

Research Article

Species diversity and major host/substrate associations of the genus Akanthomyces (Hypocreales, Cordycipitaceae)

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Abstract

Akanthomyces, a group of fungi with rich morphological and ecological diversity in Cordycipitaceae (Ascomycota, Hypocreales), has a wide distribution amongst diverse habitats. By surveying arthropod-pathogenic fungi in China and Southeast Asia over the last six years, nine Akanthomyces spp. were found and identified. Five of these were shown to represent four known species and an undetermined species of Akanthomyces. Four of these were new species and they were named A. kunmingensis and A. subaraneicola from China, A. laosensis from Laos and A. pseudonoctuidarum from Thailand. The new species were described and illustrated according to the morphological characteristics and molecular data. Akanthomyces araneogenus, which was isolated from spiders from different regions in China, Thailand and Vietnam, was described as a newly-recorded species from Thailand and Vietnam. The phylogenetic positions of the nine species were evaluated, based on phylogenetic inferences according to five loci, namely, ITS, nrLSU, TEF, RPB1 and RPB2. In this study, we reviewed the research progress achieved for Akanthomyces regarding its taxonomy, species diversity, geographic distribution and major host/substrate associations. The morphological characteristics of 35 species in Akanthomyces, including four novel species and 31 known taxa, were also compared.

Key words: Arthropod-pathogenic fungi, Cordycipitaceae, morphology, new species, phylogenetic analyses

Introduction

Akanthomyces Lebert is one of the oldest genera in the family Cordycipitaceae (Ascomycota, Hypocreales). This genus was established by Lebert in 1858 on the basis of the type species, A. aculeatus Lebert, which was found on a moth in France (Lebert 1858). Morphologically, Akanthomyces species have been characterised asexually by white, cream or flesh-coloured cylindrical, attenuated synnematal growth covered by a hymenium-like layer of phialides producing



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one-celled catenulate conidia (Mains 1950; Samson and Evans 1974; Hsieh et al. 1997). These phialides are ellipsoidal, cylindrical or narrowly cylindrical and gradually or abruptly taper to a more or less distinct neck (Hsieh et al. 1997). Owing to extensive overlap in their morphological characteristics, Akanthomyces was once considered as a synonym of Lecanicillium W. Gams & Zare, an anamorph within Cordycipitaceae with verticillium-like morphology (Gams and Zare 2001); however, many species originally described in Lecanicillium do not form a single monophyletic clade and are distributed throughout Cordycipitaceae (Wang et al. 2020). Kepler et al. (2017) phylogenetically established the genetic boundaries in Cordycipitaceae and they proposed that Lecanicillium should be rejected and, instead, could be considered as a synonym of Akanthomyces (Kepler et al. 2017). Kepler et al. (2017) also showed that the type species of Lecanicillium, L. lecanii (Zimm.) Zare & W. Gams (as Cordyceps confragosa (Mains) G.H. Sung, J.M. Sung, Hywel-Jones & Spatafora), as well as several other Lecanicillium species, namely, L. attenuatum Zare & W. Gams, L. muscarium (Petch) Zare & W. Gams and L. sabanense Chir.-Salom., S. Restrepo & T.I. Sanjuan, fall within Akanthomyces. The teleomorph of Akanthomyces was originally described as Torrubiella Boud. and it was characterised by producing superficial perithecia on a loose mat of hyphae (subiculum) or a highly reduced non-stipitate stroma (Boudier 1885). According to the most complete taxonomic treatment of Cordycipitaceae to date, this connection was verified by DNA sequencing; since Akanthomyces was described earlier than Torrubiella, the taxonomic revision recommended Akanthomyces as the name of this genus (Kepler et al. 2017).

Over the past two decades, our efforts have been applied to the investigation of Cordycipitoid fungi, especially those located in China and Southeast Asia. To date, our study team has collected over 18,000 specimens and 7,500 strains of Cordyceps Fr. sensu lato, representing more than 450 species in total (Wang et al. 2020). These specimens and strains sufficiently revealed that Cordycipitaceae is the most complex group in Hypocreales with its varied morphological characteristics and wide-ranging hosts. Some of the genera with sexual and asexual morphs, such as Akanthomyces and Hevansia Luangsa-ard, Hywel-Jones & Spatafora, share numerous similar morphological characteristics. The genus Hevansia was erected to accommodate asexual morphs on spiders that were previously described under Akanthomyces. The type species Hevansia novoguineensis (Samson & B.L. Brady) Luangsa-ard, Hywel-Jones & Spatafora, which was previously described as Akanthomyces novoguineensis Samson & B.L. Brady, differs from Akanthomyces by the immersed perithecia of the teleomorph in a disc sitting at the top of a well-formed stipe (Aini et al. 2020); however, H. novoguineensis must now be an akanthomyces-like teleomorph (Kepler et al. 2017; Aini et al. 2020). Some Akanthomyces, Samsoniella Mongkols., Noisrip., Thanakitp., Spatafora & Luangsa-ard and Cordyceps species produce similar isaria-like asexual conidiogenous structures, such as flask-shaped phialides produced in whorls and conidia with divergent chains (Wang et al. 2020; Wang et al. 2022). Due to the extensive overlap in morphological characteristics and the lack of distinctive phenotypic variation, species in many genera, Akanthomyces in particular, are not easily classified and identified. Thus, more known species and new species in the genus Akanthomyces

need to be introduced and supported by more detailed morphological and phylogenetic evidence in combination with a larger taxon sampling.

In surveys of arthropod-pathogenic fungi from different regions in Yunnan and Hunan Province, China; Chiang Mai Province, Thailand; Nghe An Province, Vietnam; and Oudomxay Province, Laos, over the last six years, approximately nine *Akanthomyces* spp. were collected and identified. In this study, we aimed to: 1) reveal the hidden species diversity of the genus *Akanthomyces* according to phylogenetic analyses and morphological observation and 2) systematically review the geographical distribution and major host/substrate associations of *Akanthomyces* species by surveying the literature to the greatest extent possible and combining the results with those generated in our study.

Materials and methods

Soil and specimen collection

All of the soil samples were collected from Yunnan Province in China. Fungal specimens were obtained from six locations between 2017 and 2022, namely, two different locations in Yunnan Province, China, one location in Hunan Province, China, one location in Chiang Mai Province, Thailand, one location in Nghe An Province, Vietnam and one location in Oudomxay Province, Laos. Soil samples and specimens were noted and photographed in the field and then they were carefully put in plastic containers at a low temperature. After that, they were brought to the laboratory and stored at 4 °C prior to examination and isolation.

Fungal isolation and culture

The Akanthomyces strains were isolated from the soil samples, based on the methods described by Wang et al. (2015) and Wang et al. (2023b). Briefly, 2 g of soil were added to a flask containing 20 ml sterilised water and glass beads. The soil suspension was shaken for about 10 min and then diluted 100 times. Subsequently, 200 µl of the diluted soil suspension was spread on Petri dishes with solidified onion garlic agar (OGA: 20 g of grated garlic and 20 g of onion were boiled in 1 litre of distilled water for 1 h; the boiled biomass was then filtered-off and 2% agar was added). Czapek yeast extract agar (CYA, Advanced Technology and Industrial Co., Ltd., China) and potato dextrose agar (PDA, Difco, USA) were used and all media had 50 mg/l rose Bengal and 100 mg/l kanamycin added. Conidia developing on invertebrate cadavers were transplanted on to plates of PDA and cultured at 25 °C. Colonies of the isolated filamentous fungi appearing in the culture were transferred on to fresh PDA media. Each purified fungal strain was transferred to PDA slants and cultured at 25 °C until its hyphae spread across the entire slope. The emerging fungal spores were washed with sterile physiological saline to form a suspension containing 1×10^3 cells/ml. To obtain monospore cultures, a sample of the spore suspension was placed on PDA on a Petri dish utilising a sterile micropipette and then the dish was incubated at 25 °C. Voucher specimens and the corresponding isolated strains were deposited in the Yunnan Herbal Herbarium (YHH) and the Yunnan Fungal Culture Collection (YFCC), respectively, of Yunnan University, Kunming, China.

Morphological observations

The specimens were examined with an Olympus SZ61 stereomicroscope (Olympus Corporation, Tokyo, Japan). Fungal structures of the specimens, such as synnemata, phialides and conidia, were mounted on glass slides with a drop of lactophenol cotton blue solution. Cultures on PDA slants were transferred to PDA plates and then they were incubated at 25 °C for 14 d. For morphological evaluation, microscope slides were prepared by placing mycelia from the cultures on PDA medium blocks (5 mm diameter) and then overlaid with a coverslip. Micro-morphological observations and measurements were performed with a light microscope (CX40, Olympus Corporation, Tokyo, Japan) and a scanning electron microscope (Quanta 200 FEG, FEI Company, Hillsboro, USA). The individual length and width measurements were recorded for 30–100 replicates and included the absolute minima and maxima.

DNA extraction, PCR and sequencing

The specimens and axenic living cultures were prepared for DNA extraction. Genomic DNA was extracted utilising a Genomic DNA Purification kit (Qiagen GmbH, Hilden, Germany), based on the manufacturer's instructions. The primer pair ITS5/ITS4 was used to amplify a fraction of the internal transcribed spacer regions of the rDNA (ITS rDNA) (White et al. 1990). Primer pair LR5/LR0R (Vilgalys and Hester 1990; Rehner and Samuels 1994) was used to amplify a fraction of the nuclear ribosomal large subunit (nrLSU) and EF1-983F/EF1-2218R primers (Rehner and Buckley 2005) were used to amplify translation elongation factor 1α (*TEF*). For amplification of the largest and second largest subunits of RNA polymerase II (RPB1 and RPB2), PCR primer pairs RPB1-5'F/ RPB1-5'R and RPB2-5'F/RPB2-5'R (Bischoff et al. 2006; Sung et al. 2007) were employed. All of the PCR reactions were performed in a final volume of 50 µl and contained 25 µl of 2 × Taq PCR Master Mix (Tiangen Biotech Co., Ltd., Beijing, China), 0.5 µl of each primer (10 µM), 1 µl of genomic DNA and 23 µl of RNase-free water. Target gene amplification and sequencing were performed, based on the methods detailed in our prior study (Wang et al. 2020).

Phylogenetic analyses

The phylogenetic analyses were based on five genes, namely, ITS, nrLSU, TEF, *RPB1* and *RPB2*, sequences. The sequences were retrieved from GenBank (http:// www.ncbi.nlm.nih.gov/, accessed on 1 March 2023) and combined with those generated in our study. Taxon information and GenBank accession numbers are listed in Table 1. Sequences were aligned with MAFFT v.7 (http://mafft.cbrc.jp/ alignment/server/, accessed on 1 March 2023). The aligned sequences were then manually corrected when necessary. After alignment, the sequences of the genes were concatenated. Conflicts amongst the five genes were resolved with PAUP* 4.0b10 (Swofford et al. 2002). The results showed that the phylogenetic signals for the five loci were congruent (P = 0.02). The data partitions were defined for the combined dataset with PartitionFinder v.1.1.1 (Lanfear et al. 2012). Phylogenetic analyses were conducted utilising Bayesian Inference (BI) and Maximum Likelihood (ML) methods, respectively. The model selected for BI

0	Voucher	11		GenBan	k accession I	numbers		Deferrer
Species	information	Host/Substrate	ITS	nrLSU	TEF	RPB1	RPB2	Reference
Akanthomyces aculeatus	HUA 186145	-	-	MF416520	MF416465	-	-	Kepler et al. (2017)
Akanthomyces aculeatus	TS772	Lepidoptera; Sphingidae	KC519371	KC519370	KC519366	_	_	Sanjuan et al. (2014)
Akanthomyces araneicola	GY29011 [⊤]	Araneae; spider	MK942431	-	MK955950	MK955944	MK955947	Chen et al. (2019)
Akanthomyces araneogenus	GZUIF DX2 [⊤]	Araneae; spider	MH978179	_	MH978187	MH978182	MH978185	Chen et al. (2018)
Akanthomyces araneogenus	YFCC 1811934	Araneae; spider	OQ509518	OQ509505	OQ506281	OQ511530	OQ511544	This study
	YFCC 2206935	Araneae; spider	OQ509519	OQ509506	OQ506282	OQ511531	OQ511545	This study
Akanthomyces araneosus	KY11341 [⊤]	Araneae; spider	ON502826	ON502832	ON525443	-	ON525442	Chen et al. (2022)
Akanthomyces attenuatus	CBS 170.76 [⊤]	Lepidoptera; Carpocapsa pomonella	MH860970	OP752153	OP762607	OP762611	OP762615	Manfrino et al. (2022)
Akanthomyces bashanensis	CQ05621 [™]	Araneae; spider	OQ300412	OQ300420	OQ325024	-	OQ349684	Chen et al. (2023)
Akanthomyces beibeiensis	CQ05921 [™]	Araneae; spider	OQ300415	OQ300424	OQ325028	-	OQ349688	Chen et al. (2023)
Akanthomyces coccidioperitheciatus	NHJ 6709	Araneae; spider	JN049865	EU369042	EU369025	EU369067	EU369086	Kepler et al. (2012)
Akanthomyces dipterigenus	CBS 126.27	Hemiptera; Icerya purchasi	AJ292385	KM283797	KM283820	KR064300	KM283862	Kepler et al. (2017)
Akanthomyces dipterigenus	YFCC 2107933	Soil	OQ509520	OQ509507	OQ506283	OQ511532	OQ511546	This study
Akanthomyces kanyawimiae	TBRC 7242	Araneae; spider	MF140751	MF140718	MF140838	MF140784	MF140808	Mongkolsamrit et al. (2018)
	TBRC 7243	Unidentified	MF140750	MF140717	MF140837	MF140783	MF140807	Mongkolsamrit et al. (2018)
Akanthomyces	YFCC 1708939	Araneae; spider	OQ509521	OQ509508	OQ506284	OQ511533	OQ511547	This study
kunmingensis	YFCC 1808940 [⊤]	Araneae; spider	OQ509522	OQ509509	OQ506285	OQ511534	OQ511548	This study
Akanthomyces laosensis	YFCC 1910941 [⊤]	Lepidoptera; Noctuidae	OQ509523	OQ509510	OQ506286	OQ511535	OQ511549	This study
	YFCC 1910942	Lepidoptera; Noctuidae	OQ509524	OQ509511	OQ506287	OQ511536	OQ511550	This study
Akanthomyces lecanii	CBS 101247	Hemiptera; Coccus viridis	JN049836	AF339555	DQ522359	DQ522407	DQ522466	Kepler et al. (2012)
Akanthomyces Iepidopterorum	GZAC SD05151⊺	Lepidoptera (pupa)	MT705973	-	_	-	MT727044	Chen et al. (2020b)
Akanthomyces muscarius	CBS 455.70B	-	-	MH871560	-	-	-	Kepler et al. (2017)
Akanthomyces neoaraneogenus	GZU1031Lea [⊤]	Araneae; spider	KX845703	-	KX845697	KX845699	KX845701	Chen et al. (2017)
Akanthomyces	GY11241 [⊤]	Coleoptera	MN093296	-	MN097813	MN097816	MN097812	Chen et al. (2020a)
neocoleopterorum	GY11242	Coleoptera	MN093298	-	MN097815	MN097817	MN097814	Chen et al. (2020a)
Akanthomyces noctuidarum	BCC 36265 [⊤]	Lepidoptera; Noctuidae	MT356072	MT356084	MT477978	MT477994	MT477987	Aini et al. (2020)
	BCC 47498	Lepidoptera; Noctuidae	MT356074	MT356086	MT477980	MT477996	MT477988	Aini et al. (2020)
	BCC 28571	Lepidoptera; Noctuidae	MT356075	MT356087	MT477981	MT478009	MT478006	Aini et al. (2020)
Akanthomyces pissodis	CBS 118231 [™]	Coleoptera; Pissodes strobi	-	KM283799	KM283822	KM283842	KM283864	Chen et al. (2020b)
Akanthomyces pseudonoctuidarum	YFCC 1808943 [™]	Lepidoptera; Noctuidae	OQ509525	OQ509512	OQ506288	0Q511537	OQ511551	This study
	YFCC 1808944	Lepidoptera; Noctuidae	OQ509526	OQ509513	OQ506289	OQ511538	0Q511552	This study
Akanthomyces pyralidarum	BCC 28816 [⊤]	Lepidoptera; Pyralidae	MT356080	MT356091	MT477982	MT478000	MT478007	Aini et al. (2020)
	BCC 32191	Lepidoptera; Pyralidae	MT356081	MT356092	MT477983	MT478001	MT477989	Aini et al. (2020)

Table 1. Specimen information and GenBank accession numbers for sequences used in this study.

	Voucher			GenBan	k accession	numbers		5.6
Species	information	Host/Substrate	ITS	nrLSU	TEF	RPB1	RPB2	Reference
Akanthomyces sabanensis	ANDES-F 1023	Hemiptera; Pulvinaria caballeroramosae	KC633237	-	KC633267	KC875222	-	Kepler et al. (2017)
	ANDES-F 1024	Hemiptera; Pulvinaria caballeroramosae	KC633232	KC875225	KC633266	_	KC633249	Kepler et al. (2017)
Akanthomyces sp.	YFCC 945	Soil	OQ509531	-	OQ506294	OQ511543	OQ511557	This study
Akanthomyces	YFCC 2107937 [™]	Araneae; spider	OQ509527	OQ509514	OQ506290	OQ511539	OQ511553	This study
subaraneicola	YFCC 2107938	Araneae; spider	OQ509528	OQ509515	OQ506291	OQ511540	OQ511554	This study
Akanthomyces sulphureus	TBRC 7248 [⊤]	Araneae; spider	MF140758	MF140722	MF140843	MF140787	MF140812	Mongkolsamrit et al. (2018)
	TBRC 7249	Araneae; spider	MF140757	MF140721	MF140842	MF140786	MF140734	Mongkolsamrit et al. (2018)
Akanthomyces sulphureus	YFCC 1710936	Araneae; spider	OQ509529	OQ509516	OQ506292	OQ511541	OQ511555	This study
Akanthomyces thailandicus	TBRC 7245 [⊤]	Araneae; spider	MF140754	-	MF140839	_	MF140809	Mongkolsamrit et al. (2018)
Akanthomyces	KY11571 [⊤]	Araneae; spider	ON502848	ON502825	ON525447	_	ON525446	Chen et al. (2022)
tiankengensis	KY11572	Araneae; spider	ON502821	ON502827	ON525449	_	ON525448	Chen et al. (2022)
Akanthomyces tortricidarum	BCC 72638 [⊤]	Lepidoptera; Tortricidae	MT356076	MT356088	MT478004	MT477997	MT477992	Aini et al. (2020)
	BCC 41868	Lepidoptera; Tortricidae	MT356077	MT356089	MT477985	MT477998	MT478008	Aini et al. (2020)
Akanthomyces tuberculatus	HUA 186131	Lepidoptera (adult moth)	-	MF416521	MF416466	_	_	Kepler et al. (2017)
Akanthomyces	KACC 44066	Rust	-	KM283784	KM283808	KM283830	KM283850	Park et al. (2016)
uredinophilus	KACC 44082 [⊤]	Rust	-	KM283782	KM283806	KM283828	KM283848	Park et al. (2016)
	KUN 101466	Insect	MG948305	MG948307	MG948315	MG948311	MG948313	Park et al. (2016)
	KUN 101469	Insect	MG948306	MG948308	MG948316	MG948312	MG948314	Park et al. (2016)
Akanthomyces waltergamsii	TBRC 7251	Araneae; spider	MF140747	MF140713	MF140833	MF140781	MF140805	Mongkolsamrit et al. (2018)
	TBRC 7252 [™]	Araneae; spider	MF140748	MF140714	MF140834	MF140782	MF140806	Mongkolsamrit et al. (2018)
Akanthomyces waltergamsii	YFCC 883	Araneae; spider	OQ509530	OQ509517	OQ506293	OQ511542	OQ511556	This study
Akanthomyces zaquensis	HMAS 246915 [™]	Fungi; Ophiocordyceps sinensis	MT789699	MT789697	MT797812	MT797810	-	Wang et al. (2023a)
	HMAS 246917	Fungi; Ophiocordyceps sinensis	MT789698	MT789696	MT797811	MT797809	-	Wang et al. (2023a)
Samsoniella aurantia	TBRC 7271 [™]	Lepidoptera	MF140764	MF140728	MF140846	MF140791	MF140818	Mongkolsamrit et al. (2018)
Samsoniella inthanonensis	TBRC 7915 [™]	Lepidoptera (pupa)	MF140761	MF140725	MF140849	MF140790	MF140815	Mongkolsamrit et al. (2018)

Boldface: data generated in this study. Ex-type materials are marked with "T".

analysis was from jModelTest version 2.1.4 (Darriba et al. 2012). The following models were implemented in the analysis: GTR + I + G for partitions of ITS, nr*L-SU* and *TEF* and GTR + I for partitions of *RPB1* and *RPB2*. The BI analysis was executed on MrBayes v.3.2.7a for five million generations (Ronquist et al. 2012). GTR + FO + G was selected as the optimal model for ML analysis and 1000 rapid bootstrap replicates were performed on the dataset. ML phylogenetic analyses were conducted in RAxML 7.0.3 (Stamatakis et al. 2008). Additional ML analyses were performed using IQ-TREE v. 2.1.3 with ultrafast bootstrapping for the estimation of branch support (Minh et al. 2020). Further, ML analysis (IQ-TREE) was applied to single-locus genealogies for ITS, nr*LSU*, *TEF*, *RPB1* and *RPB2*.

Identification of host arthropods

The host arthropods of *Akanthomyces* spp. were identified on the basis of morphological characteristics and they were further identified utilising molecular analyses according to the mitochondrial cytochrome oxidase I gene (*cox1*) and mitochondrial cytochrome b gene (*cytb*). Genomic DNA was extracted from the head and leg areas of the cadavers of the hosts by utilising the CTAB method (Liu et al. 2001). The *cox1* and *cytb* loci were amplified with the primer pair Hep-cox1F/Hep-cox1R and Hep-cytbF/Hep-cytbR, respectively (Simon et al. 1994). Sequences were analysed with MEGA v.6.06 software (Tamura et al. 2013) and processed by Standard Nucleotide BLAST (GenBank, NCBI nucleotide database) to assess similarity with reported arthropod sequences.

Results

Sequencing and phylogenetic analyses

The five DNA loci (ITS, nrLSU, TEF, RPB1, RPB2) were readily amplified and sequenced and there was a fairly high success rate in this study. Preliminary phylogenetic analyses, based on the combined five-gene sequences from 116 fungal taxa Cordycipitaceae and Trichoderma Pers., confirmed the presence and positions of Akanthomyces and related genera within Cordycipitaceae. The concatenated five-gene dataset consisted of 4,453 bp (ITS = 639 bp, nrLSU = 921 bp, TEF = 1,044 bp, RPB1 = 758 bp and RPB2 = 1,091 bp). Ten well-supported clades were recognized, which accommodate species of the genera Akanthomyces, Ascopolyporus Möller, Beauveria Vuill., Blackwellomyces Spatafora & Luangsa-ard, Cordyceps, Gibellula Cavara, Hevansia, Samsoniella, Simplicillium W. Gams & Zare and Trichoderma (Suppl. material 1: fig. S1). The phylogenetic analyses also revealed the species diversity of the genus Akanthomyces. This suggested that the group should be genetically composed of at least 30 species (Suppl. material 1: fig. S1). The further phylogenetic analyses, based on combined partial ITS+nrLSU+TEF+RPB1+RPB2 sequences consisting of 56 fungal taxa (Table 1), resolved the majority of the Akanthomyces lineages into separate terminal branches (Fig. 1). The dataset consisted of 4,401 bp of sequence data (ITS = 619 bp, nrLSU = 896 bp, TEF = 1,022 bp, RPB1 = 731 bp and RPB2 = 1,133 bp). Samsoniella aurantia Mongkols., Noisrip., Thanakitp., Spatafora & Luangsa-ard (strain TBRC 7271) and S. inthanonensis Mongkols., Noisrip., Thanakitp., Spatafora & Luangsa-ard (strain TBRC 7915) within Cordycipitaceae were used as the outgroup sequences for this dataset. This revealed a similar tree and cluster topology, as shown in Suppl. material 1: fig. S1. Amongst the hosts of Akanthomyces, Araneae (spider) and Lepidoptera (adult moth) are the two major orders. Most of the spider pathogens form a monophyletic clade, separated from the pathogens of moths, themselves forming also an apparent monophyletic clade (Fig. 1). The phylogenetic analyses also suggested the existence of distinct species in the spider pathogens and adult moth entomopathogens clade that we proposed as new species: A. kunmingensis and A. subaraneicola, which were found in the spider pathogens clade; and A. laosensis and A. pseudonoctuidarum, which were found in the adult moth entomopathogens clade (Fig. 1).



0.01

Figure 1. Phylogenetic tree of *Akanthomyces* species, based on combined partial ITS + nr*LSU* + *TEF* + *RPB1* + *RPB2* sequences. Numbers at the branches indicate support values (BI-PP/IQ-TREE-BS/ RAxML-BS) above 0.7/70%/70%. Ex-type materials are marked with "T". Isolates in bold type are those analysed in this study.

Despite differing topologies between individual loci (ITS, nrLSU, TEF, RPB1 and RPB2), the newly-proposed species usually stood out as distinct clades to other known species. Some novel species always recovered the sister relation-

ship to a particular known species for all loci. For example, the newly-discovered species *A. kunmingensis* had a close genetic relationship with *A. waltergamsii*. They were regarded as different species with strong support from ITS, nrLSU, TEF, RPB1 and RPB2 (Suppl. material 1: figs S2–S6). The new species *A. subaraneicola* was sisters to *A. araneicola* and this relationship received significant bootstrap support from ITS, TEF, RPB1 and RPB2 (Suppl. material 1: figs S2, S4–S6). Meanwhile, *A. laosensis* was inferred to form a sister clade to either *A. pyralidarum* (ITS, RPB1 and RPB2) or *A. tortricidarum* (nrLSU and TEF). Similarly, despite the differing position of *A. pseudonoctuidarum* between different markers, it always formed a clade that could be distinguished from its closely-related species, *A. noctuidarum* and *A. tortricidarum*.

Morphological features

The morphological characteristics of the five species, as well as photomicrographs of morphological structures, are shown in Figs 2–6. The detailed fungal morphological descriptions are supplied in the Taxonomy section.

Taxonomy

Akanthomyces kunmingensis Hong Yu bis, Y. Wang & Z.Q. Wang, sp. nov. MycoBank No: 848307

Fig. 2

Etymology. Named after the location, Kunming City, where the species was collected.

Type. CHINA. Yunnan Province, Kunming City, Wild Duck Lake Forest Park (25.2181°N, 102.8503°E, 2100 m above sea level), on a spider on a dead stem, 14 August 2018, collected by Yao Wang (holotype: YHH 16988; ex-type living culture: YFCC 1808940).

Description. *Sexual morph*: Undetermined. *Asexual morph*: Synnemata arising from spider body, cream to light yellow, erect, irregularly branched, producing a mass of conidia at the upper apex, powdery and floccose. Colonies on PDA reaching 15–20 mm in diameter after 14 days at 25 °C, circular, white and fluffy mycelium, middle bulge, reverse pale yellow to light brown. Hyphae smooth-walled, branched, septate, hyaline, $0.5-2.8 \mu m$ wide. Conidiophores smooth-walled, cylindrical, solitary, sometimes verticillate, $4.3-9.5 \times 1.2-2.0 \mu m$ (n = 30). Phialides consisting of a cylindrical, somewhat inflated base, verticillate on conidiophores, usually in whorls of 4–5 or solitary on hyphae, $6.2-29.4 \times 1.1-2.5 \mu m$ (n = 30). Conidia smooth and hyaline, ellipsoidal to long oval, one-celled, $1.9-3.5 \times 1.1-1.8 \mu m$ (n = 50), often in chains. Size and shape of phialides and conidia similar in culture and on natural substratum.

Host. Spider (Araneae).

Habit. On spiders on dead stems.

Distribution. Kunming City, Yunnan Province, China.

Other material examined. CHINA. Yunnan Province, Kunming City, Songming County, Dashao Village (25.3924°N, 102.5589°E, 2700 m above sea level), on a spider on a dead stem, 12 August 2017, Yao Wang (YHH 2301006; living culture: YFCC 1708939).



Figure 2. Morphology of *Akanthomyces kunmingensis* **A** the type specimen (YHH 16988) **B** culture character on PDA medium **C** conidiogenous structures on the host **D**–**H** conidiophores, conidiogenous cells and conidia I conidia. Scale bars: 3 mm (**A**); 10 mm (**B**); 10μ (**C**, **E**, **F**); 5μ (**D**); 2μ (**G**–**I**).

Commentary. In regard to phylogenetic relationships, *Akanthomyces kunmingensis* forms a distinct lineage in the genus *Akanthomyces* with high credible support (1/100%/100%) and it is closely related to *A. sulphureus* and *A. waltergamsii* (Fig. 1). Morphologically, *A. kunmingensis* is so similar to *A. waltergamsii* that it was once referred to as *A. waltergamsii* by Wang et al. (2020); however, a morphological observation revealed a significant difference of conidia shapes between *A. kunmingensis* and *A. waltergamsii*. *Akanthomyces kunmingensis* usually produces a variety of shapes of conidia (viz. spherical, ellipsoidal to long oval or fusiform), while *A. waltergamsii* produces only ellipsoidal and fusiform conidia. Moreover, *A. kunmingensis* can be distinguished from *A. sulphureus* and *A. waltergamsii* by its longer phialides (6.2–29.4 µm) and smaller conidia (1.9–3.5 × 1.1–1.8 µm) (Table 3).

Akanthomyces laosensis Hong Yu bis & Y. Wang, sp. nov.

MycoBank No: 848308 Fig. 3

Etymology. Named after the location, Laos, where the species was collected.

Type. LAOS. Oudomxay Province, Muang Xay County, Nagang Village (20.7143°N, 102.0957°E, 698 m above sea level), on the adult of Noctuidae on the underside of a dicotyledonous leaf, 5 October 2019, collected by Yao Wang (holotype: YHH 2301008; ex-holotype living culture: YFCC 1910941).

Description. Sexual morph: Undetermined. Asexual morph: Specimens examined in this study can be found on the underside of dicotyledonous leaves.



Figure 3. Morphology of Akanthomyces laosensis A, B fungus on adult moth C long synnemata D culture character on PDA medium E-H conidiophores, conidiogenous cells and conidia I conidia from long synnemata. Scale bars: 10 mm (A, B); 5 mm (C); 20 mm (D); 20 µm (E-G); 10 µm (H); 5 µm (I).

Synnemata arose at the head and in the middle of the host body, white, up to 15.6 mm long and 0.6–1.3 mm wide, rarely branched, feathery to clavate with acute or blunt ends. Colonies on PDA moderately fast-growing at 25 °C, reaching 23–26 mm in diameter in 14 days, circular, flat, white in the middle with a light yellow edge, reverse light yellow. Hyphae smooth-walled, branched, septate, hyaline, 0.8–3.5 μ m wide. Conidiogenous cells monophialidic, produced along the synnemata or solitary on hyphae in culture. Phialides smooth-walled, hyaline, cylindrical, 11.5–30.0 × 2.0–4.2 μ m (n = 30). Conidia smooth and hyaline, cylindrical or long oval, one-celled, 4.1–9.8 × 2.3–4.2 μ m (n = 30). Size and shape of phialides and conidia similar in culture and on natural substratum.

Host. Adult moth (Noctuidae, Lepidoptera).

Habit. On the adults of Noctuidae sp. on the underside of leaves of plants.

Distribution. Muang Xay County, Oudomxay Province, Laos.

Other material examined. LAOS. Oudomxay Province, Muang Xay County, Nam Kit Park (20.6651°N, 102.0007°E, 695 m above sea level), on an adult moth on the underside of a leaf, 1 October 2019, Yao Wang (YHH 2301000; living culture: YFCC 1910942).

Commentary. Phylogenetically, *Akanthomyces laosensis* forms a distinct lineage and is closely related to *A. pyralidarum* with strong statistical support (1/99%/89%) (Fig. 1). Morphologically, *A. laosensis* is distinctly different from *A. pyralidarum* because of its longer synnemata (up to 15.6 mm). Furthermore, *A. laosensis* was determined to occur on an adult of Noctuidae sp., while *A. pyralidarum* was located on an adult of Pyralidae sp. In fact, the species is easily distinguished from other known species in the genus of *Akanthomyces* by its longer phialides (11.5–30.0 µm) and larger conidia (4.1–9.8 × 2.3–4.2 µm) (Table 3).

Akanthomyces pseudonoctuidarum Hong Yu bis & Y. Wang, sp. nov.

MycoBank No: 848309 Fig. 4

Etymology. Referring to macromorphological resemblance of *A. noctuidarum*, but *A. pseudonoctuidarum* is phylogenetically distinct.

Type. THAILAND. Chiang Mai Province, Chiang Mai City, Sansai District, Maejo Farm (18.9177°N, 99.0520°E, 317 m above sea level), on the adult of Noctuidae on the underside of a dicotyledonous leaf, 22 August 2018, collected by Hong Yu (holotype: YHH 2301010; ex-type living culture: YFCC 1808943).

Description. Sexual morph: Undetermined. Asexual morph: Synnemata arising from moth body, cream to light yellow, erect, simple, cylindrical to clavate, $800-2000 \times 120-350 \mu m$. Conidia and reproductive structures on natural substratum not observed. Colonies on PDA moderately fast-growing at 25 °C, reaching a diameter of 25–28 mm within 14 days, circular, flat to raised, white and fluffy mycelium, reverse cream to pale yellow. Hypha smooth-walled, hyaline, septate, $1.0-2.9 \mu m$ wide. Conidiophores smooth-walled, cylindrical, solitary, $6.5-13.8 \times 1.8-3.6 \mu m$ (n = 30). Conidiogenous cells monophialidic or polyphialidic. Phialides verticillate, usually in whorls of 2–3 or solitary on hyphae, cylindrical with papillate end, hyaline, $6.8-26.0 \times 2.1-3.6 \mu m$ (n = 30).



Figure 4. Morphology of *Akanthomyces pseudonoctuidarum* **A** adult moth infected by *A. pseudonoctuidarum* **B, C** culture character on PDA medium **D–H** conidiophores, conidiogenous cells and conidia I conidia. Scale bars: 2 mm (**A**); 20 mm (**B, C**); 10 µm (**D–I**).

Conidia smooth and hyaline, ellipsoidal to long oval, one-celled, $2.6-6.4 \times 1.5-2.2 \ \mu m$ (n = 30).

Host. Adult moth (Noctuidae, Lepidoptera).

Habit. On the adults of Noctuidae sp. on the underside of leaves of plants. **Distribution.** Chiang Mai City, Chiang Mai Province, Thailand.

Other material examined. THAILAND, Chiang Mai Province, Chiang Mai City, Mae Rim District, Queen Sirikit Botanic Garden (18.8990°N, 98.8605°E, 536 m above sea level), on an adult of Noctuidae, 26 August 2018, collected by Yao Wang (YHH 2301011; living culture: YFCC 1808944).

Commentary. Akanthomyces pseudonoctuidarum is similar to its phylogenetically closely-related species *A. noctuidarum* in macromorphology. They have the same hosts (the adults of Noctuidae sp.) and *Isaria*-like asexual conidiogenous structures, producing cream or light yellow synnemata. However, *A. pseudonoctuidarum* is easily recognised by its larger synnemata (800– 2000 × 120–350 µm), longer phialides (6.8–26.0 µm) and larger conidia (2.6– 6.4 × 1.5–2.2 µm) (Table 3). It was easily distinguished phylogenetically from *A. noctuidarum* (Fig. 1; 1/97%/85%). Both the morphological study and phylogenetic analyses of combined ITS, nr*LSU*, *TEF*, *RPB1* and *RPB2* sequence data supported that this fungus is a distinct species in the genus *Akanthomyces*.

Akanthomyces subaraneicola Hong Yu bis, Y. Wang & Z.Q. Wang, sp. nov.

MycoBank No: 848310 Fig. 5

Etymology. "Subaraneicola" refers to morphologically resembling *A. araneicola*, but phylogenetically distinct.

Type. CHINA. Hunan Province, Huaihua City, Zhongpo National Forest Park (27.5724°N, 109.9664°E, 615 m above sea level), on a spider emerging from leaf litter on the forest floor, 10 July 2021, collected by Yao Wang (holotype: YHH 2301004; ex-type living culture: YFCC 2107937).

Description. *Sexual morph*: Undetermined. *Asexual morph*: Mycosed hosts covered by white to pale yellow mycelia, producing numerous powdery conidia, synnemata not observed. Colonies on PDA reaching 24–28 mm in diameter within 14 days at 25 °C, circular, white and fluffy mycelium in the centre, cottony with a raised mycelial density at the outer ring, reverse white to pale yellow. Hyphae smooth-walled, branched, septate, hyaline, $1.6-3.2 \mu m$ wide. Conidio-phores smooth-walled, cylindrical, solitary, sometimes verticillate, $6.5-12.3 \times 1.6-3.5 \mu m$ (n = 30). Conidiogenous cells monophialidic or polyphialidic. Phialides consisting of a cylindrical, somewhat inflated base, verticillate on conidio-phores, usually in whorls of 2–5, or solitary on hyphae, $12.1-38.2 \times 1.3-3.2 \mu m$ (n = 30). Conidia smooth and hyaline, ellipsoidal to long oval, one-celled, $3.0-5.4 \times 1.8-3.4 \mu m$ (n = 50), often in chains. Size and shape of phialides and conidia similar in culture and on natural substratum.

Host. Spider (Araneae).

Habit. On spiders on dead stems or emerging from leaf litter on the forest floor. **Distribution.** Hunan and Yunnan Province, China.

Other material examined. CHINA, Yunnan Province, Kunming City, Wild Duck Lake Forest Park (25.1244°N, 102.8716°E, 1900 m above sea level), on a spider on a dead stem, 28 July 2021, Yao Wang (YHH 2301005; living culture: YFCC 2107938).

Commentary. Morphologically, *Akanthomyces subaraneicola* resembles the phylogenetic sister species *A. araneicola*. They were found to be parasitic on spiders (Araneae) and they are easily recognised by having white to pale yellow mycelia covering the hosts with a mass of conidia; however, our morphological observation revealed a significant difference in the shape and size of conidia between *A. subaraneicola* and *A. araneicola*. *Akanthomyces subaraneicola* usually produces large ellipsoidal to long oval conidia ($3.0-5.4 \times 1.8-3.4 \mu m$), while *A. araneicola* produces small fusiform conidia ($2.5-5.0 \times 1.3-1.9 \mu m$) (Table 3). In addition, molecular phylogenetic analyses indicated that they are distinct species (Fig. 1; 1/100%/100%).



Figure 5. Morphology of *Akanthomyces subaraneicola* **A**, **B** fungus on spider **C** culture character on PDA medium **D**–**H** conidiophores, conidiogenous cells and conidia I conidia. Scale bars: 10 mm (**A**); 5 mm (**B**); 20 mm (**C**); 30 μm (**D**); 20 μm (**E**); 10 μm (**F**–**I**).

Akanthomyces araneogenus Z.Q. Liang, W.H. Chen & Y.F. Han, Phytotaxa 379(1): 69 (2018)

MycoBank No: 816114 Fig. 6

Akanthomyces tiankengensis W.H. Chen, Y.F. Han, J.D. Liang & Z.Q. Liang, Microbiology Spectrum 10(5): e01975-22, 6 (2022). Synonym.

Description. *Sexual morph*: Undetermined. *Asexual morph*: Mycosed hosts covered with white to pale yellow mycelia, occasionally several synnemata aris-

ing from all of the parts of the host. Colonies on PDA moderately fast-growing at 25 °C, reaching a diameter of 25–36 mm in 14 days at 25 °C, circular, middle bulge, white to yellowish, reverse yellowish. Hyphae smooth-walled, branched, septate, hyaline, 0.5–2.9 μ m wide. Conidiophores smooth-walled, cylindrical, solitary, 10.6–22.4 × 1.3–2.6 μ m (n = 30). Phialides consisting of a cylindrical, somewhat inflated base, verticillate on conidiophores, usually in whorls of 2–3 or solitary on hyphae, 8.1–17.8 × 1.1–3.6 μ m (n = 30). Conidia smooth and hyaline, one-celled, globose, 1.6–2.4 μ m in diameter or ellipsoidal to fusiform, 2.2–4.1 × 1.1–2.3 μ m (n = 50), often in chains. Size and shape of phialides and conidia similar in culture and on natural substratum.

Host. Spider (Araneae).

Habit. On the spiders on dead stems or emerging from leaf litter.

Distribution. Guizhou and Yunnan Province, China; Chiang Mai Province, Thailand; Nghe An Province, Vietnam.



Figure 6. Morphology of Akanthomyces araneogenus A fungus on spider B conidiogenous structures on the host C,D culture character on PDA medium E-H conidiophores, conidiogenous cells and conidia I conidia. Scale bars: 5 mm (A); 30 µm (B); 30 mm (C, D); 10 µm (E); 5 µm (F–I).

Material examined. THAILAND, Chiang Mai Province, Chiang Mai City, Queen Sirikit Botanic Garden (18.8990°N, 98.8604°E, 547 m above sea level), on a spider on a dead stem, 20 November 2018, Yao Wang (YHH 2301001; living culture: YFCC 1811934). VIETNAM, Nghe An Province, Pu Mat National Park (18.9292°N, 104.5889°E, 621 m above sea level), on spiders emerging from leaf litter on the forest floor, 28 April 2017, Yao Wang (YHH 2301007, YHH 2301012; living culture: YFCC 1704946, YFCC 1704947). CHINA, Yunnan Province, Dai Autonomous Prefecture of Xishuangbanna, Mengla County (21.1817°N, 101.7252°E, 875 m above sea level), on a spider on a dead stem, 12 June 2022, Zhi-Qin Wang (YHH 2301002; living culture: YFCC 2206935).

Commentary. In our phylogenetic analyses, *Akanthomyces araneogenus* extype strain (GZUIF DX2) and *A. tiankengensis* ex-type isolate (KY11571) and our two samples isolated from the spiders formed a well-supported clade (Fig. 1). From a phylogenetic point of view, *A. tiankengensis* could not be distinguished from *A. araneogenus*, being inside the clade of the latter. Previous morphological observations revealed several differences in the characteristics between *A. araneogenus* and *A. tiankengensis* (Chen et al. 2018; Chen et al. 2022); however, our samples from different regions showed diversity of morphology in this study. The colony colour and the shape and size of the phialides and conidia of *A. araneogenus* and *A. tiankengensis*, amongst other morphological features, have been noted in our samples. There is reason to believe that distinguishing the two species is difficult because of the extensive overlap in morphological characteristics. Thus, we propose that *A. tiankengensis* is a synonym of *A. araneogenus*.

Discussion

In this study, *Akanthomyces* comprised at least 36 species with a cosmopolitan distribution (Table 2). A collection of 31 isolates of unknown identity were shown to represent four known species, four new species and an undetermined species of *Akanthomyces*. The phylogenetic positions of the four known species were evaluated, based on phylogenetic inferences according to five loci, namely, ITS, nr*LSU, TEF, RPB1* and *RPB2*, including *A. araneogenus* from China, Thailand and Vietnam, *A. dipterigenus* and *A. waltergamsii* from China and *A. sulphureus* from Vietnam (see Table 2 and Fig. 1). The four new species, given the names *A. kunmingensis* and *A. subaraneicola* from China, *A. laosensis* from Laos and *A. pseudonoctuidarum* from Thailand, were recognised according to morphological characteristics and molecular data. The isolate YFCC 945 from China represented an unknown species in the genus *Akanthomyces*. Unfortunately, the isolate did not produce conidia or reproductive structures when grown on PDA and other media and they were, thus, tentatively treated as an undetermined species of *Akanthomyces*, pending further investigation.

The highest species diversity of *Akanthomyces* occurred in subtropical and tropical regions, especially in China and Southeast Asia (see Table 2). Based on our update, there are at least 17 *Akanthomyces* species in China and Yunnan Province has the most. There is also high species diversity of *Akanthomyces* in Southeast Asia, where more than 11 species have been recorded (Table 2). Thailand, Vietnam and Laos are located in tropical regions with extremely rich biodiversity in Southeast Asia. The forests exhibit a significant variety of plant and animal life attributed to the tropical monsoon climate, characterised by high temperatures

and rainfall (Lao et al. 2021). These have created a favourable environment for the development of arthropod-pathogenic fungi, including *Akanthomyces* spp.

Akanthomyces species inhabit diverse hosts/substrates that range from eight orders of Arthropoda, namely, Acari, Araneae, Coleoptera, Hemiptera, Hymenoptera, Lepidoptera, Orthoptera and Thysanoptera, to plants, other fungi, peat, water and rusts (see Table 2). Amongst the hosts of *Akanthomyces*, Araneae and Lepidoptera are the two major orders. Our study also found that the majority of *Akanthomyces* species are spider pathogens or adult moth entomopathogens, with the exception of a few other entomopathogens and generalists that have a remarkably broad host/substrate range (Table 2 and Fig. 1). In this study, we identified an extension of the host/substrate range to also include soil, as shown in Fig. 1. The family Cordycipitaceae has been shown to evolve from an ancestor which is ecologically versatile and most probably inhabit the soil/environment and diversified into groups of entomopathogens and mycoparasites (Sung et al. 2007; Kepler et al. 2017; Wang et al. 2020; Zhou et al. 2022). *Akanthomyces* have been principally shown to be arthropod-pathogenic fungi in this study. The fact that *Akanthomyces* can be found in soil might suggest some kind of convergence/reversion.

Due to the difficulty of isolation and the limitation of cultivation conditions, studies on the development and application of Akanthomyces species are still currently limited. As generalists that have a remarkably broad host/substrate range, A. gracilis and A. muscarius have a high potential for interspecific transmission and biological control of pest insects (Samson and Evans 1974; Zare and Gams 2001; Kuchár et al. 2019; Nicoletti and Becchimanzi 2020). Akanthomyces lecanii is an effective mycoparasite of several rust fungi, green mould and fungi causing root rot diseases (Pythium ultimum), as well as of several powdery mildew pathogens and it is receiving increasing attention as a versatile biocontrol agent of a number of plant pathogens (Benhamou and Brodeur 2001). The members of the genus Akanthomyces contain species ranging from specialists with very narrow host ranges to generalists that attack a wide range of arthropods and they might be used as an ideal model system for research on fungal arthropod pathology and fungal-pathogen speciation and host adaptation (Hu et al. 2014). Coleopterans, lepidopterans and spiders are the major host groups of arthropod-pathogenic fungi within Hypocreales (Shrestha et al. 2019). The findings indicate that the majority of the hosts of Akanthomyces are distributed in lepidopterans and spiders, with a few in coleopterans (see Table 2). These arthropod-pathogenic fungi with special nutritional preferences are more likely to produce numerous distinctive bioactive compounds. It is hoped that this study will generate continued interest amongst mycologists, arachnologists and related experts and researchers to use such fungal resources through in vitro growth and extraction of useful bio-active secondary metabolites (extrolites).

Fungal species diversity and their host/substrate associations are important aspects of fungal ecology. A strong taxonomic basis that is dependent on advances in nucleic acid sequence technology is one of the main fundamental needs in fungal ecology (Zhang et al. 2021) and is even crucial to studies on species diversity and their host-substrate associations. However, it is regrettable that a growing number of researchers have relied heavily on molecular biology techniques to the complete exclusion of fungal isolation and characterisation utilising classical methods (Walker et al. 2019; Zhang et al. 2021). Although fungal research has entered the molecular era, phenotypic and culture-based

Species	Host/Substrate	Know distribution	References
lleatus	Adult moth (Noctuidae; Sphingidae)	USA (Connecticut; Washington; Ontario); Brazil (Salvador); Amazon countries	Mains (1950); Sanjuan et al. (2014)
gustispora	Coleopterous larva	USA (Nashville)	Mains (1950)
nearum	Spider (Araneae)	USA (North Carolina; Maine); Ceylon; Netherlands; Ghana (Begoro); China	Mains (1950); Samson and Evans (1974); Hsieh et al. (1997); Zare and Gams (2001)
neicola	Spider (Araneae)	China (Guizhou)	Chen et al. (2019)
aneogenus	Spider (Araneae)	China (Guizhou; Yunnan); Thailand (Chiang Mai); Vietnam (Nghe An)	Chen et al. (2018); This study
neosus	Spider (Araneae)	China (Guizhou)	Chen et al. (2022)
enuatus	Cydia pomonella (Lepidoptera, Tortricidae); leaf litter of Acer saccharum; Symplocarpus foetidus (plants); Astrocaryum sciophilum (plants)	Poland; USA; Canada; French	Zare and Gams (2001); Ellsworth et al. (2013); Barthélemy et al. (2019)
vata	Hapithus agitator (Orthoptera, Gryllidae)	USA (Florida)	Mains (1950)
ccidioperitheciatus	Spider (Araneae)	Japan	Kepler et al. (2017); Johnson et al. (2009)
pterigenus	Hemiptera: Icerya purchasi (Coccidae); Myzus persicae (Aphididae); Macrosiphoniella sanborni (Aphididae); Citrus aphid (Aphididae); soil	UK; Sri Lanka; Peru; China (Yunnan)	Kepler et al. (2017); Zare and Gams (2001); This study
agilis	Orthopterous larva	Trinidad; Guiana; Brazil	Mains (1950); Petch (1937)
acilis	Hymenoptera, Formicidae (Paltothyreus tarsatus; Platythyrea conradti, Polyrhachis militaris; Polyrhachis monista; Polyrhachis decemdentata; Camponotus brutus; Oecophylla Ionginoda; Crematogaster bequarti; Crematogaster clariventris; Macromischoides inermis; Macromischoides aculeatus; Dorylus sp.); Coleoptera (beetle larvae, beetle imago); Lepidoptera larva; Hemiptera (Pyrrhocoridae; Cercopidae)	Ghana (Begoro); China (Guizhou)	Samson and Evans (1974); Liang et al. (2013)
hnsonii	Leaf and stem (Arctium sp., Begonia sp., Coffea sp., Dianthus sp., Ipomoea sp., Kalanchoe sp., Lycopersicon sp., Peperomia sp., and Sargassum sp.); often associated with species of Botryosporium	Ghana; Indonesia; Australia (Great Barrier Reef); UK; USA; Canada	Vincent et al. (1988)
nyawimiae	Spider (Araneae)	Thailand (Phetchabun; Chanthaburi)	Mongkolsamrit et al. (2018)
nmingensis	Spider (Araneae)	China (Yunnan)	This study
osensis	Adult moth (Lepidoptera, Noctuidae)	Laos (Oudomxay)	This study
canii	Hemiptera, Coccidae: Pulvinaria floccifera; Coccus viridis; scale insect. Tetranychus urticae (Acari: Tetranychidae); Pistacia vera (plants); Ammophila arenaria (plants); Dactylis glomerata (plants); Deschampsia flexuosa (plants); Elymus farctus (plants); Laretia acaulis (plants); Pinus sylvestris (plants); Shorea thumbuggaia (plants); Taxus baccata (plants)	W. Indies; Dominican Republic; Peru; Jamaica; USA; Sri Lanka; Indonesia; Turkey; China; Iran; Spain; Finland; Chile; Italy; Poland; India	Kepler et al. (2017); Zare and Gams (2001); Dash et al. (2018); Dolatabad et al. (2017); Nicoletti and Becchimanzi al. (2017); Nicoletti and U
oidopterorum	Pupa of Lepidoptera	China (Guizhou)	Chen et al. (2020b)
Iscarius	<i>Trialeurodes vaporariorum</i> (Hemiptera, Aleyrodidae); <i>Brachycaudus helichrysi</i> (Hemiptera, Aphididae); <i>Cecidophyopsis ribis</i> (Acari, Eriophyidae); <i>Cossus cossus</i> (Lepidoptera, Cossidae); <i>Zyginidia pullula</i> (Hemiptera, Cicadellidae); <i>Thrips tabaci</i> (Thysanoptera, Thripidae); peat, contaminated pesticide solution; <i>Pteridium aquilinum</i> (Pteridophyta); leaves of <i>Nypa fruticans</i> (Plants), <i>Hemileia vastatrix</i> (Fungi); water from domestic supply, laboratory glyphosate solution; <i>Acer campestre</i> (plants); <i>Prurus</i> <i>nobilis</i> (plants), <i>Myrtus communis</i> (plants); cabbage plants cerasus (plants); cabbage plants	UK; Italy; New Caledonia; Thailand; New Zealand	Kepler et al. (2017); Zare and Gams (2001); Nicoletti and Becchimanzi (2020); Vinit et al. (2018); Aghdam and Fotouhifar 2017; Kuchár et al. (2019)
oaraneogenus	Spider (Araneae)	China (Guizhou)	Chen et al. (2017); Mains (1949)

Species	Host/Substrate	Know distribution	References
Akanthomyces neocoleopterorum	Ladybug (Coleoptera)	China (Guizhou)	Chen et al. (2020a)
Akanthomyces noctuidarum	Adult moth (Lepidoptera, Noctuidae)	Thailand (Narathiwat; Nakhon Ratchasima; Kamphaeng Phet)	Aini et al. (2020)
Akanthomyces pissodis	Adult of Pissodes strobi (Coleoptera, Curculionidae)	Canada	Cope and Leal (2005)
Akanthomyces pseudonoctuidarum	Adult moth (Lepidoptera, Noctuidae)	Thailand (Chiang Mai)	This study
Akanthomyces pyralidarum	Adult moth (Lepidoptera, Pyralidae)	Thailand (Kanchanaburi; Chiang Mai; Phetchabun)	Aini et al. (2020)
Akanthomyces ryukyuenis	Spider (Araneae)	Japan	Kobayasi and Shimizu (1982)
Akanthomyces sabanensis	Pulvinaria caballeroramosae (Hemiptera, Coccidae)	Colombia	Chiriví-Salomón et al. (2015)
Akanthomyces subaraneicola	Spider (Araneae)	China (Hunan; Yunnan)	This study
Akanthomyces sulphureus	Spider (Araneae)	Thailand (Nakhon Ratchasima; Surat Thani); Vietnam (Nghe An)	Mongkolsamrit et al. (2018); This study
Akanthomyces thailandicus	Spider (Araneae)	Thailand (Chiang Mai)	Mongkolsamrit et al. (2018)
Akanthomyces tiankengensis	Spider (Araneae)	China (Guizhou)	Chen et al. (2022)
Akanthomyces tortricidarum	Adult moth (Lepidoptera, Tortricidae)	Thailand (Nakhon Ratchasima; Kamphaeng Phet)	Aini et al. (2020)
Akanthomyces tuberculatus (= A. pistillariaeformis)	Adult moth (Lepidoptera); Hymenoptera, Formicidae; Hemiptera, Pyrrhocoridae	China (Zhejiang; Yunnan); Begoro; Trinidad	Mains (1950); Samson and Evans (1974); Liang et al. (2007)
Akanthomyces uredinophilus	Rust; decayed insect	Korea (Gangwon; North Chungcheong); China (Yunnan)	Park et al. (2016); Wei et al. (2018)
Akanthomyces waltergamsii	Spider (Araneae)	Thailand (Saraburi; Naknon Ratchasima); China (Yunnan)	Mongkolsamrit et al. (2018); This study
Akanthomyces zaquensis	The stroma and the sclerotium of Ophiocordyceps sinensis (Fungi)	China (Qinghai)	Wang et al. (2023a)

	References	Mains (1950)	Mains (1950)	Chen et al. (2019)	Chen et al. (2018)	Chen et al. (2022)	Mains (1950)	Zare and Gams (2001); Kepler et al. (2017)	Mains (1950)	Zare and Gams (2001); Kepler et al. (2017)	Mains (1950)	Samson and Evans (1974)
	Conidia (µm)	Broadly ellipsoidal or obovoid often acute at the lower end, 3–6 × 2–3	Narrowly obclavate often acute at the lower end, narrowing upwards, rounded or obtuse at the upper end, 8–14 × 1.5–3	Mostly fusiform, 2.5-5.0 × 1.3-1.9	Globose, 1.3–2.4 in diam, or ellipsoidal, 2.1–3.3 × 1.1–1.6	Fusiform, 3.1–5.0 × 1.0–1.8	Narrowly clavate, 4.5–6 × 1.2–1.4	Cylindrical with attenuate base, occasionally 2-celled, 4.5-6.5 × 1.5-2.0	Ellipsoidal to oblong, 4.5–8.5 × 2.1–2.5	Ellipsoidal to oblong- oval, 5.0–10.5 × 1.5–2.5	Subcylindrical, somewhat narrowed and rounded at the ends, 6.5–9 × 1.5	Ellipsoidal to fusiform, 2.5-3 × 1-1.6
	Phialides (µm)	Subcylindrical or narrowly ellipsoidal, 6–16 × 2.5–4, narrowing above to an acute apex, terminated by a short sterigma up to 4 long	Obovoid or ellipsoidal 6–12 × 4–8, rounded above and abruptly narrowing into a short sterigma, asperulate	Cylindrical, somewhat inflated base, 8.1–16.9 × 1.3–1.9, tapering to a thin neck	Cylindrical, somewhat inflated base, 4.3–17.3 × 0.9–3.1, tapering to a thin neck	Solitary or in groups of two, 16.9– 18.1 × 1.3–1.9 with a cylindrical basal portion and tapered into a short, distinct neck	Oblong or narrowly ellipsoidal, 6–14 × 3–4, narrowing above into an acute apex terminated by a short sterigma	9-15.5×1-2	Subcylindrical, 17.1–21.4 × 2.8–4.3, narrowing above to acute apices, terminated by short sterigmata	$20-40 \times 1.2-2.7$, tapering towards the apex	Subcylindrical to narrowly clavate, 7-10 × 2.5-3, verrucose in the upper portions	Cylindrical basal part tapering to a slender neck, 7–10 × 1.5–2.5
	Conidiophores (µm)			Mononematous, with single phialide or whorls of two to six phialides or <i>Penicillium</i> - like from hyphae directly	Mononematous or synnematous, 21.6–48 × 1.2–2.2, <i>Penicillium</i> -like from hyphae directly	Erect conidiophores usually arose from the aerial hyphae						
cies.	Synnemata (mm)	Arising from various parts of the insect, terete, narrowing upwards, 1–8 × 0.1–0.5, yellowish	Arising from all parts of the host, cylindrical to clavate, 0.8–10 × 0.1– 0.2, simple or occasionally slightly branched, brown	Synnemata not observed	Occasionally several white synnemata arise from all parts of the host	Synnemata not observed	Arising from the body and head of the host, simple or branched, 8–13 × 0.2–0.6, flesh coloured		Numerous, arising from various parts of the host, light brown, clavate, 0.5-2.0 × 0.06-0.25		Numerous arising from all parts of the host, clavate, 0.7–1.5 x 0.03–0.09	Arising from the natural body openings and intersegmental and appendage joints, usually white to yellow-brown, cylindrical, 0.7–3 × 0.1–0.5
omyces spec	Part-spores (µm)											
on of <i>Akanth</i>	Asci (µm)											
ological comparise	Perithecia (µm)											
Table 3. Morphc	Species	Akanthormyces aculeata	Akanthomyces aranearum	Akanthomyces araneicola	Akanthomyces araneogenus	Akanthomyces araneosus	Akanthormyces angustispora	Akanthomyces attenuatus	Akanthormyces clavata	Akanthomyces dipterigenus	Akanthomyces fragilis	Akanthormyces gracilis

Species	Perithecia (µm)	Asci (µm)	Part-spores (µm)	Synnemata (mm)	Conidiophores (µm)	Phialides (µm)	Conidia (µm)	References
i				Gregarious, white, 0.4–4 tall, with a stipe 0.025–0.1 wide, subulate to cylindrical	Unbranched or with metulae arising at right angles to the stipe hyphae, $4-6 \times 2-3$	10–20 long, ellipsoidal to cylindrical body 2.5–4 wide, tapering into a narrow neck $3-5$ x $1-1.5$	Broadly fusoid with more or less truncate poles with minute frills, $3-4 \times (-)1.5-2$	Vincent et al. (1988)
myces miae				Up to 1.5 long, up to 0.4 wide, covered by dense white to cream mycelia	Erect, verticillate with phialides in whorls of two to five	$(8-)9-12(-15) \times 2-3, with cylindrical basal portion, tapering into a long neck, (2-)3-5.5(-7) \times 1-1.5$	Cylindrical to ellipsoidal, (2-)2.5-3.5(-5) × (1.5-)2(-3)	Mongkolsamrit et al. (2018)
myces ensis				Cream to light yellow, erect, irregularly branched	Cylindrical, solitary, sometimes verticillate, with phialides in whorts of four to five 4.3–9.5 x 1.2–2.0	Cylindrical, somewhat inflated base, 6.2-29.4 × 1.1-2.5	Ellipsoidal to long oval, 1.9–3.5 × 1.1–1.8	This study
myces s				Arising at the head and in the middle of the host body, white, up to 15.6 long, 0.6–1.3 wide, feathery to clavate with acute or blunt end	Monophialidic, produced along the symemata or solitary on hyphae in culture	Cylindrical, 11.5-30.0 × 2.0-4.2	Cylindrical or long oval, 4.1–9.8 × 2.3–4.2	This study
myces	Ovoid, 350-650 × 200-375	200-350 × 3.5-4				Relatively short, 11–20 (–30) × 1.3–1.8, aculeate and strongly tapering	Typically short- ellipsoidal, 2.5–3.5 (–4.2) × 1–1.5	Kepler et al. (2017); Zare and Gams (2001); Shrestha et al. (2019)
myces erorum				Synnemata not observed	Mononematous, with single phialide or two phialides	Cylindrical, somewhat inflated base, 12.7–25.8 × 1.4–1.7, tapering to a thin neck	Mostly cylindrical, 3.5– 5.6 × 1.4–2.1, forming mostly globose heads	Chen et al. (2020b)
myces us						(15-)20-35 × 1.0-1.7	Ellipsoidal to subcylindrical, $(2-)2.5-5.5(-6) \times 1-1.5(-1.8)$	Kepler et al. (2017); Zare and Gams (2001)
myces eogenus				Synnemata not observed	Moderately branched, with (1-)2-6 (-8) phialides	30-64 × 1.1-3.2	Forming mostly globose heads, cylindrical, 3.2-8.6 × 1.3-1.6	Chen et al. (2017); Mains (1949)
myces opterorum				Synnemata not observed	Mononematous, with single phialide or whorls of two to five phialides, or <i>Verticillium</i> - like from hyphae directly	Cylindrical, somewhat inflated base, 19.9–29.6 × 1.6–2.0, tapering to a thin neck	Mostly cylindrical, 3.3–6.6 × 1.5–1.8	Chen et al. (2020a)
myces arum	Ovoid, (530–)623– 993(–1000) × (290–)308–413(–425)	(170–)196– 423(–550) × (2–)2.7–3.8(–4)	(6-)7- 10.7(-13) ×1	Arising from moth body and wing veins, white to cream, erect, cylindrical to clavate, (650–)668–1191(–1500) × (50–)53.4–102(–120) µm	Monophialidic or polyphialidic	Cylindrical with papillate end, hyaline, $(5-)6.8-9(-10) \times (1.8-)2-2.4(-3)$	Cylindrical with round end, $(3-)3.5-4.5(-6) \times 1$	Aini et al. (2020)
myces				Synnemata not observed			Cylindrical to ovoid or oval, 4–9.2 × 1.6–2.4	Cope and Leal (2005)
nyces octuidarum				Arising from moth body, cream to light yellow, erect, cylindrical to clavate, 0.8-2 × 0.12-0.35	Cylindrical, solitary, 6.5– 13.8 × 1.8–3.6	Cylindrical with papillate end, 6.8–26.0 × 2.1–3.6	Ellipsoidal to long oval, 2.6–6.4 × 1.5–2.2	This study

References	Aini et al. (2020)	Kobayasi and Shimizu (1982)	5 Chiriví- 5 Salomón et al. (2015); Kepler et al. (2017)	l, Mongkolsamrit et al. (2018)	, This study	Mongkolsamrit et al. (2018)	Chen et al. (2022)	: Aini et al. (2020)	l Mains (1950)	Park et al. (2016)	Mongkolsamrit et al. (2018)	st Wang et al. (2023a)
Conidia (µm)	Not observed		Ellipsoidal to ovoid, usually straight, 3.5–4.5 × 1.5–2.0	Cylindrical to ellipsoidal $(4-)4.5-5.5(-6) \times 2-3$	Ellipsoidal to long oval _, 3.0–5.4 × 1.8–3.4	Cylindrical to ellipsoida (3-)4-6(-7) × 1.5-2	Fusiform, 2.3–3.0 × 1.5–2.3	Fusoid, long synnemats (1-)2.5-3(-3.2) × (0.8-)1-1.4(-2), short synnemata: (1-)1.8- $2.7(-3) \times 1-2$	Fusoid to subcylindrica narrowing at the ends, 2.5-5 × 1-1.5	Cylindrical, oblong, or ellipsoid, 3–9 × 1.8–3	Ellipsoidal or fusiform, (2-)3.5(-4) × 2-3	Long-ellipsoidal to almos cylindrical, (1.5–)3.0–
Phialides (µm)	Not observed		Solitary or in whorls of 2–4, 13– 19 long, from 1.0–2.0 gradually tapering to 0.5–1.0	$(10-)16(-20) \times 2-2.5$, with a cylindrical basal portion, tapering into a thin neck, 1×0.5	Cylindrical, somewhat inflated base, 12.1–38.2 × 1.3–3.2	(12-)13.5-21(-30) × 1-2, awl- shaped, <i>lecanicillium</i> -like	Solitary or in groups of two, 13.9– 17.1 × 1.1– 1.6 with a cylindrical basal portion and tapering into a short, distinct neck	Long synnemata: $(5-)6-8(-10)$ × $(1,8-)2-2.7(-3)$, short synnemata: $(5-)6.2-8.3(-10)$ × (1.8-)2-2.5(-3), cylindrical to ellipsoidal with papillate end	Subcylindrical, 6–10 × 2–3, narrowing above into an acute apex terminated by a short sterigma 2–3 long	Produced singly or in whorls of up to 3-4(-5) on prostrate hyphae, 20-60 × 1-2.5(-3)	$(10-)16(-22) \times (1-)1.5(-2)$, with cylindrical to ellipsoidal basal portion, tapering into a thin neck, $1-3 \times 1$	8.0-40.0 long, rarely over 100, 0.6-1.2 at the base, tapering to
Conidiophores (µm)	Not observed		Generally arising from submerged hyphae, moderately branched	Erect, verticillate with phialides in whorls of two to three	Cylindrical, solitary or verticillate with phialides in whorls of two to five, 6.5-12.3 × 1.6-3.5	Erect, forming verticillate branches with solitary phialides	Erect, usually aring from the aerial hyphae	Monophialidic or polyphialidic			Usually forming verticillate branches with phialides in whorls of two to five	
Synnemata (mm)	Synnemata not observed		Synnemata not observed	Synnemata not observed	Synnemata not observed	Synnemata not observed	Synnemata not observed	Long synnemata aring at the head and in the middle of the host body up to 5 long, 0.12–0.15 wide, cylindrical to clavate, short synnemata aring on moth body, wings and legs, (197–)200–267(–300) × (15–)17.7– 31.6(40–)µm, white to cream	Arising from all parts of the moths, clavate, 0.4-1.0 long, the stipe 0.025-0.05 thick	Synnemata not observed	Arising on legs of spider, erect, up to 1.5 long, 0.1–0.12 wide	Synnemata not observed
Part-spores (µm)	(5-)5.9- 9.4(-12) ×1	1 × 1-4		(300-)336(-450) × 1-1.5		4-6×1-1.5			2-6×0.5-1			
Asci (µm)	(170-)222- 329(-360) × (2-)2.53(-4)	5 wide, cap 3 wide		Up to 500 long, 2–3 wide		Up to 550 long, 5-7 wide			300-600 × 4-5			
Perithecia (µm)	Ovoid to obpyriform, (290–)342–580(– 650) × (150–)186– 291(–340)	Pyriformia, 570- 630 × 170-250		Narrowly ovoid, (650–)676(–680) × (240–)324.5(–330)		Narrowly ovoid, (700–)752–838(– 850) × (300–)305– 375(–400)			Narrowly ovoid or conoid, 420–900 × 180–370			
Species	Akanthomyces pyralidarum	Akanthomyces ryukyuenis	Akanthomyces sabanensis	Akanthomyces sulphureus	Akanthomyces subaraneicola	Akanthomyces thailandicus	Akanthomyces tiankengensis	Akanthomyces tortricidarum	Akanthomyces tuberculatus (= A. pistillariaeformis)	Akanthomyces uredinophilus	Akanthomyces waltergamsii	Akanthomyces zaquensis

studies are still an invaluable tool for fungal biology and ecology exploration (Walker et al. 2019). In addition to molecular data, morphological and ecological characteristics have a pivotal role in taxonomy and phylogenetic identification of fungi. In our work, we surveyed the literature to the greatest extent possible, combined that with the results of those obtained by morphological methods (optical microscope and electron microscope) in our study, to list and compare the morphological characteristics of 35 Akanthomyces species (Table 3). The morphological comparison revealed obvious differences in the size of ascospores and asci, morphology of the synnemata, conidiogenous structures and conidial shape and size, although the morphological features generally overlapped. Our statistics showed that at least 20 Akanthomyces species are specialists with narrow host ranges and they are either spider pathogens or adult moth entomopathogens (Table 2). They cause mortality of spiders and adult moths by nature. The cadavers are usually found attached to the underside of leaves or on tree trunks, barks, decaying logs, branches, grass, leaf litter and forest floors (Shrestha et al. 2019). These ecological characteristics are phylogenetically informative for distinguishing species of Akanthomyces and they contribute to the timely discovery of new Akanthomyces species in nature.

Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

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Data availability

All of the data that support the findings of this study are available in the main text or Supplementary Information.

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Supplementary material 1

Supplementary information

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Data type: docx

- Explanation note: **fig. S1.** Phylogenetic relationships among the genus *Akanthomyces* and its allies in Cordycipitaceae based on Bayesian inference (BI) and maximum likelihood (ML) analyses of a five-locus (ITS, nr*LSU, TEF, RPB1*, and *RPB2*) dataset. **fig. S2.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the ITS sequences. Statistical support values (≥70%) are shown at the nodes for ML boostrap support. **fig. S3.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the nr*LSU* sequences. **fig. S4.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *nrEF* sequences. **fig. S5.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB1* sequences. **fig. S6.** Phylogenetic tree of *Akanthomyces* based on Maximum Likelihood (IQ-TREE) analysis from the *RPB2* sequences.
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