980).

Results

Overview Malagasy species groups

The *Tetramorium* ant fauna of the Malagasy region can be divided into 19 species groups that represent different major lineages within this hyper-diverse genus. However, not all groups are native to the region. A proper assessment of the biogeographical affinities of a region, such as Madagascar and its surrounding South West Indian Ocean island systems, is only possible if comprehensive knowledge on the origin and distribution of each species group is available. This is of special importance when dealing with hyper-diverse genera that possess hundreds of species and dozens of evolutionary lineages throughout most or all zoogeographical regions. Fortunately, in the case of Malagasy *Tetramorium* we are able to assess the whole fauna and classify the groups into native, exotic, or shared with the Afrotropical region. Six of the 19 groups are either completely exotic, contain partly global tramps, or species of African origin. Two of these, the *T. bicarinatum* and *T. obesum* groups, are only present in the region with a few very successful global tramp species that very likely originated in the Oriental and Indo-Australian regions (Bolton 1977, 1979; McGlynn 1999; Wetterer 2010; Hita Garcia and Fisher 2011). Four other groups are clearly Afrotropical in origin: the *T. sericeiventris*, *T. setigerum*, *T. simillimum*, and *T. weitzackeri* groups. The Malagasy representatives of these groups are either species that have likely been

recently transferred from eastern and/or southern Africa by humans, or species that are Malagasy endemics that have evolved from much older colonisation events from Africa, or combinations of both. The *T. simillimum* group in the Malagasy region is a good example of the latter. It contains two global tramp species of African origin (*T. caldarium* (Roger) and *T. simillimum* (Smith)), one very widespread non-tramp species of African origin (*T. delagoense* Forel), as well as some species endemic to the Malagasy region (*T. anodontion* Bolton, *T. scytalum* Bolton, and a few undescribed species). The *T. sericeiventris* group has one species that is widespread in the southern Palearctic, Afrotropical, and Malagasy regions (*T. sericeiventris* Emery) and one species endemic to Madagascar (*T. mahafaly* Hita Garcia & Fisher). The *T. weitzckeri* group is represented by *T. humbloti* Forel, a widespread species distributed in eastern and southern Africa that is probably a more recent introduction to the Malagasy region. The *T. setigerum* group, here recorded for the first time from the Malagasy region, contains one recently discovered species endemic to Madagascar that we describe below.

Most of the abovementioned, mostly non-native, groups possess twelve-segmented antennae and a triangular to dentiform sting appendage (except the *T. weitzckeri* group, which has eleven-segmented antennae and a spatulate sting appendage). The only other Malagasy species group with twelve-segmented antennae is the *T. tosii* group, which seems to be endemic to Madagascar (Bolton 1979; Hita Garcia and Fisher 2011). In Hita Garcia and Fisher (2011) we discussed the very strong morphological similarities between the *T. tosii* group and some Afrotropical members of the *T. setigerum* group, which was not known from Madagascar at the time. At present, we propose to keep both groups separate until more data becomes available, but provide a more thorough discussion in the species group treatment of the *T. setigerum* group below.

One intriguing finding of the recent revisions (Hita Garcia and Fisher 2011, 2012a, 2012b, 2014b) is that the vast majority of native species groups (*T. bessonii*, *T. bonibony*, *T. dysalum*, *T. kelleri*, *T. marginatum*, *T. naganum*, *T. plesiarum*, *T. ranarum*, *T. schaufussi*, *T. severini*, *T. tsingy*, and *T. tortuosum*) described from Madagascar share the three following key characters: eleven-segmented antennae, anterior clypeal margin notched, and sting appendage spatulate. These twelve groups together contain more than 110 species that are endemic to Madagascar, plus three species found only on Mayotte or the Comoros. The only other group with eleven-segmented antennae, notched anterior clypeal margin, and spatulate sting appendage is the *T. weitzckeri* group. However, as noted above, *T. humbloti* represents a recent arrival from the Afrotropical region and did not evolve independently in Madagascar (Hita Garcia and Fisher 2011). Not considering the latter group, most native species groups could have originated from one or a few ancient colonisation events, with subsequent adaptive radiation that is most pronounced in the humid forests of eastern and northern Madagascar. Assessing the geographic origins of the first colonists with eleven-segmented antennae appears challenging at first glance. A number of species groups have eleven-segmented antennae and a spatulate sting appendage in the Afrotropical, Oriental, and Indo-Australian regions. However, the species morphologically closest to the Malagasy groups are mostly found in Africa.

The most closely related ants seem to belong to the comparatively species-rich *T. weitzackeri* species group (Bolton 1980; Hita Garcia et al. 2010). The group is very widespread and ecologically successful in sub-Saharan Africa, and has both eleven-segmented antennae and a spatulate sting appendage. Many of its species resemble a number of members of Malagasy groups, such as the *T. bessonii* group, parts of the *T. bonibony* group, parts of the *T. dysalum* group, and parts of the *T. marginatum* group. The older species of these groups were initially even placed in the *T. weitzackeri* group (Bolton 1979) until recent rearrangements of the Malagasy species group system (Hita Garcia and Fisher 2011, 2012a, 2012b, 2014b). In Hita Garcia and Fisher (2011) we proposed to treat the Malagasy members (with the exception of *T. humbloti* that stayed in the group) as independent developments from the *T. weitzackeri* group and created the new *T. bessonii*, *T. bonibony*, *T. dysalum*, and *T. marginatum* groups. These groups contain a high degree of morphological diversity often differing from the Afrotropical *T. weitzackeri* group. One reason for the distinctiveness of the Malagasy groups was the shape of the petiolar node. We stated that in the *T. weitzackeri* group the node is often squamiform with anterior and posterior faces approximately parallel, whereas in Madagascar this form of node is only present in few species of the *T. dysalum* group. Most other species of the groups in question have petiolar nodes that are anteroposteriorly compressed, often very strongly so, but with a more triangular to cuneiform shape, the node narrowing towards the dorsum, or the anterodorsal margin much more angled than the posterodorsal margin causing the dorsum to strongly taper backward posteriorly. The dorsum of the node is greatly reduced in a number of species, especially from the *T. bonibony* and *T. marginatum* groups. Despite being absent in the *T. weitzackeri* group, this more triangular node shape is also found in the Afrotropical *T. squaminode* group, which is also predominantly East and South African. This group also has a spatulate sting appendage but twelve-segmented antennae, and despite this difference seems to be closely related to the *T. weitzackeri* group (Bolton 1980; Hita Garcia et al. 2010). Nevertheless, the argument about the difference in node shape is less valid now after the recent description of *T. mpala* Hita Garcia & Fischer (Hita Garcia and Fischer 2014). This interesting species from Kenya belongs to the *T. weitzackeri* group but has a more triangular squamiform node shape like in the Malagasy groups mentioned above. However, instead of reuniting several species groups with more than 70 species, we prefer to wait until more data is available. Morphological similarities in as diverse a genus as *Tetramorium* can be misleading, and better taxonomic resolution on a supraspecific level can only be achieved with a large-scale analysis that combines morphology with informative molecular phylogenetic or phylogenomic data. Regardless, the Malagasy *T. bessonii*, *T. bonibony*, *T. dysalum*, and *T. marginatum* groups are very likely part of a larger *Tetramorium* lineage that also includes the Afrotropical *T. weitzackeri* and *T. squaminode* groups, even though the relationships remain unclear at present.

The Malagasy *T. naganum*, *T. schaufussi*, and *T. severini* groups (and parts of the *T. dysalum* group) also appear to have a strong African influence since they share a spatulate sting appendage, high nodiform petiolar node shape, and a lack of any sculpture

on the waist segments with the South African *T. grassii* group, even though the latter group has twelve-segmented antennae. Bolton (1980) stated that the *T. grassii* group is related and possibly ancestral to the *T. weitzckeri* and *T. squaminode* groups. We concur that these three groups are very likely closely related. Based on their unique morphology, the *T. plesiarum*, *T. ranarum*, and *T. tsingy* groups seem to be independent Malagasy developments since there are no species groups with similar morphology in any region. However, they are very probably also part of the same larger lineage as the other species groups with eleven-segmented antennae and a spatulate sting appendage. This leads us to hypothesise that there is a larger Afrotropical and Malagasy *Tetramorium* clade/lineage that contains the following groups: *T. bessonii*, *T. bonibony*, *T. dysalum*, *T. grassii*, *T. marginatum*, *T. naganum*, *T. plesiarum*, *T. ranarum*, *T. schaufussi*, *T. severini*, *T. squaminode*, *T. tsingy*, and *T. weitzckeri* species groups. The situation for the *T. kelleri* and *T. tortuosum* groups is less clear. These two groups are very close and were separated recently on the basis of their distinctiveness in Madagascar (Hita Garcia and Fisher 2012b). However, the *T. tortuosum* group is present in the Neotropical, Afrotropical, Malagasy, Oriental, and Indo-Australian regions, and displays great differences in morphological diversity and species richness from region to region (Bolton 1977, 1979, 1980; Hita Garcia and Fisher 2012b, 2013). It is difficult to assess if all the species now listed as *T. tortuosum* group indeed belong to one very widespread, possibly very old, monophyletic clade, or some have evolved independently to share several key morphological characters. Consequently, we cannot assess with any certainty whether the Malagasy *T. kelleri* and *T. tortuosum* groups are more closely related to the African or Asian members of the *T. tortuosum* group, or represent a more independent lineage.

In summary, we were able to identify a highly unique Malagasy *Tetramorium* fauna. We recognise 12 of the 19 species groups and an astonishing 113 of the 125 species as Malagasy endemics. This results in an endemism rate of 93%, which is more or less in agreement with the published value for the whole Malagasy ant fauna (ca. 96% in Fisher 2003). In Table 1 we provide an overview of the Malagasy species groups with data on their biogeography, key characters, taxonomic revisions, and preferred habitats.

Identification key to *Tetramorium* species groups in the Malagasy region (workers)

The species group key presented here is based on the one published in Hita Garcia and Fisher (2011). Although the key in that publication still works for most species in Madagascar, it does not accommodate them all. The recent revisions of most species groups, with the establishment of some new groups (Hita Garcia and Fisher 2012a, 2012b, 2014b), require an updated and improved key. The following key applies to the 19 groups we currently recognise, which contains around 125 species (the *T. ranarum* and *T. simillimum* groups will be revised in a future publication and the species count for these two groups is temporary).

Table 1. Overview of all 19 Malagasy species groups recognised in this study. For each group we provide number of Malagasy species, zoogeographical affinities, number of antennal segments, shape of sting appendage and anterior clypeal margin, the last taxonomic revision, and habitat preferences. The following abbreviations are used for zoogeographical affinities: AFR=Afrotropical, INA=Indo-Australian, MAD=only Madagascar, MAL=Malagasy (Madagascar plus islands of the Southwest Indian Ocean), NEA=Nearctic, ORI=Oriental, T=panglobal tramp.

Number Malagasy spp.	Species group name	Zoogeography	Antennal segments	Sting appendage	Anterior clypeal margin	Recent taxonomic revision	Habitat preferences
6	<i>bessonii</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2012a	dry forests, savanna, grassland, anthropogenic habitats
8	<i>bonibony</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2012a	dry forests, savanna, grassland, anthropogenic habitats
10	<i>dysalum</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2012a	predominantly lowland or montane rainforests
2	<i>kelleri</i>	MAL	11	spatulate	notched	Hita Garcia and Fisher 2012a	dry and humid forests
6	<i>marginatum</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2012a	lowland or montane rainforests
5	<i>naganum</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2014	lowland or montane rainforests
5	<i>plesiarum</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2014	dry forests, savanna, grassland
21	<i>ranarum</i>	MAD	11	spatulate	notched	in preparation	predominantly lowland or montane rainforests
20	<i>schaufussi</i>	MAL	11	spatulate	notched	Hita Garcia and Fisher 2014	mostly lowland or montane rainforests, rarely dry forests or open habitats
1	<i>severini</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2014	lowland or montane rainforests
2	<i>tsingy</i>	MAD	11	spatulate	notched	Hita Garcia and Fisher 2012a	dry forest
22	<i>tortuosum</i>	NEA, AFR, MAD, ORI & INA	11	spatulate	notched	Hita Garcia and Fisher 2012b	predominantly lowland or montane rainforests
1	<i>weitzeckeri</i>	AFR & MAL	11	spatulate	notched	Hita Garcia and Fisher 2011	dry forests, savanna, grassland, anthropogenic habitats
3	<i>bicarinatum</i>	AFR, ORI, INA & MALT	12	triangular to dentiform	notched	Hita Garcia and Fisher 2011	habitat generalist
1	<i>obesum</i>	ORI, INA & MAL, T	12	triangular to dentiform	notched	Hita Garcia and Fisher 2011	habitat generalist
2	<i>sericeiventre</i>	AFR & MAL	12	triangular to dentiform	entire	Hita Garcia and Fisher 2011	anthropogenic habitats, spiny forest, thicket, coastal and littoral forests, woodland
1	<i>setigerum</i>	AFR & MAD	12	triangular to dentiform	entire	in this study	dry forest
7	<i>simillimum</i>	AFR, MAL & MAL, T	12	triangular to dentiform	entire	in preparation	habitat generalist
2	<i>tosii</i>	MAD	12	triangular to dentiform	entire	Hita Garcia and Fisher 2011	lowland or montane rainforests
125							

- 1 Species with distinctly branched hairs, usually a mixture of simple, bifid, and trifid hairs (Fig. 2A, B).....***T. obesum* group**
- Species without branched hairs; hairs present neither bifid nor trifid, either with simple pilosity (Fig. 2C), or with reduced pilosity but short appressed pubescence (Fig. 2D).....**2**

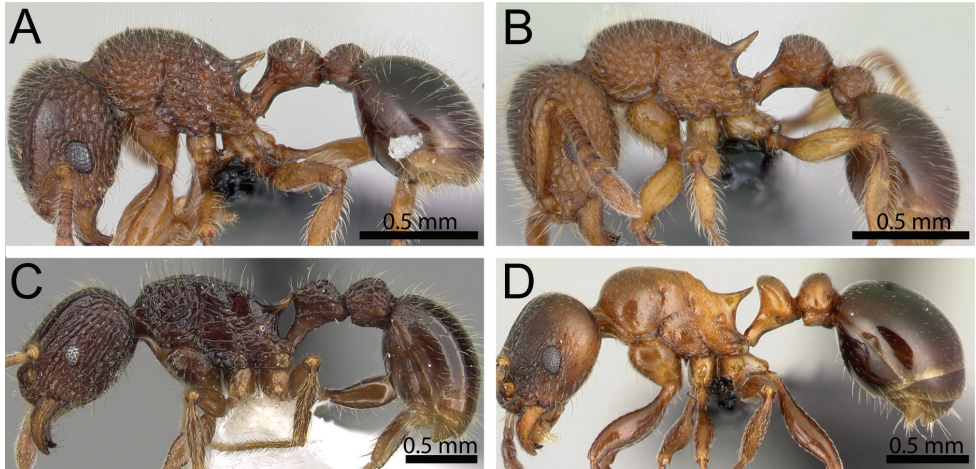


Figure 2. Body in profile. **A, B** *T. lanuginosum* (CASENT0060515; CASENT0125328) **C** *T. singletonae* (CASENT0247161) **D** *T. wardi* (CASENT0475483).

- 2 Antennae 12-segmented (Fig. 3A); sting appendage triangular to dentiform, acute apically (Fig. 3B)**3**
- Antennae 11-segmented (Fig. 3C); sting appendage spatulate, broadened apically (Fig. 3D)**7**

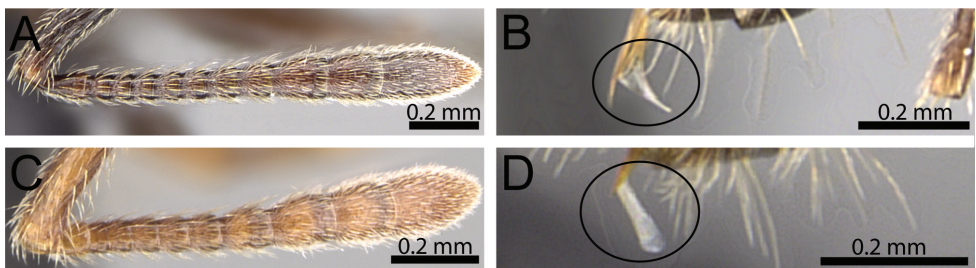


Figure 3. Antennal funiculus and sting appendage (within black ellipses). **A, B** *T. tosii* (CASENT0249662) **C** *T. jedi* (CASENT0043578) **D** *T. hobbit* (CASENT0019207).

- 3 Anterior clypeal margin with distinct median impression (Fig. 4A)..... *T. bicarinatum* group
- Anterior clypeal margin always entire and convex, never with distinct median impression (Fig. 4B, C) 4

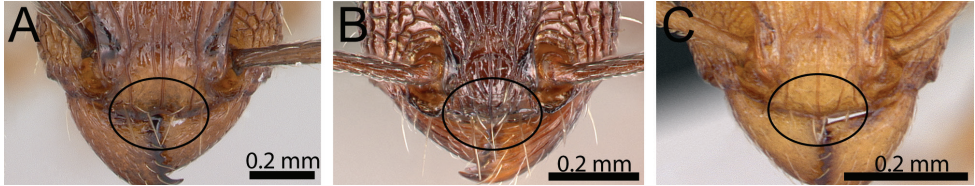


Figure 4. Anterior head in dorsal view (anterior clypeal margin within black ellipses). **A** *T. bicarinatum* (CASENT0060334) **B** *T. mahafaly* (CASENT0448984) **C** *T. simillimum* (CASENT0135001).

- 4 Propodeum armed with long to extremely long spines (PSLI 30–49), at least 2 to 3 times longer than metapleural lobes (Fig. 5A)..... *T. tosii* group
- Propodeum either unarmed (Fig. 5B), armed with small triangular teeth or denticles, or armed with medium-sized spines (Fig. 5C), propodeal spines at most only as long or weakly longer than metapleural lobes, more often propodeal spines distinctly shorter than metapleural lobes 5
- 5 Lateral portion of clypeus prominent, raised to a tooth or denticle in full-face view (Fig. 4B); propodeal spines medium-sized and spinose, approximately of same length as metapleural lobes (Fig. 5C) *T. sericeiventre* group
- Lateral portion of clypeus never modified as above (Fig. 4C); propodeum either unarmed (Fig. 5B) or armed with small triangular teeth or denticles that are shorter than metapleural lobes (Fig. 6E, F)..... 6

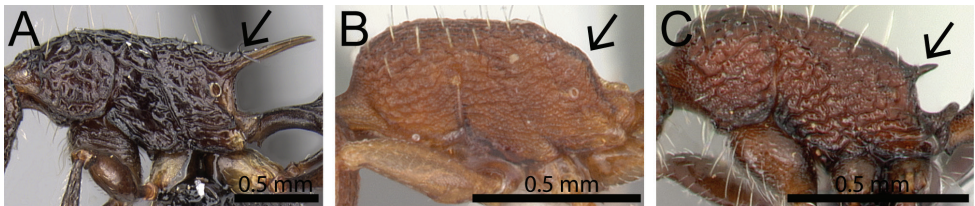


Figure 5. Mesosoma in profile (black arrows indicate propodeal spines/teeth area). **A** *T. tosii* (CASENT0249662) **B** *T. anodontion* (CASENT0102334) **C** *T. mahafaly* (CASENT0449159).

- 6 Head in full-face view relatively thin (CI < 80) and antennal scapes very long (SI > 120) (Fig. 6A); in general appearance head, antennae, and legs elongate and slender (Fig. 6E) *T. setigerum* group
- Head in full-face view relatively thicker (CI > 85) and antennal scapes conspicuously much shorter (SI < 92) (Fig. 6B, C, D); in general appearance more compact species with thicker heads and shorter antennae and legs (Fig. 6F)..... *T. simillimum* group

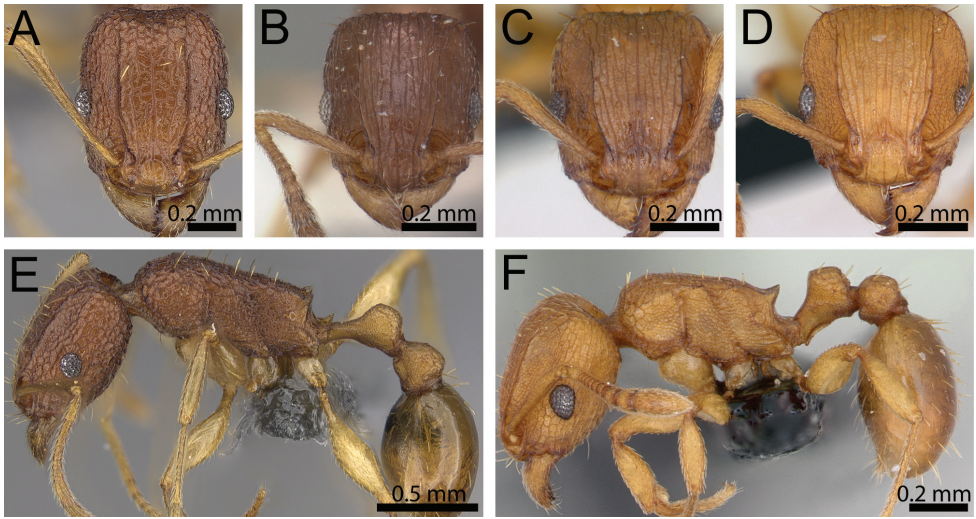


Figure 6. Head in full-face view and body in profile. **A, E** *T. cavernicola* (CASENT0373132) **B** *T. scytalum* (CASENT0102337) **C** *T. caldarium* (CASENT0125225) **D, F** *T. simillimum* (CASENT0135001).

- 7 Petiolar node and postpetiole strongly squamiform, petiolar node with anterior and posterior faces parallel and well developed, straight dorsum; petiole and postpetiole always completely unsculptured, smooth, and shining; standing pilosity scarce or absent on dorsal mesosoma and waist segments, first gastral tergite without standing pilosity (Fig. 7A)..... ***T. weitzackeri* group**
- Character combination never as above; petiole and postpetiole variably shaped, especially postpetiole never squamiform as above (Fig. 7B, C, D, E, 8C, D); if petiole squamiform and postpetiole weakly squamiform, then petiolar dorsum reduced and strongly tapering backwards posteriorly (Fig. 7F).....**8**

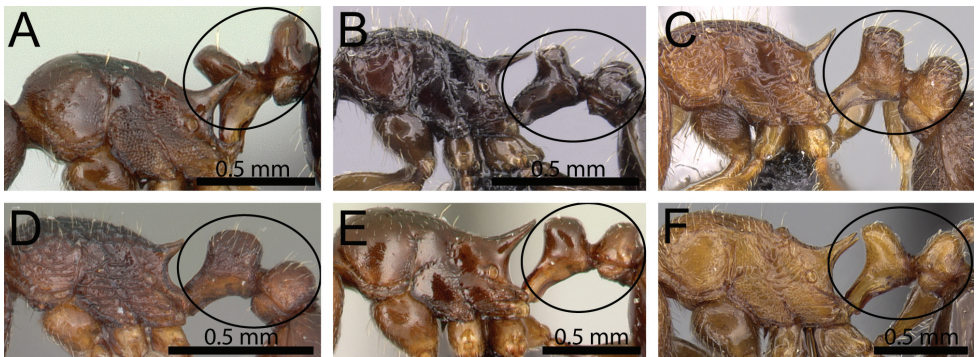


Figure 7. Mesosoma and waist segments in profile (waist segments within black ellipses). **A** *T. humbloti* (CASENT0134851) **B** *T. ambatovy* (CASENT0124721) **C** *T. gollum* (CASENT0074974) **D** *T. quasirum* (CASENT0102353) **E** *T. malagasy* (CASENT0449550) **F** *T. bessonii* (CASENT0247550).

- 8 Pronotum anterodorsally with distinct protuberance or bulge (Fig. 8A, B) ...
..... ***T. bonibony* group (in part)**
- Pronotum anterodorsally without any protuberance or bulge (Fig. 8C, D)...9

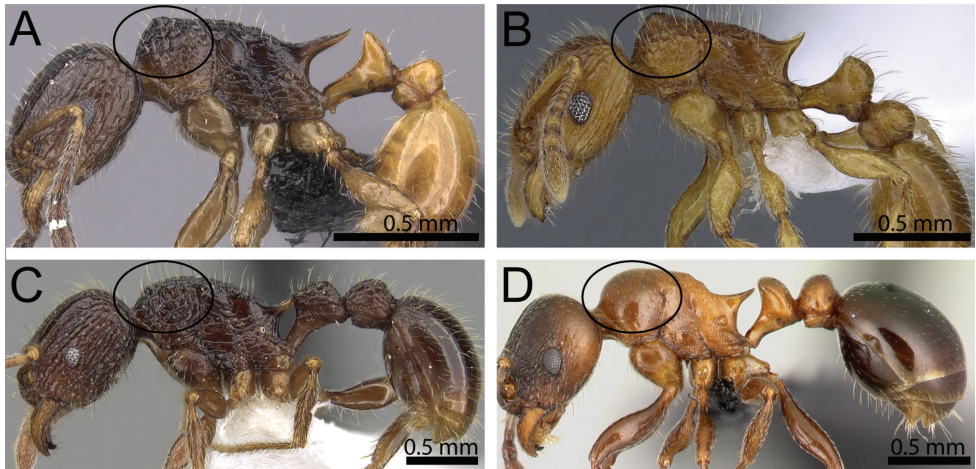


Figure 8. Body in profile (anterodorsal pronotum within black circles). **A** *T. bonibony* (CASENT0486252) **B** *T. trafo* (CASENT0404104) **C** *T. singletonae* (CASENT0247161) **D** *T. wardi* (CASENT0475483).

- 9 First gastral tergite with strongly appressed pubescence of varying length and without any standing hairs (Fig. 9A, B), or with short appressed to erect pilosity without any long, erect to suberect hairs (Fig. 9C) **10**
- First gastral tergite usually with long, erect to suberect pilosity (Fig. 9D, E, F) **17**

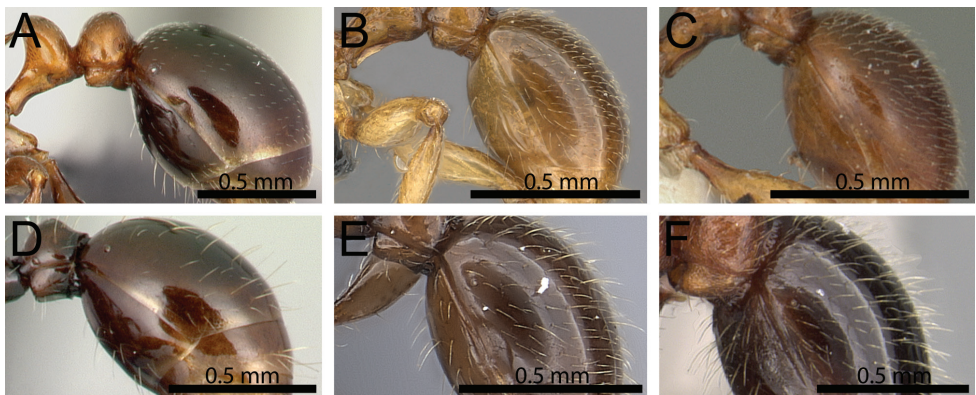


Figure 9. First gastral tergite in profile. **A** *T. wardi* (CASENT0475483) **B** *T. cognatum* (CASENT0067891) **C** *T. naganum* (CASENT0102345) **D** *T. silvicola* (CASENT0042828) **E** *T. alperti* (CASENT0042547) **F** *T. hobbit* (CASENT0019207).

- 10 Antennal scrobes well developed with sharply defined posterior and ventral margins (Fig. 10A, B) ***T. ranarum* group (in part)**

- Antennal scrobes usually weakly developed, never with well-defined posterior and ventral margins (Fig. 10C, D, E)..... 11

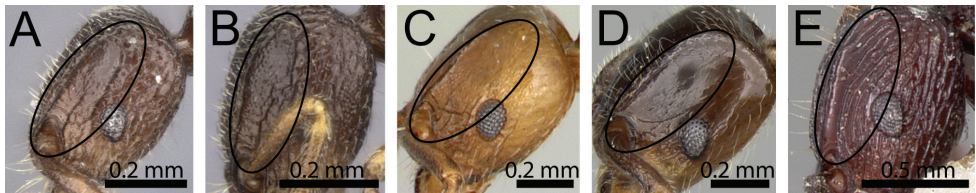


Figure 10. Lateral head in profile (antennal scrobe area within black ellipses). **A** *T. ibycterum* (CASENT0056460) **B** *T. fhg-bilb* (CASENT0448625) **C** *T. artemis* (CASENT0481732) **D** *T. tyriorum* (CASENT0249085) **E** *T. latreillei* (CASENT0101292).

- 11 In profile petiolar node rectangular nodiform with sharply angled anterodorsal and posterodorsal margins; both waist segments strongly sculptured (Fig. 11A, B)..... 12
- Petiolar node rectangular nodiform with conspicuously rounded anterodorsal and/or posterodorsal margins (Fig. 11C), high rounded nodiform (Fig. 11D), or squamiform and strongly anteroposteriorly compressed (Fig. 11E); waist segments usually completely unsculptured, smooth, and shining, rarely with very weak (Fig. 11D, E), superficial sculpture (Fig. 11C)..... 13

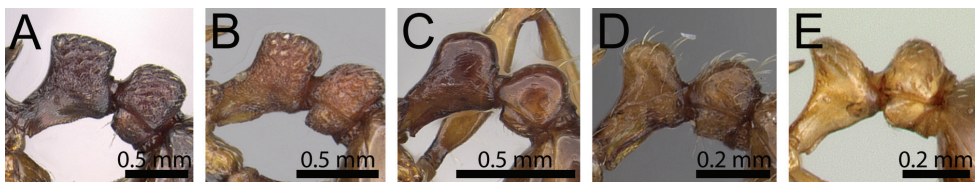


Figure 11. Petiole and postpetiole in profile. **A** *T. fhg-ants* (CASENT0248302) **B** *T. fhg-anub* (CASENT0404158) **C** *T. nassonowii* (CASENT0195504) **D** *T. naganum* (CASENT0280584) **E** *T. artemis* (CASENT0481732).

- 12 Propodeum armed with short to moderately long spines (PSLI 18–25); dorsum of head and mesosoma without any standing pilosity (Fig. 12A, B)..... *T. ranarum* group (in part)
- Propodeum armed with very long spines (PSLI 50–53); dorsum of head and mesosoma with standing pilosity (Fig. 12C) .. *T. tortuosum* group (in part)

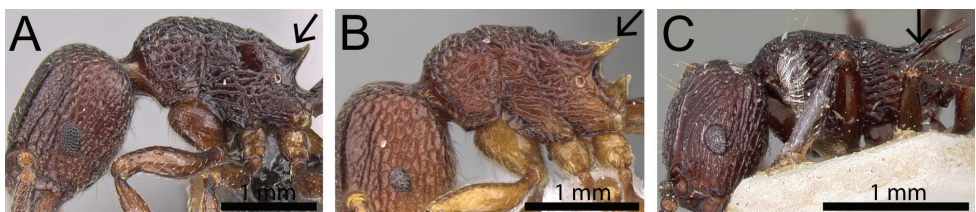


Figure 12. Posterior head and mesosoma in profile (black arrows indicate propodeal spines/teeth). **A** *T. fhg-ants* (CASENT0248302) **B** *T. fhg-anub* (CASENT0404158) **C** *T. latreillei* (CASENT0101292).

- 13 Larger species (HW 0.82–1.03; WL 1.14–1.48); mesosoma comparatively long and slender (LMI 35–37) without distinct margination between lateral and dorsal mesosoma; propodeal spines always very long (PSLI 38–43); body colouration always dark brown to black (Fig. 13A) ***T. severini* group**
- Character combination never as above, especially mesosoma either with strong margination between sides and dorsum (Fig. 13B, C, E, F) or relatively shorter, high, and compact (Fig. 13D); usually much smaller species (HW 0.43–0.87; WL 0.54–1.22) with almost always brighter body colour, but if body size in range of above, then either mesosoma clearly higher and more compact (LMI 42–49) (Fig. 13D) or propodeal spines/teeth much shorter (PSLI 12–28) (Fig. 13B) 14

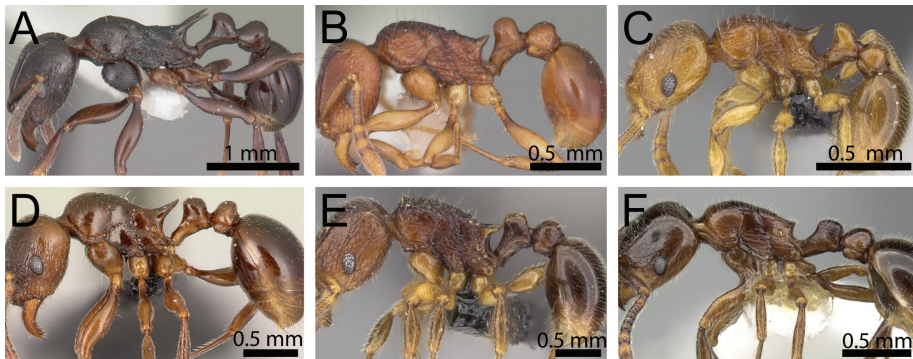


Figure 13. Body in profile. **A** *T. severini* (CASENT0102397) **B** *T. proximum* (CASENT0102342) **C** *T. rumo* (CASENT0073025) **D** *T. malagasy* (CASENT0449550) **E** *T. dalek* (CASENT0038402) **F** *T. tyrion* (CASENT0249085).

- 14 Dorsum of mesosoma generally completely unsculptured (Fig. 14A), very rarely with few superficial, weak rugulae laterally, but median area of pronesonotum always unsculptured and shiny (Fig. 14B, C); mandibles always conspicuously sculptured 15
- Mesosoma usually strongly sculptured (Fig. 14E, F), very rarely with weak rugulose sculpture including median area of pronesonotum (Fig. 14D), but then mandibles completely unsculptured and very smooth and shiny 16

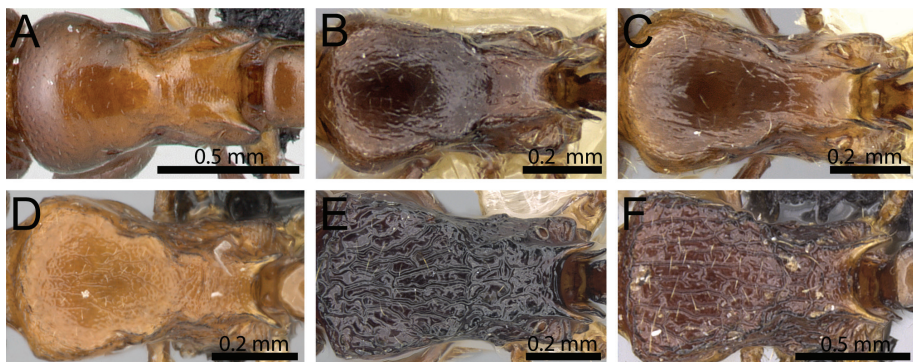


Figure 14. Mesosoma in dorsal view. **A** *T. wardi* (CASENT0475483) **B** *T. tyrion* (CASENT0249085) **C** *T. tsingy* (CASENT0426807) **D** *T. freya* (CASENT0466944) **E** *T. camelliae* (CASENT0247496) **F** *T. dalek* (CASENT0038402).

- 15 Propodeal spines long and metapleural lobes short (Fig. 15A); petiolar node in dorsal view distinctly wider than long (Fig. 15B)..... ***T. bessonii* group (in part)**
 – Propodeal spines/teeth comparatively short and metapleural lobes of almost similar size (Fig. 15C); petiolar node in dorsal view longer than wide (Fig. 15D)..... ***T. tsingy* group**

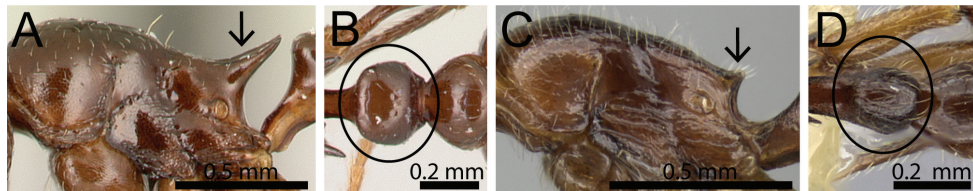


Figure 15. Mesosoma in profile (black arrows indicating propodeal spines/teeth) and petiole in dorsal view (within black ellipse). **A, B** *T. malagasy* (CASENT0449550) **C, D** *T. tyrior* (CASENT0249085).

- 16 Mandibles always unsculptured, smooth, shining (Fig. 16A, B); waist segments always without long, erect to suberect pilosity (Fig. 16E, F) ***T. schaufussii* group (in part)**
 – Mandibles variably sculptured (Fig. 16C, D); waist segments usually with long, erect to suberect pilosity (Fig. 16G); if pilosity absent (Fig. 16H), then mandibles conspicuously sculptured (Fig. 16D) ***T. naganum* group (in part)**

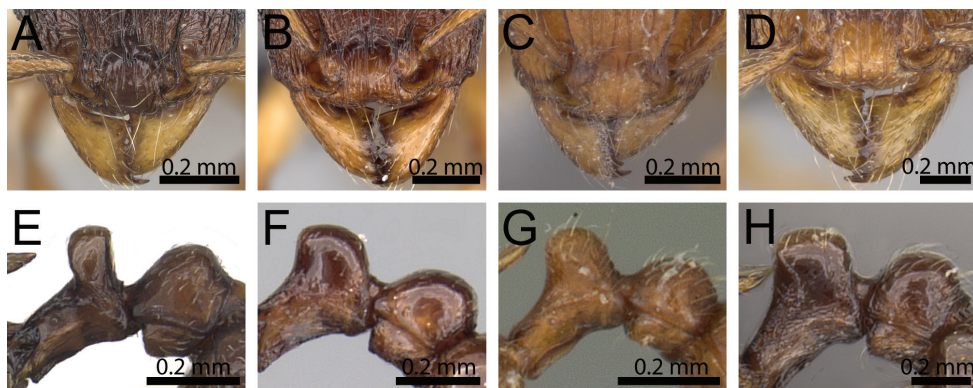


Figure 16. Anterior head in dorsal view and waist segments in profile. **A, E** *T. camelliae* (CASENT0247496) **B, F** *T. myrmidon* (CASENT0028635) **C, G** *T. naganum* (CASENT0102345) **D, H** *T. dalek* (CASENT0038402).

- 17 Antennal scrobes very well developed and distinctly impressed with sharply defined posterior and ventral margins; scrobes with very conspicuous median longitudinal scrobal carina, carina always ending between posterior eye margin and posterior margin of scrobe (Fig. 17A, C); petiolar node in dorsal view always noticeably broader than long (Fig. 17B), and in profile with more or less rounded antero- and posterodorsal margins (Fig. 17C); dorsal mesosoma with longitu-

- dinally rugose sculpture (Fig. 17B); all dorsal surfaces of body with abundant, usually dense, long, and suberect to erect hairs (Fig. 17C) ... *T. plesiarum* group
- Character combination never as above; usually antennal scrobes either almost absent (Fig. 17D, E) to weakly developed (Fig. 17F), or moderately developed and distinctly impressed, but without sharply defined posterior and ventral margins or a strong median longitudinal carina (Fig. 17G); if antennal scrobes well developed with sharp margin all around and median scrobal carina developed (Fig. 17H, I), then either dorsum of mesosoma reticulate-rugose (Fig. 17J) and/or petiolar node in profile rectangular nodiform with sharply angled antero- and posterodorsal margins (Fig. 17K)..... **18**

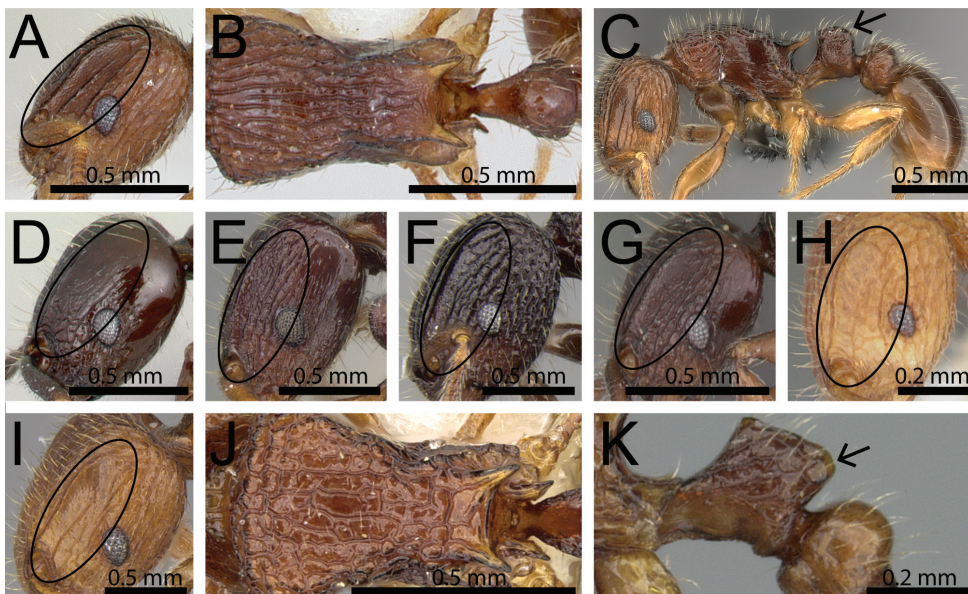


Figure 17. Head (without mandibles) in profile (antennal scrobe area within black ellipses); mesosoma in dorsal view; body and waist segments in profile. **A, B** *T. plesiarum* (CASENT0172831) **C** *T. mars* (CASENT0474279) **D** *T. silvicola* (CASENT0042828) **E** *T. nassonowii* (CASENT0195504) **F** *T. adamsi* (CASENT0247296) **G** *T. dysalum* (CASENT0102348) **H, J, K** *T. zenatum* (CASENT0102355; CASENT0344941) **I** *T. fhg-vazi* (CASENT0422522).

- 18 Sculpture on head, mesosoma, and waist segments strongly reduced: head usually very weakly sculptured (especially posteriorly) (Fig. 18A, B, C, D), dorsum of mesosoma generally completely unsculptured (Fig. 18K), very rarely with very few weak, superficial rugulae laterally (Fig. 18L), and waist segments always completely unsculptured..... **19**
- Sculpture never as strongly reduced as above, head always and to a great extent sculptured (Fig. 18E, F, G, H, I, J), dorsum of mesosoma always completely sculptured, (Fig. 18M, N), and waist segments variably sculptured..... **20**

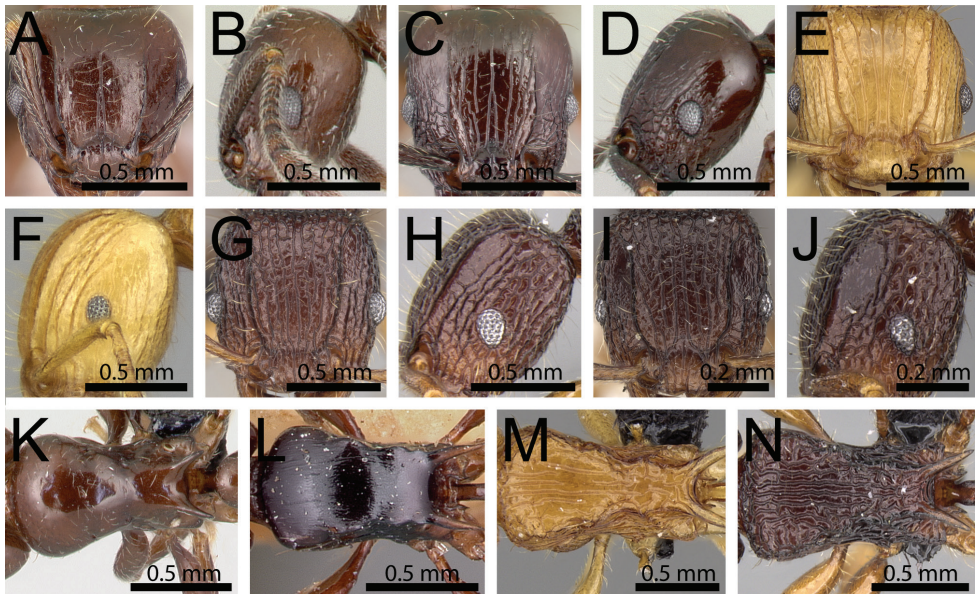


Figure 18. Head (without mandibles) in dorsal and lateral view; Mesosoma in dorsal view. **A, B** *T. ryanphelanae* (CASENT0454495) **C, D** *T. silvicola* (CASENT0042828) **E, F, M** *T. elf* (CASENT0045788) **G, H, N** *T. jedi* (CASENT0043578) **I, J** *T. quasirum* (CASENT0280585) **K** *T. ryanphelanae* (CASENT0454495) **L** *T. marginatum* (CASENT0101287).

- 19 Mesosoma only weakly marginate between lateral and dorsal mesosoma, instead sides of mesosoma generally rounding more or less smoothly onto the dorsum (Fig. 19A), mesosoma also relatively high and compact (LMI 43–48); mesosoma and first gastral tergite with relatively dense, short to moderately long, appressed to decumbent pubescence intermixed with relatively scarce, much longer erect pilosity (Fig. 19A)..... ***T. bessonii* group (in part)**
- Mesosoma usually with strong margination between lateral and dorsal mesosoma (Fig. 19B), but if mesosoma less marginate (Fig. 19C), then generally more elongate and slender (LMI 37–42); mesosoma and first gastral tergite with few to abundant long, standing hairs, often mixed with substantially fewer and (mostly) shorter appressed to subdecumbent pilosity, pubescence very scarce to absent (Fig. 19B, C)..... ***T. marginatum* group (in part)**



Figure 19. Mesosoma, waist segments, and first gastral tergite in profile. **A** *T. ryanphelanae* (CASENT0454495) **B** *T. valky* (CASENT0496394) **C** *T. silvicola* (CASENT0042828).

- 20 Relatively large species (HW 0.85–0.97; WL 1.21–1.48); SI relatively high (SI 89–104); propodeal spines very long to extremely long (PSLI 35–68); petiolar node in profile clublike, elongate and longer than high, posterodorsal angle situated higher than anterodorsal (Fig. 20A, B); dorsum of mesosoma conspicuously reticulate-rugose (Fig. 20H); whole body covered by numerous very long, fine, standing hairs (Fig. 20A, B)..... ***T. kelleri* group**
- Character combination never as above, most species much smaller with lower SI and shorter propodeal spines, a differently shaped petiolar node, and with less abundant and shorter pilosity (Fig. 20C, D, E); if species in size range of above and with similarly long propodeal spines and dense, long pilosity, then SI usually lower (SI 65–93), petiolar node more or less rectangular nodiform, and dorsum of mesosoma longitudinally rugose (Fig. 20F, G, I).....21

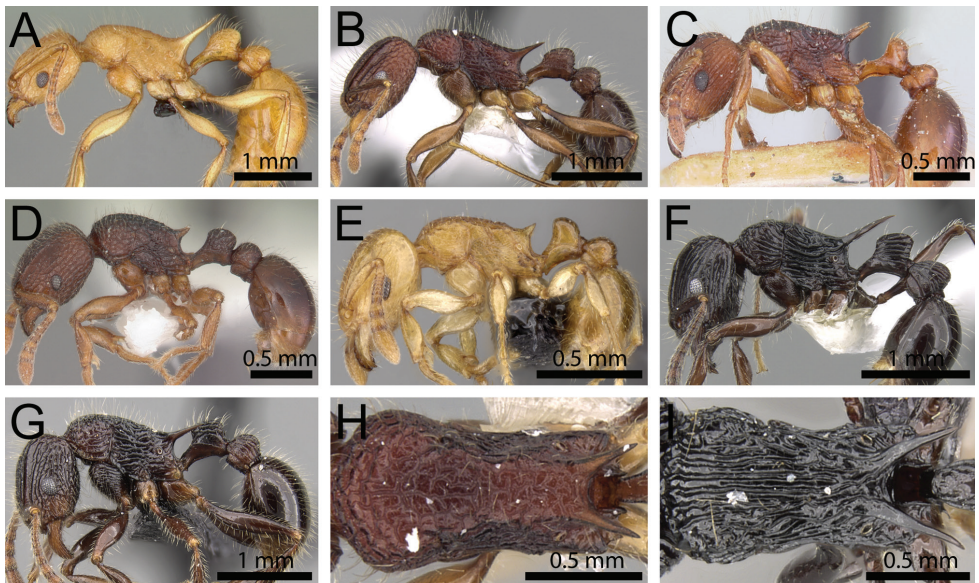


Figure 20. Mesosoma and waist segments in profile; mesosoma in dorsal view. **A** *T. kelleri* (CASENT0467063) **B, H** *T. ankarana* (CASENT0247543) **C** *T. steinbeili* (CASENT0101258) **D** *T. ranarum* (CASENT0102392) **E** *T. rala* (CASENT0162115) **F** *T. ambanizana* (CASENT0189238) **G** *T. nazgul* (CASENT0028625) **I** *T. smaug* (CASENT0121244).

- 21 Mesosomal outline in profile relatively flat, comparatively low and elongated (LMI 35–39) (Fig. 21A, B, C); in profile petiolar node rounded nodiform to high rounded nodiform with well-rounded margins (Fig. 21A, B), rarely high cuneiform or squamiform (Fig. 21C); propodeum usually with short teeth/denticles and rarely with spines of moderate length (PSLI 7–25, usually below 20); mandibles and waist segments always unsculptured, smooth and shining (Fig. 21A, B, C) ***T. schaufussii* group (in part)**
- Character combination never as above; mesosoma usually more compact and higher (LMI usually conspicuously above 40, very rarely below) (Fig. 21D, E, F,

- G, H), if LMI < 40, then propodeal spines long to very long (PSLI 27–72, usually above 35) and waist segments weakly to conspicuously sculptured (Fig. 21I) ... **22**
- 22 Mesosoma strongly marginate from sides to dorsum (Fig. 21G); dorsum of promesonotum weakly (irregularly) longitudinally rugulose with few, larger unsculptured patches medially, propodeal dorsum either fully unsculptured or only partly irregularly rugulose (Fig. 22A, B); both waist segments always completely unsculptured, smooth and shiny; body colour uniformly yellow (Fig. 21G)..... ***T. marginatum* group (in part)**
- Character combination never as above, especially dorsum of mesosoma usually with conspicuous sculpture along its entire length (Fig. 22C, D, E, F) **23**

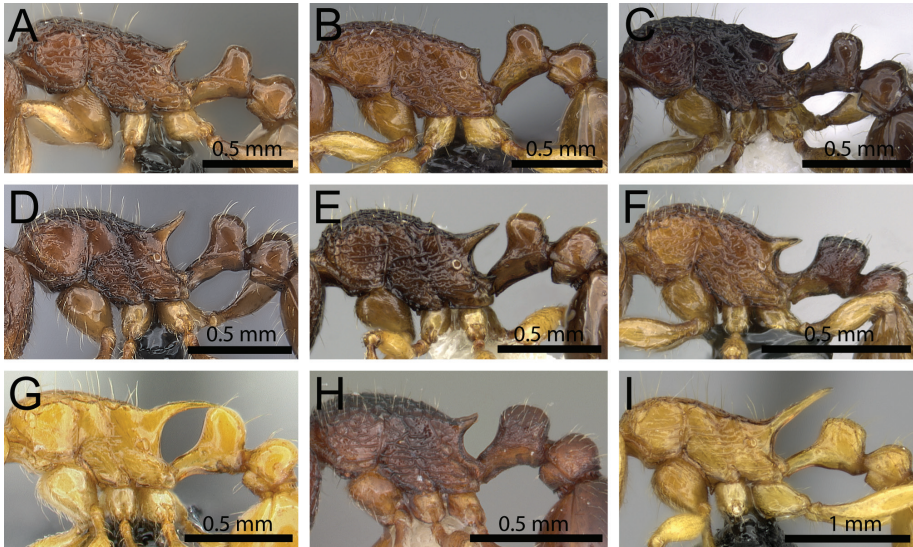


Figure 21. Mesosoma and waist segments in profile. **A** *T. pseudogladius* (CASENT0153605) **B** *T. merina* (CASENT0437226) **C** *T. scutum* (CASENT0189116) **D** *T. alperti* (CASENT0042547) **E** *T. vohitra* (CASENT0189167) **F** *T. fhg-forc* (CASENT0150949) **G** *T. shamshir* (CASENT0467696) **H** *T. quasirum* (CASENT0102353) **I** *T. elf* (CASENT0045788).

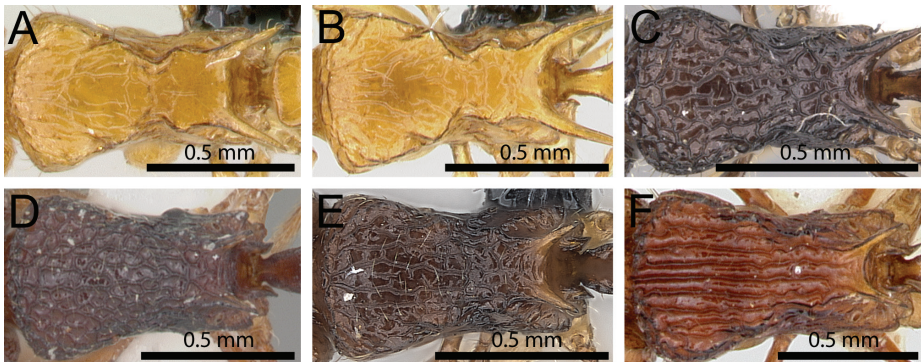


Figure 22. Mesosoma in dorsal view. **A** *T. norvigi* (CASENT0489037) **B** *T. shamshir* (CASENT0467696) **C** *T. nosybe* (CASENT0422207) **D** *T. ranarum* (CASENT0102392) **E** *T. alperti* (CASENT0042547) **F** *T. isectum* (CASENT0172829).

- 23 Petiolar node in profile dorsally conspicuously anteroposteriorly compressed and strongly narrowing towards apex, giving node a triangular or sharply cuneiform appearance; both waist segments always completely unsculptured, smooth and shiny (Fig. 23A, B, C, D) 24
- Petiolar node variably shaped: rectangular nodiform (Fig. 23E), high nodiform (Fig. 23F), squamiform (Fig. 23G), or broadly/weakly cuneiform (Fig. 23H), in the latter case node not strongly narrowing dorsally and petiole and/or postpetiole distinctly sculptured (Fig. 23H)..... 25

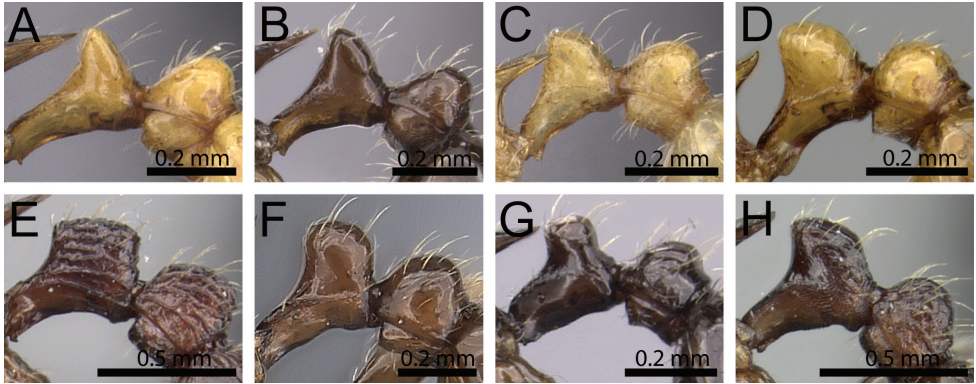


Figure 23. Petiole and postpetiole in profile. **A** *T. sada* (CASENT0443274) **B** *T. nosybe* (CASENT0422207) **C** *T. olana* (CASENT0044485) **D** *T. mackae* (CASENT0189093) **E** *T. aberni* (CASENT0045755) **F** *T. alperti* (CASENT0042547) **G** *T. ambatovy* (CASENT0124721) **H** *T. avaratra* (CASENT0445167).

- 24 Dorsum of mesosoma longitudinally rugose/rugulose (Fig. 24A, B)..... *T. dysalum* group (in part)
- Dorsum of mesosoma conspicuously reticulate-rugose, especially anteriorly (Fig. 24C, D, E, F) *T. bonibony* group (in part)

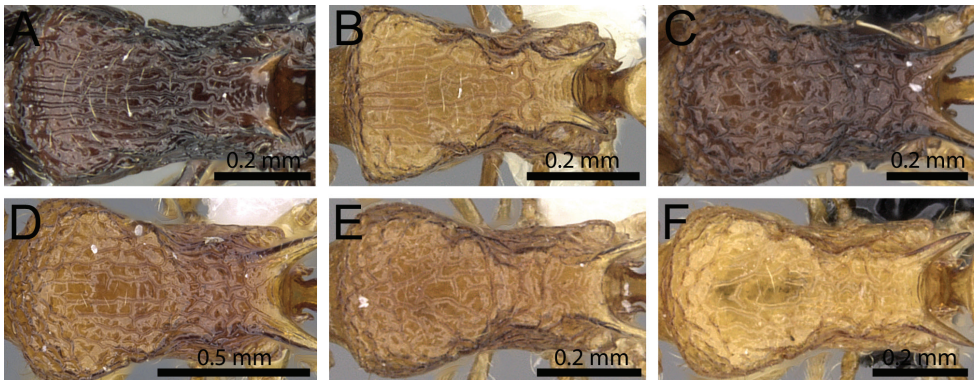


Figure 24. Mesosoma in dorsal view. **A** *T. orc* (CASENT0487093) **B** *T. mackae* (CASENT0189093) **C** *T. sada* (CASENT0443274) **D** *T. vony* (CASENT0404310) **E** *T. kali* (CASENT0235221) **F** *T. olana* (CASENT0044485).

- 25 Dorsum of mesosoma conspicuously reticulate-rugose throughout its length (Fig. 25A, B, C); petiolar node either squamiform, unsculptured, smooth, and shining or nodiform and usually conspicuously sculptured (Fig. 26A, B, C) 26
- Dorsum of mesosoma longitudinally rugose/rugulose (Fig. 25D, E), sometimes irregularly arranged but still conspicuously longitudinal in nature (Fig. 25F), if the latter, then petiolar node never as above (Fig. 27A, B, C, D, E, F, G, H)..... 27

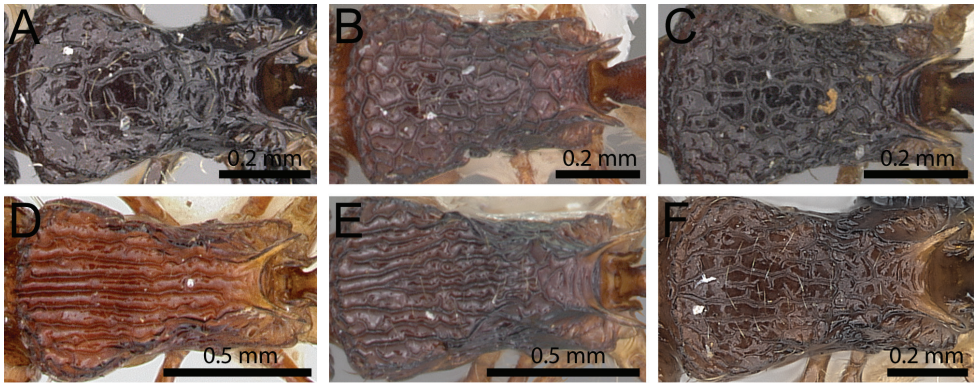


Figure 25. Mesosoma in dorsal view. **A** *T. ambatovy* (CASENT0124721) **B** *T. quasirum* (CASENT0102353) **C** *T. coillum* (CASENT0235219) **D** *T. isectum* (CASENT0172829) **E** *T. steinbeili* (CASENT0102394) **F** *T. alperti* (CASENT0042547).

- 26 Eyes relatively large (OI 25–26); petiolar node in profile distinctly squamiform and anteroposteriorly compressed (Fig. 26A); waist segments always completely unsculptured, smooth, and shining (Fig. 26A) ***T. dysalum* group (in part)**
- Eyes smaller than above, usually significantly so; petiolar node weakly cuneiform to rectangular nodiform and variably sculptured (Fig. 26B, C), but never squamiform and unsculptured as above (Fig. 26B, C) ***T. ranarum* group (in part)**



Figure 26. Body in profile (black arrows indicate eyes; waist segments within black ellipses). **A** *T. ambatovy* (CASENT0124721) **B** *T. fhg-mogw* (CASENT0056452) **C** *T. zenatum* (CASENT0344941).

- 27 Petiolar node usually rectangular nodiform with more or less sharply angled anterodorsal and posterodorsal margins (Fig. 27A, B, C), if petiolar node weakly cuneiform (Fig. 27D), then base of first gastral tergite sculptured, at least weakly so; both waist segments always distinctly sculptured, usually distinctly rugose, rarely rugulose (Fig. 27A, B, C, D).....
 ***T. tortuosum* group (in part)**
- Petiolar node usually squamiform (Fig. 27E, F), high rounded nodiform (Fig. 27G) or weakly cuneiform (Fig. 27H), if petiolar node weakly cuneiform, then first gastral tergite completely unsculptured, smooth, and shining; petiole and postpetiole fully unsculptured, smooth, and shiny in most species (Fig. 27E, F, G), a few species with one or both waist segments conspicuously sculptured (Fig. 27H)28

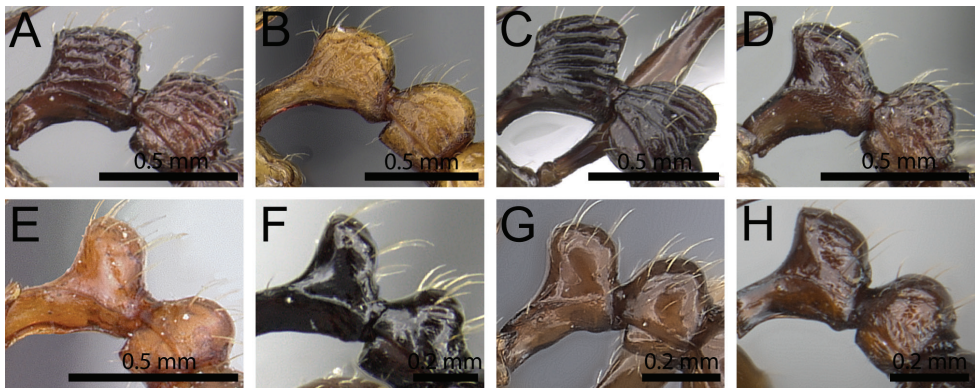


Figure 27. Petiole and postpetiole in profile. **A** *T. aberni* (CASENT0045755) **B** *T. voasary* (CASENT0247162) **C** *T. ambanizana* (CASENT0189238) **D** *T. avaratra* (CASENT0445167) **E** *T. steinheili* (CASENT0101258) **F** *T. yammer* (CASENT0042832) **G** *T. alperti* (CASENT0042547) **H** *T. dysalum* (CASENT0037931).

- 28 Pilosity and pubescence on first gastral tergite usually consisting of abundant, long, erect to suberect hairs on top of scarce, much shorter, appressed to decumbent pubescence (Fig. 28A, B, C), very rarely with long decumbent to subdecumbent pilosity on top of scarce appressed pubescence (Fig. 28D).....
 ***T. dysalum* group (in part)**
- Pilosity and pubescence on first gastral tergite variable: either with few moderately long, appressed to decumbent pubescence in combination with several much longer, fine, and erect hairs (Fig. 28E), or with a mix of short to moderately long, abundant, decumbent to suberect pilosity, pilosity appearing disorganized due to varying degrees of inclination and hair length (Fig. 28F, G), or with short, abundant, subdecumbent to suberect pilosity, and without any appressed to decumbent pubescence or long, fine erect hairs (Fig. 28H).....
 ***T. naganum* group (in part)**

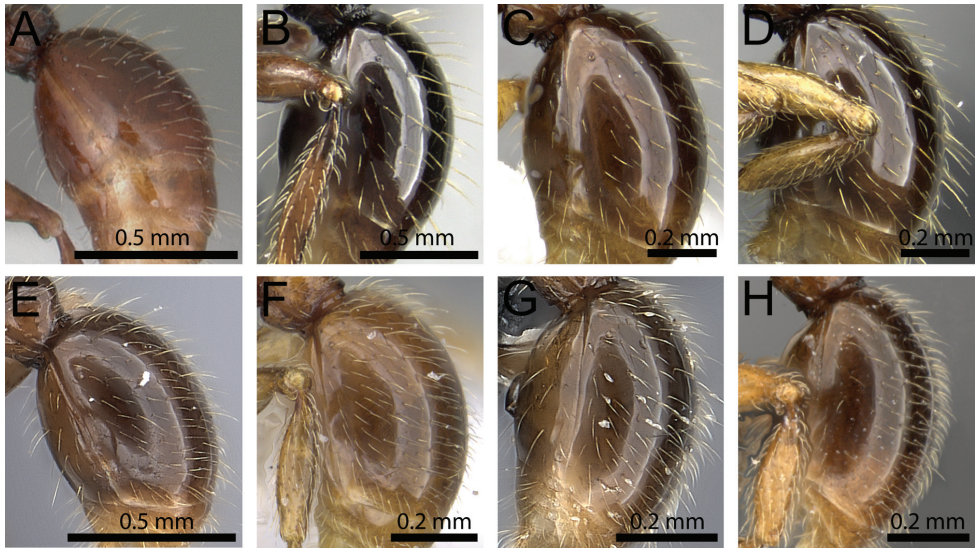


Figure 28. First gastral tergite in profile. **A** *T. dysalum* (CASENT0102349) **B** *T. yammer* (CASENT0042832) **C** *T. vohitra* (CASENT0189167) **D** *T. sargina* (CASENT0487390) **E** *T. alperti* (CASENT0042547) **F** *T. gilgamesh* (CASENT0247312) **G** *T. gilgamesh* (CASENT0163251) **H** *T. enkidu* (CASENT0045673).

Revision of the *Tetramorium setigerum* species group in Madagascar

Synopsis of the *T. setigerum* species group in Madagascar

Tetramorium cavernicola Hita Garcia & Fisher, sp. n.

Diagnosis of the *T. setigerum* group in Madagascar

Twelve-segmented antennae; antennal scapes very long (SI 120–123); anterior clypeal margin entire and clearly convex; frontal carinae well-developed, ending at or approaching posterior head margin; eyes moderate (OI 23–26); anterior face of mesosoma weakly developed, no distinct margination between lateral and dorsal mesosoma; propodeum armed with short triangular to elongate-triangular teeth (PSLI 7–11), propodeal lobes moderately developed, triangular to elongate-triangular, slightly longer and broader than propodeal teeth; petiolar node relatively small, nodiform, with weakly angled anterodorsal and posterodorsal margins, and comparatively long peduncle, petiolar dorsum flat to very weakly convex, node in profile between 1.2 to 1.4 times higher than long (LPeI 73–79), node in dorsal view between 1.2 to 1.3 times longer than wide (DPeI 121–127); postpetiole in profile approximately globular, around 1.0 to 1.1 times higher than long (LPpI 90–98); mandibles striate; clypeus longitudinally rugose/rugulose with well-developed median ruga and usually one or two weaker, sometimes irregular, lateral rugae/rugulae on each side; sculpture on cephalic dorsum irregularly longitudinally rugose to reticulate-rugose; mesosoma laterally irregularly

rugulose, dorsally reticulate-rugulose to irregularly rugulose; petiole and postpetiole conspicuously rugulose; ground sculpture on mesosoma and waist segments distinctly reticulate-punctate, much weaker on head; gaster unsculptured, smooth, and shiny; all dorsal surfaces of body with short to moderately long, thick, and apically blunt pilosity; sting appendage triangular to dentiform.

Taxonomic and biogeographic notes on the group

Prior to this study, the *T. setigerum* species group appeared endemic to the Afrotropical region where it is widely distributed. Of the 13 species recognised by Bolton (1980), most are found in more arid areas of eastern and southern Africa, a few are distributed in the rainforests of Central Africa, while two species are also found in Ethiopia and the southwestern Arabian Peninsula. The recent finding of *T. cavernicola* in Madagascar was unexpected since there was no previous indication of the presence of the group on Madagascar or any of the surrounding islands of the South West Indian Ocean. However, as outlined above, considering the strong biogeographical affinities of the *Tetramorium* ant fauna of Madagascar with the Afrotropical region, this is less of a surprise. Indeed, the *T. setigerum* group has its highest abundance and diversity in South and Southeast Africa, which is geographically comparatively close to Madagascar. As outlined above, other species or species groups that made it from Africa to Madagascar are often of predominantly eastern and southern African origin; examples include *T. humbloti* from the *T. weitzackeri* group and *T. delagoense* from the *T. simillimum* group.

The *T. setigerum* group cannot be mistaken for any other Malagasy species group. Its possession of twelve-segmented antennae, an entire and convex clypeal margin, and simple pilosity distinguish it from most other groups, except the *T. sericeiventre*, *T. simillimum*, and *T. tosii* groups. In the *T. sericeiventre* group the clypeus is distinctly modified, with the lateral portion being very prominent and raised into a tooth/denticle in full-face view while the clypeus of the *T. setigerum* group lacks such a tooth/denticle. Also, the species of the *T. simillimum* group possess much shorter antennal scapes (SI always much shorter than 100) than the *T. setigerum* group (SI over 120). The differentiation of the latter from the *T. tosii* group is more problematic. Despite the fact that the only representative of the *T. setigerum* group in Madagascar and the two species of the *T. tosii* group are easily separable (see key couplets 4 to 6), only a few morphological characters separate both groups if one also considers all members of the *T. setigerum* group from the Afrotropical region. Nevertheless, we prefer to keep both groups separate for the following reasons. First, the shape of the petiolar node is low, elongate, clublike, and always longer than high in the *T. tosii* group (Fig. 29A), whereas it is variably nodiform in the *T. setigerum* group, but usually higher than long and never low and elongate (Fig. 29B, C, D, E, F). Second, the standing pilosity in the *T. tosii* group consists of long, fine, acute hairs (Fig. 29A), whereas the pilosity in most members of the *T. setigerum* group is thick, short to moderately long, and usually blunt apically (Fig. 29B, E, F). Nevertheless, this is not the case in *T. metactum* Bolton and *T.*

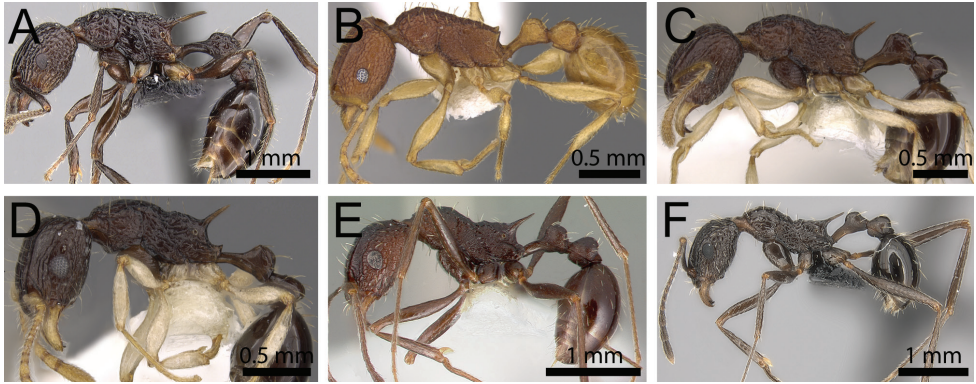


Figure 29. Body in profile. **A** *T. tosii* (CASENT0249662) **B** *T. cavernicola* (CASENT0247028) **C** *T. metactum* (CASENT0901193) **D** *T. youngi* (CASENT0901192) **E** *T. dolichosum* (CASTYPE13388) **F** *T. perlongum* (CASENT0135293).

youngi Bolton since they have long and fine pilosity (Fig. 29C, D). This may seem contradictory, but leads to our next argument. Third, we strongly suspect that the *T. setigerum* group is not a monophyletic group, but might be composed of different lineages that share a number of morphological characters that have evolved convergently. For example, the morphology of *T. cavernicola* from Madagascar is certainly closer to the species complex around *T. setigerum* Mayr and allies (Bolton 1980) than to *T. metactum* or *T. youngi*. In sum, the relationships between the *T. tosii* and the *T. setigerum* groups, as well as within the latter group, remain unclear, and we prefer to keep the groups as they are until additional data can provide better resolution of the groupings.

***Tetramorium cavernicola* Hita Garcia & Fisher, sp. n.**

<http://zoobank.org/FBE17724-6130-4C3B-8790-1C60C7278D62>

Figs 6A, E, 29B, 30

Type material. Holotype, pinned worker, MADAGASCAR, Antsiranana, Réserve Spéciale d'Ankarana, Andrafiabe, 12.92968 S, 49.05983 E, 59 m, in cave, ground nest, collection code BLF32473, 26.XI.2013 (*B. Fisher et al.*) (CASC: CASENT0247028). **Paratypes**, 15 pinned workers with same data as holotype (CASC: CASENT0247022; CASENT0247023; CASENT0247024; CASENT0247025; CASENT0247026; CASENT0247027; CASENT0247028; CASENT0247357; CASENT0247358; CASENT0248742; CASENT0248745; CASENT0248746; CASENT0373132; HLMD: CASENT0247029; MHNG: CASENT0248743; NHMB: CASENT0248744); and three pinned workers with same data as holotype except collection code BLF32472 and collected as ground foragers (BMNH: CASENT0247021; CASC: CASENT0247020; MCZ: CASENT0373133).

Non-type material. MADAGASCAR: Antsiranana, Réserve Spéciale d'Ankarana, Andrafiabe, 12.92968 S, 49.05983 E, 59 m, 26.XI.2013 (*B. Fisher et al.*).

Diagnosis. *Tetramorium cavernicola* differs from all other Malagasy congeners by the following combination of characters: 12-segmented antennae; anterior clypeal margin entire and convex; lateral clypeus not modified into tooth or denticle; antennal scape very long (SI 120–123); mesosoma in profile relatively low and slender (LMI 35–36); and propodeum armed with very short teeth/spines (PSLI 7–11).

Worker measurements (N=15). HL 0.74–0.78 (0.76); HW 0.58–0.61 (0.60); SL 0.71–0.75 (0.72); EL 0.14–0.15 (0.14); PH 0.32–0.35 (0.33); PW 0.45–0.48 (0.46); WL 0.92–0.99 (0.95); PSL 0.06–0.08 (0.07); PTL 0.17–0.19 (0.18); PTH 0.22–0.24 (0.23); PTW 0.21–0.23 (0.22); PPL 0.22–0.24 (0.23); PPH 0.24–0.25 (0.25); PPW 0.26–0.28 (0.27); CI 77–79 (78); SI 120–123 (122); OI 23–26; DMI 47–50 (49); LMI 35–36 (35); PSLI 7–11 (9); PeNI 47–49 (47); LPeI 73–79 (77); DPeI 121–127 (123); PpNI 56–60 (58); LPpI 90–98 (94); DPpI 113–123 (117); PPI 120–127 (124).

Worker description. Head much longer than wide (CI 77–79); posterior head margin weakly to moderately concave. Anterior clypeal margin entire and convex. Frontal carinae strongly developed, moderately raised, usually becoming weaker after posterior eye level, approaching or ending at posterior head margin; antennal scrobes very weak to absent. Antennal scapes very long, weakly surpassing posterior head margin (SI 120–123). Eyes moderately large (OI 23–26). Mesosomal outline in profile relatively flat, elongate and low (LMI 35–36), weakly marginate from lateral to dorsal mesosoma; promesonotal suture and metanotal groove absent. Propodeum armed with short, triangular teeth (PSLI 7–11), propodeal lobes moderately developed, triangular to elongate-triangular, slightly longer and broader than propodeal teeth. Petiolar node nodiform with moderately rounded antero- and posterodorsal margins, in profile between 1.2 and 1.4 times higher than long (LPeI 73–79), anterior and posterior faces not parallel, node weakly narrowing towards dorsum, anterodorsal and posterodorsal margins situated at about same height and both weakly to moderately angled, petiolar dorsum flat to very weakly convex; node in dorsal view around 1.2 to 1.3 times wider than long (DPeI 121–127), in dorsal view pronotum around 2.0 to 2.1 times wider than petiolar node (PeNI 47–49). Postpetiole in profile approximately globular, around 1.0 to 1.1 times higher than long (LPpI 90–98); in dorsal view around 1.1 and 1.2 times wider than long (DPpI 113–123), pronotum around 1.7 to 1.8 times wider than postpetiole (PpNI 56–60). Postpetiole in profile appearing distinctly more voluminous than petiolar node, postpetiole in dorsal view around 1.2 to 1.3 times wider than petiolar node (PPI 120–127). Mandibles striate; clypeus longitudinally rugose/rugulose with well-developed median ruga and usually one or two weaker, sometimes irregular, lateral rugae/rugulae on each side; cephalic dorsum between frontal carinae anteriorly towards posterior clypeal margin with three or four distinct but irregularly shaped longitudinal rugae with numerous cross-meshes, halfway between eye level and posterior head margin fluent transition to well-developed rugoreticulum ranging to posterior head margin; scrobal area only weakly sculptured, remainder of lateral head clearly reticulate-rugose. Mesosoma laterally and dorsally conspicuously reticulate-ru-

gose; forecoxae unsculptured, smooth, and shining. Petiole and postpetiole irregularly rugulose, better developed on dorsum than sides. First gastral tergite unsculptured, smooth, and shiny. Ground sculpture on cephalic dorsum between frontal carinae weak, distinctly reticulate-punctate on lateral head, mesosoma, and waist segments, absent from gaster. All dorsal surfaces of body with short to moderately long, thick, and apically blunt pilosity; appressed pubescence on first gastral tergite strongly reduced to absent. Anterior edges of antennal scapes and dorsal (outer) surfaces of hind tibiae with decumbent to suberect hairs. Head and mesosoma reddish brown; waist segments lighter in colour, usually orange brown; mandibles, antennae, and legs yellowish brown.

Etymology. The name of the new species is a Latin noun and means “cave dweller” or “cave inhabitant”. It refers to the microhabitat where the type series was collected. The species epithet is a nominative noun in apposition.

Distribution and biology. Currently, *T. cavernicola* is only known from Ankarana (Fig. 30D), where it was collected from a cave. The collection locality and the fact that the species is not known from outside the cave imply that *T. cavernicola* might be a specialised, cave-adapted ant. The generally very slender body and very long antennae and legs also support cave specialisation. Nevertheless, we do not consider the new species an obligate cave inhabitant. Arthropods that have evolved a cave-obligate lifestyle usually have a distinct set of morphological adaptations: reduction or loss of eyes, pigments, and wings; thinning of the cuticle; elongate antennae and legs; and slender body (Christiansen 1962; Culver 1982; Barr 1985). Yet the eyes, pigment, and wings in *T. cavernicola* are clearly not reduced since its eyes are always well developed, as are the wings in the queen and male castes, and the body colouration is brownish. In addition, we cannot detect any thinning of the cuticle. The slender gestalt and long antennae and legs could argue for cave adaptation, but are actually very typical of most species in the *T. setigerum* group. The antennae and legs of *T. dolichosum* Bolton and *T. perlongum* Santschi (Fig. 29E, F) are much longer than in *T. cavernicola*, even though these species do not live in caves. *Tetramorium cavernicola* appears to nest in the ground since most of the type series was collected from a ground nest, but no additional natural history data exists for this species.

Discussion. *Tetramorium cavernicola* is a very distinctive element of the Malagasy *Tetramorium* fauna and cannot be mistaken for any other congener based on the diagnosis provided above. There are some morphological similarities to the two species of the *T. tosii* group, as outlined earlier, but the distinction between these is easily found by comparing the shape of the head, the length of the antennal scapes, and the propodeal spines. In *T. cavernicola* the head is noticeably thinner (CI 77–79) and the antennal scapes are much longer (SI 120–123) while the propodeal spines are reduced to short teeth (PSLI 7–11). By contrast, the species in the *T. tosii* group have a thicker head (CI 85–91), much shorter scapes (SI 79–104), and much longer propodeal spines (PSLI 30–49).

Variation. Since *T. cavernicola* is only known from the type locality, there is no observable intraspecific variation.

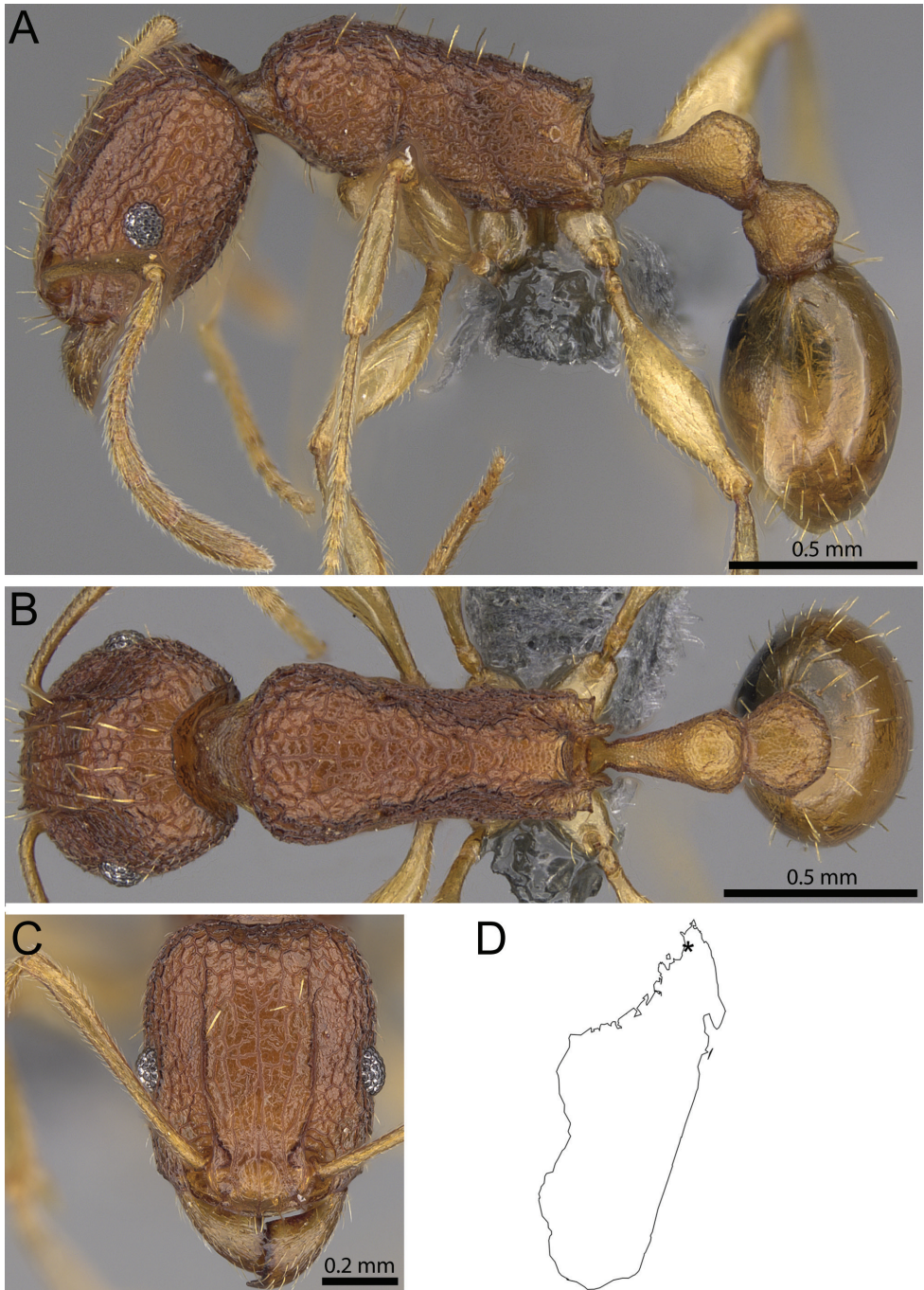


Figure 30. *Tetramorium cavernicola* sp. n. paratype (CASENT0373132). **A** Body in profile **B** Body in dorsal view **C** head in full-face view **D** map of Madagascar showing the type locality (black star symbol).

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