

Research Article

Morphological and phylogenetic analyses reveal three new species and one new record of *Tubeufia* (Tubeufiales, Tubeufiaceae) from southern China

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Abstract

During an investigation of helicosporous fungi in China, a total of seven helicosporous hyphomycetes were obtained from decaying wood in the southern region of the country. Based on phylogenetic analyses using a combined LSU, ITS, *tef1a*, and *rpb2* sequence matrix, in conjunction with morphological comparisons, these taxa were classified with-in *Tubeufia* (Tubeufiaceae, Tubeufiales) and were recognized as three new species, *viz*. *Tubeufia guttulata*, *T. hainanensis*, and *T. muriformis*, as well as one new distribution record, *viz*. *T. cocois*. Evidence for these new taxa and the new record, descriptions, illustrations, notes, and phylogenetic evidence are provided for the newly collected helicosporous species.

Key words: asexual morph, new taxa, phylogeny, taxonomy

Introduction

Tubeufia was introduced by Penzig and Saccardo (1897) with the type species *T. javanica* and two other species (*T. anceps* and *T. coronata*). *Tubeufia* species are widely distributed in tropical and temperate regions, including freshwater and terrestrial habitats, occurring primarily on well-rotted wood (Barr 1979; Rossman 1987; Ho et al. 2002; Tsui and Berbee 2006; Tsui et al. 2007; Zhao et al. 2007; Boonmee et al. 2011, 2014; Chaiwan et al. 2017; Dai et al. 2017; Doilom et al. 2017; Lu et al. 2017, 2018a, 2018b, 2022, 2023; Tian et al. 2022). There are currently 56 accepted species in *Tubeufia*, as listed in Table 1. These species exhibit a wide range of morphological characteristics in both their sexual and asexual morphs. For example, although *Tubeufia bambusicola*, *T. javanica*, and *T. latispora* shared a close phylogenetic affinity, they possess black ascomata that are densely covered with long, flexuous setae. These ascomata contain sessile asci, each containing eight fusiform ascospores. In contrast, *T. javanica* has white ascomata without seta and short pedicellate asci with eight filiform ascospores (Boonmee



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Copyright: © Jian Ma et al. This is an open access article distributed under terms of the Creative Commons Attribution License (Attribution 4.0 International – CC BY 4.0). et al. 2014; Lu et al. 2017, 2018b). The asexual morphs of *Tubeufia* encompass both helicosporous and dictyosporous hyphomycetes. The helicosporous asexual morphs of *Tubeufia* display diverse shapes, with certain species exhibiting abundant, elongated conidiophores, while others possess extremely short conidiophores. For instance, *T. abundata* is characterized by abundant, branched, and long conidiophores, *T. arctata* exhibits rarely unbranched, short conidiophores, and *T. machaerinae* does not possess conidiophores (Lu et al. 2018b).

No.	Species	Distribution	Habitat	Molecular data	Reference
1	T. abundata	Thailand	Freshwater	Available	Lu et al. 2018b
2	T. acaciae	India	Terrestrial	Not available	Tilak and Kale 1969
3	T. aciculospora	Japan	Terrestrial	Not available	Katsumoto and Harada 1979
4	T. aquatica	China, Thailand	Freshwater	Available	Luo et al. 2017; Lu et al. 2018b
5	T. bambusicola	Thailand	Terrestrial	Available	Lu et al. 2018b
6	T. brevis	Thailand	Freshwater	Available	Lu et al. 2018b
7	T. brevispina	USA	Terrestrial	Not available	Barr and Rogerson 1983; Crane et al. 1998
8	T. brunnea	Thailand	Freshwater	Available	Lu et al. 2018b
9	T. chiangmaiensis	Thailand	Terrestrial	Available	Boonmee et al. 2014
10	T. chlamydospora	Thailand	Freshwater	Available	Lu et al. 2018b
11	T. claspisphaeria	China	Freshwater	Not available	Kodsueb et al. 2004
12	T. cocois	Thailand	Freshwater/ Terrestrial	Available	Tian et al. 2022
13	T. cylindrothecia	Thailand, USA	Freshwater	Available	Luo et al. 2017
14	T. dactylariae	China	Terrestrial	Not available	Chang 2003
15	T. dentophora	China	Terrestrial	Not available	Lu et al. 2018b
16	T. dictyospora	Thailand	Freshwater/ Terrestrial	Available	Lu et al. 2018b
17	T. eccentrica	China	Freshwater	Available	Lu et al. 2018b
18	T. entadae	Thailand	Terrestrial	Available	Jayasiri et al. 2019
19	T. eugeniae	India	Terrestrial	Not available	Pande 2008
20	T. fangchengensis	China	Freshwater	Available	Lu et al. 2018b
21	T. filiformis	Thailand	Freshwater	Available	Lu et al. 2017
22	T. freycinetiae	Thailand	Terrestrial	Available	Tibpromma et al. 2018
23	T. garugae	India	Terrestrial	Not available	Pande 2008
24	T. geniculata	China	Freshwater	Available	Lu et al. 2018b
25	T. guangxiensis	China	Freshwater	Available	Chaiwan et al. 2017
26	T. hechiensis	China	Freshwater	Available	Lu et al. 2018b
27	T. hyalospora	Thailand	Freshwater	Available	Hyde et al. 2016
28	T. inaequalis	Thailand	Freshwater	Available	Lu et al. 2018b
29	T. javanica	Thailand	Terrestrial	Available	Boonmee et al. 2014
30	T. krabiensis	Thailand	Freshwater	Available	Lu et al. 2018b
31	T. latispora	Thailand	Freshwater	Available	Lu et al. 2017
32	T. laxispora	Thailand	Freshwater	Available	Lu et al. 2017
33	T. lilliputea	Australia, China, India, Japan, USA	Terrestrial	Available	Lu et al. 2018b

Table 1. Checklist of accepted Tubeufia species.

No.	Species	Distribution	Habitat	Molecular data	Reference
34	T. liyui	China	Freshwater	Available	Lu et al. 2023
35	T. longihelicospora	China, Thailand	Freshwater	Available	Boonmee et al. 2021; Tian et al. 2022
36	T. longiseta	Thailand	Terrestrial	Available	Dai et al. 2017
37	T. machaerinae	China, USA	Freshwater/ Terrestrial	Available	Lu et al. 2018b
38	T. mackenziei	Thailand	Freshwater	Available	Lu et al. 2017
39	T. nigroseptum	China	Freshwater	Available	Li et al. 2022
40	T. minuta	Denmark, Sweden	Terrestrial	Not available	Munk 1966
41	T. pachythrix	Brazil	Terrestrial	Not available	Rossman 1979
42	T. pandanicola	Thailand	Terrestrial	Available	Tibpromma et al. 2018
43	T. parvispora	Thailand	Terrestrial	Available	Tibpromma et al. 2018
44	T. parvula	Britain, Sweden	Terrestrial	Not available	Dennis 1975
45	T. roseohelicospora	Thailand	Freshwater	Available	Hyde et al. 2016
46	T. rubra	Thailand	Freshwater	Available	Lu et al. 2018b
47	T. sahyadriensis	India	Terrestrial	Available	Rajeshkumar et al. 2019
48	T. sessilis	Thailand	Terrestrial	Available	Lu et al. 2018b
49	T. silentvalleyensis	India	Terrestrial	Not available	Pande 2008
50	T. sympodihylospora	China	Freshwater	Available	Lu et al. 2018b
51	T. sympodilaxispora	China	Freshwater	Available	Lu et al. 2018b
52	T. sympodiophora	China, Peru	Freshwater/ Terrestrial	Not available	Lu et al. 2018b
53	T. taiwanensis	China	Freshwater	Available	Lu et al. 2018b
54	T. tectonae	Thailand	Freshwater/ Terrestrial	Available	Doilom et al. 2017; Lu et al. 2018b
55	T. tratensis	Thailand	Freshwater	Available	Lu et al. 2018b
56	T. xylophila	China, India	Freshwater/ Terrestrial	Available	Lu et al. 2018b

The helicosporous taxa of Tubeufia represent a promising biological resource capable of producing novel bioactive secondary metabolites. Fan et al. (2019) discovered that T. rubra exhibited a significant antifungal impact against seven plant pathogenic fungi, viz. Alternaria solani (ZYB), Botryosphaeria dothidea (B12), Fusarium graminearum (CM), Magnaporthe grisea (DWB), Phoma sp. (HGHM), Phytophthora parasitica (HJB) and Rhizoctorzia solani (WB). The inhibition rate of T. rubra against these fungal pathogens exceeded 60%, indicating its potential as an effective antifungal agent. Fan et al. (2019) also discovered that the ethyl acetate extract from the fermentation broth of T. machaerinae inhibited the human cervical cancer cell line (HeLa) and human prostate cancer cell line (PC-3) by (98.92 ± 0.15%) and (97.86 ± 0.18%), respectively. According to Chen et al. (2020), both T. hechiensis and T. rubra exhibited the ability to reverse multidrug resistance in tumor cells. Zeng et al. (2022) discovered a newly isolated compound called Rubricin A from T. rubra, which demonstrated the capacity to inhibit the expression of P-glycoprotein (P-gp) and effectively reverse multidrug resistance in tumor cells. Qian et al. (2023) reported the isolation of two novel compounds, viz. Rubracin D and E, and sixteen known glyceroglycolipids from T. rubra. Rubracin D and E demonstrated significant multidrug resistance reversal activities on MCF-7/ADM, K562/ADM, and A549/ADM cell lines.

In this study, seven helicosporous taxa were collected from the southern provinces of Hainan and Guizhou in China. Based on morphological evidence and phylogenetic analyses, three novel species were introduced and designated as *Tubeufia guttulata*, *T. hainanensis* and *T. muriformis*, and one new distribution record, *viz. T. cocois*. The new species are described with detailed morphological descriptions and illustrations.

Materials and methods

Sample collection, specimen examination, and isolation

From August 2021 to March 2022, decaying wood samples were collected from Hainan and Guizhou provinces in southern China. The collected information includes locations, dates, altitudes, latitudes, and longitude. Fresh specimens were placed in zip-lock bags and sterile, moist plastic boxes, and then incubated at room temperature for a period of two weeks. Using a stereomicroscope (SMZ-168, Nikon, Japan), the fungal colonies growing on decaying wood surfaces were carefully examined, observed, and documented through photography. Morphological features such as conidiophores, conidiogenous cells, and conidia were further captured using an ECLIPSE Ni compound microscope (Nikon, Tokyo, Japan) in conjunction with a Canon 90D digital camera. Measurements were taken with the Tarosoft (R) Image Frame Workprogram. Photoplates were developed with Adobe PhotoShop CC 2019 (Adobe Systems, USA).

According to the method described by Senanayake et al. (2020), helicosporous strains were isolated and purified on water agar (WA), and germinated conidia were aseptically transferred to fresh potato dextrose agar (PDA) plates. Mycelium was cultured on Potato Dextrose Agar (PDA) and incubated at a temperature of 25 °C for a period of 6-7 weeks. During this time, various morphological characteristics including color, shape, and size were carefully observed and recorded. Subsequently, dried specimens of the fungal cultures were deposited in the Herbarium of Kunming Institute of Botany, Chinese Academy of Sciences (Herb. HKAS), Kunming, China, and the Herbarium of Guizhou Academy of Agriculture Sciences (Herb. GZAAS), Guiyang, China. Cultures were deposited in the China General Microbiological Culture Collection Center (CGMCC; https://www. cgmcc.net/english/), Beijing, China, and the Guizhou Culture Collection, China (GZCC), Guiyang, China for future reference and scientific documentation. Faces of fungi (FOF; https://www.facesoffungi.org) numbers were obtained in accordance with Jayasiri et al. (2015) and MycoBank numbers of the new species were registered in the MycoBank database (https://www.mycobank.org/).

DNA extraction, PCR amplification, and sequencing

Fresh fungal hyphae were gently scraped and transferred to a 1.5-mL microcentrifuge tube using a sterile toothpick. Genomic DNA was extracted using the Biospin Fungus Genomic DNA Extraction Kit (BioFlux, China) and following the manufacturer's protocol for DNA extraction. Specific forward and reverse primers, namely ITS5/ITS4, LR0R/LR5, fRPB2-5F/fRPB2-7cR, and EF1-983F/EF1-2218R, were employed to amplify the internal transcribed spacer (ITS, White et al. 1990), large ribosomal subunit (LSU, Vilgalys and Hester 1990), and RNA polymerase II second-largest subunit (*rpb2*, Liu et al. 1999) and translation elongation factor 1- α gene (*tef1a*, Rehner and Buckley 2005) sequence fragments, respectively. The polymerase chain reaction (PCR) conditions employed were in accordance with the reaction conditions outlined in the publications by Lu et al. (2017, 2018a). PCR amplification reactions were conducted in a 50-µL reaction volume containing 44 µL of 1.1×T3 Supper PCR Mix (Qingke Biotech, China), 2 µL of forward and reverse primers, and 2 µL of DNA template. The products were detected by 1% agarose gel electrophoresis following PCR amplification, with primers and PCR product sequencing provided by Beijing Qingke Biotechnology Co., Ltd.

Phylogenetic analyses

The original sequences were examined with BioEdit v 7.0.5.3 (Hall 1999) and assembled with SeqMan v. 7.0.0 (DNASTAR, Madison, WI, USA; Swindell and Plasterer 1997). The taxa used in this study (Table 2) were downloaded based on high levels of identity (> 90%) and selected outgroups (https://blast.ncbi.nlm.nih.gov/Blast.cgi). The polygenic dataset was aligned by MAFFT v.7.473 (https://mafft.cbrc.jp/alignment/server/, Katoh and Standley 2013; Katoh et al. 2019). The data were trimmed using trimAl.v1.2rev59 software (Capella-Gutiérrez et al. 2009). The multigenic sequences (LSU-ITS-*tef1a-rpb2*) were merged using SequenceMatrix-Windows-1.7.8 software (Vaidya et al. 2011). The aligned Fasta file was converted to Nexus format file for Bayesian inference (BI) analyses using AliView v. 1.27 (Daniel et al. 2010).

Tayan	Strain	GenBank Accessions				
Taxon		ITS	LSU	tef1a	rpb2	
Acanthohelicospora aurea	GZCC 16-0060	KY321323	KY321326	KY792600	MF589911	
A. guianensis	UAMH 1699	AY916479	AY856891	_	_	
T. abundata	MFLUCC 17-2024 ^T	MH558769	MH558894	MH550961	MH551095	
T. aquatica	MFLUCC 16-1249 ^T	KY320522	KY320539	KY320556	MH551142	
T. aquatica	DLUCC 0574	-	KY320538	KY320555	MH551141	
T. aquatica	MFLUCC 17-1794	MH558770	MH558895	MH550962	MH551096	
T. bambusicola	MFLUCC 17-1803 [™]	MH558771	MH558896	MH550963	MH551097	
T. brevis	MFLUCC 17-1799 [™]	MH558772	MH558897	MH550964	MH551098	
T. brunnea	MFLUCC 17-2022 [™]	MH558773	MH558898	MH550965	MH551099	
T. chiangmaiensis	MFLUCC 17-1801	MH558774	MH558899	MH550966	MH551100	
T. chiangmaiensis	MFLUCC 11-0514 ^T	KF301530	KF301538	KF301557	-	
T. chlamydospora	MFLUCC 16-0223 [™]	MH558775	MH558900	MH550967	MH551101	
T. cocois	MFLUCC 22-0001 [™]	OM102541	OL985957	OM355486	OM355491	
T. cocois	GZCC 22-2038	OR030844	OR030837	OR046681	-	
T. cylindrothecia	BCC 3559	-	AY849965	_	-	
T. cylindrothecia	BCC 3585	AY916482	AY856908	-	-	
T. cylindrothecia	DLUCC 0572	KY320520	KY320537	KY320554	-	
T. cylindrothecia	MFLUCC 16-1253	KY320519	KY320536	KY320553	-	
T. cylindrothecia	MFLUCC 16-1283	KY320518	KY320535	KY320552	MH551143	

Table 2. Taxa used in this study and their GenBank accession numbers of DNA sequences.

Tayan	Otroin	GenBank Accessions				
Taxon	Strain	ITS	LSU	tef1a	rpb2	
T. cylindrothecia	MFLUCC 17-1792	MH558776	MH558901	MH550968	MH551102	
T. dictyospora	MFLUCC 17-1805 [⊤]	MH558778	MH558903	MH550970	MH551104	
T. dictyospora	MFLUCC 16-0220	MH558777	MH558902	MH550969	MH551103	
T. eccentrica	GZCC 16-0048	MH558780	MH558905	MH550972	MH551106	
T. eccentrica	GZCC 16-0084	MH558781	MH558906	MH550973	MH551107	
T. eccentrica	MFLUCC 17-1524 ^T	MH558782	MH558907	MH550974	MH551108	
T. eccentrica	GZCC 16-0035	MH558779	MH558904	MH550971	MH551105	
T. entadae	MFLU 18-2102	MK347727	MK347943	-	-	
T. fangchengensis	MFLUCC 17-0047 ^T	MH558783	MH558908	MH550975	MH551109	
T. filiformis	MFLUCC 16-1128 ^T	_	KY092407	KY117028	MF535284	
T. filiformis	MFLUCC 16-1135	KY092416	KY092411	KY117032	MF535285	
T. filiformis	MFLUCC 16-0236	_	MH558938	MH550976	MH551110	
T. freycinetiae	MFLUCC 16-0252 ^T	MH275089	MH260323	MH412786	-	
T. geniculata	BCRC FU30849 [™]	LC335817	_	-	_	
T. geniculata	NCYU U2-1B	LC335816	_	-	_	
T. guangxiensis	GZCC 16-0054	MG012027	MG012020	MG012006	MG012013	
T. guangxiensis	GZCC 16-0090	MG012029	MG012022	MG012008	MG012015	
T. guangxiensis	GZCC 16-0091	MG012028	MG012021	MG012007	MG012014	
T. guangxiensis	MFLUCC 17-0038	MG012026	MG012019	MG012005	MG012012	
T. guangxiensis	MFLUCC 17-0045 ⁺	MG012025	MG012018	MG012004	MG012011	
T. guangxiensis	MFLUCC 17-0046	MH558784	MH558909	MH550977	MH551111	
T. guangxiensis	GZCC 16-0041	MG012030	MG012023	MG012009	MG012016	
T. guttulata	GZCC 23-0404 [™]	OR030841	OR030834	OR046678	OR046684	
T. guttulata	GZCC 23-0590	OR066413	OR066420	OR058859	OR058852	
T. hainanensis	GZCC 22-2015 [™]	OR030842	OR030835	OR046679	OR046685	
T. hainanensis	GZCC 23-0589	OR066414	OR066421	OR058860	OR058853	
T. hechiensis	MFLUCC 17-0052 [™]	MH558785	MH558910	MH550978	MH551112	
T. hyalospora	MFLUCC 15-1250 ^T	MH558786	MH558911	MH550979	-	
T. inaequalis	GZCC 16-0079	MH558787	MH558912	MH550980	MH551113	
T. inaequalis	GZCC 16-0087	MH558788	MH558913	MH550981	MH551114	
T. inaequalis	MFLUCC 17-0053 [™]	MH558789	MH558914	MH550982	MH551115	
T. inaequalis	MFLUCC 17-1989	MH558790	MH558915	MH550983	MH551116	
T. inaequalis	MFLUCC 17-1998	MH558791	MH558916	MH550984	MH551117	
T. inaequalis	BCC 8808	AY916481	AY856910	-	-	
T. javanica	MFLUCC 12-0545	KJ880034	KJ880036	KJ880037	-	
T. krabiensis	MFLUCC 16-0228 [™]	MH558792	MH558917	MH550985	MH551118	
T. latispora	MFLUCC 16-0027 ^T	KY092417	KY092412	KY117033	MH551119	
T. laxispora	MFLUCC 16-0013	MH558793	MH558918	MH550986	MH551120	
T. laxispora	MFLUCC 16-0219	KY092414	KY092409	KY117030	MF535286	
T. laxispora	MFLUCC 16-0232 [⊤]	KY092413	KY092408	KY117029	MF535287	
T. laxispora	MFLUCC 17-2023	MH558794	MH558919	MH550987	MH551121	
T. lilliputea	NBRC 32664	AY916483	AY856899	_	-	
T. liyui	GZCC 22-2030 [™]	OP888466	OP888465	-	-	

Tayon	Otroin	GenBank Accessions					
Taxon	Strain	ITS	LSU	tef1a	rpb2		
T. longihelicospora	MFLUCC 21-0814	OM331690	OM331688	-	-		
T. longihelicospora	MFLUCC 21-0815	OM331691	OM331705	-	-		
T. longihelicospora	MFLUCC21-0151	OL606156	OL606149	OL964520	OL964526		
T. longihelicospora	MFLUCC 16-0753 [™]	NR_182938	-	_	_		
T. longiseta	MFLUCC 15-0188 [™]	KU940133	-	_	_		
T. machaerinae	MFLUCC 17-0055 [™]	MH558795	MH558920	MH550988	MH551122		
T. mackenziei	MFLUCC 16-0222 [™]	KY092415	KY092410	KY117031	MF535288		
T. muriformis	GZCC 22-2039 [™]	OR030843	OR030836	OR046680	OR046686		
T. muriformis	GZCC 23-0591	OR066415	OR066422	OR058861	OR058854		
T. nigroseptum	CGMCC 3.20430 [™]	MZ092716	MZ853187	OM022002	OM022001		
T. pandanicola	MFLUCC 16-0321 ^T	MH275091	MH260325	-	_		
T. parvispora	MFLUCC 17-1992	MH558796	MH558921	MH550989	MH551123		
T. parvispora	MFLUCC 17-2003	MH558797	MH558922	MH550990	MH551124		
T. parvispora	MFLUCC 17-2009	MH558798	MH558923	MH550991	MH551125		
T. roseohelicospora	MFLUCC 16-0230	MH558799	MH558924	MH550992	MH551126		
T. roseohelicospora	MFLUCC 17-1797	MH558800	MH558925	MH550993	MH551127		
T. roseohelicospora	MFLUCC 15-1247 ^T	KX454177	KX454178	-	MH551144		
T. rubra	GZCC 16-0083 [™]	MH558802	MH558927	MH550995	MH551129		
T. rubra	GZCC 16-0081	MH558801	MH558926	MH550994	MH551128		
T. sahyadriensis	NFCCI 4252/RAJ 99.1 [⊤]	MH033849	MH033850	MH033851	-		
T. sahyadriensis	NFCCI RAJ 99.2	MN393081	MN393082	MN393083	-		
T. sessilis	MFLUCC 16-0021 [™]	MH558803	-	MH550996	MH551130		
T. sympodihylospora	GZCC 16-0051	MH558805	MH558929	MH550998	MH551132		
T. sympodihylospora	MFLUCC 17-0044 ^T	MH558806	MH558930	MH550999	MH551133		
T. sympodihylospora	GZCC 16-0049	MH558804	MH558928	MH550997	MH551131		
T. sympodilaxispora	BCC 3580	_	DQ296554	-	-		
T. sympodilaxispora	GZCC 16-0058 [™]	MH558807	MH558931	MH551000	MH551134		
T. sympodilaxispora	MFLUCC 17-0048	MH558808	MH558932	MH551001	MH551135		
T. taiwanensis	BCRC FU30844 [⊤]	LC316605	-	-	_		
T. tectonae	MFLUCC 16-0235	MH558809	MH558933	MH551002	MH551136		
T. tectonae	MFLUCC 17-1985	MH558810	MH558934	MH551003	MH551137		
T. tectonae	MFLUCC 12-0392 [™]	KU144923	KU764706	KU872763	-		
T. tratensis	MFLUCC 17-1993 ^T	MH558811	MH558935	MH551004	MH551138		
T. xylophila	MFLUCC 17-1520	MH558813	MH558937	MH551006	MH551140		
T. xylophila	GZCC 16-0038	MH558812	MH558936	MH551005	MH551139		
Tubeufiaceae sp.	BCC 3512	AY916484	AY856905	-	-		
Tubeufiaceae sp.	BCC 3381	-	AY787932	-	-		
Note: "T" denotes ex-type strain. Newly generated sequences are indicated in bold. "-" means no data available in GenBank.							

Maximum likelihood (ML) analysis was carried out using the IQ Tree online website (http://iqtree.cibiv.univie.ac.at/), employing Bayesian Information Criteria (BIC) as the criterion for model selection, as described by Nguyen et al. (2015). The server automatically conducted tests to determine the appropriate substitution model for the analysis. Bayesian inference (BI) analysis was conducted in MrBayes on XSEDE (3.2.7a) (Ronquist et al. 2012). The best-fit substitution model GRT + I +G was decided for LSU, ITS, *tef1a* and *rpb2* matrix by MrModeltest 2.3 under the Akaike Information Criterion (AIC) (Nylander et al. 2008). Four simultaneous Markov chains were run for 10,000,000 generations, and trees were sampled every 1000th generations. Burn-in phase was set at 25% and the remaining trees were used for calculating posterior probabilities (PP).

Phylogenetic trees were visualized and edited using FigTree v. 1.4.4 and Adobe Illustrator CC 2019v. 23.1.0 (Adobe Systems, USA). In addition, Adobe PhotoShop CC 2019 (Adobe Systems, USA) was used to create the photo-plates.

Phylogenetic results

The phylogenetic position of the newly isolated taxa was determined in this study using partial LSU-ITS-*tef1a-rpb2* nucleotide sequences. The concatenated sequence matrix consisted of LSU (1–845 bp), ITS (846–1440 bp), *tef1a* (1441–2352 bp), and *rpb2* (2353–3397 bp) for a total of 97 taxa, including two outgroup taxa, resulting in a matrix of 3,397 characters. Maximum likelihood (ML) and Bayesian inference (BI) analyses were conducted on the concatenated datasets of LSU, ITS, *tef1a*, and *rpb2*, both yielding similar tree topologies, and the ML tree is shown in Fig. 1.

Based on the multigene phylogenetic tree depicted in Fig. 1, our study revealed that the seven new collections belong to three distinct species within the genus *Tubeufia*. *Tubeufia guttulata* is a significantly distinct lineage from *T. cocois* with 100% ML/1.00 PP supports. *T. hainanensis* establishes a sister lineage to Tubeufiaceae BCC 3512 with 85% ML/0.99 PP supports. *T. muriform-is* can be distinguished from other related *Tubeufia* species by its distinct muriform conidia. Based on phylogenetic and morphological evidence, our new isolate, GZCC 22–2038, is recognized as *T. cocois*.

Taxonomy

Tubeufia guttulata J. Ma & Y.Z. Lu, sp. nov.

MycoBank No: 900504 Facesoffungi Number: FoF14265 Fig. 2

Etymology. The epithet *"guttulata"* refers to the guttulate conidia of this taxon. **Holotype.** HKAS 128936

Description. *Saprobic* on decaying wood in a terrestrial habitat. *Sexual morph* Undetermined. *Asexual morph* Hyphomycetous, helicosporous. *Colonies* on natural substrate superficial, effuse, gregarious, white. *Mycelium* partly immersed, hyaline to pale brown, septate, branched hyphae, smooth, with masses of crowded, glistening conidia. *Conidiophores* macronematous, mononematous, flexuous, cylindrical, branched or unbranched, septate, 101–247 µm long, 5.5–8 µm wide ($\overline{x} = 165 \times 7 \mu m$, n = 20), the lower part pale brown and the upper part hyaline, smooth-walled. *Conidiogenous cells* holoblastic, mono- to polyblastic, integrated, sympodial, intercalary or terminal, cylindrical, with a denticulate protrusion, truncate at apex after co-



Figure 1. Phylogenetic tree generated from maximum likelihood (ML) analysis based on a combined LSU, ITS, tef1a, and rpb2 sequence data. The bootstrap support values of ML are equal to or greater than 75%, and Bayesian posterior probabilities (PP) equal to or greater than 0.95 are given near the nodes as ML/PP, respectively. Acanthohelicospora aurea GZCC 16-0060 and A. guianensis UAMH 1699 were used as outgroup taxa. The new species are indicated in red bold and newly generated sequences are indicated in black bold. "T" denotes ex-type strain. Symbols after generic names in Tubeufia indicate the habitats of taxa as explained in the phylogram.



Figure 2. *Tubeufia guttulata* (HKAS 128936, holotype) **a**, **b** colonies on the host surface **c**-**e** conidiophores, conidiogenous cells and conidia **f**-**i** conidiogenous cells bearing conidia **j** germinated conidium **k**-**p** conidia **q**, **r** colonies on PDA, **q** from above **r** from below. Scale bars: 50 μ m (**c**-**e**); 20 μ m (**h**-**p**); 10 μ m (**f**, **g**).

nidial secession, 9–16 µm long, 4–6 µm wide ($\bar{x} = 12 \times 5$ µm, n = 25), hyaline to pale brown, smooth-walled. **Conidia** solitary, acropleurogenous, helicoid, rounded at tip, 25–34 µm diam and conidial filament 4–6 µm wide ($\bar{x} = 29 \times 5$ µm, n = 30), 170–220 µm long ($\bar{x} = 189$ µm, n = 30), indistinctly septate, coiled $1^{1}/_{4}$ – $2^{1}/_{4}$ times, becoming uncoiled in water, guttulate, hyaline, smooth-walled.

Culture characteristics. Conidia germinating on water agar and producing germ tubes within 8 h. Colonies on PDA circular with umbonate surface and undulate edge. Growth rate 21 mm diam in 42 days at 25 °C, with a brown center with pale brown edges on PDA.

Material examined. CHINA, Hainan Province, Wuzhishan City, Shui Man Town, Wuzhishan National Nature Reserve, 18°92'N, 109°63'E, on rotting wood in a terrestrial habitat, 26 December 2021, Jian Ma, WZS70 (HKAS 128936, holotype; GZAAS 23–0408, isotype), ex-type living cultures CGMCC, GZCC 23–0404; *Ibid.*, WZS99 (GZAAS 23–0593, paratype), living culture GZCC 23–0590.

Notes. *Tubeufia guttulata* is a sister species to *T. cocois* with 100% ML/1.00 PP supports, however, the phylogenetic tree shows that they are distinct species. Morphologically, *Tubeufia guttulata* differs from *T. cocois* in that it has longer conidiophores (101–247 μ m vs. 38–123 μ m) and larger conidia (170–220 μ m vs. 116–136 μ m). In addition, the helicoid conidia of *T. guttulata* become uncoiled in water, while *T. cocois* are coiled (Tian et al. 2022).

Tubeufia hainanensis J. Ma & Y.Z. Lu, sp. nov.

MycoBank No: 900505 Facesoffungi Number: FoF14266 Fig. 3

Etymology. The epithet "hainanensis" refers to the collecting site.

Holotype. HKAS 125884.

Description. *Saprobic* on decaying wood in moist ground. *Sexual morph* Undetermined. *Asexual morph* Hyphomycetous, helicosporous. *Colonies* on natural substrate superficial, effuse, gregarious, white. *Mycelium* superficial, partly immersed, hyaline to pale brown, septate, branched hyphae, smooth. *Conidiophores* macronematous, mononematous, straight to slightly flexuous, cylindrical, unbranched, septate, 44–56 µm long, 4–5 µm wide, pale brown to hyaline, smooth-walled. *Conidiogenous cells* holoblastic, mono- to polyblastic, integrated, sympodial, intercalary or terminal, cylindrical, with a denticulate protrusion, truncate at apex after conidial secession, 8–14 µm long, 3–5 µm wide ($\bar{x} = 9.5 \times 4 \mu m$, n = 20), hyaline to pale brown, smooth-walled. *Conidia* solitary, acropleurogenous, helicoid, rounded at tip, 16–21 µm diam conidial filament 1.5–4 µm wide ($\bar{x} = 19 \times 3 \mu m$, n = 30), 127–175 µm long ($\bar{x} = 144 \mu m$, n = 30), indistinctly septate, coiled $3^{1}/_{2}$ – $3^{3}/_{4}$ times, becoming uncoiled in water, guttulate, hyaline, smooth-walled.

Culture characteristics. Conidia germinating on water agar and producing germ tubes within 8 h. Colonies on PDA circular with umbonate surface and undulate edge. Growth rate 43 mm diam in 50 days at 25 °C, with a pale brown surface.



Figure 3. *Tubeufia hainanensis* (HKAS 125884, holotype) **a**, **b** colonies on the host surface **c**-**e** conidiophores, conidiogenous cells and conidia **f**-**h** conidiogenous cells bearing conidia **j**-**o** conidia **i** germinated conidium **p**, **q** colonies on PDA, **p** from above **q** from below. Scale bars: 20 μm (**c**-**e**, **i**-**l**); 10 μm (**f**-**h**, **m**-**o**).

Material examined. CHINA, Hainan Province, Haikou City, Xiuying District, Ecological leisure trail, 20°01'N, 110°25'E, on decaying wood in terrestrial habitat, 10 August 2021, Jian Ma, HK1 (HKAS 125884, holotype; GZAAS 22–2015, isotype), ex-type living cultures CGMCC, GZCC 22–2015; *Ibid.*, HK1-2 (HKAS 125883, paratype), living culture GZCC 23–0589.

Notes. *Tubeufia hainanensis* resembles *T. parvispora* morphologically, with solitary, acropleurogenous, hyaline, helicoid conidia. However, *Tubeufia hainanensis* can be distinguished from *T. parvispora* by its unbranched conidiophores (Lu et al. 2018b). *Tubeufia hainanensis* formed a sister clade to Tubeufiaceae (BBC 3512) with 85% ML/0.99 PP supports (Fig. 1), and the phylogeny indicated that it is distinct species.

Tubeufia muriformis J. Ma & Y.Z. Lu, sp. nov.

MycoBank No: 900506 Facesoffungi Number: FoF14267 Fig. 4

Etymology. The epithet "*muriformis*" refers to the multi-septate conidia of this taxon.

Holotype. HKAS 128853.

Description. Saprobic on decaying bamboo in a terrestrial habitat. Sexual morph Undetermined. Asexual morph Hyphomycetous, helicosporous. Colonies on natural substrate superficial, effuse, gregarious, white. Mycelium superficial, partly immersed, hyaline to pale brown, septate, branched hyphae, smooth, with masses of crowded, glistening conidia. Conidiophores macronematous, mononematous, straight or flexuous, simple, cylindrical, branched or unbranched, indistinctly septate, 13-36 µm long, 3.5-7.5 µm wide, hyaline, smooth-walled. Conidiogenous cells holoblastic, monoblastic, integrated, sympodial, terminal, cylindrical, truncate at apex after conidial secession, hyaline, smooth-walled. Conidia solitary, acrogenous, muriform, curved, $23-25 \mu m$ diam and conidial filament $11-15 \mu m$ wide ($\overline{x} = 24 \times 13 \mu m$, n = 20), 16.5–58.5 μ m long (\overline{x} = 49 μ m, n = 20), composed of two rows of cells with pale, multi-septate, apical cells cylindrical, basal cells truncate, constricted at septae, tapering toward base and top of conidia, coiled $\frac{1}{4}$ -1 times, not becoming loose in water, guttulate, hyaline to pale brown, thick-walled, smooth-walled.

Culture characteristics. Conidia germinating water agar and producing germ tubes within 12 h. Growth on PDA with a circular shape and umbonate surface and entire edge. Growth rate 42 mm diameter in 40 days at 25 °C with a pale brown surface.

Material examined. CHINA, Guizhou Province, Qianxinan Prefecture, Xianheping National Forest Park, 24°97'N, 105°63'E, on submerged decaying wood in a freshwater stream, 16 March 2022, Jian Ma, XHP38 (HKAS 128853, holotype; GZAAS 22-2039, isotype), ex-type living cultures CGMCC, GZCC 22–2039; *Ibid.*, XHP64 (GZAAS 23–0594, paratype), living culture GZCC 23–0591.

Notes. *Tubeufia muriformis* shares morphological similarities with *Xenosporium helicominum*, characterized by the presence of mononematous, straight or flexuous conidiophores, monoblastic, terminal, cylindrical conidiogenous cells,



Figure 4. *Tubeufia muriformis* (HKAS 128853, holotype) **a**, **b** colonies on the host surface **c**-**e** conidiophores, conidiogenous cells and conidia **d**, **f**-**i** conidiogenous cells **k**-**o** conidia **j** germinated conidium **p**, **q** colonies on PDA, **p** from above **q** from below. Scale bars: 20 μ m (**c**-**f**, **j**-**k**); 10 μ m (**g**-**i**, **l**-**o**).

and muriform, curved, hyaline to pale-brown conidia. However, *T. muriformis* can be distinguished from *X. helicominum* with its branched conidiophores and larger conidia $(23-25 \times 11-15 \,\mu\text{m vs.} 14-16.5 \times 5.5-6.5 \,\mu\text{m}; Zhao et al. 2007)$. Moreover, *T. muriformis* differs from other *Tubeufia* species in that it has muriform, curved conidia (Lu et al. 2018b).

The phylogenetic analysis indicated that *Tubeufia muriformis* formed a closely related clade with *T. bambusicola*, supported by ML bootstrap value of 100% and PP of 1.00. This phylogenetic relationship confirms that *Tubeufia muriformis* and *T. bambusicola* are distinct species, as demonstrated by the phylogenetic tree.

Tubeufia cocois X.G. Tian & Tibpromma et al. Journal of Fungi 8: 22 (2021). MycoBank No: 555070 Facesoffungi Number: FoF10576

Fig. 5

Description. Saprobic on submerged decaying wood in a freshwater stream. Sexual morph Undetermined. Asexual morph Hyphomycetous, helicosporous. Colonies on natural substrate superficial, effuse, gregarious, white. Mycelium superficial and partly immersed, hyaline, septate, branched hyphae, smooth, with glistening conidia. Conidiophores macronematous, mononematous, straight or slightly flexuous, cylindrical, unbranched, septate, 33-85 μ m long, 5-7.5 μ m wide (\bar{x} = 48.5 × 6 μ m, n = 20), the lower part pale brown and the upper part hyaline, smooth-walled. Conidiogenous cells holoblastic, polyblastic, integrated, sympodial, terminal, cylindrical, denticulate, with a tooth-like protrusion, 1.5-4 µm long, 1.5-2.5 µm wide, truncate at apex after conidial secession, 4.5-10.5 µm long, 4.5-6 µm wide $(\bar{x} = 8.5 \times 5.5 \,\mu\text{m}, \text{n} = 20)$, hyaline, smooth-walled. **Conidia** solitary, acropleurogenous, helicoid, rounded at tip, 23-29 µm diam and conidial filament 4–6.5 μ m wide (\bar{x} = 26 × 5.5 μ m, n = 30), 100.5–138 μ m long (\bar{x} = 118 μ m, n = 25), indistinctly septate, coiled $2-2^{1}/_{2}$ times, not becoming loose in water, guttulate, hyaline, smooth-walled.

Culture characteristics. Conidia germinating on water agar and producing germ tubes within 8 h. Colonies on PDA circular with flat surface and undulate edge. Growth rate 26 mm diameter in 35 days at 25 °C, with a dark brown to black surface.

Material examined. CHINA, Hainan Province, Qiongzhong Li and Miao Autonomous County, Baihualing Rainforest cultural tourism area, 18°98'N, 109°82'E, on rotting wood in a freshwater stream, 29 December 2021, Jian Ma, BH5 (GZAAS 22–2038), living culture GZCC 22–2038.

Notes. Tian et al. (2022) introduced *Tubeufia cocois* from the dead leaves of *Cocos nucifera* in Thailand. Our newly isolated strain (GZCC 22–2038) clusters with *T. cocois* (MFLUCC 22–0001) with 93% ML/1.00 PP support. Morpholog-ically, our new isolate shares the same morphological characteristics with the holotype (MFLU 21–0192) of *T. cocois*, thus we identified our new isolate as *T. cocois*. This is the first discovery of this species in a freshwater habitat and the first discovery of this species in China.



Figure 5. *Tubeufia cocois* (GZAAS 22–2038) **a**, **b** colonies on the host surface **c**–**f** conidiophores and conidiogenous cells **g**–**j** conidiogenous cells **I**–**q** conidia **k** germinated conidium **r**, **s** colonies on PDA, **r** from above **s** from below. Scale bars: 20 μm (**c**, **d**, **f**); 10 μm (**e**, **g**–**q**).

Discussion

In this study, a total of seven helicosporous hyphomycetous taxa were collected from the southern Chinese provinces of Guizhou and Hainan. By utilizing a combination of multigene phylogenetic analysis and morphological evidence, three previously unknown species were characterized and designated as *Tubeufia guttulata*, *T. hainanensis*, and *T. muriformis*. Additionally, an additional taxon, *T. cocois*, was documented for the first time in this study.

Tubeufia is the largest genus within the family Tubeufiaceae. Currently, this genus contains 59 species (Lu et al. 2018b, 2023; Rajeshkumar et al. 2019; Boonmee et al. 2021; Li et al. 2022; Tian et al. 2022), including the newly introduced species in this study, of which 29 are found in freshwater habitat, 24 in terrestrial habitats, and six from both freshwater and terrestrial habitats (Barr 1979; Rossman 1987; Ho et al. 2002; Tsui and Berbee 2006; Tsui et al. 2007; Zhao et al. 2007; Boonmee et al. 2011, 2014; Chaiwan et al. 2017; Dai et al. 2017; Doilom et al. 2017; Lu et al. 2017, 2018b, 2022, 2023; Tian et al. 2022). Among them, 39 species produce a helicosporous conidial state.

It should be noted that the morphological features of helicosporous fungi belonging to the genus Tubeufia exhibit distinct differences compared to other helicosporous genera. Summarizing the morphological characteristics of Tubeufia at the genus level is challenging due to the absence of similarity in conidiophores and conidia among its species. Such as the morphology of the newly discovered species Tubeufia muriformis resembles Xenosporium rather than Tubeufia (Goos 1990; Zhao et al. 2007). This discovery expands the asexual morphological characteristics of the genus Tubeufia, which is distinguished by conidia that are curved dorsoventrally. Lu et al. (2018b) reported dictyosporous conidia in Tubeufia, while Rajeshkumar et al. (2019) discovered a new dictyosporous asexual morph, indicating that this genus may contain many undiscovered species. In addition, the sexual morphs of Tubeufia exhibit considerable diversity, indicating a rapid evolutionary rate within this genus. The wide range of morphological variations observed may be related to the adaptation of species to environmental changes. Given the recent research progress in uncovering new structurally active compounds in Tubeufia species, there is a compelling need for further investigations into the taxonomy and secondary metabolites of this genus (Chen et al. 2020; Zeng et al. 2022; Qian et al. 2023).

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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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Author contributions

Morphological data, photo-plates and phylogenetic analyzes were completed by Jian Ma, Li-Juan Zhang and Xing-Juan Xiao. The original draft was written by Jian Ma, and Saranyaphat Boonmee, Ning-Guo Liu, Yuan-Pin Xiao, Zong-Long Luo, Yong-Zhong Lu revised the paper.

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Data availability

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

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