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Research article

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Phylogenetic analysis and systematic position of two new species of the ant genus *Crematogaster* (Hymenoptera, Formicidae) from Southeast Asia

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Abstract. Two distinct new species of the ant genus *Crematogaster*, *C. khmerensis* sp. nov. and *C. pfeifferi* sp. nov., are described from Cambodia and Malaysia, respectively. The two species are unique among Asian *Crematogaster* in that they have vertically directed propodeal spines, but their systematic positions have not been determined based on morphological characters alone. Molecular phylogenetic analysis of 89 *Crematogaster* taxon matrices previously published plus *C. khmerensis* sp. nov., using nuclear genes, reveals that *C. khmerensis* sp. nov. is nested within the Australo-Asian *Crematogaster* clade. Morphological assignment of the developed pronotal shoulders implies a close relationship between *C. khmerensis* sp. nov. and the *C. tetracantha*-group. Based on molecular and morphological evidence, we erect a new species group, *C. khmerensis*-group, to contain *C. khmerensis* sp. nov. and *C. pfeifferi* sp. nov. Divergence time estimates using MCMCTree shows that the root node of the *C. khmerensis* sp. nov. terminal is estimated to be of middle Miocene age at 15 million years old. The position of the *C. khmerensis*-group well supports the Oriental- to Australian-region dispersal history that has been proposed for the Australo-Asian *Crematogaster* clade.

Keywords. Biogeography, divergence time estimation, morphological assignment, nuclear genes, phylogenetic position.

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Introduction

Ants (Formicidae) are one of the most familiar insect groups and they play important ecological roles in the ecosystems they inhabit (Hölldobler & Wilson 1990). They are abundant in terms of biomass, are distributed worldwide and form a diverse family, particularly in the tropics. Approximately 13 000 ant

species have been described to date (Bolton 2014), and it is estimated that this number will increase to approximately 20000 species in future (Hölldobler & Wilson 1990).

Ant taxonomy has been extensively investigated in recent years, with most taxonomic treatments typically presented as regional or systematic revisions. However, species with unique morphological characters are occasionally described separately from those comprehensive revisions. For example the *Carebara phragmotica* species clade and *Tetraponera phragmotica* have plug-shaped heads in the major worker caste (Ward 2006; Fischer *et al.* 2015); *Crematogaster masukoi* and *C. myops* have small compound eyes consisting of 5–6 ommatidia in the worker caste (Hosoishi *et al.* 2010). The possession of a phragmotic head suggests that the head is used to plug the nest entrance, and reduced compound eyes suggest a subterranean mode of life. Such derived states, as inferred by comparison with congenerics, are probably adaptations for particular life histories (e.g., predatory behavior, nesting habitat selection). Consequently, studies of distinct morphological characteristics are important contributions to ant taxonomy. While these distinct morphological forms (autapomorphies) can easily be distinguished from other more typical forms, and can therefore be described as new taxa, it can be difficult to determine their phylogenetic position. Thus, while uniqueness makes it easy to distinguish one taxon from others, elucidating the systematics of such taxa can be difficult.

Recent molecular analyses have revealed that some traditional taxa are not monophyletic e.g., (Cerapachyinae in Moreau *et al.* 2006, subgenera *Paracrema* and *Physocrema* in Blaimer 2012c, *Aphaenogaster* and *Tetramorium* by Ward *et al.* 2015). Indeed, since molecular data can resolve uncertainties in classification based on morphological data, an integrated approach that uses morphological data in conjunction with molecular phylogenetic analysis should be undertaken if fresh material is available.

The genus *Crematogaster* (Myrmicinae: Crematogastrini) is one of the hyperdiverse groups of ants, containing more than 400 species (Blaimer 2012b). The taxonomy has typically been examined as part of regional or systematic revisions (Longino 2003; Hosoishi & Ogata 2009; Blaimer & Fisher 2013), but some *Crematogaster* taxa with unique features have been recognized; for example, the *C. borneensis*-group has reduced 10-segmented antennae and has mutualistic relationships with the ant-plant genus *Macaranga* (Feldhaar *et al.* 2016), several species in the subgenus *Orthocrema* have reduced compound eyes (Hosoishi *et al.* 2010), *C. paradoxa* and related species have distinctly elongated propodeal spines (Creighton 1945), *C. reticulata* has a strongly reticulated body surface (Hosoishi 2009), and the queen of *C. cylindriceps* has a phragmotic head (Wheeler 1927; Yamane *et al.* 2011). The systematic positions of these taxa are usually proposed based on molecular or morphological data (Blaimer 2012b; Hosoishi & Ogata 2016), but the taxonomic affiliations are not always clear in some species.

In the course of a recent examination of *Crematogaster* specimens collected from Southeast Asia, we found two distinct species that appear to be new to science. None of the authors have ever seen any morphologically related species in a museum collection before. While the new species clearly belong to the subgenus *Crematogaster sensu stricto* (Blaimer 2012b), assignment to any of the species groups was not possible. The two species are considered to be distinct in that they both have vertically directed propodeal spines, which clearly distinguishes them from other Asian species of *Crematogaster*; however, their systematic positions are unknown.

The aims of this study were therefore to (1) clarify the phylogenetic position of these distinct *Crematogaster* ants using molecular phylogenetic analysis; (2) describe the two taxa as new species; (3) present the systematics of the two new species using morphological characters and molecular phylogenetic analysis. Morphological data in conjunction with molecular phylogenetic analysis will provide more comprehensive systematics of the two 'unique' species.

Material and methods

Sources of material

Type and non-type specimens were examined and/or deposited in the collections listed below. The species of *Crematogaster* from the Australian region are reviewed based on type and non-type specimen images available on AntWeb (https://www.antweb.org). Codes for public institutions generally follow those in Brandão (2000).

BMNH = Natural History Museum, London, UK

CASC = California Academy of Sciences, San Francisco, USA

ITBC = Institute for Tropical Biology and Conservation, Universiti Malaysia Sabah, Kota

Kinabalu, Malaysia

KUEC = Institute of Tropical Agriculture, Kyushu University, Fukuoka, Japan

MHNG = Musée d'Histoire naturelle, Geneva, Switzerland

SMNK = Staatliches Museum für Naturkunde Karlsruhe, Germany

THNHM = Thailand Natural History Museum, Technopolis, Khlong Luang, Pathum Thani,

Thailand

Observation

Most observations were made on a Leica M205C stereo microscope. Images were taken using a Canon EOS 50D with a Canon MP-E 65 mm $1-5 \times$ Macro lens, then processed using Combine ZM.

Measurements and indices

Measurements were made under a Leica M205C stereo microscope using ocular micrometers. All measurements are expressed in millimeters, recorded to the second decimal place. The measurements for petiole and postpetiole follow Longino (2003).

HW = Head Width; maximum width of head in full-face view.

HL = Head Length; perpendicular distance from vertex margin to line tangent anteriormost projections of clypeus in full-face view.

CI = Cephalic Index; $HW/HL \times 100$.

SL = Scape Length; length of the first antennal segment, excluding the neck and basal condyle.

SI = Scape Index; $SL/HW \times 100$.

EL = Eye Length; maximum length of the compound eye.

PW = Pronotal Width; maximum width of the pronotum in dorsal view.

WL = Weber's Length of the mesosoma; diagonal length, measured in lateral view from the anterior margin of the pronotum (excluding the collar) to the posterior extremity of the propodeal lobe.

PSL = Propodeal Spine Length; measured from tip of propodeal spine to closest point on outer rim of propodeal spiracle.

PtL = Petiole Length; length of the petiole in lateral view.

PtW = Petiole Width; maximum width of petiole in dorsal view.

PtH = Petiole Height; height of the petiole in lateral view.

PpL = Postpetiole Length; length of the postpetiole in lateral view (Longino 2003, fig. 2).

PpW = Postpetiole Width; maximum width of postpetiole in dorsal view, excluding the helcium.

PtHI = Petiole Height Index; $PtH/PtL \times 100$.

PtWI = Petiole Width Index; $PtW/PtL \times 100$.

PpWI = Postpetiole Width Index; PpW/PpL × 100.

WI = Waist Index; $PpW/PtW \times 100$.

Molecular data collection

Genomic DNA was extracted using a DNeasy Blood & Tissue kit (Qiagen, Maryland, USA). We sequenced fragments of five nuclear genes for one species, *Crematogaster khmerensis* sp. nov.: arginine kinase (ArgK, 402 bp), carbamoylphosphate synthase (CAD, 312 bp), long wavelength rhodopsin (LW Rh, 507 bp), DNA topoisomerase 1 (Top1, 795 bp) and wingless (Wg, 285 bp). The total number of base pairs for all genes was 2,301. Five new sequences were generated for this study, and the remainders were taken from Blaimer (2012c). Ninety taxa of *Crematogaster* (89 taxa of the subgenus *Crematogaster* analyzed in Blaimer (2012c) and *C. khmerensis* sp. nov.) were selected for molecular phylogenetic analysis. Primers, amplification and sequencing procedures followed Ward & Downie (2005) and Blaimer (2012a). The sequence data for *Crematogaster khmerensis* sp. nov. were deposited at DNA Data Base of Japan (DDBJ) with accession numbers: LC171383 (ArgK); LC171382 (CAD); LC171384 (LWRh); LC171381 (Top1); LC171380 (Wg).

Reactions were carried out at $10~\mu l$ volumes in a PCR Thermal Cycler MP (TaKaRa Bio Inc.) under the following conditions: first 40~cycles of $95^{\circ}C$ for 30~s, annealing at $50–58^{\circ}C$ for 30~s, and $72^{\circ}C$ for 90~s, then 1~cycle of $95^{\circ}C$ for 1~min, and finally $72^{\circ}C$ for 3~min. PCR products were visualized on a 1% agarose E-Gel 96-well system (Invitrogen), and then purified with 1.0~ul of ExoSAP-IT (GE Healthcare Life Sciences). All products were sequenced in both directions using BigDye Terminator v3.1 (Applied Biosystems) on an ABI 3100~Avant~DNA Sequencer (Applied Biosystems) at the Faculty of Science, Kyushu University, Fukuoka. Contigs were made using Vector NTI Advance TM ver. 11~(Invitrogen~Corp.) and subsequently aligned by eye. Sequence data were assembled and edited in the program Vector NTI Advance Tm ver. 11~(Invitrogen~Corp.) and MEGA 5~(Tamura~et~al.~2011).

Phylogenetic inference

The dataset was analyzed using maximum parsimony (MP), maximum likelihood (ML), and Bayesian inference (BI) to explore the strength of the phylogenetic signal under different optimality criteria.

The MP analysis was performed using TNT 1.1 program (Goloboff *et al.* 2008) with outgroup rooting, default consensus options, Tree Bisection and Reconstruction (TBR) branch swapping, and the default 'traditional search' mode. The tree search employed a parsimony ratchet with 10 000 interactions per run. Parsimony analyses were completed under conditions of equal weighting. Tree analyses were performed with WinClada version 1.00.08 software (Nixon 2002), and consensus cladograms generated from equally parsimonious trees were generated using the same program. Trees were rooted by the outgroup taxon *Crematogaster osakensis* belonging to the subgenus *Orthocrema*.

For ML and BI analyses, we first used PartitionFinder v.1.1.1 (Lanfear *et al.* 2012) to determine the best-fitting partition and substitution models for the phylogenetic analysis. For ML, under 'models = all', 'model selection = bic', and 'search = greedy', PartitionFinder identified five partitions (Table 1), which were employed in the ML analysis. Tree searches were conducted in GARLI v.2.0 (Zwickl 2006), with the following modification from default setting: genthreshfortopoterm = 100000, after Ward & Fisher (2016). Bootstrap values were calculated in PAUP*4.0b10 (Swofford 2002) using 100 bootstrap replicates from trees built in GARLI.

The BI analysis was performed using MrBayes v.3.1.2 (Ronquist & Huelsenbeck 2003). For the BI, model selection was limited to those that could be implemented in MrBayes, using the function 'model = mrbayes' in PartitionFinder. The best-fitting partition and substitution models chosen for BI analysis were shown in Table 1. We ran two independent runs with four chains each for 10 million generations, sampling every 1000 generations. All runs reached stationarity, as judged by the average standard deviation of split frequencies (ASDSF) approaching 0.01. The program Tracer v.1.6 (Rambaut *et al.* 2014) was used to evaluate convergence to the stationary distribution and effective sample size (ESS).

Table 1. Partitions and models identified by PartitionFinder and used in the maximum likelihood (ML) and Bayesian (BI) analyses.

Blocks	Partition	Model
ML		
ArgK_pos1, CAD_pos1, LWRh_pos2		K80 + I
ArgK_pos2, CAD_pos2, LWRh_pos1, Top1_pos1, Top1_pos2	p2	HKY + I + G
ArgK_pos3, LWRh_pos3, Wg_pos3	p3	K80 + G
CAD_pos3, Top1_pos3	p4	TrNef + G
Wg_pos1, Wg_pos2	p 5	K80 + I
BI		
ArgK_pos1, CAD_pos1, LWRh_pos2, Wg_pos1	p1	K80 + I + G
ArgK_pos2, CAD_pos2, LWRh_pos1, Top1_pos1, Top1_pos2, Wg_pos2	p2	HKY + I + G
ArgK_pos3, CAD_pos3, LWRh_pos3, Top1_pos3, Wg_pos3		K80 + G

The likelihood values converged to relative stationarity after ~1 million generations. ESS values for all parameters were well above 2000, providing evidence that convergence had been reached. Approximately 25% of samples were discarded as burn-in before parameters and trees were summarized. Posterior probabilities are given as statistical branch support.

Divergence time estimation

The concatenated five gene sequences from 24 myrmicine taxa (17 taxa of the Australo-Asian *Crematogaster* clade, *C. longispina*, *C. torosa*, *Aphaenogaster occidentalis*, *Leptothorax* cf. *muscorum*, *Stenamma dyscheres* and *Temnothorax rugatulus* analyzed in Blaimer (2012c), and *Crematogaster khmerensis* sp. nov.) and the monophyly-constrained topology which Blaimer (2012c) inferred were used for molecular dating analyses. Four fossil calibrations were incorporated following Blaimer (2012c). Divergence times were estimated using the Bayesian method implemented in MCMCTree of PAML4.7 (Yang 2007). Using the approximate likelihood calculation method, the gradient g and Hessian H with BASEML using the GTR + G substitution model were calculated (dos Reis & Yang 2011). The independent rate model for the molecular clock and the GTR + G model for nucleotide substitutions were set in the mcmctree.ctl control file, with the following modification from default setting: substitution rate per time unit = 0.106023; rgene gamma = 1 10; sigma2 gamma = 1 4.5.

Results

Class Hexapoda Blainville, 1816
Order Hymenoptera Linnaeus, 1758
Suborder Apocrita Latreille, 1810
Infraorder Aculeata Latreille, 1802
Superfamily Vespoidea Latreille, 1802
Family Formicidae Latreille, 1809
Subfamily Myrmicinae Lepeletier de Saint-Fargeau, 1835

Genus Crematogaster Lund, 1831

Phylogenetic position and divergence time estimation

The five nuclear gene sequences consisted of 2301 bp and contained 652 variable characters (VC) and 420 parsimony informative characters (PIC) (Table 2). The MP and ML analyses resulted in similar

Table 2. Data on number of bases, number of variable characters (VC) and number of parsimony informative characters (PIC).

Gene	No. bases	No. VC	No. PIC
ArgK	402	125	74
CAD	312	100	67
LWRh	507	140	93
Top1	795	215	142
Wg	285	72	44
Total	2301	652	420

topologies for the position of *C. khmerensis* sp. nov., but the BI analysis differed in topology from the MP and ML analyses. As our results from the BI analysis had similar results to the tree inferred by Blaimer (2012c), we use the BI tree to illustrate all our results (Fig. 1).

In MP analysis, the concatenated five-gene dataset yielded 1800 equally parsimonious trees (length = 1407, consistency index = 0.48, retention index = 0.77), of which the strict consensus tree is well resolved. The MP analysis recovered that C. khmerensis sp. nov. was nested within node A with high bootstrap support (100%) (Fig. 2). Further splitting of the node resulted in a trichotomy having branches with C. khmerensis sp. nov., C. borneensis-group and node C.

The ML analysis recovered that *C. khmerensis* sp. nov. was nested within node A with high bootstrap support (100%) (Fig. 2). Similarly the node displayed a trichotomy among *C. khmerensis* sp. nov., *C. borneensis*-group and node C.

The BI recovered that *C. khmerensis* sp. nov. was nested within node A (Fig. 2). Despite the relatively low level of support (0.88 posterior probability), *C. khmerensis* sp. nov. at node B was a sister to node C (Fig. 2).

Estimations of divergence time showed that the common ancestor of *C. khmerensis* sp. nov. and clade C began to diverge approximately 15 million years ago (Fig. 3).

Diagnosis of the Crematogaster khmerensis-group

the *Crematogaster khmerensis*-group is easily distinguished from other Asian species of *Crematogaster* by the vertically directed propodeal spines and large propodeal spiracles.

Species included in the *Crematogaster khmerensis*-group:

- C. khmerensis Hosoishi & Ogata sp. nov.
- C. pfeifferi Hosoishi & Ogata sp. nov.

Key to species based on the worker caste

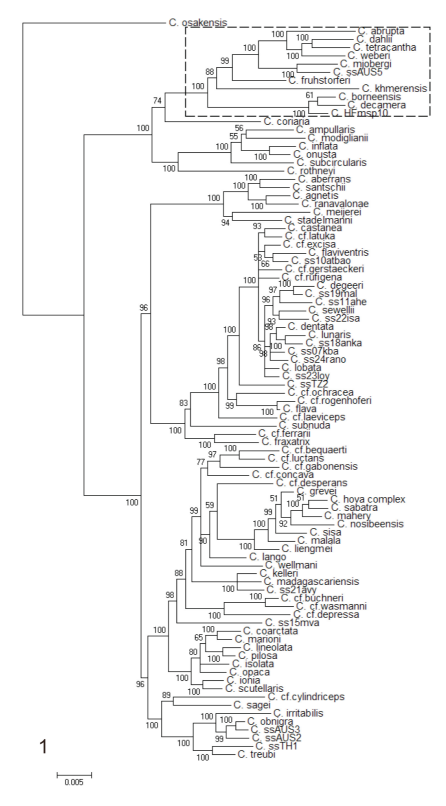


Fig. 1. Bayesian majority rule consensus tree reconstructed for 90 taxa using five genes (ArgK, CAD, LWRh, Top1, Wg) in a MrBayes analysis. Above node numbers indicate posterior probability. Data were partitioned by PartitionFinder v.1.1.1 and analyzed using a best fit model for each gene and codon position, with 10 million generations and a burn-in of 25 %. Area enclosed by dashed lines is enlarged on Fig. 2.

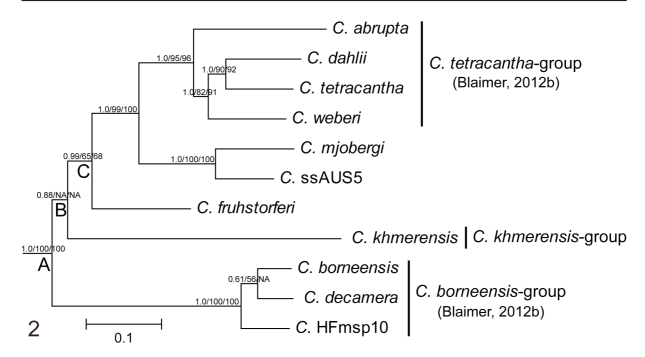


Fig. 2. Bayesian majority rule consensus tree reconstructed for 90 taxa using five genes (ArgK, CAD, LWRh, Top1, Wg) in a MrBayes analysis. Most of the outgroups are not shown. Above node numbers indicate posterior probability, bootstrap value for MP, and bootstrap value for ML. Data were partitioned by PartitionFinder v.1.1.1 and analyzed using a best fit model for each gene and codon position, with 10 million generations and a burn-in of 25 %.

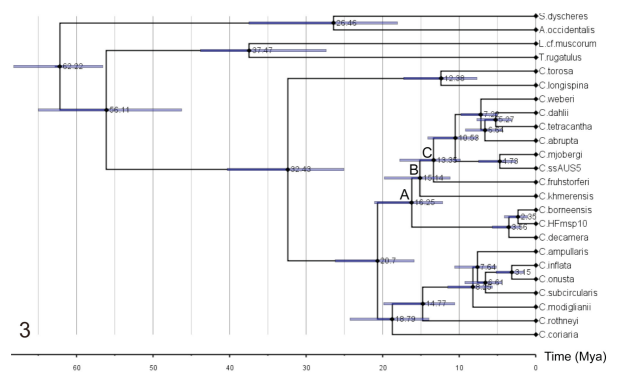


Fig. 3. Posterior estimates of divergence time of 24 taxa on the phylogenetic tree. Blue bars depict the 95% highest posterior density (HPD). Estimations were performed with MCMCTree using the independent rate model.

Crematogaster khmerensis sp. nov.

urn:lsid:zoobank.org:act:D2CC1731-9989-4708-9513-36E1458CE012

Figs 4A–D

Diagnosis

Crematogaster khmerensis sp. nov. belongs to the subgenus *Crematogaster sensu stricto* (Blaimer 2012b). This small-sized species of *Crematogaster* ants (HW 0.6–0.66; WL 0.63–0.69) is similar to *C. pfeifferi* sp. nov., but can be distinguished by the propodeal spiracles touching metapleural gland bulla, short propodeal spines (PSL 0.08–0.09), petiole broader anteriorly than posteriorly and undeveloped subpetiolar process.

Etymology

The specific name refers to the former kingdom name of Cambodia, the Khmer Empire.

Type specimens

Holotype

CAMBODIA: worker, Kampong Thom Province, 24 Nov. 2010, S. Hosoishi leg., (TUS arboreal SF3-1) (KUMANT040: deposited at THNHM).

Paratypes

CAMBODIA: 3 workers, same locality data as for holotype (TUS arboreal SF3-3, TUS arboreal SF3-6) (KUMANT041: deposited at BMNH; KUMANT042: deposited at CASC; KUMANT043: deposited at KUEC).

Measurements and indices

HW 0.60–0.66; HL 0.60–0.67; CI 97–102; SL 0.43–0.50; SI 69–76; EL 0.14–0.15; PW 0.35–0.40; WL 0.63–0.69; PSL 0.08–0.09; PtL 0.22–0.23; PtW 0.20–0.25; PtH 0.12–0.14; PpL 0.12–0.14; PpW 0.20–0.23; PtHI 52–64; PtWI 87–114; PpWI 154–167; WI 92–100 (holotype and three paratype workers measured).

Description

Worker

Workers presumably monomorphic.

Head subquadrate in full-face view, with rounded posterior corners and subparallel sides. Occipital carinae developed. Mandible with four teeth, apical tooth large. Anterior margin of clypeus convex; anterolateral margins of clypeus protruded anteriorly; posterior margin of clypeus rounded between frontal lobes. Frontal carinae almost parallel. Antennae 11-segmented; antennal club 3-segmented. Scape reaching posterior corner of head. Basal flagellar segment (antennal segment III) as broad as long. Compound eyes slightly projecting beyond lateral margins of head in full face view.

Pronotal shoulders angulate, but without distinct rugulae laterally. Mesonotal dorsum convex in lateral view. Pronotum and mesonotum not forming same dorsal outline in lateral view. Mesothoracic spiracle forms large pit concealed by posterior pronotal lobe. Metapleural gland opening slit-shaped. Propodeal spiracle large and circular, situated at posterolateral corners, touching metapleural gland bulla. Metanotal groove straight in dorsal view, deep and forming concave region between mesonotum and propodeum. In dorsal view, feeble longitudinal rugulae connecting between mesonotum and propodeum; the boundary distinct. Propodeal spines developed and directed vertically in lateral view.

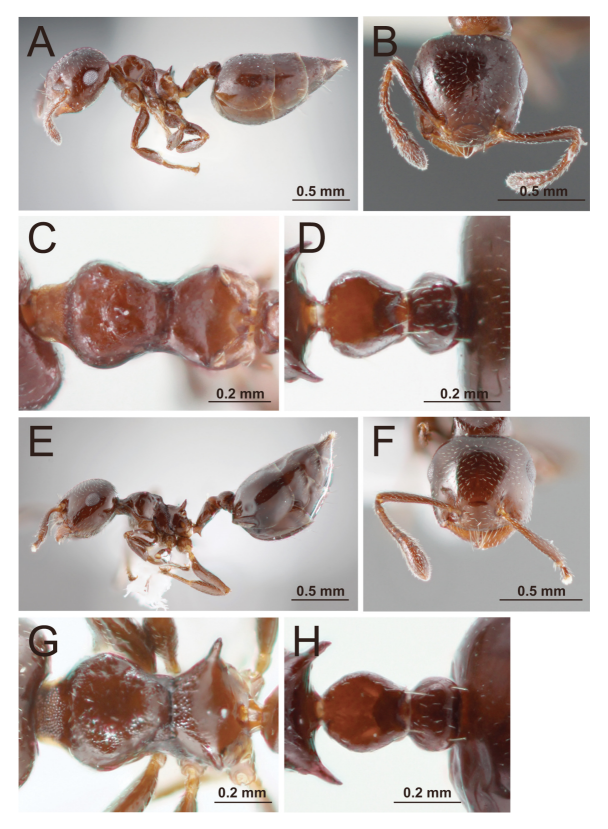


Fig. 4. A–D. *Crematogaster khmerensis* sp. nov., worker. A. Body in lateral view. B. Full-face view of head. C. Dorsal view of mesosoma. D. Petiole and postpetiole in dorsal view. — E–H. — *Crematogaster pfeifferi* sp. nov., worker. E. Body in lateral view. F. Full-face view of head. G. Dorsal view of mesosoma. H. Petiole and postpetiole in dorsal view.

Petiole scoop shaped, but flattened broader anteriorly, longer than broad; spiracle situated anteriorly midway between dorsal and ventral margin of petiole in lateral view, directed laterally. Subpetiolar process developed as a short lamellate denticle. Postpetiole without longitudinal median sulcus, but bilobed behind; spiracle situated anteriorly on lateral surface.

Dorsal surface of head mostly smooth or weakly shagreened. Clypeus mostly smooth and shining. Pronotal collar areolately sculptured. Promesonotum weakly punctate. Lateral surface of pronotum smooth and shining. Mesopleuron mostly smooth and shining, but sculptured on surrounding area. Anterodorsal surface of propodeum weakly punctate; posteior half smooth and shining. Lateral surface of propodeum smooth and shining. Dorsal and lateral surfaces of petiole smooth and shining. Dorsal and lateral surfaces of postpetiole smooth and shining.

Standing pilosity sparse. Dorsal face of head with appressed setae; one pair of erect setae on frontal lobes. Clypeus with one pair of erect setae. Anterior clypeal margin with two pairs of long setae mixed with short setae laterally. Scape with decumbent setae. Mesontum with appressed setae sparsely. Petiole with one pair of suberect setae posteriorly. Postpetiole with one pair of suberect setae posteriorly. Fourth abdominal tergite with short appressed setae sparsely.

Body color brown.

Queen and male

Queens and males of this species are unknown.

Distribution

This species is known only from the type locality in Cambodia (Fig. 5).

Crematogaster pfeifferi Hosoishi & Ogata sp. nov. urn:lsid:zoobank.org:act:BE4FEAC9-A8EE-4C65-B00D-B62263746F70 Figs 4E—H

Diagnosis

Crematogaster pfeifferi sp. nov. belongs to the subgenus *Crematogaster* sensu stricto (Blaimer 2012b). This species is small-sized *Crematogaster* ants (HW 0.65–0.68; WL 0.69–0.7). This species is similar to *C. khmerensis* sp. nov., but can be distinguished by the propodeal spiracles apart from metapleural gland bulla, long propodeal spines (PSL 0.11–0.12), elliptical petiole and acutely developed subpetiolar process.

Etymology

The specific name is dedicated to Dr. Martin Pfeiffer, who collected the type material.

Type specimens

Holotype

MALAYSIA: worker, Borneo, Golden Hope Table, 4°23 N, 117°51 E, xa178 611, 8 Oct. 2004, M. Pfeiffer leg., (KUMANT044: deposited at ITBC).

Paratypes

MALAYSIA: 3 workers, same data as for holotype, xa160 610 (KUMANT045: deposited at BMNH; KUMANT046: deposited at KUEC; KUMANT047: deposited at SMNK).

Measurements and indices

HW 0.65–0.68; HL 0.62–0.64; CI 105–106; SL 0.49–0.51; SI 75–76; EL 0.13–0.14; PW 0.37–0.39; WL 0.69–0.70; PSL 0.11–0.12; PtL 0.24–0.25; PtW 0.22–0.24; PtH 0.13–0.15; PpL 0.14–0.15; PpW 0.22–0.24; PtHI 52–63; PtWI 92–96; PpWI 157–164; WI 92–100 (holotype and three paratype workers measured).

Description

Worker

Workers presumably monomorphic.

Head subquadrate in full-face view, with rounded posterior corners and subparallel sides. Occipital carinae developed. Mandible with four teeth, apical tooth large. Anterior margin of clypeus convex; anterolateral margins of clypeus protruded anteriorly; posterior margin of clypeus rounded between frontal lobes. Frontal carinae almost parallel. Antennae 11-segmented; antennal club 3-segmented.

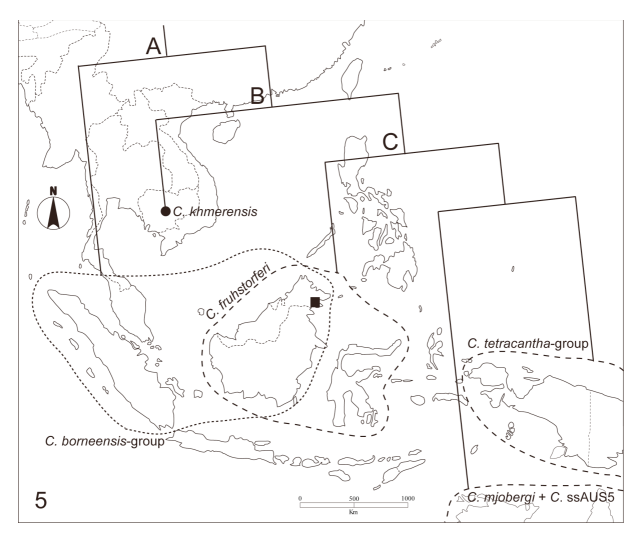


Fig. 5. Phylogeny and distribution of the *Crematogaster borneensis*-group, *C. khmerensisi* sp. nov., *C. fruhstorferi*, *C.* ssAUS5, *C. mjobergi* and the *C. tetracantha*-group. Closed circle indicates *C. khmerensis* sp. nov., closed square indicates *C. pfeifferi* sp. nov. It is noted that *C. pfeifferi* sp. nov. is not represented in the phylogeny.

Scape reaching posterior corner of head. Basal flagellar segment (antennal segment III) slightly longer than broad. Compound eyes slightly projecting beyond lateral margins of head in full-face view.

Pronotal shoulders angulate, but without distinct rugulae laterally. Mesonotal dorsum convex in lateral view. Pronotum and mesonotum not forming same dorsal outline in lateral view. Mesothoracic spiracle forms large pit concealed by posterior pronotal lobe. Metapleural gland opening slit-shaped. Propodeal spiracle large and circular, situated at posterolateral corners, apart from metapleural gland bulla. Metanotal groove straight in dorsal view, deep and forming concave region between mesonotum and propodeum. In dorsal view, feeble longitudinal rugulae connecting between mesonotum and propodeum; the boundary distinct. Propodeal spines developed and directed vertically in lateral view.

Petiole elliptical with convex sides, longer than broad; spiracle situated anteriorly midway between dorsal and ventral margin of petiole in lateral view, directed laterally. Subpetiolar process developed acutely. Postpetiole without longitudinal median sulcus, but bilobed behind; spiracle situated anteriorly on lateral surface.

Dorsal surface of head mostly smooth or weakly shagreened. Clypeus mostly smooth and shining. Pronotal collar areolately sculptured. Promesonotum weakly punctate. Lateral surface of pronotum smooth and shining. Mesopleuron mostly smooth and shining, but sculptured on surrounding area. Anterodorsal surface of propodeum weakly punctate; posterior half smooth and shining. Lateral surface of propodeum smooth and shining. Dorsal and lateral surfaces of petiole smooth and shining. Dorsal and lateral surfaces of postpetiole smooth and shining.

Standing pilosity sparse. Dorsal face of head with decumbent setae; one pair of erect setae on frontal lobes. Clypeus with one pair of erect setae. Anterior clypeal margin with two to three pairs of long setae mixed with short setae laterally. Scape with appressed setae. Mesonotum with appressed setae sparsely. Petiole with one pair of suberect setae posteriorly. Postpetiole with one pair of suberect setae posteriorly. Fourth abdominal tergite with short appressed setae sparsely.

Body color brown.

Queen and male

Queens and males of this species are unknown.

Distribution

This species is known only from the type locality in Malaysia (Borneo) (Fig. 5).

Discussion

Phylogenetic position of C. khmerensis sp. nov.

The MP and ML analyses consistently placed *C. khmerensis* sp. nov. nested within node A, and forming an unresolved trichotomy with *C. borneensis*-group and node C (Fig. 2). The reason for this is not clear, but might be due to the large amount of missing data. The BI analysis recovered *C. khmerensis* sp. nov. nested within node A, and sister to node C, albeit with a low support value. The node A corresponds to node 231 (Blaimer 2012c, fig. 3) of the Australo-Asian *Crematogaster* clade (Blaimer 2012c, fig. 2, III), the node C corresponds to node 230 (Blaimer 2012c, fig. 3).

In this study, the consensus tree obtained by the BI analysis was used to infer the phylogenetic relationships as our results (Fig. 1) from the BI analysis had similar results to the tree inferred by Blaimer (2012c). The position of *C. khmerensis* sp. nov. is recovered as a sister to clade C, which

comprises of *C. fruhstorferi* + *C.* ss_AUS5 + *C. mjobergi* + the *C. tetracantha*-group, with the basal *C. borneensis*-group (Fig. 2).

Morphological assignment of species

While we did not find conclusive evidence of synapomorphies uniting *C. khmerensis* sp. nov. and clade C in this study, several potential characters are suggested below. *Crematogaster fruhstorferi* has short, stout propodeal spines and large propodeal spiracles. The propodeal spines of *C. fruhstorferi* are diverged posteriorly, but not directed vertically as in *C. khmerensis* sp. nov. The propodeal spiracles of *C. fruhstorferi* are as large as the base of the propodeal spines, but not 2–3 times as large as in *C. khmerensis* sp. nov. These morphological data suggest affinity, but are inconclusive. The *C. tetracantha*-group has laterally-developed pronotal shoulders (Blaimer 2012b), especially *C. dahlii* and *C. tetracantha*, which have pronotal shoulders distinctly developed as spines. The presence of laterally-developed pronotal shoulders suggests a close affiliation between *C. khmerensis* sp. nov. and the members of the *C. tetracantha*-group, even though *C. fruhstorferi*, *C. mjobergi* and *C.* ssAUS5 do not have developed pronotal shoulders. The phylogenetic relationship inferred by molecular analysis indicates that laterally-developed pronotal shoulders evolved independently and in two steps, i.e., once in *C. khmerensis* sp. nov. and also at base of the *C. tetracantha*-group, or it is also parsimonious that the shoulders were gained in the ancestor of the clade B then lost in the *C. mjobergi* and C. ssAUS5 clade.

Based on molecular and morphological evidence, we treat *C. khmerensis* sp. nov. as its own species group, the *Crematogaster khmerensis*-group. In this study, no obvious synapomorphies were identified among *C. khmerensis* sp. nov. and clade C. Since the existence of undescribed and/or unanalyzed species may decrease the resolution of higher taxonomic levels, the inclusion of more species and morphological characters in both the queen and male castes is considered necessary to corroborate its validity. Further studies should therefore be conducted on closely related taxa to more accurately clarify the position of the *C. khmerensis*-group.

Although we do not have fresh material of *C. pfeifferi* sp. nov. for molecular phylogenetic analysis, the species is considered to be most closely related to *C. khmerensis* sp. nov. and is therefore assigned as a sister species.

Dispersal history of the Australo-Asian Crematogaster clade

Blaimer (2012c, fig. 3c) proposed that the Australo-Asian *Crematogaster* clade evolved in Southeast Asia before colonizing the adjacent Australasian region. Clade C, comprising *C. fruhstorferi* + *C.* ss_AUS5 + *C. mjobergi* + the *C. tetracantha*-group, is mostly distributed eastward of Wallace's line, although *C. fruhstorferi* is also known from Borneo (Blaimer 2012c, CASENT0193728 specimen). The position of *C. khmerensis* sp. nov., as sister to *C. fruhstorferi* and the Australasian clade, supports Blaimer's hypothesis of dispersal from the Oriental to the Australian regions (Figs 2, 5), suggesting that sequential dispersal events occurred in different lineages and that they were widespread.

The divergence time estimation and phylogenetic analysis suggest that the most recent common ancestor of *C. khmerensis* sp. nov. and clade C was inferred to be 15 million years old (Middle Miocene), when the Sundaic region was connected to the extended Indochinese Peninsula (Lohman *et al.* 2011, fig. 2d). The common ancestor of the *C. khmerensis*-group was likely distributed over parts of the Sundaic region during periods of low sea level, which would have allowed it to disperse into Indochina and Sundaland before diverging into different taxa as a result of vicariance. The remaining taxa may have dispersed to Sulawesi and Australasian regions east of Wallace's line and diverged. Our divergence time estimations were approximately 5 million years younger than those attained by Blaimer (2012c). While her analysis estimated the node 245 (Blaimer 2012c, fig. 3) at ca 27 Mya, our analysis estimated the same node at ca

21 Mya (Fig. 3). This incongruence might be due to the analysis with a subset of her taxon matrix and different method.

Despite extensive field surveys by myrmecologists on the Indochinese Peninsula and Borneo, only two series of the *C. khmerensis*-group have ever been recorded from Cambodia and Borneo. The members of the *C. khmerensis*-group are therefore considered to be rare and relatively old taxa with restricted distribution ranges.

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