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Title	First fossil trichophyine rove beetle from mid-Cretaceous amber of northern Myanmar (Coleoptera : Staphylinidae : Trichophyinae)
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Citation	Cretaceous Researchc, 127, 104951 https://doi.org/10.1016/j.cretres.2021.104951
Issue Date	2021-11
Doc URL	http://hdl.handle.net/2115/90159
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Rights(URL)	https://creativecommons.org/licenses/by-nc-nd/4.0/
Туре	article (author version)
File Information	Cretac. Res127_104951.pdf



1	First fossil trichophyine rove beetle from mid-Cretaceous amber of
2	northern Myanmar (Coleoptera: Staphylinidae: Trichophyinae)
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13	
14	Abstract.
15	Trichophyinae is one of the smallest subfamilies of the megadiverse rove-beetle family
16	Staphylinidae, with only 18 species in the single extant genus Trichophya Mannerheim. Here
17	we describe the first fossil representative of Trichophyinae in mid-Cretaceous amber from
18	Kachin, northern Myanmar. †Trichophya minor sp. nov. is unusual in having non-filiform
19	thick antennae, similar to T. antennalis Cameron, 1932 from India, but this new species can
20	readily be distinguished from the latter by having much smaller and smoother body. Our
21	finding indicates that the subfamily had been originated by the Albian-Cenomanian
22	boundary, also showing a potential Gondwanan distribution at that time. We also briefly
23	discuss a miniaturization trend in the Burmese amber beetles.
24	
25	Key words: Staphylinidae; Trichophyinae; Trichophya; Kachin amber; Cenomanian; new
26	species
27	
28	1. Introduction
29	
30	With over 65,000 species in one extinct and 33 extant subfamilies (Bouchard et al., 2011;
31	Tihelka et al., 2020; Yamamoto, 2021), the rove beetles (Staphylinidae) are the largest animal
32	and metazoan family, showing amazing morphological and ecological diversity in terrestrial
33	ecosystems (Grebennikov and Newton, 2009; Thayer, 2016). Fossil records are vital

information for understanding the origin and evolutionary history of each taxonomic category.
In rove beetles, fossils are known for most subfamilies, with only a few exceptions.

Trichophyinae and its presumably sister subfamily Habrocerinae are among the few 36 subfamilies without any fossil record until now (Ashe and Newton, 1993; Ashe, 2005; Cai et 37 al., 2017: supplemental information; Yamamoto and Maruyama, 2018; Orlov et al., 2020). 38 The trichophyines are one of the smallest subfamilies in the mega-diverse Staphylinidae. In 39 fact, it contains only a single genus Trichophya Mannerheim with 18 extant species native to 40 the Northern Hemisphere. Although the biology and ecology are largely unknown in 4142Trichophyinae, they are found in various forest environments, e.g., in forest leaf litter, under bark and in squirrel middens (Legner and Moore, 1977; Newton et al., 2000; Shibata, 2001), 43or on fungi (Newton, 1984; Miyashita, 1997). They are typically small beetles, usually 2-3 44millimetres in body length. Among the rove beetles, one of the morphological characters 4546which define adult Trichophyinae is the presence of extremely slender, filiform, and verticillate antennae (Newton et al., 2000). This feature is known only in or within a very 4748limited number of other staphylinid groups, namely: Dasycerinae, Habrocerinae, and Scaphidiinae (tribe Scaphisomatini only). Further notable adult characters include the 4950presence of a strong neck constriction, five-segmented maxillary palpus including a minute and aciculate pseudosegment mounted on the spindle-shaped fourth maxillary palpomere, 51elytral epipleural keel or ridge absent, protrochantin exposed, mesothoracic spiracle in large 5253well-sclerotized triangular peritreme, and two pairs of abdominal paratergites present on most segments (Ashe and Newton, 1993; Newton et al., 2000; Ashe, 2005). 54

The mid-Cretaceous amber Kachin amber, previously known as Burmese amber, from 55northern Myanmar is well known for exceptionally diverse and abundant inclusions of 56organisms. In fact, it currently encompasses the richest biota in Mesozoic amber by far, with 57nearly 1,900 named taxa (Ross, 2019, 2020, 2021), despite the fact that most studies have 58only been made in the last decade. In Staphylinidae, a total of 19 subfamilies have currently 59been recorded including recently reported subfamilies such as Piestinae (Yamamoto et al., 60 612019), Paederinae (Żyła et al., 2019), Pseudopsinae (Liu et al., 2020c), and Protopselaphinae (Liu et al., 2020a, d). Here we report the first fossil occurrence of Trichophyinae from Kachin 6263 amber. Our finding sheds new light on the Mesozoic origin and morphological evolution of the subfamily, and this discovery eliminates a huge gap in the fossil record of rove beetles. 64 65

66

67 2. Material and methods

Amber material here studied was obtained from mines at the summit of Noije Bum Hill 68 (26°20'N 96°36'E) in the Hukawng Valley, Kachin State, northern Myanmar (mapped in 69 Cruickshank and Ko 2003). The amber-bearing horizon has been dated with 98.8 ± 0.6 Ma 7071(earliest Cenomanian) as the minimum age based on the Uranium-lead dating of zircon crystals in the volcanic sedimentary matrix (Shi et al., 2012). A slightly older age for Kachin 7273amber has been implied by Mao et al. (2018) and Balashov (2021). Based on these studies and 74the discovery of an ammonite fossil in Kachin amber (Yu et al., 2019), a mid-Cretaceous age is adopted here. The fossil resin was most likely formed by Metasequoia trees (Cupressaceae) 7576 in a tropical forest located near a seashore (Grimaldi and Ross 2017; Mao et al., 2018; Yu et 77al., 2019).

78 The amber specimen was cut using a hand saw, ground with emery papers of different grain sizes, and finally polished with a polishing cloth. Observation was conducted with a 7980 Leica MZ16 stereomicroscope. Photographs (Figs 1-4, Supplementary Figs 1-3) were taken with three different methods. The habitus and body parts of the fossil and extant material 81 82 (Figs 1, 2F, 4, Supplementary Figs 1–2) were photographed using a Canon EOS 80D digital 83 camera, mounted on a Canon MP-E 65 mm macro lens (F2.8, 1-5X), and with an attached 84 Canon MT-24EX twin flash as a light source. In addition, the images of several body parts of the fossil and slide specimen of the extant material (Figs 2B, E, 3, Supplementary Fig. 3) were 85 acquired with an Olympus DP26 digital camera mounted on an Olympus BX50 86 stereomicroscope. Finally, a few images of the enlarged body parts of the fossil (Fig. 2A, C, 87 D) were obtained using a Dun Ink BK PLUS Lab System, mounted on a Canon EOS 6D 88 89 digital camera with a $10 \times$ lens. During the imaging sessions, the amber specimen was completely submerged in clove oil to enhance the quality of the outputs. The software Helicon 90 Focus 7.5.4 was used for focus stacking based on the acquired images. All images were 9192edited, arranged, and assembled with Adobe Photoshop Elements 15 software. The sole 93 specimen is deposited in the entomological collection of the Gantz Family Collections Center, Field Museum of Natural History (FMNH), Chicago, IL, USA, under the registered number 9495FMNHINS-4357787. All the examined extant specimens are deposited in FMNH, and listed in Appendix 1. The Appendix 2 includes the supplementary figures used in the texts cited as 96 97 Supplementary Figs 1–3. The supplementary data comprised of Appendices 1 and 2 have been deposited in the Zenodo repository (https://doi.org/10.5281/zenodo.4784783; accessed 98 99 on 25 May 2021). This published work and the nomenclatural acts it contains have been 100 registered in ZooBank, the proposed online registration system for the International Code of 101 Zoological Nomenclature (ICZN). The ZooBank LSIDs (Life Science Identifiers) can be

102	resolved and the associated information viewed through any standard web browser by
103	appending the LSID to the prefix 'http://zoobank.org/'. The LSIDs for this publication are:
104	urn:lsid:zoobank.org:pub:5C59F363-D3DD-450C-BDC6-14A56A492865;
105	urn:lsid:zoobank.org:act:C375811E-115A-4D84-9647-E504799A8DB2.
106	
107	
108	3. Systematic paleontology
109	
110	Order Coleoptera Linnaeus, 1758.
111	Superfamily Staphylinoidea Latreille, 1802.
112	Family Staphylinidae Latreille, 1802.
113	Subfamily Trichophyinae Thomson, 1858.
114	
115	Genus <i>Trichophya</i> Mannerheim, 1830: 73.
116	Type species: Aleochara pilicornis Gyllenhal, 1810.
117	
118	Remarks. The native distribution of Trichophya seems to be the Palearctic, Oriental and
119	Nearctic regions, but several more undescribed species are known from Mexico through
120	Nicaragua (Newton et al., 2000).
121	
122	† <i>Trichophya minor</i> sp. nov.
123	(Figs 1–3)
124	LSID: <u>urn:lsid:zoobank.org:act:C375811E-115A-4D84-9647-E504799A8DB2</u> .
125	
126	Type material. Holotype, FMNHINS-4357787, a complete, well-preserved female adult;
127	deposited in FMNH. The fossil specimen is embedded in an irregularly flattened, rectangular
128	piece of transparent yellowish amber ($10.5 \times 3.9 \times 1.9$ mm).
129	
130	Etymology. Latin adjective minor meaning small, referring to the minute body of this extinct
131	species.
132	
133	Locality and horizon. Burmese (Kachin) amber from the Hukawng Valley (26°20'N 96°36'E),
134	Kachin State, northern Myanmar; upper Albian to lower Cenomanian (mid-Cretaceous).
135	

136 *Differential diagnosis.* †*Trichophya minor* sp. nov. is morphologically closely similar to *T*.

137 *antennalis* Cameron, 1932 (Fig. 4B, Supplementary Fig. 1B), known from northern India

138 (Shimla Hills), in having relatively short and thick antennae, in contrast to the very long and

139 slender, verticillate antennae of all other extant species of the genus, but it can be

140 distinguished from *T. antennalis* by the much smaller body (ca. 1.3 mm versus 2.2 mm),

141 shorter elytra, more strongly tapered abdomen, and smoother body surface (cf. Figs 1A and

142 4B).

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Description. Female. Body small (1.28 mm, measured from dorsal view), narrowly elongate,
tachyporine-like. Color uniformly dark reddish brown; mouthparts, legs, and gonocoxites
slightly paler. Surface uniformly weakly punctate, densely pubescent with inconspicuous
microsetae.

148Head (Figs 1, 2A, B) moderately deflexed, large (0.19 mm long, 0.23 mm wide), slightly smaller than pronotum, widest across eyes, moderately produced anteriorly; clypeus 149150narrow, less than half width of maximum width of head; frontoclypeal suture (Fig. 2A, fcs) gently arcuate, lacking midcranial suture; neck constriction (see Supplementary Fig. 2B, C) 151152slightly visible, with basal carina (Fig. 2A, ch). Labrum entire, transverse. Eyes (Figs 1, 2A, 153B) large, bulging, strongly projecting laterally (Fig. 2A). Antennal insertions (Fig. 2A, B, ai) partly exposed, distant from anterior margins of eyes (Fig. 2A). Antennae (Figs 1, 2A, B, 3A) 154with 11 antennomeres, slender, fili-moniliform, slightly longer than head and pronotum 155combined, each antennomere moderately setose with microsetae: antennomere 1 (a1) robust, 156157nearly subparallel sided, only weakly dilated apically, slightly curved, about 1.4 times longer than a2; a2 strongly dilated apically, thicker than a1; a3 elongate, rather slender, much 158narrower than a2, less wide than a1; a4–9 small, spherical, each antennomere similar in shape 159160 and size; a10 very slightly elongate, slightly longer than a9; a11 elongate, nearly twice as long 161 as wide, widest near middle, much larger than a10. Mandibles inconspicuous, details not visible. Maxillary palpi (Figs 2A, B, 3D) five-segmented including pseudosegment, moderate 162163 in size and length; palpomere 1 (*mp1*) small; mp2 narrowly elongate, slightly curved; mp3 robust, much shorter than mp2, strongly dilated apically; mp4 spindle-shaped, much longer 164165than mp3, densely covered with microsetae (Fig. 3D); mp5 (pseudosegment) minute, aciculate, hyaline (Fig. 3D). Mentum trapezoidal, moderately transverse, with truncate 166 anterior margin. Labial palpi (Fig. 2A) inconspicuous, not well visible. Gular sutures 167 168 complete, widely separated, rather strongly widened toward base, each forming straight line.

Pronotum (Fig. 1A) small (0.22 mm long, 0.33 mm wide), slightly larger than head, 169170strongly transverse (width/length = 1.5), widest slightly before base; sides arcuate, relatively strongly narrowed anteriorly; anterior margin nearly truncate; posterior margin gently 171172rounded; anterolateral angles obtusely rounded; posterolateral angles rounded; surface with dense and fine microsetae. Prosternum (Fig. 2B, E, ps) small, transverse; prosternal process 173174short, sharply pointed. Pronotal hypomeron narrow (Fig. 2B, E), relatively weakly folded inward; postcoxal process absent. Procoxal cavities opened behind. Mesoscutellum (Fig. 2C, 175sc) sub-triangular, longer than wide, with pointed apex. Elytra (Figs 1A, 2C, D) short (right 176177elytron 0.32 mm long, 0.20 mm wide), but moderately longer than pronotum; each elytron 178moderately elongate (width/length = 1.6), with sinuate posterior margin (Fig. 2D, *spe*), along 179with a row of posteriorly directed setae (Fig. 2D). Elytral epipleural keel or ridge absent. 180 Hindwings (Figs 1, 2F) fully developed, probably functional, with setae forming fringe along 181margin. Mesoventrite (Fig. 2E, msv) very short, strongly transverse; mesoventral process (Fig. 1822E) short, sharply pointed. Metaventrite (Fig. 2E, *mtv*) large, weakly transverse; metaventral 183process between mesocoxae (Fig. 2E) short, seemingly pointed.

Legs (Figs 1B, 2B, E) relatively short, slender, lacking spines on tibiae. Protrochantins 184185(Fig. 2B, E, *pt*) large, well exposed; procoxae (Fig. 2B, E, *pc*) conical, small, and contiguous; protrochanters small; profemora clavate, robust; protibiae (Fig. 1B) rod-like, very slender; 186protarsi 5-segmented, less than half length of protibiae, basal four protarsomeres small and 187short, together as long as protarsomere 5. Mesocoxae (Fig. 2E, msc) obliquely oval, short, at 188least partially contiguous with each other; mesofemora and mesotibiae similar to those of 189190 forelegs in shape, but slightly longer; mesotarsi (Fig. 3C) 5-segmented, about half length of 191 mesotibiae, basal four mesotarsomeres combined moderately longer than mesotarsomere 5, 192tarsomere 1 (Fig. 3C, *mst1*) much longer than mesotarsomere 2. Metacoxae (Fig. 2E, *mtc*) transverse, sub-triangular, and contiguous; metacoxal lamella not developed; metatrochanters 193 194 (Fig. 2E) fusiform, large; metafemora clavate, longer than mesofemora; metatibiae rod-like, very slender; metatarsi (Fig. 3E, mtt) 5-segmented, more than half length of metatibiae, 195196 metatarsomere 1 (Fig. 3E, *mtt1*) longest and about twice as long as tarsomere 2, and therefore, basal four metatarsomeres combined much longer than metatarsomere 5. Claws (Fig. 3C, E) 197198simple, lacking conspicuous basal teeth.

Abdomen (Figs 1, 2F) triangular, widest in segment 4, with six visible sterna; sides evenly strongly tapering from segment 4 to apex. Tergites (Figs. 1A, 2F) lacking both pruinose spots and basolateral ridges; tergite III (Fig. 2F, *t3*) partly visible; tergites IV–VI of almost same length, each markedly transverse; tergite VII (Fig. 2F, *t7*) longest; tergite VIII 203 (Fig. 2F, t8) largely exposed, elongate, narrowing apically, with rounded posterior margin.

204 Abdominal segments III–VII each with paratergites (one or two pairs, undetermined).

205 Sternites III–VI (Fig. 1B) subequal in length; sternite VII slightly longer; sternite VIII (Fig.

3B, s8) elongate, narrowing apically, with narrowly rounded apex. Gonocoxites (Figs 1, 2F,

- 3B, gc) present, large, cylindrical; surface covered with scattered macrosetae. Styli minute,
- 208 inconspicuous, much smaller than gonocoxites.

Male. Unknown.

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- 210
- 211

212 **4. Discussion**

213Among one extinct and 33 extant subfamilies (Bouchard et al., 2011; Tihelka et al., 2142020; Yamamoto, 2021), †*Trichophya minor* sp. nov. can unambiguously be placed in the 215subfamily Trichophyinae based on the following criteria (Newton et al., 2000): body compact and fusiform, anteriorly located antennal insertion which is partly visible dorsally, head with 216217well-developed neck constriction, maxillary palpus 5-segmented (fourth maxillary palpomere 218is large and spindle-shaped, whereas the fifth one is minute and hyaline), pronotum without 219postcoxal processes, protrochantin well exposed, tarsal formula 5-5-5, and abdomen with six 220visible sterna. One of the most distinctive morphological features is the presence of a specialized antenna, i.e., extremely slender, filiform, and verticillate antennae (Figs 4A, 221Supplementary Figs 1A, 3C; Newton et al., 2000). However, †T. minor sp. nov. lacks such a 222peculiar antenna as figured (Figs 1, 2A, B, 3A). Interestingly, a single extant Trichophya 223224species, namely T. antennalis from northwestern India, is known to have a morphologically similar antenna (Fig. 4B, Supplementary Fig. 1B; Ashe and Newton, 1993). In addition, this 225species shares several important characters with $\dagger T$. *minor* sp. nov. such as a smaller body, 226 227 shorter antennae, and somewhat strongly transverse pronotum. Despite their similarities in 228morphology, $\dagger T$. *minor* sp. nov. can be clearly differenciated from T. antennalis by having the much smaller body and smoother body surface. In this study, we refrain from establishing a 229230new genus which contains these two species because of the general morphological similarities with the other extant Trichophya members. The only notable difference between these two 231232species and the rest of others are the structure of the antennae, but it is considered here merely 233a primitive and ancestral character state, rather than a distinctive derived generic feature. 234Another remarkable feature of $\dagger T$. *minor* sp. nov. is its markedly small body size; it is about 2351.28 mm, whereas the remaining congeners range from 1.7 mm to 3.4 mm, with the exception 236of T. minuta Cameron, 1950 (1.2 mm) from Peninsular Malaysia. Several fossil occurrences

have been documented that imply a miniaturization trend for the Kachin amber beetles (e.g., 237Yamamoto and Takahashi, 2019; Żyła et al., 2019; Liu et al., 2020d; Li et al., 2020). 238239Although it is difficult to assume a particular reason with evidence for this phenomenon, if 240they are not just a taphonomic artefact, then there are two hypotheses for explanation: namely, random variation, or paleoenvironmental effect (Liu et al., 2020a). In the latter case, higher 241ambient temperature and/or lower atmospheric oxygen concentration could be one of the most 242important factors for contributing to such a miniaturization (Liu et al., 2020a). Alternatively, 243it could be possible to speculate that these fossils actually indicate a "maximization" trend in 244245comparison with the Recent fauna, since (if we accept the concept of linear time) the Burmese 246fossils existed long before the modern fauna.

The paleofauna of the Kachin amber is considered to have an affinity with Gondwana. 247Indeed, the West Burma Block where the amber producing forests in Kachin once located was 248249actually connected to a marginal area of the ancient Australian landmass and was gradually uplifted northwards during the Late Triassic to Late Jurassic (Heine and Müller, 2005; Poinar, 2502512019; Westerweel et al. 2019). At the time of the mid-Cretaceous, when the amber was 252formed, the Block was an isolated island in the Tethys Sea (Westerweel et al. 2019), resulting 253in the presence of a rich Gondwanan fauna that is currently endemic to the Southern Hemisphere, specifically Australasia or South America (e.g., Cai et al., 2019; Liu et al., 2542020b). Of note, the Indian endemic leiodid subgenus Pentacolonellus Peck, 1997 of the 255genus Colonellus Szymczakowski (Peck, 1997) has been found from Kachin amber (Cai and 256Huang, 2017). Thus, our finding of a Trichophya species with a possible affinity to India is 257significant when considering such a Gondwanan association. The current distribution of 258Trichophyinae, or *Trichophya*, is rather cosmopolitan, known from the Nearctic, northern 259Neotropical, Palaearctic, Oriental, and Australian regions (Ashe and Newton, 1993; Thayer, 2602612016). However, the native range of this group is narrower and confined to the northern 262hemisphere; the distribution in Australia should be considered as a result of human introduction (Thayer, 2016). Our discovery shows that Trichophya was indigenous to the 263264northern part of Myanmar during the mid-Cretaceous, providing implications for the native distribution of Trichophyinae. Finally, the extant members of Trichophya have usually been 265266found from mountaneous areas, whereas the Kachin amber species inhabited in a tropical forest very close to a seashore (Mao et al., 2018; Yu et al., 2019). This may suggest a different 267ecological strategy of Trichophya beetles at that time in comparison with the Recent ones. 268269

271 **5.** Conclusions

A new species, *†Trichophya minor* sp. nov., is herein described from mid-Cretaceous 272273Burmese (Kachin) amber based on a well-preserved adult female. It represents the first fossil 274record of the subfamily Trichophyinae and increases the total number of staphylinid subfamilies known from Kachin amber from 19 to 20. Our finding also indicates that the 275extant genus Trichophya had originated by the mid-Cretaceous. This new fossil species 276277provides insights into several trends known in Kachin amber, i.e., a long-term morphological stability, Gondwanan affinity, and miniaturization (see Liu et al., 2020a), together with 278279evolutionary insight into a certain specialized body part, namely antennae in our case. The discovery of $\dagger T$. minor sp. nov. may be important as a key fossil calibration point for future 280dating of molecular phylogenetic trees as there is no fossil record from the presumed sister 281subfamily Habrocerinae (Cai et al., 2017: supplementary table 2). 282283284285**Declaration of interests** The authors declare that they have no known competing financial interests or personal 286287relationships that could have appeared to influence the work reported in this paper. 288Acknowledgments 289290We would like to thank Rebekah S. Baquiran (FMNH) for proving the specimen number for the holotype. We also thank two anonymous reviewers and the Editor-in-Chief 291292Eduardo Koutsoukos. The first author (SY) was supported by a JSPS Overseas Research Fellowship (No: 29-212) and a Grant-in-Aid for JSPS Fellows (JSPS KAKENHI Grant 293Number 20J00159) given by the Japan Society for the Promotion of Science (JSPS), Tokyo, 294295Japan. 296 297298References 299Ashe, J.S., 2005. Phylogeny of the tachyporine group subfamilies and 'basal' lineages of the 300

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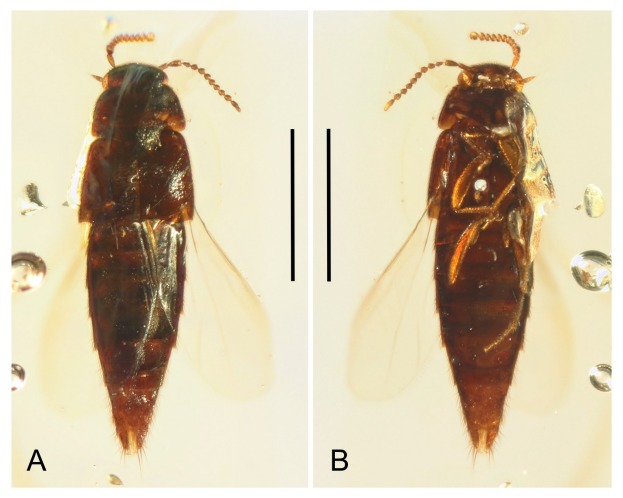
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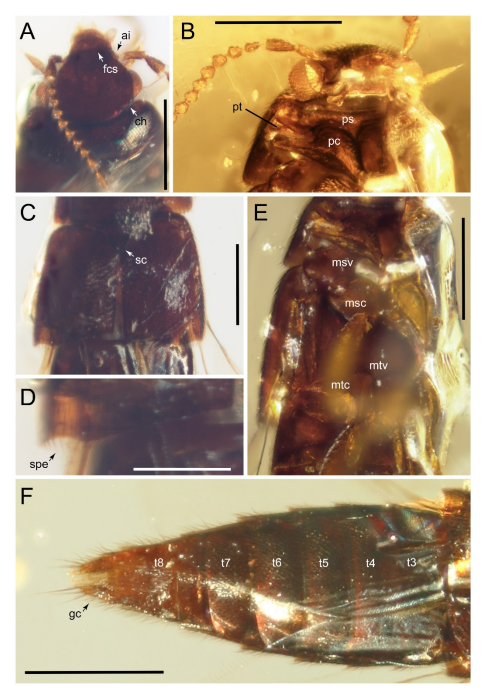
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- 444 Supplementary data to this article can be found online at https://doi.org/
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<u>Figures</u>



- **Fig. 1.** Habitus photographs of holotype (FMNHINS-4357787) of †*Trichophya minor* sp. nov.
- 453 in mid-Cretaceous Kachin amber from northern Myanmar. A. dorsal view; B. ventral view.
- 454 Scale bars: 0.5 mm.



457

458 **Fig. 2.** Morphological details of holotype (FMNHINS-4357787) of *†Trichophya minor* sp.

459 nov. in mid-Cretaceous Kachin amber from northern Myanmar. A. head and left antenna,

dorsal view; B. head, prothorax, and procoxae, ventral view; C. Elytra and mesoscutellum,

dorsal view; D. posterior margin of left elytron, dorsal view; E. Meso- and metathorax,

462 ventral view; F. Abdomen, dorsal view. Abbreviations: ai, antennal insertion; ch, basal carina

463 on head; fcs, frontoclypeal suture; gc, gonocoxite; msc, mesocoxa; msv, mesoventrite; mtc,

464 metacoxa; mtv, metaventrite; pc, procoxa; ps, prosternum; pt, protrochantin; sc,

465 mesoscutellum; spe, sinuate posterior margin of elytron; t3–8, tergites 3–8. Scale bars: 0.2

466 mm (A–C, E), 0.1 mm (D), 0.3 mm (F).

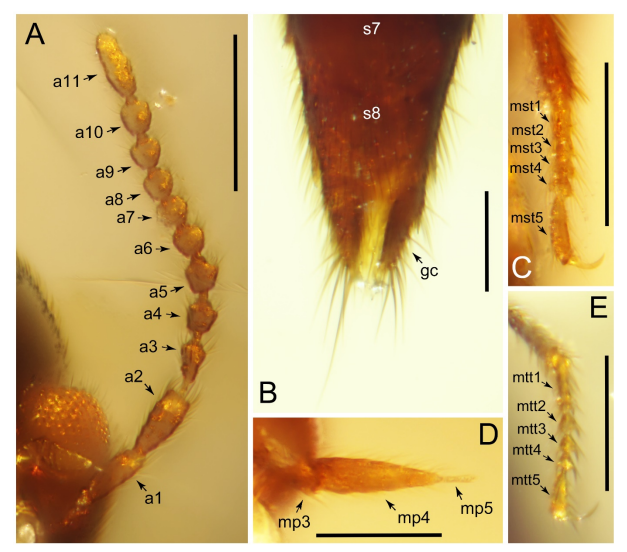
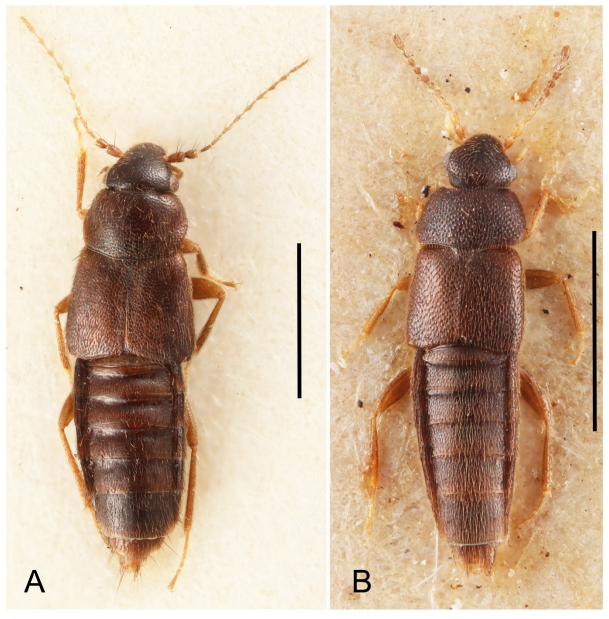


Fig. 3. Morphological details of holotype (FMNHINS-4357787) of †*Trichophya minor* sp.
nov. in mid-Cretaceous Kachin amber from northern Myanmar. A. right antenna, ventral
view; B. abdominal terminalia, ventral view; C. left mesotarsus, ventral view; D. left
maxillary palpus; E. left metatarsus, ventral view. Abbreviations: a1–11, antennomeres 1–11;
gc, gonocoxite; mp3–5, maxillary palpomeres 3–5; mst1–5, mesotarsomeres 1–5; mtt1–5,
metatarsomeres 1–5; s7–8, sternites 7–8. Scale bars: 0.1 mm (A–C, E), 0.05 mm (D).



477 Fig. 4. Extant species of Trichophyinae. A. *Trichophya pilicornis* (Gyllenhal) from Austria;

478	B. T. antennalis	Cameron fro	m northwestern	India.	Scale bars:	1.0 mm.

488	Highlights
489	• The first fossil Trichophyinae is described from mid-Cretaceous Burmese amber and placed
490	in the extant genus Trichophya.
491	
492	• <i>†Trichophya minor</i> sp. nov. is closely related to the extant species <i>T. antennalis</i> from India
493	based on antennae and body size.
494	
495	• Our finding indicates that Trichophyinae had originated by the mid-Cretaceous.
496	
497	• Our discovery is also congruent with the hypothesis of the Gondwanan origin of Burmese
498	(Kachin) amber.
499	
500	