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OPEN Multigene phylogenetics of Polycephalomyces (Ophiocordycipitaceae, Hypocreales), with two new species from Thailand

Yuan-Pin Xiao^{1,2,3}, Ting-Chi Wen¹, Sinang Hongsanan², Rajesh Jeewon⁵, J. Jennifer Luangsaard⁴, Siraprapa Brooks³, Dhanushka Nadeeshan Wanasinghe^{2,3}, Feng-Yao Long¹ & Kevin D. Hyde^{2,3}

Polycephalomyces (Ophiocordycipitaceae) species are found in subtropical regions and are parasitic or hyperparasitic on insects. Two new species, P. aurantiacus and P. marginaliradians, parasitic on Ophiocordyceps barnesii and larva of Cossidae respectively, are introduced in this paper. Morphological comparison with extant species and DNA based phylogenies from analyses of a multigene dataset support the establishment of the new taxa. Polycephalomyces aurantiacus, exhibiting a hyperparasitic lifestyle on Ophiocordyceps barnesii, differs from other species in producing orange conidia in mass and have longer β -phialides in culture. *Polycephalomyces marginaliradians* differs from other Ophiocordyceps species by producing single stromata with a stipe, smaller perithecia and branched α -phialides and catenate α -conidia and is parasitic on Cossidae. A combined nrSSU, nrLSU, ITS, tef-1a, rpb1 and rpb2 sequence data was analysed phylogenetically including Ophiocordyceps and Polycephalomyces taxa. The new species described herein are clearly distinct from other species in Polycephalomyces. We provide a key to the species of Polycephalomyces and discuss relevant interspecies relationships.

The genus Polycephalomyces was introduced by Kobayasi¹ to accommodate P. formosus Kobayasi (1941), based on its asexual characteristics² and it is presently accommodated in Ophiocordycipitaceae³. Phylogenetic placement of Polycephalomyces has always been a debate within the clavicipitoid fungi as the taxonomic hypotheses based on host substrate and sexual morph affinities were controversial^{1,4,5}. Kepler et al.⁵ amended the taxonomic circumscription of Polycephalomyces and accepted twelve species (i.e. P. cuboideus, P. cylindrosporus, P. ditmarii, P. formosus, P. kanzashiznus, P. nipponicus, P. paracubiodeus, P. prolificus, P. ramosopulvinatus, P. ramosus, P. ryogamiensis and P. tomentosus) in Ophiocordycipitaceae based on phylogenetic analyses. Later, P. sinensis⁶, P. lianzhouensis⁷, P. yunnanensis⁸, P. agaricus⁹ and P. onorei¹⁰ were introduced as new species within Polycephalomyces based on morphology and DNA sequence data. Then, Liang et al. 11 introduced a new species P. ponerae in the genus Polycephalomyces. Based on recent morphological studies and DNA based phylogenetic analyses, Polycephalomyces taxa have been segregated in two sister clades within Ophiocordycipitaceae^{5,12}. Matočec et al. 12 considered those two clades as two different genera, viz. Perennicordyceps Matočec & I. Kušan and Polycephalomyces. Perennicordyceps comprises four species (i.e. Pe. cuboidea, Pe. paracuboidea, Pe. prolifica, and Pe. ryogamiensis), which are characterized by superficial perithecia, and hirsutella-like or acremonium-like

¹Engineering Research Center of Southwest Bio-Pharmaceutical Resources, Ministry of Education, Guizhou University, Guiyang, Guizhou Province, 550025, China. ²Center of Excellence in Fungal Research, Mae Fah Luang University, Chiang Rai, 57100, Thailand. ³School of Science, Mae Fah Luang University, Chiang Rai, 57100, Thailand. ⁴Microbe Interaction and Ecology Laboratory, BIOTEC, NSTDA, 113 Thailand Science Park, Pathum Thani, 12120, Thailand. ⁵Department of Health Sciences, Faculty of Science, University of Mauritius, Reduit, 80837, Mauritius. Correspondence and requests for materials should be addressed to T.-C.W. (email: tingchiwen@yahoo.com)

asexual morphs^{12,13}. *Polycephalomyces* comprised eight species¹². Maharachchikumbura *et al*.^{14,15} and Wijayawardene *et al*.³ maintained *Polycephalomyces* within the Ophiocordycipitaceae.

The sexual morph of *Polycephalomyces* has been recorded as fertile, capitulate, globose, tuberiform to pulvinate stromata and immersed, elongated pyriform perithecia¹², while the asexual morph has branched or unbranched synnemata, ending up with clavate to spherically flared, hymeniform aggregations of conidiophores, which produce large masses of conidia united in collective globular mucus¹². There is only one species (*P. lianzhouensis*) reported with both sexual and asexual morphs. *Ophiocordyceps* fungi are ecologically important host species for *Polycephalomyces*, and to date five *Ophiocordyceps* species have been reported to be associated with *Polycephalomyces*^{6,8,9,16–18}. At the Engineering Research Center of Southwest Bio-Pharmaceutical Resources (Guizhou University, China) in collaboration with the Center of Excellence in Fungal Research (Mae Fah Luang University, Thailand), we are investigating diversity of microfungi associated with insects in the tropics and clarify their taxonomy based on morphology and multigene phylogeny^{14,15,19–24}.

Results

Molecular phylogeny. Table S1 comprises 39 taxa (including the seven newly collected taxa) analysed herein and their accession numbers. DNA sequence data of the new species have been submitted to GenBank. A concatenated sequence data-set was analyzed comprising 5003 characters with gaps (SSU: 971, LSU: 813, ITS: 667, TEF: 875, RPB1: 659, RPB2: 1018).

The RAxML analysis of the combined dataset yielded a best scoring tree (Fig. 1) with a final ML optimization likelihood value of -23008.064357. The matrix had 1750 distinct alignment patterns, with 39.76% of undetermined characters or gaps. Parameters for the GTR model of the concatenated data set were as follows: Estimated base frequencies; A = 0.236157, C = 0.276387, G = 0.278052, T = 0.209403; substitution rates AC = 1.217462, AG = 3.345152, AT = 0.776221, CG = 1.574418, CT = 6.177706, GT = 1.000; gamma distribution shape parameter α = 0.251124. The Bayesian analysis resulted in 20001 trees after 2000000 generations. The first 4000 trees, representing the burn-in phase of the analyses, were discarded, while the remaining 16001 trees were used for calculating posterior probabilities in the majority rule consensus tree.

The genus *Polycephalomyces* currently includes 15 species and only 11 species have available DNA sequence data in GenBank (Table S1), excluding the new taxa described in this study. Our multigene phylogenetic analyses herein reveal that our new taxa constitute a strongly supported monophyletic subclade and nested in between other *Polycephalomyces* species (Fig. 1). In particular it is noted that *Polycephalomyces aurantiacus* and *P. marginaliradians* share a close phylogenetic affinity to *P. nipponicus* and *P. kanzashianus* (Fig. 1).

In this paper, we illustrate a collection of *Ophiocordyceps barnesii*, which was parasitized by a *Polycephalomyces* species. Two new species of *Polycephalomyces*, one from *Ophiocordyceps barnesii*, and one from a Cossidae host are also introduced. A phylogenetic tree based on multigene sequence analyses for *Ophiocordyceps* (11 species) and *Polycephalomyces* (13 species), is also provided.

Description of *Ophiocordyceps barnesii* (Thwaites) G.H. Sung, J.M. Sung, Hywel-Jones & Spatafora, in Sung, Hywel-Jones, Sung, Luangsa-ard, Shrestha & Spatafora, Stud. Mycol. 57: 40 (2007). Index Fungorum number: 504230; Facesoffungi number: FoF 03810 (Fig. 2).

Parasitic on larvae (Coleopteran), buried in the soil. Sexual morph: *Host* 2–2.5 long × 0.5–1 cm wide, brown to dark brown without hyphae on the surface. *Stromata* 13–20 long × 0.5–1 cm diam., mostly single, stipitate, unbranched or branched into 2 or 3 fertile head, arising from between the head and thorax of larva (fusiformis), dark-brown, fleshy, cylindrical, often flexuous or angularly crooked. *Stipe* 1–2 cm long, 2–3 mm diam., brown, with a fertile apex. *Fertile head* 2–5 cm long, 1–3.5 mm diam., single or branched more than 2, cylindrical, apically tapered, brown with orange mycelium cover on the surface. *Perithecia* 308–389 × 98–132 μm (\overline{x} = 349 × 115 μm, n = 60), immersed, brown, elongated pyriform or flask-shaped, thick-walled. *Peridium* 12–21 μm (\overline{x} = 16 μm, n = 60) wide, brown, *textura angularis* to *textura globulosa* to *textura prismatica*. *Asci* 195–229 × 6–9 μm (\overline{x} = 212 × 7.5 μm, n = 90), 8-spored, hyaline, filiform, with a thin apex. *Apical cap* 3.1–4.2 × 4.1–5.4 μm (\overline{x} = 3.6 × 4.7 μm, n = 60), with a small channel in the center. *Ascospores* 155–200 × 2.2–2.7 μm (\overline{x} = 178 × 2.5 μm, n = 60), 3-septate, easily breaking into 4 part-spores, filiform, tapered at each end. *Secondary ascospores* 31.6–41.6 × 2.2–2.7 μm (\overline{x} = 36.6 × 2.5 μm, n = 90) cylindrical, thickening at each end or tapered at one end, straight, hyaline, smooth-walled. Asexual morph: undetermined.

Notes: We collected *Ophiocordyceps barnesii* in this study which was colonized by an orange hyperparasite which we introduce below as *Polycephalomyces aurantiacus*. This species may be important in future industrial production of *Cordyceps* species, which are increasingly being produced because of their medicinal properties and biopesticides potential^{25,26}. The specimen was deposited in MFLU Herbarium (MFLU 17-1393).

Description of *Polycephalomyces aurantiacus* Y.P. Xiao, T.C. Wen & K.D. Hyde, *sp. nov*. Index Fungorum number: IF553936; Facesoffungi number: FoF 03811 (Figs 3, 4).

Etymology: The specific epithet refers to the color of conidia in mass in the specimen and colony. Holotype: MFLU 17-1393

Hyperparasite on Ophiocordyceps barnesii (Ophiocordycipitaceae), buried in the soil. Sexual morph: undetermined. Asexual morph: Synnemata solitary or not solitary, arising from the fertile head of the stromata, flat-shaped, orange color. Phialides $9.9-14.3\times0.7-1.4\,\mu\text{m}$ ($\overline{x}=12.1\times1.1\,\mu\text{m}$, n=90) hyaline. Conidia $2-2.6\times1.4-2.1\,\mu\text{m}$ ($\overline{x}=2.3\times1.8\,\mu\text{m}$, n=90), oval to globose shape, hyaline, one-celled, smooth-walled, orange in mass.

Colonies on PDA medium, growing slowly, attaining 4 cm in 17 days at 25 °C, white, reverse yellow. Synnemata emerging after 30 days, solitary or not solitary, branched or unbranched, $1.3-2.2 \mu m \log (\bar{x} = 60)$, showing 1-2

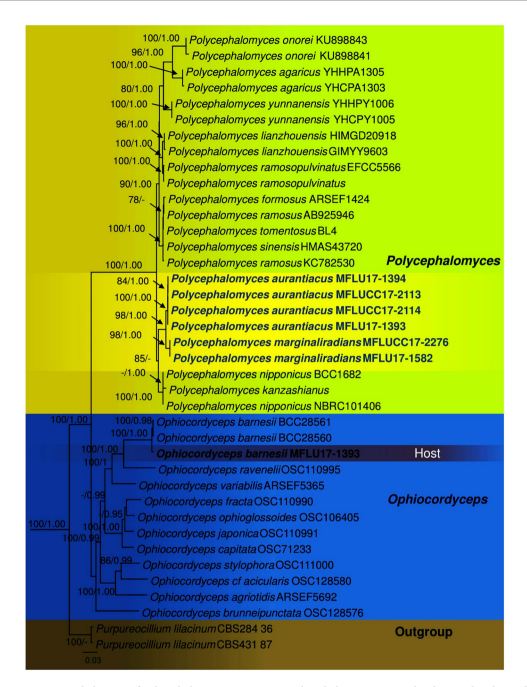


Figure 1. Phylogram of Polycephalomyces aurantiacus, Polycephalomyces marginaliradians and Ophiocordyceps barnesii generated from maximum likelihood analysis of ITS, SSU, LSU, RPB1, RPB2 and $TEF1\alpha$ sequence data. Purpureocillium lilacinum CBS 284.36 and Purpureocillium lilacinum CBS 431.87 were used as outgroup taxon. Maximum likelihood bootstrap values greater than 70% and Bayesian posterior probabilities over 0.9 are indicated above the nodes. The new species were indicated in blue. The host of Polycephalomyces aurantiacus is indicated in bold.

radiating ring-like distributions. Conidial masses generating from the apex of the synnemata or covering the surface of the colony (Fig. 4). Hyphae hyaline, branched, smooth-walled, 0.3–50 mm ($\overline{x}=20$) wide. Conidiophores undetermined, not clear. Phialides existing in two types: α -and β -phialides. α -phialides 10.4– 18.3×0.8 – $1.8\,\mu$ m ($\overline{x}=14.4\times1.3\,\mu$ m, n=90) hyaline, narrow slender, smooth. β -phialides 22.9– 64.2×1 – $1.5\,\mu$ m ($\overline{x}=43.6\times1.3\,\mu$ m, n=90) solitary, growing from hyphae, narrow slender, catenateblasto conidia, smooth. α -conidia 1.8– 2.2×1.4 – $1.9\,\mu$ m ($\overline{x}=2\times1.7\,\mu$ m, n=90) globose to subglobose, occurring in the conidial mass on the agar or on the final portion of synnemata, one-celled, smooth-walled, yellow slimy in mass. β -conidia 3.2– 3.9×1.4 – $1.8\,\mu$ m ($\overline{x}=3.5\times1.6\,\mu$ m, n=90) fusiform, and produced on the surface mycelium of colony or on the top of the synnemata, one-celled, smooth-walled, hyaline, usually in chains on a phialide.



Figure 2. Ophiocordyceps barnesii MFLU 17-1393 (host). (a) Habitat. (b) Overview of the host and stromata. (c) Host. (d) Stroma. (e) Cross section of stroma. (f) Cross sections showing the immersed perithecia. (g) Perithecia. (h-k) Asci. (l-o) Secondary ascospores. Scale Bars: b=5 cm, c=5 mm, d=2 cm, e, f=500 μ m, p=200 μ m, g-k=100 μ m, l-o, s=20 μ m, q=10 μ m, r=5 μ m.

Material examined: THAILAND, Prachuap Khiri Khan. On dead larvae (Coleopteran), 29 July 2015, YuanPin Xiao, BK15072907 (MFLU 17-1393, holotype); BK15072902, BK15072906 (MFLU 17-1394, HKAS100693, paratypes); ex-type living cultures, MFLUCC 17-2113, MFLUCC 17-2114, MFLUCC 17-2115, KUMCC 17-0256, KUMCC 17-0257.

Description of *Polycephalomyces marginaliradians* Y.P. Xiao, T.C. Wen & K.D. Hyde, *sp. nov.* Index Fungorum number: IF553937; Facesoffungi number: FoF 03812 (Figs 5, 6).

Etymology: The specific epithet refers to the feature of the colonies on the culture. Holotype: MFLU 17-1582

Parasitic on a Cossidae larva (Lepidoptera), buried in the soil. Sexual morph: Thallus within host. Host 3.2–3.5 long \times 0.4–0.6 cm wide, yellow to brown, without hyphae on the surface. Stromata 3–3.5 long \times 0.2–0.45 cm diam., mostly single, stipitate, cylindrical, unbranched or branched, arising from the head of larva, brown to yellow. Stipe 1–2 cm long, 2–3 mm diam., cylindrical, yellow to brown, with one or two lateral fertile head. Fertile head 0.40–0.42 cm long, 0.3–0.45 mm diam., capitate, lateral, globose to subglobose, pale yellow to yellow, with protruding ostiolar necks. Ascomata 676–803 \times 246–328 μ m (\bar{x} = 739 \times 287 μ m, n = 60), immersed, yellow,

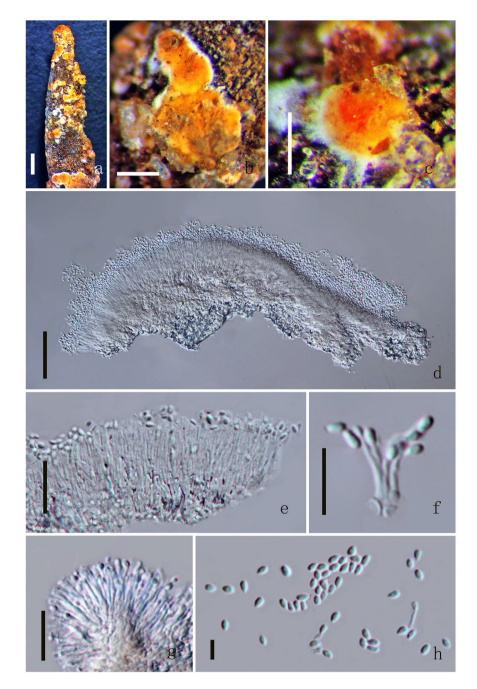


Figure 3. Polycephalomyces aurantiacus MFLU 17-1393. (a) Mycelium on the surface of stroma. (b,c) Conidiomata. (d) Section of conidiomat. (e-g) Phialides. (h) Conidia. Scale Bars: $a=1000\,\mu\text{m},\,b=500\,\mu\text{m},\,c=200\,\mu\text{m},\,d=50\,\mu\text{m},\,e=20\,\mu\text{m},\,f=g=10\,\mu\text{m},\,h=5\,\mu\text{m}.$

flask-shaped, thick-walled. *Peridium* 11–19 μ m ($\overline{x}=15\,\mu$ m, n=60) wide, brown, textura angularis to textura globulosa to textura prismatica. *Asci* 459–556 × 3.1–4.3 μ m ($\overline{x}=508\times3.7\,\mu$ m, n=90), 8-spored, hyaline, filiform, with a thin apex. *Apical cap* 1.4–2.5 × 2.2–3.2 μ m ($\overline{x}=2\times2.7\,\mu$ m, n=60), with a small channel in the center. *Ascospores* as long as the asci, easily breaking into part-spores, filiform. *Secondary ascospores* 3.2–4.2 × 1.3–1.7 μ m ($\overline{x}=3.8\times1.5\,\mu$ m, n=90) cylindrical, straight, hyaline, smooth. Asexual morph: *Synnemata* solitary or not solitary, arising from the fertile head of the host, cylindrical, pale yellow. *Phialides* 11–14.4 × 1.2–1.8 μ m ($\overline{x}=12.7\times1.5\,\mu$ m, n=90) hyaline. *Conidia* 3.6–4.9 × 1.8–2.5 μ m ($\overline{x}=4.2\times2.1\,\mu$ m, n=90), fusiform, hyaline, one-celled, smooth-walled.

Colonies on PDA medium, circular, attaining 4 cm in 10 days at 25 °C, white, reverse yellow. Synnemata emerging after 14 days in the margin of the colony, single or branched into 2 or 3 branched, 3200.8–4566.3 \times 142.9–661.8 μ m ($\overline{x}=3883.5 \times 402.3 \,\mu$ m, n = 30), showing 1–2 radiating ring-like distributions. Conidial masses generating from the middle of the synnemata or covering the surface of the colony, pale yellow to yellow, with hyaline to pale yellow exucate. Hyphae hyaline, branched, smooth-walled, 1.8–2.7 μ m ($\overline{x}=2.2$) wide.

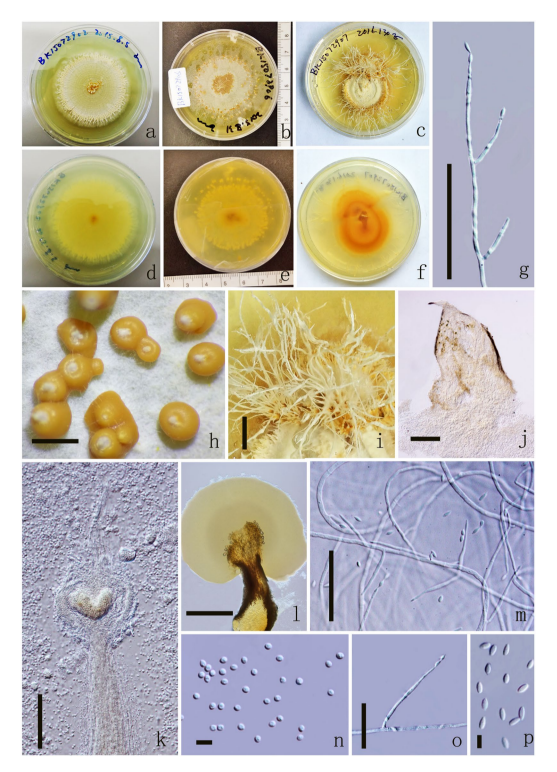


Figure 4. Polycephalomyces aurantiacus MFLUCC 17-2113. (a-c) Upper side of the culture. (d-f) Reverse side of the culture. (g,o) β -phialides. (h-l) Synnemata growing on PDA medium. (m) β -phialides with hyphae. (n) α -conidia. (p) β -conidia. Scale Bars: $h=1000\,\mu m$, $i=2000\,\mu m$, $j=200\,\mu m$, $l=500\,\mu m$, $g,k,m=50\,\mu m$, $o=20\,\mu m$, $n,p=5\,\mu m$.

Conidiophores undetermined, not clear. Phialides existing in two types: α -and β -phialides. α -phialides $11-14.4\times1.2-1.8\,\mu\text{m}$ ($\overline{x}=12.7\times1.5\,\mu\text{m}$, n=90), hyaline, smooth, elongated lageniform, caespitose, palisade-like, crowed, monoverticillate, mostly branched into 2 phialides, 3 branched on one metula. β -phialides $12.8-23.9\times1.8-2.7\,\mu\text{m}$ ($\overline{x}=18.3\times2.2\,\mu\text{m}$, n=90), hyaline, smooth, solitary, growing from hyphae, narrow slender to narrow lageniform, with or without metula at the base. α -conidia $1.9-2.6\,\mu\text{m}$ ($\overline{x}=2.3\,\mu\text{m}$, n=90) diam, globose, catenate, occurring in the conidial mass on the middle of synnemata, one-celled, smooth-walled, pale yellow

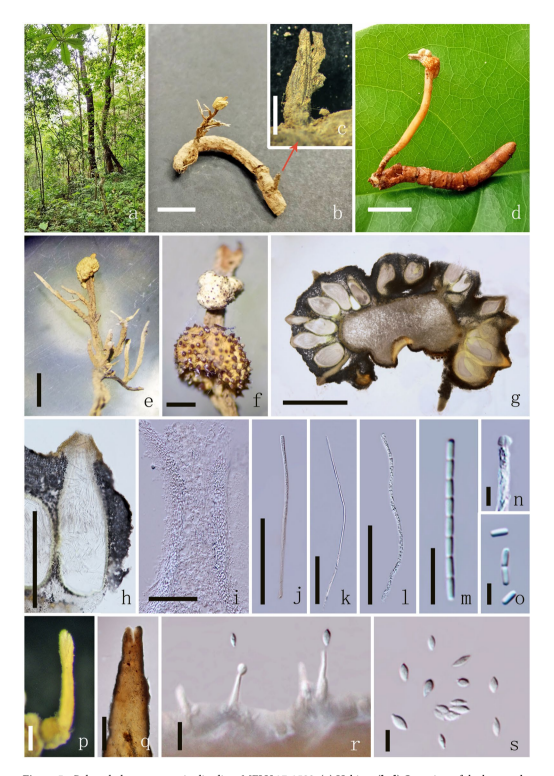


Figure 5. Polycephalomyces marginaliradians MFLU 17-1582. (a) Habitat. (b,d) Overview of the host and stromata. (c) Part of the stroma. (e,f) Stroma. (g) Cross sections showing the immersed perithecia. (h) Perithecia. (j-l) Asci. (m) Part of the ascospores. (n) Apical cap. (o) Secondary ascospores. (p,q) Synnemata. (r) Phialide. (s) Conidia. Scale Bars: b-d=1 cm, e,f=2000 μ m, g=1000 μ m, h=500 μ m, p,q=200 μ m, j-l=100 μ m, i=50 μ m, m,r=10 μ m, n,o,s=5 μ m.

slimy in mass. β -conidia $3.1-3.9 \times 1.6-2.1 \,\mu m$ ($\overline{x}=3.5 \times 1.8 \,\mu m$, n=90) fusiform, and produced on the surface mycelium of colony or on the branch of the synnemata, one-celled, smooth-walled, hyaline.

Material examined: THAILAND, Chiang Mai, The Mushroom Research Center. On dead Cossidae larvae (Lepidoptera), 11 June 2017, Yuan Pin Xiao, MRC170611 (MFLU 17-1582, holotype); CM48 (MFLU 17-1583

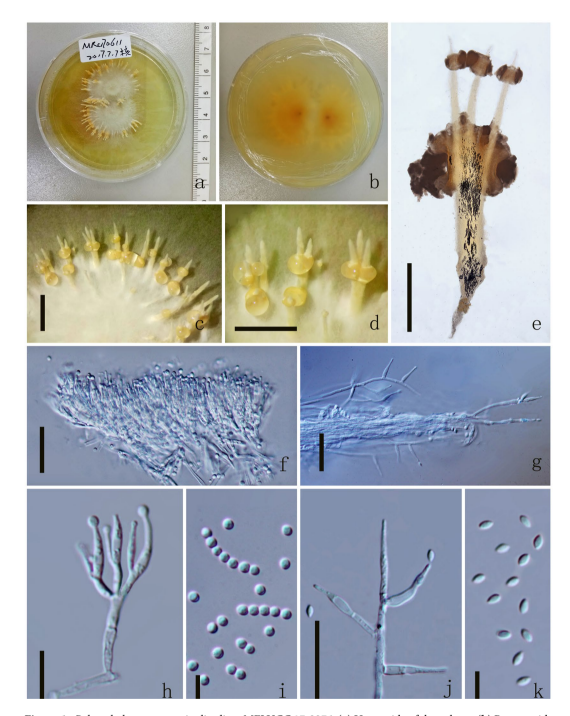


Figure 6. Polycephalomyces marginaliradians MFLUCC 17-2276. (a) Upper side of the culture. (b) Reverse side of the culture. (c,d) Synnemata growing on PDA medium. (e) Synnemata. (f,h) α -phialides. (g,j) β -phialides with hyphae. (i) α -conidia. (k) β -conidia. Scale Bars: c, d = 5000 μm, e = 1000 μm, f, g, j = 20 μm, h = 10 μm, i, k = 5 μm.

MFLU 17-1584, HKAS100694, paratypes); ex-type living cultures, MFLUCC 17-2276, MFLUCC 17-2277, MFLUCC 17-2278, KUMCC 17-0258, KUMCC 17-0259.

Discussion

Studies based on morphology and DNA sequence analyses have provided insights into the phylogeny of *Polycephalomyces* to resolve generic delimitation. Species of this genus are commonly known to exhibit a parasitic mode of life on insects and other fungi^{2,5,12}. Our fungal diversity studies on entomophagous fungi have led to the discovery of two species, new to science, which we accommodate in *Polycephalomyces*. Molecular data also reveals that our new genus belongs to the family Ophiocordycipitaceae as circumscribed by Matočec *et al.*¹². Species which exist in their sexual state display morphs such as fertile, capitulate, globose, tuberiform to pulvinate stromata and immersed, elongated pyriform perithecia while the asexual morphs occur as branched or unbranched

synnemata, ending up with clavate to spherically flared, hymeniform aggregations of conidiophores, and produce large masses of conidia united in collective globular mucus¹². To date, six species, including P. $ramosus^{16,17}$, P. $sinensis^{6,18}$, and P. $agaricus^9$ are considered as parasites of entomogenous fungi, while six species are recorded as entomogenous^{6,7,9,11}. Some species such as P. lianzhouensis and P. yunnanensis colonise both entomogenous fungi and insects^{7,8}. Because of their economic importance, species of this genus have been the subject for various research. The most recent new species introduced is P. yunnanensis and multigene phylogeny reveals a close relationship to P. formosus, P. ramosopulvinatus and P. sinensis based on 5-loci (nrSSU, nrLSU, tef- 1α , rpb1 and rpb2) phylogenetic analyses⁸.

Our taxonomic investigations herein reveal two new species of Polycephalomyces, P. aurantiacus and P. marginaliradians. Our morphological examination suggests that both species fit clearly within the generic concept of Polycephalomyces and both species produce two types of conidia. However, they exhibit different mode of life and there are sufficent morphological differences that can justify their segregation into two species. These two new species similar to P. agaricus, P. formosus, P. ponerae, P. sinensis, P. ramosus, and P. yunnanensis have produce two types of conidia, while P. ditmarii, P. lianzhouensis, P. paludosus and P. tomentosus have only one type of conidia. Polycephalomyces aurantiacus and P. marginaliradians have two types of phialides, while P. formosus and P. sinensis have only one type of phialide. Polycephalomyces ponerae also differs from P. aurantiacus and P. marginaliradians by producing Akanthomyces-like β-phialides and parasitic on ant (Ponera Latreille). Polycephalomyces agaricus differs from P. aurantiacus and P. marginaliradians by producting agaric shaped synnemata and parasitic on Ophiocordyceps sp. Polycephalomyces yunnanensis is distinct from P. aurantiacus and P. marginaliradians as it produces longer α -conidia fusiform, catenate or clump together β -conidia and parasitic on O. nutans. Polycephalomyces ramosus differs from P. aurantiacus and P. marginaliradians in having longer synnemata, shorter β-phialides and parasitic on Hirsutella guignardii. Polycephalomyces aurantiacus is distinct from P. marginaliradians as it is parasitic on the fungus Ophiocordyceps barnesii and produces longer synnemata, shorter \(\beta\)-phialides and catenate α -conidia, wheras *P. marginaliradians* is parasitic on insect and produces shorter synnemata, longer β -phialides and catenate β -conidia (further morphological differences are outlined in Table S2).

Phylogeny based on our concatenated dataset recovered also support that our two new species belong to Polycephalomyces and are distinct from each other (Fig. 1). A close relationship is observed between the two species, but both constitute independent and strongly supported monophyletic subclades indicative of two phylogenetically distinct speces. To further compare our two species, we delved in pairwise nucleotide sequence comparison and noted sufficient differences to justify them as independent taxa²⁷. ITS pairwise nucleotide sequence comparison between P. aurantiacus and P. marginaliradians revealed striking differences in 15 base pairs that justifies that both are different from each other and hence can be considered as two distinct species. There are also 6, 21, 4, 15 and 13 differences in the nrSSU, $ref-1\alpha$, rpb1, and rpb2 DNA sequence data respectively. Two species not considered in our phylogenetic sampling are P. ditmarii and P. paludosus due to the unavailability of sequence data. However these two are different from our new species with respect to one type of conidia occurring on its natural substrate and under cultural conditions^{28,29}. The hosts from which our new species have been recovered are also different. Polycephalomyces ponerae was not considered in our phylogenetic sampling as the DNA (ITS) sequence is too short, ambiguous and did not align well with other species. However, Polycephalomyces ponerae is morphologically different from our new species with respect to Akanthomyces-like β-phialides and parasitic on ant (Ponera Latreille). Further morphological differences among species are detailed in Tables S2 and S3.

Our multigene phylogeny derived herein also provides robust and well-resolved intergeneric relationships between Polycephalomyces and Ophiocordyceps. Members of both genera are clearly distinct from each other and we managed to successfully identify and sequence Ophiocordyceps barnesii, the host from which P. aurantiacus was isolated. Further interspecies taxonomic relationships are also elucidated in our molecular phylogeny. All Polycephalomyces species currently analysed constitute a strongly supported monophyletic lineage (Fig. 1), which corroborates with previous taxonomic schemes^{5,12}. In particular, a robust relationship in observed between P. onorei and P. agaricus sharing P. yunnanensis as sister taxa. These three species are also markedly different in terms of morphological characters. Polycephalomyces yunnanensis is clearly distinct from P. onorei and P. agaricus in terms of being parasitic on Ophiocordyceps nutans (Pat.) G.H. Sung, J.M. Sung, Hywel-Jones & Spatafora, longer synnemata, cylindrical to subulate α -phialides and subglobose or ellipsoidal α -conidia^{8,10}. Polycephalomyces ramosopulvinatus is closely related to P. lianzhouensis, but each species is positioned in different well-supported subclades and hence merit species status. Polycephalomyces ramosopulvinatus is also different from P. lianzhouensis with respect to being parasitic on nymph of Cicada and characterised by a long stipe and pseudo-immersed, pyriform perithecia. While phylogeny resolves our new species into well-segregated subclades, we note that relationships of P. formosus, P. tomentosus, P. ramosus and P. sinensis are still obscure and the concatenated dataset used herein did not provide adequate species resolution. A similar phylogenetic scenario is observed for P. nipponicus and P. kazanshianus. Whether these species are conspecific warrants further taxonomic investigations. The latter two species do share some morphological resemblances to P. marginaliradians especially with respect to the yellow cylindrical stipe with capitate lateral fertile part (known from their sexual morph). However, P. marginaliradians differs in having a capitate stromata with stipe, smaller perithecia and parasitic on Cossidae, while *P. nipponicus* and *P. kanzashianus* have polycephalous stromata and parasitic on Cicadidae. Meanwhile, *P.* onorei and P. ramosopulvinatus are distinct from P. marginaliradians by producing bigger perithecia and parasitic on caterpillar (Arctinae) and nymph of Cicada respectively. Phylogenies retrieved herein also support them as separate taxonomic entities.

1. Stromata arising from insect
5. α -conidia globose to subglobose (1.4–3.2 × 1.2–2.2) μ m6
5. α -conidia ovoid (2.4–3.2 × 1.6–2.4) μ m
6. β-phialides <i>Akanthomyces</i> -like, inflated at base, slenderneck at top
6. β-phialides lanceolate or narrowly lageniform or subulate
7. β-phialides lanceolate or narrowly lageniform, 22.9–64.2 μm length
7. β-phialides narrowly lageniform or subulate, 7–30 μm length8
8. α -conidia subglobose, not catenate, β -conidia fusiform, catenate
marginaliradians
8. α -conidia globose, catenate, β -conidia fusiform, not catenate
9. Phialides lanceolate or narrowly lageniform, 12.5–66 µm length
9. Phialides cylindrical, subulate, 10–15 μm length
10. Host is insect
10. Host is myxomycetes
11. Conidia globose to subglobose or cylindrical12
11. Conidia obovoid, covered by a mucus, agglutinating
12. Conidia globose to subglobose, 2.2–3.4 × 1.3–1.6um
12. Conidia subglobose to cylindrical, $5-7 \times 1.3-1.6$ um
13. Host is Cicada or nymph of Cicada14
13. Host is neither Cicada nor nymph of Cicada16
14. Stipe less than 80 mm
14. Stipe more than 90 mm
15. Perithecia flask-shaped, 900–1050 × 270–300 um
15. Perithecia flask-shaped or ovioid, 800–950 × 300–370 um
16. Stromata numerous17
16. Stromata single
17. Fertile part narrowly ovoid $355-473 \times 158-197 \mu\text{m}$
17. Fertile part pyriform, 854–950 × 330–395 um

Materials and Methods

Collection, isolation, and morphology study. Four fresh specimens were collected from southern Thailand (Prachuap Khiri Khan Province), and two from northern Thailand in the soil. The specimens were noted and photographed in the field and transported to the laboratory individually in plastic boxes and stored at 4°C until examined. Strains were isolated from single spore isolation from both stomata and synnemata following the protocol described in Chomnunti *et al.*³⁰. Cultures were incubated at 18 °C for 14–25 days on potato extract agar (PDA) as outlined by Vijaykrishna *et al.*³¹. Herbarium material is deposited at MFLU herbarium and HKAS herbarium and Facesoffungi numbers and Index Fungorum numbers are provided as in Jayasiri *et al.*³² and Index Fungorum³³. New species are based on recommendations outlined by Jeewon & Hyde²⁷.

DNA extraction, PCR amplification and determination of DNA sequences. DNA was extracted from both dried specimens and cultures by using E.Z.N.A.TM Fungal DNA MiniKit (Omega Biotech, CA, USA) according to the manufacturer's protocols. The primers used in PCR amplification were (Table S4); ITS4/ITS5 for internal transcribed spacer gene region (ITS)³⁴, NS1/NS4 for partial small subunit ribosomal RNA gene region (SSU)³⁴, LROR/LR5 for partial large subunit rDNA gene region (LSU)³⁵. 983 F/2218 R for partial translation elongation factor 1-alpha gene region (TEF-1 α)³⁶, CRPB1A/RPB1Cr for partial RNA polymerase II largest subunit gene region (RPB1)³⁷. RPB2-5F/RPB2-5R for partial RNA polymerase II second largest subunit gene region (RPB2)³⁷. PCR amplifications were conducted as outlined by Jeewon *et al.*^{38,39} and PCR products were sequenced by GenScript Biotechnology Co., Nanjing, China.

Phylogenetic analyses. All reference sequences were obtained from GenBank based on previously published data (Table S1). MAFFT v.7⁴⁰ (http://mafft.cbrc.jp/alignment/server/) was used to align combined datasets of ITS, SSU, LSU, TEF1 α and RPB1. BioEdit⁴¹ was used to check alignment manually. Gaps were treated as missing data. Two strains of *Perennicordyceps prolifica* (Kobayasi) Matočec & I. Kušan, in Matočec *et al.*¹² were selected as outgroup taxa.

ML trees were estimated by using the software RAxML 7.2.8 Black Box^{42,43} in the CIPRES Science Gateway platform⁴⁴. MrModeltest v.2.3⁴⁵ was used to determine the best-fit model of evolution for Bayesian analyses. MrBayes v.3.1.2⁴⁶ was used to evaluate posterior probabilities (PP)^{47,48} by Markov Chain Monte Carlo sampling (BMCMC). Six simultaneous Markov chains were run for 2,000,000 generations and trees were sampled every 100th generation and 20,000 trees were obtained. The first 20% of trees were discarded, which representing the burn-in phase of the analyses, while the remaining trees were used for calculation posterior probabilities in the majority rule consensus tree (critical values for the topological convergence diagnostic is 0.01). Pylogenetic trees were also constructed based on parsimony analyses as detailed by Cai *et al.*⁴⁹ and Jeewon *et al.*^{50,51}. Trees were

figured in FigTree v1.4.0 program⁵². Bayesian Posterior Probabilities (BYPP) equal to or great than 0.95 were given⁵³⁻⁶³ below each node (Fig. 1).

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Author Contributions

Y.P.X. and T.C.W. designed the study. Y.P.X. and F.Y.L. conducted all the experiments. Y.P.X., S.H., J.J.L., S.B., D.N., R.J. and K.D.H. analysed the result. Y.P.X., S.H., R.J., J.J.L., S.B., D.N., T.C.W. and K.D.H. edited the manuscript. All authors reviewed the manuscript and approved the manuscript for publication.

Additional Information

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